To our students, for their constant challenge and stimulation
This book is designed to provide a comprehensive review of both general and organ-specific pathology through multiple choice questions with explanations of the answers. The source materials are the ninth editions of Robbins and Cotran Pathologic Basis of Disease (PBD9) and Robbins Basic Pathology (BP9), and in several chapters, Robbins and Cotran Atlas of Pathology (AP3). The questions in this review book follow the chapters and topics in these source materials to facilitate ongoing self-assessment as students work their way through a curriculum to gain and then apply their understanding of key concepts. This book is intended to be a useful resource for students in a variety of health science training programs.

In keeping with recommended question writing style for licensing examinations, we have included single best-answer questions, most with a clinical vignette, followed by a series of homogenous choices. This approach emphasizes an understanding of pathophysiologic mechanisms and manifestations of disease in a clinical context. We have incorporated relevant laboratory, radiologic, and physical diagnostic findings in the questions to emphasize clinicopathologic correlations. Although this adds to the extent of individual questions, the thoroughness reinforces learning, as a review should. Each answer includes a succinct explanation of why a particular choice is “correct” and the other choices are “incorrect.” Each answer is referenced by page numbers to both Robbins and Cotran Pathologic Basis of Disease and Robbins Basic Pathology (both the current ninth edition and the previous eighth edition of each), and in several cases, to figures in the third edition of Robbins and Cotran Atlas of Pathology, to facilitate and encourage a more complete reading of topics targeted for further review. Pathology is a visually oriented discipline; hence full-color images accompany many of the questions. The illustrations are taken mainly from the Robbins textbooks, so students can reinforce their study of the figures in the texts with questions that utilize the same or similar images.

The revisions in this fourth edition reflect new topics and new understanding of disease processes reflected in the most recent editions of the Robbins textbooks. The questions are intentionally written to be fairly difficult, with the purpose of “pushing the envelope” of students’ understanding of pathology. We are pushing it even further with a comprehensive final examination section that includes questions drawn from challenging topics covered in the entire book.

Mastery of this book will better prepare the student for further challenges. Many of the questions require the student to engage in a “multi-step” process: first, to interpret the information presented to arrive at a diagnosis, and then to solve a problem based on that diagnosis. This reinforces the clinical reasoning skills needed in delivery of health care. We must hasten to add that no review book is a substitute for textbooks and other course materials provided by individual instructors within the context of a curriculum. This book should be used in conjunction with thorough study of Robbins and Cotran Pathologic Basis of Disease and/or Robbins Basic Pathology and curricular materials. Finally, we hope that both students and their faculty will find this review book to be a useful adjunct to the learning of pathology.

Edward C. Klatt, MD
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Although medical knowledge has increased exponentially over the past 100 years, the desire to learn and apply this knowledge to the service of others has not changed. The study and practice of the healing arts requires persistence more than brilliance. By continuing as a lifelong student, it is possible to become a better health practitioner with the passage of time.

Use this book to find where you are on the pathway to excellence and be inspired to continue down that path. We provide a guide to light the way toward knowledge in pathology within the welcoming environment of this book.

Common mistakes made by students in answering questions result from failure to read and analyze information carefully by: (1) relying on a single finding as an exclusionary criterion, and (2) ignoring important diagnostic information. Medicine is mostly analogue, not digital, and the information you obtain is applicable across a continuum of probability. In selecting the best answer, remember these four key elements: (1) read the question thoroughly, (2) define the terms (use your vocabulary), (3) rank possible answers from common to uncommon, and (4) recognize key diagnostic information that differentiates the answers.

There are no magic formulas for academic achievement. The most important thing you can do is to spend some time each day in a learning process. Learning requires modification of synaptic interfaces at the dendritic level in the brain, and for learning to occur, there are a finite number of synaptic modifications that can be established per unit time, above which total comprehension is reduced. Increasing the rate or length of information delivery diminishes the efficiency of learning. Lack of break periods or engaging in “all nighters” presage onset of diminished performance, particularly when least desirable—during an examination. There is also decay of learning over time, with inevitable random loss of data elements. The key branch points in learning, where review with reinforcement can reduce data loss, occur at 20 to 40 minutes (transfer to intermediate memory) and at 24 to 48 hours (transfer to long-term memory) following initial learning.

Develop methods for filtering information from quality sources. We live in an age of information overload. Stay on task and avoid distractions. Identify the important data and underlying concepts. Develop a specific, personalized plan for approaching, reviewing, and preparing for assessments of your knowledge. Seek quality feedback, both positive to provide motivation for your commitment to further learning, as well as negative to focus on your rate of progress toward competency.

We hope, therefore, that this review will be useful not only in preparing for examinations but also for courses you take throughout your career. It is our sincere hope that this review book will make you a better health practitioner in your chosen career.
We are very grateful to Laura Schmidt, content development specialist, and William Schmitt, executive content strategist, at Elsevier, for their support of this project. Special thanks is due Louise King, project manager, for her understanding of the needs of the authors, for providing good advice, and for her willingness to accommodate multiple changes. Nhu Trinh at The University of Chicago is acknowledged for crucial secretarial support to one of us. We are grateful to our families and colleagues for graciously accepting this additional demand on our time.

The authors also are indebted to the pioneers in pathology education for the Robbins and Cotran series, starting with the founding author, Dr. Stanley Robbins, and continuing with Dr. Ramzi Cotran. These lead authors have set the standard of excellence that characterizes the series. There continue to be numerous contributing authors who have made the Robbins and Cotran series a valuable educational tool.

Edward C. Klatt
Vinay Kumar
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UNIT I
A study of peripheral blood smears shows that neutrophil nuclei of women have a Barr body, whereas those of men do not. The Barr body is an inactivated X chromosome. Which of the following forms of RNA is most likely to play a role in Barr body formation?

A. lncRNA  
B. mRNA  
C. miRNA  
D. siRNA  
E. tRNA

In an experiment, a nuclear chromosomal gene is found to be actively transcribing messenger RNA (mRNA) that is transported into the cell cytoplasm. However, there is no observed protein product from translation of this mRNA. How is the silencing of this active gene’s mRNA most likely to occur?

A. Absence of tRNA  
B. Binding to miRNA  
C. Methylation of DNA  
D. Mutation of mRNA  
E. Upregulation of mtDNA

A proponent of Chilean Malbec, Syrah, and Merlot wines (all reds) touts their contribution to longevity, but this wine aficionado also controls his dietary caloric content so that his body mass index is <22. This lifestyle promotes increased insulin sensitivity and glucose utilization. He fully expects to live longer because he has read that caloric restriction prolongs life. In this man, which of the following intracellular substances will most likely mediate the effect of calorie restriction upon increased longevity?

A. Caspase  
B. Glutathione  
C. Sirtuins  
D. Telomerase  
E. Ubiquitin

A 40-year-old woman has had chronic congestive heart failure for the past 3 years. In the past 2 months, she developed a cough productive of rust-colored sputum. A sputum cytology specimen now shows numerous hemosiderin-laden macrophages. Which of the following subcellular structures in these macrophages is most important for the accumulation of this pigment?

A. Chromosome  
B. Endoplasmic reticulum  
C. Golgi apparatus  
D. Lysosome  
E. Ribosome

An experiment is conducted in which cells in tissue culture are subjected to high levels of ultraviolet radiant energy. Electron microscopy shows cellular damage in the form of increased cytosolic aggregates of denatured proteins. In situ hybridization reveals that protein components in these aggregates also are found in proteasomes. Which of the following substances most likely binds to the denatured proteins, targeting them for catabolism by cytosolic proteasomes?

A. Adenosine monophosphate  
B. Calcium  
C. Caspase  
D. Granzyme B  
E. Hydrogen peroxide  
F. Ubiquitin

At the site of a surgical incision, endothelial cells elaborate vascular endothelial growth factor. There is sprouting with migration of endothelial cells into the wound to establish new capillaries. Which of the following intracellular proteins is most important in facilitating movement of endothelial cells?

A. Actin  
B. Cytokeratin  
C. Desmin  
D. Lamin  
E. Myosin
In an experiment, release of epidermal growth factor into an area of denuded skin causes mitogenic stimulation of the skin epithelial cells. Which of the following proteins is most likely to be involved in transducing the mitogenic signal from the epidermal cell membrane to the nucleus?

A. Cyclic AMP  
B. Cyclin D  
C. Cyclin-dependent kinase  
D. G proteins  
E. RAS proteins

Various soluble mediators are added to a cell culture containing epidermal cells to determine which of the mediators might be useful for promoting epidermal cell growth. When epidermal growth factor (EGF) is added, it binds to epidermal cell surface receptors, with subsequent transcription factor translocation and DNA transcription. This effect in the epidermal cells is most likely to be mediated through which of the following intracellular pathways?

A. Calcium ion channel  
B. Cyclic AMP  
C. Cyclin-dependent kinase  
D. JAK/STAT system  
E. Mitogen-activated protein (MAP) kinase

An experiment involves factors controlling wound healing. Skin ulcerations are observed, and the factors involved in the healing process are analyzed. Which of the following factors is most likely to be effective in promoting angiogenesis?

A. Basic fibroblast growth factor  
B. Endostatin  
C. Epidermal growth factor  
D. Interleukin-1  
E. Platelet-derived growth factor

In an experiment, surgical incisions are made in a study group of laboratory rats. Observations about the wounds are recorded over a 2-week period using various chemical mediators. Which of the following steps in the inflammatory-repair response is most likely affected by neutralization of transforming growth factor β (TGF-β)?

A. Chemotaxis of lymphocytes  
B. Increase in vascular permeability  
C. Leukocyte extravasation  
D. Migration of epithelial cells  
E. Production of collagen

A 62-year-old man has had increasing knee pain with movement for the past 10 years. The knee joint surfaces are eroded and the joint space narrowed. There is loss of compressibility and lubrication of articular cartilaginous surfaces. Loss of which of the following extracellular matrix components has most likely occurred in this man?

A. Elastin  
B. Fibronectin  
C. Hyaluronan  
D. Integrin  
E. Laminin

An experiment is conducted involving cellular aspects of wound healing. Components of the extracellular matrix are analyzed to determine their sites of production and their binding patterns to other tissue components. Which of the following molecules synthesized by fibroblasts can best bind to cellular integrins and extracellular collagen and attach epidermal basal cells to basement membrane?

A. Dermatan sulfate  
B. Fibronectin  
C. Heparin  
D. Hyaluronic acid  
E. Procollagen

An experiment analyzes factors involved in the cell cycle during growth factor-induced cellular regeneration in a tissue culture. Cyclin B synthesis is induced; the cyclin B binds and activates cyclin-dependent kinase 1 (CDK1). The active kinase produced by this process is most likely to control progression in which of the following phases of the cell cycle?

A. G0 to G1  
B. G1 to S  
C. S to G2  
D. G2 to M  
E. M to G1

In an experiment, the role of low-density lipoprotein (LDL) receptors in uptake of lipids in the liver is studied. A mouse model is created in which the LDL receptor gene is not expressed in the liver. For creating such a knockout mouse, which of the following cells would be most useful?

A. Adult bone marrow mesenchymal progenitor cells  
B. Embryonic stem cells in culture  
C. Hematopoietic stem cells  
D. Hepatic oval cells  
E. Regenerating hepatocytes

Dermal fibroblasts are harvested from the skin biopsy specimen of an adult man. These fibroblasts are transduced with genes encoding for transcription factors including SOX2 and MYC. Under appropriate culture conditions these cells are then able to generate endodermal, mesodermal, and ectodermal cells. Into which of the following kinds of stem cell have these fibroblasts been transformed?

A. Embryonic  
B. Lineage-committed  
C. Mesenchymal  
D. Pleuripotent
ANSWERS

1. A There are forms of noncoding RNA that play a role in gene expression. Long noncoding RNA (IncRNA) segments greater than 200 nucleotides in length can bind to chromatin to restrict access of RNA polymerase to coding segments. The X chromosome transcribes XIST, an IncRNA that binds to and represses X chromosome expression. However, not all genes on the “inactive” X chromosome are switched off. The RNA transcribed from nuclear DNA that directs protein synthesis through translation is mRNA. MicroRNAs (miRNAs) are noncoding RNA sequences that inhibit the translation of mRNAs. Gene-silencing RNAs (small interfering RNAs [siRNAs]) have the same function as miRNAs, but they are produced synthetically for experimental purposes. Transfer RNA (tRNA) participates in the translation of mRNA to proteins by linking to specific amino acids.

PBD9 5–6  BP9 217–218  BP8 150–152  BP8 235–237

2. B MicroRNAs (miRNA) are encoded by about 5% of the human genome. miRNAs do not encode for proteins, but bind to and inactivate or cleave to mRNA, preventing translation of proteins by mRNA, effectively silencing gene expression without affecting the gene directly. There is abundant tRNA present in the cytoplasm that is not a rate-limiting step to translation. DNA methylation, particularly at CG dinucleotides, is a way of suppressing gene expression directly, as is seen with genomic imprinting. Mutations that occur in genes in DNA may result in reduced mRNA production or abnormal protein production, but mRNA itself is not mutated. Mitochondrial DNA (mtDNA) encodes for proteins mainly involved in oxidative phosphorylation metabolic pathways.

PBD9 4–5  BP9 217–218  BP8 137

3. C The one sure way to increase life span is calorie restriction. But why do without the things we like, only to do without them longer? Dietary excesses lead to increased morbidity with reduced quality of life, as well as mortality, from chronic diseases such as diabetes mellitus. The activity of sirtuins on histone acetylation and deacetylation may promote transcription of genes encoding for proteins that increase metabolic activity and inhibit effects of free radicals. Red wines have been shown to increase sirtuins, but don’t drink too much! Moderation is the key. Glutathione promotes free radical breakdown, although chronic excessive alcohol consumption depletes hepatocyte glutathione. Caspases trigger apoptosis and cell death. Telomerase aid in promoting continued cell division, but cannot be altered by lifestyle, and turning them on is one feature of neoplasia. Ubiquitin is a peptide that is part of the ubiquitin-proteasome pathway of protein degradation seen with nutrient deficiencies, so when you eat less, be sure to eat a balanced diet.

PBD9 3–4, 68  BP9 26–27  BP8 41, 444  BP8 28

4. D Heterophagocytosis by macrophages requires that endocytosed vacuoles fuse with lysosomes to degrade the engulfed material. With congestive heart failure, extravasation of RBCs into alveoli occurs, and pulmonary macrophages must phagocytose the RBCs, breaking down the hemoglobin and recycling the iron by hemosiderin formation. The other listed options are components that play a role in cell synthetic functions.

PBD9 10, 13  BP9 22–23  BP8 52–53  BP8 12

5. F Heat-shock proteins provide for a variety of cellular “housekeeping” activities, including recycling and restoration of damaged proteins and removal of denatured proteins. Ubiquitin targets denatured proteins and facilitates their binding to proteasomes, which then break down the proteins to peptides. ADP increases when ATP is depleted, helping to drive anaerobic glycolysis. Cytosolic calcium levels may increase with cell injury that depletes ATP; the calcium activates phospholipases, endonucleases, and proteases, which damage the cell membranes, structural proteins, and mitochondria. Caspases are enzymes that facilitate apoptosis. Granzyme B is released from cytotoxic T lymphocytes and triggers apoptosis. Hydrogen peroxide is one of the activated oxygen species generated under conditions of cellular ischemia, producing nonspecific damage to cellular structures, particularly membranes.

PBD9 13–14  BP9 21–22  BP8 37–38  BP8 22

6. A Actin is a microfilament involved with cell movement. The other possibilities listed in B to D are intermediate filaments, which are larger than actin but smaller than myosin (a thick filament interdigitating with actin, required for muscle movement). Cytokeratins form cytoskeletal elements of epithelial cells. Desmin forms the scaffold in muscle cells on which actin and myosin contract. Lamin is associated with the nuclear membrane.

PBD9 10–11  BP8 50

7. E RAS proteins transduce signals from growth factor receptors, such as epidermal growth factor, that have intrinsic tyrosine kinase activity. G proteins perform a similar function for G protein–linked, seven-transmembrane receptors. Cyclic AMP is an effector in the G protein signaling pathway. Cyclins and cyclin-dependent kinases regulate the cell cycle in the nucleus.

PBD9 17  BP9 179  BP8 90–92  BP8 64, 66

8. E The MAP kinase cascade is involved in signaling from activation via cell surface receptors for growth factors. This pathway is particularly important for signaling of EGF and fibroblast growth factor. Ligand binding, such as occurs with acetylcholine at a nerve-muscle junction, alters the conformation of ion channel receptors to allow flow of specific ions such as calcium into the cell, changing the electric potential across the cell membrane. Cyclic AMP is a second messenger that is typically activated via ligand binding to receptors with seven transmembrane segments that associate with GTP-hydrolyzing proteins; chemokine receptors...
function in this fashion. Cyclin-dependent kinases act within the nucleus. JAK/STAT pathways typically are recruited by cytokine receptors.

9 A Basic fibroblast growth factor is a potent inducer of angiogenesis. It can participate in all steps of angiogenesis. Endostatin is an inhibitor of angiogenesis. Epidermal growth factor and interleukin-1 have no significant angiogenic activity. Platelet-derived growth factor plays a role in vascular remodeling.

10 E TGF-β stimulates many steps in fibrogenesis, including fibroblast chemotaxis and production of collagen by fibroblasts, while inhibiting degradation of collagen. All of the other steps listed are unaffected by TGF-β.

11 C He has osteoarthritis, or degenerative joint disease, with loss of articular hyaline cartilage. Hyaluronan (hyaluronic acid) is a large mucopolysaccharide, one form of proteoglycan, which forms a hydrated, compressible gel contributing to the shock-absorbing function of joint surfaces. Elastin is a fibrillar protein that provides recoil in tissues such as skin, arterial walls, and ligaments that need to stretch and return to their original shape. Fibronectin is a form of glycoprotein that serves an adhesive function. Integrins are glycoproteins that serve as cellular receptors for extracellular matrix components; they can link to intracellular actin so that cells can alter their shape and mobility. Laminins are a form of glycoprotein that help to anchor epithelial surfaces in basement membranes.

12 B Fibronectin is a key component of the extracellular matrix and has a structure that looks like a paper clip. Fibronectin can be synthesized by monocytes, fibroblasts, and endothelium. Dermatan sulfate, a glycosaminoglycan, forms a gel that provides resilience and lubrication. Heparin that is infused has an anticoagulant function. Hyaluronic acid binds water to form a gelatinous extracellular matrix. Procollagen produced by fibroblasts is formed into ropelike strands of collagen, which provide tensile strength.

13 D CDK1 controls an extremely important transition point, the G2 to M transition, during the cell cycle, which can be regulated by CDK inhibitors. The other checkpoints listed are regulated by a distinct set of proteins.

14 B Embryonic stem (ES) cells are multipotent and can give rise to all cells, including hepatocytes. Gene targeting to produce knockout mice is done in cultures of ES cells, which are then injected into mouse blastocysts and implanted into the uterus of a surrogate mother. Mesenchymal stem cells also are multipotential, but they are not useful for gene targeting. Hematopoietic stem cells can give rise to all hematopoietic cells, but not other types of cells. Hepatocytes and oval cells within the liver can give rise only to liver cells.

15 D These transformed cells are designated iPS cells because they have been induced to become pluripotent. This transformation process gets around the problem of using embryonic stem (ES) cells derived from manipulation of human embryos, which raises ethical and religious concerns. Embryonic stem cells are totipotent, but they become pluripotent cells that can further divide into many different cell lines, yet maintain themselves in a replicating pool. Thus pluripotent cells are the next best thing compared to embryonic cells for deriving human cells that could replace damaged or diseased tissues. Further differentiation of pluripotent cells gives rise to cells with more restricted developmental capacity, such as mesenchymal stem cells that can give rise to tissues such as muscle and cartilage but not to endodermal or ectodermal cells.
A 77-year-old woman has chronic renal failure. Her serum urea nitrogen is 40 mg/dL. She is given a diuretic medication and loses 2 kg (4.4 lb). She reduces the protein in her diet and her serum urea nitrogen decreases to 30 mg/dL. Which of the following terms best describes cellular responses to disease and treatment in this woman?

A. Adaptation  
B. Apoptosis  
C. Necroptosis  
D. Irreversible injury  
E. Metabolic derangement

A 53-year-old woman with no prior illnesses has a routine checkup by her physician. On examination she has a blood pressure of 150/95 mm Hg. If her hypertension remains untreated for years, which of the following cellular alterations would most likely be seen in her myocardium?

A. Apoptosis  
B. Dysplasia  
C. Fatty change  
D. Hemosiderosis  
E. Hyperplasia  
F. Hypertrophy  
G. Metaplasia

A 22-year-old woman becomes pregnant. A fetal ultrasound examination at 13 weeks’ gestation shows her uterus measures 7 × 4 × 3 cm. At delivery of a term infant, her uterus measures 34 × 18 × 12 cm. Which of the following cellular processes has contributed most to the increase in her uterine size?

A. Endometrial glandular hyperplasia  
B. Myometrial fibroblast proliferation  
C. Endometrial stromal hypertrophy  
D. Myometrial smooth muscle hypertrophy  
E. Vascular endothelial hyperplasia

A 20-year-old woman breastfeeds her infant. On examination, her breasts are slightly increased in size. Milk can be expressed from both nipples. Which of the following processes that occurred in her breasts during pregnancy enables her to breastfeed the infant?

A. Ductal metaplasia  
B. Epithelial dysplasia  
C. Intracellular lipid deposition  
D. Lobular hyperplasia  
E. Stromal hypertrophy

A 16-year-old boy sustained blunt trauma to his abdomen when he struck a bridge abutment at high speed while driving a motor vehicle. Peritoneal lavage shows a hemoperitoneum, and at laparotomy, a small portion of the left lobe of the injured liver is removed. Two months later, a CT scan of the abdomen shows that the liver has nearly regained its size before the injury. Which of the following processes best explains this CT scan finding?

A. Apoptosis  
B. Dysplasia  
C. Hyperplasia  
D. Hydropic change  
E. Steatosis

A 71-year-old man has had difficulty with urination, including hesitancy and increased frequency, for the past 5 years. A digital rectal examination reveals that his prostate gland is palpably enlarged to twice normal size. A transurethral resection of the prostate is performed, and the microscopic appearance of the prostate “chips” obtained is that of nodules of glands with intervening stroma. Which of the following pathologic processes has most likely occurred in his prostate?

A. Apoptosis  
B. Dysplasia  
C. Fatty change  
D. Hyperplasia  
E. Hypertrophy  
F. Metaplasia
7. A 29-year-old man sustains a left femoral fracture in a motorcycle accident. His leg is placed in a plaster cast. After his left leg has been immobilized for 6 weeks, the diameter of the left calf has decreased in size. This change in size is most likely to result from which of the following alterations in his calf muscles?

A. Aplasia  
B. Atrophy  
C. Dystrophy  
D. Hyalinosis  
E. Hypoplasia

8. A 34-year-old obese woman has experienced heartburn from gastric reflux for the past 5 years after eating large meals. She undergoes upper gastrointestinal endoscopy, and a biopsy specimen of the distal esophagus is obtained. Which of the following microscopic changes, seen in the figure, has most likely occurred?

A. Columnar metaplasia  
B. Goblet cell hyperplasia  
C. Lamina propria atrophy  
D. Squamous dysplasia  
E. Mucosal hypertrophy

9. An 11-year-old girl becomes infected with hepatitis A and experiences mild nausea for 1 week. On physical examination, she has minimal right upper quadrant tenderness and scleral icterus. Laboratory findings include a serum AST of 68 U/L, ALT of 75 U/L, and total bilirubin of 5.1 mg/dL. Her laboratory findings most likely result from which of the following changes in her hepatocytes?

A. Cell membrane defects  
B. Lysosomal autophagy  
C. Mitochondrial swelling  
D. Nuclear chromatin clumping  
E. Ribosomal disaggregation

10. A 33-year-old woman has had increasing lethargy and decreased urine output for the past week. Laboratory studies show her serum creatinine is 4.3 mg/dL and urea nitrogen 40 mg/dL. A renal biopsy is performed, and the specimen is examined using electron microscopy. Which of the following morphologic cellular changes most likely suggests a diagnosis of acute tubular necrosis?

A. Chromatin clumping  
B. Mitochondrial swelling

11. A 50-year-old man has experienced an episode of chest pain for 6 hours. A representative histologic section of his left ventricular myocardium is shown in the figure. There is no hemorrhage or inflammation. Which of the following conditions most likely produced these myocardial changes?

A. Arterial thrombosis  
B. Autoimmunity  
C. Blunt chest trauma  
D. Protein-deficient diet  
E. Viral infection

12. A 38-year-old woman has experienced severe abdominal pain over the past day. On examination she is hypotensive and in shock. Laboratory studies show elevated serum lipase. From the representative gross appearance of the mesentery shown in the figure, which of the following events has most likely occurred?

A. Acute pancreatitis  
B. Gangrenous cholecystitis  
C. Hepatitis B virus infection  
D. Small intestinal infarction  
E. Tuberculous lymphadenitis
13. A 68-year-old woman suddenly lost consciousness and on awakening 1 hour later, she could not speak or move her right arm. Two months later, a head CT scan showed a large cystic area in the left parietal lobe. Which of the following pathologic processes has most likely occurred in her brain?

- A. Apoptosis
- B. Coagulative necrosis
- C. Fat necrosis
- D. Karyolysis
- E. Liquefactive necrosis

14. A screening chest radiograph of an asymptomatic 37-year-old man shows a 3-cm nodule in the middle lobe of his right lung. The nodule is excised with a pulmonary wedge resection, and sectioning shows a sharply circumscribed mass with a soft, white center. The microscopic appearance is shown in the figure. The serum interferon gamma release assay is positive. Which of the following pathologic processes has most likely occurred in this nodule?

- A. Apoptosis
- B. Caseous necrosis
- C. Coagulative necrosis
- D. Fat necrosis
- E. Fatty change
- F. Gangrenous necrosis
- G. Liquefactive necrosis

15. An experimental drug administered to a tissue preparation is found to inhibit cellular oxidative phosphorylation when given in high doses, and ATP production drops to 5% of normal. Cell membrane function is diminished. Which of the following substances is most likely to be present at increased concentration in culture fluid bathing the tissue?

- A. Calcium
- B. Glucose
- C. Ketones
- D. Potassium
- E. Sodium

16. A 47-year-old woman has poorly controlled diabetes mellitus and develops coronary artery disease. She now has decreasing cardiac output with blood pressure of 80/40 mm Hg and ejection fraction of 18%. An increase in which of the following substances in her blood is most indicative of reversible cell injury from decreased systemic arterial perfusion of multiple organs and tissues?

- A. Carbon dioxide
- B. Creatinine
- C. Glucose
- D. Lactic acid
- E. Troponin I

17. A tissue preparation is experimentally subjected to a hypoxic environment. The cells in this tissue begin to swell, and chromatin begins to clump in cell nuclei. ATPases are activated, and ATP production decreases. Which of the following ions accumulating in mitochondria and the cytosol contributes most to these findings and to eventual cell death?

- A. Ca²⁺
- B. Cl⁻
- C. HCO₃⁻
- D. K⁺
- E. Na⁺
- F. PO₄³⁻

18. In an experiment, a large amount of a drug is administered to experimental organisms and is converted by cytochrome P-450 to a toxic metabolite. Accumulation of this metabolite leads to increased intracellular lipid peroxidation. Depletion of which of the following intracellular substances within the cytosol exacerbates this form of cellular injury by this mechanism?

- A. ADP
- B. Glutathione
- C. NADPH oxidase
- D. Nitric oxide synthase
- E. mRNA
- F. Sodium

19. In an experiment, metabolically active cells are subjected to radiant energy in the form of x-rays. This results in cell injury caused by hydrolysis of water. Which of the following intracellular enzymes helps to protect the cells from this type of injury?

- A. Endonuclease
- B. Glutathione peroxidase
- C. Lactate dehydrogenase
- D. Phospholipase
- E. Protease

20. A 5-year-old child ingests 50 iron tablets, each with 27 mg of iron. Within 6 hours the child develops abdominal pain and lethargy. On physical examination he is hypotensive. Laboratory studies show metabolic acidosis. Through formation of which of the following compounds is the cell injury in this child most likely mediated?

- A. Ascorbic acid
- B. Hemosiderin
- C. Hydroxyl radical
- D. Nitric oxide
- E. Superoxide dismutase
A 63-year-old man has a 2-year history of worsening congestive heart failure. An echocardiogram shows mitral valve stenosis with left atrial dilation. A mural thrombus is present in the left atrium. One month later, he experiences left flank pain and notes hematuria. Laboratory testing shows an elevated serum AST. The representative microscopic appearance of the lesion is shown in the figure. Which of the following patterns of tissue necrosis is most likely to be present in this man?

A  Caseous
B  Coagulative
C  Fat
D  Gangrenous
E  Liquefactive

A 54-year-old man experienced severe substernal chest pain for 3 hours. An ECG showed changes consistent with an acute myocardial infarction. After thrombolytic therapy with tissue plasminogen activator (t-PA), his serum creatine kinase (CK) level increased. Which of the following tissue events most likely occurred in the myocardium after t-PA therapy?

A  Cellular regeneration
B  Drug toxicity
C  Increased synthesis of CK
D  Myofiber atrophy
E  Reperfusion injury

On day 28 of her menstrual cycle, a 23-year-old woman experiences onset of menstrual bleeding that lasts for 6 days. She has had regular cycles since menarche. Which of the following processes most likely occurs in her endometrial cells to initiate the onset of menstrual bleeding?

A  Apoptosis
B  Atrophy
C  Caseous necrosis
D  Heterophagocytosis
E  Liquefactive necrosis

An experiment introduces a knockout gene mutation into a cell line. The frequency of shrunken cells with chromatin clumping, karyorrhexis, and cytoplasmic blebbing is increased compared with a cell line without the mutation. Overall survival of the mutant cell line is reduced. Which of the following genes is most likely to be affected by this mutation?

A  BAX
B  BCL2
C  C-MYC
D  FAS
E  p53

A 22-year-old woman with leukemia undergoes bone marrow transplantation and receives a partially mismatched donor marrow. One month later, she has a scaling skin rash. A skin biopsy is obtained, and on microscopic examination, it has the cellular change shown in the figure. This change most likely results from which of the following biochemical reactions?

A  Activation of caspases
B  Elaboration of lipases
C  Increase in glycolysis
D  Peroxidation of lipids
E  Reduction of ATP synthesis

A 47-year-old man has a lung carcinoma with metastases. He receives chemotherapy. A month later, histologic examination of a metastatic lesion shows many foci in which individual tumor cells appear shrunken and deeply eosinophilic. Their nuclei exhibit condensed aggregates of chromatin under the nuclear membrane. The pathologic process affecting these shrunken tumor cells is most likely triggered by release of which of the following substances into the cytosol?

A  BCL2
B  Catalase
C  Cytochrome c
D  Lipofuscin
E  Phospholipase
27 In a study of viral hepatitis infection, it is observed that cytotoxic T lymphocytes (CTLs) induce death in virally infected hepatocytes. The CTLs release perforin to allow entry of their granules. Which of the following substances is found in those granules that directly activates programmed cell death?
   A. BCL2
   B. Endonuclease
   C. Granzyme B
   D. Nitric oxide
   E. p53

28 An experimental study of steatohepatitis in metabolic syndrome reveals that hepatocyte cell membrane injury with necrosis occurs in response to increased amounts of tumor necrosis factor (TNF). When a pharmacologic agent inhibiting caspases is administered, cell necrosis still occurs. Which of the following substances forms a supramolecular complex that increases the generation of reactive oxygen species?
   A. Catalase
   B. Cytochrome c
   C. Interleukin 1-beta converting enzyme
   D. Receptor-interacting protein
   E. Ubiquitin ligase

29 A 71-year-old man diagnosed with pancreatic cancer is noted to have decreasing body mass index. His normal connective tissues undergo atrophy by sequestering organelles and cytosol in a vacuole, which then fuses with a lysosome. However, the cancer continues to increase in size. Which of the following processes is most likely occurring in the normal cells but is inhibited in the cancer cells of this man?
   A. Aging
   B. Apoptosis
   C. Autophagy
   D. Hyaline change
   E. Karyorrhexis

30 A new drug is developed that binds to cellular microtubules. The function of the microtubules is diminished, so that mitotic spindle formation is inhibited. Which of the following is the most likely use for this drug?
   A. Antimicrobial therapy
   B. Chemotherapy
   C. Pain management
   D. Prevention of atherosclerosis
   E. Weight reduction

31 A 46-year-old man has noted increasing abdominal size for the past 6 years. On physical examination his liver span is increased to 18 cm. An abdominal CT scan shows an enlarged liver with diffusely decreased attenuation. Laboratory findings include increased total serum cholesterol and triglyceride levels, increased prothrombin time, and a decreased serum albumin concentration. The representative microscopic appearance of his liver is shown in the figure. Which of the following activities most likely led to these findings?
   A. Drinking beer
   B. Ingesting aspirin
   C. Injecting heroin
   D. Playing basketball
   E. Smoking cigarettes

32 A 69-year-old woman has had transient ischemic attacks for the past 3 months. On physical examination, she has an audible bruit on auscultation of the neck. A right carotid endarterectomy is performed. The curetted atheromatous plaque has a grossly yellow-tan, firm appearance. Microscopically, which of the following materials can be found in abundance in the form of crystals within cleftlike spaces?
   A. Cholesterol
   B. Glycogen
   C. Hemosiderin
   D. Immunoglobulin
   E. Lipofuscin

33 A 45-year-old woman has had worsening dyspnea for the past 5 years. A chest CT scan shows panlobular emphysema. Laboratory studies show a deficiency of α1-antitrypsin (AAT). Her AAT genotype is PiZZ. A liver biopsy specimen examined microscopically shows abundant PAS-positive globules within periportal hepatocytes. Which of the following molecular mechanisms is most likely responsible for this finding in her hepatocytes?
   A. Decreased catabolism of AAT in lysosomes
   B. Excessive hepatic synthesis of AAT
   C. Impaired dissociation of AAT from chaperones
   D. Inability to metabolize AAT in Kupffer cells
   E. Retained misfolded AAT in endoplasmic reticulum
34. At autopsy, the heart of a 63-year-old man weighs only 250 g (normal 330 g) and has small right and left ventricles. The myocardium is firm, with a dark chocolate-brown color throughout. The coronary arteries show minimal atherosclerotic changes. An excessive amount of which of the following substances, shown in the figure, would most likely be found in the myocardial fibers of this heart?

- A Bilirubin
- B Glycogen
- C Hemosiderin
- D Lipofuscin
- E Melanin

35. A 69-year-old woman has had a chronic cough for the past year. A chest radiograph shows a 6-cm mass in the left lung. A needle biopsy specimen of the mass shows carcinoma. A pneumonectomy is performed, and examination of the hilar lymph nodes reveals a uniform, dark black cut surface. Which of the following factors most likely accounts for the appearance of these lymph nodes?

- A Aging effects
- B Bleeding disorder
- C Cigarette smoking
- D Liver failure
- E Multiple metastases

36. A 22-year-old woman from Albania has a congenital anemia requiring multiple transfusions of RBCs for many years. On physical examination, her skin has a bronze color. Liver function tests show reduced serum albumin. Which of the following findings would most likely appear in a liver biopsy specimen?

- A Amyloid in portal triads
- B Bilirubin in canaliculi
- C Glycogen in hepatocytes
- D Hemosiderin in hepatocytes
- E Steatosis in hepatocytes

37. A 72-year-old man died suddenly from congestive heart failure. At autopsy, his heart weighed 580 g (normal 330 g) and showed marked left ventricular hypertrophy and minimal coronary arterial atherosclerosis. A serum chemistry panel ordered before death showed no abnormalities. Which of the following pathologic processes best accounts for the appearance of the aortic valve seen in the figure?

- A Amyloidosis
- B Dystrophic calcification
- C Hemosiderosis
- D Hyaline change
- E Lipofuscin deposition

38. A 70-year-old man with hypercalcemia died suddenly. At autopsy, microscopic examination showed noncrystalline amorphous deposits of calcium salts in gastric mucosa, renal interstitium, and alveolar walls of lungs. Which of the following underlying conditions would most likely explain these findings?

- A Chronic active hepatitis
- B Diffuse parathyroid hyperplasia
- C Disseminated tuberculosis
- D Generalized atherosclerosis
- E Normal aging process
- F Pulmonary emphysema

39. An experiment analyzes cells for enzyme activity associated with sustained cellular proliferation. Which of the following cells is most likely to have the highest telomerase activity?

- A Endothelial cells
- B Erythrocytes
- C Germ cells
- D Neurons
- E Neutrophils

40. A study of aging shows that senescent cells have accumulated damage from toxic byproducts of metabolism. There is increased intracellular lipofuscin deposition. Prolonged ingestion of which of the following substances is most likely to counteract this aging mechanism?

- A Antioxidants
- B Analgesics
- C Antimicrobials
- D Antineoplastic agents
- E Glucocorticoids
when the liver has attained its normal size. Hepatocytes can reenter the cell cycle and proliferate to regenerate the liver; they do not just hypertrophy (increase in size). Apoptosis is single cell death and frequently occurs with viral hepatitis. Dysplasia is disordered epithelial cell growth that can be premalignant. Hydropic change, or cell swelling, does not produce regeneration. Steatosis (fatty change) can lead to hepatomegaly, but not as a regenerative process. It is the result of toxic/metabolic hepatocyte injury.

**ANSWERS**

1 A Normal cells handle physiologic demands and maintain metabolic functions within narrow ranges, termed *homeostasis*. Under disease conditions with stress on cells, there is adaptation to a new steady state. In this case, the loss of renal function leads to a higher urea nitrogen level as well as retention of fluid. The diuretic induces loss of the excess fluid to yield a new steady state. The protein restriction reduces urea nitrogen excretion, which also leads to a new steady state. Both are adaptations. Apoptosis refers to single cell necrosis in response to injury. An irreversible injury leads to cell death, but the changes described here are not evidence for cellular necrosis. The metabolism of cells is maintained for adaptation, with response to the diuretic and to protein restriction.

2 F The pressure load on the left ventricle results in an increase in myofilaments in the existing myofibers, so they enlarge. The result of continued stress from hypertension is eventual heart failure with decreased contractility. Apoptosis would lead to loss of cells and diminished size. Dysplasia is not a diagnosis made for the heart. Hemosiderin deposition in the heart is a pathologic process resulting from increased iron stores in the body. Though hyperplasia from proliferation of myofibroblasts is possible, this does not contribute significantly to cardiac size. Metaplasia of muscle does not occur, although loss of muscle occurs with aging and ischemia as myofibers are replaced by fibrous tissue.

3 D The increase in uterine size is primarily the result of an increase in the size of myometrial smooth muscle cells. The endometrium also increases in size, mainly via hyperplasia, but it remains as a thin lining to the muscular wall and does not contribute as much to the change in size. There is little stroma in myometrium and a greater proportion in endometrium, so stroma contributes a smaller percentage to the gain in size than muscle. The vessels are a minor but essential component in this increase in size, but not the largest component.

4 D Breast lobules have an increased number of cells under hormonal influence (mainly progesterone) to provide for normal lactation. Ductal metaplasia in the breast is a pathologic process. Epithelial dysplasia denotes disordered growth and maturation of epithelial cells that may progress to cancer. Accumulation of fat within cells is a common manifestation of sublethal cell injury or, uncommonly, of inborn errors in fat metabolism. The breast stroma plays no role in lactation and may increase with pathologic processes.

5 C The liver is one of the few organs in the human body that can partially regenerate. This is a form of compensatory hyperplasia. The stimuli to hepatocyte mitotic activity cease

6 D Nodular prostatic hyperplasia (also known as benign prostatic hyperplasia [BPH]) is a common condition in older men that results from proliferation of both prostatic glands and stroma. The prostate becomes more sensitive to androgenic stimulation with age. This is an example of pathologic hyperplasia. Apoptosis results in a loss of, not an increase in, cells. Dysplasia refers to disordered epithelial cell growth and maturation. Fatty change in hepatocytes may produce hepatomegaly. Although BPH is often called “benign prostatic hypertrophy,” this term is technically incorrect; it is the number of glands and stromal cells that is increased, rather than the size of existing cells. A change in the glandular epithelium to squamous epithelium around a prostatic infarct would be an example of metaplasia.

7 B Reduced workload causes cell to shrink through loss of cell substance, a process called *atrophy*. The cells are still present, just smaller. Aplasia refers to lack of embryonic development; hypoplasia describes poor or subnormal development of tissues. Dystrophy of muscles refers to inherited disorders of skeletal muscles that lead to muscle fiber destruction, weakness, and wasting. Hyaline change (hyalinosis) refers to a nonspecific, pink, glassy eosinophilic appearance of cells.

8 A Inflammation from reflux of gastric acid has resulted in replacement of normal esophageal squamous epithelium by intestinal-type columnar epithelium with goblet cells. Such conversion of one adult cell type to another cell type is called *metaplasia*, and it occurs when stimuli reprogram stem cells. Goblet cells are not normal constituents of the esophageal mucosa, and they are a minor part of this metaplastic process. The lamina propria has some inflammatory cells, but it does not atrophy. The squamous epithelium does not become dysplastic from acid reflux, but the columnar metaplasia may progress to dysplasia, not seen here, if the abnormal stimuli continue. These cells are not significantly increased in size (hypertrophic).

9 A Irreversible cell injury is associated with loss of membrane integrity. This allows intracellular enzymes such as AST and ALT to leak into the serum. All other morphologic
UNIT I  General Pathology

changes listed are associated with reversible cell injury, in which the cell membrane remains intact and the cells do not die.

PBD9 38–39  BP9 7–8  PBD8 11–12  BP8 8–10

10 C Cell death occurs with loss of the cell nucleus, and tubular cells become necrotic. All other cellular morphologic changes listed represent forms of reversible cellular injury. The plasma membrane and intracellular organelles retain some function unless severe damage causes loss of membrane integrity.

PBD9 39, 42  BP9 8–9  PBD8 12  BP8 6, 9

11 A The figure shows deep eosinophilic staining, loss of myocardial fiber nuclei, and loss of cell structure consistent with an early ischemic injury, resulting in coagulative necrosis. Myocardial ischemia and infarction are typically caused by loss of coronary arterial blood flow. An immunological process may produce focal myocardial injury. Blunt trauma produces hemorrhage. Lack of protein leads to a catabolic state with gradual decrease in cell size, but it does not cause ischemic changes. Viral infection could cause focal necrosis of the myocardium, but this is usually accompanied by an inflammatory infiltrate consisting of lymphocytes and macrophages.

PBD9 42–43  BP9 9–10  PBD8 15–16  BP8 2, 7, 10

12 A The many focal, chalky white deposits in the mesentery, composed mainly of adipocytes, are areas of fat necrosis. The deposits result from the release of pancreatic enzymes such as lipases in a patient with acute pancreatitis. Gangrenous necrosis is mainly coagulative necrosis, but occurs over an extensive area of tissues. Viral hepatitis does not cause cell necrosis in organs other than liver, and hepatocyte necrosis from viral infections occurs mainly by means of apoptosis. Intestinal infarction is a form of coagulative necrosis. Infection with tuberculosis leads to caseous necrosis.

PBD9 43–44  BP9 10–11  PBD8 16–17  BP8 11

13 E The high lipid content of central nervous system tissues results in liquefactive necrosis as a consequence of ischemic injury, as in this case of stroke. Apoptosis affects single cells and typically is not grossly visible. Coagulative necrosis is the typical result of ischemia in most solid organs. Fat necrosis is seen in breast and pancreatic tissues. Karyolysis refers to fading away of cell nuclei in dead cells.

PBD9 43  BP9 10–11  PBD8 16–17  BP8 10–11

14 B The grossly cheeselike appearance gives this form of necrosis its name—caseous necrosis. The figure shows amorphous pink acellular material at the upper right surrounded by epithelioid macrophages, and a Langhans giant cell is visible at the upper left. In the lung, tuberculosis and fungal infections are most likely to produce this pattern of tissue injury. Apoptosis involves individual cells, without grossly apparent extensive or localized areas of tissue necrosis.

Coagulative necrosis is more typical of ischemic tissue injury. Fat necrosis most often occurs in the breast and pancreas. Fatty change is most often a feature of hepatocyte injury, and the cell integrity is maintained. Gangrene characterizes extensive necrosis of multiple cell types in a body region or organ. Liquefactive necrosis is seen in neutrophilic abscesses or ischemic cerebral injury.

PBD9 43–44  BP9 10–11  PBD8 16  BP8 10

15 D Reduction in oxidative phosphorylation leads to reduction in synthesis of ATP and diminished activity of the plasma membrane sodium pump, which maintains high intracellular potassium concentration. Loss of ATP leads to efflux of intracellular potassium, while net influx of sodium and water promote cell swelling. A marked rise in plasma potassium can indicate significant cell damage or death (such as skeletal muscle crush injury or hemolysis). When cells are not consuming glucose via oxidative metabolism, the glucose is metabolized via other pathways, and glucose is maintained within normal ranges. Though cell membranes are composed of lipid, dysfunction or disruption of those membranes does not significantly alter plasma lipid concentrations.

PBD9 45–46  BP9 12–13  PBD8 14–15  BP8 14–15

16 D Decreased tissue perfusion from hypotensive shock leads to hypoxemia and depletion of ATP when cell metabolism shifts from aerobic to anaerobic glycolysis. This shift causes depletion of glycogen stores and increased production and accumulation of lactic acid, reducing intracellular pH. Creatinine would increase with reduced renal function from decreased renal perfusion, but this would not explain the changes in other tissues. An increased glucose level would be indicative of poorly controlled diabetes mellitus, not decreased perfusion. Carbon dioxide is likely to be cleared via normal lungs, which are still sufficiently perfused by a failing heart. An increase in troponin I suggests irreversible myocardial injury.

PBD9 45–46  BP9 12–13  PBD8 14–15  BP8 14, 18

17 A Irreversible cellular injury is likely to occur when cytoplasmic calcium increases. Calcium can enter cells and also accumulate in mitochondria and endoplasmic reticulum. The excess calcium activates ATPases, phospholipases, proteases, and endonucleases, which injure cell components. Mitochondrial permeability is increased to release cytochrome c, which activates caspases leading to apoptosis. Of the other ions listed, sodium enters the cell, while potassium diffuses out when the sodium pump fails as ATP levels fall; but this is potentially reversible.

PBD9 46–47  BP9 13–14  PBD8 18–20  BP8 15

18 B The drug acetaminophen can be converted to toxic metabolites in this manner. Glutathione in the cytosol helps to reduce cellular injury from many toxic metabolites and free radicals. ADP is converted to ATP by oxidative and glycolytic cellular pathways to provide energy that drives
cellular functions, and a reduction in ATP leaves the cell vulnerable to injury. NADPH oxidase generates superoxide, which is used by neutrophils in killing bacteria. Nitric oxide synthase in macrophages produces nitric oxide, which aids in destroying organisms undergoing phagocytosis. Protein synthesis in cells depends on mRNA for longer survival and recovery from damage caused by free radicals. Failure of the sodium pump leads to increased cytosolic sodium and cell swelling with injury.

**23** A The onset of menstruation is orderly, programmed cell death (apoptosis) through hormonal stimuli, an example of the intrinsic (mitochondrial) apoptotic pathway. As hormone levels drop, the endometrium breaks down, sloughs off, and then regenerates. With cellular atrophy, there is often no visible necrosis, but the tissues shrink, something that occurs in the endometrium after menopause. Caseous necrosis is typical of granulomatous inflammation, resulting most commonly from mycobacterial infection. Heterophagocytosis is typified by the clearing of an area of necrosis through macrophage ingestion of the necrotic cells. Liquefactive necrosis can occur in any tissue after acute bacterial infection or in the brain after ischemia.

**24** B These histologic findings are typical of apoptosis. The BCL2 gene product inhibits cellular apoptosis by binding to Apaf-1. Hence, the knockout removes this inhibition. The BAX gene product promotes apoptosis, and a knockout would protect against apoptosis. The C-MYC gene is involved with oncogenesis. The FAS gene encodes for a cellular receptor for Fas ligand that signals apoptosis. Activity of the p53 (TP53) gene normally stimulates apoptosis, but mutation favors cell survival.

**25** A There is an apoptotic cell (arrow) that is shrunken and has been converted into a dense eosinophilic mass. There is a surrounding inflammatory reaction with cytotoxic lymphocytes. This pattern is typical of apoptosis. Caspase activation is a universal feature of apoptosis, regardless of the initiating cause. Apoptosis induced in recipient cells from donor lymphocytes occurs with graft-versus-host disease. Lipases are activated in enzymatic fat necrosis. Reduced ATP synthesis and increased glycolysis occur when a cell is subjected to anoxia, but these changes are reversible. Lipid peroxidation occurs when the cell is injured by free radicals.
though there is cellular proliferation, it occurs over many affected by endothelial damage and lipid accumulation, and release of mediators of inflammation. Atheroma formation is do not have microtubules. Pain is produced largely through Vigor involves the release of proteolytic enzymes that cause necrosis, and maintain a survival advantage even as the patient is dying. There is slow autophagy with aging, but autophagy is accelerated with stressors such as malnutrition and chronic disease. Hyaline is a generic term for intracellular or extracellular protein accumulations appearing pink and homogeneous with H&E staining. Karyorrhexis is nuclear fragmentation in a necrotic cell. 

Lipofuscin is a “wear-and-tear” pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes. Lipofuscin is a golden brown pigment that increases with aging in cell cytoplasm, mainly in cardiac myocytes and in hepatocytes.
35 C Lung and hilar lymph nodes accumulate anthracotic pigmentation when carbon pigment is inhaled from polluted air. The tar in cigarette smoke is a major source of such carbonaceous pigment. Older individuals generally have more anthracotic pigment, but this is not inevitable with aging—individuals living in rural areas with good environmental air quality have less pigment. Resolution of hemorrhage can produce hemosiderin pigmentation, which imparts a brown color to tissues. Hepatic failure may result in jaundice, characterized by a yellow color in tissues. Metastases are mass lesions that impart a tan-to-white appearance to tissues.

PBD9 64 BP9 24 PBD8 36 BP8 25

36 D Each unit of blood contains about 250 mg of iron. The body has no mechanism for getting rid of excess iron. About 10 to 20 mg of iron per day is lost with normal desquamation of epithelia; menstruating women lose slightly more. Any excess iron becomes storage iron, or hemosiderin. Over time, hemosiderosis involves more and more tissues of the body, particularly the liver, but also skin. Initially, hemosiderin deposits are found in Kupffer cells and other mononuclear phagocytes in the bone marrow, spleen, and lymph nodes. With great excess of iron, liver cells also accumulate iron. Amyloid is an abnormal protein derived from a variety of precursors, such as immunoglobulin light chains. Bilirubin, a breakdown product of blood, can be excreted in the bile so that a person does not become jaundiced. Glycogen storage diseases are inherited and present in childhood. Steatosis usually occurs with ingestion of hepatotoxins, such as alcohol.

PBD9 64–65 BP9 24 PBD8 36 BP8 26

37 B The valve is stenotic because of nodular deposits of calcium. The process is “dystrophic” because calcium deposition occurs in damaged tissues. The damage in this patient is a result of excessive wear and tear with aging. Amyloid deposition in the heart typically occurs within the myocardium and the vessels. Hereditary hemochromatosis is a genetic defect in iron absorption that results in extensive myocardial iron deposition (hemosiderosis). Hyaline change is a descriptive term used by histologists to describe protein deposits that are glassy and pale pink. The amount of lipofuscin increases within myocardial fibers (not valves) with aging.

PBD9 65 BP9 25–26 PBD8 38 BP8 26–27

38 B The microscopic findings suggest metastatic calcification, with deposition of calcium salts in tissues that have physiologic mechanisms for losing acid, creating an internal alkaline environment that favors calcium precipitation. Hypercalcemia can have a variety of causes, including primary and secondary hyperparathyroidism, bone destruction secondary to metastases, paraneoplastic syndromes, and, less commonly, vitamin D toxicity or sarcoidosis. Chronic renal disease reduces phosphate excretion by the kidney, resulting in an increase in serum phosphate. Because the solubility product of calcium and phosphorus must be maintained, the serum calcium is depressed, triggering increased parathyroid hormone output to increase the calcium level, which promotes calcium deposition. Chronic hepatitis leads to hyperbilirubinemia and jaundice. The granulomas of tuberculosis have caseous necrosis with dystrophic calcification. Another form of dystrophic calcification occurs when atherosclerotic lesions calcify. Dystrophic calcification is seen more often in the elderly, but it is the result of a lifetime of pathologic changes, not aging itself. Pulmonary emphysema can lead to respiratory acidosis that is compensated by metabolic alkalosis, with the result that the serum calcium level remains relatively unchanged.

PBD9 65 BP9 25–26 PBD8 38–39 BP8 26–27

39 C Germ cells have the highest telomerase activity, and the telomere length can be stabilized in these cells. This allows testicular germ cells to retain the ability to divide throughout life. Normal somatic cells have no telomerase activity, and telomeres progressively shorten with each cell division until growth arrest occurs. Erythrocytes do not even have a nucleus.

PBD9 67 BP9 26–27 PBD8 39–40 BP8 28–29

40 A Antioxidants may counteract the effects of reactive oxygen species (ROS) that may accumulate acutely and chronically within cells as a consequence of environmental insults and pathologic processes. Certainly, health food stores promote this concept with sales of products such as vitamin E. However, cellular damage is multifactorial, and proving that one compound has a significant effect is difficult. Analgesics ameliorate the perception of pain from cellular damage, but they do not prevent or diminish cell damage; they only mask it. Antimicrobials may help the body’s own immune defenses against infectious agents and shorten and/or diminish tissue damage. However, long-term use of antimicrobials is discouraged because it may alter the body’s own useful microbial flora, and it can promote development of drug-resistant strains that pose a serious health risk for the general population. (As Mr. Spock noted, “The needs of the many outweigh the needs of the few.”) Antineoplastic agents are given for malignancies and rarely have benefit for cancer prevention. Glucocorticoids provide short-term improvement in well-being, but when used for longer periods, they have deleterious effects.

PBD9 66–67 BP9 26–27 PBD8 40–41 BP8 28–29
1. An 11-year-old child falls and cuts his hand. The wound becomes infected. Bacteria extend into the extracellular matrix around capillaries. In the inflammatory response to this infection, which of the following cells removes the bacteria?

A. B lymphocyte  
B. Fibroblast  
C. Macrophage  
D. Mast cell  
E. T lymphocyte

2. A 53-year-old woman has had a high fever and cough productive of yellowish sputum for the past 2 days. Her vital signs include temperature of 37.8° C, pulse 103/min, respirations 25/min, and blood pressure 100/60 mm Hg. On auscultation of the chest, crackles are audible in both lung bases. A chest radiograph shows bilateral patchy pulmonary infiltrates. The microscopic appearance of her lung is shown in the figure. Which of the following inflammatory cell types is most likely to be seen in greatly increased numbers in her sputum specimen?

A. Langhans giant cells  
B. Macrophages  
C. Mast cells  
D. Neutrophils  
E. T lymphocytes

3. A 4-year-old child has had a high-volume diarrhea for the past 2 days. On examination she is dehydrated. A stool sample examined by serologic assay is positive for rotavirus. She is treated with intravenous fluids and recovers. Which of the following components is found on intestinal cells and recognizes double-stranded RNA of this virus to signal transcription factors that upregulate interferon production for viral elimination?

A. Caspase-1  
B. Complement receptor  
C. Lectin  
D. T cell receptor  
E. Toll-like receptor
4. A 72-year-old man with severe emphysema has had worsening right ventricular failure for the past 5 years. For the past 4 days, he has had fever and increasing dyspnea. A chest radiograph shows an accumulation of fluid in the pleural spaces. Fluid obtained by thoracentesis has a specific gravity of 1.030 and contains degenerating neutrophils. The most likely cause of this fluid accumulation is due to changes in which of the following?
   A. Colloid osmotic pressure
   B. Leukocytic diapedesis
   C. Lymphatic pressure
   D. Renal sodium retention
   E. Vascular permeability

5. A 35-year-old man has had increasing dyspnea for the past 24 hours. A chest radiograph shows large, bilateral pleural effusions. Thoracentesis yields 500 mL of slightly cloudy yellow fluid from the right pleural cavity. Cytologic examination of the fluid shows many neutrophils, but no lymphocytes or RBCs. Which of the following mechanisms contributes most to the pleural fluid accumulation?
   A. Arteriovenous vasoconstriction
   B. Endothelial contraction
   C. Inhibition of platelet adherence
   D. Lymphatic obstruction
   E. Neutrophil release of lysosomes

6. A 6-year-old child has a history of recurrent infections with pyogenic bacteria, including *Staphylococcus aureus* and *Streptococcus pneumoniae*. The infections are accompanied by a neutrophilic leukocytosis. Microscopic examination of a biopsy specimen obtained from an area of soft tissue necrosis shows microbial organisms, but very few neutrophils. An analysis of neutrophil function shows a defect in rolling. This child’s increased susceptibility to infection is most likely caused by a defect involving which of the following molecules?
   A. Complement C3b
   B. Integrins
   C. Leukotriene B4
   D. NADPH oxidase
   E. Selectins

7. In an experiment, bacteria are introduced into a perfused tissue preparation. Leukocytes leave the vasculature and migrate to the site of bacterial inoculation. The movement of these leukocytes is most likely to be mediated by which of the following substances?
   A. Bradykinin
   B. Chemokines
   C. Complement C3a
   D. Histamine
   E. Prostaglandins

8. A 12-month-old boy with a 6-month history of repeated infections has had a fever and cough for the past 3 days. A Gram stain of sputum shows many gram-positive cocci in chains. CBC shows neutrophilia. Laboratory studies show that the patient’s neutrophils phagocytose and kill organisms promptly in the presence of normal human serum, but not in his own serum. The neutrophils migrate normally in a chemotaxis assay. Which of the following is the most likely cause of this boy’s increased susceptibility to infection?
   A. Abnormality of selectin expression
   B. Diminished opsonization
   C. Defective neutrophil generation of hydrogen peroxide
   D. Deficiency of integrins
   E. Phagocytic cell microtubular protein defect

9. A 5-year-old child has a history of recurrent bacterial infections, including pneumonia and otitis media. Analysis of leukocytes collected from the peripheral blood shows a deficiency in myeloperoxidase. A reduction in which of the following processes is the most likely cause of this child’s increased susceptibility to infections?
   A. Hydrogen peroxide (H2O2) elaboration
   B. Hydroxy-halide radical (HOCl) formation
   C. Failure of migration resulting from complement deficiency
   D. Phagocytic cell oxygen consumption
   E. Prostaglandin production

10. In an experiment, neutrophils collected from peripheral blood are analyzed for a “burst” of oxygen consumption. This respiratory burst is an essential step for which of the following events in an acute inflammatory response?
    A. Attachment to endothelial cells
    B. Generation of microbicidal activity
    C. Increased production in bone marrow
    D. Opsonization of bacteria
    E. Phagocytosis of bacteria

11. A 4-year-old girl has had numerous infections with *Staphylococcus aureus* since infancy. Genetic testing shows a defect leading to a lack of β2 integrin production. Which of the following abnormalities of neutrophil function is most likely responsible for these clinical symptoms?
    A. Decreased generation of hydroxy-halide radicals (HOCl)
    B. Diminished phagocytosis of bacteria opsonized with IgG
    C. Failure of migration to the site of infection
    D. Inadequate adhesion on cytokine-activated endothelium
    E. Reduced respiratory burst after phagocytosis

12. In an experiment, peripheral blood cells are isolated and placed into a culture medium that preserves their metabolic activity. Interferon-γ is added to this culture, along with viable *Escherichia coli* organisms. Which of the following blood cell types in this medium is the most likely to have bactericidal activity against *E. coli*?
    A. Basophil
    B. B lymphocyte
    C. CD4+ lymphocyte
    D. CD8+ lymphocyte
    E. Monocyte
    F. Natural killer cell
    G. Neutrophil
13 | In an experiment, T lymphocytes from peripheral blood are placed in a medium that preserves their function. The lymphocytes are activated by contact with antigen and incubated for 4 hours. The supernatant fluid is collected and is found to contain a substance that is a major stimulator of monocytes and macrophages. Which of the following substances released into this fluid medium is most likely to stimulate macrophages?

A  Histamine  
B  Interferon-γ  
C  Leukotriene B₄  
D  Nitric oxide  
E  Phospholipase C  
F  Tumor necrosis factor (TNF)

14 | A woman who is allergic to cats visits a neighbor who has several cats. During the visit, she inhales cat dander, and within minutes, she develops nasal congestion with abundant nasal secretions. Which of the following substances is most likely to produce these findings?

A  Bradykinin  
B  Complement C₅a  
C  Histamine  
D  Interleukin-1 (IL-1)  
E  Phospholipase C  
F  Tumor necrosis factor (TNF)

15 | In a 6-month randomized trial of a pharmacologic agent, one group of patients receives a cyclooxygenase-2 (COX-2) inhibitor, and a control group does not. Both groups of adult males had mild congestive heart failure and bilateral symmetric arthritis of small joints. Laboratory measurements during the trial show no significant differences between the groups in WBC count, platelet count, hemoglobin, and creatinine. The group receiving the drug reports subjective findings different from those of the control group. Which of the following findings was most likely reported by the group receiving the drug?

A  Increased ankle swelling  
B  Increased susceptibility to bruising  
C  Increased bouts of asthma  
D  Reduced severity of urticaria  
E  Numerous febrile episodes  
F  Reduced arthritis pain

16 | A 19-year-old woman develops a sore throat and fever during the past day. Physical examination shows pharyngeal erythema and swelling. Laboratory findings include leukocytosis. She is given naproxen. Which of the following features of the acute inflammatory response is most affected by this drug?

A  Chemotaxis  
B  Emigration  
C  Leukocytosis  
D  Phagocytosis  
E  Vasodilation

17 | A 35-year-old woman takes acetylsalicylic acid (aspirin) for arthritis. Although her joint pain is reduced with this therapy, the inflammatory process continues. The aspirin therapy alleviates her pain mainly through reduction in the synthesis of which of the following mediators?

A  Complement C₁q  
B  Histamine  
C  Leukotriene E₄  
D  Nitric oxide  
E  Prostaglandins

18 | A 77-year-old woman experiences a sudden loss of consciousness, with loss of movement on the right side of the body. Cerebral angiography shows an occlusion of the left middle cerebral artery. Elaboration of which of the following mediators will be most beneficial in preventing further ischemic injury to her cerebral cortex?

A  Bradykinin  
B  Leukotriene E₄  
C  Nitric oxide  
D  Platelet-activating factor  
E  Thromboxane A₂

19 | In an experiment, bacteria are inoculated into aliquots of normal human blood that have been treated with an anticoagulant. It is observed that the bacteria are either phagocytized by neutrophils or undergo lysis. Which of the following blood plasma components is most likely to facilitate these effects?

A  Complement  
B  Fibrin  
C  Kallikrein  
D  Plasmin  
E  Thrombin

20 | Patients with extensive endothelial injury from *Escherichia coli* sepsis have consumption of coagulation factors as well as an extensive inflammatory response. Administration of activated protein C is most likely to decrease this inflammatory response by reducing the amount of which of the following substances?

A  Complement  
B  Fibrin  
C  Kallikrein  
D  Plasmin  
E  Thrombin

21 | A 95-year-old woman touches a pot of boiling water. Within 2 hours, she has marked erythema of the skin of the fingers of her hand, and small blisters appear on the finger pads. This has led to which one of the following inflammatory responses?

A  Fibrinous inflammation  
B  Granulomatous inflammation  
C  Purulent inflammation  
D  Serous inflammation  
E  Ulceration

22 | A 24-year-old, sexually active woman has experienced lower abdominal pain for the past day. Her temperature is 37.9°C, and on palpation, the left lower abdomen is markedly tender. Laboratory findings include a total WBC count of 29,000/mm³ with 75% segmented neutrophils, 6% bands, 14% lymphocytes, and 5% monocytes. Laparotomy reveals a distended, fluid-filled, reddened left fallopian tube that is about to rupture. A left salpingectomy is performed. Which of the following is most likely to be seen on microscopic examination of the excised fallopian tube?

A  Fibroblastic proliferation  
B  Langhans giant cells  
C  Liquefactive necrosis  
D  Mononuclear infiltrates  
E  Squamous metaplasia
23 A 68-year-old man has had worsening shortness of breath for the past week. On physical examination, his temperature is 38.3°C. On percussion, there is dullness over the left lung fields. Thoracentesis performed on the left pleural cavity yields 800 mL of cloudy yellow fluid that has a WBC count of 2500/mm³ with 98% neutrophils and 2% lymphocytes. A Gram stain of the fluid shows gram-positive cocci in clusters. Which of the following terms best describes the process occurring in his left pleural cavity?

A Abscess  
B Chronic inflammation  
C Edema  
D Fibrinous inflammation  
E Purulent exudate  
F Serous effusion

24 An 87-year-old woman has had a cough productive of yellowish sputum for the past 2 days. On examination her temperature is 37°C. A chest radiograph shows bilateral patchy infiltrates. Her peripheral blood shows leukocytosis. A week later she is afebrile. Which of the following is the most likely outcome of her pulmonary disease?

A Chronic inflammation  
B Fibrous scarring  
C Neoplasia  
D Resolution  
E Ulceration

25 A 53-year-old woman has experienced abdominal pain for 2 weeks. She is afebrile. There is mild upper abdominal tenderness on palpation, and bowel sounds are present. An upper gastrointestinal endoscopy is performed. The figure shows microscopic examination of a biopsy specimen of a duodenal lesion. Which of the following pathologic processes is most likely present?

A Abscess  
B Caseating granuloma  
C Chronic inflammation  
D Purulent exudate  
E Serous effusion  
F Ulceration

26 A 92-year-old woman is diagnosed with Staphylococcus aureus pneumonia and receives a course of antibiotic therapy. Two weeks later, she no longer has a productive cough, but she still has a temperature of 38.1°C. A chest radiograph shows the findings in the figure. Which of the following terms best describes the outcome of the patient’s pneumonia?

A Abscess formation  
B Complete resolution  
C Fibrous scarring  
D Chronic inflammation  
E Tissue regeneration

27 A 29-year-old woman with a congenital ventricular septal defect has had a persistent temperature of 38.6°C and headache for the past 3 weeks. A head CT scan shows an enhancing 3-cm, ring like lesion in the right parietal lobe of her brain. Which of the following actions by inflammatory cells has most likely produced this CT finding?

A Elaboration of nitric oxide by macrophages  
B Formation of immunoglobulin by B lymphocytes  
C Generation of prostaglandin by endothelium  
D Production of interferon-γ by T lymphocytes  
E Release of lysosomal enzymes from neutrophils

28 A 37-year-old man has had midepigastric pain for the past 3 months. An upper gastrointestinal endoscopy shows a 2-cm, sharply demarcated, shallow ulceration of the gastric antrum. Microscopic examination of a biopsy from the ulcer base shows angiogenesis, fibrosis, and mononuclear cell infiltrates with lymphocytes, macrophages, and plasma cells. Which of the following terms best describes this pathologic process?

A Acute inflammation  
B Chronic inflammation  
C Fibrinous inflammation  
D Granulomatous inflammation  
E Serous inflammation
29. A 65-year-old man develops worsening congestive heart failure 2 weeks after an acute myocardial infarction. An echocardiogram shows a markedly decreased ejection fraction. Now, capillaries, fibroblasts, collagen, and inflammatory cells have largely replaced the infarcted myocardium. Which of the following inflammatory cell types in this lesion plays the most important role in the healing process?

A. Eosinophils
B. Epithelioid cells
C. Macrophages
D. Neutrophils
E. Plasma cells

30. A 9-year-old boy has had a chronic cough and fever for the past month. A chest radiograph shows enlargement of hilar lymph nodes and bilateral pulmonary nodular interstitial infiltrates. A sputum sample contains acid-fast bacilli. A transbronchial biopsy specimen shows granulomatous inflammation with epithelioid macrophages and Langhans giant cells. Which of the following mediators is most likely to contribute to giant cell formation?

A. Complement C3b
B. Interferon-γ
C. Interleukin-1 (IL-1)
D. Leukotriene B4
E. Tumor necrosis factor (TNF)

31. A 32-year-old woman has had a chronic cough with fever for the past month. On physical examination, her temperature is 37.5° C. A chest radiograph shows many small, ill-defined nodular opacities in all lung fields. A transbronchial biopsy specimen shows interstitial infiltrates with lymphocytes, plasma cells, and epithelioid macrophages. Which of the following infectious agents is the most likely cause of this appearance?

A. Candida albicans
B. Cytomegalovirus
C. Enterobacter aerogenes
D. Mycobacterium tuberculosis
E. Plasmodium falciparum
F. Staphylococcus aureus

32. One month after an appendectomy, a 25-year-old woman palpates a small nodule beneath the skin at the site of the healed right lower quadrant sutured incision. The nodule is excised, and microscopic examination shows macrophages, collagen deposition, small lymphocytes, and multinucleated giant cells. Polarizable, refractile material is seen in the nodule. Which of the following complications of the surgery best accounts for these findings?

A. Abscess formation
B. Chronic inflammation
C. Exuberant granulation tissue
D. Granuloma formation
E. Healing by second intention

33. A 43-year-old man has had a cough and fever for the past 2 months. A chest CT scan shows the findings in the figure (A). A transbronchial lung biopsy is performed, yielding a specimen with the microscopic appearance shown in the figure (B). Which of the following chemical mediators is most important in the pathogenesis of this lesion?

A. Bradykinin
B. Complement C5a
C. Interferon-γ
D. Nitric oxide
E. Prostaglandins

34. An 8-year-old girl has had difficulty swallowing for the past day. On examination, her pharynx is swollen and erythematous with an overlying yellow exudate. Laboratory studies show neutrophilia. Streptococcus pyogenes (group A streptococcus) is cultured from her pharynx. Which of the following substances is most likely to increase in response to pyrogens released by this organism?

A. Hageman factor
B. Immunoglobulin E
C. Interleukin-12 (IL-12)
D. Nitric oxide
E. Prostaglandins
35 A 41-year-old man has had a severe headache for the past 2 days. On examination, his temperature is 39.2°C. A lumbar puncture is performed, and the cerebrospinal fluid obtained has a WBC count of 910/mm³ with 94% neutrophils and 6% lymphocytes. Which of the following substances is the most likely mediator for the fever observed in this man?

A Bradykinin  
B Histamine  
C Leukotriene B₄  
D Nitric oxide  
E Tumor necrosis factor (TNF)

36 A 43-year-old man with a ventricular septal defect has had a cough and fever for the past 2 days. On examination, he has a temperature of 37.6°C and a cardiac murmur. A blood culture grows *Streptococcus*, viridans group. His erythrocyte sedimentation rate (ESR) is increased. Microbial cells are opsonized and cleared. Which of the following chemical mediators is most important in producing these findings?

A Bradykinin  
B C-reactive protein  
C Interferon-γ  
D Nitric oxide  
E Prostaglandin  
F Tumor necrosis factor (TNF)

37 In an experiment, a group of test animals is infected with viral hepatitis. Two months later, complete recovery of the normal liver architecture is observed microscopically. A control test group is infected with bacterial organisms, and after the same period of time, fibrous scars from resolving hepatic abscesses are seen microscopically. Which of the following factors best explains the different outcomes for the two test groups?

A Extent of damage to the biliary ducts  
B Extent of the hepatocyte injury  
C Injury to the connective tissue framework  
D Location of the lesion within the liver  
E Nature of the injurious etiologic agent

38 A 51-year-old woman tests positive for hepatitis A antibody. Her serum AST level is 275 U/L, and ALT is 310 U/L. One month later, these enzyme levels have returned to normal. Which phase of the cell cycle best describes the hepatocytes 1 month after her infection?

A G₀  
B G₁  
C S  
D G₂  
E M

39 A 54-year-old man undergoes laparoscopic hernia repair. In spite of the small size of the incisions, he has poor wound healing. Further history reveals that his usual diet has poor nutritional value and is deficient in vitamin C. Synthesis of which of the following extracellular matrix components is most affected by this deficiency?

A Collagen  
B Elastin  
C Fibronectin  
D Integrin  
E Laminin

40 In an experiment, glass beads are embolized into the coronary arteries of rats, resulting in myocardial injury. After 7 days, sections of the myocardium are studied using light microscopy. The microscopic appearance of one of these sections is shown in the figure. Which of the following mediators is most likely being expressed to produce this appearance?

A Epidermal growth factor  
B Interleukin-2 (IL-2)  
C Leukotriene B₄  
D Thromboxane A₂  
E Tumor necrosis factor (TNF)  
F Vascular endothelial growth factor

41 A 20-year-old woman undergoes cesarean section to deliver a term infant, and the lower abdominal incision is sutured. The sutures are removed 1 week later. Which of the following statements best describes the wound site at the time of suture removal?

A Collagen degradation exceeds synthesis  
B Granulation tissue is still present  
C No more wound strength will be gained  
D Type IV collagen predominates  
E Wound strength is 80% of normal tissue
A 24-year-old man with acute appendicitis undergoes surgical removal of the inflamed appendix. The incision site is sutured. A trichrome-stained section representative of the site with blue appearing collagen is shown in the figure. How long after the surgery would this appearance most likely be seen?

A  1 day  
B  2 to 3 days  
C  4 to 5 days  
D  2 weeks  
E  1 month

43 A 40-year-old man underwent laparotomy for a perforated sigmoid colon diverticulum. A wound infection complicated the postoperative course, and surgical wound dehiscence occurred. Primary closure was no longer possible, and the wound “granulated in.” Six weeks later, the wound is only 10% of its original size. Which of the following processes best accounts for the observed decrease in wound size over the past 6 weeks?

A  Elaboration of adhesive glycoproteins  
B  Increase in synthesis of collagen  
C  Inhibition of metalloproteinases  
D  Myofibroblast contraction  
E  Resolution of subcutaneous edema

44 In an experiment involving observations on wound healing, researchers noted that intracytoplasmic cytoskeletal elements, including actin, interact with the extracellular matrix to promote cell attachment and migration in wound healing. Which of the following substances is most likely responsible for such interaction between the cytoskeleton and the extracellular matrix?

A  Epidermal growth factor  
B  Fibronectin  
C  Integulin  
D  Platelet-derived growth factor  
E  Type IV collagen  
F  Vascular endothelial growth factor

45 A 23-year-old woman receiving corticosteroid therapy for an autoimmune disease has an abscess on her upper outer right arm. She undergoes minor surgery to incise and drain the abscess, but the wound heals poorly over the next month. Which of the following aspects of wound healing is most likely to be deficient in this patient?

A  Collagen deposition  
B  Elaboration of VEGF  
C  Neutrophil infiltration  
D  Reepithelialization  
E  Serine proteinase production

46 An 18-year-old man lacerated his left ear and required sutures. The sutures were removed 1 week later. Wound healing continued, but the site became disfigured over the next 2 months by the process shown in the figure. Which of the following terms best describes the process that occurred in this man?

A  Dehiscence  
B  Keloid formation  
C  Organization  
D  Resolution  
E  Secondary union

47 A 58-year-old man had chest pain persisting for 4 hours. A radiographic imaging procedure showed an infarction involving a 4-cm area of the posterior left ventricular free wall. Laboratory findings showed serum creatine kinase of 600 U/L. Which of the following pathologic findings would most likely be seen in the left ventricular lesion 1 month later?

A  Chronic inflammation  
B  Coagulative necrosis  
C  Complete resolution  
D  Fibrous scar  
E  Nodular regeneration
Sodium and water retention helps drive transudation of fluid. The lymphatics scavenge exuded proteinaceous fluid and driven to emigrate from the vessels by chemotactic factors. Leukocytosis alone increases, and extravascular colloid osmotic pressure increases, incurred, the protein content of the extravascular space in -

**ANSWERS**

1. **C** Macrophages in tissues derived from circulating blood monocytes are phagocytic cells that respond to a variety of stimuli, and they represent the janitorial crew of the body. The other cells listed are not phagocytes. B cells can differentiate into plasma cells secreting antibodies to neutralize infectious agents. Fibroblasts form collagen as part of a healing response. Mast cells can release a variety of inflammatory mediators. T cells are a key part of chronic inflammatory processes in cell-mediated immune responses.

2. **D** These signs and symptoms suggest acute bacterial pneumonia. Such infections induce an acute inflammation dominated by neutrophils that fill alveoli, as shown in the figure, and are coughed up, which gives the sputum its yellowish, purulent appearance. Langhans giant cells are seen with granulomatous inflammatory responses. Macrophages become more numerous after initiation of acute events, cleaning up tissue and bacterial debris through phagocytosis. Mast cells are better known as participants in allergic and anaphylactic responses. Lymphocytes are a feature of chronic inflammation.

3. **E** Nonhuman microbial substances such as double-stranded RNA of viruses, bacterial DNA, and bacterial endotoxin, can be recognized by Toll-like receptors (TLRs) on human cells as part of an innate defense mechanism against infection. Caspase-1 is activated by an inflammasome complex of proteins responding to bacterial organisms, and produces biologically active interleukin-1 (IL-1). Complement receptors on inflammatory cells recognize complement components that aid in triggering immune responses through co-stimulatory signals. Lectins found on cell surfaces can bind a variety of substances, such as fungal polysaccharides, that trigger cellular defenses. T cell receptors respond to peptide antigens to trigger a cell-mediated immune response.

4. **E** The formation of an exudate containing a significant amount of protein and cells depends on the “leakiness” of blood vessels, principally venules. When exudation has occurred, the protein content of the extravascular space increases, and extravascular colloid osmotic pressure increases, causing extracellular fluid accumulation. Leukocytosis alone is insufficient for exudation because the leukocytes must be driven to emigrate from the vessels by chemotactic factors. The lymphatics scavenge exuded proteinaceous fluid and reduce the amount of extracellular and extracellular fluid. Sodium and water retention helps drive transudation of fluid.

5. **B** Exudation of fluid from venules and capillaries is a key component of the acute inflammatory process. Several mechanisms of increased vascular permeability have been proposed, including formation of interendothelial gaps by contraction of endothelium. This contraction can be caused by mediators such as histamine and leukotrienes. The vessels then become more “leaky,” and the fluid leaves the intravascular space to accumulate extravascularly, forming effusions in body cavities or edema within tissues. Arteriolar vasoconstriction is a transient response to injury that helps diminish blood loss. Platelets adhere to damaged endothelium and promote hemostasis. Lymphatic obstruction results in the accumulation of protein-rich lymph and lymphocytes, producing a chylous effusion within a body cavity. After neutrophils reach the site of tissue injury outside of the vascular space, they release lysosomal enzymes that promote liquefaction.

6. **E** Leukocyte rolling is the first step in transmigration of neutrophils from the vasculature to the tissues. Rolling depends on interaction between selectins (P-selectin and E-selectin on endothelial cells, and L-selectin on neutrophils) and their sialylated ligands (e.g., sialylated Lewis X). Integrins are involved in the next step of transmigration, during which there is firm adhesion between neutrophils and endothelial cells. Complement C3b acts as an opsonin to facilitate phagocytosis. Leukotriene B4 is a chemoattractant. NADPH oxidase is involved in phagocytic microbicidal activity.

7. **B** Chemokines include many molecules that are chemotactic for neutrophils, eosinophils, lymphocytes, monocytes, and basophils. Bradykinin causes pain and increased vascular permeability. Complement C5a causes increased vascular permeability by releasing histamine from mast cells. Histamine causes vascular leakage. Prostaglandins have multiple actions, but they do not cause chemotaxis.

8. **B** This immunoglobulin deficiency prevents opsonization and phagocytosis of microbes. Deficiency of integrins and selectins, or a defect in microtubules, would prevent adhesion and locomotion of neutrophils. \( \text{H}_2\text{O}_2 \) production is part of the oxygen-dependent killing mechanism. This mechanism is intact in this patient because the neutrophils are able to kill bacteria when immunoglobulins in normal serum allow phagocytosis.

9. **B** Myeloperoxidase is present in the azurophilic granules of neutrophils. It converts \( \text{H}_2\text{O}_2 \) into \( \text{HOCl}^- \), a powerful oxidant and antimicrobial agent. Degranulation occurs as phagolysosomes are formed with engulfed bacteria in phagocytic vacuoles within the neutrophil cytoplasm. Oxygen consumption with an oxidative or respiratory burst after phagocytosis is aided by glucose oxidation and activation of neutrophil NADPH oxidase, resulting in generation of...
superoxide that is converted by spontaneous dismutation to \( \text{H}_2\text{O}_2 \). In contrast, prostaglandin production depends on a functioning cyclooxygenase pathway of arachidonic acid metabolism.

**PBD9 79–80  BP9 38–39  PBD8 53  BP8 39**

10 B The respiratory, or oxidative, burst of neutrophils generates reactive oxygen species (e.g., superoxide anion) that are important in destruction of engulfed bacteria. This burst can be quantitated by flow cytometric analysis. Neutrophil attachment to endothelium is aided by adhesion molecules on both the endothelium and the neutrophil surface. These molecules include selectins and integrins. Myeloperoxidase does not depend on generation of superoxide. Bacteria are opsonized by complement C3b and IgG, allowing the bacteria to be more readily phagocytosed.

**PBD9 79  BP9 38–39  PBD8 53  BP8 39**

11 D During acute inflammation, in the first stage of extravasation, the neutrophils “roll over” the endothelium. At this stage, the adhesion between the neutrophils and endothelial cells is weak. Rolling is mediated by binding of selectins to sialylated oligosaccharides. The next step, firm adhesion, is mediated by binding of integrins on the leukocytes to their receptors, intercellular adhesion molecule-1 or vascular cell adhesion molecule-1 (VCAM-1), on endothelial cells. Integrins have two chains, \( \alpha \) and \( \beta \). A genetic lack of \( \beta \) chains prevents firm adhesion of leukocytes to endothelial cells. This process depends on adhesion molecules expressed on the neutrophils and endothelial cells. Formation of HOCI requires myeloperoxidase released from neutrophil granules. Phagocytosis of opsonized organisms depends on engulfment, which requires contractile proteins in the neutrophil cytoplasm. Neutrophil migration to a site of infection depends on the presence of chemotactic factors such as complement C5a that bind to the neutrophil and activate phospholipase C to begin a series of events that culminate in the influx of calcium, which triggers contractile proteins. The respiratory burst to kill phagocytized organisms depends on NADPH oxidase, and a deficiency of this enzyme leads to chronic granulomatous disease.

**PBD9 75–76  BP9 35–36  PBD8 49–50  BP8 36–37**

12 E Monocytes transforming to macrophages contain cytokine-inducible nitric oxide synthase (iNOS), which generates nitric oxide. Nitric oxide, by itself and on interaction with other reactive oxygen species, has antimicrobial activity. CD4 or CD8 lymphocytes can be the source for interferon-\( \gamma \) (IFN-\( \gamma \)), which stimulates macrophage production of NOS. Endothelial cells contain a form of NOS (eNOS) that acts to promote vasodilation. B lymphocytes produce immunoglobulins that can opsonize bacteria. Basophils release histamine and arachidonic acid metabolites, which participate in the acute inflammatory process. Natural killer cells have Fc receptors and can lyse IgG-coated target cells; they also generate IFN-\( \gamma \). Neutrophils can phagocytize microbes, but they use NADPH oxidase and enzymes other than NOS to kill the microbes.

**PBD9 79–80  BP9 54  PBD8 54  BP8 40**

13 B Interferon-\( \gamma \) secreted from lymphocytes stimulates monocytes and macrophages, which secrete their own cytokines that further activate lymphocytes. Interferon-\( \gamma \) also is important in transforming macrophages into epithelioid cells in a granulomatous inflammatory response. Histamine released from mast cells is a potent vasodilator, increasing vascular permeability. Leukotriene B\(_4\), generated in the lipoxygenase pathway of arachidonic acid metabolism, is a potent neutrophil chemotactic factor. Nitric oxide generated by macrophages aids in destruction of microorganisms; nitric oxide released from endothelium mediates vasodilation and inhibits platelet activation. Binding of agonists such as epinephrine, collagen, or thrombin to platelet surface receptors activates phospholipase C, which catalyzes the release of arachidonic acid from two of the major membrane phospholipids, phosphatidylinositol and phosphatidylcholine. Tumor necrosis factor (TNF), produced by activated macrophages, mediates many systemic effects, including fever, metabolic wasting, and hypotension.

**PBD9 94–95  BP9 56  PBD8 52  BP8 55–56**

14 C Histamine is found in abundance in mast cells, which are normally present in connective tissues next to blood vessels beneath mucosal surfaces in airways. Binding of an antigen (allergen) to IgE antibodies that have previously attached to the mast cells by the Fc receptor triggers mast cell degranulation, with release of histamine. This response causes increased vascular permeability and mucous secretions. Bradykinin, generated from the kinin system on surface contact of Hageman factor with collagen and basement membrane from vascular injury, promotes vascular permeability, smooth muscle contraction, and pain. Complement C5a is a potent chemotactic factor for neutrophils. Interleukin-1 (IL-1) and tumor necrosis factor (TNF), both produced by activated macrophages, mediate many systemic effects, including fever, metabolic wasting, and hypotension. Phospholipase C, which catalyzes the release of arachidonic acid, is generated from platelet activation.

**PBD9 83  BP9 55–56  PBD8 57–58  BP8 32, 34**

15 F The COX-2 enzyme is inducible with acute inflammatory reactions, particularly in neutrophils, in synovium, and in the central nervous system. The cyclooxygenase pathway of arachidonic acid metabolism generates prostaglandins, which mediate pain, fever, and vasodilation. Ankle swelling is most likely to result from peripheral edema secondary to congestive heart failure. Increased susceptibility to bruising results from prolonged glucocorticoid administration, which also causes leukopenia. Asthma results from bronchoconstriction mediated by leukotrienes that are generated by the lipoxygenase pathway of arachidonic acid metabolism. Inhibition of histamine released from mast cells helps reduce urticaria. Fever can be mediated by prostaglandin release, not inhibition.

**PBD9 84–85  BP9 46–47  PBD8 58–60  BP8 47–48**

16 E Naproxen, a nonsteroidal anti-inflammatory drug, targets the cyclooxygenase pathway of arachidonic acid metabolism and leads to reduced prostaglandin generation.
Prostaglandins promote vasodilation at sites of inflammation. Chemotaxis is a function of various chemokines, and complement C3b may promote phagocytosis, but neither is affected by aspirin. Leukocyte emigration is aided by various adhesion molecules. Leukocyte release from bone marrow can be driven by the cytokines interleukin-1 (IL-1) and tumor necrosis factor (TNF).

**17 E** Prostaglandins are produced through the cyclooxygenase pathway of arachidonic acid metabolism. Aspirin and other nonsteroidal anti-inflammatory drugs block the synthesis of prostaglandins, which can produce pain. Complement C1q is generated in the initial stage of complement activation, which can eventually result in cell lysis. Histamine is mainly a vasodilator. Leukotrienes are generated by the lipoxygenase pathway, which is not blocked by aspirin. Nitric oxide released from endothelium is a vasodilator.

**18 C** Endothelial cells can release nitric oxide to promote vasodilation in areas of ischemic injury. Bradykinin mainly increases vascular permeability and produces pain. Leukotriene E_{2}, platelet-activating factor, and thromboxane A_{2} have vasoconstrictive properties.

**19 A** Activation of complement may occur via microbial cell wall components such as polysaccharides (alternative pathway) or mannose (lectin pathway), or antibody attached to surface antigens (classic pathway). A variety of complement components are generated, including complement C5a, a neutrophil chemoattractant; complement C3b, an opsonin; and complement C5-9, the membrane attack complex. The remaining options are more closely associated with coagulation. Fibrin is generated by the coagulation system, but not with anticoagulation. Kallikrein may aid in generation of bradykinin and plasmin, but participates just in complement C5a generation. Plasmin is generated from plasminogen and helps lyse clots. Thrombin is generated by the coagulation cascade.

**20 E** Ongoing activation of coagulation generates an inflammatory response that further amplifies coagulation, creating a vicious cycle. Protein C antagonizes coagulation factor V, which catalyzes activation of prothrombin to thrombin, thereby breaking the cycle of thrombin generation. Complement components can become activated by plasmin (C3) and kallikrein (C5), forming anaphylatoxins (C5a and C5a) that promote inflammation. Fibrin, the end product of coagulation pathways, forms a meshwork entrapping platelets and creating a plug. Kallikrein is generated by activation of Hageman factor (XII) and leads to formation of bradykinin. Plasmin is generated from plasminogen activated by thrombosis to promote clot lysis.

**21 D** Serous inflammation is the mildest form of acute inflammation. A blister is a good example of serous inflammation. It is associated primarily with exudation of fluid into the subcorneal or subepidermal space. Because the injury is mild, the fluid is relatively protein-poor. A protein-rich exudate results in fibrin accumulation. Granulomatous inflammation is characterized by collections of transformed macrophages called epithelioid cells. Acute inflammatory cells, mainly neutrophils, exuded into a body cavity or space form a purulent (suppurative) exudate, typically associated with liquefactive necrosis. Loss of the epithelium leads to ulceration.

**22 C** This patient is experiencing an acute inflammatory response, with edema, erythema, and pain of short duration. Neutrophils form an exudate and release various proteases, which can produce liquefactive necrosis, starting at the mucosa and extending through the wall of the tube. This mechanism results in perforation. Fibroblasts are more likely participants in chronic inflammatory responses and in healing responses, generally appearing more than 1 week after the initial event. Langhans giant cells are a feature of granulomatous inflammation. Mononuclear infiltrates are more typical of chronic inflammation of the fallopian tube, in which rupture is less likely. Epithelial metaplasia is most likely to occur in the setting of chronic irritation with inflammation.

**23 E** Bacterial infections often evoke an acute inflammatory response dominated by neutrophils. The extravasated neutrophils attempt to phagocytose and kill the bacteria. In the process, some neutrophils die, and the release of their lysosomal enzymes can cause liquefactive necrosis of the tissue. This liquefied tissue debris and both live and dead neutrophils comprise pus, or purulent exudate. Such an exudate is typical of bacterial infections that involve body cavities. Another term for purulent exudate in the pleural space is empyema. An abscess is a localized collection of neutrophils within tissues. Chronic inflammation occurs when there is a preponderance of mononuclear cells, such as lymphocytes, macrophages, and plasma cells, in a process that has gone on for more than a few days—more likely weeks or months—or that accompanies repeated bouts of acute inflammation. Edema refers to increased cellular and interstitial fluid collection within tissues, leading to tissue swelling. In fibrinous inflammation, exudation of blood proteins (including fibrinogen, which polymerizes to fibrin) gives a grossly shaggy appearance to surfaces overlying the inflammation. A serous effusion is a watery-appearing transudate that resembles an ultrafiltrate of blood plasma, with a low cell and protein content.

**24 D** If inflammation is limited and brief, and the involved tissue can regenerate, then resolution is the likely outcome, without significant loss of function. In older persons this may take longer, but can still occur. Multiple bouts of acute inflammation, or ongoing inflammation, can become chronic, and there tends to be loss of some tissue function. If
significant tissue destruction occurs, there is likely to be formation of a fibrous scar in the region of the tissue loss. Acute inflammation is not a preneoplastic event. Ulceration refers to loss of an epithelial surface with acute inflammation; if the epithelium regenerates, then there is resolution.

28 B One outcome of acute inflammation with ulceration is chronic inflammation. This is particularly true when the inflammatory process continues for weeks to months. Chronic inflammation is characterized by tissue destruction, mononuclear cell infiltration, and repair. In acute inflammation, the healing process of fibrosis and angiogenesis has not begun. In fibrinous inflammation, typically involving a mesothelial surface, there is an outpouring of protein-rich fluid that results in precipitation of fibrin. Granulomatous inflammation is a form of chronic inflammation in which epithelioid macrophages form aggregates. Serous inflammation is an inflammatory process involving a mesothelial surface (e.g., lining of the pericardial cavity), with an outpouring of fluid having little protein or cellular content.

25 F Inflammation involving an epithelial surface may cause such extensive necrosis that the surface becomes eroded, forming an ulcer. If the inflammation continues, the ulcer can continue to penetrate downward into submucosa and muscularis. Alternatively, the ulcer may heal, or it may remain chronically inflamed. An abscess is a localized collection of neutrophils in tissues. A caseating granuloma is granulomatous inflammation with central necrosis; the necrosis has elements of both liquefaction and coagulative necrosis. Chronic inflammation occurs when there is a preponderance of mononuclear cells, such as lymphocytes, macrophages, and plasma cells, in a process that has gone on for more than a few days—more likely weeks or months—or that accompanies repeated bouts of acute inflammation. Pus, or a purulent exudate, appears semiliquid and yellowish because of the large numbers of granulocytes present. A serous effusion is a watery-appearing transudate that resembles an ultrafiltrate of blood plasma, with a low cell and protein content.

26 A The rounded density in the right lower lobe of the lung has liquefied contents that form a central air-fluid level. There are surrounding infiltrates. The formation of a fluid-filled cavity after infection with Staphylococcus aureus suggests that liquefactive necrosis has occurred. The cavity is filled with tissue debris and viable and dead neutrophils (pus). Localized, pus-filled cavities are called abscesses. Some bacterial organisms, such as S. aureus, are more likely to be pyogenic, or pus-forming. With complete resolution, the structure of the lung remains almost unaltered. Scarring or fibrosis may follow acute inflammation as the damaged tissue is replaced by fibrous connective tissue. Most bacterial pneumonias resolve, and progression to continued chronic inflammation is uncommon. Lung tissue, in contrast to liver, is incapable of regeneration, except for epithelium and endothelium.

27 E This patient has an infective endocarditis with septic embolization, producing a cerebral abscess. The tissue destruction that accompanies abscess formation as part of acute inflammatory processes occurs from lysosomal enzymatic destruction, aided by release of reactive oxygen species. Nitric oxide generated by macrophages aids in destruction of infectious agents. Immunoglobulin formed by B cells neutralizes and opsonizes infectious agents. Prostaglandins produced by endothelium promote vasodilation. Interferon-γ released from lymphocytes plays a major role in chronic and granulomatous inflammatory responses.

29 C Macrophages, present in such lesions, play a prominent role in the healing process. Activated macrophages can secrete various cytokines that promote angiogenesis and fibrosis, including platelet-derived growth factor, fibroblast growth factor, interleukin-1 (IL-1), and tumor necrosis factor (TNF). Eosinophils are most prominent in allergic inflammations and in parasitic infections. Epithelioid cells, which are aggregations of activated macrophages, are typically seen with granulomatous inflammation, and the healing of acute inflammatory processes does not involve granulomatous inflammation. Neutrophils are most numerous within the initial 48 hours after infarction, but are not numerous after the first week. Plasma cells can secrete immunoglobulins and are not instrumental to healing of an area of tissue injury.

30 B Interferon-γ is secreted by activated T cells and is an important mediator of granulomatous inflammation. It causes activation of macrophages and their transformation into epithelioid cells and then giant cells. Complement C3b acts as an opsonin in acute inflammatory reactions. Interleukin-1 (IL-1) can be secreted by macrophages to produce various effects, including fever, leukocyte adherence, fibroblast proliferation, and cytokine secretion. Leukotriene B4 induces chemotaxis in acute inflammatory processes. Tumor necrosis factor (TNF) can be secreted by activated macrophages and induces activation of lymphocytes and proliferation of fibroblasts, which are other elements of a granuloma.

31 D These findings suggest a granulomatous inflammation, and tuberculosis is a common cause. Candida is often a commensal organism in the oropharyngeal region and rarely causes pneumonia in healthy (non-immunosuppressed) individuals. Viral infections tend to produce a mononuclear interstitial inflammatory cell response. Bacteria such as Enterobacter and Staphylococcus are more likely to produce acute inflammation. Plasmodium produces malaria, a parasitic infection without a significant degree of lung involvement.
inflammation. Bacterial organisms often lead to fever accompanying infection through release of exogenous pyrogens that induce inflammatory cells to release endogenous pyrogens such as tumor necrosis factor (TNF) and interleukin-1 (IL-1). The pyrogens stimulate prostaglandin synthesis in the hypothalamus to “reset the thermostat,” so that fever occurs as a sign of the acute inflammatory response. Prostaglandins are mainly involved in the causation of vasodilation and pain in acute inflammatory responses.

33 C  Figure A shows diffuse reticulonodular pulmonary densities, and Figure B shows noncaseating granulomas with many epithelioid cells and two prominent large Langhans giant cells. If special stains and/or cultures for organisms (usually mycobacteria or fungi) are negative, then this is likely sarcoidosis. Macrophage stimulation and transformation to epithelioid cells and giant cells are characteristic of granuloma formation. Interferon-γ promotes the formation of epithelioid cells and giant cells. Bradykinin is released in acute inflammatory responses and results in pain. Complement C5a is chemotactic for neutrophils. Although occasional neutrophils are seen in granulomas, neutrophils do not form a major component of granulomatous inflammation. Macrophages can release nitric oxide to destroy other cells, but nitric oxide does not stimulate macrophages to form a granulomatous response. Prostaglandins are mainly involved in the causation of vasodilation and pain in acute inflammatory responses.

34 E  The findings here are those of strep throat with acute inflammation. Bacterial organisms often lead to fever accompanying infection through release of exogenous pyrogens that induce inflammatory cells to release endogenous pyrogens such as tumor necrosis factor (TNF) and interleukin-1 (IL-1). The pyrogens stimulate prostaglandin synthesis in the hypothalamus to “reset the thermostat,” so that fever occurs as a sign of the acute inflammatory response. Hageman factor with collagen and basement membrane from vascular injury, promotes vascular permeability, smooth muscle contraction, and pain. Histamine released from mast cells is a potent vasodilator, increasing vascular permeability. Leukotriene B_4 generated in the lipoygenase pathway of arachidonic acid metabolism, is a potent neutrophil chemotactic factor. Nitric oxide generated by macrophages aids in destruction of microorganisms; nitric oxide released from endothelium mediates vasodilation and inhibits platelet activation.

35 E  Fever is produced by various inflammatory mediators, but the major cytokines that produce fever are interleukin-1 (IL-1) and tumor necrosis factor (TNF), which are produced by macrophages and other cell types. IL-1 and TNF can have autocrine, paracrine, and endocrine effects. They mediate the acute phase responses, such as fever, nausea, and neutrophil release from bone marrow. Bradykinin, generated from the kinin system on surface contact of Hageman factor with collagen and basement membrane.

36 B  This acute inflammatory process leads to production of acute-phase reactants, such as C-reactive protein (CRP), fibrinogen, and serum amyloid A (SAA) protein. These proteins, particularly fibrinogen, and immunoglobulins increase RBC rouleaux formation to increase the erythrocyte sedimentation rate (ESR), which is a nonspecific indicator of inflammation. CRP production is upregulated by interleukin-6 (IL-6), whereas fibrinogen and SAA are upregulated mainly by tumor necrosis factor (TNF) and interleukin-1 (IL-1). Interferon-γ is a potent stimulator of macrophages. Nitric oxide can induce vasodilation or can assist in microbial killing within macrophages. Prostaglandins are vasodilators.

37 C  Hepatocytes are stable cells with an extensive ability to regenerate. The ability to restore normal architecture of an organ such as the liver depends on the viability of the supporting connective tissue framework. If the connective tissue cells are not injured, hepatocyte regeneration can restore normal liver architecture. This regeneration occurs in many cases of viral hepatitis. A liver abscess associated with liquefactive necrosis of hepatocytes and the supporting connective tissue heals by scarring. The other options listed may explain the amount of liver injury, but not the nature of the response.

38 A  Hepatocytes are quiescent (stable) cells that can reenter the cell cycle and proliferate in response to hepatic injury, enabling the liver to regenerate partially. Acute hepatitis results in hepatocyte necrosis, marked by elevations in AST and ALT. After the acute process has ended, cells return to the G0 phase, and the liver becomes quiescent again.

39 A  Vitamin C deficiency leads to scurvy, with reduced lysyl oxidase enzyme activity that helps cross-link fibrillar collagens to provide tensile strength. Though elastin is a fibrillar protein, it tends to regenerate poorly in scar tissue, even with the best of nutrition, explaining why a scar does not stretch like the skin around it. The other listed choices are glycoproteins that have an adhesive quality and are not vitamin C dependent.
40 F The figure shows a subacute infarction with granulation tissue formation containing numerous capillaries stimulated by vascular endothelial growth factor, representing a healing response. Epidermal growth factor aids in reepithelialization of a surface wound. Interleukin-2 (IL-2) mediates lymphocyte activation. Leukotriene B4 mediates reepithelialization of a surface wound. Interleukin-2 (IL-2) mediates lymphocyte activation.

41 A At 1 week, wound healing is incomplete, and granulation tissue is still present. More collagen is synthesized in the following weeks. Wound strength peaks at about 80% by 3 months. Type IV collagen is found in basement membranes.

42 E The figure shows dense collagen with some remaining dilated blood vessels, typical of the final phase of wound healing, which is extensive by the end of the first month. On day 1, the wound is filled only with fibrin and inflammatory cells. Macrophages and granulation tissue are seen 2 to 3 days postoperatively. Neovascularization is most prominent by days 4 and 5. By week 2, collagen is prominent, and fewer vessels and inflammatory cells are seen.

43 D Wound contraction is a characteristic feature of healing by second intention that occurs in larger wounds. Collagen synthesis helps fill the defect, but does not contract it. Adhesive glycoproteins such as fibronectin help to maintain a cellular scaffolding for growth and repair, but they do not contract. The inhibition of metalloproteinases leads to decreased degradation of collagen and impaired connective tissue remodeling in wound repair. Edema diminishes over time, but this does not result in much contraction.

44 C Integrins interact with the extracellular matrix proteins (e.g., fibronectin). Engagement of integrins by extracellular matrix proteins leads to the formation of focal adhesions where integrins link to intracellular cytoskeletal elements such as actin. These interactions lead to intracellular signals that modulate cell growth, differentiation, and migration during wound healing. Epidermal growth factor stimulates epithelial cell and fibroblast proliferation. Platelet-derived growth factor (PDGF) can be produced by endothelium, macrophages, smooth muscle cells, and platelets; PDGF mediates migration and proliferation of fibroblasts and smooth muscle cells and migration of monocytes. Type IV collagen is found in basement membranes on which cells are anchored. Vascular endothelial growth factor promotes angiogenesis (capillary proliferation) through endothelial cell proliferation and migration in a healing response.

45 A Glucocorticoids inhibit wound healing by impairing collagen synthesis. This is a desirable side effect if the amount of scarring is to be reduced, but it results in the delayed healing of surgical wounds. Angiogenesis driven by vascular endothelial growth factor (VEGF) is not significantly affected by corticosteroids. Neutrophil infiltration is not prevented by glucocorticoids. Reepithelialization, in part driven by epidermal growth factor, is not affected by corticosteroid therapy. Serine proteinases are important in wound remodeling.

46 B The healing process sometimes results in an exuberant production of collagen, giving rise to a keloid, which is a prominent raised, nodular scar, as shown in the figure. This tendency may run in families. Dehiscence occurs when a wound pulls apart. Organization occurs as granulation tissue is replaced by fibrous tissue. If normal tissue architecture is restored, resolution of inflammation has occurred. Secondary union describes the process by which large wounds fill in and contract.

47 D The elevated creatine kinase level indicates that myocardial necrosis has occurred. A fibrous scar gradually replaces the area of myocardial necrosis. Chronic inflammation is typically driven by ongoing stimuli such as persistent infection, autoimmunity, or irritation from endogenous or exogenous chemical agents, and it is not a feature of ischemic myocardial injury. Coagulative necrosis is typical of myocardial infarction, but after 1 month, a scar would be present. The destruction of myocardial fibers precludes complete resolution. Nodular regeneration is typical of hepatocyte injury because hepatocytes are stable cells.
A 45-year-old woman who works while standing for long periods notices at the end of her 8-hour shift that her lower legs and feet are swollen, although there was no swelling at the beginning of the day. There is no pain or erythema associated with this swelling. She is otherwise healthy and takes no medications; laboratory testing reveals normal liver and renal function. Which of the following mechanisms best explains this phenomenon?

A  Excessive free water intake  
B  Hypoalbuminemia  
C  Increased hydrostatic pressure  
D  Lymphatic obstruction  
E  Secondary aldosteronism

A 56-year-old woman diagnosed with cancer in her left breast underwent a mastectomy with axillary lymph node dissection. Postoperatively, she develops marked swelling of the left arm that has persisted for 6 months. Now on physical examination, her temperature is 36.9°C. Her left arm is not tender or erythematous, and it is not painful with movement or to touch, but it is enlarged with a doughy consistency. Which of the following is the most likely mechanism for these findings?

A  Cellulitis  
B  Congestive heart failure  
C  Decreased plasma oncotic pressure  
D  Lymphedema  
E  Sodium and water retention  
F  Phlebothrombosis

A 37-year-old woman has noticed a lump in her left breast over the past 2 months. On physical examination, the skin overlying the left breast is thickened, reddish orange, and pitted. Mammography shows a 3-cm underlying density. A fine-needle aspirate of the density is performed and on microscopic examination shows carcinoma. Which of the following mechanisms best explains the gross appearance of the skin of her left breast?

A  Chronic inflammation  
B  Chronic passive congestion  
C  Ischemic necrosis  
D  Lymphatic obstruction  
E  Venous thrombosis

A 7-year-old boy has had increasing lethargy for a week. On physical examination, he has periorbital edema and pitting edema at the ankles, but is normotensive and afebrile. Laboratory studies show marked albuminuria. He is given a thiazide diuretic and his urine output increases and his edema resolves. Which of the following changes most likely potenti- ated his edema?

A  Decreased aldosterone  
B  Decreased renin  
C  Increased albumin  
D  Increased cortisol  
E  Decreased antidiuretic hormone  
F  Increased salt retention
A 94-year-old woman has dyspnea and an increasing cough with frothy sputum production for the past month. She is afebrile. A chest radiograph shows the findings in the figure. Which of the following is the most likely mechanism for development of her pulmonary infiltrates?

A. Decreased sodium intake  
B. Hypoalbuminemia  
C. Increased hydrostatic pressure  
D. Inflammation  
E. Pulmonary venous obstruction

A 50-year-old man suffers an infarction of the anterior left ventricular wall. He receives therapy with anti-arrhythmic and pressor agents. He is in stable condition until he develops severe breathlessness 3 days later. An echocardiogram shows a markedly decreased ejection fraction. Representative chest radiographic findings are shown in the figure above. Which of the following microscopic changes is most likely to be present in his lungs?

A. Alveolar neutrophilic exudate  
B. Alveolar transudate  
C. Alveolar wall fibrosis  
D. Pleural fibrosis  
E. Pleural space neutrophilic exudate  
F. Pleural space transudate

A 58-year-old man with pulmonary emphysema has a 10-year history of congestive heart failure. On physical examination, he has lower leg swelling with grade 2 pitting edema to the knees and prominent jugular venous distention to the level of the mandible. His serum levels of AST and ALT are increased. The representative gross appearance of his liver is shown in the figure. Which of the following underlying conditions is most likely to be present in this man?

A. Chronic renal failure  
B. Common bile duct obstruction  
C. Congestive heart failure  
D. Portal vein thrombosis  
E. Thrombocytopenia

An 85-year-old man falls in the bathtub and strikes the back of his head. Over the next 24 hours, he becomes increasingly somnolent. A head CT scan shows an accumulation of fluid beneath the dura, compressing the left cerebral hemisphere. Which of the following terms best describes this collection of fluid?

A. Congestion  
B. Ecchymosis  
C. Hematoma  
D. Petechiae  
E. Purpura
An autopsy study is performed to correlate patterns of hemorrhage with underlying causes. Patients with the gross appearance of hemorrhage shown in the figure had minimal blood volume loss, but an appearance similar to this in many other organs. Which of the following terms best describes this pattern?

A. Congestion  
B. Ecchymosis  
C. Hematoma  
D. Petechiae  
E. Purpura

A superficial puncture wound from a needlestick injury leads to a small amount of bleeding in a healthy person. Seconds after this injury occurs, the bleeding stops. Which of the following mechanisms is most likely to stop small arteriolar blood loss from this injury?

A. Fibrin polymerization  
B. Neutrophil chemotaxis  
C. Platelet aggregation  
D. Protein C activation  
E. Vasoconstriction

A 15-year-old girl incurs a cut to the sole of her foot after stepping on a piece of broken glass. On examination, a superficial 0.5-cm laceration ceases to bleed within 5 minutes after application of local pressure. Which of the following substances is released by endothelium and is most likely to counteract platelet aggregation near this site of injury?

A. Glycoprotein IIb/IIIa  
B. Platelet-activating factor  
C. Prostacyclin  
D. Tissue-type plasminogen activator  
E. Thrombomodulin  
F. Thromboxane A₂

In an experiment, thrombus formation is studied in areas of vascular damage. The propagation of a thrombus in an area of vascular injury to adjacent normal arteries is prevented. Which of the following substances diminishes thrombus propagation by activating protein C?

A. Calcium  
B. Fibrin  
C. Platelet factor 4  
D. Prothrombin  
E. Thrombomodulin  
F. Tumor necrosis factor (TNF)

A 26-year-old woman has a history of frequent nosebleeds and increased menstrual blood flow. On physical examination, petechiae and purpura are present on the skin of her extremities. Laboratory studies show normal partial thromboplastin time (PTT), prothrombin time (PT), and platelet count, but decreased von Willebrand factor activity. This patient most likely has a derangement in which of the following steps in hemostasis?

A. Fibrin polymerization  
B. Platelet adhesion  
C. Platelet aggregation  
D. Prothrombin generation  
E. Prothrombin inhibition  
F. Vasoconstriction

A 59-year-old woman with a history of diabetes mellitus had a myocardial infarction 3 months ago. Her BMI is 35. She is now taking a low dose of aspirin to reduce the risk for recurrent arterial thrombosis. On which of the following steps in hemostasis does aspirin have its greatest effect?

A. Adhesion of platelets to collagen  
B. Aggregation of platelets  
C. Production of tissue factor  
D. Synthesis of von Willebrand factor  
E. Synthesis of antithrombin III

In an experiment, platelet function is analyzed. A substance is obtained from the dense body granules of normal pooled platelets from healthy blood donors. When this substance is added to platelets obtained from patients with a bleeding disorder, no platelet aggregation occurs. Adding the substance to platelets from a normal control group induces platelet aggregation. Which of the following substances is most likely to produce these effects?

A. Adenosine diphosphate  
B. Antithrombin III  
C. Fibronectin  
D. Fibrinogen  
E. Plasminogen  
F. Thromboxane A₂
16 A 12-year-old boy has a 10-year history of multiple soft tissue hemorrhages and acute upper airway obstruction from hematoma formation in the neck. On physical examination, he has decreased range of motion of the large joints, particularly the knees and ankles. He has no petechiae or purpura of the skin. Laboratory studies show normal prothrombin time, elevated partial thromboplastin time (PTT), and normal platelet count, but markedly decreased factor VIII activity. Which of the following mechanisms best describes the development of his disease?
   A Decrease in production of thrombin
   B Decrease in membrane phospholipid
   C Failure of platelet aggregation
   D Failure of fibrin polymerization
   E Inability to neutralize antithrombin III
   F Inability of platelets to release thromboxane A₂

17 A 58-year-old man has had episodes of prolonged epi-staxis in the past 6 months. On examination he has occult blood detected in his stool. Coagulation studies show that his prothrombin time is elevated, but his partial thromboplastin time (PTT), platelet count, and platelet function are all normal. When his plasma is mixed with an equal amount of normal plasma, the prothrombin time corrects to normal. Which of the following underlying diseases is most likely to be associated with these findings?
   A Antiphospholipid syndrome
   B Factor V Leiden mutation
   C Hemophilia A
   D Scurvy
   E Sepsis with *Escherichia coli*
   F Vitamin K deficiency

18 A 66-year-old woman has the sudden onset of chest pain that radiates to her neck and left arm. On examination 30 minutes later, she is diaphoretic and hypotensive. Her serum troponin I level is elevated. Which of the following drugs is most likely to be administered emergently as thrombolytic therapy for this woman?
   A Acetylsalicylic acid (aspirin)
   B Low-molecular-weight heparin
   C Nitric oxide
   D Tissue plasminogen activator
   E Vitamin K

19 A 71-year-old man with a history of diabetes mellitus died of an acute myocardial infarction. At autopsy, the aorta, opened longitudinally and with the superior aspect of the kidneys below the forceps, appeared as shown in the figure. Which of the following complications associated with this aortic disease would most likely have been present during his life?
   A Edema of the left leg
   B Gangrene of the foot
   C Pulmonary thromboembolism
   D Renal infarction
   E Thrombocytopenia

20 A 21-year-old woman has had multiple episodes of deep venous thrombosis during the past 10 years and one episode of pulmonary thromboembolism during the past year. Laboratory tests show that her prothrombin time (PT), partial thromboplastin time (PTT), platelet count, and platelet function studies all are normal. Which of the following risk factors is the most common cause for such a coagulopathy?
   A Antithrombin III deficiency
   B Factor V mutation
   C Hyperhomocysteinemia
   D Mutation in protein C
   E Occult malignancy
   F Oral contraceptive use
   G Smoking cigarettes

21 A 23-year-old woman has had altered consciousness and slurred speech for the past 24 hours. A head CT scan shows a right temporal hemorrhagic infarction. Cerebral angiography shows a distal right middle cerebral arterial occlusion. Within the past 3 years, she has had an episode of pulmonary embolism. A pregnancy 18 months ago ended in miscarriage. Laboratory studies show a false-positive serologic test for syphilis, normal prothrombin time (PT), elevated partial thromboplastin time (PTT), and normal platelet count. Which of the following is the most likely cause of these findings?
   A Antiphospholipid antibody
   B Disseminated intravascular coagulation
   C Factor V mutation
   D Hypercholesterolemia
   E Von Willebrand disease

22 A 71-year-old man with a history of diabetes mellitus died of an acute myocardial infarction. At autopsy, the aorta, opened longitudinally and with the superior aspect of the kidneys below the forceps, appeared as shown in the figure. Which of the following complications associated with this aortic disease would most likely have been present during his life?
A 55-year-old woman following major abdominal surgery has had discomfort and swelling of her left leg for the past week. On physical examination, the leg is slightly difficult to move, and on palpation there is tenderness. A Doppler sonogram shows thrombosis of deep left leg veins. Which of the following mechanisms is most likely to contribute to her condition?

A  Hypercalcemia  
B  Immobilization  
C  Ingestion of aspirin  
D  Nitric oxide release  
E  Turbulent blood flow

A 75-year-old man is hospitalized after falling and fracturing his left femoral trochanter. Two weeks later, the left leg is swollen, particularly below the knee. He experiences pain on movement of the leg; on palpation, there is swelling and tenderness. Which of the following complications is most likely to occur in this man?

A  Disseminated intravascular coagulation  
B  Fat embolism syndrome  
C  Gangrenous necrosis of the foot  
D  Hematoma of the thigh  
E  Pulmonary thromboembolism

A 65-year-old woman sustained fractures of the right femur, pelvis, and left humerus in a motor vehicle collision. The fractures were stabilized, and the patient’s recovery was uneventful. During a physical examination 3 weeks later, while still in the hospital, she has swelling and warmth in the left leg, and there is local pain and tenderness in the left thigh. Which of the following processes, as shown in the figure, is most likely occurring in her left femoral vein?

A  Atherosclerosis  
B  Chronic passive congestion  
C  Inflammation  
D  Mural thrombosis  
E  Phlebothrombosis  
F  Vegetation

A 59-year-old woman with hyperlipidemia has had anginal pain for the past 24 hours. Laboratory findings show no increase in serum troponin I or creatine kinase-MB. She is in stable condition 2 weeks later and has no chest pain, but a small artery in the epicardium has undergone the changes seen in the figure. Which of the following terms best describes this finding in this epicardial artery?

A  Air embolus  
B  Cholesterol embolization  
C  Chronic passive congestion  
D  Fat embolism syndrome  
E  Mural thrombosis  
F  Organization with occlusion  
G  Phlebothrombosis

A 77-year-old woman has a brief fainting episode. She was diagnosed 1 year ago with pancreatic adenocarcinoma. On auscultation of her chest, a heart murmur is heard. Echocardiography shows a 1-cm nodular lesion on the superior aspect of an intact anterior mitral valve leaflet. A blood culture is negative. Which of the following terms best describes this mitral valve lesion?

A  Atheroma  
B  Chronic passive congestion  
C  Mural thrombus  
D  Myxoma  
E  Phlebothrombosis  
F  Vegetation
28 A 70-year-old man was hospitalized 3 weeks ago for a cerebral infarction. He is now ambulating for the first time. Within minutes of returning to his hospital room, he has sudden onset of dyspnea with diaphoresis. He cannot be resuscitated. The gross appearance of the hilum of the left lung at autopsy is shown in the figure. Which of the following risk factors most likely contributed to this finding?

- A Antiphospholipid antibody
- B Bronchopneumonia
- C Factor V mutation
- D Leg vein thrombosis
- E Pulmonary arterial atherosclerosis

29 A 32-year-old man is involved in a vehicular accident and sustains fractures of the right femur and tibia and the left humerus. The fractures are stabilized surgically. He is in stable condition for 2 days, but then suddenly becomes severely dyspneic. Which of the following complications from his injuries is the most likely cause of his sudden respiratory difficulty?

- A Cardiac tamponade
- B Fat embolism
- C Pulmonary edema
- D Pulmonary infarction
- E Right hemothorax

30 A 22-year-old woman with an uncomplicated pregnancy develops sudden dyspnea with cyanosis and hypotension intrapartum during routine vaginal delivery of a term infant. She has a generalized seizure and becomes comatose. Her condition does not improve over the next 2 days. Which of the following findings is most likely to be present in her peripheral pulmonary arteries?

- A Aggregates of platelets
- B Amniotic fluid
- C Fat globules
- D Gas bubbles
- E Thromboemboli

31 A 31-year-old man is on a scuba diving trip and descends to a depth of 50 m in the Blue Hole off the coast of Belize. After 30 minutes, he has a malfunction in his equipment and quickly returns to the boat on the surface. He develops difficulty breathing within 5 minutes, with dyspnea and substernal chest pain, followed by a severe headache and vertigo. An hour later, he develops severe, painful myalgias and arthralgias. These symptoms abate within 24 hours. Which of the following occluding his arterioles is the most likely cause of his findings?

- A Fat globules
- B Fibrin clots
- C Nitrogen gas bubbles
- D Platelet thrombi
- E Ruptured atheromatous plaque

32 A 53-year-old man with congestive heart failure develops pulmonary Streptococcus pneumoniae infection after a bout of influenza. After recuperating for 2 weeks, he notes pleuritic chest pain. The pain is caused by the development of the lesion shown in the figure. Which of the following events has most likely occurred in this man?

- A Acute pulmonary congestion
- B Chronic pulmonary congestion
- C Pulmonary edema
- D Pulmonary infarction
- E Pulmonary venous thrombosis
A 44-year-old man with dilated cardiomyopathy and heart failure develops left atrial mural thrombosis. He develops the complication shown in the figure, manifested by hematuria. Which of the following is the best term for this complication?

A  Abscess  
B  Ischemic infarct  
C  Liquefactive necrosis  
D  Multiorgan failure  
E  Venous thrombosis

A 28-year-old woman with a 15-year history of recurrent thrombosis from a prothrombin gene mutation develops septicemia after a urinary tract infection with *Pseudomonas aeruginosa*. She develops multiple infarcts and organ failure over the next 2 weeks. Which of the following organs is most likely to be spared from the effects of ischemic injury in this woman?

A  Brain  
B  Heart  
C  Kidney  
D  Liver  
E  Spleen

An 80-year-old woman with dysuria for 1 week now has a fever. On examination, her temperature is 37.9°C, pulse 103/min, and blood pressure 80/40 mm Hg. She has right flank pain. A urinalysis shows numerous WBCs. Her plasma lactate is increased. Urine culture and blood culture grow *Escherichia coli*. Which of the following is most likely to contribute to her cardiovascular collapse by triggering Toll-like cell receptors?

A  Complement C3b  
B  Lipopolysaccharide  
C  Nitric oxide  
D  Platelet-activating factor  
E  Toxic shock syndrome toxin-1

A 63-year-old woman has had a fever and felt faint for the past 2 days. On physical examination, her temperature is 38.4°C, pulse is 101/min, respirations are 17/min, and blood pressure is 85/40 mm Hg. She has marked peripheral vaso-dilation. The serum lactic acid level is 6.8 mg/dL. Which of the following laboratory findings is most likely related to the cause of her clinical condition?

A  Blood culture positive for *Citrobacter*  
B  Decreased hematocrit  
C  Elevated serum creatine kinase  
D  Increased blood urea nitrogen  
E  Reduced Po2 on blood gas measurement

A 20-year-old man develops palpitations within 1 hour after a gunshot wound to the abdomen. On examination his heart rate is 112/minute and blood pressure 80/30 mm Hg. His skin is cool and clammy to the touch. Which of the following organ-specific changes is most likely to occur within 2 days after this injury?

A  Acute hepatic infarction  
B  Cerebral basal ganglia hemorrhage  
C  Gangrenous necrosis of the lower legs  
D  Pulmonary diffuse alveolar damage  
E  Renal passive congestion

A 30-year-old man is cutting wood alone in the forest and incurs a deep cut to his leg from his chain saw. He loses a large amount of blood. He is not found until the next day. A marked increase in which of the following blood analytes is most likely to indicate that he has reached an irreversible stage of shock?

A  Antidiuretic hormone  
B  Bicarbonate  
C  Catecholamines  
D  Lactate dehydrogenase  
E  Prothrombin
ANSWERS

1 C The hydrostatic pressure exerted from standing upright leads to edema in dependent parts of the body. In a healthy patient, normal renal function would be sufficient to clear free water ingested orally. Hypoalbuminemia leads to more generalized edema, although the effect is more pronounced in dependent areas. Lymphatic obstruction from infection or tumor can lead to lymphedema, but this is a chronic process. Secondary aldosteronism results from congestive heart failure and renal hyperfusion, but this is a generalized process.

2 D The surgery disrupted lymphatic return, resulting in functional lymphatic obstruction and lymphedema of the arm. The lymphatic channels are important in scavenging fluid and protein that have leaked into the extravascular tissues from the intravascular compartment. Although the amount of fluid that is drained through the lymphatics is not great, it can build up gradually. Cellulitis is caused by an infection of the skin and subcutaneous tissue, and displays erythema, warmth, and tenderness. Congestive heart failure can lead to peripheral edema, which is most marked in dependent areas such as the lower extremities and over the sacrum (in bedridden patients). Decreased plasma oncotic pressure from hypoalbuminemia, or sodium and water retention with heart or renal failure, leads to more generalized edema. Thromboembolism leads to swelling with pain and tenderness, but it is uncommon in the upper extremities.

3 D Spread of the cancer to the dermal lymphatics produces a peau d’orange appearance of the breast. Because the breast has an extensive venous drainage, cancer or other focal mass lesions are unlikely to cause significant congestion and edema of the breast. Chronic inflammation is rare in breast tissue and is not associated with cancer. Passive congestion does not involve the breast. Ischemia is rare in the breast because of the abundant arterial supply.

4 F This child has nephrotic syndrome with loss of albumin into the urine and hypoalbuminemia that decreases plasma oncotic pressure, leading to movement of intravascular water into the extravascular compartment to produce edema. In response, hypovolemia with renal hyperfusion induces increased production of renin, angiotensin, and aldosterone, which all promote sodium and water retention, further exacerbating his edema. Thiazide diuretics increase renal excretion of sodium. Hypovolemia would increase antidiuretic hormone output. Though corticosteroids are used to treat nephrotic syndrome caused by minimal change disease, the effect is probably to diminish abnormal T-cell function that is driving the glomerular damage. Cortisol leads to sodium retention, but not in response to hypovolemia.

5 C She has congestive heart failure, with bilateral diffuse pulmonary infiltrates due to edema. The left side of her heart has failed, which increases hydrostatic pressure in the pulmonary vasculature. The higher pressure forces fluid to leak into the alveoli, and it is then coughed up. Note the enlarged heart with prominent border of the left atrium in the figure. This fluid is a transudate because there is little protein or cells present. The kidneys can generally compensate for decreased salt intake, but in modern society we suffer from too much dietary salt, not too little. Edema from hypoalbuminemia is not limited to the lungs, and it is more likely to appear in areas of loose connective tissue, such as the periorbital region. She is afebrile, so the pulmonary infiltrates are unlikely to represent pneumonia that could locally produce edema in regions of inflammation. Pulmonary venous obstruction is typical for pulmonary thromboembolism that may lead to right-sided heart failure and peripheral edema if chronic.

6 B Acute left ventricular failure after a myocardial infarction causes venous congestion in the pulmonary capillary bed and increased hydrostatic pressure, which leads to pulmonary edema by transudation in the alveolar space. Neutrophils and fibrin would be found in cases of acute inflammation of the lung (i.e., pneumonia). Fibrosis and hemosiderin-filled macrophages (heart failure cells) would be found in long-standing, not acute, left ventricular failure. Purulent exudate in the pleural space (empyema) or drainage from bronchi results from bacterial infection, not heart failure. No pleural effusions are present in this radiograph. Fluid collections are likely to be transudates (few cells and minimal protein) in noninflammatory conditions.

7 C The figure shows a so-called nutmeg liver caused by chronic passive congestion from congestive heart failure. The elevated enzyme levels suggest that the process is so severe that hepatic centrilobular necrosis has also occurred. The physical findings suggest right-sided heart failure that can occur with pulmonary emphysema and pulmonary arterial hypertension. Biliary tract obstruction would produce bile stasis (cholestatic) with icterus. Hepatic congestion is not directly related to renal failure, and hepatorenal syndrome has no characteristic gross appearance. A portal vein thrombus would diminish blood flow to the liver, but it would not be likely to cause necrosis because of that organ’s dual blood supply. The regular pattern of red lobular discoloration seen in the figure is unlikely to occur in hemorrhage from thrombocytopenia, characterized by petechiae and ecchymoses.

8 C He has a subdural hematoma from this traumatic injury causing tearing of cerebral veins. A hematoma is a collection of blood in a potential space or within tissue. Congestion occurs when there is vascular dilation with pooling of venous blood within an organ. Ecchymoses are
Thrombin helps activate platelets. Calcium is a cofactor in this process, which leads to the activation of factor XIII, which cross-links fibrin. Normally, thrombin activates factors V, VIII, and IX, and also stabilizes the secondary hemostatic plug by activating factor XIII, which cross-links fibrin. Thrombin helps activate platelets. Calcium is a cofactor that assists clotting in the coagulation cascade (ethylene-diaminetetraacetic acid [EDTA] in some blood collection tubes binds calcium to prevent clotting). Fibrin protein forms a meshwork that is essential to thrombus formation. Platelet factor 4 is released from the α granules of platelets and promotes platelet aggregation during the coagulation process. Prothrombin is converted to thrombin in the coagulation cascade. Tumor necrosis factor (TNF) is not significantly involved in coagulation.

Note the small punctate 1- to 2-mm petechial hemorrhages on the epicardial surface shown in the figure. Such hemorrhages most often result from reduced numbers of platelets or reduced platelet function. It is platelets that plug small vascular defects. Congestion occurs when there is vascular dilation with pooling of blood within an organ. A hematoma is a collection of blood in a potential space or within tissue. Purpura denotes blotchy hemorrhage on skin, serosal surfaces, or mucous membrane surfaces larger than 3 mm; areas from 1 to 2 cm in size are called ecchymoses.

The initial response to arteriolar injury is vasoconstriction, since there is smooth muscle in the vessel wall. But this is transient, and the coagulation mechanism must be initiated to maintain hemostasis. Fibrin polymerization is part of secondary hemostasis after the vascular injury is initially closed. Neutrophils are not essential to hemostasis. Platelet aggregation occurs with release of factors such as ADP, but this takes several minutes. Protein C is involved in anticoagulation to counteract clotting.

Endothelial injury releases glycoprotein tissue factor (factor III) that drives the coagulation process and activates platelets. Adjacent intact endothelium generates prostacyclin (PGI₂) via arachidonic acid metabolism to inhibit clot propagation beyond where it is needed. PGI₂ and nitric oxide are powerful vasodilators and inhibitors of platelet aggregation. This limits thrombus formation just to the area of injury. Glycoprotein IIb/IIIa, which induces shape change; phospholipid, which binds fibrinogen and calcium; and platelet-activating factor are procoagulants that drive thrombosis and platelet activation. Tissue-type plasminogen activator promotes fibrinolytic activity after a thrombus has formed. Thrombomodulin binds to thrombin to form an anticoagulant that activates protein C, which then cleaves activated factor V and factor VIII. Thromboxane is generated via arachidonic acid metabolism in platelets to promote platelet activation and vasoconstriction.

Thrombomodulin is present on intact endothelium and binds thrombin, which then inhibits coagulation by activating protein C. Normally, thrombin activates factors V, VIII, and IX, and also stabilizes the secondary hemostatic plug by activating factor XIII, which cross-links fibrin. Thrombin helps activate platelets. Calcium is a cofactor that assists clotting in the coagulation cascade (ethylene-diaminetetraacetic acid [EDTA] in some blood collection tubes binds calcium to prevent clotting). Fibrin protein forms a meshwork that is essential to thrombus formation. Platelet factor 4 is released from the α granules of platelets and promotes platelet aggregation during the coagulation process. Prothrombin is converted to thrombin in the coagulation cascade. Tumor necrosis factor (TNF) is not significantly involved in coagulation.

Aspirin blocks the cyclooxygenase pathway of arachidonic acid metabolism and generation of eicosanoids in platelets, including thromboxane A₂, to block vasoconstriction and inhibit platelet aggregation. Platelet adhesion to extracellular matrix is mediated by interactions with von Willebrand factor. Tissue factor (factor III), produced by injured endothelium as well as subendothelial smooth muscle and fibroblasts, is released with tissue injury and is not platelet dependent. Endothelial cells produce von Willebrand factor independent of platelet action. Antithrombin III has anticoagulant properties because it inactivates several coagulation factors, but its function is not affected by aspirin.

ADP is released from the platelet-dense granules and is a potent stimulator of platelet aggregation. ADP also stimulates further release of ADP from other platelets. Many other substances involved in hemostasis, such as fibrinogen, fibronectin, and factors V and VIII, are stored in the α granules of platelets. Thromboxane A₂, another powerful aggregating factor of platelets, is synthesized by the cyclooxygenase pathway. Fibronectin forms part of the extracellular matrix between cells that “glues” them together. Plasminogen is activated to promote thrombolysis. Platelet aggregation requires active platelet metabolism; platelet stimulation by agonists such as ADP, thrombin, collagen, or epinephrine; the presence of calcium or magnesium ions and specific plasma proteins such as fibrinogen or von Willebrand factor (vWF); and a platelet receptor, the glycoprotein IIb/IIIa (GPIIb/IIIa) complex. Platelet stimulation results in the generation of intracellular second messengers that transmit the stimulus back to the platelet surface, exposing protein-binding sites on GPIIb/IIIa. Fibrinogen then binds to GPIIb/IIIa and cross-links adjacent platelets to produce platelet aggregates; vWF binds to drive shape change and granule release. The patients in this experiment could have Glanzmann thrombasthenia, in which platelets are deficient or defective in the GPIIb/IIIa
complex, do not bind fibrinogen, and cannot form aggregates, although the platelets can be stimulated by ADP, can undergo shape change, and are of normal size.

PBD9 117–118  BP9 82  PBD8 116–118  BP8 87–89

16 A He has hemophilia A. Factor VIII, tissue factor (III), and factor V act as cofactors or reaction accelerators in the clotting cascade leading to thrombin production. Factor VIII acts as a reaction accelerator for the conversion of factor X and factor Xa. The platelet surface provides phospholipid for assembly of coagulation factors. Platelet aggregation is promoted by thromboxane A2 and ADP. Thromboxane A2 is released when platelets are activated during the process of platelet adhesion. Fibrin polymerization is promoted by factor XIII. Antithrombin III inhibits thrombin to prolong the prothrombin time.

PBD9 121–122  BP9 83–84  PBD8 119–120  BP8 88–89, 91

17 F His elevated prothrombin time that corrects with normal plasma points to coagulation factor deficiency, and factors II, VII, IX, and X are synthesized in the liver and affect this “extrinsic” in vitro coagulation pathway. They are vitamin K dependent and therefore may also be affected by warfarin therapy or by parenchymal liver disease. Antiphospholipid syndrome has an inhibitory effect upon vitamin K activity in vivo. Antiphospholipid antibodies are directed against phospholipid-protein complexes such as β2-glycoprotein I and thrombin and are sometimes called lupus anticoagulant because they are present in some patients with systemic lupus erythematosus (SLE) or other autoimmune conditions. Patients with antiphospholipid syndrome have recurrent arterial and venous thrombosis and repeated miscarriages. In vitro, these antibodies inhibit coagulation by interfering with the assembly of phospholipid complexes, and a “mixing study” with normal serum will not correct the PTT (which primarily measures factors II, V, VIII, IX, X, XI, and XII and fibrinogen in the “intrinsic pathway” of in vivo coagulation). In vivo, these antibodies induce a hypercoagulable state by unknown mechanisms. Disseminated intravascular coagulopathy occurs.

PBD9 119, 121  BP9 83–84  PBD8 118–120  BP8 90–91

18 D Tissue plasminogen activator (t-PA) is a thrombolytic agent that promotes generation of plasmin, which cleaves fibrin to dissolve clots. Aspirin prevents formation of new thrombi by inhibiting platelet aggregation and works best as a preventive therapy. Heparin prevents thrombosis by activating antithrombin III. Nitric oxide is a vasodilator. Vitamin K is required for synthesis of clotting factors II, VII, IX, and X and facilitates the ability to form clots.

PBD9 120–121  BP9 85  PBD8 120–121  BP8 87–88

19 B Atherosclerotic damage to vascular endothelium is the most common cause of arterial thrombosis; this damage accumulates almost imperceptibly over many years. Diabetes mellitus types I and II accelerate atherosclerosis. Inhibitors to coagulation, such as antiphospholipid antibodies, typically prolong the partial thromboplastin time (PTT), the prothrombin time (PT), or both. Decreased levels of antithrombin III and mutation in the factor V gene are inherited causes of hypercoagulability; they are far less common than atherosclerosis of coronary vessels. Decreased production of tissue plasminogen activator from intact endothelial cells may occur in anoxia of the endothelial cells in veins with sluggish circulation. Stasis of blood flow is important in thrombosis within the low-pressure venous circulation.

PBD9 122–123  BP9 86  PBD8 125  BP8 94

20 B Recurrent thrombotic episodes at such a young age strongly suggest an inherited coagulopathy. The factor V (Leiden) mutation affects 2% to 15% of the population, and more than half of all individuals with a history of recurrent deep venous thrombosis have such a defect. Inherited deficiencies of the anticoagulant proteins antithrombin III and protein C can cause hypercoagulable states, but these are much less common than factor V mutation. Hyperhomocysteinemia is a less common cause of inherited risk of thrombosis than is factor V mutation. It also is a risk factor for atherosclerosis that predisposes to arterial thrombosis. Although some cancers elaborate factors that promote thrombosis, this patient is unlikely to have cancer at such a young age; a 10-year history of thrombosis is unlikely to occur in a patient with cancer. Oral contraceptive usage contributes to risk for thrombosis, but mainly in older women, particularly past age 40 years. Smoking promotes atherosclerosis with arterial thrombosis.

PBD9 123–124  BP9 87  PBD8 122–123  BP8 95

21 A These findings are characteristic of a hypercoagulable state. The patient has antibodies that react with cardiolipin, a phospholipid antigen used for the serologic diagnosis of syphilis. These so-called antiphospholipid antibodies are directed against phospholipid-protein complexes such as β2-glycoprotein I and thrombin and are sometimes called lupus anticoagulant because they are present in some patients with systemic lupus erythematosus (SLE) or other autoimmune states. Patients with antiphospholipid syndrome have recurrent arterial and venous thrombosis and repeated miscarriages. In vitro, these antibodies inhibit coagulation by interfering with the assembly of phospholipid complexes, and a “mixing study” with normal serum will not correct the PTT (which primarily measures factors II, V, VIII, IX, X, XI, and XII and fibrinogen in the “intrinsic pathway” of in vitro coagulation). In vivo, the antibodies induce a hypercoagulable state by unknown mechanisms. Disseminated intravascular coagulopathy is an acute consumptive coagulopathy characterized by elevated PT and PTT, and decreased platelet count. The PT and PTT are normal in patients with factor V (Leiden) mutation. Hypercholesterolemia promotes atherosclerosis over many years, and the risk of arterial thrombosis increases. Von Willebrand disease affects platelet adhesion and leads to a bleeding tendency, not to thrombosis.

PBD9 124–125  BP9 87–88  PBD8 122–123  BP8 94–95

22 B The figure shows a mural thrombus at the right, filling an atherosclerotic aortic aneurysm below the renal arteries. Diabetes mellitus accelerates and worsens atherosclerosis, including peripheral muscular arteries. One possible complication of mural thrombosis is embolization, which occurs
when a small piece of the clot breaks off. The embolus is carried distally and may occlude the popliteal artery, which is an end artery for the lower leg. A venous thrombus, not arterial, produces leg swelling from edema. Because the thrombus is in the arterial circulation, an embolus would not travel to the lungs. The thrombus is below the renal arteries. Although platelets contribute to the formation of thrombi, the platelet count does not decrease appreciably with formation of a localized thrombus, and a generalized process such as disseminated intravascular coagulation is needed to consume enough platelets to cause thrombocytopenia.

23 B The most important and the most common cause of venous thrombosis is vascular stasis, which often occurs with immobilization. Calcium is a cofactor in the coagulation pathway, but an increase in calcium has minimal effect on the coagulation process. Aspirin inhibits platelet function and limits thrombosis. Nitric oxide is a vasodilator and an inhibitor of platelet aggregation. Turbulent blood flow may promote thrombosis, but this risk factor is more common in fast-flowing arterial circulation.

24 E He has deep venous thrombosis as a consequence of venous stasis from immobilization. The large, deep thrombi in leg veins can embolize to the lungs, leading to death. Disseminated intravascular coagulation is not a common complication in patients with thrombosis of the extremities or in patients recovering from an injury. Fat embolism can occur with fractures, but pulmonary problems typically appear 1 to 3 days after the traumatic event. Gangrene typically occurs from arterial, not venous, occlusion in the leg. Vessels with thrombi typically stay intact; if a hematoma had developed as a consequence of the trauma from the fall, it would be organizing and decreasing in size after 2 weeks.

25 E Venous stasis favors the development of phlebothrombosis (venous thrombosis), particularly in the leg and pelvic veins. This is a common complication in hospitalized patients who are bedridden. The obstruction may produce local pain and swelling, or it may be asymptomatic. Such deep thrombi in large veins create a risk for pulmonary thromboembolism. Phlebothrombosis occurs when stasis in large veins promotes thrombosis formation, typically in leg and pelvic veins; because there is often clinically apparent swelling, warmth, and pain, the term thrombophlebitis is often employed regardless of whether true vascular inflammation is present. The figure shows alternating pink platelet-fibrin and RBC layers (lines of Zahn) at the left. After a thrombus has formed, it may become organized with ingrowth of capillaries, fibroblast proliferation, and macrophage infiltration that eventually clears part or most of the clot, forming one or more new lumens (recanalization) as shown at the right of the figure. Atherosclerosis occurs in arteries, not veins. Chronic passive congestion refers to capillary, sinusoidal, or venous stasis of blood within an organ such as the lungs or liver. Vasculitis typically involves the vascular wall with inflammatory cell infiltrates. Mural thrombi are thrombi that form on the surfaces of the heart or large arteries. A vegetation is a localized thrombus formation on cardiac endothelium, typically a valve.

26 F The figure shows an organizing thrombus in a small artery. Such a peripheral arterial occlusion was insufficient to produce infarction, as evidenced by the lack of enzyme elevation. Thrombi become organized over time if they are not dissolved by fibrinolytic activity. After a thrombus has formed, it may become organized with ingrowth of capillaries, fibroblast proliferation, and macrophage infiltration, which eventually clears part or most of the clot; there can be formation of one or more new lumens (recanalization). Air emboli are uncommon and usually the result of trauma. Air emboli on the arterial side can cause ischemia through occlusion even when very small, whereas on the venous side, more than 100 mL of air trapped in the heart may reduce cardiac output. When gases that became dissolved in tissues at high pressure bubble out at decompression with lower pressure in blood and tissues, then air emboli form. Cholesterol emboli can break off from atherosmas in arteries and proceed distally to occlude small arteries; however, because these emboli are usually quite small, they are seldom clinically significant. Chronic passive congestion refers to capillary, sinusoidal, or venous stasis of blood within an organ such as the lungs or liver. Fat emboli are globules of lipid that are most likely to form after traumatic injury, typically to long bones. Mural thrombi are thrombi that form on the surfaces of the heart or large arteries.

27 F A thrombotic mass that forms on a cardiac valve (or, less commonly, on the cardiac mural endocardium) is known as a vegetation. Such vegetations may produce thromboemboli. Vegetations on the right-sided heart valves may embolize to the lungs; vegetations on the left embolize systemically to organs such as the brain, spleen, and kidney. A so-called paradoxical embolus occurs when a right-sided cardiac thrombus crosses a patent foramen ovale and enters the systemic arterial circulation. Patients with malignant neoplasms may have a hypercoagulable state (Trousseau syndrome) that favors the development of arterial and venous thromboses. Atherosclerosis occurs within arteries, not the chambers of the heart. Endocardial metastases are quite rare. Chronic passive congestion refers to capillary, sinusoidal, or venous stasis of blood within an organ such as the lungs or liver. Mural thrombi are thrombi that form on the surfaces of the heart or large arteries. The term typically is reserved for large thrombi in a cardiac chamber or dilated aorta or large aortic branch; it is not used to describe thrombotic lesions on cardiac valves. A myxoma is a primary neoplasm of the heart that usually arises on an atrial surface; it is not associated with malignancies elsewhere. Phlebothrombosis occurs when stasis in large veins promotes thrombosis formation.
28 D The figure shows a large pulmonary thromboembolus. The most common risk factor is immobilization leading to venous stasis with phlebothrombosis, often called thrombo- phlebitis, since the lower extremities may be swollen and tender, but there is minimal inflammation. These thrombi form in the large deep leg or pelvic veins, not in the pulmonary arteries. Coagulopathies from acquired or inherited disorders, such as those from lupus anticoagulant (antiphospholipid antibodies) or factor V (Leiden) mutation, are possible causes of thrombosis, but they usually manifest at a younger age. These causes also are far less common risks for pulmonary thromboembolism than venous stasis. Local inflammation from pneumonia may result in thrombosis of small vessels in affected peripheral areas of lung. Pulmonary atherosclerosis occurs with long-standing pulmonary hypertension, not the factors driving systemic arterial atherosclerosis.

32 D The figure shows a dark red hemorrhagic infarction extending to the pleura, a typical finding when a medium-sized thromboembolus lodges in a peripheral pulmonary artery branch. The infarct is hemorrhagic because the bronchial arterial circulation in the lung (derived from the systemic arterial circulation and separate from the pulmonary arterial circulation) continues to supply a small amount of blood to the interstitium in the affected area of infarction. Persons with underlying heart or lung disease are at greater risk for pulmonary infarction. Passive congestion, whether acute or chronic, is a diffuse process, as is edema, which does not impart a red color. Pulmonary venous thrombosis is rare.

29 B The mechanism for fat embolism is unknown, in particular, why onset of symptoms is delayed 1 to 3 days after the initial injury (or up to 1 week for cerebral symptoms). The cumulative effect of many small fat globules filling peripheral pulmonary arteries is the same as one large pulmonary thromboembolus. Cardiac tamponade and hemothorax would be immediate complications after traumatic injury, not delayed events. Pulmonary edema severe enough to cause dyspnea would be unlikely to occur in hospitalized patients because fluid status is closely monitored. Pulmonary infarction may cause dyspnea, but pulmonary thromboembolism from deep venous thrombosis is typically a complication of a longer hospitalization.

30 B Amniotic fluid embolism rarely occurs in pregnancy, but it has a high mortality rate. The fluid reaches torn uterine veins through ruptured fetal membranes. Aggregates of platelets represent localized thrombosis, an unlikely event in the lungs. Fat globules are seen in fat embolism, usually after severe trauma. Gas bubbles in vessels from air embolism can be a rare event in some obstetric procedures, but it is an unlikely event in natural deliveries. Peripheral pulmonary thromboemboli are most likely to produce chronic pulmonary hypertension and develop over weeks to months.

31 C These findings are characteristic of decompression sickness (the bends), a form of air embolism. At high pressures, such as occur during a deep scuba dive, nitrogen is dissolved in blood and tissues in large amounts. Ascending too quickly does not allow for slow release of the gas, and formation of small gas bubbles causes symptoms from occlusion of small arteries and arterioles. Fat globules in pulmonary arteries are a feature of fat embolism, which usually follows trauma. Fibrin thrombi may form with widespread activation of coagulation, as with DIC. Platelet thrombi may form with microangiopathic hemolytic anemia, such as thrombotic thrombocytopenic purpura. Atherosclerosis typically occurs in elastic arteries, not arterioles.

33 B The figure shows a pale ischemic infarction of the renal cortex extending nearly to the renal capsule, a typical finding when a medium-sized arterial thromboembolus lodges in a peripheral renal artery branch. The infarct is wedge-shaped, typical for many parenchymal organs, because there is minimal collateral circulation. An abscess is a form of liquefactive necrosis from a localized collection of neutrophils in association with infection, and though it could be yellowish, it is likely to be round. Liquefactive necrosis from arterial occlusion and infarction occurs in the brain. Multiorgan failure occurs with shock, and whole organs are affected by ischemia. Venous thrombosis tends to produce hemorrhagic lesions.

34 D The liver has a dual blood supply, with a hepatic arterial circulation and a portal venous circulation. Infarction of the liver caused by occlusion of hepatic artery is uncommon. Cerebral infarction typically produces liquefactive necrosis. Infarcts of most solid parenchymal organs such as the kidney, heart, and spleen exhibit coagulative necrosis, and emboli from the left heart often go to these organs.

35 B The patient has septic shock from infection with gram-negative organisms that have lipopolysaccharide, which binds along with other microbe-derived substances containing pathogen-associated molecular patterns (PAMPs) to cells via Toll-like receptors (TLRs). Binding initiates release of various cytokines such as tumor necrosis factor (TNF) and interleukin-1 (IL-1) that produce fever. Macrophages are stimulated to destroy the organisms. Nitric oxide is released, promoting vasodilation and circulatory collapse. Complement C3b generated by bacteria via the alternative pathway acts as an opsonin. Platelet-activating factor mediates many features of acute inflammation and in large quantities can cause vasoconstriction and bronchoconstriction. Toxic shock syndrome toxin-1 is a superantigen released by staphylococcal organisms that is a potent activator of T lymphocytes, inducing cytokine release with septic shock.
Progressive septic shock with poor tissue perfusion is evidenced by the high lactate level. Vasodilation is a feature of septic shock, typically as a result of gram-negative endotoxemia. Decreased hematocrit suggests hypovolemic shock from blood loss. Elevated creatine kinase suggests an acute myocardial infarction, which produces cardiogenic shock. Increased blood urea nitrogen concentration is a feature of renal failure, not the cause of renal failure. Reduction in Po2 suggests a problem with lung ventilation or perfusion.

In the nonprogressive phase of shock, neurohumoral mechanisms include catecholamine release, activation of the renin-angiotensin axis, antidiuretic hormone (ADH) release, and generalized sympathetic stimulation with tachycardia, peripheral vasoconstriction, and renal conservation of fluid to support tissue perfusion. As shock progresses, there is hypotension with diminished tissue perfusion, anaerobic glycolysis, and lactic acidosis. An irreversible stage of shock is marked by tissue injury, hypoxia, marked metabolic acidosis, activation of coagulation pathways with consumption of coagulation factors, multiple organ failure, and leakage of lysosomal and cytosolic enzymes, such as lactate dehydrogenase (LDH).
Multiple members of a family have a disease that is associated with a genetic change that involves substitution of adenine for thymine involving one base pair on homologous chromosomes. What is the best term to describe this finding?

A. Copy number variation  
B. Deletion  
C. Epigenetic change  
D. Single nucleotide polymorphism  
E. Trinucleotide repeat mutation  
F. RNA alteration

A 15-year-old girl has developed multiple nodules on her skin over the past 10 years. On physical examination, there are 20 scattered, 0.3-cm to 1-cm, firm nodules on the patient’s trunk and extremities. There are 12 light brown macules averaging 2 to 5 cm in diameter on the skin of the trunk. Slit-lamp examination shows pigmented nodules in the iris. A sibling and a parent are similarly affected. Genetic analysis shows a loss-of-function mutation. Which of the following inheritance patterns is most likely to be present in this family?

A. Autosomal dominant  
B. Autosomal recessive  
C. Mitochondrial  
D. Multifactorial  
E. X-linked recessive

A female infant born at term shows failure to thrive and failure to achieve developmental milestones. A pedigree reveals only this child is affected out of four generations on both sides of the family. Tissue fibroblasts obtained from this child shows a 46,XX karyotype. Cultured fibroblasts show accumulation of an intermediate product in a metabolic pathway in which multiple enzymes are involved. What is the most likely recurrence risk for this condition in siblings of this infant?

A. 3%  
B. 8%  
C. 15%  
D. 25%  
E. 50%  
F. 100%

An 8-year-old girl experiences sudden severe dyspnea. On examination, she has upper airway obstruction from soft tissue swelling in her neck. A radiograph shows a hematoma compressing the trachea. Laboratory studies show her prothrombin time (PT) is normal, but her partial thromboplastin time (PTT) is increased. Further testing reveals less than 1% of normal factor VIII activity. Both parents and two female siblings are unaffected by this problem, but a male sibling has experienced a similar episode. Which of the following genetic abnormalities is most likely to account for the findings in this girl?

A. Autosomal dominant mutation  
B. Genomic imprinting  
C. Germline mosaicism  
D. Random X inactivation  
E. Spontaneous new mutation

A 66-year-old man has been prescribed clopidogrel to help prevent future acute coronary events. He states that his father, aunt, and brother took this drug and had adverse side effects, including excessive bleeding, when prescribed the recommended dose. Which of the following genetic tests will help determine the most appropriate dose of clopidogrel for this man?

A. Enumeration of tandem repeat sequences  
B. Expression profiling of mRNA  
C. FISH analysis for the karyotype  
D. Measurement of an enzyme level  
E. Probing for a cyp450 polymorphism
6. A 22-year-old man has a sudden loss of vision in the right eye. On physical examination, there is a subluxation of the right crystalline lens. On auscultation of the chest, a mid-systolic click is audible. An echocardiogram shows a floppy mitral valve and dilated aortic arch. The patient’s brother and his cousin are similarly affected. He is prescribed a beta-blocker. A genetic defect involving which of the following substances is most likely to be present in this patient?

A. Collagen  
B. Dystrophin  
C. Fibrillin-1  
D. NFI protein  
E. Spectrin

7. An 11-year-old child has exhibited poor wound healing, even with minor trauma, since infancy. On examination she has hyperextensible joints and fragile, extremely stretchable skin. A diaphragmatic hernia was repaired soon after birth. One parent and one of three siblings are also affected. A mutation in a gene encoding for which of the following types of proteins is most likely causing this child’s disease?

A. Enzyme  
B. Growth regulation  
C. Ion channel  
D. Receptor  
E. Structural support

8. A 4-year-old girl has sudden onset of right hip pain. On examination, the child’s right hip is dislocated. The child can bend her thumb backward to touch the forearm. Her skin is noted to be extraordinarily stretchable. Radiographs of her spine show marked lateral and anterior curvature. She develops retinal detachments later in childhood. A sibling is similarly affected. A mutation in tenascin-X is identified. Which of the following mutations most likely occurred?

A. Three-base pair deletion  
B. Point  
C. Nonsense  
D. Frameshift  
E. Trinucleotide repeat

9. A clinical study is undertaken with subjects from families in which complications of atherosclerotic cardiovascular disease and tendinous xanthomas occurred before age 30 years. Some of the children in these families are observed to have early atheroma formation. These affected individuals benefit from treatment with pharmacologic agents that inhibit HMG-CoA reductase. Affected individuals in these families are most likely to have a mutation in a gene encoding a cell surface receptor for which of the following?

A. Cortisol  
B. Insulin  
C. LDL cholesterol  
D. Leptin  
E. TGF-α

10. A 1-year-old female infant has failure to thrive, poor neurologic development, and poor motor function. Physical examination shows a “cherry red” spot on the macula of the retina. The infant’s muscle tone is poor. Both parents and a brother and sister are healthy, with no apparent abnormalities. One brother with a similar condition died at the age of 18 months. This genetic disorder most likely resulted from a mutation involving a gene encoding for which of the following?

A. Mitochondrial enzyme  
B. Lysosomal enzyme  
C. Cell surface receptor protein  
D. Structural protein

11. An infant born to a family living in Belarus appeared normal at birth, but at 6 months is noted to have worsening motor incoordination and mental obtundation. On examination, the infant has retinal pallor with a prominent macula. Flaccid paralysis develops by 1 year of age. Based on the figure, which of the following mutations most likely occurred?

A. Frameshift  
B. Nonsense  
C. Point  
D. Three-base pair deletion  
E. Trinucleotide repeat

12. A 22-year-old woman delivers an apparently healthy female infant after an uncomplicated pregnancy. By 4 years of age, the girl has progressive, severe neurologic deterioration. Physical examination shows marked hepatosplenomegaly. A bone marrow biopsy specimen shows numerous foamy vacuolated macrophages. Analysis of which of the following factors is most likely to aid in the diagnosis of this condition?

A. Level of α₁-antitrypsin in the serum  
B. Level of glucose-6-phosphatase in hepatocytes  
C. Level of sphingomyelinase in splenic macrophages  
D. Number of LDL receptors on hepatocytes  
E. Rate of synthesis of collagen in skin fibroblasts

13. A 2-year-old child has had failure to thrive since birth, with progressive neurologic deterioration. On physical examination, the child has hepatosplenomegaly and lymphadenopathy. Laboratory studies show pancytopenia. A bone marrow biopsy is obtained and microscopically shows numerous pale phagocytic cells filled with fine vacuoles (secondary lysosomes). An abnormality in genetic encoding for which of the following types of protein is most likely causing this child’s disease?

A. Enzyme  
B. Growth regulation  
C. Ion channel  
D. Receptor  
E. Structural support
A 10-year-old child has had recurrent otitis media for the past 8 years. On physical examination, there is hepatosplenomegaly. No external anomalies are present. Laboratory findings include anemia and leukopenia. A bone marrow biopsy is performed, and high magnification of the sample shows the findings depicted in the figure. An inherited deficiency of which of the following enzymes is most likely to produce these findings?

A. Alpha-1, 4-glucosidase  
B. Glucocerebrosidase  
C. Glucose-6-phosphatase  
D. Hexosaminidase A  
E. Lysyl hydroxylase

The parents of a male infant report that male children over three generations in the mother’s family have been affected by a progressive disorder involving multiple organ systems. These children have had coarse facial features, corneal clouding, joint stiffness, hepatosplenomegaly, and mental retardation, and many died in childhood. At autopsy, some of the children had subendothelial coronary arterial deposits that caused myocardial infarction. Laboratory testing of the infant now shows increased urinary excretion of mucopolysaccharides. Bone marrow biopsy is performed, and the accumulated mucopolysaccharides are found in macrophages (“balloon cells” filled with minute vacuoles). Which of the following enzyme deficiencies is most likely to be seen in this infant?

A. Adenosine deaminase  
B. α-1-Liduronidase  
C. Glucocerebrosidase  
D. Glucose-6-phosphatase  
E. Hexosaminidase A  
F. Lysosomal glucosidase  
G. Sphingomyelinase

A 2-year-old child with failure to thrive since infancy now exhibits a seizure. Physical examination shows hepatomegaly and ecchymoses of the skin. Laboratory studies show a blood glucose level of 31 mg/dL. A liver biopsy specimen shows cells filled with clear vacuoles that stain positive for glycogen. Which of the following conditions is most likely to produce these findings?

A. Hurler syndrome  
B. McArdle disease  
C. Pompe disease  
D. Tay-Sachs disease  
E. Von Gierke disease

A 25-year-old woman stops going to her aerobic exercise class because of severe muscle cramps that have occurred during every session for the past 2 months. Four hours after each session, she notices that her urine is a brown color. On physical examination, she has normal muscle development and strength. An inherited defect in which of the following substances is most likely to explain her findings?

A. Dystrophin  
B. Fibrillin  
C. Glucose-6-phosphatase  
D. Lysosomal glucosidase  
E. Muscle phosphorylase  
F. Spectrin

A 6-month-old male infant has failure to thrive and abdominal enlargement. His parents are concerned that he has shown minimal movement since birth. On physical examination, the infant has marked muscle weakness and hepatosplenomegaly. A chest radiograph shows marked cardiomegaly. He dies of congestive heart failure at age 19 months. The microscopic appearance of myocardial fibers at autopsy is shown in the figure. A deficiency of which of the following enzymes is most likely to be present in this infant?

A. Glucocerebrosidase  
B. Glucose-6-phosphatase  
C. Hexosaminidase A  
D. Homogentisic acid oxidase  
E. Lysosomal glucosidase  
F. Sphingomyelinase
A 13-year-old boy has been drinking large quantities of fluids and has an insatiable appetite. He is losing weight and has become more tired and listless for the past month. Laboratory findings include normal CBC and fasting serum glucose of 175 mg/dL. His parents, two brothers, and one sister are healthy. A maternal uncle is also affected. Which of the following is the probable inheritance pattern of his disease?

- A  Autosomal dominant
- B  Autosomal recessive
- C  Mitochondrial DNA
- D  Multifactorial
- E  X-linked recessive

A healthy 20-year-old woman, G3, P2, Ab1, has previously given birth to a liveborn infant and a stillborn infant, both with the same karyotypic abnormality. On physical examination, she is at the 50th percentile for height and weight. She has no physical abnormalities noted. Which of the following karyotypic abnormalities is most likely to be present in this woman?

- A  Deletion of q arm - del(22q)
- B  Isochromosome - 46,X,i
- C  Paracentric inversion - inv(18)
- D  Ring chromosome - r(13)
- E  Robertsonian translocation - t(14;21)

An 11-year-old mentally retarded boy is able to carry out activities of daily living, including feeding and dressing himself. On physical examination, he has brachycephaly and oblique palpebral fissures with prominent epicanthal folds. A transverse crease is seen on the palm of each hand. On auscultation of the chest, there is a grade III/VI systolic murmur. Which of the following diseases is he most likely to develop by age 20 years?

- A  Acute leukemia
- B  Acute myocardial infarction
- C  Aortic dissection
- D  Chronic renal failure
- E  Hepatic cirrhosis

A 23-year-old woman, G2, P1, gives birth at 37 weeks to a small-for-gestational-age male infant. The left hand of an infant showed an endocardial cushion defect and polyhydramnios from probable duodenal atresia. Which of the following chromosomal abnormalities is most likely to be present?

- A  45,X
- B  47,XX,+21
- C  47,XY,+18
- D  69,XXY
- E  47,XXY

A 39-year-old woman gives birth to a term infant with an umbilical hernia, Brushfield spots on the iris, macroglossia, low-set ears, oblique palpebral fissures, and a heart murmur. The infant survives to childhood and exhibits only mild mental retardation. Which of the following chromosomal abnormalities, affecting autosomes, is most likely to be present in the somatic cells of this child?

- A  Haploidy
- B  Monosomy
- C  Mosaicism
- D  Tetraploidy
- E  Triploidy

A 38-year-old woman gives birth at 35 weeks’ gestation to a female infant. Physical examination of the infant soon after delivery shows rocker-bottom feet, a small face and mouth, and low-set ears. On auscultation of the chest, a heart murmur is detected. The appearance of the infant’s hands is shown in the figure. The infant dies at 4 months of age. Which of the following karyotypes was most likely present in this infant?

- A  45,X
- B  46,XX
- C  47,XX,+18
- D  47,XX,+21
- E  48,XXX
**25** A 36-year-old woman gives birth at 34 weeks' gestation to a male infant who lives for only 1 hour after delivery. On physical examination, the infant is at the 30th percentile for height and weight. Anomalies include microcephaly, a cleft lip and palate, scalp defects, and the finding shown in the figure. Which of the following karyotypes is most likely to be present in this infant?

- A 45,X
- B 46,XY
- C 47,XXY
- D 47,XY,+13
- E 47,XY,+18
- F 69,XXY

**26** A 27-year-old primigravida has a fetal ultrasound performed at 18 weeks' gestation. The male fetus is mildly growth retarded. Multiple congenital anomalies are present, including ventricular and atrial septal defects, horseshoe kidney, and omphalocele. Amniocentesis is performed, and the fetal cells obtained are examined using FISH analysis using a probe for one of the chromosomes. Based on the findings shown in the figure, which of the following karyotypic abnormalities is most likely to be present in this fetus?

- A 45,X/46,XX
- B 46,XY,del(22q11)
- C 46,XY,der(14;21)(q10.0),+21
- D 47,XY,+18
- E 47,XXY

**27** A 12-year-old boy has a cough and earache for the past 2 days. He has a history of recurrent infections, including otitis media, diarrhea, and pneumonia. Physical examination shows an erythematous right tympanic membrane, a cleft palate, and murmur suggestive of congenital heart disease. A thoracic CT scan shows a small thymus. Results of laboratory studies suggest mild hypoparathyroidism. Which of the following diagnostic studies is most likely to be helpful in diagnosing this patient's condition?

- A Adenosine deaminase assay in lymphocytes
- B Branched DNA assay for HIV-1 RNA level
- C FISH analysis with a probe for chromosome 22q11.2
- D Lymph node biopsy
- E PCR analysis for trinucleotide repeats on the X chromosome

**28** A 9-month-old infant has had numerous viral and fungal infections since birth. On physical examination, no congenital anomalies are noted. Laboratory studies show hypocalcemia. FISH analysis of the infant's cells is performed. A metaphase spread is shown in the figure, with probes to two different regions on chromosome 22. Which of the following cytogenetic abnormalities is most likely to be present?

- A Hyperdiploidy
- B Deletion
- C Inversion
- D Monosomy
- E Translocation

**29** A 27-year-old man and his 24-year-old wife have been trying to conceive a child for 6 years. Physical examination shows he has bilateral gynecomastia, reduced testicular size, reduced body hair, and increased length between the soles of his feet and the pubic bone. A semen analysis indicates oligosperma. Laboratory studies show increased follicle-stimulating hormone level and slightly decreased testosterone level. Which of the following karyotypes is this man most likely to have?

- A 46,X,i(Xq)
- B 47,XY
- C 47,XXY
- D 46,XX/47,XX,+21
- E 46,XY,del(22q11)
30. A 25-year-old woman with amenorrhea has never had menarche. On physical examination, she is 145 cm (4 ft 9 in) tall. She has a webbed neck, a broad chest, and widely spaced nipples. Strong pulses are palpable in the upper extremities, but there are only weak pulses in the lower extremities. On abdominal MR imaging, her ovaries are small, elongated, and tubular. Which of the following karyotypes is she most likely to have?

A. 45,X/46,XX  
B. 46,X,X(fra)  
C. 47,XXY  
D. 47,XXX  
E. 47,XX,+16

31. A 22-year-old primigravida notes absent fetal movement for 2 days. The fetus is delivered stillborn at 19 weeks' gestation. The macerated fetus shows marked hydrops fetalis and a large posterior cystic hygroma of the neck. At autopsy, internal anomalies include aortic coarctation and a horseshoe kidney. Which of the following karyotypes is most likely to be present in cells obtained from this fetus?

A. 45,X  
B. 47,XX,+18  
C. 47,XX,+21  
D. 47,XY  
E. 69,XXX

32. A 23-year-old woman gives birth to a term infant after an uncomplicated pregnancy. On physical examination, the infant has ambiguous external genitalia. The parents want to know the infant’s sex, but the physician is hesitant to assign a sex without further information. A chromosomal analysis indicates a karyotype of 46,XX. An abdominal CT scan shows bilaterally enlarged adrenal glands, and the internal genitalia appear to consist of uterus, fallopian tubes, and ovaries. This clinical picture is most consistent with which of the following conditions?

A. Androgen insensitivity syndrome  
B. Excessive trinucleotide repeats  
C. Female pseudohermaphroditism  
D. Mitochondrial DNA mutation  
E. Nondisjunctional event with loss of Y chromosome

33. Mental retardation has affected several generations of a family, and most of the affected individuals have been males. The severity of mental retardation has increased with each passing generation. Genetic testing is performed, and about 20% of the males who have the genetic abnormality are unaffected. Which of the following mechanisms is most likely to produce this genetic condition?

A. Frameshift mutation  
B. Missense mutation  
C. Mitochondrial DNA mutation  
D. Point mutation  
E. Trinucleotide repeat mutation

34. A pregnant woman with a family history of fragile X syndrome undergoes prenatal testing of her fetus. PCR analysis to amplify the appropriate region of the FMR1 gene is attempted using DNA from amniotic fluid cells, but no amplified products are obtained. Which of the following is the most appropriate next step?

A. Routine karyotyping of the amniotic fluid cells  
B. Routine karyotyping of the unaffected father  
C. Southern blot analysis of DNA from the amniotic fluid cells  
D. PCR analysis of the mother’s FMR1 gene  
E. No further testing

35. A 19-year-old man has experienced failing eyesight and progressive muscle weakness for the past 7 years. Family history reveals that several of the patient’s male and female relatives have similar symptoms. His mother, her brother and sister, and two of the aunt’s children are affected, but the uncle’s children are not. Which of the following types of genetic disorders is most likely to be present in this patient?

A. Trinucleotide repeat expansion  
B. Genetic imprinting  
C. X-linked inheritance pattern  
D. Mitochondrial mutation  
E. Uniparental disomy

36. A 3-year-old boy has had progressive developmental delay, ataxia, seizures, and inappropriate laughter since infancy. The child has a normal karyotype of 46,XY, but DNA analysis shows that he has inherited both of his number 15 chromosomes from his father. These findings are most likely to be indicative of which of the following genetic mechanisms?

A. Genomic imprinting  
B. Maternal inheritance pattern  
C. Mutation of mitochondrial DNA  
D. Trinucleotide repeat expansion  
E. X-linked inheritance pattern

37. Three female children in a family with the pedigree shown in the figure are noted to have histories of multiple fractures along with dental problems and hearing impairment. On examination, they are of normal height and weight for age, but have steel gray to blue sclerae. Both parents are unaffected by these abnormalities. Which of the following genetic abnormalities is most likely to account for the findings in these children?

A. Genomic imprinting  
B. Gonadal mosaicism  
C. Multifactorial inheritance  
D. Random X inactivation  
E. Spontaneous new mutation
A study of families with fragile X syndrome reveals that less than 1 in 10,000 is a heterozygote. Sensitivity of mutations among ethnic groups limits detection and is expensive. Fluorescence in situ hybridization is labor-intensive and expensive. Both copies of the gene must be abnormal for detection. Fluorescence in situ hybridization is labor-intensive and expensive. Frequency of mutations among ethnic groups limits sensitivity. Less than 1 individual in 10,000 is a heterozygote. Most mutations cannot be detected by PCR.

A 32-year-old woman has had three pregnancies, all ending in stillbirths in the first trimester. On physical examination, she and her only spouse for all pregnancies have no abnormalities. Which of the following laboratory tests is most appropriate to perform on this woman for elucidating potential causes for recurrent fetal loss?
- A Genome-wide association study
- B Fluorescence in situ hybridization
- C Karyotyping
- D PCR analysis
- E Tandem mass spectroscopy

A case-control study is performed involving persons diagnosed with essential hypertension. Genetic analysis reveals linkage disequilibrium. Haplotypes of affected persons differ from the controls in the chromosome containing the angiotensinogen gene. Which of the following types of genetic analysis is most likely to yield this information?
- A Fluorescence in situ hybridization
- B Giemsa banded karyotyping
- C RNA expression
- D Single nucleotide polymorphisms
- E Southern blotting

A pedigree reveals that multiple family members over four generations have been affected by the onset of congestive heart failure within the first four decades of life. A cardiomyopathy is suspected, but specific features of the disease are not known, and no prior genetic testing has been performed. Which of the following techniques involving DNA sequencing is most likely to identify a specific mutation in a cost-effective manner?
- A Pyrosequencing
- B Sanger sequencing
- C Targeted sequencing
- D Whole exome sequencing
- E Whole genome sequencing

ANSWERS

1. D Single nucleotide polymorphisms (SNPs) are found in less than 0.5% of the genome, and only 1% of these are found in coding regions that affect protein synthesis. Some of these account for point mutations that may be associated with disease conditions. C number variations (CNVs) involve variations in large contiguous regions of DNA from 1000 to a million base pairs. Epigenetic changes involve modulation of gene expression without any change in the DNA. Trinucleotide repeats involve increased numbers of base pairs. RNA alterations may modulate DNA expression, such as noncoding micro RNAs.

2. A Neurofibromatosis type 1 (NF-1) is characterized by the development of multiple neurofibromas and pigmented skin lesions. Neurofibromas are most numerous in the dermis but also may occur in visceral organs. Patients with NF-1 also may develop a type of sarcomatous neoplasm known as a malignant peripheral nerve sheath tumor (MPNST). NF-1 is a tumor suppressor that appears with an autosomal dominant pattern of inheritance, though some cases result from spontaneous new mutations (no prior family members with the mutation). NF-1 exhibits variable expressivity, because the manifestations (location and types of neoplasms) are not the same in all patients. The other forms of inheritance listed are not associated with tumor suppressor genes.

3. D Most inborn errors of metabolism involve mutations in genes encoding for enzymes. Because one active allele produces...
half the needed enzyme, this is likely sufficient to avoid disease. Inheritance of two mutant alleles, one from each parent, is required for appearance of disease, so the pattern is autosomal recessive, and the recurrence risk is 25%. Most autosomal recessive genes are infrequent in the population, so a family history is unlikely. Even if 1 in 10 persons carries the mutant recessive gene, a homozygote will be 1 in 400. The standard recurrence risk for any pregnancy is 3%. The recurrence risk is increased to 7% in diseases such as diabetes mellitus, or when a syndrome is identified without a defined inheritance pattern, or with multifactorial inheritance. Autosomal dominant conditions usually result from mutations in genes encoding for structural genes and have a recurrence risk of 50%.

4 D This girl has features of hemophilia A. This X-linked recessive condition is expected to occur in males who inherit the one maternal X chromosome with the genetic mutation, and they do not have another X chromosome with a normal functional allele, as is the case in her brother. Hemophilia in a female can be explained by the Lyon hypothesis, which states that only one X chromosome in a female is active (the “turned off” X chromosome is the Barr body) for most genes, but this inactivation is a random event. Some unlucky females are out on the tail end of the Poisson distribution of random events and have few active X chromosomes with the normal allele, leading to markedly diminished factor VIII activity. The other choices do not explain this phenomenon.

5 E Pharmacogenomic testing may reveal polymorphisms that affect drug metabolism. The cytochrome P-450 system in hepatocytes is involved in metabolism of many substances, including drugs. Detection of polymorphisms can provide information about modification of drug dosing, or the need for a different drug. Trinucleotide repeat mutations may affect gene expression, such as the huntingtin gene, but unlike those involved in drug metabolism. Changes in DNA lead to alterations in mRNA expression, but detecting the DNA alterations directly tends to be a more effective testing strategy. FISH analysis is useful in identifying altered regions of chromosomes, not single genes abnormalities. Autosomal recessive conditions may affect enzyme levels, such as G6PD, but polymorphisms may not affect enzyme or substrate levels that can be reliably detected.

6 C Marfan syndrome is an autosomal dominant condition that is most often caused by qualitative defects in fibrillin from missense mutations in the fibrillin (FBN1) gene. An abnormal collagen gene can cause osteogenesis imperfecta and Ehlers-Danlos syndrome. Genetic mutations in the dystrophin gene are involved in Duchenne and Becker muscular dystrophies. The NFI protein is abnormal in neurofibromatosis type 1. Disordered spectrin causes hereditary spherocytosis.

7 E This classical form of Ehlers-Danlos syndrome (EDS) results from an abnormality in collagen synthesis, and lack of normal collagen affects connective tissues in skin, bone, eye, and vasculature. There are multiple forms of EDS from different mutations affecting different aspects of collagen synthesis. The inheritance pattern described here is autosomal dominant, typical for inherited defects in structural proteins. Inborn errors of metabolism involving enzymes typically have an autosomal recessive pattern of inheritance, because half of gene function is sufficient to prevent disease. Genes involved in growth regulation are typically protooncogenes and tumor suppressor genes, and mutations may underlie development of malignancies. Cystic fibrosis results from mutations in the CFTR gene that encodes chloride ion channels. Familial hypercholesterolemia is an example of a disease resulting from an abnormal LDL receptor.

8 B The joints are frequently involved in most variants of Ehlers-Danlos syndrome (EDS), and tensile strength is reduced so that skin is hypextensible, and joints are hypermobile. Deficiency of the enzyme lysyl hydroxylase can lead to defects in types I and III collagen and is inherited as an autosomal recessive disorder. Kyphoscoliosis and ocular problems also are present in this type of EDS. When EDS-like features are present, but no collagen gene mutations identified, then abnormal tenascin-X, a large multimeric protein of extracellular matrix that affects synthesis and fibril formation of type VI and type I collagens may be present. Congenital syphilis can produce abnormalities of bone, such as saber shin from periostitis and perichondritis, but does not affect the skin. Vitamin D deficiency in childhood producing rickets is accompanied by bowing deformities of long bones, but not skin abnormalities. The pattern of findings here suggests a structural gene defect leading to development of abnormalities, and not congenital anomalies without a specific cause. Battered children typically have multiple contusions and fractures, but the skin and bone structure are normal.

9 C Familial hypercholesterolemia results from mutations in the LDL receptor gene, causing plasma LDL cholesterol to increase because it is not catabolized or taken up by the liver. It is an autosomal dominant disorder with a carrier rate of 1 in 500, so the frequency of homozygosity is 1 in 1 million. Heterozygotes have total serum cholesterol levels twice normal; homozygotes have levels even higher, with death from myocardial infarction by the second decade. The statin drugs inhibit the HMG-CoA reductase and reduce cholesterol levels in heterozygotes. Steroid hormone receptors, such as those for cortisol, are located in the cell nucleus. Insulin receptors play a role in glucose metabolism and glycemic control that may be part of diabetes mellitus with risk for atherosclerosis; statin drugs have no effect on diabetes mellitus. Abnormal leptin receptors may play a role in some forms of obesity. TGF-α is a growth factor with a role in inflammation, cell proliferation, and repair.
10 B The findings listed suggest a severe inherited neurologic disease, and the pattern of inheritance (e.g., normal parents, an affected sibling) is consistent with an autosomal recessive disorder. This inheritance pattern and the cherry red spot in the retina are characteristic of Tay-Sachs disease, caused by mutations in the gene that encodes a lysosomal enzyme hexosaminidase A. Mitochondrial genes have a maternal pattern of transmission. Mutations in genes affecting receptor proteins and structural proteins typically give rise to an autosomal dominant pattern of inheritance.

PBD9 149–152  BP9 229–230  PBD8 150–152  BP8 235–237

11 A The infant has Tay-Sachs disease, an autosomal recessive condition that has a gene frequency higher in some populations, including Ashkenazi Jews and Quebecois. The gene that encodes for the enzyme hexosaminidase A in the diagram has a four-base pair insertion, leading to an altered reading frame (frameshift) and appearance of a stop codon that prematurely terminates reading to produce a nonfunctional enzyme. A frameshift mutation changes the remaining sequence of amino acids in a protein. A point mutation may change the codon to the sequence of a stop codon that truncates the protein being synthesized, typically leading to degradation of the protein. A point mutation typically is a missense mutation that leads to replacement of just one amino acid for another in the protein chain; this can lead to abnormal conformation and function of the protein. A deletion of three base pairs leads to loss of a single amino acid in a protein. A three-base pair deletion, as occurs in cystic fibrosis, results in a frameshift involving just a single amino acid. A trinucleotide repeat is the inheritance pattern for fragile X syndrome, which is caused by triple repeat expansions in the FMR1 gene.

PBD9 151–152  BP9 229–230  PBD8 150–152  BP8 235–237

12 C The clinical features of this child—neurologic involvement, hepatosplenomegaly, and accumulation of foamy macrophages—suggest a lysosomal storage disorder. One such disorder, with which the clinical history is quite compatible, is Niemann-Pick disease type A. It is characterized by lysosomal accumulation of sphingomyelin owing to a severe deficiency of sphingomyelinase. Globules of α₁-antitrypsin are seen in the liver cells of individuals with inherited deficiency of α₁-antitrypsin. The glycogen storage disease known as von Gierke disease results from glucose-6-phosphatase deficiency. In familial hypercholesterolemia, there are fewer LDL receptors on hepatocytes, leading to early and accelerated atherosclerosis by young adulthood. Collagen synthesis is impaired in individuals with Ehlers-Danlos syndrome.

PBD9 152–153  BP9 230–231  PBD8 152–153  BP8 237

13 A This child has Niemann-Pick disease type A, the worst form. Death typically ensues by age 3 years. This inborn error of metabolism occurs in a degradation pathway for sphingomyelin, and accumulation of the abnormal lipid intermediate leads to lysosomal storage disease, mainly affecting tissues of the mononuclear phagocyte system, as well as the central nervous system. Genes involved in growth regulation are typically proto-oncogenes and tumor suppressor genes, and mutations may underlie development of malignancies. Cystic fibrosis results from mutations in the CFTR gene that encodes chloride ion channels. Familial hypercholesterolemia is an example of a disease resulting from an abnormal LDL receptor. Many autosomal dominant conditions, such as Marfan syndrome from fibrillin-1 gene mutations, involve structural protein abnormalities.

PBD9 152–153  BP9 230–231  PBD8 150–152  BP8 235–237

14 B Gaucher disease type 1, seen in this child, accounts for 99% of cases and does not involve the central nervous system (CNS). It is caused by a deficiency of glucocerebrosidase, and infusion with this enzyme reduces severity and progression. Type 2 involves the CNS and is lethal in infancy. Type 3 also involves the CNS, although not as severely as type 2. A deficiency of alpha-1,4-glucosidase is a feature of Pompe disease. Von Gierke disease results from deficiency of glucose-6-phosphatase. Sphingomyelinase deficiency leads to Niemann-Pick disease types A and B. Type A, the more common form, is associated with severe neurologic deterioration. Type B, the less common form, may resemble the findings in this case, but the appearance of macrophages is different: they contain many small vacuoles. Tay-Sachs disease involves a deficiency of hexosaminidase A and is associated with severe mental retardation and death before 10 years of age. Lysyl hydroxylase deficiency is found in one form of Ehlers-Danlos syndrome.


15 B Hunter syndrome, one of the mucopolysaccharidoses (MPS), results from deficiency of the lysosomal enzyme α-l-iduronidase. The glycosaminoglycans that accumulate in MPS include dermatan sulfate, heparan sulfate, keratan sulfate, and chondroitin sulfate. All of the MPS variants are autosomal recessive except for Hunter syndrome, which is X-linked recessive. Adenosine deaminase deficiency is a cause of severe combined immunodeficiency (SCID), an immunodeficiency state in which multiple recurrent infections occur after birth. Glucocerebrosidase deficiency is seen in Gaucher disease; in the most common form of the disease, there is no neurologic impairment, and patients have splenomegaly and skeletal disease as a consequence of increased lysosomal glucocerebrosides in cells of the mononuclear phagocyte system. Glucose-6-phosphatase deficiency leads to von Gierke disease, characterized by hepatomegaly, remegaly, and impaired gluconeogenesis leading to hypoglycemia and hyperlipidemia. Hexosaminidase A deficiency occurs in Tay-Sachs disease; affected individuals manifest severe neurologic impairment, poor motor development, and blindness beginning in infancy. Lysosomal glucosidase deficiency, seen in Pompe disease, is associated with marked cardiomegaly and heart failure beginning in infancy. Sphingomyelinase deficiency occurs in Niemann-Pick disease type A, characterized by hepatosplenomegaly, lymphadenopathy, and severe motor and mental impairment.

PBD9 154–155  BP9 232  PBD8 154–155  BP8 238–239

16 E With von Gierke disease, from deficiency of glucose-6-phosphatase, stored glycogen is not metabolized readily to glucose. Affected individuals have severe hypoglycemia,
which leads to convulsions. Intracytoplasmic accumulations of glycogen occur mainly in the liver and kidney. In Hurler syndrome, the enzyme α-L-iduronidase is deficient. Affected children have skeletal deformities and a buildup of mucopolysaccharides in endocardium and coronary arteries, leading to heart failure. Another form of glycogen storage disease, McArdle disease, results from a deficiency of muscle phosphorylase and leads to muscle cramping. Cardiomegaly and heart failure mark Pompe disease, the type II form of glycogen storage disease. Tay-Sachs disease is characterized by a deficiency in hexosaminidase A and results in severe neurologic deterioration.

17 E McArdle disease is a form of glycogen storage disease in which a deficiency of muscle phosphorylase enzyme causes glycogen to accumulate in skeletal muscle. Onset is in young adulthood. Because strenuous exercise requires glycogenolysis and use of anaerobic metabolism, muscle cramps ensue, but the blood lactate level does not rise. Myoglobinuria is seen in about half of cases. A lack of dystrophin, a protein that stabilizes muscle membrane, characterizes Duchenne muscular dystrophy. A fibrillin gene mutation can lead to Marfan syndrome. Glucose-6-phosphatase deficiency leads to von Gierke disease, characterized by hepatomegaly, renomegaly, and impaired gluconeogenesis leading to hypoglycemia and hyperlipidemia. Lysosomal glucosidase deficiency is seen in Pompe disease, characterized by marked cardiomegaly and heart failure beginning in infancy. Abnormal spectrin, a RBC membrane cytoskeletal protein, leads to a condition known as hereditary spherocytosis.

18 E Pompe disease is a form of glycogen storage disease that results from a deficiency in lysosomal glucosidase (alpha-1,4-glucosidase). The glycogen stored in the myocardium results in massive cardiomegaly and heart failure within 2 years. Glucocerebrosidase deficiency occurs in Gaucher disease. In the most common form of the disease, there is no neurologic impairment, and patients have splenomegaly and skeletal disease as a consequence of increased lysosomal glucocerebrosides in cells of the mononuclear phagocyte system. Glucose-6-phosphatase deficiency leads to von Gierke disease, characterized by hepatomegaly, renomegaly, and impaired gluconeogenesis leading to hypoglycemia and hyperlipidemia. Hexosaminidase A deficiency occurs in Tay-Sachs disease and is associated with severe neurologic impairment, poor motor development, and blindness beginning in infancy. Homogentisic acid oxidase deficiency leads to alkaptonuria with ochronosis or to deposition of a blue-black pigment in joints, resulting in arthropathy. Sphingomyelinase deficiency occurs in Niemann-Pick disease; affected individuals with type A have hepatosplenomegaly, lymphadenopathy, and severe motor and mental impairment.

19 D Type 1 diabetes mellitus has an increased frequency in some families, but the exact mechanism of inheritance is unknown. The risk is increased for offspring when first-order relatives are affected. HLA-linked genes and other genetic loci and environmental factors are considered important. This pattern of inheritance is multifactorial. The other listed inheritance patterns are not seen with most cases of diabetes mellitus.

20 E Almost all of the normal genetic material is present in the case of a Robertsonian translocation because only a small amount of the p arm from each translocated chromosome is lost. The paternal karyotype is 45,XX,t(14;21). Statistically, one of six fetuses in a mother who carries a Robertsonian translocation will also be a carrier. In balanced reciprocal translocation, the same possibility of inheriting the defect exists. The other listed structural abnormalities are likely to result in loss of significant genetic material, reducing survivability, or to interfere with meiosis.

21 A Down syndrome (trisomy 21) is one of the trisomies that can result in a live-born infant. Although children with Down syndrome can function well, they often have many associated congenital anomalies. Among the more common is congenital heart disease, including ventricular septal defect. There is also a tenfold to twentyfold increased risk of acute leukemia. Virtually all individuals with Down syndrome who live to age 40 years have evidence of Alzheimer disease. Myocardial infarction at a young age suggests familial hypercholesterolemia. Aortic dissection is seen in individuals with Marfan syndrome. Chronic renal failure may be seen in genetic disorders that produce polycystic kidneys. Hepatic cirrhosis is a feature of galactosemia.

22 B The figure shows a single palmar flexion crease and a single flexion crease on the fifth digit, both features of trisomy 21. Although there is an increased risk of Down syndrome with increasing maternal age, most infants with Down syndrome are born to younger women because there are far more pregnancies at younger maternal ages. Monosomy X may be marked by a short fourth metacarpal. With trisomy 18, the fingers are often clenched, with digits 2 and 5 overlapping digits 3 and 4. Triploidy may be marked by syndactyly of digits 3 and 4. There are no characteristic hand features in males with Klinefelter syndrome.

23 C Though these features are characteristic of trisomy 21, the child is not severely affected, which suggests mosaicism. In mosaic individuals, greater numbers of potentially normal cells having the proper chromosomal complement are present, which may allow infants with abnormalities of chromosome number to survive to term and beyond. Haploidy is present in gametes. Loss of an autosomal chromosome is devastating; the only monosomy associated with possible survival to term is Turner syndrome (monosomy X). Most aneuploid conditions (trisomies and monosomies) lead to
fetal demise; fetuses with trisomy 21 are the most likely to survive to term. Triploid fetuses rarely survive beyond the second trimester and are virtually never live-born. Likewise, tetraploidy accounts for many first-trimester fetal losses and is not survivable.

24 C Trisomy 18 (Edwards syndrome), in which survival is shortened significantly, has this spectrum of findings. Turner syndrome (45,X) is associated with the presence of cystic hygroma and hydrops fetalis. The severe anomalies described in this case make it unlikely that a normal 46,XX karyotype is present. Down syndrome (47,XX,+21) is associated with longer survival than described in this case, and the external features can be quite subtle at birth. The “superfemale” karyotype (XXX) leads to mild mental retardation. Generally, abnormal numbers of sex chromosomes are tolerated better than abnormalities of autosomes.

25 D Features of trisomy 13 (Patau syndrome) include cleft lip and palate, along with microcephaly, scalp defects, and postaxial polydactyly (an extra digit, shown in the figure). These infants also commonly have severe heart defects, and may also have cyclopia and holoprosencephaly. For monosomy X (45,X) to be considered, the infant must be female. The severe anomalies described in this case occur with a normal karyotype (46,XY), but the spectrum of findings, particularly the polydactyly, suggests trisomy 13. Klinefelter syndrome (47,XXY) results in phenotypic males who are hard to distinguish from males with a 46,XY karyotype. Infants with trisomy 18 lack polydactyly and are more likely to have micrognathia than are infants with trisomy 13. Triploidy with 69 chromosomes leads to stillbirth in virtually all cases.

26 D The infant has findings associated with trisomy 18. In the FISH analysis shown, the chromosomes in each cell have been painted with a marker for chromosome 18. In this case, there are three markers per cell, consistent with a trisomy. In reality, many cells would have to be counted to allow for artifacts in preparation. In most cases, trisomy 18 results from nondisjunctional events. Most infants with trisomy 18 are stillborn, and survival beyond 4 months is rare. The other listed options do not account for this FISH analysis or for this spectrum of anomalies.

27 C DiGeorge syndrome is an immunodeficiency characterized by infection, a small thymus, congenital malformations, and hypoparathyroidism. This cluster is characteristic of the 22q11.2 deletion syndrome, readily diagnosed by FISH. Adenosine deaminase deficiency can cause immunodeficiency, but it is not associated with congenital malformations. Branched DNA assay can detect HIV infection that can lead to AIDS, but no congenital anomalies are associated with this condition. A lymph node biopsy may show a reduction in T cells or B cells associated with various forms of immunodeficiency, but this is not a specific test that can aid in confirming a specific diagnosis. Trinucleotide repeats of the X chromosome, detected by PCR, are seen in fragile X syndrome, which manifests with mental retardation in males.

28 B The infant has DiGeorge syndrome, resulting from a chromosome 22q11.2 microdeletion. This is indicated in the metaphase spread by the presence of only three dots because this region is deleted on one chromosome 22, but both number 22 chromosomes are present. Hyperdiploidy is more than 46 chromosomes. With aneuploidy, there is an abnormal number of chromosomes (trisomy, monosomy), and loss of autosomes tends to produce fetal loss except for some cases of trisomies 13, 18, and 21 and monosomy X. A chromosome inversion would shift the marked region to a different part of the same chromosome. In monosomy, only one of a pair of chromosomes is present. A translocation is the swapping of genetic material between two chromosomes.

29 C Klinefelter syndrome is a relatively common, chromosomal abnormality that occurs in about 1 of 660 live-born males. The findings can be subtle. The 46,XY,i(Xq) karyotype is a variant of Turner syndrome (seen only in females), caused by a defective second X chromosome. The 47,XXY karyotype occurs in about 1 in 1000 live-born males and is associated with taller-than-average stature. A person with a mosaic such as 46,XX/47,XX,+21 has milder features of Down syndrome than a person with the more typical 47,XX,+21 karyotype. The 22q11 deletion syndrome is associated with congenital defects affecting the palate, face, and heart and, in some cases, with T cell immunodeficiency.

30 A The features described are those of classic Turner syndrome. Individuals who reach adulthood may have mosaic cell lines, with some 45,X cells and some 46,XX cells. A female carrier of the fragile X syndrome, X(fra), is less likely to manifest the disease than a male, but the number of triple repeat sequences (CGG) increases in her male offspring. The 47,XY karyotype occurs in Klinefelter syndrome; affected individuals appear as phenotypic males. The “superfemale” karyotype (XXX) leads to mild mental retardation. Trisomy 16 is a cause of fetal loss early in pregnancy.

31 A The findings listed are characteristic of Turner syndrome (monosomy X), which accounts for many first-trimester fetal losses. The hygroma is quite suggestive of this disorder. Fetuses with this finding are rarely live-born. Trisomy 18 can be marked by multiple anomalies, but overlapping fingers and a short neck are more typical features. Down syndrome (47,XX,+21) may be accompanied by a hygroma and hydrops, but ventricular septal defect is more frequent than coarctation, and horseshoe kidney is uncommon.
The 47,XY karyotype (Klinefelter syndrome) does not result in stillbirth, but these males have no major congenital defects. Triploidy with 69 chromosomes typically leads to fetal loss, but hydrops and hygroma are not features of this condition.

**32 C** Physicians must be cautious in assigning sex to an infant with ambiguous genitalia; changing one’s opinion is about as popular as an umpire changing the call. True hermaphroditism, with ovarian and testicular tissue present, is very rare. This infant has female pseudohermaphroditism, resulting from exposure of the fetus to excessive androgenic stimulation, which in this case is due to congenital adrenal hyperplasia. The gonadal and the karyotypic sex are female. Male pseudohermaphroditism has various forms, but the most common is androgen insensitivity; affected individuals are phenotypically females, but have testes and a 46,XY karyotype. Nondisjunctional events lead to monosomies or trisomies, and most result in fetal loss. Trinucleotide repeats are seen in males with fragile X syndrome. Abnormalities of mitochondrial DNA have a maternal transmission pattern and do not involve sex chromosomes or sexual characteristics.

**33 E** Fragile X syndrome is a condition in which there are 250 to 4000 tandem repeats of the trinucleotide sequence CGG. Generally, as the number of trinucleotide repeats increases, the manifestations of the associated conditions worsen or have an earlier onset. The trinucleotide mutations are dynamic; because their number increases during oogenesis, subsequent male offspring have more severe disease compared with earlier generations. With a frameshift mutation, one, two, or three nucleotide base pairs are inserted or deleted. As a result, the protein transcribed is abnormal. A missense mutation results from a single nucleotide base substitution, and it leads to elaboration of an abnormal protein. Abnormalities of mitochondrial DNA, typically involving genes associated with oxidative phosphorylation, are transmitted on the maternal side. A point mutation of a single base pair may affect a single protein.

**34 C** Failure to find amplified product by PCR analysis in such a case could mean that the fetus is not affected, or that there is a full mutation that is too large to be detected by PCR. The next logical step is a Southern blot analysis of genomic DNA from fetal cells. Routine karyotyping of the amniotic fluid cells is much less sensitive than a Southern blot analysis. Karyotyping of the unaffected father cannot provide information about the status of the FMR1 gene in the fetus because amplification of the trinucleotide occurs during oogenesis. For the same reason, PCR analysis of the mother’s FMR1 gene is of no value.

**35 D** This is a classic pattern of maternal inheritance resulting from a mutation in mitochondrial DNA. Males and females are affected, but affected males cannot transmit the disease to their offspring. Because mitochondrial DNA encodes many enzymes involved in oxidative phosphorylation, mutations in mitochondrial genes exert their most deleterious effects on organs most dependent on oxidative phosphorylation, including the central nervous system and muscles. The other listed options do not exhibit strict maternal inheritance.

**36 A** This child has features of Angelman syndrome, and the DNA analysis shows uniparental disomy. The Angelman gene encoded on chromosome 15 is subject to genomic imprinting. It is silenced on the paternal chromosome 15, but is active on the maternal chromosome 15. If the child lacks maternal chromosome 15, there is no active Angelman gene in the somatic cells. This gives rise to the abnormalities typical of this disorder. The same effect occurs when there is a deletion of the Angelman gene from the maternal chromosome 15. The other listed options do not occur in uniparental disomy.

**37 B** The appearance of multiple siblings with a similar condition known to be autosomal dominant, such as osteogenesis imperfecta type I, when both parents are phenotypically normal, suggests that one parent has a mutation confined to gonadal germ cells. In this case the affected parent is a mosaic, and somatic cells do not carry the mutation so there is no phenotypic expression. Genomic imprinting in uniparental disomy is a feature of Angelman syndrome. Multifactorial inheritance is a feature of diseases without a defined inheritance pattern, not osteogenesis imperfecta. Random X inactivation may account for females exhibiting features of an X-linked recessive condition, such as hemophilia A, but this is unlikely to occur three times in the same generation. A spontaneous new mutation can account for one child having an autosomal dominant condition that a parent does not, but the rarity of this event makes such events in a generation highly unlikely.

**38 E** FMR1 protein (FRMP) normally acts as a translation regulator to reduce protein synthesis at synaptic junctions. Thus a reduction in FMRP (loss of function) in the classic fragile X syndrome results in increased protein translation from mRNAs to alter synaptic activity and cause mental retardation. In the fragile X tremor/ataxia syndrome, the FMR1 gene is not methylated and silenced but continues to be transcribed, and CGG-containing FMR1 mRNAs accumulate in the neuronal nucleus and form intranuclear inclusions of aggregated mRNA that sequester RNA-binding proteins, leading to events that are toxic to the cell. Inactivation of the X chromosome occurs when more than one X chromosome is present, such as in a normal female or Klinefelter male.
39 **C** Single-nucleotide polymorphisms occur at a frequency of approximately 1 nucleotide in every 1000–base pair stretch and can be used in linkage analysis for identifying haplotypes associated with disease. Proto-oncogenes are genes encoding for proteins involved in cellular growth; mutant alleles of proto-oncogenes are called oncogenes and play a role in neoplasia. Robertsonian translocations involve portions of two chromosomes that trade places, but are not completely lost, and are balanced; carriers may not be affected, but gametes have the potential to produce monosomies and trisomies. Frameshift deletions are a form of mutation that can lead to abnormal proteins and to conditions such as cystic fibrosis. Trinucleotide repeats found in DNA are three-nucleotide sequences that repeat multiple times and can be amplified to cause some disease conditions, such as fragile X syndrome.

PBD9 174–178  BP9 264–265  PBD8 177, 452  BP8 276–277

40 **C** When a genetic disease (e.g., cystic fibrosis) is caused by many different mutations, with different frequencies among populations, there is no simple screening test that can detect all the mutations. Although 70% of patients with cystic fibrosis have a 3–base pair deletion that can be readily detected by PCR (the ΔF508 mutation), the remaining 30% have disease caused by several hundred allelic forms of CFTR. To detect all would require sequencing of the CFTR genes. This prohibits mass screening. The other listed options do not apply.

PBD9 174–178  BP9 264–267  PBD8 466  BP8 264

41 **C** Recurrent fetal loss suggests a parental cause. A chromosomal abnormality such as a Robertsonian translocation may account for stillbirths, particularly in the first trimester, when many fetal losses result from chromosomal abnormalities. A genome-wide association study applies to populations, not individuals, and establishes linkages of common diseases such as hypertension and diabetes mellitus to polygenic risks. Fluorescence in situ hybridization (FISH) analysis is useful for establishing chromosome number and morphology, such as translocations, inversions, and deletions, but is not as definitive as a karyotype. PCR analysis is useful for identifying specific gene defects, not whole chromosomal abnormalities. Tandem mass spectroscopy is used in newborn screening for biochemical inborn errors of metabolism, such as phenylketonuria.

PBD9 174–178  BP9 263–268

42 **D** The human genome can be divided into blocks of DNA with varying numbers of contiguous single nucleotide polymorphisms (SNPs) that form haplotypes and can cluster from linkage disequilibrium, so that similar haplotypes suggest shared inheritance. The use of chips with more than a million SNPs can identify small variations in DNA from person to person, and linkage of these polymorphisms to disease can help narrow the search for candidate genes whose altered function may relate to a disease. Fluorescence in situ hybridization (FISH) probes aid in identifying specific chromosome regions and can identify abnormalities such as chromosomal deletions and translocations. The standard karyotype with banding provides information about chromosome number and major structural alterations, but does not identify specific genes or their loci. Analysis of mRNA expression provides a roundabout way of determining derivative DNA alterations, but DNA is easier to work with directly. Southern blotting has been largely supplanted by other techniques, but is useful for detection of trinucleotide repeat expansions and clonal gene rearrangements.

PBD9 174–178  BP9 263–268  PBD8 176–178  BP8 277

43 **D** Whole exome sequencing limits the search for the roughly 2% of the genome that consists of protein-encoding exons responsible for as many as 80% of mendelian diseases, and the cost is significantly reduced compared to whole genome sequencing. Pyrosequencing is a more sensitive variation of Sanger sequencing that is most often used when testing for particular sequence variants in a background of normal alleles, such as tumor cells admixed with large numbers of stromal cells. Sanger sequencing is used together with PCR to allow sequencing of any known segment of DNA. Targeted sequencing is useful for identification of a single gene or panel of genes, either by subselecting relevant clones from a whole genome library via custom complementary probes, or by alternate preparations from genomic DNA such as multiplex PCR.

PBD9 180–182  BP9 266–267
1. A 33-year-old man has experienced nausea and vomiting and has become mildly icteric over the past week. On physical examination, his temperature is 37.4°C. Laboratory studies show serum AST of 208 U/L and ALT of 274 U/L. Serologic findings for HBsAg and HBeAb are positive. A liver biopsy specimen examined microscopically shows focal death of hepatocytes with a portal inflammatory cell infiltrate. Which of the following is the most likely mechanism by which his liver cell injury occurs under these conditions?

A. Activated macrophage cytokine release
B. Antibody-mediated destruction of HBsAg-expressing liver cells
C. CD4+ lymphocyte recognition of circulating HBsAg
D. CD8+ lymphocyte recognition of viral peptide presented by MHC class I molecules
E. NK cell recognition of viral peptide presented by MHC class II molecules

2. An inflammatory reaction occurs when uric acid crystals are deposited in soft tissues, leading to cellular necrosis. Inflammatory cells have an inflammasome complex of proteins that include a cytosolic receptor recognizing the crystals, thereby releasing caspase-1 and cleaving interleukin-1 to an active form. Which of the following is the receptor in these inflammasomes?

A. C-type lectin receptors
B. Mannose receptors
C. NOD-like receptors
D. RIG-like receptors
E. Toll-like receptors

3. A 13-year-old girl has a gastrointestinal viral infection. Within the cytoplasm, viral proteins are processed to peptides. These peptides are displayed with MHC I molecules on the surface of macrophages. A receptor heterodimer made up of an α and a β polypeptide chain recognizes the peptide. Which of the following cells has this receptor?

A. B cell in bone marrow
B. CD8 cell in draining lymph nodes
C. Dendritic cell in skin
D. Natural killer cell in spleen
E. CD4 cells in gastrointestinal submucosa

4. A 20-year-old man steps into an elevator full of people with influenza who are coughing and sneezing. The influenza viral particles that he inhales attach to respiratory epithelium, and viral infection reduces MHC class I molecules displayed on these epithelial cells. Which of the following immune cells is most likely to rapidly destroy the virally infected cells?

A. CD4+ cell
B. Dendritic cell
C. Macrophage
D. Natural killer cell
E. Neutrophil

5. In an experiment, a cell line derived from a human malignant neoplasm is grown in culture. A human IgG antibody is added to the culture, and the tumor cells become coated by the antibody, but they do not undergo lysis. Next, human cells are added that are negative for CD3, CD19, and surface immunoglobulin, but are positive for CD16 and CD56. The tumor cells are observed to undergo lysis. Which of the following cell types is most likely to have killed the tumor cells?

A. B cell
B. CD4+ cell
C. CD8+ cell
D. Dendritic cell
E. Macrophage
F. Natural killer cell
6. During heterosexual intercourse, seminal fluid containing HIV contacts vaginal squamous mucosa. Cells capture virions and transport the virus via lymphatics to regional lymph nodes. Within the germinal centers of these lymph nodes, the virions infect CD4+ lymphocytes and proliferate, causing CD4+ cell lysis with release of more virions, which are taken up on the surface of cells having Fc receptors, allowing continued infection by HIV of more CD4+ cells passing through the nodes. Which of the following types of cells is most likely to capture HIV on its surface via Fc receptors?

A. B lymphocyte  
B. CD8+ cytotoxic lymphocyte  
C. Follicular dendritic cell  
D. Natural Killer cell  
E. Langhans giant cell  
F. Macrophage  
G. Mast cell

7. In a study that examines granuloma formation in the lung in response to infection with *Mycobacterium tuberculosis*, it is observed that some cells within the granuloma express MHC class II antigens. These class II antigen–bearing cells are most likely derived from which of the following peripheral blood leukocytes?

A. Basophil  
B. CD4+ T lymphocyte  
C. Monocyte  
D. Natural Killer cell  
E. Neutrophil

8. Within 5 minutes after a bee sting, a 15-year-old girl suddenly has difficulty breathing, with marked inspiratory stridor from laryngeal edema. She experiences marked urticaria and notes swelling of the hand that was stung. Which of the following is the best pharmacologic agent to treat her signs and symptoms?

A. Cyclosporine  
B. Epinephrine  
C. Glucocorticoids  
D. Methotrexate  
E. Penicillin

9. A laboratory worker who is “allergic” to fungal spores is accidentally exposed to a culture of the incriminating fungus on a Friday afternoon. Within 1 hour, he develops bouts of sneezing, watery eyes, and nasal discharge. The symptoms seem to subside within a few hours of returning home, but reappear the next morning, although the laboratory fungus is not present in his home environment. The symptoms persist through the weekend. Which of the following cells is most likely to predominate on microscopic examination of the patient’s nasal discharge?

A. Dendritic cells  
B. Eosinophils  
C. Macrophages  
D. Mast cells  
E. NK cells

10. A 28-year-old man has had hemoptysis and hematuria for the past 2 days. On physical examination, his temperature is 36.8°C, pulse is 87/min, respirations are 19/min, and blood pressure is 150/90 mm Hg. Laboratory studies show creatinine of 3.8 mg/dL and urea nitrogen of 35 mg/dL. Urinalysis shows 4+ hematuria, 2+ proteinuria, and no glucose. A renal biopsy specimen examined microscopically shows glomerular damage and linear immunofluorescence with labeled complement C3 and anti-IgG antibody. Which of the following autoantibodies has the greatest specificity for this patient’s condition?

A. Anti–basement membrane  
B. Anticardiolipin  
C. Anti–double-stranded DNA  
D. Antihistone  
E. Anti–SS-A  
F. Anti–U1-ribonucleoprotein

11. Laboratory tests are ordered for two hospitalized patients. During the phlebotomy procedure, the samples drawn from these patients are mislabeled. One of the patients receives a blood transfusion later that day. Within 1 hour after the transfusion of packed RBCs begins, the patient becomes tachycardic and hypotensive and passes pink-colored urine. Which of the following mechanisms is most likely to be responsible for the clinical picture described?

A. Antibody-dependent cellular cytotoxicity by natural killer cells  
B. Antigen-antibody complex deposition in glomeruli  
C. Complement-mediated lysis of red cells  
D. Mast cell degranulation with release of biogenic amines  
E. Release of tumor necrosis factor alpha into the circulation

12. A 35-year-old man has experienced increasing muscular weakness over the past 5 months. This weakness is most pronounced in muscles that are used extensively, such as the levator palpebrae of the eyelids, causing him to have difficulty vision with vision by the end of the day. After a night’s sleep, his symptoms have lessened. On physical examination, he is afebrile. No skin rashes are noted. Muscle strength is 5/5 initially, but diminishes with repetitive movement. A CT scan of his chest shows thymic enlargement. Which of the following is the most likely mechanism for muscle weakness in this patient?

A. Antibody-mediated dysfunction of neuromuscular junction  
B. Delayed hypersensitivity reaction against muscle antigens  
C. Immune complex deposition in muscle capillaries  
D. Lysis of muscle cells by CD8+ lymphocytes  
E. Secretion of cytokines by activated macrophages

13. A 26-year-old man has had myalgias and a fever for the past week. On physical examination, his temperature is 38.6°C. He has diffuse muscle tenderness, but no rashes or joint pain on movement. Laboratory studies show elevated serum creatine kinase and peripheral blood eosinophilia. Larvae of *Trichinella spiralis* are present within the skeletal muscle fibers of a gastrocnemius biopsy specimen. Two years later, a chest radiograph shows only a few small calcifications in the diaphragm. Which of the following immunologic mechanisms most likely contributed to the destruction of the larvae?

A. Abscess formation with neutrophils  
B. Antibody-mediated cellular cytotoxicity (ADCC)  
C. Complement-mediated cellular lysis  
D. Formation of Langhans giant cells  
E. Synthesis of leukotriene C4 in mast cells
14. A 29-year-old man has developed marked joint pain beginning 12 days after receiving snake antivenom injection. On physical examination, there is diffuse joint pain with movement. The stool is negative for occult blood. Laboratory studies show a serum creatinine level of 4.4 mg/dL and urea nitrogen level of 42 mg/dL. Microscopic examination of a renal biopsy specimen shows focal fibrinoid necrosis of the small arterial and arteriolar vascular media and intravascular microthrombi. Scattered neutrophils are seen in these areas of necrosis. Which of the following laboratory findings in the blood is most likely present in this patient?

A. CD4+ lymphocytosis  
B. Hypocomplementemia  
C. Increased IgE  
D. Neutropenia  
E. Thrombocytopenia

15. In an experiment, antigen is used to induce an immediate (type I) hypersensitivity response. Cytokines are secreted that are observed to stimulate IgE production by B cells, promote mast cell growth, and recruit and activate eosinophils in this response. Which of the following cells is most likely to be the source of these cytokines?

A. CD4+ lymphocytes  
B. Dendritic cells  
C. Macrophages  
D. Natural killer cells  
E. Neutrophils

16. Persons with sensitivity to body jewelry undergo skin patch testing with nickel compounds. Within 24 to 48 hours there is focal erythema and induration of their skin, accompanied by itching and pain. Histologically, there is an accumulation of neutrophils and monocytes with some CD4 T cells. Which of the following cytokines produced by T cells is most likely to be involved in mediating this reaction?

A. IL-2  
B. IL-5  
C. IL-10  
D. IL-12  
E. IL-17

17. A 40-year-old laboratory technician accidentally injects a chemical into his skin. The next day, he notes that an area of erythematous, indurated skin is forming around the site of injection. Two days later, the induration measures 10 mm in diameter. A microscopic section from this area, with immunostaining using antibody to CD4, shows many positive lymphocytes. Which of the following immunologic reactions is most consistent with this appearance?

A. Arthus reaction  
B. Graft-versus-host disease  
C. Delayed-type hypersensitivity  
D. Localized anaphylaxis  
E. Serum sickness

18. A 12-year-old girl has complained of a sore throat for the past 3 days. On physical examination, she has a temperature of 38.4°C and pharyngeal erythema with minimal exudate. A throat culture grows group A β-hemolytic Streptococcus. The pharyngitis resolves, but 3 weeks later, the girl develops fever and chest pain. Her anti-streptolysin O titer is 1:512. Which of the following immunologic mechanisms has most likely led to the chest pain?

A. Breakdown of T cell anergy  
B. Failure of T cell-mediated suppression  
C. Molecular mimicry  
D. Polyclonal lymphocyte activation  
E. Release of sequestered antigens

19. A 24-year-old woman has had increasing malaise; facial skin lesions shown in the figure are exacerbated by sunlight exposure; and arthralgias and myalgias for the past month. On physical examination she has mild pedal edema. On auscultation, a friction rub is audible over the chest. Laboratory findings include pancytopenia and serum creatinine 3 mg/dL. Urinalysis shows hematuria and proteinuria. A serologic test for syphilis yields a false-positive result. A renal biopsy shows granular deposits of IgG and complement in the mesangium and along the basement membrane. Which of the following mechanisms is most likely involved in the pathogenesis of her disease?

A. Antiphospholipid antibodies  
B. Defective clearance of apoptotic nuclei  
C. Increased production of IFN-γ  
D. Molecular mimicry  
E. Superantigen activation of T cells
A 29-year-old woman has had fever and arthralgias for the past 2 weeks. On physical examination, she has a temperature of 37.6°C and an erythematous malar skin rash. Initial laboratory studies are positive for ANA at 1:1600 and anti-double-stranded DNA antibodies at 1:3200, along with false positive serologic test for syphilis. Serum creatinine is markedly elevated, and serum complement levels are decreased. In vitro tests of coagulation (prothrombin time and partial thromboplastin time) are prolonged. For which of the following conditions is she at greatest risk?

A  Acute pyelonephritis  
B  Cerebral hemorrhage  
C  Renal cell carcinoma  
D  Malignant hypertension  
E  Recurrent thrombosis

A 33-year-old woman develops a skin rash on her face when she is outside in the sun for more than 1 hour. Laboratory studies show hemoglobin, 10.2 g/dL; hematocrit, 31.3%; platelet count, 126,800/mm³; and WBC count, 3211/mm³. Urinalysis shows no glucose, but there is 3+ proteinuria and 2+ blood. Her ANA test result is positive with a titer of 1:2048 and a diffuse homogeneous immunofluorescent staining pattern. Which of the following complications is most likely to occur in this patient?

A  Bronchoconstriction  
B  Cerebral lymphoma  
C  Hemolytic anemia  
D  Keratoconjunctivitis  
E  Sacroiliitis  
F  Sclerodactyly

A 31-year-old woman has had increasing peripheral edema, pleuritic chest pain, and an erythematous rash for the past 6 months. Laboratory studies show increasing serum creatinine, and urinalysis shows proteinuria with RBC casts. A renal biopsy is performed, and the light microscopic appearance of the PAS-stained specimen is shown in the figure, compared with normal. Which of the following autoantibodies when present is specific for this patient’s condition?

A  Centromere  
B  Cyclic citrullinated polypeptide  
C  DNA topoisomerase I  
D  Smith  
E  U1-RNP

A 26-year-old woman has had bouts of joint pain for the past 2 years. She also has a rash on the cheeks and bridge of the nose. On physical examination, there is no joint swelling or deformity, although generalized lymphadenopathy is present. Laboratory studies indicate anemia, leukopenia, a polyclonal gammopathy, proteinuria, and hematuria. The serum ANA test result is positive at a titer of 1:1024 with a rim pattern identified by immunofluorescence. The light microscopic and immunofluorescent (with antibody to IgG) appearances of a skin biopsy specimen are shown in the figure. Which of the following is the best information to give this patient about her disease?

A  Blindness is likely to occur within 5 years  
B  Avoid exposure to cold environments  
C  Joint deformities will eventually occur  
D  Chronic renal failure is likely to develop  
E  Cardiac valve replacement will eventually be required

A 33-year-old woman develops a skin rash on her face when she is outside in the sun for more than 1 hour. Laboratory studies show hemoglobin, 10.2 g/dL; hematocrit, 31.3%; platelet count, 126,800/mm³; and WBC count, 3211/mm³. Urinalysis shows no glucose, but there is 3+ proteinuria and 2+ blood. Her ANA test result is positive with a titer of 1:2048 and a diffuse homogeneous immunofluorescent staining pattern. Which of the following complications is most likely to occur in this patient?
24. A 60-year-old woman exposed to tuberculosis is found to have a positive tuberculin skin test. She receives prophylactic therapy that includes isoniazid. She develops arthralgias, myalgias, and a malar erythematous rash 9 months later. Laboratory findings include an ANA titer of 1:2560 in a diffuse pattern. Anti–double-stranded DNA antibodies are not present. Which of the following autoantibodies has the greatest specificity for her condition?

A. Anti-Sm
B. Anticentromere
C. Antihistone
D. Anti–Jo-1
E. Anti–U1-ribonucleoprotein
F. Anti–SS-A

25. A 22-year-old woman has had increasing malaise and swelling of her feet for the past week. On physical examination, she has 2+ pitting edema to the knees and puffiness around the eyes. Laboratory studies show serum creatinine of 4.6 mg/dL and urea nitrogen of 42 mg/dL. A renal biopsy specimen shows positive immunofluorescent staining for immunoglobulin and complement C3 within the glomeruli. The electron microscopic appearance of a glomerulus is shown in the figure. Which of the following immunologic mechanisms has most likely produced the renal damage seen in this patient?

A. Antibody-dependent cell-mediated cytotoxicity
B. Granulomatous inflammation
C. Immune complex-mediated hypersensitivity
D. Localized anaphylaxis
E. T cell-mediated cytotoxicity

26. A 19-year-old woman has a skin rash involving her face and scalp for the past 9 months. On examination there are 0.5- to 1.5-cm plaques with erythema and edema. A punch biopsy is taken and on microscopic examination shows follicular plugging. There is positive immunofluorescence at the dermal-epidermal junction with staining for IgG and complement C3. Which of the following pathogenic mechanisms result from TLR engagement by self nucleic acids?

A. Mast cell degranulation
B. Molecular mimicry
C. Release of sequestered antigens
D. TH17 elaboration
E. Type I interferon elaboration

27. A 54-year-old woman has been bothered by a chronic, dry cough for the past 5 years. She has had increasing difficulty with blurred vision for the past year. On physical examination, she has a perforated nasal septum, bilateral mild corneal scarring, and fissuring of the tongue and corners of her mouth. Laboratory studies show antibodies to SS-A and SS-B. Her serum creatinine is 2.5 mg/dL and urea nitrogen 25 mg/dL. A renal biopsy specimen examined microscopically shows tubulointerstitial nephritis but no glomerular involvement. Which of the following is the most serious condition likely to occur in this patient?

A. Endocarditis
B. Non-Hodgkin lymphoma
C. Renal failure
D. Salivary gland cancer
E. Esophageal dysmotility
F. Urethritis

28. A 47-year-old woman has had an ocular burning sensation with increasing blurring of vision for the past 5 years. On physical examination, she has keratoconjunctivitis. Atrophy of the oral mucosa, with buccal mucosal ulcerations, also is present. A biopsy specimen of the lip shows marked lymphocytic and plasma cell infiltrates in minor salivary glands. Antibodies to which of the following are most likely to be identified on laboratory testing?

A. Centromere
B. Double-stranded DNA
C. DNA topoisomerase (Scl-70)
D. SS-B
E. U1-RNP

29. A 43-year-old woman has experienced increasing difficulty in swallowing over the past year. She also has experienced diarrhea with a 5-kg weight loss in the past 6 months. She reports increasing dyspnea during this time. On physical examination, crackles are auscultated in all lung fields, but heart sounds are faint. Echocardiography shows a large pericardial effusion. The ANA test result is positive at 1:512 with a nucleolar pattern. The anti–DNA topoisomerase antibody titer is 1:1024. Which of the following serious complications is most likely to occur in patients with this disease?

A. Hepatic failure
B. Malignant hypertension
C. Meningitis
D. Perforated duodenal ulcer
E. Squamous cell carcinoma
30. A 44-year-old woman has had tightening of the skin of her fingers for the past 2 years, making them difficult to bend. She has had increasing difficulty swallowing for the past 8 months. During the past winter, her fingers became cyanotic and painful on exposure to cold. On physical examination, the skin on her face, neck, hands, and forearms appears firm and shiny. Her blood pressure is 200/130 mm Hg. A chest radiograph shows prominent interstitial markings, and lung function tests indicate moderately severe restrictive pulmonary disease. The result of her DNA topoisomerase I antibody test is positive. Which of the following conditions is she most likely to have?

A. Ankylosing spondylitis  
B. Diffuse scleroderma  
C. Discoid lupus erythematosus  
D. Limited scleroderma  
E. Rheumatoid arthritis  
F. Systemic lupus erythematosus

31. A 48-year-old woman has fingers that are tapered and claw-like, with decreased motion at the small joints. There are no wrinkle lines on her facial skin. The microscopic appearance of the skin is shown in the figure. The patient also has diffuse interstitial fibrosis of the lungs, with pulmonary hypertension and cor pulmonale. Which of the following dermal inflammatory cells is the most likely initiator of the process that is the cause of her skin disease?

A. CD4+ lymphocyte  
B. Macrophage  
C. Mast cell  
D. Neutrophil  
E. Natural killer cell

32. A 29-year-old woman has had increasing weakness over the past year, and now has difficulty climbing a single flight of stairs. Her muscles are sore most of the time. She has little difficulty writing or typing, however. During the past 3 months, she has had increasing difficulty swallowing. She has experienced chest pain for the past week. On physical examination, she is afebrile. Her blood pressure is 115/75 mm Hg. Muscle strength is 4/5 in all extremities. No rashes are present. She has 2+ pitting edema to the knees. Rales are auscultated over lower lung fields. Laboratory studies show serum creatine kinase level of 458 U/L and Jo-1 antibodies. Which of the following additional complications of her disease is she most likely to have?

A. Bony ankylosis  
B. Myocarditis  
C. Pericarditis  
D. Sclerodactyly  
E. Urethritis  
F. Xerophthalmia

33. A 51-year-old woman has had bilateral diffuse pain in her thighs and shoulders for the past 6 weeks. She has difficulty rising from a chair and climbing steps. She has a faint violaceous rash around the orbits and on the skin of her knuckles. On physical examination, she is afebrile. Muscle strength is 4/5 in all extremities. Laboratory studies show serum creatine kinase of 753 U/L, and the ANA test result is positive with a titer of 1:160. Which of the following serologic tests is most specific for the diagnosis of her underlying condition?

A. Anti–cyclic citrullinated peptide  
B. Anti–double-stranded DNA antibodies  
C. Anti–histone antibodies  
D. Anti–Jo-1 antibodies  
E. Anti–U1-ribonucleoprotein antibodies

34. A 22-year-old woman has been bothered by dryness in her mouth for the past 2 years. During this time, erythematous rashes have appeared on the skin of her face and upper neck. In the past 6 months, she has developed arthralgias. The fingers of her hands become pale and painful upon exposure to cold. On physical examination, she is afebrile. There are no joint deformities. Laboratory findings include a positive ANA test result, with a speckled pattern, and high titers of antibodies to U1-ribonucleoprotein (RNP). The serum creatinine is 1.1 mg/dL, and the urea nitrogen is 17 mg/dL. Which of the following diseases is most likely to produce these findings?

A. Dermatomyositis  
B. Discoid lupus erythematosus  
C. Limited scleroderma  
D. Mixed connective tissue disease  
E. Reactive arthritis  
F. Sjögren syndrome  
G. Systemic lupus erythematosus

35. A 45-year-old man with chronic renal failure received a kidney transplant from his brother 36 months ago. For the next 30 months, he had only minor episodes of rejection that were controlled with immunosuppressive therapy. During the past 6 months, he has had increasing serum creatinine and urea nitrogen levels. On physical examination, he is afebrile. Microscopic examination of a urinalysis specimen shows no WBCs. A renal scan shows that the allograft is reduced in size with reduced blood flow. Which of the following immunologic processes most likely accounts for these findings?

A. Complement-mediated cell lysis  
B. Delayed hypersensitivity  
C. Macrophage-mediated cell lysis  
D. Release of leukotriene C4 from mast cells  
E. Vascular intimal immunologic injury
A 19-year-old woman with chronic renal failure received a cadaveric renal transplantation. One month later, she experienced increasing serum creatinine and urea nitrogen levels, and a renal biopsy was performed. She was treated with increased immunosuppressive therapy including corticosteroids, and her renal function improved. Which of the following changes was most likely seen in the biopsy specimen before corticosteroid therapy was initiated?

A. Fibrinoid necrosis of renal arterioles with thrombotic occlusion  
B. Glomerular deposition of serum amyloid-associated protein  
C. Interstitial infiltration by eosinophils with epithelial damage  
D. Markedly thickened blood vessels with fibrosis of interstitium and glomeruli  
E. Tubular epithelial damage by CD3+ lymphocytes

A 35-year-old woman with myeloblastic leukemia received an allogenic hematopoietic stem cell transplant. A month later she has now developed an extensive, scaling rash. She also has jaundice and watery diarrhea. A skin biopsy specimen shows keratinocyte apoptosis along the dermal-epidermal junction, with upper dermal lymphocytic infiltrates. Which of the following is the most likely immunologic mechanism for these complications of her stem cell transplant?

A. Acute graft-versus-host disease  
B. Antibody-dependent cell mediated cytotoxicity  
C. Delayed-type hypersensitivity reaction  
D. Immune complex formation  
E. Mast cell release of cytokines

A 4-year-old boy has had almost continuous infections since he was 6 months old. These infections have included otitis media, pneumonia, and impetigo. Organisms cultured include *Haemophilus influenzae*, *Streptococcus pneumoniae*, and *Staphylococcus aureus*. He also has had diarrhea, with *Giardia lamblia* cysts identified in stool specimens. The family history indicates that an older brother with a similar condition died because of overwhelming infections. Which of the following laboratory findings would most likely be seen in this boy?

A. Absence of IgA  
B. Agammaglobulinemia  
C. Decreased complement C3  
D. High titer of HIV-1 RNA  
E. Positive ANA test result

A 12-year-old boy has had multiple recurrent infections for the past 10 years, including *Pneumocystis jiroveci* pneumonia, *Streptococcus pneumoniae* otitis media, and *Pseudomonas aeruginosa* urinary tract infection. On physical examination, he has a temperature of 38.5°C and pharyngeal erythema with exudate. Laboratory studies show hemoglobin, 9.1 g/dL, hematocrit, 27.6%; platelet count, 130,900/mm³; and WBC count, 3440/mm³ with 47% segmented neutrophils, 3% bands, 40% lymphocytes, and 10% monocytes. Serum immunoglobulin levels show very low IgG, very high IgM, and undetectable IgA. A peripheral blood smear shows nucleated RBCs. Which of the following immunologic defects is most likely to produce this disease?

A. Absence of adenosine deaminase  
B. Abnormal CD40-CD40L interaction  
C. Deletion of chromosome 22q11  
D. HIV infection  
E. Lack of IgA production by B lymphocytes  
F. Mutation in the *BTK* gene

A 4-year-old previously healthy boy has had pharyngitis with fever and malaise for a week. On physical examination he has lymphadenopathy and hepatosplenomegaly. Serologic studies demonstrate the presence of Epstein-Barr virus infection. Flow cytometry shows presence of NK cells, but they are found to be nonfunctional when tested in vitro. There is hypogammaglobulinemia. He develops a B cell non-Hodgkin lymphoma. His 2-year-old brother follows a similar course starting at age 5 years. Which of the following underlying immunologic conditions best explains his findings?

A. DiGeorge syndrome  
B. HIV infection with AIDS  
C. Severe combined immunodeficiency  
D. Wiskott-Aldrich syndrome  
E. X-linked lymphoproliferative disorder

A 30-year-old woman gives birth at term to a normally appearing infant girl. One hour after birth, the neonate exhibits tetany. On physical examination, she is at the 55th percentile for height and weight. Laboratory studies show serum calcium of 6.3 mg/dL and phosphorus of 3.0 mg/dL. Over the next year, she experienced increasing serum creatinine and urea nitrogen levels show very low IgG, very high IgM, and undetectable IgA. Positive ANA test result

A. Acquisition of maternal HIV infection at delivery  
B. Failure of differentiation of pre-B cells into B cells  
C. Impaired maturation of B cells into plasma cells  
D. Lack of the gene encoding for adenosine deaminase  
E. Malformation of third and fourth pharyngeal pouches
43 A 3-month-old boy has had recurrent infections of the respiratory, gastrointestinal, and urinary tracts since birth. The infectious agents have included Candida albicans, Pneumocystis jiroveci, Pseudomonas aeruginosa, rotavirus, and cytomegalovirus. Despite intensive treatment with antibiotics and antifungal drugs, he dies at age 5 months. At autopsy, lymph nodes are small with very few lymphocytes and no germinal centers. The thymus, Peyer patches, and tonsils are hypoplastic. There is no history of other males with similar findings. Which of the following immunologic alterations best describes the abnormality that caused this patient’s illness?

A. Deficiency of CD4 cells due to congenital HIV infection
B. Deletion involving chromosome 22q11
C. Mutation in the common γ chain of cytokine receptors
D. Mutation in the Bruton tyrosine kinase (BTK) gene
E. Mutation in CD40 ligand

44 A 14-month-old child has had multiple infections since birth, including pneumonia with Pseudomonas aeruginosa, adenovirus, and Aspergillus fumigatus; diarrhea with Isospora belli; otitis media with Haemophilus influenzae; and urinary tract infection with Candida albicans. Laboratory studies show hemoglobin, 13.2 g/dL; hematocrit, 39.7%; platelet count, 239,100/mm³; and WBC count, 3450/mm³ with 85% segmented neutrophils, 6% bands, 2% lymphocytes, and 7% monocytes. Serum immunoglobulin levels are IgG, 118 mg/dL; IgM, 14 mg/dL; and IgA, 23 mg/dL. The child dies of pneumonia. At autopsy, a hypoplastic thymus, small lymph nodes that lack germinal centers, and scant gut-associated lymphoid tissue are seen. Which of the following is the most likely cause of this disease?

A. Abnormal CD40 ligand
B. Adenosine deaminase deficiency
C. BTK gene mutation
D. Complement C2 deficiency
E. Congenital HIV infection

45 A 39-year-old woman sees her physician because of acute onset of severe dyspnea. On physical examination, she is afibrile and has marked laryngeal stridor and severe airway obstruction. The medical history indicates that she has had similar episodes since childhood and episodes of colicky gastrointestinal pain. Her mother and her brother are similarly affected. There is no history of severe or recurrent infections. She does not have urticaria. Laboratory studies show normal WBC count, hematocrit, and platelet count. A deficiency in which of the following plasma components is most likely to produce these findings?

A. β₂-Microglobulin
B. C1 inhibitor
C. C3
D. 5-Hydroxytryptamine
E. IgA
F. IgE

46 A 48-year-old man has been healthy all of his life, bothered only by an occasional mild diarrheal illness. On physical examination, his temperature is 37.1°C, and blood pressure is 125/85 mm Hg. Laboratory studies show a total WBC count of 6900/mm³ with 72% segmented neutrophils, 3% bands, 18% lymphocytes, and 7% monocytes. Serum immunoglobulin levels are IgG, 1.9 g/dL; IgM, 0.3 g/dL; and IgA, 0.01 g/dL. The ANA test result is negative. The skin test result for Candida antigen is positive. This patient is at greatest risk of infection from which of the following agents?

A. Aspergillus flavus
B. Hepatitis B virus
C. Herpes simplex virus
D. Pneumocystis jiroveci
E. Streptococcus pneumoniae

47 A 4-year-old boy has had recurrent respiratory infections with multiple bacterial and viral pathogens for the past 3 years. On physical examination, he has eczema involving the trunk and extremities. Laboratory findings include a platelet count of 71,000/µL and WBC count of 3800/mm³ with 88% segmented neutrophils, 6% bands, 3% lymphocytes, and 3% monocytes. Serum immunoglobulin levels show normal IgG, low IgM, and high IgA. This patient is at an increased risk of developing which of the following conditions?

A. Dementia
B. Hypocalcemia
C. Glomerulonephritis
D. Malignant lymphoma
E. Rheumatoid arthritis

48 A 28-year-old woman with a 9-year history of injection drug use has developed a chronic watery diarrhea that has persisted for the past week. On physical examination, she is afibrile and has mild muscle wasting. Her body mass index is 18. Laboratory studies of her stool show cysts of Cryptosporidium parvum. One month later, she develops cryptococcal meningitis, which is treated successfully. Oral candidiasis is diagnosed 1 month later. This patient is at greatest risk of developing which of the following neoplasms?

A. Cerebral astrocytoma
B. Cervical clear cell carcinoma
C. Cerebral non-Hodgkin lymphoma
D. Pulmonary adenocarcinoma
E. Retroperitoneal sarcoma

49 A 41-year-old man has been infected with HIV for the past 8 years. He then began receiving antiretroviral therapy, continued for the past 18 months with a regimen that includes multiple drugs. His HIV-1 RNA level initially decreased to less than 50 copies/µL after initiation of therapy; the current level is 5120 copies/µL. A mutation in the gene for which of the following molecules is most likely to have occurred?

A. CD40 ligand
B. Chemokine receptor
C. Cytokine receptor γ chain
D. p24 antigen
E. Protein tyrosine kinase
F. Reverse transcriptase
50. At 19 years of age, a previously healthy woman had an acute illness with fever, myalgia, sore throat, and mild erythematous rash over the abdomen and thighs. These symptoms abated after 1 month. She then remained healthy for 10 years. Now she has decreased visual acuity and pain in the right eye. Funduscopic examination shows findings of cytomegalovirus retinitis. Examination of her oral cavity shows thrush (candidiasis). Which of the following laboratory findings would most likely be present after her ocular problems began to appear?
   A. ANA titer 1:1024
   B. Anticentromere antibody titer 1:512
   C. CD4+ lymphocyte count 102/μL
   D. Positive HLA-B27
   E. Total serum globulin level 650 mg/dL

51. In epidemiologic studies of HIV infection and AIDS, investigators noticed that certain individuals did not develop HIV infection despite known exposure to the virus under conditions that caused HIV disease in all other individuals similarly exposed. When CD4+ lymphocytes from resistant individuals are incubated with HIV-1, they fail to become infected. Such resistance to infection by HIV is most likely caused by a mutation affecting genes for which of the following cellular components?
   A. CD28 receptor
   B. Chemokine receptor
   C. Fc receptor
   D. Interleukin-2 receptor
   E. T cell receptor

52. An epidemiologic study is conducted to determine risk factors for HIV infection. The study documents that individuals with coexisting sexually transmitted diseases such as chancroid are more likely to become HIV-positive. It is postulated that an inflamed mucosal surface is an ideal location for the transmission of HIV during sexual intercourse. Which of the following cells in these mucosal surfaces is most instrumental in transmitting HIV to CD4+ T lymphocytes?
   A. CD8+ cells
   B. Langerhans cells
   C. Natural killer cells
   D. Neutrophils
   E. Plasma cells

53. A 34-year-old woman infected with HIV begins to have difficulty with activities of daily living. She has memory problems and decreased ability to perform functions that require fine motor control, such as writing and painting. Her CD4+ lymphocyte count currently is 150/μL. Which of the following cell types is most important for the dissemination of the infection into the central nervous system?
   A. CD8+ lymphocyte
   B. Langerhans cell
   C. Macrophage
   D. Natural killer cell
   E. Neutrophil

54. A 45-year-old man has had a fever, cough, and worsening dyspnea for the past few days. On physical examination, his temperature is 39.2°F. Auscultation of the chest shows decreased breath sounds over all lung fields. A bronchoalveolar lavage is performed, and the fluid obtained yields cysts of Pneumocystis jiroveci. Laboratory studies show a CD4+ lymphocyte count of 135/μL; total serum globulin concentration of 2.5 g/dL; and WBC count of 7800/mm³ with 75% segmented neutrophils, 8% bands, 6% lymphocytes, 10% monocytes, and 1% eosinophils. Which of the following serologic laboratory findings is most likely to be positive in this patient?
   A. Antibodies to HIV
   B. Anti–double-stranded DNA antibody
   C. Anti–neutrophil cytoplasmic autoantibody
   D. Anti–streptolysin O
   E. Antibodies to lymphocytes

55. A 17-year-old boy has been sexually active for the past 3 years. He has had fever, lymphadenopathy, and pharyngitis for the past 3 weeks. Serologic testing shows that he is HIV-positive. He is now currently healthy and is not an intravenous drug user. Which of the following is the most likely outcome of his disease within the next year?
   A. Appearance of an extranodal non-Hodgkin lymphoma
   B. Development of cryptococcal meningitis
   C. Seronegativity with repeat HIV testing
   D. Transmission of infection with unprotected sex
   E. Worsening cognitive and motor function

56. A 37-year-old man who is HIV-positive has noticed an increasing number and size of skin lesions on his face, trunk, and extremities, as shown in the figure, over the past 18 months. Some of the larger lesions appear to be nodular. Molecular analysis of the spindle cells found in these skin lesions is likely to reveal the genome of which of the following viruses?
   A. Adenovirus
   B. Cytomegalovirus
   C. Epstein-Barr virus
   D. HIV-1
   E. Human herpesvirus-8
57. A 40-year-old man has been infected with HIV for the past 10 years. During this time, he has had several bouts of oral candidiasis, but no major illnesses. He is now diagnosed with Kaposi sarcoma involving the skin. He has had a 7-kg weight loss in the past 6 months. Laboratory studies show the HIV-1 RNA viral load is currently 60,000 copies/mL. Which of the following types of cells is most depleted in his lymph nodes?

A. CD4+ lymphocyte  
B. CD8+ lymphocyte  
C. CD19+ lymphocyte  
D. Macrophage  
E. Natural killer cell  
F. Plasma cell

58. A 63-year-old man has had increasing malaise and back pain for the past 4 months. On physical examination, he is afebrile and has mild muscle wasting. Laboratory studies show serum creatinine of 4.5 mg/dL and urea nitrogen of 44 mg/dL. Urine dipstick analysis shows no blood, protein, or glucose, but a specific test for Bence Jones proteins yields a positive result. A radiograph of his spine shows rounded lucent lesions. The microscopic appearance of his liver with Congo red stain (A) and with polarized light (B) is shown in the figure. Which of the following underlying conditions is most likely to be present in this patient?

A. Ankylosing spondylitis  
B. Common variable immunodeficiency  
C. Multiple myeloma  
D. Rheumatic fever  
E. Systemic sclerosis

59. A 63-year-old man has had chronic arthritis for the past 15 years. Physical examination shows ulnar deviation with bony ankylosis producing swan neck deformities of the fingers. Laboratory studies show 4.2 g of protein in a 24-hour urine collection, serum creatinine of 3.1 g/dL, and urea nitrogen of 3 g/dL. Level of C-reactive protein is markedly elevated. A rectal biopsy is performed, which shows deposition of amorphous pink material with H&E staining in the mucosa. The material stains positive with Congo red. Which of the following proteins is the most likely precursor to this material in the mucosa?

A. Acute-phase reactant  
B. β2-Microglobulin  
C. λ light chains  
D. Transthyretin  
E. Rheumatoid factor  
F. C-reactive protein

60. A 79-year-old man has experienced worsening congestive heart failure and pulmonary and peripheral edema for the past 4 years. On physical examination, his temperature is 36.9°C, pulse is 70/min, respirations are 16/min, and blood pressure is 120/75 mm Hg. Echocardiography shows cardiomegaly with four-chamber dilation. All laboratory studies, including serum protein electrophoresis and examination of bone marrow smear, are normal. An endomyocardial biopsy specimen shows Congo red-positive interstitial deposits of amorphous material. Which of the following proteins is most likely to be a precursor for this material?

A. α-Fetoprotein  
B. β2-Microglobulin  
C. Calcitonin  
D. IgE  
E. Light chains  
F. Transthyretin  
G. Troponin T
CHAPTER 6  Immune System Diseases

1 D  Virus-infected cells are recognized and killed by cytotoxic CD8+ T cells. The T cell receptor on the CD8+ T cells binds to the complex of viral peptide and MHC class I molecules displayed on the surface of the infected cell. Natural killer (NK) cells also recognize MHC class I molecules with self-peptides, but this self-recognition inhibits NK cell killing. Viruses that inhibit MHC I expression of peptides may hide from cytotoxic cells, but not from NK cells. The other listed options are not the major immune response to hepatitis viral infection.

PBD9 186, 192  BP9 104  PBD8 188  BP8 108, 113

2 C  The NLR-inflammasome pathway plays a role in the innate immune system recognition of urate crystals and promoting the inflammation associated with gout. These receptors may also contribute to inflammation of atherosclerosis. C-type lectin receptors (CLRs) expressed on the plasma membrane of macrophages and dendritic cells detect fungal glycans and elicit inflammatory reactions to fungi. Mannose receptors on phagocytes recognize microbial sugars with terminal mannose residues and induce microbial phagocytosis. RIG-like receptors (RLRs) are located in the cytosol of most cell types and detect nucleic acids of viruses that replicate in the cytoplasm of infected cells to stimulate production of antiviral cytokines. Toll-like receptors (TLRs) in the plasma membrane and endosomal vesicles activate transcription factors that stimulate synthesis and secretion of cytokines and expression of adhesion molecules to recruit and activate leukocytes.

PBD9 188  BP9 105

3 B  The adaptive immune system requires presentation of antigen to effector cells. Intracellular pathogens such as viruses are displayed by MHC I molecules to both CD4 and CD8 T cells with $\alpha\beta$ T cell receptors. Some CD8 cells secrete cytokines, but most are cytotoxic. B lymphocytes express immunoglobulin receptors that primarily react to extracellular antigens, such as those derived from bacteria, as an adaptive humoral immune response. Dendritic cells are a type of antigen-presenting cell. Natural killer cells are part of innate immunity and react against cells when they do not display MHC molecules. A subset of T cells beneath epithelia display yô receptors that are part of an innate immune response.


4 D  Natural killer (NK) cells have the ability to respond without prior sensitization. They carry receptors for MHC class I molecules, which inhibit their lytic function. When expression of class I MHC molecules is reduced on a cell surface by viral interference, the inhibitory receptors on NK cells do not receive a negative signal, and the targeted cell is killed. NK cells are often the first line of defense against viral infection. CD4+ cells are helper T cells that assist other cells, such as NK cells, macrophages, and B cells, in the immune response. Dendritic cells aid in antigen presentation. Macrophages can phagocytize necrotic cells, then process and display any foreign antigens within those cells. Neutrophils provide a nonspecific immune response, primarily to bacterial infections and not to intracellular viral infections.

PBD8 187, 243  BP8 113

5 F  Natural killer (NK) cells have CD16, an Fc receptor that allows them to bind to opsonized cells and lyse them. This is a form of type II hypersensitivity with antibody-mediated disease. NK cells comprise 10% to 15% of circulating lymphocytes. NK cells also may lyse human cells that have lost MHC class I expression as a result of viral infection or neoplastic transformation. B cells have surface immunoglobulin, are CD19 positive, and participate in humoral immunity. CD4+ cells are T lymphocytes that are “helper” cells; they have T cell receptors and are CD3 positive. Likewise, CD8+ cells have T cell receptors and mark with CD3, but they act as cytotoxic T lymphocytes. Dendritic cells are a form of antigen-presenting cell that expresses large amounts of MHC class II molecules. Macrophages express MHC II and act as antigen-presenting cells to CD4+ cells; they can phagocytize opsonized cells.

PBD9 192  BP9 104, 115  PBD8 188  BP8 108, 113, 125

6 C  Dendritic cells in epithelia are known as Langhans cells, and those within germinal centers are called follicular dendritic cells (FDCs). The FDCs may become infected but not killed by HIV. They have cell surface Fc receptors that capture antibody-coated HIV virions through the Fc portion of the antibody. These virions attached to the FDCs can infect passing CD4+ lymphocytes. Dendritic cells elaborate type I interferons that up-regulate antiviral proteins in neighboring cells. B cells are a component of humoral immunity, and antibody to HIV does not serve a protective function, but allows serologic detection of infection. CD8+ cells are cytotoxic lymphocytes that lack the receptor necessary for infection by HIV. Because they survive selectively, the CD4+:CD8+ ratio is reversed so that it is typically less than 1 with advanced HIV infection. Innate lymphoid cells resemble NK cells, but shape further lymphoid reactions. Langhans giant cells are “committees” of activated macrophages that are part of a granulomatous response. Macrophages are a type of antigen-presenting cell that can become infected by HIV without destruction. Mast cells have surface-bound IgE, which can be cross-linked by antigens (allergens) to cause degranulation and release of vasoactive amines, such as histamine, as part of anaphylaxis with type I hypersensitivity.

PBD9 186, 191–192  BP9 104, 145–147  PBD8 187  BP8 113

7 C  Blood monocytes expressing MHC class II antigens can migrate into tissues and become longer-lived macrophages. In tuberculosis, these macrophages further transform into epithelioid cells that can contribute to granulomatous inflammation. Macrophages play an important role in delayed hypersensitivity reactions associated with cell-mediated immunity. Basophils are circulating counterparts of mast cells
and may play a role in IgE-mediated immune responses. B cells form plasma cells that secrete immunoglobulin on stimulation and are essential to humoral immunity. Natural killer cells can function without prior sensitization. Neutrophils are important mainly in acute inflammatory responses, although some neutrophils may be present within a granulomatous reaction.

8 B Her systemic anaphylactic reaction results from an immediate type I hypersensitivity reaction with antigen triggering mast cell–bound IgE with release of preformed mediators such as histamine. Epinephrine is the fastest acting agent to treat this life-threatening condition. Cyclosporine is used to minimize lymphocyte-mediated transplant rejection. Glucocorticoids can reduce immune inflammatory reactions, although this occurs over days to weeks, not minutes. Methotrexate is useful in the treatment of graft-versus-host disease and for some malignancies. Penicillin is an antibiotic that can induce a type I hypersensitivity reaction in sensitized persons.

9 B This history is typical of the late-phase reaction in type I hypersensitivity. The initial rapid response is largely caused by degranulation of mast cells. The late-phase reaction follows without additional exposure to antigen and is characterized by more intense infiltration by inflammatory cells, such as neutrophils, eosinophils, basophils, monocytes, and T<sub>H</sub>2 CD4<sup>+</sup> lymphocytes. There is more tissue destruction in this late phase. The most characteristic cell in secretion from an allergic response is the eosinophil. Dendritic cells, lymphocytes, and mast cells remain in the epithelium. NK cells are not part of the typical allergic response.

10 A He has Goodpasture syndrome, in which an autoantibody is directed against type IV collagen in basement membranes of the glomeruli and in the lung. This is a form of type II hypersensitivity reaction. The antibodies attach to the basement membrane and fix complement, damaging the glomeruli. Anticardiolipin, along with anti–β<sub>2</sub>-glycoprotein “lupus anticoagulant,” are found with antiphospholipid syndrome, which may appear in systemic lupus erythematosus (SLE). These patients have coagulopathies with thrombosis or bleeding, or both. Anti–double-stranded DNA antibodies have specificity for SLE, whereas antihistone antibodies are characteristic of drug-induced SLE. Anti–SS-A antibody is seen in Sjögren syndrome. The anti–U1-ribonucleoprotein antibody is seen in mixed connective tissue disease (MCTD).

11 C A major transfusion reaction results from a type II hypersensitivity reaction. The patient’s serum contains naturally occurring antibodies to the incompatible donor RBCs. They attach to the donor RBCs and induce complement activation that results in generation of the C5-9 membrane attack complex. Major transfusion reactions are rare, and most result from clerical errors. Natural killer cell lysis is seen with antibody-mediated diseases. Antigen-antibody complex formation is typical of a type III hypersensitivity reaction. Mast cells degranulate with antigen attachment to IgE in type I hypersensitivity reactions. Tumor necrosis factor α is not part of hypersensitivity reactions.

12 A Myasthenia gravis is a form of type II hypersensitivity reaction in which antibody is directed against cell surface receptors. Thymic hyperplasia or thymoma is likely to be present. Antibodies to acetylcholine receptors impair the function of skeletal muscle motor end plates, leading to muscular weakness. B cells produce these antibodies; macrophages are not a significant part of this hypersensitivity reaction; and there is little or no inflammation of the muscle in myasthenia gravis. Delayed-type hypersensitivity reactions are more likely in parasitic infestations of muscles. Immune complex–mediated injury is a feature of dermatomyositis. Muscle lysis by CD8<sup>+</sup> T cells occurs in polymyositis.

13 B When antibody is directed at a parasitic infection, there is Fc receptor–mediated inflammation and phagocytosis, characteristic for ADCC. IgG and IgE antibodies bearing Fc receptors coat the parasite. Macrophages, natural killer cells, and neutrophils can then recognize the Fc receptor and destroy the antibody-coated target cells. Acute inflammatory reactions with abscess formation have little effect against tissue parasites. Complement-mediated lysis is most typical of immune destruction of RBCs with hemolysis. Langhans giant cells are seen in granulomatous inflammation, a form of type IV hypersensitivity. Leukotriene C<sub>4</sub> is a potent agent that promotes vascular permeability and bronchial smooth muscle contraction in type I hypersensitivity reactions.

14 B In the localized immune complex reaction (Arthus reaction) at the site of injection, there can be activation and depletion of complement C3. The reaction described here is serum sickness in response to the injected foreign protein, and produced more widespread antigen-antibody complex deposition, particularly in the kidneys. CD4<sup>+</sup> lymphocytes assist in various antibody-mediated and cell-mediated immune reactions, but their numbers in peripheral blood do not change appreciably. IgE concentration is increased in individuals with atopy and the potential for type I hypersensitivity. Although neutrophils are being recruited locally to the inflammatory reaction in this case, they are not depleted systemically, and they may be increased in the circulation. Thrombocytopenia could be seen with a thrombotic microangiopathy such as thrombotic thrombocytopenic purpura, but not typically in an Arthus reaction.

15 A CD4<sup>+</sup> cells of the T<sub>H</sub>2 type are essential to the induction of type I hypersensitivity because they can secrete cytokines, such as interleukin (IL)-3, IL-4, IL-5, and
granulocyte-macrophage colony-stimulating factor, which are required for the growth, recruitment, and activation of mast cells and eosinophils. Dendritic cells trap antigen and aid in antigen presentation. Macrophages are also antigen-presenting cells, and they can secrete various cytokines, but they are not essential to type I hypersensitivity. Natural killer cells can lyse other cells, such as virus-infected cells, without prior sensitization. Neutrophils are recruited by cytokines to participate in acute inflammatory reactions.

PBD9 198, 202  BP9 118–120  PBD8 195  BP8 120–122

16 E The Th17 subset of CD4 cells plays a role in delayed-type hypersensitivity reactions. Many persons react to nickel, particularly with body piercing jewelry. IL-17 may also be useful in recruiting neutrophils to fight bacterial as well as fungal infections such as aspergillosis and candidiasis. IL-2 acts as an autocrine growth factor promoting T cell proliferation. IL-5 activates eosinophils as part of a Th2 response. IL-10 is an immunosuppressive cytokine that diminishes lymphocyte activation. NK cells may secrete interferon-γ in response to stimulation by IL-12.

PBD9 198, 208–209  BP9 119–120  PBD8 195, 206  BP8 128–130

17 C Perivascular accumulation of T cells, particularly CD4+ cells, is typical of delayed hypersensitivity skin reactions, driven by a Th1 response mediated largely by release of the cytokine interleukin-2. The tuberculin skin test also works through this pathway. Anaphylaxis (type I hypersensitivity) typically occurs within minutes to hours after an encounter with an antigen to which sensitization has occurred; the localized form may occur following ingestion of the foreign protein antigen. Systemic and localized immune complex diseases (Arthus reactions and serum sickness) are type III hypersensitivity reactions; they often exhibit vascu- litis. Graft-versus-host disease is characterized by epidermal apoptosis and rash in persons receiving an allogeneic hematopoietic stem cell transplant.

PBD9 209–211  BP9 119–120  PBD8 205–207  BP8 128–130

18 C Streptococcal M proteins cross-react with cardiac glycoproteins, resulting in rheumatic heart disease, a form of autoimmunity. The other listed options are not major immune responses to streptococcal infection. Breakdown of T cell anergy usually occurs when localized tissue damage and inflammation cause up-regulation of co-stimulatory molecules on the target tissues. This is a possible mechanism of autoimmunity in the brain and in pancreatic islet β cells. Failure of T cell–mediated suppression has not yet been shown to cause any autoimmune disease; it remains a potential mechanism. Microbial products such as endotoxin or bacterial superantigens may cause polyclonal lymphocyte activation. Release of sequestered antigens can cause autoimmunity; this mechanism is likely in autoimmune uveitis (sympathetic ophthalmitis) following eye trauma.

PBD9 207, 216  BP9 124  PBD8 212  BP8 119, 124

19 B This young woman has a classic picture of systemic lupus erythematosus (SLE)—the erythematous malar facial skin rash shown in the figure, and renal failure with proteinuria and hematuria from immune complex deposition in the glomeruli. Defective clearance and hence increased burden of nuclear apoptotic bodies in thymic lymphocyte development is considered a fundamental mechanism that underlies SLE. This along with loss of self-tolerance to nuclear antigens gives rise to the pathogenic DNA–anti DNA immune complexes, as measured by the antinuclear antibody test. Antiphospholipid antibodies may be present with SLE, but lead to coagulopathy. IFN-γ is a product of CD4+ T cells and NK cells. There is no evidence of delayed hypersensitivity or NK cell dysfunction in SLE. Molecular mimicry occurs when a microbial antigen cross-reacts with a normal tissue as in rheumatic fever. Widespread and non-specific activation of T cells by superantigens occurs in toxic shock syndrome.

PBD9 218–226  BP9 125–130  PBD8 215–218  BP8 139–144

20 E The serologic features of systemic lupus erythematosus (SLE) include the more sensitive ANA test and the more specific anti-dsDNA test. The abnormal coagulation tests suggest the presence of antiphospholipid antibodies. These antibodies against phospholipid-protein complexes (antiphospholipid antibodies) also are called lupus anticoagulants because they interfere with in vitro clotting tests. In vivo, they are thrombogenic. Hence these patients can have recurrent thrombosis. Lupus anticoagulants also can occur in the absence of SLE. The other listed options are unlikely to be common or associated complications of SLE.

PBD9 218–221  BP9 125–130  PBD8 217–221  BP8 139–144

21 D Many patients with systemic lupus erythematosus (SLE) have glomerulonephritis, as evidenced by proteinuria with hematuria, and eventually develop renal failure. Blindness is uncommon in SLE. Raynaud phenomenon is associated with many autoimmune diseases, but it is most troublesome in scleroderma. Although synovial inflammation is common in SLE, joint deformity is rare. Libman-Sacks endocarditis associated with SLE tends to be nondeforming and limited, so there is minimal valve damage. It is now uncommon because of the use of corticosteroid therapy in the treatment of SLE.

PBD9 222–224  BP9 125–130  PBD8 217–219  BP8 139–144

22 C Patients with systemic lupus erythematosus (SLE) can develop anti-RBC antibodies, which can cause hemolytic anemia. Cytopenias, including leukopenia, thrombocytopenia, and anemia, are also common. Bronchoconstriction is a feature of bronchial asthma and can occur in allergies as a predominantly type I hypersensitivity reaction. Cerebral lymphomas are rare, but may occur in immunodeficient patients, particularly patients with AIDS. Keratoconjunctivitis can be seen in Sjögren syndrome as a result of decreased tear production from lacrimal gland inflammation. Sacroiliitis is a feature of many of the spondyloarthropathies, such as ankylosing spondylitis. Sclerodactyly is seen in scleroderma.
When extensive, it is usually part of the spectrum of findings associated with diffuse scleroderma; when it involves only a few areas of the skin (e.g., just the hands), it is more likely to indicate limited scleroderma (CREST syndrome).

23 D The figure shows the so-called wire loop glomerular capillary lesions of lupus nephritis. Anti-Smith and anti-double-stranded DNA are more specific for systemic lupus erythematosus, but sensitivity is low: anti-Smith is present in only 25% of SLE cases. Anticentromere antibody is seen most often with limited scleroderma, whereas anti-DNA topoisomerase I is found with diffuse scleroderma. Cyclic citrullinated polypeptide and rheumatoid factor are found most often with rheumatoid arthritis. Anti-U1-RNP can be found with mixed connective tissue disease.

24 C A drug-induced systemic lupus erythematosus (SLE)-like condition may be caused by drugs such as isoniazid, procarbamide, and hydralazine. Test results for ANA are often positive, but test results for anti–double-stranded DNA are negative. Anthistone antibodies are present in many cases. Characteristic signs and symptoms of SLE may be lacking, and renal involvement is uncommon. Remission occurs when the patient stops taking the drug. Anti-Sm antibody shows specificity for SLE. Anti-Jo-1 antibody has specificity for polymyositis/dermatomyositis. Anti-U1-ribonucleotide protein has specificity for mixed connective tissue disease. Anticentromeric antibody is most likely to be present with limited scleroderma (CREST syndrome). Anti-SS-A antibody is most characteristic of Sjögren syndrome.

25 C The figure shows extensive electron-dense deposits within glomerular basement membrane characteristic of immune complex–mediated glomerulonephritis. The immune complexes activate complement and result in acute inflammation. Granulomatous inflammation and T cell cytotoxicity are features of type IV hypersensitivity. Antibody–dependent cell-mediated cytotoxicity is initiated when IgG or IgE coats a target to attract cells that affect lysis; immune complexes do not form. Localized anaphylaxis is a type I hypersensitivity reaction that is mediated by IgE antibody.

26 E She has discoid lupus erythematosus, with skin lesions similar to those of systemic lupus erythematosus (SLE), but with systemic involvement much less likely to occur. Self nucleic acids mimic their microbial counterparts, and in conjunction with TLRs, they incite type I interferon production to activate dendritic cells and B cells. They also promote a Th1 response, which contributes to autoimmunity. Mast cell degranulation with type I hypersensitivity may produce an urticarial rash, but without autoimmune antigen-antibody complex deposition. Molecular mimicry may follow an infection and lead to polyclonal B cell activation, but autoimmune diseases involve self antigens, not exogenous antigens. Release of sequestered antigens is the mechanism for sympathetic ophthalmia. A Th17 response may be present in chronic inflammatory conditions such as inflammatory bowel diseases with macrophage and neutrophilic infiltrates.

27 B Sjögren syndrome is characterized by immunologically mediated destruction of salivary and lacrimal glands and other exocrine glands lining the respiratory and gastrointestinal tracts. Dryness and crusting of the nose can lead to perforation of the nasal septum. In 25% of cases, extraglandular tissues, such as lung, skin, kidney, and muscles, may be involved. The immune dysregulation that accompanies autoimmune diseases increases the risk for B cell lymphoid malignancies, such as MALT lymphoma. Libman-Sacks endocarditis is most often a feature of systemic lupus erythematosus (SLE). Renal failure is more likely to occur with SLE from glomerulonephritis. Esophageal dysmotility is a feature of scleroderma. When not extensive, it typically indicates limited scleroderma (CREST syndrome); when extensive, it indicates diffuse scleroderma, which has a poorer prognosis. Salivary gland cancers are unlikely to be associated with autoimmune diseases. Nongonococcal urethritis is seen in reactive arthritis, along with conjunctivitis and arthritis.

28 D Sjögren syndrome primarily involves salivary and lacrimal glands. Antibodies to SS-B are found in the majority of these patients. Anti–double-stranded DNA is a specific autoantibody for systemic lupus erythematosus. Anticentromere antibody is seen most often in limited systemic sclerosis, whereas anti–DNA topoisomerase most often appears with diffuse systemic sclerosis. U1-RNP is a marker for mixed connective tissue disease (MCTD).

29 B Diffuse systemic sclerosis (scleroderma) initially has widespread skin involvement but rapidly progresses to visceral organ involvement, including the small arteries of the kidney. These are damaged by a hyperplastic arteriolar sclerosis that can be complicated by very high blood pressure and renal failure. Meningitis and liver failure are not typical features of autoimmune diseases. With scleroderma, the gastrointestinal tract undergoes fibrosis with obstruction and malabsorption, without any tendency to perforation or ulceration. Risk factors for esophageal squamous carcinoma include smoking and alcohol abuse.

30 B This patient has cutaneous and visceral manifestations of diffuse systemic sclerosis (diffuse scleroderma). Raynaud phenomenon, skin changes, and esophageal dysmotility also can occur in limited scleroderma (the former CREST syndrome), but lung and renal involvement typically do not. In diffuse systemic sclerosis, the anti–DNA topoisomerase I antibody is often present, and patients can develop interstitial lung disease and renal disease with hyperplastic
CHAPTER 6  Immune System Diseases

peptide is present in most patients with rheumatoid arthritis, which is accompanied by inflammatory destruction of joints, not muscle, although muscle may atrophy secondary to diminished movement. Anti–double-stranded DNA is specific for systemic lupus erythematosus (SLE), in which there can be myositis without significant inflammation or necrosis. Antihistone antibodies are associated with drug-induced SLE. The anti–U1-ribonucleoprotein antibodies suggest a diagnosis of mixed connective tissue disease, a condition that can overlap with polymyositis.

PBD9 231, 1238–1239  BP9 135, 805  PBD8 225, 1273  BP8 151

Mixed connective tissue disease (MCTD) can have some features of systemic lupus erythematosus (SLE), myositis, rheumatoid arthritis, scleroderma, and Sjögren syndrome. In contrast to SLE or diffuse scleroderma, serious renal disease is unlikely. Dermatomyositis causes muscle pain, and the rash is typically a subtle heliotrope rash with a violaceous appearance to the eyelids; Jo-1 antibody is a more typical finding. Discoid lupus erythematosus (DLE) is characterized by a rash similar to SLE, but with immune complex deposition only in sun-exposed areas of the skin, a positive ANA test result in a few cases, absence of anti-Smith or anti–double-stranded DNA antibodies, and absence of serious renal disease. Some cases of DLE can progress to SLE. In limited scleroderma (previously described as CREST syndrome), anticientromere antibody is often present. In reactive arthritis (with conjunctivitis, arthritis, and nongonococcal urethritis), there is often a positive serology for HLA-B27 and a history of either nongonococcal urethritis with Chlamydia or gastrointestinal infection with Campylobacter, Salmonella, or Shigella organisms. In Sjögren syndrome, keratoconjunctivitis predominates, and antibodies to SS-A and SS-B are often present. The anti-Smith or anti–double-stranded DNA antibodies are more specific for SLE.


Scleroderma being an autoimmune disease, inflammation is present. The antibody to cyclic citrullinated peptide (anti-CCP) is more specific for SLE. The anti-Smith or anti–double-stranded DNA antibodies are more specific for systemic lupus erythematosus, and renal disease in these patients is most likely due to glomerulonephritis.


Muscle weakness in polymyositis tends to be symmetric, and proximal muscles are involved first. This condition differs from dermatomyositis in that there is no skin involvement, and polymyositis typically affects adults. On biopsy, the skeletal muscle shows infiltration by lymphocytes along with degeneration and regeneration of muscle fibers. The lymphocytes are cytotoxic CD8+ cells. Some patients may have myocarditis, vasculitis, or pneumonia; but in contrast to dermatomyositis, the risk of cancer is equivocal. Bony ankylosis is a feature of progressive or recurrent joint inflammation with rheumatoid arthritis. Pericarditis is most likely to be a feature of systemic lupus erythematosus or diffuse systemic sclerosis. Sclerodactyly is a feature of scleroderma. When not extensive, it typically indicates limited scleroderma (CREST syndrome); when extensive, it indicates diffuse scleroderma, which has a poorer prognosis. Nongonococcal urethritis, conjunctivitis, and arthritis are seen with reactive arthritis. Xerophthalmia (usually with accompanying xerostomia) is seen in Sjögren syndrome.

PBD9 231, 1238–1239  BP9 135, 805  PBD8 225, 1273  BP8 151

Acute rejection of kidney transplants occurs weeks, months, or even years after transplantation. It is characterized by infiltration with CD3+ T cells that include the CD4+ and CD8+ subsets. These cells damage tubular epithelium by direct cytotoxicity and by release of cytokines, such as interferon-γ, which activate macrophages. The reaction is called acute cellular rejection, and it can be readily treated with corticosteroids. Fibrinoid necrosis and thrombosis are more typical of hyperacute rejection, which occurs within minutes.
of placement of the transplant into the recipient, and is rare because of ABO blood group and MHC allotypic matching. Amyloid derived from serum amyloid-associated protein can occur in chronic infections and inflammation. Eosinophils accumulate in acute interstitial nephritis owing to drug reactions. Interstitial and glomerular fibrosis and blood vessel thickening occur in chronic rejection, which is generally not reversible.

PBD9 231–234  BP9 138  PBD8 228–229  BP8 131–134

37 A Her graft-versus-host disease (GVHD) is produced when the engrafted marrow is not completely matched for major histocompatibility loci and thus contains immunocompetent donor cells that can proliferate, recognize host cells as foreign, and attack host tissues. The skin, liver, and gastrointestinal epithelium are typically affected. The localized apoptosis with GVHD does not produce significant substrate proteins for amyloid formation. Delayed-type hypersensitivity is more likely seen with contact dermatitis. Immune complex formation is more typical of type III hypersensitivity with autoimmune antibody formation. Type I hypersensitivity with systemic anaphylaxis occurs with exposure to an antigen, such as penicillin, after prior exposure.

PBD9 236  BP9 139  PBD8 230  BP8 134

38 F X-linked agammaglobulinemia of Bruton is a condition in which B cell maturation stops after the rearrangement of heavy-chain genes, and light chains are not produced. Complete immunoglobulin molecules with heavy and light chains are not assembled and transported to the cell membrane. The lack of immunoglobulins predisposes the child to recurrent bacterial infections after maternally derived antibodies diminish following infancy. Because T cell function remains intact, viral, fungal, and protozoal infections are uncommon. CD4+ and CD8+ lymphocytes differentiate from precursors in the thymus, which is not affected by the BTK gene mutation that gives rise to Bruton agammaglobulinemia. Follicular dendritic cells are a form of antigen-presenting cell that is not affected by B cell and T cell disorders. Monocytes may leave the circulation to become tissue macrophages, a process not dependent on B cell maturation. Natural killer (NK) cells are part of the innate immune system and respond to antibody-coating abnormal cells—a process diminished by reduced antibody production—but the NK cells themselves are not directly affected by lack of immunoglobulin. Lack of stem cell differentiation is incompatible with life.

PBD9 239–240  BP9 140–141  PBD8 231–232  BP8 152–153

39 B This boy most likely has Bruton agammaglobulinemia, an X-linked primary immunodeficiency marked by recurrent bacterial infections that begin after maternal antibody levels diminish. Selective IgA deficiency is marked by a more benign course, with sinopulmonary infections and diarrhea that are not severe. Deficiency of complement C3 is rare; it leads to greater numbers of infections in children and young adults, but Giardia infections are not a feature of this disease. Lack of cell-mediated immunity is more likely to be seen in HIV infection in children. Although some patients with Bruton agammaglobulinemia can develop features of systemic lupus erythematosus, they generally do not have a positive test result for ANA.

PBD9 239–240  BP9 140–141  PBD8 231–232  BP8 152–153

40 B These are features of the hyper-IgM syndrome, which results from lack of isotype switching from IgM to other immunoglobulins. Patients are particularly susceptible to Pneumocystis and to bacterial infections. The abnormal IgM antibodies in excess can attach to circulating cells and lead to cytopenias. An absence of adenosine deaminase characterizes a form of severe combined immunodeficiency. The deletion of chromosome 22q11 is a feature of the DiGeorge anomaly, which affects T cell differentiation and maturation. HIV infection can be accompanied by opportunistic infections, particularly Pneumocystis, but abnormal immunoglobulin production generally is not seen. A lack of just IgA production alone is seen with selective IgA deficiency. Mutations in the BTK gene account for Bruton agammaglobulinemia with reduction in levels of all immunoglobulins.

PBD9 241  BP9 141  PBD8 232–233  BP8 153–154

41 E These findings point to mutations in the gene encoding an adaptor molecule called SLAM-associated protein (SAP) that binds to a family of cell surface molecules involved in the activation of NK cells and T and B lymphocytes, and result in increased susceptibility to viral infections, particularly Epstein-Barr virus (EBV). DiGeorge syndrome manifests in infancy with failure of cell-mediated immunity from lack of functional T cells. HIV infection is marked by failure of cell-mediated immunity. Individuals with severe combined immunodeficiency would not live as long as this patient with such mild infections. Wiskott-Aldrich syndrome is associated with eczema and thrombocytopenia.

PBD9 242  BP9 142–143

42 E DiGeorge syndrome can involve the thymus, parathyroids, aorta, and heart. T cell function is deficient, resulting in recurrent and multiple fungal, viral, and protozoal infections. The size of the 22q11.2 deletion determines the severity of the disease. HIV infection does not explain the hypocalcemia at birth. Failure of pre-B cell maturation results in Bruton agammaglobulinemia. Impaired B cell maturation to plasma cells is a mode of development of common variable immunodeficiency. Some cases of severe combined immunodeficiency are caused by lack of adenosine deaminase.

PBD9 239, 241  BP9 141  PBD8 234  BP8 153–154

43 C Severe combined immunodeficiency (SCID) has both deficient T cell and B cell arms of the immune system, so that there are severe and recurrent infections with bacteria, viruses, and fungi. When the family history indicates males are affected, then SCID is most likely X-linked and due to mutations in the common γ chain that is a part of many cytokine receptors, such as interleukin (IL)-2, IL-4, IL-7, and IL-15. These cytokines are needed for normal B cell and T cell
development, so both humoral and cell-mediated immunity are affected. The marked lymphoid hypoplasia is not typical of HIV infection. A deletion involving chromosome 22q11 is seen in DiGeorge syndrome affecting cell-mediated immunity. BTK gene mutations give rise to Bruton agammaglobulinemia, with loss of humoral immunity. Mutation in the CD40 ligand is responsible for hyper-IgM syndrome.

44 B Severe combined immunodeficiency (SCID) can be treated with allogeneic bone marrow transplantation. The transplanted stem cells in the bone marrow give rise to normal T and B cells. Half of SCID cases are caused by an X-linked mutation in the common γ chain for cytokine receptors, and the rest are due to autosomal recessive mutations in the gene encoding for adenosine deaminase, which leads to accumulation of metabolites toxic to lymphocytes. An abnormal CD40 ligand interaction with CD40 leads to lack of isotype switching in patients with hyper-IgM syndrome. The BTK gene product is required for differentiation of pro–B cells and pre–B cells, and a mutation leads to agammaglobulinemia. The 22q11 deletion is seen in infants with DiGeorge anomaly and results in lack of T cell development. Individuals lacking complement component C2 have some increase in infections, but mainly develop a disease resembling systemic lupus erythematosus. HIV infection leads to many opportunistic infections, which sometimes occur in infancy and early childhood, but it is mainly CD4+ lymphocytes that are diminished.

45 B Hereditary angioedema is a rare autosomal-recessive disorder of the complement pathway in which there is a deficiency of antigenic or functional C1 inhibitor, resulting in recurrent episodes of edema. Of the remaining choices, only C3 and IgA have a deficiency state. C3 deficiency is accompanied by recurrent infections with pyogenic bacteria. IgA deficiency leads to mild recurrent gastrointestinal and respiratory tract infections and predisposes to anaphylactic transfusion reaction. β2-Microglobulin is a component of MHC class I; it can be increased with HIV infection and can be a substrate for amyloid fibrils in patients receiving long-term hemodialysis. 5-Hydroxytryptamine (serotonin) has an effect similar to histamine, which drives vasodilation and edema. IgE participates in localized or systemic anaphylaxis with edema.

46 E Individuals with a selective (isolated) IgA deficiency are bothered by minor recurrent sinopulmonary infections and by diarrhea. IgA is useful in innate immunity against bacterial organisms in the respiratory and gastrointestinal tracts. IgA antibodies present in their serum can lead to anaphylactic transfusion reaction with IgA in donor serum. Immune reactions against fungal and viral infection are mediated mainly by T cells. Viral infections tend to be handled by T lymphocyte responses. Hepatitis infections are not directly related to immunodeficiency states, although AIDS patients with a history of injection drug use are often infected with hepatitis B or C. Herpetic infections with HIV disease tend to be nuisances and not life-threatening. Pneumocystis infections are seen in patients with more severe acquired or inherited immunodeficiency disorders, particularly patients with AIDS, which affect cell-mediated immunity.

47 D The X-linked disorder known as Wiskott-Aldrich syndrome (WAS) is characterized by thrombocytopenia, eczema, and decreased IgM. IgA may be increased. As in many immunodeficiency disorders, there is immune dysregulation, and in WAS there is an increased risk of non-Hodgkin lymphoma. Dementia can be seen in patients with AIDS. Hypocalcemia is seen in neonates with DiGeorge syndrome that affects parathyroid glands. Rheumatoid arthritis can complicate isolated IgA deficiency and common variable immunodeficiency, conditions with survival to adulthood. A deficiency of complement C3 may be complicated by immunecomplex glomerulonephritis.

48 C Opportunistic infections in an injection drug user suggest a diagnosis of AIDS from HIV infection. The most common neoplasms seen in association with AIDS are B cell non-Hodgkin lymphomas. Opportunistic infections of the brain and central nervous system lymphomas are common in patients with AIDS, but glial neoplasms are not. Cervical dysplasias and carcinomas are increased in women with HIV infection, but such lesions are less frequent than lymphoma, and clear cell carcinomas of the female genital tract are associated with maternal diethylstilbestrol exposure. Lung cancers at this woman’s age are uncommon in any circumstance. Kaposi sarcoma is the one sarcoma associated with adult HIV infection. A rare tumor associated with AIDS in children is leiomyosarcoma.

49 F The reverse transcriptase gene of HIV undergoes mutation on average once per 2000 replications, a very high rate, which can account for the appearance of drug resistance. The drugs listed are reverse transcriptase inhibitors that are part of initial combination therapy for HIV infection. The absence of CD40 ligand interaction with CD40 explains the hyper-IgM syndrome. Chemokine receptors are important in facilitating initial HIV entry into cells, and mutations in these receptors may help explain variable susceptibility to and progression of HIV infection. The cytokine receptor γ chain is abnormal in severe combined immunodeficiency. The p24 antigen is a component of the HIV virion and is used to detect infection, but it is not a target of drug therapy. Protein tyrosine kinases are involved in signal transduction; they can be abnormal in conditions such as Bruton agammaglobulinemia, but not in patients receiving HIV drug therapy. None of the listed drugs is an HIV protease inhibitor.


**UNIT I  General Pathology**

50 C This woman’s original symptoms, although nonspecific, are characteristic for acute retroviral syndrome seen in more than half of adults with acute HIV infection. The average time to development of AIDS is 8 to 10 years; increased risk for opportunistic infections occurs as the CD4+ cell count decreases below 200/μL. Spondyloarthropathies (HLA-B27) and autoimmune diseases with autoantibodies, such as systemic lupus erythematosus (high titer ANA) or scleroderma (anticentromeric antibody), are unlikely to have such a long interval between illnesses, and are not as likely to manifest opportunistic infections without immunosuppressive therapy. Individuals with AIDS may have a polyclonal gammopathy, but not marked hypogammaglobulinemia. PBD9 250–251

BP9 145–146  TBD8 243–244  BP8 157–160

51 B Entry of HIV into cells requires binding to the CD4 molecule and co-receptor molecules, such as CCR5 and CXCR4. These HIV co-receptors are receptors for chemokines on the surface of T cells and macrophages. Mutations in genes encoding these co-receptor molecules cause individuals to be resistant to the effects of HIV infection because HIV cannot readily enter lymphocytes and macrophages. The other cell surface receptors are not relevant for HIV entry into cells. The p24 antigen that is contained within the HIV virion is not part of cell entry mechanisms, although its presence aids in detection of HIV infection. PBD9 246–247

BP9 145–146  TBD8 217–221  BP8 157–160

52 B Three types of cells are most likely to carry HIV: dendritic cells (Langerhans cells in epithelium, follicular dendritic cells in lymph nodes), monocyte-macrophages, and CD4+ T cells. Mucosal dendritic cells can bind to the virus and transport it to CD4+ cells in the lymph nodes. Whether the virus is internalized by mucosal dendritic cells is unclear. Monocyte-macrophages and CD4+ T cells express CD4 surface receptor and the chemokine co-receptors CCR5 and CXCR4; HIV can enter these cells. Follicular dendritic cells are distinct from mucosal or epithelial dendritic cells; they trap antibody-coated HIV virions by means of their Fc receptors. The other listed cells cannot be infected by HIV. PBD9 246–247

BP9 147–149  TBD8 242–243  BP8 158

53 C Macrophages can become infected with HIV and are not destroyed as CD4+ cells are. Instead, macrophages survive to carry the infection to tissues throughout the body, particularly the brain. HIV infection of the brain can result in encephalitis and dementia. CD8+ lymphocytes cannot be infected with HIV, but may aid in clearance of infected cells. Langerhans cells in mucosal surfaces may aid in initial HIV infection of CD4+ lymphocytes. Natural killer cells and neutrophils, as part of the innate immune system, play no significant role in HIV infection. PBD9 249–250

BP9 148–149  TBD8 242  BP8 161–162

54 A Pneumocystis jiroveci pneumonia is a common finding in patients with HIV infection progressing to AIDS. This patient’s low CD4+ count is characteristic of AIDS. The ANA test result is positive in various autoimmune diseases, but a decrease in CD4+ count is not typical of such conditions. Anti–neutrophil cytoplasmic autoantibody (C-ANCA or P-ANCA) can be seen in some patients with vasculitis. The anti–streptolysin O (ASO) titer is elevated in patients with rheumatic fever, but there is no serious immunosuppression. Detectable antibodies to a variety of cells and tissues can be found in a variety of autoimmune diseases such as SLE, and may also increase the risk for opportunistic infections. PBD9 253

BP9 151  TBD8 245–246  BP8 162–164

55 D Individuals infected with HIV are infected for life. They can transmit the virus to others via sexual intercourse even if they appear to be well, and particularly early and late in the course of infection when viremia is higher. His symptoms suggest recent infection and acute retroviral syndrome. The average time for the development of AIDS after HIV infection is 8 to 10 years. Seroreversion in HIV infection does not occur. HIV infection affects mainly CD4+ lymphocytes, with declining CD4+ counts presaging the development of clinically apparent AIDS. Progression of HIV disease is monitored by levels of HIV-1 mRNA in the blood and by CD4+ cell counts. When HIV-1 RNA increases and the CD4+ cell count declines, then the risk for opportunistic infections, neoplasms, and neurologic decline is increased.

PBD9 250–251  BP9 150–152  TBD8 244  BP8 162–163

56 E This patient has AIDS, with plaque like to nodular reddish purple lesions. Kaposi sarcoma is associated with a herpesvirus agent that is sexually transmitted: human herpesvirus 8 (HHV-8), also called the Kaposi sarcoma herpesvirus. Other herpesviruses are not involved in the pathogenesis of Kaposi sarcoma, although infection with these viruses can occur frequently in individuals with AIDS. HIV, although present in the lymphocytes and monocytes, is not detected in the spindle cells that proliferate in Kaposi sarcoma. With the exception of varicella-zoster virus, which is associated with dermatomally distributed skin vesicles known as shingles, skin lesions are uncommon manifestations of herpesviruses, which include cytomegalovirus, Epstein-Barr virus (EBV), and adenovirus.

PBD9 253–254  BP9 152  TBD8 246–247  BP8 164–165

57 A As HIV infection progresses, there is continuing, gradual loss of CD4+ cells. In the clinically latent phase of the infection, CD4+ cells continue to be replaced, but there is ongoing loss, and the stage of clinical AIDS is reached when the peripheral CD4+ count decreases to less than 200/μL, which usually occurs over 8 to 10 years. At this point, the risk of development of opportunistic infections and neoplasms typical of AIDS increases greatly. The extent of viremia also is an indication of the progression of HIV infection; an increase in HIV-1 RNA levels is seen as immunologic containment of HIV fails. In HIV infection, the numbers of CD8+ lymphocytes tend to be maintained. Cells of the granulocytic series are relatively unaffected, although patients with AIDS may have cytopenias. Follicular dendritic cells can be infected by...
HIV and pass the virions to CD4+ cells and macrophages, but the follicular dendritic cells and the macrophages are not destroyed in large numbers by the virus and can become a reservoir for infection. Natural killer cells, CD19+ B cells, and plasma cells are not directly affected by HIV.

Amyloid exhibits “apple-green” birefringence under polarized light after staining with Congo red. Amyloidosis is most often caused by excessive light chain production with plasma cell dyscrasias such as multiple myeloma (AL amyloid). His spinal lesions are focal collections of neoplastic plasma cells. Chronic inflammatory conditions, such as rheumatic fever, ankylosing spondylitis, and systemic sclerosis, also may result in amyloidosis (AA amyloid). His spinal lesions are focal collections of neoplastic plasma cells. Chronic inflammatory conditions, such as rheumatic fever, ankylosing spondylitis, and systemic sclerosis, also may result in amyloidosis (AA amyloid). His spinal lesions are focal collections of neoplastic plasma cells. Chronic inflammatory conditions, such as rheumatic fever, ankylosing spondylitis, and systemic sclerosis, also may result in amyloidosis (AA amyloid). His spinal lesions are focal collections of neoplastic plasma cells.

In chronic inflammatory conditions such as rheumatoid arthritis, the serum amyloid-associated (SAA) precursor protein forms the major amyloid fibril protein AA. SAA is an acute phase reactant that increases with inflammatory conditions. C-reactive protein also is an acute phase reactant whose level is increased in inflammatory conditions; unlike SAA, however, this does not form amyloid. Amyloid is deposited in interstitial locations. All amyloid shows characteristic “apple-green” birefringence under polarized light microscopy after Congo red staining—anything else would not be amyloid. Amyloid derived from β2-microglobulin occurs with hemodialysis-associated amyloidosis. Amyloid derived from light chains in association with multiple myeloma has AL fibrils. Cardiac and heredofamilial forms of amyloidosis have fibrils derived from prealbumin transthyretin (TTR).

These findings are characteristic of cardiac amyloidosis. Because of the patient’s age, a senile cardiac amyloidosis, resulting from deposition of transthyretin, is most likely. α-Fetoprotein is present during fetal life, but it is best known in adults as a serum tumor marker. β2-Microglobulin contributes to the development of amyloidosis associated with long-term hemodialysis. Calcitonin forms the precursor for amyloid deposited in thyroid medullary carcinomas. IgE is not a component of amyloid. Amyloidosis associated with plasma cell dyscrasias results from light chain production. Although the heart is commonly involved in light chain amyloidosis, the normal laboratory values and absence of plasma cell collections in the marrow argue against a plasma cell dyscrasia. Troponin proteins in cardiac muscle are useful markers for acute coronary syndromes, but are not involved in amyloid formation.
1. A 40-year-old man has a positive stool guaiac test during a routine physical examination. A colonoscopy is performed and a 0.9-cm, circumscribed, pedunculated mass on a short stalk is found in the upper rectum. Which of the following terms best describes this lesion?
   A. Adenoma
   B. Carcinoma
   C. Choristoma
   D. Hamartoma
   E. Hyperplasia
   F. Sarcoma

2. A 32-year-old woman has experienced dull pelvic pain for the past 2 months. Physical examination shows a right adnexal mass. An abdominal ultrasound scan shows a 7.5-cm cystic ovarian mass. The mass is surgically excised. The surface of the mass is smooth, and it is not adherent to surrounding pelvic structures. On gross examination, the cystic mass is filled with hair. Microscopically, squamous epithelium, tall columnar glandular epithelium, cartilage, and fibrous connective tissue are present and resemble normal tissue counterparts. Which of the following is the most likely diagnosis?
   A. Adenocarcinoma
   B. Fibroadenoma
   C. Glioma
   D. Hamartoma
   E. Mesothelioma
   F. Rhabdomyosarcoma
   G. Teratoma

3. A 62-year-old man has had several episodes of hematuria in the past week. He has a 48 pack-year history of smoking cigarettes. On physical examination, there are no abnormal findings. A urinalysis shows 4+ hematuria, and cytologic examination of the urine shows that atypical cells are present. A cystoscopy is performed and a 4-cm sessile mass with a nodular, ulcerated surface is seen in the dome of the bladder. Which of the following terms best describes this lesion?
   A. Adenoma
   B. Carcinoma
   C. Fibroma
   D. Papilloma
   E. Sarcoma

4. A 23-year-old woman has noted a nodule on the skin of her upper chest. She reports that the nodule has been present for many years and has not changed in size. It is excised, and the microscopic appearance is shown in the figure. Which of the following neoplasms is this lesion most likely to be?
   A. Fibroadenoma
   B. Hemangioma
   C. Leiomyoma
   D. Lipoma
   E. Melanoma
   F. Nevus
A 50-year-old woman undergoes screening colonoscopy as part of a routine health maintenance work-up. An isolated 1-cm pedunculated polyp is found in the sigmoid colon. The excised polyp histologically shows well-differentiated glands with no invasion of the stalk. Which of the following investigational research procedures can distinguish most clearly whether the polyp represents hyperplasia of the colonic mucosa or a tubular adenoma?

A. Flow cytometry to quantitate cells in the S phase
B. Histochemical staining for mucin
C. Immunohistochemical staining for keratin
D. Molecular marker of clonality

A 39-year-old woman underwent a routine health maintenance examination for the first time in many years. A Pap smear was obtained, and the result reported was abnormal. On pelvic examination, a red, slightly raised, 1-cm lesion on the anterior ectocervix at the 2 o’clock position was excised and biopsied. The microscopic appearance on medium-power magnification is shown in the figure. Which of the following is most characteristic of this patient’s condition?

A. Primary neoplasm in the endometrium
B. Elevated CA-125 level in the serum
C. Positive HSV-2 molecular test in the lesion
D. Pulmonary nodules on a chest radiograph
E. No recurrence following local excision

A 53-year-old woman has noticed increasing malaise. On physical examination, there are no abnormal findings, but a stool guaiac test is positive. Her hemoglobin level is 7.9 g/dL. A colonoscopy is performed, and a 3-cm sessile mass is found in the cecum. A biopsy specimen of the mass shows a moderately differentiated adenocarcinoma confined to the mucosa. An abdominal CT scan shows no lymphadenopathy or hepatic lesions. Given this information, which of the following is the best course of action?

A. Administer a multiagent chemotherapeutic regimen
B. Observe the lesion for further increase in size
C. Remove the entire colon to prevent a recurrence
D. Resect the tumor and some normal surrounding tissue
E. Search for a primary malignancy in another organ

A 57-year-old woman has experienced an increasing feeling of fullness in her neck along with a 3-kg (7-lb) weight loss over the past 3 months. On physical examination, there is a firm, fixed mass in a 3 x 5 cm area in the right side of the neck. A CT scan shows a solid infiltrating mass in the region of the right lobe of the thyroid gland. A biopsy of the mass is performed and the microscopic appearance is shown in the figure. All areas of the tumor have similar morphology. Which of the following terms best describes this neoplasm?

A. Anaplastic
B. Apoptotic
C. Dysplastic
D. Metaplastic
E. Well-differentiated

A Pap smear obtained from a 29-year-old woman during a routine health maintenance examination is abnormal. She is currently asymptomatic. She has a history of multiple sexual partners. Cervical biopsy specimens are obtained and the microscopic appearance is shown in the figure. Which of the following is the most likely diagnosis?

A. Adenocarcinoma
B. Carcinoma in situ
C. Dysplasia
D. Hamartoma
E. Metaplastic transformation
F. Squamous cell carcinoma
A 44-year-old woman feels painless lumps in her armpit, which were not present a month ago. On examination, right axillary lymphadenopathy is present. The nodes are painless but firm. Which of the following is the most likely lesion in her right breast?

A  Acute mastitis with abscess  
B  Fibroadenoma  
C  Infiltrating lobular carcinoma  
D  Intraductal carcinoma  
E  Leiomyosarcoma

A 69-year-old woman has experienced increasing malaise and a 10-kg weight loss over the past year. She dies of massive pulmonary thromboembolism. The gross appearance of the liver at autopsy is shown in the figure. Which of the following best describes the lesions seen in her liver?

A  Invasive angiosarcoma  
B  Hepatocellular carcinoma  
C  Leukemic infiltration  
D  Metastatic adenocarcinoma  
E  Multifocal hepatic adenomas

A 66-year-old man with chronic cough has an episode of hemoptysis. On physical examination, there are no abnormal findings. A chest radiograph shows a 6-cm mass in the right lung. A sputum cytologic analysis shows neoplastic squamous cells. Metastases from his lung lesion are most likely to be found at which of the following sites?

A  Cerebral hemisphere  
B  Chest wall muscle  
C  Hilar lymph nodes  
D  Splenic red pulp  
E  Vertebral bone marrow

An epidemiologic study of cancer deaths recorded in the last half of the 20th century is conducted. The number of deaths for one particular cancer had increased markedly in developed nations. More than 30% of cancer deaths in men, and more than 24% of cancer deaths in women, were caused by this neoplasm in 1998. In some nations, prevention strategies reduced deaths from this cancer. Which of the following neoplasms was most likely identified by this study?

A  Cerebral glioma  
B  Bronchogenic carcinoma  
C  Hepatocellular carcinoma  
D  Colonic adenocarcinoma  
E  Pancreatic adenocarcinoma  
F  Skin melanoma

An epidemiologic study analyzes health care benefits of cancer screening techniques applied to persons more than 50 years of age. Which of the following diagnostic screening techniques used in health care is most likely to have the greatest impact on reduction in cancer deaths in Europe and North America?

A  Chest radiograph  
B  Mammography  
C  Pap smear  
D  Serum tumor markers  
E  Stool guaiac  
F  Urine cytology

A 38-year-old woman has abdominal distention that has been worsening for the past 6 weeks. An abdominal CT scan shows bowel obstruction caused by a 6-cm mass in the jejunum. At laparotomy, a portion of the small bowel is resected. Flow cytometric analysis of a portion of the tumor shows a clonal population of B lymphocytes with high S phase. Translocation with activation of which of the following nuclear oncogenes is most likely to be present in this tumor?

A  APC  
B  EGF  
C  MYC  
D  p53  
E  RAS

A 50-year-old woman has had easy fatigability and noted a dragging sensation in her abdomen for the past 5 months. Physical examination reveals that she is afebrile. She has marked splenomegaly, but no lymphadenopathy. Laboratory studies show her total WBC count is 250,000/mm³ with WBC differential count showing 64% segmented neutrophils, 11% band neutrophils, 7% metamyelocytes, 5% myelocytes, 4% myeloblasts, 3% lymphocytes, 2% basophils, 2% eosinophils, and 2% monocytes. A bone marrow biopsy is performed, and karyotypic analysis of the cells reveals a t(9;22) translocation. Medical treatment with a drug having which of the following modes of action is most likely to produce a complete remission in this patient?

A  Activating cellular caspases  
B  Antibody binding to EGF receptors  
C  Delivering normal p53 into cells with viral vectors  
D  Inhibiting tyrosine kinase activity  
E  Preventing translocation of β-catenin to the nucleus
A 54-year-old woman notes a lump in her right breast. Physical examination shows a 2-cm mass fixed to the underlying tissues beneath the areola and three firm, nontender, lymph nodes palpable in the right axilla. There is no family history of cancer. An excisional breast biopsy is performed, and microscopic examination shows the findings in the figure. Over the next 6 months, additional lymph nodes become enlarged, and CT scans show nodules in the lung, liver, and brain. Which of the following molecular abnormalities is most likely to be found in her carcinoma cells?

A. Amplification of the ERBB2 (HER2) gene
B. Deletion of one RB gene copy
C. Fusion of BCR and C-ABL genes
D. Inactivation of one BRCA1 gene copy
E. Mutation of one p53 gene copy

The mother of a 5-year-old boy notices that his abdomen has enlarged in the past 6 months. On physical examination, there is an ill-defined abdominal mass. An abdominal CT scan shows a 9-cm mass in the region of the right adrenal gland. The mass is removed and microscopically shows primitive hyperchromatic cells. Cytogenetic analysis of tumor cells shows many double minutes and homogeneously staining regions. Which of the following genes is most likely to have undergone alterations to produce these findings?

A. BCL1 (cyclin gene)
B. BCL2 (anti-apoptosis gene)
C. IL2 (growth factor gene)
D. K-RAS (GTP-binding protein gene)
E. Lyn (tyrosine kinase gene)
F. N-MYC (transcription factor gene)

A 34-year-old sexually active woman undergoes a routine physical examination. There are no abnormal findings. A Pap smear is obtained as part of the pelvic examination. Cytologically, the cells obtained on the smear from the cervix show severe epithelial dysplasia (high-grade squamous intraepithelial lesion). Which of the following therapeutic options is most appropriate for this woman?

A. Antibiotic therapy
B. Excision
C. Ovarian removal
D. Screening of family members
E. Watchful waiting

An epidemiologic study investigates the potential morphologic and molecular alterations that may contribute to the development of cancers in a population. Data analyzed from resected colonic lesions show changes that suggest the evolution of a sporadic colonic adenoma into an invasive carcinoma. Which of the following best describes the mechanism producing these changes leading to colonic malignancies?

A. Activation of proto-oncogenes by chromosomal translocation
B. Extensive regeneration of tissues increasing the mutation rate in regenerating cells
C. Inheritance of defects in TP53 genes that increase the susceptibility to develop cancer
D. Overexpression of growth factor receptor genes
E. Stepwise accumulation of multiple proto-oncogene and tumor suppressor gene mutations

A 61-year-old woman has noted a feeling of pelvic heaviness for the past 6 months. On physical examination, there is a palpable nontender lower abdominal mass. An abdominal ultrasound scan shows a 12-cm solid mass in the uterine wall. A total abdominal hysterectomy is performed. The mass has the microscopic appearance of a well-differentiated leiomyosarcoma. One year later, a chest radiograph shows a 4-cm nodule in her right lower lung. Cytologic analysis of a fine-needle biopsy specimen of the nodule shows a poorly differentiated sarcoma. The patient’s medical history indicates that she has smoked cigarettes most of her adult life. Which of the following mechanisms best explains these findings?

A. Continued cigarette smoking by the patient
B. Development of a second primary neoplasm
C. Inheritance of a defective RB gene
D. Immunodeficiency with HIV infection
E. Metastasis from an aggressive tumor subclone

A 70-year-old woman reported a 4-month history of a 4-kg weight loss and increasing generalized icterus. On physical examination, she has midepigastric tenderness on palpation. An abdominal CT scan shows a 5-cm mass in the head of the pancreas. Fine-needle aspiration of the mass is performed. On biochemical analysis, the neoplastic cells show continued activation of cytoplasmic kinases. Which of the following genes is most likely to be involved in this process?

A. APC
B. MYC
C. p53
D. RAS
E. RET
F. sis
24 A 22-year-old man has a raised, pigmented lesion on his forearm that has increased in size and become more irregular in color over the past 4 months. Physical examination shows a 0.5 × 1.2 cm black-to-brown asymmetric lesion with irregular borders. An excisional biopsy specimen shows clusters of pleomorphic pigmented cells that extend into the reticular dermis. Family history indicates that the patient’s maternal uncle died from a similar tumor. His grandfather required enucleation of the right eye because of a “dark brown” retinal mass. Which of the following genes is most likely to have undergone mutation to produce these findings in this family?

- A BCL2 (anti-apoptosis gene)
- B c-MYC (transcription factor gene)
- C IL2 (growth factor gene)
- D Lyn (tyrosine kinase gene)
- E p16 (cell cycle inhibition)
- F p53 (DNA damage response gene)

25 A 3-year-old child has exhibited difficulty with vision in her right eye. On physical examination, there is leukocoria of the right eye, consistent with a mass in the posterior chamber. MR imaging shows a mass that nearly fills the globe. The child undergoes enucleation of the right eye. Which of the following molecular abnormalities is most likely to have been altered by mutation to produce these findings?

- A BCR-ABL
- B BCL2
- C hMSH2
- D K-RAS
- E NF1
- F p53
- G RB

26 A 76-year-old man has experienced abdominal pain for the past year. On physical examination, there is an epigastric mass. An abdominal CT scan shows a 10-cm mass in the body of the pancreas. A fine-needle biopsy specimen of this mass shows a moderately differentiated adenocarcinoma. Mutational analysis of the carcinoma cells shows inactivation of cyclin-dependent kinase inhibitor with loss of growth-suppression. Regulatory pathways controlled by which of the following genes are most likely altered in this man’s carcinoma?

- A BCL2
- B β-Catenin
- C MYC
- D p53
- E TGF-β

27 A 55-year-old man has had hemoptysis and worsening cough for the past month. On physical examination, wheezes are auscultated over the right lung posteriorly. A chest radiograph shows a 6-cm right perihilar mass. A fine-needle aspiration biopsy is performed and yields cells with the microscopic appearance of non-small cell bronchogenic carcinoma. Molecular analysis of the neoplastic cells shows a p53 gene mutation. Which of the following mechanisms has most likely produced the neoplastic transformation?

- A Inability to hydrolyze GTP
- B Growth factor receptor activation
- C Loss of cell cycle arrest
- D Microsatellite instability
- E Transcriptional activation

28 A 26-year-old man with a family history of colon carcinoma undergoes a surveillance colonoscopy. It reveals hundreds of polyps in the colon, and two focal 0.5-cm ulcerated areas. A biopsy specimen from an ulcer reveals irregularly shaped glands that have penetrated into the muscular layer. Which of the following molecular events is believed to occur very early in the evolution of his colonic disease process?

- A Activation of the WNT signaling pathway
- B Inability to hydrolyze GTP-bound RAS
- C Loss of heterozygosity affecting the p53 gene
- D Mutations in mismatch repair genes.
- E Translocation of BCL2 from mitochondria to cytoplasm

29 A 63-year-old man has a cough with hemoptysis for 10 days. He has a 65 pack-year history of smoking. A chest CT scan shows a 5-cm right hilar mass. Bronchoscopy is performed, and lung biopsy specimens show small cell anaplastic lung carcinoma. His family history shows three first-degree maternal relatives who developed leukemia, sarcoma, and carcinoma before age 40 years. Which of the following gene products is most likely to have been altered by mutation to produce these findings?

- A APC (tumor suppressor)
- B BCL2 (anti-apoptosis)
- C K-RAS (GTP binding)
- D NF1 (GTPase activation)
- E p53 (DNA damage response)

30 A 30-year-old man has a 15-year history of increasing numbers of benign skin nodules. On physical examination, the firm, nontender, subcutaneous nodules average 0.5 to 1 cm. Further examination shows numerous oval 1- to 5-cm flat, light brown skin macules. Ophthalmoscopic examination shows hamartomatous nodules on the iris. A biopsy specimen of one skin nodule shows that it is attached to a peripheral nerve. Which of the following molecular abnormalities is most likely related to his clinical presentation?

- A Decreased susceptibility to apoptosis
- B Impaired functioning of mismatch repair
- C Increased production of epidermal growth factor
- D Lack of nucleotide excision repair
- E Persistent activation of the RAS gene
- F Reduced expression of RB protein

31 A 53-year-old man diagnosed with oral cancer and treated with radiation and chemotherapy 1 year ago now has a positron emission tomography (PET) scan of his neck that shows a single focus of increased uptake. This focus is resected and microscopic examination shows that it is a metastasis. Molecular analysis of this cancer shows p53, PTEN, and c-MYC gene mutations. Which of the following metabolic pathways is most likely up-regulated to promote his cancer cell survival and proliferation?

- A Aerobic glycolysis
- B Gluconeogenesis
- C Hexose monophosphate shunt
- D Oxidative phosphorylation
- E Purine degradation
**32** A 49-year-old man has a lump near his right shoulder that has been increasing in size for the past 8 months. On physical examination, a 4-cm, firm, nontender mass is palpable in the right supraclavicular region. The mass is excised, and microscopically it shows a lymphoid neoplasm. Karyotypic analysis of the cells shows a chromosomal translocation, t(14;18), bringing the immunoglobulin heavy chain gene together with another gene. Which of the following genes is most likely activated by this translocation?

- A $APC$ (tumor suppressor gene)
- B $BCL2$ (anti-apoptosis gene)
- C $BRCA1$ (DNA repair gene)
- D $c-MYC$ (transcription factor gene)
- E $IL2$ (growth factor gene)
- F $K-RAS$ (GTP-binding protein gene)

**33** A 40-year-old man notices an increasing number of lumps in his groin and armpit over the past 5 months. On physical examination, he has generalized nontender lymph node enlargement and hepatosplenomegaly. A chest CT scan reveals multiple bilateral pulmonary “cannonball” nodules. A left breast biopsy is performed, and on microscopic examination shows abnormal lymphoid cells with many mitotic figures and many apoptotic nuclei. Which of the following mechanisms has most likely produced this lymphoid neoplasm?

- A Diminished apoptosis
- B Gene amplifications
- C Increased tyrosine kinase activity
- D Loss of cell cycle inhibition
- E Reduced DNA repair

**34** In an experiment, cells from human malignant neoplasms explanted into tissue culture medium continue to replicate. This allows development of “immortal” tumor cell lines that are extremely useful for the study of tumor biology and responses to therapeutic modalities. Activation of which of the following molecular components is most likely to endow these tumor cells with limitless replicative ability in vivo and in vitro?

- A Hypoxia-induced factor 1
- B $BCL2$ gene
- C Cyclin-dependent kinase gene methylation
- D DNA replication repair
- E Telomerase

**35** A 60-year-old man has noted a nodule in his neck that has increased rapidly in size over the past 2 months. On physical examination, there is a firm, nontender, 10-cm mass in the left lateral posterior neck that appears to be fused cervical lymph nodes. Hepatosplenomegaly is noted. A head CT scan reveals a mass in the Waldeyer ring near the pharynx. A biopsy of the neck mass is performed, and on microscopic examination shows abnormal lymphoid cells with many mitotic figures and many apoptotic nuclei. He is treated with a cocktail of cell cycle–acting chemotherapeutic agents. The cervical and oral masses shrink dramatically over the next month. Based on his history and response to treatment, the tumor cells are most likely to have which of the following features?

- A Diminished vascularity
- B Evolution of polyclonality
- C High growth fraction
- D Limited capacity to metastasize
- E Strong expression of tumor antigens

**36** A 30-year-old man has a pheochromocytoma of the left adrenal gland; a sibling had a cerebellar hemangioblastoma. He undergoes adrenalectomy, and on microscopic examination there is extensive vascularity of the neoplasm. Mutational analysis of the neoplastic cells shows that both allelic copies of a gene have been lost, so that a protein that binds to hypoxia-inducible factor 1-alpha is no longer ubiquitinated, but instead translocates to the nucleus and activates transcription of $VEGF$. Which of the following genes is most likely mutated in this man?

- A $APC$
- B $BCL2$
- C $EGF$
- D $HER2$
- E $HST1$
- F $MYC$
- G $VHL$

**37** A 48-year-old woman notices a lump in her left breast. On physical examination, she has a firm, nonmovable, 2-cm mass in the upper outer quadrant of the left breast. There are enlarged, firm, nontender lymph nodes in the left axilla. A fine-needle aspiration biopsy is performed, and the cells present are consistent with carcinoma. A lumpectomy with axillary lymph node dissection is performed, and carcinoma is present in two of eight axillary nodes. Reduced expression of which of the following molecules by the tumor cells is most likely responsible for the lymph node metastases?

- A Estrogen receptors
- B $ERBB2$ (HER-2)
- C E-cadherin
- D Progesterone receptors
- E Tyrosine kinases

**38** A 55-year-old woman has felt an enlarging lump in her left breast for the past year. A hard, irregular 5-cm mass fixed to the underlying chest wall is palpable in her left breast. Left axillary nontender lymphadenopathy is noted. There is no hepatosplenomegaly. A chest CT scan reveals multiple bilateral pulmonary “cannonball” nodules. A left breast biopsy is performed, and on microscopic examination shows high-grade infiltrating ductal carcinoma. The appearance of the nodules in her lungs is most likely related to which of the following?

- A Internal mammary artery invasion by carcinoma cells
- B Lymphatic connections between the breast and the pleura
- C Overexpression of estrogen receptors within the carcinoma cell nuclei
- D Proximity of the breast carcinoma to the lungs
- E Pulmonary chemokines that bind carcinoma cell chemokine receptors
39 A study of colonic polyps is performed. Malignant cells localized to the polyg are compared to those from polyps showing invasion of the stalk. Molecular analysis shows up-regulation of certain molecules in the invasive malignant cells. Invasive lesions are more likely to exhibit lymphatic metastases. Which of the following markers is most likely to have increased expression in the invasive malignant epithelial cells?

A BCL2  
B CD44  
C EGFR  
D RAS  
E Vimentin

40 In a clinical trial, patients diagnosed with malignant melanoma are treated by infusion of autologous CD8+ T cells grown in vitro. These CD8+ T cells are known to kill melanoma cells, but not normal cells. Which of the following target antigens in the tumor cells are most likely recognized by these CD8+ T cells?

A Class I MHC molecules with a melanoma cell peptide  
B Class I MHC molecules with a peptide from normal melanocytes and melanoma cells  
C Class I MHC molecules plus a peptide derived from carcinoembryonic antigen  
D Class II MHC molecules with a melanoma cell peptide  
E Class II MHC molecules with a peptide from normal melanocytes and melanoma cells  
F Class II MHC molecules with laminin receptors on melanoma cells

41 An experiment involving carcinoma cells grown in culture studies the antitumor surveillance effects of the innate immune system. These carcinoma cells fail to express MHC class I antigens. It is observed, however, that carcinoma cells are lysed when an immune cell that has been activated by IL-2 is added to the culture. Which of the following immune cells is most likely to function in this manner?

A CD4+ lymphocyte  
B CD8+ lymphocyte  
C Macrophage  
D Neutrophil  
E NK cell  
F Plasma cell

42 A 33-year-old man has experienced occasional headaches for the past 3 months. He suddenly has a generalized seizure. CT scan of the head shows a periventricular 3-cm mass in the region of the right thalamus. A stereotactic biopsy of the mass yields large lymphoid cells positive for B cell markers. Which of the following underlying diseases is most likely to be found in this patient?

A Diabetes mellitus  
B HIV infection  
C Hypertension  
D Multiple sclerosis  
E Tuberculosis

43 An investigational study reviews cells harvested from patients 30 to 50 years of age who had right-sided colon cancer with multiple polyps present. These patients typically develop multiple malignant lesions of the colon during middle age. Molecular analysis of the cells from the lesions shows changes in hPMS1, hPMS2, and hMLH1 genes. Which of the following principles of carcinogenesis is best illustrated by this study?

A Carcinogenesis is a multistep process  
B Inability to repair DNA is carcinogenic  
C Many oncogenes are activated by translocations  
D Tumor initiators are mutagenic  
E Tumor promoters induce proliferation

44 A 12-year-old girl and a 14-year-old boy have developed skin nodules in predominantly sun-exposed areas of their skin over the past 5 years, but their six siblings have not. On physical examination, both children are of appropriate height and weight. The skin lesions are 1- to 3-cm maculopapular nodules that are erythematous to brown-colored and have areas of ulceration. Microscopic analysis of biopsy specimens of the skin lesions shows squamous cell carcinoma. The children have no history of recurrent infections, and their parents and other relatives are unaffected. Which of the following mechanisms is most likely to produce neoplasia in these children?

A Inherited mutation of the p53 gene  
B Chromosomal translocation  
C Failure of nucleotide excision repair of DNA  
D Ingestion of food contaminated with Aspergillus flavus  
E Infection with human papillomavirus

45 A 26-year-old woman has a lump in her left breast. On physical examination, she has an irregular, firm, 2-cm mass in the upper inner quadrant of the breast. No axillary adenopathy is noted. A fine-needle aspirate of the mass shows anaplastic ductal cells. The patient’s 30-year-old sister was recently diagnosed with ovarian cancer, and 3 years ago her maternal aunt was diagnosed with ductal carcinoma of the breast and had a mastectomy. Mutation involving which of the following genes is most likely present in this family?

A BCL2 (anti-apoptosis gene)  
B BRCA1 (DNA repair gene)  
C ERBB2 (growth factor receptor gene)  
D HST1 (fibroblast growth factor gene)  
E IL2 (growth factor gene)  
F K-RAS (GTP-binding protein gene)

46 In a study of patients with non-Hodgkin B cell lymphoma, a nuclear gene is found to be actively transcribed to mRNA and is transported into the cell cytoplasm. A protein is translated from this mRNA, with up-regulation of BCL2. In a control group without lymphoma, translation of the mRNA does not occur. How is the silencing of this active gene’s mRNA most likely to occur?

A Absence of tRNA  
B Binding to mRNA  
C Methylation of DNA  
D Mutation of mRNA  
E Up-regulation of mtDNA
A study of patients treated with chemotherapy protocols for cancer shows that 10% of them subsequently develop a second cancer, a much higher incidence compared with a control group not receiving chemotherapy. These chemotherapy protocols included the alkylating agent cyclophosphamide. What is the most likely mechanism by which this agent causes carcinogenesis in these treated cancer patients?

A. Activation of protein kinase C  
B. Activation of endogenous viruses  
C. Blockage of TGF-β pathways  
D. Direct DNA damage  
E. Inhibition of DNA repair  
F. Inhibition of telomerase

A 51-year-old man who works in a factory that produces plastic pipe has experienced weight loss, nausea, and vomiting over the past 4 months. On physical examination, he has tenderness to palpation in the right upper quadrant of the abdomen, and the liver span is increased. Laboratory findings include serum alkaline phosphatase, 405 U/L; AST, 45 U/L; ALT, 30 U/L; and total bilirubin, 0.9 mg/dL. An abdominal CT scan shows a 12-cm mass in the right lobe of the liver. A liver biopsy is performed, and microscopic examination shows a malignant neoplasm of endothelial cells. The patient has most likely been exposed to which of the following agents?

A. Arsenic  
B. Asbestos  
C. Benzene  
D. Beryllium  
E. Nickel  
F. Vinyl chloride  
G. Naphthalene

A 56-year-old woman has had vaginal bleeding for 1 week. Her last menstrual period was 10 years ago. On physical examination, a lower abdominal mass is palpated. An endometrial biopsy is performed and shows endometrial carcinoma. An abdominal CT scan shows a 6-cm mass in the left ovary. A total abdominal hysterectomy is performed. Microscopically, the ovarian mass is a granulosa-theca cell tumor producing estrogen. Which of the following best describes the relationship between these two neoplasms?

A. Genetic susceptibility to tumorigenesis  
B. Mutational inactivation of a tumor suppressor gene  
C. Paraneoplastic syndrome  
D. Promotion of carcinogenesis  
E. Tumor heterogeneity

A 42-year-old man is concerned about a darkly pigmented “mole” on the back of his hand. The lesion has enlarged and bled during the past month. On physical examination, there is a slightly raised, darkly pigmented, 1.2-cm lesion on the dorsum of the right hand. The lesion is completely excised. The microscopic appearance is shown in the figure. Which of the following factors presents the greatest risk for the development of this neoplasm?

A. Allergy to latex  
B. Asbestos exposure  
C. Chemotherapy  
D. Smoking tobacco  
E. Ultraviolet radiation

A 33-year-old woman with multiple sexual partners has had vaginal bleeding and discharge for the past 5 days. On physical examination, she is afebrile. Pelvic examination shows an ulcerated lesion arising from the squamocolumnar junction of the uterine cervix. A cervical biopsy is performed and microscopic examination reveals an invasive tumor containing areas of squamous epithelium, with pearls of keratin. In situ hybridization shows the presence of human papillomavirus type 16 (HPV-16) DNA within the tumor cells. Which of the following molecular abnormalities in this tumor is most likely related to infection with HPV-16?

A. Functional inactivation of the RB protein  
B. Increased expression of epidermal growth factor receptor  
C. Epigenetic silencing of the RB gene  
D. Inability to repair DNA damage  
E. Trapping of the RAS protein in a GTP-bound state
A 66-year-old woman has worked all of her life on a small family farm on the Kantō Plain near Tokyo. She has had no previous major illnesses, but has been feeling increasingly tired and weak for the past year. On physical examination, she is afebrile, but appears pale. Laboratory studies show hemoglobin, 11.3 g/dL; hematocrit, 33.8%; platelet count, 205,200/mm³; and WBC count, 64,000/mm³. Immunophenotyping of her leukocytes yields the findings shown in the figure. Assuming that the dominant cell population is clonal, which of the following microbial agents is most likely involved in this patient’s disease process?

A  Epstein-Barr virus  
B  Hepatitis B virus  
C  HIV-1  
D  Helicobacter pylori  
E  Human T cell lymphotropic virus type 1

A 40-year-old man has a history of intravenous drug use. Physical examination shows needle tracks in his left antecubital fossa. He has mild scleral icterus. Serologic studies for HBsAg and anti-HCV are positive. He develops hepatocellular carcinoma 15 years later. Which of the following viral characteristics best explains why this patient developed hepatocellular carcinoma?

A  Viral integration in the vicinity of proto-oncogenes  
B  Viral capture of proto-oncogenes from host cellular DNA  
C  Viral inflammatory changes with genomic damage  
D  Viral inactivation of RB and p53 gene expression  
E  Viral infection of inflammatory cells with host immunosuppression

A 61-year-old man with a history of chronic viral hepatitis has noted a 6-kg weight loss over the past 5 months. Physical examination shows no masses or palpable lymphadenopathy. An abdominal CT scan shows a nodular liver with a 10-cm mass in the right lobe. A stool guaiac test result is negative. An elevation in which of the following laboratory tests is most likely to be present in this man?

A  Alpha-fetoprotein  
B  CA-19-9  
C  Calcitonin  
D  Carcinoembryonic antigen  
E  Immunoglobulin M

A 49-year-old man experiences an episode of hemoptysis. On physical examination, he has puffiness of the face, pedal edema, and systolic hypertension. A chest radiograph shows an irregular perihilar 5-cm mass of the right lung. Laboratory studies show normal serum electrolytes. A transbronchial biopsy is performed, and the microscopic findings are shown in the figure. A bone scan shows no metastases. Immunohistochemical staining of the tumor cells is most likely to be positive for which of the following?

A  Antidiuretic hormone  
B  Corticotropin  
C  Erythropoietin  
D  Insulin  
E  Parathyroid hormone-related peptide

A 59-year-old man has noticed blood in his urine for the past week. Cystoscopy shows a 4-cm exophytic mass involving the right bladder mucosa near the trigone. After biopsy specimens are obtained, he undergoes a radical cystectomy. Examination of the excised specimen shows an anaplastic carcinoma that has infiltrated the bladder wall. Which of the following techniques applied to the cells from his neoplasm is most likely to categorize the cell of origin?

A  Chromosomal karyotyping  
B  Cytologic smear  
C  DNA microarray  
D  Flow cytometric analysis  
E  Immunohistochemistry
A 69-year-old man has noted a chronic cough for the past 3 months. On physical examination, there is mild stridor on inspiration over the right lung. A chest radiograph shows a 5-cm right hilar lung mass, and a fine-needle aspiration biopsy specimen of the mass shows cells consistent with squamous cell carcinoma. If staging of this neoplasm is denoted as T2N1M1, which of the following findings is most likely in this man?

A. Brain metastases  
B. Elevated corticotropin level  
C. Infiltration of the chest wall  
D. Obstruction of a mainstem bronchus  
E. Poorly differentiated tumor cells

A 76-year-old woman has reported a change in the caliber of her stools during the past month. On physical examination, there are no abnormal findings, but a stool sample is positive for occult blood. A colonoscopy is performed and a constricting mass involving the lower sigmoid colon is found. She undergoes a partial colectomy. Which of the following techniques used during surgery can best aid the surgeon in determining whether the resection is adequate to reduce the probability of a recurrence?

A. Electron microscopy  
B. Fine-needle aspiration  
C. Flow cytometry  
D. Frozen section  
E. Radiologic imaging  
F. Serum carcinoembryonic antigen assay

ANSWERS

1. A  

A discrete small mass such as that described is probably benign. Adenomas arise from epithelial surfaces. Though adenocarcinoma may arise from a colonic adenoma, such malignant lesions tend to be larger and more irregular. A choristoma is a benign neoplasm composed of tissues not found at the site of origin. A hamartoma is a rare benign mass composed of tissues usually found at the site of origin. A hyperplastic colonic lesion tends to be smaller and flatter. A sarcoma is a malignant neoplasm arising in mesenchymal tissues, not in epithelium.

PBD9 266–268 BP9 162–163 PBD8 260 BP8 174–175

2. G  

A teratoma is a neoplasm derived from totipotential germ cells that differentiate into tissues that represent all three germ layers: ectoderm, endoderm, and mesoderm. When the elements all are well differentiated, the neoplasm is “mature” (benign). Adenocarcinomas have malignant-appearing glandular elements. Fibroadenomas have a benign glandular and stromal component; they are common in the breast. Gliomas are found in the central nervous system. Hamartomas contain a mixture of cell types common to a tissue site; the lung is one site for this uncommon lesion. A mesothelioma arises from the lining of thoracic and abdominal body cavities. A rhabdomyosarcoma comprises cells that poorly resemble striated muscle; most arise in soft tissues.

PBD9 267–268 BP9 163–164 PBD8 261–262 BP8 175

3. B  

A large, irregular, ulcerated mass such as that described is most likely malignant, and the epithelium of the bladder gives rise to carcinomas. Urothelial carcinomas are associated with smoking. An adenoma is a benign epithelial neoplasm of glandular tissues. A fibroma is a benign mesenchymal neoplasm. A papilloma is a benign, localized mass that has an exophytic growth pattern. A sarcoma is derived from cells of mesenchymal origin; sarcomas are much less common than carcinomas.


4. B  

The small, discrete nature of this mass and its slow growth with nearly unchanged size suggest a benign neoplasm. The red color is consistent with vascularity. A hemangioma is a common benign lesion of the skin. Fibroadenomas arise in the breast. Leiomyomas, which are white, arise from smooth muscle and are most common in the uterus. Lipomas are yellow fatty tumors that can occur beneath the epidermis. Melanomas are malignant and tend to increase in size quickly; many are darkly pigmented. The benign counterpart to the melanoma is the nevus, which is quite common, but nevi are usually light to dark brown.

PBD9 268, 271 BP9 162, 164 PBD8 263 BP8 174–176

5. D  

A true neoplasm is a monoclonal proliferation of cells, whereas a reactive proliferation of cells is not monoclonal. Molecular genetic analysis, such as allelotype analysis with microsatellite markers, shows clonality. Reactive and neoplastic cellular proliferations may have similar histochemical and immunohistochemical staining patterns based on the type of cells and proteins that are present. Flow cytometry is effective at indicating the DNA content, aneuploidy, and growth fraction, but does not indicate clonality.

PBD9 267–270, 334 BP9 167, 212–213 PBD8 276–278 BP8 185–186

6. E  

The figure shows an in situ carcinoma of the squamous cervical epithelium with neoplastic growth only above the basement membrane. Tissue damage with repair and regeneration may give rise to metaplasia, which may progress to dysplasia, considered premalignant. In situ cancers, limited to the epithelium, are noninvasive, and local excision has a 100% cure rate. In situ lesions do not give rise to metastases and have not arisen elsewhere. This lesion is related to human papillomavirus (HPV) infection, not herpes simplex virus (HSV). CA-125 is most often a tumor marker for ovarian cancer.

PBD9 271–272 BP9 166 PBD8 264 BP8 177–178
7 D A malignant epithelial neoplasm arises in the mucosa but has a tendency to invade locally. A benign neoplasm is often well circumscribed, and compressed normal surrounding tissue appears to form a discrete border. This localized lesion can be resected easily, with adequate margins. Without evidence for spread outside the colon, chemotherapy is unlikely to be of benefit. The biopsy specimen shows a malignant lesion; it must be removed before it increases in size and invades locally or metastasizes. If there is no family history, a familial cancer with high risk of recurrence from multiple polyps is unlikely; local excision is adequate. Such a solitary mucosal lesion is unlikely to represent a metastasis.

PBD9 268–272 BP9 167–168 PBD8 268–269 BP8 174–177

8 A The cells shown in the figure show marked pleomorphism and hyperchromatism (anaplasia), and it is difficult to discern the cell of origin because no differentiation is noted. A bizarre tripolar mitotic figure is present. This degree of anaplasia is consistent with an aggressive, high-grade malignancy called anaplastic carcinoma. Apoptosis is single cell necrosis, but the cells shown appear viable and not fragmented. Dysplasia refers to changes within an epithelium that presage a neoplasm. Metaplasia with one epithelial cell type substituted for another may presage dysplasia and malignancy. Well-differentiated neoplasms tend to be less aggressive and slower growing and resemble the cell of origin.

PBD9 269–270 BP9 164–166 PBD8 262–265 BP8 177–178

9 F In the figure the disorderly, atypical epithelial cells involve the entire thickness of the epithelium. They extend through the underlying basement membrane and into the underlying stroma as rounded nests at the right, a process known as invasion. The ectocervix and the squamous metaplasia of endocervix give rise to dysplasia from which squamous cell carcinoma can arise. Carcinoma in situ is confined to the epithelium; if the basement membrane is breached, the lesion is no longer in situ, but rather invasive. An adenocarcinoma is a malignant neoplasm arising from glandular epithelium, such as the endocervix or endometrium, not the ectocervix. A dysplastic process could precede development of carcinoma in situ and squamous carcinoma; dysplasia involves only part of the thickness of the epithelium. A hamartoma contains a mixture of cell types common to a tissue site. Metaplasia can occur in response to persistent infection with human papillomavirus (HPV) and other inflammatory conditions. Metaplasia can be the precursor to dysplasia.

PBD9 271–272 BP9 167–168 PBD8 265 BP8 177–179

10 C Lymphatic spread, especially to regional lymph nodes draining from the primary site, is typical of a carcinoma. An intraductal carcinoma has not extended beyond the basement membrane, but an infiltrating carcinoma has acquired the ability to invade and spread via metastasis. The primary site may be difficult to detect if small or deep, and hence the need for radiologic imaging, such as mammography. A fibroadenoma is a benign neoplasm and cannot invade or metastasize. Infection from a breast abscess can spread to the lymph nodes, but the resulting nodal enlargement is typically associated with pain—a cardinal sign of acute inflammation. Sarcomas uncommonly metastasize to lymph nodes, and a leiomyosarcoma of breast is rare.

PBD9 272–273 BP9 168–169 PBD8 268–270 BP8 180

11 D The figure shows the appearance of multiple variably sized tan metastatic lesions in the liver from hematogenous spread of carcinoma. Adenocarcinomas from abdominal primary sites such as colon, pancreas, and stomach are most likely. Thromboembolism suggests a hypercoagulable state such as a paraneoplastic syndrome. Angiosarcomas of the liver are uncommon. A primary malignancy typically appears as a dominant mass, not multiple masses. Although some benign tumors, such as leiomyomas of the uterus, can be multiple, this is not the rule in the liver, and hepatic adenomas are rare. Although hepatocellular carcinomas can have “satellite” nodules, widespread nodules such as those seen in the figure are more characteristic of metastases. Leukemic infiltrates typically do not produce large mass lesions, though some lymphomas may do so. Resection of multiple metastases is usually futile.

PBD9 274 BP9 168–169 PBD8 269 BP8 179–181

12 C Carcinomas metastasize through lymphatics most often, usually to regional nodes first. Hematogenous metastases are possible, however, to sites such as bone marrow, liver, or the opposite lung. About half of all cerebral metastases arise from lung primary carcinomas. Soft-tissue metastases to muscle, fat, and connective tissues are rare, as are splenic metastases.

PBD9 273–274 BP9 168–169 PBD8 269 BP8 179–181

13 B The decrease in the number of gastric cancers may be related to reduced numbers of dietary carcinogens or a decrease in the prevalence of Helicobacter pylori infection; however, the exact reason is obscure. Cerebral gliomas are not as common as carcinomas; an urban legend links them to cell phone use, but legitimate epidemiologic studies have not made this link. Angiosarcomas of the liver are quite rare; they are epidemiologically linked to vinyl chloride exposure. Leukemias and lymphomas are not as common as carcinomas. Pulmonary small cell carcinomas are related to smoking, and the numbers have decreased in many countries with campaigns to reduce smoking; the death rate is typically high because the prognosis for lung cancer is so poor.

PBD9 275–277, 279 BP9 170, 204 PBD8 272, 315 BP8 214

14 B Incidence of lung cancers increased dramatically in the 20th century because of the popularity of cigarette smoking. As the number of individuals in a population who smoke increases, so do the number of lung cancers. Some cancers of the urinary tract, oral cavity, esophagus, and pancreas also are causally related to smoking. Breast, prostate, and colon cancers remain common in developed nations, but the number of cases has not increased sharply. Pap smear screening and human papillomavirus (HPV) vaccination markedly decreases numbers of cervical cancers. There has
been an increase in the incidence of melanomas worldwide, but there are still far fewer cases of melanomas than of lung cancers. Hepatic and intracranial neoplasms in adults are far less common than lung cancers.

PBD9 276–277  BP9 170  PBD8 273  BP8 182–183

15 B A screening program should reliably detect early cancers with higher incidence. Breast cancer affects up to 1 in 9 women in these regions. Mammography may aid in detection of small cancers that have a better prognosis. A chest radiograph is an insensitive technique for detecting early lung cancers. Because Pap smear screening can detect dysplasias and in situ carcinomas that can be treated before progression to invasive lesions, deaths from cervical carcinoma have steadily decreased since this screening method became widely available in the last half of the 20th century. The introduction of human papillomavirus (HPV) vaccination will diminish the numbers of cervical cancers even further. Serum tumor markers have not proved useful as general screening techniques, although they are useful in selected circumstances. Use of stool guaiac has had a minimal effect on rates of death from colorectal carcinomas, but physicians are cautioned not to indicate “rectal deferred” on the physical examination report, and hence contribute to the problem. Urine cytology is better than urinalysis for detection of urothelial malignancies, but it does not have a high sensitivity.

PBD9 276–279  BP9 170–171  PBD8 324  BP8 182

16 C The MYC oncogene is commonly activated in Burkitt lymphoma because of a t(8;14) translocation. The MYC gene binds DNA to cause a transcriptional activation of growth-related genes such as that for cyclin D1, resulting in activation of the cell cycle. EGF (such as HER2 in breast cancers) encodes the epithelial growth factor receptor located on the cell surface. p53 and APC are tumor suppressor genes that are inactivated in many cancers, including colon cancer. RAS oncogene encodes a GTP-binding protein that is located under the cell membrane.

PBD9 284, 287–288  BP9 175  PBD8 284  BP8 190–191

17 D This patient has a classic history and t(9;22) translocation with chronic myelogenous leukemia. The translocation causes uncontrolled nonreceptor tyrosine kinase activity of the BCR-ABL fusion gene. These patients undergo remission with drugs such as imatinib that inhibit tyrosine kinases. Agents that activate caspases theoretically may help in many cases, especially when apoptosis is blocked as in tumors with BCL2 overexpression. Antibodies to epithelial growth factor receptors, such as ERBB2 (HER2) receptors, are beneficial in certain breast tumors with amplification of this gene. Delivery of p53 into cells by viral vectors has not yet been proven to be valuable in cancer treatment, and it is not used in chronic myelogenous leukemia. Translocation of β-catenin to the nucleus occurs in colon cancers when there is mutational loss of APC genes.

PBD9 284, 287  BP9 174  PBD8 283–284  BP8 174

18 A Infiltrating ductal and intraductal carcinoma are present in the figure. Increased expression of ERBB2 (HER2) can be detected immunohistochemically and by fluorescence in situ hybridization (FISH) in the biopsy specimen. One third of breast cancers may show this change. Such amplification is associated with a poorer prognosis. Detection of a specific gene product in the tissue has value for determination of treatment and prognosis. BRCA1 and p53 mutations, if inherited in the germ line, can predispose the patient to breast cancer and other tumors. With BRCA1, there is familiar history of breast cancer, often at a young age. The tumor suppressor gene p53 mutations predispose to many types of cancers. An inherited deletion of RB gene predisposes to retinoblastoma. The BCR-ABL fusion product, seen in chronic myeloid leukemia, often results from t(9;22).

PBD9 284–285  BP9 175  PBD8 281, 324  BP8 188

19 F Double minutes and homogeneously staining regions seen on a karyotype represent gene amplifications. Amplification of the N-MYC gene occurs in 30% to 40% of neuroblastomas, and this change is associated with a poor prognosis. The BCL1 and BCL2 genes are mutated in some non-Hodgkin lymphomas. The IL2 mutation may be present in some T cell neoplasms. K-RAS mutations are present in many cancers, but not typically childhood neoplasms. The Lyn mutation is seen in some immunodeficiency states.

PBD9 289  BP9 175  PBD8 306  BP8 190, 208

20 B Epithelial dysplasias, especially severe dysplasias, can be precursors of carcinomas. This is a key reason for Pap smear screening. The incidence of cervical carcinoma decreases when routine Pap smears are performed. Colposcopy with biopsy is indicated to determine the extent of the lesion for removal. Though related to human papillomavirus (HPV) infection, severe dysplasias are not amenable to antibiotic therapy. Ovarian neoplasms are not related to cervical dysplasias or carcinomas. In general, cervical cancers are not related to hereditary syndromes, and cervical dysplasias are not hereditary. Screening of family members is appropriate for those who have risk factors, such as multiple sexual partners. Regression of a severe dysplasia is unlikely.

PBD9 271, 333  BP9 210  PBD8 265  BP8 177–178

21 E Development of colonic adenocarcinoma typically takes years, during which time multiple mutations occur within the mucosa, including mutations involving such genes as APC (adenomatous polyposis coli), K-RAS, and p53. The accumulation of mutations, rather than their occurrence in a specific order, is most important in the development of a carcinoma. Activation of proto-oncogenes, extensive regeneration, faulty TP53 genes, and amplification of growth factor receptor genes all contribute to the development of malignancies, but they are not sufficient by themselves alone to produce a carcinoma from an adenoma of the colon. Inherited loss of wild-type TP53 contributes to multiple cancers, but not to sporadic adenomas.

22 E Although neoplasms begin as monoclonal proliferations, additional mutations occur over time, leading to subclones of neoplastic cells with various aggressive properties. This subcloning may allow metastases, greater invasiveness, resistance to chemotherapy, and morphologic differences to occur. Because sarcomas of the lung are rare, the lung mass is statistically a metastasis. Though second primary malignancies do arise, particularly in persons who have already had a malignancy, the odds favor a metastasis in a person with a prior malignancy. Sarcomas are not related to smoking tobacco. Inheritance of a mutant RB gene is most likely to lead to childhood retinoblastomas and osteosarcomas. Kaposi sarcoma is the sarcoma most often associated with AIDS from HIV infection.

PBD9 281–282, 321–322 BP9 177 PBD8 279 BP8 185–186

23 D The RAS oncogene is the most common oncogene involved in the development of human cancers. Mutations of the RAS oncogene reduce GTase activity, and RAS is trapped in an activated GTP-bound state. RAS then signals the nucleus through cytoplasmic kinases. The APC gene can cause activation of the WNT signaling pathway. The MYC oncogene is a transcriptional activator that is overexpressed in many tumors. The p53 tumor suppressor gene encodes a protein involved in cell cycle control. The RET proto-oncogene encodes a receptor tyrosine kinase involved in neuroendocrine cells of the thyroid, adrenal medulla, and parathyroids. The sis oncogene encodes platelet-derived growth factor receptor-β, which is overexpressed in certain astrocytomas.

PBD9 284–287 BP9 179–180 PBD8 282–283 BP8 188–190

24 E A family history of malignant melanoma is present. Familial tumors often are associated with inheritance of a defective copy of one of several tumor suppressor genes. In the case of melanomas, the implicated gene is called p16, or INK4a. The product of the p16 gene is an inhibitor of cyclin-dependent kinases. Germline mutations in CDKN2A may also underlie familial melanomas. With loss of control over cyclin-dependent kinases, the cell cycle cannot be regulated, favoring neoplastic transformation. BCL2 is present in some lymphoid neoplasms. The c-MYC gene is mutated in various carcinomas, but is not known to be specifically associated with melanomas. The IL2 mutation is associated with some T cell neoplasms. The Lyn mutation is seen in immunodeficiency states. p53 mutations occur in many cancers, but not specifically in familial melanomas.

PBD9 291, 293 BP9 176, 182 PBD8 286–287 BP8 181–182

25 G The RB gene is a classic example of the two-hit mechanism for loss of tumor suppression. About 60% of these tumors are sporadic, whereas the rest are familial from inheritance of a mutated copy of the RB gene. Loss of the second copy in retinoblasts leads to the occurrence of retinoblastoma in childhood. The RB gene controls the G1 to S transition of the cell cycle; with loss of both copies, this important checkpoint in the cell cycle is lost. The BCR-ABL fusion gene in chronic myelogenous leukemia is an example of overexpression of a gene product producing neoplasia. The BCL2 gene is an inhibitor of apoptosis. The hMSH2 gene is present in most cases of hereditary nonpolyposis colon cancer and functions in DNA repair. Many cancers have the K-RAS gene, which acts as an oncogene. The NFI gene product acts as a tumor suppressor; this is a component of neurofibromatosis (which usually does not involve the eye), and the neoplasms typically appear at a later age. Many cancers have the p53 tumor suppressor gene mutation, but this is not typical of childhood ocular neoplasms.

PBD9 290–293 BP9 182–184 PBD8 288–290 BP8 192–194

26 E TGF-β inhibits cell proliferation by activation of growth-inhibiting genes, such as the CDKIs. All pancreatic cancers and 83% of colon cancers have at least one mutation in the TGF-β pathway. The BCL2 family of genes acts as a regulator of apoptosis. The β-catenin pathway seen with the APC gene is involved with growth regulation; loss of the APC gene loci leads to failure in destruction of β-catenin, which translocates to the cell nucleus, where it functions as a transcription factor promoting growth. The MYC gene is a target of the activated RAS pathway. The p53 protein is involved in tumor suppression.

PBD9 285, 293 BP9 187 PBD8 294 BP8 197–198

27 C A p53 mutation involving both wild type alleles is one of the most common genetic alterations in human cancers, including the most common cancers—lung, colon, and breast. The loss of this tumor suppressor activity indicates that the cell cycle is not properly arrested in the late G1 phase, and when DNA damage occurs, DNA repair cannot be completed before the cell proliferates. Inability to hydrolyze GTP is a result of RAS oncogene activation. Growth factors such as EGF are activators of the cell cycle to promote cell growth. Microsatellite instability occurs with mutation in genes, such as hMSH2, that repair DNA damage. Transcriptional activation is a feature of the MYC proto-oncogene.

PBD9 293–296 BP9 185–187 PBD8 290–292 BP8 195–196

28 A The patient has a classic history of familial adenomatous polyposis with numerous adenomatous polyps and malignant transformation. The earliest event in the APC → adenocarcinoma sequence is loss of APC gene function. This prevents the destruction of β-catenin in the cytoplasm, which translocates to the nucleus and coactivates transcription of several genes. The APC → β-catenin sequence is a component of the WNT signaling pathway. RAS activation occurs after the malignant transformation sequence is initiated by the APC (gatekeeper) gene. Loss of cell cycle G1 arrest occurs with p53 loss late in the sequence. Mutations in mismatch repair genes give rise to hereditary nonpolyposis colon cancer syndrome from loss of ability to repair DNA damage. The BCL2 gene is not involved in the transition from adenoma to carcinoma.

PBD9 296–297 BP9 188 PBD8 292–293 BP8 197–198

29 E p53 is the most common target for genetic alterations in human neoplasms. Most are sporadic mutations,
although some are inherited. The inheritance of one faulty p53 suppressor gene predisposes to a “second hit” that eliminates the remaining p53 gene. Homozygous loss of the p53 genes dysregulates the repair of damaged DNA, predisposing individuals to multiple tumors, as in this case. The APC gene is mutated in sporadic colon cancers and in familial polyposis coli. The BCL2 gene is mutated in some non-Hodgkin lymphomas. The HER2 gene is one of the EGF receptor family members amplified in some breast cancers. The EGF mutation is most often seen in squamous cell carcinomas of the lung. K-RAS mutations are present in many cancers, but not typically in lymphoid malignancies. The NFI gene mutation is seen in neurofibromatosis type 1.

**30** E This patient has clinical features of neurofibromatosis type 1. The NFI gene encodes for a GTPase-activating protein that facilitates the conversion of active (GTP-bound) RAS to inactive (GDP-bound) RAS. Loss of NFI prevents such conversion and traps RAS in the active signal-transmitting stage that drives cell proliferation. Thus, the wild type NFI gene acts as a tumor suppressor. All other listed mechanisms also are involved in carcinogenesis, but in different tumor types.

**31** A The PET scan is based upon selective uptake of a glucose derivative into tumor cells. The Warburg effect occurs when cancer cells shift their metabolism to aerobic glycolysis for selective growth advantage under harsh circumstances. Glycolysis also yields pyruvate for anabolic demands of increased tumor doublings. The p53 and c-MYC genes favor this metabolic change, whereas PTEN inhibits tumor cell autophagy, giving cancer cells an edge in growth. Cancer cells are less differentiated than normal cells and thus have decreased ability to do many complex biochemical processes, so they favor a simple one—glycolysis. Gluconeogenesis is a function of hepatocytes in response to decreased caloric intake. The HMP shunt and Krebs cycle are more useful to normal cells maintaining themselves at the status quo. Neoplasms generate large amounts of purines from cell divisions and cell turnover that must be eliminated as uric acid, but neoplastic cells do not perform this task.

**32** B This is an example of chromosomal translocation that brings BCL2, an anti-apoptosis gene, close to another gene (immunoglobulin heavy chain gene). The BCL2 gene becomes subject to continuous stimulation by the adjacent enhancer element of the immunoglobulin gene, leading to overexpression. The APC gene is mutated in sporadic colon cancers and cancers associated with familial polyposis coli. The BRCA1 gene mutation is seen in some breast cancers. The c-MYC gene is found on chromosome 8, and the t(8;14) translocation seen in many Burkitt lymphomas leads to MYC overexpression. The IL2 mutation may be present in some T cell neoplasms. K-RAS mutations are present in many cancers, but not typically lymphoid malignancies.

**33** A The BCL2 gene controls production of a protein that inhibits apoptosis, and overexpression of this gene allows accumulation of abnormal cells in lymphoid tissues. Gene amplifications typically affect the ERBB2 (HER2) and MYC oncogenes. Increased tyrosine kinase activity results from mutations affecting the ABL oncogene. Loss of cell cycle inhibition results from loss of tumor suppressor genes such as p53. Reduced DNA repair occurs in the inherited disorder xeroderma pigmentosum.

**34** E Chromosomal telomere shortening in normal human cells limits their replicative potential and gives rise to replicative senescence. This occurs because most somatic cells lack the enzyme telomerase. Normal human stem cells do express telomerase. By contrast, 90% or more of human tumor cells show activation of telomerase, explaining continued tumor growth in the body and “immortalized” cell lines in culture. All other pathways listed cannot affect telomerase shortening, which is the rate-limiting step in indefinite replication of cells.

**35** C Some neoplasms, including certain lymphomas, have a high proportion of cells in the replicative pool (i.e., have high growth fraction). They grow rapidly and respond rapidly to drugs that kill dividing cells. Poor vascularity would not favor rapid growth, and many neoplasms elaborate growth factors that promote vascular proliferation. Monoclonality rather than polyclonality is typical of malignant tumors, though subclones of neoplastic cells do arise over time. Aggressive neoplasms tend to be more likely to metastasize. Tumors that are highly antigenic are likely to be controlled by the immune system and not to be rapidly growing.

**36** G Angiogenesis is a key feature of neoplasms because the growing tumor needs a blood supply, and up-regulation of factors such as VEGF and FGF help to keep the cancer growing. VEGF may be up-regulated by activation of hypoxia-inducible factor 1-alpha (HIF-1-alpha). The von Hippel-Lindau (VHL) gene acts as a tumor suppressor, and it normally produces a protein that binds to hypoxia-inducible factor 1-alpha so that it is cleared. VHL mutation leads to loss of this binding protein and activation of angiogenesis factors. Individuals with VHL syndrome have various neoplasms, including pheochromocytomas, renal cell carcinomas, and hemangioblastomas. The other listed genes have products that do not directly act on angiogenesis pathways.

**37** C Several pathologic mechanisms play a role in the development of tumor metastases. The tumor cells first must become discohesive and detach from the primary site, degrade the basement membrane and interstitial connective tissue, and then attach elsewhere to become metastases. Reduced expression of adhesion molecules such as
E-cadherins promotes metastases. Tumor cells can elaborate, not reduce, proteases such as metalloproteinases to promote discohesiveness. Expression of estrogen and progesterone receptors predicts breast cancer responsiveness to antihormone therapy, and there is a monoclonal antibody, trastuzumab, that targets HER-2, a form of epidermal growth factor receptor. Tyrosine kinase receptors within cells aid in signaling cell growth.

39 B Malignant transformation includes many genetic changes, including those rendering the malignant cells capable of invasion and metastases. CD44 plays a role in cell adhesion and enables malignant cells to metastasize. Solid tumors can express CD44 to enhance their spread to lymph nodes and other metastatic sites. Though such properties may have been present with the initial clone of malignant cells, the growth of the cancer increases the number of cells and the risk for spread. BCL2 plays a role in apoptosis. Growth factor receptor expression may make malignant cells susceptible to environmental influences, such as hormones, that drive growth. RAS gene mutations are present in many cancers and lead to loss of growth control. Vimentin is best known as a protein expressed in mesenchymal neoplasms, such as sarcomas, but it is also up-regulated in EMT.

40 A All human nucleated cells express MHC class I antigens. CD8+ T cells recognize peptides presented by MHC class I antigens. In many tumors, especially melanomas, the tumor cells produce peptides that can be presented by MHC class I molecules. Such tumor-specific peptides are not produced by other cells, so the CD8+ T cells specific for such peptides lyse melanoma cells, but not normal melanocytes or other normal cells.

41 E Several types of immune cells can recognize and help destroy tumor cells. Tumor antigens that are displayed via MHC class I molecules can be recognized by cytotoxic CD8+ cells. Normal human cells should display MHC class I antigens, but many cancers do not display their antigens well, and when MHC class I molecules are not displayed, NK cells are triggered to target these cells for lysis. Macrophages may work in concert with CD8+ cells and NK cells to phagocytize and kill tumor cells when up-regulated by interferon-γ. CD4+ “helper” cells do not play a direct anticancer role. Neutrophils are ineffective against cancer cells, but may be attracted to areas of tumor necrosis. Plasma cells may produce antibodies directed against tumor antigens, but such antibodies are ineffective in controlling tumors.

42 B Primary or secondary immunodeficiency diseases carry an increased risk of neoplasia, particularly lymphomas. B cell lymphomas of the brain are 1000-fold more common in patients with AIDS from HIV infection than in the general population. Patients with diabetes mellitus can experience various vascular and infectious complications, although not neoplasia. Hypertension can lead to central nervous system hemorrhages (strokes). Multiple sclerosis is a demyelinating disease of CNS white matter and carries no significant risk of neoplasia. Tuberculosis as a chronic infection may lead to amyloidosis, not neoplasia.
45 B Approximately 5% to 10% of breast cancers are familial, and 80% of these cases result from mutations in the BRCA1 and BRCA2 genes. Onset of these familial cancers occurs earlier in life than the sporadic cancers. The protein products of these genes are involved in DNA repair. BCL2 is overexpressed in some lymphoid neoplasms. The epithelial growth factor receptor ERBB2 (HER2) overexpression is present in some sporadic breast cancers; other EGF alterations can be seen in lung, bladder, gastrointestinal, ovarian, and brain neoplasms. The IL2 mutation is seen in some gastric cancers. IL2 overexpression is associated with some T cell neoplasms. K-RAS overexpression is seen in many cancers, including some breast cancers, but the early age of onset and family history in this case strongly suggest BRCA mutations.

46 B MicroRNAs (miRNAs) are encoded by about 5% of the human genome. miRNAs do not encode for proteins, but bind and inactivate or cleave mRNA, preventing translation of proteins by mRNA. This effectively silences gene expression without affecting the gene directly. There is abundant transfer RNA (tRNA) present in the cytoplasm that is not a rate-limiting step to translation. DNA methylation, particularly at CG dinucleotides, is a way of suppressing gene expression directly, as is seen with genomic imprinting. Mutations that occur in genes in DNA may result in reduced mRNA production or abnormal protein production, but mRNA itself is not mutated. Mitochondrial DNA (mtDNA) encodes for proteins that are mainly involved in oxidative phosphorylation metabolic pathways.

47 D Chemical carcinogens can have highly reactive electrophilic groups that can directly damage DNA, leading to mutations. Direct-acting agents, such as alkylating chemotherapy drugs, do not require conversion to a carcinogen. Some environmental toxic agents, such as polycyclic hydrocarbons, require metabolic conversion to a carcinogen and are called indirect-acting agents. Phorbol esters are examples of promoters of chemical carcinogenesis that cause tumor promotion by activating protein kinase C. This enzyme phosphorylates several substrates in signal transduction pathways, including those activated by growth factors, and the cells divide. Forced cell division predisposes the accumulation of mutations in cells previously damaged by exposure to a mutagenic agent (initiator). The TGF-β pathways work via growth inhibition. Proteins such as p53 that function in DNA repair pathways can become nonfunctional through mutation. Viral infections such as hepatitis B and C tend to promote growth by binding to p53 and inactivating its protective function. Telomerase activity is not affected by carcinogens.

48 F Vinyl chloride is a rare cause of liver cancer. This causal relationship was easy to show, however, because hepatic angiosarcoma is a rare neoplasm. Arsenic is a risk factor for skin cancer. Asbestos exposure is linked to pleural malignant mesothelioma and to bronchogenic carcinomas in smokers. Benzene exposure is linked to leukemias. Beryllium exposure can produce interstitial lung disease and lung cancer. Nickel exposure increases the risk of respiratory tract cancers. Exposure to naphthalene compounds is a risk factor for cancers of the urinary tract.

49 D Estrogen, similar to many other hormones and drugs, by itself is not carcinogenic, but it is responsible for stimulation of endometrial growth (hyperplasia), which has a promoting effect when cellular mutations occur to produce carcinoma. Inherited susceptibility can never be completely excluded when an individual has two tumors; this can occur in patients with inherited mutations in the p53 gene. In this case, however, there is a clear hormonal basis for the second tumor. Faulty tumor suppressor genes are not involved in hormonal promotion of a neoplasm. A paraneoplastic syndrome results from ectopic secretion of a hormone by tumor (e.g., lung cancer cells producing corticotropin). Tumor heterogeneity does not refer to two separate kinds of neoplasms; it refers to heterogeneity with a given tumor or metastasis.

50 E Radiation is oncogenic, and the risk increases with higher dosages. Cancers of thyroid and bone as well as leukemias may develop years following environmental radiation exposure. Dosages of therapeutic radiation are carefully controlled, but risk for subsequent malignancy is still increased. The Chernobyl nuclear reactor disaster affecting persons in Belarus and Ukraine is a cautionary tale regarding environmental exposure to radiation. Trauma is not a risk factor for development of cancer, although traumatic episodes often are recalled and irrationally associated with subsequent health problems. Lack of iodine leads to goiter but not to thyroid neoplasia. Hepatocellular carcinomas can arise in cirrhosis caused by chronic alcoholism. Ataxia telangiectasia is an inherited syndrome that carries an increased risk of development of leukemias and lymphomas. Arsenic exposure, which is uncommon, leads to lung and skin cancers.

51 E Worldwide, increasing numbers of skin cancers occur because of sun exposure. The ultraviolet light damages the skin and damages cellular DNA, leading to mutations that can escape cellular repair mechanisms. Allergic reactions do not promote cancer. Asbestos exposure increases lung carcinoma risk in smokers and can lead to rare mesotheliomas of pleura. Chemotherapeutic agents have carcinogenic potential, particularly alkylating agents such as cyclophosphamide, but leukemias and lymphomas are the usual result. Smoking tobacco is related to many cancers, but skin cancers are not typically associated with this risk factor.

52 A The oncogenic potential of human papillomavirus (HPV), a sexually transmissible agent, is related to products
of two early viral genes—E6 and E7. E7 protein binds to RB protein to cause displacement of normally sequestered transcription factors, which nullifies tumor suppressor activity of the RB protein. E6 protein binds to and inactivates the p53 gene product. Increased epidermal growth factor receptor expression is a feature seen in many pulmonary squamous cell carcinomas, and the related ERBB2 (HER2) receptor is seen in some breast carcinomas. Epigenetic modifications include DNA methylation and histone modifications which, depending on their nature, may enhance or inhibit gene expression. Inability to repair DNA damage plays a role in some colon and skin cancers. Trapping of GTP-bound RAS protein can occur in many tumors but is not related to HPV infection.

PBD9 326–327  BP9 202  PBD8 313  BP8 194, 212

53 E The largest cell population in the figure, determined to be clonal, is marking for CD4. This patient has a T cell leukemia, which develops in approximately 1% of individuals infected with human T cell lymphotropic virus type 1. Infection with Epstein-Barr virus is associated with various cancers, including Burkitt lymphoma and nasopharyngeal carcinoma. Infection with hepatitis B virus may result in hepatic cirrhosis, in which hepatocellular carcinoma may arise. HIV-1 infection causes AIDS, with a diminished CD4+ cell count. H. pylori promotes chronic gastritis with increased risk for gastric adenocarcinomas and B cell lymphomas.

PBD9 325–326  BP9 201  PBD8 312–313  BP8 211

54 C Although the hepatitis B virus (HBV) and hepatitis C virus (HCV) genomes do not encode for any transforming proteins, the regenerating hepatocytes are more likely to develop mutations, such as inactivation of p53. HBV does not have a consistent site of integration in the liver cell nuclei, and it does not contain viral oncogenes. Many DNA viruses, such as human papillomavirus (HPV), inactivate tumor suppressor genes, but there is no convincing evidence that HBV or HCV can bind to p53 or RB proteins. Also, the HBV-encoded regulatory element, called HBx, disrupts normal growth of infected hepatocytes. Neither HBV nor HCV infects immune cells.

PBD9 328, 337–338  BP9 203  BP8 213  PBD8 315, 327

55 A Some chronic hepatitis B and C viral infections progress to hepatocellular and/or cholangiolar carcinoma. a-Fetoprotein (AFP) is an oncofetal protein that is a tumor marker for hepatocellular carcinomas and some testicular carcinomas. AFP is normally present in fetal life but not in adults. A serum immunoglobin level with protein electrophoresis aids in the diagnosis of myeloma. Gastrointestinal tract adenocarcinomas, including those arising in the stomach, colon, and pancreas, as well as some lung carcinomas, may be accompanied by elevations in the serum carcinoembryonic antigen level. CA-19-9 is a tumor marker for colonic and pancreatic carcinomas. Some thyroid carcinomas produce calcitonin. Unfortunately, the sensitivity and specificity of tumor marker tests for detection of cancers, when they are small, is not high.

PBD9 328–329  BP9 203, 206, 211  PBD8 327  BP8 216, 221

56 A Cachexia is a common finding in advanced cancers, and weight loss without dieting in an adult is a “red flag” for malignancy. The exact cause for this is unknown, but increases in circulating factors such as tumor necrosis factor (TNF) may play a role. Cardiac murmurs may occur in the development of nonbacterial thrombotic endocarditis, a feature of a hypercoagulable state that may occur with advanced malignancies. Icterus is most likely to occur when there is obstruction of the biliary tract by a mass (e.g., as in pancreatic cancer), but metastases are unlikely to cause such an obstruction. Neurologic abnormalities may occur in local tumor growth impinging on nerves, but dull constant pain is the most likely abnormality in malignant neoplasms that invade nerves. Metastases to the spleen are uncommon. Tympany is uncommon in cancer because obstruction by a mass tends to be incomplete and to develop over a long time. (Hint: an empty beer keg is tympanitic when percussed.)

PBD9 330  BP9 208  PBD8 320–321  BP8 217–218

57 B The small cells have scant cytoplasm but marked hyperchromatism, consistent with small cell anaplastic carcinoma. This patient has Cushing syndrome resulting from ectopic corticotropin production by the tumor, a form of paraneoplastic syndrome common to small cell carcinomas of the lung. Such small cell carcinomas are of neuroendocrine derivation. A syndrome of inappropriate antidiuretic hormone (SIADH) secretion from small cell carcinomas is also common, but leads to hyponatremia as well as edema. Erythropoietin production with polycythemia is more likely to be associated with a renal cell carcinoma. Insulin and gastrin production are most often seen in islet cell tumors of the pancreas. Hypercalcemia from a parathyroid hormone-related peptide (PTHrP) is more typically associated with pulmonary squamous cell carcinomas.


58 E Histologic sections from malignant neoplasms are frequently assessed with a panel of immunostains to detect antigenic characteristics, such as protein expression, to aid in characterizing the cell of origin, as well as provide information in selection of treatment protocols. In this case, the immunostains are likely to reveal that this neoplasm is a high-grade urothelial carcinoma. A cytology smear shows light microscopic findings helpful to screen for malignancy, but the findings often fall short of diagnosing a specific cell type. The other listed techniques are best for determination of treatment and prognosis.

PBD9 333–335  BP9 210  PBD8 323–324  BP8 220–221

59 A The M1 designation indicates that distant metastases are present. N1 means local lymph nodes are positive for carcinoma. Elevated corticotropin levels indicate secretion of an ectopic hormone that may produce a paraneoplastic syndrome, but this is not part of staging. A T2 designation indicates that the overall size of the tumor is not large; it is still within the lung parenchyma and not impinging upon margins of the lung. The TNM system is used for staging, not microscopic grading of cellular differentiation.

PBD9 332  BP9 208–209  PBD8 322–323  BP8 218–220
The rapid frozen section of resection margins helps to determine whether enough of the colon has been resected. Electron microscopy requires at least 1 day to perform, and helps to determine the cell type, but it has largely been supplanted by immunohistochemistry. Fine-needle aspiration is used for preoperative diagnosis to find a malignancy. Flow cytometry can be performed in several hours, but it is useful mainly for prognostic information and is not a “stat” procedure. Radiologic imaging aids in preoperative diagnosis and assessment of possible metastases. Serum tumor markers may aid in preoperative diagnosis or postoperative follow-up of neoplasms.
1. A 45-year-old Bangladeshi woman with atrophic gastritis has sudden onset of severe, profuse, watery diarrhea. Over the next 3 days, she becomes severely dehydrated. On physical examination, she is afebrile, but has poor skin turgor. Laboratory studies of the diarrheal fluid show microscopic flecks of mucus, but no blood and few WBCs. A blood culture is negative. The woman is hospitalized and receives intravenous fluid therapy for 1 week. Which of the following is the most likely diagnosis?

A. Amebiasis  
B. Aspergillosis  
C. Cholera  
D. Filariasis  
E. Hydatid disease  
F. Typhoid fever

2. A 5-year-old boy has had diarrhea for a week, averaging six low volume stools per day, which appear mucoid and sometimes blood-tinted. On physical examination, his temperature is 37.4°C. He has mild lower abdominal tenderness, but no masses. A stool culture is positive for *Shigella sonnei*. Which of the following microscopic findings would most likely be seen in this child’s colon?

A. Epithelial disruption with overlying neutrophilic exudate  
B. Extensive scarring of lamina propria with stricture formation  
C. Intranuclear inclusions within small intestinal enterocytes  
D. Multiple granulomas throughout the colon wall  
E. Slight increase in lymphocytes and plasma cells in lamina propria

3. A 6-month-old infant has abrupt onset of vomiting followed by profuse, watery diarrhea. On physical examination, the infant has a temperature of 38.3°C. Development is normal for age, and the only abnormal finding is poor skin turgor. Laboratory studies show serum Na⁺ of 153 mmol/L, K⁺ of 4.4 mmol/L, Cl⁻ of 113 mmol/L, CO₂ of 28 mmol/L, and glucose of 70 mg/ dL. Examination of a stool specimen shows mucus, but no RBCs or WBCs. Which of the following mechanisms best accounts for this diarrhea?

A. Decreased absorption of sodium and water  
B. Decreased breakdown of lactose to glucose and galactose  
C. Increased secretion of potassium and water  
D. Mutation in CFTR gene  
E. Presence of Yop virulence plasmid

4. A 60-year-old man has had persistent bloody diarrhea, abdominal cramps, and fever for the past week. On physical examination, his temperature is 38.1°C. He has mild diffuse abdominal pain. A stool sample is positive for occult blood. Colonoscopy shows marked mucosal erythema with focal ulceration from the rectum to the ascending colon. The ulcers do not penetrate the muscularis propria. A biopsy is performed, and the microscopic appearance of the specimen is shown in the figure. Which of the following infectious organisms is most likely to produce these findings?

A. *Bacillus cereus*  
B. *Entamoeba histolytica*
5 In an epidemiologic study of individuals who died in a worldwide pandemic after World War I, many individuals were shown to have contracted a virulent form of influenza pneumonia. At the beginning of the 21st century, a similar epidemic is still possible from such a strain of influenza for which no vaccine may be readily available. Molecular analysis of samples of tissues showed changes in the virus responsible for these virulent forms of influenza. Which of the following changes most likely occurs in this virus to increase its virulence?

A Ability to elaborate exotoxins  
B Acquisition of antibiotic resistance genes  
C Increased binding to intercellular adhesion molecule-1 (ICAM-1) receptor  
D Mutations in DNA encoding envelope proteins  
E Recombination with RNA segments from animal viruses

6 An epidemiologic study is conducted with children ages 1 to 5 years who are infected with HIV-1. Most of these children have CD4+ lymphocyte counts above 500/mm³ and undetectable plasma HIV-1 RNA levels. On physical examination they have no abnormal findings. What is the most likely mode of transmission by which these children get infected?

A Breast-feeding  
B Inhalation of droplet nuclei  
C Fecal-oral contact  
D Transfer across placenta  
E Sexual abuse

7 In an experiment, phagocytosis of bacteria by neutrophils is studied. Bacteria are introduced into plasma containing neutrophils. It is observed that phagocytosis of *Streptococcus pneumoniae, Haemophilus influenzae*, and *Neisseria meningitidis* is reduced, compared with Enterobacteriaceae. Which of the following immune evasion mechanisms in these three bacterial species best explains this finding?

A Antimicrobial peptide binding  
B Carbohydrate capsule formation  
C Interferon homologue production  
D MHC protein down-regulation  
E Surface antigen switching

8 A 26-year-old man is an injection drug user and has developed fever over the past day. On examination, he has a heart murmur. A blood culture is positive for *Staphylococcus aureus* and antibiotic sensitivity testing shows resistance to methicillin due to presence of the mecA gene in these organisms. Through which of the following adaptations are these bacteria most likely to acquire their methicillin resistance?

A Biofilm formation  
B Exotoxin release  
C Pathogenicity island transfer  
D Superantigen stimulation  
E Surface adhesin expression

9 A study of nosocomial infections involving urinary catheters is performed. The study shows that the longer an indwelling urinary catheter remains, the higher the rate of symptomatic urinary tract infections (UTIs). Most of these infections are bacterial. Which of the following properties of these bacteria increase the risk for nosocomial UTIs?

A Biofilm formation  
B Enzyme elaboration  
C Exotoxin release  
D Quorum sensing  
E Superantigen stimulation

10 A 91-year-old woman is hospitalized with sepsis. On examination she has fever and hypotension. Laboratory studies show positive blood cultures. She has disseminated intravascular coagulopathy and pulmonary diffuse alveolar damage with respiratory distress. Analysis of the microbiology laboratory findings shows that the organisms cultured are gram-negative bacilli. Which of the following substances elaborated by these organisms is most likely to cause this complex of clinical findings?

A Endotoxin  
B Exotoxin  
C Mycolic acid  
D RNA polymerase  
E Superantigen  
F Tumor necrosis factor

11 A 6-year-old girl has a blotchy, reddish-brown rash on her face, trunk, and proximal extremities that developed over the course of 3 days. On physical examination, she has 0.2-cm to 0.5-cm ulcerated lesions on the oral cavity mucosa and generalized tender lymphadenopathy. A cough with minimal sputum production becomes progressively worse over the next 3 days. Which of the following viruses is most likely to produce these findings?

A Epstein-Barr  
B Mumps  
C Rubella  
D Rubeola  
E Varicella zoster

12 An 8-year-old girl has developed a mild febrile illness with a sore throat over the past 2 days. On physical examination, her temperature is 38.4°C, and she has a mild pharyngitis. The girl’s symptoms subside in 1 week without therapy. Over the next 2 months, she has increasing right-sided facial drooping with inability to close the right eye. Which of the following infectious organisms is most likely to produce these findings?

A *Cryptococcus neoformans*  
B *Cytomegalovirus*  
C *Listeria monocytogenes*  
D *Poliovirus*  
E *Toxoplasma gondii*
13. A 6-year-old girl who lives in the Yucatán peninsula has developed a high fever over the past 3 days. On physical examination, she has a temperature of 39.6°C and marked tenderness in all muscles. Laboratory studies show WBC count of 2950/mm³ with 12% segmented neutrophils, 4% bands, 66% lymphocytes, and 18% monocytes. Over 1 week, she becomes more lethargic, with a decreased level of consciousness, and petechiae and purpura develop over the skin. Further laboratory studies show thrombocytopenia with markedly prolonged prothrombin time and partial thromboplastin time. CT scan of the brain shows a hemorrhage in the right parietal lobe. Which of the following is most likely the vector of the agent causing infection in this patient?

A. Louse  
B. Mosquito  
C. Pig  
D. Snail  
E. Tick  
F. Tsetse fly

14. A 22-year-old woman has had recurrent vesicular lesions on her labia majora and perineum for 5 years. On physical examination, she is afebrile. Clusters of clear, 0.2-cm to 0.5-cm vesicles are present on the labia, with some surrounding erythema. The figure shows the representative microscopic appearance at low magnification of one of the lesions. Which of the following cellular changes is most likely to be seen under higher magnification of this lesion?

A. Dysplastic epithelial cells containing human papillomavirus sequences  
B. Mononuclear infiltrates containing protozoal organisms  
C. Multinucleated (syncytial) cells containing pink-to-purple intranuclear inclusions  
D. Neutrophils containing ingested gram-negative diplococci  
E. Perivascular lymphoplasmacytic infiltrate surrounding arterioles, with endothelial proliferation

15. A 6-year-old boy developed a rash over his chest that began as 0.5-cm reddish macules. Within 2 days, the macules became vesicles. Three days later, the vesicles ruptured and crusted over. Over the next 2 weeks, crops of the lesions spread to the face and extremities. Which of the following clinical manifestations of this infection is most likely to appear decades later?

A. Chronic arthritis  
B. Congestive heart failure  
C. Infertility  
D. Paralysis  
E. Shingles

16. A 31-year-old HIV-positive man has had increasing respiratory difficulty for the past 2 days. On physical examination, crackles are auscultated over all lung fields. A chest radiograph shows bilateral interstitial infiltrates. Laboratory studies show 26,800 copies of HIV-1 RNA/mL. A transbronchial biopsy is performed; the microscopic appearance of the specimen is shown in the figure. Which of the following is the most likely causative organism of his pulmonary disease?

A. Adenovirus  
B. Cytomegalovirus  
C. Epstein-Barr virus  
D. Herpes zoster virus  
E. Respiratory syncytial virus

17. A 14-year-old boy presents with fever, sore throat, and cervical lymphadenopathy. He then develops hepatomegaly and splenomegaly lasting for 2 months. His peripheral blood smear shows leukocytosis with “atypical” lymphocytes. Which of the following cell types is most likely to eliminate the virally infected cells?

A. Cytotoxic CD8 cells  
B. Epithelioid macrophages  
C. Helper CD4+ cells  
D. IgG-secreting plasma cells  
E. Polymorphonuclear neutrophils
18. A 75-year-old woman has a postoperative wound infection responding poorly to antibiotic therapy. Over the next 3 days she develops confusion, nausea, vomiting, diarrhea, chills, and myalgias. On examination, she is febrile; the wound site is erythematous with necrotic, purulent exudate. Laboratory studies show neutrophilia with left shift. Gram stain of the wound exudate shows gram-positive cocci in clusters. Which of the following substances is most likely being elaborated by the infectious organisms?
   A. Lactoferrin
   B. Lipopolysaccharide
   C. Phage-encoded A-B toxin
   D. Pili proteins
   E. Superantigen

19. A 46-year-old woman has had a high fever and swelling, warmth, and tenderness of the right leg for the past 3 days. On physical examination, she has a temperature of 39.4°C and the facial appearance shown in the figure. She receives macrolide antibiotic therapy with which she recovers. Infection with which of the following organisms has most likely produced these findings?
   A. Clostridium botulinum
   B. Escherichia coli
   C. Neisseria gonorrhoeae
   D. Staphylococcus epidermidis
   E. Streptococcus pyogenes

20. A 52-year-old man has a fever and cough productive of thick, gelatinous sputum that worsens over 4 days. On physical examination, his temperature is 38.2°C. On auscultation of the chest, diffuse cracksles are heard at the right lung base. Laboratory studies show WBC count, 13,240/mm³ with 71% segmented neutrophils, 8% bands, 15% lymphocytes, and 6% monocytes. A sputum gram stain shows gram-negative bacilli with mucoid capsules. His condition improves after a course of gentamicin therapy. Which of the following complications of this infection is he most likely to develop?
   A. Abscess formation
   B. Adenocarcinoma
   C. Bullous emphysema
   D. Cavitary granulomas
   E. Gas gangrene

21. A 20-year-old woman has had increasing delirium for 2 days and is admitted to the hospital. On physical examination, she has acute pharyngitis with an overlying dirty-white, tough mucosal membrane. Paresthesias with decreased vibratory sensation are present in the extremities. On auscultation, there is an irregular cardiac rhythm. A chest radiograph shows cardiomegaly. A Gram stain of the pharyngeal membrane shows numerous small, gram-positive rods within a fibrinopurulent exudate. Which of the following is the most likely mechanism for development of cardiac disease in this patient?
   A. Exotoxin-induced cell injury
   B. Granulomatous inflammation
   C. Lipopolysaccharide-mediated hypotension
   D. Microabscess formation
   E. Vasculitis with thrombosis

22. A 33-year-old primigravida at 18 weeks’ gestation develops nausea with vomiting for 3 days and then a severe headache and neck stiffness. On physical examination, her temperature is 38.2°C. She has no papilledema. A lumbar puncture is performed, and a Gram stain of the CSF obtained shows many short, gram-positive rods, and with a wet mount, the organisms demonstrate tumbling motility. In culture, this organism grows at 25°C. She most likely acquired this illness through which of the following mechanisms?
   A. Ingestion of contaminated food
   B. Inhalation of droplet nuclei
   C. Inoculation through a cut on the skin
   D. Sharing infected needles
   E. Using a friend’s toothbrush

23. When Pharaoh did not heed Moses and let the Hebrews go, a series of plagues fell on Egypt. In the fifth plague, large domesticated mammals, including cattle, horses, and sheep, died. This was followed by a plague in which the Egyptians developed cutaneous boils that probably appeared as 1-cm to 5-cm areas of erythema with central necrosis forming an eschar. Some Egyptians also may have developed a mild, non-productive cough associated with fatigue, myalgia, and low-grade fever over 72 hours, followed by a rapid onset of severe dyspnea with diaphoresis and cyanosis. Vital signs might have included temperature of 39.5°C, pulse of 105/min, respirations of 25/min, and blood pressure of 85/45 mm Hg. On auscultation of the chest, cracksles would be heard at the lung bases. A chest radiograph would show a widened mediastinum and small pleural effusions. “Legacy” laboratory findings would include a CBC with WBC count of 13,130/mm³, hemoglobin of 13.7 g/dL, hematocrit of 41.2%, MCV of 91 μm³, and platelet count of 244,000/mm³. Despite antibiotic therapy with anachronistic ciprofloxacin and doxycycline, many of those affected would die. Which of the following organisms is most likely to produce these findings?
   A. Bacillus anthracis
   B. Herpes simplex virus
   C. Mycobacterium leprae
   D. Staphylococcus aureus
   E. Variola major
   F. Yersinia pestis
24 A 42-year-old HIV-positive man has had a fever and cough for the past month. On physical examination, his temperature is 37.5°C. On auscultation of the chest, decreased breath sounds are heard over the right posterior lung. A chest radiograph shows a large area of consolidation with a central air-fluid level involving the right middle lobe. A transbronchial biopsy specimen contains gram-positive filamentous organisms that are weakly acid-fast. His course is complicated further by empyema and acute onset of a headache. A head CT scan shows a 4-cm discrete lesion of the right hemisphere with ring enhancement. Which of the following infectious agents is most likely causing his disease?

A) Aspergillus fumigatus  
B) Mucor circinelloides  
C) Mycobacterium avium complex  
D) Nocardia asteroides  
E) Staphylococcus aureus

25 A 50-year-old man with a neurodegenerative disease has had a fever and cough productive of yellow sputum for the past 3 days. On physical examination, there is dullness to percussion at the left lung base. A chest radiograph shows areas of consolidation in the left lower lobe. Despite antibiotic therapy, the course of the disease is complicated by abscess formation, and he dies. At autopsy, there is a bronchopleural fistula surrounded by a pronounced fibroelastic reaction. Small, yellow, 1-cm to 2-mm “sulfur granules” are grossly visible within the area of abscess formation. Which of the following organisms is most likely to produce these findings?

A) Actinomyces israelii  
B) Blastomyces dermatitidis  
C) Chlamydia pneumoniae  
D) Klebsiella pneumoniae  
E) Mycobacterium kansasii

26 A 22-year-old man has had more than 6 episodes of urethritis in the past 4 years since becoming sexually active. Each time, gram-negative coccii are identified in the neutrophilic exudate. Which of the following components in these organisms undergo change that prevents development of lasting protective immunity?

A) Chitin  
B) Envelope  
C) Lipopolysaccharide  
D) Peptidoglycan  
E) Pilli  
F) Teichoic acid

27 A 25-year-old woman has had pelvic pain, fever, and vaginal discharge for 3 weeks. On physical examination, she has lower abdominal adnexal tenderness and a painful, swollen left knee. Laboratory studies show WBC count of 11,875/mm³ with 68% segmented neutrophils, 8% bands, 18% lymphocytes, and 6% monocytes. She receives ceftriaxone therapy, but is not adherent with this therapy. She undergoes a work-up for infertility 5 years later. Which of the following infectious agents is most likely to produce these findings?

A) Candida albicans  
B) Gardnerella vaginalis  
C) Herpes simplex virus-2  
D) Neisseria gonorrhoeae  
E) Treponema pallidum  
F) Trichomonas vaginalis

28 A 4-year-old child develops a runny nose and cough. After the cough persists for 2 weeks she exhibits paroxysms of coughing so severe she becomes cyanotic. On physical examination, her temperature is 37.4°C. Her mouth and pharynx reveal no erythema or swelling. On auscultation of the chest, her lungs show crackles bilaterally. She has spasmodic coughing, with a series of coughs on a single breath, bringing up mucus plugs, followed by labored inspiration. The pathogenesis of her disease most likely results from disabling of which of the following following?

A) Ciliary movement  
B) Complement lysis  
C) Immunoglobulin secretion  
D) NK cell activation  
E) Phagolysosome formation

29 A 66-year-old man incurs extensive thermal burns to his skin and undergoes skin grafting procedures in the surgical intensive care unit. Two weeks later, he has increasing respiratory distress. Laboratory studies show hemoglobin, 13.1 g/dL; hematocrit, 39.2%; platelet count, 222,200/mm³; and WBC count, 4520/mm³ with 15% segmented neutrophils, 3% bands, 67% lymphocytes, and 15% monocytes. A chest radiograph shows extensive bilateral infiltrates with patchy areas of consolidation. Bronchoscopy is performed, and microscopic examination of a transbronchial biopsy specimen shows pulmonary vasculitis and surrounding areas of necrosis with sparse inflammatory exudate. Which of the following infectious agents is most likely to produce these findings?

A) Adenovirus  
B) Histoplasma capsulatum  
C) Mycobacterium tuberculosis  
D) Pseudomonas aeruginosa  
E) Pneumocystis jiroveci  
F) Streptococcus pneumoniae

30 In October 1347, a Genoese trading ship returning from the Black Sea docked at Messina, Sicily. The ship’s crew had been decimated by an illness marked by a short course of days from onset of inguinal lymph node enlargement with overlying skin ulceration to prostration and death. A small, ulcerated pustule ringed by a rosy rash was seen on the lower extremities of some of the crew. Within days, more than half of the population of the port city had died. Which of the following insect vectors was most likely responsible for the rapid spread of this disease?

A) Fleas  
B) Mosquitoes  
C) Reduviid bugs  
D) Sand flies  
E) Ticks

31 A study of sexually transmitted infections identifies an organism most commonly found in tropical and subtropical regions. This organism is associated with painful ulcerating genital papules in HIV-infected persons. Microscopic examination of lesional exudate with silver stain shows coccobacilli. Which of the following organisms is most likely to produce these findings?

A) Chlamydia trachomatis  
B) Haemophilus ducreyi  
C) Klebsiella granulomatis  
D) Neisseria gonorrhoeae  
E) Treponema pallidum
32. A 23-year-old man from Irian Jaya, Indonesia, has a lesion on his penis that has enlarged over the past 4 months. On physical examination there is a painless 2-cm papular lesion of the dorsum of his penis that evolves into a beefy red expansile ulceration that bleeds easily. No inguinal lymphadenopathy is present. A biopsy of the lesion is taken and examined microscopically, showing pseudoepitheliomatous hyperplasia and mixed inflammatory infiltrate. Giemsa stain shows coccobacilli within vacuoles in macrophages. Social history reveals multiple sexual partners. Which of the following is the most likely diagnosis?

A. Balanoposthitis
B. Chancroid
C. Granuloma inguinale
D. Lymphogranuloma venereum
E. Secondary syphilis

33. A 31-year-old man has had cough with a low-grade fever and a 4-kg weight loss over the course of 3 months. On physical examination, his temperature is 37.5°C. Laboratory studies show anemia of chronic disease. A bone marrow biopsy is performed, and the microscopic appearance is shown in the figure. An acid-fast stain of this tissue is positive. The causative infectious agent is most likely being destroyed by which of the following mechanisms?

A. Complement-mediated lysis
B. Elaboration of nitric oxide by macrophages
C. Generation of NADPH-dependent oxygen free radicals
D. Phagocytosis by eosinophils
E. Superoxide formation within phagolysosomes

34. A 5-year-old child is exposed to *Mycobacterium tuberculosis*. A month later the child’s tuberculin skin test is positive. The child then develops fever, inspiratory stridor, and nonproductive cough. Which of the following findings is most likely to be present on the chest radiograph of this child?

A. Hilar lymphadenopathy
B. Miliary pulmonary nodules
C. Pneumonic consolidation
D. Upper lobe cavitation
E. Vertebral lytic lesions

35. A 41-year-old man has had worsening fever, cough, and dyspnea for 2 weeks. On examination, he has rales and diminished breath sounds on auscultation of his chest. A chest radiograph shows scattered infiltrates in both lungs. A tuberculin skin test shows 6 mm of induration. A sputum sample is negative, but bronchoalveolar lavage is positive, for acid-fast bacilli. His WBC count is 4600/mm³ with differential count of 80% neutrophils, 10% lymphocytes, and 10% monocytes. Which of the following is the most likely risk factor for his pulmonary disease?

A. Alcohol abuse
B. Diabetes mellitus
C. HIV infection
D. Scurvy
E. Smoking

36. A 32-year-old man has maculopapular and nodular skin lesions, mainly involving his face, elbows, wrists, and knees. The nodular lesions have slowly enlarged over the past 10 years and are now beginning to cause deformity. The lesions are not painful, but he has hypoesthesia to anesthesia in these areas. The figure shows a microscopic acid-fast stain of a biopsy specimen of a nodular skin lesion. Which of the following is the most likely diagnosis?

A. Anthrax
B. Chagas disease
C. Hansen disease
D. Leishmaniasis
E. Lyme disease
F. Onchocerciasis

37. A 20-year-old man who has multiple sexual partners and does not use barrier precautions has had a nontender ulcer on his penis for the past week. On physical examination, the 0.6-cm lesion has a firm, erythematous base and sharply demarcated borders. The lesion is scraped, and microscopic darkfield examination is positive for motile spirochetes. Which of the following inflammatory processes is most likely to accompany this infection?

A. Acute inflammation with abscess formation
B. Granulomatous inflammation with caseation
C. Gummatous inflammation with necrosis
D. Perivascular inflammation with plasma cells
38 A longitudinal study of men and women who have developed aortic root dilation and aortic insufficiency in adulthood is performed. They have a history of unprotected sexual intercourse with multiple partners. Which of the following laboratory tests is most likely to yield a positive result in these persons?

A. Blood culture  
B. Darkfield microscopy of lymph node  
C. Fluorescent treponemal antibody-absorption (FTA)  
D. Rapid plasma reagin (RPR)  
E. Venereal disease research laboratory (VDRL)

39 An infant born at term to a 33-year-old woman is severely hydropic. On physical examination, there is a diffuse rash with sloughing skin on the palms and soles. Within 2 days, the infant dies of respiratory distress. At autopsy, there is marked hepatosplenomegaly. Microscopic examination of the femur and vertebrae shows periosteitis and osteochondritis. The lungs have nodular masses with central necrosis surrounded by mononuclear leukocytes, palisading macrophages, and fibroblasts. A serologic test result for which of the following agents is most likely to be positive in the infant’s mother?

A. Cytomegalovirus  
B. Herpes simplex type 2  
C. HIV  
D. Syphilis  
E. Toxoplasma gondii

40 A 44-year-old woman notices an erythematous papule on her left lower leg that develops into a ring-like rash and then subsides over 3 weeks. Over the next 5 months, she has migratory joint and muscle pain, substernal chest pain, and an irregular heart rhythm. These problems subside, but 2 years after the initial rash appeared, she develops a chronic arthritis involving the hips, knees, and shoulders. Which of the following is the most likely diagnosis?

A. Chagas disease  
B. Dengue fever  
C. Leishmaniasis  
D. Leprosy  
E. Lyme disease  
F. Syphilis

41 A radical group commissions scientists to develop a Category A bioterrorism agent. They want an agent that will paralyze victims within hours and be disguised within innocuous-appearing cans of split pea soup. Which of the following organisms best meets the requirements stated?

A. Chlamydia psittaci  
B. Clostridium botulinum  
C. Ebola virus  
D. Hantavirus  
E. Yersinia pestis

42 A 25-year-old soldier incurs multiple skin wounds that get infected and produce extensive tissue damage within a day. Culture of necrotic tissue from deep inside one of the wounds reveals anaerobic spore-forming gram-positive rods. Which of the following microscopic pathologic reactions are the toxins produced by these organisms most likely to cause?

A. Abscess formation  
B. Cytopathic effects with apoptosis  
C. Fibrous scarring  
D. Gangrenous necrosis  
E. Granulomatous inflammation  
F. Lymphocytic infiltrates

43 A 43-year-old man cuts the skin over his shin while repairing a fence on his farm. The wound heals without any complications. Four days later, he develops muscle spasms of the face and extremities. These spasms worsen to the point of severe contractions. Which of the following actions of the microbial toxins is most likely responsible for the clinical features in this case?

A. Cleavage of synaptobrevin in synaptic vesicles of neurons  
B. Degradation of muscle cell membranes by phospholipase C  
C. Inhibition of acetylcholine release at neuromuscular junctions  
D. Release of cytokine by T lymphocytes  
E. Stimulation of adenylate cyclase production in myofibers

44 A 27-year-old man is involved in a rollover accident in which he is ejected from the vehicle. He sustains a compound fracture of the right humerus and undergoes open reduction with internal fixation of the humeral fracture. Three days later, he has marked swelling of the right arm and palpable crepitus. A Gram stain of necrotic exudate from the wound site has the appearance shown in the figure. Through which of the following mechanisms is this organism most likely causing extensive tissue necrosis?

A. Elaboration of lipopolysaccharide  
B. Inhibition of phagocytic cell function  
C. Resistance to multiple antibiotics  
D. Superinfection with Candida albicans  
E. Toxin-mediated lecithin degradation
45. A 24-year-old man who is sexually active with multiple partners has had pain during urination for the past 4 days. On physical examination, there are no lesions on the penis. He is afebrile. Urinalysis shows no blood, ketones, protein, or glucose. Microscopic examination of the urine shows few WBCs and no casts or crystals. What infectious agent is most likely to produce these findings?

A. *Candida albicans*
B. *Chlamydia trachomatis*
C. Herpes simplex virus
D. *Mycobacterium tuberculosis*
E. *Treponema pallidum*

46. A 50-year-old woman residing in Port-au-Prince has observed a small vesicle on her right labium majus for the past 4 days. She is sexually active. On physical examination, the 0.5-cm vesicle is filled with purulent exudate. Tender inguinal lymph nodes are palpable. She was diagnosed with non-Hodgkin lymphoma 10 years ago. A biopsy of one of the lymph nodes is performed and microscopically shows multiple abscesses in which central necrosis is surrounded by palisading histiocytes. These clinical and pathologic findings are most likely caused by which of the following conditions?

A. *Candida albicans* vaginitis
B. *Chlamydia trachomatis* cervicitis
C. *Gardnerella vaginalis* vaginosis
D. Herpes simplex virus infection of the perineum
E. *Treponema pallidum* infection of the external genitalia

47. A 15-year-old boy has developed a small eschar on his left forearm around the site of a tick bite he received 6 days ago. A hemorrhagic rash involving the trunk, extremities, palms, and soles then develops over the next 3 days. Over the past day, small, 0.2-cm to 0.4-cm foci of skin necrosis have developed on his fingers and toes. His temperature is 39° C. He is treated with doxycycline and improves over the next 2 weeks. Which of the following organisms is most likely to produce these findings?

A. *Borrelia burgdorferi*
B. *Leishmania braziliensis*
C. *Mycobacterium leprae*
D. *Rickettsia rickettsii*
E. *Yersinia pestis*

48. A 10-year-old girl with leukemia undergoes hematopoietic stem cell transplantation. She has poor engraftment, and 1 month later she develops fever and dyspnea. On physical examination, her temperature is 39° C. On auscultation of the chest, wheezes and crackles are heard in both lungs. A chest CT scan shows nodular lesions accompanied by cavitation, hemorrhage, and infarction. Laboratory studies show hemoglobin, 8.8 g/dL; hematocrit, 26.5%; platelet count, 91,540/mm³; and WBC count, 1910/mm³ with 10% segmented neutrophils, 2% band neutrophils, 74% lymphocytes, and 14% monocytes. A bronchoalveolar lavage is performed; the fluid was stained with Gomori methenamine silver stain and analyzed microscopically, as shown in the figure. Which of the following infectious agents is most likely to produce these findings?

A. *Aspergillus fumigatus*
B. *Coccidioides immitis*
C. *Corynebacterium diphtheriae*
D. *Histoplasma capsulatum*
E. *Mycobacterium tuberculosis*
F. *Pneumocystis jiroveci*
49. A 24-year-old woman has noted worsening pain on the right side of her face for the past 24 hours. On examination, there is marked tenderness and swelling inferior to the zygomatic arch and lateral to the nasolabial fold on the right. Laboratory studies show WBC count, 9900/mm³; serum creatinine, 2 mg/dL; sodium, 151 mmol/L; potassium, 5.4 mmol/L; chloride, 119 mmol/L; bicarbonate, 8 mmol/L; and glucose, 483 mg/dL. A head CT scan shows soft tissue swelling and bony destruction around the right maxillary sinus. A biopsy is performed; the figure shows the findings on microscopic examination. Which of the following organisms is the most likely causative agent for this patient’s infection?

A. Aspergillus niger  
B. Actinomyces israelii  
C. Candida albicans  
D. Clostridium perfringens  
E. Cryptococcus neoformans  
F. Mucor circinelloides

50. An 11-year-old boy from Liberia has had episodic fevers for 2 weeks. He developed a severe headache a week ago and has become progressively more somnolent. On funduscopic examination, he has papilledema. The representative microscopic appearance of a cerebral vein is shown. Which of the following organs is most likely to serve as the reservoir for proliferation of the infectious agent producing this disease?

A. Brain  
B. Heart  
C. Liver  
D. Lymph nodes  
E. Spleen

51. A 19-year-old woman goes on a camping trip to a wooded area in New England (USA) with lots of insects, but has forgotten to bring insect repellant. A month later, she has increasing malaise, low-grade fever, headaches, and myalgias. On physical examination, she has hepatosplenomegaly. Laboratory studies show hemoglobin, 10.4 g/dL; WBC count, 5820/mm³; and platelet count, 205,000/mm³. Her peripheral blood smear is shown in the figure. Which of the following infectious agents is most likely to produce these findings?

A. Babesia microti  
B. Borrelia burgdorferi  
C. Giardia lamblia  
D. Rickettsia rickettsii  
E. Wuchereria bancrofti

52. A 45-year-old man experiences malaise and fatigue, which slowly become more noticeable over a 2-month period. He returned from a vacation along the Costa del Sol near Barcelona 10 months ago. He now has occasional diarrhea and a low-grade fever. His abdominal discomfort worsens over the next month. On physical examination, his vital signs include temperature of 38.3°C. He has pronounced splenomegaly, an increased liver span, and generalized lymphadenopathy. Laboratory studies show hemoglobin, 11.8 g/dL; hematocrit, 34.9%; platelet count, 89,000/mm³; and WBC count, 3350/mm³ with 29% segmented neutrophils, 5% bands, 48% lymphocytes, and 18% monocytes. His total serum protein is 7.6 g/dL, albumin is 3.2 g/dL, AST is 67 U/L, ALT is 51 U/L, alkaline phosphatase is 190 U/L, and total bilirubin is 1.3 mg/dL. A stool sample is negative for occult blood. Which of the following is the most likely diagnosis?

A. Borreliosis  
B. Echinococcosis  
C. Leishmaniasis  
D. Lyme disease  
E. Schistosomiasis  
F. Typhus
53. A 24-year-old soldier stationed in the Middle East has noted the appearance of a 0.5-cm papule on his left forearm. It becomes a 1-cm nodule with a central depression, and then ulcerates over the next month. On physical examination, the 2-cm ulcerated lesion has an indurated border, and there are three smaller satellite lesions. There is no hepatosplenomegaly, but he has left axillary lymphadenopathy. Laboratory studies show hemoglobin, 14.1 g/dL; hematocrit, 42.5%; platelet count, 233,200/mm³; and WBC count, 6270/mm³. Which of the following infectious organisms is most likely to produce these findings?

A. *Borrelia recurrentis*
B. *Brugia malayi*
C. *Leishmania major*
D. *Listeria monocytogenes*
E. *Mycobacterium leprae*
F. *Trypanosoma gambiense*

54. A 22-year-old man with extensive travel history is bitten by an insect and has developed a rubbery, red, 1-cm chancre on his right forearm over the past week. Three months later, he develops splenomegaly and lymphadenopathy. Two months later, he exhibits progressive wasting with cachexia and decreased mentation. His peripheral blood smear has the appearance shown in the figure. Where is his disease most likely to have been acquired?

A. Central America
B. Polynesia
C. Southeast Asia
D. Southern Europe
E. West Africa

55. A 9-year-old child who is living in a mud hut in Paraguay has a sore persisting on her face for 4 days. Physical examination shows an indurated area of erythema and swelling just lateral to the left eye, accompanied by posterior cervical lymphadenopathy. She has unilateral painless edema of the palpebrae and periorcular tissues. Two days later, she has malaise, fever, anorexia, and edema of the face and lower extremities. On physical examination 1 week later, there is hepatosplenomegaly and generalized lymphadenopathy. Which of the following pathologic findings is most likely to develop in this patient?

A. Cerebral abscesses
B. Chronic arthritis

56. A 28-year-old woman from rural Guyana with a history of rheumatoid arthritis develops painful swelling of her hands and feet. She is treated with corticosteroid therapy. A month later, she develops profuse, watery diarrhea along with fever and cough. On examination, she has a temperature of 37.3°C. Laboratory studies show WBC count, 12,900/mm³; and the WBC differential count shows 57% segmented neutrophils, 5% bands, 16% lymphocytes, 8% monocytes, and 14% eosinophils. Microscopic examination of a stool specimen shows ova and small rhabditoid larvae. Similar larvae are present in a sputum specimen. Which of the following infectious diseases is most likely to produce these findings?

A. Cysticercosis
B. Onchocerciasis
C. Schistosomiasis
D. Strongyloidiasis
E. Trichinosis

57. A 17-year-old boy has had generalized muscle pain with fever for 1 week. Over the past 2 days, he has developed increasing muscular weakness and diarrhea. On physical examination, his temperature is 38°C. All of his muscles are tender to palpation, but he has a normal range of motion, and no significant decrease in muscle strength. Laboratory findings include hemoglobin, 14.6 g/dL; hematocrit, 44.3%; MCV, 90 μm³; platelet count, 275,000/mm³; and WBC count, 16,700/mm³ with differential of 68% segmented neutrophils, 6% bands, 10% lymphocytes, 4% monocytes, and 12% eosinophils. What is the most likely diagnosis?

A. Hemorrhagic fever
B. Influenza
C. Poliomyelitis
D. Scrub typhus
E. Trichinosis

58. A 29-year-old man has had hematuria for the past month. On physical examination, he is afebrile. There is diffuse lower abdominal tenderness, but no palpable masses. An abdominal radiograph shows a small bladder outlined by a rim of calcification. Cystoscopy is performed, and the entire bladder mucosa is erythematous and granular. Biopsy samples are taken. Which of the following histologic findings is most likely to be seen in these samples?

A. Acid-fast bacilli of *Mycobacterium avium* complex
B. Eggs of *Schistosoma haematobium*
C. Larvae of *Trichinella spiralis*
D. Migrating *Ascaris lumbricoides*
E. *Taenia solium* cysts
In a study of individuals living in a subtropical region in which an irrigation project has been completed, it is noted that rice farmers have experienced an increased rate of an infectious illness since the project began. Investigators determine that the infection is acquired through cercariae that penetrate the skin. The cercariae are released from snails living in the irrigation canals. Infected individuals develop progressive ascites. Which of the following pathologic findings is most likely to be present in these infected individuals as a consequence of the infection?

A. Dilated cardiomyopathy  
B. Scrotal elephantiasis  
C. Hepatic fibrosis  
D. Mucocutaneous ulcers  
E. Urinary bladder carcinoma

A 40-year-old man has had progressive enlargement of the right leg for the past 6 years, leading to the appearance shown in the figure. On physical examination, he is afebrile. He has inguinal lymphadenopathy and scrotal edema. Infection with which of the following organisms is most likely to be present in this man?

A. *Echinococcus granulosus*  
B. *Leishmania tropica*  
C. *Schistosoma mansoni*  
D. *Trichinella spiralis*  
E. *Wuchereria bancrofti*

Persons living in southern Africa where black flies are common and who have developed blindness are studied to identify a potential infectious cause. These persons are found to have a chronic dermatitis that preceded their blindness. Skin lesions are pruritic, scaling, and hypopigmented. Ocular lesions include punctate keratitis and focal corneal opacities, sclerosing keratitis, iridocyclitis with glaucoma, and retinitis. Which of the following morphologic forms of the infectious agent is most likely to be found in skin biopsies of these persons?

A. Acid-fast bacilli  
B. Elementary bodies  
C. Intracellular diplococci  
D. Intranuclear inclusions  
E. Microfilariae

Within the same day, an emergency department is visited by 20 individuals, all of whom work in the same building. Over the past day, they all experienced the sudden onset of high fever, headache, backache, and malaise. On examination, they are febrile. They do not have lymphadenopathy or hepatosplenomegaly. Over the next 2 days, they develop a maculopapular rash on the face, forearms, and mucous membranes of the oropharynx. Despite supportive care, a third of these patients die. Which of the following organisms is the most likely causative agent?

A. *Chlamydia psittaci*  
B. *Francisella tularensis*  
C. Hantavirus  
D. *Mycobacterium kansasii*  
E. *Rickettsia typhi*  
F. *Variola major*
Yop plasmid confers infectivity to *Yersinia* secretory diarrhea with sodium chloride and water loss. The organism, which potentiates *Vibrio cholerae* giving rise to meconium ileus in infants. Decreased breaks in the CFTR gene lead to formation of thick mucus plugs, also produce diarrhea by inhibiting the intestinal absorption of sodium chloride and water loss. Most rotavirus infections are subclinical or cause mild gastrointestinal infections. The development of IgA antibodies from secretory immunity in the bowel to rotavirus surface antigens provides older children and adults a relative resistance to rotavirus infection. Such antibodies are present in maternal milk and confer some degree of resistance to infants who breast-feed. Rotavirus infection occurs worldwide. By the age of 3 years, virtually every individual has been infected by rotaviruses at least once. Most rotavirus infections are subclinical or cause mild gastrointestinal illnesses that do not require hospitalization. The first infection is the most likely to be symptomatic; subsequent infections are often mild or asymptomatic. Many enteroviruses also produce diarrhea by inhibiting the intestinal absorption of intraluminal sodium and water, but not as severe. Mutations in the CFTR gene lead to formation of thick mucus plugs, giving rise to meconium ileus in infants. Decreased breakdown of lactose occurs in disaccharidase deficiency and gives rise to an osmotic diarrhea. Cholera is the result of secretion of an exotoxin by the *Vibrio cholerae* organism, which potentiates the epithelial cell production of adenylate cyclase and causes secretory diarrhea with sodium chloride and water loss. The Yop plasmid confers infectivity to *Vibriostia* organisms.

Amebiasis is a common cause of dysentery in developing nations. The figure shows two single-cell protozoa invading tissue with inflammatory cells. *Entamoeba histolytica* organisms are resistant to gastric acid and can invade the colonic submucosa via contact-dependent cytolyis. The amebae not only produce local necrosis with ulceration and hemorrhage, but also gain access to the venules of the portal system, which drains to the liver. Amebic liver abscess is an uncommon complication of amebiasis. The colonic lesions typically have disappeared by the time the liver lesions appear. In some cases, there can be extensive mucosal involvement with characteristic flask-shaped (similar to an Erlenmeyer flask) ulcerations similar to those seen in other severe inflammatory bowel diseases. *Bacillus cereus* is a cause of food poisoning (most often as a contaminant in reheated fried rice) and has a short incubation time. Giardiasis tends to involve the small intestine and produces variable inflammation, but no ulceration. Salmonellosis more typically involves the small intestine and in most cases produces self-limiting enteritis, although more severe disease with dissemination to other organs can occur with *Salmonella typhi* infection. Shigellosis can produce bloody dysentery with irregular superficial colonic mucosal ulceration, but the organisms typically do not invade beyond the lamina propria. Cholera is characterized by massive, secretory diarrhea without intestinal mucosal invasion or necrosis.

The lack of stomach acid in this woman predisposes to enteric infections. *Vibrio cholerae* organisms are noninvasive. Instead, they produce severe diarrhea by elaboration of an enterotoxin, called cholera toxin, that acts on bowel mucosal cells to cause persistent activation of adenylate cyclase and high levels of intracellular cyclic AMP that drives massive secretion of sodium, chloride, and water. The fluid loss is life-threatening because of resultant dehydration. Amebiasis tends to produce dysentery, with a bloody diarrhea, because the organisms can invade the mucosa. Aspergillosis is seen in immunocompromised patients, particularly patients with neutropenia, and is a rare cause of a diarrheal illness. Filariasis involves the lymphatics and produces elephantiasis. Hydatid disease caused by *Echinococcus* produces space-occupying cystic lesions in viscera. Typhoid fever produces diarrhea, and the organisms can invade mucosa and disseminate to produce many systemic symptoms.

**ANSWERS**

1. **C** The lack of stomach acid in this woman predisposes to enteric infections. *Vibrio cholerae* organisms are noninvasive. Instead, they produce severe diarrhea by elaboration of an enterotoxin, called cholera toxin, that acts on bowel mucosal cells to cause persistent activation of adenylate cyclase and high levels of intracellular cyclic AMP that drives massive secretion of sodium, chloride, and water. The fluid loss is life-threatening because of resultant dehydration. Amebiasis tends to produce dysentery, with a bloody diarrhea, because the organisms can invade the mucosa. Aspergillosis is seen in immunocompromised patients, particularly patients with neutropenia, and is a rare cause of a diarrheal illness. Filariasis involves the lymphatics and produces elephantiasis. Hydatid disease caused by *Echinococcus* produces space-occupying cystic lesions in viscera. Typhoid fever produces diarrhea, and the organisms can invade mucosa and disseminate to produce many systemic symptoms.

2. **A** Shigellosis results in bloody dysentery because *Shigella* is highly virulent, resistant to gastric acid, and can invade and destroy the colonic mucosa. There is typically a mononuclear infiltrate extending to the lamina propria, with a neutrophilic exudate overlying the ulcerated areas. Stricture formation may follow intestinal tuberculosis. Intracellular inclusions in enterocytes point to infection with DNA viruses, such as herpesviruses like cytomegalovirus. Granulomatous inflammation may be seen with granulomatous colitis (Crohn disease) and intestinal tuberculosis (rare). An increase in mononuclear inflammatory cells may be seen with milder forms of enterocolitis caused by viruses, *Giardia*, and *Salmonella* spp.

3. **A** Rotavirus, an encapsulated RNA virus, is a major cause of diarrhea in infancy. The small intestinal villous destruction with atrophy leads to decreased absorption of sodium and water. The development of IgA antibodies from secretory immunity in the bowel to rotavirus surface antigens provides older children and adults a relative resistance to rotavirus infection. Such antibodies are present in maternal milk and confer some degree of resistance to infants who breast-feed. Rotavirus infection occurs worldwide. By the age of 3 years, virtually every individual has been infected by rotaviruses at least once. Most rotavirus infections are subclinical or cause mild gastrointestinal illnesses that do not require hospitalization. The first infection is the most likely to be symptomatic; subsequent infections are often mild or asymptomatic. Many enteroviruses also produce diarrhea by inhibiting the intestinal absorption of intraluminal sodium and water, but not as severe. Mutations in the CFTR gene lead to formation of thick mucus plugs, giving rise to meconium ileus in infants. Decreased breakdown of lactose occurs in disaccharidase deficiency and gives rise to an osmotic diarrhea. Cholera is the result of secretion of an exotoxin by the *Vibrio cholerae* organism, which potentiates the epithelial cell production of adenylate cyclase and causes secretory diarrhea with sodium chloride and water loss. The Yop plasmid confers infectivity to *Vibriostia* organisms.

4. **B** Amebiasis is a common cause of dysentery in developing nations. The figure shows two single-cell protozoa invading tissue with inflammatory cells. *Entamoeba histolytica* organisms are resistant to gastric acid and can invade the colonic submucosa via contact-dependent cytolyis. The amebae not only produce local necrosis with ulceration and hemorrhage, but also gain access to the venules of the portal system, which drains to the liver. Amebic liver abscess is an uncommon complication of amebiasis. The colonic lesions typically have disappeared by the time the liver lesions appear. In some cases, there can be extensive mucosal involvement with characteristic flask-shaped (similar to an Erlenmeyer flask) ulcerations similar to those seen in other severe inflammatory bowel diseases. *Bacillus cereus* is a cause of food poisoning (most often as a contaminant in reheated fried rice) and has a short incubation time. Giardiasis tends to involve the small intestine and produces variable inflammation, but no ulceration. Salmonellosis more typically involves the small intestine and in most cases produces self-limiting enteritis, although more severe disease with dissemination to other organs can occur with *Salmonella typhi* infection. Shigellosis can produce bloody dysentery with irregular superficial colonic mucosal ulceration, but the organisms typically do not invade beyond the lamina propria. Cholera is characterized by massive, secretory diarrhea without intestinal mucosal invasion or necrosis.
Bacteria that produce capsules are more resistant to phagocytosis, and help them to avoid an initial innate immune response with neutrophils. This enables them to establish infection (respiratory tract, and possible spread to meninges) and become more virulent. Antimicrobial peptides produced by epithelial cells can bind to bacterial organisms and form pores in the cell walls to kill them by osmotic lysis. Interferons are produced against viral organisms, not bacteria. Some viruses, such as herpesviruses, can impair expression of MHC class I molecules so that viral antigens are not effectively displayed to CD4+ and CD8+ cells. Surface antigen switching also helps organisms such as trypanosomes to evade an adaptive immune response.

Organisms such as Staphylococcus aureus can express superantigens that nonselectively stimulate many T cell clones, leading to unregulated cytokine release and toxic shock. Adhesins aid in microbial binding to host cells.

Bacteria have multiple mechanisms for exchanging genetic material that affords selective growth advantages. Pathogenicity islands are bacterial chromosomal elements carrying virulence genes, such as those involving antibiotic resistance. Presence of the mecA gene imparts resistance to methicillin (methicillin-resistant Staphylococcus aureus, or MRSA) and other β-lactam antibiotics. Additional bacterial genetic transfer mechanisms include plasmids, transposons, and integrons. Microbes on tissue surfaces form biofilms of sticky polysaccharide goo to isolate themselves from immune attack. Exotoxins impart virulence through tissue damage. Organisms such as S. aureus can express superantigens that nonselectively stimulate many T cell clones, leading to unregulated cytokine release and toxic shock. Adhesins aid in microbial binding to host cells.

Microbes form biofilms of sticky polysaccharide goo that adheres particularly well to artificial surfaces such as catheters. The biofilm helps isolate the organisms from inflammatory cells and limit penetration of antibiotics. Bacterial enzymes, such as the hyaluronidases and streptokinases of streptococcal organisms, promote spread through tissues. Exotoxins of gram-positive organisms impart virulence through tissue damage. Bacteria have multiple mechanisms for exchanging genetic material that afford selective growth advantages. When the number of bacterial organisms increases, they sense this (quorum sensing) and turn on virulence genes. Organisms such as Staphylococcus aureus can express superantigens that nonselectively stimulate many T cell clones, leading to unregulated cytokine release and toxic shock.

Gram-negative sepsis is classically mediated by endotoxins, particularly the lipopolysaccharide component of the outer cell wall. With sepsis from gram-positive organisms there is release of exotoxins, such as tetanospasmin released by Clostridium tetani organisms. Mycolic acids found in the lipid wall of mycobacteria aids in the resistance of these organisms to degradation by acute inflammatory responses, leading to granulomatous inflammation. RNA polymerase is found in negative-sense RNA viruses and produces a positive-sense messenger RNA (mRNA) that directs the host cell to produce viral components. Superantigens may produce findings similar to lipopolysaccharide-induced septic shock; the best known is toxic shock syndrome toxin, which is elaborated by some staphylococcal organisms. Tumor necrosis factor (TNF) is elaborated by human inflammatory cells, not by microorganisms, but by the release of TNF by the action of endotoxins on macrophages that can mimic gram-negative sepsis.

Poliomyelitis is an entrovirus spread through fecal-oral contamination. The virus often infects the oropharynx first. It then spreads to bulbax nuclei and/or lower motor neurons in the anterior horn of the spinal cord to produce the muscular paralysis typical of polio. In places where vaccination is routinely available, this disease is rare. Cryptococcosis is a fungal disease that most often involves the lungs and meninges. Cytomegalovirus infection can be congenital; in immunocompromised adults, it can involve many organs, principally the gastrointestinal tract, brain, and lungs. Listeriosis is most often acquired via contaminated food or water; in most adults, it produces mild diarrheal illness, but in some adults and children, and in fetuses, it can produce meningitis or dissemination with microabscess (microgranuloma) formation. Toxoplasmosis can be a congenital infection. In immunocompromised adults, it can produce inflammation in multiple tissues, but most often, it causes chronic abscessing inflammation in brain.

Dengue fever, one form of hemorrhagic fever, is caused by an arbovirus of the Flavivirus group. This organism can be devastating because it produces bone marrow suppression, and because any antibodies to the virus enhance cellular viral uptake. It is transmitted by the mosquito vector Aedes aegypti. Louse-borne infections include rickettsial diseases. The pig can be involved in the life cycle of Taenia solium and of Trichinella spiralis. T. spiralis can produce marked muscle pain, but typically not disseminated intravascular coagulopathy. Some snails can serve as an intermediate host for Schistosoma organisms. Ticks can transmit typhus and Lyme disease. The tsetse fly can transmit sleeping sickness, which is endemic to Africa.
14 C The figure shows a vesicle that has resulted from herpes simplex virus (HSV) infection. Most genital infections are caused by HSV-2, whereas HSV-1 is responsible for most cases of herpetic gingivostomatitis. The viral cytopathic effect results in formation of intranuclear inclusions, multinucleated cells, and cell lysis with vesicle formation in the epithelium. Cervical dysplasias do not produce vesicular lesions and are the result of another sexually transmitted disease—human papillomavirus infection. Protozoal infection with trichomoniasis, typically involving the vagina, may produce small blisters or papules, but these are often self-limited and not typically recurrent. Gram-negative diplococci are characteristic of Neisseria gonorrhoeae infection, also a sexually transmitted disease. Lymphoplasmacytic infiltrates may be seen in chancrels caused by Treponema pallidum, the causative agent of syphilis.

PBD9 357–358  BP9 310, 552, 681  PBD8 352  BP8 322, 708–709

15 E The skin lesions are typical of chickenpox, a common childhood infection caused by varicella-zoster virus infection. The infection can remain dormant for years in dorsal root ganglia, only to reactivate when immune status is diminished. The virus, now designated herpes zoster (or varicella-zoster), spreads from the ganglion to the skin in the dermalomotorial distribution of the corresponding sensory nerve, and it causes vesicular lesions with chronic, burning pain that is difficult to stop. A chronic arthritis can be seen with Lyme disease after Borrelia burgdorferi infection. Rheumatic heart disease can appear after group A β-hemolytic streptococcal infection. Infertility is a complication of mumps orchitis. Paralysis can complicate poliovirus infection.

PBD9 358  BP9 310  PBD8 353  BP8 322–323, 877

16 B This patient has a high HIV-1 RNA level consistent with the diagnosis of AIDS. Although patients with AIDS are susceptible to many microbes, infections with cytomegalovirus are particularly common. The biopsy specimen shows an enlarged cell containing a large, distinct intranuclear inclusion and ill-defined dark cytoplasmic inclusions, which are typical of cytomegalovirus infection. Adenovirus is a viral pathogen in both immunocompromised and immunocompetent adults that may produce a clinically significant pneumonia, and intranuclear inclusions may be present, but the cells are not large, and cytoplasmic inclusions are absent. Epstein-Barr virus infection is seen frequently in patients with HIV infection, but there are no distinct pulmonary lesions associated with it. Herpes zoster infections are most likely to affect the peripheral nervous system, rarely can become disseminated to affect the lungs in immunosuppressed patients, and produce a different appearance than that shown. Respiratory syncytial virus infections are seen in children, but rarely in adults.


17 A The features described fit with infectious mononucleosis. EBV infection involves B cells that are activated to elaborate a variety of cytokines that promote viral proliferation and reduced immune response. IL-10 normally secreted by phagocytes activates T cells (the “atypical” lymphocytes), but the virally induced homologue does not. In general, viral infections are intracellular, and a cytotoxic CD8 T cell response is required to clear virus by eliminating infected cells. Epithelioid macrophages are most important in granulomatous inflammatory responses that control mycobacterial and fungal infections. Helper T cells may be infected by EBV, but do not clear the virus. Immunoglobulin responses are most important as an innate immune response directed against extracellular organisms such as bacteria.


18 E Staphylococcal toxic shock syndrome (TSS) results from elaboration of superantigens that stimulate up to 20% of T lymphocytes and generate a marked release of cytokines and an extensive inflammatory response. Increasingly, staphylococci have acquired the meca gene that imparts resistance to many penicillin (methicillin) and cephalosporin antibiotics has come to be associated with methicillin-resistant Staphylococcus aureus (MRSA). Most TSS cases occur in women, because of the relationship to vaginitis. Lactoferrin is a substance secreted by human cells that binds iron needed by bacteria, and is thus part of innate immunity. Lipopolysaccharides are elaborated by gram-negative organisms and produce endotoxin shock. Phage-encoded A-B toxin is elaborated by Corynebacterium diphtheriae. Pili proteins are characteristic for Neisseria gonorrhoeae to provide attachment to target cells in the genital tract.

PBD9 362–363  BP9 321  PBD8 344, 357  BP8 334

19 E The rash and edema are manifestations of streptococcal erysipelas, which is usually caused by group A or group C streptococci. Streptolysins elaborated by these organisms aid in the spread of the infection through subcutaneous tissues. Over 900 years ago the Order of St. Anthony was founded to treat persons with this illness, then known as St. Anthony’s fire. Clostridium botulinum elaborates an exotoxin that, when ingested, results in paralysis. Escherichia coli produces various infections, but skin infections are uncommon. Neisseria gonorrhoeae is best known as a sexually transmitted disease, and a rash is possible, although usually there is no pronounced swelling. Staphylococcus epidermidis is usually considered a contaminant in cultures.

PBD9 364–365  PBD8 359–360

20 A Bacterial infections with predominantly neutrophilic response are marked by suppurative inflammation, and a virulent organism such as Klebsiella can lead to tissue destruction with abscess formation. Carcinomas are not sequelae of bacterial infections. Infections of the lung do not result in emphysema, but may complicate emphysema. Granulomatous inflammation is characteristic of mycobacterial or fungal infections. Gas-forming bacteria, such as anaerobic Clostridium organisms, are unusual as a cause of respiratory infections.

PBD9 352  BP9 323–324  PBD8 347  BP8 334
21 A This woman has diphtheria. The Corynebacterium diphtheriae organisms proliferate in the inflammatory membrane that covers the pharynx and tonsils. These gram-positive organisms elaborate an exotoxin that circulates and produces myocarditis and neuropathy. The organisms do not disseminate to cause inflammation, abscesses or vasculitis elsewhere in the body. Granulomatous inflammation is more typical of mycobacterial and fungal infections. Endotoxins such as lipopolysaccharide tend to be elaborated by gram-negative bacterial organisms.

PBD9 365 BP9 321 PBD8 360–361 BP8 333

22 A The results of the Gram stain and culture are diagnostic for Listeria monocytogenes, an organism that is more likely to produce disseminated disease in individuals who are immunocompromised or pregnant, and it can produce a congenital infection. Since the organism grows readily at room temperature, it easily contaminates food and water. Unpasteurized dairy products are most often implicated. Listeriosis is not known to be acquired parenterally, or by the other listed routes.

PBD9 366 BP9 320 PBD8 361 BP8 256, 333, 874

23 A The features are those of cutaneous and respiratory anthrax. Bacillus anthracis forms spores that resist environmental degradation. The spores can be transmitted by aerosols, making this organism an ideal terror weapon. Similar to many gram-positive organisms, B. anthracis produces disease via elaboration of exotoxins that have an active A subunit and a binding B subunit. None of the other choices involve outbreaks in domestic animals. Herpetic infections form clear vesicles that can rupture to shallow ulcers. Mycobacterium leprae can produce a faint rash early in its course, but involvement of peripheral nerves with loss of sensation predisposes to repeated trauma with deformity. Staphylococcus aureus can produce impetigo, typically on the face and hands. Variola major is the agent for smallpox, which is characterized by skin pustules, and pneumonia is the most likely cause of death. Yersinia pestis produces plague, which can have bubonic and pneumonic forms, characterized by ulcerating lymph nodes surrounded by a rosy rash.

PBD9 366–367 BP9 321 PBD8 361–362 BP8 321

24 D Although nocardiosis typically begins in the lungs, it often becomes disseminated, particularly to the central nervous system. These infections are most often seen in immunocompromised patients. Aspergillosis also can affect immunocompromised individuals, particularly those with neutropenia, but the fungal hyphae are easily distinguishable on hematoxylin and eosin stains. Mucor organisms have broad, nonseptate hyphae and are seen most often in patients with diabetic ketoacidosis or burn injuries. Mycobacterium avium complex infections are seen in individuals with AIDS, but these are short, acid-fast rods that produce poorly formed granulomas. Bacterial pneumonias also should be considered in immunocompromised patients, and septicemia can complicate them, but Staphylococcus aureus organisms form clusters of gram-positive cocci.

PBD9 367 BP9 312 PBD8 362–363 BP8 324, 311

25 A Actinomyces that can produce chronic abscessing pneumonia, particularly in immunocompromised patients, include Actinomyces israelii and Nocardia asteroides. Persons with neurodegenerative diseases are at risk for aspiration of oropharyngeal secretions that may contain these organisms. Sulfur granules, formed from masses of the branching, filamentous organisms, are more likely to be seen in Actinomyces. Blastomyces dermatitidis infections tend to produce a granulomatous inflammatory process. Chlamydial infections produce an interstitial pattern similar to that of most viruses. Klebsiella infections, similar to other bacterial infections, can result in abscess formation, although without distinct sulfur granules. Mycobacterium kansasi infections are similar to Mycobacterium tuberculosis infections in that granulomatous inflammation is prominent.

PBD9 367 BP9 312 PBD8 362–363 BP8 324, 311

26 E Pili are cell wall structures in gram-negative bacteria, such as the Neisseria gonorrhoeae in this case, that facilitate attachment to host cells. Pili proteins are altered by genetic recombination, forming a “moving target” for host immunity, so reinfection can occur. Chitin is a prominent cell wall component of fungi. Envelopes aid attachment of viruses to their target host cells. Lipopolysaccharide in gram-negative bacterial cell walls acts as an endotoxin. Peptidoglycan forms part of the bacterial cell wall, and a greater amount of it imparts gram-positive staining. Teichoic acid is a prominent feature of gram-positive bacterial cell walls.

PBD9 368 BP9 317, 320 PBD8 343, 364 BP8 325, 332

27 D This patient has pelvic inflammatory disease (PID), which may occur as a result of infection with Neisseria gonorrhoeae or Chlamydia trachomatis. Both organisms cause sexually transmitted diseases, and chronic inflammation may lead to PID. Complications of PID include peritonitis, adhesions with bowel obstruction, and sepsis with endocarditis, meningitis, arthritis, and infertility. Of the remaining organisms listed, Candida can produce vaginitis with a curd-like discharge, but it does not typically produce PID. Gardnerella produces a whitish discharge that has a “fishy” odor with bacterial vaginosis, which tends to remain localized. Herpes simplex virus-2 (HSV-2), the most common agent of genital herpes, can produce painful vesicles, usually on the external genitalia, and is often recurrent. Treponema pallidum, the causative agent of syphilis, produces a hard chancre on skin and mucosal surfaces. Trichomoniasis may also lead to infertility, but this protozoan is not treated with cephalosporins, and it generally does not produce disseminated disease.

PBD9 368, 383 BP9 317, 695 PBD8 363–364 BP8 324–326, 727–728

28 A Bordetella pertussis is the causative agent for whooping cough. These infections occur infrequently when there is widespread childhood vaccination against this organism. This cocobacillary organism is difficult to culture, and direct fluorescent antibody (DFA) testing is the fastest and most reliable way to diagnose the infection. Nasopharyngeal aspirates and swabs are the best specimens because the organisms attach to ciliated respiratory epithelium. The toxin paralyses cilia. Complement lysis is most useful against circulating infectious agents. Immunoglobulins that circulate can bind organisms, but secretion is an adaptive immune response taking days to
weeks. NK cells attack host cells with MHC signaling turned off by intracellular infectious agents such as viruses. Mycobacterial organisms inhibit phagolysosome formation to reduce their intracellular destruction in macrophages.

29 D *Pseudomonas aeruginosa* can infect the skin following burn injuries and spread to the lungs. These organisms secrete several virulence factors, as follows: exotoxin A, which inhibits protein synthesis; exoenzyme S, which interferes with host cell growth; phospholipase C, which degrades pulmonary surfactant; and iron-containing compounds, which are toxic to endothelial cells. These virulence factors result in extensive vasculitis with necrosis. Neutrophic patients are particularly at risk. *Histoplasma capsulatum* yeasts can produce pulmonary disease resembling that of *Mycobacterium tuberculosis*, with granulomatous inflammation. Pneumocystis pneumonia is more likely to occur in patients with weak cell-mediated immunity. Pneumococcal infections produce alveolar exudates without significant vascular involvement.

30 A This incident marks the first appearance of the Black Death in Europe, a disease that persisted during the 14th and 15th centuries. The plague spread through Italy and across the European continent. By the following spring, it had reached as far north as England, and within 5 years, it had killed 25 million people, one third of the European population. Rodents form the reservoir of infection (and cats weren’t as popular as lion people, one third of the European population. Rodents were the key cellular component within granulomas, and they cannot destroy mycobacteria. NADPH-dependent reactive oxygen species are important in the lysis of bacteria by neutrophils. *M. tuberculosis* organisms reside in phagosomes, which are not acidified into phagolysosomes.

31 B The causative organism of chancroid is *Haemophilus ducreyi*, which is difficult to grow in culture and often obscured by superinfecting organisms in the ulcerated lesions. Chancroid is most common in Africa and Southeast Asia, is a co-factor in transmission of HIV, and its features overlap those of granuloma inguinale, but there is often lymph node involvement in the former, and lack of the Donovan bodies in macrophages. Lymphogranuloma venereum is caused by *Chlamydia trachomatis*, which cannot be seen with Gram stain. *Klebsiella granulomatis* (formerly *Calymmatobacterium granulomatis*) is also common in tropical and subtropical regions. It causes granuloma inguinale, and the infection may progress to scarring with urethral and lymphatic obstruction. Gonorrhea tends to produce a urethritis in men and a cervicitis in women acutely, without genital ulceration. Syphilis is marked by a chancre in the primary state and a maculopapular rash of palms and soles in the secondary stage, with a prominent lymphoplasmacytic infiltrate; the causative agent is *Treponema pallidum*, and these spirochetes cannot be identified by Gram stain.

32 C The causative organism is *Klebsiella granulomatis* (formerly *Calymmatobacterium granulomatis*), the disease is most common in tropical and subtropical regions, and the infection may progress to scarring with urethral and lymphatic obstruction. Balanoposthitis is localized inflammation of the glans penis and prepuce, typically caused by *Candida, Gardnerella*, or *Staphylococcus* spp. Chancroid caused by *Haemophilus ducreyi* has features that overlap those of granuloma inguinale, but there is often lymph node involvement in the former, and lack of the Donovan bodies in macrophages. Lymphogranuloma venereum is caused by *Chlamydia trachomatis*, which cannot be seen with Gram stain. Secondary syphilis is marked by a maculopapular rash on the palms and soles, with a prominent lymphoplasmacytic infiltrate; the causative agent is *Treponema pallidum*, and these spirochetes cannot be identified by Gram stain.

33 B The figure shows a granuloma. Activated macrophages are the key cellular component within granulomas that form to control persistent organisms such as *Mycobacterium tuberculosis*. As part of delayed type hypersensitivity with a Th1 immune response, CD4+ cells secrete interferon-γ, which activates macrophages to kill organisms with reactive nitrogen intermediates. Complement-mediated lysis is not involved in the destruction of intracellular bacteria such as *M. tuberculosis*. Complement activation on the surface of *M. tuberculosis* can opsonize the bacteria, however, for uptake by macrophages. Eosinophils are not a major component of most granulomas, and they cannot destroy mycobacteria. NADPH-dependent reactive oxygen species are important in the lysis of bacteria by neutrophils. *M. tuberculosis* organisms reside in phagosomes, which are not acidified into phagolysosomes.

34 A The child has primary tuberculosis. Most healthy persons have subclinical disease, and a minority develop clinical manifestations; of those, most have limited pulmonary involvement without dissemination. Primary tuberculosis is marked by the Ghon complex, which is a small subpleural granuloma at mid-lung along with prominent enlarged hilar lymph nodes. These nodes may impinge upon central airways. When the cell-mediated immune response is poor, then there can be numerous small granulomas scattered throughout the lungs, or disseminated to other organs, as a miliary pattern (granulomas that are the size of millet seeds). Progressive primary tuberculosis can lead to more extensive lung involvement with pnemonic infiltrates. Upper lobe cavitary disease is characteristic for secondary tuberculosis (reactivation or reinfection) in persons who have previously mounted an immune response. One pattern of disseminated tuberculosis is Pott disease of the spine, sometimes as an isolated finding.
**UNIT I  General Pathology**

**35 C** Anergy (less than the 10 mm of induration expected for a positive tuberculin test), sputum negativity despite extensive pulmonary disease, and radiographic evidence of infiltrates resembling bacterial pulmonary consolidation all point to a poor cell-mediated immune response. HIV infection depletes the body of the CD4+ lymphocytes (explaining his lymphopenia) essential for a T_{H1} immune response required to contain mycobacterial infection. The debilitation accompanying alcohol abuse is more likely to lead to typical secondary tuberculosis, but more florid. Diabetes mellitus predisposes to bacterial infections, but pulmonary disease is not characteristic for diabetic complications. Scurvy may affect connective tissues but not lung specifically. Smoking diminishes pulmonary innate immune defenses, mainly against bacterial pathogens.

PBD9 373–376   BP9 324  PBD8 347–348  BP8 334–336

**36 C** Hansen disease (leprosy) is caused by the small, acid-fast organism *Mycobacterium leprae*, which chronically infects peripheral nerves and skin. This organism cannot be cultured in artificial media. Diagnosis is made by biopsy of a skin lesion. There are two polar forms of leprosy. In the tuberculoid form, a delayed type of hypersensitivity reaction, with a T_{H1} immune response driven by interferon-γ and interleukin-2 (IL-2) cytokines, gives rise to granulomatous lesions that resemble tuberculosis; acid-fast bacilli are rare in such lesions. In contrast, in the lepromatous form, shown in the figure, T cell immunity is markedly impaired, a T_{H2} immune response is driven by IL-4 and IL-10, and granulomas are poorly formed. Instead, there are large aggregates of lipid-filled macrophages that are stuffed with acid-fast bacilli. Leprosy is poorly transmissible through aerosols (not from direct contact); it probably requires some genetic susceptibility, such as genetic variations in IL-10 and Toll-like receptors; and, similar to most diseases acquiring alcohol abuse is more likely to lead to typical secondary tuberculosis, but more florid. Diabetes mellitus predisposes to bacterial infections, but pulmonary disease is not characteristic for diabetic complications. Scurvy may affect connective tissues but not lung specifically. Smoking diminishes pulmonary innate immune defenses, mainly against bacterial pathogens.

PBD9 373–376   BP9 324  PBD8 347–348  BP8 334–336

**37 D** Infection with *Treponema pallidum* can lead to syphilitic chancres in the primary stage of syphilis. The chancres are characterized by lymphoplasmacytic infiltrates and by an obliterative endarteritis. Similar lesions also may appear with secondary syphilitic mucocutaneous lesions. Acute inflammation with abscess formation is characteristic of bacterial infections such as gonorrhea. Caseating granulomatous inflammation is more characteristic of tuberculosis or fungal infections. Gummatous inflammation can be seen in adults with tertiary syphilis or in congenital syphilis.

PBD9 378–381   BP9 672–674  PBD8 374–375  BP8 701–703

**38 C** Untreated infection with *Treponema pallidum* can lead to tertiary syphilis years later. The most common manifestations of tertiary syphilis include aortitis (typically in the thoracic portion), neurosyphilis, and gummatous nerosis of skin, soft tissue, bone, and joint (Charcot joint). This organism cannot be cultured. The spirochetes are best identified by darkfield microscopy in exudates from primary chancres, but the organisms are hard to find in the tertiary stage of the disease. Serologic testing is useful for screening and confirmation of syphilis. The nontreponemal tests (RPR, VDRL) are sensitive to a cardiolipin found in the more numerous spirochetes earlier in the disease; but these tests are not specific because the presence of cardiolipin in human tissues is associated with other diseases, causing false-positive results. The FTA test has specificity for *T. pallidum*.

PBD9 378–381   BP9 672–674  PBD8 374–375  BP8 701–703

**39 D** These are findings of congenital syphilis with nodules of gummatous nerosis. Because the spirochetes can cross the placenta in the third trimester, early stillbirths do not occur. Infants who survive have features similar to adult secondary syphilis, with rash. With survival, late complications of the periosteitis and perichondritis include bone and teeth deformities (e.g., saber shin). Herpes infections in the neonate usually are not initially obvious because most of these infections are acquired by passage through the birth canal. Most infants born with HIV infection have no initial gross or microscopic pathologic findings. Congenital toxoplasmosis and cytomegalovirus produce severe cerebral disease.

PBD9 378–381   BP9 312, 671–674  PBD8 375  BP8 592

**40 E** The acute stage of Lyme disease is marked by the appearance of erythema chronicum migrans of the skin. As the *Borrelia burgdorferi* organisms proliferate and disseminate, systemic manifestations of carditis, meningitis, and migratory arthralgias and myalgias appear. Arthritis involving the large joints occurs 2 to 3 years after initial infection. Rheumatoid arthritis can mimic Lyme disease but is not preceded by the skin lesions described. Chagas disease may be associated with acute and chronic myocarditis leading to heart failure; some patients have esophageal involvement, but arthritis and rash are not features of the disease. Hemorrhagic fever, or dengue fever, caused by an arbovirus, can produce myositis and bone marrow suppression. Mucocutaneous ulcers may be seen with *Leishmania braziliensis* infection, which is transmitted via sand flies. The area of the tick bite that introduces *Borrelia burgdorferi* spirochetes, the cause of Lyme disease, may manifest erythema chronicum migrans. Onchocerciasis occurs as a result of infection with the filarial nematode *Onchocerca volvulus* and leads to formation of a subcutaneous nodule.

PBD9 377–378   BP9 312  PBD8 372–373  BP8 324, 337

**41 B** The spores of *Clostridium botulinum* will survive the canning process in such non-acidic foods as peas when they are not heated sufficiently, so that organisms grow and elaborate a neurotoxin. However, the plot fails when everyone prefers to eat junk food and not vegetables. Perhaps the terrorists should promote trans fats, which, when combined with
lack of exercise, will increase morbidity and mortality from atherogenesis to a greater extent than any infectious agent. Chlamydia psittaci is a Category B agent that is airborne and causes pneumonia. Treponema pallidum, the Black Death, which produces lymphadenitis, pneumonia, and sepsis; the vector is the rat flea.

42 D Clostridia such as Clostridium perfringens represent one type of gram-positive rod like bacteria that produce powerful exotoxins, causing extensive tissue necrosis so quickly that the acute inflammatory response lags. Abscences are formed of neutrophils responding to the inflammatory agent, often a bacterial organism, but the liquefactive necrosis is mainly produced by enzymes released from the neutrophils. Fibrous scarring can certainly be part of the healing phase of inflammatory responses, but is less prominent with bacterial infections than with agents producing more chronic inflammation. Granulomatous inflammation typically develops in weeks to months from persistent infection from agents such as mycobacteria. Lymphocytic infiltrates are most typical for chronic and viral infections, and there tends to be minimal necrosis.

43 A This man has tetanus. The contamination of a wound with Clostridium tetani can result in the elaboration of a potent neurotoxin. This toxin is a protease that cleaves synaptobrevin, a major transmembrane protein of the synaptic vesicles in inhibitory neurons. Clostridium perfringens elaborates a variety of toxins, one of which (alpha) is a phospholipase causing myonecrosis. Inhibition of acetylcholine release is not a feature of infection. The toxin of Staphylococcus aureus is an enterotoxin that acts as a superantigen and stimulates T cell cytokine release. Cholera is produced when the toxin elaborated by Vibrio cholerae stimulates epithelial cell adenylate cyclase.

44 E The large, gram-positive rods seen in the figure are characteristic of Clostridium perfringens, which can contaminate open wounds and produce gas gangrene. Clostridial organisms can elaborate multiple toxins. C. perfringens alpha-toxin acts as a phospholipase C that degrades lecithin in cellular membranes. Lipopolysaccharides are found in gram-negative organisms. Inhibition of phagocytes is a feature of organisms such as Mycobacterium tuberculosis. Antibiotic resistance is increasing in frequency, but is not the main mechanism for clostridial virulence. Though devitalized tissues can have polymicrobial infection, Candida is typically not the most virulent among superinfecting agents.

45 B The most common cause of nongonococcal urethritis in men is Chlamydia trachomatis. The condition is a nuisance in men without significant sequelae; however, the behavior that led to the infection can place the patient at risk for other sexually transmitted diseases. Candida infections typically occur in immunocompromised patients or in patients receiving long-term antibiotic therapy. Herpes simplex can produce painful vesicles on the skin. Tuberculosis of the urinary tract is uncommon. A syphilitic chancre on the penis, not present here, is an indicator of Treponema pallidum infection.

46 B Infection with Chlamydia trachomatis is a common sexually transmitted disease. Most cases produce only urethritis and cervicitis; however, some strains of C. trachomatis can produce lymphogranuloma venereum, a chronic ulcerative disease that is more endemic in Asia, Africa, and the Caribbean. In this disease, there is a mixed granulomatous and neutrophilic inflammatory reaction, as seen in this patient. In contrast, herpes simplex virus produces clear mucocutaneous vesicles with no exudates and is unlikely to involve lymph nodes. Candidiasis can produce superficial inflammation with an exudate, but it is rarely invasive or disseminated in non-immunosuppressed individuals. Bacterial vaginosis due to Gardnerella produces a whitish discharge that has a “fishy” odor. Treponema pallidum, the causative agent of syphilis, produces a hard chancre on skin and mucosal surfaces.

47 D This patient has Rocky Mountain spotted fever, which occurs sporadically in the United States, mostly in areas other than the Rocky Mountains. Rickettsial diseases produce signs and symptoms from damage to vascular endothelium and smooth muscle similar to a vasculitis. The most common vector is the wood tick Dermacentor andersoni. Thrombosis of the affected blood vessels is responsible for foci of skin necrosis. Headache and abdominal pain are often prominent. Lyme disease, caused by Borrelia burgdorferi, can produce an erythema chronicum migrans of skin at the site of a tick bite. Mucocutaneous leishmaniasis mainly involves the nasal and oral regions. Hansen disease (leprosy), produced by Mycobacterium leprae, results in skin anesthesia that predisposes to recurrent injury. Plague, caused by Yersinia pestis, can produce focal skin necrosis at the site of a flea bite, and ulceration over infected lymph nodes (bubos).

48 A Aspergillus, Candida, and Mucor infections may become disseminated in the setting of neutropenia. Vascular invasion can occur with fungal infections, particularly with Aspergillus and Mucor. The branching septate hyphae are shown in the figure projecting from a fruiting body of Aspergillus. After these organisms gain a foothold (hyphae-hold) in tissues, they are very difficult to eradicate. Coccioidioides immitis and Histoplasma capsulatum are fungi that can produce pulmonary disease resembling that of Mycobacterium tuberculosis, with granulomatous inflammation. They do not have a propensity for vascular invasion. Corynebacterium diphtheriae produces upper respiratory tract disease, mainly in children who are not vaccinated against it. Pneumocystis pneumonia is not typically accompanied by vascular changes.
49 F This patient is in diabetic ketoacidosis, which is a significant risk factor for mucormycosis. Note the broad, nonseptated hyphae more easily visible with H&E stain than special stains, unlike other fungi. In contrast, *Aspergillus* organisms have thinner hyphae with acute angle branching and septations. *Actinomyces* organisms are long, filamentous gram-positive bacilli. *Candida* infections are typically superficial and have gram-positive budding cells with pseudo hyphae. Large, gram-positive rods are characteristic of *Clostridium perfringens*, which can contaminate open wounds and produce gas gangrene.

PBD9 389  BP9 313, 829  PBD8 385–386  BP8 324, 337, 527–528

50 C This boy had cerebral malaria, the worst form of malaria. After the infective mosquito bite, *Plasmodium falciparum* sporozoites invade liver cells and reproduce asexually. When the hepatocytes rupture, they release thousands of merozoites that infect RBCs. The infected RBCs circulate and can bind to endothelium in the brain. Small cerebral vessels become plugged with the RBCs, resulting in ischemia. The other listed options also could be secondarily involved by vascular thromboses in the setting of malaria, but are not extraerythrocytic sites for asexual reproduction.


51 A This patient’s travel history suggests an insect-borne disease. The figure shows the characteristic tetrad and ring forms of *Babesia microti*, within erythrocytes. Babesiosis is, an uncommon malaria-like protozoan disease. The northeastern United States is an endemic area. The vector is the deer tick, just as with Lyme disease from *Borrelia burgdorferi*, which is a spirochete. Giardiasis typically produces self-limited, watery diarrhea. *Rickettsia rickettsii* causes Rocky Mountain spotted fever, which occurs sporadically in the United States in areas other than the Rocky Mountains and produces signs and symptoms from damage to vascular endothelium and smooth muscle similar to a vasculitis. *Wuchereria bancrofti* is a form of filariasis that can cause elephantiasis, owing to lymphatic obstruction in the presence of an inflammatory reaction to the adult filarial worms.

PBD9 392  PBD8 388  BP8 321, 328

52 C Visceral leishmaniasis (kala-azar) is caused by protozoa in the *Leishmania donovani* complex. Of these, only *L. donovani infantum* is endemic to southern Europe and the Mediterranean area. It is transmitted to humans by the sand fly (*Phlebotomus*). Pancytopenia implies bone marrow involvement, possibly enhanced by the enlarged spleen, and the liver function abnormalities suggest liver involvement. Borreliosis causes relapsing fever and is transmitted via body lice. Echinococcal disease is caused by ingestion of tapeworm eggs and can lead to cyst formation in visceral organs. *Borrelia burgdorferi* infection is transmitted via ticks and can cause Lyme disease, characterized by erythema chronicum migrans, meningoencephalitis, and chronic arthritis. Schistosomiasis, which is transmitted via snails, can produce hepatic cirrhosis (*Schistosoma mansoni* or *Schistosoma japonicum*) or bladder disease (*Schistosoma haematobium*). Typhus is a louse-borne rickettsial disease with skin rash that may proceed to skin necrosis.

PBD9 392–393  BP9 310, 313  PBD8 388–390  BP8 324, 328

53 C This patient has cutaneous leishmaniasis, and the original papule was at the site of the sand fly vector bite. Leishmaniasis is endemic in the Middle East, South Asia, Africa, and Latin America. The organisms proliferate within macrophages in the mononuclear phagocyte system and can cause regional lymphadenopathy. The cutaneous form does not have bone marrow involvement and splenic enlargement, so pancytopenia is not present. Borreliosis causes relapsing fever and is transmitted via body lice. *Brugia malayi* is a nematode transmitted by mosquitoes that leads to filariasis involving lymphatics to produce elephantiasis. *Leishmania donovani* is transmitted by sand flies and leads to infection of macrophages, which produces hepatosplenomegaly, lymphadenopathy, and bone marrow involvement with pancytopenia. Listeriosis is most often acquired via contaminated food or water. In most adults, it produces mild diarrheal illness, but in some adults and children, and in fetuses, it may produce meningitis or dissemina tion with microabscess (microgranuloma) formation. *Mycoplasma pneumoniae* causes Hansen disease (leprosy), with infection of peripheral nerves and skin. In individuals with a strong immune response, the tuberculoid form of this disease results in granuloma formation; in individuals with a weak immune response, the lepromatous form occurs, characterized by large numbers of macrophages filled with short, thin, acid-fast bacilli. African trypanosomiasis produces sleeping sickness.

PBD9 392–393  BP9 310, 313  PBD8 388–390  BP8 324, 328

54 E The findings are consistent with African trypanosomiasis, or sleeping sickness. The eradication of the tsetse fly vector has been a priority for decades in many African countries. Filarial worms endemic in parts of Central America, Southeast Asia, and Polynesia also can appear in blood, but are smaller in size and do not lead to chronic wasting. Filariasis is not endemic in Europe.

PBD9 394  BP9 313  PBD8 390  BP8 322, 325, 328

55 C This child is infected with *Trypanosoma cruzi*, resulting in Chagas disease, endemic to Central and South America. The vector is the reduviid (triatomid) bug. The organisms can damage the heart by direct infection or by inducing an autoimmune response that affects the heart because of the existence of cross-reactive antigen. Acute myocarditis rarely occurs, but most deaths from acute Chagas disease are due to heart failure. In 20% of infected individuals, cardiac failure can occur 5 to 15 years after the initial infection. The affected heart is enlarged, and all four chambers are dilated. A cerebral abscess or acute meningitis is typically a complication of a bacterial infection with septicemia. Chronic arthritis can be seen in Lyme disease, which is transmitted by deer ticks. Mucocutaneous ulcers may be seen in *Leishmania braziliensis* infection, which is transmitted via sand flies. Paranasal sinus infection may be caused by *Mucor circinelloides*.

PBD9 394–395  BP9 402  PBD8 391  BP8 322, 414–415
56 D The rhabditoid larvae of Strongyloides stercoralis can become invasive filariform from autoinfection in immunocompromised hosts, so-called hyperinfection with involvement of multiple organs. Immunocompetent hosts typically have only diarrhea. Parasites, particularly worms, crawling through tissues incite a marked eosinophilia. Cysticercosis from eating uncooked pork can result in the release of larvae that penetrate the gut wall and disseminate hematogenously, often settling in gray and white cerebral tissue, where they develop into cysts. Onchocerciasis occurs as a result of infection with the filarial nematode Onchocerca volvulus and leads to formation of a subcutaneous nodule. Schistosoma mansoni or Schistosoma japonicum infections have adult female worms in the portal venous system that release eggs that can produce hepatic fibrosis; Schistosoma haematobium worms live in veins near the bladder and release eggs that result in hematuria. Eating infected meat, typically uncooked pork, can lead to trichinosis: Trichinella encysts in striated muscle to produce fever and myalgias.

57 E Acute muscle pain with fever and eosinophilia suggests a parasitic infestation of the skeletal muscles, most likely trichinosis; this results from ingesting poorly cooked meat infected with Trichinella spiralis larvae. Hemorrhagic fever can be a mild disease with myalgia, but can be severe with extensive vascular endothelial damage. Influenza A and B infection may produce myositis in childhood, with more severe, focal, and later onset than the diffuse myalgias of typical flu. Poliomyelitis can lead to muscle weakness, but via neurogenic atrophy, through loss of motor neurons. Scrub typhus caused by Orientia tsutsugamushi can have myalgia along with eschar and lymphadenopathy in the region of a chigger bite.

58 B Schistosoma haematobium is a parasitic infection most often seen in Africa, particularly the Nile Valley, in areas where irrigation has expanded the range of the host snails. The adult worms live in veins adjacent to the bladder and release eggs with a sharp spine to cut their way through the wall of the urinary bladder, causing severe granulomatous inflammation, fibrosis, and calcification. Mycobacterial infections of the urinary tract are uncommon and do not cause bladder fibrosis. Trichinella spiralis infects striated muscle. Ascariasis involves the lower gastrointestinal tract, and the worms reside in the lumen. Cysticercosis from the pork tapeworm Taenia solium can have a wide tissue distribution, but the brain is most often affected.

59 C These farmers are infected with either Schistosoma mansoni or Schistosoma japonicum. Female worms reside in the portal venous system and release eggs that cut their way into the liver and incite a granulomatous inflammatory reaction. With time, the portal granulomas undergo fibrosis, compressing the portal veins. This gives rise to severe portal hypertension, splenomegaly, and ascites. A dilated cardiomyopathy may occur with Chagas disease, in which the Trypanosoma cruzi organisms are transmitted through the reduviid (triatomid) bug. Elephantiasis is a complication of filariasis, which is transmitted via mosquitoes. Mucocutaneous ulcers may be seen in Leishmania braziliensis infection, which is transmitted via sand flies. Squamous cell carcinomas may be seen in the bladder in chronic Schistosoma haematobium infection.

56 E The marked soft tissue enlargement and deformity is called elephantiasis, which results from lymphatic obstruction in the presence of an inflammatory reaction to the adult filarial worms Wuchereria bancrofti. Echinococcus produces hydatid disease of the liver, lungs, or bone. Leishmania tropica can involve the skin, causing ulceration, and can enlarge parenchymal organs. Schistosomiasis from Schistosoma mansoni may affect the liver most severely. Trichinella larvae from ingested, poorly cooked meat encyst in striated muscle.

57 E Onchocerciasis is caused by Onchocerca volvulus with inflammation induced by the microfilaria. Both insect abatement programs and campaigns to treat affected populations with ivermectin have helped to reduce the prevalence of this disease. Though ivermectin kills microfilariae, it does not kill the adult worms, and treatment with doxycycline will eliminate the worm that symbiotic Wolbachia bacteria need for reproduction. Acid-fast Mycobacterium leprae organisms may be seen in skin lesions of lepromatous leprosy, but the eye is not involved. Elementary bodies can be identified in conjunctival scrapings with Chlamydia trachomatis infection, but additional eye components are not involved, and there are no skin lesions. Intracellular diplococci of Neisseria gonorrhoeae can cause neonatal blindness. Intraneuronal inclusions can be seen with herpes simplex keratitis, which can cause perforation through corneal ulcerations.

58 B The Centers for Disease Control and Prevention has classified microbes into several categories based on the danger they pose as agents for bioterrorism on the basis of their ease of production, dissemination, and production of serious illness. Variola major is the causative agent for smallpox and has a mortality rate of 30%. Francisella tular-ensis is very infectious; only 10 to 50 organisms can cause disease. As a weapon, the bacteria can be made airborne for exposure by inhalation. Infected individuals experience life-threatening pneumonia. Chlamydia psittaci can cause psittacosis, which also can produce pneumonitis, but the course is more variable. Hantavirus can produce a severe pneumonia, but the prodrome is longer, and the vector is the deer mouse. Mycobacterium kansasii produces findings similar to Mycobacterium tuberculosis. Rickettsia typhi is the causative agent for murine typhus with headache and rash.
1. In an experiment, the effects of xenobiotic activation of the compound benzo[a]pyrene, a chemical carcinogen present in cigarette smoke, are studied in various tissues. Investigators determine that formation of a secondary metabolite, which binds covalently to DNA, increases the frequency of lung cancers. Genetic polymorphisms are found among persons that affect this frequency. Which of the following is the most likely metabolic pathway for generation of this xenobiotic?

A. Biomethylation  
B. Cytochrome P-450  
C. Flavin-containing monooxygenase  
D. Glucuronidation  
E. Glutathione reduction  
F. Peroxidase-dependent cooxidation

2. An environmental study shows that a gas that exists in nature is present at increased levels in polluted city air. In the upper atmosphere, this gas blocks harmful ultraviolet radiation. At ground level, this gas generates free radicals in the lower respiratory tract. What is this gas?

A. Argon  
B. Carbon dioxide  
C. Nitrogen dioxide  
D. Ozone  
E. Sulfur dioxide

3. A textbook author standing knee-deep in a rising tide next to his home notes that this phenomenon is becoming more frequent. He surmises that the risk for illnesses from infections in populations living in such coastal areas is increasing. Which of the following is most likely to be affected by illness in these populations?

A. Cardiovascular system  
B. Central nervous system  
C. Gastrointestinal tract  
D. Hematopoietic system  
E. Respiratory tract  
F. Urinary tract

4. A 75-year-old man who lives alone in a poorly ventilated house without central heating uses a portable unvented kerosene heater to warm the house during the winter months. One morning, a neighbor finds him in an obtunded state. On physical examination, he appears cyanotic. Results of blood gas measurement on room air are Po2, 90 mm Hg; Pco2, 35 mm Hg; and pH, 7.3. Pulse oximetry shows low oxygen saturation. Exposure to which of the following is most likely to have produced this man’s illness?

A. Beryllium  
B. Carbon monoxide  
C. Nitrous oxide compounds  
D. Oxygen  
E. Ozone  
F. Sulfur dioxide

5. An 8-year-old girl exhibits lethargy and somnolence with dizziness and weakness after a 6-hour bus ride returning from a week at summer camp. On physical examination, her temperature is 37°C, pulse is 107/min, respiratory rate is 28/min, and blood pressure is 130/85 mm Hg. Breath sounds are audible in all lung fields with no wheezes or crackles. Arterial blood gas analysis shows pH, 7.35; Po2, 95 mm Hg; Pco2, 37 mm Hg; and HCO3−, 20 mEq/L. Pulse oximetry shows an oxygen saturation of 90%, but the spectrophotometrically measured oxyhemoglobin saturation is 60%. Her blood lactic acid is 8 mmol/L, and total creatine kinase is 445 U/L. She is given 100% oxygen in a hyperbaric chamber and improves in 20 minutes. She is most likely to have experienced poisoning with which of the following?

A. Aspirin (acetylsalicylic acid)  
B. Carbon monoxide  
C. Iron sulfate  
D. Lead  
E. Methanol  
F. Organophosphate insecticide
Several children between the ages of 3 and 6 years have been admitted to a local hospital because of encephalopathic crisis. They have lived in the same community all their lives. All have previously exhibited retarded psychomotor development. On physical examination, the children have diffuse abdominal pain and are experiencing nausea and vomiting. Head CT scans show marked cerebral edema. Laboratory studies show microcytic anemia. An investigator sent to the housing project where the children live finds a rundown apartment complex with extensive water damage, poor plumbing and ventilation. Toxic exposure to which of the following substances best accounts for these findings?

A Ethylene glycol  
B Kerosene  
C Lead  
D Methanol  
E Sodium hypochlorite

A previously healthy, 27-year-old agricultural worker develops nausea, vomiting, abdominal cramps, cough with wheezing and dyspnea, salivation, and lacrimation 1 hour after leaving work in a field. On examination, he is afebrile with a heart rate of 50/min and blood pressure of 90/55 mm Hg. Laboratory studies show plasma cholinesterase activity less than 50% of normal. Atropine is given intravenously, and his condition improves. He was most likely exposed to which of the following substances?

A Aflatoxin  
B Cyclodiene  
C Dioxin  
D Ergot alkaloid  
E Organophosphate  
F Pyrethrin

An epidemiologic study of a community shows an increased incidence for basal cell and squamous cell carcinomas in adults. These lesions are distributed predominantly on palms and soles. Analysis of groundwater pumped from wells for drinking shows increased levels of a heavy metal used in pressure-treated lumber (with a greenish hue), insecticides, and herbicides. Which of the following metals is most likely implicated by this study?

A Arsenic  
B Beryllium  
C Cadmium  
D Lead  
E Mercury

A 36-year-old man is the owner of a radiator repair shop, where he works cleaning, cutting, polishing, and welding metals. Over 6 months, he develops worsening malaise with headache and abdominal pains and has difficulty holding his tools. CBC indicates microcytic anemia, and basophilic stippling of RBCs is seen on the peripheral blood smear. An elevated blood level of which of the following would be most useful in determining the toxic exposure causing his illness?

A Alanine aminotransferase  
B Calcium  
C Creatine kinase  
D Selenium  
E Zinc protoporphyrin

Increasing numbers of children younger than 5 years are observed to exhibit cerebral palsy, deafness, blindness, and mental retardation in a coastal community. A coal-burning power plant was built in this community 5 years ago. Affected families have a high consumption of seafood. Which of the following compounds ingested during pregnancy by the mothers of these children is most likely to be responsible for the illness affecting the children?

A Bisphenol A  
B Cadmium  
C Methyl mercury  
D Organochlorine  
E Vinyl chloride

A 72-year-old man has had increasing dyspnea for the past year. Decreased breath sounds are heard on auscultation of the right side of the chest. A chest radiograph shows a large pleural mass that nearly encases the right lung. Exposure to which of the following metals is most likely to be associated with these findings?

A Arsenic  
B Asbestos  
C Beryllium  
D Chromium  
E Nickel

A 61-year-old woman living in a city with very poor air quality has developed a worsening cough over the past 7 months. She has an 80 pack-year history of smoking cigarettes. One week ago she had an episode of hemoptysis. A chest radiograph shows a 7-cm infiltrative, perihilar mass in the right lung. Exposure to which of the following airborne agents is most likely to be associated with these findings?

A Carbon monoxide  
B Nicotine  
C Nitrous oxide compounds  
D Ozone  
E Polycyclic aromatic hydrocarbons  
F Dusts containing silica  
G Sulfur dioxide

While attending a party, a 19-year-old university student drinks 2 L of mixed alcoholic beverages containing 50% ethanol by volume over 30 minutes. He usually does not drink much alcohol. His major use of drugs consists of acetaminophen for headaches. Which of the following complications is most likely to prove lethal within the first 12 hours?

A Acute pancreatitis  
B Brainstem depression  
C Hepatic cirrhosis  
D Variceal bleeding  
E Wernicke disease
14. Ingested ethanol is observed to be metabolized by alcohol dehydrogenase to an intermediate compound that, upon further metabolism, depletes reduced nicotinamide adenine dinucleotide (NAD<sup>−</sup>) in the cytoplasm of hepatocytes. Fatty acid oxidation is subsequently reduced. Population studies show some persons have deficient enzyme activity to metabolize this intermediate compound. What is this intermediate compound?

A. Acetaldehyde  
B. α-Tocopherol  
C. Formic acid  
D. Glutathione  
E. Hydrogen peroxide

15. A case-control study seeks to identify long-term effects of hormone replacement therapy (HRT) in postmenopausal women receiving exogenous estrogens coupled with progestins compared with a control group not receiving this therapy. The medical records of the women in the study are reviewed after 20 years. Which of the following complications is most likely to be observed in the women receiving HRT?

A. Cervical carcinoma  
B. Chronic ulcerative colitis  
C. Hepatic cirrhosis  
D. Pulmonary emphysema  
E. Thromboembolism

16. A 36-year-old woman has been using low-dose estrogen-containing oral contraceptives for the past 20 years. She has smoked one pack of cigarettes per day for the past 18 years. She is G2, P2, and both pregnancies ended with term live-born infants of low birth weight, but no anomalies. On physical examination, no abnormal findings are noted. Her BMI is 24. She is at increased risk for developing which of the following conditions?

A. Breast carcinoma  
B. Cholecystitis  
C. Dementia with Lewy bodies  
D. Endometrial carcinoma  
E. Myocardial infarction  
F. Ovarian carcinoma

17. A 22-year-old star football player suddenly collapses during practice and has a cardiac arrest and cannot be resuscitated. The medical examiner investigating this sudden death finds marked coronary atherosclerosis and histologic evidence of hypertension in the renal blood vessels at autopsy. Use of which of the following substances by this man most likely led to these findings?

A. Amphetamine  
B. Barbiturate  
C. Benzodiazepine  
D. Cocaine  
E. Ethanol  
F. Heroin  
G. Marijuana

18. A 26-year-old woman with a 6-month history of depression accompanied by active suicidal ideation ingests 35 g of acetaminophen. She quickly experiences nausea and vomiting. Within 1 day, she becomes progressively obtunded. On physical examination, her temperature is 36.9° C, pulse is 75/min, respirations are 15/min, and blood pressure is 100/65 mm Hg. She is treated with N-acetylcysteine. Depletion of which of the following is most likely to accentuate her organ damage?

A. Glutathione (GSH)  
B. Amylase  
C. Creatine kinase  
D. Ketone bodies  
E. Potassium

19. A 27-year-old, previously healthy man suddenly collapses at a party where both legal and illicit drugs are being used. En route to the hospital, he requires resuscitation with defibrillation to establish a normal cardiac rhythm. On physical examination, his temperature is 40° C; respirations, 30/min; heart rate, 110/min; and blood pressure, 175/90 mm Hg. He has dilated pupils, a perforated nasal septum, and a prominent callus on the right thumb. CT scan of the head shows an acute right frontal lobe hemorrhage. Which of the following substances is most likely responsible for these findings?

A. Amphetamine  
B. Barbiturate  
C. Cocaine  
D. Ethanol  
E. Heroin  
F. Marijuana  
G. Phencyclidine

20. A 63-year-old man with a history of chronic arthritis has had pronounced tinnitus and episodes of dizziness for the past 6 months. He has had a headache and nausea for the past 2 days. Physical examination shows he is afebrile but has a respiratory rate of 35/min and heart rate of 100/min. He has scattered petechiae over the skin of the upper extremities. There is no apparent bone conduction or nerve hearing loss. A stool guaiac test result is positive. Laboratory studies show serum lactate is 6 mmol/L. An arterial blood gas analysis shows a pH of 7.25; PO<sub>2</sub>, 95 mm Hg; P<sub>CO2</sub>, 35 mm Hg; and HCO<sub>3</sub>−, 15 mEq/L. Acute and chronic toxicity from which of the following drugs best explains these findings?

A. Acetaminophen  
B. Aspirin  
C. Chlorpromazine  
D. Iron  
E. Morphine  
F. Quinidine  
G. Tetracycline
21 A 20-year-old man is brought to the hospital emergency department by a friend who found him unconscious in his apartment after trying to contact him for 3 days. On arrival, the patient is in a state of respiratory depression. He experiences convulsions for 2 minutes, followed by cardiac arrest. Advanced cardiac life support measures are instituted, and he is stabilized and intubated. On physical examination, there are needle tracks in the left antecubital fossa, miosis, and a loud diastolic heart murmur. His temperature is 39.2° C. Use of which of the following substances by this man most likely produced these findings?

A  Cocaine  
B  Ethanol  
C  Flurazepam  
D  Heroin  
E  Meperidine  
F  Phencyclidine  
G  Lysergic acid

22 A clinical trial involves patients with a diagnosis of cancer who have intractable nausea as a result of chemotherapy. The patients are divided into two groups; the group receiving the drug is found to have reduced self-reported nausea and diminished weight loss compared with the placebo group. The patients receiving the drug seem to have no major adverse side effects. Of the following agents, classified as drugs of abuse in parts of the world, which is most likely to have the beneficial effects found in this study?

A  Barbiturate  
B  Cocaine  
C  Heroin  
D  Marijuana  
E  Methylphenidate  
F  Methamphetamine  
G  Phencyclidine

23 A 7-year-old boy falls off his bicycle while riding down the street at 5 km/hr. The skin of his right calf and right arm scrape along the pavement, and the top layer of epidermis is removed. Which of the following terms best describes this injury?

A  Abrasion  
B  Burn  
C  Contusion  
D  Incision  
E  Laceration

24 During a qualifying match for the World Cup, the goalkeeper is hit in the chest by a soccer ball (football) kicked from 10 m away. He stays in the game. Which of the following injury patterns is most likely to be seen over the chest of the goalkeeper?

A  Abrasion  
B  Contusion  
C  Incision  
D  Laceration  
E  Puncture

25 A 14-year-old boy was taken to the emergency department of a local hospital, where he died 2 hours later. The principal finding on examination is shown in the figure. Laboratory studies showed that the hematocrit was 17%. Which of the following terms best describes this injury?

A  Blast injury  
B  Electrocution injury  
C  Gunshot wound  
D  Incised wound  
E  Laceration

26 A 33-year-old man incurs thermal burn injuries to 40% of his total body surface area in an accidental fire while repairing a fuel storage tank. On physical examination, the skin of the trunk, neck, and face is pink, shows blister formation, and is painful when touched. The skin of the arms is white and anesthetic. Skin grafting is necessary on the arms, but not on other injured areas of skin. Which of the following tissue components, when absent, will necessitate skin grafting?

A  Collagen fibers  
B  Dermal appendages  
C  Epidermal stratum spinosum  
D  Keratin  
E  Macrophages  
F  Nerve endings
27. A 5-year-old boy is brought to the physician for examination. A police agency suspects child abuse. On physical examination, the child’s skin has the appearance shown in the figure. Which of the following terms best describes this injury? 
   A. Abrasion  
   B. Contusion  
   C. Hypothermia  
   D. Kwashiorkor  
   E. Laceration  
   F. Thermal burn

28. While touring the grounds of the Imperial Palace in Kyoto, a 75-year-old woman collapses suddenly. She remains conscious, but says that she feels weak and light headed. On physical examination by the nurse, her temperature is 35.2°C, pulse is 93/min, respirations are 17/min, and blood pressure is 95/50 mm Hg. The temperature in the shade is 34° C with 90% humidity. One hour after drinking cool green tea, the woman revives and is able to return to her hotel. Which of the following terms best describes these findings? 
   A. Heat cramps  
   B. Heat exhaustion  
   C. Heat stroke  
   D. Malignant hyperthermia  
   E. Thermal inhalation injury

29. A 20-year-old man is trying to repair an old electrical appliance in his garage. While testing the function of the appliance, his right hand comes in contact with a frayed electrical cord carrying 120-V, 10-amp alternating current. Which of the following is most likely to develop as a consequence of this electrical injury? 
   A. Bronchoconstriction  
   B. Cerebral artery thrombosis  
   C. Heat stroke  
   D. Gastric hemorrhage  
   E. Ventricular fibrillation

30. A 30-year-old woman is found dead in her hotel room 3 hours after firemen extinguish a fire on the floor below. Investigation of the scene shows no signs of fire within her room, and the medical examiner observes no external findings on the body. Which of the following conditions best explains the woman’s death? 
   A. Acute myocardial infarction  
   B. Cerebral hemorrhage  
   C. Malignant hyperthermia  
   D. Pseudomonas aeruginosa septicemia  
   E. Pulmonary edema

31. “It was terribly cold and nearly dark on the last evening of the old year, and the snow was falling fast. In the cold and the darkness, a poor little girl, with bare head and naked feet, roamed through the streets. So the little girl went on with her little naked feet, which were quite red and blue with the cold. Shivering with cold and hunger, she crept along.” Which of the following pathologic findings in soft tissues is most likely to be present in this girl? 
   A. Acute inflammation  
   B. Apoptosis  
   C. Edema  
   D. Hemorrhage  
   E. Nuclear karyorrhexis

32. A 76-year-old man has noticed a gradually enlarging nodule on the right lower eyelid for 3 years. On physical examination, the 0.8-cm umbilicated nodule is firm and has a small central area of ulceration. The nodule is excised, and plastic repair to the eyelid is performed. Which of the following forms of radiation most likely played the greatest role in the development of this man’s lesion? 
   A. Gamma rays  
   B. Infrared rays  
   C. Ultraviolet rays  
   D. Visible rays  
   E. X-rays

33. A 56-year-old man with a 60 pack-year history of smoking has been diagnosed with a squamous cell carcinoma of the larynx. He receives radiation therapy, 40 Sv (4000 cGy) in divided doses, to treat the carcinoma. One year later, endoscopy shows no gross evidence of residual carcinoma. Which of the following adverse effects is most likely to be present in this patient as a result of this radiotherapy? 
   A. Azoospermia  
   B. Cerebral atrophy  
   C. Colonic ulceration  
   D. Marrow aplasia  
   E. Vascular fibrosis

34. The firemen who initially responded to fight the fires from the Chernobyl nuclear reactor accident were exposed to high radiation levels. Some of the men received doses exceeding 50 Sv (5000 cGy). Within hours, many became extremely ill. Damage to which of the following tissues most likely led to this finding? 
   A. Bone marrow  
   B. Cerebrum  
   C. Heart  
   D. Lungs  
   E. Small intestine
35 A 3-year-old boy has had a succession of respiratory infections during the past 6 months. On physical examination, the child appears chronically ill, listless, and underdeveloped. He is 50% of ideal body weight and has marked muscle wasting. Laboratory findings include hemoglobin, 9.4 g/dL; hematocrit, 27.9%; MCV, 75 μm³; platelet count, 182,000/mm³; WBC count, 6730/mm³; serum albumin, 4.1 g/dL; total protein, 6.8 g/dL; glucose, 52 mg/dL; and creatinine, 0.3 mg/dL. Which of the following is most likely to explain these findings?

A Bulimia  
B Folate deficiency  
C Kwashiorkor  
D Lead poisoning  
E Marasmus

36 A 26-year-old woman has had amenorrhea for the past 8 years. She fractured her right wrist 1 year ago after a minor fall to the ground. On physical examination, she is 175 cm (5 ft 7 in) tall and weighs 52 kg (BMI 17). She has normal secondary sex characteristics. There are no abnormal findings. Radiographic measurement of bone density by dual-energy x-ray absorptiometry shows a bone mineral density that is 1.5 standard deviations below the young adult reference range. Laboratory findings include anemia and hypoalbuminemia. Which of the following is the most likely diagnosis?

A Anorexia nervosa  
B Bulimia  
C Kwashiorkor  
D Rickets  
E Scurvy

37 A 5-year-old child has had recurrent upper respiratory infections for the past 2 months. The child is at the 55th percentile for height and the 38th percentile for weight. Physical examination shows generalized edema, ascites, muscle wasting, and areas of desquamating skin over the trunk and extremities. Laboratory studies are most likely to show which of the following findings?

A Abetalipoproteinemia  
B Hypoalbuminemia  
C Hypocalcemia  
D Hyperglycemia  
E Megaloblastic anemia

38 An epidemiologic study observes increased numbers of respiratory tract infections among children living in a community in which most families are at the poverty level. The infectious agents include *Streptococcus pneumoniae*, *Haemophilus influenzae*, and *Klebsiella pneumoniae*. Most of the children have had pneumonitis and rubeola infection. The study documents increased rates of keratomalacia, urinary tract calculi, and generalized papular dermatosis in these children as they reach adulthood. These children are most likely to have a deficiency of which of the following vitamins?

A Vitamin A  
B Vitamin B₁  
C Vitamin E  
D Vitamin D  
E Vitamin K

39 A 3-year-old boy does not appear to be developing normally. On physical examination, the child has the appearance shown in the figure. The results from vision testing are normal. There are no petechiae or areas of purpura on the skin. The abdomen is not enlarged. Which of the following is the most likely diagnosis?

A Beriberi  
B Kwashiorkor  
C Rickets  
D Pellagra  
E Scurvy

40 A 3-year-old child has erosion of a roughened corneal surface caused by xerophthalmia. Keratomalacia results in corneal scarring with eventual blindness after 4 years. This ocular damage is now less common because of a United Nations initiative to treat a dietary deficiency of which of the following nutrients?

A Iron  
B Niacin  
C Protein  
D Vitamin A  
E Vitamin K
41. In a study of lifestyle influences on health, investigators observe that sending children outside to play instead of letting them sit for hours in front of the television can have long-term health benefits. Which of the following tissues is most likely to be in better condition by middle age from this lifestyle change?

A. Bone fractures
B. Ocular cataracts
C. Urinary tract calculi
D. Pulmonary emphysema
E. Skin cancers

42. A 48-year-old woman injured her right wrist in a fall down a flight of stairs. On physical examination, she has marked pain on palpation of the wrist and does not want to move the hand. A radiograph of the right hand and arm shows marked osteopenia and a fracture of the radial head. Which of the following underlying diseases is most likely to contribute to the risk of fracture in this patient?

A. Atrophic gastritis
B. Chronic lymphocytic leukemia
C. Coronary atherosclerosis
D. Primary biliary cirrhosis
E. Pulmonary emphysema

43. A 75-year-old woman lives alone and eats sparingly because of her low fixed retirement income. For the past 2 weeks, she has noticed pain in her right leg. On physical examination, there is marked tenderness to palpation over the lateral aspect of the right shin, a poorly healed cut on the right hand, and a diffuse hyperkeratotic skin rash. A radiograph shows a right tibial diaphyseal subperiosteal hematoma. Laboratory studies show a hemoglobin level of 11.3 g/dL. A deficiency of which of the following nutrients is most likely to explain these findings?

A. Ascorbic acid
B. Folate
C. Niacin
D. Riboflavin
E. Vitamin A
F. Vitamin K
G. Vitamin B12

44. A 55-year-old woman has had worsening problems with memory and the ability to carry out tasks of daily living over the past year. She has had watery diarrhea for the past 3 months. Physical examination shows red, scaling skin in sun-exposed areas. Deep tendon reflexes are normal, and sensation is intact. Which of the following diseases is she most likely to have?

A. Beriberi
B. Cheilosis
C. Hypothyroidism
D. Marasmus
E. Pellagra

45. A 9-month-old infant has failure to thrive following a premature birth with low birth weight. The infant has chronic cholestatic hepatobiliary disease. The infant is now at the 40th percentile for height and the 25th percentile for weight. On examination, he exhibits absent deep tendon reflexes, decreased vibration and pain sensation, muscle weakness, and abnormalities of eye movement. Laboratory studies show hemoglobin, 9.2 g/dL; hematocrit, 27.6%; MCV, 86 μm³; platelet count, 208,000/mm³; WBC count, 6080/mm³; total protein, 6.4 g/dL; albumin, 3.4 g/dL; glucose, 70 mg/dL; and creatinine, 0.3 mg/dL. A deficiency of which of the following vitamins is most likely to contribute to these findings?

A. Vitamin A
B. Vitamin B₁
C. Vitamin B₃
D. Vitamin B₁₂
E. Vitamin C
F. Vitamin E
G. Vitamin K

46. A 52-year-old woman with a long history of ethanol abuse has had increasing congestive heart failure for the past year. For the past month, she has experienced increasing confusion, disorientation, and difficulty ambulating. Physical examination shows nystagmus, ataxia of gait, and decreased sensation in the lower extremities. Laboratory studies show hemoglobin, 13.1 g/dL; hematocrit, 39.3%; MCV, 90 μm³; platelet count, 269,300/mm³; and WBC count 7120/mm³. A long-term dietary deficiency of which of the following nutrients is most likely to produce these findings?

A. Folate
B. Niacin
C. Pyridoxine
D. Riboflavin
E. Thiamine

47. A 41-year-old woman had a chronic cough for 7 months. She had a positive tuberculin skin test result. A chest radiograph showed multiple cavitary lesions in the upper lobes of the right and left lungs. She was given isoniazid, rifampin, pyrazinamide, and ethambutol therapy. Now on physical examination 6 months later, a peripheral neuropathy is observed. Administration of which of the following nutrients would most likely have prevented the neuropathy?

A. Ascorbic acid
B. Calciferol
C. Calcium
D. Cobalamin (vitamin B₁₂)
E. Niacin
F. Pyridoxine
G. Riboflavin

48. An infant born at term has Apgar scores of 8 and 9 at 1 and 5 minutes. The infant appears healthy, but 3 days after birth, there is bleeding from the umbilical cord stump, and ecchymoses are observed over the buttocks. Seizures soon develop. Which of the following nutrients is most likely deficient in this infant?

A. Folic acid
B. Iodine
C. Iron
D. Vitamin E
E. Vitamin K
49 Sir Robert Falcon Scott reaches the South Pole on January 17, 1912, barely 1 month after Roald Amundsen achieves this goal with a more experienced and prepared expeditionary party. Scott’s dejected party must now make the long trip back to their base, but they are weak and running low on supplies, and the weather is unusually cold, even for Antarctica. Finally, they can go no further because of severe storms. Months later, a rescue team finds the bodies of the men. All have a hyperkeratotic, papular rash; ecchymoses; and severe gingival swelling with hemorrhages. Which of the following was most likely a contributing cause of death in these men?

A Beriberi  
B Kwashiorkor  
C Pellagra  
D Pernicious anemia  
E Rickets  
F Scurvy

50 An 18-year-old pregnant woman receives no prenatal care, eats a diet containing mostly carbohydrates and fats, and does not take prenatal vitamins with iron. She feels increasingly tired and weak during the third trimester. The infant is born at 35 weeks’ gestation and is listless during the first week of life. Laboratory studies show markedly decreased serum ferritin levels in the infant and the mother. Which of the following findings from a nutritional deficiency is most likely to be present in both the infant and the mother?

A Dermatitis  
B Diffuse goiter  
C Microcytic anemia  
D Peripheral neuropathy  
E Skeletal deformities  
F Soft tissue hemorrhages

51 An epidemiologic study evaluates the rate of dental caries and tooth abscesses among children living in communities within a metropolitan area. Investigators discover that the rate is high among children living in an upper middle class community, but low in children living in a community below the poverty level. The levels of trace elements in the water supplies for those communities are measured. A higher level of which of the following minerals in the water is most likely to be associated with a lower rate of dental decay among the children living in the poor community?

A Copper  
B Fluoride  
C Iodine  
D Selenium  
E Zinc

52 A poorly funded epidemiologic study is conducted, and the results appear in a publication available at the supermarket checkout counter. The study analyzes the diet of textbook authors. Which of the following is determined to be the most likely dietary deficiency in this population?

A Calcium  
B Chocolate  
C Folate  
D Iron  
E Vitamin C

53 A 55-year-old woman has been steadily gaining weight for the past 30 years. She underwent a cholecystectomy for cholelithiasis 5 years ago. She does not smoke. She is now 164 cm (5 ft 4 in) tall and weighs 126 kg (BMI 47). On physical examination, she has decreased range of motion with pain on movement of the knees. Laboratory studies show a serum glucose level of 176 mg/dL. This patient is at greatest risk of developing which of the following neoplasms?

A Colonic adenocarcinoma  
B Endometrial carcinoma  
C Hepatocellular carcinoma  
D Pulmonary adenocarcinoma  
E Renal cell carcinoma

54 In an epidemiologic study of individuals whose BMI is greater than 35, data on lifestyle and disease patterns are collected. Investigators observe that a subset of obese individuals has a consistently high caloric intake because they lack a feeling of satiety when eating. These individuals have diminished responsiveness of a hypothalamic receptor for which of the following molecules?

A Adenosine  
B Glucagon  
C Glucose  
D Insulin  
E Leptin

55 It is 1:00 am and a hard-working second-year medical student is intent on finishing her pathology reading assignment. Soon she begins to note that her concentration is fading because 7 hours have passed since she had dinner, and she is feeling famished. Having studied the chapter on ischemic heart disease, she decides to be prudent and forgoes her favorite chocolate cookies, and instead devours two apples, gulping them down with a glass of low-fat milk. Of the following substances, which one was most likely to have increased rapidly when she became hungry and decreased promptly after she finished her healthy snack?

A α-MSH  
B Corticotropin-releasing factor (CRF)  
C Ghrelin  
D Leptin  
E Thyrotropin-releasing hormone (TRH)

56 A clinical study of adults with a body mass index of at least 30 is undertaken. About 8% of these individuals do not have hyperphagia, but are found to have normal levels of leptin and ghrelin, along with a diminished basal metabolic rate. A mutation in which of the following genes is most likely present in these individuals?

A OB-R  
B MC4R  
C OB  
D POMC  
E PPARγ
A case-control study of adult men and women is performed to determine the relationship between obesity and cancer. The data indicate an increased risk for cancers of the esophagus and kidney in subjects with a body mass index above 25. Which of the following substances is most likely to contribute to the development of cancer in these subjects?

A. Adiponectin  
B. Aflatoxin  
C. Insulin-like growth factor 1 (IGF-1)  
D. Leptin  
E. Selenium  
F. Trans fats

A 45-year-old man, whose mother, father, brother, and uncle all had a history of heart disease, asks his physician about ways to reduce his risk of developing coronary artery disease. The patient is 171 cm (5 ft 6 in) tall, weighs 91 kg, and has a blood pressure of 125/80 mm Hg. His blood glucose concentration is 181 mg/dL. Which of the following is the best dietary advice to give this patient?

A. Avoid adding salt to food  
B. Drink more water  
C. Increase dietary fiber  
D. Reduce intake of saturated fat  
E. Take vitamin A supplements

A 40-year-old man notes a family history of colon carcinoma. He asks his physician how best to reduce his risk of developing this type of cancer. Which of the following dietary practices should he be advised to follow each day?

A. Consume more beef  
B. Drink a glass of red wine  
C. Eat more vegetables  
D. Have a bowl of ice cream  
E. Reduce intake of chocolate

**ANSWERS**

1. B. The cytochrome P-450-dependent monooxygenase, or mixed function oxidase, system is found in smooth endoplasmic reticulum, particularly in hepatocytes, and normally functions to detoxify endogenous hormones. It also can serve to activate xenobiotics to carcinogens. Biomedical researchers have investigated the potential role of environmental microorganisms of inorganic mercury dumped into bodies of water can lead to accumulation of toxic methyl mercury, which can work its way up the food chain to humans. Flavin-containing monooxygenase found in endoplasmic reticulum can oxidize nicotine. Glucuronidation can convert naphthalene to a carcinogen that causes urinary tract cancers. Reduced glutathione helps to break down free radicals produced by xenobiotics such as cytochrome P-450; xenobiotic metabolism can deplete glutathione and enhance free radical cellular injury. The peroxidase-dependent cooxidation pathway can metabolize 2-naphthylamine to a carcinogen that causes urinary tract cancers.  

PBD9 407–409  BP9 271  PBD8 402–403  BP8 281

2. D. Air pollutants form a toxic soup of chemicals that diminish lung function. They are generally tolerated by healthy persons, but reduce the quality of life for persons with existing respiratory conditions. Decreasing levels of ozone in the upper atmosphere have been linked to fluorocarbon release (from refrigerants) and increase the risk for skin cancers. Vehicular exhaust is the major contributor to nitrogen dioxide and ozone emissions that react to form smog. Argon is a nonreactive noble gas that constitutes 1% of the Earth's atmosphere. Carbon dioxide is a greenhouse gas that is increasingly contributing to global climate change, but more is breathed out than is in the atmosphere. Sulfur dioxide released from industrial processes and burning coal forms acid rain and can contribute to chronic obstructive pulmonary disease.  

PBD9 407–409  BP9 272–273

3. C. Climate change is bringing challenges for public health. In coastal regions, more flooding leads to increased potential for water contamination and gastrointestinal infections. The saltwater kills the mosquitoes, one advantage of the high tide, so arboviral diseases are less likely. The rise in temperature is not enough to significantly affect the cardiovascular system, except in sporadic but increasingly frequent heat waves. The other listed options are not affected.  

PBD9 270  BP9 273  PBD8 411  BP8 282–283

4. B. Heating devices that burn hydrocarbons, such as petroleum products, generate carbon monoxide, which can build up to dangerous levels in unventilated or poorly ventilated houses. Chronic carbon monoxide poisoning produces central nervous system damage. Carbon monoxide binds much more tightly to hemoglobin than does oxygen, resulting in hypoxia. Decreased mental functioning generally begins at carboxyhemoglobin levels greater than 20%, and death is likely at levels greater than 60%. The classic “cherry red” lividity is rare and more typical for severe acute poisoning. Acute exposure to beryllium may cause pneumonitis; chronic exposure results in a sarcoid like pulmonary disease. Nitrous oxide compounds and ozone are found in smog and may cause respiratory discomfort with diminished respiratory function in very young individuals, very old individuals, and individuals with underlying respiratory diseases; but such exposure is not immediately life-threatening. In very high concentrations, oxygen can be toxic to the lungs and can promote diffuse alveolar damage. Sulfur dioxide emissions are a component of smog and promote acid rain; they may increase the risk of chronic bronchitis.  

PBD9 409  BP9 273  PBD8 411  BP8 282–283

5. B. Blame the bus. Carbon monoxide (CO) binds to hemoglobin 200 times more avidly than does oxygen, leading to reduced oxygen saturation; CO binds to cardiac myoglobin even more avidly than to hemoglobin. Tissue hypoxia results in lactic acidosis. Cardiac and skeletal muscle begin to break down, releasing creatine kinase. The brain requires high levels of oxygen delivery, so neurologic findings are often the first signs of CO poisoning. The CO toxicity is exacerbated
by exercise. The other listed options do not account for the decreased oxygen saturation. Aspirin poisoning can lead to metabolic acidosis. Iron toxicity leads to nausea and abdominal pain with fluid loss and hypovolemia, metabolic acidosis, and hyperglycemia. Lead poisoning is usually chronic and results in encephalopathy, anemia, and abdominal pain. Methanol is metabolized to formic acid and formaldehyde, causing a metabolic acidosis and damage to the central nervous system and eye. Organophosphates are irreversible cholinesterase inhibitors that produce acute neurotoxicity.

6 C Old flaking paint that is lead-based has a sweet taste, attracting small children to ingest it. The major risk to children from lead ingestion is neurologic damage. Venous blood lead levels should normally be less than 10 μg/dL. Ethylene glycol is found in antifreeze and can produce acute renal tubular necrosis. Kerosene, a hydrocarbon used in some household heating devices, can generate carbon monoxide fumes if not properly ventilated and can cause gastrointestinal and respiratory toxicity when ingested. Methanol ingestion can cause acute central nervous system depression, acidosis, and blindness. Household bleach (sodium hypochlorite) is a local irritant and is not likely to be found in the living conditions of these children.

7 E The muscarinic effects of acute organophosphate poisoning are counteracted by the atropine. Organophosphates are powerful pesticides with neurotoxicity to humans. In contrast, cyclodienes, such as DDT, have low toxicity to humans, but build up in the environment, working their way up the food chain. Dioxins are defoliants that are potentially immunosuppressive, carcinogenic, and teratogenic. Pyrethrins (derived from chrysanthemum flowers) are pesticides with neurotoxicity when ingested. Methanol ingestion can cause acute central nervous system depression, acidosis, and blindness. House- hold bleach (sodium hypochlorite) is a local irritant and is not likely to be found in the living conditions of these children.

8 A Chronic exposure to arsenic-containing compounds can increase the risk for nonmelanoma skin cancers in a distribution different from those linked to the most common cause, ultraviolet light exposure. Beryllium inhaled as a gas can produce an acute toxicity to the respiratory tract; chronic exposure can produce sarcoid like granulomas. Cadmium toxicity can lead to renal tubular necrosis, osteopenia, and chronic pulmonary disease. Lead toxicity acutely produces renal and gastrointestinal damage, and chronically it causes anemia, neuropathy, and encephalopathy. Mercury poisoning leads to central nervous system damage.

9 E This man has experienced occupational exposure to lead and shows symptoms and signs of lead toxicity. The concentration of zinc protoporphyrin is elevated in chronic lead poisoning, in anemia of chronic disease, and in iron deficiency anemia. Lead interferes with heme biosynthesis and inhibits the incorporation of iron into heme; as a result, zinc is used instead. Hepatic damage with elevation of liver enzymes ALT and AST is not a major feature of lead poisoning, but acute increases in these enzymes could be seen with acetaminophen toxicity. The muscle enzyme creatine kinase is not elevated because muscle is not directly damaged by lead, although a neuropathy can occur. Lead can damage renal tubules and cause renal failure, but specific alterations in electrolytes with decreased calcium are not specific for lead-induced renal failure. Selenium is a micronutrient.

10 C Methyl mercury is toxic to the developing brain. It is incorporated into organic compounds within bacteria and works its way up the food chain to humans. Victims of cadium poisoning primarily exhibit complications involving kidney and lung. Bisphenol A is used in the manufacture of epoxy resins found in nearly every bottle and can; it is an endocrine disruptor. Organochlorines such as the insecticide DDT also get into the food chain and mainly disrupt endocrine metabolism. Non-pesticide organochlorines include polychlorinated biphenyls (PCBs) that can cause chloracne, and an initial target of the environmental movement spearheaded by Pete Seeger sailing the Clearwater along the Hudson River. Vinyl chloride used in the manufacture of polyvinyl chloride plastics is linked to hepatic angiosarcomas.

11 B Asbestos fibers can cause pulmonary interstitial fibrosis with restrictive lung disease, and there is an increased risk of malignancy. Individuals who have been exposed to asbestos and who smoke have a greatly increased incidence of bronchogenic carcinoma. Mesothelioma is uncommon, even in individuals with asbestos exposure, but virtually all occurrences of malignant mesothelioma are in individuals who have been exposed to asbestos. Arsenic exposure is a risk factor for skin cancer. Chronic beryllium exposure may lead to sarcoid like granuloma formation. Chromium exposure increases the risk of carcinomas of the upper respiratory tract and lung. Nickel exposure is associated with cancers of the respiratory tract.

12 E The infiltrative perihilar mass suggests lung cancer. Polycyclic hydrocarbons and nitrosamines, found in tobacco smoke, are the key contributors to the development of lung cancer. The carbon monoxide levels of smokers are increased, but this promotes hypoxemia, not cancer. The nicotine in cigarette smoke has a stimulant effect on the central nervous system, but it does not play a major role in cancer development. Nitrous oxide compounds and ozone are found in smog, but they are not closely associated with the risk of lung cancer. Sili- ca dust exposure slightly increases the risk of lung cancer, most often in individuals with prolonged occupational exposure. Sul- fur dioxide emissions are a component of smog and promote acid rain; they may increase the risk of chronic bronchitis.
13 B A large amount of ethanol ingested over a short time can elevate blood ethanol to toxic levels with CNS depression because the alcohol dehydrogenase in liver metabolizes ethanol by zero-order kinetics. The combination of acetaminophen and ethanol increases the likelihood of hepatic toxicity with hepatic necrosis and acute liver failure. Pancreatitis is a potential complication of chronic ethanol abuse. Cirrhosis is a long-term complication of chronic ethanolism. Hematemesis from gastritis and gastric ulceration is more typically seen with chronic ethanolism, and variceal bleeding is a complication of hepatic cirrhosis. Wernicke disease occurs rarely, even in alcoholics, and probably results from concomitant chronic thiamine deficiency.

PBD9 417–419, 422 BP9 280–281 BPD8 412–414 BP8 290–292

14 A Acetaldehyde contributes to acute toxicity of ethanol. In some populations, such as Asian, there is a variant form of acetaldehyde dehydrogenase that is less effective at metabolizing acetaldehyde to acetic acid. α-Tocopherol is vitamin E, an antioxidant that protects cells from injury. Formic acid is one of the toxic metabolites, along with formaldehyde, that are produced when methanol is metabolized by alcohol dehydrogenase. Glutathione is an antioxidant that protects cells from free radical injury. Hydrogen peroxide is metabolized to water in peroxisomes as ethanol is converted to acetaldehyde by alcohol dehydrogenase.

PBD9 418–419 BP9 280–281 BPD8 413–414 BP8 290–292

15 E Hormone replacement therapy (HRT) increases the risk of thromboembolic disease, as is the case with oral contraceptive therapy. HRT was long considered to have a protective effect against cardiovascular disease because exogenous estrogens increase HDL and decrease LDL. This is not true of all women, however, and progestins tend to have the opposite effect. The risk of cervical carcinoma is more closely related to a lifestyle that increases the likelihood of human papillomavirus infection. Inflammatory bowel disease is not associated with HRT. Hepatic cirrhosis in men may cause decreased degradation of circulating estrogens, leading to testicular atrophy. Pulmonary emphysema typically is related to cigarette smoking, not to HRT.

PBD9 421 BP9 282–283 BPD8 414–415 BP8 292

16 E With low estrogen–containing oral contraceptives currently in use, there is no increase in risk for coronary atherosclerosis or myocardial infarction in nonsmoking women younger than 45 years. The risk is increased, however, in women older than 35 years who smoke. Smoking during pregnancy increases the likelihood for low-birth-weight infants. Oral contraceptives do not increase the risk for breast cancer, and they decrease the likelihood of ovarian and endometrial cancer; the risk for these latter two cancers is increased by postmenopausal hormone replacement therapy (HRT). The risk for cholecystitis increases with postmenopausal HRT. There is no evidence for risk of dementia with oral contraceptives or HRT.

PBD9 421 BP9 283 BPD8 415 BP8 292

17 D Cocaine is a powerful vasoconstrictor, and the cardiac complications of its use include acute arterial vasoconstriction with ischemic injury and arrhythmias. Atherosclerosis, affecting small, peripheral branches of the coronary arteries, can be marked with chronic use. Amphetamines may also induce cardiac arrhythmias. Barbiturates are depressants and can cause respiratory failure. Benzodiazepines can produce respiratory failure. Acute ethanol poisoning can cause central nervous system depression and coma. Heroin overdoses can be accompanied by pulmonary edema and respiratory failure. Marijuana is a minor tranquilizer; no serious physiologic effects are associated with its use.


18 A Acetaminophen toxicity leads to hepatic necrosis, indicated by rising ALT and AST levels. If death is not immediate, hyperbilirubinemia also can be seen. N-Acetylcysteine augments glutathione by contributing a sulfhydryl group for binding to toxic metabolites. Elevated serum amylose is seen in pancreatitis. Elevated serum creatinine is seen with injury to skeletal and cardiac muscle. Ketonuria is a feature of absolute insulin deficiency in diabetes mellitus; it also is a feature of starvation. Hypokalemia can be a feature of renal diseases and of glucocorticoid deficiency.

PBD9 422 BP9 284 BPD8 416–417 BP8 294, 297

19 C Cocaine is a powerful vasoconstrictor and has various vascular effects, including ischemic injury to the nasal septum following the route of administration, which is inhalation in this case. Many complications result from the cardiovascular effects, which include arterial vasoconstriction with ischemic injury to the heart, arrhythmias, and central nervous system (CNS) hemorrhages. Hyperthermia is another complication with “excited delirium” in some cases of cocaine intoxication. The callus is caused by flicking a lighter for use of a crack cocaine pipe. Amphetamines are CNS stimulants. Barbiturates are CNS depressants. Acute ethanolism may lead to CNS depression, but it does not have serious immediate cardiac effects. Opiates can depress CNS and respiratory function, and heroin may produce acute pulmonary edema. Marijuana (with active agent delta-9-tetrahydrocannabinol) has no serious acute toxicities. Phencyclidine (PCP) produces an acute toxicity that mimics psychosis.


20 B Chronic aspirin (acetylsalicylic acid) toxicity (<3 g/day) can result in various neurologic problems. Aspirin also inhibits platelet function by suppressing the production of thromboxane A₂, promoting bleeding. Acute aspirin toxicity can cause metabolic acidosis with respiratory compensation. Acute toxicity initially produces a pure respiratory alkalosis due to stimulation of medullary respiratory centers. Increasing absorption of the salicylic acid drives metabolic acidosis, further exacerbated by poisoning of mitochondrial oxidative phosphorylation leading to anaerobic glycolysis and lactic acidosis. Acetaminophen toxicity primarily targets the liver. Chlorpromazine causes cholestatic jaundice. Iron toxicity typically has a component of gastroenteritis with fluid and
blood loss with hypovolemia. Morphine and other opiates tend to produce respiratory depression with respiratory acidosis. Quinidine therapy may lead to hemolytic anemia. Tetracycline can discolor the teeth of children; it can lead to photo toxicity with sunburn.

21 D Heroin is an opiate narcotic that is a derivative of morphine. Opiates are central nervous system (CNS) depressants, and overdoses are accompanied by respiratory depression, convulsions, and cardiac arrest. The typical mode of administration is by injection. An infection, such as an endocarditis explaining his heart murmur, often results from such use because nonsterile injection technique is employed. Cocaine is most often inhaled rather than injected, and acutely produces a state of excited delirium. Ethanol is typically ingested and, in excess, can lead to coma and death. Flurazepam is most often ingested; excessive use can lead to respiratory depression. Meperidine is an analgesic that can cause respiratory depression and bradycardia in overdosage. Phencyclidine (PCP) is a schizoprenomimetic and is usually ingested; users have a history of erratic behavior. Lysergic acid (LSD) is a hallucinogen.

22 D The medical uses of marijuana include intractable nausea and glaucoma. Marijuana, with the active substance tetrahydrocannabinol (THC), has a sedative effect on the central nervous system. Barbiturates are sedatives, but have no effect on severe nausea. Cocaine, methylenidate, and methamphetamine are stimulants. Heroin is an opioid; morphine sulfate, an opioid derivative, has a beneficial effect in treating intractable pain in cancer patients. Phencyclidine (PCP) has a schizoprenomimetic effect.

23 A A scraping injury produces an abrasion, but does not break through the skin. A burn injury causes coagulative necrosis without mechanical disruption. A contusion is a bruise with extravasation of blood into soft tissues. Lacerations break the skin or other organs in an irregular pattern. An incised wound is made with a sharp instrument such as a knife, leaving clean edges.

24 B A blow with a blunt object produces soft tissue hemorrhage without breaking the skin. An abrasion scrapes away the superficial epidermis. A laceration is an irregular tear in the skin or other organ. An incised wound is made by a sharp object and is longer than a puncture, which has a rounded outline and is deeper than it is wide.

25 C The figure shows the entry site of a gunshot wound made at close range. There is a sharply demarcated skin defect, and the surrounding skin shows some stippling of unburned gunpowder. A blast injury can produce various findings, depending on what struck the body or what the body struck, but the focal stippling seen in the figure is not likely to be present. An electrocution injury often produces minimal findings; the site of entry of the current may be quite small. An incised wound is made by a sharp instrument that produces clean-cut edges, whereas a laceration is an irregular disruption in the skin and has torn edges.

26 B The patient has a full-thickness burn injury to the arms, but only partial-thickness burns of other areas. In a full-thickness burn, all structures from which reepithelialization could occur are lost, including dermal appendages such as sweat glands and hair follicles. Loss of only the epidermal basal layer would not prevent reepithelialization from the skin appendages. Fibroblasts can produce more collagen, although elastic fibers do not regenerate; this is why burned skin tends to lose its elasticity, requiring additional grafting procedures in growing children. The superficial layer of keratin serves a protective function. Its loss in burns increases fluid and electrolyte loss, but it reforms if reepithelialization occurs. Blood monocytes can migrate into tissues to become macrophages, which help in remodeling the damaged tissues. Loss of the nerve endings in full-thickness burns leads to the loss of sensation noted on physical examination in this patient, but this process does not govern reepithelialization.

27 A The figure shows superficial tears in the epidermis, with underlying superficial dermal hemorrhage, typical of abrasions made by a scraping type of injury. A contusion is a bruise characterized by breakage of small dermal blood vessels and bleeding. Hypothermia (exposure to cold) has no characteristic appearance. Kwashiorkor, resulting from lack of protein in the diet, may lead to flaking of the skin and irregular pigmentation. A laceration is a cut with torn edges. Thermal burns may produce erythema, blistering, and cracking of skin, but not a scraped appearance.

28 B Heat exhaustion results from failure of the cardiovascular system to compensate for hypovolemia caused by body water depletion. It is readily reversible by replacing lost intravascular volume. Vigorous exercise with electrolyte loss can produce muscle cramping typical of heat cramps. Heat stroke is associated with organ damage when core body temperature rises above 36° C. Malignant hyperthermia occurs from a metabolic derangement such as thyroid storm, or from administration of certain drugs such as succinylcholine; it may also result from excited delirium associated with cocaine use. Inhalation injury is seen when a fire occurs in an enclosed space, and hot, toxic gases are inhaled.
29 E Electric current, especially alternating current, disrupts nerve conduction and electrical impulses, particularly in the heart and brain. Heart function is often affected when electrical current passes through the body to ground. This can lead to severe arrhythmias, especially ventricular fibrillation. These are immediate effects. The amount of tissue injury from standard (U.S.) household current is generally not great, and there may be just a small thermal injury at the site of entry or exit of the current on the skin. The other listed options are not a direct consequence of an electrical injury.

PBD9 427–428 BP9 289 PBD8 422 BP8 299–300

30 E Fires in enclosed spaces produce hot, toxic gases. The inhalation of these gases can lead to death from pulmonary edema even when there is no injury from flames. An acute myocardial infarction is possible, but not probable in a woman of this age. Cerebral hemorrhage is more likely due to trauma, such as a fall or blow to the head. Malignant hyperthermia, which occurs when core body temperature is greater than 40° C, is produced by metabolic disorders such as hyperthyroidism and drugs such as succinylcholine and cocaine. Infections occur days to weeks after a burn injury because of the loss of an epithelial barrier to infectious agents.

PBD9 427 BP9 288–289 PBD8 421 BP8 298

31 C The short story “The Little Match Girl” by Hans Christian Andersen provides a description of hypothermia. Slowly developing and prolonged exposure to cold leads to peripheral vasoconstriction along with edema caused by increased vascular permeability. As her core body temperature dropped to 32.2° C (90° F), she had involuntary shivering and then envisioned her loving grandmother just before loss of consciousness, followed by bradycardia. Death ensues before an inflammatory response to cellular injury can occur. Apoptosis is single cell necrosis, and the hypothermia described in the story would affect large regions of the body. Though there is vascular permeability, vascular integrity prevents hemorrhage. Soft tissues with low metabolic rate are resistant to ischemia and infarction, so findings of coagulative necrosis, such as nuclear fragmentation, are unlikely to be seen acutely.

PBD9 427 BP9 289 PBD8 422 BP8 299

32 C Exposure to sunlight is a risk factor for developing malignancies involving the skin (basal cell carcinoma, squamous cell carcinoma, and malignant melanoma). Ultraviolet (UV) rays, mainly the UVB component, are the major causative agent of these malignancies. Infrared radiation causes mainly thermal injury. Visible light has minimal effects. The ambient x-radiation and gamma rays that filter through the earth’s atmosphere are minimal and have no significant health effects. X-radiation from radiologic imaging is certainly a concern when patients receive multiple imaging studies, but the cumulative dosages remain small.

PBD9 1155 BP9 863–864 PBD8 404 BP8 309–312

33 E Therapeutic doses of radiation can cause acute vascular injury, manifested by endothelial damage and an inflammatory reaction. With time, these vessels undergo fibrosis and severe luminal narrowing. There is ischemia of the surrounding tissue and formation of a scar. The radiation used in therapeutic dosages is carefully delivered in a limited field to promote maximal tumor damage, while reducing damage to surrounding tissues. Whole-body irradiation affects the bone marrow, gonads, gastrointestinal tract, and brain, but therapeutic radiation is carefully focused on the neoplasm to prevent widespread tissue damage.

PBD9 428–432 BP9 290–292 PBD8 423–425 BP8 300–303

34 B These firemen are true heroes whose selfless actions prevented a much greater disaster. The cerebral syndrome occurs within hours in individuals exposed to a massive total-body radiation dose. Doses of >2 Sv (>200 cGy) can be fatal because of injury to radiosensitive marrow and the gastrointestinal tract, but death occurs after days to weeks, not hours. Cardiac and skeletal muscle tissue is relatively radioresistant. Early findings in radiation-induced lung injury include edema; interstitial fibrosis develops over years in individuals who survive the injury.


35 E Body weight less than 60% of normal with muscle wasting is consistent with marasmus, a form of protein-energy malnutrition that results from a marked decrease in total caloric intake. In kwashiorkor, protein intake is reduced more than total caloric intake, and body weight is usually 60% to 80% of normal. Hypoalbuminemia is a key laboratory finding. Malignancies can promote wasting, but not to this degree. This child’s problems are far more serious than a single vitamin deficiency; a lack of folate could account for the child’s anemia, but not for the wasting. Bulimia is an eating disorder of adolescents and adults that is characterized by binge eating and self-induced vomiting. Lead poisoning can lead to anemia and encephalopathy, but it does not cause severe wasting.

PBD9 433–434 BP9 294 PBD8 428–429 BP8 304–306

36 A The decreased food intake from self-imposed dieting in a woman can lead to changes such as hormonal deficiencies (e.g., follicle-stimulating hormone, luteinizing hormone, thyroxine). The result is diminished estrogen synthesis, which promotes osteoporosis, as in the postmenopausal state. Bulimia with bingeing and purging can be accompanied by electrolyte disturbances, and weight tends to be maintained in most cases. Kwashiorkor is a disease mainly of children who have reduced protein intake. Rickets is a specific deficiency of vitamin D that causes skeletal deformities in children. Scurvy, which results from vitamin C deficiency, does not affect hormonal function.

PBD9 435 BP9 295–296 PBD8 427 BP8 304, 306

37 B The findings are consistent with kwashiorkor, a nutritional disorder predominantly of decreased protein in the diet. Hypoalbuminemia is characteristic of this condition.
Abetalipoproteinemia is a rare disorder that causes vitamin E deficiency. Hypocalcemia can occur as a consequence of vitamin D deficiency. Hyperglycemia occurs in diabetes mellitus; the wasting associated with this disease affects adipose tissue and muscle, and edema is not a feature. Megaloblastic anemia is a feature of specific deficiencies of vitamin B₁₂ or folate.

**38 A** Vitamin A is important in maintaining epithelial surfaces. Deficiency of this vitamin can lead to squamous metaplasia of respiratory epithelium, predisposing to infection. Increased keratin buildup leads to follicular plugging and papular dermatosis. Desquamated keratinaceous debris in the urinary tract forms the nidus for stones. Ocular complications of vitamin A deficiency include xerophthalmia and corneal scarring, which can lead to blindness. Vitamin B₁ (thiamine) deficiency causes problems such as Wernicke disease, neuropathy, and cardiomyopathy. Vitamin D deficiency in children causes rickets, characterized by bone deformities. Vitamin E deficiency occurs rarely; it causes neurologic symptoms related to degeneration of the axons in the posterior columns of the spinal cord. Vitamin K deficiency can result in a bleeding diathesis.

**39 C** Rickets, which is caused by vitamin D deficiency, is characterized by skeletal deformity such as the bowing of the legs seen in this boy. Lack of bone mineralization (osteopenia) leads to this deformation. Beriberi, from thiamine deficiency, can result in heart failure and peripheral edema. A diet containing insufficient protein can result in kwashiorkor, characterized by areas of flaking, depigmented skin. Pellagra, resulting from niacin deficiency, is characterized by dermatitis in sun-exposed areas of skin. Scurvy, resulting from vitamin C deficiency, can produce bone deformities, particularly at the epiphyses, because of abnormal bone matrix, not abnormal calcification. The absence of hemorrhages in this child makes this unlikely, however.

**40 D** Vitamin A is essential to maintain epithelia. The lack of vitamin A affects the function of lacrimal glands and conjunctival epithelium, promoting keratomalacia. Dr. Alfred Sommer’s research convinced UNICEF that it would cost just pennies per child to eliminate vitamin A deficiency in 250 million children on earth. Iron is essential for production of heme, which is needed to manufacture hemoglobin in RBCs. Niacin is involved with nicotinamide in many metabolic pathways, and deficiency leads to diarrhea, dermatitis, and dementia. Dietary protein is essential for building tissues, particularly muscle, but it has no specific effect in maintaining ocular structures. Vitamin K is beneficial for synthesis of coagulation factors by the liver to prevent bleeding problems.

**41 A** Vitamin D can be synthesized endogenously in skin with exposure to ultraviolet (UV) light. Together, vitamin D and calcium help build and maintain growing bone. Exercise helps build bone mass, which protects against osteoporosis later in life, particularly in women. Renal function is not greatly affected by environment. There are some deleterious effects on the eye (cataracts) and the skin (cancer, elastosis) from increased exposure to UV radiation in sunlight. Increased air pollution in many cities has led to an increased incidence of pulmonary diseases, and children are particularly at risk.

**42 D** The osteopenia in this patient can result from osteomalacia, the adult form of vitamin D deficiency. Vitamin D is a fat-soluble vitamin, and it requires fat absorption, which can be impaired by chronic cholestatic liver disease, biliary tract disease, and pancreatic disease. Atrophic gastritis affects vitamin B₁₂ absorption. Leukemias do not tend to erode bone. Heart disease caused by atherosclerosis does not affect bone density. Empysema can result in a hypertrophic osteoarthropathy, but not osteopenia.

**43 A** Signs and symptoms of scurvy can be subtle. The diet must contain a constant supply of vitamin C (ascorbic acid) because none is produced endogenously. Older individuals with an inadequate diet are as much at risk as younger individuals. Folate deficiency can lead to anemia, but it does not cause capillary fragility with hematoma formation or skin rash. Niacin deficiency can lead to an erythematous skin rash in sun-exposed areas, but not to anemia. Riboflavin deficiency can lead to findings such as glossitis, cheilosis, and neuropathy. Vitamin A deficiency can produce a skin rash, but it does not cause anemia. Vitamin K is important in maintaining proper coagulation, but a deficiency state is not associated with anemia or skin rash.

**44 E** Pellagra is caused by a deficiency of niacin. The classic presentation includes the “3 D’s”: diarrhea, dermatitis, and dementia. Beriberi, resulting from thiamine (vitamin B₁) deficiency, can lead to heart failure, neuropathy, and Wernicke disease. Cheilosis describes the fissuring at the corners of the mouth that accompanies riboflavin (vitamin B₂) deficiency. In hypothyroidism, which could be due to iodine deficiency, the skin tends to be coarse and dry. Marasmus describes the severe wasting that occurs in individuals with a diet that is markedly deficient in all nutrients.

**45 F** Vitamin E deficiency is uncommon, but it may be seen in low-birth-weight infants with poor hepatic function and fat malabsorption. The neurologic manifestations are similar to those seen in vitamin B₁₂ deficiency; affected infants may have anemia, but it is not of the megaloblastic type. Vitamin A deficiency in infants and children can lead to blindness from keratomalacia. It is the most common cause of preventable blindness in this population group. Vitamin B₃ (niacin) (thiamine) deficiency can lead to beriberi. Vitamin B₃ (niacin)
deficiency can lead to pellagra. Vitamin C deficiency leads to scurvy, which can be accompanied by anemia from bleeding and from decreased iron absorption. Vitamin K deficiency leads to bleeding problems.

**46** E Individuals with a history of chronic alcoholism are often deficient in thiamine and other nutrients (ethanol provides empty calories). Thiamine deficiency can lead to neuropathy, cardiomyopathy, and Wernicke disease. Alcoholic individuals often have folate deficiency, with resultant macrocytic anemia, but this finding is not present here. Niacin deficiency leads to pellagra. Pyridoxine and riboflavin deficiencies can lead to neuropathy, but do not produce cerebral findings.

**47** F Isoniazid is a pyridoxine (vitamin B₆) antagonist. Individuals receiving isoniazid therapy for tuberculosis may need supplementation to prevent vitamin B₆ deficiency. Ascorbic acid (vitamin C) is antiscorbutic (prevents scurvy). Calciferol (vitamin D) helps maintain calcium levels. Calcium intake helps maintain bone mass and serum calcium level; hypocalcemia can lead to neural excitability with muscular contractions. Cobalamin (vitamin B₁₂) deficiency may produce a macrocytic anemia and a peripheral neuropathy, but it does not result from isoniazid therapy. Niacin deficiency causes pellagra (diarrhea, dermatitis, dementia). Riboflavin deficiency may produce neuropathy, glossitis, and cheilosis.

**48** E Coagulation factors II, VII, IX, and X synthesized by the liver require vitamin K for their production. Hemorrhagic disease of the newborn can occur in infants who lack sufficient intestinal bacterial flora to produce this nutrient. Breastfeeding transiently potentiates this effect, because *Lactobacillus* found in breast milk does not synthesize vitamin K. Routine intramuscular injection of vitamin K soon after birth prevents this complication. Iron deficiency leads to anemia, not to bleeding. Vitamin E is an antioxidant and is rarely deficient to a degree that would cause serious illness. Folic acid helps to prevent macrocytic anemia. Iodine is needed in small quantities for thyroid hormone synthesis.

**49** F Humans do not generate vitamin C endogenously, so they must have a continuous dietary supply. The lack of fresh fruits and vegetables containing vitamin C led to scurvy in many sailors and explorers in centuries past. Beriberi leads to heart failure and results from thiamine deficiency. Kwashiorkor results from protein deficiency. Pellagra, characterized by the “3 D’s” of diarrhea, dermatitis, and dementia, is seen in niacin deficiency. Pernicious anemia from vitamin B₁₂ deficiency can be complicated by neurologic deterioration in severe cases. Rickets is seen in children who are deficient in vitamin D.

**50** C Iron deficiency, which gives rise to microcytic anemia, is common in women of reproductive age because of menstrual blood loss and in children with a poor diet. During pregnancy, women have greatly increased iron needs. Low serum ferritin is indicative of iron deficiency. Dermatitis can be seen in pellagra (niacin deficiency). Goiter results from iodine deficiency, but this is a rare occurrence today because of newborn testing and widely available foods with iodine. Peripheral neuropathy is more characteristic of beriberi (thiamine deficiency) and deficiencies in riboflavin (vitamin B₂) and pyridoxine (vitamin B₆). Bowing of the long bones and epiphyseal widening can be seen in rickets (vitamin D deficiency). Soft tissue hemorrhages can be seen in scurvy (vitamin C deficiency).

**51** B Water in some areas naturally contains fluoride, and dental problems in children are fewer in these areas because tooth enamel is strengthened. Fluoride can be added to drinking water, but opposition to this practice, from ignorance or fear, is common. Copper deficiency can produce neurologic defects. Iodine deficiency can predispose to thyroid goiter. Selenium is a trace mineral that forms a component of glutathione peroxidase; deficiency may be associated with myopathy and heart disease. Serious illnesses from trace element deficiencies are rare. Zinc is a trace mineral that aids in wound healing; a deficiency state can lead to stunted growth in children and a vesicular, erythematous rash.

**52** B This vignette is just as imaginative as those appearing in many public media sources, so beware the claims and apply principles of evidence-based medicine. Some would agree that there is never quite enough chocolate, and much of the world’s population must get by without it. However, countries with the highest chocolate consumption have produced the most Nobel laureates! Serious dietitians would probably choose option A (iron deficiency), which is a deficiency most likely to be seen in menstruating women, in pregnant women, and in children. Calcium is most important in growing children for building bones. Folate deficiency leads to macrocytic anemia and is most likely to occur in adults with an inadequate diet, such as individuals with chronic alcoholism. Vitamin C deficiency occurs in individuals who do not eat adequate amounts of fresh fruits and vegetables.

**53** B This patient is morbidly obese. The extra weight puts a strain on joints, particularly the knees, increasing the risk for osteoarthritis. Although the overall risk of cancer increases with obesity, the relationship between endometrial carcinoma and obesity is well established. About 80% of individuals with type 2 diabetes mellitus are obese. The relationship of diet and obesity to colon cancer is not as well established. Worldwide, most hepatocellular carcinomas arise in individuals infected with hepatitis B; chronic alcoholism also is a risk factor. Pulmonary adenocarcinoma is the least likely
bronchogenic cancer to be associated with smoking. Some renal cell carcinomas are associated with smoking.

54 E Leptin signaling from adipocytes that have taken up an adequate supply of fatty acids ordinarily feeds back to the hypothalamus, which decreases synthesis of neuropeptide Y. This neurotransmitter acts as an appetite stimulant, and a decrease in its synthesis causes satiety. Adenosine is a nucleoside used to treat cardiac dysrhythmias. Glucagon opposes insulin by increasing hepatic glycogen storage. An increasing blood glucose level results in an increased release of insulin to promote glucose uptake into connective tissues, muscle, and adipose tissue.

55 C Appetite and satiety are controlled by a complex system of short-acting and long-acting signals. The levels of ghrelin produced in the stomach increase rapidly before every meal and decrease promptly after the stomach is filled. Leptin released from adipocytes exerts long-term control by activating catabolic circuits and by inhibiting anabolic circuits. α-MSH is an intermediate in leptin signaling. TRH and CRF are among the efferent mediators of leptin signaling, and they increase energy consumption.

56 B There are “obesity” genes that may play a role in metabolic pathways. About 5% to 8% of obese adults have a mutation in the MC4R gene, and even though there are abundant fat stores and plenty of leptin, the lack of MC4R to drive energy consumption leads to weight gain. Leptin is the product of the ob gene, and mutations reduce leptin levels that signal satiety, but such mutations are rare. Mutations in OB-R encoding the leptin receptor are seen in about 3% of cases of early-onset obesity with hyperphagia in children. POMC is in a catabolic pathway stimulated by leptin, but mutations are rare, and affected individuals typically have childhood onset with hyperphagia. Peroxisome proliferator-activated receptor gamma (PPARγ) is stimulated by the thiazolidinedione drugs and leads to a reduction in free fatty acids, reduction in resistin, and decreased insulin resistance.

57 C IGF-1 increases in response to the hyperinsulinemia of obese persons, who are also more likely to have metabolic syndrome and type 2 diabetes mellitus. IGF-1 promotes cell growth as well as increased synthesis of estrogens and androgens that favor neoplastic transformation in cells. Adiponectin that is elaborated by adipocytes acts as an insulin-sensitizing agent that prevents hyperinsulinemia. Aflatoxin from growth of Aspergillus on foods such as cereals acts as a carcinogen via mutation of the TP53 gene. Leptin normally decreases when fat stores are high, and derangements in its receptor may underlie some forms of obesity. Selenium is a trace metal thought to have antioxidant properties protective against cancer. Trans fats are derived from artificial hydrogenation of dietary fats and are atherogenic by increasing LDL cholesterol while decreasing HDL cholesterol.

58 D These findings suggest a diagnosis of diabetes mellitus. The patient is obese and most likely has type 2 diabetes mellitus. Type 1 and type 2 diabetes mellitus greatly increase the risk of early and accelerated atherosclerosis. Decreasing total caloric intake, particularly saturated fat, helps reduce the risk of coronary artery disease. Vegetable and fish oils are preferable to animal fat as sources of dietary lipid for prevention of atherosclerosis. Reducing dietary sodium helps to decrease blood pressure. Increased fluid intake aids renal function. Dietary fiber helps to reduce the incidence of diverticulosis. Vitamin A has no significant effect on atherogenesis.

59 C More fruits and vegetables are recommended in the diet to help prevent colon cancer. Vitamins C and E have an antioxidant and antimutagenic effect. Red wine in moderation may have a beneficial antiatherogenic effect. Ice cream can include animal fat that may promote cancer, as would the animal fat of beef. Chocolate includes vegetable fat, which is not as harmful as animal fat, and persons consuming chocolate perform at a higher level.
1. A 31-year-old woman, G3, P2, has had an uneventful pregnancy except for lack of any fetal movement. She has a spontaneous abortion at 20 weeks’ gestation and delivers a stillborn boy. On examination at birth, the fetus has an abdominal wall defect lateral to the umbilical cord insertion; a short umbilical cord; marked vertebral scoliosis; and a thin, fibrous band constricting the right upper extremity. A radiograph is shown in the figure. None of the woman’s other pregnancies, which ended in term births, were similarly affected. Which of the following is the most likely cause of these findings?

   A  Congenital cytomegalovirus infection
   B  Early amnion disruption
   C  Oligohydramnios
   D  Maternal fetal Rh incompatibility
   E  Trisomy 18

2. A 16-year-old primigravida in her 18th week of pregnancy has not felt any fetal movement, and an ultrasound is performed. The amniotic fluid index is markedly decreased. Both fetal kidneys are cystic, and one is larger than the other. There is no fetal cardiac activity. The pregnancy is terminated, and a fetal autopsy is performed. Findings include multicystic renal dysplasia, hemivertebra, anal atresia, tracheoesophageal fistula, and lungs that are equivalent in size to 14 weeks’ gestation. Which of the following errors in morphogenesis most likely led to the appearance of the fetal lungs?

   A  Agenesis
   B  Aplasia
   C  Deformation
   D  Disruption
   E  Malformation
   F  Teratogenesis
3. A 31-year-old woman, G1, P0, has noticed very little fetal movement during her pregnancy. At 36 weeks’ gestation, she gives birth to an infant with the facial features and positioning of extremities shown in the figure. Soon after birth, the infant develops severe respiratory distress. Which of the following conditions affecting the infant best explains these findings?

A. Congenital rubella infection
B. Bilateral renal agenesis
C. Maternal diabetes mellitus
D. Hyaline membrane disease
E. Trisomy 13

4. A 28-year-old high school teacher, who is pregnant, presents with severe acne on the face and is concerned that these will disfigure her face. Before she visited the doctor, she does an Internet search and finds that a compound related to Vitamin A has proven effective in the treatment of acne. However, her obstetrician is reluctant to prescribe it because he thinks that it may cause fetal malformations. The teratogenic effect of this medication is most likely related to which of the following processes?

A. Disruption of the pattern of expression of homeobox genes
B. Increase in the risk of maternal infections
C. Increase in the likelihood of aneuploidy during cell division
D. Promotion of abnormal development of blood vessels in the placenta
E. Reduction in the resistance of the fetus to transplacental infections

5. A healthy, 44-year-old woman, G2, P1, has a screening ultrasound at 18 weeks’ gestation that shows no fetal anomalies. There is decreased fetal movement at 32 weeks’ gestation, and ultrasound shows fetal growth restriction with relative sparing of the fetal head. The placenta appears normally positioned in the lateral fundus, but appears small, and the amniotic fluid index is reduced. Maternal blood pressure is normal. Which of the following conditions is most likely to be present?

A. Uteroplacental insufficiency
B. Congenital Treponema pallidum infection
C. Galactosemia
D. Preeclampsia
E. Rh incompatibility
F. Trisomy 21

6. A 22-year-old primigravida gives birth to a boy at 38 weeks’ gestation. On physical examination, the infant appears normal except for an abnormal right hip click with the Ortolani maneuver. Ligamentous laxity of the hip is noted. The mother asks if there is any risk that future children will be born with a similar malformation. What is the most likely recurrence risk for future pregnancies?

A. <1%
B. 5%
C. 25%
D. 50%
E. 90%

7. A term infant has initial Apgar scores of 8 and 10 at 1 and 5 minutes. On auscultation of the chest, a heart murmur is audible. There is hepatosplenomegaly. Cataracts of the crystalline lens are noted. The infant is at the 30th percentile for height and weight. Echocardiography shows a patent ductus arteriosus. Which of the following events is the most likely risk factor for the findings in this infant?

A. Congenital rubella infection
B. Dietary folate deficiency
C. Dispermy at conception
D. Erythroblastosis fetalis
E. Maternal thalidomide use
F. Paternal meiotic nondisjunction

8. In the year 1000 CE, an infant with difficulty nursing since birth is brought to Abul Qasim al-Zahrawi, who works in Cordoba, Andalusia (present-day Spain). The boy infant is the product of a normal term pregnancy. On physical examination the only abnormality is the lack of fusion between the lateral nasal prominence and the maxillary prominence. The cleft is repaired. There is no family history of birth defects, and the mother’s other two children are healthy with no apparent abnormalities. This clinical picture is most likely the result of which of the following conditions?

A. Chromosomal anomaly
B. Early amnion disruption
C. Maternal malnutrition
D. Multifactorial inheritance
E. Single gene defect
F. Teratogenicity
9. A newborn boy delivered at 38 weeks is small for gestational age. Physical examination shows microcephaly, frontal bossing, long and narrow forehead, hypotelorism, maxillary and mandibular hypoplasia, narrow palpebral fissures, thin elongated philtrum, vermilion border of the upper lip, dental malocclusion, saddle nose, tooth enamel hypoplasia, and uvular hypoplasia. Ocular problems include microphthalmia, corneal clouding, coloboma, nystagmus, strabismus, and ptosis. A systolic murmur is heard on auscultation, and echocardiography shows a membranous ventricular septal defect. Which of the following conditions is most likely to produce these findings?

A. Congenital rubella  
B. Fetal alcohol syndrome  
C. Maternal diabetes mellitus  
D. Placenta previa  
E. Trisomy 21

10. A 25-year-old woman is G5, P0, Ab4. All of her previous pregnancies ended in spontaneous abortion in the first or second trimester. She is now in the 16th week of her fifth pregnancy and has had no prenatal problems. Laboratory findings include maternal blood type of A positive, negative serologic test for syphilis, and immunity to rubella. Which of the following laboratory studies would be most useful for determining a potential cause of recurrent fetal loss in this patient?

A. Amniocentesis with chromosomal analysis  
B. Genetic analysis of the CFTR gene  
C. Maternal serum antibody screening  
D. Maternal serum α-fetoprotein determination  
E. Maternal serologic test for HIV

11. A 17-year-old primigravida gives birth at 37 weeks’ gestation. Her Hgb A1C is 6%. The birth weight is 2350 g. On physical examination, the infant is normally developed with no anomalies. On prenatal course, the infant has multiple congenital abnormalities including single umbilical artery, cleft lip, and ptosis. A systolic murmur is heard on auscultation, and echocardiography shows a membranous ventricular septal defect. Which of the following is the most likely risk factor for these findings?

A. Infection  
B. Gestational diabetes  
C. Single gene defect  
D. Teratogen  
E. Unknown

12. A 25-year-old primigravida has an uncomplicated prenatal course. She gives birth to a 4500-g boy whose Apgar scores are 8 and 10 at 1 minute and 5 minutes. Shortly after birth, he develops irritability with seizure activity. On examination, the infant is normally developed with no anomalies. The lungs are clear to auscultation. Laboratory studies show serum Na+, 145 mmol/L; K+, 4.2 mmol/L; Cl-, 99 mmol/L; CO2, 25 mmol/L; urea nitrogen, 0.4 mg/dL; and glucose, 18 mg/dL. Which of the following pathologic findings is most likely to be present in the pancreas of this infant?

A. Acute pancreatitis  
B. Amyloid deposition  
C. Adenocarcinoma  
D. Fatty replacement  
E. Insulitis  
F. Islet hyperplasia

13. A 25-year-old woman, G3, P2, is in her 39th week of pregnancy. She has felt no fetal movement for 1 day. The infant is stillborn on vaginal delivery the next day. On physical examination, there are no external anomalies. Microscopic examination of the placenta shows acute chorioamnionitis. Which of the following infectious agents is most likely responsible for these events?

A. Cytomegalovirus  
B. Herpes simplex virus type 2  
C. Streptococcus agalactiae (group B)  
D. Toxoplasma gondii  
E. Treponema pallidum

14. A 17-year-old primigravida has an uneventful pregnancy until 25 weeks, when she notes absence of fetal movement for a day. On examination, her temperature is 37° C, pulse is 80/min, and blood pressure is 115/80 mm Hg. No fetal heart tones are present. An ultrasound examination shows no apparent fetal abnormalities. The next day she develops premature rupture of fetal membranes, and the next day a stillborn fetus is delivered. Examination of the placenta shows normal size for gestational age, but microscopic analysis shows that neutrophils have infiltrated the chorion and amnion. What is the most likely diagnosis?

A. Cystic fibrosis  
B. Rh incompatibility  
C. Sudden infant death syndrome  
D. Toxemia of pregnancy  
E. Trisomy 16  
F. Ureaplasma urealyticum infection

15. A 33-year-old woman in the 32nd week of pregnancy notices lack of fetal movement for 3 days. On physical examination, no fetal heart tones can be auscultated. The fetus is stillborn. At autopsy, scattered microabscesses are seen in the liver, spleen, brain, and placenta. No congenital anomalies are present. Investigation by Centers for Disease Control (CDC) reveals that similar fetal losses have occurred in the same community for the past 3 months and food contamination is suspected. Congenital infection with which of the following organisms is most likely to produce these findings?

A. Cytomegalovirus  
B. Group B streptococcus  
C. Herpes simplex virus type 2  
D. Listeria monocytogenes  
E. Parvovirus  
F. Toxoplasma gondii

16. A neonate born at 36 weeks’ gestation manifests severe hydrops fetalis, hepatosplenomegaly, generalized icterus, and scattered ecchymoses of the skin. Laboratory studies show a hemoglobin concentration of 9.4 g/dL and platelet count of 67,000/mm³. Ultrasound of the head shows ventricular enlargement. Death occurs 14 days after birth. At autopsy, there is extensive subependymal necrosis, with microscopic evidence of encephalitis. Within the areas of necrosis, there are large cells containing intranuclear inclusions. Congenital infection with which of the following organisms is most likely to produce these findings?

A. Cytomegalovirus  
B. Herpes simplex virus  
C. HIV  
D. Parvovirus  
E. Rubella virus
17 A 31-year-old woman, G3, P2, is in the second trimester. Her prior pregnancies ended with delivery of normal term infants who are still living. She has an ultrasound examination because of lack of fetal movement by 18 weeks, and it shows microcephaly with periventricular leukomalacia and calcifications. Her HIV test is positive; her serologic test for syphilis is negative. TORCH titers are performed on maternal blood:

<table>
<thead>
<tr>
<th></th>
<th>IgG</th>
<th>IgM</th>
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<tbody>
<tr>
<td>CMV</td>
<td>1.2 (&lt;0.9)</td>
<td>0.7 (&lt;0.9)</td>
</tr>
<tr>
<td>HSV type 1</td>
<td>1.2 (&lt;0.9)</td>
<td>0.6 (&lt;0.9)</td>
</tr>
<tr>
<td>HSV type 2</td>
<td>0.6 (&lt;0.9)</td>
<td>0.5 (&lt;0.9)</td>
</tr>
<tr>
<td>Rubella</td>
<td>15 (&lt;10)</td>
<td>0.4 (&lt;0.9)</td>
</tr>
<tr>
<td>Toxoplasma</td>
<td>11 (&lt;9)</td>
<td>3.3 (&lt;0.9)</td>
</tr>
</tbody>
</table>

Which of the following is the most likely risk factor for this fetal infection?
A Ingestion of contaminated meat
B Inhalation of droplet nuclei
C Injection drug use
D Mosquito bite
E Previous blood transfusion
F Sexual intercourse

18 A 20-year-old woman gives birth at term to an infant weighing 1900 g. On physical examination, the infant’s head size is normal, but the crown-heel length and foot length are reduced. There are no external malformations. Throughout infancy, developmental milestones are delayed. Which of the following conditions occurring during gestation would most likely produce these findings?
A Congenital cytomegalovirus
B Down syndrome
C Erythroblastosis fetalis
D Maternal diabetes mellitus
E Pregnancy-induced hypertension

19 A healthy 21-year-old primigravida gives birth at 36 weeks following rupture of membranes. Apgar scores are 8 at 1 minute and 9 at 5 minutes. The infant weighs 2000 g (4.4 lb) and on examination has no congenital anomalies noted. The head size is normal but other body measurements are decreased. Which of the following risk factors during pregnancy most likely led to these findings?
A Cigarette smoking
B Confluent placental mosaicism
C Preeclampsia
D Toxoplasmosis
E Triploidy

20 A 38-year-old primigravida has premature rupture of membranes at 36 weeks’ gestation, necessitating delivery. At birth, the infant is noted to have a two-vessel umbilical cord, a cleft lip, a heart murmur, and spina bifida. Which of the following factors is most likely to increase the risk of hyaline membrane disease in the infant?
A Chorioamnionitis
B Diabetes mellitus, type 1
C Maternal corticosteroid therapy
D Oligohydramnios
E Pregnancy-induced hypertension

21 An infant is born prematurely at 32 weeks’ gestation to a 34-year-old woman with gestational diabetes. On physical examination, the infant is at the 50th percentile for height and weight and there are no congenital anomalies. The infant requires 3 weeks of intubation with positive pressure ventilation and dies of sepsis at 4 months of age. At autopsy, the lungs show bronchial squamous metaplasia with peribronchial fibrosis, interstitial fibrosis, and dilation of airspaces. Which of the following conditions best explains these findings?
A Cystic fibrosis
B Bronchopulmonary dysplasia
C Pulmonary hypoplasia
D Sudden infant death syndrome
E Ventricular septal defect

22 In a study of lung maturation, the amount of surfactant at different gestational ages is measured. Investigators find that the amount of surfactant in the developing lung increases between 26 and 32 weeks’ gestation, with progression of lung architecture to a saccular alveolar configuration. This increase in surfactant is most likely related to which of the following developmental events?
A Apoptosis in interlobular mesenchymal cells
B Development of ciliated epithelium in airspaces
C Differentiation of alveoli from embryonic foregut
D Increased density of pulmonary capillaries
E Maturation of type II alveolar epithelial cells

23 A 25-year-old woman gives birth at 28 weeks’ gestation. The newborn girl has initial Apgar scores of 5 and 6 at 1 and 5 minutes, but within 1 hour, she experiences severe respiratory distress and appears cyanotic. A chest radiograph shows a bilateral ground-glass appearance in the lungs. She is treated with assisted ventilation and nutritional support and seems to improve for 24 hours, but then becomes progressively more cyanotic, develops seizures, and dies 4 days after birth. At autopsy, the newborn’s lungs have the microscopic appearance shown in the figure. Which of the following conditions initiates the development of these findings?
A Congenital toxoplasmosis
B Immaturity of lungs
C Marked fetal anemia
D Maternal toxemia of pregnancy
E Oligohydramnios
A 17-year-old primigravida gives birth at 34 weeks’ gestation to a male infant of low birth weight. The infant is given exogenous surfactant and does not develop respiratory distress. On the third day of life, physical examination reveals hypotension, abdominal distention, and absent bowel sounds, and there is bloody stool in the diaper. A radiograph shows pneumatosis intestinalis and abdominal free air. Surgical removal of the bowel is performed. Which of the following conditions is most likely to be present in this infant?

A. Duodenal atresia  
B. Hirschsprung disease  
C. Meckel diverticulum  
D. Meconium ileus  
E. Necrotizing enterocolitis  
F. Pyloric stenosis

A 19-year-old woman, G2, P1, has a screening fetal ultrasound at 20 weeks’ gestation that shows no abnormalities. Premature labor leads to an emergent vaginal delivery at 31 weeks. Soon after birth, the neonate develops respiratory distress requiring intubation with positive pressure ventilation. Which of the following prenatal diagnostic tests could have best predicted this neonate’s respiratory distress?

A. Chromosomal analysis with karyotyping  
B. Coombs test on cord blood  
C. Genetic analysis for cystic fibrosis gene  
D. Phospholipid level in amniotic fluid  
E. Maternal serum α-fetoprotein determination

An 18-year-old woman gives birth to a term infant after an uncomplicated pregnancy and delivery. Over the first 2 days of life, the infant becomes mildly icteric. On physical examination, there are no morphologic abnormalities. Laboratory studies show a neonatal bilirubin concentration of 4.9 mg/dL. The direct Coombs test of the infant’s RBCs yields a positive result. The infant’s blood type is A negative, and the mother’s blood type is O positive. Based on these findings, which of the following conditions is most likely to be present in this infant?

A. Complete recovery  
B. Failure to thrive  
C. Hemolytic anemia throughout infancy  
D. Kernicterus  
E. Respiratory distress syndrome

A 20-year-old woman, G3, P2, has a screening ultrasound at 18 weeks’ gestation that shows hydrops fetalis but no malformations. The woman’s two previous pregnancies ended at term in live births. The current pregnancy results in a live birth at 36 weeks. Physical examination shows marked hydrops of the neonate and placenta. Laboratory studies show a cord blood hemoglobin level of 9.2 g/dL and total bilirubin concentration of 20.2 mg/dL. Which of the following laboratory findings is most likely to be present in this case?

A. Diminished glucocerebrosidase activity in fetal cells  
B. Elevated maternal serum α-fetoprotein level  
C. Positive Coombs test result on cord blood  
D. Positive maternal hepatitis B surface antigen  
E. Positive placental culture for Listeria monocytogenes

A 23-year-old woman has reduced fetal movement at 19 weeks’ gestation. Fetal ultrasound scan is performed and shows marked soft tissue fluid collections. The maternal Coombs test is negative. TORCH titers are unremarkable. Which of the following additional findings is most likely to be observed with this ultrasound examination?

A. Congenital neuroblastoma  
B. Endocardial cushion defect  
C. Leukomalacia  
D. Low-lying placenta  
E. Splenomegaly

A 34-year-old woman, G4, P3, is large for dates at 16 weeks’ gestation. Fetal ultrasound examination shows one large plethoric twin with hydrops fetalis as well as polyhydramnios and one small anemic twin with oligohydramnios. The maternal Coombs test is negative. What is the most likely explanation for these findings?

A. α-Thalassemia major  
B. Chromosomal aneuploidy  
C. Erythroblastosis fetalis  
D. Parvovirus B19 infection  
E. Placental vascular anastomosis

The risk for fetal and neonatal accumulation of edema fluid in soft tissues is increased in Mayan mothers whose spouse is of European ancestry. Firstborn infants are affected, and the risk increases with subsequent pregnancies. Neonatal Coombs test is positive, with both IgG and IgM antibodies present. Which of the following blood groups are these Mayan women most likely to have?

A. A  
B. B  
C. O  
D. D  
E. P

A 19-year-old primigravida who has had an uncomplicated pregnancy undergoes a screening ultrasound at 16 weeks’ gestation that shows no abnormalities. At 18 weeks, the woman develops a mild rash on her face. She gives birth to a stillborn, severely hydropic male infant at 33 weeks. At autopsy, there are no congenital malformations, but cardiomegaly is present. From the histologic appearance of the bone marrow shown in the figure, which of the following is the most likely cause of these findings?

A. Chromosomal anomaly of the fetus  
B. Congenital neuroblastoma  
C. Inheritance of two abnormal CFTR genes  
D. Infection with parvovirus B19  
E. Maternal IgG crossing the placenta
32 An 18-month-old, light-skinned African American child has a developmental delay characterized by mental retardation and inability to walk. The child’s urine has a distinctly “mousy” odor. On physical examination, there is no lymphadenopathy or hepatosplenomegaly. Laboratory studies show hemoglobin, 14 g/dL; platelet count, 302,700/mm³; WBC count, 7550/mm³; glucose, 80 mg/dL; total protein, 7.1 g/dL; albumin, 5 g/dL; and creatinine, 0.5 mg/dL. A mutation involving a gene that encodes which of the following enzymes is most likely to be present in this child?

A Adenosine deaminase  
B α1-Antitrypsin  
C Galactose-1-phosphate uridyltransferase  
D Glucose-6-phosphatase  
E Lysosomal acid maltase  
F Phenylalanine hydroxylase  
G Sphingomyelinase

33 A 20-year-old woman misses a menstrual period, and her pregnancy test is positive. This pregnancy is planned. She was diagnosed with phenylalanine hydroxylase deficiency at birth and was placed on a special diet until she was 12 years old. Her physical development has been normal and she has obtained a degree from Universidad de Buenos Aires. Which of the following treatment plans is most appropriate for this pregnancy?

A Abortion to prevent birth of a mentally retarded baby  
B Enzyme infusion into the fetus  
C Gene therapy for the fetus  
D Pharmacotherapy for the mother  
E Special diet for the mother

34 An infant appears normal at birth following an uncomplicated pregnancy and begins to nurse. Within 3 days the infant exhibits vomiting and diarrhea. A week later jaundice is evident, and the infant becomes febrile. On physical examination hepatomegaly is present. Laboratory studies show abnormal reducing substances in the urine, and Escherichia coli is cultured from blood. This infant most likely has a deficiency in which of the following enzymes?

A Galactose-1-phosphate uridyltransferase  
B Glucocerebrosidase  
C Phenylalanine hydroxylase  
D Sphingomyelinase  
E Uridine diphosphate glucuronosyltransferase

35 A 21-year-old primigravida gives birth to a term infant after an uncomplicated pregnancy. The infant is of normal height and weight, and no anomalies are noted. The infant fails to pass meconium. Laboratory studies show an elevated sweat chloride level. Genetic testing indicates that a critical protein coded by a gene is missing one phenylalanine amino acid in the protein sequence.

Normal sequence:

- Ile  Ile  Phe  Gly  Val  
- ...T  ATC  ATC  TTT  GGT  GTT...

Altered sequence in this child:

- Ile  Ile  Gly  Val  
- ...T  ATC  ATT  GGT  GTT...

Which of the following types of gene mutations is most likely to produce these findings?

A Frameshift  
B Nonsense (stop codon)  
C Point  
D Three-base pair deletion  
E Trinucleotide repeat

36 A 29-year-old woman has a history of steatorrhea and recurrent pulmonary infections since childhood. She experiences a fracture, and the radiograph shows osteopenia. Laboratory studies show an abnormal sweat chloride level. Neither parents nor siblings are affected. Genetic studies show a mutation in a gene encoding for chloride ion channel. Which of the following inheritance patterns is her disease most likely to have?

A Autosomal dominant  
B Autosomal recessive  
C Mitochondrial DNA  
D Multifactorial  
E X-linked recessive

37 An 11-year-old child has had increasing episodes of diarrhea for the past 3 years. The child’s stools are bulky and foul-smelling. The child also has a history of multiple respiratory tract infections. On physical examination, vital signs include temperature of 38.1°C, pulse of 80/min, respiration of 20/min, and blood pressure of 90/55 mm Hg. On auscultation, diffuse crackles are heard over both lungs. Laboratory findings include quantitative stool fat of greater than 10 g/day. Sputum cultures have grown Pseudomonas aeruginosa and Burkholderia cepacia. Which of the following inborn errors of metabolism is most likely present in this child?

A Abnormal fibrillin production by fibroblasts  
B Galactose-1-phosphate uridyltransferase deficiency  
C Impaired epithelial cell transport of chloride ion  
D Phenylalanine hydroxylase deficiency  
E Reduced numbers of LDL receptors

38 A 15-year-old girl has had a fever and productive cough for the past 4 days. She was born at term, but developed abdominal distention in the first week of life from meconium ileus. She has had frequent bouts of pneumonia with cough productive of thick mucoid sputum for 8 years. Sputum cultures have consistently grown Pseudomonas aeruginosa. Each new bout of pneumonia is longer and increasingly nonresponsive to antibiotics. On physical examination, her temperature is now 38.2°C. Which of the following is the most important factor causing increasing frequency and severity of her Pseudomonas infections?

A Abnormal folding of CFTR protein  
B Coexistent mutations in the TGFβ1 gene  
C Defect in ciliary action from dynein arm malfunction  
D Formation of respiratory epithelial biofilms  
E Loss of bicarbonate secretion into bronchioles
39. A clinical study is performed involving male subjects with infertility. Some of these patients are found to have bilateral agenesis of the vas deferens along with a history of recurrent respiratory tract infections, steatorrhea, and biliary cirrhosis. Genetic analysis reveals polymorphisms of mannose-binding lectin 2 (MBL2) and transforming growth factor beta 1 (TGFβ1) genes. An abnormality involving which of the following laboratory tests is most likely to be found in these infertile men?

A. Blood hemoglobin A1c  
B. Urinary amino acids  
C. Serum phenylalanine  
D. Sweat chloride  
E. Urine vanillylmandelic acid (VMA)

40. A 5-month-old male infant died suddenly and unexpectedly. Scene investigation reveals the infant was prone with no airway obstruction or evidence of trauma. Gross and microscopic examination at autopsy reveals only petechiae of mesothelial surfaces. Which of the following neurotransmitters that involves signaling within brainstem medullary arousal centers is most likely implicated in the pathogenesis of this infant’s death?

A. Acetylcholine  
B. Dopamine  
C. γ-Aminobutyric acid  
D. Norepinephrine  
E. Serotonin

41. An 4-month-old African American boy is found dead in his crib one morning. The distraught parents, both factory workers, are interviewed by the medical examiner and indicate that the child had not been ill. She finds no gross or microscopic abnormalities at autopsy, and the results of all toxicologic tests are negative. The medical examiner tells the parents she believes the cause of death is sudden infant death syndrome (SIDS). This conclusion is most likely based on which of the following factors?

A. Age of the child  
B. Sex of the child  
C. Race of the child  
D. Parental socioeconomic status  
E. No abnormalities at autopsy

42. A 3-year-old child’s mother notes that a large port wine stain on the left side of the child’s face has not diminished in size since birth. This irregular, slightly raised, red-blue area is not painful, but is very disfiguring. Histologically, this lesion is most likely composed of a proliferation of which of the following tissue components?

A. Capillaries  
B. Fibroblasts  
C. Lymphatics  
D. Lymphoblasts  
E. Neuroblasts

43. A longitudinal study of pediatric neoplasia is performed. Infants born at term with no abnormalities and no family history of malignancies are identified and followed. Which of the following neoplasms is most likely to be diagnosed in the first year of life in this cohort?

A. Adenocarcinoma  
B. Ewing sarcoma  
C. Hepatoblastoma  
D. Hodgkin lymphoma  
E. Squamous cell carcinoma

44. A 28-year-old woman, G1, P0, has an uncomplicated pregnancy until 28 weeks’ gestation, when she develops uterine contractions and has premature rupture of membranes. An ultrasound reveals a lesion with the representative gross appearance shown in the figure. Which of the following is the most likely diagnosis of this lesion?

A. Hamartoma  
B. Hemangioma  
C. Lymphangioma  
D. Neuroblastoma  
E. Teratoma

45. The parents of a 2-year-old boy are concerned because their son seems to have no vision in his right eye. On examination there is strabismus and a whitish appearance to the pupil on the right, with tenderness to orbital palpation. Vision on the left appears to be intact. An enucleation of the right eye is performed, followed by radiation and chemotherapy. There is no recurrence on the right, but at age 5 years, a similar lesion develops in the left eye. At age 12 years, the boy develops an osteosarcoma of the left distal femur. Which of the following genetic mechanisms is most likely to produce these findings?

A. Aneuploidy  
B. Chromosomal translocation  
C. Trinucleotide repeat mutation  
D. Germline mutation  
E. Multifactorial inheritance  
F. Uniparental disomy  
G. X-linked gene defect
46. The mother of a 6-month-old boy notices that he has a palpable abdominal mass. On physical examination, the infant has a temperature of 37.8°C, and he is at the 33rd percentile for weight. An abdominal CT scan shows a solid 5.5-cm mass involving the right adrenal gland. Laboratory studies show that 24-hour urine levels of homovanillic acid (HVA) and vanillylmandelic acid (VMA) are increased. The adrenal gland is excised surgically; the histologic appearance of the mass is shown in the figure. Which of the following features of this lesion is most likely associated with a poor prognosis?

A. Age younger than 1 year
B. Hyperdiploidy
C. Malformations of the kidney
D. MYCN (NMYC) gene amplification
E. Presence of many ganglion cells

47. A 2-year-old boy is noted to have absence of the iris bilaterally, cryptorchidism, hypospadias, and mental retardation. His mother has observed that the child has an enlarging abdomen over the past 3 months. On physical examination, there are palpable abdominal masses. An abdominal CT scan shows bilateral adrenal enlargement and pancreatic enlargement. There is a 6-cm solid mass in the left kidney. Which of the following congenital disorders is the most likely diagnosis?

A. Edwards syndrome
B. Marfan syndrome
C. Klinefelter syndrome
D. Patau syndrome
E. Turner syndrome
F. WAGR syndrome

48. A 3-year-old child has become less active over the past 2 months. On physical examination an abdominal mass is palpable on the left. A urinalysis shows hematuria. Radiographic studies show an 8-cm mass in the left retroperitoneal space. The mass is excised and on microscopic examination shows a triphasic combination of blastemal, stromal, and epithelial cell types. The child is doing well 10 years later. Which of the following most likely related to the pathogenesis of this child’s neoplasm?

A. Anaplastic epithelium
B. Ganglion cells
C. Nephrogenic rests
D. Pseudorosettes
E. Sarcomatous component

ANSWERS

1. B. This is a classic example of an embryonic disruption that leads to the appearance of congenital abnormalities. Fibrous bands and possible vascular insults may explain such findings, which fall within the spectrum of a limb–body wall complex that includes amniotic band syndrome. Various malformations may occur as a result of congenital infections, but amniotic bands are not among them. Oligohydramnios with diminished amniotic fluid leads to deformations, not disruptions. Rh incompatibility can give rise to erythroblastosis fetalis, which may manifest as hydrops fetalis. Fetuses affected by hydrops have widespread edema and intense jaundice. In trisomy 18 and other chromosomal abnormalities, an omphalocele centered on the umbilicus is the most common abdominal wall defect.

PBD9 452–453  BP9 246  PBD8 448–449  BP8 253–254

2. C. The lungs are hypoplastic (small) because of deformation caused by an oligohydramnios sequence. The malformation (anomaly) that initiated the sequence in this case was multicystic renal dysplasia because the kidneys formed little fetal urine, which is passed into the amniotic cavity to form the bulk of the amniotic fluid. There was no disruption in this case. Teratogens may produce anomalies, but these are uncommon. The spectrum of findings in this case is consistent with the VATER association (vertebral defects, imperforate anus, tracheoesophageal fistula, and radial and renal dysplasia).

PBD9 452–454  BP9 246–247  PBD8 449–450  BP8 254

3. B. The flattened face and deformed feet of this infant suggest oligohydramnios resulting from renal agenesis. Fetal kidneys produce urine that becomes the amniotic fluid. The lack of this fluid constrains the developing fetus and restricts pulmonary growth. Pulmonary hypoplasia is the rate-limiting step to survival. Congenital rubella can lead to various malformations, but not deformations. Infants born to diabetic mothers have an increased risk of congenital anomalies without a specific pattern. Fetal lung maturity is typically achieved at 34 to 35 weeks’ gestation, and hyaline membrane disease is unlikely at 36 weeks. Trisomy 13 is accompanied by various malformations, including malformations affecting the kidneys. The external features are quite different from those seen in this case, however, and affected infants almost always have microcephaly and midline defects, such as cleft lip and palate.

PBD9 454  BP9 246–247  PBD8 449–450  BP8 254

4. A. Retinoic acid embryopathy, which is characterized by cardiac, neural, and craniofacial defects, is believed to result from the ability of retinoids to down-regulate the
transforming growth factor beta (TGF-β) signaling pathway and affect the expression of homeobox (HOX) genes. These genes are important in embryonal patterning of limbs, vertebrae, and craniofacial structures. Embryonic rhombomeres can be affected in their development. The other listed options do not apply to retinoic acid.

PBD9 454, 456 BP9 247 PBD8 453 BP8 255–256

5 A The findings are typical for fetal growth restriction in later term, owing to relative sparing of the fetal brain. The findings suggest uteroplacental insufficiency from an intrinsically placental abnormality. One cause is a cytogenetic abnormality in the developing embryo that occurs just in the trophoblast, and only placenta is affected. Congenital syphilis occurs in the third trimester, when spirochetes can cross the placenta; there is typically placental and fetal hydrops. Inborn errors of metabolism such as galactosemia usually manifest after birth because maternal metabolism clears any intermediate products of fetal enzyme deficiencies. Preeclampsia is marked by maternal hypertension. An Rh incompatibility leads to fetal and placental hydrops. Trisomy 21 is more frequent with advancing maternal age, but in this case no fetal anomalies were noted.

PBD9 456–457 BP9 247 PBD8 455 BP8 256–257

6 B This is developmental dysplasia of the hip. Most congenital malformations, particularly malformations that are isolated defects, have no readily identifiable cause. Most defects are believed to be caused by the interaction of a certain number of genes and by the interaction of those genes with environmental factors. Their transmission follows the rules for multifactorial inheritance. The recurrence rate is believed to be 2% to 7% and is the same for all first-degree relatives, regardless of sex and relationship to the index case. Some populations may have more carriers of these genes.

PBD9 454–455 BP9 247 PBD8 448–450 BP8 240–241

7 A Rubella infection in the first trimester, when organogenesis is occurring (3 to 9 weeks’ gestation), can lead to embryopathy with cardiac, ocular, skin, central nervous system, and hepatic abnormalities. Rubella is rare with routine vaccinations. Folate deficiency is most likely to be associated with neural tube defects. Dispermy is rare and can lead to triploidy, a condition that rarely results in a live birth. Erythroblastosis fetalis leads to fetal anemia with congestive heart failure and hydrops, but not to malformations. In the past, thalidomide use was a cause of malformations (almost invariably prominent limb deformities). Nondisjunctional events during meiosis in the maternal ova account for trisomies and monosomies, many of which have associated cardiac defects, including ventricular septal defect (e.g., trisomy 21). This mechanism is unlikely in paternal sperm, however, which are constantly being produced in large numbers throughout life.

PBD9 454 BP9 247 PBD8 451–452 BP8 255–256

8 D This infant has a cleft lip, one of the most common congenital anomalies. The absence of other defects suggests that this is an isolated anomaly, and not part of a syndrome or disease with a defined genetic cause. Most congenital malformations, including cleft lip, are not determined by a single gene and may be conditioned by environmental influences. Multiple genetic susceptibilities, modified by environmental influences, can underlie the appearance of 20% to 25% of all birth defects. Of the remaining options, all are more likely to produce multiple defects and to reduce fetal growth. Most chromosomal anomalies are not compatible with survival; the few fetuses (e.g., with sex chromosome aneuploidies and autosomal triploidies such as 13, 18, and 21) that do survive to term and beyond manifest multiple anomalies. Early amniocentesis may result in clefts, but more severe defects are present (e.g., gastroschisis and missing digits or limbs), and stillbirth is the usual consequence. Maternal malnutrition typically results in an infant who is small for gestational age. Single gene defects account for less than 10% of anomalies noted at birth. Teratogens account for no more than 1% of congenital anomalies. Albucasis (963-1013 CE) is the “father of surgery” whose notable work, Al-Tasrif, was composed of 30 volumes on medical science, including 3 volumes on surgery in which he described many surgical procedures. He designed dozens of surgical instruments (including the bulb syringe we have all used) and was the first to use silk sutures.

PBD9 453, 455 BP9 248 PBD8 450–452 BP8 240–241

9 B Alcohol is one of the most common environmental teratogens affecting fetuses, although the effects can be subtle. There is no threshold amount of alcohol consumption by the mother to produce fetal alcohol syndrome; no amount is safe. Children with fetal alcohol syndrome tend to be developmentally impaired throughout childhood, but the physical anomalies tend to become less apparent as the child matures. Vertebral abnormalities, including scoliosis, can be present. The liver can have fatty metamorphosis with hepatomegaly and elevated serum transaminases. The major effects of congenital rubella occur during organogenesis in the first trimester and result in more pronounced defects, including congenital heart disease. Maternal diabetes often results in a larger infant, and malformations may be present. Placenta previa, a low-lying placenta at or near the cervical os, can cause significant hemorrhage at the time of delivery or uteroplacental insufficiency with growth retardation before delivery. Placental causes of intrauterine growth retardation result in asymmetric growth retardation with sparing of the brain. The findings of trisomy 21 are subtle at birth, but typically include brachycephaly, not microcephaly.

PBD9 454–455 BP9 247–248 PBD8 452 BP8 255

10 A Multiple fetal losses earlier in gestation suggest the likelihood of a chromosomal abnormality—the mother or father may be the carrier of a balanced translocation. Cystic fibrosis transmembrane conductance regulator (CFTR) gene mutations may lead to cystic fibrosis, and like most inborn errors of metabolism, with appearance of findings postnatally. The maternal serum level of α-fetoprotein can help identify fetal neural tube defects, but these defects are not a cause of early fetal loss. Maternal HIV infection is not a cause of
significant fetal loss. Because the mother is blood type A positive, fetal loss with erythroblastosis fetalis is unlikely, although other blood group incompatibilities potentially may result in erythroblastosis fetalis.

11 E Though a specific genetic abnormality, such as a chromosomal abnormality or single gene defect, may be sought, unknown causes still account for 40% to 60% of birth defects. This is particularly true with more common birth defects, such as those listed in this case, and when they do not fit with a recognized pattern of defects suggesting a specific syndrome. Environmental factors related to birth defects include infections, maternal disease such as diabetes mellitus (not gestational diabetes), and teratogens, accounting for up to 10% of birth defects. Multifactorial causes from the influence of multiple gene predispositions is the second most likely cause for birth defects, at 20% to 25%.

12 F Maternal diabetes can result in hyperplasia of the fetal islets because of the maternal hyperglycemic environment. This can occur if gestational diabetes is present, or if the mother has had diabetes mellitus before pregnancy. After birth, the hyperplastic islets continue to overfunction, resulting in neonatal hypoglycemia. Infants of diabetic mothers also tend to exhibit macrosomia because of the growth-promoting effects of increased insulin levels. The other listed findings are more characteristic of pancreatic disorders in adults. Neutrophilic infiltration with necrosis and hemorrhage are characteristic of acute pancreatitis. A mass with irregular glands and abnormal nuclear features could be an adenocarcinoma. Amyloid deposition in islets may be seen in some cases of type 2 diabetes mellitus. Extensive fibrosis and fatty replacement of the pancreas is seen in patients with cystic fibrosis surviving for decades. Infiltration of T cells into islets occurs with the insulinis that presages overt clinical type 1 diabetes mellitus.

13 C The acute inflammation suggests a bacterial infection, and group B streptococcus, which can colonize the vagina, is a common cause for congenital infection at term. The infection can develop quickly. Cytomegalovirus, syphilis, and toxoplasmosis are congenital infections that can cause stillbirth, but they are more likely to be chronic and develop earlier in gestation. Herpetic infections are most likely to be acquired by passage through the birth canal.

14 F Preterm premature rupture of membranes accounts for 30% to 40% of preterm deliveries, and infection is often the cause, with ascending infection often resulting in chorioamnionitis and funisitis. This organism is found in the genital tract of sexually active adults. Intrauterine infection results in release of collagenases and elastases that promote rupture of membranes and release of prostaglandins that induce smooth muscle contraction and labor. Cystic fibrosis and inborn errors of metabolism are usually not manifest until after birth. Rh incompatibility is unlikely in a primigravida, unless she received a prior incompatible blood transfusion, and results in fetal hydrops. Sudden infant death syndrome occurs in infants 1 month to 1 year of age, and there are no associated pathologic abnormalities. Maternal hypertension and convulsions should be present for a diagnosis of toxemia. Trisomy 16 can be seen with first-trimester losses, and is not associated with inflammation.

15 D Listeriosis can be a congenital infection. Although pregnant women may have only a mild diarrheal illness, the organism can prove devastating to the fetus or neonate. Mini-epidemics of listeriosis are often linked to a contaminated food source, such as dairy products, chicken, or hot dogs. Neonatal meningitis can be caused by *Listeria monocytogenes*. Cytomegalovirus and toxoplasmosis are most likely to produce severe central nervous system damage. Group B streptococcal infections most often infect the fetus near term or peripartum. These organisms release a factor that inhibits the neutrophilic chemotactic factor complement C5a, inhibiting a suppressive response. Herpetic congenital infections typically are acquired via passage through the birth canal. Parvovirus infection may cause a severe fetal anemia.

16 A About 10% of cytomegalovirus-infected neonates have extensive infection with inclusions found in many organs. Severe anemia and myocardial injury cause hydrops, and the brain is often involved. The renal tubular epithelium can be infected, and large cells with inclusions can be seen with urine microscopic examination in some cases. Cytomegalovirus manifested in neonates may have been acquired transplacentally, at birth, or in breast milk. Herpes simplex virus is usually acquired via passage through the birth canal and does not cause a periventricular leukomalacia. HIV infection in utero does not produce marked organ damage. Parvovirus infection may cause a severe fetal anemia. Congenital rubella manifests in the first trimester, often with cardiac defects.

17 A The only increased IgM titer, indicating recent infection, is that for *Toxoplasma*. The fetal central nervous system findings are consistent with congenital toxoplasmosis. Her increased IgG titers for CMV and HSV type 1 suggest past infection. Her positive rubella titer is consistent with past immunization. Congenital HIV does not have significant effects on the fetus in utero. The likely route for her recent *Toxoplasma* infection is ingestion of poorly cooked, contaminated meat containing cysts of *Toxoplasma gondii*. The other routes of infection listed are not characteristic for toxoplasmosis.

18 E The infant is small for gestational age because of intrauterine growth retardation (IUGR). The asymmetric growth
with normal head size suggests a maternal or placental cause in later gestation. Fetal problems, such as chromosomal abnormalities (Down syndrome), infections (cytomegalovirus), and erythroblastosis, are likely to produce symmetric growth retardation. Fetal hydrops can accompany congenital infections and erythroblastosis, which may artificially increase fetal weight. Infants born to diabetic mothers are likely to be larger than normal (macrosomia) for gestational age.

**19 A** This infant is small for gestational age (SGA). Given that there are no other problems noted prepartum or postpartum, then maternal cigarette smoking is the likely culprit, and smoking also increases the risk for preterm premature rupture of membranes (PPROM). Placental causes for fetal growth restriction (FGR) tend to spare the infant brain. Genetic mosaicism of the placenta may lead to stillbirth as well as FGR. Preeclampsia complicates the pregnancy via maternal hypertension, edema, and proteinuria. Toxoplasmosis is a congenital infection that tends to cause abnormalities of the fetal brain, becoming apparent by the second trimester. Trisomy and other fetal chromosomal abnormalities lead to multiple congenital anomalies.

**20 B** The hyperinsulinism in the fetus of a diabetic mother suppresses pulmonary surfactant production. Corticosteroids stimulate surfactant production in type II pneumocytes. Infection may increase the risk of premature birth, but it does not significantly affect surfactant production. The risk for major malformations, called diabetic embryopathy, is 6% to 10% in infants born to mothers with diabetes mellitus; but there is no set pattern to the anomalies, and similar embryopathies are not seen in gestational diabetes. So the cause for anomalies here is unknown. Macrosomia is the result of growth-promoting effects of insulin in the fetus. Oligohydramnios leads to constriction in utero that culminates in pulmonary hypoplasia, not decreased surfactant. Maternal hypertension may reduce placental function and increase growth retardation, but it typically does not have a significant effect on the production of surfactant.

**21 B** Sustained, high-dose oxygen therapy delivered with positive pressure ventilation can cause injury to immature lungs, leading to the chronic lung disease known as bronchopulmonary dysplasia. Pulmonary manifestations of cystic fibrosis are not seen at birth or in infancy. Mortality from pulmonary hypoplasia is greatest at birth. In sudden infant death syndrome, no anatomic abnormalities are found at autopsy. A ventricular septal defect eventually could lead to pulmonary hypertension from the left-to-right shunt.

**22 E** Surfactant is synthesized by type II pneumocytes that line the alveolar sacs. They begin to differentiate after the 26th week of gestation. These cells can be recognized on electron microscopy by the presence of lamellar bodies filled with phospholipids. Surfactant production increases greatly after 35 weeks’ gestation. Other structures in the lung do not synthesize the phosphatidylcholine and phosphatidylglycerol compounds that are important in surfactant production for reducing alveolar surface tension.

**23 B** The immaturity of the fetal lungs before 35 to 36 weeks’ gestation can be complicated by lack of sufficient surfactant to enable adequate ventilation after birth. This can result in hyaline membrane disease, shown in the figure. Tests on amniotic fluid before birth, including lecithin-sphingomyelin ratio, fluorescence polarization, and lamellar body counts, are useful in predicting the degree of pulmonary immaturity. Fetal anemia leads to heart failure and pulmonary congestion. Maternal toxemia and congenital infections such as toxoplasmosis may lead to hyaline membrane disease if the birth occurs prematurely as a consequence of these conditions, but they do not directly affect lung maturity. Oligohydramnios may result in neonatal respiratory distress through the mechanism of pulmonary hypoplasia.

**24 E** Necrotizing enterocolitis is a complication of prematurity that is related to various factors, including intestinal ischemia, enterocyte apoptosis induced by platelet-activating factor, bacterial overgrowth, and formula feeding. If severe, the wall of the intestine becomes necrotic and perforates, necessitating surgical intervention. Duodenal atresia is an uncommon congenital anomaly most often associated with trisomy 21; it leads to upper gastrointestinal obstruction and vomiting. Hirschsprung disease is a congenital condition resulting from an aganglionic segment of distal colon; it leads to obstruction with distention, but not bloody diarrhea. A Meckel diverticulum is a common anomaly and is seen in about 2% of individuals. Typically, it is an incidental finding, although later in life it may be associated with gastrointestinal tract bleeding if ectopic gastric mucosa is present within the diverticulum. Meconium ileus is seen in the setting of cystic fibrosis and can lead to obstruction, but the infant typically does not pass stool. Pyloric stenosis manifests at 3 to 6 weeks of life with projectile vomiting.

**25 D** The neonate most likely has hyaline membrane disease from fetal lung immaturity and lack of surfactant. Surfactant produced by type II pneumocytes consists predominantly of phospholipids-like lecithin contained in lamellar bodies released into alveoli. The adequacy of surfactant production can be gauged by the phospholipid content of amniotic fluid because fetal lung secretions are discharged into the amniotic fluid. Chromosomal analysis may help to predict problems after birth or the possibility of fetal loss. The Coombs test may help to determine the
The presence of erythroblastosis fetalis. Cystic fibrosis does not cause respiratory problems at birth. The maternal serum α-fetoprotein level is useful to predict fetal neural tube defects and chromosomal abnormalities.

PBD9 457–458  BP9 250–251  PBD8 457  BP8 257–259

26 A This mild hemolytic anemia is most likely due to an ABO incompatibility with maternal blood type O, which results in anti-A antibody coating fetal cells. Most anti-A and anti-B antibodies are IgM. In about 20% to 25% of pregnancies, there are also IgG antibodies, which cross the placenta in sufficient titer to produce mild hemolytic disease in most cases. The bilirubin concentration in the term infant in this case is not high enough to produce kernicterus. ABO incompatibilities are not likely to have such serious consequences for subsequent pregnancies as does Rh incompatibility. As the infant matures, the level of maternal antibody diminishes, hemolysis abates, and the infant develops normally. Respiratory distress is unlikely at term because of appropriate fetal lung maturity.

PBD9 461–462  BP9 254–255  PBD8 460–461  BP8 261–262

27 C This infant has erythroblastosis fetalis, which results when prior sensitization to a fetal blood cell antigen leads to alloantibodies in maternal blood that can cross the placenta. The Rh blood group system is most often implicated. The maternal antibody coats fetal RBCs, causing hemolysis. The Coombs test is positive. The fetal anemia leads to congestive heart failure and hydrops. Hemolysis results in a very high bilirubin level. A high maternal serum level of α-fetoprotein suggests a fetal neural tube defect; such defects are not associated with hydrops. Viral hepatitis is not a perinatal infection. Diminished glucocerebrosidase activity causes Gaucher disease, and this condition does not lead to perinatal liver failure or anemia. Listeriosis or other congenital infections may produce fetal hydrops and anemia, although not of the severity described in this case.

PBD9 461–462  BP9 254–256  PBD8 460–461  BP8 261–262

28 B Nonimmune hydrops fetalis may be due to cardiac failure in utero, with cardiac defects leading to high-output congestive heart failure. Cardiac defects may occur in association with other anomalies and may be part of chromosomal aneuploidies (chromosomes 13, 18, 21, and XO). A mass lesion with increased blood flow, such as a hemangioma, could lead to hydrops, but a neuroblastoma is unlikely to have markedly increased flow, or lead to a para-neoplastic effect with anemia. Cerebral lesions such as leukomalacia may be related to hydrops when due to a congenital infection such as cytomegalovirus or toxoplasmosis (“T,” respectively, in the TORCH mnemonic), but the TORCH titer results were negative in this case. A low-lying placenta may lead to uteroplacental insufficiency with intrauterine growth retardation, and predispose to placenta previa at the time of birth. Splenomegaly is more likely to accompany immune hydrops, with marked extramedullary hematopoiesis.

PBD9 462–463  BP9 254–255  PBD8 461–462  BP8 262–264

29 E Twin–twin transfusion syndrome is the result of a vascular anastomosis between halves of a monochorionic placenta. The donor twin typically is smaller, with reduced organ function, including kidneys, and reduced amount of amniotic fluid. The recipient twin becomes plethoric with hydrops and increased amniotic fluid. The donor may also become hydropic because of high-output heart failure from the increased work of pumping blood into both twins. A chromosomal aneuploidy is likely to affect just one of fraternal twins and could lead to hydrops, but the other twin might not be affected. The other listed options should affect both twins similarly.

PBD9 462  BP9 254–255  PBD8 461  BP8 261

30 D This is immune hydrops fetalis due to ABO incompatibility. Native populations of Mesoamerica and South America are almost exclusively blood group O. Although there are naturally occurring IgM isohemagglutinins against A and B blood group antigens, some women produce IgG antibodies as well, which cross the placenta and bind to fetal cells. Persons of European heritage are predominantly blood group A. In the clash of Old and New worlds in the 16th century, populations of the latter fared far worse. Rh incompatibility is more severe, but occurs after the first pregnancy, when a fetal-maternal bleed can immunize the mother. However, the effects can be prevented with Rho(D) immune globulin administered at birth. The P blood group antigen system may play a role in some hemolytic anemias.

PBD9 461–462  BP9 254–255  PBD8 460–461  BP8 261–262

31 D The erythroid precursors exhibit large, pink, intranuclear inclusions typical of parvovirus infection. In adults, such an infection typically causes fifth disease, which is self-limiting. This is one of the “O” infections in the TORCH mnemonic, however, describing congenital infections (toxoplasmosis, other infections, rubella, cytomegalovirus infection, and herpes simplex virus or HIV infection). Parvovirus infection in the fetus can lead to a profound fetal anemia with cardiac failure and hydrops fetalis. Results of the Coombs test are negative, because no anterythrocyte antibodies are involved. Congenital tumors are an uncommon cause of hydrops, and they would produce a mass lesion, which was not described in this case. Cystic fibrosis does not affect erythropoiesis. Although various chromosomal anomalies—monosomy X, in particular—may lead to hydrops, malformations are typical. Erythroblastosis fetalis from maternal IgG antibodies directed at fetal RBCs is unlikely to occur in a first pregnancy, and only erythroid expansion would be present, not erythroid inclusions.

PBD9 460, 462  BP9 256  PBD8 459  BP8 262–263

32 F Phenylketonuria (PKU) is caused by the absence of functional phenylalanine hydroxylase genes and gives rise to hyperphenylalaninemia, which impairs brain development and can lead to seizures. The block in phenylalanine metabolism results in decreased pigmentation of skin
and hair. It also results in the formation of intermediate compounds, such as phenylacetic acid, which are excreted in urine and impart to it a “mousy” odor. Although PKU is rare, because of the devastating consequences of this inherited disorder, and because it can be treated with a phenylalanine-free diet, this is one of the diseases for which screening is performed at birth. Adenosine deaminase deficiency is a cause of severe combined immunodeficiency, which is characterized by multiple recurrent severe infections from birth. α1-Antitrypsin deficiency may produce cholestasis in children, but chronic liver disease develops later. Galactose-1-phosphate uridyltransferase deficiency causes galactosemia, which is characterized by severe liver disease and cataracts. Glucose-6-phosphatase deficiency causes type I glycogenosis (von Gierke disease), which leads to liver failure. Lysosomal acid maltase deficiency causes Pompe disease, with features that include cardiomegaly and heart failure. Sphingomyelinase deficiency causes Niemann-Pick disease; affected infants have marked hepatosplenomegaly and neurologic deterioration.

33 E Many persons with classic phenylketonuria (PKU) have survived to adulthood because of newborn screening and dietary therapy. Insufficient phenylalanine hydroxylase leads to accumulation of phenylalanine, which is toxic to the nervous system. Regardless of whether the developing fetus has this same deficiency, high maternal levels of phenylalanine can affect the fetus. Thus a special maternal diet is indicated, before and during pregnancy, and it has been suggested that persons with PKU may remain on the special diet even after childhood to reduce neurologic damage. The incidence of PKU in live births is 1 in 10,000 to 20,000 in the United States. Incidence is probably a little higher in Argentina because the population is derived from the Mediterranean region, where the prevalent mutation in the gene for phenylalanine hydroxylase is different from the most common mutation in populations of Northern Europe. The highest incidence is in Turkey, at 1 in 2600. Because this is a treatable disorder in a planned pregnancy, abortion is not the best option. Some inborn errors of metabolism may be treated with enzyme replacement following birth, but this is very expensive. Gene therapy is experimental; bone marrow transplantation has been used in some cases of inherited enzyme disorders. There is no drug therapy for PKU.

34 A This infant has galactosemia, an inborn error of metabolism with an autosomal recessive pattern of inheritance. Lactose in milk is a disaccharide that is converted by lactase to glucose and galactose. An excess of galactose is toxic to liver, brain, and eye. Affected infants begin to develop abnormalities as soon as they begin to ingest milk. Glucocerebrosidase deficiency leads to Gaucher disease. Phenylalanine hydroxylase deficiency leads to the classic form of phenylketonuria. Sphingomyelinase deficiency leads to Niemann-Pick disease. UDP-glucuronosyltransferase deficiency is present in Gilbert syndrome.

35 D In cystic fibrosis the elevated sweat chloride level is related to a defect in the transport of chloride ions across epithelia. The most common genetic defect is a deletion of three base pairs at the ΔF508 position coding for phenylalanine in the cystic fibrosis transmembrane conductance regulator (CFTR) gene. A frameshift mutation involves one or two base pairs, not three, and changes the remaining sequence of amino acids in a protein. A point mutation may change the codon to the sequence of a “stop” codon, which truncates the protein being synthesized, typically leading to degradation of the protein. A point mutation typically is a missense mutation that leads to replacement of one amino acid for another in the protein chain; this can lead to abnormal conformation and function of the protein. A trinucleotide repeat sequence mutation leads to amplification of repeats of three nucleotides, so-called tandem repeats, which prevent normal gene expression.

36 B She has cystic fibrosis, due to inheritance of two abnormal alleles for the cystic fibrosis transmembrane conductance regulator (CFTR) gene. There are multiple mutations, which may have different degrees of severity. The incidence in the United States is about 1 in 2500 live births, from a carrier rate of 1 in 25 (this example makes the math easy). Autosomal dominant mutations tend to involve 50% of family members, and structural proteins tend to be affected. Mitochondrial dominant mutations tend to involve 50% of family members, and structural proteins tend to be affected. Mitochondrial DNA is inherited on the maternal side and affects mainly genes associated with oxidative phosphorylation. Multifactorial inheritance does not have a well-defined recurrence risk, but tends to run in families, and is more characteristic for diseases such as diabetes mellitus or schizophrenia. X-linked recessive disorders are most likely to appear in males born to female carriers.

37 C With cystic fibrosis the abnormal chloride ion transport results in abnormal mucous secretions in pancreatic ducts. The secretions cause plugging with subsequent acinar atrophy and fibrosis leading to malabsorption, particularly of lipids. Abnormal fibrillin in elastic tissues is a feature of Marfan syndrome. Galactose-1-phosphate uridyltransferase deficiency gives rise to galactosemia. Patients with this condition have liver damage, but no pancreatic abnormalities. Phenylketonuria results from a deficiency of phenylalanine hydroxylase. Abnormalities of the LDL receptor in familial hypercholesterolemia lead to accelerated atherogenesis.

38 D With her classic history for cystic fibrosis (CF), genetic testing is most likely to reveal homozygosity for the ΔF508 mutation of the cystic fibrosis transmembrane conductance regulator (CFTR) gene. Biofilm formation in the airspaces has emerged as a major factor in the ability of P. aeruginosa to evade antibodies and antibiotics. The mucoid airway secretions create a hypoxic microenvironment in which the Pseudomonas organisms produce alginate, a mucoid polysaccharide capsule that permits formation of a biofilm that lines airspaces and hides bacteria inside it. Abnormal folding of CFTR is important in
the overall pathogenesis of CF and is not specific to lung infection. Mutations in the transforming growth factor beta 1 (TGFβ1) gene, when present along with CFTR mutations, are associated with several pulmonary phenotypes but are not the cause of increased pathogenicity of Pseudomonas infections. CFTR also controls bicarbonate transport, particularly in the pancreas, but does not regulate bicarbonate secretion in lungs. Primary ciliary defects occur in Kartagener syndrome but not in CF; in CF ciliary defects are secondary to viscid mucus production.

39 D These are manifestations of cystic fibrosis (CF), from mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene. The chloride channel defect in sweat ducts leads to concentration of sodium and chloride in sweat. Note that some CFTR mutations lead to a “forme fruste” (attenuated manifestation) of CF in which azoospermia may be the only manifestation. An elevated hemoglobin A1c is a feature of hyperglycemia with diabetes mellitus, but this is a late finding in CF, because the islets of Langerhans are not affected to the same degree as exocrine pancreas. Aminoaciduria suggests galactosemia, which is most likely to become apparent as hepatic failure in infancy. Hyperphenylalaninemia is a feature of phenylketonuria (PKU) manifested by mental retardation. VMA is increased in the urine of patients with neuroblastoma.

40 E The most likely cause of death here is sudden infant death syndrome (SIDS). Other causes of death must be excluded by careful examination and interviews with caretakers. By definition, there are no significant gross or microscopic autopsy findings. Infants with congenital anomalies or infections are unlikely to appear healthy, feed well, or gain weight normally. Though the cause for SIDS is unknown, abnormalities of serotonergic neural pathways in the medulla oblongata, and a respiratory tract stressor (sleeping prone) may put vulnerable infants at risk. Abnormalities in medullary centers that regulate responses to noxious stimuli such as hypoxia, hypercarbia, and thermal stress may lead to absence of cardiorespiratory and reflexive responses that normally maintain homeostasis. The other neurotransmitters listed are not implicated. Acetylcholine acts at all preganglionic and postganglionic parasympathetic neurons and at preganglionic sympathetic neurons. Dopaminergic neurons are found in the substantia nigra. GABAergic neurons have an inhibitory function on other neurons in the central nervous system. Noradrenergic neurons originate in the locus ceruleus, and they comprise postganglionic sympathetic neurons as well.

41 E The cause of sudden infant death syndrome (SIDS) is unknown, but certain demographic risk factors are well established. Among these is age. SIDS most often occurs between 1 month and 1 year of age, and 90% of SIDS deaths occur during the first 6 months of life. Along with age, other factors listed increase the risk of SIDS. Male sex, African American race, low socioeconomic background of parents, lack of underlying medical problems, and absence of anatomic abnormality all favor the likelihood of SIDS. The key factor that argues for SIDS in this case is the lack of autopsy findings that support another diagnosis. SIDS may relate to abnormalities of serotonergic neural pathways in the medulla oblongata, and a respiratory tract stressor (sleeping prone) may put vulnerable infants at risk.

42 A The most common tumor of infancy is a hemangioma, and these benign neoplasms form a large percentage of childhood tumors as well. Although benign, they can be large and disfiguring. Fibromatoses are fibromatous proliferations of soft tissues that form solid masses. Lymphangioma is another common benign childhood tumor seen in the neck, mediastinum, and retroperitoneum. Lymphoblasts as part of leukemic infiltrates or lymphomas are not likely to be seen in skin, but mediastinal masses may be seen. A proliferation of neuroblasts occurs in neuroblastoma, a common childhood neoplasm in the abdomen.

43 C Pediatric neoplasms are different from those in adults and are far less common. The earlier in life the malignancy arises, the more likely it has features of primitive, embryonic cells (blast cells). Hence, a tumor with the suffix -blastoma is likely a congenital or early childhood neoplasm. Adult neoplasia is often driven by environmental influences (carcinogens, chronic infections) acting over many years, and the most common neoplasms arise in epithelia (squamous and glandular). Sarcomas comprise a greater percentage of pediatric than adult neoplasms, but they are unlikely to be congenital. Hematologic malignancies such as acute lymphocytic leukemia can be found in childhood.

44 E Teratomas are benign neoplasms composed of tissues derived from embryonic germ layers (ectoderm, mesoderm, endoderm). Teratomas occur in midline locations, and the sacrococcygeal area pictured is the most common. Less common immature, or frankly malignant, teratomas with neuroblastic elements can occur. Hamartomas are masses composed of tissues normally found at a particular site, and they are rare. Hemangiomas form irregular, red-blue skin lesions that are flat and spreading. Lymphangiomias in childhood are most often located in the lateral head and neck region. Neuroblastomas are malignant childhood tumors that most often arise in the adrenal glands.

45 D This child has inherited an abnormal RB1 gene, and early in life the other allele is lost, leading to loss of tumor
suppression and development of a retinoblastoma. About 60% to 70% of retinoblastomas are associated with germline mutations. Aneuploidy usually results in fetal loss, although monosomy X and trisomies 13, 18, and 21 occasionally may lead to live births. Chromosomal translocations may be seen with other tumors, such as chronic myelogenous leukemia, acute promyelocytic leukemia, and Burkitt lymphoma. Trinucleotide repeats are typically not associated with neoplasms. Multifactorial inheritance may be associated with complex diseases, such as diabetes mellitus, hypertension, and bipolar disorder. Uniparental disomy is a mechanism for chromosomal imbalance, and neoplasia is unlikely. X-linked single-gene defects are unlikely to lead to neoplasia.

46 D Amplification of the MYCN (NMYC) oncogene occurs in about 25% of neuroblastomas, and the greater the number of copies, the worse the prognosis. This amplification tends to occur in neuroblastomas with a higher stage or with chromosome 1p deletions. Hyperdiploidy or near-triploidy is usually associated with lack of MYCN amplification, absence of 1p deletion, and high levels of nerve growth factor receptor TrkA expression. All of these are associated with good prognosis. Renal malformations are not related to neuroblastomas. The presence of ganglion cells is consistent with a better differentiation and better prognosis; some tumors may differentiate over time and become ganglioneuromas under the influence of TrkA.

47 F WAGR syndrome is an uncommon condition that carries an increased risk of development of Wilms tumor, a childhood neoplasm arising in the kidney. There is a deletion of the short arm of chromosome 11 and contiguous loss of neighboring genes. Loss of one allele of the Wilms tumor suppressor gene (WT1) leads to genitourinary defects and is the first “hit” in development of Wilms tumor. Deletion of one copy of the PAX6 gene leads to aniridia and to central nervous system maldevelopment with mental retardation. Renal anomalies, such as horseshoe kidney, can be seen in Edwards syndrome (trisomy 18), but not neoplasms. Likewise, Marfan syndrome is not associated with an increased risk of malignancy. The 47,XXX karyotype of Klinefelter syndrome does not carry an increased risk of renal tumors. Patau syndrome (trisomy 13) is associated with many anomalies, among them postaxial polydactyly and midline defects that include cleft lip and palate, cyclopia, and holoprosencephaly. Turner syndrome (monosomy X) occurs in females and can be associated with multiple anomalies, including cystic hygroma, aortic coarctation, and renal anomalies.

48 C Nephrogenic rests can be found in the renal parenchyma adjacent to a Wilms tumor. They are presumed precursor lesions because they may share genetic alterations with the adjacent Wilms tumor. There is an increased risk of developing Wilms tumors in the contralateral kidney if such rests are present. Anaplasia in Wilms tumor suggests a poor prognosis, but overall the cure rate is 90%. Ganglion cells and pseudorosettes are features of neuroblastoma. Long-term survivors of Wilms tumor are at increased risk for another malignancy, including a sarcoma, but the sarcoma is not part of the original tumor.
1. A malignant neoplasm is found within the cranial cavity. A biopsy is obtained and microscopic examination shows malignant cells within vascular lumina. Which of the following types of vessels is most likely to be invaded by this malignancy?
   A. Arterioles
   B. Capillaries
   C. Lymphatics
   D. Muscular arteries
   E. Veins

2. A 44-year-old woman is found to have a blood pressure of 150/100 mm Hg on a routine physical examination following a transient ischemic attack (TIA). Laboratory studies show normal serum potassium, sodium, and natriuretic peptide levels but increased plasma renin activity. Her urinary fractional excretion of sodium is normal. Which of the following vascular abnormalities is most likely to be seen on renal arterial angiography?
   A. Arterial dissection
   B. Arteriovenous fistula
   C. Berry aneurysm
   D. Focal stenosis and dilation
   E. Vascularized mass lesion

3. A dermal venule with endothelial cells in the basal state is stressed by release of biogenic amines from mast cells following anaphylaxis. The extravascular fluid compartment increases in size. Which of the following is the most likely change occurring in these activated endothelial cells?
   A. Cytoplasmic contraction
   B. Free radical formation
   C. mRNA translation
   D. Thromboxane synthesis
   E. VEGF elaboration

4. A 73-year-old healthy man has experienced light-headedness with episodes of fainting for the past 10 days. On examination, his blood pressure changes from 135/90 mm Hg when lying down to 100/60 mm Hg when he assumes a sitting position. Laboratory studies show his serum electrolyte levels are normal. Which of the following regulatory mechanisms for blood pressure is most likely altered in this man?
   A. Aldosterone release
   B. Cardiac output
   C. Intravascular volume
   D. Renin synthesis
   E. Sympathetic tone
   F. Vasoconstriction

5. A 55-year-old woman has had dull, episodic headaches for the past year, but otherwise she feels fine and has had no major medical illnesses or surgical procedures during her lifetime. On physical examination, her temperature is 37°C, pulse is 70/min, respirations are 14/min, and blood pressure is 166/112 mm Hg. Her lungs are clear on auscultation, and her heart rate is regular. An abdominal ultrasound scan shows that the left kidney is smaller than the right kidney. A renal angiogram shows a focal stenosis of the left renal artery. Which of the following laboratory findings is most likely to be present in this patient?
   A. Anti-double-stranded DNA titer 1:512
   B. C-ANCA titer 1:256
   C. Increased cryoglobulins in blood
   D. Plasma glucose level 200 mg/dL
   E. Marked elevation in HIV RNA
   F. Plasma renin 15 mg/mL/hr
6 A clinical study is performed that includes a group of subjects whose systemic blood pressure measurements are consistently between 145/95 mm Hg and 165/105 mm Hg. They are found to have increased cardiac output and increased peripheral vascular resistance. Renal angiograms show no abnormal findings, and CT scans of the abdomen show no masses. Laboratory studies show normal levels of serum creatinine and urea nitrogen. The subjects take no medications. Which of the following laboratory findings is most likely to be present in this group of subjects?
   A Decreased urinary sodium
   B Elevated plasma renin
   C Hypokalemia
   D Increased urinary catecholamines
   E Lack of angiotensin-converting enzyme

7 A 61-year-old woman has smoked two packs of cigarettes per day for the past 40 years. She has experienced increasing dyspnea for the past 6 years. On physical examination, her vital signs are temperature, 37.1° C; pulse, 60/min; respirations, 18/min and labored; and blood pressure, 130/80 mm Hg. On auscultation, expiratory wheezes are heard over the chest bilaterally. Her heart rate is regular. A chest radiograph shows increased lung volume, with flattening of the diaphragms, greater lucency to all lung fields, prominence of pulmonary arteries, and a prominent border on the right side of the heart. Laboratory studies include blood gas measurements of PO2 of 80 mm Hg, PCO2 of 50 mm Hg, and pH of 7.35. Which of the following morphologic findings is most likely to be present in her pulmonary arteries?
   A Amyloid deposition
   B Atheromatous plaques
   C Intimal tears
   D Medial calcific sclerosis
   E Necrotizing vasculitis
   F Phlebothrombosis

8 A population health study shows that the prevalence of essential hypertension has been increasing for the past 10 years and now is found in one fourth of all adults. A strategy is adopted to decrease the prevalence of hypertension through lifestyle changes. A policy to reduce dietary intake of which of the following is most likely to support this goal?
   A Alcohol
   B Animal protein
   C Coffee
   D Eggs
   E Milk products
   F Salt
   G Sugar

9 A 61-year-old man has reported increasing fatigue over the past year. Laboratory studies show a serum creatinine level of 4.7 mg/dL and urea nitrogen level of 44 mg/dL. An abdominal ultrasound scan shows that his kidneys are symmetrically smaller than normal. The representative high-magnification microscopic appearance of the kidneys is shown in the figure. These findings are most likely to indicate which of the following underlying conditions?
   A Adenocarcinoma of the colon
   B Escherichia coli septicemia
   C Polyarteritis nodosa
   D Syphilitic endarteritis
   E Systemic hypertension

10 A 45-year-old man has had poorly controlled hypertension ranging from 150/90 mm Hg to 160/95 mm Hg for the past 11 years. Over the past 3 months, his blood pressure has increased to 250/125 mm Hg. On physical examination, his temperature is 36.9° C. His lungs are clear on auscultation, and his heart rate is regular. There is no abdominal pain on palpation. A chest radiograph shows a prominent border on the left side of the heart. Laboratory studies show that his serum creatinine level has increased during this time from 1.7 mg/dL to 3.8 mg/dL. Which of the following vascular lesions is most likely to be found in this patient’s kidneys?
   A Fibromuscular dysplasia
   B Granulomatous arteritis
   C Renal arterial stenosis
   D Necrotizing arteriolitis
   E Polyarteritis nodosa
11 A 57-year-old woman experiences mild intermittent right hip pain after falling down a flight of stairs. Physical examination shows a 3-cm contusion over the right hip. The area is tender to palpation, but she has full range of motion of the right leg. A radiograph of the pelvis and right upper leg shows no fractures, but does show calcified, medium-sized arterial branches in the pelvis. This radiographic finding is most likely to represent which of the following?

A  An incidental observation
B  Benign essential hypertension
C  Increased risk for gangrenous necrosis
D  Long-standing diabetes mellitus
E  Unsuspected hyperparathyroidism

12 A study of risk factors for atherogenesis in adults is performed. Chemical factors are found that are associated with reduction in serum cholesterol. Which of the following substances is most likely to reduce serum cholesterol?

A  C-reactive protein
B  Homocysteine
C  Lipoprotein(a)
D  Omega-3 fatty acids
E  trans-unsaturated fats

13 A cohort study is performed involving healthy adult men and women born 20 years ago. They are followed to assess development of atherosclerotic cardiovascular diseases. Multiple laboratory tests are performed yearly during this study. An increase in which of the following is most likely to indicate the greatest relative risk for development of one of these diseases?

A  Anti–proteinase 3 (PR3)
B  C-reactive protein (CRP)
C  Cryoglobulin
D  Erythrocyte sedimentation rate (ESR)
E  Platelet count

14 A 58-year-old man had a myocardial infarction 1 year ago, which was the first major illness in his life. He now wants to prevent another acute coronary event and is advised to begin a program of exercise and to change his diet. A reduction in the level of which of the following serum laboratory findings 1 year later would best indicate the success of his diet and exercise regimen?

A  Calcium
B  Cholesterol
C  Glucose
D  Potassium
E  Renin

15 A 35-year-old woman has a reputation as a perfectionist. She has angina pectoris of 6 months’ duration. On physical examination, her blood pressure is 135/85 mm Hg. She is 168 cm (5 ft 5 in) tall and weighs 82 kg (BMI 29). Her Hgb A1C is 9% and fasting serum glucose is 143 mg/dL. Coronary angiography shows 75% narrowing of the anterior descending branch of the left coronary artery and 70% narrowing of the right coronary artery. Angioplasty with stent placement is performed. Which of the following is her greatest risk factor for the causation of her disease?

A  Age
B  Diabetes mellitus
C  Obesity
D  Sedentary lifestyle
E  Type A personality

16 An experiment studies early atheroma development. Lipid streaks on arterial walls are examined microscopically and biochemically to determine their cellular and chemical constituents and the factors promoting their formation. Early lesions show increased attachment of monocytes to endothelium. The monocytes migrate subendothelially and become macrophages; these macrophages transform themselves into foam cells. Which of the following substances is most likely to be responsible for the transformation of macrophages?

A  C-reactive protein
B  Homocysteine
C  Lp(a)
D  Oxidized LDL
E  Platelet-derived growth factor
F  VLDL

17 A 29-year-old man has had angina for the past year. There is a family history of cardiovascular disease. On examination, his blood pressure is 120/80 mm Hg. Laboratory studies show total serum cholesterol 185 mg/dL and glucose 85 mg/dL. A mutation involving a gene encoding for which of the following is most likely present in this man’s family?

A  Angiotensin
B  Apolipoprotein
C  Endothelin
D  Factor VIII
E  Von Willebrand factor

18 A study is conducted involving persons with LDL cholesterol levels above 160 mg/dL. They are found to have increased oxidized LDL deposited in their arteries. As a consequence the arterial lumen, particularly at branch points, is decreased in size. Which of the following is the most likely pathologic change that develops initially in these areas of arterial narrowing?

A  Endothelial cell disruption
B  Intimal thickening
C  Lymphocytic infiltrates
D  Platelet aggregation
E  Smooth muscle hypertrophy

19 A 50-year-old man has a 2-year history of angina pectoris that occurs during exercise. On physical examination, his blood pressure is 135/75 mm Hg, and his heart rate is 79/min and slightly irregular. Coronary angiography shows a fixed 75% narrowing of the anterior descending branch of the left coronary artery. He has several risk factors for atherosclerosis: smoking, hypertension, and hypercholesterolemia. Which of the following is the earliest event resulting from the effects of these factors?

A  Alteration in vasomotor tone regulation
B  Conversion of smooth muscle cells to foam cells
C  Dysfunction from endothelial injury
D  Inhibition of LDL oxidation
E  Modification of hepatic lipoprotein receptors
20. A 12-year-old boy died of complications of acute lymphocytic leukemia. There is no family history of cardiovascular disease. The gross appearance of the aorta at autopsy is shown in the figure. Histologic examination of the linear pale markings is most likely to show which of the following features?

A. Cap of smooth muscle cells overlying a core of lipid debris
B. Collection of foam cells with necrosis and calcification
C. Granulation tissue with a lipid core and areas of hemorrhage
D. Lipid-filled foam cells and small numbers of T lymphocytes
E. Cholesterol clefts surrounded by proliferating smooth muscle cells and foam cells

21. A study of atheromatous plaques shows that release of growth factors, including PDGF, FGF, and TNF-α leads to increased extracellular matrix production. As a result, the size of the plaques increases. Which of the following cells is most likely to release these growth factors in the plaques?

A. Endothelium
B. Fibroblast
C. Platelet
D. Smooth muscle
E. T lymphocyte

22. A 58-year-old woman has experienced chest pain at rest for the past year. On physical examination, her pulse is 80/min and irregular. The figure shows the microscopic appearance representative of her left anterior descending artery. Which of the following laboratory findings is most likely to be involved in the pathogenesis of the process illustrated?

A. Elevated platelet count
B. Low HDL cholesterol
C. Low Lp(a)
D. Low plasma homocysteine
E. Positive VDRL

23. In an experimental model of atherosclerosis, plaques are shown to change slowly but constantly in ways that can promote clinical events, including acute coronary syndromes. In some cases, changes were not significantly associated with acute coronary syndromes. Which of the following plaque alterations is most likely to have such an association?

A. Calcium deposition
B. Hemorrhage into the plaque substance
C. Intermittent platelet aggregation
D. Thinning of the media
E. Ulceration of the plaque surface

24. A 59-year-old woman with type II diabetes mellitus experiences an episode of chest pain with exercise. On examination, her BMI is 30. Angiography reveals proximal coronary arterial narrowing with up to 70% stenosis. Which of the following pharmacologic agents ingested in low doses daily is most appropriate for reducing her risk for myocardial infarction?

A. Acetaminophen
B. Aspirin
C. Ibuprofen
D. Paclitaxel
E. Propranolol
25 An 84-year-old man with a lengthy history of smoking survived a small myocardial infarction 2 years ago. He now reports chest and leg pain during exercise. On physical examination, his vital signs are temperature, 37.1°C; pulse, 81/min; respirations, 15/min; and blood pressure, 165/100 mm Hg. Peripheral pulses are poor in the lower extremities. There is a 7-cm pulsating mass in the midline of the lower abdomen. Laboratory studies include two fasting serum glucose measurements of 170 mg/dL and 200 mg/dL. Which of the following vascular lesions is most likely to be present in this patient?

A  Aortic dissection  
B  Arteriovenous fistula  
C  Atherosclerotic aneurysm  
D  Polyarteritis nodosa  
E  Takayasu arteritis  
F  Thromboangiitis obliterans

26 A 41-year-old man has had worsening abdominal pain for the past week. On physical examination, his vital signs are temperature, 36.9°C; pulse, 77/min; respirations, 16/min; and blood pressure, 140/90 mm Hg. An abdominal CT scan shows the findings in the figure. Which of the following is the most likely underlying disease process in this patient?

A  Diabetes mellitus  
B  Marfan syndrome  
C  Polyarteritis nodosa  
D  Systemic lupus erythematosus  
E  Syphilis

27 A 77-year-old man has had progressive dementia and gait ataxia for the past 9 years. He succumbs to bronchopneumonia. Autopsy shows that the thoracic aorta has a dilated root and arch, giving the intimal surface a “tree-bark” appearance. Microscopic examination of the aorta shows an obliterative endarteritis of the vasa vasorum. Which of the following laboratory findings is most likely to be recorded in this patient’s medical history?

A  Antibodies against Treponema pallidum  
B  Double-stranded DNA titer positive at 1:512  
C  Ketonuria of 4+  
D  P-ANCA positive at 1:1024  
E  Sedimentation rate of 105 mm/hr

28 A 23-year-old man experiences sudden onset of severe, sharp chest pain. On physical examination, his temperature is 36.9°C, and his lungs are clear on auscultation. A chest radiograph shows a widened mediastinum. Transesophageal echocardiography shows a dilated aortic root and arch, with a tear in the aortic intima 2 cm distal to the great vessels. The representative microscopic appearance of the aorta with elastic stain is shown in the figure. Which of the following diseases is the most likely cause of these findings?

A  ANCA-associated vasculitis  
B  Diabetes mellitus, maturity onset type  
C  Marfan syndrome  
D  Scleroderma, diffuse  
E  Systemic hypertension  
F  Takayasu arteritis

29 A 59-year-old man experiences sudden severe chest pain that radiates to his back. On physical examination his blood pressure is 170/110 mm Hg. Heart sounds are distant. Pulsus paradoxus is observed. A pericardiocentesis is performed and yields blood. Which of the following pathologic findings has most likely occurred in his aorta?

A  Aneurysm  
B  Arteriolosclerosis  
C  Dissection  
D  Thrombosis  
E  Vasculitis

30 A 75-year-old man has experienced headaches for the past 2 months. On physical examination, his vital signs are temperature, 37°C; pulse, 68/min; respirations, 15/min; and blood pressure, 130/85 mm Hg. His right temporal artery is prominent, palpable, and painful to the touch. His heart rate is regular, and there are no murmurs. His erythrocyte sedimentation rate is 100 mm/hr. A temporal artery biopsy is performed, and the segment of temporal artery excised is grossly thickened and shows focal microscopic granulomatous inflammation. He responds well to corticosteroid therapy. Which of the following complications of this disease is most likely to occur in untreated patients?

A  Blindness  
B  Gangrene of the toes  
C  Hemoptysis  
D  Malignant hypertension  
E  Renal failure
31 A 32-year-old woman has had coldness and numbness in her arms and decreased vision in the right eye for the past 5 months. On physical examination, she is afebrile. Her blood pressure is 100/70 mm Hg. Radial pulses are not palpable, but femoral pulses are strong. She has decreased sensation and cyanosis in her arms, but no warmth or swelling. A chest radiograph shows a prominent border on the right side of the heart and prominence of the pulmonary arteries. Laboratory studies show serum glucose, 74 mg/dL; creatinine, 1 mg/dL; total serum cholesterol, 165 mg/dL; and negative ANA test result. Her condition remains stable for the next year. Which of the following is the most likely diagnosis?

A Aortic dissection  
B Kawasaki disease  
C Microscopic polyangiitis  
D Syphilis  
E Takayasu arteritis  
F Thromboangiitis obliterans

32 A 43-year-old man has experienced malaise, fever, and a 4-kg weight loss over the past month. On physical examination, his blood pressure is 145/90 mm Hg, and he has mild diffuse abdominal pain, but no masses or hepatosplenomegaly. Laboratory studies include a serum urea nitrogen concentration of 58 mg/dL and a serum creatinine level of 6.7 mg/dL. Renal angiography shows right renal arterial thrombosis, and the left renal artery and branches show segmental luminal narrowing with focal aneurysmal dilation. During hemodialysis 1 week later, he experiences abdominal pain and diarrhea and is found to have melena. Which of the following serologic laboratory test findings is most likely to be positive in this patient?

A ANA  
B C-ANCA  
C HIV  
D HBsAg  
E Scl-70  
F RPR

33 A 3-year-old child from Osaka, Japan, has developed a fever and a rash and swelling of her hands and feet over the past 2 days. On physical examination, her temperature is 37.8° C. There is a desquamative skin rash, oral erythema, erythema of the palms and soles, edema of the hands and feet, and cervical lymphadenopathy. The child improves after a course of intravenous immunoglobulin therapy. Which of the following is the most likely to be a complication of this child’s disease if it is untreated?

A Asthma  
B Glomerulonephritis  
C Intracranial hemorrhage  
D Myocardial infarction  
E Pulmonary hypertension

34 A 50-year-old man has had a chronic cough for the past 18 months. Physical examination shows nasopharyngeal ulcers, and the lungs have diffuse crackles bilaterally on auscultation. Laboratory studies include a serum urea nitrogen level of 75 mg/dL and a creatinine concentration of 6.7 mg/dL. Urinalysis shows 50 RBCs per high-power field and RBC casts. His serologic titer for C-ANCA (proteinase 3) is elevated. A chest radiograph shows multiple, small, bilateral pulmonary nodules. A transbronchial lung biopsy specimen shows a necrotizing inflammatory process involving the small peripheral pulmonary arteries and arterioles. Which of the following is the most likely diagnosis?

A Granulomatosis with polyangiitis  
B Fibromuscular dysplasia  
C Granuloma pyogenicum  
D Kaposi sarcoma  
E Polyarteritis nodosa  
F Takayasu arteritis

35 A 50-year-old man with muscle pain and fever for a month now notes darker colored urine for the past 2 weeks. On physical examination he has palpable purpuric lesions of his skin. Urinalysis shows hematuria and proteinuria. Serum laboratory findings include a mixed cryoglobulinemia with a polyclonal increase in IgG, as well as a high titer of anti-neutrophil cytoplasmic autoantibodies, mainly antmyeloperoxidase (MPO-ANCA, or P-ANCA). A skin biopsy is performed. Which pathologic finding is most likely to be observed in this biopsy?

A Giant cells and macrophages  
B Medial fibrinoid necrosis  
C Microabscesses  
D Myotic aneurysms  
E Perivascular eosinophilic infiltrates

36 A 33-year-old man has smoked two packs of cigarettes per day since he was a teenager. He has had painful thromboses of the superficial veins of the lower legs for 1 month and episodes during which his fingers become blue and cold. Over the next year, he develops chronic, poorly healing ulcera tions of his feet. One toe becomes gangrenous and is amputated. Histologically, at the resection margin, there is an acute and chronic vasculitis involving medium-sized arteries, with segmental involvement. Which of the following is the most appropriate next step in treating this patient?

A Antibiotic therapy  
B Corticosteroid therapy  
C Hemodialysis  
D Insulin therapy  
E Smoking cessation

37 An 8-year-old child has had abdominal pain and dark urine for 10 days. Physical examination shows blotchy purple skin lesions on the trunk and extremities. Urinalysis shows hematuria and proteinuria. Serologic test results are negative for MPO-ANCA (P-ANCA) and PR3-ANCA (C-ANCA). A skin biopsy specimen shows necrotizing vasculitis of small dermal vessels. A renal biopsy specimen shows immune complex deposition in glomeruli, with some IgA-rich immune complexes. Which of the following is the most likely diagnosis?

A Giant cell arteritis  
B Henoch-Schönlein purpura  
C Polyarteritis nodosa  
D Takayasu arteritis  
E Telangiectasias
38. A 35-year-old woman with a history of injection drug use has developed a high fever over the past day. On examination her temperature is 37.6° C. A heart murmur is auscultated. Blood culture is obtained and grows *Staphylococcus aureus*. She develops a severe headache. A head CT scan shows an intracranial hemorrhage. Which of the following vascular complications has most likely occurred in her brain?

- A. Bacillary angiomatosis
- B. Hyperplastic arteriolosclerosis
- C. Lymphangitis
- D. Mycotic aneurysm
- E. Phlebothrombosis

39. A 60-year-old woman noted the change seen in the figure while she was driving to work one morning. There was associated pain and numbness. Within 20 minutes after entering the warm office building, these problems disappeared. What pathologic process has most likely led to these findings?

- A. Calcification
- B. Hypertension
- C. Thrombosis
- D. Vasculitis
- E. Vasoconstriction

40. A 21-year-old healthy woman, who is in week 34 of a normal pregnancy, complains of itching with burning pain in the perianal region for the past 4 months. She noted a small amount of bright red blood on toilet paper last week. Which of the following underlying conditions is most likely to be present in this patient?

- A. Filariasis
- B. Polyarteritis nodosa
- C. External hemorrhoids
- D. Micronodular cirrhosis
- E. Rectal adenocarcinoma

41. A 69-year-old woman has been bedridden while recuperating from a bout of viral pneumonia complicated by bacterial pneumonia for the past 2 weeks. Physical examination now shows some swelling and tenderness of the right leg, which worsens when she raises or moves the leg. Which of the following terms best describes the condition involving her right leg?

- A. Disseminated intravascular coagulation
- B. Lymphedema
- C. Thromboangiitis obliterans
- D. Thrombosis of deep veins
- E. Varicose veins

42. A healthy 54-year-old woman has noted the increasing prominence of unsightly dilated superficial veins over both lower legs for the past 5 years. Physical examination shows temperature of 37° C, pulse of 70/min, respirations of 14/min, and blood pressure of 125/85 mm Hg. There is no pain, swelling, or tenderness in either lower leg. Which of the following complications is most likely to occur as a consequence of her condition?

- A. Atrophy of the lower leg muscles
- B. Disseminated intravascular coagulation
- C. Gangrenous necrosis of the lower legs
- D. Pulmonary thromboembolism
- E. Stasis dermatitis with ulceration

43. A 48-year-old woman has developed persistent swelling and puffiness in the left arm after a mastectomy with axillary node dissection for breast cancer 1 year ago. She developed cellulitis in the left arm 3 months ago. Physical examination shows firm skin over the left arm and “doughy” underlying soft tissue. The arm is not painful or discolored. Which of the following terms best describes these findings?

- A. Lymphedema
- B. Subclavian arterial thrombosis
- C. Thrombophlebitis
- D. Tumor embolization
- E. Vasculitis

44. A 50-year-old man cuts his right index finger on a sharp metal shard while cleaning debris out of the gate in an irrigation canal. The cut stops bleeding within 3 minutes, but 6 hours later he notes increasing pain in the right arm and goes to his physician. On physical examination, his temperature is 38° C. Red streaks extend from the right hand to the upper arm, and the arm is swollen and tender when palpated. Multiple tender lumps are noted in the right axilla. A blood culture grows group A β-hemolytic streptococcus. Which of the following terms best describes the process that is occurring in this patient’s right arm?

- A. Capillaritis
- B. Lymphangitis
- C. Lymphedema
- D. Phlebothrombosis
- E. Polyarteritis nodosa
- F. Thrombophlebitis
- G. Varices

45. A 46-year-old man has noted increasing abdominal enlargement over the past 15 months. Physical examination shows multiple skin lesions on the upper chest that have central pulsatile cores and measure, from core to periphery, 0.5 to 1.5 cm. Pressing on a core causes a radially arranged array of subcutaneous arterioles to blanch. Laboratory studies show serum glucose of 112 mg/dL, creatinine of 1.1 mg/dL, total protein of 5.8 g/dL, and albumin of 3.4 g/dL. Which of the following underlying diseases is most likely to be present in this patient?

- A. AIDS
- B. Diabetes mellitus
- C. Granulomatosis with polyangiitis
- D. Marfan syndrome
- E. Micronodular cirrhosis
A 21-year-old woman is in the third trimester of an uncomplicated pregnancy. She has noted an enlarging nodule in her mouth for the past 2 weeks. On physical examination there is a 1-cm red nodule on the left lateral gingiva below the first molar. The nodule regresses following delivery. What is this nodule most likely to be?

A  Bacillary angiomatosis  
B  Capillary hemangioma  
C  Cavernous lymphangioma  
D  Glomus tumor  
E  Kaposi sarcoma

A 10-year-old boy has a 2-cm spongy, dull red, circumscribed lesion on the upper outer left arm. The parents state that this lesion has been present since infancy, and the appearance has not appreciably changed. The lesion is excised, and its microscopic appearance is shown in the figure. Which of the following is the most likely diagnosis?

A  Angiosarcoma  
B  Hemangioma  
C  Kaposi sarcoma  
D  Lymphangioma  
E  Telangiectasia

A 6-year-old previously healthy child has had increasing size of his neck for the past year. Physical examination reveals an ill-defined, soft mass deforming the left side of his neck, but no other abnormalities. Surgical resection of the 10 cm mass is attempted, but the borders of the lesion are not discrete. Pathologic examination of the resected tissue shows dilated spaces filled with milky fluid and bounded by thin connective tissue walls. Microscopically, the spaces are lined by flattened endothelium and surrounded by collagenous tissue and smooth muscle with collections of small lymphocytes. Which of the following is the most likely outcome associated with this child’s lesion?

A  Distant metastases  
B  Local recurrence  
C  More neoplasms elsewhere  
D  Opportunistic infection  
E  Sarcomatous transformation

A 35-year-old man is known to have been HIV-positive for the past 10 years. Physical examination now shows multiple reddish purple, nodular skin lesions with the microscopic appearance shown in the figure. These lesions have been slowly increasing for the past year. Which of the following risk factors is most likely to play a role in the development of these skin lesions?

A  Antiretroviral therapy  
B  Epstein-Barr virus infection  
C  Hyperlipidemia  
D  *Mycobacterium avium* complex infection  
E  Sexual intercourse

A 67-year-old woman with glucose intolerance, hypertension, central obesity, and hyperlipidemia has increasing dyspnea from worsening congestive heart failure. Echocardiography shows a left ventricular ejection fraction of 25%. Percutaneous transluminal coronary angioplasty is performed with placement of a left anterior descending arterial stent containing paclitaxel. Which of the following long-term complications in her coronary artery is most likely to be prevented by paclitaxel?

A  Angiosarcoma  
B  Bacillary angiomatosis  
C  Cystic medial degeneration  
D  Giant cell arteritis  
E  Proliferative restenosis  
F  Thromboangiitis obliterans

A 73-year-old woman with hyperhomocystinemia has noted poor circulation in her left leg for the past 2 years. On examination her dorsalis pedis and posterior tibial pulses are barely palpable. CT angiography shows 90% stenosis of the left iliac artery. Which of the following procedures would be most effective in treating this woman?

A  Arterial bypass graft  
B  Clopidogrel therapy  
C  Endovascular stent  
D  Synthetic graft  
E  Venous bypass graft
CHAPTER 11  Blood Vessels

1 E Veins have thin walls compared to their diameter, with less distinct layers, and slow blood flow. These characteristics make them more prone to compression and invasion by malignancies. This is a route for metastases via hematogenous spread. Of the remaining options, lymphatic invasion is an excellent choice, but the intracranial location chosen for this question excludes lymphatics, which are extracranial in the head. Neoplasms can elaborate growth factors to promote endothelial proliferation and capillary formation to form a stromal vascular supply for continued growth. Invasion of a muscular artery by a neoplasm is a rare event.

2 D Fibromuscular dysplasia (FMD) typically involves renal and carotid arteries. Renal arterial stenosis activates the renin-angiotensin system, leading to hypertension. On angiography, FMD appears as a “string of beads” caused by focal medial hyperplasia with thickened fibromuscular ridges adjacent to less involved areas of the arterial wall. This is a surgically correctible cause for hypertension. Arterial dissection is most likely to occur in the proximal aorta with risk factors of hypertension and cystic medial degeneration. Arteriovenous fistulas and berry aneurysms are likely to be intracranial and may rupture and bleed. Renal cell carcinomas may be highly vascular, but a focal mass effect is unlikely to obstruct the main renal artery, and does not account for the TIA. Hypertension could eventually drive atherosclerosis with arterial stenotic lesions lacking focal dilation.

3 A Activated endothelium can undergo many changes. In the acute situation of anaphylaxis, endothelial contraction within minutes leads to exudation of intravascular fluid into the extravascular compartment to produce edema, but this is also quickly reversible, because no permanent vascular injury has occurred. Free radical formation is more typical for endothelial dysfunction when stressors have acted for months to years, as is the case with atherogenesis. Endothelial injury from inflammation over days may lead to up-regulation of protein synthesis and release of regulators of thrombogenesis. Vascular injury may promote vascular endothelial growth factor (VEGF) formation, but in the basal state, procoagulants, anticoagulants, cytokines, and growth factors are elaborated at low levels and balance each other.

4 C Orthostatic hypotension is described. Diminished intravascular volume from fluid loss (sweating, urination, vomiting, diarrhea) or from decreased intake of fluids leads to dehydration. The other options listed cannot act quickly or forcefully enough to compensate for lack of intravascular fluid volume. Subtle changes in total body water and salt that play a role in essential hypertension are not reflected in altered electrolytes. Vomiting and diarrhea could alter acid-base status and electrolytes, but that is not the case here. Autoregulation by sympathetic tone and arteriolar vasoconstriction can adjust blood pressure and flow quickly, though the speed and magnitude of this response is diminished with aging. Adjustments in the synthesis and release of hormones such as renin and aldosterone take days.

5 F This is a classic example of a secondary form of hypertension for which a cause can be determined. In this case, the renal artery stenosis reduces glomerular blood flow and pressure in the afferent arteriole, resulting in renin release by juxtaglomerular cells. The renin initiates angiotensin II-inducred vasoconstriction, increased peripheral vascular resistance, and increased aldosterone, which promotes sodium reabsorption in the kidney, resulting in increased blood volume. Anti-double-stranded DNA is a specific marker for systemic lupus erythematosus. Anti-neutrophil cytoplasmic autoantibodies (ANCA) are markers for forms of vasculitis, such as microscopic polyangiitis or ANCA-associated vasculitis. Some patients with hepatitis B or C infection can develop a mixed cryoglobulinemia with a polyclonal increase in IgG. Renal involvement in such patients is common, and cryoglobulinemic vasculitis then leads to skin hemorrhages and ulceration. Hyperglycemia is a marker for diabetes mellitus, which accelerates the atherogenic process and can involve the kidneys, usually bilaterally, promoting the development of hypertension. HIV infection is not related to hypertension.

6 A The term essential hypertension (now applied when there is no obvious cause for moderate hypertension) was coined before hormonal control of blood pressure was understood, because it was thought that the high pressure was essential to force blood through narrowed arteries. Essential hypertension has several postulated theories for its cause. One theory is that there are defects in renal sodium homeostasis that reduce renal sodium excretion. The kidney retains sodium with water, increasing intravascular fluid volume, which drives increased cardiac output. The increased cardiac output is compensated by increasing peripheral vascular resistance, causing an increase in blood pressure. If angiotensin-converting enzyme (ACE) was absent, blood pressure would decrease because angiotensin I would not be converted to angiotensin II (drugs that act as ACE inhibitors are antihypertensives). An elevated plasma renin level is typical of renovascular hypertension, which can occur with narrowing of a renal artery. Hypertensive patients with hypokalemia also can have hyperaldosteronemia, which can be caused by an aldosterone-secreting adrenal adenoma. Increased urinary catecholamines can indicate increased catecholamine output from a pheochromocytoma.

7 B The pulmonary vasculature is under much lower pressure than the systemic arterial circulation and is much less likely to have endothelial damage, which promotes
atherosclerosis. Atherosclerosis in systemic arteries is most likely to occur where blood flow is more turbulent, a situation that occurs at arterial branch points, such as in the first few centimeters of the coronary arteries or in the abdominal aortic branches. Factors driving systemic arterial atherosclerosis (e.g., hyperlipidemias, smoking, diabetes mellitus, and systemic hypertension) do not operate in the pulmonary arterial vasculature. Pulmonary hypertension is the driving force behind pulmonary atherosclerosis, and it occurs as pulmonary vascular resistance increases, when the pulmonary vascular bed is decreased by either obstructive (e.g., emphysema, as in this patient) or restrictive (e.g., as in scleroderma with pulmonary interstitial fibrosis) diseases. Amyloid deposition usually occurs in small vessels and is driven by underlying conditions such as multiple myeloma or chronic inflammatory diseases. Intimal tears lead to arterial dissection. Mönckeberg arteriosclerosis is typically an incidental finding in systemic arteries. Necrotizing vasculitis occurs in pulmonary vasculature in association with anti-neutrophil cytoplasmic autoantibody (ANCA)-associated granulomatous vasculitis. Phlebothrombosis occurs in peripheral veins and is a risk factor for pulmonary thromboembolism.

**8 F** It is very difficult to keep dietary intake of sodium under 2 g per day because processed and packaged foods have considerable salt for both preservation and flavoring. Just 30 g of potato chips can have more than 150 mg of sodium. Even a single “healthy” can of soup may have half a gram of sodium! Reduced renal sodium excretion may initiate essential hypertension. Decreased sodium excretion may lead sequentially to an increase in fluid volume, increased cardiac output, and peripheral vasoconstriction, thereby elevating blood pressure. Increased salt intake potentiates hypertension. Although alcohol intake may reduce atherogenesis, there is an increased risk for cardiomyopathy as well as liver disease. Reducing protein intake may help patients with chronic renal failure. The caffeine in coffee has neuroendocrine effects, but it is the milk and sugar added that may be most deleterious. Eggs have cholesterol. Milk is a good source of calcium, but it is the milk and sugar added that may be most deleterious. Phlebothrombosis occurs in peripheral veins and is a risk factor for pulmonary thromboembolism.

**9 E** The figure shows an arteriole with marked hyaline thickening of the wall, indicative of hyaline arteriolosclerosis. Diabetes mellitus also can lead to this finding, which is most often seen in kidneys. Sepsis can produce disseminated intravascular coagulation with arteriolar hyaline thrombi. The debilitation that accompanies cancer tends to diminish the vascular disease caused by atherosclerosis. Syphilis can cause a vasculitis involving the vasa vasorum of the aorta. Polyarteritis can involve large to medium-sized arteries in many organs, including the kidneys; the affected vessels show fibrinoid necrosis and inflammation of the wall (vasculitis).

**10 D** Malignant hypertension can suddenly complicate and be superimposed on less severe, benign essential hypertension. The arterioles undergo concentric thickening and luminal narrowing with malignant hypertension, called hyperplastic arteriolosclerosis, and fibrinoid necrosis is a prominent feature. Fibromuscular dysplasia can involve the main renal arteries, with medial hyperplasia producing focal arterial obstruction. This process can lead to hypertension, but not typically malignant hypertension. A granulomatous arteritis is most characteristic of anti-neutrophil cytoplasmic autoantibody (ANCA)-associated granulomatous vasculitis, which often involves the kidney but typically involves lung and other organs. Hyaline arteriolosclerosis is seen with long-standing essential hypertension of moderate severity. These lesions give rise to benign nephrosclerosis. The affected kidneys become symmetrically shrunk and granular because of progressive loss of renal parenchyma and consequent fine scarring. Polyarteritis nodosa produces a vasculitis that can involve the kidney.

**11 A** Older adults with radiographic evidence of calcified arteries often have Mönckeberg arteriosclerosis, beginning in the internal elastic lamina. This is a benign process that is a form of arteriosclerosis, often with no serious sequelae. The distal extremities, pelvis, thyroid, and breast regions are the most common locations. Such focal peripheral arterial calcification is far less likely to be a consequence of atherosclerosis, with diabetes mellitus or with hypercalcemia. Hypertension is most likely to affect small renal arteries, with hyaline or hyperplastic arteriolosclerosis, and calcification is not a major feature, although hypertension also is a risk factor for atherosclerosis.

**12 D** Think about a diet containing teleost fish, which contain oils that may be associated with lower cholesterol and therefore may be protective against atherogenesis. The other listed options are associated with risk for atherosclerosis. C-reactive protein (CRP) is an acute phase reactant that increases in response to inflammatory cytokines, including those released from cells within atheromatous plaques. Hyperhomocysteinemia is a risk factor independent of lipids for atheroma formation. Lipoprotein(a) is a small LDL-like lipid particle that forms an insudate in arterial walls to drive atherogenesis. Trans fats chemically altered via hydrogenation make lipids last longer on the shelf and taste better, but are bad for you by increasing the ratio of LDL to HDL cholesterol.

**13 B** CRP is an acute phase reactant that increases in response to inflammation. It causes endothelial cell activation, promotes thrombosis, and increases leukocyte adheriveness in developing atheromas. Because atherogenesis is partly an inflammatory process, CRP is an independent predictor of
cardiovascular risk. PR3 is one type of anti-neutrophil cytoplasmic autoantibody (ANCA) associated with some vasculitides such as microscopic polyangiitis. Cryoglobulins may be found with some forms of immune complex-mediated vasculitis. The ESR (“sed rate”) is a nonspecific indicator of inflammation and therefore the internist’s least favorite test; the ESR is best known to be markedly elevated with giant cell arteritis. Though platelets play a role in atheroma formation, the actual number of platelets is not a predictor of atherogenesis.

14 B Reduced cholesterol, particularly LDL cholesterol, with the same or increased HDL cholesterol level, indicates a reduced risk of atherosclerotic complications. Atherosclerosis is multifactorial, but modification of diet (i.e., reduction in total dietary fat and cholesterol) with increased exercise is the best method of reducing risk for most individuals. Atherosclerosis is multifactorial, but modification of diet (i.e., reduction in total dietary fat and cholesterol) with increased exercise is the best method of reducing risk for most individuals. Atherosclerotic plaques that have formed can be reduced, albeit over months to years, but plaque regression yield an outcome better than drug or surgical therapy. Glucose is a measure of control of diabetes mellitus. Potassium, calcium, and renin values can be altered with some forms of hypertension, and free radicals increase degradation of LDL.

15 B Diabetes mellitus, suggested here by hyperglycemia, is a significant risk factor for early, accelerated, and advanced atherosclerosis. If a premenopausal woman or a young man has severe coronary atherosclerosis, diabetes mellitus or other metabolic disease such as hyperlipidemia must be suspected as a predisposing factor. “Soft” risk factors that can play a lesser role in the development of atherosclerosis include obesity, stress, and lack of exercise.

16 D Oxidized LDL can be taken up by a special “scavenger” pathway in macrophages; it also promotes monocyte chemotaxis and adherence. Macrophages taking up the lipid become foam cells that begin to form the fatty streak. Smoking, diabetes mellitus, and hypertension all promote free radical formation, and free radicals increase degradation of LDL to its oxidized form. About one third of LDL is degraded to the oxidized form; a higher LDL level increases the amount of oxidized LDL available for uptake into macrophages. C-reactive protein is a marker for inflammation, which can increase with more active atheroma and thrombus formation and predicts a greater likelihood of acute coronary syndromes. Increased homocysteine levels promote atherogenesis through endothelial dysfunction. Lp(a), an altered form of LDL that contains the apo B-100 portion of LDL linked to apo A, promotes lipid accumulation and smooth muscle cell proliferation. Platelet-derived growth factor promotes smooth muscle cell proliferation. VLDL is formed in the liver and transformed in adipose tissue and muscle to LDL.

17 B Lipoprotein(a) is an altered form of LDL cholesterol that has the apolipoprotein B-100 portion of LDL linked to apolipoprotein A, and an increase in Lp(a) is independently associated with a risk for endothelial dysfunction and atherosclerosis. Apolipoprotein E promotes metabolism and clearance of LDL. Drugs such as statins that affect LDL receptor activity do not affect Lp(a) concentration. Early in the course of atheroma formation, angiotensin and its receptor may play a role in development of hypertension, which then becomes a risk factor for atherosclerosis. Endothelin is a vasoconstrictor with no known role in atherogenesis. Decreased factor VIII leads to abnormal bleeding. Von Willebrand factor is required for normal platelet adhesion to collagen, and its absence leads to abnormal bleeding.

18 B The initial response of an arterial wall to injury is intimal thickening with neointimal smooth muscle cell proliferation and production of increased intimal extracellular matrix. Note that “injury” can be caused by inflammation, immune reactions, and toxins as well as the local physical trauma from hypertension and abnormal flow. The trauma does not produce immediate injury with endothelial cellular disruption, but a response of endothelial dysfunction that signals smooth muscle cell migration. This process takes years to show changes of vascular narrowing. As atheromatous plaques progress, there is participation by lymphocytes producing cytokines, as well as monocytes that are transformed to macrophages that accumulate lipid to evolve into foam cells. Eventually plaque disruption may incite platelet aggregation.

19 C Atherosclerosis is thought to result from an initial endothelial injury and the subsequent chronic inflammation and repair of the arterial intima. All risk factors, including smoking, hyperlipidemia, and hypertension, cause biochemical or mechanical injury to the endothelium with resulting dysfunction that initiates smooth muscle migration with proliferation, as well as lymphocyte and monocyte-macrophage infiltration. Formation of foam cells occurs after the initial endothelial injury. Vasomotor tone does not play a major role in atherogenesis. Inhibition of LDL oxidation should diminish atheroma formation. Although lipoprotein receptor alterations can occur in some inherited conditions, these account for only a fraction of cases of atherosclerosis, and other lifestyle conditions do not affect their action.

20 D The slightly raised, pale lesions shown in the figure are called fatty streaks and are seen in the aorta of almost all children older than 10 years. They are thought to be precursors of atheromatous plaques. T cells are present early in the pathogenesis of atherosclerotic lesions and are believed to activate monocytes, endothelial cells, and smooth muscle cells by secreting cytokines. T cells adhere to VCAM-1 on activated endothelial cells and migrate into the vessel wall. These T cells, activated by some unknown mechanism, secrete...
various proinflammatory molecules that recruit and activate monocytes and smooth muscle cells and perpetuate chronic inflammation of the vessel wall. Fatty streaks cause no disturbances in blood flow and are discovered incidentally at autopsy. All of the other lesions described are seen in fully developed atheromatous plaques. The histologic features of such plaques include a central core of lipid debris that can have cholesterol clefts and can be calcified. There is usually an overlying cap of smooth muscle cells. Hemorrhage is a complication seen in advanced atherosclerosis. Foam cells, derived from smooth muscle cells or macrophages that have ingested lipid, can be present in all phases of atherogenesis.

PBD9 496–497  BP9 340–341  PBD8 502  BP8 346–350

21 D Growth factor release from activated platelets, macrophages, and vascular wall cells induces smooth muscle cell recruitment, medial smooth muscle migrate into the intima, proliferate, and synthesize extracellular matrix (ECM) in much the same way that fibroblasts fill in a wound. Endothelial injury initiates atherogenesis, but endothelial cells do not form a significant part of an atheroma. Platelets do not synthesize ECM. T cells secrete inflammatory cytokines that activate macrophages, endothelial cells, and smooth muscle cells.


22 B The figure shows an arterial lumen that is markedly narrowed by atheromatous plaque complicated by calcification. Hypercholesterolemia with elevated LDL and decreased HDL levels is a key risk factor for atherogenesis. Although platelets participate in forming atheromatous plaques, their number is not of major importance. Thrombocytosis can result in thrombosis or hemorrhage. Levels of Lp(a) and homocysteine, if elevated, increase the risk of atherosclerosis. Syphilis (positive VDRL test result) produces endarteritis obliterans (Buerger disease) is a rare condition in which muscular arteries become occluded in the lower extremities in smokers.


23 D Atheromatous plaques can be complicated by various pathologic alterations, including hemorrhage, ulceration, thrombosis, and calcification. These processes can increase the size of the plaque and narrow the residual arterial lumen. Although atherosclerosis is a disease of the intima, in advanced disease, the expanding plaque compresses the media. This causes thinning of the media, which weakens the wall and predisposes to aneurysm formation.

PBD9 498–501  BP9 342–343  PBD8 503–504  BP8 348–351

24 B Aspirin (acetylsalicylic acid) inhibits the cyclooxygenase pathway of arachidonic acid metabolism and inhibits platelets that participate in thrombogenesis. When atheromatous plaques have progressed to the point of symptomatic occlusion and subsequent angina, they are likely to be unstable plaques that may rupture, ulcerate, or erode to promote thrombosis. A “baby” aspirin containing 80 mg (325 mg in the “adult” tablet) taken once a day may reduce this thrombotic risk, and also prevent the significantly increased risk for hemorrhage that is associated with higher doses. Acetaminophen is primarily analgesic in action. Ibuprofen is a nonsteroidal anti-inflammatory drug (NSAID) that has no significant effect upon atheroma progression. Paclitaxel is one type of drug (initially used as an anticancer agent) used in drug-eluting stents placed in coronary arteries following angioplasty to deter restenosis. Propranolol is a beta-blocker that is employed as an antihypertensive agent.

PBD9  BP9 342–343  PBD8 498–499  BP8 347–348

25 C Abdominal aneurysms are most often related to underlying aortic atherosclerosis. This patient has multiple risk factors for atherosclerosis, including diabetes mellitus, hypertension, and smoking. When the aneurysm reaches this size, there is a significant risk of rupture. An aortic dissection is typically a sudden, life-threatening event with dissection of blood out of the ascending aortic lumen, typically into the chest, without a pulsatile abdominal mass. The risk factors for atherosclerosis and hypertension also underlie aortic dissection. An arteriovenous fistula can produce an audible bruit on auscultation. Classic polyarteritis nodosa (PAN) can produce small microaneurysms in small arteries, most often renal and mesenteric. Takayasu arteritis typically involves the aortic arch and branches in children. Thromboangiitis obliterans (Buerger disease) is a rare condition in which muscular arteries become occluded in the lower extremities in smokers.

PBD9 501–503  BP9 344–346  PBD8 707  BP8 357–359

26 A His abdominal CT scan shows a 6-cm diameter enlargement of the abdominal aorta, which is an atherosclerotic abdominal aortic aneurysm. Diabetes mellitus, an important risk factor for atherosclerosis, must be suspected if a younger man or premenopausal woman has severe atherosclerosis. Marfan syndrome is a risk for aortic dissection starting in a dilated ascending aorta. Polyarteritis nodosa does not typically involve the aorta. Osteitis “soft” risk factor for atherosclerosis, also contributes to type 2 diabetes mellitus; however, the extent of atherosclerotic disease in this patient suggests early-onset diabetes mellitus, which is more likely to be type 1. Systemic lupus erythematosus produces small arteriolar vasculitis. Syphilitic arthritis, a feature of tertiary syphilis, most often involves the thoracic aorta, but it is rare, and most thoracic aortic aneurysms nowadays are likely to be caused by atherosclerosis.


27 A This description is most suggestive of syphilitic aortitis, a complication of tertiary syphilis, with characteristic involvement of the thoracic aorta. The history also suggests tabes dorsalis and neurosyphilis. Obliterative endarteritis is not a feature of other forms of vasculitis. High-titer double-stranded DNA antibodies are diagnostic of systemic lupus erythematosus, and the test result for P-ANCA (antibodies mainly directed at myeloperoxidase) is positive in various vasculitides, including microscopic polyangiitis. Ketonuria can occur in individuals with diabetic ketoacidosis. A high sedimentation rate is a nonspecific marker of inflammatory diseases.

PBD9 502, 504  BP9 345–346  PBD8 507–508  BP8 359–360
Patients may have polymyalgia rheumatica. Between hypertension and giant cell arteritis, but some patients have other complications of giant cell arteritis. There is no association between hypertension and giant cell arteritis and Takayasu arteritis may do so, although they are unlikely to weaken the wall to the point of rupture. 

A sudden tear in the proximal aortic intima allows blood to enter the space between layers within the wall of the aorta. This blood may pass through the aortic wall, around great vessels, and into the pericardial cavity, as in this case with cardiac tamponade. Blood may enter the chest cavity, causing hemotorax. Hypertension is the most common risk factor for aortic dissection. In contrast, a false aneurysm is characterized by formation of a hematoma by extravasated blood, but it communicates with the vascular lumen; a true aneurysm includes all three layers of the arterial wall. Atherosclerosis can be associated with hypertension, but it involves arterioles, typically in kidneys, not the aorta. Thrombosis of extravasated blood from a dissection can occur, but this is not the primary lesion. Vasculitis does not often involve the aorta, but giant cell arteritis and Takayasu arteritis may do so, although they are unlikely to weaken the wall to the point of rupture. 

Giant cell (temporal) arteritis typically involves large to medium-sized external carotid artery branches in the head (especially temporal arteries), but also vertebral and ophthalmic arteries. Involvement of the latter can affect vision. Because involvement of the kidney, lung, and peripheral arteries of the extremities is much less common, renal failure, hemoptysis, and gangrene of toes are unusual complications of giant cell arteritis. There is no association between hypertension and giant cell arteritis, but some patients may have polymyalgia rheumatica. 

Takayasu arteritis leads to “pulseless disease,” because of involvement of the aorta (particularly the arch) and branches such as coronary, carotid, and renal arteries, which results in granulomatous inflammation, aneurysm formation, and dissection. Fibrosis is a late finding, and the pulmonary arteries also can be involved. Aortic dissection is an acute problem that, in older adults, is driven by atherosclerosis and hypertension, although this patient is within the age range for complications of Marfan syndrome, which causes cystic medial degeneration of the aorta. Kawasaki disease affects children and is characterized by an acute febrile illness, coronary arteritis with aneurysm formation and thrombosis, skin rash, and lymphadenopathy. Microscopic polyangiitis affects arterioles, capillaries, and venules with a leukocytoclastic vasculitis that appears at a similar stage in multiple organ sites (in contrast to classic polyarteritis nodosa, which causes varying stages of acute, chronic, and fibrosing lesions in small to medium-sized arteries). Tertiary syphilis produces an endoarteritis with proximal aortic dilation. Thromboangiitis obliterans (Buerger disease) affects small to medium-sized arteries of the extremities and is strongly associated with smoking. 

Classic polyarteritis nodosa (PAN) has segmental involvement of medium-sized arteries with aneurysmal dilation in the renal and mesenteric vascular beds (e.g., abdominal pain, melena). PAN can affect many organs at different times. Although the cause of PAN is unknown, about 30% of patients have hepatitis B surface antigen that presumably forms immune complexes that damage vascular walls. In contrast to microscopic polyangiitis, PAN has less of an association with anti-neutrophil cytoplasmic autoantibody (ANCA). A collagen vascular disease with a positive ANA test result, such as systemic lupus erythematosus, may produce a vasculitis, but not in the pattern seen here; the affected vessels are smaller. Vasculitis with HIV infection is uncommon. The Sc1-70 autoantibody is indicative of scleroderma, which can produce renal failure. The rapid plasma reagin (RPR) is a serologic test for syphilis; an endoarteritis of the vasa vasorum can occur in syphilis. 

Muco cutaneous lymph node syndrome, or Kawasaki disease, involves large, medium-sized, and small arteries. Cardiovascular complications occur in 20% of cases and include thrombosis, ectasia, and aneurysm formation of coronary arteries. Asthma can be seen in association with Churg-Strauss vasculitis. Glomerulonephritis is a feature of anti-neutrophil cytoplasmic autoantibody (ANCA)-associated granulomatous vasculitis and of some autoimmune diseases such as systemic lupus erythematosus. Intracranial hemorrhage can occur with septic emboli to peripheral cerebral arteries, producing mycotic aneurysms that can rupture. Pulmonary hypertension can also complicate Takayasu arteritis, but is not as life-threatening as the coronary artery disease. 

Anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis (granulomatosis with polyangiitis) is a form of hypersensitivity reaction to an unknown antigen characterized by necrotizing granulomatous inflammation that typically involves small to medium-sized vessels, although many organ sites may be affected. Pulmonary and renal involvement can be life-threatening. C-ANCA (antibodies mainly directed against neutrophil proteinase 3) are found in more than 90% of cases. Fibromuscular dysplasia is a hyperplastic medial disorder, usually involving renal and carotid arteries; on angiography, it appears as a “string of beads” caused by thickened
fibromuscular ridges adjacent to less involved areas of the arterial wall. Granuloma pyogenicum is an inflammatory response that can produce a nodular mass, often on the gingiva or the skin. Kaposi sarcoma can produce plaquelike to nodular masses that are composed of irregular vascular spaces lined by atypical-appearing endothelial cells; skin involvement is most common, but visceral organ involvement can occur. Polyarteritis nodosa most often involves small muscular arteries, and sometimes veins. It causes necrosis and microaneurysm formation followed by scarring and vascular occlusion, mainly in the kidney, gastrointestinal tract, and skin of young to middle-aged adults. Takayasu arteritis is seen mainly in children and involves the aorta (particularly the arch) and branches such as the coronary and renal arteries, with granulomatous inflammation, aneurysm formation, and dissection.

**PBD9 511–512**  **BP9 353–354**  **PBD8 516–517**  **BP8 363, 367–368**

**35 B** Microscopic polyangiitis involves small vessels, typically capillaries. Kidneys and lungs are commonly involved, but many organs can be affected. There may be an underlying immune disease, chronic infection, or drug reaction. Giant cell arteritis typically involves arterial branches of the external carotid, most often the temporal artery. Microabscesses may be present with an infectious process, or with thromboangiitis obliterans (Buerger disease), which typically involves lower extremities. Mycotic aneurysms occur when a focus of infection, often from a septic embolus, weakens an arterial wall so that it bulges out. Perivascular eosinophilic infiltrates may be seen with Churg-Strauss syndrome, which typically involves the lungs.

**PBD9 510–511**  **BP9 353**  **PBD8 515–516**  **BP8 366-367**

**36 E** Thromboangiitis obliterans (Buerger disease), which affects small to medium-sized arteries of the extremities, is strongly associated with smoking. This disease may eventually involve adjacent peripheral veins and nerves. Syphilis can be treated with antibiotics, but it mainly produces an aortitis. Immunosuppressive therapy is not highly effective. Renal involvement does not occur. Although peripheral vascular disease with atherosclerosis is a typical finding in diabetes mellitus, vasculitis is not.

**PBD9 512**  **BP9 354**  **PBD8 517**  **BP8 368**

**37 B** In children, Henoch-Schönlein purpura is the multisystemic counterpart of the IgA nephropathy seen in adults. The immune complexes formed with IgA produce the vascular response that affects mainly arterioles, capillaries, and venules in skin, gastrointestinal tract, and kidney. In older adults, giant cell arteritis is seen in external carotid branches, principally the temporal artery unilaterally. Polyarteritis nodosa is seen most often in small muscular arteries and sometimes veins, with necrosis and microaneurysm formation followed by scarring and vascular occlusion. This occurs mainly in the kidney, gastrointestinal tract, and skin of young to middle-aged adults. Takayasu arteritis is seen mainly in children and involves the aorta (particularly the arch) and branches such as coronary and renal arteries, with granulomatous inflammation, aneurysm formation, and dissection. Telangiectasias are small vascular arborizations seen on skin or mucosal surfaces.

**PBD9 512–513**  **BP9 349**  **PBD8 512, 517**  **BP8 366**

**38 D** Infectious endocarditis is likely present in this woman, and portions of the vegetations often dislodge and embolize. These are septic emboli carrying organisms that can start growing and produce local arterial destruction wherever they are carried. The destruction of a vascular wall by an infectious process is uncommon, but can result from spread of local infection or via embolization. A so-called mycotic aneurysm can be due to any infectious agent that weakens an arterial wall so that it bulges out—an aneurysmal dilation. Basilar angiomegaly produces a focal vascular proliferation, typically on the skin, of an immunocompromised person infected with *Bartonella* spp. Hyperplastic arteriolar sclerosis is most often found in the kidneys of persons with malignant hypertension, some of whom may have underlying systemic sclerosis. Lymphatic channels are not found within the brain. Phlebothrombosis is typically found in large veins of the legs and pelvis, most often following prolonged immobilization.

**PBD9 502, 513**  **BP9 345, 355**  **PBD8 517**  **BP8 357, 369**

**39 E** The “red, white, and blue” changes shown represent Raynaud phenomenon, which can be a primary exaggerated vasomotor response with no underlying disease. In older persons, an underlying disease, such as an autoimmune disease, should be sought. Hyperviscosity may underlie this phenomenon. In younger persons it is “primary” and likely vasomotor hyperreactivity. Calcification with medial calcific sclerosis tends to involve arteries that are small and muscular, but larger than those of hands or feet; it is often an incidental finding on a radiograph. Hypertension may drive atherosclerosis, but not marked vasoconstriction. Thrombosis is unlikely to develop and subside so quickly. Vasculitis likewise is not an evanescent phenomenon.

**PBD9 513**  **BP9 355**  **PBD8 518**  **BP8 369–370**

**40 C** The hemorrhoidal veins can become dilated from venous congestion. They are derived from ecoderm, covered by squamous epithelium, and innervated by somatic sensory nerves. External hemorrhoids are most common in patients with chronic constipation, but a pregnant uterus presses on pelvic veins to produce similar congestion, which promotes hemorrhoidal vein dilation. Filarial infections can affect lymphatics, including those in the inguinal region, and produce lymphedema. Polyarteritis does not affect veins. Portal hypertension with cirrhosis is most likely to dilate submucosal esophageal veins, but internal hemorrhoidal veins occasionally can be affected. Cirrhosis would be rare at this patient’s age. Carcinomas are also uncommon at this age, and they are not likely to obstruct venous flow.

**PBD9 514**  **BP9 356**  **PBD8 518–519**  **BP8 370**

**41 D** Phlebothrombosis (but most often called thrombophlebitis) is a common problem that results from venous
stasis with prolonged immobilization. Phlebothrombosis may be a better (but less often used) term because there is little or no inflammation, but the former term is well established. Disseminated intravascular coagulation more often results in hemorrhage, and edema is not the most prominent manifestation. Lymphedema takes longer than 2 weeks to develop and is not caused by bed rest alone. Thromboangitis obliterans is a rare form of arteritis that results in pain and ulceration of extremities. Varicose veins are dilated, tortuous superficial veins and can thrombose, but they are not related to bed rest, and they do not predispose to pulmonary thromboembolism, as does thrombosis of deeper, larger veins.

42 E Chronic peripheral venous stasis results in hemosiderin deposition and dermal fibrosis with brownish discoloration and skin roughening. Focal ulceration can occur over the varicosities, but extensive gangrene similar to that seen in arterial atherosclerosis does not occur. Hippocrates described varicosities circa 370 BCE and recommended compression for ulceration. In 1896 Australian surgeon Jerry Moore described ligation of the great saphenous vein, a procedure still in use. The varicosities involve only the superficial set of veins, which can thrombose, but they are not the source of thromboemboli, as are the larger, deep leg veins. The varicosities do not affect muscle; however, diminished muscular support for veins to “squeeze” blood out for venous return can predispose to formation of varicose veins. The thromboses in superficial leg veins do not lead to disseminated intravascular coagulation, because there is no acute vascular injury to promote ongoing coagulopathy. Venous infarction is uncommon, because of collateral deep and superficial veins for venous return.

43 A A mastectomy with axillary lymph node dissection leads to disruption and obstruction of lymphatics in the axilla. Such obstruction to lymph flow gives rise to lymphedema, a condition that can be complicated by cellulitis. Arterial thrombosis produces ischemia distal to the obstruction. Thrombophlebitis from venous stasis is a complication seen more commonly in the lower extremities. An arterial thrombosis can lead to a cold, blue, painful extremity. Tumor emboli are generally small but uncommon. Vasculitis is not a surgical complication.

44 B The red streaks represent lymphatic channels through which the acute infection is draining to axillary lymph nodes, and these nodes drain to the right lymphatic duct and into the right subclavian vein (lymphatics from the lower body and left upper body drain to the thoracic duct). Capillaritis is most likely to be described with inflammation involving the lungs. Lymphedema occurs with blockage of lymphatic drainage and develops over a longer period without significant acute inflammation. Phlebothrombosis and thrombophlebitis describe thrombosis in veins with stasis and minimal inflammation, typically in the pelvis and lower extremities. Classic polyarteritis nodosa (PAN) involves small to medium-sized muscular arteries, typically the renal and mesenteric branches. Varices are veins dilated from obstruction to venous drainage.

45 E Spider telangiectasias are a feature of micronodular cirrhosis, typically as a consequence of chronic alcohol abuse. They are thought to be caused by hyperestrogenism (estrogen excess) that results from hepatic damage with reduced clearance of circulating steroids. The most common vascular skin lesion in patients with AIDS is Kaposi sarcoma, which is a neoplasm that manifests as one or more irregular, red-to-purple patches, plaques, or nodules. Diabetes mellitus, with its accelerated atherosclerosis, is most likely to result in ischemia or gangrene. Vasculitis does not tend to produce skin telangiectasias. The vascular involvement in Marfan syndrome is primarily in the aortic arch with cystic medial degeneration.

46 B A so-called pyogenic granuloma is described. Half of them are related to trauma and represent an exuberant repair reaction with granulation tissue. They can ulcerate and bleed. Bacillary angiomatosis represents a vascular proliferation that occurs in immunocompromised patients infected with Bartonella organisms. Cavernous lymphangiomas can be deforming mass lesions in the neck region of children, and though technically benign they do not have well-defined borders and can be difficult to excise. A glomus tumor typically presents as a painful nodule beneath the fingernail. There are various forms of Kaposi sarcoma (classic, endemic, AIDS-related) but they all appear as reddish purple patches, plaques, or nodules on the skin and are associated with Kaposi sarcoma herpes virus (KSHV), also called human herpesvirus 8 (HHV8).

47 B The figure shows dilated, endothelium-lined spaces filled with RBCs. The circumscribed nature of this lesion and its long, unchanged course suggest its benign nature. The vascular spaces of a hemangioma may be small, resembling capillaries, or large and cavernous. Angiosarcomas are large, rapidly growing malignancies in adults. Kaposi sarcoma is uncommon in its endemic form in childhood, and it is best known as a neoplastic complication associated with HIV infection. Lymphangiomas, seen most often in children, tend to be more diffuse and are not blood-filled. A telangiectasia is a radial array of subcutaneous dilated arteries or arterioles surrounding a central core that can pulsate.

48 B The lesion is a cavernous lymphangioma. Although histologically benign, such lesions have a tendency to become large and extend around adjacent structures, complicating removal. Lymphangiomas are unlikely to be associated with syndromes involving multiple neoplasms, and
there is no association with infections. The cystic hygroma of Turner syndrome has a similar appearance, but is bilateral, nonprogressive, and may account for the “web neck” appearance.

PBD9 517  BP9 358–359  PBD8 522  BP8 373

49 E Infection by HHV8, also called Kaposi sarcoma herpes virus (KSHV), is associated with Kaposi sarcoma (KS) and can be acquired as a sexually transmitted disease. KS is a complication of AIDS. Individuals with HIV infection can be infected with various viruses, including cytomegalovirus (CMV) and Epstein-Barr virus (EBV), but these viruses have no etiologic association with KS. EBV is a factor in the development of non-Hodgkin lymphoma, and CMV can cause pneumonitis or retinitis, or it can be disseminated. Hyperlipidemia may complicate antiretroviral therapy, but does not produce skin nodules. MAC infection can be seen in HIV-infected patients as well, but it produces small granulomas, mainly in tissues of the mononuclear phagocyte system.

PBD9 518–519  BP9 360–361  PBD8 523–524  BP8 375–376

50 E She has metabolic syndrome, a risk for coronary atherosclerosis. Following angioplasty, there is often intimal thickening that causes restenosis. The wire stent holds the lumen open and the paclitaxel limits smooth muscle hyperplasia. Atherosclerosis is not a risk factor for neoplasia. Bacillary angiomatosis is caused by Bartonella organisms (cat-scratch disease) and most often produces a red skin nodule. Cystic medial degeneration is a feature of Marfan syndrome with aortic dissection. Giant cell arteritis most often involves external carotid branches such as temporal arteries; it is not related to atherosclerosis. Thromboangiitis obliterans is a rare form of vasculitis involving lower extremities in smokers.

PBD9 520–521  BP9 338, 362  PBD8 528  BP8 378

51 C Endovascular stent placement can be done without major surgery, because the graft can be deployed percutaneously. Arterial bypass grafting is most often used from left internal mammary (thoracic) artery to left anterior descending coronary artery, or by harvesting a portion of radial artery to bypass a stenotic coronary artery. Clopidogrel is an antiplatelet agent to help prevent thrombosis, and patients with such a risk, including severe atherosclerosis or those with stent or graft placement, may receive such a drug. Endovascular stents have largely replaced synthetic grafts, which need major surgery for placement. Saphenous vein bypass grafts are often used for coronary artery bypass grafting. All of these grafts may be complicated by thrombosis, restenosis, or both.

PBD9 520–521  BP9 362–363  PBD8 526  BP8 377–378
1. An 82-year-old woman has had increasing fatigue for the past 2 years. During this time, she has experienced paroxysmal dizziness and syncope. On physical examination, she is afebrile. Her pulse is 44/min, respirations are 16/min, and blood pressure is 100/65 mm Hg. On auscultation, the lungs are clear, and no murmurs are heard. An echocardiogram shows a normal-sized heart with normal valve motion and estimated ejection fraction of 50%. After parasympathetic (vagal) stimulation, the heart rate slows and becomes irregular. An abnormality involving which of the following is most likely to be present in this patient?

A. Atrioventricular node
B. Bundle of His
C. Left bundle branch
D. Parasympathetic ganglion
E. Right bundle branch
F. Sinoatrial node
G. Sympathetic ganglion

2. A neonate developing normally has a newborn checkup. On physical examination, there is a systolic murmur. Echocardiography reveals a muscular defect of the intraventricular septum. A checkup 30 years later fails to reveal either a murmur or a flow defect between the ventricles. Which of the following cells most likely proliferated and led to disappearance of the defect?

A. Adipocytes
B. Conduction cells
C. Endothelial cells
D. Fibroblasts
E. Mesothelial cells
F. Stem cells

3. A 66-year-old man has had cough and worsening shortness of breath for 3 years. On examination, there is dullness to percussion at both lung bases and poorly audible breath sounds. On physical examination, pulse is 77/min and BP is 110/80 mm Hg. He does not have anginal pain. His liver span is increased to 14 cm. He has pitting edema to his knees. Jugular venous distention is noted to the angle of the jaw at 45-degree elevation of his head while lying down. Which of the following is most likely causing his heart disease?

A. Atrial myxoma
B. Essential hypertension
C. Hyperlipidemia
D. Rheumatic fever
E. Smoking

4. A 62-year-old woman has had increasing dyspnea for the past 2 years. She now awakens at night with air hunger and cough productive of frothy sputum. On examination, she has rales in all lung fields. Her point of maximal impulse is strong and displaced laterally. Echocardiography shows a decreased ejection fraction of 30% with concentric increase in left ventricular wall size. The valves appear normal. Which of the underlying diseases does she have?

A. Amyloidosis
B. Cardiomyopathy
C. Hypertension
D. Myocarditis
E. Pericarditis

5. A 41-year-old woman has been awakened at night with “air hunger” for the past year. She notes sleeping better while sitting up in bed. Her serum B-type natriuretic peptide is >400 pg/mL (very high). What cardiac disease best explains her condition?

A. Atrial myxoma
B. Fibrinous pericarditis
C. Giant cell myocarditis
D. Libman-Sacks endocarditis
E. Rheumatic valvulitis
6. A 50-year-old man has had increasing abdominal discomfort and swelling of his legs for the past 2 years. He has smoked cigarettes for 35 years. On physical examination, he has jugular venous distention, even when sitting up. The liver is enlarged and tender and can be palpated 10 cm below the right costal margin. Pitting edema is observed on the lower extremities. A chest radiograph shows bilateral diaphragmatic flattening, pleural effusions, and increased lucency of lung fields. Thoracentesis on the right side yields 500 mL of clear fluid with few cells. Which of the following is most likely to be the underlying disease in this patient?
   A. Acute myocardial infarction  
   B. Chronic bronchitis  
   C. Primary pulmonary hypertension  
   D. Pulmonary valve stenosis  
   E. Tricuspid valve stenosis

7. An infant born at term is noted to have cyanosis during the first week of life. On examination a heart murmur is auscultated. Abnormal findings with echocardiography include an overriding aorta, ventricular septal defect, right ventricular thickening, and pulmonic stenosis involving the fetal heart. This infant is most likely to have an inherited mutation involving which of the following genes?
   A. β-Myosin heavy chain (β-MHC)  
   B. Fibrillin 1 (FBN1)  
   C. KCNQ1  
   D. NOTCH2  
   E. Transthyretin (TTR)

8. Following an uncomplicated pregnancy, a term infant appears normal at birth, but at 1 day of life the infant develops respiratory distress. On physical examination the infant has tachypnea, tachycardia, and cyanosis. There is an S1 ejection click and a split S2 with prominent P sound. A radiograph shows normal heart size but prominent hilar vascular markings. Echocardiography shows a small left atrium, large right atrium, normally sized ventricles, widely patent foramen ovale, and normally positioned aorta and pulmonary trunk. What type of congenital heart disease does this infant most likely have?
   A. Atrial septal defect  
   B. Coarctation of the aorta, preductal type  
   C. Patent ductus arteriosus  
   D. Tetralogy of Fallot  
   E. Total anomalous pulmonary venous connection

9. A 77-year-old woman fell and fractured her ankle. She has spent most of her time in bed for the past 16 days. She develops sudden chest pain, dyspnea, and diaphoresis. On examination she has left thigh swelling and tenderness. A chest CT shows areas of decreased attenuation in the right and left pulmonary arteries. A day later she has difficulty speaking. MR angiography shows focal occlusion of a left middle cerebral artery branch. Which of the following cardiac abnormalities is she most likely to have?
   A. Atrial myxoma  
   B. Infective endocarditis  
   C. Nonbacterial thrombotic endocarditis  
   D. Patent foramen ovale  
   E. Ventricular aneurysm

10. A 5-year-old child is not as active as other children his age. During the past 8 months, the child has had multiple episodes of respiratory difficulty following exertion. On physical examination, his temperature is 37°C, pulse is 81/min, respiration are 19/min, and blood pressure is 95/60 mm Hg. On auscultation, a loud holosystolic murmur is audible. There are diffuse crackles over the lungs bilaterally, with dullness to percussion at the bases. A chest radiograph shows a prominent left heart border, pulmonary interstitial infiltrates, and blunting of the costodiaphragmatic recesses. The representative gross appearance of the child’s heart is shown in the figure. Which of the following additional pathologic conditions would most likely develop in this child?
   A. Aortic regurgitation  
   B. Coronary atherosclerosis  
   C. Nonbacterial thrombotic endocarditis  
   D. Pulmonary hypertension  
   E. Restrictive cardiomyopathy

11. A 3-year-old child is developing normally. Physical examination reveals a low-pitched cardiac murmur. An echocardiogram shows the presence of an ostium secundum, with a 1-cm defect. Which of the following abnormalities is most likely to be found in this child?
   A. Cyanosis at rest  
   B. Left-to-right shunt  
   C. Mural thrombosis  
   D. Pericardial effusion  
   E. Pulmonary hypertension
A 2-year-old child had an illness 1 year ago characterized by a high fever. *Staphylococcus epidermidis* was cultured from the blood. The child was given antibiotic therapy and recovered. Now, on physical examination, a harsh, waxing and waning, machinery-like murmur is heard on auscultation of the upper chest. A chest radiograph shows prominence of the pulmonary arteries. Echocardiography shows all cardiac valves to be normal in configuration. Laboratory studies show normal arterial oxygen saturation level. Which of the following congenital heart diseases is most likely to explain these findings?

A  Aortic atresia
B  Aortic coarctation
C  Atrial septal defect
D  Patent ductus arteriosus
E  Tetralogy of Fallot
F  Total anomalous pulmonary venous return

A 5-year-old girl who is below the 5th percentile for height and weight for age has exhibited easily fatigability since infancy. On physical examination, she appears cyanotic. Her temperature is 37° C, pulse is 82/min, respirations are 16/min, and blood pressure is 105/65 mm Hg. Pulse oximetry shows decreased oxygen saturation. One month later, she has fever and obtundation. A cerebral CT scan shows a right parietal, ring-enhancing, 3-cm lesion. Which of the following congenital heart diseases is the most likely diagnosis?

A  Atrial septal defect
B  Bicuspid aortic valve
C  Coarctation of the aorta
D  Patent ductus arteriosus
E  Truncus arteriosus
F  Ventricular septal defect

In a clinical study of tetralogy of Fallot, patients are examined before surgery to determine predictors observed on echocardiography that correlate with the severity of the disease and the need for more careful monitoring. A subset of patients is found to have more severe congestive heart failure, poor exercise tolerance, and decreased arterial oxygen saturation levels. Which of the following is most likely to predict a worse clinical presentation for these patients?

A  Degree of pulmonary stenosis
B  Diameter of the tricuspid valve
C  Presence of an atrial septal defect
D  Size of the ventricular septal defect
E  Thickness of the left ventricle

A 27-year-old woman gives birth to a term infant after an uncomplicated pregnancy and delivery. The infant is cyanotic at birth. Two months later, physical examination shows the infant to be at the 37th percentile for height and weight. The representative gross appearance of the infant's heart is shown in the figure. What is the most likely diagnosis?

A  Aortic stenosis
B  Pulmonic stenosis
C  Tetralogy of Fallot
D  Transposition of the great vessels
E  Truncus arteriosus

A 15-year-old boy complains of pain in his legs when he runs more than 300 m. Physical examination shows temperature, 36.8° C; pulse, 76/min; respirations, 22/min; and blood pressure, 165/90 mm Hg. The radial pulses are 4+, and the dorsalis pedis pulses are 1+. Arterial blood gas measurement shows a normal oxygen saturation level. Which of the following congenital cardiovascular anomalies is most likely to be present in this patient?

A  Aortic valve stenosis
B  Coarctation of the aorta
C  Patent ductus arteriosus
D  Transposition of the great arteries
E  Tricuspid valve atresia
17 A 21-year-old primigravida gives birth at term to a 2800-g infant with no apparent external anomalies. The next day, the infant develops increasing respiratory distress and cyanosis. Echocardiography reveals a slitlike left ventricular chamber, small left atrium, and atretic aortic and mitral valves. Through which of the following structures could blood from the lungs most likely have reached the infant’s systemic circulation?

A Anomalous venous return
B Foramen ovale
C Patent ductus arteriosus
D Right fourth aortic arch
E Truncus arteriosus
F Ventricular septal defect

18 A 60-year-old man has had angina on exertion for the past 6 years. A coronary angiogram performed 2 years ago showed 75% stenosis of the left circumflex coronary artery and 50% stenosis of the right coronary artery. For the past 3 weeks, the frequency and severity of his anginal attacks have increased, and pain sometimes occurs even when he is lying in bed. On physical examination, his blood pressure is 110/80 mm Hg, and pulse is 85/min with irregular beats. An ECG shows ST segment elevation. Laboratory studies show serum glucose, 188 mg/dL; creatinine, 1.2 mg/dL; and troponin I, 1.5 ng/mL. Which of the following is most likely to explain these findings?

A Atheromatous plaque fissure with thrombosis
B Constrictive pericarditis with calcification
C Endomyocardial fibrosis
D Extensive myocardial fiber hypertrophy
E Left ventricular mural thrombosis
F Mitral valve prolapse with regurgitation

19 A 37-year-old woman has the sudden onset of chest pain. On examination she is afebrile but tachycardic and hypotensive. An ECG shows ST segment elevation and pathologic Q waves. The representative microscopic appearance of her left circumflex artery is shown in the figure. Which of the following underlying conditions is she most likely to have?

A Acute myelogenous leukemia
B Chronic alcoholism
C Diabetes mellitus
D Marfan syndrome
E Polyarteritis nodosa

20 A 56-year-old man experiences episodes of severe substernal chest pain every time he performs a task that requires moderate exercise. The episodes have become more frequent and severe over the past year, but they can be relieved by sublingual nitroglycerin. On physical examination, he is afebrile, his pulse is 78/min and regular, and there are no murmurs or gallops. Laboratory studies show creatinine, 1.1 mg/dL; glucose, 130 mg/dL; and total serum cholesterol, 223 mg/dL. Which of the following cardiac lesions is most likely to be present in this man?

A Calcific aortic stenosis
B Coronary atherosclerosis
C Restrictive cardiomyopathy
D Rheumatic mitral stenosis
E Serous pericarditis
F Viral myocarditis

21 A retrospective study of myocardial infarction is performed to analyze patterns of cardiac injury. One pattern of injury involves the posterior left ventricular wall and septum. Which of the following pathologic abnormalities is most likely to produce this pattern?

A Ascending aortic dissection
B Left anterior descending arterial plaque rupture
C Left circumflex arterial vasculitis
D Right coronary sinus embolization
E Right posterior descending arterial thrombosis

22 A 48-year-old woman has had increasing dyspnea for the past 2 days. She experiences sudden cardiac arrest. The representative light microscopic appearance of her left ventricular free wall is shown in the figure. Which of the following is the most likely diagnosis?

A Acute rheumatic myocarditis
B Cardiomyopathy
C Myocardial infarction
D Septic embolization
E Viral myocarditis
A study of ischemic heart disease analyzes cases of individuals hospitalized with acute chest pain in which myocardial infarction was documented at autopsy. The gross and microscopic appearances of the hearts are correlated with the degree of coronary atherosclerosis and its complications, clinical symptoms, and therapies given before death. Hemorrhage and contraction bands in necrotic myocardial fibers are most likely to be seen with infarction in which of the following settings?

A. Acute coronary vasculitis  
B. Anti-arrhythmic drug usage  
C. Angioplasty with stent placement  
D. Septic embolization  
E. Thrombolytic therapy

A 50-year-old man with diabetes mellitus and hypertension has had pain in the left shoulder and arm for the past 12 hours. Over the next 6 hours, he develops shortness of breath, which persists for 2 days. On day 3, he visits the physician. On physical examination, his temperature is 37.1° C, pulse is 82/min, respirations are 18/min, and blood pressure is 160/100 mm Hg. Laboratory studies show total creatine kinase (CK) activity within reference range, but the troponin I level is elevated. He continues to experience dyspnea for the next 3 days. On day 7 after the onset of shoulder pain, he has a cardiac arrest and is resuscitated. Cardiac imaging now shows a large fluid collection around the heart. Which of the following complications has he most likely developed?

A. Aortic valvular perforation  
B. Hemopericardium  
C. Left ventricular aneurysm  
D. Papillary muscle rupture  
E. Pericarditis

A 45-year-old man experiences crushing substernal chest pain after arriving at work one morning. Over the next 4 hours, the pain persists and begins to radiate to his left arm. He becomes diaphoretic and short of breath, but waits until the end of his 8-hour shift to go to the hospital. An elevated serum value of which of the following laboratory tests would be most useful for diagnosis of this patient on admission to the hospital?

A. ALT  
B. AST  
C. CK-MB fraction  
D. C-reactive protein  
E. LDH-1  
F. Lipase

A 45-year-old man suddenly collapses and is brought to the emergency department. His vital signs are temperature, 37.1° C; pulse, 84/min; respirations, 18/min; and blood pressure, 80/40 mm Hg. Laboratory findings include hemoglobin, 13.5 g/dL; platelet count, 252,000/mm³; WBC count, 7230/mm³; serum glucose, 73 mg/dL; and creatinine, 1.2 mg/dL. The total creatine kinase (CK) level is elevated, with a CK-MB fraction of 10%. Which of the following underlying conditions is most likely to be present in this patient?

A. DiGeorge syndrome  
B. Down syndrome  
C. Familial hypercholesterolemia  
D. Hereditary hemochromatosis  
E. Marfan syndrome

A 69-year-old man with metabolic syndrome had chest pain and an elevated serum troponin I level 1 year ago. He was treated in the hospital with anti-arrhythmic agents for 1 week. An echocardiogram showed an ejection fraction of 28%. He now has markedly reduced exercise tolerance. On physical examination, his temperature is 37° C, pulse is 68/min, respirations are 17/min, and blood pressure is 130/80 mm Hg. Diffuse crackles are heard on auscultation of the lungs. The representative gross appearance of his heart is shown in the figure. Which of the following complications of this disease is the patient most likely to develop?

A. Atrial myxoma  
B. Cardiac tamponade  
C. Constrictive pericarditis  
D. Hypertrophic cardiomyopathy  
E. Infective endocarditis  
F. Systemic thromboembolism

A 72-year-old man with poorly controlled diabetes mellitus has worsening exercise tolerance for 5 years. For the past year he has had chest pain with minimal exertion. On physical examination he has bilateral pulmonary rales and pitting edema of his legs. He has an irregular heart rate. A chest radiograph shows prominent right and left heart borders. Echocardiography shows decreased left ventricular ejection fraction (25%) with diminished wall motion. Laboratory studies show an elevated serum B-type natriuretic peptide. Which of the following pathologic findings is most likely present in this man?

A. Critical coronary stenosis  
B. Left atrial mural thrombus  
C. Hypertrophic cardiomyopathy  
D. Mitral and tricuspid valve thickening  
E. Pericardial fibrinohemorrhagic exudate
29. A 68-year-old woman has had increasing dyspnea and orthopnea for the past year. She does not report any chest pain. On physical examination, her temperature is 37°C, pulse is 77/min, respirations are 20/min, and blood pressure is 140/90 mm Hg. On auscultation of the chest, diffuse crackles are heard in all lung fields. No murmurs or gallops are heard, and the heart rate is regular. A chest radiograph shows prominent right and left heart borders. Coronary angiography shows 90% occlusion of the left anterior descending artery. Echocardiography shows no valvular abnormalities, but there is decreased left ventricular wall motion and an ejection fraction of 32%. Laboratory studies show serum glucose of 81 mg/dL, creatinine of 1.6 mg/dL, total cholesterol of 280 mg/dL, triglyceride of 169 mg/dL, and troponin I of 1 ng/mL. Which of the following pharmacologic agents is most likely to be beneficial in the treatment of this patient?

A. Amiodarone  
B. Alteplase  
C. Glyburide  
D. Nitroglycerin  
E. Propranolol  
F. Simvastatin

30. A 50-year-old man has sudden onset of severe substernal chest pain that radiates to the neck. On physical examination, he is afebrile, but has tachycardia, hyperventilation, and hypotension. No cardiac murmurs are heard on auscultation. Emergent coronary angiography shows a thrombotic occlusion of the left circumflex artery and areas of 50% to 70% narrowing in the proximal circumflex and anterior descending arteries. Which of the following complications of this disease is most likely to occur within 1 hour of these events?

A. Myocardial rupture  
B. Pericarditis  
C. Valvular insufficiency  
D. Ventricular fibrillation  
E. Thromboembolism

31. A study of persons receiving emergent medical services is conducted. It is observed that 5% of persons with sudden cardiac arrest who receive cardiopulmonary resuscitation survive. Which of the following is the most likely mechanism for cardiac arrest in these survivors?

A. Arrhythmia  
B. Infarction  
C. Inflammation  
D. Valve failure  
E. Ventricular rupture

32. A 59-year-old man has experienced chronic fatigue for the past 18 months. On physical examination, he is afebrile. A chest radiograph shows bilateral pulmonary edema and a prominent left heart border. The representative gross appearance of his heart is shown in the figure. Laboratory studies show serum glucose, 74 mg/dL; total cholesterol, 189 mg/dL; total protein, 7.1 g/dL; albumin, 5.2 g/dL; creatinine, 6.1 mg/dL; and urea nitrogen, 58 mg/dL. What is the most likely diagnosis?

A. Chronic alcoholism  
B. Diabetes mellitus  
C. Hemochromatosis  
D. Pneumoconiosis  
E. Systemic hypertension

33. A 56-year-old man has worsening cough and orthopnea for the past 2 years. On physical examination, he has dullness to percussion at both lung bases and diffuse crackles in the upper lung fields. He is afebrile. Echocardiography shows marked left ventricular hypertrophy and severe aortic stenosis. The representative gross appearance of the opened heart is shown in the figure. A coronary angiogram shows no significant coronary arterial narrowing. Which of the following underlying conditions best accounts for his findings?

A. Congenital anomaly  
B. Diabetes mellitus  
C. Infective endocarditis  
D. Marfan syndrome  
E. Systemic hypertension
34. A 73-year-old woman had an episode a week ago in which she became disoriented, had difficulty speaking, and had persisting weakness on the right side of her body. On physical examination, she is now afebrile with pulse of 68/min, respirations of 15/min, and blood pressure of 130/85 mm Hg. On auscultation, the lungs are clear, the heart rate is irregular, and there is a midsystolic click. A chest CT scan shows a focus of bright attenuation within the heart. An echocardiogram shows that one valvular leaflet appears to balloon upward. The ejection fraction is estimated to be 55%. Laboratory findings show serum creatine kinase (CK), 100 U/L; glucose, 77 mg/dL; creatinine, 0.8 mg/dL; calcium, 8.1 mg/dL; and phosphorus, 3.5 mg/dL. Which of the following is the most likely diagnosis?

A. Carcinoid heart disease  
B. Hyperparathyroidism  
C. Infective endocarditis  
D. Mitral annular calcification  
E. Rheumatic heart disease  
F. Senile calcific stenosis

35. A 77-year-old woman has had episodes of syncope with exertion for the past month. On physical examination, she is afebrile. Her pulse is 66/min, respirations are 14/min, and blood pressure is 125/85 mm Hg. On auscultation, a systolic ejection murmur is heard. There are a few crackles over the lung bases posteriorly. From the representative gross appearance of the opened aorta shown in the figure, which of the following most likely contributed to the development of this lesion?

A. Aging  
B. Atherosclerosis  
C. Chromosomal aneuploidy  
D. Hypercalcemia of malignancy  
E. Systemic lupus erythematosus  
F. Tertiary syphils

36. A 65-year-old healthy woman has a check of her health status and the only finding is a midsystolic click on auscultation of the heart. Within 5 years she has increasing dyspnea. Echocardiography now shows mitral regurgitation from prolapse of a leaflet. Which of the following pathologic changes is most likely present in this valve?

A. Destructive vegetations  
B. Dystrophic calcification  
C. Fibrinoid necrosis  
D. Myxomatous degeneration  
E. Rheumatic fibrosis

37. A 35-year-old woman has had palpitations, fatigue, and worsening chest pain during the past year. On physical examination, she is afebrile. Her pulse is 75/min, respirations are 15/min, and blood pressure is 110/70 mm Hg. Auscultation of the chest indicates a midsystolic click with late systolic murmur. A review of systems indicates that the patient has one or two anxiety attacks per month. An echocardiogram is most likely to show which of the following?

A. Aortic valvular vegetations  
B. Mitral valve prolapse  
C. Patent ductus arteriosus  
D. Pulmonic stenosis  
E. Tricuspid valve regurgitation

38. An 11-year-old boy had a sore throat, no cough, tonsillar exudates, and 38.3°C fever 3 weeks ago, and a throat culture was positive for group A β-hemolytic Streptococcus. On the follow-up examination, the child is afebrile. His pulse is 85/min, respirations are 18/min, and blood pressure is 90/50 mm Hg. On auscultation, a diastolic mitral murmur is audible, and there are diffuse rales over both lungs. Over the next 2 days he has several episodes of atrial fibrillation accompanied by signs of acute left ventricular failure. Which of the following pathologic changes occurring in this child’s heart is most likely to be the cause of the left ventricular failure?

A. Amyloidosis  
B. Fibrinous pericarditis  
C. Mitral valve fibrosis  
D. Myocarditis  
E. Tamponade  
F. Verrucous endocarditis
A 14-year-old girl has fever and chest pain 2 weeks after having a mild upper respiratory tract infection. On physical examination, her temperature is 37°C, pulse is 90/min, respiration are 20/min, and blood pressure is 85/45 mm Hg. A friction rub is audible on auscultation of the chest. A chest radiograph shows pulmonary edema. An echocardiogram shows small vegetations at the closure line of the mitral and aortic valves. An endomyocardial biopsy shows focal interstitial aggregates of mononuclear cells enclosing areas of fibrinoid necrosis. Her condition improves over the next month. The representative gross appearance of the affected heart is shown in the figure. Which of the following cardiac abnormalities is most likely to occur in this patient?

A. Constrictive pericarditis  
B. Dilated cardiomyopathy  
C. Left ventricular aneurysm  
D. Myxoma  
E. Valvular stenosis

A 10-year-old girl develops subcutaneous nodules over the skin of her arms and torso 3 weeks after a bout of acute pharyngitis. She manifests choreiform movements and begins to complain of pain in her knees and hips, particularly with movement. A friction rub is heard on auscultation of her chest. An abnormality detected by which of the following serum laboratory findings is most characteristic of the disease affecting this girl?

A. Antistreptolysin O antibody titer  
B. Antinuclear antibody titer  
C. Creatinine level  
D. Rapid plasma reagin test  
E. Troponin I level

A 25-year-old man was found dead at home by the apartment manager, who had been called by the decedent’s employer because of failure to report to work for the past 3 days. An external examination by the medical examiner showed splinter hemorrhages under the fingernails and no signs of trauma. The gross appearance of the heart at autopsy is shown in the figure. Which of the following laboratory findings is most likely to provide evidence for the cause of his disease?

A. Elevated anti-streptolysin O titer  
B. Positive ANCA serology  
C. Increased creatine kinase–MB (CK-MB) fraction  
D. High double-stranded DNA autoantibody titer  
E. Positive blood culture for *Staphylococcus aureus*

A 22-year-old previously healthy man undergoes a tooth extraction, and 4 days later he develops a fever. On physical examination his temperature is 37.6°C. A high-pitched systolic murmur is auscultated. In which of the following locations is the congenital anomaly in this man most likely found?

A. Ascending aorta  
B. Atrial appendage  
C. Chordae tendineae  
D. Cusps of valves  
E. Muscular septum

A 26-year-old woman has had a fever for 5 days. On physical examination, her temperature is 38.2°C, pulse is 100/min, respirations are 19/min, and blood pressure is 90/60 mm Hg. A cardiac murmur is heard on auscultation. Her sensorium is clouded, but there are no focal neurologic deficits. Laboratory findings include hemoglobin, 13.1 g/dL; platelet count, 233,300/mm³; and WBC count, 19,200/mm³. Blood cultures are positive for gram-positive bacteria. Urinalysis shows hematuria. An echocardiogram shows a 1.5-cm vegetation on the mitral valve. Which of the following conditions is this patient most likely to develop?

A. Cerebral arterial mycotic aneurysm  
B. Dilated cardiomyopathy  
C. Myxomatous mitral valve degeneration  
D. Pericardial effusion with tamponade  
E. Pulmonary abscess
44. A 19-year-old woman with rheumatic heart disease has increased fatigue. She receives a bileaflet tilting disk mechanical mitral valve prosthesis. After surgery, he is stable, and an echocardiogram shows no abnormal valvular or ventricular function. Which of the following pharmacologic agents should he receive regularly after this surgical procedure?

A. Aspirin  
B. Ciprofloxacin  
C. Cyclosporine  
D. Digoxin  
E. Propranolol  
F. Warfarin

45. A 71-year-old woman has had a 10-kg weight loss accompanied by severe nausea and vomiting of blood for the past 8 months. On physical examination, she is afebrile. Laboratory studies show hemoglobin, 8.4 g/dL; platelet count, 227,100/mm³; and WBC count, 6180/mm³. Biopsy specimens obtained by upper gastrointestinal endoscopy show adenocarcinoma of the stomach. CT scan of the abdomen shows multiple hepatic masses. CT scan of the head shows a cystic area in the right frontal lobe. Her condition is stable until 2 weeks later, when she develops severe dyspnea. A chest CT scan shows areas of decreased pulmonary arterial attenuation. Which of the following cardiac lesions is most likely to be present in this patient?

A. Calcific aortic valvular stenosis  
B. Constrictive pericarditis  
C. Epicardial metastatic carcinoma  
D. Left ventricular mural thrombosis  
E. Nonbacterial thrombotic endocarditis

46. A 41-year-old woman has had increasing dyspnea for the past week. On physical examination, temperature is 37.3°C, pulse is 85/min, respirations are 20/min, and blood pressure is 150/95 mm Hg. There is dullness to percussion over the lung bases. A chest radiograph shows large bilateral pleural effusions and a normal heart size. Laboratory findings include serum creatinine, 3.1 mg/dL; urea nitrogen, 29 mg/dL; troponin I, 0.1 ng/mL; WBC count, 3760/mm³; hemoglobin, 11.7 g/dL; and positive ANA and anti–double-stranded DNA antibody test results. Which of the following cardiac lesions is most likely to be present in this patient?

A. Calcific aortic stenosis  
B. Hemorrhagic pericarditis  
C. Nonbacterial thrombotic endocarditis  
D. Libman-Sacks endocarditis  
E. Mural thrombosis  
F. Rheumatic verrucous endocarditis

47. A 44-year-old woman with rheumatic heart disease with aortic stenosis undergoes valve replacement with a bioprosthesis. She remains stable for the next 8 years and then develops diminished exercise tolerance. Which of the following complications involving the bioprosthesis has most likely occurred?

A. Embolization  
B. Hemolysis  
C. Myocardial infarction  
D. Paravalvular leak  
E. Stenosis

48. A 50-year-old man with a history of infective endocarditis has increasing fatigue. He receives a bileaflet tilting disk mechanical mitral valve prosthesis. After surgery, he is stable, and an echocardiogram shows no abnormal valvular or ventricular function. Which of the following infectious agents is most likely to be cultured from this patient’s blood?

A. Coxsackievirus B  
B. Mycobacterium tuberculosis  
C. Pseudomonas aeruginosa  
D. Viridans streptococci  
E. Trypanosoma cruzi

49. A 44-year-old, previously healthy man has experienced worsening exercise tolerance accompanied by marked shortness of breath for the past 6 months. On physical examination, his vital signs are normal. He has diffuse rales in all lung fields and pitting edema to the knees. Laboratory studies show serum sodium, 130 mmol/L; potassium, 4 mmol/L; chloride, 102 mmol/L; CO₂ 25 mmol/L; creatinine, 2 mg/dL; and glucose, 120 mg/dL. A 100-mL urine sample is collected. There is 1.3 mmol sodium and 40 mg creatinine in the urine sample. A chest radiograph shows cardiomegaly and pulmonary edema with pleural effusions. An echocardiogram shows four-chamber cardiac enlargement and mitral and tricuspid valvular regurgitation, with an ejection fraction of 30%. A coronary angiogram shows less than 10% narrowing of the major coronary arteries. Which of the following is the most likely diagnosis?

A. Amyloidosis  
B. Hypercholesterolemia  
C. Familial cardiomyopathy  
D. Rheumatic heart disease  
E. Trypanosoma cruzi infection

50. A 56-year-old man has experienced increased fatigue and decreased exercise tolerance for the past 2 years. On physical examination, his temperature is 37°C, pulse is 75/min, respirations are 17/min, and blood pressure is 115/75 mm Hg. On auscultation, diffuse crackles are audible. The abdomen is distended with a fluid wave, and there is bilateral pitting edema to the knees. A chest radiograph shows pulmonary edema, pleural effusions, and marked cardiomegaly. An echocardiogram shows mild tricuspid and mitral regurgitation and reduced right and left ventricular wall motion, with an ejection fraction of 30%. He experiences cerebral, renal, and splenic infarctions over the next year. Chronic use of which of the following substances has most likely produced these findings?

A. Acetaminophen  
B. Cocaine  
C. Ethanol  
D. Lisinopril  
E. Nicotine  
F. Propranolol
51 A 25-year-old man suffers a sudden cardiac arrest. He is resuscitated. On examination his vital signs are normal. Echocardiography shows that the left ventricle is normal but there is marked thinning with dilation of the right ventricle. MR imaging of his chest shows extensive fibrofatty replacement of the myocardium, but no inflammation. Which of the following is the most likely cause for his findings?

A Cardiomyopathy  
B Chagas disease  
C Hypertension  
D Long QT syndrome  
E Radiation therapy

52 A 10-year-old girl who is normally developed has chronic progressive exercise intolerance. On physical examination, temperature is 37.1 °C, pulse is 70/min, respirations are 14/min, and blood pressure is 100/60 mm Hg. A chest radiograph shows cardiomegaly and mild pulmonary edema. An echocardiogram shows severe left ventricular hypertrophy and a prominent interventricular septum. The right ventricle is slightly thickened. During systole, the anterior leaflet of the mitral valve moves into the outflow tract of the left ventricle. The ejection fraction is abnormally high, and the ventricular volume and cardiac output are both low. Which of the following is the most likely cause of the cardiac abnormalities in this patient?

A Autoimmunity against myocardial fibers  
B β-Myosin heavy chain gene mutation  
C Deposition of amyloid fibrils  
D Excessive iron accumulation  
E Latent enterovirus infection

53 A 17-year-old girl jumps up for a block in the third match of a volleyball tournament and suddenly collapses. She requires cardiopulmonary resuscitation. A similar episode occurs a month later. She had been healthy all her life and complained only of limited episodes of chest pain in games during the current school year. Which of the following pathologic findings of the heart is most likely to be present in this girl?

A Extensive myocardial hemosiderin deposition  
B Haphazardly arranged hypertrophied septal myocytes  
C Large, friable vegetations with destruction of aortic valve cusps  
D Mitral valvular stenosis with left atrial enlargement  
E Tachyzoites within foci of myocardial necrosis and inflammation

54 An 86-year-old man has had increasing dyspnea and reduced exercise tolerance for the past 7 years. On physical examination, he is afebrile and has a blood pressure of 135/85 mm Hg. An irregularly irregular heart rate averaging 76/min is audible on auscultation of the chest. Crackles are heard at the bases of the lungs. A chest radiograph shows mild cardiomegaly and mild pulmonary edema. Echocardiography shows slight right and left ventricular wall thickening with reduced left and right ventricular wall motion, reduced left ventricular filling, and an ejection fraction estimated to be 25%. An endomyocardial biopsy specimen shows amorphous pink-staining deposits between myocardial fibers, but no inflammation and no necrosis. Echocardiography would most likely show which of the following functional cardiac disturbances?

A Dynamic obstruction to ventricular outflow  
B Impaired ventricular diastolic filling  
C Increased end-systolic volume  
D Mitral and tricuspid valvular insufficiency  
E Reduced ejection fraction

55 A 33-year-old woman from Victoria, British Columbia, goes to the physician because of increasingly severe dyspnea, orthopnea, and swelling of the legs for the past 2 weeks. She has no previous history of serious illness or surgery. On physical examination, her temperature is 37.8 °C, pulse is 83/min, respirations are 20/min, and blood pressure is 100/60 mm Hg. An ECG shows episodes of ventricular tachycardia. An echocardiogram shows right and left ventricular dilation, but no valvular deformities. An endomyocardial biopsy shows focal myocyte necrosis and lymphocytic infiltrate. Which of the following organisms most likely caused the infection?

A Coxsackievirus A  
B Mycobacterium kansasii  
C Viridans streptococci  
D Staphylococcus aureus  
E Toxoplasma gondii  
F Trypanosoma cruzi

56 A 68-year-old man has become increasingly lethargic and weak for the past 7 months. On physical examination, his temperature is 36.9 °C, pulse is 70/min, respirations are 15/min, and blood pressure is 160/105 mm Hg. On auscultation of his chest, a friction rub is audible. There are no other remarkable findings. The representative gross appearance of the heart is shown in the figure. Which of the following laboratory findings is most likely to be reported for this patient?

A Elevated serum anti-streptolysin O titer  
B Elevated plasma renin level  
C Increased blood urea nitrogen level  
D Increased serum CK-MB level  
E Positive ANA with “rim” pattern  
F Positive viral serology
57. A 52-year-old woman has had a chronic cough for the past 2 years, accompanied by a small amount of occasionally blood-streaked, whitish sputum. On physical examination, her temperature is 37.9°C, pulse is 72/min, respirations are 22/min, and blood pressure is 125/80 mm Hg. Crackles are heard on auscultation over the upper lung fields. Heart sounds are faint, and there is a 15 mm Hg inspiratory decline in systolic arterial pressure. The chest radiograph shows prominent heart borders with a “water bottle” configuration. Pericardiocentesis yields 200 mL of bloody fluid. Infection with which of the following organisms is most likely to produce these findings?

A. Candida albicans
B. Coxsackievirus B
C. Group A streptococcus
D. Mycobacterium tuberculosis
E. Staphylococcus aureus

58. A 31-year-old man experienced chest pain, became increasingly dyspneic and nauseated, and lost consciousness multiple times. Seven days after the appearance of these symptoms, he was found dead in his sleep. External examination of the body by the medical examiner shows no evidence of trauma. The body is 166 cm (5 ft 5 in) in height and weighs 75 kg (BMI 27). The gross appearance of the chest cavity at autopsy with the pericardial sac opened is shown in the figure. What is the most likely underlying cause of his death?

A. Coronary atherosclerosis
B. Dilated cardiomyopathy
C. Disseminated tuberculosis
D. Systemic sclerosis
E. Malignant melanoma
F. Marfan syndrome
G. Takayasu arteritis

59. A 73-year-old woman has had episodes of chest pain during the past week. She is afebrile. Her pulse is 80/min, respirations are 16/min, and blood pressure is 110/70 mm Hg. On auscultation of the chest, heart sounds seem distant, but the lung fields are clear. Neck veins are distended to the angle of the jaw, even while sitting. There is a darkly pigmented, irregular, 1.2-cm skin lesion on the right shoulder. A chest radiograph shows prominent borders on the left and right sides of the heart. Pericardiocentesis yields bloody fluid. Laboratory findings include a serum troponin I level of 0.3 ng/mL. Which of the following lesions is the most likely cause of these findings?

A. Calcific aortic stenosis
B. Coronary atherosclerosis
C. Epicardial metastases
D. Mitral valvulitis
E. Tuberculous pericarditis

60. A 48-year-old, previously healthy woman reports having suddenly lost consciousness four times in the past 6 months. In three instances, she was unconscious for only a few minutes. After the fourth episode 1 month ago, she was unconscious for 6 hours and had weakness in her right arm and difficulty speaking. On physical examination, she is afebrile, and her blood pressure is normal. No murmurs are auscultated. She has good carotid pulses with no bruits. Which of the following cardiac lesions is most likely to be present in this woman?

A. Bicuspid aortic valve
B. Coronary artery thrombosis
C. Left atrial myxoma
D. Mitral valve stenosis
E. Pericardial effusion

61. A 55-year-old man undergoes orthotopic cardiac transplantation. Two months later, an endomyocardial biopsy specimen shows focal myocardial cell death with scattered perivascular lymphocytes and plasma cells. He is treated with sirolimus. Which of the following pathologic processes best accounts for these biopsy findings?

A. Autoimmunity
B. Autophagy
C. Ischemia
D. Infection
E. Rejection

62. A 45-year-old man receives a cardiac allograft for dilated cardiomyopathy. He has no problems with rejection, but 5 years later he has worsening exercise tolerance with increasing dyspnea and peripheral edema. Echocardiography shows a reduced ejection fraction of 35%. Which of the following pathologic abnormalities has he most likely developed in the allograft?

A. Amyloidosis
B. Constrictive pericarditis
C. Coronary arteriopathy
D. Non-Hodgkin lymphoma
E. Toxoplasmosis
1. F The pacemaker for the heart is the sinoatrial (SA) node, with a natural rhythm near 70/min and a normal range of 60/min to 100/min. Other parts of the cardiac conduction system pass along this rate. Rates less than 60/min are defined as bradycardia, and rates greater than 100/min are defined as tachycardia. Bradyarrhythmias less than 50/min suggest an SA node disorder. SA node dysfunction may worsen with cardioactive drugs, such as cardiac glycosides, β-adrenergic blockers, calcium channel blockers, and amiodarone. An increase in sinus rate results from an increase in sympathetic tone acting via β-adrenergic receptors or a decrease in parasympathetic tone acting via muscarinic receptors, or both. Abnormalities involving the other listed options are unlikely to produce such a pronounced and consistent bradycardia.

PBD9 524–525  BP9 365–367  BP8 531

2. F Native cardiac stem cells can proliferate and replace cardiac myocytes throughout life, but these stem cells are most active in neonates. They have the potential to respond to injury. Some ventricular septal defects do close during life. Research is ongoing regarding methods for inducing stem cell proliferation. Of the remaining cells listed, adipocytes enlarge with aging. Endothelial cells may proliferate to produce more coronary collateral channels in response to exercise training, and they can resurface vascular grafts. Fibroblasts respond to injury by producing collagenous scar tissue that reduces contractility.

PBD9 525  BP8 537

3. E These findings are consistent with right-sided congestive heart failure leading to peripheral edema, body cavity effusions (pleural effusions in this case), passive congestion of the liver, and jugular venous distension. Pure right-sided failure is less common than left-sided failure, and the former most often follows pulmonary disease (cor pulmonale). Chronic obstructive pulmonary disease (COPD) is more common than restrictive lung disease, and smoking leads to COPD. Atrial myxomas are uncommon, more often on the left side of the heart, and may produce intermittent valvular obstruction. Essential hypertension is systemic and places a pressure load on the left side of the heart. Hyperlipidemia is a risk factor for ischemic heart disease that is more likely to involve the left side of the heart. Rheumatic heart disease may produce heart failure, but it is more often left-sided.

PBD9 526–527, 530  BP9 365  BP8 545, 558  BP8 388, 390

4. C Left-sided congestive heart failure leads to pulmonary congestion and edema. Systemic hypertension is common and leads to pressure load with predominantly concentric left ventricular hypertrophy with systolic dysfunction. The other listed options are more likely to produce diastolic dysfunction and decrease of both right and left ventricular function.

PBD9 527–529  BP9 366–367  BP8 545, 558  BP8 388, 390

5. E Paroxysmal nocturnal dyspnea is a feature of left-sided congestive heart failure, and rheumatic heart disease most often involves the mitral, aortic, or both valves, and left-sided valvular disease leads to pulmonary edema. While upright, pulmonary edema fluid is more concentrated at lung bases, which helps improve breathing. Rheumatic heart disease was more common before antibiotic therapy for group A β-hemolytic streptococcal infections was available, and multiple bouts beginning in childhood led to valvular damage over decades. An atrial myxoma usually occurs on the left side of the heart, but the obstruction is often intermittent. Fibrinous pericarditis can produce chest pain, but the amount of accompanying fluid is often small so that cardiac function is not impaired. Giant cell myocarditis is a rare cause of cardiac failure. Libman-Sacks endocarditis, seen in systemic lupus erythematosus, typically does not impair valvular or ventricular function.

PBD9 529  BP9 367, 387  BP8 535–536  BP8 381

6. B The findings point to pure right-sided congestive heart failure. Rarely, this can be caused by right-sided cardiac valvular lesions, such as tricuspid or pulmonic stenosis. Pulmonary hypertension resulting from obstructive lung diseases, such as emphysema or chronic bronchitis, most often caused from smoking cigarettes, is much more common. Primary pulmonary hypertension also can cause right-sided heart failure, but it is a much less common cause than obstructive lung diseases. Because acute myocardial infarction usually affects the left ventricle, left-sided heart failure would be more common in these patients. Chronic left-sided heart failure eventually can lead to right-sided heart failure.

PBD9 530  BP9 368, 388  BP8 536–537  BP8 381–382

7. D There are a number of mutations in genes linked to congenital heart disease that encode proteins in transcription or signaling pathways. The NOTCH pathway plays a role in modulation of vascular development, including cardiac outflow tracts. NOTCH2 mutations are associated with tetralogy of Fallot, as in this infant. β-MHC gene mutations are associated with some cases of hypertrophic cardiomyopathy. Fibrillin-1 (FBN1) gene mutations underlie Marfan syndrome. KCNQ1 mutations may be seen with long QT syndrome. One form of cardiac amyloidosis is linked to transthyretin (TTR) gene mutations.

PBD9 532–533  BP9 369  BP8 538–539

8. E This shunt results from abnormal confluence of pulmonary veins leading to the right atrium (or systemic veins), and not the left. Obstruction is often present, as in this case, with pulmonary congestion. Deoxygenated systemic and oxygenated pulmonary venous blood mix in the right atrium. There must be an atrial septal defect for blood to reach the left atrium, but an ASD by itself does not explain this case. A coarctation is not associated with cyanosis; a preductal coarctation is life-threatening. Patent ductus arteriosus is a
left-to-right shunt without cyanosis. Tetralogy of Fallot can lead to cyanosis, but mixing of blood occurs at an overriding aorta.

PBD9 531–533  BP9 369  PBD8 543  BP8 384

9 D This is the infamous “paradoxical embolus” that has appeared far more often in question sets than in real life. She started with thrombophlebitis that led to pulmonary embolism, but there must be an explanation for the “stroke” that then occurred. Pulmonary emboli can obstruct the pulmonary arterial circulation, raising right atrial pressure, and opening a patent foramen ovale that normally remains closed because of higher left atrial pressure. The remaining choices do not explain pulmonary thromboembolism. A left atrial myxoma can embolize to the brain; lesions of endocarditis are most often on the left side of the heart and could produce cerebral emboli; a ventricular aneurysm is virtually always on the left side of the heart because it results from a healed infarction, and can be filled with mural thrombus that can embolize.

PBD9 533–534  BP9 370–371  PBD8 541  BP8 385

10 D The figure shows a large ventricular septal defect. By the age of 5 years, such an uncorrected defect causes marked shunting of blood from left to right, causing pulmonary hypertension (Eisenmenger complex). The left and right ventricular chambers undergo hypertrophy and some dilation, but the functioning of the cardiac valves is not greatly affected. In most cases, congenital heart disease is not an antecedent to ischemic heart disease. Nonbacterial thrombotic endocarditis most often occurs secondary to a hypercoagulable state in adults. Restrictive cardiomyopathy may occur from conditions such as amyloidosis or hemochromatosis.

PBD9 535  BP9 370–371  PBD8 541  BP8 384–385

11 B A persistent ostium secundum is the most common form of atrial septal defect. Because atrial pressures are low, the amount of shunting from the left atrium to the right atrium is small, and this lesion may remain asymptomatic for many years. Eventually, pulmonary hypertension can occur, with reversal of the shunt. Cyanosis is a feature of a right-to-left shunt. A dilated heart with enlarged atria predisposes to mural thrombosis and embolism. Pericardial effusions may occur much later, if congestive heart failure develops.

PBD9 534  BP9 370–371  PBD8 541  BP8 383–384

12 D Although often not causing a large shunt defect, a patent ductus arteriosus can produce a significant murmur and predispose to infection. This left-to-right shunt may eventually result in pulmonary hypertension. An atretic valve has no flow across it and does not produce a murmur, but there would be a murmur across a shunt around the atretic valve. Aortic atresia is not compatible with continued survival, as seen in hypoplastic left heart syndrome. Aortic coarctations by themselves produce no shunting and no pulmonary hypertension. An atrial septal defect is unlikely to produce a loud murmur because of the minimal pressure differential between the atra. Because pulmonic stenosis is a component of tetralogy of Fallot, no pulmonary hypertension results, and the right-to-left shunting can lead to cyanosis with decreased arterial oxygen saturation. Total anomalous pulmonary venous return is not accompanied by a murmur because of the low venous pressure.

PBD9 535  BP9 371–372  PBD8 541–542  BP8 385

13 E Cyanosis at this early age suggests a right-to-left shunt. Truncus arteriosus, transposition of the great arteries, and tetralogy of Fallot are the most common causes of cyanotic congenital heart disease. The cerebral lesion suggests an abscess as a consequence of septic embolization from infective endocarditis, which can complicate congenital heart disease. Atrial septal defect, patent ductus arteriosus, and ventricular septal defect initially lead to left-to-right shunts, though the shunt may reverse with development of pulmonary hypertension. Coarctation is not accompanied by a shunt and cyanosis. In most cases, a bicuspid valve is asymptomatic until adulthood, and there is no shunt.

PBD9 535–537  BP9 372–373  PBD8 542–544  BP8 383

14 A In tetralogy of Fallot, the severity of the obstruction to the right ventricular outflow determines the direction of flow. If the pulmonic stenosis is mild, the abnormality resembles a ventricular septal defect, and the shunt may be from left to right with no cyanosis. With significant pulmonic outflow obstruction, the right ventricular pressure may reach or exceed systemic vascular resistance, and the blood would be shunted from right to left, producing cyanotic heart disease. Even if pulmonic stenosis is mild at birth, the pulmonary orifice does not expand proportionately as the heart grows, and cyanotic heart disease supervenes.

PBD9 535–536  BP9 372–373  PBD8 543  BP8 385–386

15 D The figure shows that the aorta emerges from the right ventricle, and the pulmonic trunk exits the left ventricle, consistent with complete transposition of the great vessels. Unless there is another anomalous connection between the pulmonary and systemic circulations, this condition is incompatible with extrauterine life. The most common additional anomalous connections would be ventricular septal defect, patent ductus arteriosus, and patent foramen ovale (or atrial septal defect). In pulmonic and aortic stenosis, the great arteries are normally positioned, but small. In tetralogy of Fallot, the aorta overrides a ventricular septal defect, but is not transposed. In truncus arteriosus, the spiral septum that embryologically separates the great arteries does not develop properly.

PBD9 536–537  BP9 372–373  PBD8 543  BP8 386–387

16 B In children and adults, the coarctation is typically postductal, and collateral branches from the proximal aorta supply the lower extremities, leading to the large pulse differential between upper and lower extremities. Collaterals often involve intercostal arteries whose enlargement produces
"rib notching" on chest radiographs. Diminished renal blood flow below the coarctation increases renin production and promotes hypertension. Aortic valve stenosis causes left-sided heart failure and no pressure differential in the extremities. A patent ductus arteriosus produces a small left-to-right shunt. Transposition results in a right-to-left shunt with cyanosis. Tricuspid valve atresia affects the right side of the heart.

**PBD9 537 BP9 373–374 PBD8 544 BP8 387–388**

**17 C** These findings are compatible with hypoplastic left heart syndrome, which may have varying degrees of severity, ranging from severe (as in this case, with virtually no function on the left side of the heart) to milder degrees of hypoplasia. Most of the oxygenated blood returning to the left atrium is shunted across the foramen ovale back to the lungs, increasing pulmonary flow and decreasing oxygenation. Less oxygenated blood exiting the right ventricle into the pulmonic trunk can shunt through a patent ductus arteriosus to the aorta to supply the systemic circulation. Anomalous venous return does not generally connect to the aorta, and there still must be a connection from the lungs to the aorta. The right fourth aortic arch rarely persists. Truncus arteriosus is an anomalous, incomplete separation of the pulmonic and aortic trunks. If there is virtually no left ventricular chamber, a ventricular septal defect would not provide any significant flow.

**PBD9 531, 538 BP9 369 PBD8 544**

**18 A** Marked coronary artery occlusion with this degree of stenosis prevents adequate perfusion of the heart when myocardial demand is increased during exertion. He has angina on exertion and recently developed unstable angina, which is manifested by increased frequency and severity of the attacks and angina at rest. The ST segment elevation suggests a developing acute coronary syndrome with myocardial ischemia, but the lack of cardiac enzyme elevation suggests infarction has not yet occurred. In most patients, unstable angina is induced by disruption of an atherosclerotic plaque followed by a mural thrombus and possibly distal embolization, vasospasm, or both. An acute myocardial infarction (MI) can lead to focal fibrinous pericarditis, but it is unlikely to lead to extensive scarring that surrounds the heart. Fibrosis is a late finding from healing of infarction. Hypertrophy of the heart is unlikely to progress significantly in this case because there is neither hypertension nor a valvular lesion. Mural thrombosis may develop on the endocardial surface overlying an infarction, and may fill a ventricular aneurysm following an MI. An acute MI may be complicated by papillary muscle rupture with mitral valve insufficiency.

**PBD9 538–540 BP9 374–376 PBD8 546, 558 BP8 388–390**

**19 C** The figure shows a coronary artery with marked luminal narrowing caused by atheromatous plaque, complicated by a recent thrombus filling the narrowed lumen. Atherosclerosis is accelerated with diabetes mellitus. When a premenopausal woman develops severe atherosclerosis, as in this case, underlying diabetes mellitus or a lipid disorder must be strongly suspected. Patients with leukemias may have reversal of any atheromas, but can develop hypercoagulable states. When this occurs, there is widespread thrombosis in normal blood vessels. Individuals with chronic alcoholism often have less atherosclerosis than individuals of the same age who do not consume large amounts of alcohol. The cystic medial necrosis that occurs in Marfan syndrome most often involves the ascending aorta and predisposes to dissection that could involve coronary arteries, although with external compression. Polyarteritis nodosa can involve coronary arteries and give rise to coronary thrombosis when the arterial wall is necrotic and inflamed.

**PBD9 540–543 BP9 374–376 PBD8 547–549 BP8 389–390**

**20 B** Angina pectoris typically occurs when coronary artery narrowing exceeds 75%. His risk factors include hyperglycemia (diabetes mellitus) and hypercholesterolemia. Calcific aortic stenosis leads to left-sided congestive heart failure (CHF), and the extra workload of the left ventricle may cause angina pectoris. Calcific aortic stenosis (in the absence of a congenital bicuspid aortic valve) is rarely symptomatic at 50 years of age, however. Cardiomyopathies result in heart failure, but without chest pain. Patients with rheumatic heart disease are affected by slowly worsening CHF. Pericarditis can produce chest pain, although not in relation to exercise, and it is not relieved by nitroglycerin. Viral myocarditis may last for weeks, but not for 1 year, and pain may be present at rest.

**PBD9 539–540 BP9 376 PBD8 545, 558 BP8 388, 390**

**21 E** Myocardial infarction results from occlusion of large coronary arterial branches, and in most cases an occluding thrombus is present. The posterior left ventricle and septum are supplied by the posterior descending artery. The left circumflex artery supplies the lateral left ventricular wall, whereas the left anterior descending artery supplies the anterior left ventricle. An aortic dissection that extends proximally may cause tamponade, compressing the heart, great vessels, and even coronary arteries, but this is much less likely a cause for myocardial infarction than atherosclerotic coronary arterial disease. The coronary sinus is where venous blood from the myocardium drains into the right atrium.

**PBD9 540–543 BP9 379 PBD8 547–549 BP8 388–390**

**22 C** The figure shows intensely eosinophilic myocardial fibers with loss of nuclei, all are indicative of coagulative necrosis. The deeply red-stained transverse bands are called contraction bands. Neutrophils infiltrate between myocardial fibers. This pattern is most likely caused by a myocardial infarction (MI) that is approximately 24 to 48 hours old. Chest pain is present in most but not all cases of MI. Rheumatic myocarditis is characterized by minimal myocardial necrosis with foci of granulomatous inflammation (Aschoff bodies). There is no significant inflammation with restrictive cardiomyopathies such as amyloidosis or hemochromatosis. Septic emboli result in focal abscess formation. In viral myocarditis, there is minimal focal myocardial necrosis with round cell infiltrates.

**PBD9 544–546 BP9 379–381 PBD8 553–554 BP8 391–395**
**23 E**  Reperfusion of an ischemic myocardium by spontaneous or therapeutic thrombolysis changes the morphologic features of the affected area. Reflow of blood into vasculature injured during the period of ischemia leads to mitochondrial dysfunction, followed by leakage of blood into the tissues (hemorrhage). Contraction bands are composed of closely packed hypercontracted sarcomeres. They are most likely produced by exaggerated contraction of previously injured myofibrils that are exposed to a high concentration of calcium ions from the plasma. The damaged cell membrane of the injured myocardial fibers allows calcium to penetrate the cells rapidly. Free radical formation and release of leukocyte enzymes further potentiate myocardial cell death. Hemorrhage would not be a prominent feature in the other listed options. Vasculitides involving the heart are uncommon; Takayasu arteritis can involve coronary arteries, but is most often a rare pediatric condition. Drugs used to control arrhythmias during acute coronary syndromes are unlikely to have hemorrhage as an adverse event. Angioplasty per se does not increase the risk for hemorrhage, and stents help to keep the artery open longer. Septic embolization from infected valvular vegetations to a coronary artery is uncommon, although such emboli may produce focal necrosis and hemorrhage.

PBD9 545–547  BP9 381–382  PBD8 553–555  BP8 394–395

**24 B**  CK activity begins to increase 2 to 4 hours after an MI, peaks at about 24 to 48 hours, and returns to normal within 72 hours. Troponin I levels begin to increase at about the same time as CK and CK-MB, but remain elevated for 7 to 10 days. Total CK activity is a sensitive marker for myocardial injury in the first 24 to 48 hours. CK-MB offers more specificity, but not more sensitivity. The risk for myocardial rupture is greatest from 4 to 7 days after transmural myocardial necrosis. This patient had an MI on the day of the shoulder pain. When he saw the physician on day 3, the CK levels had returned to normal, but troponin I levels remained elevated. Three days later, the infarct ruptured, and blood filled the pericardial cavity. Cardiac valves are essentially avascular and not subject to ischemic injury. Ventricular aneurysm formation is a late complication of a healed MI. Papillary muscles are at risk for rupture, just like the free wall, but the consequence would be acute valvular insufficiency, not hemopericardium. A transmural MI may lead to pericarditis, often with some accompanying pericardial effusion, but the acute event here in the time frame described suggests rupture.

PBD9 547–548  BP9 382–383  PBD8 555  BP8 395

**25 C**  Of the enzymes listed, CK-MB is the most specific for myocardial injury from the acute myocardial infarction (MI) described in this patient. The levels of this enzyme begin to increase within 2 to 4 hours of ischemic myocardial injury. ALT elevation is more specific for liver injury. AST is found in various tissues; elevated levels are not specific for myocardial injury. The elevation of lactate dehydrogenase (LDH)-1 compared with LDH-2 suggests myocardial injury, but LDH activity peaks 3 days after an MI. C-reactive protein is elevated with inflammatory processes, but is nonspecific; it has been used as a predictor of acute coronary syndromes. Lipase is a marker for pancreatitis.

PBD9 547–548  BP9 382–383  PBD8 555  BP8 390–396

**26 C**  The laboratory findings suggest an acute myocardial infarction. Individuals with familial hypercholesterolemia have accelerated and advanced atherosclerosis, even by the second or third decade. DiGeorge syndrome can be associated with various congenital heart defects, but survival with this syndrome is usually limited by infections resulting from cell-mediated immunodeficiency. Down syndrome (trisomy 21) is often accompanied by endocardial cushion defects, not ischemic heart disease. Hereditary hemochromatosis may result in an infiltrative cardiomyopathy with iron overload, more typically by the fifth decade. Marfan syndrome may result in aortic dissection or floppy mitral valve.

PBD9 547–548  BP9 382–383  PBD8 545–547  BP8 388–393

**27 F**  The figure shows an enlarged and dilated heart with a large ventricular aneurysm with a thin wall and white fibrous endocardial surface. Such an aneurysm most likely results from weakening of the ventricular wall at the site of a prior healed myocardial infarction. Because of the damage to the endocardial lining, with stasis and turbulence of blood flow in the region of the aneurysm, mural thrombi are likely to develop. When detached, thrombi in the left side of the heart embolize to the systemic circulation and can cause infarcts elsewhere. An atrial myxoma is the most common primary cardiac neoplasm, but it is rare and is not related to ischemic heart disease. Cardiac rupture with tamponade is most likely to occur 5 to 7 days after an acute myocardial infarction. Constrictive pericarditis follows a previous suppurative or tuberculous pericarditis. Hypertrophic cardiomyopathy is not related to ischemic heart disease, but 50% of cases are familial and may be related to genetic mutations in genes encoding for cardiac contractile elements. Infective endocarditis is more likely to complicate valvular heart disease or septal defects.

PBD9 549  BP9 383–384  PBD8 556–557  BP8 396–397

**28 A**  The history of diabetes mellitus and the chest pain put ischemic heart disease at the top of the differential diagnosis list for this man, who has findings with both right and left ventricular failure and enlargement, suggesting ischemic cardiomyopathy. An elevated serum B-type natriuretic peptide (which is measured instead of atrial natriuretic peptide) is consistent with heart failure. Occlusive coronary atherosclerosis may lead to multiple infarctions, or may silently cause progressive myofiber loss, but the end stage is ischemic cardiomyopathy. Atrial mural thrombus formation may occur with aortic valve dysfunction and with dysrhythmias. Myofiber disarray is characteristic for hypertrophic cardiomyopathy, which affects the interventricular septum preferentially and is usually symptomatic by young adulthood. Although rheumatic valvulitis with thickening may involve both the left and right sides of the heart, this is unusual, and it is not associated with coronary artery disease. Pericardial fluid collection may constrict heart motion, without an enlarged heart, and hemopericardium may acutely occur with ventricular rupture.

PBD9 550  BP9 384–385  PBD8 558  BP8 397

**29 F**  An ischemic cardiomyopathy can result from coronary atherosclerosis, but she does not have an acute coronary syndrome. The major identifiable risk factor in this case is
hypercholesterolemia, and the HMG-CoA reductase inhibitors (the statin drugs) are helpful to lower cholesterol, specifically LDL cholesterol. Amiodarone is used to treat intractable arrhythmias. Glyburide is used in the treatment of type 2 diabetes mellitus, but this patient is not hyperglycemic. Nitroglycerin is a vasodilator used to treat angina. Propranolol is a β-blocker that has been used to treat hypertension, and it may exacerbate bradycardia and congestive heart failure. Alteplase (tissue plasminogen activator) is used early in treatment of coronary thrombosis to help reestablish coronary blood flow.

30 D In the period immediately after coronary thrombosis, arrhythmias are the most important complication and can lead to sudden cardiac death. It is believed that, even before ischemic injury manifests in the heart, there is greatly increased electrical irritability predisposing to dysrhythmias. Myocardial rupture, valvular insufficiency from papillary muscle involvement, and pericarditis occur several days later. Another complication is a left ventricular aneurysm, a late complication of the healing of a large transmural infarction; a mural thrombus may fill an aneurysm and become a source of emboli. If portions of the coronary thrombus break off and embolize, they enter smaller arterial branches in the distribution already affected by ischemia. Valvular insufficiency from a ruptured papillary muscle would occur later in the course.

31 A The most common cause for sudden cardiac arrest is ischemic heart disease. The risk for sudden death is increased with worsening atherosclerotic coronary arterial narrowing. However, the first event with an acute coronary syndrome is typically an arrhythmia, and this is why resuscitation, including defibrillation, can be successful, and survivors may have no ECG or enzyme changes to suggest myocardial infarction has occurred. Inflammation with infarction or infection takes days to develop. A sudden valvular incompetence from papillary muscle rupture, or wall rupture, may complicate an infarction 3 to 7 days following the initial event.

32 E The markedly thickened left ventricular wall is characteristic of myocardial fiber hypertrophy caused by increased pressure load from hypertension, which often is associated with chronic renal disease. Left ventricular failure leads to pulmonary edema. Chronic alcoholism is most often associated with dilated cardiomyopathy. Diabetes mellitus accelerates atherosclerosis, leading to ischemic heart disease and myocardial infarction; the normal glucose level does not fit with diabetes mellitus. Hemochromatosis leads to dilated cardiomyopathy. Pneumonioses produce restrictive lung disease with cor pulmonale and predominantly right ventricular hypertrophy.

33 A The bicuspid valve shown has a tendency to calcify with aging, which eventually can result in stenosis, left ventricular hypertrophy, and left-sided heart failure with pulmonary edema. In individuals with congenitally bicuspid valves, symptoms often appear by 50 to 60 years of age. By contrast, calcific aortic stenosis of tricuspid valves manifests in the seventh or eighth decade. Ischemic heart disease, expected with diabetes mellitus, does not lead to valvular stenosis. In infective endocarditis, the patient would have an infection, and the valve would tend to be destroyed, leading to insufficiency. In Marfan syndrome, loss of elastic tissue in the media leads to aortic root dilation, producing aortic valvular insufficiency. Systemic hypertension accounts for left ventricular hypertrophy, but the aortic valve is not affected.

34 D Mitral annular calcification is often an incidental finding on chest radiograph, echocardiography, or at autopsy. Larger accumulations of calcium in the mitral ring can impinge on the conduction system, however, causing arrhythmias or disrupting the endocardium to provide a focus for infective endocarditis and thrombus formation (which can embolize and cause a stroke, as in this patient). Some cases are associated with mitral valve prolapse. Carcinoid heart disease leads to endocardial and valvular collagenous thickening. Hyperparathyroidism can cause metastatic calcification, which usually does not involve the heart, and deposits would not be so focal; this patient does not have hypercalcemia. Infective endocarditis is a destructive process, and healing may lead to fibrosis, but not to nodular calcium deposition. The most common infiltrative cardiomyopathies are hemochromatosis and amyloidosis. Rheumatic heart disease can lead to scarring with some calcium deposition, but the valve leaflets undergo extensive scarring, with shortening and thickening of the chordae tendineae that preclude upward prolapse. Senile calcific stenosis involves the aortic valve; in this case, there is no evidence of stenosis.

35 A Note that this valve has three cusps, but the nodular deposits of calcium interfere with cusp movement to cause calcific aortic stenosis. This is a degenerative change that may occur in a normal aortic valve with aging. Syncope may occur upon exertion because the stenotic valve prevents stroke volume from increasing in the presence of systemic vasodilation, resulting in hypotension. Atherosclerosis does not produce valvular disease from involvement of the valve itself. Congenital anomalies with chromosomal aneuploidies (e.g., trisomy 21) are unlikely to be associated with aortic stenosis or a bicuspid valve. Hypercalcemia may cause metastatic calcification, but it is unlikely in cardiac valves; it is more likely to cause arrhythmias. Systemic lupus erythematosus may give rise to small sterile vegetations on mitral or tricuspid valves, but these rarely cause valve disease. In syphilis, the aortic root dilates, and aortic insufficiency results.

36 D Myxomatous mitral valve degeneration can be primary from a connective tissue disorder such as Marfan syndrome or secondary to chronic hemodynamic forces
(later in life), the chordae tendineae become elongated and can rupture to produce acute valvular incompetence. Destructive vegetations occur with infective endocarditis, and develop over days to weeks. Dystrophic calcification in older persons can occur in the mitral annulus or aortic valve; the former is typically incidental and the latter may produce symptomatic stenosis. Fibrinoid necrosis is most typical of hyperplastic arteriolosclerosis, not cardiac valves. Rheumatic heart disease leads to valvular scarring with shortening and thickening of the chordae tendineae, not thinning and elongation.

PBD9 556–557  BP9 390  PBD8 563–564  BP8 402–403

37 B A floppy (prolapsed) mitral valve is usually asymptomatic. When symptomatic, it can cause fatigue, chest pain, and arrhythmias. Some cases are linked to clinical depression and anxiety, and others are associated with Marfan syndrome. Valvular vegetations suggest endocarditis, and a murmur is likely to be heard with infective endocarditis causing valvular insufficiency. A patent ductus arteriosus causes a shrill systolic murmur. Pulmonic stenosis is most often a congenital heart disease. Tricuspid regurgitation is accompanied by a rumbling systolic murmur.


38 D This boy developed acute left ventricular failure, an uncommon but serious complication of acute rheumatic fever. Pancarditis with pericarditis, endocarditis, and myocarditis develop during the acute phase. Myocarditis led to dilation of the ventricle so severe that the mitral valve became incompetent. Rheumatic heart disease is now uncommon, and the number of children that require prophylactic antibiotic therapy to prevent just one case is >10,000. Chronic inflammatory conditions may produce reactive systemic amyloidosis, but this is unlikely to occur given the limited and episodic nature of the streptococcal infection that causes rheumatic heart disease. Fibrous pericarditis can produce an audible friction rub, but it is not constrictive, and the amount of fluid and fibrin are not great, so no tamponade occurs. Myocardial necrosis associated with myocarditis is patchy, and the ventricle does not rupture to produce tamponade. Fibrosis and fusion of the mitral valve leaflets develop over weeks to months and indicate chronic rheumatic valvulitis. Verrucous vegetations are small and may produce a murmur, but they do not interfere greatly with valve function and do not tend to embolize.

PBD9 557–559  BP9 391–392  PBD8 566  BP8 403–406

39 E The mitral valve in the figure shows shortening and thickening of the chordae tendineae typical of chronic rheumatic valvulitis, and the small verrucous vegetations (arrowheads) are characteristic of superimposed acute rheumatic fever. Valvular scarring can follow years after initial group A streptococcal infection. Rheumatic heart disease develops after the immune response directed against the bacterial antigens (similar to cardiac antigens, and thus a form of molecular mimicry) damages the heart because streptococcal antigens cross-react with the heart. The mitral and aortic valves are most commonly affected, so right ventricular dilation from tricuspid involvement is less likely. In almost all cases, the fibrinous pericarditis seen during the acute phase with friction rub resolves without significant scarring, and constrictive pericarditis does not typically develop. Although there is myocarditis with acute rheumatic fever, it does not lead to dilated cardiomyopathy. A left ventricular aneurysm is a complication of ischemic heart disease. Primary cardiac neoplasms, including myxoma, are rare and not related to infection.

PBD9 557–559  BP9 391–392  PBD8 565–566  BP8 403–406

40 A Acute rheumatic fever can involve any or all layers of the heart. Because rheumatic fever follows group A streptococcal infections, the antihyaluronidase, anti-DNase, and anti-streptolysin O (ASO) titers are often elevated. The strains of group A streptococci that lead to acute rheumatic fever are less likely to cause glomerulonephritis, so an elevated creatinine level is unlikely. The ANA level could be elevated in systemic lupus erythematosus, which is most likely to produce a serous pericarditis. A positive rapid plasma reagin test suggests syphilis, but the clinical features here are not those of syphilis, and cardiovascular syphilis is one form of tertiary syphilis that develops decades after initial infection. Cardiac troponins are markers for ischemic myocardial injury. Although their levels may be elevated because of the acute myocarditis that occurs in rheumatic fever, this change is not a characteristic of rheumatic heart disease.

PBD9 557–559  BP9 391–392  PBD8 565–566  BP8 403–405

41 E The aortic valve shown has large, destructive vegetations. The probe passes through a perforated leaflet, typical of infective endocarditis caused by highly virulent organisms such as Staphylococcus aureus. The verrucous vegetations of acute rheumatic fever are small and nondestructive, and the diagnosis is suggested by an elevated anti-streptolysin O titer. A positive ANCA determination suggests a vasculitis, which is unlikely to involve cardiac valves. An elevated creatine kinase-MB level suggests myocardial, not endocardial, injury. A positive double-stranded DNA finding suggests systemic lupus erythematosus, which can produce nondestructive Libman-Sacks endocarditis.

PBD9 559–561  BP9 393–394  PBD8 567–568  BP8 406–407

42 E Infective endocarditis is present in this man. If there is a known cardiovascular congenital anomaly, then dental procedures may be preceded by prophylactic antibiotic therapy. The systolic murmur suggests a left-sided lesion, with left-to-right shunt. A small ventricular septal defect (VSD) may not lead to significant shunting of blood and remain subclinical, but it still represents a risk for endocarditis. Only a murmur may provide a clue, and the higher pitch goes with a smaller defect. Most VSDs occur in the membranous septum, but about 10% are in the muscular septum, and this difference may be due to closure of many muscular defects during life. The remaining listed options represent less common sites for anomalies or areas where infectious endocarditis could develop.

PBD9 559–561  BP9 393–394  PBD8 545, 558  BP8 388, 390
43 A She developed bacterial septicemia followed by infective endocarditis of the mitral valve. Thus she has a high risk for developing complications of infective endocarditis. Such valvular vegetations are destructive of the valve. The impaired functioning of the mitral valve (most likely regurgitation) would give rise to left atrial dilatation and left ventricular failure with pulmonary edema. Septic emboli from the mitral valve vegetation could reach the systemic circulation and give rise to abscesses. Infection of an arterial wall can weaken the wall, resulting in aneurysm formation and the potential for rupture. Dilated cardiomyopathy may be due to chronic alcoholism, or it may be idiopathic. It may be familial, or it may follow myocarditis, but it is not a direct complication of infective endocarditis. Myxomatous degeneration of the mitral valve results from a defect in connective tissue, whether well defined or unknown; the mitral valve leaflets are enlarged, hooded, and redundant. Lesions on the right side of the valve can produce septic emboli that involve the lungs, but vegetations on the left side embolize to the systemic circulation, producing lesions in the spleen, kidneys, or brain. Pulmonary abscesses can occur from right-sided infective endocarditis, because septic emboli pour into the pulmonary arterial circulation.

PBD9 559–561  BP9 393–394  PBD8 567–568  BP8 406–407

44 D Prolonged fever, heart murmur, mild splenomegaly, and splinter hemorrhages suggest a diagnosis of infective endocarditis. The valvular vegetations with infective endocarditis are friable and can break off and embolize. The time course of weeks suggests a subacute form of bacterial endocarditis resulting from infection with a less virulent organism, such as viridans streptococci. Group A streptococci are better known as a cause for rheumatic heart disease, with noninfectious vegetations. Pseudomonas aeruginosa is more likely to cause an acute form of bacterial endocarditis that worsens over days, not weeks; this organism is more common as a nosocomial infection or it may occur in injection drug users. Coxsackievirus B and Trypanosoma cruzi are causes of myocardiitis. Tuberculosis involving the heart most often manifests as pericarditis.

PBD9 559–561  BP9 393–394  PBD8 567–568  BP8 406–407

45 E So-called marantic vegetations may occur on any cardiac valve, but tend to be small and do not damage the valves. They have a tendency to embolize, however. They can occur with hypercoagulable states that accompany certain malignancies, especially mucin-secreting adenocarcinomas. Thrombosis can occur anywhere, but is most common in leg veins, predisposing to pulmonary thromboembolism. This paraneoplastic state is known as Trousseau syndrome. Calcific aortic stenosis occurs at a much older age, usually in the eighth or ninth decade, and produces obstruction but not embolism. Cardiac metastases are uncommon, and they tend to involve the epicardium; they do not explain embolism with cerebral infarction in this case. A metastatic tumor can encase the heart to produce constriction, but this is rare. Mural thromboses occur when cardiac blood flow is altered, as occurs in a ventricular aneurysm or dilated atrium, but persons with malignancies likely have no or minimal ischemic heart disease.

PBD9 561–562  BP9 394–395  PBD8 568–569  BP8 407–408

46 D Libman-Sacks endocarditis is an uncommon complication of systemic lupus erythematosus (SLE) that has minimal clinical significance because the small vegetations, although they spread over valves and endocardium, are unlikely to embolize or cause functional flow problems. Calcific aortic stenosis may be seen in older individuals with tricuspid valves, or it may be a complication of bicuspid valves. Although pericardial effusions are common in active SLE, along with pleural effusions and ascites from serositis, they are usually serious effusions, and no significant hemorrhage or scarring occurs. The vegetations of nonbacterial thrombotic endocarditis are prone to embolize. Mural thrombi are most likely to form when cardiac chambers are dilated, or there is marked endocardial damage. Rheumatic heart disease is an immunologic disease based on molecular mimicry; serologic tests would be positive for anti-streptolysin O (ASO), not ANA.

PBD9 562  BP9 395  PBD8 569  BP8 408–409

47 E Bioprostheses made from pig valves are subject to wear and tear. The leaflets may calcify, resulting in stenosis, or they may perforate or tear, leading to insufficiency. Thrombosis with embolization is unlikely to occur with bioprostheses that are indicated for persons who cannot receive anticoagulant therapy; it is an uncommon complication of mechanical prostheses, lessened by anticoagulant therapy. Hemolysis is not seen in bioprostheses and is rare in modern mechanical prostheses. Myocardial infarction from embolization or from a poorly positioned valve is rare. Paravalvular leaks are rare complications of the early postoperative period.

PBD9 563  BP9 395–396  PBD8 570–571  BP8 409

48 F Anticoagulant therapy is necessary for patients with mechanical prostheses to prevent potential thrombotic complications. If the patient is unable to take anticoagulants, use of a bioprosthesis (porcine valve) may be considered. Aspirin in low doses is used to reduce the risk for acute coronary syndromes. Antibiotic therapy with agents such as ciprofloxacin is not indicated, unless the patient has an infection or requires prophylactic antibiotic coverage for surgical or dental procedures. Cyclosporine or other immunosuppressive agents are not indicated because allogeneic tissue was not transplanted (a bioprosthesis also is essentially immunologically inert). Digoxin is not indicated because the patient’s cardiac function has improved. A β-blocker such as propranolol is not needed in the absence of chronic cardiac failure.

PBD9 563  BP9 395–396  PBD8 570–571  BP8 409

49 C Congestive heart failure with four-chamber dilation is suggestive of dilated cardiomyopathy; implicated in causation are genetic factors (in 20% to 50% of cases), myocarditis, and alcohol abuse. The patient’s fractional excretion of sodium is less than 1%, consistent with prerenal azotemia.
Many cases of dilated cardiomyopathy have no known cause. Dilation is more prominent than hypertrophy, although both are present, and all chambers are involved. Amyloidosis produces restrictive cardiomyopathy. Hypercholesterolemia would predispose to atherogenesis with coronary artery narrowing and ischemic heart disease. Rheumatic heart disease would most often produce some degree of valvular stenosis, often with some regurgitation, and the course usually is more prolonged. Chagas disease from *T. cruzi* infection affects the right ventricle more often than the left.

50 C The findings point to dilated cardiomyopathy (DCM) with both right-sided and left-sided heart failure. The most common toxin producing DCM is alcohol, and individuals with chronic alcoholism are more likely to have DCM than to have ischemic heart disease. Acetaminophen ingestion can be associated with hepatic necrosis and analgesic nephropathy. Cocaine can produce ischemic effects on the myocardium. Lisinopril is an angiotensin-converting enzyme inhibitor that is used to treat hypertension. Nicotine in cigarette smoke is a risk factor for atherosclerosis. Propranolol is a β-blocker that has been used to treat hypertension, and it may exacerbate bradycardia and congestive heart failure.

51 A Arrhythmogenic right ventricular cardiomyopathy (arrhythmogenic right ventricular dysplasia) is most likely an autosomal dominant inherited condition with abnormal desmosomal adhesion proteins in myocytes. Infections of the heart are accompanied by inflammation, though a late finding in Chagas disease is ventricular fibrosis with ventricular wall thinning. Hypertension leads to ventricular hypertrophy. There is no characteristic gross or microscopic finding with long QT syndrome caused by myocyte channelopathies. Prior radiation therapy results in fibrosis, but it is not likely to be localized to the right ventricle; improving techniques that focus the beam and synchronize it with breathing motion reduce cardiac damage when treating chest cancers.

52 B Hypertrophic cardiomyopathy is familial in >70% of cases and is usually transmitted as an autosomal dominant trait. The mutations affect genes that encode proteins of cardiac contractile elements. The most common mutation in the inherited forms affects the β-myosin heavy chain. Autoimmune conditions are unlikely to involve the myocardium. Amyloidosis causes restrictive cardiomyopathy. Hemochromatosis can give rise to cardiomyopathy, but it occurs much later in life. Viral infections produce generalized inflammation and cardiac dilation.

53 B Hypertrophic cardiomyopathy is the most common cause of sudden unexplained death in young athletes. There is asymmetric septal hypertrophy that reduces the ejection fraction of the left ventricle, particularly during exercise. Histologically, haphazardly arranged hypertrophic myocardial fibers are seen. Arrhythmias can occur. If persons with this condition survive to adulthood, chronic heart failure may develop. Hemochromatosis gives rise to a cardiomyopathy in middle age. Valve destruction with vegetations is seen in infective endocarditis and would be accompanied by signs of sepsis. Rheumatic heart disease with chronic valvular changes would be unusual in a patient this age, and the course is most often slowly progressive. Tachyzoites of *Toxoplasma gondii* signify myocarditis, a process that may occur in immunocompromised individuals.

54 B Reduced cardiac chamber compliance is a feature of the restrictive form of cardiomyopathy. Cardiac amyloidosis may be limited to the heart (so-called senile cardiac amyloidosis derived from transthyretin protein) or may be part of organ involvement in systemic amyloidosis derived from serum amyloid-associated (SAA) protein or, in multiple myeloma, derived from light chains (AL amyloid). Incidental isolated atrial deposits of amyloid are derived from atrial natriuretic peptide. Myocardial fiber dysfunction markedly reduces ventricular compliance. Dynamic left ventricular outflow obstruction is characteristic of hypertrophic cardiomyopathy. Valvular insufficiency of mitral and tricuspid valves can occur with dilated cardiomyopathy, which also reduces contractility and ejection fraction with increased end-systolic volume.

55 A Focal myocardial necrosis with a lymphocytic infiltrate is consistent with viral myocarditis. This is uncommon, and many cases may be asymptomatic. In North America, most cases are caused by coxsackieviruses A and B. This illness may often be self-limited. Less often, it ends in sudden death or progresses to chronic heart failure. Mycobacterial infections of the heart are uncommon, but pericardial involvement is the most likely pattern. Septicemia with bacterial infections may involve the heart, but the patient probably would be very ill with multiple organ failure. Viridans streptococci and *Streptococcus aureus* are better known as causes of endocarditis with neutrophilic inflammatory infiltrates. *Toxoplasma gondii* may cause myocarditis with mixed inflammatory cell infiltrates in immunocompromised patients. *Trypanosoma cruzi* is the causative agent of Chagas disease, seen most often in children. This is probably the most common infectious cause of myocarditis worldwide.

56 C Fibrinous pericarditis leads to the rough, corrugated brownish surfaces of epicardium and reflected pericardial sac as shown, which is sometimes described as a “bread and butter” appearance (after dropping the buttered bread on the carpet). Friction between epicardial and pericardial surfaces yields the rub, which may disappear with fluid collection (serofibrinous pericarditis). The most common cause is uremia resulting from renal failure. Elevation of the anti-streptolysin O titer accompanies rheumatic fever. Acute rheumatic fever may produce fibrinous pericarditis, but rheumatic fever is uncommon at this age. An elevated renin level is seen in some forms of hypertension, but by itself does not indicate renal failure.
Elevation of serum creatine kinase occurs in myocardial infarction. An acute myocardial infarction may be accompanied by a fibrinous exudate over the area of infarction, not the diffuse pericarditis seen in this patient. A positive ANA test result suggests a collagen vascular disease, such as systemic lupus erythematosus, more likely associated with a serous pericarditis (without extensive fibrinous exudate). Fibrinous pericarditis is unlikely the result of an infection, but a fibrinous-purulent appearance could suggest bacterial infection.

PBD9 573–574  BP9 403–404  PBD8 581–582  BP8 396, 416

57 D The clinical features are those of pericarditis with effusion, and the most common causes of hemorrhagic pericarditis are metastatic carcinoma and tuberculosis. An effusion of this size is sufficient to produce some cardiac tamponade that diminishes cardiac output; the paradoxical drop in pressure (more than 10 mm Hg) is called pulsus paradoxus and can be caused by pericarditis and by tamponade. *Candida* is a rare cardiac infection in immunocompromised individuals. Coxsackieviruses are known to cause myocarditis. Group A streptococci are responsible for rheumatic fever; in the acute form, rheumatic fever can cause pericarditis. Other viruses that have been reported to cause pericarditis include adenovirus, coxsackievirus, mumps virus, and varicella. Coxsackievirus B produces pericarditis more commonly.

PBD9 574  BP9 403–404  PBD8 581–582  BP8 416–417

58 A The figure shows dark red blood filling the opened pericardial cavity, a massive hemopericardium with pericardial tamponade. After excluding trauma, a complication of ischemic heart disease should be suspected. Rupture of a transmural myocardial infarction typically occurs 3 to 7 days after onset, when there is maximal necrosis before significant healing of the infarct. Ischemic heart disease occurs in patients of his age, and risk factors such as obesity, smoking, diabetes mellitus, and hyperlipidemia can play a role in its development. Cardiomyopathies lead to ventricular hypertrophy or dilation, or both, but do not cause rupture. Tuberculosis can cause hemorrhagic pericarditis, particularly without tamponade. Scleroderma is most likely to produce serous effusion. Metastases from melanoma and other carcinomas can produce hemorrhagic pericarditis without tamponade. This patient does not have a marfanoid habitus, although Marfan syndrome can cause cystic medial necrosis involving the aorta, leading to aortic dissection that can cause an acute hemopericardium. Takayasu arteritis can involve coronary arteries with aneurysms and rupture, but is most often a rare pediatric condition.

PBD9 573  BP9 404  PBD8 581  BP8 396, 417

59 C Hemorrhagic pericardial effusion most commonly is caused by either tumor or tuberculosis. The most common neoplasm involving the heart is metastatic cancer, because primary cardiac neoplasms are rare. The most common primary sites are nearby — lung, breast, and esophagus. The skin lesion in this patient is likely to be a malignant melanoma, which tends to metastasize widely, including to the heart. Most cardiac metastases involve the epicardium/pericardium. (By convention, even though epicardial surfaces are often involved most severely, the term pericardial effusion is typically used when fluid is present, or pericarditis is used when inflammation is present.)

A large effusion can cause tamponade, which interferes with cardiac motion. Calcific aortic stenosis leads to left-sided congestive heart failure, with pulmonary edema as a key finding. Coronary atherosclerosis may lead to myocardial infarction, which can be complicated by ventricular rupture and hemopericardium, but the level of troponin I in this case suggests that infarction did not occur. Rheumatic heart disease mainly affects the cardiac valves, but acute rheumatic fever can produce fibrinous pericarditis. Tuberculosis is unlikely in this case because no pulmonary lesions were seen on the radiograph.

PBD9 574–576  BP9 404  PBD8 582  BP8 416–417

60 C Atrial myxoma is the most common primary cardiac neoplasm. On the left side of the heart, it can produce a ball-valve effect that intermittently occludes the mitral valve, leading to syncopal episodes and possible strokes from embolization to cerebral arteries. Calcification of a bicuspid valve can lead to stenosis and heart failure, but this condition is progressive. Coronary artery thrombosis results in an acute ischemic event, typically with chest pain. By the time left atrial enlargement with mural thrombosis and risk of embolization occurs from mitral stenosis, this patient would have been symptomatic for years. Most pericardial effusions are not large and do not cause major problems. Large effusions could lead to tamponade, but this is not an intermittent problem.

PBD9 575–576  BP9 404–405  PBD8 583–584  BP8 417–418

61 E Endomyocardial biopsies are routinely performed after cardiac transplantation to monitor possible immune rejection, and acute cellular rejection amenable to therapy with immunosuppression is described here. Turnover of cellular organelles occurs constantly by autophagy, generating lipofuscin pigment in the cells, but this slow process is not a feature of rejection. The transplant is foreign tissue to the host, so rejection is not an autoimmune process. Months to years later, coronary arteriopathy characteristic of cardiac transplantations may produce ischemic changes. Infection is a definite possibility because of the immunosuppressive drugs administered to control the rejection process, although plasma cells are not a key feature of acute infection.

PBD9 577  BP9 405  PBD8 585  BP8 418–419

62 C Nearly every allograft develops some degree of arteriopathy within 10 years, and half of patients have significant arteriopathy by 5 years following transplantation. Unlike atherosclerosis, the smaller coronary artery branches are preferentially affected, but the result is the same: ischemic damage. The inflammation that may come from rejection is not a risk for amyloid deposition. A transplanted heart does not have a functional pericardial sac, and though there may be some fibrous adhesions following surgery, they are not constrictive. Transplant recipients receiving immunosuppressive therapy have immune dysregulation that increases the risk for both carcinomas and lymphoid malignancies, but these are unlikely to involve the heart. The immunosuppression with antirejection drugs increases the risk for infection with opportunistic agents, but these are not common and they can often be treated.

PBD9 577  BP9 405  PBD8 585  BP8 418
CHAPTER 13
Hematopathology of White Blood Cells

1. A 14-year-old boy has a high fever of 10 days’ duration. Physical examination shows a temperature of 38.3°C; pulse, 100/min; respiratory rate, 28/min; and blood pressure, 80/40 mm Hg. He has scattered petechial hemorrhages on the trunk and extremities. There is no enlargement of liver, spleen, or lymph nodes. The CBC shows hemoglobin, 13.2 g/dL; hematocrit, 38.9%; MCV, 93 μm³; platelet count, 175,000/mm³; and WBC count, 1850/mm³ with 1% segmented neutrophils, 98% lymphocytes, and 1% monocytes. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia  
B. Acute myelogenous leukemia  
C. Aplastic anemia  
D. Idiopathic thrombocytopenic purpura  
E. Systemic inflammatory response syndrome

2. A 29-year-old, HIV-positive woman has developed fever, cough, and dyspnea over the past week. On physical examination, her temperature is 37.9°C. There is dullness to percussion over lung fields posteriorly. A bronchoalveolar lavage is performed, and cysts of Pneumocystis jiroveci are present. She is given trimethoprim/sulfamethoxazole. One week later, her respiratory status has improved. Laboratory studies now show hemoglobin, 7.4 g/dL; hematocrit, 22.2%; MCV, 98 μm³; platelet count, 47,000/mm³; and WBC count, 1870/mm³ with 2% segmented neutrophils, 2% bands, 85% lymphocytes, 10% monocytes, and 1% eosinophils. One week later, she experiences increasing dyspnea, and a chest CT scan shows multiple 1- to 3-cm nodules with hemorrhagic borders in all lung fields. These nodules are most likely to be caused by infection with which of the following organisms?

A. Aspergillus fumigatus  
B. Bartonella henselae  
C. Mycobacterium avium complex

3. A 41-year-old man has had fevers with chills and rigors for the past 2 weeks. On physical examination, his temperature is 39.2°C. CBC shows hemoglobin, 13.9 g/dL; hematocrit, 40.5%; MCV, 93 μm³; platelet count, 210,000/mm³; and WBC count, 13,750/mm³ with the peripheral blood smear shown in the figure. A bone marrow biopsy specimen shows hypercellularity. Which of the following is most likely to cause these findings?

A. Acute viral hepatitis  
B. Chronic myelogenous leukemia  
C. Glucocorticoid therapy  
D. Lung abscess  
E. Splenomegaly
UNIT II  Diseases of Organ Systems

4. A 23-year-old woman has noticed that she develops a skin rash if she spends prolonged periods outdoors. She has a malar skin rash on physical examination. Laboratory studies include a positive ANA test result with a titer of 1:1024 and a “rim” pattern. An anti–double-stranded DNA test result also is positive. The hemoglobin concentration is 12.1 g/dL; hematocrit is 35.5%, MCV is 89 μm³, platelet count is 109,000/mm³, and WBC count is 4500/mm³. Which of the following findings is most likely to be shown by a WBC differential count?

A  Basophilia  
B  Eosinophilia  
C  Monocytosis  
D  Neutrophilia  
E  Thrombocytosis

5. A 23-year-old man undergoing chemotherapy for acute lymphoblastic leukemia has developed a fever and abdominal pain within the past week. He now has a severe cough. On physical examination, his temperature is 38.4°C. On auscultation, crackles are heard over all lung fields. Laboratory studies show hemoglobin, 12.8 g/dL; hematocrit, 39%; MCV, 90 μm³; platelet count, 221,000/mm³; and WBC count, 16,475/mm³ with 51% segmented neutrophils, 5% bands, 18% lymphocytes, 8% monocytes, and 18% eosinophils. Infection with which of the following organisms is most likely to be complicating the course of this patient’s disease?

A  Cryptococcus neoforms  
B  Cytomegalovirus  
C  Pseudomonas aeruginosa  
D  Strongyloides stercoralis  
E  Toxoplasma gondii  
F  Varicella-zoster virus

6. A 28-year-old man is brought to the emergency department with shock that developed over the past 12 hours. On physical examination, his temperature is 38.6°C; pulse is 101/min; respirations are 26/min, and blood pressure is 80/40 mm Hg. Needle tracks are noted in the left antecubital fossa. Crackles are heard over the lower lung fields. CBC shows hemoglobin, 14.1 g/dL; hematocrit, 42.6%; MCV, 93 μm³; platelet count, 127,500/mm³; and WBC count, 12,150/mm³ with 71% segmented neutrophils, 8% bands, 14% lymphocytes, and 7% monocytes. The neutrophils show cytoplasmic toxic granulations and Döhle bodies. Which of the following is the most likely diagnosis?

A  Acute myelogenous leukemia  
B  Chronic myelogenous leukemia  
C  Infectious mononucleosis  
D  Pneumocystis jiroveci pneumonia  
E  Pseudomonas aeruginosa septicemia  
F  Pulmonary Mycobacterium tuberculosis

7. A 26-year-old man has had a fever with nonproductive cough for the past 10 weeks. On examination, his temperature is 37.4°C. A chest radiograph shows a 4-cm left upper lobe nodule. CBC shows hemoglobin, 13.3 g/dL; hematocrit, 40.5%; platelet count, 281,000/mm³; and WBC count, 13,760/mm³ with 38% segmented neutrophils, 2% bands, 45% lymphocytes, and 15% monocytes. What is the most likely diagnosis?

A  Acute lymphoblastic leukemia/lymphoma  
B  Hodgkin lymphoma, lymphocyte rich type  
C  Mycobacterium tuberculosis granuloma

8. A 36-year-old woman has a cough and fever for 1 week. On physical examination, her temperature is 38.3°C. She has diffuse crackles in all lung fields. A chest radiograph shows bilateral extensive infiltrates. CBC shows hemoglobin, 13.9 g/dL; hematocrit, 42%; MCV, 89 μm³; platelet count, 210,000/mm³; and WBC count, 56,000/mm³ with 63% segmented neutrophils, 16% bands, 7% metamyelocytes, 3% myelocytes, 1% blasts, 8% lymphocytes, and 2% monocytes. A bone marrow biopsy is obtained and shows normal maturation of myeloid cells. Which of the following is the most likely diagnosis?

A  Chronic myelogenous leukemia  
B  Hairy cell leukemia  
C  Hodgkin lymphoma, lymphocyte depletion type  
D  Leukemoid reaction  
E  Myelodysplastic syndrome

9. A 23-year-old, previously healthy man has experienced malaise and a low-grade fever and sore throat for 2 weeks. On physical examination, his temperature is 37.6°C, and he has pharyngeal erythema without exudation. There is tender cervical, axillary, and inguinal lymphadenopathy. Laboratory studies show hemoglobin, 12.2 g/dL; hematocrit, 36.6%; platelet count, 190,200/mm³; and WBC count, 8940/mm³. His peripheral blood smear is shown in the figure. Which of the following is the most likely risk factor for his illness?

A  Close personal contact (kissing)  
B  Ingestion of raw oysters  
C  Inherited disorder of globin chain synthesis  
D  Sharing infected needles for injection drug use  
E  Transfusion of packed RBCs

10. A 6-year-old boy has complained of worsening pain in the right side of his groin for the past week. Physical examination shows painful, swollen lymph nodes in the right inguinal region. An inguinal lymph node biopsy is performed, and on microscopy the node has large, variably sized, germinal centers containing numerous mitotic figures. There are numerous parafollicular and sinusoidal neutrophils. What is the most likely cause of these histologic changes?

A  Acute lymphadenitis  
B  Acute lymphoblastic leukemia  
C  Cat-scratch disease  
D  Follicular lymphoma  
E  Sarcoidosis  
F  Toxoplasmosis
11. A 39-year-old woman felt a lump in her breast 1 week ago. On physical examination, she has a firm, fixed, irregular 3-cm mass in the upper outer quadrant of the right breast and a firm, nontender lymph node in the right axilla. A lumpectomy and axillary node dissection are performed, and microscopic examination shows an infiltrating ductal carcinoma in the breast. Flow cytometric analysis of the node shows a polyclonal population of CD3+, CD19+, CD20+, and CD68+ cells with no aneuploidy. Microscopic examination of the axillary lymph node is most likely to reveal changes characteristic of which of the following conditions?

A. Acute lymphadenitis
B. Diffuse large B-cell lymphoma
C. Metastatic infiltrating ductal carcinoma
D. Necrotizing granulomas
E. Plasmacytosis
F. Sinus histiocytosis
G. Sézary syndrome

12. A 9-year-old, otherwise healthy girl has complained of pain in the right armpit for the past week. Examination shows tender, enlarged lymph nodes in the right axillary region. There are four linear and nearly healed abrasions over a 3 × 2 cm area of the distal ventral aspect of the right forearm and a single, 0.5-cm, slightly raised erythematous nodule over one of the abrasions. No other abnormalities are noted. Histologic examination of one of the lymph nodes shows stellate, necrotizing granulomas. The lymphadenopathy regresses over the next 2 months. Infection with which of the following is most likely to have produced these findings?

A. Bartonella henselae
B. Cytomegalovirus
C. Epstein-Barr virus
D. Staphylococcus aureus
E. Yersinia pestis

13. A study of persons with lymphoid malignancies reveals that there are risk factors for development of B-cell non-Hodgkin lymphomas. Which of the following is the most likely inherited condition predisposing to lymphoid malignancies?

A. Cystic fibrosis
B. Hereditary spherocytosis
C. Sickle cell disease
D. Von Willebrand disease
E. Wiskott-Aldrich syndrome

14. A 14-year-old boy complains of discomfort in his chest that has worsened over the past 5 days. On physical examination, he has generalized lymphadenopathy. A chest radiograph shows clear lung fields, but there is widening of the mediastinum. A chest CT scan shows a 10-cm mass in the anterior mediastinum. A biopsy specimen of the mass is obtained and microscopically shows effacement by lymphoid cells with lobulated nuclei having delicate, finely stippled, nuclear chromatin. There is scant cytoplasm, and many mitotic figures are seen. The cells express deoxynucleotidyl transferase negative (TdT−), CD2, and CD7 antigens. Molecular analysis reveals a point mutation in the NOTCH1 gene. The oncologist tells the parents that chemotherapy can be curative in vast majority of such cases. What is the most likely diagnosis?

A. Burkitt lymphoma
B. Follicular lymphoma
C. Hodgkin lymphoma, nodular sclerosing type
D. Lymphoblastic lymphoma
E. Mantle cell lymphoma
F. Small lymphocytic lymphoma

15. A 70-year-old man has experienced increasing fatigue for the past 6 months. On physical examination, he has nontender axillary and cervical lymphadenopathy, but there is no hepatosplenomegaly. The CBC shows hemoglobin, 9.5 g/dL; hematocrit, 28%; MCV, 90 μm³; platelet count, 120,000/mm³; and WBC count, 42,000/mm³. His peripheral blood smear shows a monotonous population of small, round, mature-looking lymphocytes. Flow cytometry shows these cells to be CD19+, CD5+, and deoxynucleotidyl transferase negative (TdT−). Cytogenetic and molecular analysis of the abnormal cells in his blood are most likely to reveal which of the following alterations?

A. Clonal rearrangement of immunoglobulin genes
B. Clonal rearrangement of T-cell receptor genes
C. t(8;14) leading to c-MYC overexpression
D. t(9;22) leading to BCR-ABL rearrangement
E. t(14;18) leading to BCL2 overexpression

16. A 69-year-old man notices the presence of lumps in the right side of his neck that have been enlarging for the past year. Physical examination shows firm, nontender posterior cervical lymph nodes 1 to 2 cm in diameter. The overlying skin is intact and not erythematous. A lymph node is biopsied and the microscopic appearance is shown in the figure. Molecular analysis of the DNA extracted from the cells reveals B cell receptor gene rearrangements. Which of the following features provides the best evidence for malignant lymphoma in this node?

A. Absence of a pattern of lymphoid follicles with germinal centers
B. Diminished sinusoidal plasma cells and immunoblasts
C. Presence of CD30+ large, multinucleated cells
D. Uniform expression of kappa, but not lambda, light chains in the lymphoid cells
E. Proliferation of small capillaries in the medullary and paracortical regions
17. A 4-year-old boy has appeared listless during the past week. He exhibits irritability when his arms or legs are touched. In the past 2 days, large ecchymoses have appeared on the right thigh and left shoulder. CBC shows hemoglobin, 9.3 g/dL; hematocrit, 28.7%; MCV, 96 μm³; platelet count, 45,000/mm³; and WBC count, 13,990/mm³. Examination of the peripheral blood smear shows blasts that lack peroxidase-positive granules, but contain PAS-positive aggregates and stain positively for deoxynucleotidyl transferase negative (TdT−). Flow cytometry shows the phenotype of blasts to be CD19+, CD3−, and sIg−. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia
B. Acute myelogenous leukemia
C. Chronic lymphocytic leukemia
D. Chronic myelogenous leukemia
E. Idiopathic thrombocytopenic purpura

18. A 7-year-old boy has complained of a severe headache for the past week. On physical examination, there is tenderness on palpation of long bones, hepatosplenomegaly, and generalized lymphadenopathy. Petechial hemorrhages are present on the skin. Laboratory studies show hemoglobin, 8.8 g/dL; hematocrit, 26.5%; platelet count, 34,700/mm³; and WBC count, 14,800/mm³. A bone marrow biopsy specimen shows 100% cellularity, with almost complete replacement by a population of large cells with scant cytoplasm lacking granules, delicate nuclear chromatin, and rare nucleoli. His oncologist is confident that chemotherapy will induce a complete remission. Which of the following combinations of phenotypic and karyotypic markers is most likely to be present in marrow cells from this boy?

A. Early pre-B CD19+ hyperdiploidy
B. Early pre-B CD20+ t(9;22)
C. Pre-B CD5+ normal karyotype
D. Pre-B CD23+ 11q deletion
E. T cell CD2+ numerous blasts
F. T cell CD3+ MLL gene translocation

19. A 66-year-old man has noted an increasing number and size of lumps over his body in the past 5 months. On examination, there is firm, nontender inguinal, axillary, and cervical lymphadenopathy. A biopsy specimen of a cervical node shows a histologic pattern of nodular aggregates of small, cleaved lymphoid cells and larger cells with open nuclear chromatin, several nucleoli, and moderate amounts of cytoplasm. A bone marrow biopsy specimen shows lymphoid aggregates of similar cells with surface immunoglobulins that are CD10+, but CD5−. Karyotyping of these lymphoid cells indicates the presence of t(14;18). What is the most likely diagnosis?

A. Acute lymphadenitis
B. Hodgkin lymphoma, nodular sclerosis type
C. Follicular lymphoma
D. Mantle cell lymphoma
E. Toxoplasmosis

20. The figure skater Sonja Henie, who won gold medals at the 1928, 1932, and 1936 Winter Olympic Games, became progressively fatigued in her late 50s. On physical examination, she had palpable nontender axillary and inguinal lymph nodes, and the spleen tip was palpable. Laboratory studies showed hemoglobin, 10.1 g/dL; hematocrit, 30.5%; MCV, 90 μm³; platelet count, 89,000/mm³; and WBC count, 31,300/mm³. From the peripheral blood smear shown in the figure, which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia
B. Chronic lymphocytic leukemia
C. Infectious mononucleosis
D. Iron deficiency anemia
E. Leukemoid reaction

21. A 37-year-old man infected with HIV for the past 10 years is admitted to the hospital with abdominal pain of 3 days’ duration. Physical examination shows abdominal distention and absent bowel sounds. An abdominal CT scan shows a mass lesion involving the ileum. He undergoes surgery to remove an area of bowel obstruction in the ileum. Gross examination of the specimen shows a firm, white mass, 10 cm long and 3 cm at its greatest depth. The mass has infiltrated through the wall of the ileum. Histologic studies show a mitotically active population of CD19+ lymphoid cells with prominent nuclei and nucleoli. Molecular analysis is most likely to show which of the following viral genomes in the lymphoid cells?

A. Cytomegalovirus
B. Epstein-Barr virus
C. HIV
D. Human herpesvirus 8
E. Human T-cell lymphotropic virus type 1
22. A 12-year-old boy has had increasing abdominal distention and pain for the past 3 days. Physical examination of his abdomen shows lower abdominal tenderness with tympany and reduced bowel sounds. An abdominal CT scan shows a 7-cm mass involving the region of the ileocecal valve. Surgery is performed and the resected mass microscopically shows sheets of intermediate-sized lymphoid cells, with nuclei having coarse chromatin, several nucleoli, and many mitotic figures. A bone marrow biopsy sample is negative for this cell population. Cytogenetic analysis of the cells from the mass shows a t(8;14) karyotype. Flow cytometric analysis reveals 40% of the cells are in S phase. The tumor shrinks dramatically after a course of chemotherapy. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia/lymphoma  
B. Burkitt lymphoma  
C. Diffuse large B-cell lymphoma  
D. Follicular lymphoma  
E. Plasmacytoma

23. A 55-year-old man felt a lump near his shoulder 1 week ago. On physical examination, there is an enlarged, nontender, supraclavicular lymph node and enlargement of the Waldeyer ring of oropharyngeal lymphoid tissue. There is no hepatosplenomegaly. CBC is normal except for findings of mild anemia. A lymph node biopsy specimen shows replacement by a monomorphous population of lymphoid cells that are twice the size of normal lymphocytes, with enlarged nuclei and prominent nucleoli. Immunohistochemical staining and flow cytometry of the node indicates that most lymphoid cells are CD19+, CD10+, CD3−, CD15−, and terminal deoxynucleotidyl transferase negative (TdT−). A BCL6 gene mutation is present. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic lymphoma  
B. Chronic lymphadenitis  
C. Diffuse large B-cell lymphoma  
D. Hodgkin lymphoma  
E. Small lymphocytic lymphoma

24. A 62-year-old man has experienced vague abdominal discomfort accompanied by bloating and diarrhea for the past 6 months. On physical examination, there is a midabdominal firm mass. The stool is positive for occult blood. An abdominal CT scan shows an 11 × 4 cm mass involving the wall of the distal ileum and adjacent mesentery. A laparotomy is performed, and the mass is removed. Microscopically, the mass is composed of sheets of large lymphoid cells with large nuclei, prominent nucleoli, and frequent mitotic figures. The neoplastic cells mark with CD19+ and CD20+ and have the BCL6 gene rearrangement. Which of the following prognostic features is most applicable to this case?

A. Aggressive, can be cured by chemotherapy  
B. Aggressive, often spreads to liver, spleen, and marrow  
C. Aggressive, often transforms to acute leukemia  
D. Indolent, can be cured by chemotherapy  
E. Indolent, often undergoes spontaneous remission  
F. Indolent, survival of 7 to 9 years without treatment

25. A 9-year-old boy living in Uganda has had increasing pain and swelling on the right side of his face over the past 8 months. On physical examination, there is a large, nontender mass involving the mandible, which deforms the right side of his face. There is no lymphadenopathy or splenomegaly, and he is afebrile. A biopsy of the mass is performed and microscopic examination shows intermediate-sized lymphocytes with a high mitotic rate. A chromosome analysis shows a 46,XY,t(8;14) karyotype in these cells. The hemoglobin concentration is 13.2 g/dL, platelet count is 272,000/mm³, and WBC count is 5820/mm³. Infection with which of the following viruses is most likely to be causally related to the development of these findings?

A. Cytomegalovirus  
B. Epstein-Barr virus  
C. Hepatitis B virus  
D. HIV  
E. Human papillomavirus  
F. Respiratory syncytial virus

26. A 61-year-old man reports back pain for 5 months. He has recently developed a cough that is productive of yellow sputum. On physical examination, he is febrile, and diffuse rales are heard on auscultation of the lungs. He has no lymphadenopathy or splenomegaly. Laboratory studies include a sputum culture growing Streptococcus pneumoniae. The serum creatinine level is 3.7 mg/dL, and the urea nitrogen level is 35 mg/dL. The figure shows a skull radiograph. A bone marrow biopsy specimen from this man is most likely to show increased numbers of which of the following?

A. Myeloblasts  
B. Small mature lymphocytes  
C. Plasma cells  
D. Reed-Sternberg cells  
E. Non-necrotizing granulomas
27 A 67-year-old man has had increasing weakness, fatigue, and weight loss over the past 5 months. He now has decreasing vision in both eyes and has headaches and dizziness. His hands are sensitive to cold. On physical examination, he has generalized lymphadenopathy and hepatosplenomegaly. Laboratory studies indicate a serum protein level of 15.5 g/dL and albumin concentration of 3.2 g/dL. A bone marrow biopsy is performed, and microscopic examination of the specimen shows infiltration by numerous small plasmacytoid lymphoid cells with Russell bodies in the cytoplasm. Which of the following additional laboratory findings is most likely to be reported for this patient?

A  Bence Jones proteinuria  
B  Hypercalcemia  
C  Karyotype with t(14;18) translocation  
D  Monoclonal IgM spike in serum  
E  WBC count of 255,000/mm³

28 A 69-year-old woman complains of increasing back pain for 1 month. On physical examination, there is tenderness over the lower back, but no kyphosis or scoliosis. A radiograph of the spine shows a partial collapse of T10 and multiple 0.5- to 1.5-cm lytic lesions with a rounded soap bubble appearance in the thoracic and lumbar vertebrae. A bone marrow biopsy is performed, and a smear of the aspirate is shown in the figure. Which of the following laboratory findings is most likely to be seen in this patient?

A  Bence Jones proteins in the urine  
B  Decreased serum alkaline phosphatase level  
C  Hypogammaglobulinemia  
D  Platelet count of 750,000/mm³  
E  t(9;22) in the karyotype of marrow cells  
F  WBC count of 394,000/mm³

29 A 48-year-old man has a routine health maintenance examination. He has no concerns other than worrying about getting older and having cancer. Physical examination shows that he is afebrile and normotensive. There is no hepatosplenomegaly or lymphadenopathy. Laboratory studies show a total serum protein level of 7.4 g/dL and albumin level of 3.9 g/dL. Serum calcium and phosphorus levels are normal. Urinalysis shows no Bence Jones proteinuria. Hemoglobin is 13.6 g/dL, platelet count is 301,500/mm³, and WBC count is 6630/mm³. The results of the serum protein electrophoresis are shown in the figure. A bone marrow biopsy specimen shows normal cellularity with maturation of all cell lines. Plasma cells constitute about 4% of the marrow. The results of a bone scan are normal, and there are no areas of increased uptake. What is the most likely diagnosis?

A  Heavy chain disease with lymphoplasmacytic lymphoma  
B  Monoclonal gammopathy of undetermined significance  
C  Multiple myeloma with IgD immunophenotype  
D  Reactive systemic amyloidosis  
E  Solitary plasmacytoma of the lung  
F  Waldenström macroglobulinemia with hyperviscosity

30 A 62-year-old man has had fever and a 4-kg weight loss over the past 6 months. On physical examination, his temperature is 38.6° C. He has generalized nontender lymphadenopathy, and the spleen tip is palpable. Laboratory studies show hemoglobin, 10.1 g/dL; hematocrit, 30.3%; platelet count, 140,000/mm³; and WBC count, 24,500/mm³ with 10% segmented neutrophils, 1% bands, 86% lymphocytes, and 3% monocytes. A cervical lymph node biopsy specimen microscopically shows a nodular pattern of small lymphoid cells. A bone marrow biopsy specimen shows infiltrates of similar small cells having surface immunoglobulins that are CD5+, but CD10−. Cytogenetic analysis indicates t(11;14) in these cells. What is the most likely diagnosis?

A  Acute lymphoblastic lymphoma  
B  Burkitt lymphoma  
C  Follicular lymphoma  
D  Mantle cell lymphoma  
E  Small lymphocytic lymphoma
A 54-year-old woman has experienced nausea with vomiting and early satiety for the past 7 months. On physical examination, she is afebrile and has no lymphadenopathy or hepatosplenomegaly. CBC shows hemoglobin, 12.9 g/dL; hematocrit, 41.9%; platelet count, 263,000/mm³; and WBC count, 8430/mm³. An upper gastrointestinal endoscopy shows loss of the rugal folds of the stomach over a 4 × 8 cm area of the fundus. Gastric biopsy specimens reveal the presence of *Helicobacter pylori* organisms in the mucus overlying superficial epithelial cells. There are extensive mucosal and submucosal monomorphous infiltrates of small lymphocytes, which are CD19+ and CD20+, but CD3−. After treatment of the *H. pylori* infection, her condition improves. What is the most likely diagnosis?

A  Acute lymphoblastic leukemia  
B  Chronic lymphocytic leukemia  
C  Diffuse large B-cell lymphoma  
D  Follicular lymphoma  
E  Hodgkin lymphoma, mixed cellularity type  
F  MALT (marginal zone) lymphoma

A 41-year-old man has experienced several bouts of pneumonia over the past year. He now complains of vague abdominal pain and a dragging sensation. Physical examination shows marked splenomegaly. CBC shows hemoglobin, 8.2 g/dL; hematocrit, 24.6%; MCV, 90 μm³; platelet count, 63,000/mm³; and WBC count, 2400/mm³. The peripheral blood smear shows many small leukocytes with the features shown in the figure. A chest radiograph shows patchy infiltrates, and a culture of sputum grows *Mycobacterium kansasii*. Which of the following laboratory findings is most characteristic of this disorder?

A  Cytoplasmic Auer rods in myeloid cells  
B  CD20 and CD11c positive lymphocytes  
C  Monoclonal IgM in serum  
D  t(9;22) translocation in stem cells  
E  Cytoplasmic toxic granulations in neutrophils

A 29-year-old, previously healthy man has had an enlarging nodular area on his arm for the past 8 months. On physical examination, there is an ulcerated, reddish violet, 3 × 7 cm lesion on his right forearm and nontender right axillary and left inguinal lymphadenopathy. A chest radiograph shows a 4-cm nodular left pleural mass. An abdominal CT scan shows a 5-cm right retroperitoneal mass. Biopsy of an inguinal node is performed, and microscopic examination shows large cells, some of which contain horseshoe-shaped nuclei and voluminous cytoplasm. The tumor cells cluster around venules and infiltrate sinuses. The patient goes into remission after chemotherapy. Which of the following markers is most likely to be positive in the tumor cells?

A  ALK protein  
B  CD10 antigen  
C  c-KIT proto-oncogene  
D  IL-2 receptor  
E  p24 antigen

A 41-year-old man has experienced several bouts of pneumonia over the past year. He now complains of vague abdominal pain and a dragging sensation. Physical examination shows marked splenomegaly. CBC shows hemoglobin, 12.9 g/dL; hematocrit, 42%; platelet count, 231,000/mm³; and WBC count, 7940/mm³ with 57% segmented neutrophils, 3% bands, 26% lymphocytes, 5% monocytes, and 9% eosinophils. A skin biopsy specimen microscopically shows the presence of lymphoid cells in the upper dermis and epidermis. These cells have cerebriform nuclei with marked infolding of nuclear membranes. Similar cells are seen on the peripheral blood smear. Which combination of the following phenotypic markers is most likely to be expressed on his abnormal lymphocytes?

A  CD3+, CD4+  
B  CD5+, CD56+  
C  CD10+, CD19+  
D  CD13+, CD33+  
E  CD19+, slg+

A 58-year-old man from Nagasaki, Japan, has noted an increasing number of skin lesions for the past 8 months. On examination, there are scaling red-brown patches on all skin surfaces. He also has generalized lymphadenopathy and hepatosplenomegaly. Laboratory studies show hemoglobin, 9.7 g/dL; hematocrit, 31%; MCV, 89 μm³; platelet count, 177,000/mm³; and WBC count, 18,940/mm³ with differential count of 35% segmented neutrophils, 2% band neutrophils, 58% lymphocytes, and 5% monocytes. His serum calcium is 11.5 mg/dL. Examination of his peripheral blood smear shows multilobated “cloverleaf” cells. Which of the following infectious agents most likely caused his illness?

A  Bartonella henselae  
B  Cytomegalovirus (CMV)  
C  Epstein-Barr virus (EBV)  
D  Helicobacter pylori  
E  HIV  
F  HTLV-1

A 26-year-old man has noted lumps in his neck that have been enlarging for the past 6 months. On physical examination, he has a group of enlarged, nontender right cervical lymph nodes. A biopsy of one of the lymph nodes microscopically shows macrophages, lymphocytes, neutrophils, eosinophils, and a few plasma cells. There are scattered CD15+ large cells with multiple nuclei or a single nucleus with multiple nuclear lobes, each with a large inclusion-like nucleolus. What is the most likely cell of origin with infectious agent for these large cells?

A  B lymphocyte, Epstein-Barr virus  
B  CD4+ cell, human T lymphotrophic virus  
C  Endothelial cell, Kaposi sarcoma herpesvirus  
D  Macrophage, human immunodeficiency virus  
E  NK cell, cytomegalovirus
37 A 34-year-old woman reports having generalized fatigue and night sweats for 3 months. Physical examination shows nontender right cervical lymphadenopathy. Biopsy of one lymph node is performed, and microscopic examination shows a pattern of thick bands of fibrous connective tissue with intervening lymphocytes, plasma cells, eosinophils, macrophages, and occasional Reed-Sternberg cells. An abdominal CT scan and bone marrow biopsy specimen show no abnormalities. Which of the following is the most likely subtype of this patient’s disease?

A Lymphocyte depletion
B Lymphocyte predominance
C Lymphocyte rich
D Mixed cellularity
E Nodular sclerosis

38 A 74-year-old man has experienced recurrent fevers and a 6-kg weight loss over the past 5 months. On physical examination, his temperature is 37.5° C, and he has splenomegaly. An abdominal CT scan shows mesenteric lymphadenopathy. A lymph node biopsy specimen shows effacement of the nodal architecture by a population of small lymphocytes, plasma cells, eosinophils, and macrophages. Which of the following additional cell types, which stains positively for CD15 and CD30, is most likely to be found in this disease?

A Epithelioid cell
B Immunoblast
C Mast cell
D Myeloblast
E Reed-Sternberg cell

39 A 63-year-old man has noticed a lump in his neck for 2 months. Examination reveals a group of three discrete nontender right posterior cervical lymph nodes, and a mass of enlarged right axillary lymph nodes. Chest and abdominal CT scans show mediastinal lymphadenopathy and hepatosplenomegaly. Microscopic examination of a cervical lymph node biopsy reveals abundant large CD15+ and CD30+ binucleate cells with prominent acidophilic nucleoli, scattered within a sparse lymphocytic infiltrate. What is molecular analysis of this lesion most likely to reveal?

A Clonal EBV integration in the large cells
B BCL6 gene rearrangements in the large cells
C Deletions of 5q in all the cells
D Helicobacter pylori infection in all the cells
E JAK2 gene mutations in the lymphocytes

40 A 22-year-old woman has experienced increasing dyspnea for the past 2 months. On physical examination, she is afebrile and normotensive. Inspiratory wheezes are noted on auscultation of the chest. A chest CT scan shows an 8 × 10 cm posterior mediastinal mass that impinges on the trachea and esophagus. A mediastinoscopy is performed, and a biopsy of the mass microscopically shows scattered large multinucleated cells, with prominent nucleoli that mark with CD15, and lymphocytes and macrophages separated by dense collagenous bands. Which of the following cells is most likely to be seen microscopically in this biopsy specimen?

A Atypical lymphocytes
B Hairy cells
C Langerhans cells
D Lacunar cells
E Lymphoblasts
F Myeloblasts

41 A 29-year-old woman has experienced fatigue, fever, night sweats, and painless lumps in the right side of her neck for the past 3 months. On physical examination, her temperature is 37.5° C. She has right cervical nontender lymphadenopathy. One of the lymph nodes is biopsied, and a histologic finding is shown at high power in the figure. A molecular analysis of large cells exemplified by the cell at the center is most likely to reveal which of the following genetic abnormalities?

A Clonal rearrangement of T-cell receptor genes
B Clonal rearrangement of immunoglobulin genes
C Integration of the HTLV-1 genome
D Integration of the human herpesvirus 8 genome
E Polyclonal rearrangement of T-cell receptor genes
F Polyclonal rearrangement of immunoglobulin genes

42 A 33-year-old man has experienced multiple nosebleeds along with bleeding gums for the past month. On examination, his temperature is 37.3° C. He has multiple cutaneous ecchymoses. Laboratory studies show hemoglobin, 8.5 g/dL; hematocrit, 25.7%; platelet count, 13,000/mm³; and WBC count, 52,100/mm³ with 5% segmented neutrophils, 5% bands, 2% myelocytes, 83% blasts, 3% lymphocytes, and 2% monocytes. Examination of his peripheral blood smear shows the blasts have delicate nuclear chromatin along with fine cytoplasmic azurophilic granules. These blasts are CD33+. Which of the following morphologic findings is most likely to be present on his peripheral blood smear?

A Auer rods
B Döhle bodies
C Hairy projections
D Heinz bodies
E Sickle cells
F Toxic granulations
43. A 38-year-old man experiences sudden onset of a severe headache. Physical examination shows no localizing neurologic signs and no organomegaly. A stool sample is positive for occult blood. Areas of purpura appear on the skin of his extremities. Laboratory studies show hemoglobin of 9.6 g/dL, hematocrit of 28.9%, platelet count of 16,400/mm³, and WBC count of 75,000/mm³. The peripheral blood smear has the appearance shown in the figure; schistocytes also are seen. The plasma D-dimer level (fibrin degradation products), prothrombin time, and partial thromboplastin time all are elevated. Cytogenetic analysis of cells from a bone marrow biopsy specimen is most likely to yield what karyotypic abnormality?

A. t(8;14)  
B. t(8;21)  
C. t(9;22)  
D. t(14;18)  
E. t(15;17)

44. A 38-year-old woman has had bleeding gums for the past 3 weeks. Physical examination shows that her gingivae are thickened and friable. She has hepatosplenomegaly and generalized nontender lymphadenopathy. CBC shows hemoglobin, 11.2 g/dL; hematocrit, 33.9%; MCV, 89 μm³; platelet count, 95,000/mm³; and WBC count, 4500/mm³ with 25% segmented neutrophils, 10% bands, 2% metamyelocytes, 5% lymphocytes, 8% monocytes, and 1 nucleated RBC per 100 WBCs. Her peripheral blood smear is shown in the figure, and these cells contain peroxidase positive granules. A bone marrow biopsy specimen shows 100% cellularity with few residual normal hematopoietic cells. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia  
B. Acute myelogenous leukemia  
C. Chronic lymphocytic leukemia  
D. Chronic myelogenous leukemia  
E. Hodgkin lymphoma

45. A 22-year-old university student reports easy fatigability of 2 months’ duration. On physical examination, she has no hepatosplenomegaly or lymphadenopathy. Mucosal gingival hemorrhages are noted. CBC shows hemoglobin, 9.5 g/dL; hematocrit, 28.2%; MCV, 94 μm³; platelet count, 20,000/mm³; and WBC count, 107,000/mm³. Her peripheral blood smear is shown in the figure, and these cells contain peroxidase positive granules. A bone marrow biopsy specimen shows 100% cellularity with few residual normal hematopoietic cells. Which of the following is the most likely diagnosis?

A. Acute lymphoblastic leukemia  
B. Acute myelogenous leukemia  
C. Acute megakaryocytic leukemia  
D. Acute monocytic leukemia  
E. Acute promyelocytic leukemia

46. A 50-year-old man with a diffuse large B-cell lymphoma underwent intensive chemotherapy, and a complete remission was achieved for 7 years. He now reports fatigue and recurrent pulmonary and urinary tract infections over the past 4 months. Physical examination shows no masses, lymphadenopathy, or hepatosplenomegaly. CBC shows hemoglobin, 8.7 g/dL; hematocrit, 25.2%; MCV, 88 μm³; platelet count, 67,000/mm³; and WBC count, 2300/mm³ with 15% segmented neutrophils, 5% bands, 2% metamyelocytes, 2% myelocytes, 6% myeloblasts, 33% lymphocytes, 35% monocytes, and 2% eosinophils. A bone marrow biopsy specimen shows 90% cellularity with many immature cells, including ringed sideroblasts, megaloblasts, hypolobated megakaryocytes, and myeloblasts. Karyotypic analysis shows 5q deletions in many cells. This clinical picture is most consistent with which of the following conditions?

A. De novo acute myeloblastic leukemia  
B. Myeloid metaplasia with myelofibrosis  
C. Myelodysplasia related to therapy  
D. Relapse of his previous lymphoma  
E. Transformation into myeloid leukemia
In an experiment, cell samples are collected from the bone marrow aspirates of patients who were diagnosed with lymphoproliferative disorders. Cytogenetic analyses are performed on these cells, and a subset of the cases is found to have the \textit{BCR-ABL} fusion gene from the reciprocal translocation t(9;22)(q34;11). The presence of this gene results in increased tyrosine kinase activity. Patients with which of the following conditions are most likely to have this gene?

A. Acute promyelocytic leukemia  
B. Chronic myelogenous leukemia  
C. Follicular lymphoma  
D. Hodgkin lymphoma, lymphocyte depletion type  
E. Multiple myeloma

A 63-year-old woman experiences a burning sensation in her hands and feet. Two months ago, she had an episode of swelling with tenderness in the right leg, followed by dyspnea and right-sided chest pain. On physical examination, the spleen and liver now appear to be enlarged. On physical examination, the spleen and liver now appear to be enlarged. CBC shows hemoglobin, 13.3 g/dL; hematocrit, 40.1%; MCV, 91 μm³; platelet count, 657,000/mm³; and WBC count, 17,400/mm³. The peripheral blood smear shows abnormally large platelets. Which of the following is the most likely diagnosis?

A. Acute myelogenous leukemia  
B. Chronic myelogenous leukemia  
C. Essential thrombocytosis  
D. Myelofibrosis with myeloid metaplasia  
E. Polycythemia vera

A 60-year-old woman has had headaches and dizziness for the past 5 weeks. She has been taking omeprazole for ulcers. On physical examination, she is afebrile and normotensive, and her face has a plethoric to cyanotic appearance. There is mild splenomegaly, but no other abnormal findings. Laboratory studies show hemoglobin, 21.7 g/dL; hematocrit, 65%; platelet count, 400,000/mm³; and WBC count, 30,000/mm³ with 85% polymorphonuclear leukocytes, 10% lymphocytes, and 5% monocytes. The peripheral blood smear shows abnormally large platelets and nucleated RBCs. Which of the following is most characteristic of the natural history of this patient’s disease?

A. Development of a gastric non-Hodgkin lymphoma  
B. Increase in monoclonal serum immunoglobulin  
C. Marrow fibrosis with extramedullary hematopoiesis  
D. Spontaneous remissions and relapses without treatment  
E. Transformation into acute B-lymphoblastic leukemia

A 50-year-old man has experienced a gradual weight loss and weakness, anorexia, and easy fatigability for 9 months. Physical examination shows marked splenomegaly. CBC shows hemoglobin, 12.9 g/dL; hematocrit, 38.1%; MCV, 92 μm³; platelet count, 410,000/mm³; and WBC count, 168,000/mm³. The peripheral blood smear is depicted in the figure. Karyotypic analysis shows the \( \Phi^1 \) chromosome. The patient undergoes chemotherapy with imatinib mesylate (tyrosine kinase inhibitor). He remains in remission for 3 years and then begins to experience fatigue and an 8-kg weight loss. CBC shows hemoglobin, 10.5 g/dL; hematocrit, 30%; platelet count, 60,000/μL; and WBC count, 40,000/μL. Karyotypic analysis shows two \( \Phi^1 \) chromosomes and aneuploidy. Flow cytometric analysis of the peripheral blood shows CD19+, CD10+, sIg−, and CD3− cells. Which of the following complications of the initial disease did this patient develop after therapy?

A. Acute myeloblastic leukemia  
B. B-cell lymphoblastic leukemia  
C. Hairy cell leukemia  
D. Myelodysplastic syndrome  
E. Sézary syndrome

A 50-year-old man has had headache, dizziness, and fatigue for the past 3 months. His friends have been commenting about his increasingly ruddy complexion. He also has experienced generalized and severe pruritus, particularly when showering. He notes that his stools are dark. On physical examination, he is afebrile, and his blood pressure is 165/95 mm Hg. There is no hepatosplenomegaly or lymphadenopathy. A stool sample is positive for occult blood. CBC shows hemoglobin, 22.3 g/dL; hematocrit, 67.1%; MCV, 94 μm³; platelet count, 453,000/mm³; and WBC count, 7800/mm³. What is the most likely diagnosis?

A. Chronic myelogenous leukemia  
B. Erythroleukemia  
C. Essential thrombocytosis  
D. Myelodysplastic syndrome  
E. Polycythemia vera
A 66-year-old man has experienced fatigue, a 5-kg weight loss, night sweats, and abdominal discomfort for 10 months. On physical examination, he has marked splenomegaly; there is no lymphadenopathy. Laboratory studies show hemoglobin, 10.1 g/dL; hematocrit, 30.5%; MCV, 89 μm³; platelet count, 94,000/mm³; and WBC count, 14,750/mm³ with 55% segmented neutrophils, 9% bands, 20% lymphocytes, 8% monocytes, 4% metamyelocytes, 3% myelocytes, 1% eosinophils, and 2 nucleated RBCs per 100 WBCs. The peripheral blood smear also shows teardrop cells. The serum uric acid level is 12 mg/dL. A bone marrow biopsy specimen shows extensive marrow fibrosis and clusters of atypical megakaryocytes. Which of the following is most likely to account for the enlargement in this patient’s spleen?

A 9-year-old boy has developed a fever over the past 15 days. He has been diagnosed and treated for otitis media multiple times in the past year. On physical examination, he has mild lymphadenopathy, hepatomegaly, and splenomegaly. There are extensive crusted papules on his skin. A CT scan of his head shows a 4-cm osteolytic mass in the mastoid bone. A biopsy of the mass is performed, with the electron micrograph shown in the figure. What is the most likely diagnosis?

A 20-year-old man is left at the door of the emergency department by “friends” after they spent an evening at a local pub. On examination, his vital signs are temperature, 37° C; pulse, 110/min; respirations, 26/min; and blood pressure, 75/40 mm Hg. He has left upper quadrant tenderness on palpation. An abdominal CT scan was obtained and is shown in the figure. What is the most likely underlying cause of this clinical picture?
A 60-year-old man with a history of chronic alcohol abuse has had increasing abdominal discomfort and fatigue for the past 9 months. He has noted easy bruising of his skin with minor trauma for the past month. On examination, he is afebrile, but his spleen is enlarged and tender. Laboratory studies show hemoglobin, 7.7 g/dL; hematocrit, 23%; platelet count, 30,000/mm³; and WBC count, 2300/mm³ with 45% polymorphonuclear leukocytes, 50% lymphocytes, and 5% monocytes. What is the most likely diagnosis?

A. Acute myelogenous leukemia
B. Infectious mononucleosis
C. Micronodular cirrhosis
D. Niemann-Pick disease
E. Metastatic adenocarcinoma
F. Systemic lupus erythematosus

A 27-year-old man has had a fever with chills and rigors for the past 10 days. On physical examination, his temperature is 37.9° C, pulse is 87/min, respirations are 21/min, and blood pressure is 100/55 mm Hg. A diastolic murmur is heard on auscultation of the chest. There is a tender, palpable spleen tip. Laboratory studies show hemoglobin, 12.8 g/dL; hematocrit, 38.4%; platelet count, 23,000/mm³; and WBC count, 12,980/mm³ with 69% segmented neutrophils, 8% bands, 1% metamyelocytes, 18% lymphocytes, and 4% monocytes. The representative gross appearance of the spleen is shown in the figure. What is the most likely underlying condition responsible for the changes in the spleen?

A. Acute myelogenous leukemia
B. Disseminated histoplasmosis
C. Hodgkin lymphoma
D. Infective endocarditis
E. Metastatic carcinoma
F. Micronodular cirrhosis
G. Rheumatic heart disease

A 14-year-old boy has developed a cough and a high fever over the past 4 days. On physical examination, he has a temperature of 39.2° C. Diffuse rales are heard over all lung fields. Laboratory studies show hemoglobin, 14.8 g/dL; hematocrit, 44.4%; platelet count, 496,000/mm³; and WBC count, 15,600/mm³. Examination of the peripheral blood smear shows RBCs with marked anisocytosis and Howell-Jolly bodies. A sputum culture grows *Haemophilus influenzae*. Which of the following is the most likely diagnosis?

A. DiGeorge syndrome
B. Galactosemia
C. Gaucher disease
D. Myeloproliferative disorder
E. Prior splenectomy
F. Trisomy 21

A clinical study is performed in which the subjects are children 1 to 4 years old who have had multiple infections with viral, fungal, and parasitic diseases. Compared with a normal control group, these children do not have thymic cells that bear markers of cortical lymphocytes. Which of the following karyotypic abnormalities is most likely to be seen in the children in this study?

A. +21
B. 22q11.2
C. t(9;22)
D. t(15;17)
E. X(fra)
F. XXY

A 49-year-old woman has experienced increasing weakness and chest pain over the past 6 months. On physical examination, she is afebrile and normotensive. Motor strength is 5/5 in all extremities, but diminishes to 4/5 with repetitive movement. There is no muscle pain or tenderness. Laboratory studies show hemoglobin, 14 g/dL; hematocrit, 42%; platelet count, 246,000/mm³; and WBC count, 6480/mm³. A chest CT scan shows an irregular 10 × 12 cm anterior mediastinal mass. The surgeon has difficulty removing the mass because it infiltrates surrounding structures. Microscopically, the mass is composed of large, spindled, atypical epithelial cells mixed with lymphoid cells. Which of the following is the most likely cause of this mass lesion?

A. Extrapulmonary tuberculosis
B. Hodgkin lymphoma
C. Lymphoblastic lymphoma
D. Malignant thymoma
E. Metastatic breast carcinoma
F. Organizing abscess
**1 E** The major finding in this patient is marked granulocytopenia. All that remains on the peripheral smear are mononuclear cells (remember to multiply the percentages in the differential by the total WBC count to get the absolute values; rather than one cell line being overrepresented, another may be nearly missing). Accelerated removal or destruction of neutrophils could account for the selective absence of granulocytes in this case. Overwhelming acute infections or other causes for widespread innate immune response can lead to increased peripheral use of neutrophils at sites of inflammation. Petechial hemorrhages also can occur in overwhelming bacterial infections, such as those caused by Neisseria meningitidis. Bleeding is unlikely to be caused by thrombocytopenia because in this case the platelet count is normal. Normal bone marrow findings exclude acute lymphoid or myeloid leukemia. In aplastic anemia, the marrow is poorly cellular, and there is a reduction in RBCs, WBCs, and platelet production.

**2 A** Her severe pancytopenia resulted from drug toxicity. This predisposed her to subsequent sepsis, with aspergillosis as the cause of pulmonary nodules, and neutropenia the significant risk factor. These fungal organisms often invade blood vessels, producing hemorrhagic lesions. Bartonellosis can produce bacillary angiomatosis, which is more likely to involve the skin. Mycobacterium avium complex is more likely to involve organs of the mononuclear phagocyte system and unlikely to produce large nodules. Escherichia coli, similar to many bacterial infections, can occur in HIV infection, but it has a pattern of acute neutrophilic infiltrates. Herpes simplex virus (type 1 or 2) is an unlikely disseminated infection in HIV. Pneumocystis pneumonia rarely produces nodular lesions. Toxoplasmosis is uncommon in the lung, even in immunocompromised individuals.

**3 D** The figures shows a neutrophilic leukocytosis, and there can also be a “left shift” from increased band neutrophils. Chronic infections and ongoing inflammatory conditions, such as lung abscesses, can lead to an expansion of the myeloid precursor pool in the bone marrow with myeloid hyperplasia. Acute viral hepatitis, in contrast to acute bacterial infections, does not cause neutrophilic leukocytosis. In chronic myelogenous leukemia, the marrow is filled with myeloid cells throughout the spectrum of maturation, and more immature forms in the peripheral blood, including metamyelocytes, myelocytes, and even a few blasts, along with increased eosinophils and basophils. Glucocorticoids can increase the release of marrow storage pool cells and diminish extravasation of neutrophils into tissues. Vigorous exercise can produce neutrophilia transiently from demargination of neutrophils. A large spleen tends to sequester peripheral blood cells, reducing their circulating numbers.

**4 C** An autoimmune disease, most likely systemic lupus erythematosus (SLE) in this patient, can be accompanied by monocytosis. Cytopenias also can occur in SLE because of autoantibodies against blood elements, a form of type II hypersensitivity. Basophilia occurs infrequently, but also can be seen in chronic myelogenous leukemia (CML). Eosinophilia is a feature more often seen in allergic conditions, tissue parasitic infestations, and CML. Neutrophilia is seen in acute infectious and inflammatory conditions. Thrombocytosis usually occurs in neoplastic disorders of myeloid stem cells, such as the myeloproliferative disorders that include CML and essential thrombocytosis.

**5 D** The eosinophilia suggests a parasitic infestation. Immunocompromised individuals can have superinfection and dissemination with strongyloidiasis. Type 1 hypersensitivity with allergic reactions may also be accompanied by eosinophilia. The other organisms listed are not known to be associated with eosinophilia.

**6 E** Toxic granulations, which are coarse and dark primary granules, and Döhle bodies, which are patches of dilated endoplasmic reticulum, represent reactive changes of neutrophils that are most indicative of overwhelming inflammatory conditions, such as bacterial sepsis. The route of infection in this case is injection drug use. Infectious mononucleosis is accompanied by an increase in “atypical” lymphocytes. Leukemia, granulomatous infections, or viral infections do not cause toxic changes in neutrophils.

**7 C** This granulomatous infection has led to leukocytosis with lymphocytosis and monocytosis. Blood monocytes become tissue macrophages that evolve into epithelioid cells and giant cells of granulomatous inflammation. The most consistent form of leukocytosis from infection is neutrophilia with acute bacterial infections, such as Staphylococcus aureus infection. Viral, mycobacterial, and fungal infections produce less consistent peripheral blood findings. An acute lymphoblastic leukemia is likely to be accompanied by a higher WBC count with circulating lymphoblasts. Hodgkin lymphomas have no consistent peripheral blood findings and are not likely to produce solitary lung nodules. Myelodysplastic syndromes are marked by the presence of immature myeloid cells and cytopenias in the peripheral blood.

**8 D** Marked leukocytosis and immature myeloid cells in the peripheral blood can represent an exaggerated response to infection (leukemoid reaction), or it can be a manifestation of chronic myelogenous leukemia (CML). Normal maturation of myeloid cells in the marrow rules out CML. Although not provided in this case, a leukocyte alkaline phosphatase (LAP) score is high in the more differentiated cell population of reactive leukocytosis, whereas in CML, the LAP score is low. The Philadelphia chromosome (present in most CML cases) is lacking in patients with leukemoid reactions. Hairy
cell leukemia is accompanied by peripheral blood leukocytes that mark with tartrate-resistant acid phosphatase. Hodgkin lymphoma is not characterized by an increased WBC count. A myelodysplastic syndrome is a stem cell maturation disorder involving all nonlymphoid cell lineages, not just granulocytes.

9 A The smear shows large “atypical” lymphocytes with abundant cytoplasm indented by red cells, and these are present most often in patients with infectious mononucleosis, and sometimes other viral infections, such as cytomegalovirus. These atypical cells are large lymphocytes with abundant cytoplasm and a large nucleus with fine chromatin. Infectious mononucleosis is caused by Epstein-Barr virus (EBV) and transmitted by close personal contact. In patients with infectious mononucleosis, the EBV genes cause proliferation and activation of multiple clones of B cells, and there is polyclonal B-cell expansion. These B cells secrete antibodies with several specificities, including antibodies that cross-react with sheep RBCs. These heterophil antibodies produce a positive monospot test result. The atypical lymphocytes are CD8+ T cells that are activated by EBV-infected B cells. There is no increase in basophils, eosinophils, or monocytes in infectious mononucleosis. Eating raw oysters is a risk factor for hepatitis A because oysters that filter polluted seawater concentrate the virus in their tissues. Disorders of globin chain synthesis affect RBCs, as in the thalassemias. Infectious mononucleosis is not known as a transfusion-associated disease. Likewise, intravenous drug use is typically not a risk for hepatitis A because infected needles are at risk for bacterial infections, HIV infection, and viral hepatitis.

10 A Painful and acute enlarged nodes suggest a reactive condition and not a neoplastic process such as a lymphoma or a leukemia. In children, enlarged tender nodes and acute lymphadenitis are common. Many infectious processes can give rise to these findings, particularly bacterial infections. Children are “antigen sponges” when exposed to the usual minor infections; they are quite active and acquire plenty of cuts and scrapes on extremities, which can become infected, with reactive hyperplasia of regional nodes. Cat-scratch disease can produce sarcoïdlike granulomas with stellate abscesses. Follicular lymphomas are B-cell neoplasms that efface the normal architecture of the lymph nodes; these tumors do not occur in children. Sarcoïdosis is a chronic granulomatous process typically seen in adults and characterized by the formation of noncaseating granulomas. Toxoplasmosis can be a congenital infection or can be seen in immunocompromised individuals; it produces a pattern of follicular hyperplasia.

11 F Lymph nodes draining from a cancer often show a reactive pattern, with dilated sinusoids that have endothelial hypertrophy and are filled with histiocytes (i.e., macrophages). Sinus histiocytosis represents an immunologic response to cancer antigens. Thus not all enlarged nodes are caused by metastatic disease in cancer patients. CD3 is a T-cell marker, CD19 and CD20 are B-cell markers, and CD68 is a macrophage (histiocyte) marker. Polyclonal proliferations are typically benign reactive processes, whereas a monoclonal proliferation suggests a neoplasm. Aneuploidy and high S phase are characteristics of malignant neoplasms; a high S phase mostly occurs in rapidly growing tumors, such as diffuse large B-cell lymphomas, and in a few carcinomas, such as small-cell anaplastic carcinoma. Inflammation would produce pain and tenderness, and the patient may be febrile. Generalized inflammatory diseases or chronic infections can increase the number of plasma cells in lymph nodes.

12 A Cat-scratch disease is a form of self-limited infectious lymphadenitis that most often is seen in children, typically “downstream” of lymphatic drainage from the site of an injury on a distal extremity. Hence axillary and cervical lymph node regions are most often involved. Cytomegalovirus infection is typically seen in immunocompromised individuals and is not a common cause of lymphadenopathy. Epstein-Barr virus (EBV) infection at this age is most often associated with infectious mononucleosis and pharyngitis, and the lymphadenopathy is nonspecific. *Staphylococcus aureus* can produce suppurative inflammation with sepsis. *Yersinia pestis*, the agent that causes bubonic plague, produces lymphadenopathy that can ulcerate and a hemorrhagic necrotizing lymphadenitis; it has a high mortality rate.

13 E Both primary and acquired immune deficiencies produce immune dysregulation from which malignancies may arise. These malignancies are most often neoplastic proliferations of white cells: lymphomas, leukemias, and Hodgkin lymphoma, but carcinomas may also occur. Of the primary immune deficiencies, Wiskott-Aldrich syndrome has the highest percentage of lymphomas among malignancies that develop. Epstein-Barr virus infection may be an additional triggering event. Conversely, persons with lymphoid malignancies may develop secondary immunodeficiency, with increased risk for opportunistic infection. The other listed options are not specifically linked to development of malignancies.

14 D His age and the mediastinal location are typical of a lymphoblastic lymphoma involving the thymus. This lesion is within the spectrum of acute lymphoblastic leukemia or lymphoma (ALL). Most cases of ALL with lymphomatous presentation are of the pre–T-cell type, supported by the expression of the T-cell markers CD2 and CD7. The *NOTCH1* gene encodes a transmembrane receptor required for T-cell development, and more than half of pre–T-cell tumors have activating point mutations. TdT is a marker of pre–T cells and pre–B cells. A Burkitt lymphoma is a B-cell lymphoma that also can be seen in adolescents, but usually is present in the jaw or abdomen. Nodular sclerosing follicular lymphomas...
and mantle cell lymphomas are B-cell tumors usually seen in older patients, and they do not involve the thymus. Hodgkin lymphoma does occur in the mediastinum, but it involves mediastinal nodes, not thymus. The histologic features of Hodgkin lymphoma include the presence of Reed-Sternberg cells, and this variant has fibrous bands intersecting the lymphoid cells. Small lymphocytic lymphoma is the tissue phase of chronic lymphocytic leukemia seen in older adults.

**15 A** The clinical history, the peripheral blood smear, and the phenotypic markers are characteristic of chronic lymphocytic leukemia (CLL), a clonal B-cell neoplasm in which immunoglobulin genes are rearranged, and T-cell receptor genes are in germline configuration. T and B-cell lymphomas are variably present in reactive conditions, but their absence can often be shown by immunostaining that reveals one light chain in the neoplastic cells. Some lymphoid neoplasms are indolent tumors that continue to accumulate cells for 7 to 9 years. The lymphoid population in acute lymphadenitis is reactive, and there is no bone marrow involvement. In Hodgkin lymphoma, there are few Reed-Sternberg cells, surrounded by a reactive lymphoid population. Mantle cell lymphoma also is a B-cell tumor; it is more aggressive than follicular lymphoma and is typified by the t(11;14) translocation, in which the cyclin D1 gene (BCL1) is overexpressed. In toxoplasmosis, there would be a mixed population of inflammatory cells and some necrosis.

**16 D** The figure shows a follicular non-Hodgkin lymphoma. All lymphoid neoplasms are derived from a single transformed cell and are monoclonal. B-cell neoplasms comprise 80% to 85% of all lymphoid neoplasms, and their monoclonality is often seen by immunostaining that reveals one light chain in the neoplastic cells. Populations of normal or reactive (polyclonal) B cells contain a mixture of B cells expressing both kappa and lambda light chains. Some lymphoid neoplasms have a follicular pattern. A normal pattern of follicles is sometimes absent if the node is involved, as in some inflammatory conditions or in immunosuppression. In keeping with this, follicular lymphomas are indolent tumors that continue to accumulate cells for 7 to 9 years. The lymphoid population in acute lymphadenitis is reactive, and there is no bone marrow involvement. In Hodgkin lymphoma, there are few Reed-Sternberg cells, surrounded by a reactive lymphoid population. Mantle cell lymphoma also is a B-cell tumor; it is more aggressive than follicular lymphoma and is typified by the t(11;14) translocation, in which the cyclin D1 gene (BCL2) is overexpressed. In toxoplasmosis, there would be a mixed population of inflammatory cells and some necrosis.

**17 A** These findings are characteristic of a childhood acute lymphoblastic leukemia of the precursor-B-cell type. The rapid expansion of the marrow caused by proliferation of blasts can lead to bone pain and tenderness. Features supporting an acute leukemia are anemia, thrombocytopenia, and the presence of blasts in the peripheral blood and bone marrow. Anemia and thrombocytopenia result from suppression of normal hematopoiesis by the leukemic clone in the marrow. The phenotype of CD19+, CD3−, and slg− is typical of pre-B cells. TdT is a marker of early T-cell-type and B-cell-type lymphoid cells. An acute myelogenous leukemia is a disease of young to middle-aged adults, and there would be peroxidase-positive myeloblasts and phenotypic features of myeloid cells. Chronic lymphocytic leukemia is a disease of older adults; patients have many small circulating mature B lymphocytes. Chronic myelogenous leukemia is a disease of adults, and the WBC count is quite high; the peripheral blood contains some myeloblasts, but other stages of myeloid differentiation also are detected. In idiopathic thrombocytopenic purpura, only the platelet count is reduced because of antibody-mediated destruction of platelets.

**18 A** These markers strongly favor a very good prognosis for acute lymphoblastic leukemia (ALL), the most common malignancy in children: early precursor-B-cell type, hyperdiploidy, and patient age between 2 and 10 years, chromosomal trisomy, and t(12;21). Marrow infiltration by the leukemic cells leads to pancytopenia. Poor prognostic markers for acute lymphoblastic leukemia/lymphoma are T-cell phenotype, patient age younger than 2 years, WBC count >100,000, presence of t(9;22) MLL gene mutations, and presentation in adolescence and adulthood. In most T-cell ALL cases in adolescents, a mediastinal mass arises in the thymus, and lymphoid infiltrates appear in tissues of the mononuclear phagocyte system. The success of a treatment plan is also aided by a caring and supportive family.

**19 C** Follicular lymphoma is the most common form of non-Hodgkin lymphoma among adults in Europe and North America. Men and women are equally affected. The neoplastic B cells mimic a population of follicular center cells and produce a nodular or follicular pattern. Nodal involvement is often generalized, but extranodal involvement is uncommon. The t(14;18) translocation, which is characteristic, causes overexpression of the BCL2 gene by juxtaposing it with the IgH locus; the cells are resistant to apoptosis. In keeping with this, follicular lymphomas are indolent tumors that continue to accumulate cells for 7 to 9 years. The lymphoid population in acute lymphadenitis is reactive, and there is no bone marrow involvement. In Hodgkin lymphoma, there are few Reed-Sternberg cells, surrounded by a reactive lymphoid population. Mantle cell lymphoma also is a B-cell tumor; it is more aggressive than follicular lymphoma and is typified by the t(11;14) translocation, in which the cyclin D1 gene (BCL2) is overexpressed. In toxoplasmosis, there would be a mixed population of inflammatory cells and some necrosis.

**20 B** Sonja Henie died from complications of chronic lymphocytic leukemia (CLL). The figure shows increased numbers of circulating small, round, mature lymphocytes with scant cytoplasm in the peripheral blood smear. The CLL cells express the CD5 marker and the pan B-cell markers CD19 and CD20. Most patients have a disease course of 4 to 6 years before death, and symptoms appear as the leukemic cells begin to fill the marrow. In some patients, the same small lymphocytes appear in tissues; the condition is then known as small lymphocytic lymphoma. Acute lymphoblastic leukemia is a disease of children and young adults, characterized by proliferation of lymphoblasts. These cells are much larger than the cells in CLL and have nucleoli. The lymphocytes seen in infectious mononucleosis are atypical lymphocytes, which have abundant, pale
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blue cytoplasm that seems to be indented by the surrounding RBCs. The RBCs in iron deficiency anemia are hypochromic and microcytic, but the WBCs are not affected. Leukemoid reactions are typically of the myeloid type, and the peripheral blood contains immature myeloid cells. The WBC count can be very high, but the platelet count is normal.

PB9 595–597  BP9 436  PB8 607–608  BP8 452–453

21 B This HIV-positive patient has an extranodal infiltrative mass, composed of B cells (CD19+), in the ileum. This is a diffuse large-cell lymphoma of B cells. These tumors contain the Epstein-Barr virus (EBV) genome, and it is thought that immune large-cell lymphoma of B cells. These tumors contain the mass, composed of B cells (CD19+), in the ileum. This is a diffuse large-cell lymphoma of B cells. These tumors contain the mass.

This is a diffuse large-cell lymphoma of B cells. These tumors contain the mass.

22 B Burkitt and Burkitt-like lymphomas can be seen sporadically (in young individuals), in an endemic form in Africa (in children), and in association with HIV infection. All forms are highly associated with translocations of the MYC gene on chromosome 8. In the African form and in HIV-infected patients, the cells are latently infected with Epstein-Barr virus (EBV), but sporadic cases are negative for EBV. This form of lymphoma is typically extranodal. Because of the high growth fraction (40% in this case), Burkitt lymphomas respond very well to chemotherapy, including agents that disrupt the cell cycle. By contrast, slow-growing tumors with a low growth fraction are more indolent and less responsive to chemotherapy. Acute lymphoblastic lymphomas can be seen in boys this age, but the mass is in the mediastinum, and the lymphocytes are T cells. Diffuse large-cell lymphomas are most common in adults, as are follicular lymphomas; they do not carry the t(8;14) translocation. Plasmacytomas appear in older adults and are unlikely to produce an abdominal mass.

PB9 597–598  BP9 436–437  PB8 607–608  BP8 453

23 C Diffuse large B-cell lymphoma occurs in older individuals and frequently manifests as localized disease with extranodal involvement, particularly of the Waldeyer ring. A third of cases have BCL6 rearrangements or mutations. The staining pattern indicates a B-cell proliferation (CD19+, CD10+). T-cell (CD3) and monocytic (CD15) markers are absent. TdT can be expressed in B lineage cells at an earlier stage of maturation. Lymphoblastic lymphoma is a T-cell neoplasm that occurs typically in the mediastinum of children. In chronic lymphadenitis, the lymph node has many cell types—macrophages, lymphocytes, and plasma cells. A monomorphous infiltrate is typical of non-Hodgkin lymphomas. Reed-Sternberg cells characterize Hodgkin lymphoma. Small lymphocytic lymphoma also is a B-cell neoplasm, but it manifests with widespread lymphadenopathy, liver and spleen enlargement, and lymphocytosis.

PB9 595–597  BP9 436  PB8 606–607  BP8 452–453

24 A Diffuse large-cell lymphoma of B cells often involve extranodal sites, show large anaplastic lymphoid cells that involve the tissues diffusely, and contain BCL6 gene rearrangements. Their clinical course is aggressive, and they become rapidly fatal if untreated. With intensive chemotherapy, however, 60% to 80% of patients achieve complete remission, and up to 50% can be cured. More aggressive lymphomas tend to be localized, whereas the indolent lymphomas tend to involve multiple nodal sites or multiple organs such as liver, spleen, and marrow.

PB9 595–597  BP9 436, 432  PB8 606–607  BP8 452–453

25 B The endemic African variety of Burkitt lymphoma is a B-cell lymphoma that typically appears in the maxilla or mandible of the jaw. This particular neoplasm is related to Epstein-Barr virus infection. Cytomegalovirus infection occurs in immunocompromised patients and can be a congenital infection, but it is not a direct cause of neoplasia. Hepatitis B virus infection can be a risk factor for hepatocellular carcinoma. HIV infection can be a risk factor for the development of non-Hodgkin lymphomas, but most of these are either diffuse large B-cell lymphomas or small noncleaved Burkitt-like lymphomas. Human papillomavirus infection is related to the formation of squamous dysplasias and carcinomas, most commonly those involving the cervix. Respiratory syncytial virus infection produces pneumonia in infants and young children, but is not related to development of neoplasms.

PB9 597–598  BP9 436–437  PB8 607–608  BP8 453

26 C Multiple myeloma produces mass lesions of plasma cells that lead to bone lysis and pain. The skull radiograph shows typical punched-out lytic lesions, produced by expanding masses of plasma cells. The Ig genes in myeloma cells always show evidence of somatic hypermutation. Hence Jones proteinuria can damage the renal tubules and give rise to renal failure. Multiple myeloma can be complicated by AL amyloid, which also can lead to renal failure. Patients with myeloma often have infections with encapsulated bacteria because of decreased production of IgG, required for opsonization. Blasts suggest a leukemic process. Nodules of small lymphocytes suggest a small-cell lymphocytic leukemia/lymphoma, which is not likely to produce lytic lesions. Reed-Sternberg cells suggest Hodgkin lymphoma. Granulomatous disease (which is not produced by pneumococcus) can involve the marrow, but usually it does not produce such sharply demarcated lytic lesions.


27 D Hyperviscosity syndrome includes visual disturbances, dizziness, headache, and Raynaud phenomena. His bone marrow is infiltrated with plasmacytoid lymphocytes that have stored immunoglobulins in their cytoplasm (Russell bodies). All of these findings are consistent with lymphoplasmacytic lymphoma (Waldenström macroglobulinemia). In this disorder, neoplastic B cells differentiate to IgM-producing
γ small (<3 g), and the patient has no Bence Jones proteinuria, the diagnosis of MGUS is made when the monoclonal spike is "spike" in the absence of any associated disease of B cells. The globulinemia would be accompanied by an IgM spike, hepato-splenomegaly, and lymphadenopathy; the protein would be very high with hyperviscosity. 

PBD9 601–602 BP9 438–440 PBD8 612 BP8 456

The characteristic punched-out bone lesions of multiple myeloma seen on radiographs result from bone destruction mediated by RANKL, a cytokine produced by the myeloma cells that activates osteoclasts. Several cytokines, most notably IL-6, are important growth factors for plasma cells. They are produced by tumor cells and by resident marrow stromal cells. High serum levels of IL-6 correlate with active disease and poor prognosis. The monoclonal population of plasma cells often produces a monoclonal serum "spike" seen in serum or urine protein electrophoresis. Myeloma is unlikely to be accompanied by leukocytosis. Patients with bone destruction and remodeling can have hypercalcemia and an increased serum alkaline phosphatase level. The neoplastic cells are generally well differentiated, with features such as a perinuclear hof, similar to normal plasma cells. The t(9;22) translocation is the Philadelphia chromosome seen in chronic myelogenous leukemia (CML), with a low leukocyte alkaline phosphatase score. Leukemias, including CML, and myeloproliferative disorders can fill the marrow spaces, sometimes are accompanied by a thrombocytosis, but are unlikely to produce mass lesions or bony destruction. 

PBD9 599–600 BP9 437–439 PBD8 610 BP8 453–456

Monoclonal gammopathy of uncertain significance (MGUS) is characterized by the presence of an M protein "spike" in the absence of any associated disease of B cells. The diagnosis of MGUS is made when the monoclonal spike is small (<3 g), and the patient has no Bence Jones proteinuria, as shown by the small (2.8 g) spike of γ-globulin, which is determined by immunoelectrophoresis to be IgG kappa. MGUS can progress to multiple myeloma in about 20% of patients over 10 to 15 years. Heavy-chain disease is a rare condition. In multiple myeloma, the spike is greater than 3 g, and usually the patient has bone lesions. In reactive systemic amyloidosis, serum amyloid-associated (SAA) protein derived from chronic inflammatory conditions is deposited as AA amyloid in visceral organs, but there is no monoclonal gammopathy. A plasmacytoma is a mass lesion that would appear on a bone scan, but they may also be extraneous. Waldenström macroglobulinemia would be accompanied by an IgM spike, hepatosplenomegaly, and lymphadenopathy; the protein would be very high with hyperviscosity. 

PBD9 599, 601 BP9 438 PBD8 611 BP8 454

The immunophenotype is characteristic for mantle cell lymphoma. Of the lesions listed, lymphoblastic lymphoma and Burkitt lymphoma occur in a much younger age group. Burkitt lymphoma has a t(8;14) translocation. The remaining three lesions occur in an older age group. Of these, small lymphocytic lymphoma manifests with absolute lymphocytosis and the peripheral blood picture of chronic lymphocytic leukemia. Follicular lymphoma has a distinct and characteristic translocation t(14;18) involving the BCL2 gene. In contrast, mantle cell lymphoma, seen in older men, has the t(11;14) translocation, which activates the cyclin D1 (BCL1) gene; these tumors do not respond well to chemotherapy, particularly when it involves the peripheral blood. 

PBD9 602–603 BP9 435 PBD8 612–613 BP8 452

Marginal zone lymphomas arise in middle-aged adults at sites of autoimmune or infectious stimulation. If the lesion is associated with lymphoid tissue, it is sometimes called a mucosa-associated lymphoid tissue tumor (MALT lymphoma, or MALToma). The most common sites are the thyroid (in Hashimoto thyroiditis), the salivary glands (in Sjögren syndrome), or the stomach (in Helicobacter pylori infection). Although monoclonal (similar to a neoplasm), these MALT lesions can regress with antibiotic therapy for H. pylori. A MALT lesion can transform to diffuse large B-cell lymphoma. The cells correspond to the marginal B cells found at the periphery of stimulated lymphoid follicles. The other conditions listed are neoplastic conditions that are not related to H. pylori, and that require chemotherapy to control. 

PBD9 603 BP9 442 PBD8 613–614 BP8 459

The figure shows mononuclear cells with reniform nuclei and pale blue cytoplasm with threadlike extensions. This patient has hairy cell leukemia, an uncommon neoplastic disorder of B cells. These CD19+, CD20+, and CD11c+ cells infiltrate the spleen and marrow. Pancytopenia results from poor production of hematopoietic cells in the marrow and sequestration of the mature cells in the spleen. There are two characteristic features of this disease: the presence of hairy projections from neoplastic leukocytes in the peripheral blood smear and coexpression of B-cell (CD19, CD20) and monocyte (CD11c) markers. In the past, staining for tartrate-resistant acid phosphatase (TRAP) was used to identify these abnormal cells as shown in the right panel of the figure. Auer rods are seen in myeloblasts in acute myeloblastic leukemia. A monoclonal IgM spike is a feature of lymphoplasmacytic lymphoma (Waldenström macroglobulinemia). The Ph1 chromosome is a distinctive feature of chronic myelogenous leukemia. Toxic granulations in neutrophils are seen most often in overwhelming bacterial infections. 

PBD9 603–604 BP9 442–443 PBD8 614 BP8 459–460

This patient has a form of T-cell neoplasm known as anaplastic large-cell lymphoma, which most often appears in children and young adults. It is often extranodal and has a characteristic gene rearrangement on chromosome 2p23 that results in production of anaplastic lymphoma kinase (ALK) with tyrosine kinase activity. CD10 is a B-cell marker. The
T-cell proliferations involving skin, known as mycosis fungoides/Sézary syndrome, are CD4+. The c-KIT proto-oncogene has been associated with some NK cell lymphomas. The IL-2 receptor is associated with lymphohistiocytosis with macrophage activation. The p24 antigen is part of HIV, which is most often associated with B-cell neoplasms.

34 A The involvement of skin and the presence of lymphocytes with complex cerebriform nuclei in the skin and the blood are features of cutaneous T-cell lymphomas. These are malignancies of CD4+ and CD3+ T cells that may produce a tumorlike infiltration of the skin (mycosis fungoides) or a leukemic picture without tumefaction in the skin (Sézary syndrome). Cutaneous T-cell lymphomas are indolent tumors, and patients have a median survival of 8 to 9 years. The other phenotypes provided here are those of CD3−, CD56+ NK cells; mature B cells with CD19+, slg+; monocytes/granulocytes with CD3+, CD13+; and neoplastic B cells in chronic lymphocytic leukemia with CD19+, CD5+.

35 F The patient lives in an area endemic for HTLV-1, which can cause leukemia/lymphoma and demyelinating disease. The neoplastic lymphoid cells can infiltrate many organs. Skin lesions resemble those of mycosis fungoides. Cat-scratch disease from Bartonella henselae infection results in lymphadenopathy with microscopic stellate necrosis. CMV is not associated with development of neoplasms, but it can complicate the course of patients with neoplasms who become immunocompromised. EBV can be associated with African Burkitt lymphoma. Helicobacter pylori can be associated with African Burkitt lymphoma. HIV is best known to be associated with non-Hodgkin lymphoma. EBV can be associated with MALT lymphoma with microscopic stellate necrosis. CMV is not as associated with Hodgkin lymphoma.

36 A The large cells are Reed-Sternberg cells, and they elaborate cytokines that promote an accompanying reactive cellular proliferation that forms the bulk of the neoplastic mass. Reed-Sternberg cells are of B-cell origin, and in many cases, Epstein-Barr virus (EBV) infection can be demonstrated in these cells. The other listed options are not part of the pathogenesis for Hodgkin lymphoma.

37 E The bands of fibrosis are typical of the nodular sclerosis type of Hodgkin lymphoma, which is most commonly seen in young adults, particularly women. Involvement of one group of lymph nodes places this in stage I. Mediastinal involvement is common. Most nodular sclerosis cases are stage I or II, and the prognosis of such early-stage cases is good. The findings do not fit with the other forms of Hodgkin lymphoma listed.

38 E The features suggest Hodgkin lymphoma (HL), mixed cellularity type, which tends to affect older men. As in all other forms of HL except the lymphocyte predominance type, the Reed-Sternberg cells and variants stain with CD15. These cells also express CD30, an activation marker on T cells, B cells, and monocytes. Clinical symptoms are common in the mixed cellular type of HL, and this histologic type tends to manifest in advanced stages. The Reed-Sternberg cells make up a relatively small percentage of the tumor mass, with most of the cell population consisting of reactive cells such as lymphocytes, plasma cells, macrophages, and eosinophils. Epithelioid cells are seen in granulomatous inflammatory reactions. Immunoblasts suggest a B-cell proliferation. Mast cells are not numerous in HL; they participate in type I hypersensitivity responses. Myeloblasts are numerous with acute myelogenous leukemia.

39 A The lymphocyte depletion variant of Hodgkin lymphoma has an abundance of Reed-Sternberg cells and a paucity of lymphocytes. Most cases present with advanced disease (stage IV in this example). Epstein-Barr virus (EBV) is present in over 90% of cases. BCL6 gene rearrangements are typical of diffuse large B-cell lymphomas. Deletions of 5q are typical of myelodysplastic syndrome. Infection with Helicobacter pylori can give rise to marginal zone lymphoma. JAK2 mutations are found in polycythemia vera and other myeloproliferative diseases.

40 D The lacunar cells and the CD15+ Reed-Sternberg cells indicate Hodgkin lymphoma, and the fibrous bands suggest the nodular sclerosis type. Lacunar cells have multilobed nuclei containing many small nucleoli. These cells have artificial retraction of the cytoplasm around the nucleus, giving the cells their distinctive appearance. The nodular sclerosis type of Hodgkin lymphoma is more common in women. Atypical lymphocytes are characteristic in the peripheral blood of individuals with infectious mononucleosis. Hairy cell leukemia often is accompanied by splenomegaly, but not a mediastinal mass, and the leukemic cells are B cells. Histiocytes with Birbeck granules are characteristic of the Langhans cell histiocytes. Lymphoblasts that mark as T cells are seen in anterior mediastinal (thymic) masses in children with acute lymphoblastic leukemia/lymphoma. Myeloblasts are characteristic of acute myelogenous leukemia, which is occasionally accompanied by soft-tissue masses.
found in cells of Kaposi sarcoma. HTLV-1 infects CD4+ T cells and gives rise to adult T-cell leukemia/lymphoma.

42 A Acute myelogenous leukemia (AML) infiltrates the marrow and reduces normal hematopoiesis to account for anemia and marked thrombocytopenia. The initial presentation may be acute. The Auer rods are condensations of the azurophilic granules. Döhle bodies, which are patches of dilated endoplasmic reticulum, and toxic granulations, which are coarse and dark primary granules, are reactive changes in mature neutrophils most indicative of marked inflammation, such as bacterial sepsis. Hairy projections are seen on the circulating B cells of hairy cell leukemia. Heinz bodies are seen in G6PD deficiency and are precipitates of denatured globin. Sickling of RBCs is a feature of sickle cell anemia, which is not related to leukemia.

43 E The peripheral blood smear is characteristic of acute promyelocytic leukemia (M3 class of acute myelogenous leukemia), with many promyelocytes containing prominent azurophilic granules and short, red, cytoplasmic, rodlike inclusions called Auer rods. Release of the granules can trigger the coagulation cascade, leading to disseminated intravascular coagulation (DIC). As in this case, many patients develop DIC. The t(15;17) translocation is characteristic of this disease; it results in the fusion of the retinoic acid receptor gene on chromosome 17 with the promyelocytic leukemia gene on chromosome 15. The fusion gene results in elaboration of an abnormal retinoic acid receptor that blocks myeloid differentiation. Therapy with retinoic acid (vitamin A) can alleviate the block and induce remission in many patients. The t(8;14) translocation can be seen in patients with Burkitt lymphoma. The t(8;21) abnormality is seen in the M2 variant of acute myelogenous leukemia. The t(9;22) translocation gives rise to Philadelphia chromosome of chronic myelogenous leukemia. A t(14;18) karyotype suggests a follicular lymphoma.

44 D This patient has an aleukemic leukemia, in which the peripheral blood count of leukocytes is not high, but the leukemic blasts fill the marrow. These blasts show features of monoblasts because they are peroxidase negative and nonspecific esterase positive. This patient has an M5 leukemia, characterized by a high incidence of tissue infiltration and organomegaly. Erythroleukemia is rare and is accompanied by dysplastic erythroid precursors. Acute lymphoblastic leukemia is typically seen in children and young adults. Acute megakaryocytic leukemia is rare, it is typically accompanied by myelofibrosis, and the blasts react with platelet-specific antibodies. The M3 variant of acute myelogenous leukemia (promyelocytic leukemia) has many promyelocytes filled with azurophilic granules, making them strongly peroxidase positive.

45 B The figure shows myeloblasts with prominent Auer rods. Along with the very high WBC count and the presence of peroxidase-positive blasts (myeloblasts) filling the marrow, these findings are characteristic of acute myelogenous leukemia (AML). This type of leukemia is most often seen in individuals 15 to 39 years old. Acute lymphoblastic leukemia occurs in children and young adults. Azurophilic, peroxidase-positive granules distinguish myeloblasts from lymphoblasts. Chronic lymphocytic leukemia is characterized by the presence of small, mature lymphocytes in the peripheral blood and bone marrow of older adults. Chronic myelogenous leukemia also is seen in adults, but this is a myeloproliferative process with a range of myeloid differentiation, and most of the myeloid cells are mature, with few blasts. Hodgkin lymphoma does not have a leukemic phase. Some cutaneous T-cell lymphomas may have circulating cells, known as Sézary cells.

46 C Myelodysplasia is characterized by a cellular marrow in which there are maturation defects in multiple lineages. This diagnosis is supported by the presence of ringed sideroblasts, megaloblasts, abnormal megakaryocytes, and myeloblasts in the marrow. Because these abnormal hematopoietic cells fail to mature normally, they are not released into the peripheral blood, leading to pancytopenia and susceptibility to infections. Myelodysplasias are clonal stem cell disorders that develop either de novo or after chemotherapy with alkylating agents, as in this case. The presence of chromosomal deletions, such as 5q, is a marker of posttherapy myelodysplasia. These morphologic abnormalities in the marrow are not seen in any of the other listed conditions.

47 B This is the Philadelphia chromosome, or Ph1, which is characteristic of patients with chronic myelogenous leukemia (CML). This karyotypic abnormality can be found using cytogenetic techniques, including fluorescence in situ hybridization (FISH). In the few cases that appear negative by karyotyping and by FISH, molecular analysis shows BCR-ABL rearrangements, and the tyrosine kinase activated via this fusion gene is the target of current therapy for CML. This rearrangement is considered a diagnostic criterion for CML. CML is a disease of pluripotent stem cells that affects all lineages, but the granulocytic precursors expand preferentially in the chronic phase. Acute promyelocytic leukemias often have the t(15;17) abnormality. Follicular lymphomas have a t(14;18) karyotypic abnormality involving the BCL2 gene. Hodgkin disease and myelomas usually do not have characteristic karyotypic abnormalities.

48 C Essential thrombocytosis is a myeloproliferative disorder. As with all myeloproliferative diseases, the transformation occurs in a myeloid stem cell. In this form of myeloproliferative disease, the dominant cell type affected is the megakaryocyte, and there is thrombocytosis. Other myeloproliferative disorders, such as chronic myelogenous leukemia, myelofibrosis, and polycythemia vera, also can be
accompanied by an increased platelet count. The diagnosis of essential thrombocytosis can be made after other causes of reactive thrombocytosis are excluded, and if the bone marrow examination shows increased megakaryocytes with no evidence of leukemia. The throbbing, burning pain in the extremities is caused by platelet aggregates that occlude small arterioles. The major manifestation of this disease is thrombotic or hemorrhagic crises. The swelling in this patient’s leg represents phlebothrombosis, followed by pulmonary embolism with infarction. The peripheral blood WBC count would be high in acute myelogenous leukemia, without thrombocytosis.

PBD9 619–620  BP9 447–448  PBD8 629–630  BP8 466

49 C The symptoms of polycythemia vera (PCV) result from an increased RBC mass with increased hematocrit and blood volume. Undetectable erythropoietin in the face of polycythemia is characteristic of PCV, a myeloproliferative disorder in which the neoplastic myeloid cells differentiate preferentially along the erythroid lineage. Other lineages also are affected, with leukocytosis and thrombocytosis in this case. These patients are Ph1 chromosome negative. Untreated, these patients die of episodes of bleeding or thrombosis—both related to disordered platelet function and the hemodynamic effects on distended blood vessels. Treatment by phlebotomy reduces the hematocrit. With this treatment, the disease in 15% to 20% of patients characteristically transforms into myelofibrosis with myeloid metaplasia. Termination in acute leukemia, in contrast to in chronic myeloid leukemia, is rare. When it occurs, it is an acute myeloid leukemia, not lymphoblastic leukemia.

PBD9 618–619  BP9 447–448  PBD8 628–629  BP8 465–466

50 B This classic history for chronic myelogenous leukemia (CML) is confirmed by the presence of different stages of myeloid differentiation in the blood and by the presence of the Philadelphia chromosome. He went into a remission but then entered a blast crisis involving B cells (CD19+). The fact that the B cells carry the original Ph1 chromosome and some additional abnormalities indicates that the B cells and the myeloid cells belong to the same original clone. The best explanation for this is that the initial transforming event affected a pluripotent stem cell, which differentiated along the myeloid lineage to produce a picture of CML. Analysis, even at this stage, indicates that the molecular counterpart of the Ph1 chromosome—the BCR-ABL rearrangement—affects all lineages, including B cells, T cells, and myeloid cells. With the evolution of the disease, additional mutations accumulate in the stem cells, which differentiate mainly along B lineages, giving rise to B-lymphoblastic leukemia; blast crisis also can affect myeloid cells, but they are not CD19+. A myelodysplastic syndrome, not present here, can precede the development of acute myelogenous leukemia. Hairy cell leukemia is an indolent disease without blasts. The Sézary syndrome has a leukemic component of CD4+ cells in addition to the skin involvement (mycosis fungoides).

PBD9 616–618  BP9 446–447  PBD8 627–628  BP8 464–466

51 E This patient has polycythemia vera (PCV), a myeloproliferative disorder characterized by an increased RBC mass, with hematocrit concentrations typically exceeding 60%. Although the increased RBC mass is responsible for most of the symptoms and signs, these patients also have thrombocytosis and granulocytosis. This occurs because, similar to other myeloproliferative disorders, PCV results from transformation of a multipotent stem cell. The high hematocrit concentration causes an increase in blood volume and distention of blood vessels. The neoplastic erythroid progenitors require extremely small amounts of erythropoietin for survival and proliferation; the levels of erythropoietin are virtually undetectable in PCV. When combined with abnormal platelet function, this condition predisposes the patient to bleeding. Abnormal platelet function also can predispose to thrombosis. The pruritus and peptic ulceration most likely are the result of the histamine release from basophils. In some patients, the disease “burns out” to myelofibrosis. A few patients “blast out” into acute myelogenous leukemia, and other patients (often with the BCR-ABL fusion gene) develop chronic myelogenous leukemia. Myelodysplastic syndromes and the other myeloproliferative disorders, such as essential thrombocytosis (with JAK2 mutations), are not accompanied by such a marked increase in RBC mass. Erythroleukemia typically is not accompanied by such a high hematocrit concentration because leukemic erythroid progenitors do not differentiate into mature RBCs.

PBD9 618–619  BP9 447–448  PBD8 628–629  BP8 465–466

52 A Myelofibrosis with myeloid metaplasia is a myeloproliferative disorder that is also a stem cell disorder in which neoplastic megakaryocytes secrete fibrogenic factors leading to marrow fibrosis. The neoplastic clone then shifts to the spleen, where it shows trilineage hematopoietic proliferation (extramedullary hematopoiesis), in which megakaryocytes are prominent. The marrow fibrosis and the extramedullary hematopoiesis in the spleen fail to regulate orderly release of leukocytes into the blood. The peripheral blood has immature RBC and WBC precursors (leukocythoblastic picture). Hematopoietic cell proliferation and turnover increases purine metabolism and uric acid production. Teardrop RBCs are misshapen RBCs that are seen when marrow undergoes fibrosis. Marrow injury also can be the result of other causes (e.g., metastatic tumors, irradiation). These causes also can give rise to a leukocythoblastic picture, but splenic enlargement with trilineage proliferation usually is not seen. The other causes mentioned—Hodgkin lymphoma, portal hypertension, and Histoplasma capsulatum infection—can cause splenic enlargement, but not marrow fibrosis. Metastases to the spleen are uncommon.

PBD9 620–621  BP9 448  PBD8 630–631  BP8 466–467

53 D Shown here are rodlike tubular Birbeck granules, with the characteristic periodicity seen in Langerhans cell proliferations. In this case, the skin eruptions, organomegaly, and lesion in the mastoid suggest infiltrates in multiple organs. The diagnosis is multifocal Langerhans cell histiocytosis, a disease most often seen in children. In half of these cases, exophthalmos occurs, and involvement of the hypothalamus and pituitary stalk leads to diabetes insipidus; these findings are called Hand-Schüller-Christian disease.
Acute lymphoblastic leukemia in children can involve the marrow, but does not produce skin or bone lesions. Tuberculosis can produce granulomatous disease with bony destruction, but the macrophages present in the granulomas are epithelioid macrophages that do not have Birbeck granules. Hodgkin lymphoma is seen in young adults and does not produce skin lesions or bone lesions. Myeloma is a disease of adults that can produce lytic bone lesions, but not skin lesions.

PBD9 621–622  BP9 449  PBD8 631  BP8 467–468

54 A Letterer-Siwe disease is a form of Langerhans cell histiocytosis, with Birbeck granules as a distinctive feature identified by electron microscopy, which are found in the cytoplasm of the Langerhans cells. Lymphoblasts that mark as T cells (CD3 positive) are seen in anterior mediastinal (thymic) masses in children with acute lymphoblastic leukemia/lymphoma. Plasma cells are seen in multiple myeloma, a disease of older adults accompanied by a monoclonal gamopathy. Reed-Sternberg cells are seen in Hodgkin lymphoma, which is an unlikely disease in children. Ringed sideroblasts can be seen in myelodysplastic syndromes. Sézary cells can be seen in peripheral T-cell lymphoma/leukemias, which often involve the skin.

PBD9 621–622  BP9 449  PBD8 631–632  BP8 467–468

55 G This patient has a splenic rupture with hematoma formation in a normal-sized spleen. Note the more darkly attenuated rim of blood adjacent to the normal-sized spleen and medial to the brightly attenuated ribs at the left in the CT image. The likelihood of acute alcohol ingestion favors trauma from falls, fights, and vehicular accidents. A "spontaneous" rupture of the spleen in the absence of trauma may occur when there is splenic enlargement from infections and neoplasms. Other causes of splenic enlargement include amyloid deposition, congestive splenomegaly from portal hypertension with cirrhosis, storage diseases such as Gaucher disease, and myeloproliferative disorders. Embolic events from endocarditis are most likely to occur with infective endocarditis and nonbacterial thrombotic endocarditis.

PBD9 625  BP9 456  PBD8 635  BP8 456

56 C The cytopenias along with the splenomegaly suggest hypersplenism as a cause for this patient’s anemia, leukopenia, and thrombocytopenia. The circulating blood cells are sequestered within the large spleen. One of the most common causes is congestive splenomegaly from portal hypertension resulting from hepatic cirrhosis. The elevation of AST more than ALT suggests alcoholic liver disease. The WBC count should be quite high with acute leukemias. Although infectious mononucleosis can lead to splenomegaly and can be marked by lymphocytosis, the course is typically not 9 months, and it is more common in younger individuals. Niemann-Pick disease is a storage disease from an inborn error of metabolism involving sphingomyelinase, and typically appears in childhood and leads to profound neurologic problems. The spleen is an uncommon location for metastases. Systemic lupus erythematosus can lead to cytopenias from reduced bone marrow function, but the spleen is not usually enlarged.

PBD9 623–624  BP9 456  PBD8 633–634  BP8 476

57 D The pale, tan-to-yellow, firm areas shown in the figure are infarcts. These lesions are either wedge-shaped and based on the capsule or are more irregularly shaped within the parenchyma. Emboli in the systemic arterial circulation most often arise in the heart, and the signs of acute infection in this case suggest emboli from vegetations on cardiac valves in a patient with infective endocarditis; these can lead to splenic infarction. Emboli exiting the aorta at the celiac axis generally take the straight route to the spleen. The kidneys and brain are other common sites for systemic emboli to lodge. Although acute myelogenous leukemia can cause enlargement of the spleen, there are typically no focal lesions—only uniform infiltration of the parenchyma—but the massive size of the spleen with chronic myelogenous leukemia predisposes to splenic infarcts. There would be scattered granulomas that are rounded and tan with granulomatous diseases of the spleen, such as histoplasmosis. In Hodgkin lymphoma, there can be focal nodules. Metastases can enlarge the spleen, but are uncommon in the spleen and are unlikely to be accompanied by signs of infection. Similarly, the congestive splenomegaly that occurs in cirrhosis and portal hypertension does not produce focal splenic lesions. In acute rheumatic fever, the verrucous vegetations are unlikely to embolize; in chronic rheumatic valvulitis, there is scarring with valve deformity, and this increases the risk of infective endocarditis.

PBD9 625  BP9 456  PBD8 634  BP8 446, 444

58 E Splenectomy in childhood reduces humoral immunity to encapsulated bacterial organisms, because splenic IgM producing B lymphocytes and splenic macrophages aid in removal of these infectious agents. The spleen is also a recycling center for RBCs, and it removes inclusions such as Howell-Jolly bodies (similar to getting the cherry pits out without damaging the cherry). About one third of all circulating platelets are pooled in the spleen, and granulocytes are marginated in splenic sinusoids, so that when the spleen is absent, the WBC and platelet counts are slightly increased. DiGeorge syndrome leads to cell-mediated immunodeficiency and increased viral, fungal, and parasitic diseases. Galactosemia results from an inborn error of metabolism, leading to liver disease and fibrosis that can cause splenomegaly. Gaucher disease leads to splenomegaly without significant immunodeficiency. Myeloproliferative disorders increase the size of the spleen. The thymus, but not the spleen, is sometimes involved in patients with Down syndrome (trisomy 21) who are immunodeficient.

PBD9 623  BP9 456  PBD8 189, 634  BP8 426

59 B These cells mark as cortical lymphocytes in the thymus of a child. An absence of such cells can be seen in DiGeorge syndrome with 22q11.2. Such patients also can have parathyroid hypoplasia and congenital heart disease.
Patients with Down syndrome (trisomy 21) can have thymic abnormalities and the T-cell dysregulation that predisposes to acute leukemia, but the thymus is typically present. The t(9;22) gives rise to the Philadelphia chromosome, which is characteristic of chronic myelogenous leukemia. The t(15;17) is seen in patients with acute promyelocytic leukemia. Individuals with fragile X syndrome usually have some form of mental retardation. Males with Klinefelter syndrome (XXY) do not have immunologic abnormalities.

Thymomas are rare neoplasms that can be benign or malignant. In one third to one half of cases, thymomas are associated with myasthenia gravis as an initial presentation (as in this case). Benign thymomas have a mixed population of lymphocytes and epithelial cells and are circumscribed, whereas malignant thymomas are invasive and have atypical cells. Thymic carcinomas resemble squamous cell carcinomas. Granulomas can have epithelioid macrophages and lymphocytes, but the thymus is an unusual location for them. Hodgkin lymphoma involves lymph nodes in the middle or posterior mediastinum, with a component of Reed-Sternberg cells. Lymphoblastic lymphoma of the T-cell variety is seen in the mediastinal region, including thymus, in children, but it has no epithelial component. Metastases to the thymus are quite unusual. An organizing abscess could have granulation tissue at its edge, with a mixture of inflammatory cell types, but not atypical cells.
1. A 77-year-old woman notices that small, pinpoint-to-blotchy areas of superficial hemorrhage have appeared on her gums and on the skin of her arms and legs over the past 3 weeks. On physical examination, she is afebrile and has no organomegaly. Laboratory studies show a normal prothrombin time and partial thromboplastin time. CBC shows hemoglobin of 12.7 g/dL, hematocrit of 37.2%, MCV of 80 μm³, platelet count of 276,000/mm³, and WBC count of 5600/mm³. Platelet function studies and fibrinogen level are normal, and no fibrin split products are detectable. Which of the following conditions best explains these findings?

A. Chronic renal failure  
B. Macronodular cirrhosis  
C. Meningococcemia  
D. Metastatic carcinoma  
E. Vitamin C deficiency

2. A healthy 19-year-old woman suffered blunt abdominal trauma in a motor vehicle accident. On admission to the hospital, her initial hematocrit was 33%, but over the next hour, it decreased to 28%. A paracentesis yielded serosanguineous fluid. She was taken to surgery, where a liver laceration was repaired, and 1 L of bloody fluid was removed from the peritoneal cavity. She remained stable. A CBC performed 3 days later is most likely to show which of the following morphologic findings in the peripheral blood?

A. Basophilic stippling of red cells  
B. Hypochromic red cells  
C. Leukoerythroblastosis  
D. Reticulocytosis  
E. Schistocytosis

3. A 65-year-old man has experienced worsening fatigue for the past 5 months. On physical examination, he is afebrile and has a pulse of 91/min, respirations of 18/min, and blood pressure of 105/60 mm Hg. There is no organomegaly. A stool sample is positive for occult blood. Laboratory findings include hemoglobin of 5.9 g/dL, hematocrit of 18.3%, MCV of 99 μm³, platelet count of 250,000/mm³, and WBC count of 7800/mm³. The reticulocyte concentration is 3.9%. No fibrin split products are detected, and direct and indirect Coombs test results are negative. A bone marrow biopsy specimen shows marked erythroid hyperplasia. Which of the following conditions best explains these findings?

A. Aplastic anemia  
B. Autoimmune hemolytic anemia  
C. Chronic blood loss  
D. Iron deficiency anemia  
E. Metastatic carcinoma

4. During the past 6 months, a 25-year-old woman has noticed a malar skin rash that is made worse by sun exposure. She also has had arthralgias and myalgias. On physical examination, she is afebrile and has a pulse of 100/min, respirations of 20/min, and blood pressure of 100/60 mm Hg. There is erythema of skin over the bridge of the nose. No organomegaly is noted. Laboratory findings include positive serologic test results for ANA and double-stranded DNA, hemoglobin of 8.1 g/dL, hematocrit of 24.4%, platelet count of 87,000/mm³, and WBC count of 3950/mm³. The peripheral blood smear shows nucleated RBCs. A dipstick urinalysis is positive for blood, but there are no WBCs, RBCs, or casts seen on microscopic examination of the urine. Which of the following laboratory findings is most likely to be present?

A. Decreased haptoglobin  
B. Decreased iron  
C. Decreased reticulocytosis  
D. Elevated D dimer  
E. Elevated hemoglobin F  
F. Elevated protoporphyrin
5 A 28-year-old woman has had a constant feeling of lethargy since childhood. On physical examination, she is afebrile and has a pulse of 80/min, respirations of 15/min, and blood pressure of 110/70 mm Hg. The spleen tip is palpable, but there is no abdominal pain or tenderness. Laboratory studies show hemoglobin of 11.7 g/dL, platelet count of 159,000/mm³, and WBC count of 5390/mm³. The peripheral blood smear shows small round erythrocytes that lack a zone of central pallor. An inherited abnormality in which of the following RBC components best accounts for these findings?

A α-Globin chain  
B β-Globin chain  
C Carbonic anhydrase  
D Glucose-6-phosphate dehydrogenase  
E Heme with porphyrin ring  
F Spectrin cytoskeletal protein

6 A 13-year-old boy has the sudden onset of severe abdominal pain and cramping accompanied by chest pain, non-productive cough, and fever. On physical examination, his temperature is 39°C, pulse is 110/min, respirations are 22/min, and blood pressure is 80/50 mm Hg. He has diffuse abdominal tenderness, but no masses or organomegaly. Laboratory studies show a hematocrit of 18%. The peripheral blood smear is shown in the figure. A chest radiograph shows bilateral pulmonary infiltrates. Which of the following is the most likely mechanism for initiation of his pulmonary problems?

A Chronic hypoxia of the pulmonary parenchyma  
B Defects in the alternative pathway of complement activation  
C Extensive RBC adhesion to endothelium  
D Formation of autoantibodies to alveolar basement membrane  
E Intravascular antibody-induced hemolysis

8 A clinical study of patients who inherit mutations that reduce the level of ankyrin, the principal binding site for spectrin, in the RBC membrane cytoskeleton shows an increased prevalence of chronic anemia with splenomegaly. For many patients, it is observed that splenectomy reduces the severity of anemia. This beneficial effect of splenectomy is most likely related to which of the following processes?

A Decrease in opsonization of RBCs and lysis in spleen  
B Decrease in production of reactive oxygen species by splenic macrophages  
C Decrease in splenic RBC sequestration and lysis  
D Increase in deformability of RBCs within splenic sinusoids  
E Increase in splenic storage of iron

9 An 18-year-old woman from Copenhagen, Denmark, has had malaise and a low-grade fever for the past week, along with arthralgias. On physical examination, she appears very pale, except for a bright red malar facial rash. She has a history of chronic anemia, and spherocytes are observed on a peripheral blood smear. Her hematocrit, which normally ranges from 35% to 38%, is now 28%, and the reticulocyte count is very low. The serum bilirubin level is 0.9 mg/dL. Which of the following events is most likely to have occurred in this patient?

A Accelerated extravascular hemolysis in the spleen  
B Development of anti-RBC antibodies  
C Disseminated intravascular coagulation  
D Reduced erythropoiesis from parvovirus infection  
E Superimposed dietary iron deficiency

10 A 10-year-old child has experienced multiple episodes of pneumonia and meningitis with septicemia since infancy. Causative organisms include Streptococcus pneumoniae and Haemophilus influenzae. On physical examination, the child has no organomegaly and no deformities. Laboratory studies show hemoglobin of 9.2 g/dL, hematocrit of 27.8%, platelet count of 372,000/mm³, and WBC count of 10,300/mm³. A hemoglobin electrophoresis shows 1% hemoglobin A₂, 7% hemoglobin F, and 92% hemoglobin S. Which of the following is the most likely cause of the repeated infections in this child?

A Absent endothelial cell expression of adhesion molecules  
B Diminished hepatic synthesis of complement proteins  
C Impaired neutrophil production  
D Loss of normal splenic function  
E Reduced synthesis of immunoglobulins
11 A 32-year-old woman from Hanoi, Vietnam, gives birth at 34 weeks’ gestation to a markedly hydropic stillborn male infant. Autopsy findings include hepatosplenomegaly and cardiomegaly, serous effusions in all body cavities, and generalized hydrops. No congenital anomalies are noted. There is marked extramedullary hematopoiesis in visceral organs. Which of the following hemoglobins is most likely predominant on hemoglobin electrophoresis of the fetal RBCs?

A Hemoglobin A1  
B Hemoglobin A2  
C Hemoglobin Bart’s  
D Hemoglobin E  
E Hemoglobin F  
F Hemoglobin H

12 A 17-year-old girl has had a history of fatigue and weakness for her entire life. She has not undergone puberty. On physical examination, secondary sex characteristics are not well developed. She has hepatosplenomegaly. CBC shows hemoglobin of 9.1 g/dL, hematocrit of 26.7%, MCV of 66 μm³, platelet count of 89,000/mm³, and WBC count of 3670/mm³. The appearance of the peripheral blood smear is shown in the figure. Additional laboratory findings include serum glucose of 144 mg/dL, TSH of 6.2 mU/mL, and ferritin of 679 ng/mL. A mutation in a gene encoding for which of the following is most likely to be present in this girl?

A Ankyrin  
B β-Globin  
C G6PD  
D HFE  
E NADPH oxidase

13 A 12-year-old boy has a history of episodes of severe abdominal, chest, and back pain since early childhood. On physical examination, he is afebrile, and there is no organomegaly. Laboratory studies show hemoglobin of 11.2 g/dL, platelet count of 194,000/mm³, and WBC count of 9020/mm³. The peripheral blood smear shows occasional sickled cells, nucleated RBCs, and Howell-Jolly bodies. Hemoglobin electrophoresis shows 1% hemoglobin A₂, 6% hemoglobin F, and 93% hemoglobin S. Hydroxyurea therapy is found to be beneficial in this patient. An increase in which of the following is the most likely basis for its therapeutic efficacy?

A Erythrocyte production  
B Overall globin chain synthesis  
C Oxygen affinity of hemoglobin  
D Production of hemoglobin A  
E Production of hemoglobin F

14 A 25-year-old woman has a 3-year history of arthralgias. Physical examination shows no joint deformity, but she appears pale. Laboratory studies show total RBC count of 4.7 million/mm³, hemoglobin of 12.5 g/dL, hematocrit of 37.1%, platelet count of 217,000/mm³, and WBC count of 5890/mm³. The peripheral blood smear shows hypochromic and microcytic RBCs. Total serum iron and ferritin levels are normal. Hemoglobin electrophoresis shows 93% hemoglobin A₁ with elevated hemoglobin A₂ level of 5.8% and hemoglobin F level of 1.2%. What is the most likely diagnosis?

A Anemia of chronic disease  
B Autoimmune hemolytic anemia  
C β-Thalassemia minor  
D Infection with Plasmodium vivax  
E Iron deficiency anemia

15 A 23-year-old African-American man passes dark reddish brown urine 3 days after taking an anti-inflammatory medication that includes phenacetin. He is surprised, because he has been healthy all his life and has had no major illnesses. On physical examination, he is afebrile, and there are no remarkable findings. CBC shows a mild normocytic anemia, but the peripheral blood smear shows precipitates of denatured globin (Heinz bodies) with supravital staining and scattered “bite cells” in the population of RBCs. Which of the following is the most likely diagnosis?

A α-Thalassemia minor  
B β-Thalassemia minor  
C Glucose-6-phosphate dehydrogenase deficiency  
D Sickle cell trait  
E Abnormal ankyrin in RBC cytoskeletal membrane  
F Warm antibody autoimmune hemolytic anemia

16 Since childhood, a 30-year-old man has been easily fatigued with minimal exercise. Laboratory studies show hypochromic microcytic anemia. Hemoglobin electrophoresis reveals decreased Hgb A₁ with increased Hgb A₂ and Hgb F. His serum ferritin is markedly increased. Which of the following mutations is most likely to be present in the β-globin gene of this man?

A New stop codon  
B Single base insertion, with frameshift  
C Splice site  
D Three-base deletion  
E Trinucleotide repeat
17 A 16-year-old boy notes passage of dark urine. He has a history of multiple bacterial infections and venous thromboses for the past 10 years, including portal vein thrombosis in the previous year. On physical examination, his right leg is swollen and tender. CBC shows hemoglobin, 9.8 g/dL; hematocrit, 29.9%; MCV, 92 μm³; platelet count, 150,000/mm³; and WBC count, 3800/mm³ with 24% segmented neutrophils, 1% bands, 64% lymphocytes, 10% monocytes, and 1% eosinophils. He has a reticulocytosis, and his serum haptoglobin level is very low. A mutation affecting which of the following gene products is most likely to give rise to this clinical condition?
A β-Globin chain
B Factor V
C Glucose-6-phosphate dehydrogenase
D Phosphatidylglycerol (PGI) A (PGI)
E Prothrombin G20210A
F Spectrin

18 A 30-year-old, previously healthy man from Lagos, Nigeria, passes dark brown urine 2 days after starting the prophylactic antimalarial drug primaquine. On physical examination, he appears pale and is afebrile. There is no organomegaly. Laboratory studies show that his serum haptoglobin level is decreased. Which of the following is the most likely explanation of these findings?
A Antibody-mediated hemolysis
B Impaired DNA synthesis
C Impaired globin chain synthesis
D Increased susceptibility to complement-induced lysis
E Mechanical fragmentation of RBCs as a result of vascular narrowing
F Oxidative injury to hemoglobin
G Reduced deformability of RBC membrane

19 A 34-year-old woman reports becoming increasingly tired for the past 5 months. On physical examination, she is afebrile and has mild splenomegaly. Laboratory studies show a hemoglobin concentration of 10.7 g/dL and hematocrit of 32.3%. The peripheral blood smear shows spherocytes and rare nucleated RBCs. Direct and indirect Coombs test results are positive at 37° C, although not at 4° C. Which of the following underlying diseases is most likely to be diagnosed in this patient?
A Escherichia coli sepsis
B Hereditary spherocytosis
C Infectious mononucleosis
D Mycoplasma pneumoniae infection
E Systemic lupus erythematosus

20 A 22-year-old woman has experienced malaise and a sore throat for 2 weeks. Her fingers turn white on exposure to cold. On physical examination, she has a temperature of 37.8° C, and the pharynx is erythematous. Laboratory findings include a positive monospot (heterophile antibody) test result. Direct and indirect Coombs test results are positive at 4° C, although not at 37° C. Which of the following molecules bound on the surfaces of the RBCs most likely accounts for these findings?
A δ2-Macroglobulin
B Complement C3b
C Fibronectin
D Histamine
E IgE

21 A 65-year-old man diagnosed with follicular non-Hodgkin lymphoma is treated with chemotherapy. He develops fever and cough of a week’s duration. On examination, there are bilateral pulmonary rales. A chest radiograph shows diffuse interstitial infiltrates. A sputum specimen is positive for cytomegalovirus. He develops scleral icterus and Raynaud phenomenon. Laboratory studies show hemoglobin, 10.3 g/dL; hematocrit, 41.3%; MCV, 101 μm³; platelet count, 7600/mm³; and platelet count, 205,000/mm³. His serum total bilirubin is 6 mg/dL, direct bilirubin is 0.8 mg/dL, and LDH is 1020 U/L. Coombs test is positive. Which of the following is the most likely mechanism for his anemia?
A Marrow aplasia caused by chemotherapy
B Vitamin K deficiency caused by cytomegalovirus hepatitis
C Megaloblastic anemia caused by folate deficiency
D Extravascular hemolysis caused by cold agglutinins
E Iron deficiency caused by metastases to colon

22 A 29-year-old woman has had fatigue with dizziness for the past 5 months. On physical examination, she has an erythematous malar rash. She has no lymphadenopathy, but there is a palpable spleen tip. She is afebrile. Laboratory studies show hemoglobin, 8.9 g/dL; hematocrit, 27.8%; MCV, 103 μm³; RBC distribution width index, 22; WBC count, 8650/mm³; platelet count, 222,000/mm³; and reticulocyte count, 3.3%. The peripheral blood smear shows polychromasia, but no schistocytes. Her serum total bilirubin is 3.2 mg/dL with direct bilirubin 0.8 mg/dL, and haptoglobin is 5 mg/dL. Antinuclear antibody and anti-double-stranded DNA tests are positive. What additional laboratory test finding is she most likely to have?
A D-dimer 10 μg/mL
B Increased RBC osmotic fragility
C Positive Coombs test
D Serum cobalamin (vitamin B12) 50 pg/mL
E Serum ferritin 240 ng/mL

23 A 29-year-old rugby player takes part in a particularly contentious game between New Zealand and South Africa. He is the forward prop in the scrums, hitting hard and being hit hard by other players. He feels better after downing several pints of beer following the game, but notes darker urine. Urinalysis is positive for blood. Which of the following pathogenic mechanisms underlies change in the color of urine?
A Complement lysis
B Intravascular disruption
C Osmotic fragility
D Sinusoidal sickling
E Splenic sequestration

24 In an epidemiologic study of anemias, the findings show that there is an increased prevalence of anemia in individuals of West African ancestry. By hemoglobin electrophoresis, some individuals within this region have increased hemoglobin S levels. The same regions also have a high prevalence of an infectious disease. Which of the following infectious agents is most likely to be endemic in the region where such anemia shows increased prevalence?
A Borrelia burgdorferi
B Clostridium perfringens
C Cryptococcus neoformans
D Plasmodium falciparum
E Treponema pallidum
F Trypanosoma brucei
An infant is born at 34 weeks' gestation to a 28-year-old woman, G3, P2. At birth, the infant is observed to be markedly hydropic and icteric. A cord blood sample is taken, and direct Coombs test result is positive for the infant's RBCs. Which of the following is the most likely mechanism for the findings in this infant?

A. Hemolysis of antibody-coated cells
B. Hematopoietic stem cell defect
C. Impaired globin synthesis
D. Mechanical fragmentation of RBCs
E. Oxidative injury to hemoglobin
F. Reduced deformability of RBC membranes

A 22-year-old woman after returning from a trip to Africa has experienced febrile episodes over the past 2 weeks. On physical examination, her temperature is 37.5°C, pulse is 82/min, respirations are 18/min, and blood pressure is 105/65 mm Hg. Laboratory studies show hemoglobin of 10.8 g/dL, hematocrit of 32.5%, platelet count of 245,700/mm³, and WBC count of 8320/mm³. The serum haptoglobin level is decreased, and direct and indirect Coombs test results are negative. The reticulocyte count is increased. The prothrombin time is 12 seconds, and the partial thromboplastin time is 31 seconds. She is observed over the next week and found to have temperature spikes to 39.1°C, with shaking chills every 48 hours. Infection with which of the following organisms is most likely to cause this patient's illness?

A. Aspergillus niger
B. Babesia microti
C. Dirofilaria immitis
D. Escherichia coli
E. Plasmodium vivax
F. Wuchereria bancrofti

A 33-year-old previously healthy man with persistent fever and heart murmur is diagnosed with infective endocarditis. He receives a high dosage of a cephalosporin antibiotic during the next 10 days. He now has increasing fatigue. On physical examination he has tachycardia and scleral icterus. Laboratory studies show hemoglobin of 7.5 g/dL, hematocrit, 28.8%; MCV, 101 μm³; platelet count, 145,000/mm³; WBC count, 6920/mm³; Na⁺, 146 mmol/L; K⁺, 5.5 mmol/L; Cl⁻, 106 mmol/L; CO₂, 26 mmol/L; creatinine, 2.3 mg/dL; urea nitrogen, 22 mg/dL; LDH, 1095 U/L; and amylase, 45 U/L. The peripheral blood smear is shown in the figure. What infectious agent is most likely to produce these findings?

A. Babesia microti
B. Borrelia burgdorferi
C. Leishmania donovani
D. Plasmodium falciparum
E. Trypanosoma brucei
F. Wuchereria bancrofti

A 7-year-old child has had worsening headaches and is obtunded for the past 2 days. Physical examination shows temperature of 39.5°C, pulse of 103/min, respirations of 18/min, and blood pressure of 90/55 mm Hg. There is bilateral papilledema on funduscopic examination. No focal neurologic deficits are noted. Palpation of the abdomen reveals hepatosplenomegaly. Laboratory findings show hemoglobin, 9.5 g/dL; hematocrit, 28.8%; MCV, 101 μm³; platelet count, 145,000/mm³; WBC count, 6920/mm³; Na⁺, 146 mmol/L; K⁺, 5.5 mmol/L; Cl⁻, 106 mmol/L; CO₂, 26 mmol/L; creatinine, 2.3 mg/dL; urea nitrogen, 22 mg/dL; LDH, 1095 U/L; and amylase, 45 U/L. The peripheral blood smear is shown in the figure. What infectious agent is most likely to produce these findings?

A. Babesia microti
B. Borrelia burgdorferi
C. Leishmania donovani
D. Plasmodium falciparum
E. Trypanosoma brucei
F. Wuchereria bancrofti

A 54-year-old, previously healthy man has experienced minor fatigue on exertion for the past 9 months. On physical examination, there are no remarkable findings. Laboratory studies show hemoglobin of 11.7 g/dL, hematocrit of 34.8%, MCV of 73 μm³, platelet count of 315,000/mm³, and total WBC count of 8035/mm³. Which of the following is the most likely cause for his anemia?

A. Dietary nutrient deficiency
B. Disseminated intravascular coagulopathy
C. Immune-mediated hemolysis
D. Infection with parvovirus
E. Inherited hemoglobinopathy
F. RBC cytoskeletal protein disorder

A. Bone marrow biopsy
B. Hemoglobin electrophoresis
C. Serum ferritin
D. Serum haptoglobin
E. Serum iron
F. Serum transferrin
30 A 73-year-old man takes no medications and has had no prior major illnesses or surgeries. For the past year, he has become increasingly tired and listless. Physical examination shows that he appears pale but has no hepatosplenomegaly and no deformities. CBC shows hemoglobin, 9.7 g/dL; hematocrit, 32.4%; MCV, 87 μm³; RBC count, 4.28 million/mm³; platelet count, 240,000/mm³; and WBC count, 6800/mm³. His peripheral blood smear is shown in the figure. Which of the following is the most likely diagnosis?

A Autoimmune hemolytic anemia  
B Chronic alcohol abuse  
C β-Thalassemia major  
D Hemophilia A  
E Occult malignancy  
F Vitamin B₁₂ deficiency

31 A clinical study is performed using adult patients diagnosed with peptic ulcer disease, chronic blood loss, and hypochromic microcytic anemia. Their serum ferritin levels average 5 to 7 ng/mL. The rate of duodenal iron absorption in this study group is found to be much higher than in a normal control group. After treatment with omeprazole and clarithromycin, study group patients have hematocrits of 40% to 42%, MCV of 82 to 85 μm³, and serum ferritin of 30 to 35 ng/mL. Measured rates of iron absorption in the study group after therapy are now decreased to the range of the normal controls. Which of the following substances derived from liver is most likely underlying condition causing this patient’s findings?

A Divalent metal transporter-1 (DMT-1)  
B Hemosiderin  
C Hepcidin  
D HLA-like transmembrane protein  
E Transferrin

32 A 39-year-old man has experienced chronic fatigue and weight loss for the past 3 months. There are no remarkable findings on physical examination. Laboratory studies show hemoglobin, 10.0 g/dL; hematocrit, 30.3%; MCV, 91 μm³; platelet count, 240,000/mm³; WBC count, 7550/mm³; serum iron 80 μg/dL; total iron-binding capacity, 145 μg/dL; and serum ferritin, 565 ng/mL. Serum erythropoietin levels are low for the level of Hb and hepcidin levels are elevated. Which of the following is the most likely diagnosis?

A Anemia of chronic disease  
B Aplastic anemia

33 A 45-year-old woman has experienced worsening arthritis of her hands and feet for the past 15 years. On physical examination, there are marked deformities of the hands and feet, with ulnar deviation of the hands and swan-neck deformities of the fingers. Laboratory studies show an elevated level of rheumatoid factor. CBC shows hemoglobin, 11.6 g/dL; hematocrit, 34.8%; MCV, 87 μm³; platelet count, 268,000/mm³; and WBC count, 6800/mm³. There is a normal serum haptoglobin level, serum iron concentration of 20 μg/dL, total iron-binding capacity of 195 g/dL, percent saturation of 10.2, and serum ferritin concentration of 317 ng/mL. No fibrin split products are detected. The reticulocyte concentration is 1.1%. What is the most likely mechanism underlying this patient’s hematologic abnormalities?

A Autoantibodies against RBC membranes  
B Impaired synthesis of β-globin chains  
C Inadequate usage of stored iron  
D Mutation in the phosphatidylinositol glycan A (PIGA) gene  
E Sequestration of RBCs in splenic sinusoids  
F Space-occupying lesions in the bone marrow

34 A 62-year-old man is taken to the emergency department in a state of inebriation. He is well known there because this scenario has been repeated many times over 15 years. On physical examination, he is afebrile and normotensive. The spleen edge is firm. Laboratory studies show hemoglobin of 8.2 g/dL, hematocrit of 25.1%, MCV of 107 μm³, platelet count of 135,000/mm³, and WBC count of 3920/mm³. The peripheral blood smear shows prominent anisocytosis and macrocytosis. Polychromatophilic RBCs are difficult to find. A few of the neutrophils show six to seven nuclear lobes. Which of the following is the most likely explanation of these findings in this peripheral blood cells?

A Diminished nuclear maturation from impaired DNA synthesis  
B Extravascular hemolysis of antibody-coated cells  
C Imbalance in synthesis of α-globin and β-globin chains  
D Increased susceptibility to lysis by complement  
E Reduced deformability of RBC membranes

35 An 83-year-old man complains of worsening malaise and fatigue over the past 5 months. On physical examination, he is afebrile and normotensive. The spleen tip is palpable. A CBC shows hemoglobin, 10.6 g/dL; hematocrit, 29.8%; MCV, 92 μm³; platelet count, 95,000/mm³; and WBC count, 4900/mm³ with 63% segmented neutrophils, 7% bands, 2% metamyelocytes, 1% myelocytes, 22% lymphocytes, 5% monocytes, and 3 nucleated RBCs per 100 WBCs. The peripheral blood smear shows occasional teardrop cells. An examination of the bone marrow biopsy specimen and smear is most likely to show which of the following findings?

A Erythroid hyperplasia  
B Extensive fibrosis  
C Fatty replacement  
D Many megaloblasts  
E Numerous myeloblasts
A clinical study is performed to assess outcomes in patients who have macrocytic anemias as a result of Vitamin 12 or folate deficiency. A comparison of laboratory testing strategies shows that the best strategy includes testing for serum homocysteine, methylmalonic acid, vitamin B12 (cobalamin), and folate. What is the most important reason for ordering these tests simultaneously?

A. Aplastic anemia can result from lack of either nutrient
B. Both nutrients are absorbed similarly
C. Neurologic injury must be prevented
D. Life-threatening thrombocytopenia can occur in both
E. Therapy for one deficiency also treats the other

A 37-year-old woman has experienced abdominal pain and intermittent low-volume diarrhea for the past 3 months. On physical examination, she is afebrile. A stool sample is positive for occult blood. A colonoscopy is performed, and biopsy specimens from the terminal ileum and colon show microscopic findings consistent with Crohn disease. She does not respond to medical therapy, and part of the colon and terminal ileum are removed. She is transfused with 2 U of packed RBCs during surgery. Three weeks later, she appears healthy, but complains of easy fatigability. On investigation, CBC findings show hemoglobin of 10.6 g/dL, hematocrit of 31.6%, RBC count of 2.69 million/μL, MCV of 118 μm³, platelet count of 140,000/mm³, and WBC count of 1400/mm³ with 20% segmented neutrophils, 1% bands, 66% lymphocytes, and 13% monocytes. The reticulocyte count is 0.1%. Which of the following historical findings would be most useful in determining the cause of her condition?

A. Dietary habits
B. Exposure to medications
C. Family history of anemias
D. Menstrual history
E. Recent bacterial infection

A 28-year-old, previously healthy man has noted increasing fatigue for the past 6 months and formation of bruises after minimal trauma. Over the past 2 days, he has developed a cough. On physical examination, his temperature is 38.9° C, and he has diffuse rales in both lungs. He has no hepatosplenomegaly and no lymphadenopathy. Laboratory findings in-clude a sputum culture positive for Streptococcus pneumoniae, hemoglobin of 7.2 g/dL, hematocrit of 21.7%, platelet count of 23,400/mm³, WBC count of 1310/mm³, prothrombin time of 13 seconds, partial thromboplastin time of 28 seconds, and total bilirubin of 1 mg/dL. The ANA test result is negative. What is the most likely explanation of these findings?

A. Hematopoietic stem cell defect
B. Hemolysis of antibody-coated cells
C. Increased susceptibility to lysis by complement
D. Metastatic adenocarcinoma to bone marrow
E. Secondary hypersplenism

In a study of idiopathic aplastic anemia, patients are found who have premature senescence of hematopoietic stem cells. Their hematopoietic cells have normal morphology, but there are fewer cells in myeloid, erythroid, and megakaryocytic cell lines. Which of the following enzymes is most likely deficient in their marrow stem cells?

A. Alkaline phosphatase
B. Metalloproteinase
C. Pyruvate kinase
D. Telomerase
E. Tyrosine kinase

A 44-year-old woman has a 2-week history of multiple ecchymoses on her extremities after only minor trauma. She also reports feeling extremely weak. Over the previous 24 hours, she has developed a severe cough productive of yellowish sputum. On physical examination, her temperature is 38.4° C, and she has diffuse crackles on all lung fields. Laboratory studies show hemoglobin, 7.2 g/dL; hematocrit, 21.4%; MCV, 88 μm³; platelet count, 35,000/mm³; and WBC count, 1400/mm³ with 20% segmented neutrophils, 1% bands, 66% lymphocytes, and 13% monocytes. The reticulocyte count is 0.1%. Which of the following historical findings would be most useful in determining the cause of her condition?

A. Recent bacterial infection
B. Menstrual history
C. Family history of anemias
D. Exposure to medications
E. Dietary habits
43 Soon after crossing the finish line in a 10-km race, a 31-year-old man collapses. On physical examination, his temperature is 40.1°C, pulse is 101/min, respirations are 22/min, and blood pressure is 85/50 mm Hg. He is not perspiring, and his skin shows decreased turgor. Laboratory studies show Na⁺, 155 mmol/L; K⁺, 4.6 mmol/L; Cl⁻, 106 mmol/L; CO₂ 27 mmol/L; glucose, 68 mg/dL; creatinine, 1.8 mg/dL; hemoglobin, 20.1 g/dL; hematocrit, 60.3%; platelet count, 230,400/mm³; and WBC count, 6830/mm³. What is the most likely diagnosis?

A  Erythroleukemia  B  Chronic obstructive pulmonary disease  C  Diabetes insipidus  D  Hemoconcentration  E  Paraneoplastic syndrome  F  Polycythemia vera

44 A 33-year-old woman, G3, P0, who has had two spontaneous abortions, is in the second trimester of her third pregnancy. An ultrasound at 18 weeks' gestation revealed symmetric growth retardation. She gives birth to a stillborn fetus at 25 weeks, and experiences sudden onset of dyspnea. A pulmonary ventilation/perfusion scan indicates a high probability of thromboembolism. Four months later, she experiences an altered state of consciousness and sudden loss of movement in the right arm. A cerebral angiogram shows occlusion of a branch of the left middle cerebral artery. Laboratory findings show hemoglobin, 13.4 g/dL; hematocrit, 40.3%; MCV, 91 μm³; platelet count, 124,000/mm³; WBC count, 5530/mm³; prothrombin time, 13 seconds; partial thromboplastin time, 46 seconds; positive antiphospholipid antibody; positive serologic test result for syphilis; and negative ANA. Which of the following best explains these findings?

A  Antiphospholipid syndrome  B  Myeloproliferative disorder  C  Thrombophlebitis  D  Treponema pallidum infection  E  Trousseau syndrome  F  Von Willebrand disease

45 A 23-year-old woman in her 25th week of pregnancy has felt no fetal movement for the past 3 days. Three weeks later, she still has not given birth and suddenly develops dyspnea with cyanosis. On physical examination, her temperature is 37°C, pulse is 106/min, respirations are 23/min, and blood pressure is 80/40 mm Hg. She has large ecchymoses over the skin of her entire body. A stool sample is positive for occult blood. Laboratory studies show an elevated prothrombin time and partial thromboplastin time. The platelet count is decreased, plasma fibrinogen is markedly decreased, and fibrin split products are detected. A blood culture is negative. Which of the following is the most likely cause of her bleeding diathesis?

A  Consumption of coagulation factors  B  Defects in platelet aggregation  C  Increased vascular fragility  D  Reduced production of platelets  E  Toxic injury to the endothelium

46 A 30-year-old man has had pain and burning on urination for the past week. On physical examination, he is febrile and has a pulse of 92/min, respirations of 18/min, and blood pressure of 80/45 mm Hg. Digital rectal examination indicates that he has an enlarged, tender prostate. There is costovertebral angle tenderness on the right. Scattered ecchymoses are present over the trunk and extremities. Laboratory studies show a blood culture positive for Klebsiella pneumoniae. The appearance of the RBCs in a peripheral blood smear is shown in the figure. Which of the following hematologic disorders is he most likely to have?

A  Autoimmune hemolytic anemia  B  Hereditary spherocytosis  C  Iron deficiency anemia  D  Megaloblastic anemia  E  Microangiopathic hemolytic anemia

47 A 37-year-old woman has noted an excessively heavy menstrual flow each of the past 6 months. She also has noticed increasing numbers of pinpoint hemorrhages on her lower extremities in the past month. Physical examination shows no organomegaly or lymphadenopathy. CBC shows hemoglobin of 14.2 g/dL, hematocrit of 42.5%, MCV of 91 μm³, platelet count of 15,000/mm³, and WBC count of 6950/mm³. On admission to the hospital, she has melena and a transfusion of platelets, her platelet count does not increase. Which of the following describes the most likely basis for her bleeding tendency?

A  Abnormal production of platelets by megakaryocytes  B  Defective platelet-endothelial interactions  C  Destruction of antibody-coated platelets by the spleen  D  Excessive loss of platelets in menstrual blood  E  Suppression of pluripotent stem cell division

48 A 9-year-old boy has developed prominent bruises on his extremities over the past week. On physical examination, he has ecchymoses and petechiae on his arms and legs. Laboratory studies show hemoglobin, 13.8 g/dL; hematocrit, 41.9%; MCV, 93 μm³; platelet count, 11,300/mm³; and WBC count, 7720/mm³. He had respiratory syncytial virus pneumonia 3 weeks ago. His condition improves with corticosteroid therapy. Which of the following abnormalities is most likely to cause his hemorrhagic diathesis?

A  Antiplatelet antibodies  B  Bone marrow aplasia  C  Glycoprotein IIb/IIIa dysfunction  D  Vitamin C deficiency  E  Von Willebrand factor metalloproteinase deficiency
49 A 21-year-old woman known to have a protein C deficiency develops recurrent pulmonary thromboembolism and is placed on anticoagulant therapy. Two weeks after initiation of this therapy, she has a sudden change in mental status and experiences difficulty speaking and swallowing. A cerebral angiogram shows a left middle cerebral artery occlusion. Laboratory studies show hemoglobin of 13 g/dL, platelet count of 65,400/mm³, WBC count of 5924/mm³, prothrombin time of 12 seconds, and partial thromboplastin time of 51 seconds. The anticoagulant therapy is discontinued. Which of the following pharmacologic agents used as an anticoagulant in this patient is most likely to have caused these findings?

- A  Acetylsalicylic acid (aspirin)
- B  Heparin
- C  Tissue plasminogen activator
- D  Urokinase
- E  Warfarin

50 A 56-year-old woman suffers the sudden onset of headache and photophobia, and her condition worsens for the next 2 days. On physical examination, she has a temperature of 38°C and is disoriented. CBC shows hemoglobin of 11.2 g/dL, hematocrit of 33.7%, MCV of 94 μm³, platelet count of 32,000/mm³, and WBC count of 9900/mm³. The peripheral blood smear shows schistocytes. The serum urea nitrogen level is 38 mg/dL, and the creatinine level is 3.9 mg/dL. Which of the following is the most likely diagnosis?

- A  Autoimmune hemolytic anemia
- B  β-Thalassemia major
- C  Disseminated intravascular coagulation
- D  Idiopathic thrombocytopenic purpura
- E  Paroxysmal nocturnal hemoglobinuria
- F  Thrombotic thrombocytopenic purpura

51 A 44-year-old woman has experienced malaise with nausea and vomiting for 3 months. On physical examination, she has scleral icterus and a yellowish hue to her skin. She has difficulty remembering three objects after 3 minutes. There are no neurologic deficits. Laboratory studies show a positive serologic test result for hepatitis C, a serum ALT of 310 U/L, AST of 269 U/L, total bilirubin of 7.6 mg/dL, direct bilirubin of 5.8 mg/dL, alkaline phosphatase of 75 U/L, and ammonia of 55 μmol/L. An abnormal result of which of the following laboratory studies of hemostatic function is most likely to be reported?

- A  Fibrin split products
- B  Immunoassay for plasma von Willebrand factor
- C  Platelet aggregation
- D  Platelet count
- E  Prothrombin time

52 A 23-year-old woman has a history of easy bruising. Physical examination shows multiple bruises ranging in color from red to blue to purple on her arms and legs. There is no organomegaly, and no deformities are noted. Laboratory studies show hemoglobin, 9.5 g/dL; hematocrit, 28.2%; platelet count, 229,300/mm³; WBC count, 7185/mm³; prothrombin time, 12 seconds; and partial thromboplastin time, 38 seconds. A 1:1 dilution of the patient’s plasma with normal pooled plasma corrects the partial thromboplastin time. Ristocetin-dependent platelet aggregation in patient plasma is markedly reduced. Factor VIII activity is 30% (reference range 50% to 150%). Which of the following is the most likely potential consequence of this disease?

- A  Bone marrow failure from aplasia
- B  Excessive bleeding after oral surgery
- C  Increasing difficulty with joint mobility
- D  Myeloproliferative disorder
- E  Recurrent deep venous thrombosis

53 A 42-year-old woman has had nosebleeds, easy bruising, and increased bleeding with her menstrual periods for the past 4 months. On physical examination, her temperature is 37°C, pulse is 88/min, and blood pressure is 90/60 mm Hg. She has scattered petechiae over the distal extremities. There is no organomegaly. Laboratory studies show hemoglobin of 12.3 g/dL, hematocrit of 37%, platelet count of 21,500/mm³, and WBC count of 7730/mm³. A bone marrow biopsy specimen shows a marked increase in megakaryocytes. The prothrombin and partial thromboplastin times are within the reference range. What is the most likely diagnosis?

- A  Disseminated intravascular coagulation
- B  Hemophilia B
- C  Immune thrombocytopenic purpura
- D  Metastatic breast carcinoma
- E  Thrombotic thrombocytopenic purpura
- F  Vitamin K deficiency
- G  Von Willebrand disease

54 A clinical study is performed involving adult patients diagnosed with microangiopathic hemolytic anemia. A subgroup of patients who had fever or diarrhea preceding the initial diagnosis of anemia were excluded. The patients had schistocytes present on peripheral blood smears. Some of these patients were found to have a deficiency of a metalloproteinase known as ADAMTS13. Which of the following conditions were the patients with this deficiency most likely to have?

- A  Disseminated intravascular coagulation (DIC)
- B  Hemolytic-uremic syndrome (HUS)
- C  Heparin-induced thrombocytopenia (HIT)
- D  Idiopathic thrombocytopenic purpura
- E  Thrombotic thrombocytopenic purpura

55 A 45-year-old woman has had episodes of blurred vision and headaches for the past 6 months. She has had worsening confusion with paresthesias over the past 3 days. On physical examination, she has a temperature of 39.6°C, pulse of 100/min, respiratory rate of 20/min, and blood pressure of 80/50 mm Hg. Petechial hemorrhages are noted over her trunk and extremities. Laboratory findings include hemoglobin, 10.9 g/dL; hematocrit, 34%; MCV, 96 μm³; platelet count, 28,000/mm³; and WBC count, 8500/mm³. Fragmented RBCs are noted on her peripheral blood smear. Blood urea nitrogen is 40 mg/dL, and serum creatinine is 3.1 mg/dL. Which of the following is the most likely underlying cause for her findings?

- A  Circulating toxin that injures capillary endothelium
- B  Decreased factor VIII activity
- C  Defective ADP-induced platelet aggregation
- D  Formation of autoantibodies to platelet glycoproteins IIb/IIIa and Ib-IX
- E  Inappropriate release of thrombopoietic substances into blood
- F  Presence of antibodies against ADAMTS13 metalloproteinase
56 A 12-year-old boy has had worsening problems with joint mobility involving his arms and legs, particularly his knees and ankles, for the past 6 years. He has been receiving therapy for this condition. His grandfather had a similar condition and died at age 25 years. On physical examination, he has no visible petechiae or areas of purpura. Laboratory studies show that prothrombin time is 12 seconds, and partial thromboplastin time is 52 seconds. After addition of an equivalent aliquot of normal plasma, the partial thromboplastin time is 30 seconds. Hemoglobin is 12.9 g/dL, platelet count is 238,500/mm³, and WBC count is 6620/mm³. His platelet function studies are normal. What is the most likely inheritance pattern for his condition?

A Autosomal dominant
B Autosomal recessive
C Confined placental mosaicism
D Germline mutation
E X-linked recessive

57 An 11-year-old child has a history of easy bruising. At age 9 years, he experienced hemorrhaging around the pharynx that produced acute airway obstruction. Family history indicates that other male relatives have similar bleeding problems. Laboratory studies show hemoglobin, 13.1 g/dL; hematocrit, 39.2%; platelet count, 228,000/mm³; WBC count, 5950/mm³; prothrombin time, 13 seconds; and partial thromboplastin time, 52 seconds. After 1:1 dilution of the patient’s plasma with normal pooled plasma corrects the partial thromboplastin time, 60 seconds. A chest radiograph shows bilateral pulmonary edema. He is most likely to have developed a transfusion reaction to which of the following components of the donor blood product?

A Red blood cells
B Platelets
C Granulocytes
D Fresh frozen plasma
E Albumin

58 A 15-year-old girl has a history of easy bruising and hemorrhages. Since menarche at the age of 13 years, she has had menometrorrhagia. On physical examination, she displays joint deformity and has decreased mobility of the ankles, knees, and wrists. Laboratory studies show hemoglobin, 11.8 g/dL; hematocrit, 35.1%; platelet count, 267,000/mm³; WBC count, 5960/mm³; prothrombin time, 13 seconds; and partial thromboplastin time, 60 seconds. A 1:1 dilution of the patient’s plasma with normal pooled plasma corrects the partial thromboplastin time. Which of the following is the most likely diagnosis?

A Antiphospholipid syndrome
B Hemophilia B
C Idiopathic thrombocytopenic purpura
D Thrombotic thrombocytopenic purpura
E Von Willebrand disease

59 A 16-year-old girl has had frequent nosebleeds since childhood. Her gums bleed easily, even with routine tooth brushing. She has experienced menorrhagia since menarche at age 13 years. On physical examination, there are no abnormal findings. Laboratory studies show hemoglobin, 14.1 g/dL; hematocrit, 42.5%; MCV, 90 µm³; platelet count, 277,400/mm³; and WBC count, 5920/mm³. Her platelets fail to aggregate in response to ADP, collagen, epinephrine, and thrombin. The ristocetin agglutination test result is normal. There is a deficiency of glycoprotein IIb/IIIa.

56 What is the most likely diagnosis?

A Disseminated intravascular coagulation
B Glanzmann thrombasthenia
C Immune thrombocytopenic purpura
D Vitamin C deficiency
E Von Willebrand disease

60 A 25-year-old man involved in a motorcycle accident incurs a laceration to his thigh. The bleeding is stabilized en route to the hospital, but on arrival he is noted to have orthostatic hypotension and his hematocrit is 21%. He receives 2 units of PRBCs. As the first unit is nearly finished transfusing, he becomes febrile and hypotensive. Urine output ceases. The serum above the clot in a red top phlebotomy tube is pink.

What is the most likely complication of transfusion that has most likely occurred in this man?

A Donor antibodies were directed against his granulocytes
B Donor blood was contaminated with hepatitis C virus
C Fluid overload led to congestive heart failure
D Mislabeled specimens were processed in the laboratory
E Foreign T lymphocytes attacked his tissues

61 A 72-year-old woman undergoes laparotomy for ruptured diverticulitis. A day later her WBC count is elevated and her blood pressure is 85/45 mm Hg. Her peripheral blood smear shows schistocytes. She receives 5 units of fresh frozen plasma. As the 5th unit is being transfused, she develops sudden severe dyspnea and begins coughing up large quantities of frothy sputum. A chest radiograph shows bilateral pulmonary edema. She is most likely to have developed a transfusion reaction to which of the following components of the donor blood product?

A Albumin
B Fibrinogen
C Granulocytes
D Platelets
E Red blood cells

62 A study of transfusion reactions reveals that some patients experienced an increase in body temperature that was greater than 1°C, accompanied by chills and hypotension. Blood cultures are positive for bacterial organisms. These patients respond to antibiotic therapy. Which of the following types of blood products did they most likely receive?

A Cryoprecipitate
B Fresh frozen plasma
C Granulocytes
D Platelets
E Red blood cells
CHAPTER 14  Hematopathology of Red Blood Cells and Bleeding Disorders

1. **E** Platelet number and function in this case are normal, and there is no detectable abnormality in the extrinsic or intrinsic pathways of coagulation as measured by the prothrombin time or partial thromboplastin time. Petechiae and ecchymoses can result from increased vascular fragility, a consequence of nutritional deficiency (e.g., vitamin C), infection (e.g., meningococcemia), and vasculitic diseases. Chronic renal failure may depress platelet function. Chronic liver disease would affect the prothrombin time. Meningococcemia is an acute illness. Metastatic disease does not directly affect hemostasis, although extensive marrow metastases could diminish platelet production.

2. **D** The acute blood loss, in this case intraperitoneal hemorrhage, results in a reticulocytosis from marrow stimulation by anemia. Basophilic stippling of RBCs suggests a marrow injury, such as with a drug or toxin. Hypochromic RBCs occur in iron deficiency and thalassemias, both associated with reduced hemoglobin synthesis. Acute blood loss does not give rise to iron deficiency if iron stores and diet are adequate. Leukoe DMA, which is a myelophthisic process in the marrow, with both immature WBCs (myelocytes) and RBCs (nucleated forms) present. Schistocytes suggest a microangiopathic hemolytic anemia, which can accompany shock or sepsis.

3. **C** The marked reticulocytosis and marrow hyperplasia indicate that the marrow is responding to a decrease in RBCs. The reticulocytes are larger RBCs that slightly increase the MCV. An aplastic marrow is very hypocellular and unable to respond to anemia; it is associated with pancytopenia. The normal Coombs test results exclude an autoimmune hemolytic anemia. Iron deficiency impairs the ability of the marrow to mount a significant and sustained reticulocytosis. Iron deficiency anemia is typically microcytic and hypochromic, but could be partially masked here by reticulocytosis, which would not be as marked if iron were not available, but his diet is supplying needed iron. Infiltrative disorders, such as metastases in the marrow, would impair the ability to mount a reticulocytosis of this degree.

4. **A** Haptoglobin is a serum protein that binds to free hemoglobin. Ordinarily, circulating hemoglobin is contained within RBCs, but hemolysis can release free hemoglobin. The haptoglobin is used up as the amount of free hemoglobin increases. Systemic lupus erythematosus (SLE) is an autoimmune disease that can result in hemolysis by means of autoantibodies directed at RBCs, and the Coombs test result is often positive. SLE is best known to afflict young women, but it has a broad age range. Decreased iron can cause a hypochromic, microcytic anemia, but with hemolysis, the RBCs are recycled. Hemolysis is often accompanied by reticulocytosis if the marrow is intact and the iron is not lost. An elevated D-dimer level suggests a microangiopathic hemolytic anemia. Autoimmune diseases do not affect globin chain synthesis. Protoporphyrin can be increased with some forms of porphyria.

5. **F** Hereditary spherocytosis is a condition in which a mutation affects one of several membrane cytoskeletal proteins. Spectrin and related proteins are cytoskeletal proteins that are important in maintaining the RBC shape. These proteins include ankyrin (most common) and band 4.2, which binds spectrin to the transmembrane ion transporter; band 3; and protein 4.1, which binds the “tail” of spectrin to another transmembrane protein, glycoporin A. The abnormal RBCs with such mutant proteins are less deformable, lack central pallor on a peripheral blood smear, and they are sequestered and destroyed in the spleen. Thalassemias with abnormal α-globin or β-globin chains are associated with hypochromic microcytic anemias. Iron deficiency affects the heme portion of hemoglobin, leading to hypochromia and to microcytosis. Carbonic anhydrase in RBCs helps to maintain buffering capacity. Glucose-6-phosphate dehydrogenase (G6PD) deficiency is an X-linked condition that most commonly affects black males. Porphyrias may affect the production of porphyrin rings and may lead to hemolytic anemia along with abdominal pain, neurologic problems, or skin findings.

6. **C** The crescent-shaped RBCs (sickled RBCs) are characteristic of hemoglobin SS. This disease is most common in individuals of African and eastern Arabian descent. The sickled RBCs are susceptible to hemolysis (mainly vascular, in the spleen), but they also can cause microvascular occlusions anywhere in the body, most commonly bone, lungs, liver, and brain, leading to ischemia and severe pain. Vascular occlusions in the lungs are often accompanied by infection and lead to “acute chest syndrome.” Abdominal pain and back pain are common and severe, requiring prompt and effective analgesia. The cell membranes of reversibly sickled cells are abnormally “sticky,” and they adhere to capillary endothelium, especially in lungs. Vasoconstriction is caused by depletion of NO by free hemoglobin. Adhesion of RBCs to endothelial retards blood flow, creates hypoxia, and precipitates local sickling and vascular occlusion. Chronic tissue hypoxia does occur in sickle cell anemia, but it produces insidious impairment of function in organs such as heart, kidneys, and lungs. Defects in the alternative pathway of complement activation predispose to infection with encapsulated bacteria, such as Haemophilus influenzae and Streptococcus pneumoniae. Autoantibodies to alveolar basement membrane can be part of Goodpasture syndrome, which also affects kidneys. The most severe intravascular hemolysis occurs with major transfusion reactions.

7. **D** Patients with hereditary spherocytosis may have an aplastic crisis precipitated by a parvovirus infection. In adults who do not have a defect in normal RBC production, such as hereditary spherocytosis or sickle cell anemia, or
who are not immunosuppressed, parvovirus infection is self-limited and often goes unnoticed. When there is an underlying RBC production defect, then RBC production is shut down by parvovirus, and there is no reticuloendotheliosis. Disseminated intravascular coagulation gives rise to thrombocytopenia, bleeding, and the appearance of fragmented RBCs in the blood smear. Reticuloendotheliosis would be prominent with hemolysis and with RBC antibodies. Iron deficiency does not occur in hemolytic anemias because the iron that is released from hemolysed cells is reused.

PBD9 632–634  BP9 410–411  PBDB 665  BP8 410–411

8 C In patients with hereditary spherocytosis, spheroidal cells are trapped and destroyed in the spleen because the abnormal RBCs have reduced deformability. Splenectomy is beneficial because the spherocytes are no longer detained by the spleen. Splenectomy has no effect on the synthesis of spectrin or RBC deformability; the RBCs in spherocytosis are not killed by opsonization. In warm antibody hemolytic anemia, opsonized RBCs are removed by the spleen. Reactive oxygen species do not play a role in anemias. Iron is not the rate-limiting step to RBC production when the iron can be recycled within the body.


9 A This patient of Mediterranean descent has β-thalassemia major. In this condition, there is a severe reduction in the synthesis of β-globin chains without impairment of α-globin synthesis. The free, unpaired α-globin chains form aggregates that precipitate within normoblasts and cause them to undergo apoptosis. The death of RBC precursors in the bone marrow is called "ineffective erythropoiesis." Not only does this cause anemia, but it also increases the absorption of dietary iron, giving rise to iron overload, which results in hemochromatosis with infiltrative cardiomyopathy, hepatic cirrhosis, and "bronze diabetes" from pancreatic islet dysfunction. The severe anemia triggers erythropoietin synthesis, which expands the erythropoietic marrow. The marrow expansion encroaches on the bones, causing maxillofacial deformities. Extramedullary hematopoiesis causes hepatosplenomegaly. In comparison, the hemolytic anemia is mild in β-thalassemia minor, and there is very little ineffective erythropoiesis. Hemochromatosis is particularly detrimental to the liver and heart. Patients with chronic anemia may require RBC transfusions, which adds even more iron to body stores. The other listed options do not lead to a marked expansion of hematopoiesis.


10 D In sickle cell anemia, the cumulative ischemic damage to the spleen results in autosplenectomy, leaving behind a small fibrotic remnant of this organ. The impaired splenic function and resultant inability to clear bacteria from the bloodstream can occur early in childhood, leading to risk for infection with encapsulated bacterial organisms. Immunodeficiency results from lack of splenic function, not from lack of immunoglobulins. Endothelium can be damaged with sickling, and adhesion between endothelial cells and RBCs is increased in sickle cell anemia. Complement proteins are part of innate immune responses in acute inflammation. There is no impairment in production or function of neutrophils.

PBD9 635–638  BP9 411–413  PBDB 647  BP8 427–428

11 C The infant had α-thalassemia major, which is most likely to occur in individuals of Southeast Asian ancestry, each of whose parents could have two abnormal α-globin genes on chromosome 16. A complete lack of α-globin chains precludes formation of hemoglobins A_1, A_2, and F. Only a tetramer of γ chains (Bart’s hemoglobin) can be made, leading to severe fetal anemia. Inheritance of three abnormal α-globin chains leads to hemoglobin H disease, with tetramers of β chains; survival to adulthood is possible. Hemoglobin E disease produces mild hemolytic anemias


12 B This patient has β-thalassemia, probably of at least intermediate severity. There is decreased β-globin chain formation, with increased hemoglobin A_2 and F to compensate. There is ineffective erythropoiesis and increased erythropoietin to drive increased iron absorption, leading to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload. Chronic anemia requiring transfusion therapy exacerbates hemochromatosis. Iron deposited in endocrine tissues can lead to gonadal, pituitary, thyroid, islet cell, and adrenal failure. Secondary hypersplenism can result from the splenomegaly, with sequestration of platelets and leukocytes. The abnormal ankyrin gene leads to hereditary spherocytosis and a mild hemolytic anemia with splenomegaly, but not to iron overload.


13 E Children and adults with sickle cell anemia may benefit from hydroxyurea therapy, which can increase the concentration of hemoglobin F in RBCs, which interferes with the polymerization of hemoglobin S. However, the therapeutic response to hydroxyurea often precedes the increase in hemoglobin F levels. Hydroxyurea also has an anti-inflammatory effect, increases the mean RBC volume, and can be oxidized by heme groups to produce nitric oxide that promotes vasodilation. Because hemoglobin F levels remain high through the first 5 to 6 months of life, patients with sickle cell anemia typically do not manifest the disease during infancy. Because both β-globin chains are affected, no hemoglobin A_1 is produced, and A_2 levels are never high. Globin synthesis overall is not going to increase, and globin synthesis must be balanced to produce normal hemoglobin. The hemolysis associated with sickling promotes erythropoiesis, but the concentration of hemoglobin S is not changed. Hydroxyurea does not significantly shift the oxygen dissociation curve or change the oxygen affinity of the various hemoglobins.

PBD9 635–638  BP9 411–413  PBDB 645–648  BP8 426–428
CHAPTER 14  Hematopathology of Red Blood Cells and Bleeding Disorders

14 C Although β-thalassemia minor and iron deficiency anemia are both characterized by hypochromic and microcytic RBCs, there is no increase in hemoglobin A₂ in iron deficiency states. A normal serum ferritin level also excludes iron deficiency. In contrast to β-thalassemia major, there is usually a mild anemia without major organ dysfunction with β-thalassemia minor. Diseases that produce hemolysis and increase erythropoiesis (e.g., autoimmune hemolytic anemia, malaria) do not alter the composition of β-globin chain production. Anemia of chronic disease may mimic iron deficiency and thalassemia minor with respect to hypochromia and microcytosis; however, anemia of chronic disease is associated with an increase in the serum concentration of ferritin.

15 C Glucose-6-phosphate dehydrogenase (G6PD) deficiency is an X-linked disorder that affects about 10% of African-American males. The lack of this enzyme subjects hemoglobin to damage by oxidants, including drugs such as primaquine, sulfonamides, nitrofurantoin, phenacetin, and aspirin (in large doses). Infection can also cause oxidative damage to hemoglobin. Heinz bodies are denatured hemoglobin, and they damage the RBC membrane, giving rise to intravascular hemolysis. The “bite cells” result from the attempts of overeager splenic macrophages to pluck out the Heinz bodies, adding an element of extravascular hemolysis. Heterozygotes with α-thalassemia (1 or 2 abnormal genes out of 4 total α-globin genes) have no major problems, but in cases of α-thalassemia major, perinatal death is the rule. Likewise, β-thalassemia minor and sickle cell trait are conditions usually with no major problems and no relation to drug usage. RBC membrane abnormalities, such as hereditary spherocytosis (caused by abnormal spectrin), typically produce a mild anemia without significant hemolysis, and there is no drug sensitivity. Some autoimmune hemolytic anemias can be drug related, but the hemolysis is predominantly extravascular.

16 C This is one mechanism for β⁺ thalassemia. Because the introns are usually involved, the flanking exons remain, and some normal splicing can occur, so that some β-globin chain synthesis can occur, but not sufficient for adequate hemoglobin production. The other listed mutations lead to a block in translation, with no functional β-globin chain synthesis, typical for β⁰ thalassemia.

17 D Paroxysmal nocturnal hemoglobinuria (PNH) is a disorder that results from an acquired stem cell membrane defect produced by a PIGA gene mutation that prevents the membrane expression of certain proteins that require a glycolipid anchor. These include proteins that protect cells from lysis by spontaneously activated complement. As a result, RBCs, granulocytes, and platelets are exquisitely sensitive to the lytic activity of complement. The RBC lysis is intravascular, so patients can have hemoglobinuria (dark urine). Defects in platelet function are believed to be responsible for venous thrombosis. Recurrent infections can be caused by impaired leukocyte functions. Patients with PNH may develop acute leukemia or aplastic anemia as complications. Mutations in the β-globin chain can give rise to hemoglobinopathies such as sickle cell anemia. Patients with factor V (Leiden) and prothrombin G20210A mutations can present with thromboses, but there is no anemia or leukopenia. Patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency have an episodic course from exposure to agents such as drugs that induce hemolysis. Spectrin mutations give rise to hereditary spherocytosis.

18 F Glucose-6-phosphate dehydrogenase (G6PD) deficiency predisposes the hemoglobin in RBCs to oxidative injury from drugs such as primaquine, and can induce hemolysis. Oxidant injury to hemoglobin produces inclusion of denatured hemoglobin within RBCs. The inclusions damage the cell membrane directly, giving rise to intravascular hemolysis. These damaged RBCs have reduced membrane deformability, and they are removed from the circulation by the spleen. The remaining mechanisms listed are not directly drug dependent. Hemolytic anemias with antibody coating RBCs can occur with autoimmune diseases, prior transfusion, and erythroblastosis fetais. Impaired RBC nuclear maturation occurs as a result of vitamin B₁₂ or folate deficiency. Impaired globin synthesis occurs in thalassemias. Complement lysis is enhanced in paroxysmal nocturnal hemoglobinuria, which results from mutations in the PIGA gene. Mechanical fragmentation of RBCs is typical of microangiopathic hemolytic anemias, such as disseminated intravascular coagulation. Reduced RBC membrane deformability is seen in patients with abnormalities in cytoskeletal proteins, such as spectrin; the latter causes hereditary spherocytosis.

19 E This patient has a warm autoimmune hemolytic anemia secondary to systemic lupus erythematosus (SLE). A positive Coombs test result indicates the presence of anti-RBC antibodies in the serum and on the RBC surface. Most cases of warm autoimmune hemolytic anemia are idiopathic, but one fourth occur in individuals with an identifiable autoimmune disease, such as SLE; in other cases, drugs are the cause. The immunoglobulin coating the RBCs acts as an opsonin to promote splenic phagocytosis. Nucleated RBCs can be seen in active hemolysis because the marrow compensates by releasing immature RBCs. Septicemia is more likely to lead to a microangiopathic hemolytic anemia. The increased RBC destruction in hereditary spherocytosis is extravascular and not immune mediated. Infections such as mononucleosis and Mycoplasma are associated with cold autoimmune hemolytic anemia (with an elevated cold agglutinin titer).

20 B Cold agglutinin disease has antibody (usually IgM) coating RBCs. The IgM antibodies bind to the RBCs at low temperature at peripheral body sites and fix complement; however, complement is not lytic at this temperature. With an increase in temperature within core internal organs, the IgM is dissociated from the cell, leaving behind C₃b. Most
of the hemolysis occurs extravascularly in the cells of the mononuclear phagocyte system, such as Kupffer cells in the liver, or splenic macrophages, because the coating of complement C3b acts as an opsonin. IgG is typically involved in warm antibody hemolytic anemia, which is chronic and is not triggered by cold. Raynaud phenomenon occurs in exposed, colder areas of the body, such as the fingers and toes. The patient probably has an elevated cold agglutinin titer. Histamine is released in type I hypersensitivity reactions. Fibronectin is an adhesive cell surface glycoprotein that aids in tissue healing. IgE is present in allergic conditions.


21 D The findings point to Coombs-positive immune hemolytic anemia with indirect hyperbilirubinemia. Cold agglutinin immunohemolytic anemia can be seen with lymphoid neoplasms and infections such as Mycoplasma, Epstein-Barr virus, HIV, influenza virus, and cytomegalovirus (CMV). IgM binds to RBCs at cooler peripheral body regions and then fixes complement. At warmer central regions, the antibody is eluted, but the complement marks the RBCs for extravascular destruction in the spleen, but there is minimal intravascular hemolysis. The increased RBC turnover increases the MCV and the bilirubin, which is mainly indirect. Chemotherapy can suppress bone marrow production, but more likely all cell lines, and without an immune component. Although this patient has CMV infection, CMV hepatitis would likely increase direct and indirect bilirubin, and not account for anemia. Folate deficiency could account for macrocytosis, but not a positive Coombs test. Non-Hodgkin lymphomas do not often involve colon, but this might account for gastrointestinal bleeding with features of iron deficiency and microcytosis.


22 C She has a circulating antibody against her RBCs leading to hemolytic anemia. The indirect antiglobulin (Coombs) test detects antibody in the plasma. The direct antiglobulin (Coombs) test detects antibody bound to RBCs. Autoimmune hemolytic anemias can be a feature of autoimmune diseases, such as systemic lupus erythematosus in this woman. Most of the hemolysis is extravascular in the spleen, but some can be intravascular. The reticulocyte count is typically increased (polychromasia) with hemolysis, and serum haptoglobin is diminished. An elevated D-dimer suggests a microangiopathic hemolytic anemia, but she has no schistocytes on the peripheral blood smear. Increased osmotic fragility is a feature of RBCs in paroxysmal nocturnal hemoglobinuria. Her mild macrocytosis indicates increased reticulocytosis, not vitamin B12 deficiency, and hemolysis is not part of pernicious anemia. Increased serum ferritin is typical for anemia of chronic disease (mild increase) or hemochromatosis (marked increase).


23 B Mechanical trauma to RBCs is possible, but typically is not severe. It can follow strenuous exercises involving repeated blows to body parts. Complement-mediated lysis is a feature of immunohemolytic anemias. Increased osmotic fragility is noted in spherocytes. Sickle cell anemia is not likely to be found in the population groups in the countries noted, and persons with this disease are not likely to be playing rugby. Splenic sequestration is a feature of hemolytic anemias due to membrane defects and antibodies.

PBD9 644  BP9 418  PBD8 654  BP8 433

24 D Throughout human history, malaria has influenced the increasing gene frequency of hemoglobin S. Individuals who are heterozygous for hemoglobin S have the sickle cell trait. They are more resistant to malaria, particularly the most malignant form caused by Plasmodium falciparum, because the parasites grow poorly or die at low oxygen concentrations, perhaps because of low potassium levels caused by potassium efflux from RBCs on hemoglobin sickling. The malarial parasite has difficulty completing its life cycle, even in cells with moderate amounts of hemoglobin S, as found in heterozygotes, giving a selective advantage to such persons living in endemic areas for falciparum malaria. Borrelia burgdorferi is the spirochete that causes Lyme disease. Clostridium perfringens may produce gas gangrene after soft-tissue injuries. Cryptococcus neoformans can cause granulomatous disease in immunocompromised individuals. Treponema pallidum is the infectious agent causing syphilis. Treponosoma brucei infection causes sleeping sickness.

PBD9 634–635, 638  BP9 418–419  PBD8 645  BP8 426–428, 433–434

25 A The infant most likely has erythroblastosis fetalis because maternal antibodies are coating fetal RBCs. A fetal-maternal hemorrhage in utero or at the time of delivery in a previous pregnancy (or with previous transfusion of incompatible blood) can sensitize the mother, resulting in production of irregular IgG antibodies. In subsequent pregnancies, these antibodies (in contrast to the naturally occurring IgM antibodies) can cross the placenta to attach to fetal cells, leading to hemolysis. In the past, most cases were caused by Rh incompatibility (e.g., Rh-negative mother, Rh-positive infant), but the use of RhoGAM administered at birth to Rh-negative mothers has eliminated almost all such cases when recognized. Other, less common blood group antigens can be involved in this process, however. The other conditions listed are not antibody mediated. A stem cell defect results in aplastic anemia and immunodeficiency. Impaired globin synthesis occurs in thalassemias. Mechanical fragmentation of RBCs is typical of microangiopathic hemolytic anemias, such as disseminated intravascular coagulation, which is more typical of pregnant women with obstetric complications. Oxidative injury to hemoglobin is typical of glucose-6-phosphate dehydrogenase (G6PD) deficiency. Reduced RBC membrane deformability is seen in patients with abnormalities in cytoskeletal proteins, such as spectrin; the latter causes hereditary spherocytosis.


26 E This is benign tertian malaria. The bite of the Anopheles mosquito introduces sporozoites, which travel to the liver to reproduce. The resulting merozoites are released into the bloodstream and infect RBCs. Asexual reproduction within
the RBCs yields trophozoites, and periodic hemolysis with release of the parasites produces the characteristic clinical findings. Aspergillus organisms invade blood vessels and cause thrombosis, but hemolysis of RBCs is inconsequential. Babesiosis is far less common than malaria, is endemic to the northeastern United States, and does not produce episodic fevers. Dirofilaria is the heartworm found in dogs, which rarely infects humans and does not cause hemolysis. Similar to other gram-negative bacteria, Escherichia coli can release lipopolysaccharide, which causes severe sepsis and possible disseminated intravascular coagulation, a microangiopathic hemolytic anemia. Wuchereria bancrofti is a nematode that prefers to live in lymphatics.

**27 C** Drug-induced hemolytic anemias are neither common nor severe enough to be recognized, since the hemolysis is mainly extravascular. However, many patients receive drugs, so the potential for a drug reaction exists, and this immune-mediated mechanism must be distinguished from other causes for anemia. Cephalosporins are the most frequent drugs implicated. Treatment consists of cessation of therapy with the drug, because most cases are due to drug-dependent antibody formation. Nutrient deficiencies reduce marrow production, so a reticulocytosis is unlikely. DIC is unlikely with a normal platelet count. Parvovirus infection may suppress erythropoiesis transiently in individuals with normal red cells, but may precipitate an aplastic crisis in those with a hemoglobinopathy. Persons with abnormal red cells are likely to have a history of anemia. Hemoglobinopathies are not Coombs positive.

**28 D** This child has cerebral malaria, and the smear shows numerous ring forms of the parasites in RBCs. Infection occurs via the bite of an Anopheles mosquito. Malarial parasites infect RBCs, causing hemolysis and anemia. Falciparum malaria is the worst form. The parasites tend to be released from cells at periodic intervals, leading to periodic fever and chills. The parasites adhere to the vascular endothelium and lead to ischemia in various organs, including the brain with consequent acute cerebral edema. There is hemolytic anemia. The liver and spleen become progressively enlarged. Babesiosis is a rare, tick-borne disease found in the northeastern United States, which can produce a hemolytic anemia, but the organisms produce a classic “tetrad” in RBCs. Lyme disease, caused by Borrelia burgdorferi, is best known for producing chronic arthritis, but meningoencephalitis, neuritis, and neuropathy may complicate this illness. Leishmaniasis, caused by Leishmania donovani, produces mainly visceral disease without cerebral findings. Sleeping sickness, caused by Trypanosoma brucei, can be a chronic disease or a more acute disease causing brain dysfunction (sleeping sickness). Wuchereria bancrofti produces lymphatic filariasis with elephantiasis.

**29 C** With RBC microcytosis, iron deficiency anemia must be considered. It could be a nutritional deficiency in children and pregnant women, but more likely is due to chronic blood loss in adults. The ferritin concentration is a measure of storage iron because it is derived from the total body storage pool in the liver, spleen, and marrow. About 80% of functional body iron is contained in hemoglobin; the remainder is in muscle myoglobin. Individuals with severe liver disease can have an elevated serum ferritin level because of its release from liver stores. A bone marrow biopsy specimen provides a good indication of iron stores because the iron stain of the marrow shows hemosiderin in macrophages, but such a biopsy is an expensive procedure. Some patients with hemoglobinopathies, such as β-thalassemia, also can have a microcytic anemia, but this is far less common than iron deficiency. The serum haptoglobin level is decreased with intravascular hemolysis, but the anemia is normocytic because the iron can be recycled. The serum iron concentration or transferrin level by itself gives no indication of iron stores because in anemia of chronic disease, the patient’s iron level can be normal to low, and the transferrin levels also can be normal to low, but iron stores are increased. Transferrin, a serum transport protein for iron, usually has about 33% iron saturation.

**30 E** The figure shows RBC hypochromia and microcytosis, consistent with iron deficiency, the most common cause for anemia worldwide, either from nutritional deficiency or chronic blood loss. The lack of iron impairs heme synthesis. The marrow response is to “downsize” the RBCs, resulting in a microcytic and hypochromic anemia. At this patient’s age, bleeding from an occult malignancy, such as a colonic adenocarcinoma, should be strongly suspected as the cause of iron deficiency. An autoimmune hemolytic anemia would appear as a normocytic anemia or as a slightly increased MCV with pronounced reticulocytosis. Macrocytosis would accompany a history of chronic alcoholism, probably because of poor diet and folate deficiency. Thalassemias may result in a microcytosis, but β-thalassemia major causes severe anemia soon after birth, and survival to age 73 years is unlikely. By this patient’s age, hemophilia A would have resulted in joint problems; because the bleeding is mainly into soft tissues without blood loss, the iron is recycled. Vitamin B₁₂ deficiency also results in a macrocytic anemia.

**31 C** Iron absorption from the gut is tightly controlled. When body iron stores are adequate, absorption of dietary iron via DMT-1 in the duodenum is retarded, and release of iron from storage pools is inhibited. When body iron stores decrease, as with chronic blood loss, iron absorption increases. The liver-derived plasma peptide hepcidin has been found to be the iron absorption regulator. Hepcidin levels increase when iron stores are high. Such fine control of iron absorption may fail, as in patients with ineffective erythropoiesis (e.g., β-thalassemia) who continue to absorb iron despite excess storage iron. Hepcidin levels are inappropriately low with
both hereditary and acquired hemochromatosis. DMT-1 is an
iron transporter that moves nonheme iron from the gut lumen to duodenal epithelium. Hemosiderin is an aggregated form of ferritin that does not circulate and is found only in tissues. Mutations in the HFE gene, which encodes an HLA-like transmembrane protein, lead to excessive absorption of dietary iron and hemochromatosis. Transferrin transports iron between plasma, iron stores, and developing erythroblasts.

32 A The increased ferritin concentration and reduced total iron-binding capacity are typical of anemia of chronic disease, such as an autoimmune disease. Increased levels of cytokines such as interleukin-6 lead to increased hepatic production of hepcidin that stops ferroportin from releasing storage iron, promoting sequestration of storage iron, with poor use for erythropoiesis. Secretion of erythropoietin by the kidney is impaired. Various underlying diseases, including cancer, collagen vascular diseases, and chronic infections, can produce this pattern of anemia. Iron deficiency would produce a microcytic anemia, with a low serum ferritin level and reduced hepcidin production. Aplastic anemia is unlikely because the platelet count and WBC count are normal. Megaloblastic anemias are macrocytic without an increase in iron stores. Microangiopathic hemolytic anemias are caused by serious acute conditions such as disseminated intravascular coagulation; these patients have thrombocytopenia caused by widespread thrombosis. Thalassemia minor is uncommon and is not associated with a positive ANA test result.

33 C The iron concentration and iron-binding capacity are low; however, in contrast to the finding in anemia of iron deficiency, the serum ferritin level is increased. This increase is typical of anemia of chronic disease. In this case, the chronic disease is rheumatoid arthritis. Underlying chronic inflammatory or neoplastic diseases increase the secretion of cytokines such as interleukin-1, tumor necrosis factor, and interferon-γ. These cytokines promote sequestration of iron in storage compartments and depress erythropoietin production. Autoantibody hemolytic anemias occur in several autoimmune diseases, such as systemic lupus erythematosus, but not usually in patients with rheumatoid arthritis, as in this case. Normal serum haptoglobin rules out intravascular hemolysis; iron is recycled at a rapid rate. Impaired synthesis of β-globin chains gives rise to β-thalassemias, also characterized by hemolysis. Complement lysis is enhanced in paroxysmal nocturnal hemoglobinuria, which results from mutations in the PIGA gene. Patients with this disorder have a history of infections. Sequestration of RBCs in the spleen occur when RBC membranes are abnormal, as in hereditary spherocytosis or sickle cell anemia, or RBCs are coated by antibodies, as in autoimmune hemolytic anemias. Metastases are space-occupying lesions (myelophthisic process) that can lead to leukoerythroblastosis, with nucleated RBCs and immature WBCs appearing on the peripheral blood smear.

34 A Chronic alcohol abuse can lead to folate deficiency, giving rise to megaloblastic anemia. Folic acid and vitamin B12 act as coenzymes in DNA synthetic pathways. A deficiency of either impairs the normal process of nuclear maturation. The hematopoietic cell nuclei remain large and primitive looking, giving rise to megaloblasts. The mature RBCs are larger than normal (macrocytes). Neutrophils often show defective segmentation, manifested by extra nuclear lobes. The nuclear maturation defect affects all rapidly dividing cells in the body. Patients with chronic alcohol abuse can have thrombocytopenia and leukopenia, often because of secondary hypersplenism (alcoholic cirrhosis, leading to splenomegaly). Polychromatophilic RBCs represent reticulocytes, and their number is reduced because of the failure of marrow to produce adequate numbers of RBCs despite anemia. Hemolytic anemias, in which antibody coats RBCs, can occur in autoimmune diseases, prior transfusion, and erythroblastosis fetalis. Hemoglobinopathies can produce a mild macrocytosis because more reticulocytes are released. An imbalance in α-globin and β-globin chain synthesis, seen in thalassemias, leads to microcytosis of RBCs. Complement lysis is enhanced in paroxysmal nocturnal hemoglobinuria, which results from mutations in the PIGA gene. Reduced RBC membrane deformability is seen in patients with abnormalities of cytoskeletal proteins, such as spectrin; the latter causes hereditary spherocytosis.

35 B Teardrop RBCs are indicative of a myelophthisic disorder (i.e., something filling the bone marrow, such as fibrous connective tissue). The leukoerythroblastosis, including immature RBCs and WBCs, is most indicative of myelofibrosis. Splenomegaly also is typically seen in myelofibrosis. A leukoerythroblastic picture also can be seen in patients with infections and metastases involving the marrow. Hyperplasia of erythroid normoblasts occurs in hemolytic anemias. Leukoerythroblastosis is not seen in hemolytic anemias. Replacement of marrow by fat occurs in aplastic anemia, which is characterized by pancytopenia. The presence of megaloblasts in the marrow indicates folate or vitamin B12 deficiency—both cause macrocytic anemia. Marrow packed with myeloblasts is typical of acute myeloid leukemia. In this condition, the peripheral blood also would show many myeloblasts and failure of myeloid maturation.

36 C Although folate and vitamin B12 deficiency both give rise to a macrocytic anemia, a deficiency of vitamin B12 also can result in demyelination of the posterior and lateral columns of the spinal cord. In some cases this deficiency will only be revealed by elevated levels of homocysteine and methylmalonic acid in the serum, because these are more sensitive indicators, particularly earlier in the disease. The anemia caused by vitamin B12 deficiency can be ameliorated by increased administration of folate; this masks the potential neurologic injury by improving the anemia. Treating vitamin B12 deficiency does not improve the anemia caused by
folate deficiency, however. An aplastic anemia is unlikely to result from a nutritional deficiency. Folate has no cofactor for absorption, but vitamin B₁₂ must be complexed to intrinsic factor and secreted by gastric parietal cells, and then the complex must be absorbed in the terminal ileum, so diseases such as atrophic gastritis and Crohn disease can affect vitamin B₁₂ absorption more than folate. The peripheral smear could appear the same and offers no means for distinguishing these deficiencies.

PBD9 648–649  BP9 422–423  PBD8 658  BP8 438

37 E The high MCV indicates a marked macrocytosis, greater than expected from reticulocytosis alone. The two best-known causes for such an anemia (also known as megaloblastic anemia when characteristic megaloblastic precursors are seen in the bone marrow) are vitamin B₁₂ and folate deficiency. Because vitamin B₁₂ complexed with intrinsic factor is absorbed in the terminal ileum, its removal can cause vitamin B₁₂ deficiency. Anemia of chronic disease is generally a normocytic anemia. Chronic blood loss and iron deficiency produce a microcytic pattern of anemia, as does dietary iron deficiency.

Hemolytic anemia is unlikely several weeks after blood transfusion. Inflammatory bowel diseases (e.g., Crohn disease) increase the risk of malignancy, but myelophthisic anemias (from space-occupying lesions of the marrow) are usually normocytic to mildly macrocytic (from reticulocytosis).


38 A Aplastic anemia leads to marked pancytopenia. Many cases are idiopathic, although some can follow toxic exposures to chemotherapy drugs or to chemicals, such as benzene. Some cases may follow viral hepatitis infections. An intrinsic defect in stem cells, or T lymphocyte suppression of stem cells, can play a role in the development of aplastic anemia. Hemolysis is unlikely because the bilirubin is normal, and there is no history of an autoimmune disease. An increased susceptibility to complement lysis occurs in paroxysmal nocturnal hemoglobinuria as a result of mutations in the PIGA gene. It is unlikely that the patient has metastatic disease at this age, with no prior illness; metastases are more likely to produce a leukoerythroblastic peripheral blood appearance.

Sequestration of peripheral blood cells in an enlarged spleen could account for mild pancytopenia, but in this case the spleen is not enlarged.

PBD9 653–655  BP9 424  PBD8 663–664  BP8 439–440

39 D Telomerase is normally present in continuously dividing cells, such as hematopoietic stem cells, to prevent shortening of chromosomal telomeres; otherwise, shortened chromosomes cannot divide properly. Both genetic and acquired forms of aplastic anemia have been found that exhibit this mechanism. Alkaline phosphatase participates in bone remodeling. A deficiency of ADAMTS13, a metalloproteinase, can lead to accumulation of large von Willebrand multimers that promote platelet microaggregate formation, resulting in thrombotic thrombocytopenic purpura (TTP). Pyruvate kinase deficiency is one rare form of congenital anemia. Tyrosine kinases participate in cell growth regulation, and are better known to be involved with myeloproliferative disorders.

PBD9 653–654  BP9 424  PBD8 663  BP8 439

40 B Her pancytopenia and absence of a reticulocytosis strongly suggest bone marrow failure. Aplastic anemia has no apparent cause in half of all cases. In other cases, drugs and toxins may be identified; drugs such as chemotherapeutic agents are best known for this effect. A preceding viral infection may be identified in some cases, but bacterial infections rarely cause aplastic anemias. Individuals with pancytopenia are subject to bleeding disorders because of the low platelet count and to infections because of the low WBC count. Dietary history would not be helpful because this patient’s clinical and laboratory picture is not characteristic of iron deficiency or folate or vitamin B₁₂ deficiency. The only known familial cause of aplastic anemia (Fanconi anemia) is rare. Menstrual history would be relevant if the patient had hypochromic microcytic anemia.

PBD9 653–654  BP9 424  PBD8 663–664  BP8 439–440

41 E His prostatic adenocarcinoma has metastasized to the bone marrow. High alkaline phosphatase, hypercalcemia, and a leukoerythroblastic pattern in the peripheral blood (immatu re WBCs and RBCs) are a consequence of the tumor acting as a space-occupying lesion. Such myelophthisic anemias also may be caused by infections. The anemia of chronic disease is mild. Aplastic anemias are unlikely to include leukoerythroblastosis. Hemolytic anemia should be accompanied by an increase in bilirubin and no abnormalities in calcium metabolism. The MCV in this case is not in the megaloblastic range.

PBD9 655  BP9 424  PBD8 665  BP8 440

42 A She has myasthenia gravis with thymoma and red cell aplasia. The edrophonium, an acetylcholinesterase inhibitor, will counteract the effect of the acetylcholine receptor antibodies of myasthenia gravis, but not improve muscle function with antibodies against voltage-gated calcium channels in Lambert-Eaton myasthenic syndrome (a paraneoplastic syndrome often associated with small cell lung carcinomas). The pulmonary findings suggest heart failure, and the tachycardia is consistent with high-output congestive heart failure from anemia. Pure red cell aplasia can be primary or arise secondarily to neoplasms, particularly thymic tumors, or autoimmune disorders. A lymphocytosis would be characteristic for lymphocytic leukemia, but this would not affect muscle strength, and lymphoblastic leukemia/lymphoma produces mediastinal masses in much younger persons. Megakaryocytic hyperplasia would be characteristic for peripheral consumption of platelets, with a disorder such as immune thrombocytopenic purpura (ITP), which is not likely to be associated with thymoma. Most thymomas act in a benign fashion and do not metastasize widely. Myelofibrosis could produce anemia with fatigue and weakness, but not predominantly with repetitive motion. Plasmacytosis is associated with myeloma that affects bone marrow, not the mediastinum.

PBD9 655  BP9 457  PBD8 664–665  BP8 439
**D** Heat stroke caused by hyperthermia and loss of perspiration from dehydration is producing hemoconcentration with a relative polycythemia (note the elevated serum sodium level). Erythroblastemia is quite rare, and patients with this disorder are too ill to run a race. Chronic obstructive pulmonary disease is a cause of secondary polycythemia from chronic hypoxemia, but it does not produce hemoconcentration. Diabetes insipidus can result from a lack of antidiuretic hormone release, which leads to free water loss and dehydration, but not to hyperthermia. Increased erythropoietin levels are seen in secondary polycythemias, including those associated with chronic hypoxemia (high altitude or lung disease) and those associated with neoplasms secreting erythropoietin (paraneoplastic syndrome). Polycythemia vera is a form of myeloproliferative disorder, in which erythropoietin levels are low, and there is no dehydration.

**A** Some patients with antiphospholipid syndrome (APS) have systemic lupus erythematosus, but others (such as this patient) do not. Arterial and deep venous thrombosis can occur in APS, with increased risk particularly of cerebral arterial thrombosis. Anticardiolipin antibody often leads to a false-positive serologic test result for syphilis (Treponema pallidum infection). Thrombocytopenia is often present. APS should be considered in women who have recurrent miscarriages. Polycythemia vera is a myeloproliferative disorder that predisposes to thrombosis, but this patient’s hemoglobin value does not support this diagnosis. Thrombophlebitis occurs more frequently in pregnancy, but this explains only venous thrombosis and not the anticardiolipin antibodies. Trousseau syndrome, a hypercoagulable state associated with an underlying malignancy, can explain venous and arterial thromboses, but not the anticardiolipin antibodies. The patient’s age argues against cancer. Von Willebrand disease is a bleeding disorder without thrombotic complications.

**A** The presence of thrombocytopenia, increased prothrombin and partial thromboplastin times, and fibrin split products, and the low fibrinogen concentration all suggest disseminated intravascular coagulation (DIC), which was most likely caused by a retained dead fetus. This obstetric complication can lead to DIC through release of thromboplastins from the fetus. The thromboplastins cause widespread microvascular thrombosis and consume clotting factors and platelets. There is no defect in platelet function. There is no damage to the vascular endothelium or vascular wall. Platelet production is normal, but platelets are consumed by widespread thrombosis of small vessels.

**E** Gram-negative sepsis with widespread endothelial damage causes disseminated intravascular coagulation. The figure shows fragmented RBCs, including “helmet cells,” typical of conditions that can produce a microangiopathic hemolytic anemia, such as disseminated intravascular coagulation, thrombocytopenia purpura, systemic lupus erythematosus, hemolytic-uremic syndrome, and malignant hypertension. Such fragmented RBCs are called schistocytes. In autoimmune hemolytic anemia, the hemolysis also is extravascular, and schistocytes are sometimes formed. Schistocytes may be present in hereditary spherocytosis, but the RBC destruction is extravascular, and fragmented RBCs do not appear in the peripheral blood. Marked anisocytosis and poikilocytosis can occur in iron deficiency and in megaloblastic anemias, but fragmentation of RBCs is not seen.

**C** This patient’s bleeding tendency is caused by a low platelet count. She most likely has idiopathic chronic immune thrombocytopenic purpura (ITP), in which platelets are destroyed in the spleen after being coated with antibodies to platelet membrane glycoproteins IIb-IIIa or Ib-IX. These antibodies coat both the patient’s platelets and any transfused platelets. Because the spleen is the major source of the antibody and the site of platelet destruction, splenectomy can be beneficial if corticosteroid therapy is not. There is no defect in the production of platelets. Platelet functions are normal in ITP. Chronic blood loss would not lead to thrombocytopenia when normal bone marrow function is present. Abnormal platelet-endothelial interactions are more likely to cause thrombosis. Suppression of pluripotent stem cells gives rise to aplastic anemia, which is accompanied by pancytopenia.

**A** Acute immune thrombocytopenic purpura (ITP) and chronic ITP are caused by antiplatelet autoantibodies, but the acute form is typically seen in children after a viral disease. If the bone marrow was aplastic, all cell lines should be reduced. Glycoprotein IIb/IIIa dysfunction/deficiency can be seen with Glanzmann thrombasthenia and chronic ITP. Scurvy caused by vitamin C deficiency leads to increased capillary fragility with ecchymoses, but not to thrombocytopenia. Von Willebrand factor metalloproteinase deficiency is a feature of thrombotic thrombocytopenic purpura.

**B** Heparin-induced thrombocytopenia affects 3% to 5% of patients treated for 1 to 2 weeks with unfractionated heparin. These patients form IgG antibodies to heparin–platelet factor 4 complexes that bind to Fc receptors on the surface of platelets, causing platelet activation and, paradoxically, thrombosis. Aspirin has antiplatelet effects that take days to occur, and bleeding (not thrombosis) is the major risk. Tissue plasminogen activator and urokinase are fibrinolytic agents, with the former used acutely to treat conditions such as coronary thrombosis, although the latter also may be used for venous clot lysis. Warfarin (Coumadin) was avoided in this patient because of the protein C deficiency; typically, the patient is switched from heparin to
warfarin. Warfarin therapy prolongs the prothrombin time by interfering with vitamin K–dependent clotting factor synthesis in the liver.

PBD9 659  BP9 453  PBD8 668–669  BP8 472

50 F The diagnosis of thrombotic thrombocytopenic purpura (TTP) is based on finding a classic pentad: transient neurologic problems, fever, thrombocytopenia, microangiopathic hemolytic anemia, and acute renal failure. The diagnosis is suggested by decreased ADAMTS13 activity. There is a deficiency of ADAMTS13, which acts as a von Willebrand factor multimer protease. Most cases of TTP are idiopathic and associated with antibodies to ADAMTS13. Hereditary TTP may result from mutations in the ADAMTS13 gene. The abnormalities are produced by small platelet-fibrin thrombi in small vessels in multiple organs. The heart, brain, and kidney often are severely affected. Of the other choices, only disseminated intravascular coagulation is a microangiopathic process often are severely affected. Of the other choices, only disseminated intravascular coagulation is a microangiopathic process.


51 E This patient has hepatitis C with severe hepatocyte damage. Many of the clotting factors that are instrumental in the intravascular measurement of the extrinsic pathway of coagulation, as measured by the prothrombin time, are synthesized in the liver. Increased fibrin split products suggest a consumptive coagulopathy, such as disseminated intravascular coagulation. Von Willebrand factor is produced by endothelial cells, not hepatocytes. Platelet aggregation is a measure of platelet function, which is not significantly affected by liver disease. The platelet count is not affected directly by liver disease.

PBD9 663–665  BP9 454  PBD8 119, 670  BP8 468–469

52 B An inherited bleeding disorder with normal platelet count and prolonged bleeding time suggests von Willebrand disease, confirmed by the ristocetin-dependent bioassay for von Willebrand factor (vWF). Von Willebrand disease is a common bleeding disorder, with an estimated frequency of 1%. In most cases, it is inherited as an autosomal dominant trait. In these cases, a reduction in the quantity of vWF impairs platelet adhesion to damaged vessel walls, and hemostasis is compromised. Because vWF acts as a carrier for factor VIII, the level of this procoagulant protein (needed for the intrinsic pathway) is diminished, as in this case. The levels of factor VIII rarely are reduced enough, however, to be clinically significant, perhaps 1 in 10,000 persons. Prolonged partial thromboplastin time corrected by normal plasma is a reflection of factor VIII deficiency. Because the disease is not a disorder of stem cells, bone marrow failure or myeloproliferative disorder is not a likely outcome. Joint hemorrhages are a feature of hemophilia A and B, not von Willebrand disease. Patients with von Willebrand disease are not prone to thrombosis, as are individuals with factor V (Leiden) mutation or other inherited disorders of anticoagulation.

PBD9 661–662  BP9 455  PBD8 671–672  BP8 473–474

53 C Reduced numbers of platelets can result from decreased production or increased destruction. Marrow examination in this case shows numerous megakaryocytes, which excludes decreased production. Accelerated destruction can be caused by hypersplenism, but there is no splenomegaly in this case. Peripheral platelet destruction is often immunologically mediated and can result from well-known autoimmune diseases such as systemic lupus erythematosus, or it can be idiopathic. When all known causes of thrombocytopenia are excluded, a diagnosis of idiopathic (immune) thrombocytopenic purpura (ITP) can be made. This patient seems to have no other symptoms or signs and has no history of drug intake or infections that can cause thrombocytopenia. ITP is most likely. Thrombotic thrombocytopenic purpura (TTP) is another entity to be considered, but TTP produces a microangiopathic hemolytic anemia (MAHA) that typically is associated with fever, neurologic symptoms, and renal failure. Disseminated intravascular coagulation (DIC) is another form of MAHA. Hemophilia B, similar to hemophilia A, leads to soft tissue bleeding, and the partial thromboplastin time is prolonged, but the platelet count is normal. Metastases can act as a space-occupying lesion in the marrow to reduce hematopoiesis, but this is unlikely to be selective with megakaryocytes, and in this case, there is a megakaryocytic hyperplasia. Vitamin K deficiency prolongs the prothrombin time initially and the partial thromboplastin time if severe, but does not affect platelets. In von Willebrand disease, bleeding is due to abnormal platelet adhesion, but platelet numbers are normal.

PBD9 659–660  BP9 453–454  PBD8 667–668  BP8 471–472

54 E A deficiency of ADAMTS13, from an acquired antibody to this metalloproteinase or a genetic mutation in the encoding gene, can lead to accumulation of large von Willebrand multimers that promote platelet microaggregate formation, resulting in TTP that is marked by a pentad of microangiopathic hemolytic anemia, fever, neurologic changes, thrombocytopenia, and renal failure. DIC results from acquired conditions that promote consumption of coagulation factors, not a metalloproteinase deficiency. HUS is very similar to TTP, but is more likely related to a preceding infectious gastroenteritis with diarrhea. HIT occurs in about 5% of individuals receiving heparin, and the most serious complication is widespread arterial and venous thrombosis. ITP is mainly complicated by bleeding from thrombocytopenia.

PBD9 659–660  BP9 453–454  PBD8 667–668  BP8 471–472

55 F The clinical features (neurologic abnormalities, fever, thrombocytopenia, microangiopathic hemolytic anemia, renal failure) point to thrombotic thrombocytopenic purpura (TTP), in which there is an inherited or acquired deficiency of the von Willebrand factor (vWF) metalloproteinase (ADAMTS13) that normally cleaves very high molecular weight multimers of vWF. The absence of ADAMTS13 gives rise to large multimers of vWF that promote widespread platelet aggregation, and the resulting microvascular occlusions in brain, kidney, and elsewhere produce organ dysfunction, thrombocytopenia, microangiopathic hemolytic anemia (MAHA), and bleeding. Circulating toxins, principally
endotoxins elaborated by Enterobacteriaceae such as Escherichia coli are important in causing endothelial injury in hemolytic-uremic syndrome (HUS). HUS has similar clinical findings to TTP, but has a different pathogenesis. Decreased factor VIII activity is a feature of hemophilia A, an X-linked disorder rare in women, characterized by bleeding into soft tissues, such as joints, and normal platelet number and function. Defective aggregation of platelets in the presence of ADP and thrombin is a feature of a rare inherited disorder of platelets called Glanzmann thrombasthenia. Release of thromboplastic substances from tumor cells or a retained dead fetus can lead to disseminated intravascular coagulation with MAHA, but this patient has no source of thromboplastins.

56 E His hemophilia A is monitored with factor VIII activity. Hemophilia B with factor IX deficiency would present similar findings, but it is much less common. Transfusion of factor VIII helps to prevent joint and soft-tissue hemorrhages. Of individuals with hemophilia A, 20% may develop an inhibitor to factor VIII, typically an IgG antibody that neutralizes activity of any infused factor VIII. In this case, the partial thromboplastin time partly corrects, indicating that some inhibitor may be present. Macrocotaneous bleeding is more typical for platelet disorders and von Willebrand disease. With hemophilia, bleeding is often into joints and soft tissues. The prothrombin time (PT) is normal, and clotting factors affecting this pathway are made in the liver; factor VIII is in the intrinsic pathway measured by partial thromboplastin time (PTT). Hemophilia A and hemophilia B are X-linked inherited conditions, so males are mainly affected. There can be rare new mutations (such as in Queen Victoria), which introduce the gene into a family.

57 B The severity of hemophilia A depends on the amount of factor VIII activity. With less than 1% activity, there is severe disease, and joint hemorrhages are common, leading to severe joint deformity and ankylosis. Mild (1% to 5%) and moderate (5% to 75%) activity is often asymptomatic except in severe trauma. Hemophilia A is typically caused by decreased factor VIII activity. The affected patient is male and has male relatives who are affected (X-linked transmission). The partial thromboplastin time (PTT) is prolonged because factor VIII is required for the intrinsic pathway; the prothrombin time (PT) is normal because the extrinsic pathway does not depend on factor VIII function. The correction of the PTT by mixing with normal plasma is important. With a deficiency of factor VIII only, the addition of normal plasma, a source of factor VIII, corrects the PTT. Failure to correct PTT by normal plasma indicates the presence of an inhibitor in the patient’s serum. About 15% of patients with hemophilia eventually develop an inhibitor to factor VIII. Petechiae, seen in patients with thrombocytopenia, are not a feature of hemophilia. Factor VIII deficiency does not affect the life span of RBCs. Because individuals with factor VIII deficiency do not depend on RBC transfusions, iron overload is not a usual consequence. The bleeding tendency of hemophilia is not associated with splenomegaly.

58 B The history in this case is similar to that in the preceding question; however, the partial thromboplastin time (PTT) is corrected by normal pooled plasma. The patient who has hemophilia B caused by factor IX deficiency, and inhibitors of factor IX are absent from the patient’s serum. How is this possible in a female patient? X-inactivation (“unfavorable lyonization”) can explain this phenomenon and could explain why female carriers of hemophilia A or B have a tendency to bleed. (“When you have eliminated the impossible, that which remains, however improbable, must be the truth,” said Sherlock Holmes in The Sign of Four.) An in vitro mixing study of patient and pooled plasma such as this usually corrects an abnormality caused by a deficiency of a procoagulant factor, but if there is a coagulation inhibitor in the patient’s plasma, the clotting test would show an abnormal result. The mixing study excludes the antiphospholipid syndrome. Idiopathic thrombocytopenic purpura is characterized by the presence of antiplatelet antibodies and thrombocytopenia. Thrombotic thrombocytopenic purpura is a microangiopathic hemolytic anemia characterized by renal failure and central nervous system abnormalities. Von Willebrand disease is caused by decreased platelet adhesion and has features resembling thrombocytopenia.

59 B Glanzmann thrombasthenia is a rare autosomal recessive disorder with defective platelet aggregation from deficiency or dysfunction of glycoprotein IIb/IIIa. The platelet aggregation studies described here are characteristic for this disorder. Disseminated intravascular coagulation results in consumption of all coagulation factors and platelets, so the prothrombin time and partial thromboplastin time are elevated with thrombocytopenia. Immune thrombocytopenic purpura is caused by antibodies to platelet membrane glycoproteins IIb/IIIa or Ib/IX. Scurvy resulting from vitamin C deficiency causes bleeding into soft tissues and skin from increased capillary fragility, but platelet number and function are normal. Von Willebrand disease is one of the most common bleeding disorders and results from qualitative or quantitative defects in von Willebrand factor.

60 D The findings are those of a hemolytic (ABO) transfusion reaction, virtually all of which result from clerical errors. You learned that proper phlebotomy procedure requires labeling the tubes just after drawing the blood, not handing the tubes to someone else for labeling and possible mix-up. This is not consistent with a transfusion-related acute lung injury (TRALI) in which there are donor-derived HLA or granulocyte-specific antibodies in the recipient’s blood product that are directed against antigens on the recipient WBCs. Though hepatitis C infection is still possible, but uncommon, from transfused blood, this infection has an incubation period of months. The two units of packed RBCs in a young person are very unlikely to lead to circulatory overload. He does not have transfusion-associated graft-versus-host disease, which is a rare but untreatable condition with a very high mortality rate.
Transfusion-related acute lung injury (TRALI) is caused when the donor plasma contains HLA or granulocyte-specific antibodies which correspond to antigens found on the recipient patient WBCs. Granulocyte enzymes are released, increasing capillary permeability and resulting in sudden pulmonary edema with respiratory distress. TRALI most often occurs with administration of blood products with plasma, such as fresh frozen plasma, and more often from donor women with more HLA antibodies from prior pregnancies. TRALI may occur when the granulocytes are “primed” by sensitization within pulmonary capillaries by preexisting lung disease. Transfusion reactions to the blood proteins albumin and fibrinogen do not occur. Platelet transfusion may cause alloimmunization and subsequent platelet destruction with future platelet transfusion. Incompatible RBCs can cause alloimmunization, but more acutely, a hemolytic transfusion reaction can occur.

Because platelets must be stored at room temperature, any contamination that occurred at the time of collection, such as bacterial skin flora of the donor, is more likely to cause sepsis in the recipient. The other products listed are refrigerated, which reduces bacterial growth. The causative agent of syphilis, *Treponema pallidum*, is often killed at refrigerator temperatures.
1 A 24-year-old primigravida has a fetal ultrasound performed at 18 weeks’ gestation that shows a normal amniotic fluid index, but there is a large echogenic region in the right lung. A term infant is delivered with no apparent external anomalies, but soon after birth the infant has respiratory distress. CT imaging shows a normal left lung, but there is an area of opacification in the right lung supplied by a vessel from the aorta. Which of the following is the most likely diagnosis?

A Extralobar sequestration
B Foregut cyst
C Hyaline membrane disease
D Oligohydramnios sequence
E Tracheoesophageal fistula

2 A 30-year-old man is hospitalized after a motor vehicle accident in which he sustains blunt trauma to his chest. On physical examination, there are contusions to the right side of the chest, but no lacerations. Within 1 hour after the accident, he develops sudden difficulty breathing and marked pain on the right side. Vital signs now show that he is afebrile; his pulse is 80/min, respirations are 30/min and shallow, and blood pressure is 100/65 mm Hg. Breath sounds are not audible, and there is tympany to percussion on the right side. Which of the following radiographic findings for this man is shown in the figure?

A Interstitial fibrosis
B Patchy infiltrates
C Pleural effusion
D Pneumothorax
E Ventilation/perfusion mismatch

3 A 58-year-old man with ischemic heart disease undergoes coronary artery bypass graft surgery under general anesthesia. Two days postoperatively, he experiences increasing respiratory difficulty with decreasing arterial oxygen saturation. On physical examination, his heart rate is regular at 78/min, respirations are 25/min, and blood pressure is 135/85 mm Hg. The hemoglobin concentration has remained unchanged, at 13.7 g/dL, since surgery. After he coughs up a large amount of mucoid sputum, his condition improves. Which of the following types of atelectasis does he most likely have?

A Compression
B Contraction
C Micro
D Relaxation
E Resorption
A 45-year-old man incurs blunt chest trauma in a boating accident. On examination he has marked right chest wall pain. A chest radiograph shows a fractured 7th rib on the right side. Over the next 2 days he has subcutaneous soft tissue swelling with non-painful crepitance on palpation of the right chest wall. Leakage of which of the following is most likely producing this swelling?

A. Acid  
B. Air  
C. Blood  
D. Lymph  
E. Pus

A 68-year-old man has had worsening dyspnea and orthopnea for the past 3 years with increased production of frothy sputum. On examination, crackles are auscultated at lung bases. A chest radiograph shows bilateral interstitial infiltrates, distinct Kerley B lines, and a prominent left heart border. Laboratory studies show Na+, 135 mmol/L; K+, 3.8 mmol/L; Cl−, 99 mmol/L; CO₂, 25 mmol/L; glucose, 76 mg/dL; creatinine, 1.5 mg/dL; and urea nitrogen, 30 mg/dL. Fractional excretion of sodium is less than 1%. Plasma renin, aldosterone, and antidiuretic hormone levels all are increased. B-type natriuretic peptide (BNP) is 200 pg/mL (normal <100 pg/mL). Which of the following pathologic findings is this man most likely to have?

A. Aldosteronoma  
B. Bilateral adrenal atrophy  
C. Chronic glomerulonephritis  
D. Ischemic heart disease  
E. Pulmonary fibrosis  
F. Small cell carcinoma

A 26-year-old woman with postpartum sepsis is afebrile on antibiotic therapy, but she has had worsening oxygenation over the past 3 days. Her chest radiograph shows scattered bilateral pulmonary opacifications. A ventilation-perfusion scan shows areas of mismatch. Which of the following microscopic findings is most likely to be present in her lungs?

A. Alveolar hyaline membranes  
B. Arterial plexiform lesions  
C. Interstitial fibrosis  
D. Lymphocytic infiltrates  
E. Respiratory bronchiolar destruction

After a hemicolectomy to remove a colon carcinoma, a 56-year-old man develops respiratory distress. He is intubated and receives mechanical ventilation with 100% oxygen. Three days later, his arterial oxygen saturation decreases to 60%. A chest radiograph shows increasing opacification in all lung fields. A transbronchial lung biopsy specimen on microscopic examination shows hyaline membranes lining distended alveolar ducts and sacs. Which of the following is the most likely mechanism underlying these morphologic changes?

A. Aspiration of oropharyngeal contents  
B. Intravascular thrombi with coagulopathy  
C. Leukocyte-mediated injury to alveolar capillaries  
D. Reduced production of surfactant  
E. Release of fibrogenic cytokines by macrophages

A 48-year-old man has gradually increasing dyspnea and a 4-kg weight loss over the past 2 years. He has smoked two packs of cigarettes per day for 20 years, but not for the past year. Physical examination shows an increase in the anteroposterior diameter of the chest. Auscultation of the chest shows decreased breath sounds. A chest radiograph shows bilateral hyperlucent lungs; the luency is especially marked in the upper lobes. Pulmonary function tests show that FEV₁ is markedly decreased, but FVC is normal, and the FEV₁/FVC ratio is decreased. Which of the following is most likely to contribute to the pathogenesis of his disease?

A. Abnormal epithelial cell chloride ion transport  
B. Decreased ciliary motility with irregular dynein arms  
C. Impaired hepatic release of α₁-antitrypsin  
D. Macrophage recruitment and release of interferon-γ  
E. Release of elastase from neutrophils

A 20-year-old, previously healthy man is jogging one morning when he trips and falls to the ground. He suddenly becomes markedly short of breath. On examination in the emergency room there are no breath sounds audible over the right side of the chest. A chest radiograph shows shift of the mediastinum from right to left. A chest tube is inserted on the right side, and air rushes out. Which of the following underlying diseases is most likely to have produced this complication?

A. Asthma  
B. Bronchiectasis  
C. Centriacinar emphysema  
D. Chronic bronchitis  
E. Distal acinar emphysema  
F. Panlobular emphysema

A 49-year-old man has had increasing dyspnea for the past 4 years. He has an occasional cough with minimal sputum production. On physical examination, his lungs are hyperresonant with expiratory wheezes. Pulmonary function tests show increased total lung capacity (TLC) with slightly increased FVC and decreased FEV₁ and FEV₁/FVC ratio. Arterial blood gas measurement shows pH of 7.35; PO₂, 65 mm Hg; and PCO₂, 50 mm Hg. Which of the following disease processes should most often be suspected as a cause of these findings?

A. Centrilobular emphysema  
B. Chronic pulmonary embolism  
C. Diffuse alveolar damage  
D. Nonatopic asthma  
E. Sarcoïdosis  
F. Silicosis
11 A 33-year-old man has had increasing dyspnea for the past 8 years. He does not smoke. On examination, there are decreased breath sounds over lower lung fields. A chest radiograph shows flattened diaphragms; his CT scan is shown in the figure. Pulmonary function tests show decreased DLCO, decreased FEV₁, and increased FVC. Arterial blood gas analysis shows Po₂, 65 mm Hg; Pco₂, 60 mm Hg; HCO₃⁻, 32 mEq/L; and pH, 7.35. A sibling is similarly affected. What is the most likely mechanism for his pulmonary disease?

A Atopy with IgE binding to mast cells  
B CFTR gene mutation  
C Increased neutrophil proteases  
D Prior infection with tuberculosis  
E Reduced antielastase activity

12 A 45-year-old man has smoked two packs of cigarettes per day for 20 years. For the past 4 years, he has had a chronic cough with copious mucoid expectoration. During the past year, he has had multiple respiratory tract infections diagnosed as “viral flu.” He has also developed difficulty breathing, tightness of the chest, and audible wheezing. His breathing difficulty is relieved by inhalation of a β-adrenergic agonist and disappears after the chest infection has resolved. Which of the following pathologic conditions is most likely responsible for his clinical condition?

A α₁-Antitrypsin deficiency with panlobular emphysema  
B Centrilobular emphysema with cor pulmonale  
C Chronic asthmatic bronchitis  
D Cystic fibrosis with bronchiectasis  
E Hypersensitivity pneumonitis with restrictive lung disease

13 A study is conducted of individuals who smoked at least one pack of cigarettes per day for 30 years. These individuals undergo pulmonary function testing, and a large subset is found to have decreased FEV₁, normal to decreased FVC, and FEV₁/FVC ratios less than 70%. Autopsy data from this subset of individuals in the study with a low FEV₁/FVC ratio are analyzed. Which of the following respiratory tract structures in the lungs is likely to be affected most by the underlying disease?

A Alveolar duct  
B Alveolar sac  
C Bronchi  
D Respiratory bronchiole  
E Terminal bronchiole

14 A 62-year-old man is a smoker with a 10-year history of cough productive of copious mucopurulent sputum. Over the past 6 months, he has developed progressive dyspnea. Physical examination shows bilateral pedal edema and a soft but enlarged liver. A chest radiograph shows bilateral pleural effusions and a prominent right heart border. Arterial blood gas values are Po₂, 60 mm Hg; Pco₂, 52 mm Hg; pH, 7.30; and HCO₃⁻, 29 mEq/L. He is intubated and placed on a ventilator, and he requires increasing amounts of oxygen. Which of the following microscopic findings is most likely to be present in the affected lungs?

A Bronchovascular distribution of granulomas  
B Carcinoma filling lymphatic spaces  
C Extensive interstitial fibrosis  
D Hypertrophy of bronchial submucosal glands  
E Mucosal infiltrates of eosinophils

15 A 12-year-old girl has a 7-year history of coughing and wheezing and repeated attacks of difficulty breathing. The attacks are particularly common in the spring. During an episode of acute respiratory difficulty, a physical examination shows that she is afebrile. Her lungs are hyperresonant on percussion, and a chest radiograph shows increased lucency of all lung fields. Laboratory tests show an elevated serum IgE level and peripheral blood eosinophilia. A sputum sample examined microscopically also has increased numbers of eosinophils. Which of the following histologic features is most likely to characterize the lung in her condition?

A Dilation of respiratory bronchioles with loss of elastic fibers  
B Inflammatory destruction of bronchial walls  
C Interstitial and alveolar edema with hyaline membrane formation  
D Patchy areas of consolidation with leukocytic exudates in alveoli  
E Remodeling of airways with smooth muscle hyperplasia
16. A 33-year-old man suddenly develops severe dyspnea with wheezing. On physical examination, his vital signs are temperature, 37°C; pulse, 95/min; respiration, 35/min; and blood pressure, 130/80 mm Hg. A chest radiograph shows increased lucency in all lung fields. Arterial blood gas analysis shows $\text{PO}_2$, 65 mm Hg; $\text{PCO}_2$, 30 mm Hg; and pH, 7.48. A sputum cytologic specimen shows Curschmann spirals, Charcot-Leyden crystals, branching septate hyphae, and eosinophils in a background of abundant mucus. What is the most likely risk factor predisposing him to this illness?

A. Cytokine gene polymorphisms
B. Foreign body aspiration
C. Inhalation of environmental inorganic dusts
D. Inheritance of a CFTR gene mutation
E. Reduced circulating $\alpha_1$-antitrypsin levels
F. Smoking cigarettes for >10 years

17. A pharmaceutical company is designing agents to treat the recurrent bronchospasm of bronchial asthma. Several agents that are antagonistic of bronchoconstriction are tested for efficacy in reducing the frequency and severity of acute asthmatic episodes. An inhaled drug reducing which of the following mediators is most likely to be effective in treating recurrent bronchial asthma?

A. Th1 cytokines
B. Vasoactive amines
C. Th2 cytokines
D. Leukotrienes
E. Prostaglandins

18. A 35-year-old man has a 5-year history of episodic wheezing and coughing. The episodes are more common during the winter months, and he has noticed that they often follow minor respiratory tract infections. In the period between the episodes, he can breathe normally. There is no family history of asthma or other allergies. On physical examination, there are no remarkable findings. A chest radiograph shows no abnormalities. A serum IgE level and WBC count are normal. Which of the following is the most likely mechanism that contributes to the findings in his illness?

A. Accumulation of alveolar neutrophilic exudate
B. Bronchial hyperreactivity to chronic inflammation
C. Emigration of eosinophils into bronchi
D. Hypermorphosis to *Aspergillus spores*
E. Secretion of interleukin (IL)-4 and IL-5 by T cells

19. A study of persons with atopic asthma reveals that they develop pathologic changes in their airways with repeated bouts. These changes include smooth muscle and mucus gland hypertrophy. It is observed that the late-phase inflammatory response to allergens potentiates epithelial cell cytokine production that promotes airway remodeling. Which of the following immune cells is most important in this excessive inflammatory response to allergens?

A. B lymphocyte
B. Cytotoxic lymphocyte
C. Natural killer cell
D. Th1 lymphocyte
E. Th2 lymphocyte
F. Th17 lymphocyte

20. A 70-year-old woman has had episodes of dyspnea with wheezing and coughing, accompanied by urticaria for the past 3 years. She has had bouts of rhinitis. She has a 10-year history of osteoarthritis. On physical examination she has nasal polyps. Use of which of the following medications is the most likely risk factor for her respiratory disease?

A. Acetaminophen
B. Aspirin
C. Gabapentin
D. Morphine
E. Prednisone

21. A 19-year-old man has a history of recurrent mucoid rhinorrhea with chronic sinusitis and otitis media since childhood. He has experienced multiple bouts of pneumonia. His temperature is 37.7°C. On examination of his chest, there is tactile fremitus, rhonchi, and rales in lower lung fields. Nasal polyps are noted. A chest radiograph shows bronchial dilation with bronchial wall thickening, focal atelectasis, and areas of hyperinflation; his heart shadow appears mainly on the right. Which of the following abnormalities is he most likely to have?

A. $\alpha_1$-Antitrypsin deficiency
B. Atopy
C. Chloride ion channel dysfunction
D. Ciliary dyskinesia
E. HIV infection

22. A 35-year-old woman has experienced multiple bouts of severe necrotizing pneumonia since childhood, with *Haemophilus influenzae*, *Staphylococcus aureus*, *Pseudomonas aeruginosa*, and *Serratia marcescens* cultured from her sputum. She now has a cough productive of large amounts of purulent sputum. On physical examination, there is dullness to percussion with decreased breath sounds over the right mid to lower lung fields. A chest radiograph shows areas of right lower lobe consolidation. A bronchogram shows marked dilation of right lower lobe bronchi. Which of the following mechanisms is the most likely cause of her disease?

A. Congenital malformation of the bronchial walls
B. Damage to bronchial mucosa by major basic protein of eosinophils
C. Diffuse infiltration by bronchogenic carcinoma
D. Recurrent inflammation with bronchial wall destruction
E. Unopposed action of neutrophil-derived elastase on bronchi
23. A 41-year-old man has a 6-year history of increasing shortness of breath and weakness. On physical examination, he is afebrile and normotensive. A radiograph of his chest shows diffuse interstitial markings. Pulmonary function tests indicate diminished FVC, decreased DLCO (diffusing capacity), and a normal FEV$_1$/FVC ratio. Which of the following sets of pathologic changes is most likely to be found in his lungs?

A. Acute inflammation of bronchial walls with prominence of eosinophils
B. Chronic inflammation with bronchial mucus gland hypertrophy
C. Dilation of airspaces distal to respiratory bronchioles
D. Honeycomb lung with extensive alveolar septal fibrosis
E. Widespread alveolar epithelial necrosis and prominent hyaline membranes

24. A 68-year-old man has had worsening dyspnea with a nonproductive cough for the past 9 months. On physical examination, he is afebrile and normotensive. A chest radiograph shows irregular opacifications throughout both lungs. A transbronchial biopsy is obtained and the microscopic findings with trichrome stain are shown in the figure. Laboratory studies include negative serologic tests for ANA, anti-DNA topoisomerase I, ANCA, and anticitrulline antibody. Despite glucocorticoid therapy, his condition does not improve, and he dies 2 years later. What is the most likely diagnosis?

A. Goodpasture syndrome
B. Hypersensitivity pneumonitis
C. Idiopathic pulmonary fibrosis
D. Sarcoidosis
E. Systemic sclerosis

25. A study of pulmonary disease in persons who are smokers shows that tobacco used in greater amounts and for longer periods is positively correlated with the degree of lung parenchymal destruction with centrilobular emphysema. However, some persons with a history of extensive tobacco use have less lung damage than persons who smoked less. Polymorphisms involving which of the following genes are most likely to explain these differences in the repair response to lung injury in smokers?

A. AAT
B. BMPR2
C. CFTR
D. GM-CSF
E. TGF-β

26. A 52-year-old woman, an electrical engineer and non-smoker, has a 3-month history of increasing dyspnea. On examination she is afebrile and normotensive. CT imaging of her chest shows lower lobe reticular opacities. A transbronchial biopsy is performed and microscopically shows patchy interstitial inflammation with lymphocytes and plasma cells. No organisms are identified. Her condition slowly worsens over the next 10 years. Which of the following is the most likely diagnosis?

A. Desquamative interstitial pneumonitis
B. Hypersensitivity pneumonitis
C. Idiopathic interstitial fibrosis
D. Nonatopic bronchial asthma
E. Nonspecific interstitial pneumonia

27. A 54-year-old woman has had a mild fever with cough for a week. Her symptoms gradually improve over the next 10 days. She then begins to have increasing fever, cough, shortness of breath, and malaise. Now, on physical examination, her temperature is 37.9° C. There are inspiratory crackles on auscultation of the chest. A chest radiograph shows bilateral, patchy, small alveolar opacities. Chest CT scan shows small, scattered, ground-glass and nodular opacities. A transbronchial biopsy specimen shows polypoid plugs of loose fibrous tissue and granulation tissue filling bronchioles, along with a surrounding interstitial infiltrate of mononuclear cells. She receives a course of corticosteroid therapy, and her condition improves. Which of the following is the most likely diagnosis?

A. Bronchiectasis
B. Cryptogenic organizing pneumonia
C. Desquamative interstitial pneumonitis
D. Hypersensitivity pneumonitis
E. Pulmonary alveolar proteinosis
28 A 63-year-old man has had progressively worsening dyspnea over the past 10 years. He has noticed a 5-kg weight loss in the past 2 years. He has a chronic cough with minimal sputum production and no chest pain. On physical examination, he is afebrile and normotensive. A chest radiograph shows extensive interstitial disease. Pulmonary function tests show diminished DLCO, low FVC, and normal FEV₁/FVC ratio. Increased exposure to which of the following pollutants is most likely to produce these findings?
   A Carbon monoxide
   B Ozone
   C Silica
   D Tobacco smoke
   E Wood dust

29 A study of persons with a history of mining occupational exposure to inhaled dusts is performed. Though found in urban air in small amounts, this dust consists of 1- to 5-micron particles that are inert and insoluble. Fibrosis occurs only with large amounts of dust accumulation, mainly in upper lobes, with nodular opacities larger than 1 cm seen on chest radiographs. What is most likely in this dust?
   A Asbestos
   B Beryllium
   C Carbon
   D Iron
   E Sulfur dioxide

30 A 76-year-old man has experienced increasing dyspnea for the past 4 years. On physical examination, he is afebrile, with a pulse of 70/min, respirations 30/min, and blood pressure 120/75 mm Hg. A chest radiograph shows increased interstitial markings, but no effusions. The right heart border and the pulmonary arteries are prominent. A transbronchial biopsy is performed; the figure shows the microscopic appearance with Prussian blue stain. Which of the following is the most likely diagnosis?
   A Anthracosis
   B Asbestosis
   C Berylliosis
   D Calciosis
   E Silicosis

31 A radiographic study of inhalational lung diseases is conducted. One pattern of involvement is seen in persons whose total lung capacity, diffusing capacity, and compliance is decreased. This pattern consists of numerous bilateral nodular opacifications on chest radiographs. Polarizable needlelike crystals are seen on microscopic examination of these nodules. What inhaled substance is most likely to produce these findings?
   A Cigarette smoke
   B Mold spores
   C Silica dust
   D Sulfur dioxide
   E Wood particles

32 A 36-year-old woman has had a low-grade fever and worsening nonproductive cough and dyspnea for the past 2 years. On examination, she has breath sounds in all lung fields. A chest CT scan shows the findings in the figure. An arterial blood gas shows pH, 7.45; Po₂, 83 mm Hg; Pco₂, 30 mm Hg; and HCO₃⁻, 19 mEq/L. Pulmonary function tests show total lung capacity 3 L (60% of predicted), FEV₁ 2.5 (66% of predicted), and DLCO 10 mL/min/mm Hg (50% of predicted). Her pulmonary compliance is reduced. What is the most likely diagnosis?
   A α₁-Antitrypsin deficiency
   B Chronic bronchitis
   C Diffuse alveolar damage
   D Goodpasture syndrome
   E Nonatopic asthma
   F Sarcoidosis

33 A 65-year-old man worked in a shipyard for 10 years, and then he worked for 5 years for a company that installed fire retardant insulation. He experienced increasing dyspnea for 11 years with progressive respiratory failure and hypoxemia. A CT scan of his chest now shows a large mass encasing the left lung. Which of the following findings is most likely to be seen on a chest radiograph in this patient?
   A Bilateral fluffy perihilar infiltrates
   B Bilateral upper lobe cavitation
   C Diaphragmatic pleural calcified plaques
   D Endobronchial mass with atelectasis
   E Pleural effusions
34 A 61-year-old woman has noted increasing dyspnea and a nonproductive cough for 5 months. On physical examination, her temperature is 37.7°C. A chest radiograph shows prominent hilar lymphadenopathy with reticulonodular infiltrates bilaterally. A transbronchial biopsy is performed, and the microscopic findings include interstitial fibrosis and small, noncaseating granulomas. One granuloma contains an asteroid body in a Langhans giant cell. The medical history indicates that she smoked cigarettes for 10 years, but stopped 5 years ago. Which of the following is the most likely cause of her illness?

A  T cell-mediated response to unknown antigen  
B  Antibody-mediated diffuse alveolar damage  
C  Deposition of immune complexes  
D  Infection with atypical mycobacteria  
E  Smoke inhalation with loss of bronchioles

35 A 64-year-old alfalfa farmer has a 15-year history of increasing dyspnea. On physical examination, his temperature is 37.6°C. A chest radiograph shows a bilateral increase in linear markings. Pulmonary function tests show reduced FVC with a normal FEV1. A transbronchial lung biopsy specimen shows interstitial infiltrates of lymphocytes and plasma cells, minimal interstitial fibrosis, and small granulomas. What is the most likely cause of this clinical and pathologic picture?

A  Autoantibodies against alveolar basement membranes  
B  Chronic inhalation of silica particles  
C  Hypersensitivity to spores of actinomycetes  
D  Infection with *Mycobacterium tuberculosis*  
E  Prolonged exposure to inorganic dusts

36 A 25-year-old man experiences acute onset of fever, cough, dyspnea, headache, and malaise a day after moving into a new apartment. His symptoms subside over 3 days when he visits a friend in another city. On the day of his return, the symptoms recur. There are no remarkable findings on physical examination. A chest radiograph also is unremarkable. Which of the following pathogenetic mechanisms is most likely to produce these findings?

A  Antigen-antibody complex-mediated injury  
B  Antibody-mediated injury to basement membrane  
C  Formation of mycolic acid as a result of tubercular infection  
D  Generation of prostaglandins by basophil recruitment  
E  Release of histamine from mast cells  
F  Toxic injury to type I pneumocytes caused by inhaled dust

37 A 46-year-old man has had increasing dyspnea with nonproductive cough for the past year. On physical examination he is afebrile and has clubbing of digits. Pulmonary function testing reveals a mild restrictive abnormality along with reduced DLCO. A transbronchial biopsy is performed and microscopic examination shows numerous alveolar macrophages, plump epithelial cells, mild interstitial fibrosis, and loss of respiratory bronchioles. Lamellar bodies and iron pigment are present within these macrophages. Which of the following is the most likely etiology for his pulmonary disease?

A  Type I hypersensitivity  
B  Cigarette smoking  
C  Ciliary dyskinesia  
D  Inhalation of mold spores  
E  Cell-mediated response to silica dust

38 A 33-year-old woman has had increasing dyspnea with cough for the past 10 days. Over the past 2 days, her cough has become productive of chunks of gelatinous sputum. On physical examination, she is afebrile. There is extensive dullness to percussion over all lung fields. A chest radiograph shows diffuse opacification bilaterally. A transbronchial biopsy is performed and the microscopic appearance with H&E staining is shown in the figure. On electron microscopy, there are many lamellar bodies. Antibody directed against which of the following substances is most likely to cause her illness?

A  α1-Antitrypsin  
B  Chloride channel protein  
C  DNA topoisomerase 1  
D  Glomerular basement membrane  
E  GM-CSF  
F  Neutrophilic myeloperoxidase
A 61-year-old man had a myocardial infarction 1 year ago and now has marked congestive heart failure that reduces his ability to ambulate. Over the past 24 hours, he has developed right-sided chest pain. On auscultation, there are lower lobe rales. He is afebrile, his pulse is 70/min, his respirations are 27/min and shallow, and his blood pressure is 130/85 mm Hg. His chest CT scan is shown in the figure. Which of the following clinical disorders is most likely to precede the appearance of the lesion shown?

A  Chronic obstructive pulmonary disease  
B  HIV infection  
C  Nonbacterial thrombotic endocarditis  
D  Deep vein thrombosis  
E  Polyarteritis nodosa  
F  Silicosis

A clinical study is performed that includes patients who are hospitalized for more than 2 weeks and who were bedridden for more than 90% of that time. These patients undergo Doppler venous ultrasound examination of the lower extremities, blood gas testing, and radiographic pulmonary ventilation and perfusion scanning. A cohort of patients is found who have abnormal ultrasound results suggestive of thrombosis, blood gas parameters with a slightly lower PO2, and small pulmonary perfusion defects. Which of the following symptoms and signs are most likely to be seen in this cohort of patients?

A  Dyspnea  
B  Hemoptysis  
C  Palpitations  
D  Pleuritic pain  
E  Orthopnea  
F  No symptoms

A 68-year-old woman had a cerebral infarction and was hospitalized for 3 weeks. Her condition improved, and she was able to get up and move about with assistance. A few minutes after walking to the bathroom, she experienced sudden onset of severe dyspnea with chest pain and diaphoresis. Despite resuscitative measures, she died 30 minutes later. The major autopsy finding is shown in the figure. Which of the following is the most likely mechanism for sudden death in this patient?

A  Bronchoconstriction  
B  Compression atelectasis  
C  Hemorrhagic infarction  
D  Interstitial edema  
E  Acute cor pulmonale

A 45-year-old man has had progressive dyspnea on exertion with fatigue for the past 2 years. On auscultation of his chest he has a prominent pulmonary component of S2, a systolic murmur of tricuspid insufficiency, and bruits over peripheral lung fields. Jugular venous distension is present to the angle of his jaw when sitting. Laboratory studies show antiphospholipid antibodies. CT angiography shows eccentric occlusions with pulmonary arteries and mosaic attenuation of pulmonary parenchyma. Which of the following is the most likely disease process causing his pulmonary disease?

A  Atherosclerosis  
B  Pneumonitis  
C  Sarcoidosis  
D  Thromboembolism  
E  Vasculitis

A 75-year-old woman has had worsening lower leg edema and dyspnea for the past 5 years. On physical examination, her temperature is 36.9° C, pulse is 74/min, respirations are 19/min, and blood pressure is 110/75 mm Hg. There is dullness to percussion at the lung bases. A low rumbling heart murmur is present. A chest radiograph shows bilateral pleural effusions. An echocardiogram shows a large (4-cm) atrial septal defect. Which of the following conditions is most likely to be present in this woman?

A  Pulmonary hypertension  
B  Granulomatous inflammation  
C  Hemorrhagic infarction  
D  Interstitial fibrosis  
E  Necrotizing vasculitis
44. A 25-year-old woman has had progressive dyspnea and fatigue for the past 2 years. On physical examination, she has pedal edema, jugular venous distention, and hepatomegaly. Lung fields are clear on auscultation. Chest CT scan shows right heart enlargement. Cardiac catheterization is performed, and the pulmonary arterial pressure is increased, without gradients across the pulmonic valve, and no shunts are noted. A transbronchial biopsy is performed, and microscopic examination shows plexiform lesions. A mutation in a gene encoding for which of the following is most likely to cause her pulmonary disease?

A. B-type natriuretic peptide (BNP)
B. Bone morphogenetic receptor 2 (BMPR2)
C. Endothelial nitric oxide synthetase (eNOS)
D. Fibrillin-1
E. Lysyl hydroxylase
F. Renin

45. A 29-year-old man who has had no major illnesses experiences acute onset of hemoptysis. On physical examination, he has a temperature of 37°C, pulse of 83/min, respirations of 28/min, and blood pressure of 150/95 mm Hg. A chest radiograph shows bilateral fluffy infiltrates. A transbronchial lung biopsy on microscopic examination shows focal necrosis of alveolar walls associated with prominent intra-alveolar hemorrhage. Two days later, he has oliguria. The serum creatinine level is 2.9 mg/dL, and urea nitrogen is 31 mg/dL. Which of the following antibodies is most likely involved in the pathogenesis of his condition?

A. Anti–DNA topoisomerase I antibody
B. Anti–glomerular basement membrane antibody
C. Antimitochondrial antibody
D. Anti–neutrophil cytoplasmic antibody
E. Antinuclear antibody

46. A 42-year-old man has had chronic sinusitis for 7 months. He now has malaise and a mild fever that have persisted for 3 weeks. On physical examination, his temperature is 37.9°C. On auscultation, a few crackles are heard over the lungs. Laboratory studies show serum urea nitrogen of 35 mg/dL; creatinine, 4.3 mg/dL; ALT, 167 U/L; AST, 154 U/L; and total bilirubin, 1.1 mg/dL. A chest radiograph shows scattered 1-cm pulmonary nodules. A transbronchial lung biopsy is performed, and microscopic examination shows necrotizing granulomatous capillaritis, a poorly formed granuloma, and intra-alveolar hemorrhage. Urinalysis shows RBCs with RBC casts. Autoantibodies to which of the following are most likely to be present in this man?

A. DNA topoisomerase I (Scl-70)
B. Double-stranded DNA (dsDNA)
C. Glomerular basement membrane (GBM)
D. Serine proteinase 3 (PR3-ANCA)
E. Thermoactinomyces vulgaris

47. A clinical study is conducted in which patients who have undergone surgical procedures with intubation, mechanical ventilation, and general anesthesia are followed to determine the number and type of postoperative complications. The study group is found to have a higher incidence of pulmonary infections in the 2 weeks following their surgical procedure than patients who were not intubated and did not receive general anesthesia. Anesthesia is most likely to produce this effect via which of the following mechanisms?

A. Decreased ciliary function
B. Diminished macrophage activity
C. Hypogammaglobulinemia
D. Neutropenia
E. Squamous metaplasia
F. Tracheal erosions

48. A 54-year-old woman has had an increasingly severe cough productive of yellowish sputum for four days. On physical examination, her temperature is 38.9°C, and diffuse crackles are heard in the left lower lung posteriorly. Laboratory studies show a WBC count of 11,990/mm³ with 71% segmented neutrophils, 9% bands, 16% lymphocytes, and 4% monocytes. The representative gross appearance of the affected lung is shown in the figure. Which of the following pathogens is most likely to be cultured from this patient’s sputum?

A. Cryptococcus neoformans
B. Mycobacterium kansasii
C. Mycoplasma pneumoniae
D. Nocardia brasiliensis
E. Pneumocystis jiroveci
F. Streptococcus pneumoniae

49. A 71-year-old woman has smoked a pack of cigarettes per day for 50 years. She has had increasing dyspnea for 12 years. Over the past 3 days she has become febrile, with a productive cough, and severe dyspnea. Auscultation of her chest reveals rales and expiratory wheezes. Laboratory studies show peripheral blood neutrophilia. Which of the following organisms is most likely to be cultured from her sputum?

A. Histoplasma capsulatum
B. Influenza A
C. Moraxella catarrhalis
D. Mycobacterium kansasii
E. Pneumocystis jiroveci
50. A 51-year-old man has a history of chronic alcohol abuse. He is found in a stuporous condition after 3 days of binge drinking. On physical examination, his temperature is 39.2°C. A few crackles are heard on auscultation of the right lung base. A chest radiograph shows a 3-cm round lesion with an air-fluid level in the right lower lobe. Which pair of the following organisms is most likely to be detected in his sputum?

A. *Cryptococcus neoformans* and *Candida albicans*
B. *Cytomegalovirus* and *Pneumocystis jiroveci*
C. *Mycobacterium tuberculosis* and *Aspergillus fumigatus*
D. *Nocardia asteroides* and *Actinomyces israelii*
E. *Staphylococcus aureus* and *Bacteroides fragilis*

51. A 20-year-old man has had a mild fever with nonproductive cough, headache, and myalgias for the past week. On physical examination he has a temperature of 37.9°C and erythema of the pharynx. Diffuse crackles are heard on auscultation of the lungs. A chest radiograph shows bilateral extensive patchy infiltrates. A sputum Gram stain shows normal flora. Cold agglutinin titer is elevated. He receives a course of erythromycin therapy, and his condition improves. Infection with which of the following organisms is most likely to produce these findings?

A. *Legionella pneumophila*
B. *Mycobacterium fortuitum*
C. *Mycoplasma pneumoniae*
D. *Nocardia asteroides*
E. Respiratory syncytial virus

52. A 26-year-old woman from East Asia developed a fever with chills over the past 4 days. Yesterday, she had increasing shortness of breath and a nonproductive cough, headache, and myalgias. On physical examination, her temperature is now 38.6°C. A chest radiograph shows right lower lobe infiltrates. Laboratory studies show hemoglobin, 13.4 g/dL; hematocrit, 40.2%; platelet count, 78,400/mm³; and WBC count, 3810/mm³ with 77% segmented neutrophils, 2% bands, 5% lymphocytes, and 16% monocytes. Over the next 2 days, she has increasing respiratory distress requiring intubation and mechanical ventilation. A repeat chest radiograph shows worsening bilateral infiltrates. Infection with which of the following is most likely to have caused this patient’s illness?

A. Coronavirus
B. *Cytomegalovirus*
C. *Ebola virus*
D. *Herpes simplex virus*
E. Respiratory syncytial virus

53. An epidemiologic study shows that a highly pathogenic strain of influenza A virus with the antigenic type H5N1 that normally causes disease in birds has been increasingly found to cause influenza in humans. Unlike other strains of influenza A virus, this H5N1 virus is associated with a 60% mortality rate. The enhanced pathogenicity of this avian flu virus is primarily due to mutation in its genome that enables it to do which of the following?

A. Elicit a weak cytotoxic T-cell response
B. Enter many types of host cells
C. Escape inactivation by macrophages
D. Infect CD4+ helper T cells
E. Spread from humans to humans

54. A 4-year-old healthy girl from Utrecht in the Netherlands has had a fever with dyspnea, tachypnea, nonproductive cough, myalgias, and rhinorrhea for 3 days. On auscultation of her chest there are inspiratory and expiratory wheezes. A chest radiograph shows bilateral diffuse interstitial infiltrates. She recovers in 2 weeks with no sequelae. Which of the following organisms is most likely to be identified by PCR of her respiratory secretions?

A. *Group A Streptococcus*
B. *Bordetella pertussis*
C. *Candida albicans*
D. *Cytomegalovirus*
E. *Haemophilus influenzae*
F. Human metapneumovirus

55. A 3-year-old boy has had a cough, headache, and slight fever for 5 days. He is now having increasing respiratory difficulty. On physical examination, his temperature is 37.8°C, pulse is 81/min, respirations are 25/min, and blood pressure is 90/55 mm Hg. On auscultation, there are inspiratory crackles, but no dullness to percussion or tympany. Respiratory syncytial virus is isolated from a sputum sample. Which of the following chest radiographic patterns is most likely to be present?

A. Hilar lymphadenopathy
B. Hyperinflation
C. Interstitial infiltrates
D. Lobar consolidation
E. Pleural effusions
F. Upper lobe cavitation

56. A study of HIV-infected persons shows that those with CD4+ lymphocyte counts below 100 cells/μL are found to be at increased risk for pulmonary infections. Some of them have concurrent hepatosplenomegaly and lymphadenopathy, as well as malabsorption with weight loss, night sweats, and fever. Bronchoalveolar lavage specimens examined microscopically show macrophages filled with acid-fast infectious organisms. Which of the following infections have these persons developed?

A. *Aspergillus niger*
B. *Candida albicans*
C. *Legionella pneumophila*
D. *Mycobacterium avium-complex*
E. *Nocardia asteroides*
F. *Pseudomonas aeruginosa*
A 46-year-old woman has a routine health maintenance examination. On physical examination, there are no remarkable findings. Her body mass index is 22. She does not smoke. A tuberculin skin test is positive. A chest radiograph shows a solitary, 3-cm left upper lobe mass without calcifications. The mass is removed at thoracotomy by wedge resection. The microscopic appearance of this lesion is shown in the figure. Which of the following is the most likely diagnosis?

A. Mycobacterium tuberculosis infection
B. Necrotizing granulomatous vasculitis
C. Poorly differentiated adenocarcinoma
D. Staphylococcus aureus abscess
E. Thromboembolism with infarction

A 44-year-old man, a nonsmoker, has experienced a 3-kg weight loss over the past 3 months. He recently developed a low-grade fever and cough with mucoid sputum production, and after 1 week, he noticed blood-streaked sputum. On physical examination, his temperature is 37.7°C. There are bilateral crackles in the left upper lobe on auscultation of the chest. Chest CT scan shows a 3-cm left upper lobe nodule with decreased attenuation centrally. Laboratory studies show hemoglobin, 14.5 g/dL; platelet count, 211,400/mm³; and WBC count, 9890/mm³ with 40% segmented neutrophils, 2% bands, 40% lymphocytes, and 18% monocytes. Which of the following findings in his sputum sample is most likely to be present?

A. Acid-fast bacilli
B. Branching septate hyphae
C. Charcot-Leyden crystals
D. Foreign body giant cells
E. Gram-negative bacilli
F. Small dark neoplastic cells

A previously healthy, 20-year-old woman has had a low-grade fever for the past 2 weeks. On physical examination, her temperature is 37.7°C; there are no other remarkable findings. The gross appearance of the lung shown in the figure is representative of her disease. Which of the following laboratory studies is most likely to report a positive result?

A. Anticentromere antibody
B. HIV serologic test
C. Interferon-γ release assay
D. Rapid plasma reagin
E. Rheumatoid factor
F. Sweat chloride

A 46-year-old man from northern Mexico has had fever, nonproductive cough, and weight loss for 2 months. On examination his temperature is 37.5°C. A chest radiograph shows a miliary pattern of small nodules in all lung fields. Bronchoalveolar lavage is performed and microscopic examination of the fluid shows organisms averaging 50 microns in diameter with thick walls and filled with endospores. Which of the following infections is he most likely to have?

A. Blastomycosis
B. Coccidioidomycosis
C. Histoplasmosis
D. Mycobacteriosis
E. Nocardiosis
F. Paracoccidioidomycosis
A 56-year-old man has had fever, night sweats, and a 3-kg weight loss over the past 4 months. In the past month, he has had episodes of hemoptysis. On physical examination there are upper lobe rales. He has hypoxemia. The appearance of his chest radiograph is shown in the figure. He is most likely to have an infection with which of the following organisms?

A  Candida albicans  
B  Influenza A  
C  Legionella pneumophila  
D  Mycobacterium tuberculosis  
E  Mycoplasma pneumoniae  
F  Nocardia asteroides

A 56-year-old man is undergoing chemotherapy for leukemia. He has developed fever, nonproductive cough, dyspnea, pleuritic chest pain, and hemoptysis over the past week. A chest CT scan shows multiple 1- to 4-cm nodular densities having surrounding areas of ground-glass infiltrate (halo sign). Bronchoalveolar lavage is performed, and microscopic examination of the fluid shows narrow branching septate hyphae. A CBC shows Hgb, 13 g/dL; Hct, 38.7%; WBC count, 2000/μL; and platelet count, 200,100/μL. He has most likely developed an infection with which of the following organisms?

A  Candida albicans  
B  Aspergillus fumigatus  
C  Cryptococcus neoformans  
D  Moraxella catarrhalis  
E  Mycoplasma pneumoniae  
F  Nocardia asteroides

A 47-year-old woman known to be HIV-positive has had a decreasing CD4+ lymphocyte count for the past year despite antiretroviral therapy. She has developed a fever with nonproductive cough over the past week. On auscultation of her chest fine crackles are present in both lungs. A chest radiograph shows infiltrates in both lungs. A transbronchial biopsy is obtained and the microscopic appearance is shown in the figure. Which of the following organisms is most likely infecting this woman?

A  Candida albicans  
B  Cryptococcus neoformans  
C  Cytomegalovirus  
D  Klebsiella pneumoniae  
E  Pneumocystis jiroveci

A 43-year-old woman has had malaise and an 8-kg weight loss over the past 3 years. She has had fever and a nonproductive cough with increasing dyspnea for the past 3 days. On physical examination, her temperature is 37.8° C. There is dullness to percussion over the lungs and diffuse crackles on auscultation. A chest radiograph shows extensive bilateral infiltrates. Bronchoalveolar lavage is done, and the fluid is stained with Gomori methenamine silver, with high-power microscopic appearance shown in the figure. Which of the following underlying conditions is most likely present in this woman?

A  Centrilobular emphysema  
B  Diabetes mellitus  
C  HIV infection  
D  Sarcoidosis  
E  Severe combined immunodeficiency  
F  Systemic lupus erythematosus
65 | A 64-year-old man, who is a chain-smoker, has had a cough and a 5-kg weight loss over the past 3 months. Physical examination shows clubbing of the fingers. He is afebrile. A chest radiograph shows no hilar adenopathy, but there is cavitation within a 3-cm lesion near the right hilum. Laboratory studies show a serum calcium of 12.3 mg/dL, phosphorus of 2.4 mg/dL, and albumin of 3.9 g/dL. Bronchoscopy shows a lesion almost occluding the right main bronchus. A surgical procedure with curative intent is attempted. Which of the following neoplasms is most likely to be present in this patient?

A | Adenocarcinoma in situ
B | Kaposi sarcoma
C | Large cell anaplastic carcinoma
D | Metastatic renal cell carcinoma
E | Non-Hodgkin lymphoma
F | Small cell anaplastic carcinoma
G | Squamous cell carcinoma

66 | A 79-year-old woman has had increasing malaise and a 5-kg weight loss over the past 5 months. She has had a chronic nonproductive cough for 3 months. On physical examination, she has no abnormal findings. Pulmonary function tests are normal. Her peripheral blood counts are normal. Her chest radiograph is shown in the figure. What is a biopsy of one of her lung lesions most likely to show?

A | Adenocarcinoma
B | Granulomatous inflammation
C | Necrotizing vasculitis
D | Organizing abscess
E | Silica crystals

67 | A 45-year-old woman, a nonsmoker, has had a chronic nonproductive cough for 6 months along with an 8-kg weight loss. On physical examination there are no remarkable findings. Her chest radiograph shows a right peripheral subpleural mass. A fine-needle aspiration biopsy is performed, and she undergoes a right lower lobectomy. The microscopic examination of the lesion shows glands invading the surrounding lung. Which of the following molecular test findings is most useful in deciding if her cancer may benefit from therapy targeting a tyrosine kinase?

A | Amplification of FGFR1 gene
B | Inactivation of CDKN2A gene
C | Loss of both copies of TP53
D | Mutation in K-RAS gene
E | Rearrangement of ALK gene

68 | A 60-year-old woman has had a chronic nonproductive cough for 4 months along with loss of appetite and a 6-kg weight loss. She does not smoke. On physical examination there are no remarkable findings. Her chest radiograph shows a right peripheral subpleural mass. A fine-needle aspiration biopsy is performed, and she undergoes a right lower lobectomy. The microscopic appearance of the lesion is shown in the figure. She receives immunotherapy directed at epithelial growth factor receptor (EGFR) and remains symptom-free for the next 10 years. Which of the following neoplasms did she most likely have?

A | Adenocarcinoma
B | Bronchial carcinoid
C | Hamartoma
D | Large cell carcinoma
E | Small cell anaplastic carcinoma
F | Squamous cell carcinoma

69 | A 50-year-old man has developed truncal obesity, back pain, and skin that bruises easily over the past 5 months. On physical examination, he is afebrile, and his blood pressure is 160/95 mm Hg. A chest radiograph shows an ill-defined, 4-cm mass involving the left hilum of the lung. Cytologic examination of bronchial washings from bronchoscopy shows round epithelial cells that have the appearance of lymphocytes but are larger. The patient is told that, although his disease is apparently localized to one side of the chest cavity, surgical treatment is unlikely to be curative. He also is advised to stop smoking. Which of the following neoplasms is most likely to be present in this patient?

A | Adenocarcinoma
B | Bronchial carcinoid
C | Bronchioloalveolar carcinoma
D | Large cell carcinoma
E | Non-Hodgkin lymphoma
F | Small cell anaplastic carcinoma
G | Squamous cell carcinoma
70 A 57-year-old woman has had a cough and pleuritic chest pain for the past 3 weeks. On physical examination, she is afebrile. Some crackles are audible over the left lower lung on auscultation. A chest radiograph shows an ill-defined area of opacification in the left lower lobe. After 1 month of antibiotic therapy, her condition has not improved, and the lesion is still visible radiographically. CT-guided needle biopsy of the left lower lobe of the lung is performed, and the specimen has the histologic appearance shown in the figure. Which of the following neoplasms is most likely to be present in this patient?

A Adenocarcinoma in situ  
B Large cell anaplastic carcinoma  
C Malignant mesothelioma  
D Metastatic breast carcinoma  
E Squamous cell carcinoma

71 A 59-year-old man who has smoked one pack of cigarettes per day for the past 43 years has developed a severe cough with hemoptysis over the past month. He has experienced a 10-kg weight loss over the past year. On physical examination, he is afebrile. Laboratory studies show a serum Na⁺ of 120 mmol/L, K⁺, 3.8 mmol/L, Cl⁻, 90 mmol/L, CO₂, 24 mmol/L; glucose, 75 mg/dL; creatinine, 1.2 mg/dL; calcium, 8.1 mg/dL; phosphorus, 2.9 mg/dL; and albumin, 4.2 g/dL. Which of the following findings is most likely to be seen on a chest radiograph?

A Bilateral upper lobe cavitation  
B Diaphragmatic pleural calcified plaques  
C Extensive areas of infiltrates  
D Invasive perihilar mass  
E Pneumothorax  
F Subpleural nodule with hilar adenopathy  
G Upper lung nodule with air-fluid level

72 A 72-year-old woman has had difficulty with vision in her right eye for 3 months. She also has pain in the right upper chest. The findings on physical examination include unilateral enophthalmos, miosis, anhidrosis, and ptosis on the right side of her face. A chest radiograph shows right upper lobe opacification and bony destruction of the right rib. Which of the following conditions is most likely to be present in her?

A Bronchopneumonia  
B Bronchiectasis  
C Bronchogenic carcinoma  
D Sarcoïdosis  
E Tuberculosis

73 A 43-year-old woman has never smoked and works as a file clerk at a university that designates all work areas as non-smoking. A routine chest radiograph shows a 3-cm, sharply demarcated mass in the left upper lobe of the lung. Fine-needle aspiration of the mass is attempted, but the pathologist performing the procedure remarks, “This is like trying to biopsy a ping-pong ball.” No tissue is obtained. Thoracotomy with wedge resection is performed. On sectioning, the mass has a firm, glistening, bluish white cut surface. A culture of the mass yields no growth. This mass most likely represents which of the following?

A Adenocarcinoma  
B Hamartoma  
C Large cell carcinoma  
D Mesothelioma  
E Non-Hodgkin lymphoma  
F Squamous cell carcinoma

74 A 40-year-old man has had an increasing cough with hemoptysis for 2 weeks. He has never smoked and is in very good health. On physical examination, his temperature is 38.2° C. A chest radiograph shows an area of consolidation in the right upper lobe. His condition improves with antibiotic therapy; however, the cough and hemoptysis persist for 2 more weeks. Chest CT scan now shows right upper lung atelectasis. Bronchoscopic examination shows a tan, circumscribed obstructive mass filling a right upper lobe bronchus. Which of the following neoplasms is most likely to produce these findings?

A Adenocarcinoma  
B Carcinoid tumor  
C Hamartoma  
D Kaposi sarcoma  
E Large cell carcinoma

75 A 24-year-old man has had increasing dyspnea for the past 10 weeks. On physical examination, he is afebrile. There is dullness to percussion over the lungs posteriorly and decreased breath sounds. A chest radiograph shows large bilateral pleural effusions and widening of the mediastinum. Thoracentesis is performed on the left side and yields 500 mL of milky white fluid. Laboratory studies of the fluid show a high protein content; microscopy shows many lymphocytes and fat globules. What is the most likely cause for these findings?

A Bacterial pneumonia with empyema  
B Congenital heart disease with congestive failure  
C Marfan syndrome with aortic dissection  
D Micronodular cirrhosis with hypoalbuminemia  
E Miliary tuberculosis with granulomatous pleuritis  
F Non-Hodgkin lymphoma with lymphatic obstruction
A 68-year-old man has had increasing dyspnea with cough productive of frothy sputum for the past 5 months. On physical examination, he is afebrile, and his blood pressure is 165/100 mm Hg. There is dullness to percussion at lung bases. He has pitting edema of the ankles. A chest radiograph shows blunting of costophrenic recesses bilaterally and cardiomegaly with prominent right and left heart borders. A right thoracentesis is performed, and 300 mL of straw-colored fluid is removed. Laboratory studies on this fluid show total protein of 2.2 g/dL (serum is 6.5 g/dL), glucose of 45 mg/dL, lactate dehydrogenase of 200 U/L (serum is 420 U/L), pH 7.2, and cell count of 100/mm3 mononuclear leukocytes, and no RBCs. What condition does he most likely have?

A  Cavitary tuberculosis
B  Congestive heart failure
C  Malignant mesothelioma
D  Non-Hodgkin lymphoma
E  Pneumococcal pneumonia
F  Small cell carcinoma

A 47-year-old woman, a non-smoker, has a 4-month history of mild but persistent right-sided chest pain. On physical examination, there are no remarkable findings. A chest radiograph shows a pleural mass on the right side. No pleural effusions are seen. Chest CT scan shows a localized, circumscribed 3 × 7 cm mass attached to the visceral pleura; the lungs and chest wall appear normal. At thoracotomy, the mass is excised. On microscopic examination, the mass is composed of spindle cells resembling fibroblasts with abundant collagenous stroma. With immunohistochemical staining, the spindle cells mark for CD34, but are cytokeratin-negative. There has been no recurrence of the lesion. Which of the following is the most likely diagnosis?

A  Bronchioloalveolar carcinoma
B  Hamartoma
C  Hodgkin lymphoma, nodular sclerosis type
D  Malignant mesothelioma
E  Metastatic breast carcinoma
F  Solitary fibrous tumor

Inhalation of which of the following pollutants is the most likely factor in the pathogenesis of this mass?

A  Asbestos
B  Bird dust
C  Coal dust
D  Cotton fibers
E  Ozone
F  Silica

A 78-year-old man has had increasing dyspnea without cough or increased sputum production for the past 4 months. On physical examination, he is afebrile. Breath sounds are reduced in all lung fields. A chest CT scan shows a dense, brightly attenuated pleural mass encasing most of the left lung. Microscopic examination of a pleural biopsy specimen shows spindle and cuboidal cells that invade adipose tissue.
worse than it feels, and within days the air is resorbed. When the leak occurs centrally, the term mediastinal emphysema can be used. The term pulmonary interstitial emphysema (PIE) describes this process in newborns, often with mechanical ventilation under positive pressure, and in the setting of respiratory distress syndrome. Rupture of the stomach is most likely to leak gastric acidic contents into the peritoneum to produce peritonitis, and be recognized by free air under the diaphragm on radiographs. Blood in the pleural space is called hemothorax; within soft tissues blood can form a hematoma. Leakage of lymph is rare, because the lymphatics have little or no pressure within them; blockage of the thoracic duct may produce a chylothorax. Pus in the pleural space is called empyema, and typically complicates an existing pneumonia.

**5 D** He has left-sided heart failure with pulmonary edema and congestion. His reduced cardiac output leads to diminished renal blood flow that stimulates the renin-angiotensin mechanism to retain salt and water to increase plasma volume. He has prerenal azotemia with a high BUN-to-creatinine ratio and low fractional excretion of sodium. The other options do not explain his pulmonary edema. An aldosterone-secreting adenoma (Conn syndrome) would increase aldosterone, but decrease the plasma renin. In chronic adrenal failure (Addison disease), there should be hyperkalemia and hypoglycemia accompanying hyponatremia. Chronic glomerulonephritis with chronic renal failure would be associated with a BUN-to-creatinine ratio around 10:1. Pulmonary fibrosis would lead to cor pulmonale and a prominent right heart border with features of right-sided congestive heart failure. The syndrome of inappropriate antidiuretic hormone (ADH) is a paraneoplastic syndrome that can occur with pulmonary small cell carcinomas, and secretion of antidiuretic hormone (SIADH) would increase ADH and cause more severe hyponatremia, but would not have a major effect on the renin-angiotensin mechanism, and sodium excretion would be higher.

**6 A** She has acute lung injury with noncardiogenic pulmonary edema and development of diffuse alveolar damage (DAD), clinically known as acute respiratory distress syndrome (ARDS). Inciting sepsis, trauma, or other forms of lung injury leads to a vicious cycle of inflammation with ongoing damage, mainly through the action of neutrophils. Plexiform lesions are characteristic for pulmonary hypertension. Though ARDS may eventually proceed to fibrosis, most patients do not survive that long. Lymphocytic infiltrates may be seen with infections such as viral pneumonias or immune-mediated lung diseases. Destruction of respiratory bronchioles is a feature of centrilobular emphysema.

**7 C** This clinical and morphologic picture is that of acute lung injury (ALI), which, when severe, leads to acute respiratory distress syndrome (ARDS). ARDS is characterized by the pathologic finding of diffuse alveolar damage (DAD), which is initiated in most cases by injury to capillary endothelium, with neutrophils and macrophages that aggregate in alveolar capillaries and release toxic oxygen metabolites, cytokines, and eicosanoids. Oxygen toxicity from high levels of inspired oxygen exacerbate DAD. The damage to the capillary endothelium allows leakage of protein-rich fluids. Eventually, the overlying alveolar epithelium also is damaged. Aspiration of bacteria causes bronchopneumonia. ARDS and disseminated intravascular coagulation (DIC) together can complicate septic shock, but DIC is not the cause of ARDS. Release of fibrogenic cytokines is an important cause of chronic diffuse pulmonary fibrosis. Reduced surfactant production causes respiratory distress syndrome with hyaline membrane disease in newborns.

**8 E** The patient’s findings are predominantly those of an obstructive lung disease—emphysema—with a centrilobular pattern of predominantly upper lobe involvement. Smoking is a major cause of this disease. The subtle but long-term inflammation that can accompany smoking leads to increased neutrophil and macrophage elaboration of elastase that is not sufficiently inhibited by the antiprotease action of α1-antitrypsin. This results in a loss of lung tissue, not fibrogenesis, over decades. Fibrogenesis is typical of restrictive lung diseases, such as pneumoconioses, that follow inhalation of dusts. Abnormal chloride ion transport is a feature of cystic fibrosis, which leads to widespread bronchiectasis. Dynein arms are absent or abnormal in Kartagener syndrome, which leads to bronchiectasis. α1-Antitrypsin deficiency is uncommon and leads to a panlobular pattern of emphysema. Macrophage recruitment and activation by interferon-γ released from T cells is a feature of chronic inflammatory conditions and pneumoconioses.

**9 E** Distal acinar (paraseptal) emphysema is localized, beneath pleura typically in an upper lung lobe, and may occur in an area of fibrosis or scar formation. Although the lesions are usually less than 2 cm in diameter, they are prone to rupture spontaneously or with minor trauma, leading to pneumothorax. They can be a cause for spontaneous pneumothorax in young adults. A “ball valve” effect can lead to air trapping in pleura, producing tension pneumothorax, as in this case. Centriacinar emphysema arises in respiratory bronchioles and is seen in smokers. Panacinar (panlobular) emphysema involves most of the lung lobule and can be seen in all lobes; α1-antitrypsin deficiency is the most likely antecedent. Asthma results from bronchoconstriction with air trapping, but is not likely to be complicated by pneumothorax. Bronchiectasis results from inflammation with destruction of bronchi; hemoptysis is the most likely complication, not pneumothorax. Chronic bronchitis is unlikely to produce a bronchopleural fistula with pneumothorax.

**10 A** These findings point to an obstructive lung disease, such as emphysema, which occurs with airway narrowing and loss of elastic recoil. It has led to compensated respiratory acidosis in this man. Chronic pulmonary embolism does not
11 E The extensive pulmonary involvement with increased lucency in all lung fields and increased anterior-posterior diameter is consistent with emphysema. The panlobular form, which can be worse in the lower lobes, can be due to a decrease in α1-antitrypsin, which is the major circulating antielastase. This deficiency is an inherited disease, typically with the PiZZ genotype; liver disease may also occur. Individuals with atopy are more likely to develop asthma, which has transient air trapping, not emphysema. The CFTR gene mutations lead to cystic fibrosis and widespread pulmonary bronchiectasis, starting in childhood. Smoking increases inflammation with neutrophils releasing proteases, mainly in upper lobes, producing the centriacinar pattern of emphysema over decades. Prior infection with tuberculosis may result in upper lobe cavitation, not emphysema.

12 C This patient’s disease meets the clinical definition of chronic bronchitis. He has had persistent cough with sputum production for at least 3 months in 2 consecutive years. Chronic bronchitis is a disease of smokers and individuals living in areas of poor air quality, which explains the chronic cough with mucoid sputum production. This patient’s episodes of bronchoconstriction set off by viral infections suggest, however, a superimposed element of nonatopic asthma. Cor pulmonale leads to pleural effusions, not to bronchoconstriction. Centrilobular emphysema and chronic bronchitis (both complications of smoking) can overlap in clinical and pathologic findings, but significant bronchoconstriction is not a feature of emphysema. The panlobular emphysema of α1-antitrypsin can be worsened by smoking, but there is no bronchoconstriction. Bronchiectasis results in airway dilation from destructive bronchial wall inflammation, but the onset of pulmonary disease with cystic fibrosis is typically in childhood. Hypersensitivity pneumonitis is marked by features of a restrictive lung disease, sometimes with dyspnea, but without mucus production, and is often episodic from intermittent antigen exposure.

13 D Centrilobular emphysema results from damage to the central part of the lung acinus, with dilation that primarily affects the respiratory bronchioles. There is relative sparing of the distal alveolar ducts and alveolar sacs. Bronchi have cartilage that is not affected by emphysema. In panacinar emphysema, the lung lobule is involved from the respiratory bronchiole to the terminal alveoli. In paraseptal emphysema, the distal acinus is involved.

14 D Chronic bronchitis can be complicated by pulmonary hypertension and cor pulmonale. There are few characteristic microscopic features of chronic bronchitis, so it is mainly defined clinically by the presence of a persistent cough with sputum production for at least 3 months in at least 2 consecutive years. Chronic bronchitis does not lead to diffuse pulmonary fibrosis. Granulomatous disease is more typical of sarcoidosis or mycobacterial infection. Lymphangitic metastases may fill lymphatic spaces and produce a reticulonodular pattern on a chest radiograph, but patients tend not to live long with such advanced cancer. Increased eosinophils are characteristic of bronchial asthma, which is an episodic disease unlikely to cause cor pulmonale.

15 E Atopic asthma is a type I hypersensitivity reaction in which there are presensitized, IgE-coated mast cells in mucosal surfaces and submucosa of airways. Contact with an allergen results in degranulation of the mast cells, with both immediate release (minutes) of mediators such as histamine to promote bronchoconstriction, and delayed release (an hour or more) of leukotrienes and prostaglandins via the arachidonic acid pathway; these attract leukocytes, particularly eosinophils, and promote bronchoconstriction. The characteristic histologic changes in the bronchi, including remodeling of airways and smooth muscle hyperplasia, result from the episodes of inflammation. Dilation of the respiratory bronchiole is a feature of centrilobular emphysema. Bronchial dilation with inflammatory destruction is a feature of bronchiectasis. Hyaline membranes are seen with acute diffuse alveolar damage. Neutrophilic exudates with consolidation are seen in pneumonic processes, typically from bacterial infections.

16 A Asthma, particularly extrinsic (atopic) asthma, is driven by a type I hypersensitivity response and is associated with an excessive Th2 and Th17 cell-mediated immune response. Genetic factors are important in the pathogenesis of atopic asthma and linkage to cytokine genes that map on 5q are strongly associated with development of asthma and other atopic allergies. The Charcot-Leyden crystals represent the breakdown products of eosinophil granules. The Curschmann spirals represent the whorls of sloughed surface epithelium within the abundant mucin. The septated hyphae are Aspergillus organisms colonizing the tracheobronchial tree (allergic bronchopulmonary aspergillosis). Foreign body aspiration may result in inflammation, but without eosinophils. Inorganic dust inhalation leads to restrictive, not obstructive, lung disease. CFTR mutations with cystic fibrosis lead to chronic widespread bronchiectasis. Inflammation with eosinophils is not a significant component of emphysema related to α1-antitrypsin deficiency or to smoking.

17 C The early, acute phase of bronchial asthma is triggered by release of chemical mediators, whereas the late phase is mediated by recruited inflammatory cells and the Th2 cytokines they release. Acute asthmatic episodes respond best to inhaled β-adrenergic agonists. Histamine released from mast cells acts during the early acute phase of type I hypersensitivity.
reactions, but antihistaminic agents are not useful for treating recurrent bouts of asthma. Th2 cytokines play an important role in recurrent asthma and antagonists of these, in particular IL-13 and IL-4, are in development. It is not clear if any one of the Th2 cytokines alone mediates recurrent bronchospasm. Among the early to late phase mediators, the leukotrienes C₄, D₄, and E₄ promote intense bronchoconstriction and mucin production. Montelukast is an agent that binds to cysteinyl leukotriene (CysLT) receptors on mast cells and eosinophils to block the lipoxygenase pathway of arachidonic acid metabolism, which generates the leukotrienes. Prostaglandin D₂ also is a bronchoconstrictor, but its role is less well defined than that of leukotrienes.

18 B This history is typical of nonatopic, or intrinsic, asthma. There is no family history of asthma, no eosinophilia, and a normal serum IgE level. The fundamental abnormality in such cases is bronchial hyperresponsiveness (i.e., the threshold of bronchial spasm is intrinsically low). When airway inflammation occurs after viral infections, the bronchial smooth muscle spasms, and an asthmatic attack occurs. Such bronchial hyperreactivity also may be triggered by inhalation of air pollutants, such as ozone, sulfur dioxide, and nitrogen oxides. Even exercise and cold air may act as a trigger. Accumulation of neutrophils is typical of bacterial pneumonia, which could follow viral infection, but lead to lung consolidation. Bronchopulmonary aspergillosis refers to colonization of asthmatic airways by *Aspergillus*, which is followed by development of additional IgE antibodies. Secretion of interleukin (IL)-4 and IL-5 by type 2 helper T cells also occurs in cases of allergic asthma.

19 E The Th₂ helper lymphocyte response drives cytokine production, such as IL-4, IL-5, and IL-13, that promotes eosinophil infiltration and IgE production by mast cells. This allergic response potentiates inflammation, which promotes the airway remodeling that facilitates additional airway reactivity and asthmatic episodes. The Th₁ response drives granulomatous inflammation. Th₁/Th₁₇ lymphocytes aid in inflammatory responses to infectious agents, but may play a role in autoimmunity. B lymphocytes produce antibodies, but mainly via the action of T helper cells. Cytotoxic lymphocytes are primarily directed at intracellular infectious agents. NK cells assist in fighting infectious agents.

20 B Drug-induced asthma is most likely to occur in older patients who develop increased sensitivity to a drug. Aspirin (acetylsalicylic acid) blocks the cyclooxygenase pathway of arachidonic acid metabolism but not the lipoxygenase pathway that potentiates bronchoconstriction. NSAIDs may have the same effect as aspirin. Angiotensin-converting enzyme (ACE) inhibitors may also induce asthmalike episodes. Acetaminophen is an analgesic that can be substituted for aspirin and is unlikely to provoke asthmatic attacks. Gaba pentin and morphine act centrally as analgesics. Prednisone is an anti-inflammatory agent that is used to treat immune-mediated diseases.

21 D He has Kartagener syndrome (sinusitis, bronchiectasis, and situs inversus associated with ciliary dyskinesia). There is an abnormality of ciliary dynein arms that diminishes the mucociliary function of the respiratory epithelium, predisposing to recurrent and chronic infections of both upper and lower respiratory tract. Bronchiectasis is ongoing destruction and dilation of bronchi with infection and airway obstruction. α₁-Antitrypsin deficiency leads to panlobular emphysema, mainly of lower lobes, and the upper respiratory tract is not involved. Atopy may be associated with nasal polyps, but leads to asthma, not bronchiectasis. Cystic fibrosis with CFTR gene mutations involving chloride ion channels can lead to widespread bronchiectasis, but generally not upper airway problems, and not situs inversus. HIV infection is marked by opportunistic infections with progression to AIDS, but usually without bronchiectasis, and no situs inversus.

22 D Bronchiectasis is a chronic obstructive airway disease from irreversible dilation of bronchi that results from inflammation and destruction of bronchial walls after prolonged infections or obstruction. Serious bouts of pneumonia can predispose to bronchiectasis. Congenital chondromalacia weakening the bronchial wall is rare. Bronchial mucosal damage by eosinophils occurs in bronchial asthma. It does not cause destruction of the bronchial wall. Bronchioalveolar carcinoma may mimic an infiltrative pneumonia because of its lepiderc pattern of spread, but it mainly produces a mass effect, and it does not start in childhood. Unopposed action of elastases damages the elastic tissue of alveoli, giving rise to emphysema.

23 D The pulmonary function data suggest a restrictive lung disease process. The progressive pulmonary interstitial fibrosis of a restrictive lung disease such as a pneumococcosis can eventually lead to dilation of remaining residual proximal airspaces, giving a honeycomb appearance. The loss of lung tissue with emphysema also leads to airspace dilation, but without alveolar wall fibroogenesis. Eosinophil infiltrates suggest atopic asthma, an episodic disease without fibrogenesis. The increase in mucous glands with chronic bronchitis leads to copious sputum production, but not fibrogenesis. Hyaline membranes, edema, inflammation, and focal necrosis are features of diffuse alveolar damage (acute respiratory distress syndrome) in the acute phase; if patients survive for weeks, diffuse alveolar damage may resolve to honeycomb change.

24 C The cause of many slowly progressive cases of restrictive lung disease is unknown, and the frustrated but empathetic health care provider can only say, “I am sorry,
but there is nothing more that we can do." These cases must be distinguished from cases with identifiable causes, such as infection, collagen vascular disease, drug use, and pneumoconioses. Idiopathic pulmonary fibrosis leads to progressive restrictive lung disease. An unknown antigen incites the T(H)2 inflammatory process with activated macrophage release of cytokines such as fibroblast growth factor and TGF-β1. TGF-β1 down-regulates telomerase activity and leads to epithelial cell apoptosis. TGF-β1 diminishes caveolin-1, a protein that inhibits fibrosis. Goodpasture syndrome is a form of type II hypersensitivity characterized by diffuse pulmonary hemorrhage superimposed upon normal lung. Hypersensitivity pneumonitis may have an element of type IV hypersensitivity with some fibrosis, but usually not as severe or rapid as in this case. Sarcoidosis is marked by granulomatous inflammation. In this case, scleroderma is less likely because of the negative serologic test result.

PBD9 685–687  BP9 472–473  PBD8 694–695  BP8 494–495

25 E The inflammatory response to lung injury may determine the nature and extent of the pulmonary disease. The cytokine TGF-β may modulate the mesenchymal cell response to lung injury. Reduced TGF-β signaling may lead to an inadequate repair response, with loss of lung parenchyma that characterizes emphysema. The PiZZ genotype of α1-antitrypsin deficiency puts persons, especially smokers with greater tobacco use, at increased risk for panlobular emphysema. BMPR2 is associated with development of primary pulmonary hypertension. Mutations in the CFTR gene are associated with development of cystic fibrosis. Mutations in GM-CSF are related to development of pulmonary alveolar proteinosis.

PBD9 686–687  BP9 473  PBD8 494–495  BP8 494–495

26 E Nonspecific interstitial pneumonia has both cellular and fibroblastic patterns, but the former has a better prognosis. Some patients have an underlying connective tissue disorder. Desquamative interstitial pneumonitis (DIP) is smoking-related. Hypersensitivity pneumonitis most often relates to episodic inhaled allergens and rarely progresses to marked interstitial disease. Idiopathic pulmonary fibrosis tends to have a more rapid course and involve more of the lungs. Nonatopic asthma is typically episodic and rarely progresses to extensive interstitial disease.

PBD9 686–687  BP9 472–473  PBD8 695  BP8 495–496

27 B Bronchiolitis obliterans is a feature of cryptogenic organizing pneumonia, an uncommon, nonspecific reaction to a lung injury, such as an infection or toxic exposure. Bronchiectasis involves ongoing inflammatory destruction with dilation of bronchi not reversed by corticosteroids. Desquamative interstitial pneumonitis (DIP) is an uncommon smoking-related interstitial disease in which monocytes gather to form intra-alveolar macrophages; DIP is not related to idiopathic pulmonary fibrosis. Hypersensitivity pneumonitis is a type III (and type IV) hypersensitivity response to an inhaled allergen. Pulmonary alveolar proteinosis is a rare idiopathic condition in which there are gelatinous alveolar proteinaceous exudates.

PBD9 687  BP9 473–474  PBD8 696, 720  BP8 496

28 C Silica crystals incite a fibrogenic response after inhalation and ingestion by pulmonary macrophages. The greater the exposure to silica dust and the longer the time of exposure, the greater is the lung injury. Silica is a major component of the earth’s crust, including sand, which contains the mineral quartz. Mining, manufacturing, farming, and construction/renovation activities generate small silica crystals that can be inhaled, and their buoyancy allows them to be carried to alveoli. There, they are ingested by macrophages, which secrete cytokines that recruit other inflammatory cells and promote fibrogenesis. Carbon monoxide readily crosses the alveolar walls and binds avidly to hemoglobin, but does not injure the lung directly. Ozone, a component of smog, has no obvious pathologic effects. Tobacco smoke leads mainly to loss of lung tissue and emphysema, not to fibrosis. Particulate matter such as wood dust is mainly screened out by the mucociliary apparatus of the upper airways, but may invoke bronchoconstriction.

PBD9 688, 690  BP9 474–476  PBD8 698–699  BP8 498–499

29 C Coal worker’s pneumoconiosis (CWP) is now less common because of workplace safeguards in mining. Coal dust is relatively inert, so that large amounts must be inhaled before a fibrogenic response occurs, but the response continues over many years. The other listed substances are more reactive. Asbestos produces ferruginous bodies. Chronic berylliosis tends to be associated with sarcoid-like granulomas. Iron produces siderosis with fibrosis. Sulfur dioxide is a gas that contributes to obstructive lung diseases, particularly chronic bronchitis and asthma.

PBD9 689  BP9 474–476  PBD8 697–698  BP8 497–498

30 B The ferruginous bodies shown in the figure are long, thin crystals of asbestos that have become encrusted with iron and calcium. The inflammatory reaction incited by these crystals promotes fibrogenesis and resultant pneumoconiosis. Anthracosis is a benign process seen in city dwellers as a consequence of inhaled carbonaceous dust. Chronic berylliosis is marked by noncaseating granulomas. Calcium deposition may rarely occur along alveolar walls when the serum calcium level is very high (metastatic calcification). silica crystals are not covered by iron and tend to result in formation of fibrous nodules (silicotic nodules).

PBD9 690–692  BP9 477–478  PBD8 700  BP8 499–500

31 C Silicotic nodules form when the silica crystals ingested by macrophages elicit a fibrogenic response as cytokines, such as tumor necrosis factor, are released. The nodules may become confluent with progressive massive fibrosis. Pneumoconioses such as silicosis lead to restrictive lung disease. Cigarette smoke contributes to loss of lung parenchyma with emphysema. Mold spores tend to elicit a hypersensitivity pneumonitis
that rarely goes on to extensive restrictive lung disease. Sulfur dioxide in polluted air tends to drive chronic obstructive lung disease. Wood dust tends to elicit an asthmatic response.

PBD9 690  BP9 476  PBD8 698–699  BP8 498–499

32 F She has a restrictive lung disease; the figure shows hilar adenopathy and a reticulonodular pattern of infiltrates characteristic of sarcoid. The blood gas values show mild hypoxemia with a compensated respiratory alkalosis. Because there is no obstructive disease, the CO₂ can be normal or low from increased respirations to compensate for the diffusion block (with low DLCO). α₁-Antitrypsin deficiency and chronic bronchitis lead to obstructive lung disease with increased total lung capacity and diminished FEV₁ and respiratory acidosis. Chronic bronchitis is defined by prolonged sputum production. Diffuse alveolar damage is an acute restrictive lung disease from a severe underlying injury, such as sepsis. Goodpasture syndrome results in pulmonary hemorrhage with hemoptysis. Asthma is an acute obstructive lung disease with dyspnea and wheezing.

PBD9 693–694  BP9 478–480  PBD8 701–703  BP8 501–502

33 C This patient has an occupational risk of asbestos exposure. The inhaled asbestos fibers become encrusted with iron and appear as the characteristic ferruginous bodies with iron stain. The firm, tan mass encasing the pleura is most likely a malignant mesothelioma. Asbestosis more commonly gives rise to pleural fibrosis and interstitial lung disease, similar to other pneumoconioses. This is seen grossly as a dense pleural plaque, which often is calcified. Asbestosis can give rise to bronchogenic carcinoma, especially in smokers. Fluffy infiltrates suggest an infectious process. Upper lobe cavitation suggests secondary tuberculosis. An endobronchial mass could be a carcinoid tumor, which is not related to asbestosis. The pleural mass likely leads to obliteration of the pleural space, with no effusion.

PBD9 690–692  BP9 477–478  PBD8 699–701  BP8 499–500

34 A The clinical and morphologic features strongly suggest sarcoidosis. This granulomatous disease has an unknown cause, but the presence of granulomas and activated T cells in the lungs indicates a delayed hypersensitivity response to some inhaled antigen. Lung involvement, occurring in about one third of cases, may be asymptomatic or may lead to restrictive lung disease. Sarcoidosis can involve multiple organs, particularly those of the mononuclear phagocyte system, especially lymph nodes. Diffuse alveolar damage is an acute lung injury seen in acute respiratory distress syndrome. Hypersensitivity pneumonitis is an immune complex disease that is triggered by inhaled allergens. This form of lung disease is characterized by acute dyspneic episodes. The disease starts with type III hypersensitivity, but there can sometimes be granulomas in the lung, and lymph node enlargement is not seen. Atypical mycobacteria cause caseating granulomas, as does Mycobacterium tuberculosis. Smoking causes chronic bronchitis and emphysema.

PBD9 693–694  BP9 478–480  PBD8 701–703  BP8 501–502

35 C Farmer’s lung is a form of hypersensitivity pneumonitis caused by inhalation of actinomycete spores in moldy hay. These spores contain the antigen that incites the hypersensitivity reaction. Because type III (early) and type IV immune hypersensitivity reactions are involved, granuloma formation can occur. The disease abates when the patient is no longer exposed to the antigen. Chronic exposure can lead to more extensive interstitial lung disease. Antibodies directed against pulmonary basement membrane are a feature of Goodpasture syndrome, which mainly produces pulmonary hemorrhage. Silicosis can produce a restrictive lung disease with fibrosis, but there are nodules of fibrosis that develop over years with minimal inflammation. Pulmonary tuberculosis can produce granulomas, but the pattern would be miliary, and it is unlikely that it would continue for 15 years. Pneumococci with exposure to dusts such as silica can produce interstitial fibrosis over many years, and the risk of neoplasia is increased slightly for silicosis and greatly for asbestosis.

PBD9 694–695  BP9 480–481  PBD8 703  BP8 503

36 A Hypersensitivity pneumonitis has acute symptoms that occur soon after exposure to an antigen, often actinomycetes or fungi (molds) growing in contaminated HVAC systems (air conditioner or ventilation ducts). The symptoms improve when the patient leaves the environment where the antigen is located. The pulmonary pathologic changes are usually minimal, with interstitial mononuclear infiltrates. It is mainly a type III hypersensitivity reaction, but with more chronic exposure to the antigen, there may be a component of type IV hypersensitivity with granulomatous inflammation and fibrosis. Attachment of antibody to basement membrane occurs in Goodpasture syndrome. Mycolic acid is a component of the cell wall of mycobacteria, and infections with these organisms are chronic, not episodic. Prostaglandins are produced by the cyclooxygenase pathway of arachidonic acid metabolism during acute inflammation, and they mediate pain and vasodilation. Histamine release is characteristic of a type I hypersensitivity reaction that more typically occurs in allergic disease. A toxic injury is more typical of inhalation of a toxic gas, such as sulfur dioxide (so-called silo filler’s disease).

PBD9 694–695  BP9 480–481  PBD8 703–704  BP8 503

37 B He has desquamative interstitial pneumonitis (DIP), one form of smoking-related interstitial lung disease. Most cases abate with cessation of smoking and corticosteroid therapy. Atopy is classically related to asthma, an acute obstructive pulmonary process. One form of primary ciliary dyskinesia is Kartagener syndrome, which leads to bronchiectasis from ongoing inflammation with infection. Inhalation of mold spores produces farmer’s lung—hypersensitivity pneumonitis. Inhalation of silicates leads to pulmonary fibrosis over years, but without large numbers of macrophages.

PBD9 695–696  BP9 481  PBD8 704  BP8 481–482, 504

38 E The acquired form of pulmonary alveolar proteinosis (PAP) is an uncommon condition of unknown etiology characterized by autoantibodies against granulocyte-macrophage...
colony-stimulating factor (GM-CSF). Ten percent of PAP cases are congenital secondary to mutations in the GM-CSF gene. Both forms of PAP have impaired surfactant clearance by alveolar macrophages, leading to accumulation of a gelatinous alveolar exudate. α1-Antitrypsin deficiency leads to panlobular emphysema with hyperlucent lungs on radiographs. CFTR gene mutations lead to cystic fibrosis and widespread bronchiectasis. Anti-DNA topoisomerase I antibodies are seen in diffuse scleroderma, which produces pulmonary interstitial fibrosis. Anti-glomerular basement membrane antibody is present in Goodpasture syndrome with extensive alveolar hemorrhage. Neutrophilic myeloperoxidase is a form of anti-neutrophil cytoplasmic autoantibody seen in ANCA-associated vasculitis.

**39 D** The CT scan shows large filling defects with decreased attenuation within large pulmonary arteries. This lesion is typical of pulmonary thromboembolism that affects patients who are immobilized in the hospital, such as patients with congestive heart failure. The source of the emboli is usually thrombi within deep pelvic or leg veins affected by phlebothrombosis. Patients with underlying cardiac or respiratory diseases that compromise pulmonary circulation are at greater risk of infarction if thromboembolism occurs. Thromboembolism is not a complication of smoking in patients with emphysema or asthma. HIV infection increases the risk of pulmonary infections, but not of thromboembolism. The small emboli from the small vegetations of nonbacterial thrombotic endocarditis are unlikely to produce such large filling defects, and most of them arise on the left side of the heart. Vasculitis of the lung typically involves arterioles, capillaries, or venules of insufficient size to produce a grossly apparent filling defect in large arteries. Pneumoconiosis with restrictive lung disease produce pulmonary fibrosis, but not a compromised vasculature or thromboembolism.

**40 F** The findings in this study suggest pulmonary thromboembolism, and most pulmonary emboli are small and clinically silent. Dyspnea can occur with medium to large emboli. Hemoptysis with pulmonary embolism is uncommon, although it may occur when a hemorrhagic infarction results from thromboembolism. Cor pulmonale can result from repeated embolization with reduction in the pulmonary vascular bed, leading to heart failure and arrhythmias with palpitations, but this is not common. Thromboembolism with infarction based upon the pleura may lead to pleuritic pain. Sudden death may occur with large emboli that occlude the main pulmonary arteries before any other changes can occur. Orthopnea from fluid-filling airspaces most likely accompanies heart failure.

**41 E** The figure shows a saddle pulmonary thromboembolus in the opened main pulmonary arteries. Sudden death occurs from hypoxemia or from acute cor pulmonale with right-sided heart failure. Because the airways are not obstructed, the lungs do not collapse, and there is no mass effect upon lung parenchyma. There is no bronchoconstriction. With such an acute course, there is not enough time for a hemorrhagic pulmonary infarction to occur. Edema is not a feature of thromboembolism.

**42 D** Over half of persons with chronic pulmonary thromboembolism with pulmonary hypertension do not have a history of recurrent pulmonary embolism. Rather than one large life-threatening embolus, chronic thromboembolism typically involves multiple smaller emboli that reduce the pulmonary vascular bed and increase pulmonary pressures, leading to cor pulmonale. Recanalization of thrombi leads to narrowing channels causing the bruits. Anti-phospholipid antibodies pose a risk for thrombosis. The risk factors for systemic arterial atherosclerosis are not operative in the pulmonary arterial tree, and pulmonary atherosclerosis is a consequence of pulmonary hypertension, not a cause of it. Pneumonitis with parenchymal inflammation reduces ventilation more than perfusion. Sarcoidosis is an idiopathic granulomatous disease that mainly affects the pulmonary parenchyma. ANCA-associated vasculitis of the pulmonary arterial tree may produce vascular occlusion, but there is usually parenchymal disease as well, along with multisystem involvement.

**43 A** This patient has findings associated with right and left heart failure. The left-to-right shunt produced by the atrial septal defect leads to increased pulmonary arterial pressure, thickening of the pulmonary arteries, and increased pulmonary vascular resistance. Eventually, the shunt may reverse, which is known as the Eisenmenger complex. The plural effusions are the result of right heart failure from cor pulmonale. The chest radiograph findings here do not include interstitial disease. Granulomatous inflammation does not occur from increased pulmonary arterial pressure, but extensive granulomatous inflammation, as with sarcoidosis, may lead to pulmonary hypertension. An infarction of the lung can occur with pulmonary embolism. Pulmonary interstitial fibrosis can be caused by diseases such as pneumoconioses, collagen vascular diseases, and granulomatous diseases; interstitial lung disease can lead to pulmonary hypertension. Pulmonary necrotizing vasculitis may be seen with immunologically mediated diseases, such as ANCA-associated vasculitis.

**44 B** The finding of pulmonary hypertension in a young individual without any known pulmonary or cardiac disease is typical for primary pulmonary hypertension. The increased pulmonary arterial pressure leads to hypertrophy of the right side of the heart. The large pulmonary arteries show atherosclerosis; the arterioles show plexogenic arteriopathy with a tuft of capillary formations producing a network, or web, that spans the lumens of dilated thin-walled arteries. BMPR2, a cell surface protein belonging to the TGF-β receptor superfamily, causes inhibition of vascular smooth muscle cell proliferation and favors apoptosis. In the absence of
CHAPTER 15  The Lung

45 B Goodpasture syndrome leads to renal and pulmonary lesions produced by an antibody directed against an antigen common to the basement membrane in glomerulus and alveolus. This leads to a type II hypersensitivity reaction. The anti-DNA topoisomerase I antibody is a marker for scleroderma. Antimitochondrial antibody is associated with primary biliary cirrhosis. C-ANCA and P-ANCA are best known as markers for various forms of systemic vasculitis. ANA is used as a general screening test for various autoimmune conditions, typically collagen vascular diseases such as systemic lupus erythematosus.

PB9 701 BP9 485 PB8 507–508

46 D Polyangitis with granulomatosis, a form of ANCA-associated vasculitis, may affect multiple organs, but the lung and kidney are most often involved. The C-ANCA test (PR3) result is often positive in granulomatosis with polyangitis, whereas a positive P-ANCA (MPO) result suggests microscopic polyangitis. Renal and pulmonary disease may be present in Goodpasture syndrome; there may be a positive result for anti-glomerular basement membrane antibody, but no C-ANCA or P-ANCA positivity. Of the collagen vascular diseases, systemic sclerosis is more likely to produce significant pulmonary disease, but hemoptysis is not a prominent feature, and the C-ANCA result is unlikely to be positive. In systemic lupus erythematosus, renal disease is far more likely than pulmonary disease, and C-ANCA or P-ANCA positivity is not expected, but anti-Smith and anti-dsDNA are most specific for SLE. Anti-GBM is a feature of Goodpasture syndrome with extensive intra-alveolar hemorrhage. In hypersensitivity pneumonitis, often the result of inhalation of thermophilic actinomycetes, an initial type III hypersensitivity response is followed by a type IV response, and renal disease is not expected.

PB9 701–710 BP8 507–508

47 A The anesthetic gases tend to reduce the ciliary function of the respiratory epithelium that lines the bronchi. The mucuscciliary apparatus helps clear organisms and particulate matter that are inhaled into the respiratory tree. Mucophagy function is not significantly affected by anesthesia. The levels of γ-globulins in serum are not reduced by the effects of anesthesia. The anesthetic gases and induction drugs do not typically result in marrow failure with neutropenia. The exposure to anesthetic gases is not prolonged enough to produce squamous metaplasia, which most typically occurs in response to chronic irritation, as from cigarette smoke. The subglottic tracheal region, where the cuff of the endotracheal tube is located, can become eroded, but this is more likely to occur when intubation is prolonged for weeks.

PB9 702, 706 BP8 486–487 PB8 710 BP8 510

48 F The productive cough suggests an alveolar exudate with neutrophils, and the course is compatible with an acute infection. Bacterial organisms should be suspected. Pneumococcus is the most likely agent to be cultured in individuals acquiring pneumonia outside of the hospital, and particularly when a lobar pneumatic pattern is present, as in this case. The primary atypical pneumonia of Mycoplasma does not usually produce purulent sputum, unless there is a secondary bacterial infection, which is a common complication of viral and Mycoplasma pneumonias. Cryptococcal and mycobacterial infections typically produce granulomatous disease. Nocardiosis also is seen in immunocompromised patients and produces chronic abscessing inflammation. Pneumocystis pneumonia is seen in immunocompromised patients and is unlikely to produce a lobar pattern of infection.

PB9 702–704 BP9 487–489 PB8 711–714 BP8 509–511

49 C The short time course and acute inflammatory response are consistent with bacterial pneumonitis. Moraxella is the only bacterial organism listed and is in the differential diagnosis of both upper and lower respiratory infections, particularly in persons with chronic obstructive pulmonary disease (COPD), such as this woman. Moraxella is an aerobic gram-negative diplococcus that is oxidase positive. Histoplasmosis is typically a granulomatous disease that develops over weeks. Influenza A viral infection is interstitial, without neutrophilia. M. kansasii produces pulmonary granulomatous disease similar to tuberculosis. Pneumocystis pneumonia is most often seen in immunocompromised persons.

PB9 703 BP9 487–489 PB8 712 BP8 512

50 E A lung abscess often results from aspiration, which can occur in individuals with a depressed cough reflex or in neurologically impaired individuals (e.g., owing to acute alcoholism, anesthesia, or Alzheimer disease). Aspiration into the right lung and the lower lobe is more common because the main bronchus to the left lung is more acutely angled. Bacterial organisms are most likely to produce abscesses, and the infection may be polymicrobial. The most common pathogen is Staphylococcus aureus, but anaerobes such as Bacteroides, Peptococcus, and Fusobacterium spp. also may be implicated. These anaerobes normally are found in the oral cavity and are readily aspirated. The purulent, liquefied center of the abscess can produce the radiographic appearance of an air-fluid level. Cytomegalovirus, Pneumocystis, and cryptococcal infections are seen in immunocompromised individuals and do not typically form abscesses. Candida pneumonia is rare. Nocardial and actinomycotic infections often lead to chronic abscesses without significant liquefaction and affect immunocompromised individuals. Tuberculosis can produce
granulomatous lesions with central cavitation that may be colonized by Aspergillus, although not over a few days.

PBD9 702, 708–709  BP9 488–489, 492  PBD8 716–717  BP8 515

51 C This primary atypical pneumonia is caused by Mycoplasma pneumoniae, a cell wall–deficient organism that is difficult to culture. Often, a diagnosis is made empirically. The findings are similar to those of other viral infections, and serologic testing shows the specific organism. Legionella can produce an extensive pneumonia with neutrophilic alveolar exudates, and the organisms are difficult to grow—they may be revealed by Dieterle silver stain. Mycobacterium fortuitum is a rare infection that is most likely to be seen in very ill or immunocompromised individuals. Nocardiosis produces chronic abscessing inflammation; it is seen mostly in immunosuppressed individuals. Respiratory syncytial virus is typically an infection of early childhood.

PBD9 704  BP9 490  PBD8 714–715  BP8 513–514

52 A Severe acute respiratory syndrome is caused by a strain of coronavirus that is much more virulent than the coronaviruses known to be associated with the common cold. Cytomegalovirus is seen in immunocompromised patients and often involves multiple organs. Ebola virus is virulent and does not cause specific respiratory findings. Herpes simplex virus is a very rare cause of pneumonia, even in immunocompromised patients. Respiratory syncytial virus causes acute respiratory illness in young children.

PBD9 707  BP9 491  PBD8 716  BP8 514

53 B The antigenic drift of influenza viruses, by mutation with minor alteration of either their hemagglutinin (H) or neuraminidase (N) genes, allows them to escape host antibodies. A shift occurs with major change in H or N or both, as happened with H1N1 in this case. Cleavage of influenza virus hemagglutinin by host proteases is essential for the virus to enter cells. The less virulent influenza viruses are cleaved by proteases that are mainly localized to the lung and hence the disease is limited to the lungs. H5N1 virus has much broader tissue tropism because its hemagglutinin can be cleaved by proteases present in many tissues. Host responses to flu virus, such as a cytotoxic T-cell response or macrophage engulfment, are not the major determinant of pathogenicity. Selective infection of CD4+ T cells is a propensity of HIV. Currently avian flu cannot be spread from human to human—but should that happen there would be an avian flu pandemic.

PBD9 346, 705–706  BP9 491  PBD8 715  BP8 514

54 F Human metapneumovirus is second only to respiratory syncytial virus as a cause for pediatric lower respiratory infection, and the two are indistinguishable clinically. She mainly has bronchiolitis. Like most viral infections, round cell infiltrates are interstitial, and those at greatest risk are the very young, elderly, and immunocompromised. Group A streptococcal infections typically produce pharyngitis. Pertussis is rare when childhood vaccinations are done; it causes whooping cough from upper airway involvement. Candida infections of the lower respiratory tract are uncommon even in immunocompromised persons. Cytomegalovirus is most likely to involve the lungs of immune compromised persons. H. influenzae is more likely to produce neutrophilic patchy infiltrates with productive cough.


55 C Respiratory syncytial virus (RSV) pneumonia is most common in children, and it can occur in epidemics. Viral, chlamydial, and mycoplasmal pneumonias are most often interstitial, without neutrophilic alveolar exudates. The diagnosis is often presumptive because viral culture is technically difficult and expensive, such as a PCR assay. Hyperinflation can accompany bronchoconstriction in asthma. Marked lymphadenopathy is more characteristic of chronic processes, such as granulomatous diseases or metastases. Lobar consolidation is more typical of a bacterial process, such as can be seen in Streptococcus pneumoniae infection. Pleural effusions can be seen in pulmonary inflammatory processes, but they are most pronounced in heart failure. Cavitation is most likely to complicate secondary tuberculosis in adults.

PBD9 705  BP9 491  PBD8 349, 714–715  BP8 511, 536

56 D Nontuberculous mycobacterial infections such as Mycobacterium avium-complex (MAC) are likely to become disseminated illnesses in immunocompromised persons. In immunocompetent persons, such infections are more likely to resemble tuberculosis. The acid-fast MAC organisms proliferate profusely in macrophages within the mononuclear phagocyte system. Extensive and severe aspergillosis and candidiasis are more likely to occur with profound neutropenia, not lymphocytopenia. Legionella produces an extensive bacterial pneumonia, not disseminated disease. Nocardiosis occurs in immunocompromised patients, and the organisms can be weakly acid-fast, but focal nodules or a chronic abscessing inflammatory response are more likely.

PBD9 376–377  BP9 499  PBD8 372–373  BP8 523

57 A The figure shows pink, amorphous tissue at the lower left, representing caseous necrosis. The rim of the granuloma has epithelioid cells and Langhans giant cells. Caseating granulomatous inflammation is most typical of Mycobacterium tuberculosis infection. Calcifications would have helped to identify this mass as an old granuloma, not likely to be a neoplasm. Necrotizing vasculitis is unlikely to produce a single nodule, and there can be hemoptysis. A carcinoma may have central necrosis, not caseation, and there would be atypical, pleomorphic cells forming the mass. A pulmonary infarct should have extensive hemorrhage. A lung abscess would have an area of liquefactive necrosis filled with tissue debris and neutrophils.

PBD9 371–376, 709  BP9 493–497  PBD8 367–372  BP8 516–519

58 A This “coin lesion” on imaging of his lungs could be an infectious granuloma, a neoplasm, or a hamartoma. His fever suggests infection, and the CT finding of decreased central attenuation in the nodule suggests necrosis in a neoplasm.
or caseous necrosis in a granuloma. The lymphocytosis and monocyctosis are consistent with tuberculosis. Nonsmokers are unlikely to develop primary lung neoplasms, and adenocarcinoma is the most common in that setting. Smokers are most likely to develop squamous cell carcinomas and small cell anaplastic carcinomas. Individuals who are immunocompromised are most likely to develop fungal infections, particularly with *Aspergillus* spp., which have branching septate hyphae. Charcot-Leyden crystals form from eosinophil granules in individuals with allergic asthma. Foreign body giant cells can be seen with lipid pneumonias. Gram stain is most useful for determining which bacterial organisms may be present, and gram-negative bacilli such as the Enterobacteriaceae produce acute pneumonias and abscesses with neutrophilia.

> **PBD9 374–376** BP9 498 PBD8 718, 730 BP8 522

**59 C** These findings represent the so-called Ghon (or primary) complex, consisting of a small subpleural granuloma with extensive hilar nodal caseating granulomas. The Ghon complex is a feature of primary tuberculosis, which is most often a subclinical disease of younger individuals. The interferon-γ release assay is positive when T lymphocytes are activated against *Mycobacterium tuberculosis* antigens, and is negative in individuals who received BCG vaccination. Individuals who are immunocompromised, such as HIV-infected patients, do not mount a good granulomatous response and have more extensive poorly formed granulomas, dissemination of tuberculosis, or both. Anticentromere antibody is characteristic of limited scleroderma, which does not have significant pulmonary involvement, in contrast to diffuse scleroderma. The rapid plasma reagin test is used to diagnose syphilis, which does not have significant pulmonary disease. Rheumatoid nodules may be seen in rheumatoid arthritis; these can be subpleural, but patients typically have arthritis. Patients with cystic fibrosis and an elevated sweat chloride level (more often elevated in children than in adults) develop widespread bronchiectasis along with infection by bacterial agents, particularly *Pseudomonas aeruginosa* and *Burkholderia cepacia*.

> **PBD9 374–375** BP9 495 PBD8 370 BP8 519

**60 B** Coccidioidomycosis is endemic to semiarid regions of the Americas. The species *Coccidioides immitis* is found in the southwestern United States, and the nearly identical *Coccidioides posadasii* is found from Mexico to South America. Dimorphic fungal diseases may produce granulomatous infection resembling tuberculosis. The military pattern described in this patient occurs with a poor immune response. Blastomyces dermatitidis organisms are smaller than 10 microns and exhibit broad-based budding. *Histoplasma capsulatum* organisms are 2 to 4 microns and are often found within phagocytic cells. Mycobacteria are acid-fast rod-shaped organisms only a micron across. *Nocardia* organisms are long, filamentous bacteria that may be weakly acid-fast. *Paracoccidioides brasiliensis* organisms can be 20 to 30 microns and exhibit multiple buds in a “captain’s wheel” configuration.

> **PBD9 710–711** BP9 499–500 PBD8 719 BP8 523

**61 D** The radiograph in the figure shows prominent upper lobe cavitation, typical of reactivation-reinfection tuberculosis in adults. *Candida* is a rare cause of lung infection. Influenza viral infections have mainly interstitial mononuclear inflammation. Bacterial organisms such as *Legionella* are more likely to produce a widespread bronchopneumonia with alveolar neutrophilic exudates. *Mycoplasma* infection produces mainly interstitial mononuclear inflammation. No-cardiosis of the lung appears mainly as chronic abscessing inflammation.

> **PBD9 374–376** BP9 496–498 PBD8 369–372 BP8 520–521

**62 A** There are several patterns of pulmonary involvement with *Aspergillus* spp. Immunocompromised patients with neutropenia may develop invasive aspergillosis. Other patterns include allergic bronchopulmonary aspergillosis in persons with asthma and an aspergilloma, or fungus ball, colonizing a cavitary lesion of tuberculosis or bronchiectasis. Candidiasis may also develop in the setting of neutropenia, but less commonly causes extensive lung involvement, appears as budding cells with pseudohyphae, and more likely produces an oral, nasal, or pharyngeal infection. Cryptococcosis can cause extensive pulmonary infections, particularly with loss of cell-mediated immunity, and the organisms have large mucoid capsules and exhibit narrow-based budding. *Monaxella* is a bacterial organism most often causing sinusitis, otitis, and upper respiratory infections. *Mucor* appears as broad, nonseptated hyphae and is most often a complication of diabetic ketoacidosis, with nasal involvement.

> **PBD9 388–389, 712** BP9 504 PBD8 384–385, 720 BP8 324, 527

**63 C** Within the airspace are multiple large cells with prominent purple intranuclear inclusions, typical for CMV. Although CMV antibodies can be found in most immunocompetent persons, the infection is subclinical. However, in immunocompromised persons, CMV infection can be a severe and systemic infection, including marked interstitial pneumonitis. The other listed organisms are much smaller. *Candida* is a rare cause for pneumonia, and this yeast appears as budding cells and pseudohyphae. Cryptococcal organisms exhibit narrow-based budding and thick mucoid capsules. *Klebsiella pneumoniae* organisms are encapsulated, but these bacteria are too small to be seen with H&E staining. *Pneumocystis* organisms are best seen with Gomori methenamine silver stain and appear as round to cup-shaped cysts.


**64 C** Although *Pneumocystis jiroveci* pneumonia can be seen with various acquired and congenital immunodeficient states (mainly those affecting cell-mediated immunity), it is most often associated with AIDS and is diagnostic of AIDS in HIV-infected individuals. Persons with diabetes mellitus and pulmonary emphysema are most prone to contract bacterial infections. Patients with autoimmune diseases may have cytopenias that predispose to infection, and if they are treated with immunosuppressive drugs, various infections are possible. Likewise, patients with sarcoidosis treated with corticosteroid therapy may have opportunistic infections. A patient with severe combined immunodeficiency is susceptible
65 G Of all lung cancers, squamous cell carcinoma is most likely to produce paraneoplastic hypercalcemia, and there is a strong association with smoking. These tumors also can undergo central necrosis—hence a cavity may form. Localized squamous cell carcinomas, in contrast to small cell carcinomas, may be cured by surgery. Adenocarcinomas and large cell carcinomas tend to produce peripheral masses and generally are less likely to be associated with paraneoplastic syndromes. Kaposi sarcoma involving visceral organs is most often seen in association with AIDS, and it is often multifocal. Renal cell carcinomas may be associated with hypercalcemia, but metastases usually appear as multiple masses (although of all metastatic tumors, renal cell carcinoma is most likely to produce solitary metastases). Non-Hodgkin lymphomas generally do not have paraneoplastic effects; they are uncommon in the lung and are not associated with smoking. Small cell carcinomas are never localized enough for curative surgery (they are usually detected at an advanced stage), although they often produce various paraneoplastic syndromes, but less likely hypercalcemia.

66 A The long history and the weight loss suggest a chronic, debilitating process. The radiograph shows multiple mass lesions. The focal nature of the lesions, with remaining pulmonary reserve capacity, means that measured lung function remains normal. The most common neoplastic process of lung is metastatic disease, because many primary sites outside of lung can gain vascular access and spread hematogenously, and lung has a rich capillary bed for tumor cell emboli to colonize. Immunohistochemical markers on this adenocarcinoma might help characterize the primary site, or they might not. Depending upon the markers, there might be specific antineoplastic therapies, or not. The lack of fever and any evidence for inflammation makes an infectious process unlikely. Necrotizing vasculitis is likely to produce hemoptysis. Silicosis is unlikely without an environmental exposure in a job setting, and the lesions lead to restrictive lung disease.

67 E The ALK gene encodes a receptor tyrosine kinase that gets activated by fusion with the EML gene. The fused ALK-EML (just as BCR-ABL in CML) can be targeted by inhibitors of tyrosine kinases. Gain-of-function mutations in multiple genes encoding receptor tyrosine kinases, including EGFR, ALK, ROS, MET, and RET, are often mutated in adenocarcinomas, which tend to be peripheral masses. Along with EGFR, ALK gene mutations can be targeted in the subset of lung adenocarcinomas that bear these molecular changes. All the other genes, although more commonly mutated in a variety of cancers, cannot be targeted by any of the available drugs.

68 A Cancers that arise in nonsmokers are pathogenetically distinct from those that occur in smokers. They may have either EGFR mutations or KRAS mutations. Most are adenocarcinomas. Twenty-five percent of lung cancers worldwide occur in nonsmokers. Primary adenocarcinomas in the lung tend to be small, peripheral masses that are amenable to surgical excision and have a better overall prognosis than other forms of lung cancer. Overall, far more metastatic adenocarcinomas involve the lung than do primary adenocarcinomas. Bronchial carcinoids are uncommon endobronchial lesions. Hamartomas are small, peripheral masses that contain benign epithelial and connective tissue elements. Large cell carcinomas are too poorly differentiated to be called adenocarcinomas or squamous cell carcinomas. The most common cancers in smokers are small cell anaplastic and squamous cell carcinomas.

69 F Cushing syndrome is a paraneoplastic syndrome resulting from ectopic corticotrophin production (most often from a pulmonary small cell carcinoma), which drives the adrenal cortices to produce excess cortisol. Small cell carcinomas are aggressive neuroendocrine tumors that tend to metastasize early. Even when they appear to be small and localized, they are not or will not remain so. Surgery is not an option for these patients. They are treated as if they have systemic disease; some chemotherapy protocols afford benefit for 1 year or more, but cure is uncommon. Adenocarcinomas and large cell carcinomas tend to be peripheral neoplasms in the lung, and they are less likely to produce a paraneoplastic syndrome. Bronchial carcinoids at the more benign end of the neuroendocrine tumor spectrum tend to be small and are not likely to produce paraneoplastic effects; rarely, they produce carcinoid syndrome. Non-Hodgkin lymphomas rarely occur within the lung, are not associated with smoking, and do not produce Cushing syndrome. Squamous cell carcinomas can be central and occur in smokers, but they are more likely to produce hypercalcemia as a paraneoplastic syndrome.

70 A Adenocarcinoma in situ (AIS), formerly termed bronchioloalveolar carcinoma, can present as a peripheral tumor that can mimic pneumonia. Most of these tumors are well differentiated. Adenocarcinomas and large cell carcinomas tend to be peripheral, but the former tend to produce a localized mass, whereas cells of the latter are large and pleomorphic and form sheets; sometimes it is difficult to distinguish among them. Mesotheliomas almost always occur in the setting of prior asbestos exposure; they are large pleural masses. Metastases tend to appear as multiple nodules. Squamous cell carcinomas occasionally can be peripheral (although most are central) and are composed of pink, polygonal cells that have intercellular bridges. If well differentiated, squamous cell carcinomas show keratin pearls.

71 D The patient probably has a small cell anaplastic (oat cell) carcinoma of the lung, which is most likely to produce
a paraneoplastic syndrome with the syndrome of inappro-
priate secretion of antidiuretic hormone (SIADH), marked
by free water retention with hyponatremia. Oat cell cancers
tend to be central masses, and they are strongly associated
with smoking. Upper lobe cavitation suggests secondary
tuberculosis. Diaphragmatic pleural plaques can be a feature of
pneumoconioses, particularly asbestosis. Infiltrates can suggest
an inflammatory process. Pneumothorax is most likely to occur
from chest trauma, not from a neoplasm. A subpleural nodule
with hilar adenopathy is the classic Ghon complex of primary
tuberculosis, which is unlikely to manifest with hemoptysis.
An air-fluid level suggests liquefaction in an abscess.

72 C Horner syndrome is a result of sympathetic au-
tonomic nerve involvement by invasive pulmonary carcinoma.
Such a neoplasm in this location with these associated find-
ings is called a Pancost tumor. Infectious processes such as
pneumonia are unlikely to impinge on structures outside the
lung. Bronchiectasis destroys bronchi within the lung. Sar
coidosis can result in marked hilar adenopathy with a mass
effect, but involvement of the peripheral nervous system is
unlikely. Likewise, tuberculosis is a granulomatous disease
that can lead to hilar adenopathy, although usually without
destruction of extrapulmonary tissues.

73 B Hamartomas are uncommon but benign peripheral les-
sions of the lung. They are composed of benign-appearing epithelial cells and connective tissue, typically with a large component of
cartilage. They are included in the differential diagnosis of a “coin lesion” that also includes carcinoma and granuloma. Adenocarcinoma is the most common primary lung malignancy in nonsmokers, and it can manifest as a coin lesion, but it is composed of gland-forming, malignant cells without cartilage. It tends to be peripheral, making surgical resection an option in many cases. Large cell carcinomas also are more likely to be peripheral, but they tend to be larger masses, with poorly differentiated cells. Malignant mesothelioma is a rare neoplasm, even in individuals who have been exposed to asbestos, and it arises on the pleura. Primary non-Hodgkin lymphomas of the lung are uncommon, but may involve hilar or mediastinal lymph nodes. Some squamous cell carcinomas can be peripheral, but they are most likely to occur in individuals who smoke.

74 B Most pulmonary carcinoids are central obstructing masses involving a large to medium-sized bronchus. These neuroendocrine tumors have unpredictable behavior, but many are localized, resectable, and follow a benign course. They typically manifest with hemoptysis and the consequences of bronchial obstruction. In this case, the pneumonia in the right upper lobe probably resulted from obstruction to drainage caused by the tumor. Obstruction may lead to peripheral resorption atelectasis. Adenocarcinomas are common lung tumors, but are typically peripheral. A hamartoma is an uncommon but benign pulmonary lesion that also is located peripherally. Kaposi sarcoma can involve the lung in some patients with AIDS, and the tumor often has a bronchovascular distribution; obstruction is not common, but bleeding on biopsy is. Large-cell carcinomas are typically large, bulky, peripheral masses.

75 F The pleural fluid findings are typical of chylothorax, which is uncommon but distinctive. Lymph fluid is rich in lymphocytes, protein, and lipid (chylomicrons). Disruption of the thoracic duct in the posterior chest is most likely to cause chylothorax, and malignant neoplasms, such as non-Hodgkin lymphoma, are most likely to do this. An empyema is composed of pus formed from neutrophilic exudation and would appear cloudy and yellow. Congenital heart disease can lead to congestive heart failure with a serious effusion. Aortic dissection is an acute condition that can produce a hemothorax. Cirrhosis is more likely to be associated with ascites or liver failure with hypoalbuminemia leading to hy-
drothorax. Miliary tuberculosis is seen as a reticulonodular pattern on a chest radiograph; tuberculosis may produce hemorrhagic effusions.

76 B The pleural fluid findings with low protein, low LDH, and low cell count are consistent with a transudate with hydrothorax. His hypertension has likely led to left ventricular failure with pulmonary edema, and longstanding left ventricular failure can lead to right ventricular failure with body cavity effusions and peripheral edema. Granulomatous diseases, including tuberculosis, along with carcinomas that are primary or metastatic to pleura, tend to produce cellular effusions with numerous RBCs. Mesothelioma is likely to shed numerous cells into the pleural cavity, although in most cases there is dense tumor that obliterates the pleural cavity. Lymphomas may block lymphatics, including the thoracic duct, to produce a milky chylothorax with lipid and leukocytes. Bacterial pneumonias spreading to the pleura are likely to produce an exudative effusion with numerous leukocytes, predominantly neutrophilic.

77 A Malignant mesothelioma is a rare tumor even in individuals with a history of asbestos exposure. The tu-
more may appear decades after exposure, and is not related to amount or length of exposure. Bronchogenic carcinoma is more common in individuals with asbestos exposure, particularly when there is a history of smoking. Bird dust inhalation can lead to hypersensitivity pneumonitis. Coal dust inhalation can lead to marked anthracosis, but without a significant risk of lung cancer. Inhalation of cotton fibers (byssinosis) leads to symptoms resembling asthma related to bronchoconstriction. Ozone and nitrogen oxides in smog can cause acute respiratory discomfort, but are not known to be promoters of neoplasia. Silicosis is typified by interstitial fibrosis and causes a slight increase in the risk of bronchogenic carcinoma.
The solitary fibrous tumor, or localized mesothelioma, of pleura (or peritoneum) is a rare neoplasm that appears as a pedunculated mass. There is no relationship to asbestos exposure or other environmental pathogens. Many do not recur or metastasize following resection, but larger tumor size, higher mitotic count, and greater patient age (>55 years) increase the risk for metastasis. Bronchioloalveolar carcinomas are peripheral (but intraparenchymal) masses with atypical epithelial cells growing along the framework of the lung. A hamartoma is a peripheral intraparenchymal mass with a significant component of fibrous connective tissue and usually with cartilage present. Hodgkin lymphoma is more likely to involve lymph nodes in the mediastinum. A malignant mesothelioma forms a pleural mass that is not circumscribed; the cells are atypical and cytokeratin positive. Metastases are typically multiple and often produce bloody effusions.
1. A 47-year-old man sees his dentist for a routine checkup. He states that his gums bleed easily on brushing his teeth. On examination, he is found to have marked gingival recession with erythema, along with extensive plaque and calculus formation over tooth surfaces. Which of the following organisms is most likely to be associated with development of his oral lesions?
   - A. Actinobacillus
   - B. Candida
   - C. Epstein-Barr virus
   - D. Herpes simplex virus
   - E. Human papillomavirus
   - F. Mucor circinelloides

2. A 17-year-old girl notices a small, sensitive, gray-white area forming along the lateral border of her tongue 2 days before the end of her final examinations. On examination by the physician’s assistant, the girl is afebrile. There is a shallow, ulcerated, 0.3-cm lesion with an erythematous rim. No specific therapy is given, and the lesion disappears within 2 weeks. The history shows that the girl does not use tobacco or alcohol. Which of the following is the most probable diagnosis?
   - A. Aphthous ulcer
   - B. Herpes simplex stomatitis
   - C. Leukoplakia
   - D. Oral thrush
   - E. Sialadenitis

3. A 55-year-old woman notes a nodule while rubbing her tongue on the side of her mouth. On physical examination by her dentist, there is a firm, nontender 0.6-cm nodule covered by pink buccal mucosa at the bite line next to the first molar on the lower right. The lesion is excised and does not recur. What is the most likely diagnosis?
   - A. Candidiasis
   - B. Fibroma
   - C. Leukoplakia
   - D. Pyogenic granuloma
   - E. Sialadenitis

4. A 23-year-old primigravida has noticed a rapidly enlarging nodule next to a tooth for the past 16 days. On physical examination there is a 1-cm, soft, reddish, pedunculated mass above a left upper bicuspid. She is advised that the lesion will likely regress. Which of the following pathologic findings is most likely found in this lesion?
   - A. Granulation tissue
   - B. Lymphoid proliferation
   - C. Neutrophilic exudate
   - D. Rhabdomyosarcoma
   - E. Squamous hyperplasia

5. A 25-year-old man notices several 0.3-cm, clear vesicles on his upper lip after a bout of influenza. The vesicles rupture, leaving shallow, painful ulcers that heal over the course of 10 days. Three months later, after a skiing trip, similar vesicles develop, with the same pattern of healing. Which of the following microscopic findings is most likely to be associated with these lesions?
   - A. Budding cells with pseudohyphae
   - B. Mononuclear inflammatory infiltrates
   - C. Neutrophils within abscesses
   - D. Squamous epithelial hyperkeratosis
   - E. Intranuclear inclusions

6. A 35-year-old, HIV-positive man complains that he has had a “bad” taste in his mouth and discoloration of his tongue for the past 6 weeks. On physical examination, there are areas of adherent, yellow-to-gray, circumscribed plaque on the lateral aspects of the tongue. This plaque can be scraped off as a pseudomembrane to show an underlying granular, erythematous base. What is the most likely diagnosis?
   - A. Aphthous ulcer
   - B. Cheilosis
   - C. Hairy leukoplakia
   - D. Herpetic stomatitis
   - E. Leukoplakia
   - F. Oral thrush
7. A 42-year-old man has had a constant bad taste in his mouth for the past month. On physical examination there are white fluffy patches on the sides of his tongue. These cannot be scraped off. A biopsy is taken and on microscopic examination shows squamous epithelial hyperkeratosis, parakeratosis, and koilocytosis. Immunohistochemical staining for Epstein-Barr virus (EBV) is positive. Which of the following is the most likely risk factor for his oral lesions?

- A Chronic alcohol abuse
- B Diabetes mellitus
- C HIV infection
- D Pernicious anemia
- E Sjögren syndrome

8. A 58-year-old man, a cigar smoker, visited his dentist for a routine dental examination. The dentist noticed lesions with the clinical (A) and histologic (B) appearance shown in the figure. The medical history showed no major medical problems. Which of the following etiologic factors most likely contributed to the development of these lesions?

- A Chronic sialadenitis
- B Dental caries
- C Eating smoked foods
- D Herpes simplex virus type 1
- E Smoking tobacco

9. A 51-year-old man from Kolkata has an area of depression in his mouth that has enlarged over the past 7 months. On oral examination, there is a 1.5 × 0.7 cm velvety, erythematous area with focal surface erosion on his left buccal mucosa. The lesion is excised and on microscopic examination there is dysplastic squamous epithelium. Which of the following is the most likely risk factor for developing this lesion?

- A Candidiasis
- B Dental malocclusion
- C Epstein-Barr virus infection
- D Immunosuppression
- E Eating hot, spicy food
- F Tobacco chewing

10. A 49-year-old man has used chewing tobacco and snuff for many years. On physical examination the lesion shown in the figure is seen on the hard palate. It cannot be removed by scraping. A biopsy is performed, and microscopic examination of the lesion shows a thickened squamous mucosa. Four years later, a biopsy specimen of a similar lesion shows carcinoma in situ. Which of the following is the most likely diagnosis?

- A Oral thrush
- B Lichen planus
- C Leukoplakia
- D Pyogenic granuloma
- E Xerostomia

11. A 54-year-old man, a nonsmoker, has a nonhealing ulceration at the base of his tongue on the right side for 2 months. On examination this lesion is 1 cm in diameter with irregular borders. Biopsy of the lesion is performed and microscopic examination shows infiltrating squamous cell carcinoma. Which of the following infectious agents is most likely to be associated with this lesion?

- A Candida albicans
- B Herpes simplex virus (HSV)
- C Human papillomavirus (HPV)
- D Prevotella intermedia
- E Group A streptococcus
12 A 19-year-old woman has noted swelling in the back of her mouth for 2 months. On dental examination, she has an area of swelling in the location of the left third molar. Dental radiographs show a radiolucent unilocular, well-circumscribed cyst surrounding the crown of the unerupted third mandibular molar. The lesion is excised, and on microscopic examination, the cyst is lined by stratified squamous epithelium and surrounded by a chronic inflammatory infiltrate. What is the most likely diagnosis?
   A Ameloblastoma  B Dentigerous cyst  C Odontogenic keratocyst  D Odontoma  E Periapical cyst/granuloma

13 A 19-year-old man noted progressive swelling on the left side of his face over the past year. On physical examination, there is painless swelling in the region of the left posterior mandible. Head CT scan shows a circumscribed multilocular cyst of the left mandibular ramus. The lesion is surgically excised with wide bone margins. On microscopic examination, the lesion shows cysts lined by stratified squamous epithelium with a prominent basal layer; no inflammation or granulation tissue is seen. What is the most likely diagnosis?
   A Ameloblastoma  B Dentigerous cyst  C Odontogenic keratocyst  D Odontoma  E Periapical cyst/granuloma

14 A 26-year-old man has had difficulty breathing through his nose for 3 years, but this problem has become progressively worse over the past 2 months. Physical examination shows glistening, translucent, polypoid masses filling the nasal cavities. Histologic examination of the excised masses shows respiratory mucosa overlying an edematous stroma with scattered plasma cells and eosinophils. Which of the following laboratory findings is most likely to be present in this patient?
   A Elevated serum hemoglobin A1c level  B Increased serum IgE level  C Nuclear staining for Epstein-Barr virus antigens  D Positive ANA test result  E Tissue culture positive for Staphylococcus aureus

15 A 39-year-old woman has been bothered by headache, facial pressure, nasal obstruction with discharge, and diminished taste sensation for the past 6 months. On physical examination there is discomfort on palpation over her left maxillary sinus. No oral lesions are noted. Rhinoscopy shows nasal erythema, marked edema, and purulent discharge. Which of the following complications is most likely to occur in this patient?
   A Mucocele  B Nasopharyngeal carcinoma  C Osteomyelitis  D Sinonasal papilloma  E T-cell lymphoma

16 On December 13, 1799, George Washington, recently retired as first President of the United States, developed a “cold” with mild hoarseness. By the next morning he had difficulty breathing and swallowing, with throat pain. He was treated with the usual therapy of the time: bloodletting. Had vital signs been recorded, they may have shown temperature of 37.8°C, pulse 115/min, respiratory rate 24/min, and blood pressure 90/60 mm Hg. Which of the following organisms most likely caused his illness?
   A Coronavirus  B Corynebacterium diphtheriae  C Haemophilus influenzae  D Parainfluenza virus  E Prevotella intermedia  F Group A streptococcus

17 A 3-year-old child has had difficulty breathing for the past 24 hours. On physical examination, the child is febrile and has a harsh cough with prominent inspiratory stridor. The lungs are clear on auscultation. An anterior-posterior neck radiograph shows the steeple sign caused by edema producing loss of normal shoulders on the subglottic larynx. The child’s oxygen saturation is normal with pulse oximetry. She improves over the next 3 days while taking nebulized glucocorticoids. Which of the following organisms is the most likely cause of the child’s condition?
   A Corynebacterium diphtheriae  B Epstein-Barr virus  C Haemophilus influenzae  D Human papillomavirus  E Parainfluenza virus  F Streptococcus, group A

18 A 9-year-old girl has had a sore throat for the past 2 days. On physical examination there is pharyngeal erythema with yellowish exudates over swollen palatine tonsils. A Gram stain of the exudate shows gram-positive cocci in chains. She is given penicillin therapy. What is the most likely complication prevented by prompt treatment of this girl?
   A Carditis  B Hepatitis  C Meningitis  D Otitis  E Pneumonitis
A 48-year-old man from Hong Kong has had difficulty breathing through his nose and has experienced dull facial pain for the past 4 months. On physical examination, there is a mass filling the right nasal cavity. CT scan of the head shows a 5-cm mass in the nasopharynx on the right that erodes adjacent bone. The mass is excised, and microscopic examination shows that it is composed of large epithelial cells with indistinct borders and prominent nuclei. Mature lymphocytes are scattered throughout the undifferentiated neoplasm. Which of the following etiologic factors most likely played the greatest role in the development of this lesion?

A  Allergic rhinitis  
B  ANCA-associated vasculitis  
C  Epstein-Barr virus infection  
D  Sjögren syndrome  
E  Smoking tobacco

A 28-year-old man who is a singer/songwriter has been experiencing hard times for the past 3 years. He has played at a couple of clubs a night to earn enough to avoid homelessness. He comes to the free clinic because he has noticed that his voice quality has become progressively hoarser over the past year. On physical examination, he is afebrile. There are no palpable masses in the head and neck area. He does not have a cough or significant sputum production, but he has been advised on previous visits to give up smoking. Which of the following is most likely to produce these findings?

A  Croup  
B  Epiglottitis  
C  Reactive nodule  
D  Squamous cell carcinoma  
E  Squamous papillomatosis

A 6-year-old boy has had increased difficulty breathing, and the character of his voice has changed over the past 3 months. Endoscopic examination shows three soft, pink excrescences on the true vocal cords and in the subglottic region. The masses are 0.6 to 1 cm in diameter. Microscopic examination of the excised masses shows fingerlike projections of orderly squamous epithelium overlying fibrovascular cores. Immunostaining for human papillomavirus 6 antigens is positive. Based on these findings, which of the following statements is the best advice to give the parents of this boy?

A  A total laryngectomy is necessary  
B  Congenital heart disease may be present  
C  The boy should not overuse his voice  
D  The lesions are likely to recur  
E  Therapy with acyclovir is indicated

A 58-year-old man bothered by increasing hoarseness for almost 6 months now has an episode of hemoptysis. On physical examination, no lesions are noted in the nasal or oral cavity. There is a firm, nontender anterior cervical lymph node. The lesion shown in the figure is identified by endoscopy. The patient undergoes biopsy, followed by laryngectomy and neck dissection. Which of the following etiologic factors most likely played the greatest role in the development of this lesion?

A  Epstein-Barr virus infection  
B  Human papillomavirus infection  
C  Repeated bouts of aspiration  
D  Smoking tobacco  
E  Type I hypersensitivity

A 5-year-old boy has had repeated bouts of earache for 3 years. Each time on examination, the bouts have been accompanied by a red, bulging tympanic membrane, either unilaterally or bilaterally, sometimes with a small amount of yellowish exudate. Laboratory studies have included cultures of Staphylococcus aureus, Pseudomonas aeruginosa, and Moraxella catarrhalis. The most recent examination shows that the right tympanic membrane has perforated. The boy responds to antibiotic therapy. Which of the following complications is most likely to occur as a consequence of these events?

A  Cholesteatoma  
B  Eosinophilic granuloma  
C  Labyrinthitis  
D  Otosclerosis  
E  Squamous cell carcinoma
**24** A 64-year-old man has had progressive difficulty hearing, particularly with the left ear, over the past 10 years. Audiometric testing shows that he has a bone conduction type of deafness. CT scan of the head shows no abnormal findings. The patient’s brother and mother are similarly affected. What is the most likely diagnosis?

- A Cholesteatoma
- B Chondrosarcoma
- C Otitis media
- D Otosclerosis
- E Schwannoma

**25** A 25-year-old woman is concerned about a lump on the left side of her neck that has remained the same size for the past year. Physical examination shows a painless, movable, 3-cm nodule beneath the skin of the left lateral neck just above the level of the thyroid cartilage. There are no other remarkable findings. Fine-needle aspiration of the mass is performed. Her physician is less than impressed by the pathology report, which notes, “Granular and keratinaceous cellular debris.” Fortunately, she has saved her Robbins pathology textbook from medical school. She consults the head and neck chapter to arrive at a diagnosis, using the data from the report. Which of the following terms best describes this nodule?

- A Branchial cyst
- B Metastatic thyroid carcinoma
- C Mucocele
- D Mucoepidermoid tumor
- E Paraganglioma
- F Thyroglossal duct cyst

**26** A 17-year-old girl is concerned about a “bump” on her neck that she has noticed for several months. It does not seem to have increased in size during that time. On physical examination, there is a discrete, slightly movable nodule in the midline of the neck just adjacent to the region of the hyoid. The nodule is excised, and microscopic examination shows a cystic mass lined by squamous and respiratory epithelium surrounded by fibrous tissue with lymphoid nodules. Which of the following additional histologic elements would most likely be located adjacent to this cyst?

- A Malignant lymphoma
- B Noncaseating granulomas
- C Serous salivary glands
- D Squamous cell carcinoma
- E Thyroid follicles

**27** A 56-year-old woman has noticed an enlarging lump on the right side of her neck for the past 7 months. On physical examination, there is a 3-cm nodule in the right upper neck, medial to the sternocleidomastoid muscle and lateral to the trachea at the angle of the mandible. CT scan shows a circumscribed, solid mass adjacent to the carotid bifurcation. Microscopic examination of the excised mass shows nests of round cells with pink, granular cytoplasm. Tests for immunohistochemical markers chromogranin and S-100 are positive. Electron microscopy shows neurosecretory granules in the tumor cell cytoplasm. The tumor recurs 1 year later and is again excised. What is the most likely diagnosis?

- A Metastatic squamous cell carcinoma
- B Metastatic thyroid medullary carcinoma
- C Mucoepidermoid carcinoma
- D Paraganglioma
- E Warthin tumor

**28** A 67-year-old man with Parkinson disease has experienced an increasingly dry mouth for the past 3 months, and this interferes with eating and swallowing. He has noted dry eyes as well. On physical examination, he has minimal tremor at rest; there are no other abnormal findings. Laboratory studies show no detectable autoantibodies. Which of the following agents is most likely to be found in this gland?

- A Alcohol ingestion
- B Anticholinergic drug use
- C Candidiasis
- D Sialadenitis with blockage of salivary duct
- E Sjögren syndrome
- F Tobacco use

**29** A 69-year-old man has a major psychosis. He has been bothered by pain on the left side of the face for 2 weeks. On physical examination, there is a tender area of swelling 4 cm in diameter beneath the skin, anterior to the left auricle above the angle of the jaw. CT scan of the head shows cystic and solid areas in the region of an enlarged left parotid gland. After a course of antibiotic therapy, there is only minimal improvement. A parotidectomy is performed. Microscopic examination of the excised gland shows acute and chronic inflammation, with fibrosis and abscess formation, duct lithiasis, and atrophy of acini. Which of the following infectious agents is most likely to be found in this gland?

- A Epstein-Barr virus
- B Human papillomavirus
- C Prevotella intermedia
- D Rubella virus
- E Staphylococcus aureus

**30** A 95-year-old man has noted swelling of his lower lip for the past month. On examination, there is a fluctuant, 1-cm nodule with a blue, translucent hue just beneath the oral mucosa on the inside of his lip. The lesion is excised, and on microscopic examination shows granulation tissue. What is the most likely etiology for this lesion?

- A Eating chili peppers
- B French kissing
- C HIV infection
- D Local trauma
- E Pipe smoking

**31** A 65-year-old woman has noticed a slowly enlarging nodule on her face for the past 3 years. On physical examination, a 3-cm, nontender, mobile, discrete mass is palpable on the left side of the face, anterior to the ear and just superior to the mandible. The mass is completely excised, and histologic examination shows ductal epithelial cells in a myxoid stroma containing islands of chondroid-like tissue and bone. This patient is most likely to have which of the following neoplasms?

- A Acinic cell tumor
- B Mucoepidermoid carcinoma
- C Pleomorphic adenoma
- D Primitive neuroectodermal tumor
- E Squamous cell carcinoma
- F Warthin tumor
A 57-year-old man notices a lump on the right side of his face that has become larger over the past year. On physical examination, a 3- to 4-cm firm, mobile, painless mass is palpable in the region of the right parotid gland. The oral mucosa appears normal. He does not complain of difficulty in chewing food or talking. The mass is completely excised, and histologic examination shows the findings in the figure. What is the most likely diagnosis?

A. Mucoepidermoid carcinoma
B. Non-Hodgkin lymphoma
C. Pleomorphic adenoma
D. Sialolithiasis
E. Sjögren syndrome
F. Warthin tumor

A 60-year-old woman noticed an enlarging “bump” beneath her tongue for the past year. She does not smoke or use alcohol. On physical examination, there is a 2.5-cm, movable, submucosal mass arising in the minor salivary glands on the buccal mucosa beneath the tongue on the right. Histologic examination of the excised mass shows that it is malignant and locally invasive. The tumor recurs within 1 year. Which of the following is the most likely diagnosis?

A. Non-Hodgkin lymphoma
B. Mucoepidermoid carcinoma
C. Primitive neuroectodermal tumor
D. Pleomorphic adenoma
E. Squamous cell carcinoma
F. Warthin tumor

ANSWERS

1. A. Periodontitis becomes more prevalent with age, often secondary to the effects of dental plaque formation driven by oral flora. The gingival recession increases the risk for dental caries. Regular dental cleanings to remove the plaque and regular gentle tooth brushing help to slow the progression of periodontitis. Some periodontitis cases arise in the setting of systemic disease. Candidiasis is seen in immunocompromised individuals and often forms an inflammatory membrane on the tongue. Epstein-Barr virus has been associated with development of hairy leukoplakia. Herpes simplex virus results in vesicles that can rupture and form superficial ulcers on oral mucosa. Human papillomavirus can drive squamous epithelial hyperplasia, dysplasia, and carcinoma. Mucor has broad, nonseptated hyphae and can result in sinusitis, particularly in the setting of ketoacidosis.

2. A. An aphthous ulcer is a common lesion that also is known as a canker sore. The lesions are never large, but are annoying and tend to occur during periods of stress. Aphthous ulcers are not infectious; they probably have an autoimmune origin. Herpetic lesions are typically vesicles that can rupture. Leukoplakia appears as white plaques on oral mucosa. Hyperkeratosis. It may be a precursor to squamous cell carcinoma in a few cases. The temperance ditty mentioned in the history is a cautionary note for all young people. Oral thrush is a superficial candidal infection that occurs in diabetic, neutropenic, and immunocompromised patients. Inflammation of a salivary gland (sialadenitis), typically a minor salivary gland in the oral cavity, may produce a localized, tender nodule.

3. B. Chronic irritation is the most likely cause for an “irritation” fibroma of the buccal mucosa, which is due to connective tissue hyperplasia. Oral thrush from candidiasis produces white-to-gray plaques on the tongue. Leukoplakia is hyperplasia of the squamous epithelium and appears as a white plaque or patch, and can be premalignant. A pyogenic granuloma is a reddish nodule of granulation tissue on the gingiva, and it often ulcerates. A minor salivary gland could become obstructed, producing a mucocele, or become inflamed and tender (sialadenitis).

4. A. A pyogenic granuloma may begin to enlarge abruptly and increase in size rapidly, which can be alarming, but the process is benign and often regresses, or resolves into fibrous connective tissue. Though there are both acute and chronic inflammatory cells within this granulation tissue, neither
predominates. Rhabdomyosarcoma is more likely to be a childhood tumor, and sarcomas in adults are more likely to occur in deep soft tissues. This reddish nodule is not leukoplakia, which is a white plaque from squamous epithelial hyperplasia.

PBD9 729–730  BP9 553  PBD8 741–742

5 E The lesions of herpes simplex virus type 1 (HSV-1), also known as cold sores or fever blisters, are common. Many individuals have been infected with HSV-1, which is latent, and the oral and perianal lesions appear during periods of stress. Recurrence of herpes labialis is the norm. Budding cells with pseudohyphae suggest a candidal infection with oral thrush. A mononuclear infiltrate is nonspecific and can be seen with aphthous ulcers. Atypical lymphocytes are seen with infectious mononucleosis. They may be accompanied by a rash, but do not produce vesicular lesions of the skin. Neutrophilic abscesses suggest bacterial infection. Leukoplakia is marked by hyperkeratosis.

PBD9 729  BP9 552  PBD8 742–743  BP8 580–581

6 F Oral thrush is a common but not life-threatening condition, resulting from oral candidiasis in immunocompromised individuals. The lesion is typically superficial. Microscopic examination shows the typical budding cells and pseudohyphae of Candida. Aphthous ulcers, or canker sores, are very common in young individuals, but can appear at any age; they tend to be recurrent superficial ulcerations. Cheilitis is fissuring or cracking of the mucosa, typically at the corners of the mouth, which may be seen with vitamin B2 (riboflavin) deficiency. Hailey leukoplakia also can be seen with HIV infection, but it is far less common than oral thrush. It occurs from marked hyperkeratosis, forming a rough “hairy” surface, and is related to Epstein-Barr virus infection. Multinucleated cells suggest a herpesvirus infection, which typically has vesicles that ulcerate. Atypical squamous epithelial cells usually arise from areas of oral leukoplakia.

PBD9 729–730  BP9 552  PBD8 743  BP8 581

7 C Oral hairy leukoplakia is seen in immunocompromised persons. It presages AIDS in persons who are HIV-positive. Chronic alcohol and/or tobacco use are associated with oral squamous cell carcinomas. Type 1 diabetes mellitus with ketoacidosis is associated with fungal sinusitis, particularly with mucormycosis. Pernicious anemia from vitamin B12 deficiency is associated with glossitis that is mainly atrophic. Sjögren syndrome leads to inflammation and atrophy of salivary glands leading to xerostomia with atrophy, fissuring, and ulcerations in the oral cavity mucosa.

PBD9 730  BP9 554  PBD8 743  BP8 581

8 E This whitish, well-defined mucosal patch on the tongue has the characteristic appearance of leukoplakia, a premalignant lesion that can give rise to squamous cell carcinoma. Use of tobacco products is implicated in the development of leukoplakia. Chronic alcohol abuse also is implicated, but the association is less strong than with tobacco. Ill-fitting dentures may lead to leukoplakia, but far less commonly than smoking. Infections and inflammation are not recognized risk factors for oral leukoplakia or oral squamous cell cancers. Dental caries is not a risk factor for leukoplakia, unless the affected tooth becomes eroded and misshapen. The type of food eaten has less of a correlation with cancer of the oral cavity than with cancer of the esophagus.

PBD9 731  BP9 553–554  PBD8 744–745  BP8 581–582

9 F Erythroplakia is a premalignant lesion that is more likely to progress to squamous carcinoma than leukoplakia, but the major risk factors are the same: tobacco, alcohol, insufficient fruit intake, and betel nut. Countries of the Indian subcontinent have the highest incidence, accounting for up to 10% of all cancers in those populations. Of the remaining options, dental malocclusion may lead to leukoplakia. The oral infections listed are not premalignant, but may be found with immunosuppression. Dietary fruit tends to mitigate the risk, but spices have no effect either way.

PBD9 731  BP9 553–554  PBD8 744–745  BP8 581–582

10 C The raised white patches suggest leukoplakia. This is a premalignant condition. Risk factors include tobacco use, particularly tobacco chewing, and chronic irritation. Human papillomavirus infection has been implicated in some lesions. Oral thrush appears most often on the tongue of immunocompromised individuals as a yellowish plaquelike area. Microscopic examination shows budding cells with pseudohyphae characteristic of Candida infection. Lichen planus in the oral cavity usually appears with similar skin lesions; it forms whitish patches that may ulcerate. The lesions have intense submucosal chronic inflammation. A pyogenic granuloma forms a painful gingival nodule of granulation tissue. Xerostomia, or “dry mouth,” is seen in Sjögren syndrome.

PBD9 731  BP9 553–554  PBD8 744–745  BP8 581–582

11 C Smoking and alcoholism are frequent etiologies for oral squamous cell carcinomas, and mutations in the TP53 gene are often present. However, in nonsmokers, HPV infection may be implicated, along with overexpression of p16. The good news: the oral carcinomas arising with HPV have a better prognosis, though they may be multifocal and recur. The better news: vaccination against HPV may help prevent this disease. Oral candidiasis (thrush) may occur in immunocompromised persons. HSV causes self-limited acute gingivostomatitis (cold sores). The genus Prevotella includes anaerobes that are associated with periodontitis and with buccal infections that become cellulitis (Ludwig angina). Strep throat is an acute exudative pharyngitis that has the immunologic complications of rheumatic heart disease or postinfectious glomerulonephritis.

PBD9 731–733  BP9 554  PBD8 746  BP8 582–583

12 B A dentigerous cyst typically occurs in young persons when teeth are erupting, particularly molars. It is benign and does not recur following complete excision. Dentigerous cysts originate around the crown of an unerupted tooth, typically the third molar, and are lined by a thin, nonkeratinizing layer
of squamous epithelium; they contain a dense chronic inflammatory infiltrate in the stroma. An odontogenic keratocyst that arises from rests of odontogenic epithelium within the jaw and is benign, but can recur if inadequately excised. Ameloblastoma and odontoma are tumors arising from odontogenic epithelium. Odontoma, the most common odontogenic tumor, shows extensive deposition of enamel and dentin. Periapical cysts/granulomas are inflammatory lesions that develop at the apex of teeth as complications of long-standing pulpitis.

13 C An odontogenic keratocyst arises from rests of odontogenic epithelium within the jaw. It is benign, but can recur if inadequately excised. Ameloblastoma and odontoma are tumors arising from odontogenic epithelium. Odontoma, the most common odontogenic tumor, shows extensive deposition of enamel and dentin. Dentigerous cysts originate around the crown of an unerupted tooth, typically the third molar, and are lined by a thin, nonkeratinizing layer of squamous epithelium; they contain a dense chronic inflammatory infiltrate in the stroma. Periapical cysts/granulomas are inflammatory lesions that develop at the apex of teeth as complications of long-standing pulpitis.

14 B Inflammatory nasal polyps can be associated with recurrent allergic rhinitis, a form of type I hypersensitivity often called hay fever. Type I hypersensitivity is associated with high IgE levels in the serum. The elevated hemoglobin A1c level indicates diabetes mellitus. Diabetes is not a risk factor for polyp formation, but ketoacidosis can lead to nasopharyngeal mucormycosis. Epstein-Barr virus infection can be found in nasopharyngeal carcinomas. Autoimmune diseases are not associated with nasal polyp formation. Staphylococcus aureus often colonizes the nasal cavity, but it usually does not cause problems.

15 C Chronic sinusitis is a common condition and may be punctuated by episodes of acute sinusitis. Lack of smell with nasal cavity inflammation often affects sensation of taste. Once the cycle of inflammation, obstruction, stasis, mucociliary damage, and polymicrobial infection is established it becomes difficult to stop. Increased pressure with inflammation in the sinus can erode into adjacent bone, causing osteomyelitis. A mucocele filled with nonpurulent secretions is more likely to occur in frontal and ethmoid sinuses. Sinusitis is not a risk factor for malignancy. Nasopharyngeal carcinomas are related to Epstein-Barr virus (EBV) infection. T-cell lymphomas typically occur in men and are EBV positive. Papillomas most often occur in men and have an exophytic growth pattern, but those that are endophytic aggressively extend into adjacent soft tissue and bone, making removal difficult.

16 C George Washington likely succumbed to an acute bacterial epiglottitis, which is now treatable but still life-threatening, particularly in children, in whom it is more common. Medical care has advanced since the year 1799, but it has been little more than a hundred years that medical care has done more good than harm. Haemophilus influenzae may cause inflammation with an abrupt onset of pain and possible airway obstruction, particularly in children. In adults, the airway is typically long enough to preclude marked obstruction. Thus, Washington’s illness was survivable, but the treatments he received at that time in history (bloodletting, purgatives, blistering agents) contributed to his demise. This cautionary tale supports the adage: if you don’t know what you’re doing, then stop. Coronavirus is best known to cause the common cold. Corynebacterium diphtheriae is the cause of diphtheria, which produces laryngitis with a characteristic dirty gray membrane that may slough and be aspirated. This infection is now rare because of routine childhood immunizations. Another cause for epiglottitis is parainfluenza virus, which has no vaccine, and is best known as the cause for croup in children. The genus Prevotella includes anaerobes that are associated with periodontitis and with buccal infections that become cellulitis (Ludwig angina). Group A streptococci produce a strep throat that is an acute exudative pharyngitis.

17 E The child has croup, a laryngotracheobronchitis that is most often caused by parainfluenza virus. The inflammation may be severe enough to produce airway obstruction. Corynebacterium diphtheriae is the cause of diphtheria, which produces laryngitis with a characteristic dirty gray membrane that may slough and be aspirated. This infection is now rare because of routine childhood immunizations. Epstein-Barr virus may be associated with infectious mononucleosis and produce pharyngitis. Epstein-Barr virus also is associated with nasopharyngeal carcinoma. Haemophilus influenzae may cause an acute bacterial epiglottitis with an abrupt onset of pain and possible airway obstruction. Human papillomavirus is associated with laryngeal papillomatosis. Group A streptococci produce an exudative pharyngitis.

18 A She has a group A β-hemolytic streptococcal pharyngitis, and the feared complication is an autoimmune response from molecular mimicry to streptococcal M proteins. Rheumatic fever results 2 to 3 weeks later from formation of antibodies directed at endocardium, epicardium, and/or myocardium (rheumatic heart disease). Poststreptococcal glomerulonephritis may also occur. The pharyngitis is unlikely to spread elsewhere or produce septicemia. Streptococcus pneumoniae is more likely to produce meningitis, otitis, and pneumonia. Streptococci are unlikely to involve liver.

19 C Nasopharyngeal carcinoma has a strong association with Epstein-Barr virus infection, which contributes to the transformation of squamous epithelial cells. Allergic rhinitis is associated with development of nasal polyps, but these do not become malignant. ANCA-associated vasculitis can involve the respiratory tract, causing granulomatous inflammation and necrotizing vasculitis, but there is no risk of malignant transformation. Sjögren syndrome is associated with
malignant lymphomas, but these typically arise in the salivary gland, not the nasal cavity. Smoking is not associated with nasopharyngeal carcinoma, although it does contribute to oral and esophageal cancers.

PBD9 737–738  BP9 513  PBD8 751–752  BP8 537

20 C Reactive nodules (vocal cord polyps, or singer’s nodules) occur most often in men who are heavy smokers or who strain their vocal cords. The nodules are generally only a few millimeters in size and have a fibrovascular core covered by hyperplastic and hyperkeratotic squamous epithelium. They are not premalignant. Group is an acute laryngotraechobronchitis that most often occurs in children and produces airway narrowing with inspiratory stridor. Epiglottitis is an acute inflammatory process that may cause airway obstruction. Squamous cell carcinomas of the pharynx and larynx form irregular, ulcerating masses, are more common in smokers, but generally are seen in individuals older than this patient. Squamous papillomatosis usually first appears in childhood; if it is extensive, it can produce airway obstruction.

PBD9 739  BP9 513  PBD8 752  BP8 537

21 D Recurrent respiratory papillomatosis is caused by human papillomavirus types 6 and 11. These lesions frequently recur after excision. They may regress after puberty. Laryngeal papillomas arising in adulthood are usually solitary and do not recur. There is no effective antiviral therapy for human papillomavirus. Although the lesions can arise throughout the airways, they are benign and do not become malignant. The occurrence of the lesions is not related to the use of the voice, as is a laryngeal nodule, which is quite small. This is not a congenital condition and is not part of a syndrome.

PBD9 739  BP9 513–414  PBD8 752  BP8 537–538

22 D The figure shows a large, fungating neoplasm that has the typical appearance of a laryngeal squamous cell carcinoma. The most common risk factor is smoking, although chronic alcohol abuse also plays a role; some patients harbor human papillomavirus sequences. Invasive cancers arise from squamous epithelial dysplasias. Epstein-Barr virus infection is associated with nasopharyngeal carcinomas. Aspiration may result in acute inflammation, but not neoplasia. Allergies with type I hypersensitivity may result in transient laryngeal edema, but not neoplasia.

PBD9 739–740  BP9 514  PBD8 753  BP8 538

23 A Cholesteatomas are not true neoplasms, but they are cystic masses lined by squamous epithelium. The desquamated epithelium and keratin degenerates, resulting in cholesterol formation and giant cell reaction. Although their histologic findings are benign, cholesteatomas can gradually enlarge, eroding and destroying the middle ear and surrounding structures. They occur as a complication of chronic otitis media. Although cholesteatomas have a squamous epithelial lining, malignant transformation does not occur. An eosinophilic granuloma of bone occasionally may be seen in the region of the skull in young children, but it is characterized by the presence of Langerhans cells. Labyrinthitis typically is caused by a viral infection and is self-limited. Otosclerosis is abnormal bone deposition in the ossicles of the middle ear that results in bone deafness in adults.

PBD9 740  PBD8 754

24 D Otosclerosis can be familial, particularly when it is severe. It results from fibrous ankylosis followed by bony overgrowth of the little ossicles (malleus, incus, stapes) of the middle ear. A cholesteatoma is typically a unilateral process that complicates chronic otitis media in a child or young adult. Uncomplicated otitis media is usually self-limited and is uncommon in adults. Chondrosarcomas may involve the skull in older adults, but are rare, solitary, bulky masses in the region of the jaw. A schwannoma typically involves the vestibulocochlear nerve and results in a nerve conduction form of deafness. Schwannomas are usually unilateral, although familial neurofibromatosis could result in multiple schwannomas.

PBD9 740–741  PBD8 754

25 A Branchial cysts, also known as lymphoepithelial cysts, may be remnants of an embryonic branchial arch or a salivary gland inclusion in a cervical lymph node. They are distinguished from thyroglossal duct cysts by their lateral location, the absence of thyroid tissue, and their abundant lymphoid tissue. Occult thyroid carcinoma, often a papillary carcinoma, may manifest as a metastasis to a node in the neck, but the microscopic pattern is that of a carcinoma. About 5% of squamous cell carcinomas of the head and neck initially manifest as a nodal metastasis, without an obvious primary site. This patient is quite young for such an event, however. Mucoceles form in minor salivary glands; mucoepidermoid tumors form in salivary glands. The nodule in this patient is in the neck. Paragangliomas are solid tumors that may arise deep in the region of the carotid body near the common carotid bifurcation.

PBD9 741  PBD8 755

26 E A thyroglossal duct (tract) cyst is a developmental abnormality that arises from elements of the embryonic thyroglossal duct extending from the foramen cecum of the tongue down to the thyroid gland. One or more remnants of this tract may enlarge to produce a cystic mass. Although lymphoid tissue often surrounds these cysts, malignant transformation does not occur. Granulomatous disease is more likely to involve lymph nodes in the typical locations in the lateral neck regions. Salivary gland choriostomas are unlikely at this site. The cysts may contain squamous epithelium, but squamous cell carcinoma does not arise from such a cyst. If there is a cystic lesion with lymphoid tissue and squamous carcinoma in the neck, it is probably a metastasis from an occult primary tumor of the head and neck.

PBD9 741  PBD8 755

27 D Paragangliomas are neuroendocrine tumors that rarely produce sufficient catecholamines to affect blood pressure, in contrast to their adrenal medullary counterpart,
The microscopic appearance of these lesions does not always correlate with their biological behavior. There is a tendency for recurrence and metastasis despite the tumor’s “bland” appearance. Metastases always should be considered in patients this age. About 5% of squamous cell carcinomas of the head and neck manifest initially as a nodal metastasis, without an obvious primary site, but the microscopic pattern here is not that of squamous cell carcinoma. Some thyroid cancers initially may manifest as a nodal metastasis, but the microscopic pattern in this case fits best with paraganglioma. A mucoepidermoid carcinoma or a Warthin tumor arises in a salivary gland.

PBD9 741–742  PB8 755–756

**28 B** The most common cause for dry mouth (xerostomia) and dry eyes (xerophthalmia) is a medication effect. Anticholinergics such as trihexyphenidyl to treat the parkinsonian tremor can be implicated, as well as antidepressants, antipsychotics, and antihistaminics. Alcohol and tobacco use are risks for precancerous lesions and squamous cancers of the oral cavity. The lack of saliva is unlikely to be associated with inflammation.

PBD9 742–743  BP9 555  PB8 756  BP8 583

- **29 E** Sialadenitis is more common in older individuals, and individuals receiving therapy for schizophrenia with "typical" antipsychotics such as haloperidol can have reduced salivary secretions, which promotes stasis and infection. Most neuroleptic drugs are dopamine receptor blockers, but they have extrapyramidal and anticholinergic side effects. The dry mouth, coupled with dehydration, favors inspissation of salivary gland secretions and stone formation to block ducts and increase the risk of inflammation and infection. *S. aureus* is the most likely organism to cause infection with supplicative inflammation. Epstein-Barr virus can be associated with hairy leukoplakia. Human papillomavirus infection may lead to the development of squamous dysplasias and carcinomas.

PBD9 743  BP9 555  PB8 756–757  BP8 583

- **30 D** The clinical and histologic features suggest a mucocele of a minor salivary gland, which is most often the result of local trauma in the very young and very old. There is either rupture or blockage of a salivary gland duct. Chili peppers contain capsaicin, which evokes a sensation of tingling and burning pain by activating a nonselective cation channel, called VR1, on vanilloid receptors of sensory nerve endings; there is no significant tissue damage. Social behavior may be a risk factor for infections such as herpes simplex virus. HIV infection is most often associated with oral thrush (candidiasis) and with herpes simplex virus infections. Oral leukoplakia may appear in various intraoral sites and on the lower lip border, and pipe smoking and tobacco chewing are implicated in the development of these white patches. Irritation from misaligned teeth or dentures also may produce leukoplakia. In some parts of the world, the chewing of betel nut is a risk factor for oral cancer.

PBD9 743  BP9 555  PB8 756–757  BP8 583

**31 C** Pleomorphic adenoma is the most common tumor of the parotid gland. These tumors are rarely malignant, although they can be locally invasive. An acinic cell tumor is composed of cells resembling the serous cells of the salivary gland; they are generally small, but about one sixth metastasize to regional lymph nodes. Mucoepidermoid tumors are less common than pleomorphic adenomas in major salivary glands. They may be high-grade and aggressive. Primitive neuroectodermal tumor, also known as an olfactory neuroblastoma, is a small, round, blue cell tumor that occurs in childhood. It is likely to arise in the nasopharyngeal region. Squamous cell carcinomas arise in the buccal mucosa and are invasive. Warthin tumors are uncommon and indolent, although they may be bilateral or multicentric.

PBD9 744–745  BP9 556–557  PB8 758–759  BP8 584–585

**32 F** Warthin tumor is the second most common salivary gland tumor, and it almost always arises within the parotid gland. These tumors tend to be slow growing. Microscopically there are spaces lined by a double layer of superficial columnar and basal cuboidal epithelial cells that are surrounding a lymphoid stroma. Mucoepidermoid carcinomas are infiltrative and form mucous cysts along with a population of squamous cells. Non-Hodgkin lymphoma may arise in patients with long-standing Sjögren syndrome. Pleomorphic adenomas are more common than Warthin tumors, but have a microscopic appearance with ductal epithelial cells in a myxoid stroma containing islands of chondroid and bone. Sialolithiasis is usually accompanied by sialadenitis and is quite painful. It may produce some gland enlargement, but usually is not a mass effect. Sjögren syndrome can produce some salivary gland enlargement, but the process is typically bilateral.

PBD9 745  BP9 556  PB8 759  BP8 584–585

**33 B** Mucoepidermoid carcinomas can arise in major and minor salivary glands. They account for most neoplasms that arise within minor salivary glands, particularly malignant neoplasms. Low-grade mucoepidermoid carcinomas may be invasive, but the prognosis is usually good, with a 5-year survival of 90%. High-grade mucoepidermoid carcinomas can metastasize and have a 5-year survival of only 50%. Non-Hodgkin lymphomas are found in adjacent cervical lymph nodes or in the Waldeyer ring of lymphoid tissue. A primitive neuroectodermal tumor, also known as an olfactory neuroblastoma, is small, round, blue cell tumor of childhood; it is likely to arise in the nasopharyngeal region. Pleomorphic adenomas are more common in the major salivary glands than are mucoepidermoid tumors, and they are more likely to be indolent. Squamous cell carcinomas are invasive and arise in the buccal mucosa. Warthin tumors are uncommon and indolent.

PBD9 745–746  BP9 557  PB8 759–760  BP8 584
1 A 23-year-old primigravida gives birth at term to a boy infant. Ultrasound examination before delivery showed polyhydramnios. A single umbilical artery is seen at the time of birth. The infant vomits all feedings, and then develops a fever and difficulty with respirations within 2 days. A radiograph shows both lungs and the heart are of normal size, but there are pulmonary infiltrates and no stomach bubble. What is the most likely diagnosis?

A Achalasia  
B Diaphragmatic hernia  
C Esophageal atresia  
D Hiatal hernia  
E Pyloric stenosis  
F Zenker diverticulum

2 A 24-year-old man has developed abdominal pain and increasing fatigue over the past 6 months. On physical examination, he is afebrile and appears pale. On palpation, there is mild pain in the right lower quadrant of the abdomen. There are no masses, and bowel sounds are active. Laboratory studies show hemoglobin, 8.9 g/dL; hematocrit, 26.7%; MCV, 74 μm³; platelet count, 255,000/mm³; and WBC count, 7780/mm³. His stool is positive for occult blood. Upper gastrointestinal endoscopy and colonoscopy showed no lesions. One month later, he continues to experience the same abdominal pain. Which of the following is most likely to cause this patient’s illness?

A Acute appendicitis  
B Angiodysplasia  
C Celiac disease  
D Diverticulosis  
E Giardia lamblia infection  
F Meckel diverticulum

3 A 23-year-old woman, G2, P1, gave birth at term to a boy of normal weight and length following an uncomplicated pregnancy. The infant initially did well, but at 6 weeks, he began feeding poorly for 1 week, and his mother noticed that much of the milk he ingested was forcefully vomited within 1 hour. Now, on physical examination, the infant is afebrile, and there are no external anomalies. A midabdominal mass is palpable. Bowel sounds are active. The medical history indicates that both the mother and her first child had the same illness during infancy. Which of the following conditions is most likely to explain these findings?

A Annular pancreas  
B Diaphragmatic hernia  
C Duodenal atresia  
D Pyloric stenosis  
E Tracheoesophageal fistula

4 A 24-year-old woman gives birth to term infant after an uncomplicated pregnancy. Apgar scores are 9 and 10 at 1 and 5 minutes after birth. The infant’s length and weight are at the 55th percentile. There is no significant passage of meconium. Three days after birth, the infant vomits all oral feedings. On physical examination, the infant is afebrile, but the abdomen is distended and tender, and bowel sounds are reduced. An abdominal ultrasound scan shows marked colonic dilation above a narrow segment in the distal sigmoid region. A biopsy specimen from the narrowed region shows an absence of ganglion cells in the muscle wall and submucosa. Which of the following is most likely to produce these findings?

A Colonic atresia  
B Hirschsprung disease  
C Intussusception  
D Necrotizing enterocolitis  
E Trisomy 21  
F Volvulus
A 3-year-old child has attained enough mobility, curiosity, and dexterity to explore places in the home that should not be accessed. The child finds a bottle with a liquid under the kitchen sink, and he drinks it. Within minutes he has chest pain. His mother takes him to the emergency department, and brings the bottle. Analysis of the residual contents reveals a pH of 12. Which of the following complications is most likely to occur following this injury?

A Pharyngeal diverticulum  
B Esophageal stenosis  
C Gastric lymphoma  
D Duodenal ulceration  
E Megacolon

A 22-year-old woman has had multiple episodes of aspiration of food associated with difficulty swallowing during the past year. On auscultation of her chest, crackles are heard at the base of the right lung. A barium swallow shows marked esophageal dilation above the level of the lower esophageal sphincter. A biopsy specimen from the lower esophagus shows an absence of the myenteric ganglia. What is the most likely diagnosis?

A Achalasia  
B Barrett esophagus  
C Plummer-Vinson syndrome  
D Sliding hiatal hernia  
E Systemic sclerosis

A 24-year-old woman living in eastern Bolivia has had increasing difficulty with swallowing both liquids and solids for the past year. She has substernal discomfort from a feeling that foods “get stuck” going down. On examination her BMI is 18. A barium swallow radiologically shows marked esophageal dilation. An endoscopic biopsy is obtained and microscopically shows reduced ganglion cells in myenteric plexus along with lymphocytic infiltration. Which of the following organisms is most likely infecting this woman?

A Bordetella pertussis  
B Candida albicans  
C Corynebacterium diphtheriae  
D Herpes simplex virus  
E Trypanosoma cruzi

A 53-year-old man consumes a very large meal, washed down with considerable alcohol. The ensuing discomfort prompts him to take an emetic, but soon afterward he develops lower chest pain. Physical examination reveals crepitus in subcutaneous tissue over his chest along with tachycardia and tachypnea. Which of the following abnormalities of the esophagus is most likely present in this man?

A Stricture  
B Achalasia  
C Ectopia  
D Rupture  
E Varices

A 30-year-old man has sudden onset of hematemesis after a weekend in which he consumed large amounts of alcohol. The bleeding stops, but he has another episode under similar circumstances 1 month later. Upper gastroesophageal endoscopy shows longitudinal tears at the gastroesophageal junction. What is the most likely mechanism to cause his hematemesis?

A Absent myenteric ganglia  
B Autoimmune inflammation  
C Herpes simplex virus infection  
D Portal hypertension  
E Vomiting  
F Widened diaphragmatic crura

A 16-year-old boy who is receiving chemotherapy for acute lymphoblastic leukemia has had pain for 1 week when he swallows food. Physical examination shows no abnormal findings. Upper gastrointestinal endoscopy shows 0.5- to 0.8-cm mucosal ulcers in the region of the mid to lower esophagus. The shallow ulcers are round and sharply demarcated, and have an erythematous base. Which of the following is most likely to produce these findings?

A Aphthous ulcerations  
B Reflux esophagitis  
C Herpes simplex esophagitis  
D Gastroesophageal reflux disease  
E Mallory-Weiss syndrome

A 44-year-old woman has had increasing difficulty swallowing liquids and solids for the past 6 months. On physical examination, her fingers have reduced mobility because of taut, nondeforming skin. A barium swallow microscopically shows reduced ganglion cells in myenteric plexus along with lymphocytic infiltration. Which of the following is most likely to produce these findings?

A Barrett esophagus  
B Hiatal hernia  
C Iron deficiency  
D Portal hypertension  
E Systemic sclerosis

A 57-year-old woman has had burning epigastric pain after meals for more than 1 year. Physical examination shows no abnormal findings. Upper gastrointestinal endoscopy shows an erythematous patch in the lower esophageal mucosa. A biopsy specimen shows basal zone squamous epithelial hyperplasia, elongation of lamina propria papillae, and scattered intraepithelial neutrophils with some eosinophils. Which of the following is the most likely diagnosis?

A Barrett esophagus  
B Esophageal varices  
C Iron deficiency  
D Reflux esophagitis  
E Systemic sclerosis
A 51-year-old man has sudden onset of massive emesis of bright red blood. On physical examination, his temperature is 36.9°C, pulse is 103/min, respirations are 23/min, and blood pressure is 85/50 mm Hg. His spleen tip is palpable. Laboratory studies show a hematocrit of 21%. The serologic test result for HBsAg is positive. He has had no prior episodes of hematemesis. The hematemesis is most likely to be a consequence of which of the following?

A. Barrett esophagus  
B. *Candida albicans* infection  
C. Esophageal varices  
D. Reflux esophagitis  
E. Squamous cell carcinoma  
F. Zenker diverticulum

A 55-year-old man has had increasing difficulty swallowing during the past 6 months. There are no significant findings on physical examination. Upper gastrointestinal endoscopy shows areas of erythematous mucosa 3 cm above the Z-line. A biopsy specimen from the lower esophagus has changes in the mucosal epithelium illustrated in the figure. Which of the following complications is most likely to occur as a consequence of this patient’s condition?

A. Achalasia  
B. Adenocarcinoma  
C. Diverticular formation  
D. Lacerations (Mallory-Weiss syndrome)  
E. Squamous cell carcinoma

A 68-year-old man from Birmingham, England, has had “heartburn” and substernal pain after meals for 25 years. For the past year, he has had increased pain with difficulty swallowing both liquids and solids. On physical examination, there are no remarkable findings. Upper gastrointestinal endoscopy shows an ulcerated lower esophageal mass that nearly occludes the lumen of the esophagus. A biopsy specimen of this mass is most likely to show which of the following neoplasms?

A. Adenocarcinoma  
B. Carcinoid tumor  
C. Leiomyosarcoma  
D. Non-Hodgkin lymphoma  
E. Squamous cell carcinoma

A 73-year-old man with a history of chronic alcoholism has had increasing difficulty swallowing and has noticed a 3-kg weight loss over the past 2 months. On physical examination, there are no remarkable findings. Upper gastrointestinal endoscopy shows a 3-cm ulcerative mass in the midesophagus that partially occludes the esophageal lumen. Esophagectomy is performed; the gross appearance of the lesion is shown in the figure. Which of the following is most likely to be seen on microscopic section of this mass?

A. Adenocarcinoma  
B. Dense collagenous scar  
C. Dilated vascular channels  
D. Multinucleated cells with intranuclear inclusions  
E. Squamous cell carcinoma

A 66-year-old man living in Tehran, Iran, has been bothered by difficulty swallowing for the past year. He is now consuming liquid food. Yesterday he regurgitated food stained with blood. On esophagoscopy, there is an ulcerated obstructing lesion 20 cm from the lips. Biopsies are taken and on microscopic show infiltrating nests of keratinized cells with distinct cell borders and hyperchromatic, angulated nuclei. Which of the following is the most likely risk factor for his disease?

A. Genetic susceptibility  
B. Autoimmunity  
C. Diet  
D. Infection  
E. Reflux
18. A 38-year-old woman has had nausea for 6 months. She reports no vomiting or diarrhea. On physical examination, there are no remarkable findings. Upper gastrointestinal endoscopy shows diffuse gastric mucosal erythema with focal mucosal erosions, but no ulcerations. The esophageal and duodenal mucosal surfaces appear normal. Gastric biopsies are obtained and microscopic examination shows focal mucosal hemorrhage, loss of the surface epithelium, and increased numbers of neutrophils, lymphocytes, and plasma cells in an edematous mucosa. No *Helicobacter pylori* organisms are seen. Laboratory studies show a normal serum gastrin level. Which of the following pharmacologic agents is most likely to produce these findings?

A. Aspirin  
B. Chlorpromazine  
C. Cimetidine  
D. Clindamycin  
E. Omeprazole

19. A 72-year-old man takes large quantities of nonsteroidal anti-inflammatory drugs (NSAIDs) because of chronic degenerative arthritis of the hips and knees. Over the past 2 weeks, he has had epigastric pain with nausea and vomiting and an episode of hematemesis. On physical examination, there are no remarkable findings. A gastric biopsy specimen is most likely to show which of the following lesions?

A. Acute gastritis  
B. Adenocarcinoma  
C. Epithelial dysplasia  
D. *Helicobacter pylori* infection  
E. Hyperplastic polyp

20. A 54-year-old, previously healthy man sustained an extensive thermal burn injury involving 70% of the total body surface area of his skin. He was hospitalized in stable condition. Three weeks after the initial burn injury, he developed melanotic stools. His blood pressure dropped to 80/40 mm Hg, and his hematocrit declined to 18%. Where are gastrointestinal ulcerations most likely to be found in this man?

A. Colon  
B. Duodenum  
C. Esophagus  
D. Ileum  
E. Stomach

21. A 51-year-old woman has been feeling increasingly tired for the past 7 months. There are no remarkable findings on physical examination. Laboratory studies include hemoglobin, 9.5 g/dL; hematocrit, 29.1%; MCV, 124 μm³; platelet count, 268,000/mm³; and WBC count, 8350/mm³. The reticulocyte index is low. Hypersegmented polymorphonuclear leukocytes are found on a peripheral blood smear. The serum gastrin is markedly increased. Antibodies to which of the following are most likely to be found in this patient?

A. Gastric H⁺,K⁺-ATPase  
B. Gliadin  
C. *Helicobacter pylori*  
D. Intrinsic factor receptor  
E. *Tropheryma whippelli*

22. A 59-year-old man has had nausea and vomiting for 5 months. He has experienced no hematemesis. On physical examination, there is no abdominal tenderness, and bowel sounds are present. Upper gastrointestinal endoscopy shows erythematous areas of mucosa with thickening of the rugal folds in the gastric antrum. The microscopic appearance of a gastric biopsy specimen with a Steiner silver stain is shown in the figure. Which of the following factors is most likely responsible for this gastric mucosal pathology?

A. Cysteine proteinase  
B. Cytotoxin-associated gene A  
C. Heat-stable enterotoxin  
D. Shiga toxin  
E. Verocytotoxin

23. A 47-year-old woman with a lengthy history of heartburn and dyspepsia experiences sudden onset of abdominal pain. On physical examination, she has severe mid epigastric pain with guarding. Bowel sounds are reduced. An abdominal plain film radiograph shows free air under the left leaf of the diaphragm. She is immediately taken to surgery, and a perforated duodenal ulcer is repaired. Which of the following organisms is most likely to have produced these findings?

A. Campylobacter jejuni  
B. Cryptosporidium parvum  
C. Giardia lamblia  
D. *Helicobacter pylori*  
E. Salmonella typhi  
F. Shigella flexneri  
G. *Yersinia enterocolitica*

24. A 35-year-old man has had epigastric pain for more than 1 year. The pain tends to occur 2 to 3 hours after a meal and is relieved if he takes antacids or eats more food. He has noticed a 4-kg weight gain in the past year. He does not smoke and drinks 1 glass of Johannisberg Riesling daily. The result of a urea breath test is positive, and a gastric biopsy specimen contains urease. He begins a 2-week course of antibiotics, but on day 4, he feels better and discontinues treatment. Three weeks later, the epigastric pain recurs. If he does not seek further treatment, which of the following complications is he most likely to develop?

A. Carcinoid syndrome  
B. Fat malabsorption  
C. Hematemesis  
D. Migratory thrombophlebitis  
E. Vitamin B₁₂ deficiency
25. A 52-year-old man notes nausea with abdominal discomfort after meals. On physical examination, there are no abnormal findings. Upper endoscopy is performed, and there are three ovoid nodules in the fundus and antrum ranging from 0.3 to 1.2 cm in size. They have rounded, smooth surfaces. Biopsies are taken and on microscopic examination there are irregular, cystically dilated and elongated foveolar glands. Which of the following treatment strategies is most appropriate for his gastric lesions?

A. Antibiotics  
B. Chemotherapy  
C. Corticosteroids  
D. Multivitamins  
E. Total gastrectomy  
F. Vagotomy

26. A 49-year-old woman has a history of peptic ulcer disease for which she has been treated with proton pump inhibitors. She has had nausea with vomiting for the past 2 months. Upper GI endoscopy reveals three circumscribed, round, smooth lesions in the gastric body from 1 to 2 cm in diameter. Biopsies are taken and microscopically show the lesions to consist of irregular glands that are cystically dilated and lined by flattened parietal and chief cells. No inflammation, Helicobacter pylori, metaplasia, or dysplasia is present. What is the most likely diagnosis?

A. Fundic gland polyps  
B. Gastric adenomas  
C. Hyperplastic polyps  
D. Hypertrophic gastropathy

27. A 53-year-old woman has had nausea, vomiting, and midepigastric pain for 5 months. On physical examination, there are no significant findings. An abdominal CT scan shows gastric outlet obstruction. Upper gastrointestinal endoscopy shows an ulcerated 2 × 4 cm bulky mass in the antrum at the pylorus. A urease test is positive. Which of the following neoplasms is most likely to be seen in a biopsy specimen of this mass?

A. Adenocarcinoma  
B. Leiomyosarcoma  
C. Neuroendocrine carcinoma  
D. Non-Hodgkin lymphoma  
E. Squamous cell carcinoma

28. A 67-year-old woman has experienced severe nausea, vomiting, early satiety, and a 9-kg weight loss over the past 4 months. On physical examination, she has muscle wasting. Upper gastrointestinal endoscopy shows that the entire gastric mucosa is eroded and has an erythematous, cobblestone appearance. An abdominal CT scan shows that the stomach is small and shrunken. Which of the following is most likely to be found on histologic examination of a gastric biopsy specimen?

A. Chronic atrophic gastritis  
B. Primary gastric lymphoma  
C. Gastrointestinal stromal tumor  
D. Granulomatous inflammation  
E. Signet ring cell adenocarcinoma

29. A 52-year-old man has had a 4-kg weight loss and nausea for the past 6 months. He has no vomiting or diarrhea. On physical examination, there are no remarkable findings. Upper gastrointestinal endoscopy shows a 6-cm area of irregular, pale fundic mucosa and loss of the rugal folds. A biopsy specimen shows a monomorphous infiltrate of lymphoid cells microscopically. Helicobacter pylori organisms are identified in mucus overlying adjacent mucosa. Cytogenetic analysis shows t(11;18)(q21;q21). He receives antibiotic therapy for H. pylori, and the repeat biopsy specimen shows a resolution of the infiltrate. What is the most likely diagnosis?

A. Autoimmune gastritis  
B. Chronic gastritis  
C. Crohn disease  
D. Diffuse large B-cell lymphoma  
E. Gastrointestinal stromal tumor  
F. Mucosa-associated lymphoid tissue tumor

30. A 26-year-old man is brought to the emergency department after sustaining abdominal gunshot injuries. At laparotomy, while repairing the small intestine, the surgeon notices a 1-cm mass at the tip of the appendix. The yellow-tan submucosal mass is removed, and the microscopic appearance of the mass is shown in the figure. Immunohistochemical staining is positive for chromogranin and synaptophysin but negative for Ki-67. Which of the following is the most likely cell of origin of this lesion?

A. Lipoblast  
B. Ganglion cell  
C. Goblet cell  
D. Neuroendocrine cell  
E. Smooth muscle cell

31. A 55-year-old man experiences episodes of diaphoresis, dyspnea, and diarrhea for 10 months. On physical examination he has midabdominal discomfort with deep palpation, and bowel sounds are reduced. There are no abnormal findings with upper endoscopy. Abdominal CT scan shows three nodules in the liver, from 1 to 3 cm in size. Laboratory studies show a high level of serum 5-hydroxyindoleacetic acid (5-HIAA). Camera endoscopy is performed, and on review of the images, there is a midjejunal mass that partially obstructs the lumen. At laparotomy a 5-cm submucosal jejunal mass is resected, and on microscopy it is composed of nests and trabeculae of round cells with pink, granular cytoplasm. The cells of this mass are most likely related to which of the following embryologic derivatives?

A. Endoderm  
B. Ectoderm  
C. Neural crest  
D. Notochord  
E. Splanchnic mesoderm
32. A 61-year-old man with increasing fatigue, early satiety, and nausea for 5 months vomited dark granular material yesterday. Endoscopy reveals a large ulcerated mass in the gastric fundus. Biopsies are taken and microscopically the mass is composed of spindle cells that are positive for c-Kit with immunohistochemical staining. Mitoses are frequent. Gastrectomy is performed, and the 10-cm circumscribed mass arises from the gastric wall. Which of the following is most likely to have produced his findings?

A. Amoxicillin
B. Azathioprine
C. Cyclophosphamide
D. Imatinib
E. Prednisone
F. Radiation

33. A 57-year-old man from Innsbruck, Austria, goes to the emergency department because of increasing abdominal pain with distention that developed over the past 24 hours. On physical examination, there is diffuse abdominal tenderness. The abdomen is tympanitic, without a fluid wave, and bowel sounds are nearly absent. There is a well-healed, 5-cm transverse scar in the right lower quadrant of the abdomen. There is no caput medusa. A stool sample is negative for occult blood. An abdominal plain film shows dilated loops of small bowel with air-fluid levels, but there is no free air. At laparotomy, the surgeon notices a 20-cm portion of reddish black ileum with changes abruptly to pink-appearing bowel on distal and proximal margins. His medical history is significant only for an appendectomy at age 25 years. Which of the following is most likely to be a useful adjunct in treatment of his disease?

A. Appendicitis
B. Cholecystitis
C. Pancreatitis
D. Intestinal infarction
E. Pseudomembranous colitis

34. An 11-month-old, previously healthy infant has not produced a stool for 1 day. The mother notices that the infant’s abdomen is distended. On physical examination, the infant’s abdomen is very tender, and bowel sounds are nearly absent. An abdominal plain film radiograph shows no free air, but there are distended loops of small bowel with air-fluid levels. Which of the following is most likely to produce these findings?

A. Duodenal atresia
B. Hirschsprung disease
C. Intussusception
D. Meckel diverticulum
E. Pyloric stenosis

35. A 61-year-old man has had severe abdominal pain and bloody diarrhea for the past day. On physical examination, his abdomen is diffusely tender, and bowel sounds are absent. Abdominal plain films show no free air. Laboratory studies show a normal CBC and normal levels of serum amylase, lipase, and bilirubin. His Hgb A<sub>1c</sub> is 10%. He develops shock. A year ago he had an acute myocardial infarction. Which of the following lesions is most likely to be found in this man?

A. Appendicitis
B. Cholecystitis
C. Pancreatitis
D. Intestinal infarction
E. Pseudomembranous colitis

36. A 71-year-old woman with a history of rheumatic heart disease is hospitalized with severe congestive heart failure. Four days after admission, she develops cramping lower abdominal pain. On physical examination, she is afebrile. The abdomen is distended and tympanic, without a fluid wave, and bowel sounds are absent. A stool sample is positive for occult blood. An abdominal plain film shows no free air. Colonoscopy shows patchy areas of mucosal erythema with some overlying tan exudate in the ascending and descending colon. No polyps or masses are found. What is the most likely diagnosis?

A. Ischemic colitis
B. Mesenteric vasculitis
C. Shigellosis
D. Ulcerative colitis
E. Volvulus

37. A 60-year-old man has had increasing fatigue for the past 8 months. On physical examination, he appears pale. On digital rectal examination, no masses are palpable, but a stool sample is positive for occult blood. Auscultation of the abdomen shows active bowel sounds, and on palpation there are no masses or areas of tenderness. Laboratory studies show hemoglobin, 8.3 g/dL; hematocrit, 24.6%; MCV, 73 μm³; platelet count, 226,000/mm³; and WBC count, 7640/mm³. Colonoscopy shows no identifiable source of the bleeding. Angiography shows a 1-cm focus of dilated and tortuous vascular channels in the mucosa and submucosa of the cecum. What is the most likely diagnosis?

A. Angiodysplasia
B. Collagenous colitis
C. Diverticulosis
D. Internal hemorrhoids
E. Mesenteric vein thrombosis

38. A 21-year-old man has had increasingly voluminous, bulky, foul-smelling stools and a 7-kg weight loss for the past year. There is no history of hematemesis or melena. He has some bloating, but no abdominal pain. On physical examination, there are no palpable abdominal masses, and bowel sounds are present. Which of the following laboratory findings is most likely to be present on examination of his stool?

A. Entamoeba histolytica trophozoites
B. Giardia lamblia cysts
C. Increased stool fat
D. Occult blood
E. Vibrio cholerae organisms
39 A 34-year-old woman is bothered by a low-volume, mostly watery diarrhea associated with flatulence. The symptoms occur episodically, but they have been persistent for the past year. She has experienced a 4-kg weight loss. She has no fever, nausea, vomiting, or abdominal pain. On physical examination, there are no significant findings. A stool sample is negative for occult blood, ova, and parasites, and a stool culture yields no pathogens. An upper gastrointestinal endoscopy is performed and a biopsy specimen from the upper part of the small bowel shows severe diffuse blunting of villi and a chronic inflammatory infiltrate in the lamina propria. Which of the following serologic tests is most likely to be positive in this patient?

A Anticentromere antibody
B Anti-DNA topoisomerase I antibody
C Anti mitochondri al antibody
D Antinuclear antibody
E Antitransglutaminase antibody

40 A 41-year-old woman has had diarrhea and fatigue with a 3-kg weight loss over the past 6 months. On physical examination, she is afebrile and has mild muscle wasting, but her motor strength is normal. Laboratory studies show no occult blood, ova, or parasites in the stool. A biopsy specimen from the upper jejunum is obtained, and microscopic findings are reviewed. The patient begins following a special diet with no wheat or rye grain products. The change in diet produces dramatic improvement. Which of the following microscopic features is most likely to be seen in the biopsy specimen?

A Crypt abscesses and mucosal ulceration
B Foamy macrophages within the lamina propria
C Lymphatic obstruction
D Noncaseating granulomas
E Villous blunting and flattening

41 An epidemiologic study of children with failure to thrive is undertaken in Guatemala. Some of these children with ages 1 to 3 years have repeated bouts of diarrhea, but do not improve with dietary supplements. Jejunal biopsies show blunted, atrophic villi with crypt elongation and chronic inflammatory infiltrates. What is the most likely factor contributing to recurrent diarrhea in these children?

A Abetalipoproteinemia
B Bacterial infection
C Chloride ion channel dysfunction
D Disaccharidase deficiency
E NOD2 gene mutations

43 A potluck lunch party is held at the office at noon. Various meats, salads, breads, and desserts that were brought in earlier that morning are served. Everyone has a good time, and most of the food is consumed. By midafternoon, the single office restroom is being used by many employees who have vomiting and acute, explosive diarrhea accompanied by abdominal cramping. Which of the following infectious agents is most likely responsible for this turn of events?

A Bacillus cereus
B Clostridium difficile
C Escherichia coli
D Salmonella enterica
E Staphylococcus aureus
F Vibrio parahaemolyticus

44 A healthy 21-year-old woman develops a profuse, watery diarrhea 1 day after a meal of raw oysters. On physical examination, her temperature is 37.5°C. A stool sample is negative for occult blood. There is no abdominal distention or tenderness, and bowel sounds are present. The diarrhea subsides over the next 3 days. Which of the following organisms is most likely to produce these findings?

A Cryptosporidium parvum
B Entamoeba histolytica
C Staphylococcus aureus
D Vibrio parahaemolyticus
E Yersinia enterocolitica

45 A 26-year-old man traveling to Ching Mai, Thailand, had fever, headache, and muscle pains for a day followed by watery diarrhea of 5 to 10 stools per day for 4 days. In the past day, the diarrhea has been bloody and accompanied by tenesmus. On physical examination there is diffuse abdominal pain. Microscopic examination of the stool shows numerous leukocytes and gram-negative curved rods. The diarrhea subsides, but 2 weeks later he has increasing weakness in his legs. Which of the following organisms is most likely to produce his disease?

A Bacillus cereus
B Campylobacter jejuni
C Clostridium perfringens
D Giardia lamblia
E Rotavirus

46 A 36-year-old man experiences cramping abdominal pain with fever and watery diarrhea 2 days after eating a chicken salad sandwich. Physical examination shows mild diffuse abdominal pain on palpation, but there are no masses. Bowel sounds are present. A stool sample is negative for occult blood. He recovers completely within 5 days without treatment. Which of the following infectious organisms is most likely to produce these findings?

A Bacillus cereus
B Entamoeba histolytica
C Escherichia coli
D Rotavirus
E Salmonella enterica
F Staphylococcus aureus
G Yersinia enterocolitica
In an epidemiologic study of infections of the gastrointestinal tract, cases of patients living in Haiti from whom definitive cultures were obtained are analyzed for clinical and pathologic findings that may be useful for diagnosis. A group of patients is identified who initially had abdominal pain and diarrhea during week 1 of their illness. By week 2, these patients had splenomegaly and elevations in serum AST and ALT levels. By week 3, they were septic and had leukopenia. At autopsy, the patients who died were found to have ulceration of Peyer patches. Which of the following infectious agents is most likely to produce these findings?

A. Campylobacter jejuni  
B. Clostridium perfringens  
C. Mycobacterium bovis  
D. Salmonella typhi  
E. Shigella sonnei  
F. Yersinia enterocolitica  

A 65-year-old woman is being treated in the hospital for pneumonia complicated by septicemia. She has required multiple antibiotics and was intubated and mechanically ventilated earlier in the course. On day 20 of hospitalization, she has abdominal distention. Bowel sounds are absent, and an abdominal radiograph shows dilated loops of small bowel suggestive of ileus. She has a low volume of bloody stool that is positive for Clostridium difficile toxin. Laboratory studies show leukocytosis and hypoalbuminemia. At laparotomy, a portion of distal ileum and cecum is resected. The gross appearance of the mucosal surface is shown in the figure. What is the most likely diagnosis?

A. Gas gangrene with myonecrosis  
B. Inflammatory bowel disease  
C. Ischemic bowel disease  
D. Pseudomembranous enterocolitis  
E. Toxic megacolon  

Over a holiday weekend, more than 100 adults at a resort hotel develop a diarrheal illness marked by voluminous, watery stools more than 10 times per day. They also report headache, abdominal cramping pain, and myalgias. On physical examination they have manifestations of dehydration and mild fever. Laboratory studies of stool samples show no increase in leukocytes or fat, and no RBCs. Their illness lasts just 1 to 3 days and resolves with no sequelae. Which of the following infectious agents is the most likely cause for their illness?

A. Cytomegalovirus  
B. Clostridium botulinum  
C. Norovirus  
D. Staphylococcus aureus  
E. Strongyloides stercoralis  
F. Vibrio cholerae  

A 5-month-old, previously healthy infant girl in Bangladesh develops a watery diarrhea that lasts for 1 week. The infant has a mild fever during the illness, but has no abdominal pain or swelling. On physical examination, her temperature is 37.7° C. A stool sample is negative for occult blood, ova, or parasites. Her parents are told to give her plenty of fluids, and she recovers fully. Which of the following organisms is most likely to produce these findings?

A. Campylobacter jejuni  
B. Cryptosporidium parvum  
C. Escherichia coli  
D. Listeria monocytogenes  
E. Norwalk virus  
F. Rotavirus  
G. Shigella flexneri  

A study of children living in rural Malawi in Africa reveals a high prevalence of iron deficiency anemia. Stool samples are positive for occult blood. Pruritus of the skin of their feet as well as cough are additional findings in many of these children. Which of the following parasitic infestations is the most likely cause for these findings?

A. Ancylostoma duodenale  
B. Ascaris lumbricoides  
C. Cryptosporidium parvum  
D. Enterobius vermicularis  
E. Schistosoma mansoni  

A 31-year-old woman had increasingly severe diarrhea 1 week after returning from a trip to Central America. Gross examination of the stools showed mucus and streaks of blood. The diarrheal illness subsided within 4 weeks, but now she has become febrile and has pain in the right upper quadrant of the abdomen. An abdominal ultrasound scan shows a 10-cm, irregular, partly cystic mass in the right hepatic lobe. Which of the following infectious organisms is most likely to produce these findings?

A. Clostridium difficile  
B. Cryptosporidium parvum  
C. Giardia lamblia  
D. Entamoeba histolytica  
E. Strongyloides stercoralis
A 27-year-old man has sudden onset of marked abdominal pain. On physical examination, his abdomen is diffusely tender and distended, and bowel sounds are absent. He undergoes surgery, and a 27-cm segment of terminal ileum with a firm, erythematous serosal surface is removed. The microscopic appearance of a section through the excised ileum is shown in the figure. Which of the following additional complications is the patient most likely to develop as a result of this disease process?

A. Adenocarcinoma
B. Enterocutaneous fistula
C. Intussusception
D. Liver abscess
E. Mesenteric artery thrombosis

A 30-year-old woman has a 5-year history of recurrent episodes marked by days of abdominal bloating with alternating constipation and diarrhea. She notes hard stools of narrow caliber, low volume mucous diarrhea, and pain in the left lower quadrant. Her symptoms are relieved by defecation, which occurs more frequently now. On physical examination there are no abnormal findings. Laboratory studies including stool for ova and parasites, bacterial pathogens, and fat show no abnormalities. An abdominal CT scan is unremarkable. What is the most likely diagnosis?

A. Cystic fibrosis
B. Diverticulitis
C. Fat malabsorption
D. Perirectal fistula formation
E. Primary biliary cirrhosis
F. Pseudomembranous colitis

A 49-year-old woman has had abdominal cramps and diarrhea with six stools per day for the past month. She has a history of similar episodes of self-limited pain and diarrhea, which have occurred multiple times during the past 20 years. Each episode lasts about 2 weeks and resolves without treatment. Findings on physical examination are unremarkable, but a stool sample is positive for occult blood. Laboratory studies show no ova or parasites in the stool. Colonoscopy shows diffuse and uninterrupted mucosal inflammation and superficial ulceration extending from the rectum to the ascending colon. Colonic biopsy specimens from the area show the findings in the figure. She is at greatest risk for developing which of the following complications?

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B. Diverticulitis
C. Fat malabsorption
D. Perirectal fistula formation
E. Primary biliary cirrhosis
F. Pseudomembranous colitis

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A 35-year-old woman has had increasing lower back pain for 5 years. During the past year she also has had arthritic pain involving the knees, hips, and wrists. A stool sample is positive for occult blood. A pelvic radiograph shows changes consistent with sacroilitis. A colonoscopy is performed, and she undergoes a total colectomy. The figure shows the gross appearance of the colectomy specimen. What is the most likely underlying mechanism of the illustrated condition?

A  Development of autoantibodies directed against tropomyosin
B  Development of antimicrobial antibodies that cross react with colonic mucosa
C  Development of T_{H}17 immune responses
D  Germline inheritance of the APC gene mutation
E  Mutations in the NOD2 gene

A 30-year-old woman has suffered intermittent bouts of lower abdominal pain and low-volume diarrhea for the past 2 years. On colonoscopy there is friable mucosa from the rectum to the ascending colon, and a perianal fistula is noted. Biopsies are taken and on microscopic examination show acute and chronic mucosal inflammation with focal erosion. Her stool is negative for ova, parasites, and bacterial pathogens. Which of the following ongoing testing procedures is most useful for long-term follow-up of this woman?

A  Abdominal CT scanning
B  Biopsy screening for dysplasia
C  Genetic mutational analysis for NOD2
D  Serologic titers for Saccharomyces
E  Stool cultures for microbiota

A 26-year-old man has had intermittent cramping abdominal pain and low-volume diarrhea for 3 weeks. On physical examination, he is afebrile; there is mild lower abdominal tenderness but no masses, and bowel sounds are present. A stool sample is positive for occult blood. The symptoms subside within 1 week. Six months later, the abdominal pain recurs with perianal pain. On physical examination, there is now a perirectal fistula. Colonoscopy shows many areas of mucosal edema and ulceration and some areas that appear normal. Microscopic examination of a biopsy specimen from an ulcerated area shows a patchy acute and chronic inflammatory infiltrate, crypt abscesses, and noncaseating granulomas. Which of the following underlying disease processes best explains these findings?

A  Amebiasis
B  Crohn disease
C  Sarcoidosis
D  Shigellosis
E  Ulcerative colitis

A clinical study of adult patients with chronic bloody diarrhea is performed. One group of these patients is found to have a statistically increased likelihood for the following: antibodies to Saccharomyces cerevisiae but not anti-neutrophil cytoplasmic autoantibodies, NOD2 gene polymorphisms, T_{H}1 and T_{H}17 immune cell activation, vitamin K deficiency, megaloblastic anemia, and gallstones. Which of the following diseases is this group of patients most likely to have?

A  Angiodysplasia
B  Crohn disease
C  Diverticulitis
D  Ischemic enteritis
E  Ulcerative colitis

A 65-year-old woman has a routine health maintenance examination. A stool sample is positive for occult blood. CT scan of the abdomen shows numerous air-filled, 1-cm outpouchings of the sigmoid and descending colon. Which of the following complications is most likely to develop in this patient?

A  Adenocarcinoma
B  Bowel obstruction
C  Pericolic abscess
D  Malabsorption
E  Toxic megacolon

The mother of a 4-year-old child notes blood when laundering his underwear. Physical examination reveals a rectal mass. On proctoscopy, there is a smooth-surfaced, pedunculated, 1.5-cm polyp. It is excised and microscopically shows cystically dilated crypts filled with mucus and inflammatory debris, but no dysplasia. What is the most likely diagnosis?

A  Familial adenomatous polyposis
B  Gardner syndrome
C  Juvenile polyp
D  Lynch syndrome
E  Peutz-Jeghers syndrome
A 53-year-old woman undergoes a routine checkup. The only abnormal finding is a stool specimen that contains occult blood. Colonoscopy shows a 1.5-cm, solitary, rounded, erythematous polyp on a 0.5-cm stalk at the splenic flexure. The polyp is removed; its histologic appearance is shown in the figure at low (A) and high (B) magnifications. Her colonic lesion is most likely associated with which of the following?

A  Low risk for development of carcinoma  
B  Inheritance of an abnormal tumor suppressor gene  
C  Presence of similar lesions in the small intestine  
D  History of iron deficiency anemia  
E  Risk for development of endometrial carcinoma  

A 70-year-old man has a routine health maintenance examination. On physical examination, there are no remarkable findings, but a stool sample is positive for occult blood. A colonoscopy is performed and shows a 5-cm sessile mass in the upper portion of the descending colon at 50 cm from the anal verge. The histologic appearance at low power of a biopsy specimen of the lesion is shown in the figure. The patient refused further workup and treatment. Five years later, he has constipation, microcytic anemia, and a 5-kg weight loss over 6 months. On surgical exploration, there is a 7-cm mass encircling the descending colon. Which of the following neoplasms is he now most likely to have?

A  Adenocarcinoma  
B  Non-Hodgkin lymphoma  
C  Carcinoïd tumor  
D  Leiomyosarcoma  
E  Mucinous cystadenoma  
F  Villous adenoma
64] A 19-year-old man is advised to see his physician because genetic screening has detected a disease in other family members. On physical examination, a stool sample is positive for occult blood. A colonoscopy is performed, followed by a colectomy. The figure shows the gross appearance of the mucosal surface of the colectomy specimen. Microscopic examination shows these lesions are tubular adenomas. Molecular analysis of this patient’s normal fibroblasts is most likely to show a mutation in which of the following genes?

A  APC  
B  MLH1  
C  KRAS  
D  NOD2  
E  p53

65] A 44-year-old woman has had increasing abdominal distention for the past 6 weeks. On physical examination, there is an abdominal fluid wave, and bowel sounds are present. Paracentesis yields 1000 mL of slightly cloudy serous fluid. Cytologic examination of the fluid shows malignant cells consistent with adenocarcinoma. Molecular analysis of these cells shows an MSH2 gene mutation with microsatellite instability. Her medical history indicates that she has had no major medical illnesses and no surgical procedures. Her sister was diagnosed with endometrial cancer and her brother had carcinoma of the stomach. Which of the following conditions is the most likely cause of this patient’s symptoms?

A  Angiodysplasia  
B  Crohn disease  
C  Diverticulosis  
D  Lynch syndrome  
E  Peptic ulcer disease

66] A 33-year-old man has a routine health maintenance examination. A stool sample is positive for occult blood. On colonoscopy, a 6-cm ulcerative lesion is seen projecting into the cecum. There are three smaller sessile lesions from 1 to 3 cm in size. The microscopic appearance of a section of the ulcerated lesion is shown in the figure. The smaller lesions are reported as sessile serrated adenomas. Which of the following molecular biological events is thought to be most critical in the development of such lesions?

A  Amplification of ERBB2 gene  
B  Defective DNA mismatch repair gene  
C  Germline transmission of a defective RB1 gene  
D  Overexpression of E-cadherin gene  
E  Translocation of retinoic acid receptor alpha gene

67] A 73-year-old man has noted a change in bowel habits for the past year. Defecation is more difficult and the caliber of stools has decreased. On physical exam, there are no abnormal findings except for stool positive for occult blood. Colonoscopy is performed for the first time in this man, followed by colonic resection with the gross appearance shown in the figure. Which of the following molecular abnormalities has most likely led to these findings?

A  Acquired APC gene mutation  
B  Homozygous loss of PTEN gene  
C  Inactivation of the RB1 protein by HPV-16  
D  Mutation in a DNA mismatch repair gene  
E  Tyrosine kinase activation with KIT mutation
68  A 20-year-old woman has increasing pain on defecation and notices bright red blood on the toilet paper. She has had no previous gastrointestinal problems. After she gives birth, the rectal pain subsides, and there is no more bleeding. Which of the following is the most likely cause of these findings?

A  Angiodyplasia
B  Hemorrhoids
C  Intussusception
D  Ischemic colitis
E  Volvulus

69  A 20-year-old woman has had nausea and vague lower abdominal pain for the past 24 hours, but now the pain has become more severe. On physical examination, the pain is worse in the right lower quadrant, and there is rebound tenderness. A stool sample is negative for occult blood. Abdominal plain film radiographs show no free air. The result of a serum pregnancy test is negative. Which of the following laboratory findings is most useful to aid in the diagnosis of this patient?

A  Entamoeba histolytica cysts in the stool
B  Hyperamylasemia
C  Hypernatremia
D  Increased serum alkaline phosphatase
E  Increased serum carcinoembryonic antigen
F  Neutrophilia with left shift

70  A 53-year-old woman has increasing abdominal girth for the past 2 years. On physical examination she has abdominal distension. An abdominal CT scan shows multiple nodules on peritoneal surfaces along with low attenuation mucinous ascites. Paracentesis is performed and cytologic examination of the fluid obtained shows well-differentiated columnar cells with minimal nuclear atypia. Where did this proliferative process most likely arise in this woman?

A  Appendix
B  Jejunum
C  Ileum
D  Pancreas
E  Stomach

71  A 59-year-old man with a lengthy history of chronic alcoholism has noticed increasing abdominal girth for the past 6 months. He has had increasing abdominal pain for the past 2 days. On physical examination, his temperature is 38.2° C. Examination of the abdomen shows a fluid wave and prominent caput medusae over the skin of the abdomen. There is diffuse abdominal tenderness. An abdominal plain film radiograph shows no free air. Paracentesis yields 500 mL of cloudy yellow fluid. Gram stain of the fluid shows gram-negative rods. Which of the following is the most likely diagnosis?

A  Appendicitis
B  Collagenous colitis
C  Diverticulitis
D  Ischemic colitis
E  Pseudomembranous colitis
F  Spontaneous bacterial peritonitis

**Answers**

1  An esophageal atresia is often combined with a fistula between the esophagus and trachea. Gastrointestinal obstruction in utero can lead to polyhydramnios. The presence of a single umbilical artery suggests additional anomalies are present. Vomiting in an infant risks aspiration with development of pneumonia. Achalasia is incomplete relaxation of the lower esophageal sphincter and is usually not manifested at birth. Absence of a diaphragmatic leaf, usually on the left, results in herniation of abdominal contents into the chest and functional gastrointestinal obstruction, but in this case normal-sized lungs suggest no herniated contents were present. A hiatal hernia from widened diaphragmatic muscular crura predisposes to gastroesophageal reflux, and obstruction is not a typical complication. pyloric stenosis is a cause for gastric outlet obstruction in an infant, but the onset is usually in the second or third week of life. A pharyngoesophageal (Zenker) diverticulum above the upper esophageal sphincter is usually a disease of adults.

2  About 2% of individuals have a Meckel diverticulum, an embryologic remnant of the omphalomesenteric duct, but only a small subset of these individuals have ectopic gastric mucosa within it, which causes intestinal ulceration. The symptoms may mimic acute appendicitis, but appendicitis should not last for 1 month or cause significant blood loss. Angiodysplasia may be difficult to detect, and it is almost always seen in patients older than 70 years, but can cause significant blood loss. Celiac disease can occur in young individuals, but it does not produce significant hemorrhage. Diverticulosis can be associated with hemorrhage, but the diverticula are almost always in the colon of older persons. Giardiasis produces a self-limited, watery diarrhea without hemorrhage.

3  The infant’s condition occurred several weeks after birth because of hypertrophy of pyloric smooth muscle. pyloric stenosis has features of multifactorial inheritance with a “threshold of liability,” above which the disease is manifested when more genetic risks are present, such as family history and twin gestation. The incidence in males is 1 in 200 and in females is 1 in 1000, reflecting the fact that more risks must be present in females for the disease to occur. Annular pancreas is a rare anomaly that can also cause obstruction of the duodenum, and has variable age of onset, but a palpable mass would not be expected. Tracheoesophageal fistula, diaphragmatic hernia, and duodenal atresia are serious conditions that are manifested at birth and are often associated with multiple anomalies. pyloric stenosis is an isolated condition that typically occurs without other anomalies.
4 B In Hirschsprung disease, seen in 1 in 5000 live births, the aganglionic segment (either a short or long segment) of the bowel wall produces a functional obstruction with proximal distention. Most familial cases and some sporadic cases have RET gene mutations affecting neural crest cell migration. Atresias are congenitally narrowed segments of bowel (usually the small intestine) that occur with other anomalies. Patients with trisomy 21 may have intestinal (usually duodenal) atresias. Complete absence of the colonic lumen lies. Pertussis is whooping cough, typically a childhood disease affecting the upper airways. Candidiasis tends to produce surface plaques with minimal erosion in immunocompromised persons. Diphtheria is most often a childhood disease of upper airways, and there can be toxin-mediated systemic disease, including myocarditis, but there is no chronic infection. Herpetic ulcers are sharply demarcated, and infection is most often found in immunocompromised persons.

5 B Caustic alkaline solutions tend to damage the esophagus, and may not even get as far as the stomach. If the esophagus is perforated, a severe mediastinitis may occur. The inflammation can resolve with scarring and stenosis, and that tends to affect swallowing of solids more than liquids, typical for mechanical obstruction. A pharyngeal Zenker diverticulum occurs at a point of weakness in the hypopharynx, most often between the inferior constrictor muscle and cricopharyngeus muscle; it is a pulsion diverticulum from motility problems. Gastric lymphomas may be related to Helicobacter pylori infection (MALTomas) and to immune dysregulation. Duodenal ulcerations are predominantly related to H. pylori infection. Megacolon results from marked colonic inflammation or motor disturbances, and swallowed substances are not likely to reach the colon unaltered.

6 A In achalasia, there is incomplete relaxation of the lower esophageal sphincter with lack of peristalsis. Most cases are “primary” or of unknown origin. They may be caused by degenerative changes in neural innervation; the myenteric ganglia are usually absent from the body of the esophagus. There is a long-term risk of development of squamous cell carcinoma. In Barrett esophagus, there is columnar epithelial metaplasia, but the myenteric plexuses remain intact. Reflux esophagitis may be associated with hiatal hernia, but myenteric ganglia remain intact. Plummer-Vinson syndrome is a rare condition caused by iron deficiency anemia; it is accompanied by an upper esophageal web. Systemic sclerosis (scleroderma) is marked by fibrosis with stricture.

7 E Chronic Chagas disease can lead to damage to not only myocardium but also tubular structures of the GI tract, especially the esophagus with secondary achalasia. The organisms are hard to find microscopically, but they elicit the inflammatory response that damages neurons to produce the motility problems. Pertussis is whooping cough, typically a childhood disease affecting the upper airways. Candidiasis tends to produce surface plaques with minimal erosion in immunocompromised persons.
CHAPTER 17 Gastrointestinal Tract

11 E Esophageal dysmotility is the E in CREST, a mnemonic that details the key findings with the limited form of systemic sclerosis (scleroderma): C = calcinosis; R = Raynaud phenomenon; E = esophageal dysmotility; S = sclerodactyly; T = telangiectasias. Although scleroderma is an autoimmune disorder that often includes formation of anticientromere antibodies, little inflammation is seen by the time the patient seeks clinical attention. There is increased collagen deposition in gastrointestinal submucosa and muscularis. Fibrosis may affect any part of the gastrointestinal tract, but the esophagus is the site most often involved. For a diagnosis of Barrett esophagus, columnar metaplasia must be seen histologically, and there is often a history of gastroesophageal reflux disease. Hiatal hernia is frequently diagnosed in individuals with reflux esophagitis and can lead to inflammation, ulceration, and bleeding, but formation of a stricture is uncommon. An upper esophageal web associated with iron deficiency anemia might produce difficulty in swallowing, but this condition is rare. Portal hypertension gives rise to esophageal varices, not fibrosis.

PBD9 750–753, BP9 560–561, PBD8 223–225, BP8 150

12 D Her ongoing inflammatory process results from reflux of acid gastric contents into the lower esophagus. Gastroesophageal reflux disease (GERD) is a common problem that stems from a variety of causes, including sliding hiatal hernia, decreased tone of the lower esophageal sphincter, and delayed gastric emptying. Patients may have a history of heartburn after eating. Barrett esophagus is a complication of long-standing GERD and is characterized by columnar metaplasia of the squamous epithelium that normally lines the esophagus. There may be inflammation and mucosal ulceration overlying varices, but this condition usually does not have heartburn as the major feature. Esophageal varices from portal hypertension can lead to marked hematemesis. A rare complication of iron deficiency is the appearance of an upper esophageal web (Plummer-Vinson syndrome). Progressive fibrosis with stenosis is found in scleroderma.

PBD9 755–756, BP9 560–561, PBD8 769–770, BP8 588

13 C Variceal bleeding is a common complication of hepatic cirrhosis, which can be an outcome of chronic hepatitis B infection. Portal hypertension leads to dilated submucosal esophageal veins that can erode and bleed profusely. Barrett esophagus is a columnar metaplasia that results from gastroesophageal reflux disease (GERD). Bleeding is not a key feature of this disease. Esophageal candidiasis may be seen in immunocompromised patients, but it most often produces raised mucosal plaques and is rarely invasive. GERD may produce acute and chronic inflammation and, rarely, massive hemorrhage. Esophageal carcinomas may bleed, but not enough to cause massive hematemesis. A Zenker diverticulum is located in the upper esophagus and results from cricopharyngeal motor dysfunction; it presents a risk for aspiration, but not for hematemesis.

PBD9 756–757, BP9 559, PBD8 771–772, BP8 587–588

14 B The biopsy specimen shows residual ulcerated squamous epithelium along with columnar metaplasia and focal dysplasia, typical of Barrett esophagus. Patients with a focus of Barrett esophagus have a higher risk of developing adenocarcinoma than the general population, particularly when high-grade dysplasia is present. Achalasia refers to the failure of the lower esophageal sphincter to relax, which gives rise to dilation of the proximal portion of the esophagus. An episodic diverticulum in the lower esophagus is not associated with Barrett mucosa, but arises from increased intraluminal pressure against lower esophageal sphincter obstruction. Mallory-Weiss syndrome is associated with vertical lacerations in the esophagus that may occur with severe vomiting and retching. Squamous cell carcinomas occur in the midesophagus, but they do not arise in association with Barrett esophagus. Instead, they are linked to smoking and alcohol consumption.

PBD9 757–758, BP9 561–562, PBD8 770–771, BP8 588–589

15 A Adenocarcinomas of the esophagus are typically located in the lower esophagus, where Barrett esophagus develops at the site of long-standing gastroesophageal reflux disease. Barrett esophagus is associated with an increased risk of developing adenocarcinoma, particularly when high-grade dysplasia is present. Columnar metaplasia may progress to dysplasia, then adenocarcinoma. Carcinoid tumors occur in different parts of the gut, including the appendix, ileum, rectum, stomach, and colon. Leiomyosarcoma of the esophagus is rare and is unrelated to a history of heartburn. Malignant lymphomas of the gastrointestinal tract do not commonly occur in the esophagus and are not related to reflux esophagitis. Squamous cell carcinomas of the esophagus are most often associated with a history of chronic alcoholism and smoking.

PBD9 758–759, BP9 562, PBD8 772–773, BP8 589–591

16 E This large, ulcerated lesion with heaped-up margins is a malignant tumor of the esophageal mucosa. There are two main histologic types of esophageal carcinomas—squamous cell carcinoma and adenocarcinoma—with distinct risk factors. Smoking and alcoholism are the primary risk factors for esophageal squamous cell carcinoma in the Western world. Adenocarcinoma is most likely to arise in the lower third of the esophagus and to be associated with Barrett esophagus. Chronic inflammation may lead to stricture and not to a localized mass. Dilated veins occur in esophageal varices; they do not produce an ulcerated mass. A dense, collagenous scar of the mid esophagus is uncommon, but it may occur after injury from ingestion of a caustic liquid. Intranuclear inclusions suggest infection with herpes simplex virus or cytomegalovirus, both of which are more likely to produce ulceration without a mass; both occur in immunocompromised patients.

PBD9 758–759, BP9 562–563, PBD8 773–774, BP8 589–591

17 C The Turkmen population around the Caspian Sea has the highest rate of esophageal cancer on earth, and most of these are squamous cell carcinomas arising in the mid-esophagus. Consuming hot tea, contamination with silicates in consumed food, micronutrient deficiencies, and family history have been implicated, as well as human papillomavirus infection. There are no specific gene mutations known to be associated with esophageal carcinoma in this population.
contrast, tobacco use and alcohol consumption are linked to esophageal cancers in Europe and North America. The main autoimmune disease affecting the esophagus, systemic sclerosis (scleroderma), is not a major risk for cancer. Infectious agents such as Candida and herpes simplex virus do not carry a risk for cancer; the role for human papillomavirus in this process is not well established. Reflux esophagitis is a risk for adenocarcinomas arising in the lower third of the esophagus.

**PBD9 759–760 BP9 563 PBD8 772–774 BP8 589–590**

18 A These findings are consistent with an acute gastritis. If significant inflammation is not present, then the term gastropathy is used. Heavy consumption of ethanol is probably the most common cause, but aspirin and nonsteroidal anti-inflammatory drugs (NSAIDs), smoking, and chemotherapy agents can produce the same findings. NSAIDs can be cofactors in peptic ulcer disease. Chlorpromazine (used to treat nausea) does not have the same association. Cimetidine and omeprazole are used to treat peptic ulcer disease by reducing gastric acid production, increasing the serum gastrin. Cimetidine is an H2 receptor blocker, and omeprazole is a proton pump inhibitor. Clindamycin is a broad-spectrum antibiotic that may alter flora in the lower gastrointestinal tract.

**PBD9 760–762 BP9 564–565 PBD8 774–775 BP8 593**

19 A Prolonged use of nonsteroidal anti-inflammatory drugs (NSAIDs) is an important cause of acute gastritis. NSAIDs inhibit cyclooxygenase-dependent synthesis of prostaglandins E2 and I2, which stimulate nearly all defense mechanisms. Excessive alcohol consumption and smoking also are possible causes. Acute gastritis tends to be diffuse and, when severe, can lead to significant mucosal hemorrhage that is difficult to control. Epithelial dysplasia may occur at the site of chronic gastritis. It is a forerunner of gastric cancer. Infection with Helicobacter pylori is not associated with acute gastritis. Hyperplastic polypos of the stomach do not result from acute gastritis, but may arise in association with chronic gastritis. Acute gastritis does not increase the risk of gastric adenocarcinoma.

**PBD9 760–762 BP9 564–565 PBD8 774–775 BP8 593**

20 E So-called stress ulcers, also known as Curling ulcers, can occur in patients with burn injuries. The ulcers are often small (<1 cm) and shallow, never penetrating the muscularis propria, but they can bleed profusely. Similar lesions can occur after traumatic or surgical injury to the central nervous system (Cushing ulcers). Duodenal ulcers are typically peptic ulcers in individuals with Helicobacter pylori infection. Esophageal varices can cause massive hematemesis, but they occur in patients with portal hypertension, caused most commonly by cirrhosis. Metaplastic columnar epithelium at the lower end of the esophagus is present in Barrett esophagus, resulting from chronic gastroesophageal reflux disease. Ileal ulcerations and colonic ulcerations are often due to inflammatory bowel disease that can be from infections such as shigellosis, or they may be idiopathic, as in Crohn disease.

**PBD9 762 BP9 564–565 PBD8 775–776 BP8 596**

21 A The high MCV is indicative of a megaloblastic anemia, most likely pernicious anemia, resulting from autoimmune atrophic gastritis. Delayed maturation of the myeloid cells leads to hypersegmentation of polymorphonuclear leukocytes. Loss of gastric parietal cells from autoimmune injury causes a deficiency of both intrinsic factor and acid. In the absence of this factor, vitamin B12 cannot be absorbed in the distal ileum. Among the various anti-parietal cell antibodies are those directed against the acid-producing proton pump enzyme H+,K+-ATPase. Antigliadin antibodies are found with celiac disease that does not affect the gastric mucosa. *H. pylori* causes chronic gastritis and peptic ulcer disease, but does not injure parietal cells. In pernicious anemia, no antibodies are directed against intrinsic factor receptor on ileal mucosal cells. Infection with *Tropheryma whipplei* causes Whipple disease, which may involve any organ, but most often affects intestines, central nervous system, and joints; malabsorption is common.

**PBD9 764–765 BP9 567 PBD8 778–779 BP8 438–439,592**

22 B Helicobacter pylori organisms shown in the figure reside in the mucus layer above the gastric mucus and are associated with various gastric disorders, ranging from chronic gastritis with erythema and thickened rugal folds, as in this case, to peptic ulcers and to adenocarcinoma. *H. pylori* organisms elaborate several toxic substances that injure the epithelium. The *H. pylori* gene from a pathogenicity island encodes cytotoxin-associated antigen (CagA) and is present in many patients with chronic gastritis and peptic ulcers; it increases the risk for gastric cancer. Cysteine proteinases produced by *Entamoeba histolytica* aid in tissue invasion. Heat-stable enterotoxin is produced by strains of *Escherichia coli* that cause traveler’s diarrhea. Shiga toxin is elaborated by *Shigella flexneri* organisms, which cause a form of bacillary dysentery. Verocytotoxin produced by some *E. coli* strains is associated with hemolytic uremic syndrome mediated by endothelial injury.

**PBD9 763–764 BP9 566–567 PBD8 776–778 BP8 592–594**

23 D Although they are not found in the duodenum, *Helicobacter pylori* organisms alter the microenvironment of the stomach, causing the stomach and duodenum to be susceptible to peptic ulcer disease. Virtually all duodenal peptic ulcers are associated with *H. pylori* infection. Ulceration can extend through the muscularis and result in perforation, as in this case. The other organisms listed are not related to peptic ulcer formation, but to infectious diarrheal illnesses. *Salmonella typhi* may produce typhoid fever with more systemic symptoms; the marked ulceration of Peyer patches may lead to perforation.

**PBD9 763–764 BP9 566–567 PBD8 776–778 BP8 494–497**

24 C The clinical symptoms in this case suggest peptic ulcer disease. In most cases, peptic ulcers are associated with *Helicobacter pylori* infection. These bacteria secrete urease, which can be detected by oral administration of urea 14C. After drinking the labeled urea solution, the patient blows into a tube. If *H. pylori* urease is present in the stomach, the
urea is hydrolyzed, and labeled carbon dioxide is detected in the breath sample. In the biopsy urease test, antral biopsy specimens are placed in a gel containing urea and an indicator, and if *H. pylori* is present, the color changes within minutes. If not properly treated, peptic ulcers can produce many complications, including massive bleeding that can be fatal. Carcinoid tumors can occur in the stomach, but they are rare and are not related to peptic ulcer disease, which this patient has. He does not have fat malabsorption because fat absorption does not occur in the stomach. Peptic ulcers rarely progress to gastric carcinoma. The stomach has numerous arterial supplies and therefore is unlikely to be affected by focal thrombosis. Vitamin B₁₂ deficiency can occur with autoimmune atrophic gastritis because intrinsic factor, which is required for vitamin B₁₂ absorption, is produced in gastric parietal cells.

25 A Gastric inflammatory/hyperplastic polyps may arise in the setting of *Helicobacter pylori* infection. They are the most common type of gastric polyp. They may be precursors to gastric adenocarcinomas, particular lesions larger than 1.5 cm and with high-grade dysplasia. The other listed options are not appropriate for an infectious etiology.

26 A There is an association of fundic gland polyps with use of proton pump inhibitors and also with familial adenomatous polyposis (FAP); increased gastrin may drive glandular hyperplasia. Gastric adenomas are most common in the antrum, have intestinal metaplasia with dysplasia, and are precursors to adenocarcinoma; they may occur with FAP. Hyperplastic polyps are associated with chronic gastritis, often from *H. pylori* infection. One form of hypertrophic gastropathy is Ménétrier disease, which results from massive secretion of transforming growth factor alpha (TGF-α) with diffuse enlargement of gastric rugae and protein-losing enteropathy.

27 A The most likely cause of a large mass lesion in the stomach is a gastric carcinoma, and this lesion is an adenocarcinoma, likely the intestinal type found in the antral region. Adenocarcinoma is related to *Helicobacter pylori* infection, with β-catenin mutation. The incidence of this type of gastric cancer has been decreasing for decades in places where food processing methods have improved. Malignant lymphomas and leiomyosarcomas are less common and tend to form bulky masses in the fundus. Neuroendocrine carcinomas are rare. Squamous cell carcinomas typically appear in the esophagus.

28 E The endoscopic and radiologic findings describe the linitis plastica (“leather bottle”) appearance of diffuse gastric carcinoma. Histologically, these carcinomas are composed of the gastric type of mucus cells that infiltrate the stomach wall diffusely. The individual tumor cells have a signet ring appearance because the cytoplasmic mucin pushes the nucleus to one side. In chronic atrophic gastritis, the rugal folds are lost, but there is no significant scarring or shrinkage. Primary gastric lymphomas are less common than adenocarcinomas; a lymphoma may be large but would not involve the stomach in a diffuse pattern. Gastrointestinal stromal tumors tend to be bulky masses. Granulomas are rare at this site.
32 D This gastrointestinal stromal tumor (GIST) is derived from the interstitial cell of Cajal, and hence of mesenchymal origin. Those arising in the stomach may be less aggressive than those arising in the intestine, but most are KIT positive and amenable to tyrosine kinase inhibitor therapy. Some GISTs may have mutations in platelet-derived growth factor receptor A (PDGFRα). Antibiotic therapy to obliterate Helicobacter pylori infection may be useful in treating MALTomas. Azathioprine and corticosteroids may be employed in treating inflammatory bowel disease, but the greatest risk for malignancy with inflammatory bowel disease is adenocarcinoma, particularly of the colon. Cyclophosphamide is a chemotherapy agent not employed in treating GISTs. Radiotherapy is not generally effective against mesenchymal malignancies.

33 B The patient has acute bowel obstruction, and the findings at surgery show bowel infarction. The most common causes in developed nations are adhesions, hernias, and metastases. Adhesions are most often the result of prior surgery, as in this case, and produce “internal” hernias, where a loop of bowel becomes trapped (incarcerated), and the blood supply is compromised. Loops of bowel that become trapped in direct or indirect inguinal hernias also can infarct. When metastases are the cause, the primary site is generally known, and the cancer stage is high. Primary adenocarcinomas of the small bowel are uncommon. Crohn disease can be focal and manifest with bowel obstruction, but it is uncommon in patients of this age. Intussusception can be focal, but it is uncommon. Abdominal tuberculosis may cause circumferential stricture of the bowel, and should be considered in regions where the prevalence of tuberculosis is high. Volvulus may involve the cecal or sigmoid regions of the colon (because of their mobility). When volvulus involves the small intestine, torsion around the mesentery generally occurs, and there is extensive (not segmental) small bowel infarction.

34 C The infant has signs and symptoms of acute bowel obstruction. Intussusception occurs when one small segment of small bowel becomes telescoped into the immediately distal segment. This disorder can have sudden onset in infants and may occur in the absence of any anatomic abnormality. Duodenal atresia (which typically occurs with other anomalies, particularly trisomy 21) and Hirschspring disease (from an aganglionic colonic segment) usually manifest soon after birth. Almost all cases of Meckel diverticulum are asymptomatic, although in some cases functional gastric mucosa is present and can lead to ulceration with bleeding. Pyloric stenosis is seen much earlier in life and is characterized by projectile vomiting.

35 D The patient’s history of myocardial infarction suggests that he had severe coronary atherosclerosis, and the elevated Hgb A1c suggests diabetes mellitus. Systemic atheromatous disease most likely involves the mesenteric vessels as well, giving rise to thrombotic occlusion of the blood vessels that perfuse the bowel. The symptoms and signs suggest infarction of the gut. Acute appendicitis rarely leads to such a catastrophic illness, unless there is perforation. (The absence of free air in the radiograph argues against perforation of any viscus.) Acute pancreatitis can be a serious abdominal emergency, but the normal levels of amylase and lipase tend to exclude it. Acute cholecystitis can produce severe abdominal pain, but bloody diarrhea and absence of bowel sounds (paralytic ileus) are unlikely. Pseudomembranous colitis develops in patients receiving broad-spectrum antibiotic therapy.

36 A Hypotension with hypoperfusion from heart failure is a common cause of ischemic bowel in hospitalized patients. The ischemic changes begin in scattered areas of the mucosa and become confluent and transmural over time. This can give rise to paralytic ileus and bleeding from the affected regions of the bowel mucosa. A mesenteric vasculitis is uncommon, but could lead to bowel infarction. Shigellosis is an infectious diarrhea that causes diffuse colonic mucosal erosion with hemorrhage. Ulcerative colitis usually produces marked mucosal inflammation with necrosis, usually in a continuous distribution from the rectum upward. Volvulus is a form of mechanical obstruction caused by twisting of the small intestine on its mesentery or twisting of the cecum or sigmoid colon, resulting in compromised blood supply that can lead to infarction of the twisted segment.

37 A Angiodysplasia refers to tortuous dilations of mucosal and submucosal vessels, seen most often in the cecum in patients older than 50 years. These lesions, although uncommon, account for 20% of cases involving significant lower intestinal bleeding. Bleeding usually is not massive, but can occur intermittently over months to years. This lesion is difficult to diagnose and is often found radiographically. The focus (or foci) of abnormal vessels can be excised. Collagenous colitis is a rare cause of a watery diarrhea that is typically not bloody. Colonic diverticulosis can be associated with hemorrhage, but the outpouchings usually are seen on colonoscopy. Hemorrhoids at the anorectal junction may account for bright red rectal bleeding, but they can be seen or palpated on rectal examination. Mesenteric venous thrombosis is rare and may result in bowel infarction with severe abdominal pain.

38 C Fat malabsorption can occur from impaired intraluminal digestion. Smelly, bulky stools containing increased amounts of fat (steatorrhea) are characteristic. Pancreatic or biliary tract diseases are important causes of fat malabsorption. Amebiasis can produce a range of findings from a watery diarrhea to dysentery with mucus and blood in the stool. Giardiasis produces mainly a watery diarrhea. Malabsorption with steatorrhea is unlikely to be associated with bleeding. Cholera results in a massive watery diarrhea.
**CHAPTER 17  Gastrointestinal Tract**

produces children worldwide. Recurrent infection sets up a cycle of infection, and damage the enterocytes expressing MIC-A. A trial of gluten-free diet is the most logical therapeutic option. Patients usually become symptom-free, and normal histologic features of the mucosa are restored. Some patients develop dermatitis herpetiformis, and a few enteropathy-associated T-cell lymphomas. Anticentromere antibody is more specific for limited scleroderma (formerly CREST syndrome) with esophageal dysmotility. The anti-DNA topoisomerase I antibody is most specific for diffuse scleroderma, in which gastrointestinal tract involvement by submucosal fibrosis may be more extensive, and malabsorption may be present. Antimitochondrial antibody is more specific for primary biliary cirrhosis. Antinuclear antibody is present in a wide variety of autoimmune diseases, but it is not characteristic of celiac sprue.

**40 E** The malabsorption responded to dietary treatment. She probably has celiac disease (gluten sensitivity) with histologic features including flattening of the mucosa, diffuse and severe atrophy of the villi, crypt hyperplasia, and chronic inflammation of the lamina propria. There is an increase in intraepithelial lymphocytes, both CD4+ and CD8+. Affected persons are HLA-DQ2 or HLA-DQ8 positive. Crypt abscesses are nonspecific and can be seen in inflammatory bowel disease. Lymphatic obstruction occurs in Whipple disease, and in addition, foamy macrophages accumulate in the lamina propria. The macrophages contain PAS-positive granules that under electron microscopy show an actinomycete called *Tropheryma whippelli*. Noncaseating granulomas are found in the intestinal wall in Crohn disease.

**41 B** Environmental enteropathy affects millions of children worldwide. Recurrent infection sets up a cycle of mucosal injury and inflammatory response that produces an appearance similar to celiac disease. There is no single infectious agent implicated, but likely there are many pathogens that cumulatively contribute to mucosal damage. Abetalipoproteinemia is a rare condition from mutations in microsomal triglyceride transfer protein that impairs enterocyte transport of lipoproteins. Cystic fibrosis results from *CFTR* gene mutations affecting chloride ion channels, but the resultant diarrhea is primarily from loss of pancreatic function. The most common disaccharidase deficiency is lactase deficiency, with milk intolerance. *NOD2* gene mutations may contribute to Crohn disease.

**42 E** Disaccharidase (lactase) deficiency, either congenital or acquired, is symptomatic when the lactose in milk products is not broken down into glucose and galactose by terminal digestion, resulting in an osmotic diarrhea and gas production from gut flora. Affected individuals do not always make the connection between diet and symptoms, or they do not consume enough milk products to become symptomatic. An autoimmune gastritis is most likely to result in vitamin B12 malabsorption. Celiac disease also is diet related and results from sensitivity to gluten in some grains. Cholelithiasis can cause biliary tract obstruction with malabsorption of fats and pain in the right upper quadrant of the abdomen. Cystic fibrosis affects the pancreas and mainly produces fat malabsorption.

**43 E** The clinical features suggest food poisoning caused by the ingestion of a preformed enterotoxin. *Staphylococcus aureus* grows in food (milk products and fatty foods are favorites) and elaborates an enterotoxin that, when ingested, produces diarrhea within hours. *Bacillus cereus* is better known for growing on reheated fried rice; it produces an exotoxin that causes acute nausea, vomiting, and abdominal cramping. *Clostridium difficile* can produce a pseudomembranous colitis in patients treated with broad-spectrum antibiotics. Some strains of *Escherichia coli* can produce various diarrheal illnesses, but without a preformed toxin. *Salmonella enterica* is most often found in poultry products, but the diarrheal illnesses develop within 2 days. *Vibrio parahaemolyticus* is found in shellfish.

**44 D** Raw or poorly cooked shellfish can be the source of *Vibrio parahaemolyticus*, which tends to produce a milder diarrhea than *Vibrio cholerae*. *Vibrio* organisms produce a toxin that increases adenylate cyclase, leading to chloride ion secretion and osmotic diarrhea. *Cryptosporidium* as a cause of watery diarrhea is most often found in immunocompromised individuals. *Entamoeba histolytica* produces colonic mucosal invasion along with exudation and ulceration; stools contain blood and mucus. *Staphylococcus aureus* can produce food poisoning through elaboration of an enterotoxin that causes an explosive vomiting and diarrhea within 2 hours after ingestion. *Yersinia enterocolitica* is invasive and can produce extraintestinal infection.

**45 B** The source of *Campylobacter jejuni* can include contaminated water, unpasteurized milk, and poorly cooked poultry. The bloody diarrhea (dysestera) and leukocytes suggest intestinal mucosal invasion by a bacterial organism. An ascending paralysis (Guillain-Barré syndrome) may complicate some *Campylobacter* infections because of cross-reactivity between human ganglioside G	extsubscript{M1} and bacterial lipopolysaccharide. *Bacillus cereus* food poisoning tends to produce abrupt onset of vomiting. *Clostridium perfringens* tends to produce gas gangrene. Giardiasis produces a watery diarrhea without dysentery or extraintestinal complications. Rotavirus infections are most common in children.
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**46 E** Infection by one of several *Salmonella enterica* (not Typhi) causes a self-limited diarrhea. This is a form of food poisoning, typically from contaminated poultry products. *Bacillus cereus* growing in foods such as reheated fried rice produces an exotoxin, which, on ingestion, can produce acute onset of nausea, vomiting, and abdominal pain. Amebiasis from *Entamoeba histolytica* can be an invasive, exudative infection. The stools contain blood and mucus. Various diseases result from contamination with different strains of *Escherichia coli*, based on the characteristics of the organisms, and whether they invade or produce an enterotoxin. Poultry products are usually not contaminated with *E. coli*. Rotavirus is most likely to produce symptomatic watery diarrhea in children, unrelated to diet. *Staphylococcus aureus* causes an acute onset of abdominal pain, bloating, and diarrhea, not by directly infecting the gastrointestinal tract, but by producing an exotoxin while growing on food that is subsequently ingested. *Yersinia enterocolitica* is most often found in contaminated milk or pork products and may disseminate to produce lymphadenitis and further extraintestinal infection.

**47 D** Typhoid fever begins as an intestinal infection, but it becomes a systemic illness. A chronic carrier state can occur in some infected individuals, with colonization of the gallbladder. *Campylobacter jejuni* may produce dysentery, but generally not systemic disease. *Clostridium perfringens* can cause gas gangrene. *Mycobacterium bovis* is now rare because of pasteurization of milk products; it was best known as a cause of bowel obstruction from circumferential ulceration and scarring of the small bowel. Shigellosis can produce dysentery, but the infection is generally limited to the colon. Infection with *Yersinia enterocolitica* can produce extraintestinal infection with lymphadenitis, but generally not dysentery.

**48 D** The opened colon shows pseudomembranes that are patches of fibrinopurulent debris attached to the mucosa. Pseudomembranous enterocolitis is a complication of broad-spectrum antibiotic therapy, which alters gut flora to allow overgrowth of *Clostridium difficile* or other organisms that are capable of inflicting mucosal injury. *Clostridium septicum* infection can lead to myonecrosis that is most often associated with malignancy or immunosuppression. An inflammatory bowel disease does not typically produce a pronounced exudate and is not associated with *C. difficile*. This gross pattern also can appear from ischemic injury that is vascular or mechanical, but this patient’s history and the time course support an iatrogenic cause. An ischemic colitis resulting from mesenteric artery thrombosis could appear similar, but it is not associated with *C. difficile*. A dilated, thinned, toxic megacolon is an uncommon complication of ulcerative colitis.

**49 C** Norovirus outbreaks result from contamination of food or water, most often in venues where multiple persons congregate. Was it the resort pool? Noroviruses, as well as the *Giardia* parasite, are resistant to chlorination. Was it the buffet? Salads, shellfish, and meats are often implicated. The voluminous diarrhea suggests small intestinal involvement. The lack of leukocytes makes bacterial infection less likely. Cytomegalovirus infections are more likely in immunocompromised persons. Botulism leads to paralysis from a neurotoxin. Staphylococcal food poisoning tends to be abrupt in onset and of short duration. Strongyloidiasis tends to persist for months to years. Cholera produces life-threatening fluid loss.

**50 F** Rotavirus is the most common cause of viral gastroenteritis in children. It is a self-limited disease that affects mostly infants and young children, who can lose a significant amount of fluid relative to their size and can quickly become dehydrated. The death rate is less than 1%. *Campylobacter jejuni* is more often seen in children and adults as a food-borne cause of fever, abdominal pain, and diarrhea. Cryptosporidiosis most often causes a watery diarrhea in immunocompromised adults. Enterohemorrhagic strains of *Escherichia coli* can produce hemolytic uremic syndrome in young children. Listeriosis can be a congenital infection that is present along with meningitis and sepsis at birth; in infants, children, and adults, it is a food-borne or water-borne infection that tends to occur in epidemics. Norwalk virus is a common cause of diarrheal illness in adults. Shigellosis produces dysentery with bloody diarrhea.

**51 A** Hookworm infections may be caused by *Ancylostoma duodenale* (Old World) or *Necator americanus* (New World) or both, because the geographic distributions may overlap, particularly in Africa and Asia. The sharp hooks of the worms penetrate the small intestinal mucosa and produce bleeding. The worms live for months to years. Infection occurs through the skin, and larval development occurs in the lungs until migration to the trachea and swallowing conduct the worms to the duodenum. Organisms listed in the remaining choices are unlikely to produce significant gastrointestinal hemorrhage.

**52 D** Diarrhea with mucus and blood in the stools may be caused by several enteroinvasive microorganisms, including *Shigella dysenteriae* and *Entamoeba histolytica*. In most cases, the diarrhea is self-limited. The initial episode of diarrhea could have been caused by one of several organisms; however, the occurrence of a liver abscess after an episode of diarrhea most likely results from infection with *E. histolytica*. Colonic mucosal and submucosal invasion by *E. histolytica* allows the organisms to access the submucosal veins draining to the portal system and the liver. *Clostridium difficile* causes pseudomembranous colitis after antibiotic therapy. Dissemination of *Cryptosporidium* and *Strongyloides* organisms may occur in immunocompromised patients. *Giardia* produces a self-limited, watery diarrhea.

**53 B** The ileum shows chronic inflammation with lymphoid aggregates. The inflammation is transmural, affecting
the mucosa, submucosa, and muscularis as shown in the figure. During surgery, inflammation is also observed in the serosa. A deep fissure extending into the muscularis is present. These histologic features are highly suggestive of Crohn disease. Extension of fissures into the overlying skin can produce enterocutaneous fistulas, although enterenteric fistulas between loops of bowel are more common. Although the risk of adenocarcinoma is increased in Crohn disease, this complication is less common than sequelae of inflammation. Intussusception may occur when there is a congenital or acquired obstruction in the bowel. Hepatic abscess can follow amebic colitis, or other infections. Mesenteric artery thrombosis, typically a complication of atherosclerosis, is unlikely in a 27-year-old man.

54 D Irritable bowel syndrome (IBS) can be difficult to diagnose because of protean manifestations found in many other conditions. No pathologic or physiologic abnormalities can be identified reliably with IBS. Patients may benefit from behavioral therapies. Placebos may work as well as pharmacotherapies. The lack of an increased stool fat in this case indicates that chronic pancreatitis and cystic fibrosis are unlikely. Diverticular disease is more likely to occur in older adults. Inflammatory bowel disease has both pathologic and radiographic findings. Viral gastroenteritis is unlikely to persist for 5 years.

55 A The figure shows a diffuse, predominantly mononuclear infiltrate in the lamina propria along with a crypt abscess. Ulcerative colitis can lead to relapsing and remitting episodes of low volume diarrhea containing blood and mucus and diffuse inflammation and ulceration of the rectal and colonic mucosa. One of the most dreaded complications of ulcerative colitis is the development of colonic adenocarcinoma. There is a twentyfold to thirtyfold higher risk in patients who have had ulcerative colitis for 10 or more years compared with control populations. Diverticulitis can produce abdominal pain and blood in the stool, but there is no association with ulcerative colitis. Fat malabsorption usually does not occur in ulcerative colitis because the ileum often is not involved. Perirectal fistula formation is more typical of Crohn disease, in which there is transmural inflammation. Ulcerative colitis is associated with several extraintestinal manifestations, including sclerosing cholangitis, but it has no relationship to primary biliary cirrhosis. Pseudomembranous colitis is caused by *Clostridium difficile* infections associated with broad-spectrum antibiotic treatment.

56 C The segment of the colon shows the diffuse and severe ulceration characteristic of ulcerative colitis. The inflammation shown is so severe that areas of mucosal ulceration have produced pseudopolyps or islands of residual mucosa. Ulcerative colitis is a systemic disease; in some patients, it is associated with migratory polyarthritis, ankylosing spondylitis, and primary sclerosing cholangitis. The pathogenesis of ulcerative colitis is unclear, but is most likely mediated by a T-cell response to an unknown antigen (but not a gut infection), leading to an imbalance between T-cell activation and regulation. The Th17 immune response has CD4+ T cells present in the lesions that secrete damaging substances. Autoantibodies against tropomyosin are present, but do not play a pathogenic role in ulcerative colitis. Mutations in the *NOD2* gene are linked to Crohn disease, not ulcerative colitis. Inheritance of a germline APC mutation causes familial adenomatous polyposis with a very high risk for colon cancer. Ulcerative colitis also increases the risk for colon cancer, but not secondary to APC gene mutation.

57 B The findings in Crohn disease and ulcerative colitis overlap, and in at least 10% of cases it may be impossible to differentiate between them—a so-called indeterminate colitis. Regardless of the exact diagnosis, there is a considerable increase in risk for development of carcinoma 8 to 10 years after disease onset. Surveillance screening can detect dysplasia as a precursor to carcinoma, but would you just remove the colon with the ongoing problem and avoid missing the possible cancer? If you remove the colon, but it turns out to be Crohn disease, it may recur. Extraintestinal manifestations may occur regardless. Doctors like to go for the win with a “cure,” but patients want to avoid potential loss of life or function. The doctor gets to walk away from any outcome, but the patient does not. There are often no easy answers in medicine.

58 B The clinical and histologic features are consistent with Crohn disease, one of the idiopathic inflammatory bowel diseases. Crohn disease is marked by segmental bowel involvement and transmural inflammation that leads to strictures, adhesions, and fistula. Ulcerative colitis has mucosal involvement extending variable distances from the rectum. In contrast to Crohn disease, the mucosal involvement is diffuse and does not show “skip areas.” Fissures and fistulas are not frequently seen in ulcerative colitis. The findings in Crohn disease and ulcerative colitis overlap, and in at least 10% of cases it may be impossible to differentiate between them—a so-called indeterminate colitis. Generally, crypt abscesses are more typical of ulcerative colitis, and granulomas are more typical of Crohn disease, but these features are not present in most biopsy specimens from patients with either condition. A story is told of an attending physician at an academic medical center who was known to berate students and residents on rounds for not definitively diagnosing ulcerative colitis and Crohn disease. When he retired, incomplete records for patients with idiopathic inflammatory bowel disease were found in his office; the records represented about one sixth of the total cases of inflammatory bowel disease that he had seen. Amebiasis and shigellosis are infectious processes that can cause mucosal ulceration, but they do not produce granulomas or fissures. Sarcoidosis can involve many organs and give rise to noncaseating granulomas; however, involvement of the intestines is uncommon, and sarcoidosis does not give rise to ulcerative disease.
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59 B These are findings of idiopathic inflammatory bowel disease most likely to be Crohn disease. The ileal involvement accounts for the vitamin K and vitamin B12 deficiencies as well as disrupted enterohepatic circulation of bile salts predisposing to gallstone formation. The inflammatory response in Crohn disease may result from inappropriate innate immune responses to gut flora, as discussed in the text. Angiodysplasia leads to bleeding from abnormal submucosal vessels, most often in the cecum of older adults. Diverticular disease is common in older persons but results from mechanical, not immune, mechanisms. Severe peripheral atherosclerosis may cause ischemic bowel disease, but this is usually an acute process.

PBD9 796–800 BP9 589–590 PBD8 808–811 BP8 611–612

60 C Colonic diverticulosis may be accompanied by intermittent minimal bleeding and, rarely, by severe bleeding. One or more diverticula may become inflamed (diverticulitis) or, less commonly, may perforate to produce an abscess, peritonitis, or both. Diverticular disease is not a premalignant condition. The diverticula project outward, and even with inflammation, luminal obstruction is unlikely. Malabsorption is not a feature of diverticular disease. Toxic megacolon is an uncommon complication of inflammatory bowel disease.

PBD9 803–804 BP9 586–587 PBD8 814–815 BP8 603–604

61 C Juvenile polyps are the most common form of hamartomatous polyp. Singly they are likely to be sporadic, and the only complication is rectal prolapse; but when multiple polyps are present, they may be the result of an autosomal dominant syndrome with risk for development of adenocarcinoma. The remaining choices include polyposis syndromes unlikely to appear at this age.

PBD9 805–806 BP9 592 PBD8 816–817 BP8 617–618

62 A The figure shows a solitary pedunculated adenoma of the colon with no evidence of malignancy. High magnification shows a small focus of dysplastic, non-mucin-secreting epithelial cells lining a colonic crypt, giving rise to “tubular” architecture. Such a small (<2 cm) solitary, tubular adenoma is unlikely to harbor a focus of malignancy; a search for metastases is unwarranted. Such colonic adenomas are more likely to occur in older persons; hence the recommendation for colonoscopy screening after age 50. Removing such an adenoma does not leave the chance for further growth of the lesion with possible development of adenocarcinoma. Individuals who inherit a mutant APC gene usually develop hundreds of polyps at a young age; this patient does not need genetic testing for a somatic mutation in the APC gene. Patients with hereditary nonpolyposis colorectal cancer, with multiple polyps present, have an increased risk of endometrial cancer and develop colon cancer at a young age. It is unlikely that the blood loss from a small polyp would be sufficient to cause iron deficiency, although the small amount of blood emanating from colonic polyps and cancers is the rationale to test for fecal occult blood. Peutz-Jehgers syndrome is associated with development of hamartomatous polyps in the small intestine.

PBD9 807–809 BP9 593–595 PBD8 819–820 BP8 618–619

63 A The figure shows a large villous adenoma. There is a high probability that large villous adenomas will progress to invasive adenocarcinoma. When they occur in the descending colon, these lesions are annular and cause obstruction. In the colon, non-Hodgkin lymphomas are far less common than adenocarcinomas, and they do not manifest as mucosal sessile masses. Carcinoid tumors are typically small and yellowish, and most grow slowly. Leiomysarcomas are rare; they produce large bulky masses, but they do not arise on the mucosa. Mucinous cystadenomas are cystic and are more likely to arise in an ovary or in the pancreas. The original lesion in this patient was a villous adenoma.

PBD9 808–809 BP9 594–595 PBD8 819–820 BP8 617–618

64 A This young patient’s colon shows hundreds of polyps. This is most likely a case of familial adenomatous polyposis (FAP) syndrome, which results from inheritance of one mutant copy of the APC tumor-suppressor gene (a few FAP cases are associated with DNA mismatch repair genes). Every somatic cell of this patient most likely has one defective copy of the APC gene. Polyps are formed when the second copy of the APC gene is lost in many colon epithelial cells. Without treatment, colon cancers arise in 100% of these patients because of accumulation of additional mutations in one or more polyps, typically before 30 years of age. Patients with a gene for hereditary nonpolyposis colorectal carcinoma, such as MLH1 and MSH2, also have an inherited susceptibility to develop colon cancer, but in contrast to patients with FAP, they do not develop numerous polyps. Sporadic colon cancers may have CpG island hypermethylation along with KRAS mutations, whereas others have p53 mutations, but the somatic cells of patients with these cancers do not show abnormalities of these genes. NOD2 mutations are linked with Crohn disease.

PBD9 809–810 BP9 595–596 PBD8 820–822 BP8 619

65 D Of the conditions listed, the one most likely to lead to adenocarcinoma in a patient of this age is hereditary nonpolyposis colorectal cancer, or Lynch syndrome. Crohn disease is unlikely because the patient has not had prior serious illness, and Crohn disease of long duration is unlikely to remain asymptomatic. Although adenocarcinoma may complicate Crohn disease, it does not occur as frequently as in ulcerative colitis. This explains why colectomy is often performed for ulcerative colitis, but bowel resections are avoided, if possible, in Crohn disease. The other conditions listed are not premalignant.

PBD9 810 BP9 596 PBD8 821–822 BP8 621–622

66 B The lesion is an adenocarcinoma, showing irregular glands infiltrating the muscle layer. Such a lesion in a 30-year-old man strongly indicates a hereditary predisposition. One hereditary form of cancer is called hereditary nonpolyposis colorectal cancer (HNPPC) and results from defective DNA mismatch repair genes. As a result, mutations accumulate in microsatellite repeats (microsatellite instability) that lead to loss of transforming growth factor beta (TGF-β) receptor-mediated control of colonic epithelial cell proliferation and...
loss of proapoptotic BAX protein enhancing survival of these transformed cells. He could have taken NSAIDs that inhibit COX-2 expressed in most colonic adenomas and carcinomas. In contrast to familial adenomatous polyposis syndrome, HNPCC does not lead to the development of hundreds of polyps in the colon. Detection of ERBB2 (HER2/NEU) expression is important in breast cancers. Germline inheritance of the tumor suppressor gene RB1 predisposes to retinoblastoma and osteosarcoma, not colon carcinoma. E-cadherin is required for intercellular adhesion; its levels are reduced, not increased, in carcinoma cells. Translocation of the retinoic acid receptor alpha gene is characteristic of acute promyelocytic leukemia.

67 D The figure shows an encircling mass that is typical of adenocarcinoma of the descending colon. Such cancers likely to obstruct, but they can also bleed a small amount over months to years, causing iron deficiency anemia. The APC gene, a negative regulator of β-catenin in the WNT signaling pathway, is associated with familial adenomatous polyposis syndrome and most sporadic colon cancers, as in this case. This pathway also is known as the adenoma-carcinoma sequence because the carcinomas develop through an identifiable series of molecular and morphologic steps. Loss of the PTEN tumor suppressor gene is seen in endometrial carcinomas not associated with colon carcinoma and with some hamartomatous polyps of the colon. Evidence for an additional cancer, such as an endometrial cancer, would suggest an inherited mutation in one of the DNA mismatch repair genes, such as MSH2 and MLH1. Homozygous loss of these genes can give rise to right-sided colon cancer and endometrial cancer. Such a mutation is typically associated with microsatellite instability. Infection with some strains of human papillomavirus leads to RB1 protein inactivation and development of cervical carcinoma. Mutation with activation of KIT tyrosine kinase activity occurs in gastrointestinal stromal tumors, which respond well to treatment with imatinib mesylate, a tyrosine kinase inhibitor also used to treat chronic myelogenous leukemia.

68 B Hemorrhoids are a common problem that can stem from any condition that increases venous pressure and causes dilation of internal or external hemorrhoidal veins above and below the anorectal junction. Angiodysplasia of the colon leads to intermittent hemorrhage, typically in older individuals. Ischemic colitis is rare in young individuals because the most common underlying cause (severe atherosclerotic disease involving mesenteric vessels) occurs in older patients. Intussusception and volvulus are rare causes of mechanical bowel obstruction; they occur suddenly in adults and are surgical emergencies.

69 F Acute appendicitis can be accompanied by an elevated WBC count with neutrophilia and left shift. This is helpful but not decisive, and the decision to operate must be based on clinical judgment. Amebiasis is most likely associated with a history of diarrhea, often with blood in the stool. Hyperamylasemia occurs in acute pancreatitis. Diarrhea with fluid loss and dehydration can lead to hypotension. The serum carcinoembryonic antigen level may be increased in patients with colonic cancers; however, this test is not specific for colon cancer. The alkaline phosphatase level may be increased in biliary tract obstruction.

70 A Pseudomyxoma peritonei (PP) is described here. It may arise from low-grade mucinous adenocarcinoma of the appendix, which may be so differentiated that it resembles an appendiceal mucocele. However, PP tends to recur. In women, PP needs to be distinguished from mucinous tumors of the ovary. Mucinous tumors may also arise in the pancreas, but are less likely to disseminate through the peritoneal cavity. Malignancies arising in the small intestine are rare. Mucin-producing malignancies of the stomach are most likely to have a signet ring cell pattern and diffusely infiltrate the gastric wall.

71 F Spontaneous bacterial peritonitis is an uncommon complication found in about 10% of adult patients with cirrhosis of the liver and ascites. The ascitic fluid provides an excellent culture medium for bacteria, which can invade the bowel wall or spread hematogenously to the serosa. Spontaneous bacterial peritonitis also can occur in children, particularly children with nephrotic syndrome and ascites. The most common organism cultured is Escherichia coli. Appendicitis has a peak incidence in younger patients; the pain is often (but not always) more localized in the right lower quadrant, and ascites is usually absent. Appendicitis is not related to alcoholism. Collagenous colitis is uncommon; it most often leads to watery diarrhea in middle-aged women. Diverticulitis with rupture could produce peritonitis, but there is typically no ascites, and diverticulitis is not related to alcoholism. Ischemic colitis may produce infarction with rupture and peritonitis, but ascites is usually lacking, and individuals with chronic alcoholism are unlikely to have marked atherosclerosis. Pseudomembranous colitis is a complication of antibiotic therapy.
1. A previously healthy, 38-year-old woman has become increasingly obtunded in the past 4 days. On physical examination, she has scleral icterus, abdominal fluid wave, and asterixis. She is afebrile, and her blood pressure is 110/55 mm Hg. Laboratory findings show a prothrombin time of 38 seconds (INR 3.1), serum ALT of 1854 U/L, AST of 1621 U/L, albumin of 1.8 g/dL, and total protein of 4.8 g/dL. Serum or blood levels of which of the following will most likely be abnormal in this patient?
   - A. Alkaline phosphatase
   - B. Ammonia
   - C. Amylase
   - D. Anti-HCV
   - E. Antinuclear antibody (ANA)

2. A pathologic study of hepatic cirrhosis is performed. There is collapse of reticulin with bridging fibrosis from deposition of collagen in the space of Disse to form fibrous septae. Which of the following cell types is activated under the influence of cytokines to give rise to collagen-producing cells?
   - A. Bile duct cell
   - B. Endothelial cell
   - C. Hepatocyte
   - D. Macrophage
   - E. Stellate cell

3. A 54-year-old woman has a long history of chronic hepatitis B infection and has had increasing malaise for the past year. She was hospitalized 1 year ago because of upper gastrointestinal hemorrhage. Physical examination now shows a firm nodular liver. Laboratory findings show a serum albumin level of 2.5 g/dL and prothrombin time of 28 seconds. Which of the following additional physical examination findings is most likely to be present in this woman?
   - A. Caput medusae
   - B. Diminished deep tendon reflexes
   - C. Distended jugular veins
   - D. Papilledema
   - E. Splinter hemorrhage

4. A 57-year-old woman has had increasing abdominal enlargement for 6 months. During the past 2 days, she developed a high fever. On physical examination, her temperature is 38.5°C. The abdomen is enlarged and diffusely tender, and there is a fluid wave. Paracentesis yields 500 mL of cloudy yellowish fluid. The cell count is 532/μL with 98% neutrophils and 2% mononuclear cells. A blood culture is positive for *Escherichia coli*. The representative gross appearance of her liver is shown in the figure. Which of the following underlying diseases most commonly accounts for these findings?
   - A. α1-Antitrypsin deficiency
   - B. Chronic alcohol abuse
   - C. Hepatitis E viral infection
   - D. Hereditary hemochromatosis
   - E. Primary sclerosing cholangitis
5 A study of patients with ascites includes measurements of serum and ascitic fluid protein levels. The serum-ascites albumin gradient (SAAG) is calculated. Some patients are found to have a high gradient, along with splenomegaly. They are found to have serum albumin less than 2.5 g/dL. Which of the following conditions is most likely to produce a SAAG greater than 1.1?

A Budd-Chiari syndrome
B Cirrhosis
C Nephrotic syndrome
D Pancreatitis
E Peritonitis

6 A 65-year-old man with a history of alcohol abuse has had hematemesis for the past day. Physical examination reveals mild jaundice, spider angiomas, and gynecomastia. He has mild pedal edema, normal jugular venous pulsation (JVP), and a massively distended abdomen. Paracentesis is performed and the fluid obtained shows accumulation of protein-poor fluid that is free of inflammatory cells. Which of the following factors is most likely to be responsible for the collection of abdominal fluid in this man?

A Congestive heart failure
B Hepatopulmonary syndrome
C Hyperbilirubinemia
D Portosystemic shunts
E Splanchnic arterial vasodilation

7 A 59-year-old man has had increasing dyspnea on exertion for the past year. His dyspnea is worse in the upright position and diminishes when he is recumbent. On physical examination he has clubbing of the fingers. Exercise induces a decrease in his Po2 that improves when he stops and lies down. Which of the following liver abnormalities is he most likely to have?

A Biliary obstruction
B Chronic inflammation
C Cirrhosis
D Metastases
E Steatosis

8 A 50-year-old man has a history of chronic alcoholism, but he stopped drinking alcohol 10 years ago. He has been taking no medications. On physical examination, he is afebrile. The abdomen is not enlarged, and there is no tenderness. The liver span is normal. Serologic test results for hepatitis A, B, and C are negative. The hematocrit is 35%. Which of the following morphologic features is most likely to be present in his liver?

A Concentric “onion-skin” bile duct fibrosis
B Hepatic venous thrombosis
C Interface hepatitis
D Massive hepatocellular necrosis
E Periportal PAS-positive globule deposition
F Portal fibrosis with regenerative nodules

9 A 58-year-old woman has experienced gradually increasing malaise, icterus, and loss of appetite for the past 6 months. On physical examination, she has generalized jaundice and scleral icterus. She has mild right upper quadrant tenderness; the liver span is normal. Laboratory studies show total serum bilirubin of 7.8 mg/dL, AST of 190 U/L, ALT of 220 U/L, and alkaline phosphatase of 26 U/L. A liver biopsy is done, and microscopic examination shows the findings in the figure, along with portal bridging fibrosis. These findings are most typical of which of the following conditions?

A Choledocholithiasis
B Congestive heart failure
C HAV infection
D HCV infection
E Hemochromatosis
F Sclerosing cholangitis

10 A 27-year-old man develops malaise, fatigue, and loss of appetite three weeks after a meal at the Trucker’s Cafe. He notes passing dark urine. On physical examination, he has mild scleral icterus and right upper quadrant tenderness. Laboratory studies show serum AST of 62 U/L and ALT of 58 U/L. The total bilirubin concentration is 3.9 mg/dL, and the direct bilirubin concentration is 2.8 mg/dL. His symptoms abate over the next 3 weeks. On returning to the cafe, he finds that the city’s health department has closed it. Which of the following serologic test results is most likely to be positive in this patient?

A Anti-HAV
B Anti-HBc
C Anti-HBs
D Anti-HCV
E Anti-HDV
In a clinical study, patients with infectious hepatitis, including viral hepatitis A, B, C, D, E, F, and G, are followed for 5 years. During that time, prothrombin time, serum AST, ALT, alkaline phosphatase, total bilirubin, and ammonia are periodically measured. A liver biopsy is performed each year, and the microscopic findings are recorded. Which of the following is most likely the best predictor of whether a patient with viral hepatitis will develop chronic liver disease that progresses to cirrhosis?

A. Degree to which hepatic transaminase enzymes are elevated
B. Length of time that hepatic enzymes remain elevated
C. Presence of chronic inflammatory cells in the portal tract
D. Presence of inflammatory cells in the hepatic sinusoids
E. Specific form of hepatitis virus responsible for the infection

A 30-year-old man had a 2-week episode of malaise, fever, and jaundice 7 years ago. On physical examination, there were needle tracks in the left antecubital fossa. Serologic test results were positive for HBsAg, HBV DNA, and IgG anti-HBc. Two years later, he was seen in the emergency department because of hematemesis and ascites. Serologic test results were similar to those reported earlier. Further serologic test results were negative for IgM anti-HAV, anti-HBc, and anti-HCV. Repeat testing 6 months later yields the same results. Which of the following is the most appropriate statement regarding the pathophysiology of this patient’s condition?

A. Chronic carrier state with no therapy indicated
B. Clinically overt hepatitis will occur within 1 year
C. Erroneous test results that need to be repeated
D. Hepatitis B vaccination series is now required
E. Infection acquired through intravenous drug use

A 42-year-old man experiences malaise and increasing icterus for 2 weeks. Physical examination shows jaundice, but there are no other significant findings. Serologic test results are positive for IgM anti-HAV and negative for anti-HCV, HBsAg, and IgM anti-HBc. Which of the following outcomes is most likely to occur in this man?

A. Chronic active hepatitis
B. Complete recovery
C. Fulminant hepatitis
D. Hepatocellular carcinoma
E. Negative serologic test results

A 41-year-old woman who works as a tattoo artist has had increasing malaise and nausea for the past 2 weeks. On physical examination, she has icterus and mild right upper quadrant tenderness. Laboratory studies show serum AST of 79 U/L, ALT of 85 U/L, total bilirubin of 3.3 mg/dL, and direct bilirubin of 2.5 mg/dL. She continues to have malaise for the next year. A liver biopsy is done, and microscopic examination shows minimal hepatocyte necrosis, mild steatosis, and minimal portal bridging fibrosis. An infection with which of the following viruses is most likely to produce these findings?

A. HAV
B. HBV
C. HCV
D. HDV
E. HEV

A study is conducted of patients who are infected with hepatitis virus A, B, C, D, E, or G. The patients are categorized according to the type of virus and are followed over the next 10 years. They receive periodic serologic testing to determine whether they are producing antibodies to the virus with which they were infected. Analysis of the data shows that a cohort of these patients developed antibodies, but subsequently did not clear the virus until treated with pegylated interferon and ribavirin. Which of the following forms of viral hepatitis was most likely to infect this subset of patients?

A. HAV
B. HBV
C. HCV
D. HDV
E. HEV

F. HGV
A 52-year-old woman has experienced worsening malaise during the past year. On physical examination, she has mild scleral icterus. There is no ascites or splenomegaly. Serologic tests are positive for IgG anti-HCV and HCV RNA and negative for anti-HAV, HBsAg, ANA, and antimitochondrial antibody. The serum AST level is 88 U/L, and ALT is 94 U/L. Her condition remains stable for months. Which of the following morphologic findings is most likely to be present in this patient’s liver?

- A Concentric “onion-skin” bile duct fibrosis
- B Copper deposition within hepatocytes
- C Granulomatous bile duct destruction
- D Hepatic venous thrombosis
- E Interface hepatitis
- F Massive hepatocellular necrosis
- G Microvesicular steatosis

A 27-year-old man with a history of intravenous drug use is known to have been infected with hepatitis B virus for the past 6 years and has not been ill. He is seen in the emergency department because he has had nausea, vomiting, and passage of dark-colored urine for the past week. Physical examination shows scleral icterus and mild jaundice. Neurologic examination shows a confused, somnolent man oriented only to person. He exhibits asterixis. Laboratory studies show total protein, 5 g/dL; albumin, 2.7 g/dL; AST, 2342 U/L; ALT, 2150 U/L; alkaline phosphatase, 233 U/L; total bilirubin, 8.3 mg/dL; and direct bilirubin, 4.5 mg/dL. Superinfection with which of the following viruses has most likely occurred in this man?

- A HAV
- B HCV
- C HDV
- D HEV
- E HGV

A 36-year-old, G3, P2, woman living in New Delhi, India, has worsening nausea and malaise for a week. On physical examination her sclerae are icteric. Her liver span is increased and the liver edge is tender. She is at 16 weeks’ gestation. Laboratory studies show her serum AST is 495 U/L and ALT is 538 U/L. She recovers and hepatic function returns to normal, but spontaneous abortion occurs at 18 weeks. Epidemiologic studies show a point source of contaminated water for infection. With which of the following viruses was she most likely infected?

- A Cytomegalovirus (CMV)
- B Epstein-Barr virus (EBV)
- C Hepatitis E virus (HEV)
- D Herpes simplex virus (HSV)
- E Yellow fever virus

A 29-year-old man has developed malaise and nausea 2 months following intercourse with a new sexual contact. He notes scleral icterus 10 days later. He now has two more sexual contacts who subsequently become ill. Serologic testing shows that he is HBsAg positive, HAV-IgM negative, and anti-HCV negative. His AST is 77 IU/L and ALT 95 IU/L. A month later his anti-HBs is positive. Which of the following is the most likely course of his illness?

- A Asymptomatic illness
- B Chronic hepatitis
- C Fulminant hepatic failure
- D Hepatitis with recovery
- E Macronodular cirrhosis

A 53-year-old woman from southern China has had fever, right upper quadrant pain, and jaundice for the past 6 months. On examination she has an increased liver span. An abdominal CT scan shows a 5-cm right hepatic tumor with a branching, infiltrative appearance. A liver biopsy is performed and on microscopic examination shows irregular invasive glands in a desmoplastic stroma. This patient is most likely to have chronic infection with which of the following?

- A Clonorchis sinensis
- B Echinococcus granulosus
- C Plasmodium vivax
- D Mycobacterium tuberculosis
- E Salmonella typhi

A 31-year-old woman has experienced increasing malaise for the past 4 months. Physical examination yields no remarkable findings. Laboratory studies show total serum protein of 6.4 g/dL, albumin of 3.6 g/dL, total bilirubin of 1.4 mg/dL, AST of 67 U/L, ALT of 91 U/L, and alkaline phosphatase of 99 U/L. Results of serologic testing for HAV, HBV, and HCV are negative. Test results for ANA, anti-liver kidney microsome-1, and anti-smooth muscle antibody are positive. A liver biopsy is done; microscopically, there are minimal portal mononuclear cell infiltrates with minimal interface hepatitis and mild portal fibrosis. What is the most likely diagnosis?

- A $\alpha_1$-Antitrypsin deficiency
- B Autoimmune hepatitis
- C Chronic alcoholism
- D HDV infection
- E Isoniazid ingestion
- F Primary biliary cirrhosis

A 66-year-old woman with a history of chronic alcohol abuse has had headaches and nausea for the past 4 days. She has become increasingly obtunded. On physical examination she has right upper quadrant tenderness, tachycardia, tachypnea, and hypotension. Laboratory studies show serum AST of 475 U/L, ALT of 509 U/L, alkaline phosphatase of 23 U/L, total bilirubin of 0.9 mg/dL, albumin of 3.8 g/dL, and total protein of 6.1 g/dL. She is treated with N-acetylcysteine. Which of the following drugs has she most likely ingested in excess?

- A Acetaminophen
- B Aspirin
- C Ibuprofen
- D Meperidine
- E Oxycodone
26 A 63-year-old man with a 30-year history of alcohol abuse notes hematemesis for the past day. On examination, he has ascites, mild jaundice, and an enlarged spleen. He also has gynecomastia, spider telangiectasias of the skin, and testicular atrophy. Rectal examination indicates prominent hemorrhoids and a normal-sized prostate. Emergent upper endoscopy shows dilated, bleeding submucosal vessels in the esophagus. Laboratory studies show total protein, 5.9 g/dL; albumin, 3.2 g/dL; AST, 137 U/L; ALT, 108 U/L; total bilirubin, 5.4 mg/dL; prothrombin time, 20 seconds; ammonia, 76 μmol/L; and hematocrit, 21%. Which of the following pathologic findings in his liver is most likely to explain the hematemesis?

A Cholangitis  
B Cholestasis  
C Cirrhosis  
D Hepatitis  
E Steatosis

27 A 52-year-old man has had increasing malaise and swelling of the lower legs for the past 4 months. On physical examination, he is afebrile and normotensive. There is pitting edema to the knees. The abdomen is slightly distended with a fluid wave, but there is no tenderness. The liver span is increased. Laboratory studies show total serum protein of 5 g/dL, albumin of 2.2 g/dL, AST of 65 U/L, ALT of 65 U/L, alkaline phosphatase of 93 U/L, and total bilirubin of 1.8 mg/dL. A liver biopsy is performed and the microscopic appearance with trichrome stain is shown in the figure. Ingestion of which of the following is most likely to have caused this illness?

A Acetaminophen  
B Allopurinol  
C Aspirin  
D Chlorpromazine  
E Ethanol  
F Isoniazid

28 A 48-year-old man has noticed increasing abdominal girth and a yellowish color to his skin over the past 5 months. On physical examination, he has scleral icterus and generalized jaundice. His abdomen is distended, and a fluid wave is present. Laboratory studies include total serum bilirubin of 5.2 mg/dL, direct bilirubin of 4.2 mg/dL, AST of 380 U/L, ALT of 158 U/L, alkaline phosphatase of 95 U/L, total protein of 6.4 g/dL, and albumin of 2.2 /dL. The prothrombin time is 18 seconds, and the partial thromboplastin time is 30 seconds. The blood ammonia level is 105 mmol/L. What is the most likely cause of these findings?

A Acute HAV infection  
B Alcoholic liver disease  
C Cholelithiasis  
D Metastatic adenocarcinoma  
E Primary biliary cirrhosis

29 A 38-year-old man feels acutely ill with nausea, upper abdominal pain, and jaundice following a heavy bout of drinking over the weekend. On physical examination, there is right upper quadrant tenderness. Laboratory studies include a total WBC count of 16,120/mm³ with 82% segmented neutrophils, 8% bands, 8% lymphocytes, and 2% monocytes. The total serum bilirubin is 4.9 mg/dL, AST is 542 U/L, ALT is 393 U/L, and alkaline phosphatase is 118 U/L. A liver biopsy is obtained and the microscopic appearance is shown in the figure. What is the nature of the accumulations seen in the biopsy?

A α1-Antitrypsin globules  
B Lysed red cells  
C Bile pigment  
D Hemosiderin deposition  
E Mallory-Denk bodies  
F Viral inclusions (HBsAg)

30 A longitudinal study is conducted of non-alcoholics with type 2 diabetes mellitus, dyslipidemia, and BMI >30. There is an increasing prevalence of liver disease in these persons over time. Which of the following microscopic pathologic findings is most characteristic for the livers of these persons?

A Apoptosis  
B Cholestasis  
C Cirrhosis  
D Hemosiderosis  
E Inflammation  
F Steatosis
A study of persons with increased risk for ischemic heart disease reveals that some of them also have liver disease. Risk factors include lack of exercise and increased consumption of fast-food products containing high fructose corn syrup. Laboratory studies show that their blood glucose averages 117 mg/dL. Serum AST and ALT are elevated. Abdominal CT imaging shows hepatomegaly with diffusely decreased attenuation but no focal lesions. Some of them go on to develop hepatocellular adenoma. Which of the following underlying disorders do these persons most likely have?

A Type 1 diabetes mellitus  
B Familial hypercholesterolemia  
C Hepatitis C virus infection  
D Hereditary hemochromatosis  
E Metabolic syndrome

A 4-year-old girl has abrupt onset of vomiting, which remains protracted for 24 hours. On arrival at the emergency department, the child is lethargic and febrile to 37.7°C. The parents state that she had a mild upper respiratory tract illness 3 days ago, but was improving, and the only medication she received was acetylsalicylic acid (aspirin). On physical examination, there is poor skin turgor, the lungs are clear, the abdomen is nontender, and the heart rate is regular. Laboratory findings show Na⁺, 150 mmol/L; K⁺, 4.5 mmol/L; Cl⁻, 93 mmol/L; CO₂, 30 mmol/L; glucose, 60 mg/dL; creatinine, 1.1 mg/dL; amylase, 25 U/L; AST, 386 U/L; ALT, 409 U/L; alkaline phosphatase, 120 U/L; total bilirubin, 1.1 mg/dL; ammonia, 80 μmol/L; and prothrombin time, 26 seconds with INR of 2. The child becomes comatose. What pathologic finding is most likely present in the liver of this girl?

A Common bile duct atresia  
B Hepatic vein thrombosis  
C Hepatoblastoma  
D Intrahepatic duct litihsis  
E Microvesicular steatosis  
F Multinucleated giant cell hepatitis

A 23-year-old man has noted a yellow color to his sclerae for the past 2 weeks. On physical examination he has generalized jaundice. He has the physique of a bodybuilder. Laboratory studies show serum total bilirubin, 5.6 mg/dL; ALT, 117 U/L; AST, 103 U/L; alkaline phosphatase, 148 U/L; albumin, 5.5 g/dL; and total protein, 7.9 g/dL. Which of the following substances is he most likely to be using?

A Acetaminophen  
B Anabolic steroid  
C Chlorpromazine  
D Ethyl alcohol  
E Isoniazid

A 25-year-old medical student from Mozambique notices a slight yellowish color on the day of the final examination. He has the physique of a bodybuilder. Laboratory studies show serum total bilirubin, 5.6 mg/dL; ALT, 117 U/L; AST, 103 U/L; alkaline phosphatase, 148 U/L; albumin, 5.5 g/dL; and total protein, 7.9 g/dL. Which of the following substances is he most likely to be using?

A Acetaminophen  
B Anabolic steroid  
C Chlorpromazine  
D Ethyl alcohol  
E Isoniazid

The prothrombin time is 13 seconds, and partial thromboplastin time is 25 seconds. Serologic test results for HAV, HBV, and HCV are negative. A chest radiograph shows no abnormal findings. What is the most likely diagnosis?

A Antiphospholipid syndrome  
B Ascending cholangitis  
C Chronic alcoholism  
D Metastatic adenocarcinoma  
E Sclerosing cholangitis

A 20-year-old primigravida gives birth at term following an uncomplicated pregnancy to a boy infant of normal weight and length. On examination no abnormalities are noted. Within the first week, the infant becomes mildly icteric. The infant receives phototherapy, and there is no more icterus after the second week of life. Which of the following mechanisms most likely led to this infant’s icterus?

A Atresia of the common bile duct  
B Congenital infection with cytomegalovirus  
C Inherited deficiency of a canalicular transporter  
D Low hepatic glucuronyl transferase activity  
E Maternally derived antibody-mediated hemolysis

A 35-year-old woman has noticed an increasing yellowish hue to her skin for the past week. On physical examination, there is no abdominal pain or tenderness, and the liver span is normal. Laboratory findings include hemoglobin, 11.7 g/dL; hematocrit, 35.2%; MCV, 98 μm³; platelet count, 207,600/mm³; WBC count, 6360/mm³; total protein, 5.5 g/dL; albumin, 3.5 g/dL; total bilirubin, 8.7 mg/dL; direct bilirubin, 0.6 mg/dL; AST, 39 U/L; ALT, 24 U/L; and alkaline phosphatase, 35 U/L. What is the most likely diagnosis?

A Cholelithiasis  
B Hemolytic anemia  
C Hepatitis A viral infection  
D Micronodular cirrhosis  
E Oral contraceptive use

A 68-year-old woman has become increasingly tired, with a 3-kg weight loss without dieting over the past 6 months. On physical examination a stool sample is positive for occult blood. Laboratory studies show total serum protein, 6.1 g/dL; albumin, 3.9 g/dL; total bilirubin, 1.1 g/dL; AST, 38 U/L; ALT, 44 U/L; alkaline phosphatase, 294 U/L; glucose, 70 mg/dL; and creatinine, 0.9 mg/dL. CBC shows hemoglobin, 8.9 g/dL; hematocrit, 26.7%; MCV, 75 μm³; platelet count, 198,400/mm³; and WBC count, 5520/mm³.
38 | A 42-year-old woman from Lisbon, Portugal, has had fever, chills, and bouts of colicky right upper quadrant pain for the past week. On physical examination, her skin is icteric, and there is scleral icterus. Laboratory studies show a total serum bilirubin concentration of 7.1 mg/dL and direct bilirubin concentration of 6.7 mg/dL. An abdominal ultrasound scan shows cholelithiasis; dilation of the common bile duct; and two cystic lesions, 0.8 cm and 1.5 cm, in the right lobe of the liver. Which of the following infectious agents is most likely to produce these findings?

A  Clonorchis sinensis  
B  Cryptosporidium parvum  
C  Cytomegalovirus  
D  Entamoeba histolytica  
E  Escherichia coli

39 | A 17-year-old woman, G2, P1, gives birth to a term infant after an uncomplicated pregnancy. The infant does well for 3 weeks, but then begins to have abdominal enlargement, light-colored stools, and dark urine. On physical examination, the infant is icteric. There is hepatomegaly, but no splenomegaly or lymphadenopathy. Laboratory studies show serum AST of 101 U/L, ALT of 123 U/L, alkaline phosphatase 20 U/L, glucose of 81 mg/dL, and creatinine of 0.4 mg/dL. A liver biopsy is done, and microscopically shows lobular disarray with focal hepatocyte necrosis, giant cell transformation, cholestasis, portal mononuclear cell infiltrates, Kupffer cell hyperplasia, and microscopically shows marked proliferation of bile ducts. Which of the following morphologic features is most likely to be present in his liver?

A  Erythroblastosis fetalis  
B  Extrahepatic biliary atresia  
C  Galactosemia  
D  Idiopathic neonatal hepatitis  
E  Primary biliary cirrhosis  
F  Von Gierke disease

40 | A 19-year-old mother notices that her 3-week-old neonate has increasing jaundice. The pregnancy was uncomplicated and ended in a normal term birth. On physical examination, the infant now exhibits generalized jaundice, hepatomegaly, and acholic stool. Laboratory studies show total serum bilirubin of 10.1 mg/dL, AST of 123 U/L, ALT of 140 U/L, glucose of 77 mg/dL, and creatinine of 0.4 mg/dL. The alkaline phosphatase is normal. A liver biopsy is done and microscopically shows marked proliferation of bile ducts, portal tract edema and fibrosis, and extensive intrahepatic and canalicular bile stasis. The infant develops progressively worsening jaundice and dies of liver failure at 9 months of age. What is the most likely diagnosis?

A  α1-Antitrypsin deficiency  
B  Cholelithiasis  
C  Congenital toxoplasmosis  
D  Extrahepatic biliary atresia  
E  Hepatoblastoma

41 | A 45-year-old woman has had increasing pruritus and icterus for 7 months. On physical examination, she has generalized jaundice. Laboratory studies show total serum protein, 6.3 g/dL; albumin, 2.7 g/dL; total bilirubin, 5.7 mg/dL; direct bilirubin, 4.6 mg/dL; AST, 77 U/L; ALT, 81 U/L; and alkaline phosphatase, 221 U/L. A liver biopsy specimen shows destruction of portal tracts, loss of bile ducts, and lymphocytic infiltrates. Which of the following additional laboratory findings is most likely to be present in this woman?

A  Decreased α1-antitrypsin level  
B  Elevated sweat chloride level  
C  Increased serum ferritin level  
D  Positive anti-HCV  
E  Positive antimitochondrial antibody

42 | A 43-year-old man has experienced progressive fatigue, pruritus, and icterus for 4 months. A colectomy was performed 5 years ago for treatment of ulcerative colitis. On physical examination, he now has generalized jaundice. The abdomen is not distended; on palpation, there is no abdominal pain and there are no masses. Laboratory studies show a serum alkaline phosphatase level of 285 U/L and an elevated titer of anti-neutrophil cytoplasmic antibodies. Cholangiography shows widespread intrahepatic biliary tree obliteration and a beaded appearance in the remaining ducts. Which of the following morphologic features is most likely to be present in his liver?

A  Concentric “onion-skin” ductular fibrosis  
B  Copper deposition in hepatocytes  
C  Granulomatous bile duct destruction  
D  Interface hepatitis  
E  Periportal PAS-positive globules  
F  Portal bridging fibrosis

43 | A 44-year-old man has had increasing arthritis pain, swelling of the feet, and reduced exercise tolerance over the past 3 years. Laboratory studies include serum glucose of 201 mg/dL, creatinine of 1.1 mg/dL, and ferritin of 893 ng/mL. A chest radiograph shows bilateral pleural effusions, pulmonary edema, and cardiomegaly. He undergoes a liver biopsy; the microscopic appearance of a biopsy specimen stained with H&E (right panel) and Prussian blue (left panel) is shown in the figure. Based on these findings, which of the following is the most appropriate therapy for this patient?

A  Cholecystectomy  
B  Interferon-α  
C  Phlebotomy  
D  Prednisone  
E  Reduce alcohol intake
44 A 46-year-old man has had gradually increasing abdominal distention along with decreased libido for the past 7 months. Physical examination reveals excessive skin pigmentation in sun-exposed areas. He has an abdominal fluid wave and modest splenomegaly. Fasting serum laboratory findings include glucose, 200 mg/dL; creatinine, 0.8 g/dL; ferritin, 650 ng/mL; total protein, 6.3 g/dL; and albumin, 2.2 g/dL. His total bilirubin, AST, ALT, and alkaline phosphatase values are normal. Hemoglobin is 13.5 g/dL, hematocrit is 40.6%, MCV is 94 μm^3, platelet count is 200,000/mm^3, and WBC count is 6570/mm^3. His prothrombin time is 20 seconds, and partial thromboplastin time is 65 seconds. A mutation in a gene leading to which of the following molecular abnormalities most likely explains his disease?

A. β_2-Microglobulin gene, preventing the binding of β_2-microglobulin to HFE
B. β-Globin gene, with ineffective erythropoiesis and excessive absorption of iron
C. Divalent metal transporter 1 gene, causing increased binding to intestinal luminal iron
D. HFE gene, reducing hepcidin synthesis and increasing iron absorption
E. Transferrin gene, with sequestration of iron in the liver

45 A 23-year-old man has had worsening congestive heart failure along with arthritis resembling pseudogout for the past 6 years. On examination his skin has a slate-grey color, his heart rate is irregular, his liver span is increased, and the spleen is palpable. Both testes appear small and atrophic. Laboratory studies show an elevated hemoglobin A1c and increased serum ferritin. A mutation involving which of the following genes is most likely to be present in this woman?

A. MHC Class I
B. ATP7B
C. HAMP
D. HNF1-α
E. DMT1

46 A 19-year-old woman is bothered by a tremor at rest, which becomes progressively worse over the next 6 months. She exhibits paranoid ideation with auditory hallucinations and is diagnosed with an acute psychosis. On physical examination, she has scleral icterus. A slit lamp examination shows corneal Kayser-Fleischer rings. Laboratory findings include total serum protein, 5.9 g/dL; albumin, 3.1 g/dL; total bilirubin, 4.9 mg/dL; direct bilirubin, 3.1 mg/dL; AST, 128 U/L; ALT, 157 U/L; and alkaline phosphatase, 56 U/L. Which of the following additional serologic test findings is most likely to be reported in this patient?

A. Decreased α_1-antitrypsin level
B. Decreased ceruloplasmin level
C. Increased α-fetoprotein level
D. Increased ferritin level
E. Positive antimitochondrial antibody
F. Positive HbsAg

47 A 28-year-old man has had increasing shortness of breath for the past year. On physical examination, he is afebrile and normotensive. Breath sounds are decreased in all lung fields. His medical history indicates that he developed marked icterus as a neonate, but he has been healthy since then. There is a family history of liver disease. A liver biopsy is performed, and the figure shows the microscopic appearance stained with PAS. This patient is most likely at a very high risk for development of which of the following conditions?

A. Acute fulminant hepatitis
B. Diabetes mellitus
C. Pulmonary emphysema
D. Systemic lupus erythematosus
E. Ulcerative colitis

48 A 39-year-old woman has had increasing abdominal girth for a month, then pain for the past day. On physical examination there is hepatomegaly and caput medusae. Laboratory studies show Hgb, 20.5 g/dL; Hct, 61.7%; platelet count, 411,000/mm^3; AST, 333 U/L; and ALT, 358 U/L. What is ultrasonography of her abdomen most likely to show?

A. Choledocholithiasis
B. Cirrhosis
C. Hepatic vein thrombosis
D. Hepatocellular carcinoma
E. Macrophage steatosis
F. Subphrenic abscess
A 50-year-old man has increasing dyspnea with idiopathic pulmonary fibrosis, which was diagnosed 18 months ago. Physical examination shows elevated jugular venous pressure and pedal edema. Laboratory studies show serum AST of 221 U/L, ALT of 234 U/L, alkaline phosphatase of 48 U/L, lactate dehydrogenase of 710 U/L, total bilirubin of 1.2 mg/dL, albumin of 3.5 g/dL, and total protein of 5.4 g/dL. The figure shows the microscopic appearance of a liver biopsy specimen. Which of the following terms best describes these findings?

- Apoptosis
- Centrilobular congestion
- Cholestasis
- Hemosiderin deposition
- Macrovesicular steatosis
- Mallory-Denk bodies
- Portal fibrosis

A 26-year-old primigravida in the third trimester has had worsening peripheral edema for the past month. On examination, she is afebrile, but her blood pressure is 165/105 mm Hg. She has pitting edema to her thighs. Laboratory studies show hemoglobin, 11.7 g/dL; platelet count, 46,000/mm³; WBC count, 7210/mm³; albumin, 3.2 g/dL; total protein, 5.8 g/dL; total bilirubin, 1.2 mg/dL; AST, 208 U/L; ALT, 241 U/L; alkaline phosphatase, 32 U/L; and haptoglobin, 5 mg/dL. Urinalysis shows 4+ proteinuria, but no glucosuria. Her prothrombin time is 30 seconds. What is the best treatment for her condition?

- Corticosteroids
- Intravenous immunoglobulin
- Liver transplantation
- Plasmapheresis
- Pregnancy termination
- Portacaval shunt
- Ribavirin and interferon

A 36-year-old woman is in the sixth month of her first pregnancy, but she is unsure of her dates because she was taking oral contraceptives at the time she became pregnant. She experiences sudden onset of severe abdominal pain. On physical examination, she is afebrile and normotensive. There is right upper quadrant tenderness on palpation. An ultrasound scan of the abdomen shows a well-circumscribed, 7-cm subcapsular hepatic mass. Paracentesis yields bloody fluid. At laparotomy, the mass in the right hepatic lower lobe, which has ruptured through the liver capsule, is removed. The remaining liver parenchyma appears to be of uniform consistency, and the liver capsule is otherwise smooth. Which of the following is the most likely diagnosis?

- Cholangiocarcinoma
- Focal nodular hyperplasia
- Hepatic adenoma
- Hepatocellular carcinoma
- Macronodular cirrhosis
- Metastatic adenocarcinoma
A 79-year-old man has had increasing abdominal discomfort and malaise for the past year. On physical examination his liver span is increased. His stool is positive for occult blood. Laboratory studies show Hgb, 8 g/dL; Hct, 22%; MCV, 72 μm³; total protein, 5.6 g/dL; albumin, 3.7 g/dL; and alkaline phosphatase, 209 U/L. The representative appearance of his liver is shown in the figure. Which of the following underlying conditions is he most likely to have?

A Chronic alcohol abuse  
B Colonic adenocarcinoma  
C Gilbert syndrome  
D Hepatitis B virus infection  
E Metabolic syndrome

A clinical study of patients with cirrhosis includes MR imaging. Patients with an intrahepatic nodule larger than 2 cm that has arterial blood flow are found to be at increased risk for hepatocellular carcinoma. Which of the following pathologic findings is most likely to characterize these nodules?

A Adenoma  
B Dysplasia  
C Regeneration  
D Siderosis  
E Steatosis

A 38-year-old man from Shanghai, China, has experienced fatigue and a 10-kg weight loss over the past 3 months. Physical examination yields no remarkable findings. Laboratory test results are positive for HBsAg and negative for anti-HCV and anti-HAV. Abdominal CT scan shows a 10-cm solid mass in the left lobe of a nodular liver. A liver biopsy of the lesion is obtained and microscopically shows hepatocellular carcinoma. Which of the following is most likely responsible for the development of this lesion?

A Co-infection with Clonorchis sinensis  
B Development of hepatic adenoma that accumulates mutations  
C Insertion of viral DNA in the vicinity of the NMYC oncogene  
D Inherited mutation in the DNA mismatch repair genes  
E Ongoing infection with liver cell necrosis and regeneration

A 41-year-old woman experienced increasing malaise and a 10-kg weight loss in the past year. She becomes increasingly obtunded and lapses into a coma. Her serum α-fetoprotein is increased. The representative gross appearance of her liver is shown in the figure. Ingestion of which of the following substances is most likely to have played a role in the development of this condition?

A Acetaminophen  
B Aflatoxin  
C Aspirin  
D Ferrous sulfate  
E Nitrites  
F Raw oysters

A 36-year-old woman has become increasingly icteric for 1 month. She has had several bouts of colicky, mid-abdominal pain for 3 years. On physical examination, she has generalized jaundice with scleral icterus. Her BMI is 32. There is tenderness in the right upper quadrant, and the liver span is normal. A liver biopsy is obtained, and microscopic examination shows bile duct proliferation and intracanalicular bile stasis, but no inflammation or hepatocyte necrosis. The level of which of the following is most likely to be increased in the patient’s serum?

A Alkaline phosphatase  
B Ammonia  
C Antimitochondrial antibody  
D Hepatitis C antibody  
E Indirect bilirubin level

A 34-year-old woman from Kobe, Japan, has experienced intermittent upper abdominal pain for 3 weeks. Physical examination yields no remarkable findings. Laboratory findings show total serum protein of 7.3 g/dL, albumin of 5.2 g/dL, total bilirubin of 7.5 mg/dL, direct bilirubin of 6.8 mg/dL, AST of 35 U/L, ALT of 40 U/L, and alkaline phosphatase of 207 U/L. Representative microscopic findings in her liver include intracanalicular cholestasis in the centrilobular regions, swollen liver cells, and portal tract edema, but no necrosis and no fibrosis. Which of the following underlying conditions is she most likely to have?

A Cardiomyopathy  
B Choledochal cyst  
C Hemoglobinopathy  
D Hepatitis B viral infection  
E Ulcerative colitis  
F Venoocclusive disease
A 54-year-old First Nations woman has had colicky right upper quadrant pain for the past week. She has nausea, but no vomiting or diarrhea. On physical examination, she is afebrile. There is marked tenderness of the right upper quadrant. The liver span is normal. Her height is 160 cm (5 feet 3 inches), and her weight is 90 kg (body mass index 33). An abdominal ultrasound scan shows calculi within the lumen of the gallbladder, and the gallbladder wall appears thickened. Intrahepatic and extrahepatic bile ducts appear normal. The patient’s gallbladder is removed by laparoscopic cholecystectomy; it has the appearance shown in the figure. Which of the following mechanisms is most likely to play the greatest role in development of her disease?

A  Antibody-mediated RBC lysis  
B  *Ascaris lumbricoides* within bile ducts  
C  Biliary hypersecretion of cholesterol  
D  Decreased renal excretion of phosphate  
E  Hepatocyte infection by HBV  
F  Ingestion of foods rich in fat  
G  Involvement of the terminal ileum by Crohn disease

A 78-year-old man has had abdominal pain for a week. On physical examination he has a temperature of 37° C with pulse 106/min and blood pressure 85/50 mm Hg. There is left lower quadrant and right upper quadrant pain. Abdominal CT imaging shows sigmoid diverticula along with gallbladder dilation and biliary sludge. A blood culture grows polymicrobial flora. Which of the following is most likely to cause his gallbladder disease?

A  Hemolysis  
B  Carcinoma  
C  Drug reaction  
D  Lithiasis  
E  Sepsis

A 55-year-old man has developed abdominal pain and jaundice over 5 weeks. On physical examination, there is right upper quadrant pain, but no abdominal distention. Abdominal CT scan shows a markedly thickened gallbladder wall. A cholecystectomy is performed, and sectioning shows an enlarged gallbladder containing a fungating, 4 × 7 cm firm, lobulated, tan mass. Which of the following risk factors is most likely associated with this mass?

A  Alcohol abuse  
B  Cholelithiasis  
C  *Clonorchis sinensis* infection  
D  Primary sclerosing cholangitis  
E  Ulcerative colitis
**ANSWERS**

1. **B** The history points to an acute liver failure from fulminating hepatitis with massive hepatic necrosis. The loss of hepatic function from destruction of 80% to 90% of the liver results in hyperammonemia from the defective hepatocyte urea cycle, and this leads to hepatic encephalopathy within 2 weeks of the onset of jaundice. An elevated alkaline phosphatase level suggests extrahepatic or intrahepatic biliary obstruction. An elevated amylase level suggests pancreatitis. Fulminant hepatitis from HCV is rare. An autoimmune hepatitis with a positive ANA finding is not likely to produce a fulminant hepatitis.

   **PBD9 825–827 BP9 604 BPDB8 853 BP8 634–635**

2. **E** Stellate cells (formerly Ito cells) may transform into myofibroblasts secreting collagen when hepatocytes are injured and elaborate free radicals and cytokines. This process takes years, but is potentially reversible to some degree if the injurious stimulus is removed. If sufficient functioning hepatic parenchyma remains, the cirrhosis may be well-compensated. The remaining choices listed do not produce collagen. The normal space of Disse contains only a small amount of type IV collagen. Trail tip: eating polar bear liver, which contains large amounts of vitamin A, may produce vitamin A toxicity.

   **PBD9 822–824 BP9 607 BPDB8 835, 837 BP8 635–636**

3. **A** Cirrhosis with portal hypertension increases venous collateral flow in esophageal submucosal veins, producing varices, and in the abdominal wall, producing caput medusae. Hyperreflexia, but not diminution of deep tendon reflexes, can occur when hepatic encephalopathy develops from decompensated cirrhosis. Right-sided heart failure, in which the liver may be enlarged because of passive congestion, is associated with distended jugular veins. Liver failure with cirrhosis may lead to hepatic coma, but brain swelling with papilledema is not a major feature. The coagulopathy from decreased liver function may lead to purpuric hemorrhages, but splinter hemorrhages of the nails are most characteristic of embolization from infective endocarditis.


4. **B** The diffuse nodularity with depressed scars between the nodules is characteristic of cirrhosis, which led to her ascites complicated by spontaneous bacterial peritonitis and septicemia. The cirrhosis may be partially decompensated until infection occurs. A common cause of cirrhosis in the Western world is alcohol abuse. α1-Antitrypsin deficiency and hereditary hemochromatosis can result in cirrhosis, but both of these diseases are uncommon. In hereditary hemochromatosis, the liver has a dark brown gross appearance caused by extensive iron deposition. Of the various forms of viral hepatitis, those caused by HBV or HCV are most likely to be followed by cirrhosis. This complication is rare or nonexistent in HAV, HGV, and HEV infections. In sclerosing cholangitis, there is portal fibrosis, but not much nodular regeneration, so the liver is green and hard and has a finely granular surface.

   **PBD9 827–830 BP9 607–609 BPDB8 837–838 BP8 635–537**

5. **B** The SAAG is calculated by subtracting the ascitic albumin level from the serum albumin level, and it correlates with portal pressure. With the architectural remodeling of cirrhosis, there is portal hypertension and increased loss of hepatic interstitial fluid with protein into the peritoneal cavity. This is a transudative ascites. The serum albumin is likely to be low with chronic liver disease because of decreased synthetic capacity. The remaining choices include conditions that lead to an exudative ascites with SAAG less than 1.1.

   **PBD9 829 BP9 609 BPDB8 839 BP8 637**

6. **E** This patient most likely has alcoholic cirrhosis with hepatic failure and portal hypertension. The ascites is caused by portal hypertension which results from two major changes: (1) mechanical obstruction to blood flow in the liver due to scarring and compression of sinusoids by regenerating nodules, and (2) splanchic arterial vasodilation giving rise to hyperdynamic circulation which leads to increased portal venous blood flow. The latter is an important factor in the pathogenesis of portal hypertension and consequent ascites. The splanchic arterial vasodilation is caused by increased nitric oxide (NO) production in the splanchic arterial bed. This patient has no signs and symptoms of congestive heart failure—notice the normal JVP. Hyperbilirubinemia in this case is due to hepatic failure. In hepatopulmonary syndrome there are pulmonary intravascular dilations due to NO synthesis in the lung, not liver. Portosystemic shunts give rise to esophageal varices that bleed to cause hematemesis.

   **PBD9 826–829 BP9 607–609 BPDB8 838–839 BP8 636–637**

7. **C** Hepatopulmonary syndrome (portopulmonary hypertension) is described. The cause is obscure, but the result is pulmonary arterial vasoconstriction and ventilation-perfusion (V/Q) mismatches that lead to hypoxemia. The remaining choices are conditions that do not cause portal hypertension. Chronic inflammation and steatosis may be seen with cirrhosis, but by themselves do not account for portopulmonary hypertension. Metastases tend to be focal, leaving residual functioning hepatic parenchyma. Biliary obstruction leads to jaundice.

   **PBD9 830 BP9 610 BPDB8 836**

8. **F** Portal bridging fibrosis and nodular hepatocyte regeneration are features of cirrhosis. If liver stem cells support hepatocyte regeneration, and ductular reactions are minimal, then cirrhosis may be less progressive, and thin septae suggest some degree of regression. The massive upper gastrointestinal bleeding suggests esophageal varices as a consequence of portal hypertension from cirrhosis. If the patient is currently not drinking alcohol, no fatty change (steatosis) would be present. The architectural changes of cirrhosis persist for decades after cirrhosis develops. Concentric bile duct fibrosis is seen in primary sclerosing cholangitis, which may be idiopathic or may appear in association with inflammatory bowel disease. Budd-Chiari syndrome in hepatic venous thrombosis leads to hepatic enlargement, and it is rare. Interface hepatitis is a
characteristic of chronic active HBV or HCV infection. Massive hepatocellular necrosis may occur rarely as a complication of HAV infection or ingestion of massive amounts of acetaminophen. α1-Antitrypsin deficiency with the PAS-positive periportal globules is associated with development of cirrhosis, but this is far less common than alcoholic cirrhosis.

PBD9 823–824, 828 BP9 611–613 BPD8 837–838 BP8 635–636

9 D The figure shows interface hepatitis (formerly called piecemeal necrosis but better termed apoptosis of hepatocytes) at the limiting plate, with a mononuclear infiltrate, and Councilman bodies. Liver disease that has persisted for 6 months, and histologic evidence of hepatic necrosis with portal inflammation and fibrosis, are features of chronic hepatitis. Of all the hepatitis viruses, HCV is most likely to produce chronic hepatitis, and HAV is the least likely to produce chronic disease. Cholecodolithiasis leads to extrahepatic biliary obstruction and an elevated alkaline phosphatase level, but it is unlikely to produce hepatocellular necrosis. Hepatic congestion with right-sided heart failure produces centrilobular necrosis, but not portal fibrosis. Hemochromatosis can produce portal fibrosis and cirrhosis, but the liver cells show prominent accumulation of golden brown hemosiderin pigment. Sclerosing cholangitis leads to inflammation and obliterative fibrosis of bile ducts.

PBD9 837–838 BP9 612–613 BPD8 847–848 BP8 645–647

10 A He most likely developed a mild, self-limited liver disease from HAV infection after a meal at a restaurant with consumption of contaminated food or water. The presence of IgM anti-HAV indicates recent infection. The IgM antibody is replaced within a few months by IgG antibodies, which impart immunity to reinfection. The incubation period for HAV infection is short, and the illness is short and mild, with no significant tendency to develop chronic hepatitis. The most common mode of infection for HAV is via the fecal-oral route. HBV and HCV infections have longer incubation periods and are most often acquired parenterally. HDV infection develops from co-infection with HBV or by superinfection in a hepatitis B carrier.

PBD9 831 BP9 614 BPD8 844–845 BP8 640–641

11 E The most important predictor of whether a patient with viral hepatitis will develop chronic liver disease is the etiologic agent that caused the hepatitis. Of all the hepatotropic viruses, infection with HCV is the most likely to progress to chronicity and ultimately to cirrhosis. HAV, HEV, and HGV almost never cause chronic hepatitis. The pattern of histologic change, the degree of transaminase elevation, and the duration of transaminase elevation are poor predictors of chronicity.

PBD9 833–835 BP9 614–618 BPD8 847–848 BP8 643–644

12 C This intravenous drug user developed chronic HBV infection, as evidenced by the persistence of HBsAg, HBV DNA, and IgG anti-HBc antibodies. Of individuals with a history of intravenous drug use, 80% to 90% are found to have serologic evidence of HBV or HCV infection. Ruptured varices and ascites suggest that this patient subsequently developed cirrhosis and portal hypertension. His final presentation of weight loss and rapid enlargement of the abdomen suggests that a hepatocellular carcinoma has developed, and in most cases is confirmed by an elevated α-fetoprotein level. The other test findings, including prolonged prothrombin time, increased ALT level, and increased ferritin level, all indicate chronic liver disease. Any mass lesion in the liver is associated with an elevated alkaline phosphatase level. An increasing blood ammonia level indicates marked liver failure.


13 B The detection of IgM anti-HAV indicates acute infection. Progression of HAV infection to chronic hepatitis does not occur, but a few cases are complicated by fulminant hepatitis. HAV viremia is transient, so blood-borne transmission of HAV is rare. HAV is spread by the fecal-oral route, such as raw shellfish from a bay in which raw sewage is dumped.

PBD9 831–833 BP9 614–616 BPD8 846–847 BP8 641–643

14 E In regions where HBV is endemic, vertical transmission produces a carrier rate of 90% to 95%. However, successful implementation of a national childhood HBV immunization program can lead to a low prevalence of HBsAg among children and adolescents. Singapore achieved the World Health Organization goal to reduce the prevalence of chronic HBV infection. Development of viral hepatitis requires an immune response against virus-infected cells. In immunocompetent individuals, HBV induces T cells specific for HBsAg that cause apoptosis of infected liver cells. During the neonatal period, immune responses are not fully developed; hepatitis does not occur. The high carrier rate is medically significant because it increases the risk of hepatocellular carcinomas 200-fold. In populations with a high carrier rate, coexistent cirrhosis may be absent in 50% of patients. In contrast, in places where HBV is not endemic, cirrhosis is present in 80% to 90% of patients who develop liver cancer. HBV infection from blood transfusion is rare because of screening of blood products. Transmission of HBV via sexual contact is uncommon and induces a carrier state in a few cases; in most cases, an immune response is elicited. Oral transmission of HAV is common (but not HBV or HCV). The risk of acquiring HBV through needlestick injury is 1% to 6%.

PBD9 831–833 BP9 614–615 BPD8 845 BP8 641–643

15 A Persistence of HBsAg in serum for 6 months or more after initial detection denotes a carrier state. Worldwide, most individuals with a chronic carrier state for HBV acquired this infection in utero or at birth. Only 1% to 10% of adult HBV infections yield a chronic carrier state. The carrier state is stable in most individuals, the so-called “inactive” carrier state, without elevation in liver enzymes, and some infected persons may eventually clear the virus. There is currently no therapy to aid this viral clearance. Vaccination is useful to prevent infection, not clear the virus, although carriers become a reservoir for infection of others.

**16 C** Necrosis with portal bridging suggests chronic hepatitis. Mild steatosis is seen in HCV infection. The incidence of chronic hepatitis is highest with HCV infection. More than 50% of individuals infected with this virus develop chronic hepatitis, and many cases progress to cirrhosis. This is partly because the IgG antibodies against HCV that develop after acute infection are not protective.

**17 C** Antibodies to hepatitis C do not confer protection against reinfection. HCV RNA remains in the circulation, despite the presence of neutralizing antibodies. Treatment strategies may also target viral polymerase and protease, similar to antiretroviral regimens for HIV infection. In infections with HAV, HBV, HDV, HEV, or HGV, development of IgG antibodies offers lifelong immunity. An HBV vaccine exists for this purpose.

**18 E** This patient has evidence of HCV infection with symptoms of liver disease for 1 year. Clinically, she has chronic hepatitis (≥6 months), which may have followed an asymptomatic acute HCV infection. The anti-HCV IgG antibody is not protective. This is supported by continued HCV viremia. Approximately 85% of cases of HCV progress to chronic hepatitis, but fulminant hepatitis is uncommon. Chronic hepatitis is characterized by apoptosis of hepatocytes at the interface between portal tracts and the liver lobule. This eventually leads to cirrhosis with portal bridging fibrosis and nodular regeneration. At this time, however, the patient has no signs or symptoms of cirrhosis. Concentric bile duct fibrosis occurs in sclerosing cholangitis, which may be idiopathic or, more commonly, is associated with inflammatory bowel disease. Copper deposition is characteristic of Wilson disease, which may be associated with chronic hepatitis and cirrhosis, but it is not related to the much more common HCV infection. Granulomatous bile duct destruction suggests primary biliary cirrhosis. Budd-Chiari syndrome in hepatic venous thrombosis leads to hepatic enlargement and necrosis and to ascites.

**19 C** HDV cannot replicate in the absence of HBV; isolated HDV infection does not occur. The evidence for chronic hepatitis B is the presence of HBsAg and anti-HBc IgG antibody in the absence of anti-HBc IgM antibody. Confirmatory serologic evidence of recent HDV infection is the presence of anti-HDV IgM antibodies. HBV and HDV infections are likely to occur in drug users who inject parenterally. When HDV infection is superimposed on chronic HBV, three outcomes are possible: mild hepatitis B may be converted to fulminant disease; acute hepatitis may occur in an asymptomatic HBV carrier; or chronic progressive disease may develop, culminating in cirrhosis. The other listed viruses can cause infection by themselves.

**20 C** HEV infections are most common in East and South Asia. Spread is by a fecal-oral route. Most persons have a subclinical infection, but 1 in 7 develops acute hepatitis; death is uncommon, except in pregnant women. HEV does not go on to chronic hepatitis. CMV and HSV can be congenital infections, but are unlikely to affect the maternal liver significantly; CMV is a significant cause for liver failure in orthotopic transplants, and both can affect immunocompromised persons. EBV can affect the liver as part of infectious mononucleosis, but the infection is typically mild. Yellow fever is seen in tropical and subtropical regions of Africa and South America and is spread via mosquitoes.

**21 D** He has hepatitis B virus (HBV) infection. The most common outcome with HBV infection is recovery. He was asymptomatic, as evidenced by icterus, malaise, and nausea. His ALT and AST were not very high. Presence of anti-HBs is consistent with recovery. However, though recovery occurs, in the acute phase of the illness beyond incubation, he is highly infective to others. Fulminant hepatitis is infrequent with HBV, <1% of cases. Only 10% of cases progress to chronic hepatitis, and a subset of those go on to cirrhosis.

**22 A** Parasitic liver flukes are endemic to East Asia and Southeast Asia. Infection may be asymptomatic for years, but can progress to a chronic phase complicated by recurrent pyogenic cholangitis and jaundice. There is risk for development of cholangiocarcinoma, the second most common primary hepatic malignancy. The other listed foils do not carry a risk for neoplasia. Echinococcosus leads to hydatid disease, but not to malignancy. The extraerythrocytic phase of malaria with plasmodium infection includes the liver. Disseminated tuberculosis produces granulomas, usually small and multifocal. Typhoid fever can be a systemic disease with liver involvement.

**23 B** Autoimmune hepatitis is a chronic liver disease of unknown cause in which antibodies to hepatocyte structural components cause progressive necrosis of hepatocytes, leading to cirrhosis and liver failure. Patients tend to improve with glucocorticoid therapy. 1-Antitrypsin deficiency and Wilson disease may be associated with chronic hepatitis and cirrhosis, but autoimmune markers are absent. Chronic alcoholism is not associated with formation of autoantibodies. Because this patient does not have evidence of HBV infection, there can be no superinfection with HDV. Isoniazid may cause an acute or chronic hepatitis, but without autoantibodies. Patients with primary biliary cirrhosis often have antimitochondrial antibody (which also can be seen in autoimmune hepatitis), but the bilirubin concentration and alkaline phosphatase level would be much higher in primary biliary cirrhosis.

**24 D** Acute hepatic injury is described. Isoniazid used to treat tuberculosis may produce hepatotoxicity, and in most
cases it is mild, but in some cases it leads to massive hepatic necrosis. There may be minimal symptoms until severe injury has occurred. The remaining choices include conditions that are more likely to produce a pattern of chronic hepatic injury.

PBD9 840–841  BP9 612–613  PBD8 856–857  BP8 652–653

25  **A**  In the setting of chronic liver disease, ingestion of acetaminophen is more likely to produce hepatotoxicity because detoxification by conjugation is exceeded. This leads to metabolism by cytochrome P-450 to the toxic metabolite \( \text{N-acetyl-p-benzoquinineimine} \) (NAPQI), which accumulates beyond the capacity of glutathione. The \( \text{N-acetylcysteine} \) increases available glutathione. Aspirin ingestion is a cause for Reye syndrome in children. Ibuprofen, which is a non-steroidal anti-inflammatory drug (NSAID), meperidine, and oxycodone do not have significant hepatotoxicity.

PBD9 422, 825–826, 841  BP9 610–611  PBD8 856–857  BP8 652–653

26  **C**  Portal fibrosis and nodular hepatocyte regeneration are typical features of chronic alcohol abuse. Spider telangiectasias (angiomata) refer to vascular lesions in the skin characterized by a central, pulsating, dilated arteriole from which small vessels radiate. These lesions result from hyperestrogenism (which also contributes to the testicular atrophy). The failing liver is unable to metabolize estrogens normally. Spider angiomata are a manifestation of hepatic failure. Ascites, splenomegaly, hemorrhoids, and esophageal varices all are related to portal hypertension from cirrhosis and the resultant collateral venous congestion and dilatation. The other listed findings do not explain portal hypertension.


27  **E**  The figure shows macrovesicular steatosis (fatty change) of the liver with early fibrosis. The most common cause of fatty liver and fibrosis is chronic alcoholism. In patients with no history of significant ethanol ingestion, a nonalcoholic steatohepatitis may be considered, with obesity, diabetes mellitus, or both as possible causes. Excessive acetaminophen ingestion can cause centrilobular necrosis or diffuse necrosis. Allopurinol toxicity can cause granuloma formation. Aspirin may be associated with a microvesicular steatosis as Reye syndrome in children. Chlorpromazine can lead to cholestasis. Isoniazid use can be complicated by acute injury and granuloma formation.


28  **B**  The elevated transaminase levels, some loss of liver function with abnormal prothrombin time, and cholestasis are not specific for a given type of liver injury. An AST level that is higher than the ALT level is characteristic, however, of liver cell injury associated with chronic alcoholism. In this patient, the disease is decompensating, as evidenced by the elevated blood ammonia level. HAV is typically a mild disease without a preponderance of direct bilirubin. Choledocholedolithiasis results in a conjugated hyperbilirubinemia, but without the high ammonia level that is evidence of liver failure. Metastases are unlikely to obstruct all biliary tract drainage or lead to liver failure severe enough to cause elevations of blood ammonia. Primary biliary cirrhosis is rare, particularly in men, and the alkaline phosphatase level would be much higher.

PBD9 844  BP9 623–624  PBD8 835–836  BP8 633–635

29  **E**  This is acute alcoholic hepatitis. The figure shows globular eosinophilic cytoplasmic inclusions called \textit{Mallory-Denk} bodies. These cytokeratin inclusions are characteristic of, but not specific for, alcoholic hepatitis. There also are areas of hepatocyte necrosis surrounded by neutrophils. Some neutrophils can be seen in the figure. Centrilobular congestion can lead to centrilobular necrosis without inflammation or Mallory-Denk bodies. Cholestasis is marked by plugs of yellow-green bile in canaliculi. Hemosiderin appears granular and brown on H&E staining, but it is blue with Prussian blue stain. Periportal PAS-positive globules are characteristic of \( \alpha_{1} \)-antitrypsin deficiency; the globules tend to be smaller than Mallory-Denk bodies, and there is no acute inflammation.

PBD9 843  BP9 623–624  PBD8 858  BP8 650

30  **F**  Nonalcoholic fatty liver disease (NAFLD) occurs in persons with metabolic syndrome/type 2 diabetes mellitus, with impaired fatty acid and lipoprotein metabolism. Some cases may go on to steatohepatitis with fibrosis and even cirrhosis (so-called cryptogenic cirrhosis when alcohol use could not be blamed, but remember to ask the CAGE questions). Though there may be hepatocyte loss and inflammation, these are not the most prominent features. Release of tumor necrosis factor alpha (TNF-\( \alpha \)) and interleukin-6 (IL-6) from dysfunctional adipocytes may drive hepatocyte apoptosis to start the process. Bile stasis is unlikely with NAFLD, and jaundice is not a key feature. Hemosiderin deposition is a function of iron absorption from diet or genetics.

PBD9 845–847  BP9 625  PBD8 860–861  BP8 654

31  **E**  Nonalcoholic fatty liver disease (NAFLD) may progress to nonalcoholic steatohepatitis (NASH) and even to cirrhosis. Risk factors of metabolic syndrome and type 2 diabetes mellitus are driven by obesity. NAFLD may be found in a third of adults, closely paralleling the prevalence of obesity. Pathologic features with NAFLD and NASH are similar to those of alcoholic liver disease. Type 1 diabetes mellitus leads to a catabolic state with weight loss. Familial hypercholesterolemia mainly drives atherosclerosis, without liver disease. Chronic viral hepatitis may have an element of steatosis, but not marked, and without vascular disease. The iron accumulation of hemochromatosis may produce cardiomyopathy as well as chronic liver disease without much steatosis.

PBD9 845–847  BP9 625  PBD8 860–861  BP8 654

32  **E**  The microvesicular steatosis characteristic of Reye syndrome rarely occurs now because the link between aspirin use in children following a fever and hepatic injury has been
recognized; it is caused by severe mitochondrial dysfunction in the brain and liver. Biliary atresia with marked hyperbilirubinemia becomes apparent in the neonatal period. Hepatic venous thrombosis leads to Budd-Chiari syndrome, which is typically a disease of adults that complicates such conditions as polycythemia or pregnancy. Hepatoblastomas may be congenital, but they are mass lesions unlikely to be associated with such marked increases in liver enzymes. Intrahepatic l illness is unlikely to occur in children and is unlikely to produce marked increases in liver enzymes. Neonatal giant cell hepatitis can produce findings of acute hepatitis in neonates, not in children.

PBD9 841  BP9 626  PBD8 856–857  BP8 657–658

33 B Anabolic androgenic steroids are injected for the purpose of increasing muscle mass to enhance athletic performance. Because performance is primarily correlated with skill and training, the potential gain from muscle mass is problematic, particularly in view of the deleterious effects, such as hepatic cholestatic hepatitis. Acetaminophen can produce hepatic necrosis, but icterus is less likely. Chlorpromazine is more likely to produce a pure cholestasis as an idiosyncratic (unpredictable) reaction. Ethanol can produce steatosis, and in excess the AST is often higher than the ALT. Isoniazid may produce hepatic necrosis in persons treated for tuberculosis.

PBD9 841  BP9 629  PBD8 822–825, 863–865  BP8 655

34 D An elevated alkaline phosphatase level suggests obstruction of the biliary tract, but this case must be focal because the bilirubin is not elevated. The microcytic anemia and the blood in the stool suggest gastrointestinal tract hemorrhage, and a colonic adenocarcinoma should be suspected as the primary site for the hepatic metastases in this case. Hepatic metastases from colon cancer are common. They appear as multiple masses within the hepatic parenchyma. Antiphospholipid syndrome predisposes to thrombosis with venous obstruction, in which case hepatic enzyme levels should be higher, and the partial thromboplastin time should be prolonged. Ascending cholangitis is typically caused by bacteria such as Escherichia coli or Klebsiella, and patients develop acute symptoms of fever, chills, jaundice, and abdominal pain. Chronic alcoholism is not accompanied by an increase in the alkaline phosphatase level, and there is often a macrocytic anemia. Sclerosing cholangitis would increase the bilirubin concentration and the alkaline phosphatase level.

PBD9 822, 853–854  BP9 605–606  PBD8 840–841  BP8 663

35 D This is neonatal physiologic jaundice that is mild and transient and is due to diminished hepatic conjugating and excreting capacity for bilirubin that is not fully functional until 2 weeks of life. Biliary atresia with obstruction produces more severe jaundice and requires surgical intervention. Neonatal hepatitis can be due to congenital infections that produce more severe jaundice that persists more than 2 weeks. The rare Dubin-Johnson syndrome can occur with autosomal recessive mutation of a gene encoding a canalicular transporter protein that impairs bilirubin excretion. Erythrobástasis fetalis from maternal sensitization to fetal RBC antigens is unlikely to affect the first pregnancy.

PBD9 853–857  BP9 605–606  PBD8 841  BP8 639

36 B An unconjugated hyperbilirubinemia can result from hemolysis. With increased RBC destruction, there is more bilirubin than can be conjugated by the hepatocytes. Obstructive jaundice with biliary tract lusis results in mostly conjugated hyperbilirubinemia. The total bilirubin concentration may be increased in patients with viral hepatitis or cirrhosis and in individuals taking drugs such as oral contraceptives. Although direct and indirect hyperbilirubinemia may occur in these conditions, conjugated hyperbilirubinemia predominates.

PBD9 853–854, 877  BP9 605–606  PBD8 841, 884  BP8 667–668

37 D Gilbert syndrome results from decreased levels of uridine diphosphate glucuronosyltransferase (UDPGT), which may be the status of 3% to 7% of the general population of the United States. Lower levels are more common in Africa but less frequent in Asia. The condition is often never diagnosed. Stress may cause transient unconjugated hyperbilirubinemia to a point that scleral icterus is detectable, when the serum bilirubin reaches about 2 to 2.5 mg/dL. Acetaminophen in small quantities can be properly detoxified, but ingestion of large quantities can produce hepatocyte necrosis. Choledochal cyst is a rare congenital anomaly producing extrahepatic biliary obstruction with conjugated hyperbilirubinemia. Primary biliary cirrhosis results in conjugated hyperbilirubinemia, as does the rare Dubin-Johnson syndrome. HAV infection can often be mild, but it is not so transient; it can be accompanied by a mild increase in conjugated and unconjugated bilirubin.

PBD9 853–854  BP9 606  PBD8 841–842  BP8 639

38 E This patient has a history of gallstones and has developed an ascending cholangitis caused by Escherichia coli. These bacteria reach the liver by ascending the biliary tree. Obstruction from lithiasis is the most common risk factor. Development of cystic lesions in the right lobe of the liver suggests that the patient has developed liver abscesses. Clonorchis sinensis is a liver fluke that is endemic to East Asia, and it is a risk factor for biliary tract cancer. Cryptosporidiosis in immunocompromised patients occasionally can occur in the biliary tract and elsewhere. Cytomegalovirus infection also can be seen in immunocompromised patients; it produces a clinical picture similar to that of hepatitis, but without biliary tract disease. A patient with amebiasis involving the liver is most likely to present with a history of diarrhea with blood and mucus.

PBD9 839, 854–855  BP9 642  PBD8 887  BP8 670

39 D Neonatal cholestasis is newborn jaundice that persists more than 2 weeks following birth. One cause is neonatal hepatitis, which is most often idiopathic, and most infants recover without specific therapy. Some cases are caused by α1-antitrypsin deficiency, and some are due to extrahepatic
biliary atresia. Patients with extrahepatic biliary atresia will have a high alkaline phosphatase and require surgery to anastomose extrahepatic ducts and prevent progressive liver damage. Patients with erythroidosis fetalis have hydrops and icterus at birth because of maternal IgG antibody directed at fetal RBCs, leading to hemolysis. Galactosemia is an inborn error of metabolism in which deficiency of galactose-1-phosphate uridylytransferase damages cells of the kidney, liver, and brain; there is hepatomegaly, splenomegaly, hypoglycemia, and eventually cirrhosis. Primary biliary cirrhosis affects adults. Von Gierke disease results from deficiency of glucose-6-phosphatase, and affected infants develop hypoglycemia, lactic acidosis, hyperuricemia, and hyperlipidemia.

40 D Extrahepatic biliary atresia is a rare condition in which some or all of the bile ducts are destroyed. If the disease spares a large enough bile duct to anastomose around the obstruction, the problem may be correctable. In many cases, such as this one, obstruction of bile ducts occurs above the porta hepatis, however, and the only option for treatment is liver transplantation. α1-Antitrypsin deficiency can produce a neonatal hepatitis that may clinically resemble extrahepatic biliary atresia, but most infants recover. A choledochal cyst may cause biliary colic in children; it is a congenital condition that produces dilations of the common bile duct. Congenital infections may involve the liver, and usually other organs as well; infants with these infections are ill from birth. Hepatoblastomas are rare and may be seen in infancy, but mass lesions in the hepatic parenchyma typically do not obstruct the biliary tree completely.

41 E Primary biliary cirrhosis is an uncommon autoimmune disorder that causes progressive intrahepatic bile duct destruction. Pruritus, conjugated hyperbilirubinemia, and increased alkaline phosphatase levels are indicative of obstructive jaundice resulting from bile duct destruction. About 90% or more of patients with this disease have antimitochondrial antibodies in the serum. α1-Antitrypsin deficiency can affect the liver, causing chronic hepatitis and cirrhosis, and causes panlobular emphysema. An elevated sweat chloride level is found in cystic fibrosis, which can cause neonatal jaundice. An increased serum ferritin level is seen in patients with hereditary hemochromatosis. Chronic hepatitis C is marked by hepatocyte necrosis, not by bile duct destruction.

42 A The major targets in primary sclerosing cholangitis are intrahepatic bile ducts, and ulcerative colitis coexists in 70% of cases. Ducts undergo a destructive cholangitis that leads eventually to periductal fibrosis and cholestatic jaundice. Eventually, cirrhosis and liver failure can occur. Copper deposition is characteristic of Wilson disease, which is associated with chronic hepatitis and cirrhosis. Granulomatous bile duct destruction occurs in primary biliary cirrhosis. Interface hepatitis is characteristic of chronic active viral hepatitis. α1-Antitrypsin deficiency with PAS-positive perportal globules is associated with cirrhosis. Portal bridging fibrosis with nodular regeneration defines cirrhosis.

43 C This patient has clinical, histologic, and laboratory features of genetic hemochromatosis. In this condition, iron overload occurs because of excessive absorption of dietary iron. The absorbed iron is deposited in many tissues, including the heart, pancreas, and liver, giving rise to heart failure, diabetes, and cirrhosis. It appears blue with Prussian blue stain, as seen in this figure. High serum ferritin concentration is an indicator of a vast increase in body iron. Genetic hemochromatosis is an autosomal recessive condition; siblings are at risk of developing the same disease. Phlebotomy removes 250 mg of iron per unit of blood, and over time can reduce iron stores.

44 D This patient has hereditary hemochromatosis from excessive iron storage leading to ascertes, splenomegaly, impaired liver function, diabetes mellitus, skin pigmentation, and elevated ferritin. Because he has no predisposing causes for increased iron absorption, the most likely diagnosis is primary, or genetic, hemochromatosis. This disease results from a mutation in the HFE gene that encodes for an HLA class I–like molecule that binds β2-microglobulin. Mutant HFE reduces hepcidin synthesis to decrease circulating hepcidin. The resulting decreased hepcidin-ferroportin interaction allows for increased ferroportin activity, increased iron efflux from enterocytes, giving rise to systemic iron overload in hereditary hemochromatosis. Because this patient is not anemic and has a normal MCV, a β-globin gene mutation with β-thalassemia is unlikely. None of the other listed mutations affect iron absorption.

45 C Although most cases of hereditary hemochromatosis result from mutations of the HFE gene, some cases may occur from mutations of genes encoding for transferrin receptors, hemojuvelin, and rarely hepcidin (encoded by the HAMP gene). However, the main regulator of iron absorption is the protein hepcidin and all the genetic causes of hereditary hemochromatosis are associated with reduced hepcidin levels. Mutations in the HAMP gene produce extremely severe disease at an early age (juvenile hemochromatosis) with cardiomyopathy and hypogonadism. Ordinarily the liver increases hepcidin production when iron stores are adequate, preventing release of iron from intestinal enterocytes and macrophages. MHC Class I proteins are involved in peptide antigen recognition for T-cell mediated immunity. ATP7B gene mutations are present with Wilson disease, a disorder of copper metabolism. Mutations of HNF1-α are seen with maturity onset diabetes of the young (MODY) and can lead to the appearance of hepatic adenomas. DMT1 is involved in enterocyte iron absorption.

46 B Wilson disease is an inherited disorder in which toxic levels of copper accumulate in tissues, particularly the
The ATP7B gene for Wilson disease encodes a copper-transporting ATPase in the hepatocytes. With mutations in this gene, copper cannot be secreted into plasma. Ceruloplasmin is an $\alpha_2$-globulin that carries copper in plasma. Because copper cannot be secreted into plasma, ceruloplasmin levels are low. Chronic liver disease and panlobular emphysema may occur in $\alpha_1$-antitrypsin deficiency. An increased $\alpha$-fetoprotein is a marker for hepatocellular carcinoma. An increased serum ferritin may indicate hereditary hemochromatosis. A positive finding for antimitochondrial antibody can be seen in primary biliary cirrhosis. A positive HBsAg result indicates HBV, which infects only the liver.

47 C The PAS-positive globules in the liver seen here are characteristic of $\alpha_1$-antitrypsin (AAT) deficiency. Approximately 10% of individuals with the homozygous deficiency (PiZZ phenotype) of AAT deficiency develop significant liver disease, including neonatal hepatitis and progressive cirrhosis. Deficiency of AAT also allows unchecked action of elastases in the lung, which destroys the elastic tissue and causes emphysema. AAT can produce a picture of chronic hepatitis in adults; it can lead to neonatal hepatitis with acute but often transient liver injury. Diabetes mellitus and heart failure are features of hemochromatosis, a condition of iron overload. Iron deposition in liver is detected by the Prussian blue stain. Systemic lupus erythematosus is an immune complex disease that may affect many organs. Liver involvement is uncommon, however. Ucercolitis coli is strongly associated with primary sclerosing cholangitis, a condition in which there is inflammation and obliterative fibrosis of bile ducts.

48 C This is Budd-Chiari syndrome, which results from hepatic venous outflow obstruction. The high hemoglobin and platelet count in this woman is consistent with polycythemia vera as part of a myeloproliferative disorder, and this is the likely cause. Cholelithiasis should lead to jaundice. Cirrhosis may also lead to caput medusae and increasing girth from ascites, but polycythemia is not seen with cirrhosis. Some hepatocellular carcinomas, arising in the setting of cirrhosis, may produce Budd-Chiari syndrome, but there is no polycythemia. Fatty change alone does not produce hepatic venous obstruction. An abscess below the diaphragm is unlikely to impinge upon the hepatic vein.

49 B The microscopic appearance is that of intense centrilobular congestion. The area around the portal tract is less congested. The restrictive lung disease leads to cor pulmonale with right-sided congestive heart failure. This causes passive venous congestion in the liver that is most pronounced in the centrilobular areas. When congestion is severe, the anoxia can cause centrilobular necrosis with transaminase elevation. Apoptosis does not produce widespread necrosis because single cells are involved, and this is most typical of viral hepatitis. Cholestasis is marked by plugs of yellow-green bile in canaluli. Hemosiderin appears granular and brown on H&E staining, but it is blue with Prussian blue stain. Large lipid droplets fill the hepatocyte cytoplasm with macrovesicular steatosis. Mallory bodies are globular red cytoplasmic structures most characteristic of alcoholism, in particular, acute alcoholic hepatitis. Portal fibrosis begins the process of cirrhosis.

50 E She has maternal HELLP syndrome (hemolysis, elevated liver enzymes, and low platelets), which is a severe complication of preeclampsia, and when a coagulopathy is apparent from the increasing prothrombin time, emergent delivery must be undertaken to save the life of the mother and fetus. Corticosteroid therapy is used for inflammatory conditions, typically those that have an autoimmune basis. Intravenous immunoglobulin may aid in treating infections, such as hepatitis B immunoglobin. Liver transplantation is indicated with liver failure from which no potential recovery is possible. Plasmapheresis aids in treating conditions such as thrombotic thrombocytopenic purpura, which have circulating antibodies, proteins, or toxins that can be removed emergently. Portacaval shunt is used to treat cirrhosis with hepatic encephalopathy. Ribavirin and interferon therapy is used for viral hepatitis C infection.

51 D She has acute fatty liver of pregnancy, an uncommon condition of variable severity. Accumulation of small droplets of fat in hepatocytes (macrovesicular steatosis) is the typical histologic finding. This feature is not seen in any of the other conditions listed. The disease may occur because of a defect in mitochondrially fatty acid oxidation. Concentric bile duct fibrosis is a feature of sclerosing cholangitis. Hereditary hemochromatosis manifests with complications in middle age after extensive iron deposition has occurred. Extrahepatic biliary atresia is a rare neonatal disease. The loss of intrahepatic bile ducts in primary biliary cirrhosis also is a rare disease of middle age. Multinucleated giant cells may be seen in neonatal giant cell hepatitis. PAS-positive globules are seen in $\alpha_1$-antitrypsin deficiency, a condition that affects adults.

52 B Focal nodular hyperplasia is a well-demarcated but unencapsulated benign lesion that is characterized by a central scar. Cholangiocarcinomas have extensive collagen deposition, but are malignant and often associated with risk factors including viral hepatitis. Hepatic adenomas can be seen with oral contraceptive use. Hepatocellular carcinomas occur after liver injury from such conditions as viral hepatitis or alcohol abuse, and they often can be multifocal. Cirrhosis involves the entire liver, not just a portion. Metastases can be solitary or multiple and can have central necrosis when large.

53 E The circumscribed mass in the liver suggests a benign tumor, such as hepatic adenoma. These tumors, which may...
develop in young women who have used oral contraceptives, can enlarge and rupture from estrogenic stimulation during pregnancy. These adenomas may have HFN1-α or β-catenin gene mutations. Cholangiocarcinomas and hepatocellular carcinomas can be related to viral hepatitis infection and alcoholism; they are large, irregular masses that tend to occur in patients who are older than this woman. Choledochal cysts of the biliary tract are rare embryonic remnants that typically become symptomatic in childhood, along with biliary colic. Molar pregnancy can include choriocarcinoma, which can metastasize and rupture, but a solitary circumscribed metastasis is unlikely. Hepatoblastomas are rare liver neoplasms found in children.

54 B Multiple masses shown in the figure are consistent with metastatic carcinoma. Metastases in liver are far more common than primary liver neoplasms. The microcytotic anemia and the stool positive for occult blood suggest colon cancer. Mass lesions cause intrahepatic biliary obstruction to elevate the alkaline phosphatase. The hepatic parenchyma has a rich vascular supply that makes it a target for hematogenous spread of malignancies. Cirrhosis and the potential for primary hepatocellular carcinoma may ensue from either alcohol abuse or viral hepatitis B, but there is likely a dominant mass lesion. Gilbert syndrome and other hereditary hyperbilirubinemias are unlikely to lead to malignancies. Metabolic syndrome and diabetes mellitus increase the risk for NAFLD, NASH, cirrhosis, and hepatocellular carcinoma.

55 B Dysplastic nodules are precursors to hepatocellular carcinomas. Regenerative nodules as part of cirrhosis tend to be smaller than 2 cm, and often smaller than 1 cm, with portal venous blood supply. As dysplasia progresses to a high-grade level, there is increased angiogenesis with arteri-alization. Hepatic adenomas tend not to undergo malignant transformation. Siderotic nodules may occur with hemo-chromatosis. Steatosis is a feature of many hepatic diseases, and is not a premalignant finding.

56 E There is a long-term risk of hepatocellular carcinoma in patients infected with HBV. This infection is more common (often from vertical transmission) in Asia than in North America and Europe, and it accounts for more cases of primary liver cancer worldwide than other causes, such as chronic alcoholism. HBV does not encode any oncogene, and it does not integrate next to a known oncogene, such as NMYC. Most likely, neoplastic transformation occurs because HBV induces repeated cycles of liver cell death and regeneration. This repeated cycling increases the risk of accumulating mutations during several rounds of cell division. Infection with the liver fluke Clonorchis sinensis predisposes to bile duct carcinoma. In contrast to colon carcinomas, hepatic carcinomas are not known to develop from adenomas. Hereditary nonpolyposis colon carcinoma syndrome is associated with inherited DNA mismatch repair genes.

57 B Aflatoxin is a hepatotoxin and is the product of the fungus Aspergillus flavus, which grows on moldy cereals. Aflatoxin can be carcinogenic, leading to hepatocellular carcinoma, as shown in the figure. Ingestion of large amounts of acetaminophen leads to hepatocellular necrosis. Aspirin has been implicated in causing Reyes syndrome in children, which results in extensive microvesicular steatosis. Prolonged and excessive intake of oral iron rarely can cause secondary hemochromatosis. Nitrites have been causally linked with cancers in the upper gastrointestinal tract. Oysters can concentrate hepatitis A virus (HAV) from seawater contaminated with sewage, and eating raw oysters can result in HAV infection.

58 A The findings suggest obstructive jaundice from biliary tract disease (e.g., gallstones). Elevation of the serum alkaline phosphatase level is characteristic of cholestasis from biliary obstruction. The alkaline phosphatase comes from bile duct epithelium and hepatocyte canalicular membrane. The blood ammonia concentration increases with worsening liver failure. When hepatic failure is sufficient to cause hyperammonemia, mental obtundation is seen. In this case, the patient has only jaundice. Primary biliary cirrhosis with an increased anti-mitochondrial antibody titer is much less common, and PBD eventually leads to bile duct destruction. Most cases of active HCV infection are accompanied by some degree of inflammation with fibrosis. In obstructive biliary tract disease, the direct bilirubin, not the indirect bilirubin, should be elevated.

59 B Intermittent upper abdominal pain, most often on the right, is a nonspecific symptom that often occurs in patients with gallstones. When a stone passes through the cystic duct and into the common bile duct, or forms in the common duct, intrahepatic cholestasis occurs. This explains the conjugated hyperbilirubinemia and the increased alkaline phosphatase level. Most reported cases of choledochal cyst, much less common than gallstones, come from Japan; the biliary stasis promotes infection with hydrolysis of bilirubin digluconides to form brown stones. Chronic passive congestion from heart failure does not typically produce hyperbilirubinemia. Hemoglobinopathies with risk for hemolysis produce unconjugated hyperbilirubinemia. Active viral hepatitis should be accompanied by some hepatocellular necrosis with liver enzyme elevation. Ulcerative colitis is associated with sclerosing cholangitis. Venoocclusive disease is rare and is accompanied by hyperbilirubinemia and cholestasis, but not by biliary tract obstruction.
The figure shows cholesterol gallstones. These stones are pale yellow, but acquire a variegated appearance by trapping bile pigments. In comparison, pigment stones are uniformly dark. Risk factors for such cholesterol stones include Native American descent, female sex, obesity, and increasing age. These factors cause secretion of bile that is supersaturated in cholesterol. Patients with RBC hemolysis develop pigment stones, whether the hemolysis is antibody-mediated (autoimmune hemolytic anemia), or whether it is caused by intrinsic RBC abnormalities (hemoglobinopathies such as sickle cell anemia). Infection of the biliary tract (Escherichia coli, Ascaris worms, or liver flukes) can lead to increased release of β-glucuronidases that hydrolyze bilirubin glucuronides, favoring pigment stone formation. Renal failure with phosphate retention can be a cause of secondary hyperparathyroidism with hypercalcemia, which increases the risk of gallstone formation; these stones are mixed stones. Viral hepatitis is not a risk factor for stone formation. Fatty foods may trigger the biliary colic, but diet does not play a direct role in stone formation. Severe ileal dysfunction, as occurs in Crohn disease, also can predispose to pigment stones.

This patient has acute acalculous cholecystitis. The tachycardia, hypotension, and sepsis with recent illness suggest that his fluid intake has been poor, with subsequent dehydration that concentrates bile and promotes stasis. Sepsis promotes hypotension that increases ischemia of the gallbladder. However, an isolated, de novo elevation of bilirubin in a hospitalized patient should suggest the possibility of sepsis. Adenocarcinoma of the pancreas may produce biliary tract obstruction, but the CT in this case did not identify a mass. The CT should also show any gallstones that are present. Intrahepatic cholestasis is more likely to be an adverse drug reaction.

Almost all gallbladder carcinomas are adenocarcinomas, and most are found in gallbladders that also contain gallstones. Although alcohol abuse is associated with liver disease, it is not a significant antecedent to gallbladder disease. Infection with the biliary tree fluke Clonorchis sinensis is a risk factor for biliary tract cancer, not gallbladder cancer. Similarly, primary sclerosing cholangitis increases the risk of developing cholangiocarcinoma. Ulcerative colitis is associated with primary sclerosing cholangitis.
A 24-year-old, G2, P1001, woman has a screening ultrasound that shows polyhydramnios at 17 weeks' gestation. She gives birth at term to a boy who on examination has no abnormal findings. Since birth, he has often vomited after feedings. An infant radiograph with contrast enhancement shows the findings in the figure. Laboratory studies show sodium, 130 mmol/L; potassium, 3.4 mmol/L; chloride, 85 mmol/L; CO₂, 32 mmol/L; glucose, 65 mg/dL; and amylase, 15 U/L. What is the most likely diagnosis?

A Acute pancreatitis
B Annular pancreas
C Chronic pancreatitis
D Islet cell adenoma
E Pancreatic adenocarcinoma
F Pyloric stenosis

A 16-year-old boy incurs a gunshot wound to the abdomen in a drive-by shooting. At exploratory laparotomy, the surgeon finds a perforated portion of jejunum. She resects this portion and palpates a mass in the jejunal submucosa. Sectioning of this lesion reveals that it is a 1.5-cm diameter circumscribed, solid, tan mass. The mass is sent for frozen section. What is the pathologist most likely to see under her microscope?

A Adenocarcinoma
B Adrenal medulla
C Gastric mucosa
D Non-Hodgkin lymphoma
E Pancreatic acini

A 33-year-old woman with Hodgkin lymphoma of cervical lymph nodes has an abdominal CT scan for staging. The scan reveals a solitary 4-cm cyst in the body of the pancreas. A fine-needle aspirate is performed and yields clear serous fluid that microscopically has low cuboidal cells with no atypia. What is this cyst most likely to be?

A Adenocarcinoma
B A component of autosomal dominant polycystic kidney disease
C Congenital malformation
D Cystadenoma, serous
E Pseudocyst
4. A clinical study of patients with acute abdomen is performed. Those patients with elevated serum lipase are identified. A decrease in which of the following analytes is most likely to predict a worse prognosis?
   A. Albumin
   B. Bilirubin
   C. Calcium
   D. Fibrinogen
   E. Haptoglobin

5. A 35-year-old man has a 1-year history of bouts of dull abdominal pain. Over the next 5 years he also develops steatorrhea and mild glucose intolerance. He does not develop jaundice. An abdominal CT scan shows specks of calcification in the midabdomen, particularly near the duodenum. One of his sisters has a similar clinical picture, but both parents are unaffected. Loss of inhibition of which of the following is the most likely cause for the disease seen in these persons?
   A. Amylase
   B. Complement
   C. Lipase
   D. Transforming growth factor beta (TGF-β)
   E. Trypsin

6. A 52-year-old man has had severe abdominal pain for the past 2 days. Physical examination shows boardlike rigidity of the abdominal muscles, making further examination difficult. There is no observable abdominal distention. The representative gross appearance of the disease process is shown in the figure. Which of the following is the mechanism most likely to produce this appearance?
   A. CFTR gene mutation
   B. Coxsackievirus B infection
   C. Dysregulation of trypsinogen inactivation
   D. Marked hypertriglyceridemia
   E. Vasculitis with acute ischemia
   F. Blunt force trauma to the abdomen

7. An 11-year-old boy has experienced multiple bouts of severe abdominal pain for the past 6 years, but no other medical problems. His father and grandfather have been similarly affected. On examination during the last episode, bowel sounds were absent, and he exhibited marked diffuse abdominal pain with abdominal wall rigidity. A radiograph of the abdomen showed dilated loops of bowel with air-fluid levels. Laboratory studies showed serum amylase of 3320 U/L. This boy is most likely to have an inherited mutation involving which of the following genes?
   A. CFTR
   B. KRAS
   C. PRSS1
   D. SMAD4
   E. SPINK1

8. A 38-year-old woman with a long history of gallbladder disease has a sudden onset of severe midabdominal pain. On physical examination, she has marked abdominal tenderness, particularly in the upper abdomen, and bowel sounds are reduced. An abdominal radiograph shows no free air, but there is marked soft tissue edema. Abdominal CT scan shows decreased attenuation with fluid density involving the pancreas. She is given intravenous fluids and nasogastric suction and recovers gradually. Which of the following serum laboratory tests is most useful for diagnosis of her disease process?
   A. Alanine aminotransferase
   B. Ammonia
   C. Bicarbonate
   D. Bilirubin
   E. Glucose
   F. Amylase

9. A 63-year-old man who had worsening congestive heart failure with cardiac dysrhythmias for the past year of his life died of pneumonia. At autopsy, his pancreas is grossly small and densely fibrotic. Microscopic examination shows extensive atrophy of the acini with residual chronic inflammation, fibrosis, and inspissated protein plugs in small, obstructed pancreatic ducts. Some of the protein plugs show calcification. The islets of Langerhans appear normal. The heart weighs 500 g, and all four chambers are dilated. Which of the following conditions is most likely to account for his findings?
   A. α1-Antitrypsin deficiency
   B. Blunt trauma to the abdomen
   C. Cholelithiasis
   D. Chronic alcoholism
   E. Cystic fibrosis
   F. Hypercholesterolemia
10 A 39-year-old man has had numerous bouts of pneumonia caused by *Pseudomonas aeruginosa* and *Burkholderia cepacia* for the past 35 years. He now has diarrhea of mild-to-moderate volume. On physical examination, he has decreased breath sounds and dullness to percussion in both lungs. His stool guaiac test is negative. Laboratory studies show the ΔF508 mutation. His quantitative stool fat is 7.5 g/day. Which of the following pathologic findings is most likely to be present in the pancreas of this patient?

A. Acute inflammation  
B. Acinar atrophy  
C. Adenocarcinoma  
D. Amyloidosis  
E. Pseudocyst

11 A 36-year-old woman with a history of pulmonary infections since childhood has had chronic abdominal pain for 4 years, along with a 6-kg weight loss. On physical examination her BMI is 18. There is pitting edema on the lower leg. Laboratory studies show serum albumin 3 g/dL, total protein 5.2 g/dL, and glucose 155 mg/dL. Abdominal CT imaging shows calcifications in the upper abdomen, posterior to the stomach. No mass lesion is noted. She is most likely to have a mutation involving which of the following genes?

A. CFTR  
B. KRAS  
C. PRSS1  
D. SPINK1  
E. VHL

12 A 65-year-old woman has had upper abdominal pain for the past month. On examination, the pain is localized to the epigastric region on palpation. Abdominal CT scan shows a well-circumscribed, 8-cm mass in the tail of the pancreas that has many small fluid-filled areas. At laparotomy, the mass is removed and on microscopic examination shows glycogen-rich, low cuboidal cells surrounding spaces filled with clear fluid. There is no recurrence of the lesion. What is the most likely diagnosis?

A. Adenocarcinoma  
B. Autosomal dominant polycystic kidney disease  
C. Chronic pancreatitis  
D. Cystic fibrosis  
E. Pseudocyst  
F. Serous cystadenoma

13 A 46-year-old woman has severe abdominal pain for 2 days. On physical examination she has marked epigastric pain, and bowel sounds are reduced. Laboratory studies show an elevated serum lipase. With supportive care, her acute condition subsides within 7 days. Which of the following complications is most likely to occur in this patient?

A. Gastric ulceration  
B. Hemoperitoneum  
C. Hyperosmolar coma  
D. Ketoacidosis  
E. Pseudocyst formation  
F. Small bowel infarction

14 A 58-year-old woman has experienced abdominal discomfort for the past year. On physical examination, she has midepigastric tenderness on palpation. Laboratory studies show a normal serum lipase. The figure shows her abdominal CT scan. The lesion is excised, and on microscopic examination has cells that show cytologic and architectural atypia, but no invasion of surrounding pancreatic parenchyma. What is the most likely diagnosis?

A. Adenocarcinoma  
B. Autosomal dominant polycystic kidney disease  
C. Mucinous cystic neoplasm  
D. Cystic fibrosis  
E. Islet cell tumor, nonfunctional  
F. Pseudocyst

15 A 79-year-old woman belongs to a religious sect that follows the dictum “if it sounds like fun, you shouldn’t do it,” and has in so (not) doing avoided many risk factors for disease and remained healthy. For the past 7 months, she has had vague abdominal pain, however, and yesterday she experienced acute chest pain with dyspnea. Chest and abdomen CT scans show a pulmonary embolus. Her abdominal CT scan is shown in the figure. Laboratory studies show elevation in CEA and CA19-9. A mutation involving which of the following genes is most likely implicated in development of this mass?

A. APC  
B. BRCA2  
C. CFTR  
D. KRAS  
E. PRSS1  
F. SPINK1
A 73-year-old woman has noticed a 10-kg weight loss in the past 3 months. She is becoming increasingly icteric and has constant vague epigastric pain, nausea, and episodes of bloating and diarrhea. On physical examination, she is afebrile. There is mild tenderness to palpation in the upper abdomen, but bowel sounds are present. Her stool is negative for occult blood. Laboratory findings include a total serum bilirubin concentration of 11.6 mg/dL and a direct bilirubin level of 10.5 mg/dL. Which of the following conditions involving the pancreas is most likely to be present?

- A Adenocarcinoma
- B Chronic pancreatitis
- C Cystic fibrosis
- D Islet cell adenoma
- E Pseudocyst

A 68-year-old man notes dull but constant abdominal pain accompanied by nausea with vomiting for the past 8 months. On physical examination he has tenderness to palpation of the upper abdomen. Abdominal CT imaging shows a 2-cm circumscribed mass in the body of the pancreas. Pancreatocystectomy is performed and microscopic examination of this mass shows tall, columnar, mucinous epithelial cells lining papillary excrescences within the main pancreatic duct. There is minimal atypia and no invasion. What is the most likely future course for this man?

- A Additional gastrointestinal neoplasms
- B Complete remission
- C Development of extraabdominal malignancies
- D Local recurrence
- E Pseudocyst formation

An 80-year-old man has increasing jaundice with abdominal pain for the past 2 weeks. He has lost 4 kg over the past 5 months. On physical examination there is tenderness with palpable gallbladder in the right upper quadrant. An abdominal CT scan shows gallbladder and common bile duct dilation, along with a 3-cm mass in the head of the pancreas. Which of the following lesions is the most likely precursor to this mass?

- A Colonic neuroendocrine carcinoma
- B Duodenal adenocarcinoma
- C Neuroendocrine tumor
- D Pancreatic intraepithelial neoplasia
- E Serous cystadenoma

**ANdWERS**

**1 B** When the embryologic ventral and dorsal pancreatic buds do not rotate and fuse properly, the duodenum can become encircled by pancreatic tissue, producing an obstruction called *annular pancreas*. Depending on the degree of obstruction, this condition may manifest at birth, in childhood, or in adulthood. Polyhydramnios should suggest upper gastrointestinal obstruction in utero. The radiograph shows a region of duodenal narrowing from the surrounding pancreas. An ultrasound scan may show the double bubble sign with gastric and duodenal bulb distention with air proximal to a region of duodenal obstruction. Acute and chronic pancreatitis should be accompanied by abdominal pain and by an elevated serum amylase; they do not generally cause mechanical bowel obstruction, although the inflammation may cause paralytic ileus. Neoplasms are focal lesions that may obstruct the biliary tree and pancreatic duct, but generally do not obstruct the duodenum. Pyloric stenosis is seen in neonates and early infants, but generally at 2 to 3 weeks after birth, with projectile vomiting and a palpable epigastric mass.

**2 E** This incidental finding is consistent with ectopic pancreas, which can be found in the bowel in 2% of individuals. It may be found within a Meckel diverticulum, although in this case no diverticulum was found, and such a diverticulum would be found in the distal ileum as a vitelline duct remnant. Adenocarcinoma is unlikely in a boy in his teens, and primary adenocarcinoma of the small intestine is rare. Adrenal cortical rests may be found in the pelvis, but usually not bowel. Ectopic gastric mucosa is most often found within a Meckel diverticulum. Non-Hodgkin lymphomas of the bowel produce larger masses and are usually seen in older individuals.

**3 C** This is an *incidentaloma*, or finding with no clinical significance, other than recognizing that it should be left alone. More diagnostic testing yields more findings that are not significant to the health of the patient. Follow the dictum in fishing: catch and release. Polycystic disease is marked by multiple cysts, which are also benign, but have a tendency to hemorrhage or become infected. Without additional cellular proliferative features, this is not likely to be a neoplasm. A pseudocyst is a complication of pancreatitis and is lined by granulation tissue.

**4 C** Acute pancreatitis is marked by release of enzymes that produce chalky areas of fat necrosis, with precipitation of calcium that lowers serum calcium. The worse the inflammation, the lower the calcium level. Hypoalbuminemia is more likely to occur as a complication of chronic pancreatitis with malabsorption. Hyperbilirubinemia can occur in cases of gallstone-induced pancreatitis, but the level of bilirubin does not give an indication of the severity of pancreatitis. Fibrinogen is an acute phase reactant that increases with inflammation, but does not correlate well with location or extent of inflammation. The serum haptoglobin decreases with hemolysis.

**5 E** This form of hereditary autosomal recessive pancreatitis results from abnormal activation of trypsin, causing a cascade of events leading to acute pancreatitis. The *serine protease inhibitor Kazal type 1 (SPINK1)* gene codes for a pancreatic secretory trypsin inhibitor. Activation of trypsin also leads to activation of Hageman factor, kallikrein, and complement that promote the vascular abnormalities in pancreatitis. Both
SPINK1 and PRSS1 mutations carry a greatly increased risk for pancreatic cancer. Both amylase and lipase are released in pancreatitis as a consequence of trypsin activation. TGF-β plays a role in the fibrogenesis of chronic pancreatitis.

6 C The figure shows acute hemorrhagic pancreatitis with foci of chalky white fat necrosis. Fundamental to the causation of acute pancreatitis is inappropriate activation of digestive enzymes in the acini and the consequent autodigestion of the pancreas. These enzymes are present as proenzymes in the acini and are activated by trypsin. Trypsin itself is derived from trypsinogen, and any abnormality that prevents regulated inactivation of trypsinogen can lead to excessive trypsin-mediated activation of other digestive enzymes, such as lipase, amylase, and elastase. Evidence for this mechanism comes from the observation that the rare disease hereditary pancreatitis, with germline mutations that affect a site on the cationic trypsinogen molecule essential for the cleavage of trypsin by trypsin itself, results in trypsinogen and trypsin that become resistant to inactivation, and the abnormally active trypsin activates other digestive proenzymes, leading to development of pancreatitis. Pancreatic duct obstruction by gallstones is the most common event precipitating trypsinogen activation. A CFTR gene mutation can give rise to chronic pancreatitis, even in the absence of cystic fibrosis. Viral infections, hypertriglyceridemia, vasculitis, and trauma are less common causes for pancreatitis.

7 C This boy has evidence for recurrent pancreatitis, marked by severe abdominal pain, paralytic ileus, and hyperamylasemia. An onset of pancreatitis in children suggests a genetic basis, and the history of an involved parent and grandparent suggests an autosomal dominant mode of transmission. The PRSS1 gene encodes for cationic trypsinogen, which, when mutated, leads to resistance of trypsin to inactivation. The SPINK1 gene encodes for a trypsin inhibitor, and mutation can lead to pancreatitis, but the inheritance is autosomal recessive. A mutation in the CFTR gene can lead to cystic fibrosis, but there should be pulmonary problems by age 11 years, and bouts of pancreatitis are usually not severe. KRAS (an oncogene) and SMAD4 (a tumor suppressor gene) are involved with development of pancreatic adenocarcinoma.

8 F The clinical features and the preexisting gallbladder disease are highly suggestive of acute pancreatitis. This is confirmed by the appearance of the pancreas on radiologic imaging. The serum amylase as well as serum and urine lipase levels are rapidly elevated after an attack of acute pancreatitis. These enzymes are released from necrotic pancreatic acini. Abnormalities in liver function test results may be seen in cases of gallstone pancreatitis but do not confirm pancreatic involvement. An increased ALT level is more characteristic of liver cell injury. Hyperammonemia is a feature of liver failure. Although the pancreatic acinar cells can produce bicarbonate, levels of this analyte do not correlate well with pancreatitis. Bilirubin may be elevated with gallstone passage down the biliary tree, but does not confirm pancreatic disease. In acute pancreatitis, the islets of Langerhans still function, but typically do not become hyperactive.

9 D This patient has chronic pancreatitis. Alcohol promotes intracellular proenzyme activation that leads to acinar cell injury. Although the exact mechanism for development of pancreatitis with alcoholism is unknown, the production of a protein-rich pancreatic secretion that forms inspissated plugs that cause ductal obstruction is observed in many cases. Ductal obstruction predisposes to acinar injury, and the ongoing or repeated injury leads to chronic pancreatitis. Patients with a history of alcohol abuse may have bouts of chronic pancreatitis that go unnoticed, only to have a superimposed case of clinically apparent acute pancreatitis. Alcohol, drugs, traumatic injury, and viral agents may have a direct injurious effect on pancreatic exocrine acinar cells. A dilated cardiomyopathy also can occur in chronic alcoholism, as in this case. A deficiency of α1-antitrypsin can produce liver disease with chronic hepatitis or cirrhosis, or both. Blunt abdominal trauma can produce hemorrhage, sometimes with a component of acute pancreatitis. Cystic fibrosis decreases bicarbonate excretion and promotes protein plugging of ducts, but the appearance is more of a subclinical chronic pancreatitis, with fat and fat-soluble vitamin malabsorption, and depending upon the CFTR mutation, other features of cystic fibrosis may be lacking in these patients. Hypercholesterolemia (typically at least 1000 mg/dL or more) can cause acute pancreatitis, as can cholelithiasis.

10 B Cystic fibrosis is an autosomal recessive condition that results from an abnormal cystic fibrosis transmembrane conductance regulator (CFTR) gene. The most common mutation is ΔF508. There are abnormal viscid secretions affecting the pancreas. This results in ductal obstruction, and this leads to a form of chronic pancreatitis with acinar atrophy. Eventually, the exocrine function is gone. Clinically symptomatic, florid acute and chronic pancreatitis and the complication of inflammation known as a pseudocyst do not typically occur in this setting. Cystic fibrosis is not a risk factor for adenoma or carcinoma of the pancreas. Amyloid deposition may be seen in the islets of Langerhans in a patient with type 2 diabetes mellitus, but generalized amyloid deposition of the pancreas is rare.

11 A She has chronic pancreatitis. One could reflexively blame this on alcohol abuse, but the real cause can be obscure, and many cases of pancreatitis have been termed idiopathic. Genetic testing may reveal a mutation in the CFTR gene. There are hundreds of such mutations, and not all of them are associated with pulmonary disease, but may have other clinical manifestations, such as pancreatitis. KRAS mutations are found with adenocarcinomas. PRSS1 and SPINK1 mutations are associated with hereditary pancreatitis as well as increased risk for pancreatic carcinoma, but neither is associated with lung disease. VHL gene mutations may be associated with serous cystadenoma.
Most pancreatic cysts are nonneoplastic, such as pseudocysts and the scattered cysts associated with autosomal dominant polycystic kidney disease (ADPKD). It is unlikely that ADPKD would involve pancreas in the absence of very large multicystic kidneys. This woman has a solitary mass with features of a benign serous cystadenoma. Adenocarcinomas are more aggressive neoplasms that are rarely cured by surgery because they are large and infiltrative at the time of diagnosis; they tend to be solid masses, although there can be areas of necrosis. Chronic pancreatitis leads to fibrosis. Cystic fibrosis can lead to chronic pancreatitis. The pancreas is usually small and fibrotic with chronic pancreatitis. A pseudocyst usually forms in the setting of extensive pancreatic destruction with acute inflammation, typically acute pancreatitis, or acute inflammation superimposed on chronic inflammation with chronic alcoholism.

During acute pancreatitis, the extent of necrosis may be so severe that a liquefied area becomes surrounded by granulation tissue, forming a cystic mass. Because there is no epithelial lining in the cyst, however, it is best called a pseudocyst. Although the pancreas is inferior and posterior to the stomach, spread of inflammation to the stomach does not typically occur. Although acute pancreatitis may be hemorrhagic, the hemorrhage is confined to the body of the pancreas and surrounding fibroadipose tissue. The islets of Langerhans usually continue to function despite marked inflammation of the parenchyma. Lack of insulin is not a typical feature of pancreatitis. The inflammation is unlikely to compromise the blood supply to abdominal organs and produce an infarction.

The CT scan shows a circumscribed, multilocular, cystic mass. Mucinous tumors of the pancreas can be completely benign, borderline, or malignant. The presence of cytologic and architectural atypia (low-grade dysplasia) indicates the lesion is not completely benign, but in the absence of clearly malignant features of invasion or metastasis, the lesion fits in the borderline category. Such “in between” terminology frustrates both health care providers and their patients, but there are often no simple answers when dealing with neoplasms. However, over time an adenocarcinoma can arise within a mucinous cystic neoplasm. Pancreatic adenocarcinomas are more aggressive solid neoplasms that have an overall poor prognosis. The cysts of ADPKD are benign and scattered throughout the pancreas without forming a mass effect; the kidneys in this CT scan are not cystic. Cystic fibrosis can lead to chronic pancreatitis, not neoplasia of the pancreas. Islet cell tumors are solid masses that are usually quite small when benign and sometimes associated with hormone production. A pseudocyst is an inflammatory structure arising in the setting of pancreatitis with extensive necrosis; it is nonneoplastic.

Mortality happens regardless. Prevalence for sporadic cancer (no identifiable inherited risks) increases with age. The figure shows an irregular mass occupying most of the body and tail of the pancreas, along with hepatic metastases. She has a pancreatic adenocarcinoma with Trousseau syndrome (a paraneoplastic hypercoagulable state) leading to pulmonary thromboembolism. The tumor markers CEA and CA19-9 are often present, but insensitive for early diagnosis of pancreatic cancer. KRAS mutations are found in more than 90% of pancreatic adenocarcinomas. APC gene mutations are associated with hereditary and sporadic forms of colon adenocarcinoma. BRCA2 mutations are found in some pancreatic cancers, but usually there is a history of familial breast and ovarian cancers. CFTR mutations are associated with cystic fibrosis, which is not a risk for pancreatic cancer. PRSSI mutations carry a risk for pancreatic adenocarcinoma, but there should be a history of pancreatitis starting early in life. SPINK1 mutations are also associated with hereditary pancreatitis, and a significant risk for pancreatic cancer.

The weight loss and pain suggest a malignant neoplasm. The jaundice (a conjugated hyperbilirubinemia) occurs because of biliary tract obstruction by a mass in the head of the pancreas. Such a carcinoma may manifest with “painless jaundice” as well, but it is more likely to invade the nerves around the pancreas, causing pain. Islet cell adenoma is not as common as pancreatic carcinoma. An adenoma located near the ampulla could have an effect similar to that of carcinoma; however, weight loss with adenoma is unlikely. Chronic pancreatitis usually does not obstruct the biliary tract. In cystic fibrosis, there is progressive pancreatic acinar atrophy without a mass effect. Most pseudocysts from pancreatitis are in the region of the body or tail of the pancreas, not the head, and they are nonneoplastic.

He has an intraductal papillary mucinous neoplasm (IPMN) of the pancreas. If they are discovered early and when small, they can be noninvasive and curable with surgery, and without recurrence. Over time, they may progress to contain more severe dysplasia and give rise to adenocarcinoma. They are not likely to be part of a syndrome with development of additional neoplasms. If the pancreas is resected, there is nothing left to form a pseudocyst.

He is most likely to have pancreatic adenocarcinoma. Diagnosis at the stage of pancreatic intraepithelial neoplasia (PanIN) is nearly impossible. The dilated palpable gallbladder represents Courvoisier sign from obstruction of the biliary tree by a neoplasm. Despite this sign, the cancer has had time to invade and is unlikely to be cured, so the prognosis of pancreatic adenocarcinoma is poor. However, there are chemotherapy agents and biologic monoclonal therapies that may slow the cancer and prolong survival. This solitary mass is unlikely to be a metastasis. Primary carcinomas arising in the small intestine are rare. Neuroendocrine tumors that may arise from islets of Langerhans are often benign when small, and they do not transform to adenocarcinomas. Serous cystadenomas of the pancreas are nearly all benign, and likely to be cystic.
1. A 71-year-old man has had decreased urine output <500 mL per day for the past 3 days. Physical examination shows vital signs with temperature 37°C, pulse 88/min, respiratory rate 18/min, and blood pressure 85/60 mm Hg. He has peripheral edema and diffuse rales on auscultation of the chest. Urinalysis shows specific gravity 1.019 and no protein, blood, glucose, ketones, WBCs, RBCs, or casts. His serum creatinine is 3.3 mg/dL, and urea nitrogen is 62 mg/dL. The fractional excretion of sodium (FENa) is <1%. Which of the following underlying conditions is he most likely to have?

A. Dilated cardiomyopathy
B. Membranous nephropathy
C. Prostatic hyperplasia
D. Systemic lupus erythematosus
E. Urothelial carcinoma

2. A 36-year-old woman has had increased malaise for 3 weeks and urine output <500 mL/day for the past 4 days. On examination, she has blood pressure 170/112 mm Hg and peripheral edema. Urinalysis shows protein 1+ and blood 3+, but no glucose or ketones. Urine microscopic analysis shows RBCs and RBC casts. Her serum urea nitrogen is 39 mg/dL, and creatinine is 4.3 mg/dL. Her serum complement C1q, C3, and C4 are decreased. A renal biopsy is performed, and immunofluorescence microscopy shows a granular pattern of staining with antibody to C3. Which of the following types of hypersensitivity reactions is most likely causing her renal disease?

A. I (IgE-mediated systemic anaphylaxis)
B. II (Antibody-dependent cell-mediated cytotoxicity)
C. III (Immune complex formation)
D. IV (Delayed-type hypersensitivity)

3. A 29-year-old man with chronic hepatitis C virus infection has noted dark urine for the past 2 weeks. On examination he is hypertensive but afebrile. Laboratory studies show serum creatinine of 3.8 mg/dL and urea nitrogen of 35 mg/dL. Cryoglobulins are detected. Urinalysis shows RBCs and RBC casts. A renal biopsy is performed and microscopically shows hypercellular glomeruli with lobulation and a double-contour appearance to split basement membranes adjacent to subendothelial immune complexes. Which of the following cell types has most likely proliferated in his glomeruli?

A. Juxtaglomerular cells
B. Mesangial cells
C. Parietal epithelial cells
D. Podocytes
E. Endothelial cells

4. A study of renal disease identifies patients with greater than 3.5 g of protein in a 24-hour urine collection, but no RBCs or WBCs. Dysfunction involving which of the following cells is most likely to be responsible for proteinuria?

A. Endothelium
B. Macula densa
C. Mesangium
D. Parietal epithelium
E. Podocytes

5. A 7-year-old boy is recovering from impetigo. Physical examination shows five honey-colored crusts on his face. The crusts are removed, and a culture of the lesions grows *Streptococcus pyogenes*. He is treated with antibiotics. One week later, he develops malaise with nausea and a slight fever and passes dark brown urine. Laboratory studies show a serum anti-streptolysin O titer of 1:1024. Which of the following is the most likely outcome of his renal disease?

A. Chronic renal failure
B. Complete recovery
C. Crescentic glomerulonephritis
D. Rheumatic heart disease
E. Streptococcal urinary tract infection
6 A 17-year-old girl living in the Congo has had a chronic febrile illness for 2 years. In the past 2 days she notes her urine is smoky brown. On physical examination her blood pressure is 145/95 mm Hg. Laboratory studies show her serum creatinine is 3.7 mg/dL, and urea nitrogen is 35 mg/dL. Urinalysis shows 4+ blood with 1+ protein, but no glucose, ketones, or leukocytes. The serum haptoglobin is decreased and Coombs test is negative. Her Hgb is 8.5 g/dL. A peripheral blood smear shows rare ring stage trophozoites. Immunofluorescence microscopy performed on renal biopsy shows granular deposition of IgG and C3 in glomerular capillary basement membranes. Electron microscopy shows electron-dense subepithelial “humps.” Which of the following renal diseases is she most likely to have?

A  Focal segmental glomerulosclerosis  
B  Hereditary nephritis  
C  IgA nephropathy  
D  Lupus nephritis  
E  Membranous nephropathy  
F  Acute proliferative glomerulonephritis

7 A 45-year-old man has experienced increasing malaise, nausea, and reduced urine output for the past 3 days. On physical examination, he is afebrile and normotensive. Laboratory findings show a serum creatinine level of 7.5 mg/dL. Urinalysis shows hematuria, but no pyuria or glucosuria. A renal biopsy is done; the light microscopic picture is shown in the figure. Which of the following additional studies is most useful for classification and treatment of this disease?

A  Antinuclear antibody  
B  Anti-glomerular basement membrane antibody  
C  HIV-1 RNA copy level  
D  Quantitative serum immunoglobulins  
E  Rheumatoid factor  
F  Urine immunoelectrophoresis

8 A 21-year-old previously healthy man has noticed blood in his urine for the past 2 days. He reports no dysuria, frequency, or hesitancy of urination. On physical examination, there are no abnormal findings. Laboratory findings show a serum urea nitrogen level of 39 mg/dL and creatinine level of 4.1 mg/dL. A renal biopsy specimen is obtained; the immunofluorescence pattern of staining with antibody against human IgG is shown in the figure. Which of the following serologic findings is most likely to be present in this patient?

A  Anti–glomerular basement membrane antibody  
B  Anti–streptolysin O antibody  
C  C3 nephritic factor  
D  Hepatitis B surface antibody  
E  HIV antibody

9 A 46-year-old woman has had worsening malaise for the past 36 hours. Her urine output is markedly diminished, and it has a cloudy brown appearance. On examination she has periorbital edema. Laboratory findings include serum creatinine of 2.8 mg/dL and urea nitrogen of 30 mg/dL. A renal biopsy is performed and on microscopic examination shows focal necrosis in glomeruli with glomerular basement membrane breaks and crescent formation. No immune deposits are identified with immunofluorescence. Which of the following autoantibodies is most likely detectable in her serum?

A  Anti–DNA topoisomerase antibody  
B  Anti–glomerular basement membrane antibody  
C  Anti–neutrophil cytoplasmic autoantibody  
D  Antinuclear antibody  
E  Anti-HBs Ag
10. A 44-year-old woman has developed a fever, nonproductive cough, and decreased urine output over the past 3 days. On physical examination, her temperature is 37.7°C, and blood pressure is 145/95 mm Hg. She has sinusitis. On auscultation, crackles are heard over all lung fields. A chest radiograph shows bilateral patchy infiltrates and nodules. The serum creatinine level is 4.1 mg/dL, and the urea nitrogen level is 43 mg/dL. The results of serologic testing are negative for ANA, but positive for C-ANCA. A renal biopsy specimen shows glomerular crescents and damage to small arteries. The result of immunofluorescence staining with anti-IgG and anti-C3 antibodies is negative. Which of the following additional microscopic findings is most likely to be seen in this biopsy?

A. Focal segmental glomerulosclerosis
B. Glomerular basement membrane thickening
C. Hyperplastic arteriolosclerosis
D. Infiltrations by neutrophils
E. Mesangial proliferation
F. Necrotizing granulomatous vasculitis

11. A 48-year-old man has had increased swelling in the extremities for 2 months. Physical examination showed generalized edema. A 24-hour urine collection yielded 4.1 g of protein (albumin and globulins). He did not respond to a course of corticosteroid therapy. A renal biopsy was done, and microscopic examination showed diffuse thickening of the basement membrane. Immunofluorescence staining with antibody to the C3 component of complement was positive in a granular pattern in the glomerular capillary loops. Two years later, he experiences increasing malaise. Laboratory studies now show serum creatinine level of 4.5 mg/dL and urea nitrogen level of 44 mg/dL. Which of the following immunologic mechanisms was most likely responsible for the glomerular changes observed in the biopsy specimen?

A. Antibodies that react with basement membrane collagen
B. Antibodies against streptococci that cross-react with the basement membrane
C. Cytotoxic T cells directed against renal antigens
D. Deposition of immune complexes on the basement membrane
E. Release of cytokines by inflammatory cells

12. A 7-year-old boy has become less active over the past 10 days. On physical examination, the boy has facial puffiness. Urinalysis shows no blood, glucose, or ketones, and microscopic examination shows no casts or crystals. The serum creatinine level is normal. A 24-hour urine collection yields 3.8 g of protein. He improves after corticosteroid therapy. He has two more episodes of proteinuria over the next 4 years, both of which respond to corticosteroid therapy. What is the most likely mechanism causing his disease?

A. Cytokine-mediated visceral epithelial cell injury
B. Cytotoxic T cell-mediated tubular epithelial cell injury
C. IgA-mediated mesangial cell injury
D. Immune complex-mediated glomerular injury
E. Verocytotoxin-induced endothelial cell injury

13. A 44-year-old man with increasing malaise for the past month now is bothered by increasing swelling in the hands and legs. On physical examination, there is generalized edema. He is afebrile, and his blood pressure is 140/90 mm Hg. Urinalysis shows a pH of 6.5; specific gravity 1.017; 4+ proteinuria; and no blood, glucose, or ketones. Microscopic examination of the urine shows no casts or RBCs and 2 WBCs per high-power field. The 24-hour urine protein level is 4.2 g. A renal biopsy specimen is obtained, and immunofluorescence staining with antibody to the C3 component of complement produces the pattern shown in the figure. Which of the following underlying disease processes is most likely to be present in this man?

A. Chronic hepatitis B virus infection
B. HIV infection
C. Multiple myeloma
D. Recurrent urinary tract infection
E. Nephrolithiasis

14. A 6-year-old girl has become increasingly lethargic over the past 2 weeks. On examination she has puffiness around the eyes. Her temperature is 36.9°C, and blood pressure is 100/60 mm Hg. Laboratory findings show serum creatinine, 0.7 mg/dL; urea nitrogen, 12 mg/dL; and cholesterol, 217 mg/dL. Urinalysis shows pH, 6.5; specific gravity, 1.011; 4+ proteinuria; and no blood or glucose. The 24-hour urine protein level is 3.8 g. The child’s condition improves after glucocorticoid therapy. Which of the following findings by electron microscopy is most likely to characterize this disease process?

A. Areas of thickened and thinned basement membrane
B. Effacement of podocyte foot processes
C. Increased mesangial matrix
D. Reduplication of glomerular basement membrane
E. Subepithelial electron-dense humps
A 12-year-old girl has experienced increasing malaise for the past 2 weeks. On physical examination, she has periorbital edema. The child is afebrile. Laboratory findings show proteinuria on dipstick urinalysis, but no hematuria or glucosuria. Microscopic examination of the urine shows numerous oval fat bodies. The serum creatinine level is 2.3 mg/dL. She receives a course of corticosteroid therapy, but does not improve. A renal biopsy is performed and microscopic examination shows that approximately 50% of the glomeruli in the specimen are affected by the lesion shown in the figure. What is the most likely diagnosis?

A. Dense deposit disease  
B. Focal segmental glomerulosclerosis  
C. Minimal change disease  
D. Nodular glomerulosclerosis  
E. Postinfectious glomerulonephritis  
F. Rapidly progressive glomerulonephritis

A 46-year-old Hispanic man has had increasing malaise with headaches and easy fatigability for the past 3 months. Physical examination reveals his blood pressure is 200/100 mm Hg. There are no palpable abdominal masses and no costovertebral tenderness. Laboratory studies show hemoglobin, 9.5 g/dL; hematocrit, 28.3%; MCV, 92 μm³; creatinine, 4.5 mg/dL; and urea nitrogen, 42 mg/dL. Urinalysis reveals 3+ hematuria and 3+ proteinuria, but no glucose or leukocytes. A renal biopsy is done; light microscopic examination of the biopsy specimen shows that approximately 50% of the glomeruli appear normal, but the rest show that a portion of the capillary tuft is sclerotic. Immunofluorescence staining shows IgM and C3 deposition in these sclerotic areas. His history is significant for repeated episodes of passing dark brown urine, which failed to respond to corticosteroid therapy. Which of the following mechanisms is most likely responsible for his disease?

A. Attachment of anti-glomerular basement membrane antibodies  
B. Deposition of immune complexes containing microbial antigens  
C. Deposition of C3 nephritic factor (C3NeF)  
D. Dysfunction of the podocyte slit diaphragm apparatus  
E. Inherited defect in the basement membrane collagen

A 38-year-old woman has been feeling lethargic for 4 months. On physical examination, she is afebrile, and her blood pressure is 140/90 mm Hg. Laboratory findings show the serum creatinine level is 5.8 mg/dL. C3 nephritic factor is present in serum, resulting in hypocomplementemia, and the ANA test result is negative. Urinalysis shows 2+ blood and 1+ protein. A renal biopsy is done; microscopic examination shows hypercellular glomeruli and prominent ribbonlike deposits along the lamina densa of the glomerular basement membrane. Which of the following forms of glomerulonephritis is most likely to be present in this patient?

A. Chronic glomerulonephritis  
B. Dense deposit disease  
C. Membranous nephropathy  
D. Postinfectious glomerulonephritis  
E. Rapidly progressive glomerulonephritis

A 25-year-old man has a 5-year history of celiac sprue. Four days after a mild upper respiratory infection, he begins passing dark red-brown urine. The dark urine persists for the next 3 days and then becomes clear and yellow, only to become red-brown again 1 month later. There are no remarkable findings on physical examination. Urinalysis shows a pH of 6.5; specific gravity, 1.018; 3+ hematuria; 1+ proteinuria; and no glucose or ketones. Microscopic examination of the urine shows RBCs, but no WBCs, casts, or crystals. A 24-hour urine protein level is 200 mg. A renal biopsy specimen from the glomeruli of this patient is most likely to show which of the following alterations?

A. Diffuse cellular proliferation and basement membrane thickening  
B. Granular staining of the basement membrane by anti-IgG antibodies  
C. Mesangial IgA staining by immunofluorescence  
D. Subepithelial electron-dense deposits  
E. Thrombosis within the glomerular capillaries

One week after a mild flulike illness, a 9-year-old boy has an episode of hematuria that subsides within 2 days. One month later, he tells his parents that his urine is red again. On physical examination, there are no significant findings. Urinalysis shows a pH of 7; specific gravity, 1.015; 1+ hematuria; 1+ proteinuria; and no ketones, glucose, or urobilinogen. The serum urea nitrogen level is 36 mg/dL, and the creatinine level is 3.2 mg/dL. Serum electrophoresis shows increased IgA1. Which of the following glomerular structures is most likely to show structural alterations in this boy?

A. Basement membranes  
B. Capillaries  
C. Mesangium  
D. Parietal epithelium  
E. Podocytes
20 A 15-year-old boy has been passing dark-colored urine for the past month. On physical examination, he has bilateral sensorineural hearing loss and corneal erosions. Urinalysis shows a pH of 6.5; specific gravity, 1.015; 1+ hematuria; 1+ proteinuria; and no ketones, glucose, or leukocytes. The serum creatinine level is 2.5 mg/dL, and the urea nitrogen level is 24 mg/dL. A renal biopsy specimen shows tubular epithelial foam cells by light microscopy. By electron microscopy, the glomerular basement membrane shows areas of attenuation, with splitting and lamination of lamina densa in other thickened areas. What is the most likely diagnosis?

A. Mutation in a gene encoding type IV collagen  
B. Increased synthesis of abnormal IgA  
C. Autoimmune destruction of pancreatic beta cells  
D. Acquired deficiency of ADAMTS13 metalloprotease  
E. Toxic injury to slit diaphragm proteins

21 A 56-year-old woman is found on health screening to have a blood pressure of 168/109 mm Hg. No other physical examination findings are noted. Urinalysis shows a pH of 7.0; specific gravity, 1.020; 1+ proteinuria; and no blood, glucose, or ketones. The ANA and ANCA test results are negative. The serum urea nitrogen level is 51 mg/dL, and the creatinine level is 4.7 mg/dL. The hemoglobin A1c concentration is within the reference range. An abdominal ultrasound scan shows bilaterally and symmetrically small kidneys with no masses. What is her most likely diagnosis?

A. Amyloidosis  
B. Autosomal dominant polycystic kidney disease  
C. Chronic glomerulonephritis  
D. Microscopic polyangiitis  
E. Nodular glomerulosclerosis

22 An autopsy study is performed involving persons with gross pathologic findings of bilaterally small kidneys (<100 g) that have a coarsely granular surface appearance. Microscopic examination shows sclerotic glomeruli, a fibrotic interstitium, tubular atrophy, arterial thickening, and scattered lymphocytic infiltrates. Which of the following clinical findings was most likely reported in these patients’ medical histories?

A. Hemoptysis  
B. Hypertension  
C. Lens dislocation  
D. Pharyngitis  
E. Skin rash

23 A 33-year-old woman has had fever and increasing fatigue for the past 2 months. Over the past year, she has noticed soreness of her muscles and joints and has had a 4-kg weight loss. On physical examination, her temperature is 38.4° C. She has a palpable spleen tip, bilateral costovertebral angle tenderness, and diastolic cardiac murmur. Laboratory findings show a serum urea nitrogen level of 15 mg/dL. Urinalysis shows 2+ hematuria, and no glucose, protein, or ketones. A blood culture is positive for *Staphylococcus aureus*. Which of the following best describes the likely gross appearance of the kidneys in this patient?

A. Enlarged, and replaced by 1- to 4-cm, fluid-filled cysts  
B. Marked pelvic and calyceal dilation with thinning of the cortices  
C. Normal size, with smooth cortical surfaces  
D. Shrunken, with uniformly finely granular cortical surfaces  
E. Slightly swollen, with scattered petechial hemorrhages  
F. Small and asymmetric, with irregular cortical scars and marked calyceal dilation  
G. Wedge-shaped regions of yellow-white cortical necrosis

24 A 33-year-old woman with a history of intravenous drug use comes to the emergency department because she has had a high fever for the past 2 days. On physical examination, her temperature is 38.4° C. She has a palpable spleen tip, bilateral costovertebral angle tenderness, and diastolic cardiac murmur. Laboratory findings show a serum urea nitrogen level of 15 mg/dL. Urinalysis shows 2+ hematuria, and no glucose, protein, or ketones. A blood culture is positive for *Staphylococcus aureus*. Which of the following best describes the likely gross appearance of the kidneys in this patient?

A. Enlarged, and replaced by 1- to 4-cm, fluid-filled cysts  
B. Marked pelvic and calyceal dilation with thinning of the cortices  
C. Normal size, with smooth cortical surfaces  
D. Shrunken, with uniformly finely granular cortical surfaces  
E. Slightly swollen, with scattered petechial hemorrhages  
F. Small and asymmetric, with irregular cortical scars and marked calyceal dilation  
G. Wedge-shaped regions of yellow-white cortical necrosis

25 A 55-year-old woman with poorly controlled hyperglycemia for many years now has had burning pain on urination for the past 3 days. Physical examination shows a 2-cm ulceration on the skin of the heel and reduced sensation in the lower extremities. Her visual acuity is 20/100 bilaterally. Urinalysis shows 1+ proteinuria; 2+ glucosuria; and no blood, ketones, or urobilinogen. A urine culture contains more than 100,000 colony-forming units/mL of Klebsiella pneumoniae. Which of the following pathologic findings is most likely to be present in both her kidneys?

A. Deposits of IgG and C3 in the glomerular basement membrane  
B. Effacement of podocyte foot processes  
C. Formation of glomerular crescents  
D. Mesangial deposits of IgA  
E. Necrotizing granulomatous vasculitis  
F. Nodular hyaline mesangial deposits
A 58-year-old man is found to have mild hypertension. Laboratory findings show a serum creatinine level of 2.2 mg/dL and urea nitrogen level of 25 mg/dL. Microalbuminuria is present, with excretion of 250 mg/day of albumin. Two years later, he remains hypertensive and has a serum creatinine level of 3.8 mg/dL, urea nitrogen level of 38 mg/dL, and 24-hour urine protein level of 2.8 g. A renal biopsy is done; the light microscopic appearance of a PAS-stained specimen is shown in the figure. Blood/serum test for which of the following is most likely to be abnormal in this patient?

A. Anti–glomerular basement membrane antibody
B. Antinuclear antibody
C. Anti–neutrophil cytoplasmic autoantibody
D. Anti–streptolysin O
E. C3 nephritic factor
F. Hemoglobin A1c
G. Hepatitis B surface antigen

A 17-year-old boy is involved in a motor vehicle accident in which he sustains severe blunt trauma to the extremities and abdomen. Over the next 3 days, he develops oliguria and dark brown urine. The urine dipstick analysis is positive for myoglobin and for blood, but microscopic examination of the urine shows no RBCs. His serum urea nitrogen level increases to 38 mg/dL, and 24-hour urine protein level of 2.8 g. A renal biopsy is done; the light microscopic appearance of a PAS-stained specimen is shown in the figure. Blood/serum test for which of the following is most likely to be abnormal in this patient?

A. Anti–glomerular basement membrane antibody
B. Antinuclear antibody
C. Anti–neutrophil cytoplasmic autoantibody
D. Anti–streptolysin O
E. C3 nephritic factor
F. Hemoglobin A1c
G. Hepatitis B surface antigen

A 63-year-old man is in stable condition after an acute myocardial infarction when he became hypotensive for 3 hours before paramedical personnel arrived. Over the next week, the serum urea nitrogen level increases to 48 mg/dL, the serum creatinine level increases to 5 mg/dL, and the urine output decreases. He undergoes hemodialysis for the next 2 weeks and then develops marked polyuria, with urine output of 2 to 3 L/day. His renal function gradually returns to normal. Release of which of the following substances most likely participated in the elevation of BUN, creatinine, and reduced urinary output?

A. Aldosterone
B. Endothelin
C. Erythropoietin
D. Natriuretic peptide
E. Vasopressin

A 19-year-old woman has had several episodes of fever and chills accompanied by right flank pain for the past 3 days. She has had two similar episodes during the past year. On physical examination, her temperature is 38.3° C; her blood pressure is 152/94 mm Hg, and there is right costovertebral angle tenderness. Laboratory findings show a serum glucose level of 77 mg/dL and creatinine level of 1 mg/dL. Urinalysis shows a pH of 6.5; specific gravity, 1.018; and no protein, blood, glucose, or ketones. Microscopic examination of the urine shows many WBCs and WBC casts. Which of the following is the most useful test to obtain on this patient?

A. Antinuclear antibody
B. Plasma renin
C. Renal biopsy
D. Urine culture
E. Abdominal CT scan

A 51-year-old woman has had dysuria and urinary frequency for the past week. On physical examination, her temperature is 38° C, and she has pain on palpation over the left costovertebral angle. Laboratory findings show glucose, 177 mg/dL; hemoglobin A1c, 9.8%; hemoglobin, 13.1 g/dL; platelet count, 232,200/mm³; and WBC count, 11,320/mm³. Urinalysis shows a pH of 6.5; specific gravity, 1.016; 2+ glucosuria; and no blood, protein, or ketones. Microscopic examination of the urine shows numerous neutrophils, and a urine culture is positive for *Escherichia coli*. Which of the following complications is most likely to develop in this patient?

A. Acute tubular necrosis
B. Crescentic glomerulonephritis
C. Hydronephrosis
D. Necrotizing papillitis
E. Renal calculi

A 53-year-old woman has had fever and flank pain for the past 2 days. On physical examination, her temperature is 38.2° C, pulse is 81/min, respirations are 16/min, and blood pressure is 130/80 mm Hg. Urinalysis shows no protein, glucose, or ketones. The leukocyte esterase test is positive. Microscopic examination of the urine shows numerous polymorphonuclear leukocytes and occasional WBC casts. Which of the following organisms is most likely to be found in the urine culture?

A. Cryptococcus neoformans
B. *Escherichia coli*
C. Group A streptococcus
D. *Mycobacterium tuberculosis*
E. *Mycoplasma hominis*
32 A 30-year-old woman has had increasing malaise with nocturia and polyuria for the past year. She has had a high fever for the past 3 days. On physical examination, her blood pressure is 170/95 mm Hg. Urinalysis shows a pH of 7.5; specific gravity, 1.010; 1+ proteinuria; positive leukocyte esterase and nitrite; and no glucose, blood, or ketones. A renal ultrasound scan shows an enlarged right kidney with pelvic and calyceal enlargement and cortical thinning; the left kidney appears normal. A right-sided nephrectomy is performed, and grossly there are large U-shaped scars at the poles with underlying blunted calyces. Microscopic examination shows inflammatory infiltrates extending from the medulla to the cortex, with tubular destruction and extensive interstitial fibrosis. Lymphocytes, plasma cells, and neutrophils are abundant. Which of the following underlying conditions is most likely to produce these findings?

A Polycystic kidney disease  
B Essential hypertension  
C Congestive heart failure  
D Systemic lupus erythematosus  
E Systemic amyloidosis  
F Vesicoureteral reflux

33 A 29-year-old woman has had a fever and sore throat for the past 3 days. On physical examination, her temperature is 38°C. The pharynx is erythematous, with yellowish tonsillar exudate. She is treated with amoxicillin and recovers fully in 7 days. Two weeks later, she develops fever and a rash, and notices a slight decrease in urinary output. Her temperature is 37.7°C, and there is a diffuse erythematous rash on the trunk and extremities. Urinalysis shows a pH of 6; specific gravity, 1.022; 1+ proteinuria; 1+ hematuria; and no glucose or ketones. Microscopic examination of the urine shows RBCs and WBCs, including eosinophils, but no casts or crystals. What is the most likely cause of her disease?

A Deposition of immune complexes with streptococcal antigens  
B Formation of antibodies against glomerular basement membrane  
C Hematogenous dissemination of septic emboli  
D Hypersensitivity reaction to ampicillin  
E Renal tubular cell necrosis caused by bacterial toxins

34 A 32-year-old man developed a fever and rash over 3 days. Five days later, he has increasing malaise. On physical examination, the maculopapular erythematous rash on his trunk has nearly faded away. His temperature is 37.1°C, and blood pressure is 135/85 mm Hg. Laboratory studies show a serum creatinine level of 2.8 mg/dL and blood urea nitrogen level of 29 mg/dL. Urinalysis shows 2+ proteinuria; 1+ hematuria; and no glucose, ketones, or nitrite. The leukocyte esterase result is positive. Microscopic examination of urine shows RBCs and WBCs, some of which are eosinophils. Which of the following most likely precipitated his renal disease?

A Antibiotic ingestion  
B Congestive heart failure  
C Eating poorly cooked ground beef  
D Streptococcal pharyngitis  
E Urinary tract infection

35 A 72-year-old man with chronic arthritis has taken more than 3 g of analgesics per day, including phenacetin, aspirin, and acetaminophen, for the past 20 years. He now has increasing malaise, nausea, and diminished mentation. On physical examination, his blood pressure is 156/92 mm Hg. Laboratory findings show serum urea nitrogen level of 68 mg/dL and creatinine level of 7.1 mg/dL. Which of the following renal diseases is he most likely to develop?

A Acute tubular injury  
B Chronic glomerulonephritis  
C Hydronephrosis  
D Renal cell carcinoma  
E Renal papillary necrosis

36 A 49-year-old man is found on physical examination to have a blood pressure of 160/110 mm Hg, but no other abnormalities. Laboratory studies show serum glucose of 75 mg/dL, creatinine of 1.3 mg/dL, and urea nitrogen of 20 mg/dL. His plasma renin is elevated. CT angiography shows marked stenosis of his renal arteries. He is treated with an angiotensin-converting enzyme inhibitor. A week later, he has a headache for which he takes ibuprofen. Over the next day, his urine output decreases. This effect on urinary output is most likely mediated by the unopposed action of which of the following chemicals?

A Aldosterone  
B Histamine  
C Nitric oxide  
D Prostaglandin  
E Tumor necrosis factor

37 A 28-year-old man is diagnosed with acute myelogenous leukemia (M2), with a total leukocyte count of 45,000/mm³, including 60% blasts. After induction with a multiagent chemotherapy protocol, he has an episode of lower abdominal pain accompanied by passage of red urine. He has no fever, dysuria, or urinary frequency. On physical examination, there are no remarkable findings. Urinalysis shows a pH of 5.5; specific gravity, 1.021; 2+ hematuria; and no protein, ketones, or glucose. There are no remarkable findings on an abdominal radiograph. Which of the following additional urinalysis findings is most likely to be reported for this patient?

A Bence Jones protein  
B Eosinophils  
C Myoglobin  
D Oval fat bodies  
E RBC casts  
F Uric acid crystals

38 A 49-year-old woman had a mastectomy of the right breast 2 years ago to remove a carcinoma. She now has bone pain, and a radionuclide scan shows multiple areas of increased uptake in the vertebrae, ribs, pelvis, and right femur. Urinalysis shows a specific gravity of 1.010, which remains unchanged after water deprivation for 12 hours. She undergoes multiple courses of chemotherapy over the next year. During this time, the serum urea nitrogen level progressively increases. Which of the following abnormal laboratory findings is most likely to be reported for this patient?

A Hypercalcemia  
B Hypercholesterolemia  
C Hypergammaglobulinemia  
D Hyperglycemia  
E Hyperuricemia
39. A 63-year-old man has noted increasing back pain for 7 months. He has had three respiratory tract infections with *Streptococcus pneumoniae* during the past year. On examination, he has pitting edema to his thighs. Laboratory studies show total serum protein, 9.6 g/dL; albumin, 3.5 g/dL; creatinine, 3 mg/dL; urea nitrogen, 28 mg/dL; and glucose, 79 mg/dL. Urinalysis shows proteinuria of 4 g/24 hr, but no glucosuria or hematuria. Abdominal CT scan shows enlarged kidneys without cysts or masses. A renal biopsy specimen stained with H&E shows deposits of amorphous pink material within glomeruli, interstitium, and arteries. Which of the following diseases is he most likely to have?

A. Analgesic nephropathy  
B. ANCA-associated granulomatous vasculitis  
C. Type 2 diabetes mellitus  
D. Membranous nephropathy  
E. Multiple myeloma  
F. Systemic lupus erythematosus

40. A 66-year-old woman has experienced five transient ischemic attacks within a week. On physical examination, the only abnormal finding is a blood pressure of 150/95 mm Hg. Urinalysis shows 1+ proteinuria, and no glucose, blood, or ketones. Microscopic examination of the urine shows no RBCs or WBCs and few oxalate crystals. On abdominal ultrasound, the kidneys are slightly decreased in size. The representative high magnification microscopic appearance of the kidneys is shown in the figure. Which of the following renal lesions is most likely to be present in this patient?

A. Acute tubular necrosis  
B. Fibromuscular dysplasia  
C. Hyaline arteriolosclerosis  
D. Interstitial nephritis  
E. Necrotizing vasculitis

41. A 45-year-old man has had headaches, nausea, and vomiting that have worsened over the past 5 days. He has started “seeing spots” before his eyes and experienced periods of mental confusion. On physical examination, his blood pressure is 270/150 mm Hg. Urinalysis shows 1+ proteinuria; 2+ hematuria; and no glucose, ketones, or leukocytes. The serum urea nitrogen and creatinine levels are elevated. He dies 2 weeks later from a cerebral bleed. Which of the following histologic findings is most likely to be seen in this patient’s kidneys at autopsy?

A. Glomerular crescents  
B. Hyperplastic arteriolosclerosis  
C. Mesangial IgA deposition  
D. Nodular glomerulosclerosis  
E. Segmental tubular necrosis

42. A 5-year-old girl develops cramping abdominal pain and diarrhea 4 days after eating a hamburger, chili, and ice cream at a home barbecue. The next day, she has decreased urine output. On physical examination, there are petechial hemorrhages on the skin. Her temperature is 37° C, pulse is 90/min, respirations are 18/min, and blood pressure is 90/50 mm Hg. A stool sample is positive for occult blood. Laboratory findings show a serum creatinine level of 2.2 mg/dL and urea nitrogen level of 20 mg/dL. CBC shows hemoglobin, 10.8 g/dL; hematocrit, 32.4%; platelet count, 64,300/mm³; and WBC count, 6480/mm³. The peripheral blood smear shows schistocytes, and the serum D-dimer level is elevated. Urinalysis shows a pH of 6; specific gravity, 1.016; 2+ hematuria; and no protein or glucose. A renal biopsy specimen shows small thrombi within glomerular capillary loops. This complication develops most commonly after infection with which of the following organisms?

A. *Candida albicans*  
B. *Clostridium difficile*  
C. *Escherichia coli*  
D. *Proteus mirabilis*  
E. *Staphylococcus aureus*

43. Adult patients with bilateral renal cystic disease are found to have defects in renal tubular function. Their kidneys are markedly increased in size. They have ciliopathy, with reduced mechanosensing of tubular fluid flow. Mutations are found in a gene encoding for a protein component of these cilia. What is this protein most likely to be?

A. Elongin  
B. Fibrocystin  
C. Nephrocystin  
D. Polycystin  
E. Tuberin
A 61-year-old woman has experienced increasing malaise and bouts of abdominal pain for the past 5 years. On physical examination, she has a blood pressure of 150/95 mm Hg. On auscultation, a midsystolic click is heard over the mitral area. Bilateral abdominal masses are palpable. An abdominal ultrasound shows enlarged kidneys with the representative gross appearance shown in the figure. Which of the following extrarenal conditions is she most likely to have?

A  Berry aneurysm  
B  Coagulopathy  
C  Hepatic cirrhosis  
D  Ischemic cardiomyopathy  
E  Pulmonary fibrosis

A 20-year-old primigravid woman is in the third trimester and has felt minimal fetal movement. An ultrasound scan shows bilaterally enlarged echogenic kidneys and a markedly decreased amniotic fluid index. She gives birth to a stillborn male fetus at 33 weeks’ gestation. At autopsy, there are deformations resulting from marked oligohydramnios, including flattening of the facies, varus deformities of the feet, and marked pulmonary hypoplasia. Microscopic examination of the liver shows multiple epithelium-lined cysts and a proliferation of bile ducts. Which of the following is the most likely renal disease in this fetus?

A  Autosomal dominant polycystic kidney disease  
B  Autosomal recessive polycystic kidney disease  
C  Medullary sponge kidney  
D  Multicystic renal dysplasia  
E  Urethral atresia

A 31-year-old woman experiences abdominal pain 1 week after noticing blood in her urine. She has had three episodes of urinary tract infection during the past year. There are no remarkable findings on physical examination. Urinalysis shows 2+ hematuria, 1+ proteinuria, hypercalciuria, and no glucose or ketones. Serum creatinine is 1.0 g/dL. Microscopic examination of the urine shows numerous RBCs and oxalate crystals. An abdominal CT scan with contrast shows linear striations radiating into the renal papillae, along with small cystic collections of contrast material in dilated collecting ducts. She is advised to increase her daily intake of fluids, and her condition improves. Which of the following renal cystic diseases is most likely to be associated with these findings?

A  Autosomal dominant polycystic kidney disease  
B  Autosomal recessive polycystic kidney disease  
C  Medullary sponge kidney  
D  Multicystic renal dysplasia

A 6-year-old child has been drinking more water, with more frequent urination, for the past 7 months. On physical examination dehydration is noted. Urinalysis findings include pH of 6.5; specific gravity, 1.010; and no protein, blood, glucose, or ketones. There are no WBCs, RBCs, or casts. Serum electrolytes show Na+, 152 mmol/L; K+, 4.6 mmol/L; Cl-, 120 mmol/L; HCO3-, 21 mmol/L; urea nitrogen, 29 mg/dL; and creatinine, 3.2 mg/dL. An ultrasound scan shows bilaterally small kidneys with barely visible medullary cysts concentrated at the corticomedullary junction. Which of the following genes is most likely mutated in this child?

A  MCKD1  
B  NPHP1  
C  PKD1  
D  PKHD1

An 83-year-old man has experienced difficulty with urination for the past 15 years. On physical examination, he has a 2-cm nonhealing ulceration on the ball of his right foot. He has decreased sensation to light touch over his lower extremities. The representative gross appearance of his right kidney is shown in the figure. The left kidney has a similar appearance. Which of the following conditions is he most likely to have?

A  Analgesic abuse  
B  Benign nephrosclerosis  
C  Chronic pyelonephritis  
D  Diabetes mellitus  
E  Prostatic adenocarcinoma
A 42-year-old man has had right flank pain for the past 2 days. On physical examination, his temperature is 37.1°C, pulse is 70/min, respirations are 14/min, and blood pressure is 130/85 mm Hg. Laboratory studies show a serum creatinine level of 1.1 mg/dL. Urinalysis shows no blood, protein, or glucose, and microscopic examination of the urine shows no WBCs or RBCs. Abdominal CT scan shows a 7-cm eccentric lesion of the upper pole of the right kidney. The lesion is well circumscribed and cystic with a thin wall and focal hemorrhage. What is the most likely diagnosis?

A  Acute pyelonephritis
B  Acute tubular injury
C  Diabetic nephropathy
D  Hydronephrosis
E  Simple renal cyst
F  Renal cell carcinoma
G  Urothelial carcinoma

A 24-year-old man has severe lower abdominal pain that radiates to the groin. On a scale of 1 to 10, the pain is at 10 and comes in waves. He then notes blood in his urine. He has no underlying illnesses and has been healthy all his life. On physical examination, he is afebrile and has a blood pressure of 110/70 mm Hg. Laboratory studies show serum Na+, 142 mmol/L; K+, 4 mmol/L; Cl−, 96 mmol/L; CO2, 25 mmol/L; glucose, 74 mg/dL; creatinine, 1.1 mg/dL; calcium, 9.1 mg/dL; and phosphorus, 2.9 mg/dL. Urinalysis shows a pH of 7; specific gravity of 1.020; and no protein, glucose, or nitrite. He is advised to drink more water. He likes iced tea and consumes large quantities over the course of a hot summer. He continues to have similar episodes. Which of the following substances is most likely to be increased in his urine?

A  Calcium oxalate
B  Cystine
C  Magnesium ammonium phosphate
D  Mucoprotein
E  Uric acid

A 28-year-old, previously healthy man suddenly develops severe abdominal pain and begins passing red urine. There are no abnormalities on physical examination. Urinalysis shows a pH of 7; specific gravity, 1.015; 1+ hematuria; and no protein, glucose, or ketones. The patient is given a device to use in straining the urine for calculi. The next day, he recovers a 0.3-cm stone that is sent for analysis. The chemical composition shows a pH of 7.5; specific gravity, 1.020; 1+ hematuria; and no protein, glucose, or ketones. Microscopic examination of the urine shows many RBCs, WBCs, and triple phosphate crystals. Which of the following renal lesions is most likely to be present?

A  Acute tubular injury
B  Malignant nephrosclerosis
C  Papillary necrosis
D  Renal cell carcinoma
E  Staghorn calculus

A 60-year-old man has had back pain and has passed dark-colored urine for the past month. On physical examination, there is tenderness over the right costovertebral angle. Urinalysis shows a pH of 6; specific gravity, 1.015; 2+ hematuria; and no protein, glucose, or ketones. Microscopic examination of the urine shows numerous RBCs, few WBCs, and no casts or crystals. The figure shows the representative gross appearance of the renal lesion. Which of the following substances is most likely to be increased in the blood of this man?

A  Cortisol
B  Globulins
C  Erythropoietin
D  Renin
E  Vasopressin

A 55-year-old man has had back pain and has passed dark-colored urine for the past month. On physical examination, there is tenderness over the right costovertebral angle. Urinalysis shows a pH of 7; specific gravity, 1.015; 1+ hematuria; and no protein, glucose, or ketones. Microscopic examination of the urine shows many RBCs, WBCs, and triple phosphate crystals. Which of the following renal lesions is most likely to be present?

A  Acute tubular injury
B  Malignant nephrosclerosis
C  Papillary necrosis
D  Renal cell carcinoma
E  Staghorn calculus
55. Members of a family with a history of renal cancers undergo ultrasound screening. Two adults are found to have multifocal and bilateral renal mass lesions. Biopsies are obtained, and microscopic examination shows a papillary pattern. A mutation involving which of the following genes is most likely to be found in this family?
   A. MET
   B. PKD1
   C. RAS
   D. TSC1
   E. WT1

56. A 60-year-old man has noted a nonproductive cough along with back pain for 4 months. He has passed darker urine for 1 month. He has a 50 pack/year history of smoking. On examination, his blood pressure is 175/110 mm Hg. He has tenderness to percussion of the upper back. Urinalysis shows 3+ blood but no casts or crystals. Chest CT imaging shows a 4-cm solid nodule in the right lower lobe of his lung, as well as 1- to 2-cm lytic lesions in thoracic vertebrae. A neoplasm is most likely to have arisen in which of the following urinary tract locations in this man?
   A. Bladder dome
   B. Calyx
   C. Penile urethra
   D. Renal cortex
   E. Urachus

57. A 4-year-old girl has complained of abdominal pain for the past month. On physical examination, she is febrile, and palpation of the abdomen shows a tender mass on the right side. Bowel sounds are present. Laboratory studies show hematuria without proteinuria. Abdominal CT scan shows a 12-cm, circumscribed, solid mass in the right kidney. A right-sided nephrectomy is performed; the gross appearance of the mass is shown in the figure. What is the most likely diagnosis?
   A. Angiomyolipoma
   B. Interstitial cell tumor
   C. Renal cell carcinoma
   D. Transitional cell carcinoma
   E. Wilms tumor

ANSWERS

1. A. He has azotemia, and the BUN-to-Cr ratio is >20:1, the low FENa, high urine specific gravity, and hypotension suggest prerenal azotemia, and cardiac failure is the most likely cause. Membranous nephropathy is a renal cause with BUN-to-Cr ratio of 10:1 or less and likely proteinuria at nephrotic levels. Prostatic hyperplasia with urinary tract obstruction may produce postrenal azotemia with BUN-to-Cr ratio between 10:1 and 20:1. Systemic lupus erythematosus is likely to produce nephritic syndrome with RBCs and RBC casts, and BUN-to-Cr ratio of 10:1 or less. Urothelial carcinoma is likely to be unilateral, with sufficient reserve renal function in the remaining kidney to prevent azotemia.

2. C. The findings fit with immune-mediated glomerular injury with antigen-antibody complex deposition, typical of an underlying disease such as SLE, and nephritic picture. Type I hypersensitivity may play a role in drug-induced acute interstitial nephritis. Type II hypersensitivity is present with anti–glomerular basement membrane diseases, such as Goodpasture syndrome. Type IV hypersensitivity plays a role in renal transplant rejection and some cases of drug-induced interstitial nephritis.
4. E The podocytes (visceral epithelial cells) of the glomerulus form a filtration barrier that depends upon both the anionic charge as well as slit diaphragms. Injuries that cause effacement, retraction, detachment, and vacuolization allow protein, principally albumin, to escape into Bowman space. Podocytes have limited capacity for regeneration and repair. The loss of 3.5 g/day of protein defines nephrotic syndrome. About a sixth of filtration occurs through the mesangium. The remaining listed options do not participate in this epithelial protein filtration barrier.

PB9 900, 908 BP9 518, 523 BP8 542, 548

5. B The strains of group A streptococci that cause post-streptococcal glomerulonephritis (GN) differ from the strains that cause rheumatic fever, and most elicit an immune response via streptococcal pyogenic exotoxin B (SpeB). Most children with poststreptococcal GN recover, although 1% develop a rapidly progressive GN characterized by crescent formation. Progression to chronic renal failure occurs in 40% of affected adults. A urinary tract infection is not likely to accompany poststreptococcal GN because the organisms that caused the immunologic reaction are no longer present when symptoms of GN appear.

PB9 910–911 BP9 529–530 BP8 554–555

6. F She has Plasmodium infection with malaria and immune complex deposition with leukocytes in glomeruli that has led to a nephritidic syndrome associated with her acute proliferative glomerulonephritis; many of these are postinfectious. The inciting infection depends upon the locale and frequency of occurrence. In the United States, Staphylococcus aureus (with IgA antibodies) is now more common than Streptococcus pyogenes as a cause for postinfectious GN in developed nations. Of the remaining choices, only membranous GN is likely to have an infectious cause, but this is more likely to produce a nephrotic syndrome.

PB9 910–911 BP9 529–530 PB8 917–918 BP8 554–555

7. B The figure shows glomeruli with epithelial crescents indicative of a rapidly progressive glomerulonephritis. Crescentic GN is divided into three groups on the basis of immunofluorescence: type I (anti-glomerular basement membrane [GBM] disease); type II (immune complex disease); and type III (characterized by the absence of anti-GBM antibodies or immune complexes). Each type has a different cause and treatment. The presence of anti-GBM antibodies suggests Goodpasture syndrome; patients with this disorder require plasmapheresis. Type II crescentic GN can occur in systemic lupus erythematosus, in Henoch-Schönlein purpura, and after infections. Causes of type III crescentic GN include granulomatosis with polyangiitis (ANCA-associated vasculitis) and microscopic polyangiitis. A positive ANA test result may be reported in patients with lupus nephritis, which uncommonly manifests with glomerular crescents. HIV nephropathy has features similar to those of focal segmental glomerulosclerosis (FSGS), which is not rapidly progressive. Quantitative serum immunoglobulins are not helpful because the important consideration is the pattern of immune deposits in the kidney. Rheumatoid factor is present in rheumatoid arthritis, which is not typically associated with renal complications. Urine immunoelectrophoresis is useful in categorizing a monoclonal gammopathy.


8. A The linear pattern of staining shown in the figure indicates the presence of anti-glomerular basement membrane antibodies (directed against the noncollagenous domain of the α3 chain of type IV collagen). Such antibodies are typically seen in Goodpasture syndrome, a form of rapidly progressive glomerulonephritis (GN) that has a bimodal pattern of occurrence in younger and older men. The anti-streptolysin O titer is increased in poststreptococcal GN, which typically has a granular pattern of immune complex deposition. The C3 nephritic factor can be a marker for dense deposit disease. Some cases of membranous nephropathy are associated with hepatitis B virus infection, but the immune complex deposition is granular, not linear. HIV infection can lead to a nephropathy that resembles FSGS, in which IgM and C3 are deposited in the mesangial areas of affected glomeruli.


9. C This is pauci-immune crescentic glomerulonephritis. The lack of immune deposits excludes anti–glomerular basement membrane disease (Goodpasture syndrome). Cases can be idiopathic, associated with ANCAs, and limited to the kidney. Rapidly progressive GN often has an abrupt onset with nephritis marked by oliguria. Anti-DNA topoisomerase antibody is seen with scleroderma, which mainly affects the vasculature. Antinuclear antibody is present with many autoimmune diseases, such as SLE, with immune complex deposition in glomeruli. Anti-HBs may occur in some cases of secondary membranous nephropathy.


10. F Granulomatosis with polyangiitis (ANCA-associated vasculitis) is a cause for rapidly progressive glomerulonephritis (GN) characterized by epithelial crescents in Bowman space. Several features differentiate it from other forms of crescentic GN (e.g., Goodpasture syndrome), including the presence of granulomatous vasculitis, the absence of immune complexes or anti–glomerular basement membrane (GBM) antibodies, and the presence of C-ANCA. Focal segmental glomerulosclerosis (FSGS) does not affect renal vessels and is unlikely to produce crescents with a rapidly progressive presentation. Goodpasture syndrome is a form of rapidly progressive GN with crescent formation, but a granulomatous vasculitis is not present, and there is anti-GBM antibody, not C-ANCA. Hyperplastic arteriolsclerosis can lead to focal hemorrhages and necrosis, but without a granulomatous component, and the blood pressure is usually quite high. Lupus nephritis, membranoproliferative GN, and postinfectious GN (with many neutrophils in glomeruli) occasionally can have a rapidly progressive course with crescent formation, but they do not produce granulomatous vasculitis. In patients with lupus, the ANA test result is often positive. Membranous nephropathy is most likely to produce nephrotic syndrome without crescents.

PB9 912 BP9 520, 532 BP8 517, 935 BP8 558
11 D Nephrotic syndrome may be produced by an idiopathic form of membranous nephropathy. Diffuse basement membrane thickening, in the absence of proliferative changes, and granular deposits of IgG and C3 are typical of this condition. It is caused by antibody targeting M-type phospholipase A2 receptor antigen, the deposition of immune complexes on the basement membrane, and complement activation. In 75% of patients with membranous nephropathy, the cause of immune complex deposition is unknown. In the remaining cases an associated systemic disease (e.g., systemic lupus erythematosus) or some known cause of immune complex formation (e.g., drug reaction) exists. Antibodies that react with basement membrane give rise to a linear immunofluorescence pattern, as in Goodpasture syndrome. Membranous nephropathy has no association with streptococcal infections. There also is no evidence of cytokine-mediated or cell-mediated damage in this disease.

PBD9 914–916 BP9 526–527 PBD8 922–923 BP8 551–552

12 A Steroid-responsive proteinuria in a child is typical of minimal change disease, in which the kidney looks normal by light microscopy, but fusion of foot processes is visible with electron microscopy. The most likely cause of foot process fusion is a primary injury to visceral epithelial cells caused by T cell–derived cytokines. Acute cellular renal transplant rejection is mediated by T cell injury with tubulitis. IgA nephropathy with mesangial IgA deposition and consequent glomerular injury causes recurrent gross or microscopic hematuria and, far less commonly, nephrotic syndrome. Immune complex deposition in membranous nephropathy can cause nephrotic syndrome, but is less common in children than in adults and is not steroid responsive. Certain verocytotoxin-producing Escherichia coli strains can cause hemolytic uremic syndrome by injury to capillary endothelium.

PBD9 917–918 BP9 524–525 PBD8 942–946 BP8 550–551

13 A One of the most common causes of nephrotic syndrome in adults is membranous nephropathy, caused by immune complex deposition, shown in the figure as extensive granular deposits with C3. About 75% of cases are idiopathic and due to autoantibodies reacting against podocyte antigens; but some cases follow infections (e.g., hepatitis, malaria), or are associated with causes such as malignancies or autoimmune diseases. In some cases of AIDS, a nephropathy resembling focal segmental glomerulosclerosis occurs. Multiple myeloma can be complicated by systemic amyloidosis, which can involve the kidney. Recurrent urinary tract infections are typically caused by bacterial organisms and can cause chronic pyelonephritis. Nephrotitis may lead to interstitial nephritis, but it does not cause glomerular injury.

PBD9 914–916 BP9 526–527 PBD8 922–923 BP8 551–552

14 B A child with nephrotic syndrome and no other clinical findings is most likely to have minimal change disease, a name that reflects the paucity of pathologic findings. There is fusion of podocyte foot processes, which can be seen only by electron microscopy. This fusion leads to selective proteinuria of low molecular weight proteins (albumin). Variability of basement membrane thickening may be seen in Alport syndrome. The mesangial matrix is expanded in some forms of glomerulonephritis (e.g., IgA nephropathy) and other diseases, such as diabetes mellitus, but not in minimal change disease. Reduplication of glomerular basement membrane may be seen with membranoproliferative GN. Subepithelial electron-dense humps represent immune complexes and are seen in postinfectious GN.

PBD9 917–918 BP9 524–525 PBD8 924–926 BP8 549–550

15 B Focal segmental glomerulosclerosis (FSGS) shows sclerosis of only a segment of the glomerulus (segmental lesion), and because only 50% of the glomeruli are affected, this is focal disease. FSGS manifests clinically with nephrotic syndrome that does not respond to corticosteroid therapy. FSGS can result from many forms of glomerular injury; some may be linked to NPHS gene mutations. In contrast, corticosteroid-responsive nephrotic syndrome in children is typically caused by minimal change disease (lipid nephrosis) that is not associated with any glomerular change seen under the light microscope. Membranoproliferative glomerulonephritis (GN) and dense deposit disease are more likely to produce a nephritic syndrome in adults. A diabetic patient with nephrotic syndrome is likely to have nodular glomerulosclerosis or diffuse thickening of the basement membrane. An acute proliferative postinfectious GN has hypercellular glomeruli with neutrophils. A rapidly progressive GN is associated with hematuria, and glomerular crescents are present.

PBD9 918–919 BP9 525–526 PBD8 916–917 BP8 550–551

16 D Corticosteroid-resistant hematuria and proteinuria leading to hypertension and renal failure is typical for focal segmental glomerulosclerosis (FSGS). FSGS is now the most common cause of nephrotic syndrome in adults in the United States. Specialized extracellular areas overlying the glomerular basement membrane between adjacent foot processes of podocytes are called slit diaphragms, and these exert control over glomerular permeability. Mutations in genes affecting several proteins, including nephrin and podocin, have been found in inherited cases of FSGS; podocyte dysfunction, possibly caused by cytokines or unknown toxic factors, may be responsible for acquired cases of FSGS. FSGS with collapsing glomerulopathy is seen in patients with HIV-associated nephropathy. Immune complexes containing microbial antigens cause postinfectious glomerulonephritis (GN). Anti-glomerular basement membrane antibodies are responsible for Goodpasture syndrome. C3NeF is an autoantibody directed against C3 convertase, and it is seen in membranoproliferative GN. Inherited defects in basement membrane collagen cause Alport syndrome, also characterized by hematuria, but other congenital abnormalities, such as deafness, are often present, and nephrotic syndrome is uncommon.

PBD9 918–919 BP9 525–526 PBD8 916–917 BP8 550–551

17 B Dense deposit disease (formerly membranoproliferative glomerulonephritis type II) usually leads to hematuria, and half of cases end in chronic renal failure. The term
chronic glomerulonephritis (GN) often is used when sclerosis of many glomeruli is present with no clear cause. Membranous nephropathy is often accompanied by proteinuria but less likely hematuria, and is characterized by thickening of only the basement membrane and small electron-dense deposits. Postinfectious GN is often characterized by a hypercellular glomerulus with infiltration of polymorphonuclear leukocytes, but no basement membrane thickening. A rapidly progressive GN is marked by crescents forming in the Bowman space.

IgA nephropathy (also known as Berger disease) can explain this nephritic condition with the presence of recurrent hematuria in a young adult. Nephrotic syndrome is not present, and mesangial IgA deposition is characteristic. The initial episode of hematuria usually follows an upper respiratory infection. IgA nephropathy occurs with increased frequency in patients with celiac disease and liver disease. It proceeds to chronic renal failure within 20 years in up to half of cases. Diffuse proliferation and basement membrane thickening denote membranoproliferative glomerulonephritis (GN), with IgG and C3 deposited in the glomeruli. Granular staining of basement membrane with IgG antibodies denotes immune complex deposition, which may occur in postinfectious GN, along with subepithelial deposits seen on electron microscopy. Patients with these changes also have nephritic syndrome. Glomerular capillary thrombosis is typical of hemolytic uremic syndrome.

IgA nephropathy is a form of glomerulonephritis that does not produce tubular epithelial changes. TTP is often associated with inherited or acquired deficiencies of ADAMTS13, leading to thrombotic microangiopathy, similar to hemolytic uremic syndrome from *Escherichia coli* Shiga toxin damaging vascular endothelium. Nodular and diffuse glomerulosclerosis are typical changes in diabetic nephropathy that can occur following loss of pancreatic islet beta cells. Toxic injuries are most likely to damage tubular cells, leading to acute tubular necrosis.

Development of recurrent hematuria after a viral illness in a child or young adult is typically associated with IgA nephropathy. A renal biopsy specimen will show diffuse mesangial proliferation and electron-dense deposits in the mesangium. In these patients, some defect in immune regulation causes excessive mucosal IgA synthesis in response to viral or other environmental antigens. IgA complexes are deposited in the mesangium and initiate glomerular injury. Defects in the structure of glomerular basement membrane are a feature of hereditary nephritis, and antibodies against type IV collagen are formed in Goodpasture syndrome. Podocytes may be affected by many forms of glomerular disease, but singularly malfunction in minimal change disease.

Alport syndrome is a form of hereditary nephritis. Hematuria is the most common presenting feature, but proteinuria is often present and may be in the nephrotic range. Patients progress to chronic renal failure in adulthood. An X-linked pattern of inheritance is present in 85% of cases, but autosomal dominant and autosomal recessive pedigrees also exist. The foamy change in the tubular epithelial cells and ultrastructural alterations of the basement membrane are characteristic features. The genetic defect results from mutation in the gene for the α5 chain of type IV collagen.

Lupus nephritis is one manifestation of systemic problems related to immune complex deposition, including fever, arthralgias, myalgias, pancytopenia, and serositis with pericarditis and pleuritis, which are characteristic of systemic lupus erythematosus (SLE). Renal disease is common in SLE, and a renal biopsy helps to determine the severity of involvement and the appropriate therapy. Antidoublestranded DNA antibody is more specific for diffuse scleroderma, which is unlikely to have renal involvement. Anti–DNA topoisomerase I antibody is more specific for diffuse scleroderma, which does have renal involvement, although usually this
manifests as vascular disease and not as glomerulonephritis. Anti-glomerular basement membrane antibody is characteristic of Goodpasture syndrome, in which IgG antibody is deposited in a linear fashion along glomerular capillary basement membranes. Anti-histone antibody may be present in drug-induced lupus. ANCAs can be seen in some forms of vasculitis, such as ANCA-associated granulomatous vasculitis or microscopic polyangiitis. Antithrombocyte protein is present in mixed connective tissue disease, which has some features of SLE, but usually does not include severe renal involvement.

24 G This patient is septic, and the heart murmur strongly suggests infective endocarditis. Cardiac lesions are the source of emboli (from valvular vegetations or mural thrombi) that can lodge in renal artery branches, producing areas of coagulative necrosis. These areas of acute infarction typically are wedge-shaped on cut section because of the vascular flow pattern. In addition, these septic emboli can produce abscesses where they lodge in the vasculature. Bilaterally enlarged, cystic kidneys are typical of autosomal dominant polycystic kidney disease. This patient’s kidneys may have been normal-sized and smooth-surfaced before this event. Small, shrunken kidneys represent an end stage of many chronic renal diseases. Petechiae and edema may be seen in hypertensive arteriolosclerosis associated with malignant hypertension. Irregular cortical scars with pelvicalyceal dilation may represent hydronephrosis complicated by infection in chronic pyelonephritis, whereas dilation alone points to obstructive uropathy, such as occurs with bladder outlet obstruction.

25 F Nodular and diffuse glomerulosclerosis is a classic lesion in diabetic nephropathy. Patients with diabetes mellitus have an elevated level of glycosylated hemoglobin (HbA1c) and may initially have microalbuminuria, which predicts development of future overt diabetic nephropathy. There is progressive loss of renal function. These patients are often hypertensive and have hyaline arteriolosclerosis. The presence of overt proteinuria suggests progression to end-stage renal disease within 5 years. Anti-glomerular basement membrane antibody is seen in Goodpasture syndrome, which manifests as a rapidly progressive glomerulonephritis (GN). The ANA test is positive in a variety of autoimmune diseases, most typically systemic lupus erythematosus, which can be accompanied by GN. The ANCA test is positive in some forms of vasculitis, which can involve the kidneys. The anti-streptolysin O titer is elevated after streptococcal infections, which may cause postinfectious GN. The C3 nephritic factor may be present in dense deposit disease. Some patients with membranous nephropathy have a positive serologic test result for HBsAg.

26 F The figure shows nodular and diffuse glomerulosclerosis that often occur in patients with long-standing diabetes mellitus, which is often complicated by urinary tract infections. Infections with bacterial organisms also occur more frequently in patients with diabetes mellitus who have an elevated Hgb A1c. Deposits of IgG and C3 in the glomerular basement membrane occur with forms of glomerulonephritis (GN) caused by immune complex deposition, including lupus nephritis and membranous nephropathy. The only abnormality observed in minimal change disease is effacement of podocyte foot processes, but this change is not specific for minimal change disease and may be seen in other disorders that produce proteinuria. Crescentic GN is not typically seen in diabetes mellitus. IgA deposition in the mesangium occurs in IgA nephropathy (Berger disease). A necrotizing granulomatous vasculitis can be present in the kidneys of patients having granulomatosis with polyangiitis (ANCA-associated vasculitis).

27 B His severe muscle injury resulted in myoglobinemia and myoglobinuria. The large amount of excreted myoglobin produces a toxic form of acute tubular injury. With supportive care, the tubular epithelium can regenerate, and renal function can be restored. During the recovery phase of acute tubular injury, patients excrete large volumes of urine because the glomerular filtrate cannot be adequately reabsorbed by the damaged tubular epithelium. An infection with pyelonephritis is unlikely to be characterized by such a short course or such a marked loss of renal function. Trauma is not a cause of malignant hypertension. Glomerulonephritis does not occur as a result of trauma. A bilateral renal vein thrombosis is uncommon and not related to muscle trauma.

28 B The most common cause of acute tubular necrosis is ischemic injury. The hypotension that develops after myocardial infarction causes decreased renal blood flow, with intrarenal vasoconstriction. Sublethal endothelial injury from reduced renal blood flow leads to the increased release of the vasconstrictor endothelin and diminished amounts of the vasodilators nitric oxide and prostaglandin. The ischemic form of acute tubular injury is often accompanied by rupture of the basement membrane (tubulorrhexis). An initiating phase that lasts approximately 1 day is followed by a maintenance phase during which progressive oliguria and increasing blood urea nitrogen levels occur, with salt and water overload. This is followed by a recovery phase, during which there is a steady increase in urinary output and hypokalemia. Eventually, tubular function is restored. Treatment of this acute renal failure results in recovery of nearly all patients. Aldosterone plays a role in sodium absorption. Erythropoietin drives RBC production. Natriuretic peptide increases when there is congestive heart failure, but does not lead to renal ischemia. Vasopressin (antidiuretic hormone) controls free water clearance.

29 D In younger persons presenting with recurrent acute pyelonephritis, a search for acquired or congenital conditions producing obstruction or reflux is extremely important.
Culture helps identify organisms resistant to antibiotic therapy. The pathogenesis of ascending urinary tract infections involves bacteria ascending from the urinary bladder into the ureter and the pelvis. Urinary tract infections generally are more common in females because of their shorter urethra, but in the absence of abnormalities of the urinary tract, the infections tend to remain localized in the urinary bladder. Older women and sexually active women are at increased risk of urinary tract infections. An ANA may be ordered in the workup of autoimmune diseases such as systemic lupus erythematosus that may involve the kidney, but are noninfectious. A renal biopsy should not be done with findings of renal infection. Measurement of renin may be part of a workup for hypertension that can cause renal vascular narrowing and ultimately impair renal function, but it does not predispose to infections. Though radiologic imaging studies may help document the extent of urinary tract abnormalities, the infection still needs to be adequately treated, hopefully before significant damage occurs.

30 D These laboratory findings are consistent with diabetes mellitus and clinical features of acute pyelonephritis caused by *Escherichia coli* infection. Necrotizing papillitis with papillary necrosis is a complication of acute pyelonephritis, and diabetic patients are particularly prone to this development. In the absence of diabetes mellitus, papillary necrosis develops when acute pyelonephritis occurs in combination with urinary tract obstruction. Papillary necrosis also can occur with long-term use of anlogesics. Acute tubular necrosis typically occurs in acute renal failure caused by hypoxia (e.g., shock) or toxic injury (e.g., mercury). Crescentic glomerulonephritis causes rapidly progressive renal failure. Hydronephrosis occurs when urinary outflow is obstructed in the renal pelvis or in the ureter. Renal calculi can complicate conditions such as gout, but they do not complicate diabetes mellitus.

31 B The clinical features in this patient are typical of urinary tract infection, and *Escherichia coli* is the most common cause. The WBCs are characteristic of an acute inflammatory process. The presence of WBC casts indicates that the infection must have occurred in the kidney because casts are formed in renal tubules. Most infections of the urinary tract begin in the lower urinary tract and ascend to the kidneys. Hematogenous spread is less common. *Cryptococcus* and *Mycoplasma* are rare urinary tract pathogens. Group A streptococcus is best known as an antecedent infection to poststreptococcal glomerulonephritis, an immunologically mediated disease in which the organisms are not present at the site of glomerular injury. *Mycobacterium tuberculosis* causes the rare sterile pyuria; however, renal tuberculosis typically does not manifest as an acute febrile illness.

32 F This gross appearance of the kidney is characteristic of chronic pyelonephritis, caused most often by reflux nephropathy. Typical features include coarse and irregular scarring resulting from ascending infection, blunting and deformity of calyces, and asymmetric involvement of the kidneys. The loss of tubules from scarring gives rise to reduced renal concentrating ability; the patient had polyuria with a low specific gravity of the urine. Urinary tract obstruction favors recurrent urinary tract infection (UTI). Vesicoureteral reflux propels infected urine from the urinary bladder to the ureters and renal pelvis and predisposes to infection; it can be unilateral. Autosomal dominant polycystic kidney disease is a bilateral process; patients usually are not symptomatic until middle age; although cysts may become infected, there is unlikely to be recurrent UTI. Benign nephrosclerosis is a vascular disease that may accompany hypertension but does not carry a risk for infection. Congestive heart failure may predispose to acute tubular injury. Lupus nephritis is associated with extensive inflammatory changes of glomeruli that are noninfectious. Amyloidosis can lead to progressive renal failure as a result of amyloid deposition in the glomeruli; however, amyloid does not evoke an inflammatory response.

33 D An acute drug-induced interstitial nephritis can be caused by ampicillin. This is an immunologic reaction, probably caused by a drug acting as a hapten. Pharyngitis with poststreptococcal glomerulonephritis with deposition of immune complexes is unlikely to be accompanied by a rash or by eosinophils in the urine. Anti–glomerular basement membrane antibodies occur in Goodpasture syndrome, with hemorrhages in lungs as well. Acute pyelonephritis is an ascending infection; it is uncommonly caused by hematogenous spread of bacteria from other sites. Acute tubular injury can cause acute renal failure. It is caused by hypoxia resulting from shock or from toxic injury caused by chemicals such as mercury, and only rarely, if ever, by bacterial toxins.

34 A Various drugs can cause drug-induced interstitial nephritis, including sulfonamides, penicillins, cephalosporins, the fluoroquinolone antibiotics ciprofloxacin and norfloxacin, and the antituberculous drugs isoniazid and rifampin. Acute tubulointerstitial nephritis also can occur with use of thiazide and loop diuretics, cimetidine, ranitidine, omeprazole, and nonsteroidal anti-inflammatory drugs. The disease manifests about 2 weeks after the patient begins to use the drug. Elements of type I (increased IgE) and type IV (skin test positivity to drug hapten) hypersensitivity are present. Congestive heart failure can lead to acute tubular injury, but it is not associated with a rash or proteinuria. Hemolytic-uremic syndrome can occur after ingestion of strains of *Escherichia coli* that may be present in ground beef. Poststreptococcal glomerulonephritis (GN) could account for the proteinuria and hematuria seen in this patient, but not for the rash, because the strains of group A β-hemolytic streptococci that cause a skin infection precede by weeks the development of GN. WBCs, but not eosinophils, may be present in the urine of a patient with a urinary tract infection.
35 E Analgesic nephropathy damages the renal interstitium and can give rise to papillary necrosis; an uncommon but feared complication is urothelial carcinoma. Hydrenephrosis is unlikely to develop because there is no urinary tract obstruction in analgesic nephropathy. The sloughed papilla is likely to pass down the ureter. The toxic injury that occurs with analgesic use is slowly progressive and not acute, in contrast to the course of acute tubular injury. Glomeruli are not specifically injured with analgesic abuse.

PBD9 936  PBD8 945–946

36 D This patient’s hypertension is due to renal vascular constriction, typical for renal arterial atherosclerosis. In the face of reduced renal blood flow, his glomerular filtration rate (GFR) is maintained by prostaglandin-mediated vasoconstriction of afferent arterioles and angiotensin II-mediated vasoconstriction of efferent arterioles. The angiotensin-converting enzyme (ACE) inhibitor decreases efferent arteriolar vasoconstriction and decreases glomerular capillary perfusion pressure. Nonsteroidal anti-inflammatory drugs (NSAIDs) such as ibuprofen inhibit prostaglandin synthesis and lead to vasoconstriction that reduces renal blood flow and reduces GFR. Aldosterone is increased with increased renin and angiotensin production and leads to reduced sodium excretion. Histamine is a vasodilator from mast cell granules that plays a role in acute inflammatory processes, but not blood pressure regulation. Nitric oxide is a vasodilator, but does not have a significant effect on capillary blood flow. Tumor necrosis factor plays a role in many inflammatory processes, but not renal blood flow.

PBD9 936  BP9 517–518  PBD8 950–951

37 F The rapid cell turnover in acute leukemias and cell death from treatment cause the release of purines from the cellular DNA breakdown. The resulting hyperuricemia can predispose to the formation of uric acid crystal precipitation in collecting ducts. If renal calculi develop they can produce colicky pain when they pass down the ureter and through the urethra, and the local trauma to the urothelium can produce hematuria. Uric acid stones form in acidic urine. In contrast to stones containing calcium, uric acid stones are harder to visualize on a plain radiograph. The urine dipstick is sensitive to albumin, but not to globulins; a separate test for Bence Jones protein may be positive, although the dipstick protein result is negative. Bence Jones proteinuria is characteristic of multiple myeloma, however, not of leukemias or lymphomas. Eosinophils may appear in the urine in drug-induced interstitial nephritis. Myoglobin can cause the dipstick reagent for blood to become positive in the absence of RBCs or hemoglobin. Myoglobinuria results most often from rhabdomyolysis, which can occur after severe crush injuries. Oval fat bodies are sloughed tubular cells containing abundant lipid; they are characteristic of nephrotic syndromes. RBC casts appear in nephritic syndromes as a result of glomerular injury.

PBD9 936–937  BP9 545  PBD8 947  BP8 571–572

38 A These findings are characteristic of nephrocalcinosis resulting from hypercalcemia. One of the most common causes of hypercalcemia in adults is metastatic disease. The hypercalcemia produces a chronic tubulointerstitial disease of the kidneys that is initially manifested by loss of concentrating ability. With continued hypercalcemia, there is progressive loss of renal function. Urinary tract stones formed of calcium oxalate also may be present. Hypercholesterolemia may be seen in some cases of minimal change disease. Hypergammaglobulinemia with a monoclonal protein (M protein) may be present in multiple myeloma, but not in breast cancer. Hyperglycemia can occur in diabetes mellitus, but patients with cancer are not at increased risk of developing diabetes mellitus. Hyperuricemia occurs in some cases of gout. It also can occur in patients with neoplasms (particularly lymphomas and leukemias) that have a high proliferation rate and are treated with chemotherapy. In these cases, extensive cell death (lysis syndrome) causes acute elevations in uric acid levels, leading to urate nephropathy.

PBD9 937  PBD8 947

39 E There is a large amount of serum globulin, back pain from lytic lesions, immunosuppression with recurrent infections, and amyloid deposition enlarging the kidneys, all consistent with multiple myeloma. This AL amyloid deposition occurs in 6% to 24% of myeloma cases, and can involve kidneys. Patients with myeloma often have Bence Jones proteinuria (not detected by the standard dipstick urinalysis), and some have cast nephropathy, which can cause acute or, more commonly, chronic renal failure. Analgesic nephropathy (e.g., aspirin, phenacetin, acetaminophen) can lead to tubulointerstitial nephritis and papillary necrosis. There can be necrotizing vasculitis and fibrinoid necrosis of renal arteries with ANCA-associated granulomatous vasculitis, but not amyloid deposition. His serum glucose is not in the range for diabetes mellitus, and the pink deposits seen with nodular or diffuse glomerulosclerosis are not amyloid. The pink-staining, thickened capillary loops of membranous nephropathy represent immune deposits, not amyloid. Systemic lupus erythematosus can result in immune deposits to produce “wire loop” thickening of glomerular capillaries from immune deposition, not amyloid.

PBD9 937–938  BP9 439  PBD8 252, 935  BP8 167

40 C The figure shows hyaline arteriolosclerosis, which typically occurs in patients with benign hypertension, and the renal parenchymal changes may be termed benign nephrosclerosis. Similar changes can be seen with aging in the absence of hypertension. Vascular narrowing causes ischemic changes that are slow and progressive. There is diffuse scarring and shrinkage of the kidneys. Blood pressure screening is an important method that can identify patients with hypertension before significant organ damage has occurred. Essential (benign) hypertension may evolve to malignant hypertension that causes distinctive renal vascular lesions, including fibrinoid necrosis and hyperplastic arteriosclerosis. Chronic hypertension predisposes to cerebrovascular disease accompanied by transient ischemic attack and stroke. Acute tubular necrosis results from anoxic or toxic injury to the renal tubules. Fibromuscular dysplasia can involve one or more renal arterial layers and produce focal stenosis. In
interstitial nephritis, more cells would be seen in the urine sediment. An ANCA-associated vasculitis may have a necrotizing component, as can hyperplastic arteriolosclerosis.

41 B Malignant hypertension may follow long-standing benign hypertension. Two types of vascular lesions with accelerated nephrosclerosis are found in malignant hypertension. Fibrinoid necrosis of the arterioles may be present; in addition, there is intimal thickening in interlobular arteries and arterioles, caused by proliferation of smooth muscle cells and collagen deposition. The proliferating smooth muscle cells are concentrically arranged, and these lesions, called hyperplastic arteriolosclerosis, cause severe narrowing of the lumen. The resultant ischemia elevates the renin level, which further promotes vasoconstriction to potentiate the injury. Glomerular crescents are a feature of a rapidly progressive glomerulonephritis; however, the blood pressure elevation is not as marked as that seen in this patient. An IgA nephropathy involves glomeruli, but not typically the interstitium or vasculature. Nodular glomerulosclerosis is a feature of diabetes mellitus that slowly progresses over many years. Segmental tubular necrosis occurs in ischemic forms of acute tubular injury.

42 C Hemolytic uremic syndrome is one of the most common causes of acute renal failure in children. It most commonly occurs after ingestion of meat infected with verocytotoxin-producing Escherichia coli, most often serotype O157:H7. This Shiga toxin damages endothelium, reducing nitric oxide, promoting vasoconstriction and necrosis, and promoting platelet activation to form thrombi in small vessels. With supportive therapy, most patients recover in a few weeks, although perhaps one fourth progress to chronic renal failure. However, hemolytic uremic syndrome may also occur in adults from Shiga toxin and drug ingestion. Thrombotic thrombocytopenic purpura (TTP) in adults can lead to similar renal thrombotic microangiopathy, but is due to abnormal ADAMTS13 metalloproteinase clearance of von Willebrand multimers. Candidal urinary tract infections typically affect the urinary bladder. Clostridium difficile is best known for causing a pseudomembranous enterocolitis, not renal lesions. Proteus is a common cause of bacterial urinary tract infections, whereas Staphylococcus aureus is a less common cause.

43 D Autosomal dominant polycystic kidney disease (ADPKD) is described. Most cases have mutations in the PKD1 gene encoding polycystin-1, whereas about 15% have PKD2 mutations encoding polycystin-2 and a more slowly progressive course. Large cysts develop over many years, culminating in renal failure in adulthood. Elongin proteins are part of a complex formed with the von Hippel-Lindau (VHL) gene activity; cysts may be found in many organs, but VHL is best known as a tumor suppressor. Fibrocystin mutations are associated with autosomal recessive polycystic kidney disease (ARPKD) that manifests in utero. Mutations in NPHP1 encoding nephrocystin are associated with nephronophthisis–medullary cystic disease complex. Tuberin encoded by TSC2 plays a role in development of cysts associated with tuberous sclerosis.

44 A These findings are characteristic of autosomal dominant polycystic kidney disease (ADPKD). As seen in the figure, multiple large cysts have completely replaced the renal parenchyma; enlarging cysts and hemorrhage into cysts can cause pain. Mitral valve prolapse may be found in a fourth of patients. About 10% to 30% of affected patients with ADPKD have an intracranial berry aneurysm, and some of these can rupture without warning. Disseminated intravascular coagulation may complicate hemolytic uremic syndrome. Although ADPKD may involve liver, there are cysts, not cirrhosis. Ischemic heart disease results from atherosclerosis, which can accompany diabetes mellitus, frequently accompanied by renal disease. Pulmonary disease does not accompany ADPKD.

45 B Autosomal recessive polycystic kidney disease (ARPKD) most often occurs in children, and in this case with the distinctive finding of congenital hepatic fibrosis; most cases have PKHD1 gene mutations encoding for fibrocystin expressed in kidney, liver, and pancreas. By contrast, autosomal dominant polycystic kidney disease (ADPKD) manifests with renal failure in adults and involves PKD1 and PKD2 gene mutations encoding for polycystin proteins found in renal tubules. Some less common forms of ARPKD are accompanied by survival beyond infancy, and these patients develop congenital hepatic fibrosis. Enlarged kidneys with 1- to 4-cm cysts are characteristic of ADPKD in adults. Perhaps the most common renal cystic disease seen in fetuses and infants is multicystic renal dysplasia (multicystic dysplastic kidney), with focal, unilateral, or bilateral from variably sized cysts, but congenital hepatic fibrosis is not present. Medullary sponge kidney is a benign condition usually found on radiologic imaging of adults. Urethral atresia would produce marked bladder dilation, hydrourere, and hydronephrosis.

46 C The congenital disorder known as medullary sponge kidney (MSK) is present to some degree in 1% of adults. In MSK, cystic dilation of 1 to 5 mm is present in the inner medullary and papillary collecting ducts. MSK is bilateral in 70% of cases. Not all papillae are equally affected, although calculi are often present in dilated collecting ducts. Patients usually develop kidney stones, infection, or recurrent hematuria in the third or fourth decade. More than 50% of patients have stones. Autosomal dominant polycystic kidney disease (ADPKD) produces much larger cysts that involve the entire kidney, eventually leading to massive renomegaly. Autosomal recessive polycystic kidney disease (ARPKD) is rare and leads to bilateral, symmetric renal enlargement manifested...
in utero, with renal failure evident at birth. Multicystic renal dysplasia may occur sporadically or as part of various genetic syndromes, such as Meckel-Gruber syndrome, in fetuses and newborns.

PBD9 948  BP9 544  PBD8 957–959

47 B The child has nephronophthisis, the most common genetic cause for end-stage renal disease in children and adolescents, and transmitted in autosomal recessive pattern. The NPHP1 to NPHP11 genes encode for proteins found in the primary cilia, attached ciliary basal bodies, or the centrosome organelle, from which the basal bodies originate. There is loss of concentrating ability and renal tubular acidosis. MCKD1 mutations are associated with adult medullary cystic disease with autosomal dominant transmission. PKD1 encodes for polycystin-1, associated with autosomal dominant polycystic kidney disease (ADPKD). PKHD1 mutations encode for fibrocystin, and is associated with autosomal recessive polycystic kidney disease (ARPKD).

PBD9 948–949  BP9 544  PBD8 959–960  BP8 570–571

48 D The pelvic and calyceal dilation results from long-standing obstruction leading to hydrourter and hydronephrosis. In some patients with diabetes mellitus, neuropathy is complicated by a neurogenic bladder, and this can lead to functional obstruction. His diabetic neuropathy is also contributing to “diabetic foot” with the ulceration. There are many renal complications of diabetes mellitus, mostly from vascular, glomerular, or interstitial injury, but there is no obstruction. The scarring that accompanies analgesic nephropathy or chronic pyelonephritis can be marked; it is associated with significant loss of renal parenchyma, but not with pelvic dilation. With benign nephrosclerosis, the kidneys become smaller and develop granular surfaces, but there is no dilation. It is unlikely he would have lived 15 years with a prostatic cancer large enough to cause urinary tract obstruction.

PBD9 950–951  BP9 545–546  PBD8 960–962  BP8 571–573

49 E Simple cysts are common in adults, and multiple cysts may occur. The cysts are not as numerous as cysts occurring in autosomal dominant polycystic kidney disease (ADPKD), and there is no evidence of renal failure. Simple cysts may be as large as 10 cm, and hemorrhage sometimes occurs into a cyst. Multiple cysts sometimes develop in patients receiving long-term hemodialysis. Acute pyelonephritis is unlikely in this patient because of the absence of fever and WBCs in the urine. Acute pyelonephritis may be associated with small abscesses, but not with cysts, although in patients with ADPKD, cysts may become infected. Acute tubular necrosis follows ischemic or toxic injury, and there is evidence of renal failure. Diabetic nephropathy includes vascular and glomerular disease, but not cysts. Hydronephrosis may produce a focal obstruction of a calyx with dilation, but it does not produce an eccentric cyst. Neoplasms usually produce solid masses, although sometimes a renal cell carcinoma is cystic. The latter is much less common than a simple cyst.

PBD9 949  BP9 542–543  PBD8 960  BP8 569

50 A Ureteral colic from the passage of a stone down the ureter produces intense pain (10 out of 10). About 70% of all renal stones are composed of calcium oxalate crystals. Patients with these stones tend to have hypercalciuria without hypercalcemia. Uric acid stones and cystine stones tend to form in acidic urine. Cystine stones are rare. Triple phosphate (magnesium ammonium phosphate) stones tend to occur in association with recurrent urinary tract infections, particularly infections caused by urease-positive bacteria, such as Proteus. Mucoproteins may coalesce into hyaline casts, which are too small to produce signs and symptoms.

PBD9 951–952  BP9 545  PBD8 962–963  BP8 571–572

51 C Calcium oxalate stones are the most common type of urinary tract stone, and approximately 50% of patients have increased excretion of calcium without hypercalcemia. The basis of hypercalciuria is unclear. Infections can predispose to the formation of magnesium ammonium phosphate stones, particularly urea-splitting Proteus organisms. Diabetes mellitus is an uncommon cause of urinary tract lithiasis. Although infections are more common in diabetics, most are not caused by urea-splitting bacteria. Hyperparathyroidism predisposes affected individuals to form stones containing calcium, but few patients with urinary tract stones have this condition. Most uric acid stones are formed in acidic urine and are not related to gout. It is thought that these patients have an unexplained tendency to excrete acidic urine. At low pH, uric acid is insoluble, and stones form.

PBD9 951–952  BP9 545  PBD8 962  BP8 571

52 E Recurrent urinary tract infections with urea-splitting organisms such as Proteus can lead to formation of magnesium ammonium phosphate stones. These stones are large, and they fill the dilated calyceal system. Because of their large size and projections into the calyces, such stones are sometimes called staghorn calculi. Cases of acute tubular necrosis typically occur from toxic or ischemic renal injuries. Malignant nephrosclerosis is primarily a vascular process that is not associated with infection. Papillary necrosis can complicate diabetes mellitus. Infections are not a key feature of renal cell carcinoma.

PBD9 951–952  BP9 545  PBD8 962  BP8 571–572

53 C The figure shows a renal cell carcinoma. About 5% to 10% of these tumors secrete erythropoietin, giving rise to polycythemia. Other substances can be secreted, including corticotropin (adrenocorticotropic hormone), resulting in hypercortisolism in Cushing syndrome; but these cases are encountered less frequently than polycythemia. Renal cell carcinomas are usually unilateral, and typically they do not destroy all of a kidney, so there is no significant loss of renal function, and the serum urea nitrogen and creatinine levels are not elevated. Hypertension from hyperreninemia can occur in patients with some renal cell carcinomas, although this is uncommon. A syndrome of inappropriate antidiuretic hormone (vasopressin) is more likely a paraneoplastic syndrome associated with small cell lung carcinomas. Globulin
would be increased with multiple myeloma and lead to amyloid deposition, a diffuse process.

54 A The clear cell form of renal cell carcinoma, the most common form of kidney cancer, often manifests with painless hematuria, most often in individuals in the sixth or seventh decade; tobacco use is a risk factor. Most sporadic clear cell carcinomas show loss of both alleles of the VHL gene. Germline inheritance of the VHL mutation can give rise to von Hippel-Lind syndrome, with peak incidence of renal cell carcinoma in the fourth decade, and they may have other tumors, including cerebellar hemangioblastomas, retinal angiomas, and adrenal pheochromocytomas. HPV-16 infection is associated with carcinomas of the uterine cervix. Microsatellite instability is a feature of Lynch syndrome, also called hereditary nonpolyposis colon cancer syndrome, characterized by right-sided colon cancer and, in some cases, endometrial cancer. Mutation of the MET gene on chromosome 7 is associated with the papillary variant of renal cell carcinoma, but trisomies are not specific to renal cell carcinomas.

55 A The second most common carcinoma of the kidney in adults is papillary renal cell carcinoma, associated with MET gene mutations that can be familial or sporadic. PKD1 mutations are associated with autosomal dominant polycystic kidney disease. RAS mutations are common in many cancers at many sites, particularly the gastrointestinal tract, but are not likely associated with familial renal cancers. TSC1 encodes hamartin, associated with tuberous sclerosis, and angiomyolipomas of the kidney. WTI mutations can be found with Wilms tumors of the kidney, seen in childhood.

56 D Carry the CT imaging down to the upper abdomen and a renal mass is likely to be found, representing a renal cell carcinoma. Smoking is a risk factor. Distant metastases may be evident before the primary tumor produces symptoms. Hematuria is common, but can be microscopic. Production of hormones by a renal cell carcinoma may lead to hypertension. Removing the primary tumor may lead to regression of metastases, dependent upon elaboration of cytokines such as VEGF by the primary renal cell carcinoma. Urothelial carcinomas, also associated with smoking, are less likely to metastasize to bone and lung, and may arise in the renal calyces and pelvis, as well as in the urinary bladder and ureters. A urachal carcinoma, usually an adenocarcinoma, is a rare cause for hematuria that arises in a urachal remnant of the embryonic allantois.

57 E Wilms tumor is the most common renal neoplasm in children, and one of the most common childhood neoplasms. A complex staging, grading, and molecular analysis formula, and surgery, chemotherapy, and radiation result in a high cure rate. The microscopic pattern of Wilms tumor (nephroblastoma) resembles the fetal kidney nephrogenic zone. Angiomyolipomas may be sporadic or part of the genetic syndrome of tuberous sclerosis. They may be multiple and bilateral and have well-differentiated muscle, adipose tissue, and vascular components. Renomedullary interstitial cell tumors (medullary fibromas) are generally smaller than 1 cm and are incidental findings. Renal cell carcinoma is rare in children, and the most common patterns are clear cell, papillary, and chromophobe. Transitional cell carcinomas arise in the urothelium in adults and microscopically resemble urothelium.
1. An infant boy has had recurrent urinary tract infections since birth. Ultrasonography reveals dilated pelvis and calyces in the left kidney; the right kidney is absent. Surgical repair of an obstruction is performed. Which of the following pathologic findings is most likely present at the site of obstruction?

A. Adenocarcinoma within bladder extrophy  
B. Diverticulum with hemorrhage in the wall of the ureter  
C. Granulomatous inflammation within a double ureter  
D. Smooth muscle discontinuity at the uteropelvic junction  
E. Urachal remnant at the dome of the bladder

2. A 73-year-old man with urinary frequency and hesitancy has had three urinary tract infections within the past year. On physical examination, his prostate is diffusely enlarged. Which of the following pathologic findings is most likely to be present in his urinary bladder?

A. Diverticulum  
B. Interstitial cystitis  
C. Malakoplakia  
D. Papilloma  
E. Schistosome ova

3. A 69-year-old man with history of recurrent pancreatitis treated with corticosteroids now has increasing fatigue for 2 years. He does not drink alcohol and has no evidence of gallbladder disease. On examination, there are no abnormalities. Laboratory studies show his serum creatinine is 5 mg/dL and urea nitrogen is 48 mg/dL. His serum IgG4 is elevated. Ultrasound imaging shows bilateral hydronephrosis. What is abdominal CT imaging most likely to show in this man?

A. Nephrolithiasis  
B. Polypoid cystitis  
C. Retroperitoneal fibrosis  
D. Renal cell carcinoma  
E. Urothelial carcinoma

4. The top of the diaper is often noted to be damp on a girl infant. Radiologic imaging with contrast enhancement shows that there is a connection from the bladder to umbilicus. What is the most likely diagnosis?

A. Congenital diverticulum  
B. Exstrophy  
C. Persistent urachus  
D. Vesicoureteral reflux  
E. Vitelline duct remnant

5. A 51-year-old woman with diabetic nephropathy receives a renal allograft. An episode of acute cellular rejection requires an increase in immunosuppressive therapy. She develops dysuria. On examination, she has suprapubic pain on palpation. A urinalysis shows hematuria. Cystoscopy is performed, and 3- to 4-cm soft, yellow, slightly raised mucosal plaques are seen. Biopsy specimens of these lesions microscopically show mucosal infiltration by foamy macrophages with abundant PAS-positive cytoplasmic granules and small, laminated mineralized concretions. Which of the following organisms is most likely to be found in her urine?

A. Adenovirus  
B. Candida albicans  
C. Chlamydia trachomatis  
D. Escherichia coli  
E. Schistosoma haematobium

6. A study of patients with urothelial carcinoma of the urinary bladder is performed. Gross, microscopic, and molecular characteristics of these malignancies are analyzed. Survival is correlated with treatment. Which of the following findings in these malignancies is most likely to require radical cystectomy to improve survival?

A. Exposure to chemical carcinogens  
B. Invasion of muscularis propria  
C. Lack of response to BCG therapy  
D. Origin from inverted urothelial papilloma  
E. TP53 gene mutation
A 58-year-old man from Shanghai has noted passing darker urine within the past 3 weeks. On physical examination, there are no abnormalities. Urinalysis shows blood is present. Cystoscopy is performed and there is a 0.5 × 1.7 cm reddish area on the dome of the bladder. Biopsies are obtained and have the microscopic appearance shown in the figure. What is the most likely risk factor for his disease?

A. Congenital urachal remnant  
B. Inherited gene mutation  
C. Obesity  
D. Schistosomiasis  
E. Smoking

A 65-year-old man recently retired after many years in a job that involved exposure to aniline dyes, including β-naphthylamine. One month ago, he had an episode of hematuria that was not accompanied by abdominal pain. On physical examination, there are no abnormal findings. Urinalysis shows 4+ hematuria, and no ketones, glucose, or protein. Microscopic examination of the urine shows RBCs that are too numerous to count, 5 to 10 WBCs per high-power field, and no crystals or casts. The result of a urine culture is negative. Cystoscopy is performed, and biopsy of a lesion reveals the microscopic appearance in the figure. Which of the following neoplasms is he most likely to have?

A. Adenocarcinoma  
B. Rhabdomyosarcoma  
C. Renal cell carcinoma  
D. Squamous cell carcinoma  
E. Urothelial carcinoma

A 78-year-old man has had increasing difficulties with urination for the past 6 years. He has difficulty starting and stopping the urine stream. On physical examination, his temperature is 37° C and blood pressure is 130/85 mm Hg. The figure shows the representative gross appearance of the bladder. Which of the following laboratory findings is most likely to be reported in this patient?

A. Positive antinuclear antibody test  
B. Hemoglobin concentration of 22.5 g/dL  
C. Prostate-specific antigen level of 5 ng/mL  
D. Schistosoma haematobium eggs in urine  
E. Positive skin test for Mycobacterium tuberculosis

A 57-year-old woman has had pain on urination for 5 months and yesterday noted blood on her underwear. On examination there is a tender red 1-cm nodule on the posterior lip of the external urethra. It is excised. What pathologic finding is most likely to be present on microscopic examination of her lesion?

A. Granulation tissue  
B. Multinucleated cells  
C. Plasma cell infiltrates  
D. Rhabdomyosarcoma  
E. Squamous carcinoma  
F. Urothelial dysplasia

A 74-year-old woman has noticed a slowly enlarging mass on her urethra for the past 6 months. The mass causes local pain and irritation and is now bleeding. Physical examination shows a 2.5-cm warty, ulcerated mass protruding from the external urethral meatus. There are no lesions on the labia or vagina. A biopsy specimen of the lesion is most likely to identify which of the following?

A. Clear cell carcinoma  
B. Embryonal rhabdomyosarcoma  
C. Leiomyoma  
D. Condyloma lata  
E. Squamous cell carcinoma
12 A 5-year-old boy has a history of recurrent urinary tract infections. Urine cultures have grown *Escherichia coli, Proteus mirabilis*, and *Enterococcus*. Physical examination now shows an abnormal constricted opening of the urethra on the ventral aspect of the penis, 1.5 cm from the tip of the glans penis. There also is a cryptorchid testis on the right and an inguinal hernia on the left. What term best describes the child’s penile abnormality?

A Balanitis  
B Bowen disease  
C Epispadias  
D Hypospadias  
E Phimosis

13 A 19-year-old man has worsening local pain and irritation with difficult urination over the past 3 years. He has become more sexually active during the past year and describes his erections as painful. Physical examination shows that he is not circumcised. The prepuce (foreskin) cannot be easily retracted over the glans penis. What is the most likely diagnosis?

A Bowenoid papulosis  
B Epispadias  
C Genital candidiasis  
D Paraphimosis  
E Phimosis

14 A 46-year-old man with a history of poorly controlled diabetes mellitus has had painful, erosive, markedly pruritic lesions on the glans penis, scrotum, and inguinal regions of the skin for the past 2 months. Physical examination shows irregular, shallow, 1- to 4-cm erythematous ulcerations. Scrapings of the lesions are examined under the microscope. Which of the following microscopic findings in the scrapings is most likely to be reported?

A Atypical cells with hyperchromatic nuclei  
B Budding cells with pseudohyphae  
C Eggs and excrement of mites  
D Enlarged cells with intranuclear inclusions  
E Spirochetes under dark-field examination

15 A 23-year-old, sexually active man has been treated for *Neisseria gonorrhoeae* infection 6 times during the past 5 years. He now comes to the physician because of the increasing number and size of warty lesions slowly enlarging on his external genitalia during the past year. On physical examination, there are multiple 1- to 3-mm sessile, nonulcerated, papillary excrescences over the inner surface of the penile prepuce. These lesions are excised, but 2 years later, similar lesions appear. Which of the following conditions most likely predisposed him to development of these recurrent lesions?

A *Candida albicans* infection  
B Circumcision  
C Human papillomavirus infection  
D *Neisseria gonorrhoeae* infection  
E Paraphimosis  
F Phimosis

16 A 56-year-old man from Fortaleza, Brazil, has noted increasing size of a penile lesion for the past 18 months. Physical examination reveals the appearance shown in the figure, following resection. What is most likely to be seen on microscopic examination?

A Acute and chronic inflammation with budding cells and pseudohyphae  
B Dysplastic urothelium above the basement membrane  
C Hyperkeratotic, acanthotic, squamous epithelium overlying ectatic blood vessels  
D Infiltrating, pleomorphic, poorly differentiated squamous cells  
E Invasive glands with tall columnar mucinous epithelium

17 A 48-year-old man has noticed a reddish area on the penis for the past 3 months. On physical examination, there is a solitary 0.8-cm, plaquelike, erythematous area on the distal shaft of the penis. A routine microbiologic culture with a Gram-stained smear of the lesion shows normal skin flora. Microscopic examination of a biopsy specimen of the lesion shows dysplasia involving the full thickness of the epithelium. What is the most likely diagnosis?

A Balanitis  
B Bowen disease  
C Condyloma acuminatum  
D Primary syphilis  
E Soft chancre

18 An 18-year-old man comes to his physician for a routine health maintenance examination. On physical examination, there is no left testis palpable in the scrotum. The patient is healthy, has had no major illnesses, and has normal sexual function. Which of the following complications will you tell this man is most likely to occur?

A Carcinoma  
B Heritability  
C Infection  
D Infertility  
E No sequelae
19. A 64-year-old man noted pain with burning on urination a week ago. He has had discomfort in his scrotum for the past 2 days. On examination, the right testis is swollen and tender. Which of the following organisms is most likely to cause this man’s illness?

A. *Escherichia coli*
B. Human papillomavirus
C. Mumps virus
D. *Mycobacterium tuberculosis*
E. *Treponema pallidum*

20. A 36-year-old man and his 33-year-old wife have tried to conceive a child for 12 years, and now they are undergoing an infertility work-up. On physical examination, neither spouse has any remarkable findings. Laboratory studies show that the man has a sperm count in the low-normal range. On microscopic examination of the seminal fluid, the sperm have a normal morphologic appearance. A testicular biopsy is done. The biopsy specimen shows patchy atrophy of seminiferous tubules, but the remaining tubules show active spermatogenesis. Which of the following disorders is the most likely cause of his findings?

A. Failure of normal testicular descent
B. Hydrocele formation with compression
C. Klinefelter syndrome
D. Past mumps virus infection
E. Prior radiation exposure

21. A 23-year-old, previously healthy man suddenly develops severe pain in the scrotum. The pain continues unabated for 6 hours, and he goes to the emergency department. On physical examination, he is afebrile. There is exquisite tenderness of a slightly enlarged right testis, but there are no other remarkable findings. The gross appearance of the right testis is shown in the figure. Which of the following conditions is most likely to cause these findings?

A. Hemorrhagic choriocarcinoma
B. Lymphatic obstruction
C. Mycobacterial infection
D. Obstruction of blood flow
E. Previous vasectomy

22. A 33-year-old man has noted asymmetric enlargement of the scrotum over the past 4 months. On physical examination, the right testis is twice its normal size and has increased tenderness to palpation. The right testis is biopsied. The epididymis and the upper aspect of the right testis have extensive granulomatous inflammation with epithelioid cells, Langhans giant cells, and caseous necrosis. Which of the following infections is the most likely cause of these findings?

A. Chancroid
B. Gonorrhea
C. Mumps
D. Syphilis
E. Tuberculosis

23. A study of testicular carcinomas in adults is performed. These neoplasms have a high frequency of karyotypic abnormalities, particularly i(12p). Pathologic findings include focal intratubular germ cell neoplasia adjacent to the malignancies. Which of the following is the most likely risk factor for these carcinomas?

A. Gonadal dysgenesis
B. Human papillomavirus infection
C. Hydrocele
D. Syphilis
E. Torsion

24. A 29-year-old man complains of a vague feeling of painless heaviness in the scrotum for the past 5 months. He is otherwise healthy. Physical examination shows that the right testis is slightly larger than the left testis. An ultrasound scan shows a solid, circumscribed, 1.5-cm mass in the body of the right testis. The representative gross appearance of the mass is shown in the figure. A biopsy is done, and microscopic examination of the mass shows uniform nests of cells with distinct cell borders, glycogen-rich cytoplasm, and round nuclei with prominent nucleoli. There are aggregates of lymphocytes between these nests of cells. Which of the following features is most characteristic of this lesion?

A. Association with 46,X(fra)Y karyotype
B. Association with 46,XXY karyotype
C. Response to radiation therapy
D. Extensive pulmonary metastases
E. Elevation of human chorionic gonadotropin level
F. Elevation of α-fetoprotein level
25 A 37-year-old man has noticed bilateral breast enlargement over the past 6 months. On physical examination, both breasts are enlarged without masses. His right testis is firm and 1.5 times larger than his left testis. His serum estrogen is increased. An ultrasound scan shows a circumscribed 2-cm mass in the body of the right testis. A right orchiectomy is performed, and grossly the mass has a uniform, brown cut surface. The microscopic appearance is shown in the figure. With electron microscopy the cells have rod-shaped crystalloids of Reinke. What is the most likely diagnosis?

A Choriocarcinoma  
B Embryonal carcinoma  
C Gonadoblastoma  
D Leydig cell tumor  
E Seminoma  
F Teratoma  
G Yolk sac tumor

26 A 28-year-old man has noticed increasing enlargement and a feeling of heaviness in his scrotum for the past year. On physical examination, the right testis is twice its normal size, and it is firm and slightly tender. An ultrasound examination shows a 3.5-cm solid right testicular mass. Abdominal CT scan shows enlargement of the para-aortic lymph nodes. Multiple lung nodules are seen on a chest radiograph. Laboratory findings include markedly increased serum levels of chorionic gonadotropin and \( \alpha \)-fetoprotein. Which of the following neoplasms is the most likely diagnosis?

A Choriocarcinoma  
B Large diffuse B-cell lymphoma  
C Leydig cell tumor  
D Metastatic prostatic adenocarcinoma  
E Mixed germ cell tumor  
F Pure spermatocytic seminoma

27 A 32-year-old man has noticed an increased feeling of heaviness in his scrotum for the past 10 months. On physical examination, the left testis is three times the size of the right testis and is firm on palpation. An ultrasound scan shows a 6-cm solid mass within the body of the left testis. Laboratory studies include an elevated serum \( \alpha \)-fetoprotein level. Which of the following cellular components is most likely to be present in this mass?

A Cytotrophoblasts  
B Embryonal carcinoma cells  
C Leydig cells  
D Lymphoblasts  
E Seminoma cells  
F Yolk sac cells

28 A 26-year-old man has occasionally felt pain in the scrotum for the past 3 months. On physical examination, the right testis is more tender than the left, but does not appear to be enlarged. An ultrasound scan shows a 1.5-cm mass within the right testis. A right orchiectomy is performed, and gross examination shows the mass to be hemorrhagic and soft. A retroperitoneal lymph node dissection is done. In sections of the lymph nodes, a neoplasm is found with extensive necrosis and hemorrhage. Microscopic examination shows that areas of viable tumor are composed of cuboidal cells intermingled with large eosinophilic syncytiat cells containing multiple dark, pleomorphic nuclei. Immunohistochemical staining of syncytial cells is most likely to be positive for which of the following?

A \( \alpha \)-Fetoprotein  
B Carcinoembryonic antigen  
C CD20  
D Human chorionic gonadotropin  
E Testosterone  
F Vimentin

29 The mother of a 2-year-old boy notices that he has had increasing asymmetric enlargement of the scrotum over the past 6 months. On physical examination, there is a well-circumscribed, 2.5-cm mass in the left testis. A left orchiectomy is performed, and histologic examination of this mass shows sheets of cells and ill-defined glands composed of cuboidal cells, some of which contain eosinophilic hyaline globules. Microcysts and primitive glomeruloid structures also are seen. Immunohistochemical staining shows \( \alpha \)-fetoprotein (AFP) in the cytoplasm of the neoplastic cells. What is the most likely diagnosis?

A Choriocarcinoma  
B Leydig cell tumor  
C Seminoma  
D Teratoma  
E Yolk sac tumor

30 A 32-year-old man has noticed increased heaviness with enlargement of the scrotum over the past 9 months. On physical examination, there is an enlarged, firm left testis, but no other remarkable findings. An ultrasound scan shows a 5-cm solid mass within the body of the left testis. An orchiectomy of the left testis is performed. Microscopic examination of the mass shows areas of mature cartilage, keratinizing squamous epithelium, and colonic glandular epithelium. Laboratory findings include elevated levels of serum human chorionic gonadotropin (hCG) and \( \alpha \)-fetoprotein (AFP). Despite the appearance of the cells in the tumor, the surgeon tells the patient that he probably has a malignant testicular tumor. The surgeon’s conclusion is most likely based on which of the following factors?

A Age of the patient at diagnosis  
B Elevation of hCG and AFP levels  
C Location of the mass in the left testis  
D Presence of colonic glandular epithelium  
E Size of the tumor
31. A 59-year-old man notices gradual enlargement of the scrotum over the course of 1 year. The growth is not painful, but produces a sensation of heaviness. He has no problems with sexual function. Physical examination shows no lesions of the overlying scrotal skin and no obvious masses, but the scrotum is enlarged, boggy, and soft bilaterally. The transillumination test result is positive. What is the most likely diagnosis?

A. Elephantiasis
B. Hydrocele
C. Orchitis
D. Seminoma
E. Varicocele

32. A 54-year-old man has had dysuria with increased frequency and urgency of urination for the past 6 months. He has sometimes experienced mild lower back pain. On physical examination, he is afebrile. There is no costovertebral angle tenderness. The prostate gland feels normal in size; no nodules are palpable. Laboratory studies show that expressed prostatic secretions contain 30 leukocytes per high-power field. What is the most likely diagnosis?

A. Acute bacterial prostatitis
B. Chronic abacterial prostatitis
C. Prostatic adenocarcinoma
D. Prostatic hyperplasia
E. Syphilitic prostatitis

33. A 65-year-old man has had multiple, recurrent urinary tract infections for the past year. *Escherichia coli* and streptococcal organisms have been cultured from his urine during these episodes, with bacterial counts of more than 10^9/mL. He has difficulty with urination, including starting and stopping the urinary stream. Over the past week, he has again developed burning pain with urination. Urinalysis now shows a pH of 6.5, and specific gravity of 1.020. No blood or protein is present in the urine. Tests for leukocyte esterase and nitrite are positive. Microscopic examination of the urine shows numerous WBCs and a few WBC casts. Which of the following is the most likely condition predisposing him to recurrent infections?

A. Epispidias
B. Nodular prostatic hyperplasia
C. Phimosis
D. Posterior urethral valves
E. Prostatic adenocarcinoma
F. Vesicoureteral reflux

34. A clinical trial of two pharmacologic agents compares one agent that inhibits 5α-reductase and diminishes dihydrotestosterone (DHT) synthesis in the prostate with another agent that acts as an α1-adrenergic receptor blocker. The subjects are 40 to 80 years old. The study will determine whether symptoms of prostate disease are ameliorated in the individuals who take these drugs. Which of the following diseases of the prostate is most likely to benefit from one or both of these drugs?

A. Acute prostatitis
B. Adenocarcinoma
C. Chronic prostatitis
D. Leiomyoma
E. Nodular hyperplasia

35. A 72-year-old man has had increasing difficulty with urination for the past 10 years. He now has to get up several times each night because of a feeling of urgency, but each time the urine volume is not great. He has difficulty starting and stopping urination. On physical examination, the prostate is enlarged to twice its normal size, but is not tender to palpation. One year ago, his serum prostate-specific antigen (PSA) level was 6 ng/mL, and it is still at that level when retested. Which of the following drugs is most likely to be effective in treatment of this man?

A. Estrogen (hormone)
B. Finasteride (5α-reductase inhibitor)
C. Mitoxantrone (chemotherapy agent)
D. Nitrofurantoin (antibiotic)
E. Prednisone (corticosteroid)

36. A 71-year-old, previously healthy man comes to his physician for a routine health examination. On palpation, there is a nodule in his normal-sized prostate. Laboratory studies show a serum prostate-specific antigen (PSA) level of 17 ng/mL. A routine urinalysis shows no abnormalities. Which of the following histologic findings is most likely to be found in a subsequent biopsy specimen of his prostate?

A. Acute prostatitis
B. Adenocarcinoma
C. Chronic abacterial prostatitis
D. Nodular hyperplasia
E. Prostatic intraepithelial neoplasia

37. An 85-year-old man has experienced urinary hesitancy and nocturia for the past year. He has had increasing back pain for the past 6 months. On digital rectal examination, there is a hard, irregular prostate gland. A bone scan shows increased areas of uptake in the thoracic and lumbar vertebrae. Laboratory studies show a serum alkaline phosphatase level of 300 U/L, and serum prostate-specific antigen (PSA) level of 72 ng/mL. The blood urea nitrogen concentration is 44 mg/dL, and the serum creatinine level is 3.8 mg/dL. Transrectal biopsy specimens of all lobes of the prostate are obtained. Microscopic examination shows that more than 90% of the tissue has a pattern of cords and sheets of cells with hyperchromatic pleomorphic nuclei, prominent nuclei, and scant cytoplasm. Which of the following is the best classification for this patient’s disease?

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<tr>
<th>Stage</th>
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A 71-year-old African American man visits his physician for a checkup because he is worried about his family history of prostate cancer. Physical examination does not indicate any abnormalities. Because of the patient’s age and family history, his prostate-specific antigen (PSA) level is measured and is 8 ng/mL. Six months later, the PSA level is 10 ng/mL. A urologist obtains transrectal biopsy specimens, and microscopic examination shows multifocal areas of glandular hyperplasia and the appearance shown in the figure. Which of the following statements applies best to this clinical and pathologic scenario?

A. Associated with increased risk for invasive cancer
B. Chronic inflammation from urinary tract obstruction
C. Normal microscopic finding of the peripheral zone
D. Related to an inherited tumor suppressor gene mutation
E. Responsive to 5α-reductase inhibitor therapy

A 45-year-old man comes to the physician for a routine health maintenance examination. On physical examination, there are no remarkable findings. Laboratory findings include serum creatinine, 1.1 mg/dL; urea nitrogen, 17 mg/dL; glucose, 76 mg/dL; alkaline phosphatase, 89 U/L; and prostate-specific antigen (PSA), 16 ng/mL. Prostate biopsies are performed and the high power microscopic appearance of a biopsy specimen is shown in the figure. Which of the following is the most likely risk factor for his disease?

A. Epigenetic hypermethylation of GSTP1 gene
B. Exposure to naphthylamine compounds
C. Overproduction of dihydrotestosterone
D. Prolonged use of smoking tobacco
E. Recurrent bacterial urinary tract infections

ANSWERS

1. D The features of vesicoureteral reflex from ureteropelvic junction (UPJ) obstruction are described. This is the most common cause for hydronephrosis in infants and children. Obstructed tubules do not reabsorb β2-microglobulin. Other anomalies, such as contralateral renal agenesis, may be present. Surgical correction of the abnormal UPJ restores proper peristaltic urine flow. Bladder extrophy is an open abdominal wall defect. Urinary diverticula may predispose to infection, but not obstruction. All or part of one or both ureters may be duplicated, but this is usually an incidental finding; granulomatous inflammation in the urinary tract is uncommon. Urachal remnants (embryologic allantois) may predispose to infection but not hydronephrosis; adenocarcinoma may arise in a urachal cyst.

2. A He has findings consistent with nodular prostatic hyperplasia with bladder outlet obstruction. Diverticula may develop in the setting of obstruction and bladder wall hypertrophy, and predispose to urinary stasis with infection; they are not true diverticula. Interstitial cystitis is a complication of recurrent bladder infection, most often in women, and characterized by chronic pain. Malakoplakia is a rare response to bacterial infection in which collections of macrophages filled with degraded bacterial products elicit formation of intracellular laminated, calcified concretions called Michaelis-Gutmann bodies. True papillomas are uncommon urothelial proliferations and asymptomatic. Schistosomiasis leads to hematuria and inflammation with bladder wall fibrosis.

3. C Bilateral hydronephrosis, without hydroureter or bladder dilation, suggests that the problem involves both ureters. Obstructive uropathy has led to chronic renal failure with uremia. The elevated IgG4 suggests a fibroinflammatory process with IgG4-secreting plasma cells in the retroperitoneum; this process is part of autoimmune pancreatitis and...
may also involve biliary tract, salivary glands, and lungs, but is not associated with the other listed choices. Nephrolithiasis could cause ureteropelvic junction obstruction, but bilaterality would be uncommon. Polypoid cystitis results from inflammation but may mimic a tumor mass, and could obstruct one or both ureteral orifices with hydroureter. Renal cell carcinoma is likely to be unilateral, but may cause only focal obstruction. Urothelial carcinomas may be multifocal, but are unlikely to obstruct both ureters simultaneously.

4 C The embryologic urachus may not become obliterated, leaving a fistulous tract or a cyst between the bladder and abdominal wall at the umbilicus. Congenital diverticula result from either focal failure in formation of bladder musculature or bladder outlet obstruction, and there is no fistulous tract. Exstrophy refers to failure in development of the lower abdominal wall, leaving an open defect to the bladder. Abnormal reflux of bladder contents into the ureter defines vesicoureteral reflux, which may be due to congenital abnormalities of bladder development, but there is no fistulous tract. A vitelloidal remnant may account for a Meckel diverticulum, or rarely a fistulous tract from small intestine to umbilicus.

5 D This woman has a peculiar form of cystitis known as malakoplakia, when macrophages have reduced phagocytic function, and the concretions within macrophages are known as Michaelis-Gutmann bodies. Malakoplakia is a reaction to chronic bacterial infections, usually Escherichia coli and Proteus species, and often in the setting of immunosuppression. The most common organism associated with cases of acute cystitis is E. coli. The other organisms listed are uncommon causes for cystitis and for malakoplakia.

6 B Local resection may suffice for lower grade urothelial carcinomas that are minimally invasive and localized. The risk for recurrence may be reduced with BCG installation into the bladder. Once the muscular wall is invaded, radical cystectomy is needed. Chemical carcinogens such as aniline dyes increase the risk for developing urothelial carcinoma, but do not dictate therapy. An inverted urothelial papilloma is benign, but a papillary urothelial neoplasm of low malignant potential (PUNLAP) is a precursor to urothelial carcinomas, which can have TP53 mutations regardless of stage.

7 E Urothelial carcinoma in situ is shown in the figure. Smokers are at increased risk, and China has a third of the world’s smokers. Urothelial malignancies can be recurrent and multifocal; they are far more likely to be sporadic than familial, although gene mutations such those in TP53 may be the same. Familial malignancies often present much earlier in life. The embryologic allantois extends from the developing bladder and may persist as a urachal remnant forming a diverticulum, cyst, or fistula to the umbilicus, and there is a low risk for adenocarcinoma. Obesity increases the risk for malignancy, but correlation with a specific malignancy is difficult to draw. Schistosomiasis can lead to squamous metaplasia and increased risk for squamous carcinoma of the bladder.

8 E Exposure to arylamines markedly increases the risk of urothelial carcinoma shown in the figure, which can occur decades after the initial exposure. After absorption, aromatic amines are hydroxylated into an active form, which is detoxified by conjugation with glucuronic acid and then excreted. Urinary glucuronidase splits the nontoxic conjugated form into the active carcinogenic form. Adenocarcinoma is a complication of the congenital condition known as exstrophy of the bladder. Rhabdomyosarcoma of the pelvis is typically a pediatric neoplasm, and is not associated with chemical exposures. Renal cell carcinomas also may manifest as painless hematuria, but exposure to aniline dyes is not a risk factor. Squamous cell carcinoma is the most common malignancy of the urethra, but it is rare and has no relation to carcinogens.

9 C Bladder hypertrophy can result from outlet obstruction. In an older man, this type of obstruction is most often caused by prostatic enlargement caused by hyperplasia or carcinoma. Mild elevations in the prostate-specific antigen (PSA) level may occur in patients with prostatic hyperplasia, and greater increases in PSA suggest carcinoma. Autoimmune conditions may be associated with interstitial cystitis, but cystitis does not cause bladder neck obstruction. Polycythemia can be the result of a paraneoplastic syndrome, but urothelial malignancies are unlikely to produce this finding; renal cell carcinoma is a more likely cause. Schistosomiasis leads to chronic inflammation and scarring. Bladder outlet obstruction can increase the risk of infection, typically with bacterial organisms such as Escherichia coli, not Mycobacterium tuberculosis.

10 A She has a urethral caruncle, which is most common in postmenopausal women as a result of urethral prolapse with atrophy from decreased estrogen. Topical estrogen creams and anti-inflammatory agents may shrink the lesion. It is not an infectious process, so multinucleated cells (herpes simplex virus) or plasma cell infiltrates (syphilis) are unlikely. It is not malignant or premalignant. Lichen sclerosus occurs at this age, but not produce a mass lesion.

11 E Carcinoma of the urethra is uncommon. It tends to occur in older women and is locally aggressive. A clear cell carcinoma occurs on the cervix and may be related to in utero exposure to diethylstilbestrol. An embryonal rhabdomyosarcoma (sarcoma botryoides) is a rare tumor that occurs in children. Benign tumors, such as a leiomyoma, are typically
well circumscribed and do not ulcerate. Condyloma lata may appear in association with secondary syphilis, but are flat and typically do not ulcerate. Condyloma acuminate are papillary lesions with acanthosis, related to HPV infection, and usually do not ulcerate.

12 D Hypospadias is a congenital condition seen in about 1 in 300 male infants. The inguinal hernia and the cryptorchidism are abnormalities that may accompany this condition. Epispadias is a congenital condition in which the urethra opens abnormally on the dorsal aspect of the penis. Bowen disease, which is squamous cell carcinoma in situ of the penis, occurs in adults. Phimosis is a constriction preventing retraction of the prepuce. It can be congenital, but more likely is the result of inflammation of the foreskin of the penis (e.g., balanitis, a form of local inflammation of the glans penis).

13 E Phimosis can be congenital, but is more often a consequence of multiple episodes of balanitis (inflammation of the glans penis or foreskin). Balanitis leads to scarring that prevents retraction of the foreskin. Forcible retraction may result in vascular compromise, with further inflammation and swelling (paraphimosis). Bowenoid papulosis is a premalignant lesion of the penile shaft resulting from viral infection. Epispadias is a congenital condition in which the penile urethra opens onto the dorsal surface of the penis. Candidiasis is most likely to produce shallow ulcerations that are intensely pruritic.

14 B Genital candidiasis can occur in individuals without underlying illnesses, but it is far more common in individuals with diabetes mellitus. Warm, moist conditions at these sites favor fungal growth. Scabies mites are more likely to be found in linear burrows in epidermis scraped from the extremities. Neoplasms with atypical cells may ulcerate, but such lesions are unlikely to be shallow or multiple without a mass lesion present. Intranuclear inclusions suggest a viral infection; however, diabetes is not a risk factor for genital viral infections. These lesions are too large and numerous to be syphilitic chancre.

15 C Condyloma acuminatum is a benign, recurrent squamous epithelial proliferation resulting from infection with human papillomavirus (HPV) infection, one of many sexually transmitted diseases that can occur in sexually active individuals. Koilocytosis is particularly characteristic of HPV infection. Candidiasis can be associated with inflammation, such as balanoposthitis, but not condylomata. Recurrent gonococcal infection indicates that the patient is sexually active and at risk for additional infections, but is not the cause for the condylomata. Gonococcal infection causes suppurative lesions in which there may be liquefactive necrosis and a neutrophilic exudate or mixed inflammatory infiltrate. Circumcision generally reduces risks for infections. Phimosis is a nonretractile prepuce, and paraphimosis refers to forcible retraction of the prepuce that produces pain and urinary obstruction.

16 D Penile squamous carcinomas such as this large ulcerated mass are likely invasive, and this lowers the 5-year survival to less than 70%; if there is nodal involvement, 5-year survival is less than 30%. Prior phimosis and human papillomavirus infection, more likely in uncircumcised men, are risk factors. Candidiasis is not a risk factor. Angiokeratomas appear as localized, benign, red or blue papules. Urothelium extends to the urethral orifice and development of urothelial carcinoma is theoretically possible at this site, but is far less common than squamous carcinomas. Adenocarcinomas are rare at this site.

17 B Bowen disease is the in situ form of squamous cell carcinoma of the penis. Similar to carcinoma in situ elsewhere, it has a natural history of progression to invasive cancer if untreated. Poor hygiene and infection with human papillomavirus (particularly types 16 and 18) are factors that favor development of dysplasias and cancer of the genital epithelia. Balanitis is an inflammatory condition without dysplasia. Condylomas are raised, whitish lesions. Syphilis is a sexually transmitted disease that produces a hard chancre, which heals in a matter of weeks. A soft chancre may be seen with *Haemophilus ducreyi* infections.

18 A Cryptorchidism results from failure of the testis to descend from the abdominal cavity into the scrotum during fetal development. One or both testes may be involved. It is associated with an increased risk of testicular cancer. An undescended testis eventually atrophies during childhood. Unilateral cryptorchidism may lead to infertility, because it may be associated with atrophy of the contralateral descended testis. Isolated cryptorchidism is a developmental defect that is usually sporadic and is not inherited in the germline. Mumps infection tends to produce patchy bilateral testicular atrophy, usually without infertility.

19 A This is acute epididymitis/orchitis, and most of these infections are secondary to ascending infections from the urinary tract. The time course suggests bacterial infection. Human papillomavirus affects squamous epithelium. Mumps orchitis is likely to be bilateral, and not associated with urinary tract infection. Tuberculosis can produce testicular infection, but the time course is likely to be weeks to months, and with preceding respiratory disease. Syphilis can lead to orchitis, but is unlikely to be preceded by urinary tract infection.
Mumps is a common childhood infection that can produce orchitis as well as parotitis. Adults who have this infection more often develop orchitis. The orchitis in children is usually not severe, and its involvement of the testis is patchy or unilateral so that infertility is not a common outcome. Mumps orchitis may be more severe in adults. Cryptorchidism results from failure of the testis to descend into the scrotum normally; the abnormally positioned testis becomes atrophic throughout. A hydrocele is a fluid collection outside the body of the testis that does not interfere with spermatogenesis. Klinefelter syndrome and estrogen therapy can cause tubal atrophy, although it is generalized in both cases. Patchy loss of seminiferous tubules indicates a local inflammatory process. Radiation as well as many chemotherapeutic agents are particularly harmful to rapidly and continuously proliferating testicular germ cells, but the effect would be diffuse within the testicular parenchyma. Radiotherapy is typically targeted to malignancies to prevent damage to normal surrounding tissues. Patients who wish to father children may want to store sperm in a sperm bank before undergoing radiation or chemotherapy.

The markedly hemorrhagic appearance in the figure results from testicular torsion that obstructs venous outflow to a greater extent than the arterial supply. Doppler ultrasound shows reduced or no vascular flow in the affected testis. An abnormally positioned or anchored testis in the scrotum is a risk factor for this condition. Testicular carcinomas do not obstruct the blood flow, and are not likely to produce an acute event. Parasitic infestation, typically filariasis, obstructs the flow of lymph, leading to gradual enlargement of the scrotum with thickening of the overlying skin. Tuberculosis can spread from the lung through the bloodstream, producing miliary nodes that organize into primitive endodermal sinuses (Schiller-Duval bodies). The infection typically starts in the epididymis and spreads to the body of the testis. Chancroid caused by Haemophilus ducreyi leads to ulcerated nodules of the external genitalia. Mumps produces patchy orchitis with minimal inflammation, which heals with patchy fibrosis. Syphilis involves the body of the testis, and there can be gummatous inflammation with neutrophils, necrosis, and some mononuclear cells. Gonococcal infections produce acute inflammation.

Mutations involving SRY or other genes involved in testicular differentiation may increase the risk for testicular cancer. Another risk factor is cryptorchidism. Infections of the testis are not generally known to be associated with neoplasia. Mechanical problems, such as torsion with ischemia, tend not to be antecedents for neoplasia. Hydrocele is a benign fluid collection.

Seminoma is the most common form of “pure” testicular germ cell tumor that may remain confined to the testis (stage I). The figure shows a homogenous mass lesion. The prognosis is good in most cases, even with metastases, because seminomas are radiosensitive. Human chorionic gonadotropin (hCG) levels may be slightly elevated in about 15% of patients with seminoma. Elevated hCG levels suggest a component of syncytial cells; very high levels suggest choriocarcinoma. α-Fetoprotein levels are elevated in testicular tumors with a yolk sac component, and many tumors with an embryonal cell component also contain yolk sac cells. Testosterone is a product of Leydig cells, not germ cells. Fragile X syndrome is associated with bilaterally enlarged testes and mental retardation. Klinefelter syndrome is associated with bilaterally decreased testicular size and reduced fertility.

Leydig cell tumors of the testis are most often small, benign masses that may go unnoticed. Some patients have gynecomastia caused by androgenic or estrogenic hormone production (or both) by the tumor. Most patients are young to middle-aged men; sexual precocity may occur in the few boys who have such tumors. Choriocarcinomas are grossly soft and hemorrhagic masses that have large bizarre syncytiotrophoblast and cytotrophoblast cells and are aggressive. Embryonal carcinomas are large, aggressive tumors that have a variegated gross appearance and primitive cells with large, hyperchromatic nuclei. Gonadoblastomas are rare testicular tumors that arise in the setting of gonadal dysgenesis. A pure seminoma can be uniformly brown on cut surface, but often has a lymphoid stroma, and is not likely to secrete androgens or estrogens. Pure teratomas are rare and contain elements of three germ layers. Yolk sac tumors have cells that organize into primitive endodermal sinuses (Schiller-Duval bodies).

Although a modest elevation of the human chorionic gonadotropin (hCG) concentration can occur when a seminoma contains some syncytial giant cells, significant elevation of the α-fetoprotein (AFP) level never occurs with pure seminomas. Elevated levels of AFP and hCG effectively exclude the diagnosis of a pure seminoma and indicate the presence of a nonseminomatous tumor of the mixed type. The most common form of testicular neoplasm combines multiple elements; the term teratocarcinoma is sometimes used to describe tumors with elements of teratoma, embryonal carcinoma, and yolk sac tumor. The yolk sac element explains the high AFP level. Mixed tumors may include seminoma. Choriocarcinomas secrete high levels of hCG, but no AFP. It is unusual for a tumor to metastasize to the testis; this patient is of an age at which a primary cancer of the testis should be considered when a testicular mass is present. Lymphomas may involve the testis, usually...
when there is systemic involvement by a high-grade lesion. Prostatic adenocarcinoma and lymphomas do not elaborate hormones. Leydig cell tumors are non-germ cell tumors derived from the interstitial (Leydig) cells; they may elabo-
rate androgens.

27 F α-Fetoprotein (AFP) is a product of yolk sac cells that can be shown by immunohistochemical testing. Pure yolk sac tumors are rare in adults, but yolk sac components are common in mixed nonseminomatous tumors. Cytotrophoblasts do not produce a serum marker, but they may be present in a choriocarcinoma along with syncytiotrophoblasts, which do produce human chorionic gonadotropin. Embryonal carcinoma cells by themselves do not produce any specific marker. Embryonal carcinoma cells are common in nonseminomatous tumors, however, and are often mixed with other cell types. Leydig cells produce androgens. Lymphomas may be seen in high-grade non-Hodgkin lymphomas, which do not produce hormones. Pure seminomas do not produce AFP.

28 D Choriocarcinoma is the most aggressive testicular carcinoma. It often metastasizes widely. The primitive syncytiotrophoblasts mimic the syncytiotrophoblast of placental tissue and stain for human chorionic gonadotropin. α-Fetoprotein is a marker that is more likely to be found in mixed tumors with a yolk sac component. Carcinoembryonic antigen (CEA) is found in a variety of epithelial neoplasms, particularly adenocarcinomas. CD20 is a lymphoid marker for B cells. Testosterone is found in Leydig cells. Vimentin is more likely to be seen in sarcomas, which are rare in the testicular region.

29 E Yolk sac tumors are typically seen in boys younger than 3 years. The primitive glomeruloid structures are known as Schiller-Duval bodies. The cells are strongly positive for AFP. Embryonal carcinomas with yolk sac cells contain AFP, but they are seen in adults. They are composed of cords and sheets of primitive cells. Choriocarcinomas contain large, hyperchromatic, syncytiotrophoblastic cells. Seminomas have sheets and nests of cells resembling primitive germ cells, often with an intervening lymphoid stroma. Leydig cell tumors act in a benign fashion and may produce androgens or estrogens or both. Teratomas contain elements of mature cartilage; bone; or other endodermal, mesodermal, or ectodermal structures.

30 B The tumor has elements of all three germ layers and is a teratoma. It is uncommon for teratomas in men to be completely benign. The most common additional histologic component is embryonal carcinoma. The elevated levels of human chorionic gonadotropin and α-fetoprotein indicate that this is a mixed tumor with elements of choriocarcinoma and yolk sac cells. The size of the tumor, age of the patient, location of the tumor (e.g., right, left, cryptorchid), and differentiation of the glandular epithelium are not markers of malignancy. On examining more histologic sections from the mass, the pathologist would find the malignant elements.

31 B Hydrocele is one of the most common causes of scrotal enlargement. It consists of a serous fluid collection within the tunica vaginalis. Most cases are idiopathic, although some may result from local inflammation. Elephantiasis is a complication of parasitic filarial infections involving the inguinal lymphatics; it is typically bilateral. Orchitis involves the body of the testis without marked enlargement, but with tenderness. A seminoma is typically a firm unilateral mass. A varicocele is a collection of dilated veins (pampiniform plexus) that may produce increased warmth, which inhibits spermatogenesis.

32 B The patient has more than 10 leukocytes per high-power field, indicating prostatitis. Chronic abacterial prostatitis is the most common form of the disorder. Patients typically do not have a history of recurrent urinary tract infections. Patients with acute bacterial prostatitis, most often caused by Escherichia coli infection, have fever, chills, and dysuria; on rectal examination, the prostate is very tender. Prostate carcinomas generally do not have a significant amount of acute inflammation, and metastases are most often associated with pain; most prostatic conditions causing dysuria are benign. Nodular prostatic hyperplasia by itself is not an inflammatory process. Syphilis is a disease of the external genitalia, although the testis may be involved.

33 B Of the diseases listed, prostatic nodular hyperplasia is the most common in older men. When it causes obstruction of the prostatic urethra, it can predispose to bacterial urinary tract infections. Epispadias is a congenital condition, observed at birth. Phimosis can occur in uncircumcised men. It may be congenital or acquired from inflammation, usually at a much younger age. Posterior urethral valves produce bladder outlet obstruction in utero, with oligohydramnios. Prostatic adenocarcinomas are less likely than hyperplasia to cause obstructive symptoms. Vesicoureteral reflux is more likely to be present at an earlier age, and it does not account for the obstructive symptoms the patient has on urination.

34 E Androgens are the major hormonal stimuli of glandular and stromal proliferation resulting in nodular prostatic hyperplasia. Although testosterone production decreases with age, prostatic hyperplasia increases, probably because of an increased expression of prostatic hormonal receptors that enhance the effect of any DHT that is present. The 5α-reductase inhibitors, such as finasteride, diminish the prostate volume, specifically the glandular component, leading to improved urine flow. The α1-adrenergic receptor
blocks, such as tamsulosin, cause smooth muscle in the bladder neck and prostate to relax, which relieves symptoms and improves urine flow immediately. The other listed conditions are not amenable to therapy with these drugs.

35 B The clinical features are typical of nodular prostatic hyperplasia causing a slight elevation of the PSA level. A PSA level that remains unchanged for 1 year, as in this case, is less likely to be found with a prostate cancer. Finasteride is a 5α-reductase inhibitor that decreases formation of dihydrotestosterone (DHT) that binds to androgen receptors in prostatic stromal and epithelial cells, driving proliferation with prostate gland enlargement. However, α1-adrenergic blockers that diminish smooth muscle tone are somewhat more effective in treating nodular hyperplasia. Estrogen therapy has been used as antihormonal therapy in prostate cancer. Mitoxantrone is a chemotherapy agent that, when given with prednisone, has been shown to be effective in treating advanced prostate cancers. Nitrofurantoin is an antibiotic that is often used in treating urinary tract infections.

36 B The prostate-specific antigen (PSA) level is significantly elevated in this patient. The large increase is likely to be indicative of carcinoma. Typically, prostatic carcinomas are adenocarcinomas that form small glands packed back to back. Many adenocarcinomas of the prostate do not produce obstructive symptoms and may not be palpable on digital rectal examination. Inflammation and nodular hyperplasias can increase the PSA level, although not to a high level that increases significantly over time. Prostatic intraepithelial neoplasia, although an antecedent to adenocarcinoma, is not likely to increase the PSA significantly over time.

37 E The presence of a hard irregular nodule, along with the extremely high prostate-specific antigen (PSA) level, points most clearly to prostate carcinoma. Modest elevations of the PSA concentration can occur in nodular hyperplasia of the prostate and prostatitis. Symptoms of urinary obstruction are more prominent in nodular hyperplasia because the nodules are in the periurethral region; but this sign is insufficient to distinguish cancer from hyperplasia. Similarly, renal failure owing to obstruction or infarction is most common with nodular hyperplasia, but can occur with cancer as well. Levels of alkaline phosphatase are elevated when prostate carcinoma gives rise to osteoblastic metastases. Although staging and grading schemes for malignant disease seem daunting, they are applied intuitively. The lowest stage is the smallest, most localized tumor; higher stages represent larger tumors or spread of the disease inside or outside of the primary organ site. Grading schemes also start with the lowest, most well-differentiated tumor, as seen with the microscope. Higher grade tumors have increasingly abnormal-appearing cells and structures so poorly differentiated that they hardly resemble their site of origin. In this case, the prostate cancer has the highest grade (it does not have glandular structures) and the highest stage (it has metastasized to the spine).

38 A Prostatic intraepithelial neoplasia (PIN) shown here has dysplastic features including hyperchromatic cells crowded into a pseudo-multilayer appearance, with preservation of gland architecture. PIN is a potential precursor of prostatic adenocarcinoma. By itself, it does not warrant therapy because only about one third of patients diagnosed with PIN develop invasive cancer within 10 years. Conversely, in about 80% of cases in which prostate cancer is present, PIN can be found in the surrounding tissue. PIN usually does not increase the PSA levels. In this case, the elevation in PSA levels may have been caused in part by the coexistent hyperplasia. Although prostatitis may increase PSA levels, no inflammation is seen here. Although PIN can be found in the peripheral zone, it is not a normal finding. Although there is a family history, specific risk factors are difficult to identify; BRCA2 mutations account for a small number of prostate cancers. Prostatic hyperplasia may respond to inhibition of DHT synthesis, but not PIN or cancer.

39 A The figure shows prostatic adenocarcinoma with back-to-back glands, prominent nucleoli, and perineural invasion. Alterations of the glutathione S-transferase (GSTP1) gene allow damage from carcinogens. TMPRSS2-ETS fusion gene and PTEN mutations are common. Other genetic abnormalities in prostate cancer include variations in CAG repeats in the androgen receptor gene, BRCA2 mutations, and translocation of ETS family transcription genes. His prostate-specific antigen (PSA) level is four times the upper limit of normal. This is worrisome, but not an absolute indication of prostate cancer. Elevated PSA levels can occur with nodular hyperplasia or prostatitis. A higher level, a level that increases over time, or an increased free PSA is more suggestive of carcinoma. Naphthylamine compounds are linked to urothelial carcinomas. Increased dihydrotestosterone output from prostatic stromal cells drives nodular hyperplasia. Tobacco use is associated with many other cancers, including urothelial carcinoma and renal cell carcinoma. Recurrent urinary tract infections and hydronephrosis are complications of obstruction more commonly from nodular prostatic hyperplasia.
A 31-year-old, sexually active woman has had a mucopurulent vaginal discharge for 1 week. On pelvic examination, the cervix appears reddened around the os, but no erosions or mass lesions are present. A Pap smear shows numerous neutrophils, but no dysplastic cells. A cervical biopsy specimen shows marked follicular cervicitis. Which of the following infectious agents is most likely to produce these findings?

A. *Candida albicans*
B. *Chlamydia trachomatis*
C. *Gardnerella vaginalis*
D. *Herpes simplex virus*
E. *Human papillomavirus*
F. *Neisseria gonorrhoeae*
G. *Trichomonas vaginalis*

A 31-year-old woman has had vulvar pruritus along with a thick, whitish, odorless, globular vaginal discharge for the past week. On pelvic examination, the cervix appears erythematous, but there are no erosions or masses. A Pap smear shows budding cells and pseudohyphae. No dysplastic cells are present. Which of the following infectious agents is most likely to produce these findings?

A. *Candida albicans*
B. *Chlamydia trachomatis*
C. *Neisseria gonorrhoeae*
D. *Trichomonas vaginalis*
E. *Ureaplasma urealyticum*

A 17-year-old sexually active girl has had pelvic pain for 1 week. A pelvic examination shows mild erythema of the ectocervix and pain on palpation of right adnexa. A Pap smear shows many neutrophils, but no dysplastic cells. A cervical culture grows *Neisseria gonorrhoeae*. If the infection is not adequately treated, she will be at increased risk for which of the following complications?

A. Cervical carcinoma
B. Dysfunctional uterine bleeding
C. Ectopic pregnancy
D. Endometrial hyperplasia
E. Endometriosis
F. Placenta previa

A 25-year-old woman has experienced discomfort during sexual intercourse for the past month. On physical examination, there are no lesions of the external genitalia. Pelvic examination shows a focal area of swelling on the left posterolateral inner labium that is very tender on palpation. A 3-cm cystic lesion filled with purulent exudate is excised. In which of the following structures is this lesion most likely to develop?

A. Bartholin gland
B. Gartner duct
C. Hair follicle
D. Urogenital diaphragm
E. Vestibular bulb

A 53-year-old postmenopausal woman is concerned about pale areas on her labia that have been slowly enlarging for the past year. The areas cause discomfort and become easily irritated. Physical examination shows pale gray to parchment-like areas of skin that involve most of the labia majora, labia minora, and introitus. The introitus is narrowed. A biopsy specimen is taken and microscopically shows thinning of the squamous epithelium, a dense band of upper dermal hyaline collagen, and scattered upper dermal mononuclear inflammatory cells. What is the most likely diagnosis?

A. Extramammary Paget disease
B. Human papillomavirus infection
C. Lichen sclerosus et atrophicus
D. Pelvic inflammatory disease
E. Vulvar intraepithelial neoplasia

A 40-year-old woman has noted pruritic patches on her vulva for the past 4 months. On physical examination there are multiple 1.5- to 3-cm white, scaly plaques on the vulva. A biopsy of one lesion is taken, and on microscopic examination, it shows epidermal thickening with hyperkeratosis and intense dermal inflammation. Mitoses are seen in keratinocytes, but they exhibit no atypia. What is the most likely diagnosis?

A. Condyloma acuminatum
B. Contact dermatitis
C. Psoriasis
D. Squamous cell hyperplasia
E. Vulvar intraepithelial neoplasia
CHAPTER 22  Female Genital Tract

7. A 36-year-old sexually active woman has noticed that warty vulvar lesions have been increasing in size and number over the past 5 years. On physical examination, there are multiple 0.5- to 2-cm, red-pink, flattened lesions with rough surfaces present on the vulva and perineum. One of the larger lesions is excised; its microscopic appearance is shown in the figure. Which of the following infectious agents is most likely to produce these lesions?

A. *Candida albicans*
B. *Chlamydia trachomatis*
C. *Haemophilus ducreyi*
D. Human papillomavirus
E. *Treponema pallidum*

8. A 57-year-old woman recently noticed a pale area of discoloration on the labia. Pelvic examination shows the presence of a 0.7-cm flat, white area on the right labia majora. A biopsy specimen is obtained and on microscopic examination shows dysplastic cells that occupy half the thickness of the squamous epithelium, with minimal underlying chronic inflammation. In situ hybridization shows human papillomavirus type 16 DNA in the epithelial cells. What is the most likely diagnosis?

A. Chronic vulvitis
B. Condyloma acuminatum
C. Lichen sclerosus et atrophicus
D. Squamous hyperplasia
E. Vulvar intraepithelial neoplasia

9. A 52-year-old woman has noted increasing size of a red, pruritic lesion on her left labium over the past 7 months. On physical examination, this rough, scaly lesion is 0.4 × 0.9 cm. The perineum appears normal; there is no lymphadenopathy, and there are no rectal lesions. A Pap smear shows no abnormal findings. The lesion is excised and on microscopic examination shows large atypical cells lying singly or in small clusters within the epidermis. These cells have abundant cytoplasm that stains with periodic acid–Schiff (PAS). What is the most likely diagnosis?

A. Condylomata acuminata
B. Extramammary Paget disease
C. Lichen sclerosus et atrophicus
D. Lichen simplex chronicus
E. Vulvar intraepithelial neoplasia

10. An 18-year-old sexually active woman has had dyspareunia followed by vaginal bleeding for the past month. On pelvic examination, a red, friable, 2.5-cm nodular mass is seen on the anterior wall of the upper third of the vagina. The microscopic appearance of a biopsy specimen is shown in the figure. Which of the following conditions is likely to have contributed most to the origin of this neoplasm?

A. Congenital adrenal hyperplasia
B. Diethylstilbestrol (DES) exposure
C. Human papillomavirus infection
D. Polycystic ovary syndrome (PCOS)
E. *Trichomonas* vaginitis

11. A 4-year-old girl is brought to the physician by her parents, who noticed bloodstained underwear and “something” protruding from her external genitalia. On physical examination, there are polypoid, grapelike masses projecting from the vagina. Histologic examination of a biopsy specimen from the lesion shows small, round tumor cells, some of which have eosinophilic straplike cytoplasm. Immunohistochemical staining shows desmin, vimentin, and myogenin in these cells. What is the most likely diagnosis?

A. Clear cell carcinoma
B. Infiltrating squamous cell carcinoma
C. Neuroblastoma
D. Sarcoma botryoides
E. Vulvar intraepithelial neoplasia

12. A healthy 30-year-old woman comes to the physician for a routine health maintenance examination. No abnormalities are found on physical examination. A screening Pap smear shows cells consistent with a low-grade squamous intraepithelial lesion (LSIL). Subsequent cervical biopsy specimens confirm the presence of cervical intraepithelial neoplasia (CIN) I. Which of the following risk factors is most likely related to her Pap smear findings?

A. Diethylstilbestrol (DES) exposure
B. Multiple sexual partners
C. Oral contraceptive use
D. Prior treatment for a malignancy
E. Vitamin B₁₂ (cobalamin) deficiency
A 33-year-old woman comes to her nurse practitioner for a routine health maintenance examination. On physical examination, there are no abnormal findings. A Pap smear shows abnormalities; colposcopy and a biopsy are performed. The figure shows the microscopic appearance of the biopsy specimen. Which of the following is the best strategy to prevent the development of this lesion?

A  Avoidance of tobacco products  
B  Consumption of a diet rich in vegetables  
C  Maintenance of an ideal body weight  
D  Use of oral contraceptives  
E  Vaccination for human papillomavirus

A 42-year-old woman has a Pap smear as part of a routine health maintenance examination. There are no remarkable findings on physical examination. The Pap smear shows cells consistent with a high-grade squamous intraepithelial lesion (HSIL) with human papillomavirus type 18. Cervical biopsy specimens are obtained, and microscopic examination confirms the presence of extensive moderate dysplasia (CIN II) along with intense chronic inflammation with squamous metaplasia in the endocervical canal. What is the most likely explanation for proceeding with cervical conization for this patient?

A  Her reproductive years are over  
B  HPV infection cannot be treated  
C  Perimenopausal state  
D  Presence of chronic cervicitis  
E  Risk for invasive carcinoma

A 28-year-old sexually active woman comes to her physician’s assistant for a routine health maintenance examination. There are no abnormal findings on physical examination. She has been taking oral contraceptives for the past 10 years. A Pap smear shows a high-grade squamous epithelial lesion (HSIL), also termed moderate dysplasia, or cervical intraepithelial neoplasia (CIN) II. What is the most likely molecular pathogenesis for this finding?

A  Estrogenic stimulation of cell proliferation  
B  Inheritance of a tumor suppressor gene mutation  
C  Recurrent gonococcal cervicitis  
D  Up-regulation of antiapoptosis genes  
E  Viral inactivation of the RB1 gene product

A 34-year-old woman has a routine Pap smear for the first time. The results indicate that dysplastic cells are present, consistent with a high-grade squamous intraepithelial lesion (HSIL), also called cervical intraepithelial neoplasia (CIN) III. She is referred to a gynecologist, who performs colposcopy and takes multiple cervical biopsy specimens that all show CIN III. Conization of the cervix shows a focus of microinvasion at the squamocolumnar junction. Based on these findings, what is the next most likely step in treating this patient?

A  Bone scan for metastatic lesions  
B  Course of radiation therapy  
C  No further therapy  
D  Pelvic exenteration  
E  Vaginal hysterectomy

A 45-year-old woman has had a small amount of vaginal bleeding and a brownish, foul-smelling discharge for the past month. On pelvic examination, there is a 3-cm lesion on the ectocervix, shown in the figure. Microscopic examination of the lesion is most likely to show which of the following?

A  Adenocarcinoma  
B  Cervical intraepithelial neoplasia  
C  Chronic cervicitis  
D  Clear cell carcinoma  
E  Extramammary Paget disease  
F  Squamous cell carcinoma
18. A 43-year-old woman has had postcoital bleeding for 6 months. She experienced menarche at age 11 years and has had 12 sexual partners during her life. She continues to have regular menstrual cycles without abnormal intermenstrual bleeding. Pelvic examination shows a focal, slightly raised area of erythema on the cervix at the 5 o’clock position. A Pap smear shows a high-grade squamous intraepithelial lesion (HSIL), also termed severe cervical intraepithelial neoplasia (CIN III). Analysis of cells from the cervix shows the presence of human papillomavirus type 16. Which of the following malignancies is she at greatest risk of developing if the lesion is not treated?

A. Clear cell carcinoma  
B. Immature teratoma  
C. Krukenberg tumor  
D. Leiomyosarcoma  
E. Sarcoma botryoides  
F. Squamous cell carcinoma

19. A 13-year-old girl began menstruation 1 year ago. She now has abnormal uterine bleeding, with menstrual periods that are 2 to 7 days long and 2 to 6 weeks apart. The amount of bleeding varies from minimal spotting to a very heavy flow. On physical examination, there are no remarkable findings. A pelvic ultrasound scan shows no abnormalities. Which of the following is most likely to produce these findings?

A. Anovulatory cycles  
B. Ectopic pregnancy  
C. Endometrial carcinoma  
D. Endometrial polyp  
E. Uterine leiomyomata

20. A 41-year-old G5, P5 woman has noticed lower abdominal pain with fever for the past 2 days. She delivered a normal term infant 1 week ago. On examination, she has a temperature of 37.4° C. There is a foul-smelling vaginal discharge. Which of the following pathologic findings is she most likely to have?

A. Cervical intraepithelial neoplasia  
B. Endometrial neutrophilic infiltrates  
C. Myometrial smooth muscle neoplasm  
D. Ovarian endometrioma  
E. Tubal granulomatous inflammation  
F. Vaginal trichomoniasis

21. A 35-year-old woman presents with infertility. She has had dysmenorrhea, dyspareunia, and pelvic pain on defecation for 4 years. Laparoscopic examination reveals red-blue nodules on the surface of the uterus and extensive adhesions between ovaries and the fallopian tubes. Histologic examination of a biopsy from one of the nodules shows hyperplastic endometrial glands and hemorrhage in the stroma. Molecular analysis of the biopsy material reveals hypomethylation of the promoter regions of the genes that encode steroidogenic factor 1 and estrogen receptor beta. There are no mutations in the PTEN, KRAS, and MLH1 genes. Which of the following is an appropriate treatment modality in this case?

A. Aromatase inhibitors  
B. Chemotherapy  
C. Estrogen  
D. Antitubercular therapy  
E. Total abdominal hysterectomy

22. A 36-year-old woman has had menorrhagia and pelvic pain for six months. She had a normal, uncomplicated pregnancy 10 years ago but has failed to conceive since then. She has been sexually active with one partner for the past 20 years and has had no dyspareunia. On pelvic examination she has a symmetrically enlarged uterus, with no apparent nodularity or palpable mass. A serum pregnancy test result is negative. What is the most likely diagnosis?

A. Adenomyosis  
B. Chronic endometritis  
C. Endometrial hyperplasia  
D. Endometriosis  
E. Leiomyoma

23. A 32-year-old woman has cyclic abdominal pain that coincides with her menses. Attempts to become pregnant have failed over the past 5 years. There are no abnormal findings on physical examination. Laparoscopic examination shows numerous hemorrhagic 0.2- to 0.5-cm lesions over the peritoneal surfaces of the uterus and ovaries. Which of the following ovarian lesions is most likely to be associated with her findings?

A. Fibroma  
B. Brenner tumor  
C. Endometriotic cyst  
D. Krukenberg tumor  
E. Metastatic choriocarcinoma  
F. Mucinous cystadenocarcinoma

24. A 49-year-old perimenopausal woman has had menometrorrhagia for the past 3 months. On physical examination, there are no remarkable findings. The microscopic appearance of an endometrial biopsy specimen is shown in the figure. The patient undergoes a dilation and curettage, and the bleeding stops, with no further problems. What condition is most likely to produce these findings?

A. Chronic endometritis  
B. Ovarian mature cystic teratoma  
C. Pregnancy with missed abortion  
D. Repeated failure of ovulation  
E. Use of oral contraceptives
25. A 52-year-old perimenopausal woman has had vaginal bleeding for a week. She has no medical problems and takes no medications. Hysteroscopy is performed and there is a single, 2-cm, smooth, soft mass protruding into the endometrial cavity. Biopsies are taken. What is microscopic examination of this lesion most likely to show?

A. Endocervical glands with squamous metaplasia  
B. Endometrial glands resembling stratum basalis  
C. Papillae with marked cellular atypia  
D. Smooth muscle cells in bundles  
E. Tubular glands lined by clear cells with glycogen

26. A 42-year-old woman has had menometrorrhagia for the past 2 months. She has no history of prior irregular menstrual bleeding, and she has not yet reached menopause. On physical examination, there are no vaginal or cervical lesions, and the uterus appears normal in size, but there is a right adnexal mass. An abdominal ultrasound scan shows the presence of a 7-cm solid right adnexal mass. Endometrial biopsy shows hyperplastic endometrium, but no cellular atypia. What is the most likely lesion that underlies her menstrual abnormalities?

A. Corpus luteum cyst  
B. Endometrioma  
C. Granulosa-theca cell tumor  
D. Mature cystic teratoma  
E. Metastasis  
F. Polycystic ovarian syndrome

27. A 62-year-old childless woman noticed a blood-tinged vaginal discharge twice during the past month. Her last menstrual period was 10 years ago. Bimanual pelvic examination shows that the uterus is normal in size, with no palpable adnexal masses. There are no cervical erosions or masses. Her body mass index is 33. Her medical history indicates that for the past 30 years she has had hypertension and type 2 diabetes mellitus. An endometrial biopsy specimen is most likely to show which of the following?

A. Adenocarcinoma  
B. Choriocarcinoma  
C. Leiomyosarcoma  
D. Malignant müllerian mixed tumor  
E. Squamous cell carcinoma

28. A study of patients with postmenopausal uterine bleeding reveals that some of them have malignant neoplasms that arise from prior atypical hyperplastic lesions. The peak incidence is between 55 and 65 years of age in women who have obesity, hypertension, and/or diabetes mellitus. Molecular analysis reveals mutations of the PTEN tumor suppressor gene in most of them. Their malignancies tend to remain localized for years before spreading to local lymphatics. Which of the following neoplasms is most likely to have these characteristics?

A. Clear cell carcinoma  
B. Endometrioid carcinoma  
C. Leiomyosarcoma  
D. Müllerian mixed tumor  
E. Stromal carcinoma

29. A 62-year-old obese, nulliparous woman has an episode of vaginal bleeding, which produces only 5 mL of blood. On pelvic examination, there is no enlargement of the uterus, and the cervix appears normal. A Pap smear shows cells consistent with adenocarcinoma. Which of the following preexisting conditions is most likely to have contributed to the development of this malignancy?

A. Adenomyosis  
B. Chronic endometritis  
C. Endometrial hyperplasia  
D. Human papillomavirus infection  
E. Use of oral contraceptives

30. A 40-year-old nulliparous woman has had menorrhagia for the past 6 months. On physical examination, her blood pressure is 154/93 mm Hg, there are no cervical lesions or adnexal masses, and the uterus is normal in size. She is 155 cm (5 feet 1 inch) tall and weighs 74.5 kg (body mass index 38). A Pap smear shows atypical glandular cells of uncertain significance. Hemoglobin A1c concentration is 9.8%. Endometrial biopsy shows complex hyperplasia with atypia; molecular analysis detects loss of PTEN gene heterozygosity and enhanced AKT phosphorylation. Which of the following metabolic pathways is most likely to be activated in this tumor?

A. Decreased glucose uptake  
B. Decreased prostaglandin synthesis  
C. Increased aerobic glycolysis  
D. Increased glycogen storage  
E. Increased oxidative phosphorylation

31. A healthy 59-year-old woman has had a feeling of pelvic heaviness for the past 11 months. There is no history of abnormal bleeding, and her last menstrual period was 8 years ago. Her physician palpates an enlarged nodular uterus on bimanual pelvic examination. A Pap smear shows no abnormalities. Pelvic CT scan shows multiple solid uterine masses; there is no evidence of necrosis or hemorrhage. A total abdominal hysterectomy is performed. Based on the gross appearance of the mass shown in the figure, what is the most likely diagnosis?

A. Adenomyosis  
B. Endometriosis  
C. Leiomyomas  
D. Metastases  
E. Tuberculosis
A 53-year-old woman whose last menstrual period was 3 years ago notes vaginal bleeding for a week. On physical examination, her uterus is markedly enlarged, but there are no adnexal masses. CT imaging reveals an irregular 8-cm mass in the body of the uterus. A total abdominal hysterectomy is performed, and microscopic examination of the soft, hemorrhagic mass shows spindle cells with atypia and numerous mitoses. There is coagulative necrosis of tumor cells. Which of the following is the most likely cell of origin for this mass?

A Cytotrophoblastic cells  
B Endometrial glandular cells  
C Germ cells  
D Smooth muscle cells  
E Squamous epithelial cells

A 69-year-old woman has passed blood per vagina for a month. On pelvic examination no abnormal findings are noted. Which of the following diagnostic procedures should be performed next?

A Endometrial biopsy  
B Magnetic resonance imaging  
C Microbiologic culture  
D Pap smear  
E Pregnancy test

A 28-year-old woman has had fever, pelvic pain, and a feeling of pelvic heaviness for the past week. Pelvic examination shows a palpable painful left adnexal mass. Laparoscopy shows an indistinct left fallopian tube that is part of a 5-cm circumscribed, red-tan mass involving the left adnexal region. Which of the following infectious agents is most likely to produce these findings?

A Chlamydia trachomatis  
B Haemophilus ducreyi  
C Herpes simplex virus  
D Mycobacterium tuberculosis  
E Treponema pallidum

A 19-year-old woman has the sudden onset of abdominal pain. On physical examination, there is pelvic pain on palpation. Her stool is negative for occult blood. The serum and urine pregnancy tests are negative. Transvaginal ultrasound shows no intrauterine gestational sac, and uterus and adnexa are normal in size. Culdocentesis yields a small amount of blood-tinged fluid. Which of the following has most likely led to these findings?

A Ectopic pregnancy  
B Endometriosis  
C Follicle cyst  
D Invasive mole  
E Pelvic inflammatory disease

A 21-year-old woman experienced menarche at age 14 years and had regular menstrual cycles for the next 3 years. For the past year, she has had oligomenorrhea and has developed hirsutism. She has noticed a 10-kg weight gain in the past 4 months. On pelvic examination, there are no vaginal or cervical lesions, the uterus is normal in size, and the adnexa are prominent. A pelvic ultrasound scan shows that each ovary is twice normal size, whereas the uterus is normal in size. Magnetic resonance imaging is shown in the figure. Which of the following conditions is most likely to be present in this woman?

A Immature teratomas  
B Krukenberg tumors  
C Ovarian cystadenocarcinomas  
D Polycystic ovarian syndrome  
E Tubo-ovarian abscesses

A 35-year-old woman has had increasing abdominal enlargement for the past 6 months. She states that she feels like she is pregnant, but results of a pregnancy test are negative. On physical examination, there is abdominal distention with a fluid wave. A pelvic ultrasound scan shows bilateral cystic ovarian masses, 10 cm on the right and 7 cm on the left. The masses are surgically removed. On gross examination, the excised masses are unilocular cysts filled with clear fluid, and papillary projections extend into the central lumen of the cyst. Microscopic examination shows that the papillae are covered with atypical cuboidal cells that invade underlying stroma. Psammoma bodies are present. What is the most likely diagnosis?

A Endometrioid tumor  
B Cystadenocarcinoma  
C Dysgerminoma  
D Granulosa cell tumor  
E Mature cystic teratoma  
F Sertoli-Leydig cell tumor
38. A 40-year-old woman has noticed progressive enlargement of the abdomen over the past 5 months, although her diet has not changed, and she has been exercising more. Physical examination shows no palpable masses, but a fluid wave is present. Paracentesis yields 500 mL of slightly cloudy fluid. Cytologic examination of the fluid shows malignant cells. An abdominal ultrasound scan shows a 15-cm multilobular mass that involves the right adnexal region. The uterus is normal in size. The mass is surgically removed; the figure shows the gross features of a section of the excised mass. What is the most likely diagnosis?

A. Choriocarcinoma  
B. Dysgerminoma  
C. Granulosa cell tumor  
D. Mucinous cystadenocarcinoma  
E. Teratoma with malignant transformation

39. A 56-year-old woman has had weight loss accompanied by abdominal enlargement for the past 5 months. There is a family history of breast and ovarian carcinoma. On physical examination, there are no lesions of the cervix, and the uterus is normal in size, but there is a left adnexal mass. An abdominal ultrasound scan shows a 10-cm cystic mass in the left adnexal region, with scattered 1-cm peritoneal nodules, and ascites. Cytologic studies of peritoneal fluid show malignant cells. Which of the following mutated genes is most likely a factor in the development of this neoplasm?

A. BRCA1  
B. ERBB2 (HER2)  
C. MYC  
D. KRAS  
E. RB1

40. A 42-year-old woman has noted dull lower abdominal pain for the past year. She reports no abnormal bleeding. On physical examination there is a large left adnexal mass. The pregnancy test is negative. Transvaginal ultrasound shows a right adnexal 10-cm cystic mass filled with fluid. The mass is removed and has the microscopic appearance shown in the figure. Which of the following is most likely to be associated with this lesion?

A. Brain metastases  
B. Endometrial hyperplasia  
C. Masculinization  
D. Peritoneal implants  
E. Sarcomatous transformation

41. A 33-year-old woman has had dull, constant abdominal pain for 6 months. On physical examination, the only finding is a right adnexal mass. CT scan of the pelvis shows a 7-cm circumscribed cystic mass on the right ovary, and it contains irregular calcifications. The right fallopian tube and ovary are surgically excised. The gross appearance of the ovary, which has been opened, is shown in the figure. Microscopic examination of this lesion is most likely to show which of the following?

A. Mature squamous epithelium  
B. Papillary structures with psammoma bodies  
C. Primitive neuroepithelium  
D. Rhabdomyoblasts in a cellular stroma  
E. Sheets of trophoblasts and syncytiotrophoblasts
A 23-year-old woman has had pelvic discomfort for 4 months. On pelvic examination, there is a large, nontender, right adnexal mass. An abdominal CT scan shows the 11-cm mass to be solid and circumscribed. On surgical removal, the mass is solid and white, with small areas of necrosis. Microscopically, it contains mostly primitive mesenchymal cells along with some cartilage, muscle, and foci of neuroepithelial differentiation. What is the most likely diagnosis?

A  Brenner tumor
B  Dysgerminoma
C  Granulosa cell tumor
D  Immature teratoma
E  Leiomyosarcoma
F  Malignant müllerian mixed tumor

A 52-year-old woman has had dull pain in the lower abdomen for the past 6 months and minimal vaginal bleeding on three occasions. Her last menstrual period was 2 years ago. Pelvic examination shows a right adnexal mass, and the uterus appears normal in size. An abdominal ultrasound scan shows an 8-cm solid mass, a small amount of ascites, and a right pleural effusion. A total abdominal hysterectomy is performed, and the mass is determined to be an ovarian fibroma. Which of the following additional lesions is most likely to be found in the excised specimen?

A  Bilateral chronic salpingitis
B  Cervical condylomata acuminata
C  Endometrial hyperplasia
D  Metastases to the uterine serosa
E  Partial mole of the uterus

A clinical study of women diagnosed with ovarian neoplasms reveals that 1 in 200 develop masculinizing signs and symptoms, including hirsutism, acne, breast atrophy, and amenorrhea. These women are found to have well-circumscribed, lobulated, firm, yellow mass lesions averaging 5 cm. Microscopically they have plump pink cells that show positive immunohistochemical staining for inhibin. Which of the following neoplasms are most likely to have these features?

A  Brenner tumor
B  Dysgerminoma
C  Endometrioid carcinoma
D  Granulosa-theca cell tumor
E  Sertoli-Leydig cell tumor

A 17-year-old girl missed a menstrual period, and her pregnancy test is positive. A month later, she notes suprapubic pain and passing blood clots from her vagina. She passes a small amount of tissue 3 days later. Pathologic examination of this tissue shows products of conception. Which of the following is the most likely cause for her pregnancy loss?

A  Bifid uterus
B  Group B streptococcus infection
C  Polycystic ovarian syndrome
D  Preeclampsia
E  Smoking cigarettes
F  Fetal trisomy 16

A 36-year-old woman has had an uneventful pregnancy for the past 37 weeks. Over the past 12 hours, she has developed lower abdominal pain. On examination, there is suprapubic tenderness. Her temperature is 37.4°C. Pelvic examination reveals a purulent cervical discharge. The infant is delivered 12 hours later. Which of the following organisms is most likely responsible for her premature labor?

A  Group B streptococcus
B  Herpes simplex virus
C  Rubella virus
D  Toxoplasma gondii
E  Treponema pallidum

A 22-year-old woman experiences sudden onset of severe lower abdominal pain. Physical examination shows no masses, but there is severe tenderness in the right lower quadrant. A pelvic examination shows no lesions of the cervix or vagina. Bowel sounds are detected. An abdominal ultrasound scan shows a 4-cm focal enlargement of the proximal right fallopian tube. A dilation and curettage procedure shows only decidua from the endometrial cavity. Which of the following laboratory findings is most likely to be reported for this patient?

A  Cervical culture positive for Neisseria gonorrhoeae
B  Detection of human chorionic gonadotropin in serum
C  69,XXY karyotype on decidual tissue cells
D  Pap smear showing pseudohyphae of Candida
E  Positive result of serologic testing for syphilis

A 36-year-old primigravida develops peripheral edema late in the second trimester. On physical examination, her blood pressure is 155/95 mm Hg. Urinalysis shows 2+ proteinuria, but no blood, glucose, or ketones. At 36 weeks, she gives birth to a normal viable but low-birth-weight infant. Her blood pressure returns to normal, and she no longer has proteinuria. Which of the following pathologic findings is most likely to be found on examination of the placenta?

A  Chorioamnionitis
B  Chronic villitis
C  Hydropic villi
D  Multiple infarcts
E  Partial mole

A 35-year-old primigravid woman at 30 weeks’ gestation develops worsening headaches along with a 3-kg weight gain over 1 week. This morning she had a generalized seizure. On physical examination, she is afebrile, but her blood pressure is 190/115 mm Hg (it was 120/80 mm Hg at a prenatal visit 1 month ago). She has peripheral edema involving her head and all extremities. Fetal heart tones of 140/min and fetal movement are present. Laboratory studies show hemoglobin, 12.5 g/dL; hematocrit, 37.6%; MCV, 92 μm³; platelet count, 199,000/mm³; serum creatinine, 1 mg/dL; potassium, 4.2 mmol/L; and glucose, 101 mg/dL. Urinalysis shows 2+ proteinuria, but no hematuria, RBCs, WBCs, or casts. Which of the following is the most likely underlying factor in the causation of her disease?

A  Adrenal cortical hyperplasia
B  Disseminated intravascular coagulation
C  Gestational trophoblastic disease
D  Ovarian neoplasm producing estrogen
E  Placental ischemia
F  Uncontrolled gestational diabetes
50. A 21-year-old G2, P1 woman is in the early second trimester. She has noted a small amount of vaginal bleeding for the past week and has had marked nausea and vomiting for 3 weeks. On physical examination, the uterus measures large for dates. An ultrasound examination shows intrauterine contents with a “snowstorm appearance,” and no fetus is identified. The gross appearance of tissue obtained by dilation and curettage is shown in the figure. Which of the following substances is most likely to be greatly increased in her serum?

A. Acetylcholinesterase  
B. α-Fetoprotein  
C. Estradiol  
D. Human chorionic gonadotropin  
E. Human placental lactogen

51. A 23-year-old woman, G3, P2, has a spontaneous abortion at 15 weeks’ gestation. The male fetus is small for gestational age and is malformed, with syndactyly of the third and fourth digits of each hand. The placenta also is small, and shows 0.5-cm grapelike villi scattered among morphologically normal villi. Chromosomal analysis of placental tissue is most likely to show which of the following karyotypes?

A. 45,X  
B. 46,XX  
C. 47,XXY  
D. 47,XY,+18  
E. 69,XXY

52. A 23-year-old woman suddenly notices a bloody, brownish vaginal discharge. The next day she has shortness of breath. On physical examination, a 3-cm, red-brown mass is seen on the lateral wall of the vagina. A chest radiograph shows numerous 2- to 5-cm nodules in both lungs. Laboratory studies show that her serum human chorionic gonadotropin levels are markedly elevated. A biopsy specimen of the vaginal mass is obtained and shown in the figure. Chromosome analysis of these cells shows a 46,XX karyotype. Which of the following cells is most likely present in this mass lesion?

A. Amnionic  
B. Rhabdomyoblast  
C. Serous epithelial  
D. Smooth muscle  
E. Syncytiotrophoblast

53. A 26-year-old woman delivered a normal neonate a month ago following an uncomplicated pregnancy. She now has vaginal bleeding. Hysteroscopy shows a nodule in the uterine fundus. Laboratory studies show hCG level of 200 mIU/mL. She is given chemotherapy but the lesion does not regress. Hysterectomy is performed. Microscopic examination of the nodule shows intermediate trophoblast cells. Immunostaining for which of the following proteins is most likely to yield positive results in this nodule?

A. α-Fetoprotein  
B. Chromogranin  
C. Desmin  
D. Human placental lactogen  
E. Neuron-specific enolase
CHAPTER 22  Female Genital Tract

1. B  The inflammatory cells in the cervical discharge with redness (erythema), and the biopsy findings indicate that the patient has cervicitis. *Chlamydia trachomatis* is the most common cause of cervicitis in sexually active women. Candidiasis, gonorrhea, and trichomoniasis also are common. Candidiasis often produces a scant, white, curdlike vaginal discharge; gonorrhea may have an associated urethritis; and *Trichomonas* may produce a profuse homogeneous, frothy, and adherent yellow or green vaginal discharge. *Gardnerella* is found in bacterial vaginosis, a common condition caused by overgrowth of bacteria. *Gardnerella* infection produces a moderate, homogenous, low-viscosity, adherent vaginal discharge that is white or gray and has a characteristic fishy odor; clue cells are seen on a wet mount. Herpetic infections are more likely to manifest as clear vesicles on the skin in the perineal region. Infection with human papillomavirus is associated with condylomata, dysplasias, and carcinoma.

2. A  The presence of the budding cells with pseudohyphae indicates a fungal infection with *Candida*. Candidal (monilial) vaginitis is common; this organism is present in about 5% to 10% of women. Recurrent episodes of vaginal candidiasis may be associated with non-albicans species. The inflammation tends to be superficial, and there is typically no invasion of underlying tissues. *Ureaplasma* is a bacterial agent, as is *Chlamydia*, and both can produce cervicitis. *Neisseria gonorrhoeae*, a gram-negative diplococcus, is the causative agent of gonorrhea. Infection with *Trichomonas vaginalis* can produce a purulent vaginal discharge, but the organisms are protozoa and do not produce hyphae.

3. C  Gonorrheal infections can lead to salpingitis and pelvic inflammatory disease with scarring of the fallopian tube. This predisposes to ectopic pregnancy, because the fertilized ovum has difficulty traversing the tube. Gonorrhea and other genital tract infections do not cause dysfunctional bleeding. Gonorrhea does not carry the risk of dysplasias or carcinomas that human papillomavirus infection does. Gonorrhea and other infections do not contribute to endometrial hyperplasia. The cause of endometriosis is not known with certainty, but infection does not seem to play a role in this process. Placenta previa results from low-lying implantation of the placenta and is not related to sexually transmitted diseases.

4. A  Bartholin glands may become obstructed, inflamed, and cystic because of abcess formation, which then produces focal pain. A Gartner duct cyst may form in the lateral vaginal wall from the remnant of a wolfian (mesonephric) duct; the cyst is filled with fluid and is usually not inflamed. Hair follicles are not present in the inner labia. The Bartholin gland lies just inferior to the fascia of the urogenital diaphragm and just anterior to the vestibular bulb, which is not glandular and does not become cystic.

5. C  Lichen sclerosus et atrophicus is most common in postmenopausal women. Although this lesion is not premalignant, there is a 1% to 5% risk that women with this condition will later develop a squamous cell carcinoma. In contrast, lichen simplex chronicus appears grossly as leukoplakia from squamous hyperplasia and is not associated with malignancy. Extramammary Paget disease is rare; it produces reddish areas of scaling and is caused by the presence of adenocarcinoma-like cells at the dermal-epidermal junction. Human papillomavirus infection is associated with condylomata acuminata and with squamous epithelial dysplasias. Pelvic inflammatory disease results from infection of internal genital organs with organisms such as *Neisseria gonorrhoeae* and *Chlamydia trachomatis*. Vulvar intraepithelial neoplasia is marked by dysplastic squamous epithelial changes.

6. D  Squamous cell hyperplasia, formerly called *lichen simplex chronicus*, is most often seen in women aged 30 to 50. It is not premalignant, but it may coexist with lichen sclerosus, and leukoplakia suggests the possibility of a squamous cell carcinoma to be distinguished on biopsy. Human papillomavirus infection is associated with condylomata acuminata and with squamous epithelial dysplasias that show keratinoctye vacuolization and minimal inflammation. Contact dermatitis produces red patches and vesicles, with intense round cell infiltrates, and tends to diminish when the offending antigen (such as a skin cream) is not used. Psoriatic lesions have extensive scaling, and microscopically show focal thinning of the epidermis with marked parakeratosis. Vulvar intraepithelial neoplasia is marked by dysplastic squamous epithelial changes.

7. D  The epithelium shows typical features of infection with human papillomavirus (HPV)—specifically, prominent perinuclear vacuolization (koilicytosis) and angulation of nuclei. These lesions, called *condylomata acuminata*, may occur anywhere on the anogenital surface, as single lesions or, more commonly, as multiple lesions. They are not precancerous. Condylomata are associated with HPV infection, often types 6 and 11. Candidal infections produce a vaginitis or cervicitis with exudate and erythema. Chlamydial infections may produce urethritis, cervicitis, and pelvic inflammatory disease. *Haemophilus ducreyi* is the agent that produces the soft chancre of chancroid. *Treponema pallidum* is the infectious agent of syphilis, characterized by the gross appearance of a hard chancre.
8 E Presence of dysplastic cells occupying half of the thickness of the epithelium suggests vulvar intraepithelial neoplasia (VIN). The incidence of these lesions has been increasing, probably because of more cases of human papillomavirus (HPV) infections. Some VIN lesions may progress to invasive cancers. Chronic inflammation alone does not produce dysplasia. A condyloma is usually a raised, nodular lesion. It also is caused by HPV, principally HPV-6 and HPV-11. Lichen sclerosus is a vulvar dystrophy characterized by thinning of the squamous epithelium and sclerosis of the dermis. Similar to VIN, squamous hyperplasia, another form of vulvar dystrophy, can appear as an area of leukoplakia, but no dysplastic changes are present.

9 B Extramammary Paget disease is a rare condition that is usually not associated with an underlying malignancy, in contrast to Paget disease of the breast. In many cases, the extramammary Paget cells remain in the epithelium, often for years, creating an annoying itchy red lesion. However, in a fourth of cases there may be an underlying neoplasm, so that local invasion and even metastases are possible. A condyloma is the result of human papillomavirus (HPV) infection and leads to koilocytic atypia, but the cells of a condyloma are not malignant. Lichen sclerosus is a white patch of epithelial thinning with dermal fibrosis and chronic inflammation that can be extensive enough to constrict the vaginal orifice; it may have an autoimmune basis, and there is an increased risk for future development of a squamous carcinoma. Lichen simplex chronicus is an area of epithelial hyperplasia that has no atypia and no association with malignancy. Vulvar intraepithelial neoplasia has neoplastic cells extending the full thickness of the epithelium; it is related to HPV infection.

10 B The microscopic appearance is that of a malignant tumor containing cells with a clear cytoplasm. Vaginal clear cell carcinomas are associated with exposure of the patient’s mother to diethylstilbestrol (DES) during pregnancy. These tumors are generally first diagnosed in the late teenage years. Congenital adrenal hyperplasia can produce masculinization in girls, manifesting in early childhood. Infection with human papillomavirus is associated with squamous epithelial dysplasias and malignancies, not with clear cell adenocarcinomas. Polycystic ovary disease can lead to hormonal imbalances from excess androgen production, but vaginal neoplasms do not arise in this setting. Trichomonal infections do not give rise to neoplasia.

11 D Embryonal rhabdomyosarcoma is an uncommon vaginal tumor that can be found in girls younger than 5 years. Because it forms polypoid, grapelike masses, it is sometimes called sarcoma botryoides. Histologically, it is a small round blue cell tumor that shows skeletal muscle differentiation in the presence of muscle-specific proteins such as desmin. Neuroblastomas are childhood tumors and are also small blue cell tumors, but they occur in the adrenal glands or extra-adrenal sympathetic chain. Clear cell carcinomas of the vagina may be related to in utero exposure to maternal diethylstilbestrol (DES), but have an onset in the second or third decades of life. Invasive squamous cell carcinomas are rare in very young patients, and they show histologic evidence of squamous epithelial differentiation, and are related to human papillomavirus (HPV) infection. Vulvar intraepithelial neoplasia is a carcinoma in situ of the vulvar skin, squamous in origin, and related to HPV infection.

12 B Cervical intraepithelial neoplasia (CIN) I represents minimal (mild) dysplasia (low-grade squamous intraepithelial lesion, or LSIL) and is a potentially reversible process. Dysplasias are preneoplastic and may progress to carcinomas if not treated. Risk factors for cervical dysplasias and carcinomas include early age at first intercourse, multiple sexual partners, and a male partner with multiple previous sexual partners. These factors all increase the potential for infection with human papillomavirus. Diethylstilbestrol (DES) exposure is a factor in the development of clear cell carcinomas of the vagina and cervix. Use of oral contraceptives, which contain very low amounts of hormonally active compounds, does not cause cervical dysplasia or carcinoma. Treatment of cancers does not typically result in dysplasias, although the atypical changes in epithelial cells from radiation and/or chemotherapy may be challenging to distinguish from cancer. A vitamin B12 deficiency may produce some megaloblastic epithelial changes, but not dysplasia.

13 E The figure shows a high-grade squamous intraepithelial lesion (HSIL) termed cervical intraepithelial neoplasia (CIN) III because the dysplasia involves the full thickness of the cervical epithelium. Such lesions arise more frequently in women who have had first intercourse at an early age, have multiple sexual partners, or have a male partner with multiple sexual partners. These factors are believed to increase the risk of infection with human papillomavirus (HPV), particularly types 16 and 18, which have high risk for dysplasias and carcinomas of the cervix. Because of the causal relationship with HPV infection, the use of HPV vaccines has been shown to prevent disease progression. Cervical squamous neoplasia has not been shown to be associated with smoking, diet, body weight, or hormonal influences.

14 E This patient’s cervical intraepithelial neoplasia (CIN) II is a high-grade squamous intraepithelial lesion (HSIL) that may progress to invasive carcinoma in several years if not treated, particularly because she has a high-risk subtype of HPV. Infection with HPV often drives this process, but the presence of HPV alone does not determine therapy. HPV infection cannot be eradicated with antibiotics, but patients may clear the virus. Chronic cervicitis with squamous metaplasia is not a malignant lesion and does not determine
therapy in this case. The conization can preserve fertility in women who are of childbearing age.

PBD9 1002–1004  BP9 686–688  PBD8 1020–1021  BP8 718–719

15 E Dysplasias of the cervix should not be ignored because they naturally progress to more severe dysplasias and to invasive carcinomas. Although not all cases progress, the physician should not take this chance. Dysplasias are strongly related to human papillomavirus (HPV) infections, and HPV DNA can be found in up to 90% of cases. Viral E6 and E7 proteins bind to Rb to up-regulate cyclin E. In about 10% to 15% of cases, there is no evidence of HPV, and other factors may play a role in the development of the dysplasia. With such HPV infection, the Pap smear may show changes of cervical intraepithelial carcinoma (CIN) I (low-grade squamous intraepithelial lesion, or LSIL). Oral contraceptives with low-dose estrogens and progestins do not increase the risk of dysplasia significantly. Though the RB1 gene is involved, this is not an inherited problem, and retinoblastomas are not seen with HPV infection. Cervicitis usually is due to bacterial or fungal organisms and is not a significant risk for dysplasia or carcinoma. Antipapoptosis genes such as BCL2 do not play a role in cervical carcinogenesis.


16 C Microinvasive squamous cell carcinomas of the cervix are stage I lesions that have a survival rate similar to that of in situ lesions. Such minimal invasiveness does not warrant more aggressive therapies. The likelihood of metastasis or recurrence is minimal.


17 F The cervical lesion shown in the figure is large and ulcerative and projects into the vagina. It is most likely an invasive squamous cell carcinoma that has infiltrated the subepithelial region. Dysplastic changes confined to the epithelium represent cervical intraepithelial neoplasia and do not form mass lesions. Glandular invasive lesions indicate an adenocarcinoma, which is much less common than squamous cell carcinoma of the cervix. Chronic cervicitis has erythema, but no mass effect. Clear cell carcinomas are uncommon and most likely found arising in the vagina. Extramammary Paget disease usually arises on the vulva, producing an eczematous lesion, not a mass, because the neoplastic cells are confined to the epithelium and to adjacent skin adnexa.

PBD9 1004–1006  BP9 684  PBD8 1021–1023  BP8 719–721

18 F This woman has several risk factors for the development of cervical squamous cell carcinoma, including multiple sexual partners, documented infection of the cervix with high-risk human papillomavirus (HPV) type 16, and diagnosis of a high-grade squamous intraepithelial lesion (HSIL). The remaining choices are not related to HPV infection. Clear cell carcinomas of the cervix are uncommon; some are associated with maternal use of diethylstilbestrol (DES) in pregnancy. An immature teratoma arises in the ovary. A Krukenberg tumor is a form of metastasis to the ovary. Leiomyosarcomas are rare and typically arise in the myometrium, although they can occur in the cervix. Sarcoma botryoides is a vaginal lesion that typically occurs in young girls.


19 A Anovulatory cycles are a common cause of dysfunctional uterine bleeding in young women who are beginning menstruation and in women approaching menopause. There is prolonged estrogenic stimulation that is not followed by secretion of progesterone. An ectopic pregnancy has acute findings and does not have a prolonged course. Endometrial carcinomas are rare in patients this age. Polyps are more common in older women. Submucosal leiomyomas are a cause of less variable bleeding and are more likely to be seen in older women.

PBD9 1007–1010  BP9 690–691  PBD8 1026–1027  BP8 723

20 B Acute endometritis in this case is the result of retained products of conception after delivery. Endometritis may also follow premature rupture of membranes with ascending infection to the uterine cavity. There is often polymicrobial infection with organisms found in the vagina. Some cases of chronic endometritis may be associated with Neisseria and Chlamydia infections and produce lymphoplasmacytic infiltrates within the endometrium. Cervical dysplasias are confined to the epithelium and are usually asymptomatic so that detection is by Pap smear. A myometrial neoplasm is unlikely to produce acute inflammation. An ovarian endometrioma is a mass lesion resulting from continued hemorrhage into a focus of endometriosis; but this mass lesion is not associated with pregnancy, and endometriosis may be a cause for infertility. Mycobacterium tuberculosis infection may spread to the female genital tract, most often the fallopian tube, but acute signs are unlikely to be present, and inflammation of the tube can be a cause for infertility. Vaginitis may produce acute inflammation with discharge, but trichomonal infections typically are associated with a watery gray-to-green discharge.

PBD9 1010  BP9 689  PBD8 1027  BP8 721

21 A In 30% to 40% of cases, endometriosis presents with infertility, menstrual irregularities, and pelvic pain. The presence of endometrial tissue in the nodules confirms this diagnosis. The glands in the nodules are hyperplastic but show no evidence of malignancy; in addition, all the genes implicated in endometrial cancer are normal. Hypomethylation of the two genes, NR5A1 (steroidogenic factor 1) and ESR2 (estrogen receptor beta) is found in endometriosis. These lead to overproduction of prostaglandins and estrogens. Aromatase inhibitors are used to suppress estrogen production. Lesions of endometriosis are not neoplastic and chemotherapy or major surgery with organ removal is not indicated. Endometriosis is not infectious, so antibiotics are not indicated.

PBD9 1010–1012  BP9 689–690  PBD8 1028–1029  BP8 722

22 A In adenomyosis, endometrial glands extend from the endometrium down into the myometrium. The process may
be superficial, but occasionally it is extensive, and the uterus becomes enlarged two to four times its normal size because of a reactive thickening of the myometrium. Chronic endometritis does not extend to the myometrium and does not increase uterine size. Endometrial hyperplasias do not increase the size of the uterus because the process is limited to the endometrium. In endometriosis, endometrial glands and stroma are found outside the uterus in such sites as peritoneum, ovaries, and ligaments. A leiomyoma is a myometrial tumor mass that, if large, produces an asymmetric uterine mass.

**23 C** Endometriosis is a condition in which functional endometrial glands are found outside the uterus. Common sites include ovaries, uterine ligaments, rectovaginal septum, and pelvic peritoneum. These endometrial glands can respond to ovarian hormones so that cyclic abdominal pain coincides with menstruation. Recurrent hemorrhages may incite scarring and the formation of fibrous adhesions in the pelvic region. This may cause distortion of the ovaries and fallopian tubes and may lead to infertility. One common variation is formation of an endometrioma, or chocolate cyst, which represents a focus of endometriosis that becomes an expanding cystic lesion as its center becomes filled with chocolate-brown sludge from the recurrent hemorrhage. The remaining choices are not associated with endometriosis, although endometrioid tumors may form in foci of endometriosis.

**24 D** Endometrial hyperplasia with numerous crowded glands as shown in the figure results from excessive estrogenic stimulation. This lesion often occurs with failure of ovulation about the time of menopause. Hyperplasias do not develop from endometritis. Estrogen-secreting ovarian tumors also may produce endometrial hyperplasia, but teratomas are not known for this phenomenon. A secretory pattern of the endometrium is seen in pregnancy, not the proliferative pattern shown in the figure. Oral contraceptives contain small doses of estrogenic compounds that do not lead to hyperplasia.

**25 B** She has an endometrial polyp, seen most often in perimenopausal and postmenopausal women. The lesion can lead to abnormal bleeding, but rarely gives rise to a malignancy. Endocervical glands with squamous metaplasia are seen most often with chronic cervicitis. Papillae with marked cellular atypia are seen with the serous type of endometrial carcinoma. Smooth muscle cells in bundles characterize a leiomyoma, which may be submucosal. Tubular glands lined by clear cells with glycogen are seen with the rare clear cell carcinoma.

**26 C** The mass is probably producing estrogen, which has led to endometrial hyperplasia. Estrogen-producing tumors of the ovary are typically sex cord tumors, such as a granulosa-theca cell tumor or a thecoma-fibroma, the former more often being functional. Teratomas can contain various histologic elements, but not estrogen-producing tissues. Endometriosis can give rise to an adnexal mass called an endometrioma, which enlarges over time. Endometrial glands are hormonally sensitive, but they do not produce hormones. Corpus luteum cysts are common, but they are unlikely to produce estrogens. Metastases to the ovary do not cause increased estrogen production. Polycystic ovarian syndrome would involve both ovaries.

**27 A** Postmenopausal vaginal bleeding is a red flag for endometrial carcinoma. Such carcinomas often arise in the setting of endometrial hyperplasia. Increased estrogenic stimulation is thought to drive this process, and risk factors include obesity, type 2 diabetes mellitus, hypertension, and infertility. Choriocarcinomas are gestational in origin. A submucosal leiomyosarcoma could produce vaginal bleeding, but the uterus would be enlarged because leiomyosarcomas tend to be large masses. Malignant müllerian mixed tumors are much less common than endometrial carcinomas, but they could produce similar findings. Malignant müllerian mixed tumors are typically uterine neoplasms that have glandular and stromal elements; the malignant stromal component can be heterologous and resemble mesenchymal cells that are not ordinarily found in the myometrium, such as cartilage. Squamous carcinomas of the endometrium are rare, and more likely to arise in the cervix.

**28 B** Most endometrial cancers have the endometrioid pattern and are classified as type I endometrial carcinomas. They arise in the setting of unopposed estrogen stimulation and may also have PTEN mutations as well as microsatellite instability. In contrast, type II endometrial carcinomas occur at an older age in the background of atrophic endometrium; they usually have a serous carcinoma pattern, but may also exhibit clear cell and müllerian mixed patterns, and TP53 mutations are common. Leiomyosarcomas and stromal sarcomas are far less common than endometrial carcinomas, and they have no known risk factors.

**29 C** Endometrial carcinomas can be associated with estrogen stimulation from anovulatory cycles, nulliparity, obesity, and exogenous estrogens (in higher amounts than found in birth control pills). These risks may initially give rise to endometrial hyperplasia that can progress to endometrial carcinoma if the estrogenic stimulation continues. Atypical endometrial hyperplasias progress to endometrial cancer in about 25% of cases. Adenomyosis increases the size of the uterus and is not a risk for endometrial carcinoma. Chronic endometritis and human papillomavirus infection (associated with squamous epithelial dysplasias and neoplasia) do not cause cancer.
30 C She has obesity, diabetes mellitus, and nulliparity — factors that contribute to development of endometrial hyperplasias and carcinomas caused by hyperestrinism. She has complex endometrial hyperplasia with atypicality of cells, which is a precursor for type I endometrial carcinoma. These lesions often have loss of PTEN tumor suppressor genes. In many if not all cancers, there is activation of aerobic glycolysis (i.e., glycolysis even in the presence of enough oxygen) — the so-called Warburg effect. This is linked to loss of PTEN and offers a growth advantage to tumor cells. When aerobic glycolysis is stimulated there is a reciprocal decrease in oxidative phosphorylation. Tumors are metabolically active, so glucose uptake and glycogen utilization is enhanced and not reduced. This uptake is the basis for positron emission tomography (PET) scans, where positron-emitting fluordeoxyglucose F 18 is preferentially taken up into foci of malignancy. In many cancers the COX-2 enzyme is up-regulated (e.g., colon cancer), and this leads to increased prostaglandins, but this is not related to PTEN loss.

PBD9 1013–1015  BP9 692–693  PBD8 1031–1032  BP8 725–727

31 C The masses shown are well circumscribed, suggesting the presence of multiple benign tumors. Leiomyomas (fibroids) can be present in one third to one half of all women. They tend to enlarge during the reproductive years, and then stop growing or involute after menopause. Although leiomyomas are often asymptomatic, leiomyomas that are submucosal may produce menometrorrhagia and chronic blood loss, leading to iron deficiency anemia. About 10% of complete moles are complicated by invasive mole, which is unlikely to produce a large, circumscribed mass. A leiomyosarcoma arises de novo, not from a leiomyoma, and is usually a larger, more irregular mass composed of more pleomorphic spindle cells with many mitoses. Decreased ovarian function after menopause accelerates bone loss, which may be severe enough to be termed osteoporosis, but this process is not related to female genital tract neoplasia. Preeclampsia with hypertension and proteinuria is associated with abnormal decidual vascularization and placental ischemia.

PBD9 1019–1020  BP9 693–694  PBD8 1036–1037  BP8 724–725

32 D Leiomyosarcomas arising in the uterine corpus account for about 5% of all GYN malignancies, and is most often present in postmenopausal women. The cellular atypia, coagulative necrosis, and numerous mitoses distinguish this neoplasm from the much more common leiomyoma (which does not give rise to leiomyosarcoma), both derived from smooth muscle. Anaplastic cytotothoblasts are seen with chorionicarcinomas. Cross striations are seen with rhabdomyosarcomas. Adenocarcinomas arise from glandular epithelium. Germ cells give rise to ovarian tumors such as teratoma and dysgerminoma. Squamous carcinomas are much more common but arise in the cervical portion of the uterus.

PBD9 1020–1021  BP9 694  PBD8 1037–1038  BP8 724–725

33 A Causes for postmenopausal uterine bleeding include endometrial atrophy, carcinoma, hyperplasia, and polyps. An early potentially curable endometrial carcinoma should not be missed. Even if the MRI is normal, a biopsy is still indicated. Infections are uncommon at this age and unlikely to cause bleeding. A Pap smear is insensitive for detection of endometrial lesions. She is postmenopausal and neither pregnancy nor gestational trophoblastic disease is probable.

PBD9 1009, 1018  BP9 690–692  PBD8 1027, 1034  BP8 723, 726

34 A Sexually transmitted diseases are the most common cause of inflammation of the fallopian tube. When the incidence of gonorrhea caused by Neisseria gonorrhoeae decreases in a population, the proportion of cases of salpingitis caused by Chlamydia and Mycoplasma increases. The fallopian tube can become distended and adherent to the ovary and may form a tubo-ovarian abscess. These are features of pelvic inflammatory disease. Haemophilus ducreyi causes chancroid, which can produce erythematous papules of the external genitalia or vagina, but grossly visible lesions may not be present in women. Herpes simplex virus most often involves the external genitalia, but it may produce vaginal or cervical lesions; it is unlikely to advance farther. Mycobacterium tuberculosis is an uncommon cause of salpingitis. Treponema pallidum infection causes syphilis, which does not produce florid inflammation with mass effect, just a chancre.

PBD9 1021  BP9 695  PBD8 1008, 1038  BP8 727–728

35 C Follicle cysts and lutein cysts of the ovary are so common that they are virtually normal findings and incidentalomas in diagnostic studies. Though most of them are less than 2 cm and asymptomatic, occasionally they can be larger (4 to 5 cm) and even enlarge a little more in response to midcycle hormones, occasionally rupturing to produce pain and bleeding. The negative pregnancy test helps to eliminate intrauterine or ectopic pregnancy. Endometriosis tends to produce more chronic pain, and though there is hemorrhage in the lesions, it tends to be contained within the lesions. The pregnancy test would be positive with an invasive mole, with uterine enlargement from the mass of grapelike villi. Pelvic inflammatory disease tends to produce chronic pain, and there is unlikely to be bleeding.

PBD9 1022  BP9 695–696  PBD8 1039  BP8 728

36 D Polycystic ovarian syndrome (PCOS) is a disorder of unknown origin that is typically associated with oligomenorrhea, obesity, and hirsutism. The MR image shows an enlarged ovary with multiple round cysts of increased signal intensity. It is thought to be caused by abnormal regulation of androgen synthesis. Teratomas are mass lesions that can be bilateral, but usually are not symmetric, and aside from struma ovarii not known for hormonal abnormalities. Krukenberg tumors represent metastatic disease involving the ovaries, usually from a primary site in the gastrointestinal tract, and are rare among patients of this age. Cystadenocarcinoma can be bilateral; however, androgen production by ovarian tumors is except by the Sertoli-Leydig cell tumors. Abscesses are usually unilateral and do not account for the hormonal changes seen in this patient.

PBD9 1022  BP9 695–696  PBD8 1039–1040  BP8 728
**37** B Cystadenocarcinomas are common ovarian tumors that are often bilateral. The serous type occurs more frequently than the mucinous type and is typically unilocular, whereas mucinous tumors are multilocular. Serous cystadenocarcinomas account for more than half of ovarian cancers. As the name indicates, they are cystic in appearance. They may be benign, borderline, or malignant. Benign tumors have a smooth cyst wall with small or absent papillary projections. Borderline tumors have increasing amounts of papillary projections. Endometrioid carcinomas resemble endometrial carcinomas and may arise in foci of endometriosis. Dysgerminomas are solid tumors of germ cell origin. Granulosa cell tumors can be solid and cystic and may produce estrogens. Mature cystic teratomas typically contain abundant hair and gooey sebaceous fluid within the cystic cavity; surrounding tissues are formed from various germ layers. Sertoli-Leydig cell tumors are rare, yellow-brown, solid masses; they may secrete androgens or estrogens.

PBD9 1023–1026  BP9 696–697  PBD8 1042–1044  BP8 730

**38** D Mucinous tumors of the ovary are of epithelial origin, are less common than serous tumors, and tend to be multiloculated. The appearance of ascites suggests metastases, which is most common with surface epithelial neoplasms of the ovary. Choriocarcinomas rarely reach this size because they metastasize early; they are typically hemorrhagic. Granulosa cell tumors and dysgerminomas tend to be solid masses. Teratomas are germ cell tumors differentiating into three germ layers; malignant transformation is rare, and is usually an element of squamous cell carcinoma from the ectodermal component.

PBD9 1026–1027  BP9 697–698  PBD8 1044–1045  BP8 730–731

**39** A Some familial cases of ovarian carcinoma (usually serous cystadenocarcinoma) are associated with the homozygous loss of the BRCA1 gene. This tumor-suppressor gene also plays a role in the development of familial breast cancers. Familial syndromes account for less than 5% of all ovarian cancers, however. The ERBB2 gene may be overexpressed in ovarian cancers; however, mutations of this gene do not give rise to familial tumors, and it is best known for an association with breast carcinomas. Mutations of the RAS and MYC oncogenes occur sporadically in many types of cancer. The RB1 gene, a tumor suppressor, can be involved in familial malignancies, including retinoblastoma and osteosarcoma.

PBD9 1024–1025  BP9 697  PBD8 1042  BP8 729–730

**40** D This is a borderline serous tumor of the ovary, and the figure shows a complex papillary projection into the cyst lumen. This is the most common serous ovarian tumor, and though most act in a benign fashion even when peritoneal implants are present, some tend to recur, particularly when KRAS or BRAF mutations are present, and the implants are invasive. Distant metastases are unlikely. Serous tumors do not have hormonal effects, either estrogenic to drive endometrial hyperplasia, or androgenic to drive masculinization. Ovarian carcinomas do not transform to sarcomas, and sarcomas at this site are rare.

PBD9 1023–1025  BP9 696–697  PBD8 1042–1043  BP8 729–730

**41** A A cystic tumor with a mass of hair in the lumen is the typical appearance of a mature cystic teratoma. This tumor also is known as a dermoid cyst because it is cystic and filled with hair and sebum derived from well-differentiated ectodermal structures. Teratomas with mature tissue elements are benign tumors of germ cell origin, and they can contain various ectodermal, endodermal, and mesodermally derived tissues. Papillary structures with psammoma bodies would characterize a cystadenocarcinoma. Primitive neuroepithelium in a more solid and less cystic mass would be consistent with an immature teratoma. Sarcomas of the ovary are uncommon; a rhabdomyosarcoma element could be part of a uterine malignant mixed Mullerian tumor. A choriocarcinoma with trophoblastic cells is usually gestational in origin and has a hemorrhagic appearance.

PBD9 1029–1030  BP9 698–700  PBD8 1047–1048  BP8 733

**42** D Immature teratomas are not cystic like mature teratomas. Tissues derived from multiple germ cell layers are present, as in all teratomas, but at least one immature tissue element is present. Often that immature element is neuroectodermal tissue. The less differentiated and more numerous the neuroepithelial elements, the higher the grade and the worse the prognosis. Adjuvant chemotherapy and radiotherapy yield a high response rate. Brenner tumors of the ovary are uncommon solid tumors that contain epithelial nests resembling transitional cells of the urinary tract; most are benign. Dysgerminomas are the female equivalent of male testicular seminomas. Granulosa cell tumors have cells that resemble those in ovarian follicles and may secrete estrogens. Leiomyosarcomas are solid tumors of smooth muscle origin that are found most often in the myometrium. Malignant Mullerian mixed tumors are typically uterine neoplasms that have glandular and stromal elements; the malignant stromal component can be heterologous and may resemble mesenchymal cells not ordinarily found in the myometrium, such as cartilage.

PBD9 1029–1030  BP9 700  PBD8 1048  BP8 733

**43** C Fibromas and thecomas are sex cord–stromal tumors that may be hormonally active and secrete estrogens that can lead to endometrial hyperplasia or even carcinoma. Fibromas can be associated with Peutz-Jeghers syndrome (ovarian tumor with ascites and right pleural effusion). Most of these tumors also are benign and do not metastasize. In most cases, chronic salpingitis is related to sexually transmitted infections, such as gonorrhea. A condyloma acuminatum is related to infection with human papillomavirus and is more likely to occur in younger, sexually active women on external genitalia and perineum. A partial mole is an uncommon form of gestational trophoblastic disease with a triploid karyotype and occurs only in reproductive-age women.

PBD9 1033  BP9 699  PBD8 1050–1051  BP8 732
CHAPTER 22  Female Genital Tract

44  E  The Sertoli cell group of ovarian neoplasms mimics testicular differentiation and may produce androgens. These neoplasms tend to be better differentiated and act in a more benign fashion. Brenner tumors are uncommon solid masses, usually act in a benign manner, and may be associated with endometrial hyperplasia, though they may not directly produce estrogenic hormones unless there are thecalike cells present. Dysgerminomas and endometrioid carcinomas tend not to produce hormonal effects. Granulosa-theca cell tumors are known for association with estrogenic effects.

PBD9 1033–1034  BP8 699  PBD8 1051–1052  BP8 732

45  F  Spontaneous abortion (miscarriage) may occur in at least a third of pregnancies, and most occur in the first trimester. Fetal problems are the most likely cause for early losses, whereas maternal problems account for most late fetal losses. Half of early abortuses have a chromosomal abnormality, many of which are incompatible with prolonged survival, such as trisomy 16. If there is recurrent early pregnancy loss, a parental germ line chromosomal anomaly may be suspected. Infections, uterine anomalies, masses such as leiomyomas, and toxemia are more likely to cause fetal loss later in pregnancy. Polycystic ovarian syndrome is more likely to be a cause for infertility. Maternal smoking is more likely to affect fetal weight, and less likely to cause early fetal loss.

PBD9 1035–1036  PBD8 1053

46  A  Placental infections are most likely to ascend from the vagina, and they are not usually homogenous. Preterm premature rupture of membranes may predispose to ascending infection, or it may be caused by prostaglandins released from acute inflammatory cells in the infection as suggested by the purulent exudate. Premature labor with delivery is likely to occur over the next 24 hours. Of the TORCH infections, the one most likely in this case is the O, including bacteria, such as group B streptococcus, whereas Listeria monocytogenes may produce more chronic inflammation.

PBD9 1036–1037  BP8 701  PBD8 1055  BP8 734

47  B  Conditions predisposing to ectopic pregnancy include chronic salpingitis (which may be caused by gonorrhea, but a culture would be positive only with acute infection), intrauterine tumors, and endometriosis. In about half of cases, there is no identifiable cause. Gestational trophoblastic disease associated with a triploid karyotype with partial mole developing outside the uterus is rare. Candida produces cervicitis and vaginitis and is rarely invasive or extensive in immunocompetent patients. Syphilis is not likely to produce a tubal mass with acute symptoms (a guma is a rare finding).

PBD9 1036  BP8 701  PBD8 1053–1054  BP8 734–735

48  D  Toxemia of pregnancy in this case is best classified as preeclampsia, because she has hypertension, proteinuria, and edema, but no seizures. The placenta tends to be small because of reduced maternal blood flow and uteroplacental insufficiency; infarcts and retroplacental hemorrhages can occur. Microscopically, the decidual arterioles may show acute atherosclerosis and fibrinoid necrosis. Chorioamnionitis is most often due to ascending bacterial infections and leads to, or follows, premature rupture of membranes. A chronic villitis is characteristic of a congenital infection such as cytomegalovirus. Placental hydrops often accompanies fetal hydrops in conditions such as infections and fetal anemias. In a partial mole, a fetus is present, but it is malformed and rarely live-born.

PBD9 1037–1039  BP8 703–704  PBD8 1055–1057  BP8 737–738

49  E  Classic features of eclampsia are defined by hypertension, edema, and proteinuria, typically with onset in the third trimester. The addition of seizures defines eclampsia. Primigravid women are at greater risk. There is no evidence in this case that primary renal disease could cause her hypertension, and the onset was sudden. Although the precise cause of preeclampsia/eclampsia is unknown, placental ischemia is believed to be the underlying mechanism. This is associated with shallow placenta and incomplete conversion of decidual vessels into high-volume channels required to perfuse the placenta adequately. Untreated patients may go on to disseminated intravascular coagulation. Cushing syndrome with adrenal cortical hyperplasia could lead to hypertension with sodium retention, but she does not have hypokalemia or hyperglycemia. Gestational trophoblastic disease predisposes patients to preeclampsia, but hydatidiform mole is excluded by the presence of a fetus, and a partial mole would be unlikely to persist into the third trimester. Functional ovarian tumors, most commonly estrogen secreting, such as granulosal cell tumor or thecoma, do not produce hypertension and proteinuria. Gestational diabetes may increase the risk for fetal loss, but in this case the glucose is normal.

PBD9 1037–1039  BP8 703–704  PBD8 1055–1057  BP8 737–738

50  D  The figure shows a hydatidiform mole, or complete mole, with enlarged, grapelike villi that form the tumor mass in the endometrial cavity. These trophoblastic tumors secrete large amounts of human chorionic gonadotropin (hCG). Moorelary pregnancies result from abnormal fertilization, with only paternal chromosomes present. Neural tube defects can be distinguished from other fetal defects (e.g., abdominal wall defects) by use of the acetycholinesterase test on amniotic fluid obtained by amniosentesis. If acetylcholinesterase and maternal serum α-fetoprotein are elevated, a neural tube defect is likely. If the acetylcholinesterase is not detectable, another fetal defect is suggested. α-Fetoprotein is a marker for some germ cell tumors that contain yolk sac elements. Estrogens can be elaborated by various ovarian stromal tumors, including thecomas and granulosa cell tumors. More ominously, a decrease in maternal serum estriol suggests incipient abortion. Human placental lactogen is produced in small quantities in the developing placenta, and serum levels typically are not measured.

PBD9 1039–1040  BP8 701–702  PBD8 1057–1059  BP8 735–736

51  E  Partial hydatidiform mole develops from triploidy (69 chromosomes). In contrast to a complete mole with only
paternal chromosomes, in which no fetus is present, a partial mole has a fetus because maternal chromosomes are present. Survival of the triploid fetus to term is rare. A partial mole may contain some grapelike villi, or none. The fetus is usually malformed, often with 3,4 syndactyly. A 46,XX karyotype could be present in a complete mole or a normal female fetus. A fetus with Turner syndrome (monosomy X) has a 45,X karyotype. Most female fetuses with loss of an X chromosome undergo spontaneous abortion, but some survive. Klinefelter syndrome has a 47,XXY karyotype, and male infants are live-born, with no placental problems. A 47,XY,+18 karyotype of trisomy 18 is associated with multiple congenital malformations, but not with a partial mole.

52 E Choriocarcinomas are aggressive, malignant trophoblastic tumors. Some of these tumors can arise without evidence of pregnancy. Metastases in the vaginal wall and lungs and a hemorrhagic appearance are characteristic. The large pleomorphic and hyperchromatic syncytiotrophoblastic cells produce human chorionic gonadotropin. Treatment with agents such as etoposide, methotrexate, actinomycin D, cyclophosphamide, and vincristine can often lead to remission and cure. Amnionic cells do not give rise to neoplasms. Rhabdomyoblasts are present in embryonal rhabdomyosarcomas of the vagina of young girls. Serous epithelium does not give rise to gestational trophoblastic disease. Smooth muscle cells give rise to leiomyomas, and rarely leiomyosarcomas, typically arising in the uterus.

53 D This is a placental site trophoblastic tumor (PSTT), the rarest of all forms of gestational trophoblastic disease. The intermediate trophoblastic cells do not produce large amounts of hCG, but do produce human placental lactogen (hPL). Most of these lesions are treated surgically and are controlled, but some recur and respond minimally to chemotherapy and radiation. α-Fetoprotein is produced by some testicular neoplasms and by hepatocellular carcinomas. Chromogranin is a marker for neuroendocrine tumors. Desmin is likely to be seen in tumors of mesenchymal origin. Neuron-specific enolase can be seen with tumors of neural and neuroendocrine differentiation.
1 A 21-year-old woman delivered a normal term infant a week ago and is now nursing the infant. She now notes a lump in her right axilla that has increased in size over the past week. On physical examination there is a rubbery, mobile, 1.5-cm mass beneath the skin at the right anterior axillary line. The mass is excised and the microscopic appearance is shown in the figure. Which of the following hormones most likely produced the greatest effect upon this tissue?

A Cortisol  
B Growth hormone  
C Oxytocin  
D Prolactin  
E Testosterone

2 A 24-year-old woman is breastfeeding 3 weeks after giving birth to a normal term infant. She notices fissures in the skin around her left nipple. Over the next 3 days, a 5-cm region near the nipple becomes erythematous and tender. Purulent exudate from a small abscess drains through a fissure. Which of the following organisms is most likely to be cultured from the exudate?

A Candida albicans  
B Lactobacillus acidophilus  
C Listeria monocytogenes  
D Staphylococcus aureus  
E Viridans streptococci

3 A 30-year-old woman sustained a traumatic blow to her right breast. Initially, there was a 3-cm contusion beneath the skin that resolved within 3 weeks, but she then felt a firm, painless lump that persisted below the site of the bruise 1 month later. What is the most likely diagnosis for this lump?

A Abscess  
B Fat necrosis  
C Fibroadenoma  
D Inflammatory carcinoma  
E Sclerosing adenosis

4 A study of mammographic findings on women of reproductive years is performed. The study identifies mammograms showing 1- to 5-cm cysts with focal microcalcifications and surrounding densities. Subsequent fine-needle aspiration yielded turbid fluid with few cells. Which of the following microscopic changes is most likely to be present in these lesions?

A Apocrine metaplasia  
B Ductal carcinoma in situ  
C Fat necrosis  
D Papillomatosis  
E Sclerosing adenosis
5 A 27-year-old woman feels a lump in her right breast. She has normal menstrual cycles, she is G3, P3, and her last child was born 5 years ago. On examination a 2-cm, irregular, firm area is palpated beneath the lateral edge of the areola. This lumpy area is not painful and is movable. There are no lesions of the overlying skin and no axillary lymphadenopathy. A biopsy specimen shows microscopic evidence of an increased number of dilated ducts surrounded by fibrous connective tissue. Fluid-filled ducts with apocrine metaplasia also are present. What is the most likely diagnosis?

A Fibroadenoma
B Fibrocystic changes
C Infiltrating ductal carcinoma
D Mammary duct ectasia
E Traumatic fat necrosis

6 A 47-year-old woman has a routine health examination. There are no remarkable findings except for a barely palpable mass in the right breast. A mammogram shows an irregular, 1.5-cm area of density with scattered microcalcifications in the upper outer quadrant. A biopsy specimen from this area is obtained and microscopically shows ductal hyperplasia. Which of the following is the most appropriate option for follow-up of this patient?

A Cessation of smoking cigarettes
B Continued screening for breast cancer
C Performing a simple mastectomy
D Testing for the BRCA1 oncogene
E Prescribing broad-spectrum antibiotic therapy

7 A 34-year-old woman has noticed a bloody discharge from the nipple of her left breast for the past 3 days. On physical examination, the skin of the breasts appears normal, and no masses are palpable. There is no axillary lymphadenopathy. She has regular menstrual cycles and is using oral contraceptives. Excisional biopsy is most likely to show which of the following lesions in her left breast?

A Acute mastitis
B Fibroadenoma
C Intraductal papilloma
D Phyllodes tumor
E Sclerosing adenosis

8 A 57-year-old man has developed bilateral breast enlargement over the past 2 years. On physical examination, the enlargement is symmetric and is not painful to palpation. There are no masses. He is not obese and is not taking any medications. Which of the following underlying conditions best accounts for his findings?

A ACTH-secreting pituitary adenoma
B Choriocarcinoma of the testis
C Chronic glomerulonephritis
D Diabetes mellitus
E Micronodular cirrhosis
F Rheumatoid arthritis

9 A 58-year-old woman sees her naturopathic health care provider for a routine health examination. There are no remarkable findings on physical examination. A screening mammogram shows a 0.5-cm irregular area of increased density with scattered microcalcifications in the upper outer quadrant of the left breast. Excisional biopsy shows atypical lobular hyperplasia. She has been on postmenopausal estrogen-progestrone therapy for the past 10 years. She has smoked 1 pack of cigarettes per day for the past 35 years. Which of the following is the most significant risk factor for the development of lobular carcinoma in patients with such lesions?

A Atypical cytologic changes
B History of smoking
C Hormone replacement therapy
D Postmenopausal age
E Underlying BRCA1 gene mutation

10 A 25-year-old Jewish woman sees her physician after finding a lump in her right breast. On physical examination, a 2-cm, firm, nonmovable mass is palpated in the upper outer quadrant. No overlying skin lesions and no axillary lymphadenopathy are present. The figure shows an excisional biopsy specimen. The family history indicates that the patient’s mother, maternal aunt, and maternal grandmother have had similar lesions. Her 18-year-old sister has asked a physician to determine whether she is genetically at risk of developing a similar disease. A mutated gene encoding for which of the following is most likely to be found in her sister?

A BRCA1
B Estrogen receptor (ER)
C HER2/neu
D TP53
E Progesterone receptor (PR)
F RB1

11 A clinical study is performed on postmenopausal women living in Paris, France, who are between the ages of 45 and 70 years. All have been diagnosed with infiltrating ductal carcinoma positive for estrogen receptor (ER) and progesterone receptor (PR), but negative for HER2 expression, which has been confirmed by biopsy and microscopic examination of tissue. None has the BRCA1 or BRCA2 mutation. Which of the following is most likely to indicate the highest relative risk of developing the carcinomas seen in this group of women?

A Age at menarche older than 16 years
B Age at menopause younger than 45 years
C First-degree relative with breast cancer
D Multiparity
E Prior diagnosis of mastitis
12 A 54-year-old woman feels a lump in her left breast. On examination there is a firm, irregular mass in the lower outer quadrant. A mammogram shows a 2-cm density with focal microcalcifications. Excisional biopsy shows intraductal and invasive carcinoma. Immunohistochemical staining is negative for estrogen receptor (ER). FISH analysis (green = HER2; red = chromosome 17 centromere) shows the findings in the figure. When combined with doxorubicin, which of the following drugs is most likely to be useful in treating this patient?

A Hydroxyurea  
B Letrozole  
C Raloxifene  
D Tamoxifen  
E Trastuzumab

13 A 66-year-old nulliparous woman received hormone replacement therapy for 7 years following menopause at age 53 years. Her BMI is 33. She now undergoes screening mammography, and an irregular mass is identified in the right breast. An excisional biopsy yields a 1.5-cm mass that microscopically has invasive cells that are positive for estrogen receptor but negative for HER2, with low proliferation markers and mutated PIK3CA gene. Following surgical removal of the mass, which of the following clinical courses will most likely occur over the next year?

A Detection of cancer in the left breast  
B Need for chemotherapy  
C Very low likelihood of recurrence  
D Need for treatment with trastuzumab  
E Occurrence of widespread metastases

14 A 63-year-old woman has a screening mammogram that shows an irregular density with microcalcifications. On physical examination, there are no lesions of the overlying skin, and there is no axillary lymphadenopathy. An excisional biopsy specimen shows no mass on sectioning. Microscopic examination shows the findings in the figure. What is the most likely diagnosis?

A Colloid carcinoma  
B Ductal carcinoma in situ  
C Infiltrating ductal carcinoma  
D Infiltrating lobular carcinoma  
E Medullary carcinoma  
F Papillary carcinoma

15 A 48-year-old woman has noticed a red, scaly area of skin on her left breast that has grown slightly larger over the past 4 months. On physical examination, there is a 1-cm area of eczematous skin adjacent to the areola. The figure shows the microscopic appearance of the skin biopsy specimen. What is the most likely diagnosis?

A Apocrine metaplasia  
B Fat necrosis  
C Inflammatory carcinoma  
D Lobular carcinoma in situ  
E Paget disease of the breast
A 54-year-old woman noticed a lump in her right breast. On examination, she has an ill-defined, 1-cm mass in the upper outer quadrant. The mass is cystic on ultrasound. An excision is done, and microscopically the mass shows predominantly fibrocystic changes, but the lesion shown in the figure also is present. Fine-needle aspirates of both breasts reveal additional foci of similar cells. Which of the following breast lesions is most likely to produce these findings?

A. Infiltrating ductal carcinoma
B. Lobular carcinoma in situ
C. Malignant phyllodes tumor
D. Medullary carcinoma
E. Mucinous (colloid) carcinoma

A 49-year-old woman felt a lump in her left breast 1 week ago. On examination, a firm, irregular mass is palpable in the upper outer quadrant of her left breast. There are no overlying skin lesions. The gross appearance of the excisional biopsy specimen is shown in the figure. Which of the following additional findings is she most likely to have on physical examination?

A. Axillary lymphadenopathy
B. Bloody discharge from the nipple
C. Chest wall tenderness
D. Cushingoid facies
E. Mass in the opposite breast

A 57-year-old woman has felt a lump in her left breast for 4 months. She has had new onset headaches associated with nausea for the past month. Her physician palpates a firm but irregular 2-cm mass in her left breast. CT imaging of her brain shows leptomeningeal enhancement. A lumpectomy with axillary node sampling is performed. Immunohistochemical staining of these cells shows absence of E-cadherin and HER2, but presence of estrogen receptor (ER) and progesterone receptor (PR). An H and E stained section is shown in the figure. No nodal metastases are present. Which of the following is the most likely diagnosis?

A. Lobular carcinoma
B. Medullary carcinoma
C. Metaplastic carcinoma
D. Metastatic glioblastoma
E. Phyllodes tumor

A 39-year-old woman has noticed an enlarging mass in her left breast for the past 2 years. The physician palpates a 4-cm firm mass. Following biopsy, a simple mastectomy is performed with axillary lymph node sampling. On gross sectioning, the mass has a soft, tan, fleshy surface. Histologically, the mass is composed of large cells with vesicular nuclei and prominent nucleoli. There is a marked lymphocytic infiltrate within the tumor, and the tumor has a discrete, noninfiltrative border. No axillary node metastases are present. The tumor cells are triple negative, for HER2, estrogen receptor (ER), and progesterone receptor (PR). What is the most likely diagnosis?

A. Colloid carcinoma
B. Infiltrating ductal carcinoma
C. Infiltrating lobular carcinoma
D. Medullary carcinoma
E. Papillary carcinoma
F. Phyllodes tumor

An epidemiologic study is conducted with male subjects who have been diagnosed with breast carcinoma. Their demographic data, medical histories, family histories, and laboratory data are examined to identify factors that increase the risk of cancer. Which of the following factors is most likely to be associated with the greatest number of male breast carcinomas?

A. Age older than 70 years
B. Asian ancestry
C. ATM gene mutation
D. Chronic alcoholism
E. Gynecomastia
21 A study of women with breast carcinoma is done to determine the presence and amount of estrogen receptor (ER) and progesterone receptor (PR) in the carcinoma cells. Large amounts of ER and PR are found in the carcinoma cells of some patients. These receptors are not present in the cells of other patients. The patients with positivity for ER and PR are likely to exhibit which of the following traits?
A Greater immunogenicity
B Greater likelihood of metastases
C Greater risk of familial breast cancer
D Higher response to therapy
E Higher tumor stage
F Higher tumor grade

22 A 26-year-old woman has felt a breast lump for the past month and is worried because she has a family history of early onset and bilateral breast cancers. On physical examination, there is a firm, 2-cm mass in the upper outer quadrant of her left breast. A biopsy is done, and the specimen microscopically shows carcinoma. Genetic analysis shows that she is a carrier of the BRCA1 gene mutation, as are her mother and sister. Which of the following histologic types of breast carcinoma has the highest incidence in families such as hers?
A Lobular carcinoma
B Medullary carcinoma
C Metaplastic carcinoma
D Papillary carcinoma
E Tubular carcinoma

23 A 79-year-old, previously healthy woman feels a lump in her right breast. The physician palpates a 2-cm firm mass in the upper outer quadrant. Nontender right axillary lymphadenopathy is present. A lumpectomy with axillary lymph node dissection is performed. Microscopic examination shows that the mass is an infiltrating ductal carcinoma. Two of 10 axillary nodes contain metastases. Flow cytometry on the carcinoma cells shows a small aneuploid peak and high S-phase. Immunohistochemical tests show that the tumor cells are positive for estrogen and progesterone receptor (ER/PR), negative for HER2/neu expression, and positive for cathepsin D expression. What is the most important prognostic factor for this patient?
A Age at diagnosis
B DNA content in the carcinoma
C Estrogen receptor positivity
D Expression of stromal proteases in the carcinoma
E Histologic subtype of carcinoma
F Lack of HER2/neu expression in the carcinoma
G Presence of lymph node metastases

24 A study of gene expression profiling involving breast biopsies showing invasive carcinoma of no specific type (NST) is performed. A subset of these cases, comprising about 15% of all cases, has the following characteristics: estrogen receptor (ER) and progesterone receptor (PR) negative, HER2/neu negative, basal keratin positive, flow cytometry showing aneuploidy and high proliferation rate, and association with BRCA1 mutations. Which of the following therapies is most likely to be effective in women with this subset of NST breast cancer?
A Chemotherapy
B Radiation
C Surgery alone
D Tamoxifen
E Trastuzumab

25 A 51-year-old woman has noticed an area of swelling with tenderness in her right breast that has worsened over the past 2 months. On physical examination, the 7-cm area of erythematous skin is tender with a rough, firm surface resembling an orange peel. There is swelling of the right breast, nipple retraction, and right axillary nontender lymphadenopathy. Excisional biopsy of skin and breast is most likely to show which of the following lesions?
A Acute mastitis
B Atypical epithelial hyperplasia
C Fat necrosis
D Infiltrating ductal carcinoma
E Sclerosing adenosis

26 A 26-year-old woman has noticed a lump in her right breast for the past year. A 2-cm, firm, circumscribed, movable mass is palpated in the lower outer quadrant. The figure shows the excised mass (A) and the mammogram (B). What is the most likely diagnosis?
A Fat necrosis
B Fibroadenoma
C Fibrocystic changes
D Infiltrating ductal carcinoma
E Mastitis
F Phyllodes tumor
A 27-year-old woman in the third trimester of her third pregnancy discovers a lump in her left breast. On physical examination, a 2-cm, discrete, freely movable mass beneath the nipple is palpable. After the birth of a term infant, the mass appears to decrease in size. The infant is breastfed without difficulty. What is the most likely diagnosis?

A Fibroadenoma  
B Intraductal papilloma  
C Lobular carcinoma in situ  
D Medullary carcinoma  
E Phyllodes tumor

A 24-year-old woman notes a lump in her right breast for the past month. She is concerned because her sister was diagnosed with a poorly differentiated “triple negative” breast cancer at age 31. Ultrasonography of the breast shows a solid mass. Fine needle aspiration is attempted but no diagnostic cells are obtained. Mammography is performed and there is a single 1-cm density with small clustered calcifications in the right breast but no lesions of the opposite breast. Which of the following is the best course of action for this patient?

A Biopsy to obtain tissue from the lesion  
B Continued monthly breast self-examination  
C Genetic testing for BRCA1 mutations  
D Hormonal therapy with tamoxifen  
E Radiologic imaging to detect metastases

A 48-year-old woman has felt a poorly defined lump in her right breast for the past year. On examination, she has a nontender, firm, 6-cm mass in the upper inner quadrant of her right breast. There are no lesions of the overlying skin and no axillary lymphadenopathy. A biopsy is performed, and microscopic examination of the specimen shows the findings in the figure. The mass is excised with a wide margin, but recurs 1 year later. After further excision, the lesion does not recur. What is the most likely diagnosis?

A Fibroadenoma  
B Fibrocystic changes  
C Lobular carcinoma  
D Phyllodes tumor  
E Tubular carcinoma

ANSWERS

1 D This is accessory breast tissue with lactational change. Prolactin secretion from the adenohypophysis increases in postpartum women to support milk production in breast lobules. Oxytocin released from the posterior pituitary stimulates myoepithelial cells to contract during nursing. The remaining hormones listed do not have a direct effect upon breast tissue. The presence of the breast tissue in the axilla represents accessory breast tissue, and can explain the origins of breast cancer in women following simple mastectomy.

2 D Staphylococcal acute mastitis typically produces localized abscesses, whereas streptococcal infections tend to spread throughout the breast, because streptococci often produce streptolysins. Acute mastitis can be associated with the first few months of breastfeeding. Candida may cause some local skin irritation, but is likely to become invasive only in immunosuppressed patients. Lactobacillus acidophilus is the organism used to produce fermented nonhuman milk. Listeriosis can be spread by contaminated food, including milk products, not by human milk.

3 B Fat necrosis is typically caused by trauma to the breast. The damaged, necrotic fat is phagocytosed by macrophages, which become lipid laden. The lesion resolves as a collagenous scar within weeks to months. The firm scar can mammographically and grossly resemble a carcinoma. An abscess may form a palpable but painful mass lesion, and often from Staphylococcus aureus infection when localized. A fibroadenoma is a neoplasm, and tumors are not induced by trauma. Inflammatory carcinoma refers to dermal lymphatic invasion by an underlying breast carcinoma, giving a rough red-to-orange appearance to the skin. Sclerosing adenosis is a feature of fibrocystic changes, a common cause of nontraumatic breast lumps.

4 A Nonproliferative cysts are quite common in the breast. When they are fluid-filled, they are unlikely to contain proliferative elements. The cells lining these cysts may be flattened cuboidal to atrophic, but often have abundant pink cytoplasm resembling apocrine change. Microcalcifications may be seen in both benign and malignant breast lesions, but in the case of cysts they represent calcified secretions. If excised, the intact cysts may have a blue to brown color. Ductal carci...
nomas are likely to be solid lesions. Fat necrosis may contain many macrophages, but also connective tissue, producing a firm lesion. Papillomatosis is a proliferative feature in fibrocystic changes that tends to form a solid lesion. Sclerosing adenosis produces a firm, fibrous lesion.

5 B Nonproliferative (fibrocystic) changes account for the largest category of breast lumps. These lesions are probably related to cyclic breast changes that occur during the menstrual cycle. In about 30% of cases of breast lumps, no specific pathologic diagnosis can be made. Fibrocystic changes include ductal proliferation, ductal dilation (sometimes with apocrine metaplasia), and fibrosis. A fibroadenoma is a discrete mass formed by a proliferation of fibrous stroma with compressed ductules. Carcinomas have proliferations of atypical neoplastic cells that fill ducts and can invade stroma. Inpsissated duct secretions may produce duct ectasia with a surrounding lymphoplasmacytic infiltrate. Trauma with subsequent fat necrosis may produce a localized, firm lesion that mimics carcinoma, but microscopically shows macrophages and neutrophils surrounding necrotic adipocytes, and healing leaves a fibrous scar.

6 B Fibrocystic changes without epithelial hyperplasia do not suggest a significantly increased risk of breast cancer. Moderate to florid hyperplasia increases the risk twofold, and atypical ductal or lobular hyperplasias increase the risk fivefold. The risk in this patient is not great enough to suggest radical or simple mastectomy at this time, but follow-up is needed. Breast cancers are not associated with tobacco use. The BRCA1 gene accounts for a small percentage of breast cancers, primarily in families in which cancer onset occurs at a young age, and genetic testing of all persons at risk for breast cancer is not warranted. These proliferative changes are not the result of infection.

7 C Intraductal papillomas are usually solitary and smaller than 1 cm. They are located in large lactiferous sinuses or large ducts, and have a tendency to bleed, though they are benign. Abscesses complicating mastitis organize with a fibrous wall. Fibroadenomas contain ducts with stroma and are not highly vascular; these lesions are not located in ducts. Phyllodes tumors also arise from intralobular stroma and can be malignant, but they do not invade ducts to cause bleeding. Sclerosing adenosis, a lesion occurring with fibrocystic changes, has abundant collagen, not vascularity.

8 E Micronodular cirrhosis is most often a consequence of chronic alcoholism and impairs hepatic estrogen metabolism, which can lead to bilateral gynecomastia. ACTH-secreting pituitary adenomas cause truncal obesity because of Cushing syndrome. Choriocarcinomas of the testis produce human chorionic gonadotropin and may cause some breast enlargement. Choriocarcinomas are highly malignant neoplasms that would not remain indolent for 2 years. Chronic renal failure is unlikely to have this consequence. Diabetes mellitus slightly increases the risk for breast cancer in women. Rheumatoid nodules can appear in various locations along with rheumatoid arthritis, but they rarely occur in the breast and are unlikely to be bilateral.

9 A Atypical lobular hyperplasia and atypical ductal hyperplasia increase the risk of breast cancer fivefold; the risk affects both breasts and is higher in premenopausal women or women who have a family history of breast cancer. Smoking and exogenous estrogen therapy are not well-established risk factors for breast cancer. The BRCA1 mutation accounts for about 10% to 20% of familial breast carcinomas and only a few percent of all breast cancers.

10 A The biopsy specimen shows an invasive breast cancer. Given the young age of the patient and the strong family history of breast cancer, it is reasonable to assume that she has inherited an altered gene that predisposes to breast cancer. There are two known breast cancer susceptibility genes: BRCA1 and BRCA2. Both are cancer suppressor genes. Specific mutations of BRCA1 are common in some ethnic groups, such as Ashkenazi Jews. Estrogen receptors are expressed in 50% to 75% of breast cancers. Their presence bodes well for therapy with hormone receptor antagonists. There is no known relationship between the structure of the estrogen receptor gene and susceptibility to breast cancer. Likewise, presence of progesterone receptors in the cancer cells indicates potential response to hormonal therapy, not risk for breast cancer. HER2/neu is a growth factor receptor gene that is amplified in certain breast cancers and is a marker of poor prognosis, not susceptibility. There is alteration of TP53 in many cancers, typically acquired and not familial, including breast carcinomas, but it does not have predictive value for risk. Inheritance of RB1 mutations increases the risk for retinoblastoma and osteosarcomas, but not breast carcinomas.

11 C The relative risk of breast cancer increases with various factors, but family history is one of the strongest. A history of bilateral breast disease and earlier age of onset of cancer increase the risk. The earlier age of onset increases the risk of identifying a BRCA1 or BRCA2 mutation. A longer reproductive life, with early menarche (<11 years old) and late menopause (>55 years old), and nulliparity increase the risk of breast cancer, probably because of increased estrogen exposure. “Soft” risk factors include exogenous estrogens and obesity. Mastitis does not affect the risk of breast cancer.

12 E The expression of HER2/neu, an epidermal growth factor receptor, suggests that biotherapy with trastuzumab may have some effectiveness. Drug names with the suffix -mab are monoclonal antibodies that target a specific
biochemical component of cells. This form of biotherapy is useful because normal breast cells do not express HER2/ neu. Doxorubicin is a standard chemotherapeutic agent that is part of various multiagent protocols. Hydroxyurea is a cycle-acting agent that is not useful in breast cancer. Letrozole is an aromatase inhibitor that is useful for treating ER-positive breast cancers. Raloxifene is a selective estrogen receptor modulator (SERM) that reduces risk for breast cancer and reduces osteoporosis. Tamoxifen is an antiestrogenic compound that has effectiveness in the treatment of breast cancers positive for ER/PR.

13 C This luminal A form comprises over half of all invasive breast cancers, and it tends to be low grade, and lack BRCA1, BRCA2, TP53, and CHEK2 familial gene mutations. It is often responsive to antiestrogen hormonal therapy, although surgery alone can be curative. Even with metastases, the course is prolonged. This patient has multiple risks for breast cancer, including nulliparity, obesity, and hormone replacement therapy. Breast cancer at her age is less likely to be familial. Trastuzumab is useful for HER2-positive breast cancers.

14 B An intraductal carcinoma, or ductal carcinoma in situ (DCIS), may not produce a palpable mass. The figure shows ducts that contain large, atypical cells in a cribriform pattern. If grossly soft, white material is extruded from small ducts when pressure is applied, then there is necrosis of the neoplastic cells in the ducts (that leads to dystrophic calcification), and the term comedocarcinoma is applicable. Intraductal carcinomas represent one fourth of all breast cancers. If not excised, such lesions become invasive. Intraductal carcinoma has several other histologic patterns, including noncomedo DCIS and Paget disease of the nipple, in which extension of the malignant cells to the skin of the nipple and areola produces the appearance of a seborrheic dermatitis. Colloid carcinomas occur about as frequently as medullary carcinomas, but they are often positive for estrogen receptor and progesterone receptor, and the prognosis is better than average. Infiltrating ductal carcinomas tend to produce irregular, firm, mass lesions because they are more invasive. Infiltrating lobular carcinomas can have a diffuse pattern without significant mass effect. Medullary carcinomas tend to be large masses; microscopically, they have nests of large cells with a surrounding lymphoid infiltrate. True papillary carcinomas are rare, although a papillary component may be present in other types of breast carcinoma.

15 E Paget cells are large cells that have clear, mucinous cytoplasm and infiltrate the skin overlying the breast. They are malignant and extend to the skin from an underlying breast carcinoma, which may be occult, so that Paget disease may be the first sign of malignancy. Apocrine metaplasia affects the cells lining the cystically dilated ducts in fibrocystic change. The macrophages in fat necrosis do not infiltrate the skin and do not have the atypical nuclei seen in the figure. Inflammatory carcinoma does not refer to a specific histologic type of breast cancer; rather, it describes the involvement of dermal lymphatics by infiltrating carcinoma, and there may be thickening and a reddish-orange appearance to the skin. In lobular carcinoma in situ, terminal ducts or acini within the breast are filled with neoplastic cells.

16 B Among primary malignancies of the breast, lobular carcinoma in situ (LCIS) is most likely to be bilateral. LCIS may precede invasive lesions by several years. Lobular carcinoma may be mixed with ductal carcinoma, and it may be difficult to distinguish them histologically. The other neoplasms listed are less likely to be bilateral and more likely to produce a single mass effect.

17 A This irregular, infiltrative mass is an infiltrating (invasive) ductal carcinoma, the most common form of breast cancer. Breast carcinomas are most likely to metastasize to regional lymph nodes. By the time a breast cancer becomes palpable, lymph node metastases are present in more than 50% of patients. A bloody discharge from the nipple most often results from an intraductal papilloma. Pain in the chest wall could be bone metastases, but less likely local invasion, and there is a margin of adipose tissue around the mass in the specimen shown. Breast cancers are associated in rare cases with ectopic corticotropin secretion or Cushing syndrome. Lobular carcinomas are more often bilateral, but they are less common than infiltrating ductal carcinomas.

18 A In this lobular carcinoma, note the pleomorphic cells infiltrating single file through the stroma. The metastatic profile of this cancer includes the carcinomatous meningitis suggested by her leptomeningeal enhancement, as well as intra-abdominal metastases. E-cadherin is an adhesion molecule that serves as a tumor suppressor, and its loss characterizes another infiltrating carcinoma—signet ring carcinoma of the stomach. Medullary carcinomas are solid masses of cells with little desmoplasia, but prominent lymphoid infiltrates. Metaplastic carcinomas are rare in humans and have a component resembling another tissue, such as squamous carcinomas. In the setting of a malignant breast mass, any brain lesion is a suspected metastatic lesion; although glioblastoma is capable of extracranial metastases, this is rare, and there should be a bulky cerebral mass present. Phyllodes tumors can be malignant, with a stromal component, but these are typically bulky masses, and there is a microscopic leaflike pattern of cystic spaces lined by epithelium.

19 D Medullary carcinomas account for about 1% to 5% of all breast carcinomas. They tend to occur in women at
youngster ages than do most other breast cancers. Despite poor prognostic indicators (such as absence of HER2, ER, and PR), medullary carcinomas have a better prognosis than most other breast cancers. Perhaps the infiltrating lymphocytes are a helpful immune response. Colloid carcinomas occur about as frequently as medullary carcinomas, but they are often positive for ER, and the prognosis is better than average. Infiltrating ductal and infiltrating lobular carcinomas tend not to produce large, localized lesions because they are more invasive, and they lack a distinct lymphoid infiltrate. True papillary carcinomas are quite rare, although other types of breast carcinoma may have a papillary component. The phyllodes tumor is typically large, but it has stromal and glandular components.

20 A Male breast cancers are rare, and they occur primarily among the elderly. Additional risk factors include first-degree relatives with breast cancer, decreased testicular function, exposure to exogenous estrogens, infertility, obesity, prior benign breast disease, exposure to ionizing radiation, and residency in Western countries. Of cases in men, 4% to 14% are attributed to germline BRCA2 mutations, less frequently for BRCA1, and ATM mutations in less than 1%. Gynecomastia does not seem to be a risk factor.

21 D The estrogen receptor and progesterone receptor (ER and PR) status helps predict whether chemotherapy with antiestrogen compounds such as tamoxifen would be effective; however, the correlation is not perfect. ER and PR do not affect immunogenicity and are not targets for immunotherapy. In contrast, immunotherapy targeted to the overexpressed HER2/neu gene is being used. The overall prognosis may be predicted from several factors, including histologic type, histologic grade, presence of metastases, degree of aneuploidy, and tumor stage. A family history and the presence of specific mutations such as BRCA1 or BRCA2 correlate with familial risk of breast cancer.

22 B Patients with the BRCA1 gene mutation have a high incidence of carcinomas with medullary features that are poorly differentiated and triple negative (do not express the HER2/neu protein, and are negative for estrogen and progesterone receptors).

23 G Many factors affect the course of breast cancer. The involvement of axillary lymph nodes is the most important prognostic factor listed. If there is no spread to axillary nodes, the 10-year survival rate is almost 80%. It decreases to 35% to 40% with 1 to 3 positive nodes, and to 15% with more than 10 positive nodes. Increasing age is a risk for breast cancer, but age alone does not indicate a prognosis, and treatment of cancers in the elderly can be successful. An increased DNA content with aneuploidy and a high S-phase suggests a worse prognosis, but staging is still a more important determinant of prognosis. Estrogen receptor positivity suggests a better response to hormonal manipulation of the tumor, whereas expression of HER2/neu suggests responsiveness to biotherapy with the monoclonal antibody trastuzumab. Some histologic types of breast cancer have a better prognosis than others, but staging is a more important factor than histologic type. The expression of stromal proteases, such as cathepsin D, predicts metastases, but in this case “the horse is out of the barn,” and metastasis has occurred.

24 A This is the basal-like subset of NST breast cancers that is triple negative for the usual immunohistochemical markers. Hence, lack of ER positivity predicts that antihormonal therapy with tamoxifen will not be of benefit, and lack of HER2/neu indicates that trastuzumab will be ineffective. The basal-like cancers are highly aggressive and tend to metastasize early, so containment with surgery or radiation is unlikely. However, some of them are cured by chemotherapy. This emphasizes the importance of gene expression profiling, so that treatment is individualized to each cancer patient for the best chance of success.

25 D The gross appearance of the skin is consistent with invasion of dermal lymphatics by carcinoma—the so-called inflammatory carcinoma, which is not a histologic type of breast cancer, but a descriptive phrase based upon the gross appearance (peau d’orange) resembling an inflammatory process. Nipple retraction and nontender axillary lymphadenopathy also suggest invasive ductal carcinoma. Atypical ductal hyperplasia may increase the risk of carcinoma, but it is not capable of invasion and does not produce visible surface skin changes. Acute mastitis may produce pain and swelling, but it is more likely to occur in association with breastfeeding, and as an inflammatory process would be more likely to produce painful lymphadenopathy. Fat necrosis on palpation can mimic that of carcinoma, but the skin is not involved. Sclerosing adenosis is a feature of benign fibrocystic changes producing breast lumps, but it has no skin involvement.

26 B Grossly and radiographically, this patient has a discrete mass that in a woman her age is most likely a fibroadenoma. Fat necrosis and infiltrating cancers are masses with irregular outlines. Fibrocystic changes are generally irregular lesions, not discrete masses. Mastitis has a more diffuse involvement, without mass effect. Phyllodes tumors are typically much larger and are far less common.

27 A Fibroadenomas are common and may enlarge during pregnancy or late in each menstrual cycle. Most intraductal
Papillomas are smaller than 1 cm and are not influenced by hormonal changes. Lobular carcinoma in situ is typically an ill-defined lesion without a mass effect. Medullary carcinomas tend to be large; they account for only about 1% of all breast carcinomas. Phyllodes tumors are uncommon and tend to be larger than 4 cm.

28 A Her age would suggest the lesion is probably benign, and even fibroadenomas and fibrocystic changes can have calcifications. The fibrous component of a fibroadenoma or fibrocystic changes can make it difficult to aspirate cells from them. However, the family history and the mammographic appearance of small clustered calcifications are concerning for carcinoma. A delay in diagnosis and treatment of breast cancer decreases survival. Although BRCA1 mutations are associated with HER2 and ER- and PR-negative breast cancers, the lesion must still be diagnosed. Based upon the histologic findings and molecular markers, a treatment plan can then be instituted that may include additional studies and pharmacologic therapies.

29 D Phyllodes tumors, although grossly and microscopically similar to fibroadenomas, occur at an older age, are larger, and are more cellular than fibroadenomas; they can recur locally following excision, but rarely metastasize. The figure shows cellular stroma protruding into spaces lined by a single layer of cuboidal epithelium. In contrast, fibrocystic changes can produce a breast lump, but usually not as large as 6 cm, and without firm areas of cellular stroma. A lobular carcinoma has malignant-appearing epithelial cells in clusters and rows and may not even produce a significant mass effect. Tubular carcinomas of the breast are uncommon, most are less than 1 cm in diameter, and most have small tubular structures in a noncellular stroma.
1. A 25-year-old woman has noted breast secretions for the past month. She is not breastfeeding and has never been pregnant. She has not menstruated for the past 5 months. Physical examination yields no abnormal findings. MRI of the brain shows a 0.7-cm mass within the sella turcica. Which of the following additional complications is she most likely to have?
   - A. Acromegaly
   - B. Cushing disease
   - C. Hyperthyroidism
   - D. Infertility
   - E. Neurologic dysfunction

2. A 20-year-old man’s closest friends tell him he looks different now than a year ago, with coarse features. He bought new shoes with his usual size and they do not fit. A year later his 23-year-old brother has similar problems. Both of them have hyperglycemia. Which of the following genetic alterations is most likely present in both brothers?
   - A. Germline mutation of AIP
   - B. Overexpression of cyclin D1
   - C. Fusion of PAX8-PPARG
   - D. Loss of function mutation in PTEN
   - E. Loss of VHL heterozygosity

3. A 39-year-old woman has had no menstrual periods for the past year, along with malaise, cold intolerance, and loss of body hair. She has had headaches for the past 5 months. On physical examination her lateral visual fields are reduced. She is most likely to have a neoplasm composed of which of the following cell types?
   - A. Beta
   - B. Chief
   - C. Chromaffin
   - D. Chromophobe
   - E. Glomerulosa
   - F. Parafollicular

4. A 41-year-old woman notices that her gloves from the previous winter no longer fit her hands. Her facial features have become coarse in the past year, and her voice seems deeper. On physical examination, her blood pressure is 140/90 mm Hg. There is decreased sensation to pinprick over the palms in the distribution of her thumb and first two fingers. A radiograph of the foot shows an increased amount of soft tissue beneath the calcaneus. A chest radiograph shows cardiomegaly. Laboratory studies indicate a fasting serum glucose level of 138 mg/dL and hemoglobin A1c level of 8.6%. Which of the following additional test results is most likely to indicate the cause of her physical and laboratory findings?
   - A. Abnormal glucose tolerance test result
   - B. Failure of growth hormone suppression
   - C. Hyperprolactinemia
   - D. Increased serum TSH level
   - E. Loss of diurnal serum cortisol levels

5. A 39-year-old G2, P2 woman, whose last pregnancy was 14 years ago, has had absent menstrual cycles for 6 months. She also reports expression of milk from her breasts. On physical examination, she is normotensive. She is 150 cm tall and weighs 63 kg (body mass index 28). Secondary sex characteristics are normal. Laboratory testing indicates that β-human chorionic gonadotropin level is normal. She has a normal growth hormone stimulation test. CT scan of the head shows no abnormalities of bone and no hemorrhage. Brain MRI shows fluid density within a normal-sized sella turcica. What is the most likely diagnosis?
   - A. Craniohypophysectomy
   - B. Empty sella syndrome
   - C. Hereditary hemochromatosis
   - D. Prader-Willi syndrome
   - E. Prolactinoma
   - F. Sheehan syndrome
6. A 21-year-old woman delivers a term infant after an uncomplicated pregnancy. The placenta cannot be delivered, however, and there is substantial hemorrhage, requiring transfusion of 10 U of packed RBCs. She must undergo a hysterectomy. Over the next 3 months, she is unable to produce sufficient milk to breastfeed her infant, and she becomes increasingly fatigued. Laboratory studies show Na⁺, 134 mmol/L; K⁺, 5.2 mmol/L; Cl⁻, 88 mmol/L; CO₂, 23 mmol/L; glucose, 59 mg/dL; calcium, 9.3 mg/dL; phosphorus, 3.5 mg/dL; and creatinine, 0.9 mg/dL. Over the next 5 months, her menstrual cycles do not return. Which of the following laboratory findings is now most likely to be reported in this woman?

A. Decreased corticotropin-releasing hormone
B. Decreased oxytocin
C. Failure of antidiuretic hormone release
D. Failure of growth hormone stimulation
E. Increased corticotropin
F. Increased dopamine

7. A 42-year-old man has had polyuria and polydipsia for the past 4 months. His medical history shows that he fell off a ladder and hit his head just before the onset of these problems. On physical examination, there are no specific findings. Laboratory findings include serum Na⁺, 155 mmol/L; K⁺, 3.9 mmol/L; Cl⁻, 111 mmol/L; CO₂, 27 mmol/L; glucose, 84 mg/dL; creatinine, 1 mg/dL; and osmolality, 350 mOsm/mL. The urine specific gravity is 1.002. This patient is most likely to have a deficiency of which of the following hormones?

A. Corticotropin
B. Melatonin
C. Oxytocin
D. Prolactin
E. Vasopressin

8. A 69-year-old man has become progressively obtunded over the past week. He has an 80 pack-year history of smoking cigarettes. A head CT scan shows no intracerebral hemorrhages. Laboratory findings include serum Na⁺ of 115 mmol/L, K⁺ of 4.2 mmol/L, Cl⁻ of 85 mmol/L, and bicarbonate of 23 mmol/L. The serum glucose is 80 mg/dL, urea nitrogen is 19 mg/dL, and creatinine is 1.7 mg/dL. Which of the following neoplasms is most likely to be present in this man?

A. Adenohypophyseal adenoma
B. Adrenocortical carcinoma
C. Pheochromocytoma
D. Small cell lung carcinoma
E. Renal cell carcinoma

9. A 23-year-old man has experienced headaches, polyuria, and visual problems for the past 3 months. On physical examination, he has bilateral temporal visual field deficits. CT scan of the head shows a large, partially calcified, cystic mass occupying the sellar and suprasellar areas. Laboratory findings show a serum prolactin concentration of 60 ng/mL and serum sodium level of 152 mEq/L. Serum calcium, phosphate, and glucose levels are normal. The mass is excised, and histologic examination shows a mixture of squamous epithelial elements and lipid-rich debris containing cholesterol crystals. Which of the following lesions is most consistent with the clinical and laboratory findings in this patient?

A. Craniopharyngioma
B. Metastases from a lung neoplasm
C. Multiple endocrine neoplasia type 1
D. Multiple endocrine neoplasia type 2
E. Prolactinoma

10. A 42-year-old woman has a sudden onset of fever with headache, nausea, diaphoresis, and palpitations. On physical examination her temperature is 39.2° C; pulse, 115/min; irregular respiratory rate, 30/min; and blood pressure, 150/85 mm Hg. Deep tendon reflexes are 4+ bilaterally. Her outstretched hands exhibit a high frequency tremor. Which of the following drugs should she receive emergently?

A. Aspirin
B. Hydrocortisone
C. Insulin
D. Propranolol
E. Tetracycline

11. A 47-year-old woman has had increasing fatigue with dyspnea and reduced exercise tolerance for the past year. On examination she has nonpitting edema of the lower extremities. Laboratory studies show a serum TSH level of 10 mU/L and T₄ level of 2 μg/dL. She is most likely to have pathologic findings affecting which of the following cells?

A. Hypophyseal basophils
B. Hypophyseal pituicytes
C. Hypothalamic glial cells
D. Hypothalamic neurons
E. Thyroid C cells
F. Thyroid follicular cells

12. A 2-year-old child has failure to thrive since infancy. Physical examination shows that the child is short and has coarse facial features, a protruding tongue, and an umbilical hernia. As the child matures, profound intellectual disability becomes apparent. A deficiency of which of the following hormones is most likely to explain these findings?

A. Cortisol
B. Insulin
C. Norepinephrine
D. Somatostatin
E. Thyroxine (T₄)
A 43-year-old woman has had increasing lethargy and weakness over the past 3 years. She has cold intolerance and wears a sweater in the summer. One year ago, she had menorrhagia, but now she has oligomenorrhea. She has difficulty concentrating, and her memory is poor. She has chronic constipation. On physical examination, her temperature is 35.5°C, pulse is 54/min, respirations are 13/min, and blood pressure is 110/70 mm Hg. She has alopecia, and her skin appears coarse and dry. Her face, hands, and feet appear puffy, with doughlike skin. Laboratory findings show hemoglobin, 13.8 g/dL; hematocrit, 41.5%; glucose, 73 mg/dL; and creatinine, 1.1 mg/dL. The representative microscopic appearance of her causative disease is shown in the figure. Which of the following serologic test findings is most likely to be positive in this woman?

A Anti-centromere antibody  
B Anti-double-stranded DNA antibody  
C Anti-Jo-1 antibody  
D Antimitochondrial antibody  
E Antiribonucleoprotein antibody  
F Anti-thyroid peroxidase antibody

A 37-year-old woman has had difficulty swallowing and a feeling of fullness in the anterior neck for the past week. She is recovering from a mild upper respiratory tract infection 1 month ago. On physical examination, her temperature is 37.4°C, pulse is 74/min, respirations are 16/min, and blood pressure is 122/80 mm Hg. Palpation of her diffusely enlarged thyroid elicits pain. Laboratory studies show an increased serum T4 level and a decreased TSH level. Two months later, she no longer has these complaints. The T4 level is now normal. Which of the following conditions is most likely to have produced these findings?

A Hashimoto thyroiditis  
B Medullary thyroid carcinoma  
C Subacute granulomatous thyroiditis  
D Toxic follicular adenoma  
E Toxic multinodular goiter

A 30-year-old woman has given birth to her second child. She develops heat intolerance and loses more weight than expected postpartum. On physical examination, her thyroid gland is enlarged but painless; there are no other remarkable findings. Laboratory studies show a serum T4 level of 12 μg/dL and a TSH level of 0.4 mU/L. A year later she is euthyroid. Which of the following is most indicative of the pathogenesis of this patient’s disease?

A Activational mutations in the RET proto-oncogene  
B Anti-thyroid peroxidase antibodies  
C Irradiation of the neck during childhood  
D Prolonged iodine deficiency  
E Recent viral upper respiratory tract infection

A 20-year-old woman and her twin sister both experience increasing diplopia. Their conditions develop within 3 years of each other. On physical examination, they have exophthalmos and weak extraocular muscle movement. The thyroid gland is diffusely enlarged but painless in each sister, and there is no lymphadenopathy in either woman. Which of the following serum laboratory findings is most likely to be reported in these sisters?

A Decreased free thyroxine level  
B Decreased thyroid-stimulating hormone level  
C High titer thyroid peroxidase autoantibodies  
D Increased thyrotropin-releasing hormone level  
E Increased triiodothyronine level

A 21-year-old woman has noted increasing fatigue and a 7-kg weight loss without dieting over the past 4 months. She also has increasing anxiety and nervousness with diarrhea. Physical examination shows a diffusely enlarged thyroid gland. Her temperature is 37.5°C, pulse is 103/min, respirations are 28/min, and blood pressure is 140/75 mmHg. A radionuclide scan of the thyroid shows a diffuse increase in uptake. The figure shows the representative microscopic appearance of the thyroid gland. What is most likely to produce these findings?

A Antibodies against TSH receptor  
B Dietary deficiency of iodine  
C Irradiation of the neck  
D Maternal deficiency in T4  
E Mutation in the RET proto-oncogene
A 40-year-old man notes weight loss, increased appetite, and double vision for 6 months. On physical examination, his temperature is 37.7°C, pulse is 106/min, respirations are 20/min, and blood pressure is 140/80 mm Hg. A fine tremor is observed in his outstretched hands. He has bilateral proptosis and corneal ulceration. Laboratory findings include a serum TSH level of 0.1 mU/L. A radioiodine scan indicates increased diffuse uptake throughout the thyroid. He receives propylthiouracil therapy, and his condition improves. Which of the following best describes the microscopic appearance of his thyroid gland before therapy?

A Destruction of follicles, lymphoid aggregates, and Hürthle cell metaplasia
B Enlarged thyroid follicles lined by flattened epithelial cells
C Follicular destruction with inflammatory infiltrates containing giant cells
D Nodules with nests of cells separated by hyaline stroma that stains with Congo red
E Papillary projections in thyroid follicles and lymphoid aggregates in the stroma

A 45-year-old woman from Kathmandu, Nepal, reports a feeling of fullness in her neck, but has no other concerns. The enlargement has been gradual and painless for more than 1 year. Physical examination confirms diffuse enlargement of the thyroid gland without any apparent masses or lymphadenopathy. Laboratory studies of thyroid function show a normal free T4 level and a slightly increased TSH level. What is the most likely cause of these findings?

A Diffuse nontoxic goiter
B Follicular adenoma
C Hashimoto thyroiditis
D Papillary carcinoma
E Subacute granulomatous thyroiditis
F Toxic multinodular goiter

A 14-year-old girl noticed gradual neck enlargement during the past 8 months. On physical examination her thyroid gland is diffusely enlarged. Her serum TSH level is normal. A dietary history is most likely to reveal that she has begun eating more of which of the following foods?

A Cabbage
B Fava beans
C Fish
D Plantains
E Rye bread

A 70-year-old man has had greater difficulty swallowing for the past 2 years. Over the past 6 months, he has lost 3 kg. On physical examination, his temperature is 37.3°C, and pulse is 102/min. There is fullness to the anterior neck, with a 5 × 10 cm irregular mass on palpation. Laboratory studies show serum TSH of 0.2 mU/L. A thyroid scintigraphic scan shows a 1.5-cm nodule with increased uptake in the right thyroid lobe, and decreased uptake into the remaining enlarged thyroid. What is the most likely diagnosis?

A follicular adenoma
B Graves disease
C Hashimoto thyroiditis
D Papillary carcinoma
E Toxic multinodular goiter

A 38-year-old woman felt a small lump on the right side of her neck 1 month ago, and it has not changed since then. Physical examination shows a 1-cm painless nodule palpable in the right lower pole of the thyroid gland. There is no lymphadenopathy. Radionuclide scanning shows that the nodule does not absorb radioactive iodine, and no other nodules are present. A fine-needle aspiration biopsy of the nodule is done, and the cytologic features are those of a follicular neoplasm. Which of the following laboratory findings is most likely to be present in this patient?

A Anti-TSH receptor immunoglobulins
B High free T4
C Low T3
D Normal TSH
E Anti-thyroid peroxidase antibodies

A 30-year-old woman has a 6-month history of weight loss (3 kg), hand tremors, mild watery diarrhea, and heat intolerance. On physical examination, vital signs are temperature, 37.3°C; pulse, 103/min with sinus rhythm; respirations, 20/min; and blood pressure, 125/85 mm Hg. She has a 1-cm firm, painless nodule palpable on the left side of her neck. There is no lymphadenopathy. No other abnormal findings are noted. Laboratory findings include a total serum T4 of 11.6 μg/dL with TSH of 0.2 mU/L. A scintigraphic scan shows more uptake of radioactive iodine into the nodule than the surrounding thyroid. A partial thyroidectomy is performed, and microscopic examination of the excised nodule shows well-differentiated thyroid follicles without vascular or capsular invasion. Molecular analysis of this nodule is most likely to reveal which of the following genetic changes?

A Activating missense mutation of GNAS1 gene
B Fusion gene formed by PAX8 and PPARG genes
C Gain of function mutation of TSH receptor gene
D Mutational activation of RET tyrosine kinase receptor
E Overexpression of cyclin D1 (CCND1) gene
F Stop codon mutation of autoimmune receptor (AIRE) gene

A 44-year-old, otherwise healthy woman feels a small lump on the left side of her neck. A firm, painless, 1.5-cm cervical lymph node is palpable. The thyroid gland is not enlarged. A chest radiograph is unremarkable. Laboratory findings include serum glucose, 83 mg/dL; creatinine, 1.2 mg/dL; calcium, 9.1 mg/dL; phosphorus, 3.3 mg/dL; thyroxine, 8.7 μg/dL; and TSH, 2.3 mU/L. The hemoglobin is 14 g/dL, platelet count is 240,400/mm³, and WBC count is 5830/mm³. A fine-needle aspiration biopsy of the thyroid gland is done. What is the most likely diagnosis?

A Anaplastic carcinoma
B Follicular carcinoma
C Medullary carcinoma
D Papillary carcinoma
E Parathyroid carcinoma
F Small lymphocytic lymphoma
25. A 45-year-old man has felt a lump on the left side of his neck for 4 months. Physical examination shows a nontender nodule on the left lobe of the thyroid gland. An adjacent cervical lymph node is enlarged and nontender. Laboratory studies show no thyroid autoantibodies in his serum, and the T4 and TSH levels are normal. A thyroidectomy is performed; the figure shows the microscopic appearance of the nodule. Which of the following etiologic factors is most likely to be involved in the pathogenesis of the thyroid nodule in this patient?

A. Autoimmunity  
B. Chronic dietary iodine deficiency  
C. Consumption of goitrogens  
D. RET gene mutation  
E. Viral infection

26. A 62-year-old woman has felt a lump on the right side of her neck for 5 months. On physical examination, she has no palpable lymphadenopathy. A fine-needle aspiration biopsy is done, and she undergoes a thyroidectomy. A 3-cm solid mass within the right thyroid lobe has the microscopic appearance shown in the figure. Six months later, she has pain in the right thigh. A radiograph shows a fracture of the right femur in an area of lytic bone destruction. A radioiodine scan shows uptake localized to the region of the fracture. Which of the following is the most likely diagnosis?

A. Papillary carcinoma  
B. Follicular carcinoma  
C. Granulomatous thyroiditis  
D. Hashimoto thyroiditis  
E. Non-Hodgkin lymphoma

27. A 44-year-old man with no previous illnesses has been bothered by progressive hoarseness, shortness of breath, and stridor for the past 3 weeks. On physical examination, he has a firm, large, tender mass involving the entire right thyroid lobe. CT scan shows extension of this mass posterior to the trachea and into the upper mediastinum. A fine-needle aspiration biopsy of the mass is done, and microscopically shows pleomorphic spindle cells. The mass is noted at surgery to have infiltrated the adjacent skeletal muscle. Four of seven cervical lymph nodes have metastases. Pulmonary metastases also are identified on a chest radiograph. Which of the following neoplasms is most likely to be present in this man?

A. Anaplastic carcinoma  
B. Follicular carcinoma  
C. Medullary carcinoma  
D. Non-Hodgkin lymphoma  
E. Papillary carcinoma

28. A 42-year-old woman has noted increasing fullness in her neck for the past 7 months. On physical examination, her thyroid gland is diffusely and asymmetrically enlarged and nodular. There is no lymphadenopathy. She undergoes thyroidectomy. Gross examination of the thyroid shows a multicentric thyroid neoplasm; microscopically, the neoplasm is composed of polygonal- to spindle-shaped cells forming nests and trabeculae. There is a prominent, pink hyaline stroma that stains positively with Congo red. Electron microscopy shows varying numbers of intracytoplasmic, membrane-bound, electron-dense granules. Immunohistochemical staining for which of the following antigens is most useful for diagnosis of this neoplasm?

A. Calcitonin  
B. CD3  
C. Cytokeratin  
D. Estrogen receptor  
E. Parathyroid hormone  
F. Vimentin

29. A 40-year-old man experiences weakness and easy fatigue of 2 months' duration. Physical examination yields no remarkable findings. Laboratory studies show serum calcium of 11.5 mg/dL, inorganic phosphorus of 2.1 mg/dL, and serum parathyroid hormone of 58 pg/mL, which is near the top of the reference range. A radionuclide bone scan fails to show any areas of increased uptake. What is the most likely cause of these findings?

A. Chronic renal failure  
B. Hypervitaminosis D  
C. Medullary thyroid carcinoma  
D. Parathyroid adenoma  
E. Parathyroid carcinoma  
F. Parathyroid hyperplasia
30. A 63-year-old woman had frequent headaches for 1 month. She now suddenly experiences a generalized seizure and becomes obtunded. She is taken to the emergency department, where a physical examination reveals an irregular heart rate. Laboratory findings include serum calcium of 15.4 mg/dL, serum phosphorus of 1.9 mg/dL, and albumin of 42 g/dL. A chest radiograph shows multiple lung masses and lytic lesions of the vertebral column. Which of the following conditions best accounts for these findings?

A. Chronic renal failure
B. Disseminated tuberculosis
C. Metastatic breast carcinoma
D. Parathyroid carcinoma
E. Vitamin D toxicity

31. A 40-year-old woman notes lethargy, weakness, and constipation for the past 6 months. On physical examination, she is afebrile and normotensive, and her heart rate is irregular. There is pain on palpation of the left third proximal finger. An ECG shows a prolonged QT (corrected) interval. Laboratory studies show glucose, 73 mg/dL; creatinine, 1.2 mg/dL; calcium, 11.6 mg/dL; phosphorus, 2.1 mg/dL; total protein, 7.1 g/dL; albumin, 5.3 g/dL; and alkaline phosphatase, 202 U/L. A radiograph of the left hand shows focal expansion by a cystic lesion of the third proximal phalanx. A technicium radionuclide scan shows a 1-cm area of increased uptake in the right lateral neck. A mutation in which of the following genes is most likely present in this woman?

A. GNAS1
B. MENI
C. TP53
D. RET
E. VHL

32. A 68-year-old man has experienced increasing malaise for 3 years. Physical examination shows no remarkable findings. Laboratory findings include a serum creatinine level of 4.9 mg/dL and urea nitrogen level of 45 mg/dL. Abdominal CT scan shows small kidneys. Which of the following endocrine glandular lesions has developed secondary to the underlying disease in this patient?

A. Adrenal atrophy
B. Insulin cell hyperplasia
C. Multinodular goiter
D. Parathyroid hyperplasia
E. Pituitary microadenoma

33. A 47-year-old woman noticed a lump in her neck 1 week ago. On physical examination, there is a 2-cm nodule in the right lobe of the thyroid gland. A fine-needle aspiration biopsy is performed, and microscopic examination of the specimen shows cells consistent with a follicular neoplasm. She undergoes a subtotal thyroidectomy. Which of the following laboratory tests should be performed on this patient in the immediate postoperative period?

A. Antithyroglobulin antibody
B. Calcitonin
C. Calcium
D. Parathyroid hormone
E. TSH

34. A 27-year-old man has controlled his diabetes mellitus for the past 10 years with insulin injections. This morning, his roommate is unable to awaken him. The man is unconscious when brought to the emergency department. On physical examination, his temperature is 37° C, pulse is 91/min, respirations are 30/min, and blood pressure is 90/65 mm Hg. Laboratory findings include a high plasma level of insulin and a lack of detectable C peptide. Urinalysis shows no blood, protein, or glucose, but 4+ ketonuria. Which of the following conditions is most likely to be present?

A. Acute myocardial infarction
B. Bacteremia
C. Hepatic failure
D. Hyperosmolar syndrome
E. Hypoglycemic coma
F. Ketoacidosis

35. Blood relatives of individuals diagnosed with type 1 or type 2 diabetes mellitus are studied for 10 years. Laboratory testing for glucose and insulin levels and autoantibody formation is performed on a periodic basis. The HLA types of the subjects are determined. A cohort of the subjects who are 8 to 22 years old has no overt clinical illnesses and no hyperglycemia; however, autoantibodies to glutamic acid decarboxylase are present. Many subjects in this cohort have the HLA-DR3 and HLA-DR4 alleles. Which of the following pancreatic abnormalities is most likely to be found in this cohort of study subjects?

A. Acinar acute inflammation and necrosis
B. Acinar fibrosis and fatty replacement
C. Islet amyloid deposition
D. Islet hyperplasia
E. Insulitis
F. Normal islets in a fibrous stroma

36. A 23-year-old woman has a routine health status examination. Her body mass index is 22. Laboratory studies show fasting plasma glucose is 130 mg/dL. Urinalysis shows mild glucosuria, but no ketonuria or proteinuria. She has no detectable insulin autoantibodies. Her father was similarly affected at age 20 years. She is most likely to have a mutation in a gene encoding for which of the following?

A. Glucagon
B. Glucokinase
C. GLUT4
D. Insulin
E. MHC DR

37. A 13-year-old girl collapses while playing basketball. On arrival at the emergency department, she is obtunded. On physical examination, she is hypotensive and tachycardic with deep, rapid, labored respirations. Laboratory studies show serum Na+, 151 mmol/L; K+, 4.6 mmol/L; Cl−, 98 mmol/L; CO2, 7 mmol/L; and glucose, 521 mg/dL. Urinalysis shows 4+ glucosuria and 4+ ketonuria levels, but no protein, blood, or nitrite. Which pathologic abnormality is most likely to be present in her pancreas at the time of her collapse?

A. Loss of islet beta cells
B. Acute inflammation of islets
C. Amyloid replacement of islet beta cells
D. Chronic inflammation of islets
E. Hyperplasia of alpha cells
F. Pancreatic neuroendocrine tumor
A study of patients more than 25 years of age with body mass index above 30, dyslipidemia, hypertension, and fasting glucose averaging 115 mg/dL is performed. They have adipose tissue abnormalities including increased nonesterified fatty acid release, altered adipokines with decreased adiponectin, greater proinflammatory cytokine release, and diminished peroxisome proliferator-activated receptor gamma (PPARγ) function. Which of the following is the best initial therapeutic intervention for these patients?

A. Adrenalectomy  
B. Caloric restriction  
C. Insulin injection  
D. l-Thyroxine  
E. Liposuction

An infant is born following premature delivery. Multiple external congenital anomalies are noted. The infant exhibits a seizure soon after birth. The blood glucose is 19 mg/dL. Which of the following maternal diseases is the most likely cause for the observed findings in this infant?

A. Cystic fibrosis  
B. Diabetes mellitus, type 2  
C. Gestational diabetes  
D. Maturity onset diabetes of the young  
E. Pancreatic neuroendocrine tumor

A clinical study is conducted in patients diagnosed with either type 1 or type 2 diabetes mellitus. Persons with either type develop complications of accelerated and advanced atherosclerosis. All untreated patients have an elevated hemoglobin A1c. Which of the following features common to patients with either type 1 or type 2 diabetes mellitus is most likely to be found by this study?

A. Association with certain MHC class II alleles  
B. High concordance rate in monozygotic twins  
C. Marked resistance to the action of insulin  
D. Nonenzymatic glycosylation of proteins  
E. Presence of islet cell antibodies

A 50-year-old man with fasting blood glucose >140 mg/dL on two occasions is put on a restricted caloric diet and started on a glucagon-like peptide-1 (GLP-1) receptor agonist. Which of the following laboratory studies is most likely to afford the best method of monitoring disease control in this man?

A. Cholesterol, total  
B. Fasting plasma glucose  
C. Glycosylated hemoglobin  
D. Microalbuminuria  
E. Random plasma glucose  
F. Serum fructosamine

A 50-year-old man has had a nonhealing ulcer on the bottom of his foot for 2 months. On examination, the 2-cm ulcer overlies the right first metatarsal head. There is reduced sensation to pinprick in his feet. His visual acuity is reduced bilaterally. Laboratory studies show serum creatinine is 2.9 mg/dL. Which of the following laboratory test findings is he most likely to have?

A. Glucosuria  
B. Hypoalbuminemia

A 52-year-old man has been concerned about a gradual weight gain over the past 30 years. He is 174 cm (5 feet 7 inches) tall and weighs 91 kg (body mass index 30). He is taking no medications. On physical examination, he has decreased sensation to pinprick and light touch over the lower extremities. Patellar reflexes are reduced. Motor strength seems to be normal in all extremities. Laboratory studies show blood glucose of 169 mg/dL, creatinine of 1.9 mg/dL, total cholesterol of 220 mg/dL, HDL cholesterol of 27 mg/dL, and triglycerides of 261 mg/dL. A chest radiograph shows mild cardiomegaly. Five years later, he has claudication in the lower extremities when he exercises. Based on these findings, which of the following complications is most likely to occur in this man?

A. Gangrene  
B. Hypoglycemic coma  
C. Ketoacidosis  
D. Mucormycosis  
E. Pancreatitis  
F. Systemic amyloidosis
A 50-year-old man has a 35-year history of diabetes mellitus. During this time, he has had hemoglobin A\textsubscript{1c} values between 7% and 10%. He now has problems with sexual function, including difficulty attaining an erection. He also is plagued by mild but recurrent low-volume diarrhea and difficulty with urination. He has delayed gastric emptying. These problems are most likely to originate from which of the following mechanisms of cellular injury?

A. Cross-linking of extracellular matrix proteins
B. Production of vascular endothelial growth factor
C. Abnormal transforming growth factor-beta signaling
D. Increased endothelial procoagulant activity
E. Nonenzymatic glycosylation
F. Polyol-induced susceptibility to oxidative stress

A 74-year-old woman is admitted to the hospital in an obtunded condition. Her temperature is 37° C, pulse is 95/min, respirations are 22/min, and blood pressure is 90/60 mm Hg. She appears dehydrated and has poor skin turgor. Her serum glucose level is 872 mg/dL. Urinalysis shows 4+ glucosuria, but no ketones, protein, or blood. Which of the following factors is most important in the pathogenesis of this patient’s condition?

A. Autoimmune insulitis
B. Glucokinase gene mutation
C. HLA-DR3/HLA-DR4 genotype
D. Peripheral insulin resistance
E. Virus-induced injury to beta cells in islets

A 40-year-old woman has experienced chest pain on exertion for the past 2 months. A month ago, she had pneumonia with \textit{Streptococcus pneumoniae} cultured from her sputum. On physical examination, she has a body mass index of 35. A random blood glucose value is 132 mg/dL. The next day, a fasting blood glucose is 120 mg/dL, followed by a value of 122 mg/dL on the following day. She is given an oral glucose tolerance test, and her blood glucose is 240 mg/dL 2 hours after receiving the standard 75-g glucose dose. On the basis of these findings, she is prescribed an oral thiazolidinedione (TZD) drug. After 2 months of therapy, her fasting blood glucose is 90 mg/dL. The beneficial effect of TZD in this patient is most likely related to which of the following processes?

A. Activation of PPAR\textgamma nuclear receptor in adipocytes
B. Decreased production of insulin autoantibodies
C. Greater density of insulin receptors in adipocytes
D. Increased half-life of circulating plasma insulin
E. Reduced secretion of glucagon by a cell in islets of Langerhans
F. Regeneration of beta cells in islets of Langerhans

A family is followed longitudinally for two generations. Four of eight children develop hyperglycemia by age 18 years. They are found to have serum islet autoantibodies. They have similar MHC I and MHC II loci. Treatment with insulin injections normalizes their Hgb A\textsubscript{1c} levels. Which of the following is the most likely mechanism leading to their disease?

A. Chloride ion channel abnormality
B. Chromosome 21 trisomy
C. Glucokinase gene mutation
D. Peripheral insulin resistance
E. Loss of T-cell tolerance

A 33-year-old woman has had several “fainting spells” over the past 6 months. Each time, she has a prodrome of light-headedness followed by a brief loss of consciousness. After each episode, she awakens and on examination has no loss of motor or sensory function. Physical examination after the current episode shows that she is afebrile, with a pulse of 72/min, respirations of 17/min, and blood pressure of 120/80 mm Hg. Imaging studies reveal a 0.5-cm lesion in the head of the pancreas. The microscopic appearance of this lesion is shown in the figure. Which of the following pancreatic disorders is most likely to be present in this patient?

A. Acute necrotizing pancreatitis
B. Adenocarcinoma
C. Cystic fibrosis
D. Neuroendocrine tumor
E. Pseudocyst

A 43-year-old man from Stockholm, Sweden, has had low-volume watery diarrhea for the past 3 months. He now has midepigastric pain. Over-the-counter antacid medications do not relieve the pain. On physical examination, he is afebrile; on palpation, there is no abdominal tenderness and no masses. An upper gastrointestinal endoscopy shows multiple 0.5- to 1.1-cm, shallow, sharply demarcated ulcerations in the first and second portions of the duodenum. He is given omeprazole. Three months later, repeat endoscopy shows that the ulcerations are still present. Which of the following analytes is most likely to be increased in his serum or plasma?

A. Gastrin
B. Glucagon
C. Insulin
D. Somatostatin
E. Vasoactive intestinal polypeptide (VIP)
**51** A 39-year-old man has had headache, weakness, and a 5-kg weight gain over the past 3 months. He has experienced mental depression during the same period. On physical examination, his face is puffy. His temperature is 36.9°C and blood pressure is 160/75 mm Hg. He has cutaneous striae over the lower abdomen and ecchymoses scattered over the extremities. A radiograph of the spine shows a compressed fracture of T11. Laboratory findings show fasting plasma glucose level of 200 mg/dL, serum Na⁺ of 150 mmol/L, and serum K⁺ of 3.1 mmol/L. The plasma cortisol level is 38 μg/dL at 8:00 am and 37 μg/dL at 6:00 pm. Administration of low and high doses of dexamethasone fails to suppress the plasma cortisol level and excretion of urinary 17-hydroxycorticosteroids. The plasma corticotropin level is 0.8 pg/mL. Which of the following lesions is most likely to be present in this man?

A. Adenohypophyseal adenoma
B. Adrenal cortical adenoma
C. Extra-adrenal pheochromocytoma
D. Small cell carcinoma of the lung
E. Thyroid medullary carcinoma

**52** A 43-year-old woman has had absent menstrual cycles along with increasing weakness and weight gain over the past 5 months. She notes low back pain for the past week. On physical examination, vital signs include blood pressure of 155/95 mm Hg. She has a prominent fat pad in the posterior neck and back. Facial plethora, hirsutism, and abdominal cutaneous striae are present. Laboratory findings include Na⁺, 139 mmol/L; K⁺, 4.1 mmol/L; Cl⁻, 96 mmol/L; CO₂, 23 mmol/L; glucose, 163 mg/dL; creatinine, 1.3 mg/dL; calcium, 8.9 mg/dL; and phosphorus, 4.1 mg/dL. Her serum ACTH level is low. A radiograph of the spine shows decreased bone density with a compression fracture at T9. Which of the following radiographic findings is most likely to be present in this patient?

A. Adrenal 10-cm solid mass with abdominal CT scan
B. Decreased radionuclide uptake in a thyroid gland nodule
C. Pulmonary 6-cm hilar mass on chest radiograph
D. Retroperitoneal 5-cm mass at the aortic bifurcation on pelvic MRI scan
E. Sella turcica enlargement with erosion on head CT scan

**53** A 73-year-old woman has experienced malaise and a 10-kg weight loss over the past 4 months. She has a 100 pack-year history of smoking cigarettes. Physical examination shows muscle wasting and 4/5 motor strength in all extremities. Abdominal CT scan shows bilaterally enlarged adrenal glands. A chest radiograph shows a 6-cm perihilar mass on the right and prominent hilar lymphadenopathy. Laboratory studies show Na⁺, 118 mmol/L; K⁺, 6 mmol/L; Cl⁻, 95 mmol/L; CO₂, 21 mmol/L; and glucose, 49 mg/dL. Her 8:00 am serum cortisol level is 9 ng/mL. What is the most likely diagnosis?

A. Amyloidosis
B. Ectopic corticotropin syndrome
C. Meningococcemia
D. Metastatic carcinoma
E. Pituitary adenoma

**54** A 28-year-old, otherwise healthy man has had headaches for the past 2 weeks. Physical examination yields no remarkable findings except for a blood pressure of 174/116 mm Hg. An abdominal CT scan shows an enlarged right adrenal gland. A right adrenalectomy is done; the figure shows the gross appearance of the specimen. Which of the following laboratory findings in his blood was most likely reported in this patient before surgery?

A. Hyperglycemia
B. Hyperkalemia
C. Hyponatremia
D. Low corticotropin level
E. Low insulin level
F. Low renin level

**55** A 40-year-old woman has experienced increasingly frequent episodes of weakness accompanied by numbness and tingling in her hands and feet for the past year. On examination her blood pressure is 168/112 mm Hg. Laboratory studies show sodium, 142 mmol/L; potassium, 2.9 mmol/L; chloride, 104 mmol/L; HCO₃⁻, 28 mmol/L; and glucose, 74 mg/dL. Her plasma renin activity is low. Which of the following radiologic findings is most likely to be present in this woman?

A. Adrenal nodular enlargement
B. Pancreatic mass
C. Retroperitoneal mass
D. Thyroid nodular enlargement
E. Mediastinal mass
A 5-year-old boy has developed features that suggest puberty over the past 6 months. On physical examination, the boy has secondary sex characteristics, including pubic hair and enlargement of the penis. Which of the following morphologic features is most likely to be seen in his adrenal glands?

A  Cortical atrophy  
B  Cortical hyperplasia  
C  Cortical nodule  
D  Medullary atrophy  
E  Medullary hyperplasia  
F  Medullary nodule

A female infant is born at term to a 41-year-old Yupik woman after an uncomplicated pregnancy. Soon after birth, the neonate develops hypotension. Physical examination shows ambiguous genitalia with a prominent clitoris. Labo-

ratory studies show Na⁺, 131 mmol/L; K⁺, 5.1 mmol/L; CO₂, 18 mmol/L; glucose, 65 mg/dL; creatinine, 0.4 mg/dL; testosterone, 50 mg/dL (normal <30 mg/dL); and cortisol, 2 μg/dL. An abdominal ultrasound scan shows bilaterally enlarged adrenal glands. Which of the following enzyme deficiencies is most likely to be present in this infant?

A  Aromatase  
B  11-Hydroxylase  
C  21-Hydroxylase  
D  17α-Hydroxylase  
E  Oxidase

A 37-year-old woman states that, although most of the time she feels fine, she has had episodes of palpitations, tachycardia, tremor, diaphoresis, and headache over the past 3 months. When her symptoms are worse, her blood pressure is measured in the range of 155/90 mm Hg. A chest radiograph shows no abnormal findings. Laboratory findings include serum Na⁺, 120 mmol/L; K⁺, 5.1 mmol/L; glucose, 58 mg/dL; urea nitrogen, 18 mg/dL; and creatinine, 0.8 mg/dL. The serum corticotropin level is 82 pg/mL. Which of the following is most likely to account for these findings?

A  Adenohypophyseal adenoma  
B  Autoimmune destruction of the adrenals  
C  Pancreatic neuroendocrine tumor  
D  Metastatic carcinoma with lung primary  
E  Neisseria meningitidis infection of the adrenals  
F  Sarcoidosis of the lung and adrenals

A 29-year-old woman with systemic lupus erythematosus has been treated with corticosteroid therapy for several years because of recurrent lupus nephritis. She undergoes an emergency appendectomy for acute appendicitis. On postoperative day 2, she becomes somnolent and develops severe hyperpigmentation of the skin. Vital signs include temperature of 36.9° C, pulse of 70/min, respirations of 14/min, and blood pressure of 95/65 mm Hg. Laboratory studies include a serum cortisol level of 3 μg/mL at 8:00 AM with a serum corticotropin level of 65 pg/mL. Which of the following diseases most often occurs in patients with this disorder?

A  Type 2 diabetes mellitus  
B  Classic polyarteritis nodosa  
C  Hashimoto thyroiditis  
D  Systemic lupus erythematosus  
E  Ulcerative colitis  
F  Sarcoidosis of the lung and adrenals

A 44-year-old woman has become increasingly listless and weak and has had chronic diarrhea and a 5-kg weight loss over the past 7 months. She also notices that her skin seems darker, although she rarely goes outside because she is too tired for outdoor activities. On physical examination, she is afibrile, and her blood pressure is 85/50 mm Hg. A chest radiograph shows no abnormal findings. Laboratory findings include serum Na⁺, 120 mmol/L; K⁺, 5.1 mmol/L; glucose, 58 mg/dL; urea nitrogen, 18 mg/dL; and creatinine, 0.8 mg/dL. The serum corticotropin level is 82 pg/mL. Which of the following is most likely to account for these findings?

A  Adenohypophyseal adenoma  
B  Autoimmune destruction of the adrenals  
C  Pancreatic neuroendocrine tumor  
D  Metastatic carcinoma with lung primary  
E  Neisseria meningitidis infection of the adrenals  
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A 5-year-old man has experienced increasing lethargy for the past 7 months. Physical examination shows hyperpigmentation of the skin. Vital signs include temperature of 36.9° C, pulse of 70/min, respirations of 14/min, and blood pressure of 95/65 mm Hg. Laboratory studies include a serum cortisol level of 3 μg/mL at 8:00 AM with a serum corticotropin level of 65 pg/mL. Which of the following diseases most often occurs in patients with this disorder?

A  Type 2 diabetes mellitus  
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C  Hashimoto thyroiditis  
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E  Ulcerative colitis  
F  Sarcoidosis of the lung and adrenals

A 19-year-old, previously healthy woman collapsed after complaining of a mild sore throat the previous day. On examination she is hypotensive and febrile with purpuric skin lesions. Her peripheral blood smear shows schistocytes. Imaging studies show her adrenal glands are enlarged, and there are extensive bilateral cortical hemorrhages. Infection with which of the following organisms best accounts for these findings?

A  Cytomegalovirus  
B  Histoplasma capsulatum  
C  Mycobacterium tuberculosis  
D  Neisseria meningitidis  
E  Streptococcus pneumoniae

A 5-year-old boy has developed features that suggest puberty over the past 6 months. On physical examination, the boy has secondary sex characteristics, including pubic hair and enlargement of the penis. Which of the following morphologic features is most likely to be seen in his adrenal glands?

A  Cortical atrophy  
B  Cortical hyperplasia  
C  Cortical nodule  
D  Medullary atrophy  
E  Medullary hyperplasia  
F  Medullary nodule

A 25-year-old woman gives birth to a term infant following an uncomplicated pregnancy. On physical examination, the newborn is found to have an enlarged abdomen, but there are no other abnormal findings except for slightly elevated blood pressure. An abdominal ultrasound scan shows a right retroperitoneal mass in the adrenal gland. An increase in which of the following substances is most likely to be associated with the lesion in this neonate?

A  Serum corticotropin (ACTH) level  
B  Serum cortisol level  
C  Serum growth hormone level  
D  Serum prolactin level  
E  Urinary free catecholamine level  
F  Urinary homovanillic acid (HVA) level

A 37-year-old woman states that, although most of the time she feels fine, she has had episodes of palpitations, tachycardia, tremor, diaphoresis, and headache over the past 3 months. When her symptoms are worse, her blood pressure is measured in the range of 155/90 mm Hg. A chest radiograph shows no abnormal findings. Laboratory findings include serum Na⁺, 120 mmol/L; K⁺, 5.1 mmol/L; glucose, 58 mg/dL; urea nitrogen, 18 mg/dL; and creatinine, 0.8 mg/dL. The serum corticotropin level is 82 pg/mL. Which of the following is most likely to account for these findings?

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A 44-year-old woman has become increasingly listless and weak and has had chronic diarrhea and a 5-kg weight loss over the past 7 months. She also notices that her skin seems darker, although she rarely goes outside because she is too tired for outdoor activities. On physical examination, she is afibrile, and her blood pressure is 85/50 mm Hg. A chest radiograph shows no abnormal findings. Laboratory findings include serum Na⁺, 120 mmol/L; K⁺, 5.1 mmol/L; glucose, 58 mg/dL; urea nitrogen, 18 mg/dL; and creatinine, 0.8 mg/dL. The serum corticotropin level is 82 pg/mL. Which of the following is most likely to account for these findings?

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B  Serum cortisol level  
C  Serum growth hormone level  
D  Serum prolactin level  
E  Urinary free catecholamine level  
F  Urinary homovanillic acid (HVA) level
A 27-year-old man has headaches that have occurred frequently for the past 3 months. On physical examination, he is afebrile, and his blood pressure is 140/85 mm Hg. There are no neurologic abnormalities and novisual defects; however, he has an enlarged thyroid. Laboratory studies show that his serum calcitonin level is elevated. A total thyroidectomy is performed, and on sectioning, the thyroid has multiple tumor nodules in both lobes. Microscopically, the thyroid nodules are composed of nests of neoplastic cells separated by amyl oid-rich stroma. The endocrinologist says that the patient’s family members could be at risk for development of similar tumors and advises that they undergo genetic screening. Which of the following morphologic findings in the adrenal glands is most likely to be present in this patient?

A  Bilateral 4-cm medullary masses  
B  Bilateral cortical atrophy  
C  Bilateral cortical nodular hyperplasia  
D  Solitary caseating granuloma  
E  Solitary 1-cm cortical mass with contralateral cortical atrophy  
F  Solitary 12-cm hemorrhagic cortical mass

A 10-year-old boy has been bothered by frequent headaches for the past 5 months. Physical examination yields no specific findings. Laboratory studies show normal electrolyte levels. CT scan of the head shows no bony abnormalities and no intracranial hemorrhage. MRI of the brain shows a 2-cm solid mass without calcifications or cystic change in the area inferior to the splenium of the corpus callosum, superior to the collicular plate, and between the right and left thalamic pulvinar regions. Because of the location, the mass is difficult to remove completely. Which of the following neoplasms is most likely to be present in this child?

A  Cranioopharyngioma  
B  Hypothalamic glioma  
C  Lymphoblastic lymphoma  
D  Metastatic carcinoma  
E  Pineoblastoma  
F  Prolactinoma

**ANSWERS**

1  **D** Prolactinomas are more common than other hormone-secreting pituitary adenomas. In addition to her galactorrhea and infertility, this patient also may have decreased libido, and her amenorrhea is secondary to the excessive prolactin secretion. Microadenomas might not have pressure effects on surrounding structures such as optic chiasm, but they can be discovered because of their hormonal effects. Acromegaly results from a growth hormone–secreting pituitary adenoma in an adult. Cushing disease occurs when there is an ACTH-secreting pituitary adenoma. A thyroid-stimulating hormone (TSH)–secreting pituitary adenoma is uncommon, but it could account for elevated levels of T4 and TSH in a hyperthyroid patient. Neurologic dysfunction with hyponatremia from the syndrome of inappropriate diuretic hormone leading to neurologic dysfunction is most often a paraneoplastic syndrome caused by a small cell anaplastic lung carcinoma, but may also be due to head trauma that damages the nerve tracts and neurohypophysis with uncontrolled release of vasopressin (antidiuretic hormone, ADH).

2  **A** The aryl hydrocarbon receptor–interacting protein (AIP) gene mutations account for a minority of growth hormone (GH)–producing pituitary adenomas, but may be present in younger persons in familial cases. GH opposes insulin, and excesses can lead to secondary diabetes mellitus. In adults with closed epiphyses, gigantism does not occur, but acromegaly of soft tissues does. Cyclin D1 mutations may be found in some follicular thyroid neoplasms. The PAX8-PPARG fusion gene may be found in some follicular thyroid neoplasms. The tumor suppressor gene PTEN may be mutated in some pancreatic neuroendocrine tumors. The VHL tumor suppressor gene may be associated with some pheochromocytomas.

3  **D** She has a nonfunctioning (null-cell) pituitary adenoma that has enlarged to compress and obliterate the normal adenohypophyseal cells resulting in hypopituitarism; the adenoma also presses on the optic chiasm, producing bitemporal hemianopsia. About a fourth of all pituitary tumors are nonfunctioning. Beta cells of the pancreatic islets of Langerhans produce insulin. Chief cells in the parathyroids produce parathormone that increases serum calcium. Chromaffin cells in the adrenal medulla produce catecholamines. Glomerulosa cells of the adrenal cortex produce corticosteroids. Parafollicular cells (C cells) of the thyroid interstitium produce calcitonin.
Failure to suppress growth hormone (GH) levels by glucose infusion suggests autonomous GH production. The patient’s symptoms suggest acromegaly, and a GH-secreting adenoma is most likely. Acromegaly causes an overall increase in soft tissue in adults because of the anabolic effects of the increase in GH. Because the epiphyses of the long bones are closed in adults, there is not the increase in height, or gigantism, that would be seen in children with a pituitary adenoma that is secreting excessive GH. Instead, the increase in soft tissue mass manifests as increasing shoe or glove size, carpal tunnel syndrome, and coarse facial features. This woman probably has an abnormal glucose tolerance test result, but this does not indicate the underlying cause of diabetes mellitus, which in this case is secondary to acromegaly. A prolactinoma would cause amenorrhea and galactorrhea in a woman. A thyroid-stimulating hormone (TSH)—secreting adenoma of the pituitary can give rise to hyperthyroidism with an increased metabolic rate that would most likely lead to weight loss, and glucose intolerance is not a feature of hyperthyroidism. Functional pituitary tumors can be detected clinically before they become large enough to cause pressure symptoms such as visual disturbances. Cushing syndrome from an adrenal cortical neoplasm producing cortisol could be accompanied by glucose intolerance, hypertension, and truncal obesity, but there is no overall increase in soft tissues.

Empty sella syndrome is a rare condition, seen most frequently in obese women, and results from herniation of the arachnoid through the diaphragma sellae. Although the increased pressure can lead to reduction in pituitary tissue through compression atrophy, there is typically adequate functional anterior pituitary to prevent hypopituitarism. This herniation can cause a “stalk section” effect, however, with loss of prolactin inhibition and hyperprolactinemia. A craniopharyngioma is a destructive tumor mass that is usually seen at a younger age. Hemochromatosis can interfere with organ function, including hypopituitarism; onset usually occurs later in women than in men (in the 60s in women compared with the 40s in men), owing to differences in physiologic iron losses (e.g., menstrual blood loss). Prader-Willi syndrome is an example of genomic imprinting with hypothalamic dysfunction seen in prepubertal boys. A prolactinoma could be a microadenoma, but MRI in this case rules this out because of the fluid density in the sella (seen with T2 weighting). If she had Sheehan syndrome after her pregnancy, she would have manifested hypopituitarism within months, not years.

Sheehan syndrome, or postpartum pituitary necrosis, is caused by pituitary enlargement during pregnancy, causing its blood supply to be more tenuous so that intrapartum hypotension with obstructive bleeding complications (e.g., the placenta accreta in this patient) predisposes to infarction. The anterior pituitary is at greater risk than the posterior pituitary. The laboratory findings in this patient suggest adrenal insufficiency, and her inability to breastfeed is caused by lack of prolactin; loss of menstrual cycles suggests that follicle-stimulating hormone and luteinizing hormone levels are deficient. If she were to have primary adrenal failure, the corticotropin (ACTH) level would be increased, but in her case, ACTH is low because of anterior pituitary failure. Because the hypothalamus is unaffected, corticotropin-releasing hormone would still be present. Oxytocin release from the posterior pituitary is probably not affected. She does not have diabetes insipidus from lack of antidiuretic hormone, because the posterior pituitary is less likely to be involved. Dopamine production in the hypothalamus is not affected.

Diabetes insipidus ensues from lack of antidiuretic hormone (ADH), also called arginine vasopressin. There is failure of resorption of free water in the renal collecting tubules—hence the increased dilute urine with higher serum osmolality and hypernatremia. Corticotropin stimulates the adrenal glands, mainly with the effect of increasing cortisol secretion. Oxytocin is involved in lactation. Prolactin and melatonin deficiencies have no identifiable specific clinical effects in men.

The syndrome of inappropriate antidiuretic hormone (SIADH) secretion results in increased free water resorption by the kidney and subsequent hyponatremia. SIADH is most often a paraneoplastic effect, and small cell (oat cell) anaplastic carcinoma of the lung (of neuroendocrine derivation and seen almost exclusively in smokers) is the most likely candidate among probable malignant neoplasms. Anterior pituitary adenomas do not produce antidiuretic hormone (ADH), which is released from the posterior pituitary. Adrenal cortical carcinomas can secrete cortisol or sex steroids, but not ADH. Pheochromocytomas secrete catecholamines. Renal cell carcinomas are known for various paraneoplastic effects, but SIADH is not a high probability.

Cranioopharyngiomas are uncommon, usually suprasellar neoplasms; they are typically found in young individuals. They are thought to arise from embryologic remnants of the Rathke pouch in the region of the pituitary. These are aggressive neoplasms that infiltrate and destroy surrounding tissues, making complete excision difficult. Despite their aggressive behavior, they are composed of benign-appearing squamous or primitive tooth structures. The increase in prolactin occurs as a “stalk section” effect, and the hypernatremia results from diabetes insipidus caused by destruction of the hypothalamus, posterior pituitary, or both. A metastasis to this location in a young individual is highly unlikely. Multiple endocrine neoplasia (MEN) type 1 includes pituitary adenomas, but not cranioopharyngiomas. MEN 2 does not involve the pituitary. Prolactinomas, similar to pituitary adenomas, can enlarge the sella when they are macroadenomas, but are not typically suprasellar or destructive of surrounding structures.
Thyroid storm is a medical emergency. There is not enough time to wait for confirmatory laboratory thyroid testing. There are increased catecholamine levels, and the β-blocker propranolol will help prevent emergent death from cardiac failure. Acetaminophen and ice packs are better choices to treat fever alone. Propylthiouracil (PTU) is the antithyroid medication with the fastest onset of action (hours), along with iodine to help reduce preformed thyroid hormone output. The fever and ancillary findings here go beyond what would be expected with an acute infection.

The normal feedback loop of peripheral thyroid hormones (T₃ and T₄) onto the basophils (thyrotrophs) of the adenohypophysis regulates TSH release (under tropic control of TRH from the hypothalamus). When patients with primary thyroid failure, the most common cause for hypothyroidism with myxedema in adults, do not have sufficient residual functioning thyroid follicular cells producing thyroid hormones, then the TSH will rise, as in this case, in conjunction with a low T₄ level. The levels of TRH from the hypothalamus are much lower and harder to measure for correlation with thyroid gland function. The measurement of TSH is also the most useful screening test for hyperthyroidism. Neurohypophyseal axons release ADH and oxytocin produced in the hypothalamus, whereas modified glial cells called pituicytes do not release hormones. Thyroid parafollicular, or “C” cells, produce calcitonin.

Cretinism is a condition that is uncommon whenever routine newborn screening is available for testing and treatment at birth for hypothyroidism. Hypothyroidism that develops in older children and adults is known as myxedema. A lack of cortisol from primary adrenal failure leads to Addison disease, or a 21-hydroxylase deficiency could produce congenital adrenal hyperplasia. An absolute deficiency of insulin leads to type 1 diabetes mellitus, but this is more likely to develop in childhood or later, and there would be weight loss. There is no deficiency state caused by a lack of norepinephrine or somatostatin.

The lymphoid follicles and the large, pink nodules of Hürthle cells in this photomicrograph are typical for Hashimoto thyroiditis. The anti–thyroid peroxidase (antimicrosomal) and antithyroglobulin antibody titers typically are increased in patients with Hashimoto thyroiditis when thyroid enlargement is still present. In the later, “burnt-out” phase of Hashimoto thyroiditis, the antibodies are sometimes undetectable—only the hypothyroidism is. The thyroid-stimulating hormone (TSH) level is an indication of whether there is a primary disease in the thyroid. If the patient appears hypothyroid and a primary thyroid disease (e.g., Hashimoto thyroiditis) is suspected, the TSH level is elevated. If the patient appears hyperthyroid and a primary thyroid disease (e.g., Graves disease) is suspected, the TSH level is decreased. Anticentromere antibody is characteristic of limited scleroderma (CREST syndrome). Anti–double-stranded DNA antibody is very specific for systemic lupus erythematosus. Anti-Jo-1 antibody can be seen in polymyositis. Increased antimitochondrial antibody indicates primary biliary cirrhosis. Antiribonucleoprotein antibodies are seen in some collagen vascular diseases, such as mixed connective tissue disease.

Subacute granulomatous thyroiditis (de Quervain thyroiditis) is a self-limited condition that can be of viral origin because many cases are preceded by an upper respiratory infection. The transient hyperthyroidism results from inflammatory destruction of the thyroid follicles and release of thyroid hormone. The released colloid acts as a foreign body, producing florid granulomatous inflammation in the thyroid. Hashimoto thyroiditis can enlarge the thyroid transiently, but there is usually no pain or hyperthyroidism. Thyroid neoplasms are not typically associated with signs and symptoms of inflammation and are rarely functional. A toxic multinodular goiter likewise produces no signs of inflammation, and does not reverse functionality.

The presence of autoantibodies in the serum in this patient with transient hyperthyroidism would suggest Hashimoto thyroiditis (“hashitoxicosis”), but the variant called subacute lymphocytic painless thyroiditis may affect 1 in 20 postpartum women, and a minority progress to hypothyroidism. Mutations in the RET proto-oncogene are associated with papillary carcinoma and medullary carcinoma of the thyroid. Irradiation of the thyroid gland can give rise to hypothyroidism, but it is unlikely that irradiation in childhood would give rise to hypothyroidism at age 60 years. Irradiation also can predispose to the development of papillary carcinoma, but these tumors usually do not affect thyroid hormone secretion. An iodine deficiency can lead to hypothyroidism, but a goiter would be present. A history of a viral infection sometimes precedes subacute granulomatous thyroiditis, which is typically a self-limited disease that lasts for weeks to 2 months.

Exophthalmos is a feature seen in about 40% of individuals with Graves disease. The hyperfunctioning thyroid gland leads to an increased T₄ level, with positive feedback from the pituitary to decrease thyroid-stimulating hormone (TSH) secretion. There is about 50% concordance of Graves disease among identical twins. The autoimmune character of this disorder is evidenced by an association with HLA-DR3 and by the presence of an autoantibody against HLA-DR3 that activates T₈ secretion. An increased thyrotropin-releasing hormone (TRH) level would increase the TSH level and increase the T₄, but feedback typically occurs at the level of the pituitary and the hypothalamus, and abnormal increases in TRH are uncommon. Anti–thyroid peroxidase antibodies
can be seen in Hashimoto thyroiditis and in Graves disease, but the highest titers occur in Hashimoto thyroiditis. T₃ levels are less likely to be elevated than T₄ levels. PBD9 1089–1090  BP9 726–727  PBD8 1114–1115  BP8 763–764

17 A The tall columnar epithelium with papillary infoldings and scalloping of the colloid is characteristic of Graves disease, which leads to hyperthyroidism. This disease is caused by autoantibodies that bind to the thyroid-stimulating hormone (TSH) receptor and mimic the action of TSH. Dietary iodine deficiency can cause diffuse compensatory enlargement of the thyroid, but it does not cause hyperthyroidism. Irradiation of the neck is a predisposing factor for papillary carcinoma of the thyroid. Maternal thyroid hormone deficiency may affect childhood mental development. Mutations in the RET proto-oncogene occur in papillary carcinoma of the thyroid and in medullary carcinomas of the thyroid. These neoplasms do not cause diffuse enlargement of the gland, hyperthyroidism, or diffuse increase in radioiodine uptake. Tumors usually produce “cold nodules” on radioiodine scans. PBD9 1089–1090  BP9 726–727  PBD8 1114–1115  BP8 763–764

18 E The clinical findings in this case point to hyperthyroidism, and the increased, diffuse uptake corroborates Graves disease as a probable cause because the thyroid-stimulating hormone (TSH) level is quite low. The thyroid-stimulating immunoglobulins that appear in this autoimmune condition result in diffuse thyroid enlargement and hyperfunction, and papillary projections lined by tall columnar epithelial cells. Destruction of thyroid follicles with lymphoid aggregates and Hürthle cell metaplasia is characteristic of Hashimoto thyroiditis. A goiter has enlarged follicles and flattened epithelial cells; most of these patients are euthyroid. Follicular destruction and the presence of giant cells occur in granulomatous thyroiditis. Nests of cells in a Congo red–positive hyaline stroma characterize a medullary carcinoma, which can be multifocal, but is not diffuse and does not lead to hyperthyroidism. PBD9 1089–1090  BP9 726–727  PBD8 1114–1116  BP8 758–759

19 A Diffuse nontoxic goiter is most often caused by dietary iodine deficiency. This condition is endemic in regions of the world where there is a deficiency of iodine (e.g., inland mountainous areas); it also may occur sporadically. As in this case, patients are typically euthyroid. A follicular adenoma rarely functions to produce excess thyroid hormone; most are “cold,” nonfunctioning nodules that do not involve the thyroid diffusely. A chronic lymphocytic thyroiditis, such as Hashimoto thyroiditis, initially can produce thyroid enlargement, but atrophy eventually occurs with resulting hypothyroidism. Papillary carcinomas most often produce a mass effect or metastases and do not affect thyroid function. Subacute granulomatous thyroiditis can lead to diffuse enlargement, and transient hyperthyroidism can occur, but the disease typically runs a course of no more than 6 to 8 weeks. Plummer disease, or toxic multinodular goiter, occurs when there is a hyperfunctioning nodule in a goiter. PBD9 1090–1091  BP9 728–729  PBD8 1116  BP8 764–765

20 A She has developed a sporadic goiter. Vegetables of the Brassicaceae family, including cabbage, turnips, and Brussels sprouts, contain glucosinolate, which can decompose to release thiocyanate, a byproduct that interferes with thyroid hormone synthesis. Thus such substances are known as goitrogens. Young persons with increased demand for thyroid hormone are at increased risk. Fava beans contain oxidizing agents that incite hemolysis in persons with glucose-6-phosphate dehydrogenase (G6PD) deficiency. In addition, beans contain long-chain sugars that are indigestible with human intestinal enzymes, leaving them to be fermented by colonic bacteria that release gas (flatus). Fish have omega-3 fatty acids that protect against atherogenesis. Plantains are starchy, as anyone mistaking them for sweet bananas soon discovers; rare food allergy develops to them. Fungus growing on moldy rye produces ergot poisoning. PBD9 1091  BP9 728  PBD8 1116–1117  BP8 764–765

21 E A long-standing diffuse goiter can evolve into a multinodular goiter, and one of the nodules can begin hyperfunctioning to cause so-called Plummer disease. This “toxic” nodule has acquired growth and functional characteristics similar to a benign neoplasm, such as a follicular adenoma, but one that is functional. Rare toxic follicular adenomas can function and produce “hot” nodules, but the remaining gland is often atrophic, not enlarged. In Graves disease, the thyroid is enlarged, but usually diffusely, without pronounced nodularity, so that there is increased uptake into the entire gland. In addition, clinical features such as dermopathy and ophthalmopathy that are lacking with Plummer disease are associated with Graves disease. There may be initial diffuse thyroid enlargement with Hashimoto thyroiditis and transient hyperfunction, but over time the thyroid atrophies, and hypothyroidism ensues. It is extremely rare for a papillary carcinoma to function, and although this would be a hot nodule, the remaining thyroid would not be enlarged. PBD9 1091–1092  BP9 728–729  PBD8 1116–1118  BP8 764–765

22 D Solitary “cold” and solid (not cystic) thyroid nodules are likely to be neoplastic, and most are benign follicular adenomas that do not affect thyroid function. If the nodule was hyperfunctioning, and produced hyperthyroidism, it would appear “hot” on the scan, and suppress thyroid-stimulating hormone (TSH). Anti-TSH receptor immunoglobulins can be seen in Graves disease, as may high T₄ and low TSH levels, but this is a diffuse disease of the thyroid. T₃ levels are rarely measured. Antimicosomal and antithyroglobulin antibodies are seen in Hashimoto thyroiditis (and Graves disease), but thyroiditis is a diffuse process and unlikely to produce a solitary nodule. As Hashimoto thyroiditis progresses, decreasing function of the thyroid can lead to decreasing T₄ level and increasing TSH level, typical of primary thyroid failure. PBD9 1092–1094  BP9 729–730  PBD8 1118–1119  BP8 767

23 C A thyroid nodule that is well differentiated, noninvasive, and hyperfunctioning suggests a “toxic” follicular adenoma, and a large proportion of these adenomas have
activating mutations of the thyroid-stimulating hormone (TSH) receptor–signaling pathway involving the TSH receptor or associated G protein. Mutation of the GNAS1 gene is seen in McCune-Albright syndrome and some pituitary adenomas. A PAX8-PPARG fusion gene is found in certain follicular carcinomas of the thyroid. Activation of the RET gene occurs in papillary thyroid carcinomas. Overexpression of the cyclin D1 gene (CCND1) is characteristic of parathyroid adenomas. The AIRE gene regulates expression of self-antigens in the thymus, and mutation in this gene causes autoimmune polyendocrinopathy affecting adrenals, parathyroids, gonads, and gastric parietal cells with destruction of these tissues and consequent hypofunction.

PBD9 1093  PBD8 1116–1118  BP8 729–730

24 D Papillary thyroid carcinomas may present initially with metastases, and local lymph nodes are the most common sites for metastases. The primary site may not be detectable as a palpable nodule. Papillary carcinoma is the most common thyroid malignancy. Metastases to the thyroid are uncommon. Anaplastic carcinomas are uncommon, but are very aggressive, locally invasive lesions. Follicular and medullary carcinomas tend to spread hematogenously. A parathyroid carcinoma is a locally infiltrative mass, and the serum calcium level is usually quite high in such patients. A small lymphocytic lymphoma, which is the tissue form of chronic lymphocytic leukemia, is uncommon at this patient’s age, usually involves multiple nodes, and is accompanied by an elevated WBC count.

PBD9 1094–1097  BP9 730–733  PBD8 1121–1122  BP8 768–769

25 D The papillary architecture in this nodule, with cells that have clear nuclei, is a pattern typical for papillary carcinoma. There is no such thing as a papillary adenoma. These nuclear changes, even if the pattern is follicular, confirm the diagnosis of papillary carcinoma. About 30% of all papillary thyroid carcinomas have mutational activation of RET or NTRK1 proto-oncogenes, which belong to the family of receptor tyrosine kinases that transduce extracellular signals for cell growth and differentiation and exert many of their downstream effects through the ubiquitous MAP kinase signaling pathway. Iodine deficiency gives rise to uniform thyroid enlargement because the secretion of thyroid-stimulating hormone (TSH) increases when there is reduced synthesis of T4. Autoimmunity plays a role in Hashimoto thyroiditis and Graves disease, and these do not progress to carcinoma, although non-Hodgkin lymphoma may develop in the former. Goitrogens interfere with thyroid hormone synthesis and have an effect similar to that of iodine deficiency, with potential hypothyroidism, but not malignancy. Viral infections can cause a subacute thyroiditis, not carcinoma.


26 B Follicular carcinoma can be difficult to differentiate from follicular adenoma, unless there is microscopic evidence of invasion, as shown in the figure, so the term follicular neoplasm may be given for a fine-needle aspiration specimen. Follicular carcinomas are often indolent, but they can metastasize, and then are easily distinguished from follicular adenomas. Follicular carcinomas are much less likely than papillary carcinomas to involve lymph nodes, but they are more likely to metastasize to distant sites, such as bone, lung, and liver. If the metastatic lesions are functional, they absorb radioactive iodine. Thyroiditis is unlikely to manifest as a mass because the entire gland is typically involved. Medullary carcinoma is less common than follicular carcinoma. There is an increased risk of non-Hodgkin B-cell lymphoma in patients with Hashimoto thyroiditis, but lymphomas are unlikely to destroy bone, although they can be present in the marrow.

PBD9 1097–1098  BP9 733–734  PBD8 1123–1124  BP8 768–769

27 A The large size, histologic features, and aggressive nature of this neoplasm are consistent with anaplastic carcinoma. The prognosis is poor. Other thyroid malignancies tend to form solitary or multifocal (in papillary and medullary carcinomas) masses without spindle cells; they are less likely to be extensively invasive, although metastases can occur, particularly to local lymph nodes in the case of papillary carcinomas or lung in the case of follicular carcinomas. Malignant lymphomas do not have spindle cells and do not tend to be infiltrative.

PBD9 1098–1099  BP9 734  PBD8 1124  BP8 771

28 A Medullary carcinomas are derived from the C cells, or parafollicular cells, of the thyroid, with embryologic origin from neural crest. Therefore they have neuroendocrine function, including synthesis of calcitonin. An amyloid stroma is a common feature of this tumor. These tumors occur sporadically in about 70% of cases, but they can be familial and part of multiple endocrine neoplasia types 2A and 2B. CD3 is a useful marker for some lymphoid neoplasms. Although various tissues may show positivity for estrogen receptors, this finding has no clinical significance in thyroid. Staining for parathyroid hormone is useful to determine if a parathyroid carcinoma is present. Vimentin is a marker for sarcomatous neoplasms, and cytokeratin is a useful marker to determine if a neoplasm is epithelial.

PBD9 1099–1100  BP9 734–735  PBD8 1124–1126  BP8 769–771

29 D When a patient develops hypercalcemia, a disorder of the parathyroid glands or a malignancy at a visceral location must be considered. The elevated parathyroid hormone (PTH) suggests primary hyperparathyroidism. The most common cause of primary hyperparathyroidism is a parathyroid adenoma. Secondary hyperparathyroidism, most commonly resulting from renal failure, is excluded when the serum inorganic phosphate level is low because phosphate is retained with chronic renal failure. Hypervitaminosis D can cause hypercalcemia because of increased calcium absorption, but in these cases, the PTH levels are expected to be near the low end of the reference range because of feedback suppression. Serum PTH levels near the high end of the reference range indicate autonomous PTH secretion unregulated by hypercalcemia. Although medullary carcinomas of the thyroid often have positive immunohistochemical staining for calcitonin, and plasma levels are sometimes increased,
30 A common cause of clinically significant hypercalcemia in adults is a malignancy. When a patient presents with hypercalcemia, a disorder of the parathyroid glands or a malignancy at a visceral location must be considered. Hypercalcemia from malignancy can be caused by osteolytic metastases or a paraneoplastic syndrome from secretion of parathyroid hormone–related protein by the tumor. Metastatic disease from common primary sites, such as the breast, lung, and kidney, is much more common than parathyroid carcinoma, which tends to be local but aggressive. Chronic renal failure causes phosphate retention, which tends to depress the serum calcium level and leads to secondary hyperparathyroidism; the serum calcium level is maintained at near-normal levels. Tuberculosis, a granulomatous disease, can be associated with hypercalcemia from up-regulation of 1,25-dihydrocholecalciferol in activated macrophages; lytic bone lesions from tuberculosis are uncommon. Parathyroid carcinomas are an uncommon cause of hyperparathyroidism, and bone metastases from parathyroid carcinomas are rare. Vitamin D toxicity theoretically can lead to hypercalcemia, but this condition is uncommon.

31 A parathyroid neoplasm, most likely an adenoma, is causing hypercalcemia complicated by osteitis fibrosa cystica of her finger. Parathyroid hyperplasia involving all the glands is less likely. Parathyroid carcinomas are rare, thankfully, because they are aggressive, and serum calcium levels may be so high that cardiac arrhythmias occur. The MEN1 mutation is the second most common mutation in parathyroid tumors, after the inversion with overexpression of CCND1, the gene encoding cyclin D1. MEN1 is a tumor suppressor gene, the loss of which occurs not only in sporadic parathyroid tumors, but also in multiple endocrine neoplasia (MEN) type 1. GNAS1 mutations are seen in 40% of cases of somatotroph (growth hormone–producing) pituitary adenomas. The TP53 mutation can be seen in anaplastic thyroid carcinomas. The RET gene mutation is seen in cases of medullary thyroid carcinoma. The VHL mutation can be seen in some pheochromocytomas in patients with von Hippel–Lindau disease.

32 Chronic kidney injury, often leading to small end-stage kidneys, with chronic renal failure can lead to secondary hyperparathyroidism resulting from decreased phosphate excretion by the kidneys. The resultant hyperparaphatemia depresses the serum calcium level and stimulates parathyroid gland activity. Because of reduced renal parenchymal function, there also is less active vitamin D, which leads to decreased dietary calcium absorption. Renal failure does not lead to any of the other endocrine lesions listed.

33 C Inadvertent removal of or damage to the parathyroid glands during thyroid surgery can cause hypocalcemia secondary to hypoparathyroidism. This is the most common cause of hypoparathyroidism. Individuals with hypocalcemia exhibit neuromuscular irritability, carpopedal spasm, and sometimes seizures. Antithyroglobulin antibody levels are of no use in diagnosing surgical diseases of the thyroid. Calcitonin quantitation is not a useful measure to determine the status of calcium metabolism. Parathyroid hormone levels decrease if the parathyroid glands are inadvertently removed during thyroid surgery, but the calcium level is the best immediate indicator of hypoparathyroidism, and this test is more readily available in the laboratory. The thyroid-stimulating hormone (TSH) concentration can increase if the patient becomes hypothyroid after surgery and is not receiving thyroid hormone replacement, but this is not an immediate problem.

34 An insulin overdose produces hypoglycemic coma. He does not have detectable C peptide, which indicates that there is no endogenous insulin production, typical for type 1 diabetes. The high insulin level is the result of the patient’s use of exogenous insulin to treat his diabetes mellitus. Because he has not eaten enough to maintain glucose at an adequate level, he has developed hypoglycemia. The ketoacidosis in this case results from decreased food intake, and anyone not consuming enough calories will develop ketosis. Ketoacidosis in type 1 diabetes mellitus would be accompanied by hyperglycemia. Acute myocardial infarction is a complication that generally occurs later in the course of diabetes when more atherosclerosis has developed. The patient has no obvious source of sepsis. Insulin is not injected into the bloodstream, and the injections are almost never complicated by infection. Hepatic failure is not a typical complication of diabetes mellitus. Hyperosmolar coma can complicate type 2 diabetes mellitus.

35 The presence of HLA-DR3 and HLA-DR4 alleles of the MHC class II region has the strongest linkage to type 1 diabetes mellitus. Autoantibodies to islet cell antigens such as glutamic acid decarboxylase are present years before overt clinical diabetes develops. An insulincausis caused by T cell infiltration occurs before the onset of symptoms or very early in the course of type 1 diabetes mellitus. The insulincausis in type 1 diabetes mellitus is associated with increased expression of class I MHC molecules and aberrant expression of class II MHC molecules on the beta cells of the islets. These changes are mediated by cytokines such as interferon-γ elaborated by CD4+ cells (along with CD8+ cells). Acute neutrophilic infiltration with necrosis and hemorrhage are characteristic of acute pancreatitis. Extensive fibrosis and fatty replacement of the pancreas is seen in patients with cystic fibrosis surviving for decades. Islet hyperplasia occurs in infants of diabetic mothers. Amyloid deposition in islets may be seen in some cases of type 2 diabetes mellitus. A fibrous stroma with minimal chronic inflammation and scattered normal islets is seen with chronic pancreatitis.
The findings are complications of diabetes with pregnancy, and the malformations suggest that hyperglycemia preceded the pregnancy, and type 2 diabetes is quite common, even now in women of childbearing age. The neonatal hyperglycemia is a consequence of excessive islet beta-cell function from having been in a hyperglycemic environment. Though cystic fibrosis is present from birth, the loss of pancreatic exocrine function takes years, and loss of islets is a late finding. Gestational diabetes refers to glucose intolerance in pregnancy, and newborns are likely to have hypoglycemia as a consequence of their own beta cell hyperfunction, but not anomalies. Maturity-onset diabetes of the young (MODY; but not in infancy) resembles type 2 diabetes and can occur from a variety of genetic defects in pathways monitoring glucose levels, but is much less common. Pancreatic neuroendocrine tumors are uncommon but could secrete glucagon with secondary diabetes.

Type 1 diabetes mellitus does not become overt until the beta cells are markedly depleted, and insulin levels are greatly reduced. In this case, the girl has ketoacidosis. Amyloid replacement of islets is a feature of type 2 diabetes mellitus; ketoacidosis is not a feature of type 2 diabetes mellitus. Acute or chronic pancreatitis diminishes exocrine pancreatic function, but rarely destroys enough islets to cause overt diabetes mellitus. Insulitis with inflammatory cells, mostly T cells, can be seen in the islets of patients with type 1 diabetes mellitus before the diabetes is clinically overt. Eosinophils are rare, however, with insulitis, but instead may be found in the islets of diabetic infants who fail to survive the immediate postnatal period. A pancreatic neuroendocrine tumor may become hormonally active and can lead to hypoglycemia if insulin is produced in excess; production of glucagon by such an adenoma may lead to secondary diabetes mellitus from having been in a hyperglycemic environment. Though cystic fibrosis is present from birth, the loss of pancreatic exocrine function takes years, and loss of islets is a late finding. Gestational diabetes refers to glucose intolerance in pregnancy, and newborns are likely to have hypoglycemia as a consequence of their own beta cell hyperfunction, but not anomalies. Maturity-onset diabetes of the young (MODY; but not in infancy) resembles type 2 diabetes and can occur from a variety of genetic defects in pathways monitoring glucose levels, but is much less common. Pancreatic neuroendocrine tumors are uncommon but could secrete glucagon with secondary diabetes.

Nonenzymatic glycosylation of proteins is a function of the level of blood glucose, rather than the cause of hyperglycemia. This is the basis for an elevated hemoglobin A1c. Type 1 and type 2 diabetes mellitus are characterized by hyperglycemia, but the underlying pathogenetic mechanisms are different. Type 1 diabetes mellitus is an autoimmune disease that is associated with certain alleles of the MHC class II molecules. It is characterized by a very high concordance rate in twins and the presence of islet autoantibodies. Insulin resistance is a key feature of type 2 diabetes mellitus.

Nonenzymatic glycosylation refers to the chemical process whereby glucose attaches to proteins without the aid of enzymes. The degree of glycosylation is proportionate to the level of blood glucose. Many proteins, including hemoglobin, undergo nonenzymatic glycosylation. Because RBCs have a life span of about 120 days, the amount of glycosylated hemoglobin is a function of the blood glucose level over the previous 120-day period. The level of glycosylated hemoglobin is not appreciably affected by short-term changes in plasma glucose levels. Random glucose testing is an immediate way for monitoring short-term adjustments with diet and medications such as insulin and oral agents. Fasting glucose testing affords a better way to diagnose diabetes mellitus initially. Measurements of cholesterol and fructosamine have no value in managing diabetes mellitus. Microalbuminuria may presage the development of diabetic renal disease. The “incretin effect” is diminished in patients with type 2 diabetes, and use of GLP-1 receptor agonists can help to restore incretin function and lead to improved glycemic control and loss of weight via increased satiety.
A Complications of diabetes mellitus are described in this man: vascular disease, neuropathy, nephropathy, and retinopathy. Hyperglycemia exceeds the capacity of renal tubular reabsorption, so glucose appears in the urine. The other listed findings involve organs that are not typically involved in diabetes mellitus: liver disease with decreased albumin synthesis, hyperaldosteronemia with hypokalemia, decreased narrow function with leukopenia, exocrine pancreatic disease with steatorrhea, or disordered uric acid metabolism.

PBD9 1116–1119  BP9 744–747  PBD8 1140–1142  BP8 782–784

A Severe peripheral atherosclerotic disease is a common complication of long-standing diabetes mellitus. Atherosclerotic narrowing of the arteries to the lower legs can cause ischemia and gangrene. The foot is often involved with gangrene, which may necessitate amputation. Diabetic neuropathy with decreased sensation increases the risk of repeated trauma, which enhances the risk of ulcers that cause infection and inflammation that promotes gangrene. Patients with type 2 diabetes mellitus or obesity, or both, are at increased risk of developing nonalcoholic steatohepatitis. Because this patient is not taking medications such as insulin, it is unlikely that he could become severely hypoglycemic. Because he is overweight, it is more likely that he has type 2 diabetes mellitus; ketoacidosis is unlikely. Infections with *Mucor circinelloides* are more likely to occur in ketoacidosis. Although pancreatic islets may have amyloid deposits, systemic amyloidosis and chronic pancreatitis (which involves the parenchymal acini) do not occur with type 2 diabetes mellitus.

PBD9 1116–1117  BP9 743–748  PBD8 1138–1139  BP8 781–782

C Nodular glomerulosclerosis, as shown in the figure, is a characteristic feature of renal involvement in diabetes mellitus and explains her proteinuria (which may progress to nephrotic syndrome). Peripheral insulin resistance is strongly linked to type 2 diabetes mellitus. Her history is classic for type 2 diabetes, as is the elevated hemoglobin A1c. Note that although premenopausal women are relatively protected from ischemic heart disease, diabetes tilts the balance and can promote development of coronary artery disease in younger women. Chronic pancreatitis typically affects exocrine pancreatic function more than endocrine function. Glucokinase gene mutations can lead to maturity-onset diabetes of the young (MODY), which is far less common than type 2 diabetes. Although localized amyloid deposition in islets can be seen with type 2 diabetes, it is not linked to conditions such as multiple myeloma that involve additional organs with amyloid deposition. T-cell mediated destruction of islet beta cells is a feature of type 1 diabetes.

PBD9 1117–1119  BP9 746–747  PBD8 1136–1137  BP8 778–780

D A complication of type 2 diabetes mellitus is hyperosmolar, nonketotic coma. In type 2 diabetes mellitus, the fundamental defect is insulin resistance, leading to an eventual decrease in plasma insulin or a relative lack of insulin, but there is still enough insulin to prevent ketosis. The resulting hyperglycemia tends to produce polyuria, leading to dehydration, which increases the serum glucose level further. If not enough fluids are ingested, dehydration drives the serum glucose to very high levels. Glucokinase gene mutations can be present with maturity-onset diabetes of the young (MODY). The HLA-DR3/HLA-DR4 genotype is a predisposing factor for type 1 diabetes mellitus. Severe loss of beta cells with insulitis, which may be triggered by viral infection, is a feature of autoimmune, or type 1, diabetes mellitus.

PBD9 1115  BP9 748  PBD8 1145  BP8 785, 787

A Complications of diabetes mellitus are described in this man: vascular disease, neuropathy, nephropathy, and retinopathy. Hyperglycemia exceeds the capacity of renal tubular reabsorption, so glucose appears in the urine. The other listed findings involve organs that are not typically involved in diabetes mellitus: liver disease with decreased albumin synthesis, hyperaldosteronemia with hypokalemia, decreased narrow function with leukopenia, exocrine pancreatic disease with steatorrhea, or disordered uric acid metabolism.

PBD9 1116–1119  BP9 744–747  PBD8 1140–1142  BP8 782–784

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PBD9 1115  BP9 748  PBD8 1145  BP8 785, 787

F Peripheral neuropathy, including autonomic neuropathy, can be caused by long-standing diabetes mellitus. It is thought that nerve cells do not require insulin for glucose uptake. In the presence of hyperglycemia, excess glucose diffuses into the cell cytoplasm and accumulates. The excess glucose is metabolized via the polyol pathway by intracellular aldose reductase enzyme to sorbitol and then to fructose, a reaction that uses NADPH as a co-factor. NADPH is also required for a reaction that regenerates reduced glutathione (GSH). GSH provides important antioxidant mechanisms in the cell. Thus persistence intracellular hyperglycemia through reduction of GSH makes the nerve cells susceptible to oxidative stress. Advanced glycosylation end products (AGEs) can directly cross-link extracellular matrix proteins to predispose vessels to shear stress and endothelial injury. Glycosylation tends to affect vascular walls and promote atherosclerosis. The downstream effects of protein kinase C activation are numerous, including production of VEGF, TGF-β, and the procoagulant protein plasminogen activator inhibitor-1 by the vascular endothelium.

PBD9 1115–1116, 1232  BP9 743–745  PBD8 1138–1140  BP8 780–781

C Nodular glomerulosclerosis, as shown in the figure, is a characteristic feature of renal involvement in diabetes mellitus and explains her proteinuria (which may progress to nephrotic syndrome). Peripheral insulin resistance is strongly linked to type 2 diabetes mellitus. Her history is classic for type 2 diabetes, as is the elevated hemoglobin A1c. Note that although premenopausal women are relatively protected from ischemic heart disease, diabetes tilts the balance and can promote development of coronary artery disease in younger women. Chronic pancreatitis typically affects exocrine pancreatic function more than endocrine function. Glucokinase gene mutations can lead to maturity-onset diabetes of the young (MODY), which is far less common than type 2 diabetes. Although localized amyloid deposition in islets can be seen with type 2 diabetes, it is not linked to conditions such as multiple myeloma that involve additional organs with amyloid deposition. T-cell mediated destruction of islet beta cells is a feature of type 1 diabetes.

PBD9 1117–1119  BP9 746–747  PBD8 1136–1137  BP8 778–780

A The clinical features of obesity with angina and glucose intolerance in this patient strongly suggest type 2 diabetes mellitus. This is confirmed by the oral glucose tolerance test (>200 mg/dL at 2 hours), useful in this case because her fasting blood glucose levels of 120 mg/dL and 122 mg/dL did not quite reach the diagnostic criterion of 126 mg/dL. The fundamental abnormality in type 2 diabetes mellitus is insulin resistance. Several adipocyte-derived molecules, such as adiponectin and resistin, have been implicated in the causation of insulin resistance, establishing the link between obesity and type 2 diabetes mellitus. The nuclear receptor peroxisome proliferator-activated receptor gamma (PPARγ) has emerged as a key molecule in the regulation of insulin resistance through its actions on adipocyte hormones. TZDs bind to and activate PPARγ in adipocytes, and increase the levels of the insulin-sensitizing hormone adiponectin and reduce the levels of free fatty acids and resistin, both of which increase insulin resistance. Insulin autoantibodies are seen with type 1 diabetes mellitus. Beta cell loss and density of insulin receptors are not major factors in the pathogenesis of type 2 diabetes mellitus. TZDs do not affect the metabolism of insulin. Glucagon excess worsens diabetes, but TZDs do not affect its secretion. Beta cells do not regenerate, but many antidiabetogenic drugs in type 2 diabetes mellitus are designed to work with the beta cells that are left.

PBD9 1111–1112, 1115  BP9 748–750  PBD8 1136–1137  BP8 778–780
Peripheral insulin resistance with obesity underlies type 2 to one form of maturity-onset diabetes of the young (MODY). Accompanied by diabetes, but having multiple children in- sometimes islet loss, in middle age. Down syndrome may be ride ion channels may lead to chronic pancreatic acinar, and sometimes islet loss, in middle age. Down syndrome may be involved is highly unlikely. Glucokinase gene mutations lead to one form of maturity-onset diabetes of the young (MODY). Peripheral insulin resistance with obesity underlies type 2 diabetes mellitus.

The plasma corticotropin level distinguishes between these two possibilities. Corticotropin levels are high if there is an ectopic source, whereas glucocorticoid secretion from an adrenal neoplasm suppresses corticotropin production by the pituitary, leading to atrophy of the contralateral adrenal cortex. A pheochromocytoma secretes catecholamines, accounting for hypertension, but not osteoporosis or the electrolyte changes noted. Medullary thyroid carcinomas have neuroendocrine cells, but are unlikely to produce corticosteroids.

The clinical and laboratory features of this case point to type 1 diabetes mellitus with childhood onset, MHC linkage, and evidence for autoimmune dysfunction mediated by T cells; the loss of islets leads to absolute lack of insulin, which requires insulin therapy. The remaining choices are not associated with autoimmunity. Cystic fibrosis with CFTR gene mutations affecting chloride ion channels may lead to chronic pancreatic acinar, and sometimes islet loss, in middle age. Down syndrome may be associated by diabetes, but having multiple children involved is highly unlikely. Glucokinase gene mutations lead to one form of maturity-onset diabetes of the young (MODY). Peripheral insulin resistance with obesity underlies type 2 diabetes mellitus.

The clinical and laboratory features of this case point to type 1 diabetes mellitus with childhood onset, MHC linkage, and evidence for autoimmune dysfunction mediated by T cells; the loss of islets leads to absolute lack of insulin, which requires insulin therapy. The remaining choices are not associated with autoimmunity. Cystic fibrosis with CFTR gene mutations affecting chloride ion channels may lead to chronic pancreatic acinar, and sometimes islet loss, in middle age. Down syndrome may be associated by diabetes, but having multiple children involved is highly unlikely. Glucokinase gene mutations lead to one form of maturity-onset diabetes of the young (MODY). Peripheral insulin resistance with obesity underlies type 2 diabetes mellitus.

The clinical findings of Cushing syndrome with masculinization suggest an adrenal cortical carcinoma, which is more likely to have endocrine function in women. A low ACTH level helps to rule out the possibility of an ACTH-secreting pituitary adenoma in the sella turcica, or ectopic ACTH from a neoplasm such as lung carcinoma. A “cold” nodule of the thyroid gland can represent a thyroid medul- lary carcinoma seen in multiple endocrine neoplasia type 2 in association with adrenal pheochromocytomas, but Cushing syndrome is not part of this complex. The location of a mass at the aortic bifurcation could be the “infamous” extra-adrenal pheochromocytoma of the obscure organ of Zuckerkandl, which explains hypertension with excess catecholamine release, but not the other features of Cushing syndrome.

The lung mass is likely a primary lung carcinoma, and the bilaterally enlarged adrenal glands can be explained by metastases to the adrenal glands. Destruction of over 90% of the adrenal cortices is responsible for adrenal failure manifested by malaise and the low serum cortisol concentration and the electrolyte disturbances. Amyloidosis can increase adrenal size, but does not produce a lung mass. In ectopic corticotropin syndrome, a lung cancer is a likely finding, and typically the adrenal glands are enlarged, but hypercortisolism also would be present. Waterhouse-Friderichsen syndrome, caused by Neisseria meningitidis infection, can increase the adrenal size secondary to marked hemorrhage (two to three times the normal size), but this does not explain the lung mass. This syndrome has an abrupt onset. A pitu- itary adenoma that is secreting corticotropin could increase adrenal size bilaterally, but hypercortisolism would occur.

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of the anterior pituitary, and corticotropin levels are not affected.

PBD9 1125–1127, 1132  BP9 753–754  PBD8 1151–1152  BP8 792

55 A Hypokalemia with neuromuscular irritability, hypertension, and low plasma renin suggests hyperaldosteronism. The most common cause for primary hyperaldosteronism is idiopathic adrenal cortical nodular hyperplasia. An insulinoma arising in the pancreas could account for episodic weakness, but the glucose level would be low. About 10% of pheochromocytomas are extra-adrenal, including para-aortic, and could account for hypertension from catecholamine excess, but there would not be hypokalemia. Thyroid enlargement could be Graves disease, though it is usually diffuse, and could account for weakness and hypertension, but with a wider pulse pressure, and without hypokalemia. A malignancy in the chest is more likely to be the cause for a paraneoplastic syndrome, but that is unlikely to be hypokalemia.

PBD9 1125–1127  BP9 755  PBD8 1151–1152  BP8 792

56 B Adrenogenital syndrome can lead to precocious puberty, which is most commonly associated with a deficiency of 21-hydroxylase. The lack of this enzyme reduces cortisol production, driving corticotropin production, which leads to adrenal hyperplasia and production of sex steroid hormones. Bilateral adrenal cortical atrophy is typically seen in cases of Addison disease or after long-term exogenous glucocorticoid therapy. A nodule in the adrenal cortex that has zona glomerulosa cells produces primary hyperaldosteronism; if it has zona fasciculata cells, it produces Cushing syndrome. Most adrenal nodules are nonfunctional and incidental findings. A nodule in the adrenal medulla, if functional, produces catecholamines, and older patients with such nodules have hypertension. Medullary atrophy is rare but might result from infections or toxins. Medullary hyperplasia is uncommon but could also produce catecholamines.

PBD9 1127–1128  BP9 756  PBD8 1152–1154  BP8 792–793

57 C A complete deficiency of 21-hydroxylase leads to the classic salt-wasting form of adrenogenital syndrome because the enzyme deficiency blocks formation of aldosterone and cortisol. Mutations may be deletions or duplications, or involve recombination between the CYP21 gene and a pseudogene. These mutations may lead to severe deficiency with salt-wasting and prenatal virilization or partial deficiency with postnatal virilization, with an apparent autosomal recessive pattern of inheritance. A deficiency of 11-hydroxylase blocks cortisol and aldosterone production as well, although intermediate metabolites with some glucocorticoid activity also are synthesized. Aromatase is involved with conversion of androstenedione to estrone, a pathway of steroid synthesis that does not affect cortisol production. A deficiency of 17α-hydroxylase would lead to reduction of both cortisol and sex steroid synthesis. Oxidase is the final enzyme in the pathway to aldosterone production.

PBD9 1127–1128  BP9 756  PBD8 1152–1154  BP8 792–793

58 D This is the typical adrenal finding in Waterhouse-Friderichsen syndrome, and meningococcemia is the most likely cause of such a rapid course. Chronic adrenocortical insufficiency can result from disseminated tuberculosis and from fungal infections, such as histoplasmosis, that involve the adrenal glands. Cytomegalovirus infections of the adrenals can be seen in immunocompromised states and can be severe enough to produce diminished adrenal function, although not acute failure. Streptococcus pneumoniae can produce septicemia, but it is unlikely to involve the adrenal glands specifically.

PBD9 1129–1130  BP9 757  PBD8 1155  BP8 794

59 B This woman has findings of acute adrenocortical insufficiency (acute addisonian crisis). Long-term corticosteroid therapy shuts off corticotropin stimulation to the adrenals, leading to adrenal atrophy. When this history is not elicited, and the patient is not continued on the corticosteroid therapy, a crisis ensues, in this case made worse by the stress of surgery. When tuberculosis is more prevalent and more severe without drug therapy, dissemination to adrenals occurs more frequently. An adrenal cortical adenoma without atrophy of the contralateral adrenal cortex could be a nonfunctioning adenoma or an aldosterone-secreting adenoma. If the contralateral cortex is grossly atrophic, the adenoma on the opposite side is probably secreting excess glucocorticoids. A carcinoma is most likely to destroy one adrenal, be nonfunctioning, and leave the remaining adrenal intact. Addison disease from granulomatous destruction of the adrenals was more common, but this is a chronic process. Hemorrhagic necrosis suggests Waterhouse-Friderichsen syndrome, which can complicate septicemia with organisms such as Neisseria meningitidis. Cortical nodular hyperplasia can be driven by an ACTH-secreting pituitary adenoma, or it can be idiopathic; in either case, hypercortisolism ensues, not Addison disease.

PBD9 1129–1130  BP9 757  PBD8 1155–1157  BP8 793–794

60 C Addison disease (primary chronic adrenocortical insufficiency) most often results from an idiopathic autoimmune condition (in areas of the world where the incidence of active tuberculosis is low). Autoimmune adrenalitis is associated with the appearance of other autoimmune diseases in about half of all cases. Such autoimmune phenomena are frequently seen in other endocrine organs, such as the thyroid gland. Other presumed autoimmune diseases, such as systemic lupus erythematosus, ulcerative colitis, and the vasculitides, are usually not forerunners to adrenal failure, although treatment of these conditions with corticosteroids can lead to iatrogenic adrenal atrophy. Type 2 diabetes mellitus, unlike type 1, does not have an autoimmune basis.

PBD9 1130–1131  BP9 757–758  PBD8 1155–1157  BP8 794–795

61 B Chronic adrenal insufficiency (Addison disease) results in decreased cortisol production and decreased mineralocorticoid activity. The skin hyperpigmentation results from increased corticotropin precursor hormone production,
which also stimulates melanocytes. The most common cause of Addison disease, in areas where tuberculosis is not endemic, is autoimmune adrenalitis. This process causes gradual destruction of the adrenal cortex, mediated most likely by infiltrating lymphocytes. A pancreatic neuroendocrine tumor (islet cell adenoma) secreting insulin could account for hypoglycemia, but not for the other metabolic changes. Metastases occasionally can destroy enough adrenal cortex to cause adrenal insufficiency, but the most common primary site is the lung, and there is no lung mass in her chest radiograph. Bilateral hemorrhages and resultant destruction of the adrenal glands are typically caused by mycog Beckwith, and this manifests as acute adenocortical insufficiency. Sarcoidosis also may involve the adrenals, but this is less common, and in this woman’s case, the normal chest radiograph helps to eliminate this possibility because hilar adenopathy is almost always present in sarcoidosis.

62 E These findings suggest a pheochromocytoma of the adrenal medulla. This is a rare neoplasm, but in cases of episodic hypertension, this diagnosis should be considered. Screening for urinary free catecholamines, metanephrine, and vanillylmandelic acid (VMA) can help to determine the diagnosis. Up to 25% of cases may be associated with an underlying tumor suppressor gene mutation, such as RET, NFI, or VHL. The level of HVA is more likely to be increased in a neuroblastoma, which is a tumor that occurs in children. The serum cortisol is increased with Addison disease, which is accompanied by hypotension. Hypoglycemia can also occur in Addison disease, as well as islet cell tumors. The serum potassium level can be decreased with aldosterone-secreting adrenal adenomas. An increased T₄ level occurs in patients with Graves disease; this disease can cause weight loss, heat intolerance, anxiety, tachycardia, tremors, and cardiac arrhythmia.

63 F Neuroblastomas are neoplasms that occur in children and may be congenital. They arise most commonly in the retroperitoneum in the adrenal glands or in extra-adrenal paraganglia. They are primitive small blue cell tumors that can produce high levels of catecholamine precursors and their metabolites such as HVA and vanillylmandelic acid (VMA). HVA is most specific. Because neuroblastomas are usually unilateral, abnormalities of ACTH or cortisol production are unlikely. Neuroblastomas do not affect serum prolactin or growth hormone levels. Adult pheochromocytomas are more likely to be detected by increased urinary free catecholamines, but the primitive cells of neuroblastoma are unlikely to produce amounts of catecholamines in the range of pheochromocytomas.

64 A These findings suggest multiple endocrine neoplasia (MEN) type 2A (Sipple syndrome) or possibly MEN type 2B (Williams syndrome). These patients have medullary carcinomas of the thyroid, pheochromocytomas, and parathyroid adenomas. This patient’s headaches could be caused by hypertension from a pheochromocytoma arising in the adrenal medulla. More than 70% of cases of pheochromocytomas are bilateral when familial. Medullary carcinoma also tends to be multifocal in this syndrome. This syndrome is associated with germ line mutations in the RET proto-oncogene. Family members who inherit the same mutation are at increased risk of developing similar cancers. Genetic screening followed by increased surveillance of affected family members is advised. Bilateral cortical atrophy from autoimmune destruction of the adrenals, leading to bilateral cortical atrophy, is now the most common cause of Addison disease. Cortical nodular hyperplasia can be driven by an ACTH-secreting pituitary adenoma, or it can be idiopathic; in either case, hypercortisolism ensues, not Addison disease. Granulomatous destruction of the adrenal glands suggests disseminated tuberculosis as a cause of Addison disease, which leads to adrenal insufficiency if bilateral, not unilateral. An adrenal cortical adenoma with atrophy of the contralateral adrenal cortex could be secreting excess glucocorticoids. A large mass with hemorrhage and necrosis in an adrenal suggests a cortical carcinoma.

65 C Multiple endocrine neoplasia (MEN) type 1 is also known as Wermer syndrome. (Remember the “three P’s” in neoplasia or hyperplasia—pancreas, pituitary, and parathyroids.) Adrenal pheochromocytomas are associated with MEN 2B. Endometrial carcinomas can arise in patients who have unopposed estrogen secretion, which can occur in estrogen-producing ovarian tumors. These are not part of MEN 1. Small cell carcinomas of the lung are known for various paraneoplastic syndromes, but not usually hypercalcemia. It also is doubtful that this patient would have lived 5 years with a small cell carcinoma. If her hypercalcemia had been a paraneoplastic syndrome, the parathyroid glands would not have been enlarged, and the serum calcium level would not have returned to normal after surgery. Medullary thyroid carcinomas are part of MEN 2A or 2B.

66 E The anatomic location of the mass is the pineal gland. In children, the most common pineal tumor is a pineoblastoma, whereas in adults, it is a pineocytoma. Both are quite rare, and their location makes them difficult to remove completely. Cranioopharyngiomas are aggressive neoplasms that are often suprasellar and difficult to remove. Hypothalamic gliomas also are suprasellar. Central nervous system lymphomas are uncommon in children and rare at this site. Metastatic carcinoma is rare in any location in children because children do not have many malignancies, and the malignancies that do have are often not carcinomas. A prolactinoma occurs in the sella turcica.

67 C Metastatic carcinoma of the adrenal can cause hypercalcemia, but in this patient, the chest radiograph shows no evidence of a lung mass. Bilateral adrenal hemorrhages and resultant destruction of the adrenals is less common, and in this woman’s case, the normal chest radiograph helps to eliminate this possibility because hilar adenopathy is almost always present in sarcoidosis.

68 F Neuroblastomas are neoplasms that occur in children and may be congenital. They arise most commonly in the retroperitoneum in the adrenal glands or in extra-adrenal paraganglia. They are primitive small blue cell tumors that can produce high levels of catecholamine precursors and their metabolites such as HVA and vanillylmandelic acid (VMA). HVA is most specific. Because neuroblastomas are usually unilateral, abnormalities of ACTH or cortisol production are unlikely. Neuroblastomas do not affect serum prolactin or growth hormone levels. Adult pheochromocytomas are more likely to be detected by increased urinary free catecholamines, but the primitive cells of neuroblastoma are unlikely to produce amounts of catecholamines in the range of pheochromocytomas.
1. A 20-year-old man has noted a cluster of small lesions on his upper lip for the past 5 days. On physical examination, there are four lesions ranging from 0.2 to 0.5 cm that are raised and filled with clear fluid. Which of the following descriptive terms best applies to his lesions?
   A. Bullae
   B. Macules
   C. Papules
   D. Pustules
   E. Vesicles

2. A 5-year-old girl has a routine health checkup. On physical examination, she has scattered 1- to 3-mm, light brown macules on her face, trunk, and extremities. The parents state that these macules become more numerous in the summer months, but fade over the winter. The lesions do not itch, bleed, or hurt. What is the most likely diagnosis?
   A. Freckle
   B. Lentigo
   C. Melasma
   D. Nevus
   E. Vitiligo

3. A 74-year-old woman has noted increasing size and number of darker brown patches on the dorsum of each hand for the past 15 years. They do not change with sun exposure, are nonpruritic, and not tender. On examination, these 0.5- to 1-cm lightly pigmented lesions are flat. Which of the following is the most likely microscopic finding in these lesions?
   A. Basal melanocytic hyperplasia
   B. Dermal nevus cells
   C. Loss of melanin in surrounding skin
   D. Mast cell proliferation
   E. Pigmented fungal hyphae

4. A 10-year-old girl has a routine check of her health status. On physical examination, a flat, uniformly brown, 2-cm skin lesion is noted just above the right buttock. The lesion has been present since birth and has not changed in appearance. The figure shows the representative microscopic appearance of the lesion. What is the most likely diagnosis of this lesion?
   A. Acanthosis nigricans
   B. Basal cell carcinoma
   C. Dysplastic nevus
   D. Lentigo
   E. Malignant melanoma
   F. Melanocytic nevus
   G. Seborrheic keratosis
5. A 17-year-old girl has hundreds of skin lesions on her body that have been forming since childhood. On physical examination, 0.4- to 1.7-cm, macular to slightly raised, plaquelike, dark brown pigmented lesions are present on sun-exposed and non-sun-exposed areas of skin. The lesions have irregular contours, and there is variability in pigmentation. She says that her 15-year-old brother has similar lesions. She is most likely to have an inherited mutation involving which of the following genes?

A. CDKN2A  
B. FGFR3  
C. NF1  
D. PTCH  
E. TSC1

6. Skin cancers are studied for epidemiologic associations with racial and environmental factors. The incidence of basal cell carcinoma, squamous cell carcinoma, and malignant melanoma has increased over the past 50 years. In which of the following countries is this incidence the highest?

A. Australia  
B. Bolivia  
C. Korea  
D. Nigeria  
E. Norway  
F. Tunisia

7. A 39-year-old woman has a nodule on her back that has become larger over the past 2 months. On physical examination, there is a 2.1-cm pigmented lesion with irregular borders and irregular brown to black areas. An excisional biopsy with wide margins is performed, and microscopic examination of the biopsy specimen shows a malignant melanoma composed of epithelioid cells that extend 2 mm into the reticular dermis. There is a band of lymphocytes beneath the lesion. Which of the following is the most important determinant of prognosis for this woman?

A. Age at diagnosis  
B. Depth of the lesion  
C. Extent of radial growth  
D. Inflammatory response  
E. Location on the skin

8. A 77-year-old man has a lesion on the right side of his face that has enlarged slowly over the past 5 years. On examination, the 3-cm lesion has irregular borders, irregular brown to black pigmentation, and a central 2-mm raised blue-black nodule. The lesion is resected and microscopically shows radial growth of large round malignant cells, some isolated and others in nests in the epidermis and superficial papillary dermis. The cells have prominent red nucleoli and dustlike cytoplasmic pigment. Which of the following mutated genes is most likely to be present in the skin lesion of this man?

A. ATM  
B. BRAF  
C. NF1  
D. TYR  
E. XPA

9. A 53-year-old man noticed enlargement over 2 months in the skin lesion on the upper, outer area of his right arm, shown in the figure. Physical examination yields no other remarkable findings. Which of the following occupations is this man most likely to have had earlier in life?

A. Auto repair mechanic  
B. Chemist in a factory  
C. Lifeguard on the beach  
D. Miner in a coal mine  
E. Radiation oncologist at a cancer center

10. A 75-year-old man has noticed slowly enlarging lesions, similar to those shown in the figure, on his trunk over the past 20 years. One of the lesions is excised, and microscopic examination shows sheets of lightly pigmented basaloid cells that surround keratin-filled cysts. This lesion is sharply demarcated from the surrounding epidermis. What is the most likely diagnosis?

A. Basal cell carcinoma  
B. Intradermal nevus  
C. Melanoma  
D. Seborrheic keratosis  
E. Squamous cell carcinoma  
F. Verruca vulgaris
A 64-year-old man has noted changes in the texture and color of skin in his armpits and groin over the past 3 months. On physical examination, there is thickened, darkly pigmented skin in the axillae and flexural areas of the neck and groin. These areas are neither painful nor pruritic. A punch biopsy specimen of axillary skin shows undulating epidermal acanthosis with hyperkeratosis and basal layer hyperpigmentation. These lesions are most likely to be cutaneous markers of which of the following underlying diseases?

A. AIDS  
B. Colonic adenocarcinoma  
C. Langerhans cell histiocytosis  
D. Mastocytosis  
E. Systemic lupus erythematosus

A 52-year-old woman notes several small, baglike lesions that have appeared on the skin in front of her armpits over the past 2 years. They are located where her bra rubs against the skin. On physical examination, five small, soft papules in the anterior axillary line are covered by wrinkled skin and attached to the skin surface by a thin pedicle. They are 0.5 to 0.8 cm in length and about 0.3 cm in diameter. One lesion has undergone torsion and is more erythematous and painful to touch than the others. What is the most likely diagnosis?

A. Fibroepithelial polyp  
B. Hemangioma  
C. Lentigo senilis  
D. Melasma  
E. Pilar cyst  
F. Xanthoma

A 31-year-old man notes a bump on the skin of the lower abdomen that has enlarged over the past 4 years and has become more painful in the past week. On physical examination, there is a subcutaneous, movable, soft nodule at the belt line anteriorly that elicits pain with pressure. The overlying skin is intact. He states that the nodule began hurting about 1 day after he vigorously squeezed it. The lesion is excised and does not recur. Which of the following is the most likely diagnosis?

A. Acne vulgaris  
B. Dermatofibroma  
C. Epidermal inclusion cyst  
D. Fibroepithelial polyp  
E. Trichoepithelioma  
F. Xanthoma

A 69-year-old woman has been bothered by a discolored area of skin on her forehead that has not faded during the past 3 years. On physical examination, there is a 0.8-cm red, rough-surfaced lesion on the right forehead above the eyebrow. A biopsy specimen examined microscopically shows basal cell hyperplasia. Some of the basal cells show nuclear atypia associated with marked hyperkeratosis and parakeratosis with thinning of the epidermis. The upper dermal collagen and elastic fibers show homogenization with elastosis. What is the most appropriate treatment option for this patient?

A. Apply hydrocortisone cream  
B. Reduce intake of dietary fat  
C. Surgically excise the lesion with wide margins  
D. Take antioxidants  
E. Wear a hat outdoors

A 53-year-old man with idiopathic dilated cardiomyopathy underwent orthotopic heart transplantation. During the next 5 years, he had two episodes of minimal cellular rejection, which were adequately treated by an increase in immunosuppressive therapy. He has developed multiple skin lesions on the face and upper trunk over the past 6 months. On physical examination, the lesions are similar to the lesion shown in the figure. Some of the larger lesions have ulcerated. A biopsy specimen is most likely to identify which of the following lesions?

A. Dermatofibroma  
B. Erythema multiforme  
C. Lichen planus  
D. Psoriasis  
E. Squamous cell carcinoma

A 6-year-old boy has had the appearance of an enlarging nodule on his ear during the past month. On physical examination, there is a 1.2-cm, flesh-colored, dome-shaped nodule on his right ear lobe. The nodule is excised and microscopically has nests of poorly differentiated islands of squamous epithelium invading the dermis. The boy continues to develop similar lesions. He is most likely to have a genetic mutation involving which of the following cellular functions?

A. Induction of apoptosis  
B. Growth factor receptor signaling  
C. Keratinocyte adhesion  
D. Nucleotide excision repair  
E. Amplification of oncogenes
17 A 76-year-old man has had a slowly enlarging nodule on his right eyelid for 4 years. On physical examination, there is a 0.3-cm pearly nodule on the upper eyelid near the lateral limbus of the right eye. The lesion is excised, but multiple frozen sections are made during the surgery to minimize the extent of the resection and preserve the eyelid. The microscopic appearance of the lesion is shown at low magnification in the figure. What is the most likely diagnosis?

A Actinic keratosis  
B Basal cell carcinoma  
C Dermatofibroma  
D Malignant melanoma  
E Nevocellular nevus

18 A 22-year-old woman has had a light brown nodule on her left shin for the past year. On examination it is a 0.7-cm firm, nontender nodule. It is excised and on microscopic examination consists of a localized, dermal, spindle cell proliferation with overlying epidermal hyperplasia and downward elongation of hyperpigmented rete ridges. What is the most likely diagnosis?

A Dermatofibroma  
B Leiomyoma  
C Merkel cell carcinoma  
D Neurofibroma  
E Schwannoma

19 A 46-year-old man has had a slowly enlarging nodule on his trunk for the past 10 months. On examination, the 5-cm lesion is in the right anterior axillary line at the level of the nipple and has a central ulcerated plaque. The lesion is widely excised and on microscopic examination is composed of dense spindle cells in a storiform pattern that invades the dermal adipose tissue. The lesion recurs within a year, and the man is treated with a tyrosine kinase inhibitor. Molecular analysis of the lesion is most likely to show activation with overexpression of which of the following genes?

A TSC1  
B PDGFB  
C FGFR3  
D KIT  
E PTCH1

20 A 9-year-old girl has been scratching a group of small bumps on the skin of her forearm for the past month. On physical examination, there are five 0.4- to 0.9-cm, small, flat to slightly papular, pale brown lesions on the volar surface. The lesions become more pruritic with swelling and erythema when rubbed. A biopsy specimen of one of the lesions examined microscopically shows an upper dermal infiltrate of large cells with abundant pink cytoplasm that stains an intense purple color with toluidine blue. Which of the following cell types is most likely to form these lesions?

A CD4+ lymphocyte  
B CD8+ lymphocyte  
C Langerhans cell  
D Macrophage  
E Mast cell  
F Merkel cell

21 A 30-year-old man is known for his large appetite. At a luncheon meeting, he notices that all the cookies contain nuts, which the other diners have ordered knowing that he would not eat them because he would develop blotchy, erythematous, slightly edematous, pruritic plaques on his skin. These plaques would form and then fade within 2 hours. If the man eats the cookies, which of the following sensitized cells would release a mediator that produces these skin lesions?

A CD4+ lymphocyte  
B Mast cell  
C Natural killer cell  
D Neutrophil  
E Plasma cells

22 A 31-year-old woman tries a new facial cosmetic product and a day later has red papules from 0.2 to 0.5 cm scattered over her skin. She sees her traditional healer 2 days later, and on physical examination some of the lesions are now vesicular and oozing clear fluid, whereas others are crusted. An herbal preparation is applied over the next 3 days and the lesions subside. Which of the following is the most likely diagnosis for this woman?

A Eczematous dermatitis  
B Erythema multiforme  
C Impetigo  
D Lichen simplex chronicus  
E Urticaria
23. A 43-year-old woman develops a red rash on her trunk and extremities 1 week after being hospitalized for treatment of an upper respiratory infection complicated by pneumonia. On physical examination, the 2- to 4-mm lesions are erythematous, papulovesicular, oozing, and crusted. The lesions begin to disappear after she is discharged from the hospital 1 week later. What is the most likely pathogenesis of these lesions?

A. Bacterial septicemia
B. Drug reaction
C. Human papillomavirus infection
D. Photosensitivity
E. Type I hypersensitivity

24. A 28-year-old HIV-infected man has had increasing fever, cough, and dyspnea for the past 3 days, which has culminated in acute respiratory failure. On admission to the hospital, his temperature is 37.8°C. On physical examination, he has a respiratory rate of 30/min and diffuse crackles with diminished breath sounds in all lung fields. He undergoes a bronchoalveolar lavage that yields *Pneumocystis jiroveci* by direct fluorescent antigen testing. Within 1 week after initiation of therapy, he develops target lesions of the skin composed of red macules with a pale, vesicular center. The 2- to 5-cm lesions are distributed symmetrically over the upper arms and chest. Which of the following drugs is most likely to be implicated in the development of these lesions?

A. Dapsone
B. Pentamidine
C. Ritonavir
D. Sulfamethoxazole
E. Zidovudine

25. A 44-year-old woman has developed skin lesions over her elbows and knees during the past year. The lesions start as 4-mm pustules with surrounding erythema but then evolve into 1- to 5-cm plaques that are covered with a silvery-white scale. The lesions appear first in areas of local trauma, but exposure to sunlight causes the lesions to regress. A biopsy of one lesion shows thinning of stratum granulosum with marked overlying parakeratotic scale containing microabscesses. Which of the following risk factors is most likely to be associated with her skin disease?

A. Atopy with elevated levels of IgE
B. Autoantibodies to epidermal desmoglein
C. Exposure of skin to plant allergens
D. Inheritance of certain HLA-Cw alleles
E. Herpes simplex virus infection

26. A 27-year-old woman has developed areas of scaling skin over the past month. On physical examination there are 1- to 3-cm light pink plaques covered with silvery scale on her arms and torso. A punch biopsy of one lesion examined microscopically shows keratinocyte nuclei retained within cells in the stratum corneum. Which of the following descriptive terms best applies to this microscopic finding?

A. Acanthosis
B. Dyskeratosis
C. Hyperkeratosis
D. Parakeratosis
E. Spongiosis

27. A 28-year-old man has had waxing and waning of the lesions shown in the figure for 10 years. The scalp, lumbosacral region, and glans penis also are affected. For the past 2 years, he has had chronic arthritis in the hips and knees. Which of the following physical findings would most likely be present in this patient?

A. Friction rub
B. Guaiac-positive stool
C. Hyperreflexia
D. Hypertension
E. Nail changes

28. A 39-year-old woman goes to her dentist for a routine checkup. The dentist notes that she has 0.2- to 1.5-cm scattered, white, reticulated areas on the buccal mucosa. The woman says that these lesions have been present for 1 year. She also has multiple 0.3-cm purple, pruritic papules on each elbow. A biopsy specimen of a skin lesion is taken and microscopically shows a bandlike infiltrate of lymphocytes at the dermal-epidermal junction with degeneration of basal keratinocytes. Which of the following is the most likely outcome for this woman?

A. Disseminated opportunistic infection
B. Multiple allergies to medications
C. Progression to chronic renal disease
D. Resolution of the lesions
E. Skin adnexal tumor development

29. A 39-year-old woman has developed vesicular skin lesions over the past week. On physical examination, she has multiple, 0.2- to 1-cm vesicles and bullae on the skin of her scalp, axillae, groin, and knees. Many lesions appear to have ruptured, and a shallow erosion with a dried crust of serum remains. A biopsy specimen of an axillary lesion examined microscopically shows epidermal acantholysis and formation of an intraepidermal blister. The basal cell layer is intact. Which of the following additional tests is most likely to explain the pathogenesis of the patient’s disease?

A. Darkfield microscopy of vesicular fluid
B. HLA genotyping
C. Immunostaining with antidesmoglein
D. Quantitation of serum IgE level
E. Viral culture of vesicular fluid
A 65-year-old man has developed pruritus followed by blistering skin lesions over the trunk, legs, and arms over the past month. On physical examination, there are 1- to 4-cm tense bullae, particularly over flexural surfaces of skin. A biopsy of one lesion is examined microscopically by direct immunofluorescence staining and shows a subepidermal bulla, with both IgG and C3 deposited linearly along the dermal-epidermal junction. He is treated with topical corticosteroids, and a month later the lesions are healed without scarring. Which of the following components of the skin has most likely been targeted by an autoantibody in this man?

- A Hemidesmosome
- B Keratinocyte cell membrane
- C Lamina densa
- D Nucleus
- E Reticulin

A 35-year-old man has had an outbreak of pruritic lesions over the extensor surfaces of the elbows and knees during the past month. He has a history of malabsorption that requires him to eat a special diet, but he has had no previous skin problems. On physical examination, the lesions are 0.4- to 0.7-cm vesicles. A 3-mm punch biopsy of one of the lesions over the elbow is performed and on microscopic examination shows accumulation of neutrophils at the tips of dermal papillae and formation of small blisters caused by separation at the dermoepidermal junction. Immunofluorescence studies show granular deposits of IgA localized to tips of dermal papillae. Laboratory studies show serum antigliadin antibodies. What is the most likely diagnosis?

- A Bullous pemphigoid
- B Contact dermatitis
- C Dermatitis herpetiformis
- D Discoid lupus erythematosus
- E Erythema multiforme
- F Pemphigus vulgaris

A study of persons developing skin lesions following sun exposure is conducted. The lesions are not found on skin protected from ultraviolet light. Biopsies of involved skin show immunoglobulin G deposition along the dermal-epidermal junction, along with vacuolization of the basal layer and a perivascular lymphocytic infiltrate. No other organ involvement is present. Which of the following diseases do these patients most likely have?

- A Bullous pemphigoid
- B Celiac disease
- C Discoid lupus erythematosus
- D Dysplastic nevus syndrome
- E Toxic epidermal necrolysis

An 18-year-old man has facial and upper back lesions that have waxed and waned for the past 6 years. On physical examination, there are 0.3- to 0.9-cm comedones, erythematous papules, nodules, and pustules most numerous on the lower face and posterior upper trunk. Other family members have been affected by this condition at a similar age. The lesions worsen during a 5-day cruise to the Adriatic. Which of the following organisms is most likely to play a key role in the pathogenesis of these lesions?

- A Group A β-hemolytic streptococcus
- B Herpes simplex virus type 1
- C Mycobacterium leprae

A 29-year-old man with a history of Crohn disease has noted the appearance of a painful red nodule on his left lower leg during the past week. On physical examination, his temperature is 37.3° C. There is a 0.4-cm, dark red, exquisitely tender nodule that has a surrounding 5-cm diameter area of paler red skin. This lesion resolves over the next 3 weeks, but another develops on the opposite calf. A skin biopsy of the second lesion is examined microscopically and shows a dermal mixed inflammatory infiltrate with neutrophils, round cells, and giant cells, affecting adipose tissue, along with pronounced edema. These lesions resolve without scarring, but more lesions develop during the next year. What is the most likely diagnosis?

- A Acne vulgaris
- B Dermatitis herpetiformis
- C Erythema nodosum
- D Impetigo
- E Molluscum contagiosum

A 13-year-old girl has two nontender lesions on her fingers that have appeared over the past 5 months. On physical examination, there are 0.5-cm slightly raised, pebbly-surfaced, gray-white papules, one on the dorsum of her distal right index finger and another periungual to her little finger. They gradually disappear over the next 18 months. Which of the following is the most likely factor in the pathogenesis of her lesions?

- A BRAF gene mutation
- B Human papillomavirus infection
- C IgA antibody deposition
- D Type IV hypersensitivity reaction
- E Vitamin B3 deficiency

A 26-year-old man has noticed slowly enlarging lesions on his hands for the past 3 years. On physical examination, the lesions appear similar to those shown in the figure. There are no other remarkable findings. The lesions have not changed in color, do not itch or bleed, and are not associated with pain. What is the most likely diagnosis?

- A Condyloma acuminatum
- B Dermatofibroma
- C Intradermal nevus
- D Molluscum contagiosum
- E Seborrheic keratosis
- F Squamous cell carcinoma
- G Verruca vulgaris
A 35-year-old man has noted a small bump on his upper trunk for the past 6 weeks. On physical examination, there is a solitary, 0.4-cm, flesh-colored nodule on the upper trunk. The dome-shaped lesion is umbilicated, and a curdlike material can be expressed from the center. This material is smeared on a slide, and Giemsa stain shows many pink, homogeneous, cytoplasmic inclusions. The lesion regresses over the next 2 months. Which of the following infectious agents most likely produced this lesion?

A  Histoplasma capsulatum  
B  Human papillomavirus  
C  Molluscum contagiosum  
D  Staphylococcus aureus  
E  Tinea corporis  
F  Varicella-zoster virus

A 6-year-old boy developed 1- to 2-cm erythematous macules and 0.5- to 1-cm pustules on his face over the past week. During the next 2 days, some of the pustules break, forming shallow erosions covered by a honey-colored crust. New lesions form around the crust. The boy’s 40-year-old uncle develops similar lesions after visiting for 1 week during the child’s illness. Removal of the crusts from the boy’s face is followed by healing within 1 week. The uncle does not seek medical care, and additional pustules form at the periphery of the crusts. Which of the following conditions most likely explains these findings?

A  Acne vulgaris  
B  Bullous pemphigoid  
C  Contact dermatitis  
D  Erythema multiforme  
E  Impetigo  
F  Lichen planus

A 23-year-old man and other members of his racquetball club have noticed more itching of their feet in the past 2 months. On physical examination, the man has diffuse, erythematous, scaling skin lesions between the toes of both feet. There are no other remarkable findings. These findings are most likely the result of infection with which of the following organisms?

A  Group A β-hemolytic streptococcus  
B  Herpes simplex virus  
C  Human papillomavirus  
D  Propionibacterium acnes  
E  Sarcoptes scabiei  
F  Staphylococcus aureus  
G  Trichophyton rubrum

A 10-year-old girl has multiple excoriations that have appeared on the skin of her hands over the past week. The child reports that she scratches her hands because they itch. Physical examination shows multiple 0.2- to 0.6-cm linear streaks in the interdigital regions. Treatment with a topical lindane lotion resolves the condition. Which of the following organisms is most likely responsible for these findings?

A  Ixodes scapularis  
B  Poxvirus  
C  Staphylococcus aureus  
D  Sarcoptes scabiei  
E  Tinea corporis

A 32-year-old woman has noticed depigmented areas on her trunk that have waxed and waned for 3 months. She says that they do not itch or bleed, and are not painful. Physical examination shows variably sized, 0.3- to 1.2-cm macules over her upper trunk. The macules are lighter colored than the surrounding skin and have a fine, peripheral scale. Infection with which of the following organisms is most likely to produce these findings?

A  Epidermophyton species  
B  Human papillomavirus  
C  Malassezia furfur  
D  Mycobacterium leprae  
E  Propionibacterium acnes  
F  Sarcoptes scabiei

An 11-year-old healthy girl has had an erythematous, scaly plaque on her upper chest for the past 2 days. When examined 2 days later, an annular 7-cm lesion has slightly elevated, peripheral erythema with overlying scale. A skin scraping of the lesion is performed, followed by topical ketoconazole therapy. Her family acquired a new German Shepherd puppy last week. What is most likely to be seen on microscopy with potassium hydroxide (KOH) on the scraping of this lesion?

A  Bacteria in long chains  
B  Budding yeast cells  
C  Molluscum bodies  
D  Septate branching hyphae  
E  Vacuolization of keratinocytes
ANSWERS

1 E Vesicles have a maximum size of 5 mm. They are often easily traumatized and unroofed to leave shallow ulcerations. In this case, herpetic vesicles are most likely present. Bullae are also fluid filled, but larger than 0.5 cm. Macules are not raised. They are flat areas less than 0.5 cm and distinguished by a difference in color; if the area is larger than 0.5 cm, then it is a patch. Pustules are elevated and less than 0.5 cm; if larger, the term nodule is applied; either may be dome-shaped or flat-topped. Pustules are raised and filled with leukocytes to impart a yellow color.

2 A Freckles are common. Individuals with a light complexion and/or red hair are more likely to have freckles. These lesions may be a cosmetic problem, but they have no other significance. A lentigo is a brown macule whose pigmentation is not related to sun exposure. Melasma is most often a masklike area of facial hyperpigmentation associated with pregnancy. Nevi do not wax and wane with sun exposure. Areas of skin with vitiligo are depigmented.

3 A Unlike a freckle that darkens upon light exposure, lentigo (pleural: lentigenes) does not. There is focal increase in melanocytes of the basal layer, along with thinning of epidermis and elongation of rete ridges. Lentigenes can occur at any age; those in older persons may be termed senile lentigenes, or age spots. A nevus typically forms a nodule and likely appears much earlier in life. Loss of melanin is termed vitiligo, and is a separate process from melanocyte hyperplasia. A collection of mast cells can produce urticaria pigmentosa, which often has a light brown appearance, but pruritus is common. Some superficial fungal infections (black piedra) may be pigmented, but they typically do not persist for years.

4 F The figure shows both junctional (along the dermal-epidermal junction) and intradermal nevus cells in a form of melanocytic nevus known as a congenital nevus. Larger nevi do have an increased risk of melanoma arising within them; however, the lack of additional lesions and the bland microscopic appearance shown here argue against dysplastic nevus syndrome. Acanthosis nigricans is an uncommon condition with hyperpigmented areas in skin folds; it may occur in association with endocrinopathies or with neoplasms. Basal cell carcinomas occur in sun-exposed skin of adults, and the cells have darker nuclei with scant cytoplasm. A lentigo is a common focal pigmented lesion that can appear at any age; it is a melanocyte hyperplasia. Melanoma occurs in adults and shows signs of rapid growth and change, with very atypical cells invading the dermis. Seborrhoeic keratoses are seen in older adults and are raised, pigmented lesions of thickened epidermis.

5 A The clinical appearance, distribution, and occurrence in two siblings suggest that these lesions represent the dysplastic nevus syndrome. Dysplastic nevi are precursors of malignant melanoma. The most important gene in familial cases is that of cyclin-dependent kinase inhibitor 2 (CDKN2A), which encodes several tumor suppressors including p16/INK4A. FGFR3 mutations may be found with seborrhoeic keratoses in older adults. The PTCH gene is implicated in the pathogenesis of sporadic basal cell carcinomas and the familial nevoid basal cell carcinoma syndrome. TSCI mutations are associated with tuberous sclerosis.

6 A Most of Australia’s population represents immigration of fair-skinned Europeans. That fact, combined with an upper atmospheric ozone loss that lets in more ultraviolet radiation, and an outdoor lifestyle in a sunny clime, has led to an increase in the number of cases. The nonmelanoma cancers do not contribute significantly to death, but the melanomas do. The other listed countries have populations with darker skin or less sun exposure.

7 B Extension downward into the reticular dermis indicates vertical (nodular) growth. When a melanoma exhibits a nodular growth pattern, rather than a radial pattern, there is increased likelihood that a clone of neoplastic cells has arisen that is more aggressive and is more likely to metastasize. For lesions deeper than 2 mm, the 5-year survival is about 65%, but it is greater than 90% if the depth of the lesion is less than 1 mm. Although there has been a lymphocytic response to this tumor, it is insufficient to destroy or contain it completely. Most melanomas are sporadic, nonfamilial, and related to sun exposure. They may occur at a variety of ages, but early diagnosis and removal is the key to better prognosis. Melanomas may occur in a variety of locations, even mucosal surfaces and the eye, but the depth of the lesion carries the most significance for prognosis.

8 B He has lentigo maligna melanoma, which exhibits an indolent radial growth phase for many years before reaching an accelerated nodular vertical growth phase. Most of these lesions arise on the skin of the head and neck of elderly persons. Activating BRAF mutations are downstream from RAS mutations and are present in the majority of malignant melanomas. ATM gene mutations underlie ataxia-telangiectasia, in which there are dilated subepidermal blood vessels forming lesions called telangiectasias. NFI mutations are associated with the appearance of many dermal neurofibromas. TYR mutations are seen in association with albinism; TYR encodes the enzyme tyrosinase that is part of the initial pathway of melanin production by melanocytes. Mutations in the XP family of genes are seen with xeroderma pigmentosum, in which defects in DNA excision repair predispose to skin
cancers (basal cell carcinoma, squamous cell carcinoma, malignant melanoma) starting in childhood from exposure to ultraviolet light.

PBD9 1149–1150  BP9 867–869  PBD8 1172–1175  BP8 855–857

9 C The figure shows a malignant melanoma with abnormal ABCs: Asymmetry, irregular Borders, and variable Color of pigmentation. A quickly developing change in a pigmented lesion suggests the possibility of melanoma. Some melanomas are familial, arising from conditions such as dysplastic nevus syndrome. Most melanomas occur sporadically, however, and are related to sun exposure, as may occur in a lifeguard. The other listed occupations have no significant sun exposure.

PBD9 1149–1150  BP9 867–869  PBD8 1172–1173  BP8 854–857

10 D Seborrheic keratoses are flat, round, pigmented, sharply demarcated lesions and benign tumors. They are composed of pigmented basaloid cells. Seborrheic keratoses are common in older individuals. The lesions gradually enlarge, but they are not painful and do not ulcerate. They mainly are a cosmetic problem. Basal cell carcinomas are nodular, slowly enlarging lesions most common on the head and trunk and are related to sun exposure. An intradermal nevus can produce a pigmented nodule, but microscopically it is composed of nests of small nevus cells, and the lesions increase minimally in size over time. Melanomas have very atypical spindle to epithelioid cells that invade the dermis; they tend to change in appearance over weeks to months, not years. The cells in a squamous cell carcinoma are atypical and may invade the dermis. A verruca vulgaris, or wart, also has a rough surface, but such lesions are more common on the hands and feet, and they may regress after several years.

PBD9 1151  BP9 862  PBD8 1175  BP8 848–849

11 B Acanthosis nigricans may be a cutaneous marker for benign and malignant neoplasms. The skin lesions often precede signs and symptoms of associated cancers. They are believed to arise from the action of epidermal growth-promoting factors produced by neoplasms. Various skin lesions are associated with AIDS, including disseminated infections and papulosquamous dermatoses, although not pigmented lesions. Involvement of the skin in Langerhans cell histiocytoses typically occurs in children; they produce reddish papules or nodules or erythematosus scaling plaques because the Langerhans cell infiltrates in the dermis. Skin lesions of mastocytosis in adults often exhibit urticaria. The erythematous rashes that develop in systemic lupus erythematosus are the result of antigen-antibody complex deposition and often exhibit photosensitivity.

PBD9 1151  PBD8 1175–1176

12 A Also known as an acrochordon or skin tag, a fibroepithelial polyp is a common lesion composed of a central core of fibrovascular connective tissue covered by normal-appearing squamous epithelium. They may become irritated but are otherwise incidental findings. A hemangioma forms a red nodule that is composed of vascular spaces in the upper dermis. A lentigo is a focus of melanocyte hyperplasia that produces a brown macule; in older individuals, lentigines are known as senile lentigines or age spots and are commonly found on the hands. Melasma is most often a masklike area of facial hyperpigmentation associated with pregnancy. A pilar (trichilemmal) cyst is an epithelial cyst that forms on the scalp. A xanthoma is a localized dermal collection of lipid-laden macrophages associated with hyperlipidemia.

PBD9 1152  PBD8 1176

13 C The epidermal inclusion cyst is the most common form of epithelial cyst, or wen, which is a cystic structure formed from downward growth of epithelium or expansion of a hair follicle. It is lined by squamous epithelium that desquamates keratinaceous debris into the center of the expanding cyst. Rupture of the cyst produces a local inflammatory reaction. Acne produces comedones, most typically on the face and upper trunk of adolescents and young adults. A dermatofibroma is typically solitary, firm, and dermally located. A fibroepithelial polyp is a skin tag that projects from the surface of the skin on a narrow pedicle. A trichoepithelioma is a form of a benign adnexal tumor, which is uncommon; it is a subcutaneous nodule usually seen on the head, neck, and upper trunk. Xanthomas are soft, yellow nodules that may form in the dermis from collections of lipid-laden macrophages in individuals with hyperlipidemia.

PBD9 1152  PBD8 1176

14 E Actinic keratoses are premalignant lesions associated with sun exposure. Older individuals are more likely to have actinic keratoses because of greater cumulative sun exposure, not because of aging alone. More extensive actinic keratoses may be treated with 5-fluorouracil applied topically. Hydrocortisone can alleviate the symptoms of many dermatologic conditions, but it cannot reverse actinic damage. Decreasing dietary fat is always a good idea, but it does not have much effect on the skin of the face. A surgical excision with wide margins is indicated for a more aggressive lesion, such as malignant melanoma. Many drugs can cause acute eczematous dermatitis and erythema multiforme, but not skin cancer. Antioxidants are touted for slowing the effects of aging, but they cannot reverse actinic damage.

PBD9 1154  BP9 862–863  PBD8 1178  BP8 850

15 E Risk factors for squamous cell carcinoma include ultraviolet light exposure, scarring from burn injury, irradiation, and immunosuppression. This patient was immunosuppressed to prevent graft rejection, and the immune dysregulation favors carcinogenesis in transplant patients. Squamous cell carcinomas also arise in rare disorders of DNA repair, such as xeroderma pigmentosum. A dermatofibroma is typically solitary, firm, and dermally located. Erythema multiforme is a hypersensitivity response to infections or drugs; the lesions have multiple forms, including papules,
macules, vesicles, and bullae. Lichen planus is a self-limited inflammatory disorder that manifests as “purple, pruritic, polygonal papules,” not as elevated ulcerated lesions. Psoriasis is an inflammatory dermatosis that can be associated with underlying arthritis, myopathy, enteropathy, or atherosclerotic heart disease.

**16 D** The appearance of a squamous cell carcinoma in a child in a sun-exposed area suggests the rare disorder xeroderma pigmentosum (XP), which is an inherited condition resulting from mutation in one of several XP genes. UV radiation causes cross-linking of pyrimidine residues and prevents normal DNA replication. This DNA damage is ameliorated by the nucleotide excision repair system. Persons with XP can have any of the malignancies related to sun exposure, so they stay indoors or use extensive sun protection. The other answers are not involved in this mechanism.

**17 B** Basal cell carcinomas arise as pearly papules on sun-exposed areas of the skin, particularly the face. They slowly infiltrate surrounding tissues, gradually enlarging. Although it rarely metastasizes, a basal cell carcinoma can have serious local effects, particularly in the area of the eye. An actinic keratosis is a premalignant lesion of the epidermis that does not invade surrounding tissue. A dermatofibroma is a benign lesion of the dermis composed of cells resembling fibroblasts. Melanomas are usually pigmented, and they are composed of polygonal or spindle cells that tend to grow in sheets and infiltrate to produce a grossly irregular border to the lesion. A nevus is a small, localized, benign lesion.

**18 A** Benign fibrous histiocytoma, or dermatofibroma, is an indolent lesion that may vary in size over time. It could be a localized response to trauma. The malignant counterpart that grows larger and is more aggressive is the dermatofibrosarcoma protubersans. Leiomyomas, benign smooth muscle neoplasms, can occur in many locations (presumably in vascular smooth muscle) but are rare in skin. Merkel cells are neuroendocrine cells derived from neural crest and can form the obscure Merkel cell tumor, which resembles a small cell carcinoma.

**19 B** Dermatofibrosarcoma protubersans is the malignant counterpart of dermatofibroma. The PDGFB gene is juxtaposed with the promoter region of COL1A1, leading to up-regulation of a growth-promoting factor. Imatinib mesylate can be employed to inhibit the PDGF receptor tyrosine kinase to control lesions that are recurrent or metastatic. FGFR3 mutations are more characteristic for seborrhic keratoses. KIT mutations are found in mast cell proliferations. Basal cell carcinomas have PTCH1 mutations. TSC1 mutations are found in tuberous sclerosis, with skin manifestations, including ash-leaf patches, shagreen patches, subungual fibromas, and angiofibromas.

**20 E** Urticaria pigmentosa is a localized form of mastocytosis. The cutaneous lesions often show the characteristic Darier sign on rubbing. Some patients may have systemic mastocytosis. Point mutations in the KIT oncogene may be present. The other immunologic cells listed, including lymphocytes, Langerhans cells, and macrophages, participate in inflammatory reactions, such as contact dermatitis. Macrophages and Langerhans cells are antigen-presenting cells. Merkel cells are neuroendocrine cells derived from neural crest and can form the obscure Merkel cell tumor, which resembles a small cell carcinoma.

**21 B** If the man eats the cookies, he will have hives, or urticaria, from an allergy to an antigen in the nuts. This causes a type I hypersensitivity reaction in which IgE antibodies are bound to the IgE receptor on mast cells. IgE-sensitized mast cells degranulate when the antigen is encountered. CD4+ helper lymphocytes are mainly part of more extended cell-mediated or humoral immune reactions in adaptive immune responses. Natural killer cells mediate antibody-dependent cell-mediated cytotoxicity and lyse major histocompatibility complex class I-deficient target cells. Neutrophils may become attracted to this site, but they are not the sensitized cells. Plasma cells secrete the IgE antibodies, but do not release the mediators for allergic reactions.

**22 A** An acute eczematous dermatitis is a common reaction to many skin irritants. The epidermis has prominent spongiosis. If the irritant is removed, the lesions resolve in a matter of days. Contact dermatitis is one form that occurs following prior exposure and sensitization. Did the herbal remedy help? Because the lesions will subside on their own, it is hard to prove. If remedies are not harmful and not expensive, then they can have benefit from a placebo effect, which can be powerful (and without adverse reactions). Traditional healers tend to spend more time with patients than do physicians, and if they can sort out the serious problems requiring pharmacologic or surgical approaches, then there is benefit for patient well-being. Erythema multiforme may also be a reaction to a chemical, often a drug, but it is a systemic effect, and the lesions often begin with a targetoid appearance. Impetigo often appears on the face, and the lesions crust, but it is due to an infection (usually staphylococci or streptococci). Eczema may turn into lichen simplex chronicus if the irritant is not removed and there is considerable scratching and rubbing of itchy lesions. Urticaria (hives) appears as wheals from a type I hypersensitivity response, typically from an insect bite or ingested food or substance.
23 B The time course of this case suggests a drug reaction producing an acute erythematous dermatitis. Sepsis rarely involves the skin as an erythematous dermatitis. Human papillomavirus infection is associated with formation of verruca vulgaris, the common wart. Photosensitivity may be enhanced by drugs, but ultraviolet light is the key component in light that produces photodermatitis, so photosensitivity is not likely to be encountered with indoor lighting in the hospital room. Most employers do not buy the more expensive light bulbs that mimic daylight, thereby increasing the prevalence of seasonal affective disorder and decreasing productivity by staff. Urticaria in type I hypersensitivity is not as severe or as long lasting.

24 D Erythema multiforme (EM) is a hypersensitivity response to certain infections and drugs, such as sulfonamides and penicillin. Other inciting factors for EM may include herpes simplex virus, mycoplasmal and fungal infections, malignant diseases, and collagen vascular diseases such as systemic lupus erythematosus. The other listed antiretroviral drugs or antimicrobials are far less likely to cause skin reactions. Acetaminophen and dapsone are less likely to produce EM, but many drugs can cause skin rashes and eruptions.

25 D She has psoriasis, and the appearance of lesions with trauma is known as the Koebner phenomenon. Though two thirds of affected persons have the HLA-Cw*0602 allele, only 10% of all persons with this allele develop psoriasis. There is abnormal CD4+ and CD8+ lymphocyte activation with release of many cytokines, including tumor necrosis factor, which mediate the skin damage. Atopy increases the risk for allergic reactions mediated by type I hypersensitivity reactions, often with urticarial skin lesions that dissipate within hours. Autoantibodies to desmoglein in pemphigus vulgaris cause suprabasal blister formation. Plant allergens such as urushiol in poison oak cause a contact dermatitis that typically fades in days. Herpes simplex virus infection can produce vesicular eruptions, or it may underlie erythema multiforme manifested by targetoid lesions.

26 D The lesions are those of psoriasis, driven by T lymphocyte elaboration of cytokines driving cellular proliferation so that the normal sequence of keratinocyte maturation is disrupted. Acanthosis refers to epidermal hyperplasia, in which the thickness of the entire epidermis is increased. Dyskeratosis refers to premature keratinization below the stratum granulosum. Hyperkeratosis is increased thickness of stratum corneum with abnormal keratin; a callus would refer to thickened stratum corneum in response to mechanical forces, such as the hands of a carpenter working with tools, or the soles of the feet of a child going barefoot. Spongiosis is edema between the cells of the epidermis.

27 E Psoriasis is a chronic skin condition with marked epithelial hyperplasia and parakeratotic scaling. CD4+ lymphocytes, as part of a T_{H1} and T_{H11} response, elaborate cytokines that promote cell proliferation, and CD8+ cells cause cellular damage. Nail changes, such as yellow-brown discoloration, pitting, dimpling, and separation of the nail plate from the nail bed (onycholysis), affect about one third of patients. Other manifestations of psoriasis include arthritis (resembling rheumatoid arthritis), myopathy, enteropathy, and atherosclerotic heart disease. A friction rub from fibrous pericarditis does not occur in psoriasis because mesothelial surfaces are not involved. Gastrointestinal mucosal involvement with hemorrhage is not a feature of psoriasis. Joint laxity with hyperextensibility is a feature of Ehlers-Danlos syndrome. Renal disease and hypertension are not typically the result of psoriasis.

28 D The classic “pruritic, purple, polygonal papules” of lichen planus are present, with the distinctive bandlike infiltrate of lymphocytes at the dermal-epidermal junction. Cytotoxic CD8+ cells are reacting to antigens in the basal layer and dermoepidermal junction. The lesions of lichen planus are typically self-limited, although the course can run for several years. Oral lesions may persist longer. There is risk for squamous cell carcinoma in chronic lesions. Although a lymphocytic infiltrate is present, an infection or autoimmunity is not implicated. A drug eruption would not last this long, and lichen planus is not a hypersensitivity reaction. Lesions of erythema multiforme are more likely to follow infections, drugs, autoimmune diseases, and malignancies. Skin adnexal tumors are typically benign; often have apocrine differentiation; and arise in areas were apocrine glands are prevalent, such as scalp and axilla. They are generally not associated with other diseases.

29 C Pemphigus vulgaris lesions are caused by IgG autoantibodies directed at an intercellular cement substance called desmoglein, giving a netlike appearance with immunofluorescence microscopy. The antibody deposition disrupts intercellular bridges, causing the epidermal cells to detach from each other (acantholysis). This action causes the formation of an intraepidermal blister. Staining with anti-IgG illuminates intercellular junctions at sites of incipient acantholysis. Darkfield microscopy is used almost exclusively to identify spirochetes in cases of syphilis. Some HLA types have an increased risk for some skin diseases, but this does not predict who will develop the disease. Type I hypersensitivity with IgE fixed to mast cells and urticaria does not produce an acantholytic vesicle. Herpes simplex viral infections can produce crops of vesicles, but such a wide distribution would be unusual.

30 A Subepidermal bullae of bullous pemphigoid usually heal without scarring. Subsequent oral lesions may appear. Most often seen in the elderly, this disease results from linear IgG deposition at the basal cell–basement membrane attach-
ment plaques (hemidesmosomes) containing bullous pemphigoid antigen (BPAG). The lamina densa of the basement membrane is not directly involved, and the actual blister of bullous pemphigoid forms in the lamina lucida. In contrast, the antibodies in pemphigus vulgaris attack the desmosomes that attach the epidermal keratinocytes. Antibodies directed against nuclear antigens are more typical for systemic autoimmune diseases such as systemic lupus erythematosus. The anti-idiotype antibodies of dermatitis herpetiformis cross-react with dermal reticulin, and there are microabscesses at the tips of dermal papillae.

31 C Dermatitis herpetiformis can accompany celiac disease. The IgA or IgG antibodies formed against the gliadin protein in gluten that is ingested (commonly in wheat, rye, and barley grains) cross-react with reticulin. Reticulin is a component of the anchoring fibrils that attach the epidermal basement membrane to the superficial dermis. This explains the localization of the IgA to the tips of dermal papillae and the site of inflammation. A gluten-free diet may relieve the symptoms. Bullous pemphigoid can occur in older individuals, with antibody directed at keratinocytes to produce flaccid bullae, but there is no association with celiac disease. Contact dermatitis is most likely to be seen on the hands and forearms. It is a type IV hypersensitivity reaction without immunoglobulin deposition and would not persist for 1 month. Discoid lupus erythematosus is seen on sun-exposed areas and has the appearance of an erythematosous rash. Erythema multiforme is a hypersensitivity response to infections and drugs; it produces macules and papules with a red or vesicular center, but it is probably mediated by cytotoxic lymphocytes and not by immunoglobulin deposition. Pemphigus vulgaris is an autoimmune disease in which IgG deposited in acantholytic areas forms vesicles that rupture to form erosions; it is not related to celiac disease.

32 C The more benign discoid lupus involves just skin, unlike systemic lupus erythematosus, but is still a form of type III hypersensitivity with antigen-antibody complex deposition along the basement membrane of the epidermis. The other listed options are not associated with sun exposure. Bullous pemphigoid lesions occur at the dermal-epidermal junction from antibody deposition targeting type VII collagen as a component of hemidesmosomes. Dermatitis herpetiformis associated with celiac disease has IgA antibodies deposited at tips of dermal papillae. Dysplastic nevi develop in relation to mutations in genes encoding for growth control proteins. Toxic epidermal necrolysis is a severe form of erythema multiforme mediated by cytotoxic CD8+ cells targeting epidermal basal cells. Early lesions of discoid lupus erythematosus appear as well-demarcated scaly purple macules or papules, and later expand into discoid plaques. Microscopically there is basal vacuolar degeneration, areas of epidermal atrophy, acanthosis, keratotic follicular plugging, basement membrane thickening, and superficial and deep perivascular lymphocytic infiltrates.

33 D Teenagers and young adults are affected by acne vulgaris more often than other age groups, and males are affected more often than females. Propionibacterium acnes organisms break down sebaceous gland oils to produce irritative fatty acids, and this may promote the process. The food on the cruise probably did not play a role, but stress causes the lesions to worsen. Staphylococcus aureus and group A streptococci are implicated in the inflammatory skin condition known as impetigo, which can include pustules and a characteristic pale yellow-brown crust. Herpes simplex virus produces vesicular skin eruptions, most often in a perioral or genital distribution. Mycobacterium leprae causes leprosy, which is a chronic condition that can produce focal depigmentation and areas of skin anesthesia.

34 C Erythema nodosum is a form of panniculitis that can be associated with infections, drug ingestion, inflammatory bowel disease, and malignancies, but an underlying condition is not always found. The inflammation primarily involves dermal adipose tissue. Acne vulgaris is most likely to appear on the face and upper trunk, centered around hair follicles; it often resolves with scarring. Celiac disease is associated with dermatitis herpetiformis, and the anti-idiotype antibodies can cross-react with dermal reticulin, producing microabscesses at the tips of dermal papillae. Impetigo is seen on the face and hands with crusting lesions from Staphylococcus aureus and β-hemolytic streptococcal infections producing subcorneal pustules. The lesions of molluscum contagiosum are firm nodules that microscopically contain pink cytoplasmic inclusions, called molluscum bodies.

35 B Verrucae, or warts, are quite common, particularly in children, and tend to persist for up to a couple of years. There are subtypes of human papillomavirus (HPV) associated with distinct clinical appearances: verruca vulgaris on the hands, verruca plana (flat wart) on the face and hands, verruca palmaris on the palms, verruca plantaris on the soles, and condyloma acuminatum (venereal wart) on genitalia. BRAF mutations can be present with melanocytic proliferations, such as dysplastic nevi. IgA deposition is found with dermatitis herpetiformis. Contact dermatitis is a form of type IV hypersensitivity. A photosensitive dermatitis may be seen with niacin deficiency.

36 G The warts in this patient are a common problem and result from infection with one of many types of human papillomavirus (HPV). They do not become malignant, and they tend to regress after several years. Condylomata acuminata, or genital warts, are caused by a type of HPV that is sexually transmitted; the lesions tend to be pink to white. A dermatofibroma forms a subcutaneous nodule, as does an intradermal nevus. Molluscum contagiosum is a self-limited condition with a nodular appearance. Seborrheic keratoses are pale brown to dark brown, slowly enlarging lesions with a rough surface that are most commonly found on the trunk and face
of older individuals. A squamous cell carcinoma continues to grow irregularly, invade adjacent tissues, and ulcerate.

40 D The small scabies mites burrow through the stratum corneum to produce the linear lesions, and the mites along with their eggs and feces produce intense pruritus. Scabies is easily transmitted by contact and typically occurs in community outbreaks. *Ixodes scapularis* is the tick that is the vector for *Borrelia burgdorferi* organisms, which cause Lyme disease and erythema chronicum migrans. *Molluscum contagiosum* is caused by a poxvirus that produces a localized, firm nodule. The erythematous macules and pustules of impetigo in children are often caused by staphylococcal and group A streptococcal infection. *Tinea corporis* is a superficial dermatophytic fungal infection that can produce erythema and crusting.

37 C The pink cytoplasmic inclusions, called *molluscum bodies*, are characteristic of this lesion. Immunocompromised individuals may have multiple, larger lesions. The infectious agent is a poxvirus. Disseminated fungal infections such as histoplasmosis involving skin are uncommon except in immunocompromised patients. Human papillomavirus (not a toad) is implicated in the appearance of verruca vulgaris, or the common wart. *Staphylococcus aureus* is a factor in the development of acne. The little hind cervical and anal squamous cell dysplasias.

38 E Impetigo is a superficial infection of skin that produces shallow erosions. These erosions are covered with exuded serum that dries to give the characteristic honey-colored crust. Cultures of the lesions of impetigo usually grow coagulase-positive *Staphylococcus aureus* or group A β-hemolytic *streptococcus*. The lesions are highly infectious. Acne vulgaris is typically seen during adolescence and produces pimples and pustules, but not crusts. Bullous pemphigoid can occur in older individuals with antibody directed at keratinocytes to produce flaccid bullae. Contact dermatitis is most likely to be seen on the hands and forearms. Erythema multiforme is a hypersensitivity response to infections and drugs that produces macules and papules with a red or vesicular center. Lichen planus appears as violaceous papules and plaques.

39 G Athlete’s foot is a common disorder resulting from superficial dermatophyte infection by various fungal species, including *Trichophyton*, *Epidermophyton*, and *Microsporum*. Infections that involve the foot produce the condition known as *tinea pedis*. Streptococcal and staphylococcal organisms cause impetigo, which is more likely to be seen on the face and hands. Herpetic infections first produce crops of clear vesicles, which may burst and form painful shallow ulcers. Human papillomavirus is best known as the cause of genital warts (condyloma acuminatum) and as a driving force behind cervical and anal squamous cell dysplasias. *Propionibacterium acnes* is a factor in the development of acne. The little eight-legged critters known as *Sarcoptes scabiei* crawl around in the stratum corneum, usually between the fingers, and cause itching, a process called *scabies*. 

41 C *Tinea versicolor* is a common condition caused by a superficial fungal infection of *Malassezia furfur*. The lesions can be lighter or darker than the surrounding skin. *Epidermophyton*, *Trichophyton*, and *Microsporum* genera are dermatophytic fungi best known as the cause of athlete’s foot and jock itch. Human papillomavirus is best known as the cause of genital warts (condyloma acuminatum) and as a driving force behind cervical and anal squamous cell dysplasias. *Mycobacterium leprae* is the cause of Hansen disease, which can manifest with areas of skin anesthesia that predispose to repeated trauma. Infection with *Propionibacterium acnes* is a factor in the development of acne. *Molluscum contagiosum* is the cause of scabies, which appears as pruritic reddish lesions.

42 D The findings are consistent with *tinea corporis*, a very common superficial fungal infection. The annular appearance with central clearing may be termed “ringworm,” but has nothing to do with helminths. The infection is often acquired from other humans, animals, and fomites. Although annoying and cosmetically displeasing, dermatophyte infections do not invade or disseminate.

The topical azoles for humans can be effective therapy for the animals, too. The dermatophyte in this case could be speciated as *Microsporum canis*, as the dog suggests. Bacterial cocci in chains are consistent with streptococcal infection that could be causing impetigo. Budding yeast cells suggest candidiasis, a common superficial infection, but are more likely found in warm, moist skin folds. *Molluscum contagiosum*, a poxvirus, produces a raised umbilicated papule. Cytoplasmic vacuolization is a typical viral effect that can be seen with common warts (verrucas).
1. When mechanical stress is placed upon bone, osteoprogenitor cells produce WNT proteins that bind to receptors on osteoblasts, increasing β-catenin. As a result, which of the following proteins is most likely to diminish osteoclast activity and increase bone formation?
   A. Bone morphogenetic protein
   B. Matrix metalloproteinase
   C. Nuclear factor kappa B
   D. Osteoprotegerin
   E. RANK ligand

2. A 39-year-old man on vacation is involved in a skiing accident in which he sustains a right tibial diaphyseal fracture. The fracture is set with open reduction and internal fixation for proper alignment. What is the most likely function of osteoclasts present at his fracture site 1 week later?
   A. Dividing mitotically
   B. Elaborating cytokines
   C. Forming collagen
   D. Resorbing bone
   E. Synthesizing osteoid

3. A 29-year-old woman, G3, P2, gives birth to an infant following an uncomplicated pregnancy. The infant’s height is below the fifth percentile. On physical examination, the infant’s torso and head size are normal, but the extremities are short. The forehead appears prominent. Radiographs show short, slightly bowed long bones, but no osteopenia. The other two children in the family are of normal height. The affected child has no difficulty with activities of daily living after modifications are made in the home and school for short stature, and later becomes a physician. Which of the following conditions is likely to be present in this child?
   A. Achondroplasia
   B. Hurler syndrome
   C. Osteogenesis imperfecta
   D. Rickets
   E. Scurvy
   F. Thanatophoric dysplasia

4. A 14-year-old girl who was normal at birth now has bilateral hearing loss. Audiometry indicates bilateral mixed conductive and sensorineural hearing loss. CT scan of the head shows maldevelopment of both middle ears with deficient ossification. Further history indicates that her dentist has tried various whiteners to diminish the yellow-brown color of her teeth, which have a slight bell-shaped appearance. The optometrist noted that her sclerae have a peculiar steel-gray color, and her vision is 20/40. At age 30 years, she falls and fractures the left femur. A radiograph shows that the femur is osteopenic. Bone densitometry reveals osteopenia of all measured sites. Which of the following molecular mechanisms is most likely to produce these findings?
   A. Deficient hypoxanthine-guanine phosphoribosyltransferase (HGPRT) activity
   B. Diminished osteoprotegerin binding to macrophage RANK receptor
   C. Failure of type I collagen formation by osteoblasts
   D. Fibroblast growth factor receptor 3 inhibition of cartilage proliferation
   E. Increased interleukin-6 production by osteoblasts
   F. Reduced number of vitamin D receptors
5 A 23-year-old primigravida notes decreased fetal movement, and a screening ultrasound at 18 weeks' gestation shows decreased fetal size. A stillborn is delivered at 25 weeks' gestation. At autopsy, a radiograph shows marked osteopenia and multiple bone fractures. Mutational analysis of fetal cells is most likely to show an abnormality involving which of the following genes?
- A COL1A1
- B EXT
- C FGFR3
- D FBN1
- E HGPR
- F RB

6 A 17-year-old primigravida gives birth prematurely to an infant small for gestational age. The infant has immediate respiratory distress. Newborn examination shows limb shortening, frontal skull bossing, and small thorax. A radiograph shows normal bone density without fractures. What is the most likely diagnosis?
- A Achondroplasia
- B Congenital syphilis
- C Osteogenesis imperfecta
- D Rickets
- E Thanatophoric dysplasia

7 A 2-year-old child has a history of multiple bone fractures with minor trauma. On examination he has hepatosplenomegaly and palsies involving cranial nerves II, VII, and VIII. Laboratory studies show pancytopenia. Radiographs reveal diffusely and symmetrically sclerotic bones with poorly formed metaphyses. Molecular analysis of his bone reveals a defect in production of carbonic anhydrase to solubilize hydroxyapatite crystal. He is treated with hematopoietic stem cell transplantation. Which of the following cells in his bones was most likely functionally deficient and replaced following transplantation?
- A Chondroblast
- B Chondrocyte
- C Osteoblast
- D Osteoclast
- E Osteocyte

8 A 77-year-old woman trips on the carpet in her home and falls to the floor. She immediately has marked pain in the right hip. On physical examination, there is shortening of the right leg with external rotation and marked pain with any movement. A radiograph shows a right femoral neck fracture. The fracture is repaired. Six months later, a dual-energy x-ray absorptiometry (DEXA) scan performed on the hip and lumbar vertebrae shows bone mineral density 2 standard deviations below the young adult reference range. Which of the following cellular processes contributes most to development of her findings?
- A Decreased secretion of interleukin-6 by monocytes
- B Increased sensitivity of osteocytes to parathyroid hormone
- C Insensitivity of bone matrix to 1,25-dihydroxycholecalciferol
- D Mutation in the fibroblast growth factor receptor 3 gene
- E Increased osteoclast activity
- F Synthesis of chemically abnormal osteoid matrix

9 A 35-year-old woman with active lupus nephritis falls forward and lands on her left hand. She has immediate pain. On examination there is crepitus at the wrist. A radiograph shows radial and navicular fractures along with marked osteopenia. Which of the following medications most likely contributed to the fracture?
- A Hydrocortisone
- B Ibuprofen
- C Lisinopril
- D Losartan
- E Methotrexate

10 An epidemiologic study of postmenopausal women is performed. The subjects undergo periodic examination by dual-energy x-ray absorptiometry (DEXA) scan performed on the hip and lumbar vertebrae to evaluate bone mineral density over the next 10 years. They respond to a survey regarding their past and present use of drugs, diet, activity levels, history of bone fractures, and medical conditions. A cohort of the subjects is identified whose bone mineral density is closest to that of the young adult reference range and in whom no bone fractures have occurred. Which of the following strategies is most likely to be supported by the study data to provide the best overall long-term reduction in risk of fracture in postmenopausal women?
- A Increasing bone mass with exercise during young adulthood
- B Limited alcohol use, and avoidance of the use of tobacco
- C Initiation of estrogen replacement therapy after a fracture
- D Supplementation of the diet with calcium and vitamin D after menopause
- E Use corticosteroid therapy for inflammatory conditions

11 A 70-year-old man complains of right hip and thigh pain of 8 months' duration. On physical examination, he has reduced range of motion in both hips, but there is no tenderness or swelling on palpation. Radiographs of the pelvis and right leg show sclerotic, thickened cortical bone with a narrowed joint space near the acetabulum. Laboratory studies show a serum alkaline phosphatase level of 173 U/L, calcium of 9.5 mg/dL, and phosphorus of 3.4 mg/dL. The figure shows the representative microscopic appearance of his pelvic bone. Which of the following conditions is most likely to produce these findings?
- A Degenerative osteoarthritis
- B Hyperparathyroidism
- C Osteochondroma
- D Paget disease of bone
- E Vitamin D deficiency
**12** An 82-year-old man has had progressively worsening lower back, bilateral hip, and right shoulder pain for the past 6 years. He reports that he has had to buy larger hats. On physical examination, there is no joint swelling, erythema, warmth, or tenderness, but the range of motion is reduced. Radiographs show narrowing of joint spaces with adjacent bony sclerosis. A skull radiograph shows thickening but diminished density of the skull bones. A bone biopsy specimen at the iliac crest shows a loss of normal trabeculae, with a mosaic pattern and increased numbers of osteoclasts and osteoblasts. Which of the following complications is the patient most likely to experience as a result of this condition?

A. Ankylosing spondylitis  
B. Enchondromatosis  
C. Fibrous dysplasia  
D. Osteoid osteoma  
E. Osteosarcoma

**13** A 38-year-old man has had chronic leg pain for the past 4 months. He passed a urinary tract calculus a month ago. On physical examination, there is local swelling with tenderness just below the right patella. A radiograph of the right lower leg shows a 4-cm cystic area in the right tibial diaphysis without erosion of the cortex or soft-tissue mass. Laboratory studies show serum calcium is 12.6 mg/dL, and phosphorus is 2.1 mg/dL. A biopsy specimen of the lesion is taken and microscopically shows increased osteoclasts and fibroblast proliferation. Which of the following underlying conditions is most likely to account for these findings?

A. Chronic glomerulonephritis  
B. Chronic osteomyelitis  
C. Giant cell tumor of bone  
D. Paget disease of bone  
E. Parathyroid adenoma

**14** A 28-year-old man flips over an all-terrain vehicle, and he lands on his leg. On physical examination there is intense pain on palpation over the right shin, but there is no shortening of the limb. The overlying skin is intact. A radiograph shows right tibial and fibular midshaft fracture into multiple bone fragments. Which of the following terms best describes these fractures?

A. Comminuted  
B. Compound  
C. Displaced  
D. Incomplete  
E. Pathologic

**15** A 26-year-old woman has had malaise, arthralgias, and myalgias for the past 2 months. On physical examination, there is no joint swelling or deformity. Laboratory studies show a serum creatinine level of 3.9 mg/dL. A renal biopsy specimen shows a proliferative glomerulonephritis. She receives glucocorticoid therapy for 3 months. She now has left hip pain with movement. On physical examination, there is no swelling or deformity. A radiograph of the left leg and pelvis shows patchy radiolucency and density of the femoral head with flattening of the bone. A total replacement of the left hip is performed, and gross examination of the sectioned femoral head shows collapse of articular cartilage over a pale, wedge-shaped, subchondral area. What is the most likely diagnosis?

A. Avascular necrosis  
B. Enchondroma  
C. Osteoarthritis  
D. Osteomyelitis  
E. Renal osteodystrophy

**16** A 9-year-old boy has had pain in the area of the right hip for the past 3 weeks. On physical examination, his temperature is 38.2° C. There is swelling with marked tenderness to palpation in the area of the right hip, pain, and reduced range of motion. Radiographs of the pelvis and legs show areas of osteolysis and cortical erosion involving the femoral metaphysis, with adjacent soft-tissue swelling extending from the subperiosteal region, and apparent abscess formation. Which of the following organisms is most likely to produce these findings?

A. *Haemophilus influenzae*  
B. *Neisseria gonorrhoeae*  
C. *Salmonella enterica*  
D. *Staphylococcus aureus*  
E. Group B streptococcus

**17** A 7-year-old boy sustained an open compound fracture of the right tibia and fibula in a fall from a barn loft to the floor below. On physical examination, the lower tibia and fibula can be seen protruding from the lower leg. The fracture is set by external manipulation, and the skin wound is sutured, but nothing more is done. One year later, he continues to have pain in the right leg, and a draining sinus tract has developed in the lateral lower right leg. A radiograph of the lower right leg is now most likely to show which of the following?

A. Cortical nidus with surrounding sclerosis  
B. Involucrum and sequestrum  
C. Osteolysis with osteosclerosis  
D. Soft-tissue hemorrhage and swelling  
E. Tumor mass with bony destruction

**18** A 39-year-old man has experienced back pain for 3 months. He has had a chronic cough for 2 years. On physical examination, there is tenderness to palpation over the lumbar vertebrae, but no warmth, swelling, or erythema. A radiograph of the spine shows a compressed fracture at the L2 level. CT scan of the abdomen shows an abscess involving the right psoas muscle. Infection with which of the following microbial agents is most likely to produce these findings?

A. *Cryptococcus neoformans*  
B. *Mycobacterium tuberculosis*  
C. *Shigella flexneri*  
D. *Staphylococcus aureus*  
E. *Streptococcus pyogenes*  
F. *Treponema pallidum*
19 A 12-year-old girl has had sudden onset of severe pain in her left knee that has awakened her from sleep on several occasions during the past 6 weeks. For each episode, her mother has given her acetylsalicylic acid (aspirin), and the pain has been relieved. On physical examination, there are no remarkable findings. A radiograph of the left knee shows a well-defined, 1-cm lucent area surrounded by a thin rim of bony sclerosis located in the proximal tibial cortex. The patient undergoes radioablation of the lesion, and the pain does not recur. What is the most likely diagnosis of this lesion?

A Enchondroma  
B Fibrous dysplasia  
C Giant cell tumor  
D Osteoblastoma  
E Osteochondroma  
F Osteoid osteoma

20 A 19-year-old man falls and strikes his leg. He feels intense pain. On physical examination there is swelling in his lower leg. MR imaging is performed and shown in the figure with T1 weighting. Which of the following is the most likely diagnosis?

A Chondrosarcoma  
B Ewing sarcoma  
C Giant cell tumor  
D Metastatic seminoma  
E Osteosarcoma

21 A 16-year-old boy has had pain around the right knee for the past 3 months. There are no physical findings except for local pain over the area of the distal right femur. A radiograph of the right leg shows an ill-defined mass involving the metaphyseal area of the distal right femur, and there is elevation of the adjacent periosteum. A bone biopsy specimen is obtained and on microscopic examination shows large, hyperchromatic, pleomorphic spindle cells forming an osteoid matrix. Which of the following tumor suppressor genes is most likely to be mutated in this boy?

A APC  
B BRCA1  
C NF1  
D PTEN  
E RB

22 A 23-year-old man has had pain in the area of the right knee for the past year. On physical examination, there is point tenderness in a 2-cm focal area just below the patella laterally over the tibia. A radiograph of the right leg shows a 3-cm, broad-based excrescence projecting from the metaphyseal region of the upper tibia. The lesion is excised. The figure shows the gross appearance of the sectioned lesion. What is the most likely diagnosis?

A Enchondroma  
B Fibrous dysplasia  
C Giant cell tumor  
D Osteoblastoma  
E Osteochondroma
23. A 38-year-old healthy man notes occasional pain in his right index finger when using a hammer. On palpation of the right proximal phalanx, there is point tenderness. A radiograph shows a 1-cm oval lucency in this phalanx, with a surrounding rim of bright, radiodense bone. The microscopic appearance of the excised lesion is shown in the figure. Which of the following cells is most likely to have given rise to this lesion?

A. Chondrocyte  
B. Giant cell  
C. Macrophage  
D. Osteoblast  
E. Plasma cell  
F. Fibroblast

24. A 30-year-old woman has experienced pain in the area of the left knee for 1 month. On physical examination, there is tenderness to palpation of the distal left thigh and knee. The area is firm, but there is no erythema or warmth. A radiograph of the left leg shows a 7-cm mass in the distal femoral epiphyseal area, with a “soap bubble” appearance. Microscopic examination of a curettage specimen of the lesion shows the findings in the figure. The lesion recurs in the next year; it is excised and does not recur again. What is the most likely diagnosis?

A. Chondrosarcoma  
B. Enchondroma  
C. Giant cell tumor  
D. Osteitis fibrosa cystica  
E. Osteoblastoma  
F. Plasmacytoma

25. A 45-year-old man has experienced pain in the area of the left hip and upper thigh for the past 7 months. On physical examination, there is tenderness on deep palpation of the left side of the groin. The range of motion at the left hip is decreased, but there is no swelling or warmth on palpation. Pelvic and left leg radiographs show an upper femoral metaphyseal mass lesion that erodes into the surrounding bone cortex. The proximal femur is excised and on sectioning has the gross appearance shown in the figure. Which of the following cell types is most likely to be proliferating in this mass?

A. Chondrocyte  
B. Osteoblast  
C. Osteoclast  
D. Plasma cell  
E. Primitive neuroectodermal cell

26. A 13-year-old, previously healthy boy has had pain in the right leg for the past month. There is no history of trauma or recent illness. On physical examination, there is warmth and tenderness to palpation of the right lower thigh anteriorly, and the circumference of the right thigh is slightly larger than that of the left. His temperature is 39°C. A radiograph of the right leg shows a 6-cm expansile mass in the diaphyseal region of the right lower femur that extends into the soft tissue and is covered by layers of reactive bone. A biopsy of the mass is done, and microscopic examination shows sheets of closely packed primitive cells with small, uniform nuclei and only scant cytoplasm. Karyotypic analysis of the tumor cells shows a t(11;22) translocation. What is the most likely diagnosis?

A. Chondrosarcoma  
B. Ewing sarcoma  
C. Giant cell tumor  
D. Metastatic carcinoma  
E. Osteosarcoma  
F. Plasmacytoma
A 14-year-old girl experiences severe pain in the right leg after performing a gymnastic floor exercise. On physical examination, there is marked pain on palpation of the right lower thigh just above the knee. Radiographs show a pathologic fracture across a 3-cm lower femoral diaphyseal lesion that has central lucency with a thin sclerotic rim. The lesion is completely intramedullary and well circumscribed. A bone biopsy specimen of the affected region is taken and microscopically shows scattered trabeculae of woven bone in a background of fibroblastic proliferation. What is the most likely diagnosis?

- A Ewing sarcoma
- B Fibrous dysplasia
- C Fracture callus
- D Osteogenic sarcoma
- E Osteoid osteoma

A 75-year-old woman has experienced increasing dull but constant pain in the back, right chest, left shoulder, and left upper thigh for the past 6 months. She has now developed a sudden, severe, sharp pain in the left thigh. On physical examination, she has intense pain on palpation of the upper thigh, and the left leg is shorter than the right. A radiograph of the left leg shows a fracture through the upper diaphyseal region of the femur in a 5-cm lytic area that extends through the entire thickness of the bone. A bone scan shows multiple areas of increased uptake in the left femur, pelvis, vertebrae, right third and fourth ribs, upper left humerus, and left scapula. Laboratory studies show serum creatinine, 0.9 mg/dL; total protein, 6.7 g/dL; albumin, 4.5 g/dL; total bilirubin, 1 mg/dL; AST, 28 U/L; ALT, 22 U/L; and alkaline phosphatase, 202 U/L. What is the most likely diagnosis?

- A Hyperparathyroidism
- B Metastatic carcinoma
- C Multiple myeloma
- D Osteochondromatosis
- E Paget disease of bone
- F Polyostotic fibrous dysplasia

A 78-year-old woman has had a constant, dull pain in her back that has persisted for more than a month. She is in no acute distress. On physical examination, there are no abnormal findings. Laboratory findings include creatinine, 0.9 mg/dL; urea nitrogen, 17 mg/dL; total protein, 6.8 g/dL; albumin, 4.2 g/dL; total bilirubin, 0.8 mg/dL; AST, 25 U/L; ALT, 29 U/L; calcium, 10.8 mg/dL; phosphorus, 2.3 mg/dL; and alkaline phosphatase, 228 U/L. What is the most likely cause of hypercalcemia?

- A Defects in articular chondrocyte function
- B Gene mutation involving the synthesis of type I collagen
- C Infiltration of the synovial membrane by activated CD4+ T cells
- D Inheritance of HLA-B27 genotype
- E Partial deficiency of hypoxanthine-guanine phosphoribosyltransferase (HGPRT)

A 83-year-old woman has pain and limitation of movement affecting the right hip joint, worsening for the past 15 years. Physical examination shows a nodular bony outgrowth can be felt in the distal interphalangeal joint of the right index finger. The right hand is normal. There is no evidence of systemic disease, and cardiovascular and respiratory findings are unremarkable. A radiograph of the affected hip shows narrowing of the joint space and subchondral sclerosis. Laboratory studies do not show rheumatoid factor or antinuclear antibodies. The serum uric acid level is 5 mg/dL. Which of the following factors is most important in the pathogenesis of her disease?

- A Defects in articular chondrocyte function
- B Gene mutation involving the synthesis of type I collagen
- C Infiltration of the synovial membrane by activated CD4+ T cells
- D Inheritance of HLA-B27 genotype
- E Partial deficiency of hypoxanthine-guanine phosphoribosyltransferase (HGPRT)

A 47-year-old woman has had increasing pain and deformities in her hands for the past 10 years. On physical examination, she has metacarpophalangeal joint swelling, erythema, and tenderness. There is a subcutaneous nodule on the ulnar aspect of the right forearm. A biopsy specimen of the nodule has the microscopic appearance depicted in the figure. Which of the following therapies is most likely to be effective in this patient?

- A Bisphosphonates
- B Broad-spectrum antibiotics
- C Anti-TNF agents
- D Uricosuric agent cell cycle inhibitors
33 A 34-year-old woman has experienced malaise, fatigue, and joint pain for the past 5 months. She has had progressive loss of joint motion, making it more difficult to walk and to use her hands. On physical examination, the joint involvement is symmetric, and most of the affected joints are in the hands and feet. The involved joints are swollen and warm to the touch. The second and third digits on the right hand have a swan neck deformity, and there is ulnar deviation of both hands. Reconstructive surgery is performed on her right hand. Microscopic views of the excised joint capsule tissue are shown in the figure. Which of the following serologic laboratory test findings is most likely to be positive in this patient?

A. Anti-DNA topoisomerase I antibody
B. Antinuclear antibody
C. Borrelia burgdorferi
D. Campylobacter jejuni
E. Citrullinated peptides

34 A 10-year-old girl has developed worsening pain in the knees and ankles for the past 3 months and now has difficulty walking. On physical examination, these joints are swollen and warm to the touch, and there is diffuse muscle tenderness. She has a temperature of 39.2°C. There is an erythematous skin rash across the bridge of her nose and on the dorsa of her hands. Hepatosplenomegaly is present, and a friction rub is audible on auscultation of her chest. A joint aspirate is obtained from the left knee, and a microbiological culture of the fluid is negative. On microscopy, the joint fluid has increased numbers of lymphocytes, but few neutrophils. Her condition improves over the next year, and she has no residual joint deformity. Which of the following laboratory findings is most likely to be found in this patient?

A. ANA serologic titer of 1:1024
B. Borrelia burgdorferi serology positive
C. Chlamydia trachomatis urine culture positive
D. Ferritin level in serum of 7245 ng/mL
E. Hemoglobin S on hemoglobin electrophoresis
F. Rheumatoid factor serologic titer of 1:512
G. Serologic test for syphilis (STS) positive
H. Uric acid level in serum of 15.8 mg/dL

35 A 35-year-old Sami man from Finland has increasing lower back pain for 6 years. On examination he has decreased mobility of his spine. Radiographs show loss of lumbar lordosis, narrowing with sclerosis of sacroiliac joints bilaterally, and syndesmophyte formation at the junction of vertebrae and annulus fibrosus of intervertebral discs. Which of the following complications is most likely to develop in this man?

A. Glomerulonephritis
B. Hepatitis
C. Malabsorption
D. Meningitis
E. Uveitis

36 A 30-year-old man has had cramping abdominal pain and bloody diarrhea for the past 4 days. On physical examination, there is diffuse tenderness on palpation of the abdomen. Bowel sounds are present. There are no masses and no organomegaly. A stool culture is positive for Shigella flexneri. The episode resolves spontaneously within 1 week after onset. Six weeks later, he has increasingly severe lower back pain. Physical examination now shows stiffness of the lumbar spine and tenderness of the sacroiliac joints. He is treated with nonsteroidal anti-inflammatory agents. Two months later, the back pain recurs, and he complains of redness of the right eye and blurred vision. Serologic testing for which of the following is most likely to be positive in this patient?

A. Borrelia burgdorferi
B. Chlamydia trachomatis
C. Epstein-Barr virus
D. HLA-B27
E. Rheumatoid factor
A 47-year-old woman has been bothered for 20 years by recurring skin lesions over the elbows, knees, scalp, and lumbar-sacral area. These skin lesions are silvery to salmon-colored, 1- to 4-cm plaques with scaling. She has had increasing pain in her left hand and in her hips, more prominent on the left, over the past 2 years. On physical examination, she has yellow-brown discoloration with pitting of the fingernails. The distal interphalangeal joints of the left hand are slightly swollen and tender. There is minimal reduction in left hip mobility and no swelling or warmth to the touch. A radiograph of the left hip shows minimal joint space narrowing and surface erosion. Bone density is normal for age. During the next 10 years, the joint pain persists, but there is no joint destruction or deformity. She continues to have the same skin lesions. Which of the following is most likely to be seen on a biopsy specimen of her skin lesions?

- A Bandlike upper dermal infiltrate of lymphocytes
- B Epidermal spongiosis with dermal edema and eosinophils
- C Epidermal thinning with hyperkeratosis and parakeratosis
- D Focal keratinocyte apoptosis
- E IgG deposited along the dermal-epidermal junction

A 15-year-old boy has been hospitalized multiple times since childhood as a result of painful abdominal crises. He has had pain in his right hip region for the past week. On physical examination, there is marked tenderness and swelling to palpation over the right hip. Laboratory studies show hemoglobin of 8.5 g/dL, hematocrit of 25.7%, platelet count of 199,900/mm³, and WBC count of 12,190/mm³. Examination of the peripheral blood smear shows sickled erythrocytes, and nucleated RBCs. A radiograph of the pelvis and right upper leg shows acute inflammatory changes in the femoral head and metaphysis of the right proximal femur. Which infectious agent is most likely responsible for his condition?

- A Treponema pallidum
- B Neisseria gonorrhoeae
- C Salmonella enterica
- D Staphylococcus aureus
- E Streptococcus pneumoniae

A 55-year-old, previously healthy man has had episodes of pain and swelling of the right first metatarsophalangeal joint for the past year. These flare-ups usually occur after consumption of alcohol, typically port wine (Six Grapes). On physical examination, there is exquisite tenderness with swelling and erythema of the right first metatarsophalangeal joint. A joint aspiration is performed, and polarized light microscopy shows the finding in the figure, and many neutrophils in a small amount of fluid. Which of the following infectious agents is most likely responsible for his condition?

- A Treponema pallidum
- B Neisseria gonorrhoeae
- C Salmonella enterica
- D Staphylococcus aureus
- E Borrelia burgdorferi

A 27-year-old man develops acute pain and swelling of the left knee 5 days after an episode of urethritis. On physical examination, the left knee is swollen, warm, and tender to the touch. No other joints are affected. Laboratory examination of fluid aspirated from the left knee joint shows numerous neutrophils. A Gram stain of the fluid shows gram-negative intracellular diplococci. No crystals are seen. Which of the following infectious agents is most likely responsible for his condition?

- A Borrelia burgdorferi
- B Haemophilus influenzae
- C Neisseria gonorrhoeae
- D Staphylococcus aureus
- E Treponema pallidum

A 48-year-old woman had chronic pain of the left shoulder and right hip for 8 months. The pain resolved within 1 month. Two months later, she developed pain in the right knee and ankle, which resolved within 6 weeks. On physical examination, she is now afebrile. There is pain on movement of the left shoulder and right hip. A radiograph of the left arm shows extensive bony erosion of the humeral head. A biopsy specimen of synovium is taken and on microscopic examination shows a marked lymphoplasmacytic infiltrate and arteritis with endothelial proliferation. Which of the following infectious agents is most likely responsible for these findings?

- A Borrelia burgdorferi
- B Group B streptococcus
- C Mycobacterium tuberculosis
- D Neisseria gonorrhoeae
- E Treponema pallidum

A 8-year-old boy complains of left leg pain for 3 days. On physical examination, his temperature is 38.9° C, and he exhibits irritability when his left leg is moved. A radiograph of the left leg shows changes suggesting acute osteomyelitis in the proximal portion of the left femur. Culture of the infected bone is most likely to grow which of the following organisms?

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- B Haemophilus influenzae
- C Neisseria gonorrhoeae
- D Staphylococcus aureus
- E Treponema pallidum
43. A 51-year-old man has endured episodes of intense local pain involving his left foot for the past 4 months. The pain may last hours to days. Physical examination identifies the right metatarsophalangeal (MP) joint as the focus of tenderness and swelling, but minimal loss of joint mobility. A painless 2-cm nodule with overlying ulcerated skin is present on the lateral aspect of the MP joint. Beneath the eroded skin is a chalky white deposit of soft material. A firm 1-cm subcutaneous nodule on the extensor surface of the left elbow is excised and has the microscopic appearance shown in the figure. Which of the following mechanisms is most important in causing joint injury in this man?

A. Activation of neutrophils by phagocytosis of urate crystals
B. Release of TNF causing acute joint inflammation
C. Deposition of serum cholesterol into the synovium
D. Granulomatous inflammation with Mycobacterium tuberculosis infection
E. Reduced metabolism of homogentisic acid

44. A 58-year-old man with a diagnosis of chronic myeloid leukemia is treated with intensive chemotherapy. He goes into remission, but develops pain in the left wrist. On physical examination, there is swelling and warmth on palpation of the wrist. Polarized light microscopy of fluid aspirated from the wrist joint shows needle-shaped crystals that display negative birefringence. Which of the following processes most likely played an important role in the pathogenesis of the patient’s wrist pain?

A. Abnormal platelet function with joint hemorrhages
B. Chemotherapeutic toxicity to the articular cartilage
C. Cytokine-induced synovial proliferation
D. Excessive production of uric acid
E. Leukemic cell infiltration of the synovium

45. A 48-year-old man has had increasing pain in the left knee for the past 4 years, but the pain has become worse in the past week. On physical examination, the left knee is slightly swollen and warm to the touch. The cell count of a joint aspirate shows increased neutrophils. The figure shows a smear preparation of the fluid examined with polarized light microscopy (arrow in axis of red compensator). The patient experiences reduced knee joint mobility over the next 5 years. He also develops congestive heart failure, diabetes mellitus, and hepatic cirrhosis. Which of the following laboratory findings is most characteristic of this disease process?

A. ANA serologic titer of 1:1024
B. Ferritin level in serum of 7245 ng/mL
C. Rheumatoid factor serologic titer of 1:512
D. Serologic test for syphilis (STS) positive
E. Uric acid level in serum of 15.8 mg/dL

46. A 33-year-old woman has been bothered by a bump on the dorsum of her left wrist for the past 4 months. On physical examination, there is a 1-cm firm but fluctuant subcutaneous nodule over an extensor tendon of the left wrist. The nodule is painful on palpation and movement. Mucoid fluid is aspirated from the nodule. What is the most likely diagnosis?

A. Ganglion cyst
B. Giant cell tumor
C. Lipoma
D. Nodular fasciitis
E. Rheumatoid nodule
F. Tophus

47. A 25-year-old man has had right knee pain with “popping and catching” for the past 2 years. On examination there is reduced range of motion at the right knee, but no tenderness or swelling. MR imaging reveals a 3-cm mass in the anterior joint space. Arthroscopic surgery with partial synovectomy is performed, and a nodular, encapsulated mass is removed. Microscopic examination of the mass shows synoviocyte-like tumor cells in a hyalinized stroma containing osteoclast-like giant cells and hemosiderin. An antagonist to which of the following growth factors is most useful in treating this lesion?

A. EGF
B. FGF
C. M-CSF
D. PDGF
E. TGF-β
F. VEGF
A 33-year-old man has noticed a lump over his right flank. The lump is painless and has enlarged slowly over the past 3 years. On physical examination, a soft 2-cm nodule is palpable in the subcutis of the right flank above the iliac crest. The lesion is excised. Grossly, it is circumscribed and has a uniformly yellow cut surface. Which of the following is the most likely prognosis for this lesion?

A  Antibiotic therapy will be needed  
B  Family members will develop similar lesions  
C  Metastases to regional lymph nodes will occur  
D  More skin lesions will occur over time  
E  No recurrence is expected

A 36-year-old woman has noted a nodule beneath the skin in her left groin since adolescence. On physical examination, the lesion has a 2-cm diameter and is nontender, soft, rubbery, and movable. Which of the following cell types is most likely to comprise this lesion?

A  Adipocyte  
B  Endothelial cell  
C  Fibroblast  
D  Skeletal muscle  
E  Smooth muscle

A 47-year-old man has had dull, constant pain in the midsection of the right thigh for the past 4 months. On physical examination, there is pain on palpation of the anterior right thigh, which worsens with movement. The right thigh appears to have a larger circumference than the left thigh. A radiograph of the right upper leg and pelvis shows no fracture, but there is an ill-defined soft-tissue mass anterior to the femur. MRI shows a 10 × 8 × 7 cm solid mass deep to the quadriceps, but it does not involve the femur. Karyotypic analysis of tumor cells reveals t(12;16)(q13;p11) with amplification of MDM2 gene. What is the most likely diagnosis?

A  Chondrosarcoma  
B  Liposarcoma  
C  Metastatic adenocarcinoma  
D  Nodular fascitis  
E  Osteosarcoma  
F  Rhabdomyosarcoma

A 26-year-old man is struck in the left arm by a swinging steel beam at a construction site. On physical examination, a 4-cm area of the lateral upper left arm exhibits swelling and redness with pain on palpation. A radiograph of the left arm shows no fracture. Three weeks later, there now is a 2-cm painful, well-circumscribed, subcutaneous mass at the site of the original injury. A radiograph shows a solid soft-tissue mass. Which of the following lesions is most likely to be present in this man?

A  Lipoma  
B  Nodular fascitis  
C  Organizing abscess  
D  Pleomorphic fibroblastic sarcoma  
E  Superficial fibromatosis

A 57-year-old woman has noticed increasing deformity and difficulty with movement involving her left hand over the past 6 months. On physical examination, there is a contracture involving the third digit of her left hand that prevents her from fully extending this finger. A firm, hard, cordlike 1 × 3 cm area is palpable beneath the skin of the left palm. Microscopically, which of the following is most likely to be seen in greatest abundance composing this lesion?

A  Atypical spindle cells  
B  Collagen  
C  Dystrophic calcification  
D  Granulation tissue  
E  Lipoblasts

A 6-year-old boy complains of discomfort in the right upper neck that has worsened over the past 6 months. On physical examination, a 5-cm firm mass is palpable in the right lateral neck. The mass is not tender or warm. A biopsy is taken and the microscopic appearance of this mass is shown in the figure. Immunohistochemical staining for which of the following antigens is most likely to be positive in the cells of this lesion?

A  CD3  
B  Cytokeratin  
C  Dystrophin  
D  Neuron-specific enolase  
E  Vimentin

A 66-year-old woman has experienced pain in the area around the left knee for the past 6 weeks. She can recall no trauma to the leg. On physical examination, no mass is palpable; there is no warmth or swelling, and there is no loss of range of motion. MRI shows a well-circumscribed 4-cm mass superior and inferior to the patella. The mass is within soft tissue, without bony erosion. A biopsy of the mass is obtained and on microscopic examination shows a biphasic pattern of spindle cells and epithelial cells forming glands. Karyotypic analysis of tumor cells shows a t(X;18) translocation. What is the most likely diagnosis?

A  Desmoid tumor  
B  Leiomyosarcoma  
C  Mesothelioma  
D  Osteoblastoma  
E  Synovial sarcoma
ANSWERS

1 D Bone remodeling is a balance between osteoblastic and osteoclastic activity. The decoy protein osteoprotegerin (OPG) can bind RANK ligand to reduce its binding to RANK and reduce the nuclear factor kappa B transcription factor and diminish osteoclast activity. The WNT/β-catenin pathway can increase OPG production to tip the balance in favor of bone formation. Bone morphogenetic proteins oppose fibroblast growth factors to decrease chondrocyte proliferation. Matrix metalloproteinases generated from osteoclasts promote bone resorption.

PBD9 1180–1183  BP9 765–766  PBD8 1206–1209  BP8 765–766

2 D Bone remodeling is accomplished when osteoblasts produce new bone, and osteoclasts resorb it. This is a normal, slow, ongoing process in concert with all bones, but the process is accelerated in fracture callus. Osteoprogenitor cells give rise to osteoblasts. Osteoclasts are derived from the same hematopoietic progenitor cells that give rise to macrophages and monocytes. Terminally differentiated osteoclasts do not undergo mitosis. Cytokines can be elaborated by macrophages and other inflammatory cells within the callus. Collagen is produced by fibroblasts. Osteoid is produced by osteoblasts.

PBD9 1180–1183  BP9 765–766  PBD8 1219–1220  BP8 765–766

3 A Achondroplasia is most often the result of a spontaneous new mutation in the fibroblast growth factor receptor 3 (FGFR3) gene, leading to abnormal cartilage proliferation at growth plates and affecting mainly endochondral bone growth. The homozygous form is lethal in utero. In Hurler syndrome, mucopolysaccharidoses type I, children are normal at birth but then develop growth retardation, mental retardation, hepatosplenomegaly, and joint stiffness. Osteogenesis imperfecta may manifest at birth with multiple fractures from severe osteopenia as a result of abnormal type I collagen synthesis. Rickets may occur in childhood from vitamin D deficiency, producing deficient bone mineralization, but dwarfism is not a feature. Vitamin C deficiency causes scurvy and can lead to abnormal bone matrix with mild deformity, but not dwarfism. Thanatophoric dysplasia is the most common form of lethal dwarfism and also results from a new mutation in FGFR3.

PBD9 1183–1185  BP9 767  PBD8 1210–1211  BP8 803

4 C The patient has osteogenesis imperfecta, most likely type I. Type II is the perinatal lethal form, which causes death in utero, at birth, or shortly thereafter. Type III is seen in children and adults and is more severe than type I. Type IV is difficult to distinguish from type III. Osteogenesis imperfecta causes osteopenia (“brittle bones”) and predisposes to fractures. Patients often have “blue” sclerae, dental abnormalities, and progressive hearing loss. Absence of HGPR, an X-linked disorder known as Lesch-Nyhan syndrome and characterized by hyperuricemia. Decreased binding of osteoprotegerin to the macrophage RANK receptor is part of the mechanism of osteoporosis. Mutations in the fibroblast growth factor receptor 3 (FGFR3) gene can be seen in achondroplasia. Increased interleukin-6 production by osteoblasts occurs in Paget disease of bone and with postmenopausal decrease in estrogen, causing increased bone loss. Decreased absorption of vitamin D in the small intestine leads to rickets in children; glucocorticoids can decrease receptor expression in many tissues, including intestine.

PBD9 1185–1186  BP9 767  PBD8 1211–1222  BP8 802–803

5 A The stillborn has evidence for the type II lethal variant of osteogenesis imperfecta, with a defect in collagen 1 formation leading to multiple fractures in utero with long bone shortening. EXT mutations are seen in cases of hereditary and sporadic osteochondromas. The FGFR3 gene is implicated in cases of achondrogenesis with long bone shortening, but without osteopenia and fractures. Fibrillin-1 (FBN1) gene mutations are seen with Marfan syndrome, which severely affects the cardiovascular system, including aortic dissection. Hypoxanthine-guanine phosphoribosyltransferase (HGPRT) is an enzyme in the purine salvage pathway, and is associated with the X-linked Lesch-Nyhan syndrome with hyperuricemia. The tumor suppressor gene RB is associated with many cases of osteosarcoma.

PBD9 1185–1186  BP9 767  PBD8 1211–1212  BP8 802–803

6 E Thanatophoric dwarfism is a rare condition resulting from FGFR3 gene mutations. The markedly reduced thoracic size leads to pulmonary hypoplasia, the rate-limiting step to survival. The other listed options can lead to appearance of bone abnormalities at birth, but with the exception of the perinatal lethal form of osteogenesis imperfecta, are not life threatening.

PBD9 1184–1185  BP9 767  PBD8 1210  BP8 803

7 D Osteopetrosis (Albers-Schönberg disease) is a rare bone disease resulting from mutations in genes that regulate osteoclast activity and bone resorption. Carbonic anhydrase generates the protons used by the H+-ATPase proton pump located on the osteoclast ruffled border. Marrow is reduced within the sclerotic bone, with subsequent extramedullary hematopoiesis in spleen and liver. Nerve compression in narrowed bony foramina leads to the palsies. The other cells listed are derived from mesenchymal stem cells, not hematopoietic stem cells. Osteopetrosis is the opposite of osteoporosis. Remember that in osteoporosis the osteoclast formation and action are enhanced when osteoblasts produce less osteoprotegerin and M-CSF, and increased RANK and RANKL interaction promote osteoclast differentiation and survival.

PBD9 1186–1187  BP9 767–768  PBD8 1212–1214  BP8 803

8 E With advancing age, the ability of osteoblasts to divide and to lay down osteoid is reduced, whereas in some persons osteoclast activity increases, giving rise to accelerated bone loss known as osteoporosis. Differentiation
of stromal progenitor cells into osteoclasts requires binding of RANK ligand on osteoclasts to RANK receptor on osteoclast precursors and stimulation by M-CSF produced by osteoblasts. Osteoprotegerin (OPG) is a “decoy receptor” for RANK ligand that slows osteoclast formation and action. When osteoclasts produce less OPG, bone loss is accelerated. Marrow stromal cells produce WNT proteins that bind to osteoblast receptors, activating β-catenin and OPG production. Postmenopausal osteoporosis is characterized by hormone-dependent acceleration of bone loss. Estrogen deficiency results in increased secretion of interleukin-1, interleukin-6, and tumor necrosis factor-α by monocytes-macrophages. These cytokines act by increasing the levels of RANK and RANKL, and decreasing the levels of OPG. In older women, bone loss is accelerated by reduced synthesis and increased resorption. Nonhormonal drugs such as bisphosphonates are designed to reduce osteoclast resorption. There are no age-associated changes in the sensitivity to vitamin D or parathyroid hormone action or composition of osteoid. Fibroblast growth factor receptor 3 gene mutations occur in dwarfism syndromes, such as achondroplasia.

9 A Corticosteroids may be needed to treat severe lupus nephritis, and continuing therapy increases the risk for Cushings syndrome with osteoporosis. Corticosteroids stimulate RANKL expression, inhibit osteoblast osteoprotegerin (OPG) synthesis, and thereby enhance osteoclast proliferation to promote bone resorption. Ibuprofen and other nonsteroidal anti-inflammatory drugs (NSAIDs) should not be used with renal failure. Lisinopril, an angiotensin-converting enzyme (ACE) inhibitor, and losartan, an angiotensin receptor blocker (ARB), may be used to treat hypertension with renal disease, but they do not directly affect bone density. Methotrexate is a chemotherapy agent that is best used with rheumatoid arthritis, not systemic lupus erythematosus, and at low doses is less likely to lead to osteoporosis.

10 A The total bone mass is an important determinant of the subsequent risk of osteoporosis and its complications. A proactive regimen of exercise that puts stress on bones to increase mass before the inevitable loss after age 30 years is most likely to reduce the subsequent risk of osteoporosis. Alcohol and tobacco use are not major risks for osteoporosis. Postmenopausal estrogen or raloxifene therapy can help preserve bone mass; however, by the time a fracture has occurred, there has already been significant bone loss. Likewise, dietary supplements after menopause are not harmful, but at best only partially slow the loss of bone that accompanies aging. Corticosteroid therapy is just one of many risk factors for osteoporosis, but short courses of corticosteroids have minimal effects on bone formation.

11 D The mosaic pattern of lamellar bone in the figure is characteristic of osteitis deformans (Paget disease of bone).

This disease has three phases. Early in the course, there is a lytic phase, followed by the more classic mixed phase of osteosclerosis and osteolysis, leading to the appearance of a “mosaic” of irregular bone. A sclerotic “burnt-out” phase then ensues. Elderly white people are most often affected, and the disease progresses over many years. Joints adjacent to affected bone manifest osteoarthritis with chronic pain from joint erosion. Osteitis fibrosa cystica is seen in hyperparathyroidism. An enchondroma is a tumorlike projection of bone capped by cartilage that protrudes from the metaphyseal region of a long bone. Osteomalacia results in osteopenia from vitamin D deficiency in an adult.
15 A Avascular necrosis of bone (osteonecrosis) represents a localized area of bone infarction, most often in a metaphyseal medullary cavity or subchondral epiphyseal location. The femoral head is most often affected. Underlying conditions associated with osteonecrosis include hemoglobinopathies (sickle cell disease in particular), fracture, barotrauma, hypercoagulable states, and hyperlipidemia. Glucocorticoid therapy decreases osteoblastogenesis to promote avascular necrosis, as in this patient with systemic lupus erythematosus and glomerulonephritis. An enchondroma is a benign tumor of hyaline cartilage that arises in the medullary space of young adults. Osteoarthritis may produce some cartilaginous erosion, but not collapse or bone infarction. Osteomyelitis typically is not so localized, and there is irregular new bone formation (involucrum). The patient’s course is quite short for renal osteodystrophy, which is mediated through chronic renal failure and produces lesions such as osteitis fibrosa cystica. PBD9 1194–1195 BP9 773 PBD8 1220–1221 BP8 809–810

16 D Pyogenic osteomyelitis may arise from hematogenous dissemination of an infection. In children with no history of previous illnesses, Staphylococcus aureus is the most common causative organism. Haemophilus influenzae and group B streptococcal infections are most common in the neonatal period. Gonorrhea occasionally may disseminate and involve the bones (osteoarthritis) or joints (septic arthritis) of sexually active individuals. Salmonella infection involving bone is infrequent, except in patients with sickle cell anemia. PBD9 1195–1196 BP9 773–774 PBD8 1221–1222 BP8 810

17 B This patient has chronic osteomyelitis. The most likely sequence of events is the occurrence of a compound fracture that became infected by direct extension of bacteria into the bone. The subsequent care for this patient was inadequate, and he developed chronic osteomyelitis. Infection of the bone and the associated vascular compromise caused bone necrosis, giving rise to a dead portion of bone, called sequestrum. With chronicity, a shell of reactive new bone, called involucrum, is formed around the dead bone. A nidus with surrounding sclerosis suggests an osteoid osteoma. Osteolysis and osteoarthritis are features of bone remodeling with Paget disease of bone. Soft-tissue hemorrhage and swelling should be minimal and resolve soon after the fracture is stabilized. A mass suggests a malignancy, and the most common neoplasm to develop in a sinus tract draining from osteomyelitis is squamous cell carcinoma, but this is uncommon. PBD9 1195–1196 BP9 773–774 PBD8 1221–1222 BP8 810

18 B The presence of a destructive lesion in the vertebrae with extension of the disease along the psoas muscle, without signs of acute inflammation (cold abscess) is characteristic of tuberculous osteomyelitis. Tuberculosis of bones usually results from hematogenous spread of an infection in the lung. Long bones and vertebrae are the favored sites for tuberculosis involving the skeletal system. Dissemination of Cryptococcus neoformans infection from the lungs occurs most commonly in immunocompromised patients, but frequently produces osteomyelitis. Shigella species are unlikely to cause osteomyelitis but may be associated with reactive arthritis. Staphylococcal osteomyelitis is more common than streptococcal osteomyelitis, but both organisms are more likely to involve long bones and produce an acute hot joint. Treponema pallidum may produce gummatus necrosis in the tertiary stage, but this involves soft tissues more than bone. PBD9 1196 BP9 774 PBD8 1222–1223 BP8 811

19 F An osteoid osteoma is a benign tumor of the bone with a central nidus of woven bone and sclerotic rim. It most often occurs in children and young adults. Pain disproportionate to the size of the tumor is characteristic. If such a lesion is larger than 2 cm, it is classified as an osteoblastoma, and most of these arise in vertebral posterior elements. It can be treated effectively by curettage. The acute pain is mediated by release of prostaglandins, so aspirin is an effective analgesic. An enchondroma is a benign tumor of hyaline cartilage that arises in the medullary space in young adults. Fibrous dysplasia is a localized area of developmental arrest of bone formation. A giant cell tumor is a benign but locally aggressive lesion that arises in the epiphysis of the long bones of young adults and has a “soap bubble” radiographic appearance. An osteochondroma is a projection of the cartilaginous growth plate to form an exostosis. PBD9 1197–1198 BP9 775–776 PBD8 1224–1225 BP8 812

20 E This osteosarcoma is a large destructive lytic and blastic mass arising in the metaphyseal region of the distal femur and extending into the surrounding soft tissue. This is the most common location. Trauma may call attention to this tumor, but is not a factor in pathogenesis. Chondrosarcomas arise over a wide age range, most often in the pelvis, shoulder, and ribs, and most patients are over 40 years of age. Ewing sarcomas arise in the diaphyseal region. Giant cell tumors also arise about the knee, but are large lytic, eccentric lesions with a thin rim of reactive, sclerotic bone. Such a large mass as that shown in the figure is unlikely to be a metastasis at any age; although testicular seminomas can occur in young men, they most often metastasize to regional lymph nodes. PBD9 1198–1199 BP9 776–777 PBD8 1225–1227 BP8 812–813

21 E The osteoid production by a sarcoma is diagnostic of osteosarcoma. The metaphyseal location in a long bone, particularly in the region of the knee, is consistent with osteosarcoma, as is the presentation in a young individual. Sporadic cases of osteosarcoma must have loss of both alleles of RB1, whereas in familial cases there is inheritance of one bad copy, and in those cases retinoblastomas are likely to appear first. Mutations of TP53 are found in many cancers, including osteosarcomas, and with the rare familial Li-Fraumeni syndrome. APC mutations are seen in association with colonic adenocarcinomas. BRCA1 mutations are associated with breast and ovarian cancers. NFI mutations lead to neurofibromatosis type 1, with many neural and soft tissue tumor types, but less likely primary bone tumors. PTEN mutations are most often associated with endometrial and prostate cancers. PBD9 1198–1199 BP9 776–777 PBD8 1225–1227 BP8 812–813
UNIT II  Diseases of Organ Systems

22 E The figure shows an osteochondroma with glistening cartilaginous cap overlying cancellous bone. This tumorlike lesion is benign and, when solitary, is essentially an incidental finding because a sarcoma rarely arises from an osteochondroma. Multiple osteochondromas can be part of an inherited syndrome, however, and onset can be in childhood, accompanied by bone deformity and an increased risk of development of a sarcoma. Both hereditary and sporadic osteochondromas have loss of heterozygosity in EXT1 or EXT2 genes. An osteochondroma is a projection of the cartilaginous growth plate with proliferation of mature bone capped by cartilage. When skeletal growth ceases, osteochondromas tend to cease proliferation as well. They may produce local irritation and pain. An enchondroma is a benign tumor of hyaline cartilage that arises in the medullary space of young adults. Fibrous dysplasia is a localized area of developmental arrest of bone formation. A giant cell tumor is a benign but locally aggressive lesion that arises in the epiphysis of the long bones of young adults and has a “soap bubble” radiographic appearance. An osteoblastoma is a large osteoid osteoma, which can arise in epiphyseal lesions and cause intense pain.

PBD9 1200  BP9 777–778  PBDB 1227  BP8 813–814

23 A This is an enchondroma. Cartilaginous lesions may be benign when they occur peripherally (hands and feet) and are localized, but a low-grade chondrosarcoma is more likely in a central location. The risk for malignancy is higher with multiple enchondromas (Ollier disease or Maffucci syndrome with mutations in isocitrate dehydrogenase genes). Giant cells are seen in many mass lesions of bone, but particularly in giant cell tumors and aneurysmal bone cysts occurring in larger bones. Macrophages may increase in lysosomal storage diseases such as Gaucher disease that involve bone marrow. Osteoblasts may be seen in an osteoid osteoma, which is more likely to occur at a younger age and produce pain at night. Plasma cells in myeloma are unlikely to occur at this age or at this site. Fibroblasts may be seen in localized fibrous lesions such as fibrous cortical defects that involve long bones.

PBD9 1201–1202  BP9 777  PBDB 1227–1228  BP8 814

24 C Giant cell tumors typically arise in the epiphyses of long bones of individuals 20 to 40 years old; there is a slight female predominance. The tumors may recur after curettage. Although most are histologically and biologically benign, with multinucleated cells in a stroma predominantly composed of spindle-shaped mononuclear cells as shown in the figure, in rare cases, a sarcoma can arise in a giant cell tumor of bone. Chondrosarcomas are typically larger destructive lesions. Enchondromas are most often peripheral skeletal lesions involving the metaphyseal region of small tubular bones of the hands and feet. An osteoblastoma usually involves the spine. Osteitis fibrosa cystica is a complication of hyperparathyroidism. A plasmacytoma composed of neoplastic plasma cells is most often one lesion of multiple myeloma, more likely to occur in older adults.

PBD9 1203–1204  BP9 208–209, 781  PBDB 1233  BP8 817

25 A The glistening, gray-blue appearance shown in the figure is typical of cartilage, and this lesion most likely represents neoplastic proliferation of chondrocytes. This chondrosarcoma has infiltrated the medullary cavity and invaded the overlying proliferation of chondrocytes. Most chondrosarcomas are low grade. They occur in a broad age range, in contrast to many other primary bone tumors that tend to occur in the first two decades of life. Most chondrosarcomas arise in the metaphyseal region of the skeleton. Osteoblast proliferation may be seen in a small tumor mass of the cortex known as ostoid osteoma, whereas a larger mass in the axial skeleton may be called an osteoblastoma. Osteosarcomas are derived from osteoblasts, but are malignant. They are usually seen in younger individuals and do not have a bluish-white appearance because they are marked by osteoid production. Giant cell tumors arise during the third to fifth decades; they involve epiphyses and metaphyses. Grossly, they are large, red-brown, cystic tumors. Giant cells resembling osteoclasts are present in giant cell tumors of bone. These tumors are believed to arise from cells of monocyte-macrophage lineage. Atypical plasma cells appear with multiple myeloma. Myelomas are dark red, rounded, lytic lesions that are often multiple. Primitive neuroectodermal cells are present in a Ewing sarcoma.

PBD9 1202  BP9 777–779  PBDB 1229–1230  BP8 814–815

26 B The histologic appearance is characteristic of a Ewing sarcoma. The radiologic appearance of the mass in this child is typical for a malignant tumor, with bone destruction and soft-tissue extension. The two most common malignant bone tumors in children are osteosarcoma and Ewing sarcoma. Osteosarcomas typically arise in the metaphyseal region, whereas Ewing sarcoma arises in the diaphyseal region of long tubular bones, as seen in this case. This tumor usually occurs in patients 10 to 15 years old. The t(11;22) translocation is present in about 85% of Ewing sarcomas and the related primitive neuroectodermal tumors (PNETs), which belong to the small round cell tumors of childhood that can be difficult to distinguish microscopically. The translocation gives rise to the EWS-FLI1 fusion gene, now considered the definitive test for the diagnosis of these tumors. Ewing sarcomas often produce tender masses with fever and leukocytosis, mimicking acute osteomyelitis. A chondrosarcoma can occur across a wide age range, in contrast to most primary malignancies arising in bone, which occur most often in the first two decades; most are sufficiently differentiated so that a cartilaginous matrix is apparent on microscopic examination. A giant cell tumor is a benign but locally aggressive lesion that arises in the epiphysis of the long bones of young adults and has a “soap bubble” radiographic appearance. Metastatic carcinoma is the most common tumor of adults involving bone because there are far more carcinomas than primary bone malignancies; childhood bone metastases are rare. An osteosarcoma typically arises in the metaphyseal region, and the malignant spindle cells produce an osteoid matrix. A plasmacytoma produces a focal lytic lesion within bone, and microscopically there are recognizable plasma cells.

PBD9 1203  BP9 780–781  PBDB 1232–1233  BP8 816–817
27 B This single focus (monostotic) fibrous dysplasia weakens the bone to the point of pathologic fracture. This benign tumorlike condition is uncommon. The histologic appearance of woven bone in the middle of benign-looking fibroblasts is characteristic. Seventy percent of cases are monostotic, and the ribs, femur, tibia, mandible, and calvaria are the most frequent sites of involvement. Local deformity and, occasionally, fracture can occur. Polyostotic fibrous dysplasia may involve craniofacial, pelvic, and shoulder girdle regions, leading to severe deformity and risk for fracture. Ewing sarcoma usually occurs in the diaphyseal region of the long bones and is identified histologically by sheets of small, round cells. A fracture callus should not be so localized within bone, and should not develop so quickly following trauma. An osteosarcoma is typically a large destructive lesion without central lucency. An osteoid osteoma has a small central nidus with surrounding sclerosis.

PBD9 1206  BP9 779–780  PBD8 1230–1231  BP8 816

28 B An elevated alkaline phosphatase level in an older adult should raise the suspicion of bone metastases, particularly when there is a “pathologic” fracture resulting from a bone lesion, rather than a fracture from trauma. Likely primary sites include the breast (in women), prostate (in men), lung (in smokers), kidney, and thyroid. Hyperparathyroidism can lead to osteitis fibrosa cystica with lytic lesions that are usually small, involve just cortex, and appear first in phalanges. Multiple myeloma can produce lytic bone lesions, but the patient’s serum gamma globulin level is not elevated. Osteochondromas are exostoses and do not produce lytic bone lesions. Paget disease of bone is characterized by osteolysis coupled with bone formation, but without lytic lesions. Fibrous dysplasia coupled with café-au-lait spots on skin and with endocrinopathies is known as McCune-Albright syndrome; this is a rare condition that occurs in young girls.

PBD9 1207  BP9 781  PBD8 1235  BP8 817–818

29 C The prevalence of cancer increases with age. In the absence of other findings, a metastatic carcinoma involving bone should be suspected. In addition, parathyroid hormone–related peptide elaborated by neoplasms can be a cause for hypercalcemia of malignancy. Chondrosarcoma can occur over a wide age range, but these focal lesions involving bone are not likely to elevate serum calcium. Metastases to bone are far more common than primary bone malignancies. Biliary tract obstruction or infiltrative diseases of the liver are usually the cause for an elevated alkaline phosphatase of liver origin. Gouty arthritis may be accompanied by some local bone destruction, but without a marked increase in serum calcium and the lesions are typically in peripheral joints. Osteoarthritis produces pain, usually involving weight-bearing joints or hands, but has no abnormal laboratory markers. Although there is accelerated bone loss with osteoporosis, alkaline phosphatase and serum calcium are normal. Paget disease of bone is associated with an increased alkaline phosphatase, but not hypercalcemia.

PBD9 330–331, 1207  BP9 781  PBD8 321, 1235  BP8 817–818

30 D Osteoarthritis is a common problem of aging, and various joints, from large, weight-bearing joints to small joints, can be involved. Joint stiffness in the morning is a common feature, but it is minimal and quickly subsides. Ankylosing spondylitis causes back pain with deformity, but does not usually affect distal extremities. Gouty arthritis occurs in patients with elevated serum levels of uric acid, whereas pseudogout arises from calcium pyrophosphate dihydrate crystal deposition; both produce acute arthritis with pain and swelling. Multiple myeloma can produce lytic lesions in bone, but does not typically involve joints. Rheumatoid arthritis can involve large or small joints. It is typically associated with symmetric involvement of small joints of the hands and feet.

PBD9 1208–1209  BP9 782–783  PBD8 1235–1237  BP8 818–820

31 A The progressive involvement of large, weight-bearing joints and osteophytes in the interphalangeal joints in an elderly woman are characteristic of osteoarthritis (OA). The absence of rheumatoid factor, and the asymmetric joint involvement, render the diagnosis of rheumatoid arthritis unlikely. OA is a multifactorial disease in which genetic predisposition and biomechanical forces affect chondrocytes. In early OA, chondrocytes proliferate. This is accomplished by changes in the cartilage matrix owing to secretion of proteases and inflammatory mediators by chondrocytes. Eventually, the ongoing inflammation and injury result in loss of the cartilage that causes reactive subchondral sclerosis. Inherited defects in type I collagen cause a group of rare disorders called osteogenesis imperfecta, which may be lethal in utero or in some cases lead to premature OA. Infiltration of the synovium with CD4+ cells is seen in rheumatoid arthritis. A partial deficiency of HGPRT gives rise to hyperuricemia and gout. Deficiency of HGPRT is implicated in some cases of gout. HLA-B27 is associated with ankylosing spondylitis and other seronegative spondyloarthropathies.

PBD9 1208–1209  BP9 782–783  PBD8 1235–1237  BP8 818–820

32 C A rheumatoid nodule is shown in the figure. Subcutaneous rheumatoid nodules such as this are typically found over extensor surfaces. Features of chronic rheumatoid arthritis (RA) include bilateral symmetric involvement of joints, destruction of joints with characteristic deformities, and presence of rheumatoid nodules. Although the pathogenesis of RA is complex, it is believed that the tissue injury is mediated by an autoimmune reaction in which CD4+ T cells secrete cytokines that have a cascade of effects on B cells, macrophages, and endothelial cells. B cells are driven to form rheumatoid factors, which form immune complexes in the joint; macrophages secrete cytokines such as tumor necrosis factor (TNF) and interleukin-1, which activate cartilage cells, fibroblasts, and synovial cells; and endothelial cell activation promotes accumulation of inflammatory cells in the synovium. Together, these processes form a pannus and eventually cause joint destruction. The central role of TNF in orchestrating joint destruction is the basis for the highly successful treatment of RA with anti-TNF therapy.
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UNIT II  Diseases of Organ Systems

Bisphosphonates diminish osteoclast activity to treat conditions with bone loss. RA is noninfectious, so antibiotics are not indicated. Cell cycle inhibitors as part of chemotherapy are reserved for malignancies; RA does not give rise to cancer. Although a gouty tophus can occur in soft tissue and resemble the lesion shown, the distribution of lesions does not fit with gout.

PBD9 1209–1212  BP9 784–786  PBD8 1237–1240
BP8 145–147, 820

33 E In addition to rheumatoid factor, serology for cyclic citrullinated peptides (CCPs) has specificity for rheumatoid arthritis and may indicate chronicity of the disease. The immunologically mediated damage leads to chronic inflammation with synovial proliferation, shown in the figure, with pannus formation that gradually erodes and destroys the joints, resulting in joint deformity. Typically, the pannus involvement is bilateral and symmetric, and small joints are often involved. Anti-topoisomerase I has specificity for scleroderma (systemic sclerosis). Antinuclear antibodies can be found in a variety of autoimmune diseases, including rheumatoid arthritis, but lack specificity. Lyme disease, caused by *Borrelia burgdorferi* infection, can produce a chronic arthritis that can destroy cartilage, but larger joints are usually involved. *Campylobacter* spp. may be related to cases of reactive arthritis.

PBD9 1210–1212  BP9 784–786  PBD8 1237–1240
BP8 145–147, 820

34 A Juvenile rheumatoid arthritis (RA), or juvenile idiopathic arthritis (JIA), in contrast to the adult type of rheumatoid arthritis, is more likely to be self-limited and non-deforming. JIA typically is rheumatoid factor negative, but ANA positive. JIA is more likely than the adult form to have systemic manifestations, such as rash, myalgia, myocarditis, pericarditis, uveitis, and glomerulonephritis. A positive serologic test for *Borrelia burgdorferi* is seen in Lyme disease, which tends to be associated with migratory arthritis of large joints. Similar to JIA, Lyme disease produces a chronic deforming arthritis in only about 10% of cases. *Chlamydia trachomatis* is typically the agent that produces the nongonococcal urethritis seen with reactive arthritis, which, similar to other spondyloarthopathies, most commonly involves the sacroiliac joint. Ferritin levels are markedly increased in hereditary hemochromatosis, in which iron deposition in joints can produce a chronic arthritis similar to osteoarthritis or pseudogout. Sickle cell disease with hemoglobin S can lead to aseptic necrosis, often of the femoral head, and to bone infarcts, with chronic arthritis secondary to bone deformity. Rheumatoid arthritis tends to be recurrent and causes progressive joint deformities, typically of hands and feet. Congenital syphilis can produce periostitis and osteochondritis with bone deformities; tertiary syphilis in adults can produce gummatus necrosis with joint destruction or loss of sensation, particularly in the lower extremities, leading to repeated trauma that deforms joints (Charcot joint). Some cases of gouty arthritis are accompanied by hyperuricemia; gouty arthritis tends to manifest as an acute attack in older individuals.

PBD9 1212  BP9 786  PBD8 1240  BP8 147

35 E Ankylosing spondyloarthritis has a very strong association with the HLA-B27 genotype that has a high prevalence among the native peoples of the circumpolar arctic and subarctic regions of Eurasia and North America. This progressive disease typically involves the lower back and pelvis. The radiographic feature of bamboo spine is characteristic. Spondyloarthopathies can have extra-articular manifestations such as anterior uveitis and ankylosing spondylitis. Glomerulonephritis is most likely a complication of collagen vascular diseases such as systemic lupus erythematosus. Autoimmune hepatitis is not related to joint diseases. Malabsorption is more likely to accompany diarrhea with enteropathic arthritis with infectious etiology. Meningitis may accompany Lyme disease.

PBD9 1213  BP9 786  PBD8 1241  BP8 147–148

36 D This patient developed enteritis-associated arthritis affecting the lumbar and sacroiliac joints several weeks after *Shigella* dysentery. He subsequently developed conjunctivitis and, most likely, uveitis. This symptom complex is a classic representation of a cluster of related disorders called seronegative spondyloarthopathies. This cluster includes ankylosing spondylitis, reactive arthritis, psoriatic arthritis, and enteropathic arthritis (as in this case). A common feature is a very strong association with the HLA-B27 genotype. Despite some similarities with rheumatoid arthritis, these patients invariably have a negative test result for rheumatoid factor. Urethritis caused by *Chlamydia trachomatis* can trigger reactive arthritis, another form of seronegative spondyloarthopathy. Such infection precedes the onset of arthritis, however. There is no relationship between infection with *Borrelia burgdorferi*, the causative agent of Lyme disease, and reactive arthritis in individuals testing positive for HLA-B27. Similarly, Epstein-Barr virus infection is not a trigger for these disorders.

PBD9 1213  BP9 786  PBD8 1241  BP8 147–148

37 C Psoriasis with psoriatic arthritis has features of rheumatoid arthritis, but without significant joint destruction. Psoriasis is common, affecting 1% to 2% of individuals, and about 5% of these have psoriatic arthritis. For the remaining choices, there is no significant association with arthritis. A bandlike dermal infiltrate is typical of lichen planus, which produces pruritic violaceous plaques or papules, but tends to abate in 1 to 2 years. Epidermal spongiosis with eosinophilic infiltrates can be seen in acute eczematous dermatitis as part of a drug reaction. Focal keratinocyte apoptosis is seen in graft-versus-host disease. IgG deposition can be seen in systemic lupus erythematosus and in bullous pemphigoid.

PBD9 1213  PBD8 1241

38 D Though *Staphylococcus aureus* infection is responsible for 80% to 90% of all cases of osteomyelitis in which an organism can be cultured. *Salmonella* osteomyelitis is especially common, however, in patients with sickle cell anemia. Group B streptococcal infections causing osteomyelitis are most common in neonates. *Klebsiella pneumoniae* osteomyelitis may rarely be seen in adults with urinary tract infections caused by this organism. Tuberculosis is a rare cause of osteomyelitis
Arthritis is quite rare. Necrosis that can involve large joints, and there can be lymphocytes or neutrophils. The inflammatory infiltrate contains a preponderance of neutrophils. Neisseria gonorrhoeae can cause an acute suppurative arthritis. In both conditions, Group B streptococcus may produce an acute osteomyelitis or arthritis in neonates. Tuberculous arthritis may involve large, weight-bearing joints, and it can be progressive, leading to ankylosis. The histologic features of tuberculous arthritis include caseating granulomas. Neisseria gonorrhoeae can cause an acute supplicative arthritis. In both conditions, the inflammatory infiltrate contains a preponderance of neutrophils. Treponema pallidum may produce gummatous necrosis that can involve large joints, and there can be lymphoplasmacytic infiltrates with endarteritis, but syphilitic arthritis is quite rare.

**40 A** Staphylococcus aureus is the most common cause of osteomyelitis in children and adults. Infections with *H. influenzae* may occur in children less than 2 years of age, but vaccination has reduced the incidence of such cases. Both organisms can cause congenital infections. Gonorrhea as a cause of acute osteomyelitis should be considered in sexually active adults, and may be passed to the fetus in the birth canal, but typically involves the eyes. Salmonella osteomyelitis is most characteristic of individuals with sickle cell anemia. Pneumococcal osteomyelitis is uncommon.

**41 C** Gonorrhea should be considered the most likely cause of an acute supplicative arthritis in sexually active individuals; in some cases multiple joints can be involved. In men, a urethritis may occur with gonorrheal infection. *Borrelia burgdorferi* causes Lyme disease, characterized by chronic arthritis that may mimic rheumatoid arthritis. *Haemophilus influenzae* is a short, gram-negative rod that can cause osteomyelitis in children. *Staphylococcus aureus* is the most common cause of osteomyelitis, but the Gram stain would show gram-positive cocci. *Treponema pallidum* infection, also a sexually transmitted disease, can lead to syphilitic gummata in the tertiary phase of syphilis that may produce joint deformity. There is no preceding urethritis, however. Tertiary syphilis may be preceded years earlier by a primary syphilitic chancre.

**42 C** Acute inflammation of the first MP joint, caused by precipitation of the needle-shaped negatively birefringent (yellow) uric acid crystals in the joint space, is typical of gout. Hyperuricemia is a sine qua non for the development of gout. Not all patients with hyperuricemia develop gout, and not all patients with gout have hyperuricemia, however. Other, ill-defined factors play a role in pathogenesis. Involvement of the big toe is classic, but other joints may be involved. Although attacks of gout are often precipitated by a heavy bout of alcohol consumption, liver damage (marked by elevated transaminases) is not a feature of gouty arthritis. Nonsteroidal anti-inflammatory drugs (NSAIDs) represent the first line of therapy for acute attacks. This is a chemically induced inflammatory reaction, not infectious, so antibiotics are not indicated. Glucocorticoids have a more pronounced effect upon chronic inflammatory conditions, and their continued use in joints will lead to degenerative arthritic changes. Methotrexate may be considered for treatment of rheumatoid arthritis. Xanthine oxidase inhibitors reduce uric acid formation and uricosuric agents help increase uric acid secretion, but they do not work acutely.

**43 A** The histologic picture is that of a central amorphous aggregate of urate crystals surrounded by reactive fibroblasts and mononuclear inflammatory cells. This is a gouty tophus. Tophi are large collections of monosodium urate crystals that can appear in joints or soft tissues of patients with gout. Large superficial tophi can erode the overlying skin. Precipitation of urate crystals into the joints produces an acute inflammatory reaction in which neutrophils and monocytes can be found. Neutrophils phagocytize urate crystals, which cannot be digested, but cause release of destructive neutrophilic lysosomal enzymes and oxygen free radicals. Release of crystals from the neutrophils perpetuates this cycle of inflammatory response. Inflammation of the joints involves different mechanisms depending on the etiology. In rheumatoid arthritis, release of tumor necrosis factor (TNF) by macrophages plays a central role, as evidenced by the dramatic relief provided by anti-TNF agents. Marked hypercholesterolemia, as occurs in familial hypercholesterolemia, can lead to deposition of cholesterol in tendons and elsewhere. When deposited in tendons, the yellowish lesions are called xanthomas; the cholesterol crystals appear microscopically as clefts in the tissue. Extra-pulmonary Mycobacterium tuberculosis infection can cause granulomatous inflammation and chronic arthritis and skin lesions, but there is caseous necrosis with epithelioid cells and no urate crystals. Tuberculous arthritis, in contrast to gouty arthritis, almost never begins in the MP joint. Reduced metabolic breakdown of homogentisic acid occurs in the inborn error of metabolism known as alkaptonuria, and deposition of homogentisic acid (ochronosis) in cartilage causes an arthritis that typically affects large joints, such as knees, intervertebral disks, hips, and shoulders, but small joints of the hands and feet are spared.

**44 D** Secondary gout can occur in patients with leukemia, especially patients with a high leukocyte count (as occurs in chronic myeloid leukemia), who are treated with chemotherapeutic agents causing massive lysis of leukemic cell nuclei,
and large amounts of urate are produced. The hyperuricemia leads to deposition of crystals in the joint space that triggers a local inflammatory response. Joint hemorrhages might be seen in patients with thrombocytopenia, but they more typically occur in patients with hemophilia, and neither induces needle-shaped crystal formation. Rhomboid cholesterol crystals are the most likely product of blood breakdown. Articular cartilage damage can be seen in any form of chronic arthritis, but is a prominent feature of osteoarthritis, not rheumatoid arthritis. Synovial proliferation also is a nonspecific change that is prominent in the inflammation of rheumatoid arthritis, not leukemia. Leukemic cells may infiltrate a variety of visceral organs, but usually not soft tissues.

**45 B** This patient shows evidence of hemochromatosis—caused skin pigmentation, heart failure, diabetes, and cirrhosis. The figure shows negatively birefringent (blue) rhomboidal calcium pyrophosphate crystals that have been deposited in the articular matrix. In progression of the disease, the crystals can seed the joint space and give rise to pseudogout, or calcium pyrophosphate dihydrate deposition (CPDD) disease, also called chondrocalcinosis. CPDD can be primary (hereditary) or, more commonly, secondary to various systemic diseases, such as hemochromatosis or, in the elderly, secondary to preexisting joint damage from other conditions. In most autoimmune diseases with a positive ANA result, such as systemic lupus erythematosus, there are arthralgias, but no arthritis, and little or no joint swelling, destruction, or deformity occurs. Rheumatoid arthritis tends to be recurrent and causes progressive joint deformities, typically of the hands and feet. Congenital syphilis can produce periostitis and osteochondritis with bone deformities; tertiary syphilis in adults, sometimes occurring after trauma. Rheumatoid nodules are firm, solid masses that typically occur in individuals who already have joint involvement with rheumatoid arthritis. A tophus is a solid mass of chalky sodium urate crystals in patients who have a history of gout.

**46 A** A ganglion cyst has a thin wall and clear, mucoid content. It arises in the connective tissue of a joint capsule or tendon sheath. The extensor surfaces of the hands and feet are the most common sites, particularly the wrist. Ganglion cysts probably arise after trauma from focal myxoid degeneration of connective tissue to produce a cystic space. They may regress. If not, and if they are painful, they can be excised. Tenosynovial giant cell tumor (villonodular synovitis) is a more diffuse form of giant cell tumor of tendon sheath (a solid mass lesion) and is a proliferation of mononuclear cells resembling synoviocytes. A lipoma is a mass of adipocytes and is not cystic. Nodular fasciitis is a solid reactive fibroblastic proliferation seen in the upper extremities and trunk of young adults, sometimes occurring after trauma. Rheumatoid nodules are firm, solid masses that typically occur in individuals who already have joint involvement with rheumatoid arthritis. A tophus is a solid mass of chalky sodium urate crystals in patients who have a history of gout.

**47 C** A tenosynovial giant cell tumor (formerly called pigmented villonodular synovitis) is the most common soft tissue tumor of the hand, but can occur around the knee as either a localized or diffuse proliferation. An acquired translocation placing the coding sequence for M-CSF adjacent to the promoter of the collagen COL6A3 gene is found in these lesions. Though histologically benign, lesions can be locally aggressive and extend into adjacent bone and soft tissue. The other listed options are not part of the pathogenesis of this lesion.

**48 E** Lipoma is the most common benign soft-tissue neoplasm. Such masses are extremely well differentiated and discrete. Multiple lipomas may be seen in some familial cases, but these are rare. These benign tumors do not metastasize; mesenchymal neoplasms do not often metastasize through lymphatics. Recurrence of some atypical lipomas or liposarcomas is possible, but benign lipomas do not recur. Secondary infection of this uncomplicated excision procedure is unlikely.

**49 A** The most common neoplasm is a lipoma, composed of mature adipocytes. They are slow growing and rarely produce problems. Endothelial cells are found in hemangiomas, which are common but firmer. In the groin region, they are likely to be on the skin surface. Fibroblasts may be found in reactive lesions such as fibromatoses, which are firm and immovable. Smooth muscle cells form firm leiomyomas. Neoplasms resembling skeletal muscle (rhabdomyomas) are quite rare.

**50 B** A large, deep soft-tissue mass suggests cancer, most likely a sarcoma. Liposarcomas are located in deep soft tissues, can be indolent, and can reach a large size. They are the most common sarcomas of adulthood. Chondrosarcomas can be seen over a wide age range, but they arise within bone. Carcinomas are far more common in adults than sarcomas, and can metastasize, but such a large lesion in soft tissue is unlikely to be a metastasis. Nodular fasciitis is a reactive fibroblastic lesion of young adults, usually on the upper extremities and trunk, and can develop several weeks after local trauma. Osteosarcomas generally occur in individuals younger than 20 years and typically arise in the metaphyseal region of long bones. Rhabdomyosarcoma occurs in children and is most often a tumor of the head and neck, genitourinary tract, or retroperitoneum.

**51 B** Nodular fasciitis is a reactive fibroblastic proliferation that is seen in the upper extremities and trunk of young adults, sometimes occurring after trauma. A lipoma is a common benign soft-tissue tumor that is not painful and does not follow trauma. A contusion is unlikely to lead to abscess formation because there is no disruption of the skin to allow entry of infectious agents. Pleomorphic fibroblastic sarcoma
is most likely to arise in the retroperitoneum and deep soft tissues of extremities in older adults. A superficial fibromatosis is a deforming lesion of fascial planes that develops over a long period, and the most common is a Dupuytren contracture involving the palm of the hand.

The patient has superficial fibromatosis that has produced a lesion best known as a Dupuytren contracture. These lesions contain mature fibroblasts surrounded by dense collagen. A hard, firm lesion of this size is unlikely to be malignant. Though some fibroblastic cells are present with a spindle shape, they are not numerous and generally not significantly atypical. Dystrophic calcification occurs in necrotic tissues; it is not commonly a localized mass. Granulation tissue from an injury would give rise to a stable scar without such severe retraction. Lipoblasts are seen in a liposarcoma, which is more likely to arise in deep soft tissues, such as thigh or retroperitoneum.

Rhabdomyosarcoma is the most common sarcoma in children. Sarcomas mark with antibody to vimentin, an intermediate cytoplasmic filament, with immunohistochemistry. Note the neoplastic spindle-shaped but markedly pleomorphic cells that have pink cytoplasm with a hint of striations mimicking skeletal muscle cells. CD3 is a T lymphocyte marker. Cytokeratin is a marker for tumors of epithelial origin (e.g., carcinomas). Dystrophin is a membrane-stabilizing protein in striated muscle; it is absent in Duchenne muscular dystrophy. Neuron-specific enolase is a marker of neoplasms with neural differentiation.

Synovial sarcomas account for 10% of all adult sarcomas and can be found around a joint or in deep soft tissues because they arise from mesenchymal cells, not synovium. Most synovial sarcomas show the t(X:18) translocation. A desmoid tumor is a fibromatosis composed of fibroblasts and collagen. Leiomyosarcomas do not have a biphasic pattern microscopically and are rarely seen in soft tissues. A mesothelioma can be biphasic, but it more typically arises in the pleura, or less commonly the mesothelial surface of peritoneum or pericardium. An osteoblastoma is a bone mass that arises in the epiphyseal region.
1. A 62-year-old woman has a slowly enlarging mass anterior to the right ear. Surgery is performed to remove a pleomorphic adenoma of the parotid gland. The tumor has infiltrated the overlying soft tissue, and the surgeon must remove a portion of the facial nerve to obtain an adequate margin. He places a 2-cm nerve graft in the excised area. Which of the following best describes the most likely outcome during the first week after surgery?
   - A. Acute inflammatory cells around the graft
   - B. Formation of a traumatic neuroma
   - C. Recurrent tumor along the nerve graft
   - D. Segmental demyelination and axonal loss of the nerve proximal to the graft
   - E. Fragmentation of distal axons and myelin sheaths

2. A 16-year-old boy has a deep laceration of the left lower thigh. The bleeding is stopped. On physical examination, he has loss of sensation in the lateral left foot and movement in the left foot. The wound is surgically repaired, including nerve, and he receives physical therapy. How long will it take him to regain the use of his left foot?
   - A. 1 day
   - B. 1 week
   - C. 1 month
   - D. 6 months

3. A 60-year-old man has pain in the left neck, shoulder, and forearm that has been worsening for a month. Over the next month there is increasing weakness of the left pectoralis major, latissimus dorsi, and triceps muscles. Paresthesias of the tips of the thumb, forefinger, and ring finger are present. There is no warmth, tenderness, or swelling. A lesion involving which of the following structures is most likely to be present in this man?
   - A. Vasculature
   - B. Axon
   - C. Ganglion
   - D. Myofiber
   - E. Schwann cell

4. A 41-year-old man had an influenza-like illness for 1 week, followed 4 days later by rapidly progressive, ascending motor weakness requiring mechanical ventilation. On physical examination, he is now afebrile and has 3/5 motor strength in his extremities. A lumbar puncture is done and yields clear, colorless cerebrospinal fluid under normal pressure. This fluid has a slightly elevated protein concentration, but a normal glucose level, and a cell count with only a few mononuclear cells. He recovers in 3 weeks. If lymphocytic infiltrates were seen in peripheral nerves along with segmental demyelination at the time he initially saw his physician, what would be the most likely diagnosis?
   - A. Amyotrophic lateral sclerosis
   - B. Guillain-Barré syndrome
   - C. Multiple sclerosis
   - D. Varicella-zoster virus infection
   - E. Vitamin B₁₂ (cobalamin) deficiency

5. A 37-year-old HIV-positive man has had a relapsing and remitting course of motor and sensory problems for the past year, including difficulty with ambulation as well as symmetric numbness and tingling in all extremities. Nerve conduction studies show findings consistent with demyelination and remyelination. He is treated with plasmapheresis. Which of the following disorders is most likely to cause this man’s neurologic disease?
   - A. Bacterial infection
   - B. Carcinoma
   - C. Hyperglycemia
   - D. Immune dysregulation
   - E. Traumatic injury
6 A 93-year-old woman has been bothered by continuing outbreaks of painful lesions on the skin of her right chest for the past year. On physical examination, there is a vesicular eruption in a 1 × 8 cm area over the right seventh rib. She is treated with acyclovir, and partial resolution of the skin lesions occurs, but the pain persists for the next 3 months. Which of the following is the most likely cause for her findings?

A. Aging
B. Diabetes mellitus
C. Multiple sclerosis
D. Somatoform pain disorder
E. Varicella-zoster virus infection
F. Vitamin B12 (cobalamin) deficiency

7 A 66-year-old man receiving hemodialysis for chronic renal failure has noted increasing loss of sensation in his legs for the past 4 years. On physical examination, there is symmetrically decreased sensation over both lower extremities. He has no decrease in strength or abnormality of gait. Which of the following is most likely to produce these findings?

A. Cerebral astrocytoma
B. Diabetes mellitus
C. Hansen disease
D. Multi-infarct dementia
E. Multiple sclerosis
F. Uremia

8 A 55-year-old man has had a foot ulcer for 2 months that has not healed. Physical examination shows a 2-cm shallow, nonhealing ulceration of the left medial malleolus. There is symmetric decreased sensation in the distal regions of the lower extremities. He has a history of multiple urinary tract infections resulting from difficulty in completely emptying the bladder. He is impotent. Which of the following pathologic findings is most likely to be present in the peripheral nerves?

A. Acute inflammation
B. Axonal neuropathy
C. Onion bulb formation
D. Segmental demyelination
E. Wallerian degeneration

9 A 24-year-old woman has had episodes of numbness and tingling in both hands for 5 months. The problem is worse near the end of the day and makes it difficult for her to use the computer keyboard. The thumb and first two fingers are most affected. There is no pain or swelling, and she does not recall any trauma to the upper extremities. On physical examination, she has a positive Tinel sign and decreased sensation to light touch and pinprick over the palmar surface of both hands in the distribution of the first three digits. Thenar muscle atrophy is present. This neuropathy is most likely due to which of the following underlying causes?

A. Acute intermittent porphyria
B. Diabetes mellitus, type 2
C. Entrapment with compression
D. Systemic lupus erythematosus
E. Varicella-zoster virus infection

10 A 41-year-old woman has noted marked pain in the right foot for the past 2 months. The pain makes it difficult for her to wear high-heeled shoes and seems to be worse at the end of the day. On physical examination, she has severe pain on palpation of the interdigital space between the second and third toes. There is no swelling or erythema of the foot. Motor strength in the lower extremities seems to be normal. What has most likely produced these findings?

A. Beriberi
B. Type 2 diabetes mellitus
C. Entrapment neuropathy
D. Lead poisoning
E. Wallerian degeneration

11 A 58-year-old man has experienced worsening double vision and eyelid drooping, particularly toward the end of the day, for 1 month. He also has had difficulty chewing his food at dinner. He was diagnosed with Sjögren syndrome more than a decade ago. On physical examination, he has 5/5 motor strength in his extremities that decreases to 4/5 strength with repetitive movement. Administration of edrophonium restores muscle strength. There is no pain on palpation and no decrease in joint mobility. Which of the following laboratory findings is most likely to be reported for this patient?

A. Acetylcholine receptor antibody positivity
B. Anti–histidyl tRNA synthetase (anti–Jo-1) titer 1:512
C. Elevated serum creatine kinase level
D. Increased serum cortisol level
E. Peripheral blood eosinophilia

12 A 72-year-old man has had a 7-kg weight loss, proximal muscle weakness, and difficulty with urination for the past 4 months. On physical examination, he has 4/5 muscle strength in his proximal extremities that does not diminish with repetitive motion. He has no muscle pain or loss of mobility. Laboratory studies show that he does not have serum antibodies to acetylcholine receptor. He was prescribed anticholinesterase agents but shows no improvement. Which of the following underlying conditions is most likely to be present in this man?

A. Chronic hepatitis C
B. Diabetes mellitus
C. Duchenne muscular dystrophy
D. Lead poisoning
E. Small cell lung carcinoma

13 A previously healthy 40-year-old man and his 42-year-old wife have had increasing blurred vision and weakness for the past day. On examination they are afebrile. Orthostatic hypotension is present. They have 3/5 muscle weakness in all extremities and difficulty breathing. They are treated with intubation and mechanical ventilation. Which of the following is the most likely cause for their paralysis?

A. Botulism
B. Cocaine ingestion
C. Diabetes mellitus
D. Guillain-Barré syndrome
E. Thymoma
14. A 42-year-old man has had increasing progressive muscle weakness in both arms and legs along with dysarthria and difficulty in swallowing for the past 2 years. He is now wheelchair-bound. Physical examination shows 3/5 motor strength in all extremities. He has no muscle pain on palpation, no deformities or loss of joint mobility, and no tremor. A biopsy specimen of the quadriceps muscle is obtained, and microscopic examination shows no inflammation, but only atrophy of the myofibers. What is the most likely diagnosis?

A. Amyotrophic lateral sclerosis
B. Becker muscular dystrophy
C. Mitochondrial myopathy
D. Myasthenia gravis
E. Werdnig-Hoffmann disease

15. A 30-year-old woman has had gradually increasing muscle weakness with myalgia for the past year. She now has difficulty getting up from a chair and climbing stairs. She does not have weakness of her hand muscles. Physical examination reveals a fine violaceous rash on her face, predominantly palpebral. Dusky, flat, red patches are present on her elbows, knees, and knuckles. Laboratory studies show serum creatine kinase of 620 U/L. A deltoid biopsy specimen is obtained, and on microscopic examination shows a mononuclear inflammatory cell infiltrate around small blood vessels and groups of atrophic myofibers at the periphery of fascicles. What mechanism is most likely responsible for her disease?

A. Antibody- and complement-mediated injury to the microvasculature
B. Expansion of CTG repeat sequences on chromosome 19q13.2
C. Mutation in a gene encoding for voltage-gated calcium channels
D. Myofiber injury by CD8+ cells directed against muscle antigens
E. T cell–mediated nerve injury induced by *Mycoplasma pneumoniae* infection

16. A 56-year-old woman has had increasing generalized muscle weakness for the past 2 months. On physical examination, she has 4/5 motor strength in upper and lower extremities. She has fat redistribution in the upper trunk and rounded facies. Ecchymoses are scattered over the extremities. She is afebrile, and her blood pressure is 155/90 mm Hg. A biopsy specimen of the gastrocnemius muscle is obtained, and histochemical staining with ATPase shows type II muscle fiber atrophy. What is the most likely diagnosis?

A. Cushing syndrome
B. McArdle disease
C. Duchenne muscular dystrophy
D. Myasthenia gravis
E. Polymyositis

17. A 71-year-old woman is receiving a drug to lower her serum cholesterol. Over the past week she has developed muscle pain and weakness unrelated to physical activity. On examination she has diffuse but mild muscle tenderness. Laboratory studies show her serum creatine kinase is 2049 U/L and creatinine is 2 mg/dL. Urine dipstick analysis is positive for blood, without RBCs on urine microscopy. Which of the following drugs is most likely to produce her findings?

A. Cholestyramine
B. Clofibrate
C. Ezetimibe
D. Lovastatin
E. Nicotinic acid

18. A 44-year-old man has had worsening exercise intolerance for the past year. On physical examination, he has 4/5 motor strength in the extremities, but has no muscle pain or loss of joint mobility. He has pitting edema to the knees. A chest radiograph shows cardiomegaly with pulmonary edema and pleural effusions. A deltoid muscle biopsy specimen is obtained. The figure shows the immunohistochemical staining pattern with antibody to dystrophin (A, normal; B, patient). What is the most likely diagnosis?

A. Amyotrophic lateral sclerosis
B. Becker muscular dystrophy
C. Myasthenia gravis
D. Polymyositis
E. Werdnig-Hoffmann disease
19 A 5-year-old boy develops increasing muscle weakness. He is unable to play with other children because he quickly becomes tired and is unable to keep up with them. On physical examination, he is afebrile. No deformities are noted. He has 4/5 muscle strength in his extremities, with more apparent weakness of the proximal muscles. Laboratory studies show a serum creatine kinase level of 689 U/L. A muscle biopsy is done and the microscopic appearance shown in the figure. Which of the following tests would be most appropriate to confirm the diagnosis in this boy?

A Absolute eosinophil count in peripheral blood  
B Immunohistochemical staining for dystrophin  
C PCR to detect expansion of CGG repeats on chromosome Xq27.3  
D Presence of oligoclonal bands of immunoglobulin in cerebrospinal fluid  
E Serum acetylcholinesterase antibody titer

20 A 16-year-old boy has had two episodes of sudden loss of motor function with residual weakness in his right arm and right leg in the past 2 years. He has had muscle weakness and a seizure disorder since childhood. During the past year, he has had difficulty with memory and performing activities of daily living. On physical examination, he has short stature. He has 4/5 motor strength in all extremities, with no muscle tenderness. Laboratory studies show glucose, 71 mg/dL; creatinine, 1.1 mg/dL; and lactic acid, 9.2 mmol/L. A gastrocnemius muscle biopsy specimen is obtained, and microscopic examination shows ragged red fibers. On electron microscopy, the myofibrils have "parking lot" inclusions. The boy’s mother and grandmother had similar findings, but his father and grandfather did not. Which of the following most likely explains the pathogenesis of his disease?

A Abnormal voltage-gated calcium channel  
B Antibodies to acetylcholine receptor  
C Cytotoxic CD8+ lymphocytes against myofibers  
D Decreased sarcolemmal dystrophin  
E Deficient mitochondrial ATP generation  
F Increased CTG repeat sequences at 19q13.2-13.3

21 A 10-year-old girl has exhibited muscular weakness since early childhood that has not worsened. She can ambulate unassisted, but does not participate in strenuous physical activities. On examination she has 4/5 motor strength in proximal muscles and 5/5 in distal muscles. There is no muscle pain on palpation. A biopsy of deltoid muscle is obtained, and with Gomori trichrome stain, microscopic analysis shows subsarcolemmal aggregates of rod-shaped intracytoplasmic inclusions. Laboratory studies show a normal serum creatine kinase. Which of the following is the most likely form of muscle disease she has?

A Channelopathy  
B Congenital myopathy  
C Glycogen storage disease  
D Inflammatory myopathy  
E Mitochondrial myopathy

22 An infant born at term exhibits difficulty with movement beginning at 1 month of age. By 1 year of age, there is flaccid paralysis. A muscle biopsy is done, and microscopically shows panfascicular atrophy of myofibers with scattered enlarged myofibers, but no inflammation. The serum creatine kinase is not elevated. What is the most likely diagnosis?

A Amyotrophic lateral sclerosis  
B Becker muscular dystrophy  
C McArdle disease  
D Myasthenia gravis  
E Myotonic dystrophy  
F Spinal muscular atrophy

23 A 40-year-old man undergoes elective laparoscopic hernia repair. He receives anesthesia with halothane and succinylcholine. His blood loss is minimal. Thirty minutes into this surgery, his temperature increases to 39.5° C, and pulse increases to 115/min. The anesthesiologist notices muscular spasms with rigidity of the extremities. Laboratory studies show an elevated serum creatine kinase and myoglobinuria. This man most likely has a mutation in a gene encoding for a protein that regulates the function of which of the following?

A Calcium ion channel  
B Motor end plate  
C Oxidative phosphorylation enzyme  
D Sarcoglycan complex  
E Thick filament

24 A 42-year-old woman has had an increasing number of subcutaneous nodules developing over the past 20 years. On physical examination the nodules range from 0.5 to 1.5 cm in size and are firm and nontender. She also has multiple café-au-lait spots on her skin. She now has pain in her posterior left thigh. MR imaging shows an 8-cm mass. A mutation encoding for which of the following proteins is most likely to be present in this woman?

A Actin  
B Cardamom  
C Caveolin  
D Dystrophin  
E Neurofibromin  
F Troponin
The axons distal to the point of injury or transection degenerate. The Schwann cells that remain can guide the regrowing nerves. Macrophages help to remove myelin-derived debris from the area of nerve injury, but acute inflammation is not a typical feature of diseases involving peripheral nerves. Traumatic neuromas may occur after transection, but careful dissection prevents this, and the purpose of the graft is to guide orderly regrowth. A tumor is unlikely to follow a nerve, although a feature of a malignant tumor is a tendency to invade nerves. Segmental demyelination is more typical for diabetic neuropathy.

Laceration of the tibial nerve results in Wallerian degeneration distal to the injury. Realignment of the nerve is accompanied by axonal sprouting. The new axons find the residual myelin sheaths and grow down at the rate of about 2 mm/day, taking 1 year to traverse the length of the calf. There can be reinnervation of the muscle, but there is type grouping of the muscle fascicles that are reinnervated. Physical therapy can aid in the interim. Complete restoration of motor and sensory functions may not occur.

The distribution is that of C7, likely from a herniated disc compressing the nerve root. The digital branch of the median nerve is purely C7. Nerve entrapment or compression typically involves the axon. If the nerve is not severed, then axonal regrowth can occur. Neurapraxia occurs with transient injury (e.g., when your foot “falls asleep”) because there is conduction loss but no axonal damage. Axonotmesis involves axonal loss and Wallerian degeneration, but without damage to the myelin sheath, so axonal regrowth can occur. Neurotmesis involves transection of the nerve, and regrowth can only occur with exact alignment of the severed nerve so axonal sprouts can find the proper myelin sheath. Vasculitis could involve an artery or supplying a nerve, but is unlikely to involve a single nerve root. The motor and sensory loss here is not consistent with injury to a ganglion. The sensory component here means more than myofibers are involved, and the myofibers supplied by C7 have neurogenic atrophy. The paresthesias, not anesthesia, suggest the nerve was not severed, and Schwann cells remained.

Guillain-Barré syndrome is an uncommon disorder that most often follows a bacterial (Campylobacter jejuni), viral (cytomegalovirus), or mycoplasmal infection and is thought to be caused by generation of myelin-reactive T cells, or by molecular mimicry, somehow triggered by the infection. The paralysis of respiratory muscles may be life-threatening, although many patients recover after weeks of ventilatory support. Amyotrophic lateral sclerosis is associated with slowly progressive muscle weakness. Various presentations are possible in multiple sclerosis, but the plaques of demyelination are generally not large or diffuse enough to cause paralysis of the respiratory muscles. Mechanical ventilation may be necessary eventually. Varicella-zoster virus infection most often involves the skin in a dermatomal distribution from a spinal nerve root. Vitamin B12 deficiency results in subacute progressive degeneration of the spinal cord plus sensorimotor disturbances in the extremities.

Chronic inflammatory demyelinating polyneuropathy (CIDP) can be seen in patients with immunologic diseases. Treatment may aid recovery, but a chronic course may ensue. Bacterial infections produce signs of acute inflammation with redness and swelling, and tend not to involve nerves specifically. Some paraneoplastic syndromes may occur with carcinomas, such as Lambert-Eaton myasthenic syndrome, with weakness. Carcinomas are less likely at his age, and are not frequent complications of HIV infection. Hyperglycemia is characteristic of diabetes mellitus, and diabetic neuropathy is likely to be progressive and unremitting. Traumatic injury is unlikely to produce such widespread findings and unlikely to be followed by a variable course.

Failing infection with chickenpox, the varicella-zoster virus (VZV) becomes dormant in dorsal root ganglia, only to reactivate later when the immune system no longer contains it. Such containment failures are more likely with age and with immunocompromised states, and this produces the classic appearances of shingles in a dermatomal distribution. She also has postherpetic neuralgia, a disabling condition of chronic pain that is difficult to manage. Aging alone does not explain disease, but older individuals are more likely to have disease conditions. Diabetes mellitus leads to neuropathies that can have loss of sensation or even pain, but there are no associated skin lesions. Multiple sclerosis can involve the spinal cord white matter with variable neurologic findings, but there are no skin lesions, and the onset of MS is at a younger age. Pain out of proportion to the pathologic findings suggests a somatoform pain disorder, but she does have significant pathology findings that explain her continued pain. Cobalamin deficiency with pernicious anemia and subacute combined degeneration of the spinal cord can lead to paresthesias and loss of function, but not to skin lesions.

The most common cause of a predominantly sensory peripheral neuropathy is diabetes mellitus. Long-standing diabetes mellitus also gives rise to nephropathy with chronic renal failure. Sensorimotor disturbances are typically not seen with intracranial mass lesions such as astrocytomas. Cerebral infarctions could lead to decreased motor activity and/or sensory loss, although not usually in a symmetric pattern; small infarcts culminating in dementia may not produce significant motor or sensory loss. Hansen disease (leprosy) produces focal anesthesia. The demyelinating
lesions of multiple sclerosis in different white matter sites and at different times can produce many signs and symptoms, but symmetric lesions should suggest another disease process. Uremic neuropathy resembles diabetic neuropathy, but it typically regresses with dialysis.

PBD9 1232–1233  BP9 799  PBD8 1265–1266  BP8 784–785, 898

8 D The features described are consistent with a peripheral neuropathy associated with diabetes mellitus. Both motor and sensory nerves can be involved, and there may be an autonomic neuropathy. Histologic examination shows an axonal neuropathy with segmental demyelination. Difficulty in emptying the urinary bladder and impotence are results of autonomic neuropathy. Longer nerves are affected first; this explains the lower leg involvement and accounts for many cases of diabetic foot, with trauma and subsequent ulceration. Acute inflammation is not generally seen in neuropathies. Lymphocytic infiltrates may be seen in Guillain-Barré syndrome. Onion bulb formation is a feature of the hereditary neuropathy known as Refsum disease. Wallerian degeneration typically occurs with traumatic transection of a nerve.

PBD9 1232–1233  BP9 799  PBD8 1265–1266  BP8 784–785, 898

9 C Carpal tunnel syndrome is a form of compression neuropathy that results from entrapment of the median nerve beneath the flexor retinaculum at the wrist. Women are more commonly affected than men, and the problem is often bilateral. The role of excessive repetitive use of the wrist in causation has been debated, but lifestyle modifications and ergonomics can be employed as initial conservative therapy. Conditions such as hypothyroidism, amyloidosis, and edema with pregnancy also diminish the space in the carpal tunnel. Acute intermittent porphyria can lead to a hereditary form of pattern of muscular atrophy. This occurs because motor neurons can be affected, resulting in a denervation process. Werdnig-Hoffman disease also is a neuropathic disease with grouped atrophy, but onset is in infancy. The autoantibodies are directed against calcium channels at motor nerve terminal membranes. Patients with chronic viral hepatitis may have generalized malaise and weakness that is not related to specific muscle disease. Diabetes mellitus also may produce peripheral neuropathy, but more often involving distal, not proximal, regions first. Duchenne muscular dystrophy is an X-linked disease that manifests early in childhood. Lead poisoning leads to peripheral neuropathy, often with central nervous system findings.

PBD9 1234  BP9 797–798  PBD8 1266–1267

10 C Fashion may have a price. The patient has a Morton neuroma, a form of compressive neuropathy in which a plantar nerve is trapped between metatarsal heads. Chronic injury leads to growth of a tangled mass of axons, fibroblasts, and perineural cells. Diabetic neuropathy, which occurs in either type 1 or type 2 disease, is bilateral. It is characterized by loss of sensation, which may be a predisposing factor for foot trauma. Toxic disorders (lead poisoning) and metabolic disorders (beriberi with thiamine deficiency) are not as focal. Wallerian degeneration is a dying-back neuropathy associated with severing of a nerve; a neuroma may form at the site of injury.

PBD9 1234  BP9 808  PBD8 1267

11 A Amyotrophic lateral sclerosis is also known as Lou Gehrig disease, named for the famous New York Yankee first baseman affected with the disorder. Lower and upper motor neurons can be affected, resulting in a denervation type of pattern of muscular atrophy. This occurs because an individual neuron innervates a group of muscle fibers. Bulbar (cranial nerve) involvement denotes a more rapid course. Werdnig-Hoffman disease also is a neuropathic disease with grouped atrophy, but onset is in infancy. The other listed options are not associated with denervation.

PBD9 1300–1301  BP9 801–802  PBD8 1324–1325  BP8 827, 896

12 E Lambert-Eaton myasthenic syndrome is a rare form of paraneoplastic syndrome. Proximal muscles tend to be involved first. Similar to many paraneoplastic syndromes, it is most often associated with small cell carcinoma of the lung. The autoantibodies are directed against calcium channels at motor nerve terminal membranes. Patients with chronic viral hepatitis may have generalized malaise and weakness.
ion channel genes give rise to various channelopathies, including hypokalemic periodic paralysis and malignant hyperthermia. The CD8+ T cells are believed to be important in the pathogenesis of polymyositis. T cell–mediated myelin injury is seen with Guillain-Barré syndrome, causing an acute ascending paralysis.

16 A Type II muscle fiber atrophy can occur with glucocorticoid excess and also after prolonged immobilization. Routine light microscopy may not distinguish type II atrophy from denervation atrophy. Histochemical staining for ATPase could be done (but a biopsy is not needed, given this history). There is a deficiency of myophosphorylase enzyme in McArdle disease, leading to muscle pain and cramping with vigorous exercise. Duchenne muscular dystrophy is an X-linked condition and is rare in females. Onset is in early childhood. Antibodies to the acetylcholine receptor cause the muscular weakness in myasthenia gravis. Polymyositis is an inflammatory condition affecting all fiber types.

17 D She has a statin-induced myopathy, with a creatine kinase level more than 10 times normal from rhabdomyolysis (myoglobin released from muscle can be detected by the urine dipstick). Statins are HMG-CoA reductase inhibitors that reduce endogenous cholesterol synthesis in liver. Cholestyramine binds bile acids in the intestine and disrupts enterohemorrhagic bile acid circulation to increase conversion of cholesterol to bile acids in the liver. Clofibrate enhances uptake and oxidation of free fatty acids in muscle. Ezetimibe interferes with intestinal lipid absorption. Nicotinic acid inhibits mobilization of peripheral free fatty acids to reduce hepatic triglyceride synthesis and secretion of VLDL.

18 B The biopsy specimen shows a reduced amount of dystrophin, but not a complete absence; this suggests Becker muscular dystrophy. In Duchenne muscular dystrophy, dystrophin is absent because of gene deletion. In keeping with the diagnosis of Becker muscular dystrophy, the patient is older and not severely affected. Both dystrophies are X-linked conditions. Amyotrophic lateral sclerosis is a denervation atrophy seen in adults, with loss of anterior horn cells in the spinal cord and cranial nerve nuclei. Myasthenia gravis results from antibodies to acetylcholine receptors, and there is minimal structural change to the muscle. Polymyositis is an autoimmune disease that results from a T cell–mediated attack on muscle fibers, causing muscle fiber degeneration with inflammation. Werdnig-Hoffmann disease is a form of spinal muscular atrophy. Onset is at birth, and it results from a genetically determined loss of anterior horn cells.

19 B The onset of muscle weakness in childhood suggests an inherited muscular dystrophy. The biopsy specimen shows variation in muscle fiber size and increased connective tissue between the fibers. This morphologic finding in a boy strongly suggests X-linked muscular dystrophy. Immunohistochemical staining for dystrophin would show an absence of dystrophin, confirming the diagnosis of Duchenne muscular dystrophy. Eosinophilia may be present in allergic or parasitic disorders, including trichinosis. Expansion of CGG repeats on Xq27.3 is diagnostic of familial mental retardation. Oligoclonal immunoglobulin bands in the cerebrospinal fluid are a feature of multiple sclerosis. Antibodies to the acetylcholine receptor are found in myasthenia gravis, which is characterized by weakness in muscles after repetitive use.

20 E ATP generation by oxidative phosphorylation in mitochondria can be affected by mitochondrial genes, which are separate from those on chromosomes in the cell nucleus. These abnormal genes can lead to mitochondrial myopathies, encephalopathies, and deafness. In this case, there is mitochondrial encephalopathy with lactic acidosis and strokelike episodes (MELAS). Mitochondrial genes have a maternal pattern of inheritance. An abnormal voltage-gated calcium channel is seen in one of the channelopathies; it causes hypokalemic periodic paralysis. Channelopathies are typically inherited in an autosomal dominant fashion. Antibodies to acetylcholine receptor cause myasthenia gravis; its sole manifestation is muscle weakness. Cytoxic CD8+ cells mediate the muscle injury in polymyositis. Decreased sarcoplasmic dystrophin is present in Becker muscular dystrophy; dystrophin is absent in Duchenne muscular dystrophy. The dystrophin gene is located on the X chromosome. Increased CTG repeats occur in myotonic dystrophy.

21 B Nemaline rod myopathy, one form of congenital myopathy, may present in infancy or childhood and may be nonprogressive or slowly progressive. Congenital myopathies are often named for their characteristic histologic features. Channelopathies typically present as periodic paralysis with abnormalities in serum potassium. The two glycogen storage diseases most often affecting striated muscle include Pompe disease (type II) and McArdle disease (type V). There are increased glycogen deposits seen with periodic acid-Schiff (PAS) stain. Neuropathies affect muscle through denervation, leading to groups of atrophic muscle fibers. Polymyositis and dermatomyositis are inflammatory myopathies that are accompanied by myalgia and fiber degeneration with increased serum creatine kinase. Mitochondrial myopathies may appear in childhood but are usually progressive, and often other organs such as heart or brain are involved.

22 F Werdnig-Hoffman disease is a form of spinal muscular atrophy resulting from loss of motor neurons in infancy, so the biopsy specimen shows grouped atrophy of myofibers. Death is inevitable by age 3 years. Amyotrophic lateral sclerosis is a progressive disease with a neurogenic form of muscle atrophy resulting from loss of motor neurons. Becker muscular dystrophy has an X-linked pattern of inheritance.
and onset in adulthood, unlike Duchenne muscular dystrophy, but in both there is a mutation in the dystrophin gene. McArdle disease is an autosomal recessive condition resulting from a deficiency in muscle phosphorylase and does not produce progressive weakness. Myasthenia gravis results from acetylcholine receptor antibody and leads to progressive weakness. Myotonic dystrophy is characterized by facial and upper body weakness, cataracts, gonadal atrophy, cardiomyopathy, and dementia.

Malignant hyperthermia is a rare but life-threatening disorder seen in only 1 in 20,000 adults, but with millions of surgical procedures under anesthesia being performed, it must be considered when the patient’s temperature increases rapidly. In patients with mutations in the ryanodine receptor, there is impaired reuptake of calcium into the sarcoplasmic reticulum with increased intracellular calcium leading to hypermetabolism. Motor end plates include acetylcholine and acetylcholine receptors that are involved with myasthenia gravis. The mitochondrial DNA genes are mainly involved in oxidative phosphorylation, and there is a maternal inheritance pattern, with possible muscular weakness, but not hyperthermia. The sarcoglycan complex includes many proteins such as dystrophin that can be involved with muscular dystrophies, but not hyperthermia. Thick filaments include the myosin contractile protein; β-myosin mutations are found in some cases of hypertrophic cardiomyopathy.

She has neurofibromatosis type 1 (NF-1) with multiple subcutaneous neurofibromas, and she has developed a larger malignant peripheral nerve sheath tumor (MPNST). Such MPNSTs may arise from a preexisting neurofibroma. Neurofibromin functions as a tumor suppressor. Actin is one of the contractile proteins of muscle and its absence is not compatible with life. Cardamom is an aromatic spice from the ginger family of plants, used in traditional medicine for oral and intestinal conditions. Caveolin is an intracellular vesicle transport protein, and mutations involving this protein may be present in some forms of limb-girdle muscular dystrophy. Dystrophin stabilizes sarcolemmal membranes, and its absence leads to Duchenne muscular dystrophy. Troponin is part of the myofiber unit and is used to detect striated muscle injury, specifically myocardial ischemia and infarction.
1. In a study of hypoglycemic shock, cellular changes in the brain are analyzed. One cell type in the hippocampus is noted to exhibit intense cytoplasmic eosinophilia, central chromatolysis, spheroidal swellings, and nuclear pyknosis. These changes appear 12 hours after blood glucose levels drop below 20 mg/dL. What is this cell type most likely to be?
   - A. Astrocytes
   - B. Endothelial cells
   - C. Microglia
   - D. Neurons
   - E. Oligodendroglia

2. A 75-year-old man has a history of transient attacks of loss of vision. The only abnormalities on physical examination are bruits over the carotids in the neck. Two days later he suddenly becomes hemiplegic and loses consciousness. He is rushed to the emergency room and a CT scan shows evidence of cerebral infarction. He is put on life support but dies 5 days later. At autopsy there is an area of necrosis and microscopically these lesions are noted to have increased numbers of cells distributed around the central zone of necrosis. Which of the following cell types is most likely to have a phagocytic function in these lesions?
   - A. Arachnoidal cells
   - B. Astrocytes
   - C. Ependymal cells
   - D. Microglia
   - E. Oligodendroglia

3. A 49-year-old woman has had a severe headache for 2 days. On physical examination, she is afebrile and normotensive. Funduscopic examination shows papilledema on the right. One day later, she has right pupillary dilation and impaired ocular movement. She then becomes obtunded. Which of the following lesions best explains these findings?
   - A. Chronic subdural hematoma
   - B. Frontal lobe abscess
   - C. Glioblastoma with edema
   - D. Hydrocephalus ex vacuo
   - E. Occipital lobe infarction
   - F. Ruptured middle cerebral berry aneurysm

4. A 16-year-old boy with no prior medical problems has complained of headaches for the past 9 months. There are no abnormal findings on physical examination. CT scan of the head shows enlargement of the lateral cerebral ventricles and third ventricle. A lumbar puncture is performed with normal opening pressure, and clear CSF is obtained, which has a slightly elevated protein, normal glucose, and no leukocytes. Which of the following intracranial lesions is most likely to cause these findings?
   - A. Aqueductal stenosis
   - B. Cerebral abscess
   - C. Cryptococcal meningitis
   - D. Ependymoma
   - E. Multiple sclerosis
   - F. Vascular malformation

5. A 61-year-old man has had worsening mental function with confusion for the past year, along with headaches. At first the headaches occurred in the morning, but for the past 3 months they have become continuous, along with nausea and blurred vision. On physical examination there is bilateral papilledema. A head CT scan shows enlargement of the entire ventricular system. Which of the following prior illnesses most likely led to his current problems?
   - A. Aqueductal stenosis
   - B. Choroid plexus papilloma
   - C. Cerebral infarction
   - D. Dandy-Walker malformation
   - E. HIV infection
   - F. Pneumococcal meningitis
6. A 67-year-old woman has had new onset headaches with nausea for the past month. She now has a worsening headache with weakness in her right leg. On physical examination she has 4/5 motor strength involving her right leg. Extraocular muscle movements are intact. In which of the following locations is a neoplasm most likely to be found in this woman?

A. Left inferior frontal lobe  
B. Left lateral temporal lobe  
C. Left superior occipital lobe  
D. Right inferior occipital lobe  
E. Right medial temporal lobe  
F. Right superior frontal lobe

7. An 81-year-old man with a history of poorly controlled atrial fibrillation suddenly collapses while watching television at home. Emergency medical services arrive promptly, but multiple attempts at cardioversion over 15 minutes are required to reestablish a stable pulse and blood pressure. Over the next day he develops bilateral papilledema, and an MRI of his brain shows an indistinct cortical gray-white junction and narrowing of ventricles. Which of the following intracranial abnormalities most likely developed in this man?

A. Acute inflammation  
B. Cytotoxic edema  
C. Metastatic adenocarcinoma  
D. Obstruction of CSF flow  
E. Subarachnoid hemorrhage

8. A 45-year-old man develops a severe headache and fever over 2 days. On physical examination, he has nuchal rigidity and bilateral papilledema. His temperature is 38.5°C. A blood culture shows gram-positive cocci in chains, and *Streptococcus pneumoniae* is identified. The figure shows the representative gross appearance of a section of his brain. Based on this appearance, which of the following complications most likely resulted from this patient’s infection?

A. Abscess formation  
B. Herniation  
C. Hydrocephalus  
D. Laminar cortical necrosis  
E. Subarachnoid hemorrhage

9. A 30-year-old woman, G3, P2, is in the third trimester of pregnancy. She has noted minimal fetal movement throughout the pregnancy. A fetal ultrasound scan shows normal amniotic fluid volume, normally implanted placenta, and the abnormality shown in the figure. Which of the following laboratory findings is most likely to be present in this woman?

A. 45, XX, t(14;21)(p11;q11) karyotype  
B. Elevated serum α-fetoprotein level  
C. High cytomegalovirus IgM titer  
D. Hyperbilirubinemia with anemia  
E. Increased hemoglobin A1c level

10. A 22-year-old primigravida had a fetal screening ultrasound study at 18 weeks showing a single large cerebral ventricle and fused thalami. On physical examination at birth at 36 weeks’ gestation, the infant is small for gestational age and has multiple anomalies, including postaxial polydactyly of hands and feet, cyclopia, microcephaly, cleft lip and palate, and rocker-bottom feet. The infant dies 1 hour after birth. Which of the following CNS abnormalities best explains these findings?

A. Anencephaly  
B. Arnold-Chiari II malformation  
C. Dandy-Walker malformation  
D. Holoprosencephaly  
E. Periventricular leukomalacia

11. A 24-year-old man incurs head and neck trauma in a motor vehicle accident. He now has impaired pain and temperature sensation from the shoulders down to his feet, but proprioception and vibratory sense is preserved. He has motor weakness with muscle atrophy starting in his hands and extending to forearms and shoulders. An MRI of the cervical spinal cord shows a transverse slit-like cavity extending from the level of C2 to C7. What is the most likely diagnosis?

A. Dandy-Walker malformation  
B. Diffuse axonal injury  
C. Rachischisis  
D. Spina bifida  
E. Syringomyelia
12. A 15-year-old girl has had progressive difficulty speaking during the past 6 months. She becomes dizzy and falls frequently. She complains of headache and facial and neck pain. During the past month, she has had decreasing bladder and bowel control. On physical examination, there is loss of pain and temperature sensation over the nape of the neck, shoulders, and upper arms, but vibration and position sensation are preserved. She has muscle wasting in the lower neck and shoulders. MRI of the spinal cord shows cervical and thoracic enlargement with a CSF collection dilating the central canal. MRI of the brain shows gross findings similar to those shown in the figure. Which of the following is the most likely diagnosis?

A. Arnold-Chiari II malformation
B. Cerebral palsy
C. Corpus callosal agenesis
D. Holoprosencephaly
E. Polymicrogyria

13. A neonate is born prematurely at 28 weeks’ gestation to a 22-year-old primigravida. The infant is initially stable, and a newborn physical examination shows no abnormalities. The infant becomes severely hypoxemic 24 hours later, and seizure activity is observed. There is poor neurologic development during infancy. CT scan of the head shows symmetrical enlarged cerebral ventricles at 8 months of age. Which of the following perinatal complications most likely produced these findings?

A. Congenital cytomegalovirus infection
B. Cerebral palsy
C. Corpus callosal agenesis
D. Holoprosencephaly
E. Polymicrogyria

14. A 21-year-old woman incurs a blow to her head from a fall while mountain biking. She then has loss of consciousness for 5 minutes. On examination her deep tendon reflexes are diminished. A head CT scan 6 hours later shows no abnormalities. She recovers over the next week, with no neurologic deficits, but cannot remember this event. During the next year she has irritability, headache, difficulty sleeping, trouble concentrating, and fatigue. Which of the following is the most likely consequence from her injury?

A. Arteriolosclerosis
B. Concussion
C. Hydrocephalus
D. Leukoencephalopathy
E. Post-traumatic dementia
F. Myelinolysis

15. An 83-year-old woman slips in the bathtub in her home and falls backward, striking her head. She is taken to the emergency department, where examination shows a 3-cm reddish, slightly swollen area over the occiput. She is arousable but somnolent. There are no motor or sensory deficits. There is no papilledema. CT scan of the head is performed. Acute hemorrhage in which of the following locations is most likely to be seen?

A. Basal ganglia
B. Basis pontis
C. Cerebral ventricle
D. Epidural space
E. Inferior frontal lobe
F. Sella turcica

16. A 19-year-old snowboarder wearing protective equipment consisting of a baseball cap, baggy shorts, and a flak jacket flew off a jump and hit a tree. He was initially unconscious, and then “came to” and wanted to try another run, but his friends thought it best to call for help. On the way to the emergency department, he became comatose. Physical examination now shows left papilledema. Skull radiographs show a linear fracture of the left temporoparietal region. This clinical picture is most consistent with which of the following lesions?

A. Acute leptomenigitis
B. Contusion of frontal lobes
C. Middle meningeal artery laceration
D. Ruptured berry aneurysm
E. Tearing of cerebral bridging veins

17. A 72-year-old woman trips and falls down the stairs. She does not lose consciousness. She develops a headache and confusion 30 hours later and is taken to the emergency department. On physical examination, she is conscious and has a scalp contusion on the occiput. What is the most likely location of an intracranial hemorrhage in this patient?

A. Basal ganglia
B. Epidural
C. Pontine
D. Subarachnoid
E. Subdural
18. An 80-year-old resident of a nursing home is admitted to the hospital because of recent onset of fluctuating levels of consciousness with headache and confusion for the past 2 days. On physical examination, she is arousable, but disoriented and irritable. Vital signs include temperature of 36.9°C and blood pressure of 130/85 mm Hg. There is papilledema on the right. CT scan of the head shows a collection of blood in the subdural space on the right. Which of the following vascular lesions most likely produced these findings?

- A. Bleeding from an arteriovenous malformation
- B. Laceration of the middle meningeal artery
- C. Rupture of a saccular aneurysm
- D. Tearing of the cerebral bridging veins
- E. Thrombosis of the middle cerebral artery

19. A 22-year-old man is caught in a rip current off Cabo San Lucas. He becomes tired and overcome by the waves. Lifeguards get him to shore, but he has no pulse. Resuscitative measures over the next 20 minutes establish a pulse. However, he does not regain consciousness. A month later an electroencephalogram (EEG) shows no brain wave activity. Which of the following cells most likely predominated in the cerebral cortex at the time of this EEG?

- A. Fibroblasts
- B. Lymphocytes
- C. Macrophages
- D. Neutrophils
- E. Oligodendrocytes
- F. Red neurons

20. A 68-year-old woman with atrial fibrillation suddenly lost consciousness and fell to the ground. When she became arousable, she was unable to move her left arm and had difficulty speaking. On physical examination, her temperature was 37°C, pulse was 81/min, respirations were 18/min, and blood pressure was 135/85 mm Hg. The figure shows the representative gross appearance of her brain in radiologic orientation. An MRI 3 months later shows a cystic space. Which of the following conditions most likely resulted from which of the following conditions?

- A. Arteriovenous malformation
- B. Embolic arterial occlusion
- C. Metastatic carcinoma
- D. Organizing subdural hematoma
- E. Superficial cortical contusion

21. A 79-year-old man with metabolic syndrome has had 6 episodes of sudden dysarthria, a feeling of weakness in his hand, and dizziness in the past 3 months. These episodes usually last less than 1 hour, and then he feels fine. Today, he suddenly lost consciousness while walking to the bathroom in his house and fell to the floor. On regaining consciousness 4 minutes later, he was unable to move his right arm. Which of the following underlying lesions is most likely to be found in his brain?

- A. Arteriovenous malformation
- B. Cerebral atherosclerosis
- C. Frontal lobe astrocytoma
- D. Meningoencephalitis
- E. Subdural hematoma

22. A study is conducted to identify causes of neuronal loss in patients 18 to 90 years old who died in the hospital from a natural manner of death and who had autopsies performed. Subsequent microscopic examination of sections revealed red, shrunken neurons, decreased numbers of neurons, or absent neurons. The hippocampal pyramidal cells, the cerebellar Purkinje cells, and the superior parasagittal neocortical pyramidal cells are affected. What condition is most likely to be the major cause of neuronal loss in these patients?

- A. Autoimmunity
- B. Chemotherapy
- C. Diabetes mellitus
- D. Global hypoxia
- E. Lead ingestion
- F. Poor nutrition

23. A 70-year-old woman had an episode 2 days earlier during which she lost consciousness for several minutes. On physical examination, there is 4/5 motor strength in the right upper extremity and decreased sensation to pinprick on the right arm and hand. There are bilateral carotid bruits. CT scan of the head shows no intracranial hemorrhage, but there is a slight midline shift; MRI of the brain shows edema near the left internal capsule. A lumbar puncture is performed with normal opening pressure. Laboratory studies on 10 mL of clear, colorless CSF show two mononuclear WBCs/mm³, no RBCs, protein concentration of 40 mg/dL, and glucose concentration of 70 mg/dL. The serum glucose concentration is 95 mg/dL. Which of the following laboratory findings is most suggestive of the risk factor for her disease?

- A. Antiphospholipid antibody
- B. Blood culture positive for Streptococcus pneumoniae
- C. Elevated serum concentration of very long chain fatty acids
- D. Hyperammonemia
- E. Hypercholesterolemia
- F. Positive serologic test for syphilis

24. A 59-year-old woman had sudden loss of consciousness 4 months ago. On physical examination, she now has left hemiplegia. CT imaging shows a large, cystic space in the right parietal region. MR angiography shows occlusion of a peripheral cerebral artery branch at the gray-white junction near the lesion. What underlying disease process is this woman most likely to have?

- A. AIDS with a low CD4+ T-lymphocyte count
- B. Chronic alcoholism with micronodular cirrhosis
- C. Chronic renal failure with hypertension
- D. Colonic adenocarcinoma with Trousseau syndrome
- E. Ischemic heart disease with left ventricular thrombosis
25 A 39-year-old man presents with headache and altered mental status of 60 hours’ duration. On examination he is afebrile and normotensive and has a reduced level of consciousness with aphasia. Lumbar puncture is performed and the CSF obtained has a lymphocytic pleocytosis with modest protein elevation. An MRI of the brain shows focal hyperintense cortical lesions. A stereotaxic brain biopsy shows chronic inflammation with granulomas involving arterioles and venules. Which of the following therapeutic options is most appropriate for this man?

A Antibiotics
B Anti-hypertensives
C Anti-pyretics agents
D Immunosuppressives
E Mannitol infusion

26 A 55-year-old man suddenly loses consciousness while driving his truck, but he is traveling at a slow speed and comes to a stop without a collision. Paramedics arrive but are unable to arouse him. On physical examination, there is bilateral papilledema. He has no spontaneous movements. The figure shows the gross appearance of the brain at autopsy. What underlying condition is most likely to have resulted in this lesion?

A Chronic alcoholism
B Metastatic carcinoma
C Multiple sclerosis
D Systemic hypertension
E Thromboembolism

28 A 72-year-old man with diabetes mellitus has had stepwise cognitive decline for the past 5 years. On multiple occasions he has had an acute event, such as loss of consciousness or confusion, followed by worsening ability to perform activities of daily living. On physical examination, he has mild right hemiparesis, ataxia, and dysarthria. Which of the following pathologic findings is most likely to be present and numerous in this man?

A Lacunes
B Lewy bodies
C Neuritic plaques
D Neurofibrillary tangles
E Pick bodies
F Plaques of demyelination

29 An 86-year-old man has become progressively unable to live independently for the past 10 years, and he now requires assistance with bathing, dressing, toileting, feeding, and transfers in and out of chairs and bed. On physical examination, he has no motor or sensory deficits. He cannot give the current date or state where he is. Six months later, he suddenly becomes comatose and dies. At autopsy, there is a large superficial left parietal lobe hemorrhage. Histologic examination of the brain shows numerous neocortical neuritic plaques and neurofibrillary tangles. The peripheral cerebral arteries and the core of each plaque stain positively with Congo red. Which of the following mechanisms is most likely responsible for his disease?

A Aggregation of Aβ peptide
B Conformational change in the prion protein (PrP)
C Dopamine deficiency
D Expansion of polyglutamine repeats
E Mutations in the tau gene

30 A 50-year-old woman develops a sudden, severe headache and is taken to the emergency department. On examination, she has nuchal rigidity. Her blood pressure is 115/83 mm Hg. A lumbar puncture is done; the CSF shows numerous RBCs, no neutrophils, a few mononuclear cells, and a normal glucose level. The Gram stain result is negative. CT imaging shows subarachnoid hemorrhage at the base of the brain. Which of the following vascular events has most likely occurred in this woman?

A Bleeding from cerebral amyloid angiopathy
B Hematoma formation from arteriolosclerosis
C Middle cerebral artery thromboembolism
D Rupture of an intracranial berry aneurysm
E Tear of subdural bridging veins

31 A 45-year-old, previously healthy man has developed headaches over the past month. There are no remarkable findings on physical examination. A cerebral MR angiogram shows a 7-mm saccular aneurysm at the trifurcation of the right middle cerebral artery. Which of the following is the most likely complication from this lesion?

A Cerebellar tonsillar herniation
B Hydrocephalus
C Epidural hematoma
D Subarachnoid hemorrhage
E Subdural hematoma
A 25-year-old man has complained of headaches for the past 5 months. During that time, family members noticed that he was not as mentally sharp as he has been in the past, and that he has become more emotionally labile. Over a 2-week period, he has 4 generalized seizures. On physical examination, he now has no papilledema or movement disorder. CT scan of the head shows a 2-cm mass in the right frontal lobe. A stereotactic biopsy specimen of this lesion shows only gliosis and evidence of recent and remote hemorrhage. The mass is removed, and histologic examination shows a conglomerate of various-sized tortuous vessels surrounded by gliosis. What is the most likely diagnosis?

A  Angiosarcoma  
B  Arteriovenous malformation  
C  Multiple sclerosis plaque  
D  Organizing abscess  
E  Prior head trauma  
F  Ruptured saccular aneurysm

A 19-year-old man has a sore throat followed a day later by sudden onset of a severe headache. Physical examination shows mild pharyngitis and nuchal rigidity. His skin shows petechial hemorrhages. His temperature is 38.8°C, pulse is 98/min, respirations are 26/min, and blood pressure is 95/45 mm Hg. The figure shows the representative gross appearance of the surface of his brain. Which of the following infectious organisms is most likely to have produced his disease?

A  Cryptococcus neoformans  
B  Mycobacterium tuberculosis  
C  Neisseria meningitidis  
D  Poliovirus  
E  Toxoplasma gondii

A 44-year-old woman who is an intravenous drug user is admitted to the hospital with increasing headache and high fever for the past 24 hours. On physical examination, her temperature is 38.4°C, pulse is 85/min, respirations are 18/min, and blood pressure is 125/85 mm Hg. CT scan of the head shows no mass lesion or midline shift. A lumbar puncture is performed. The CSF shows 70,000 neutrophils/mm³, an increased protein concentration, and a decreased glucose level. Which of the following infectious agents is most likely to produce these findings?

A  Herpes simplex virus  
B  JC polyomavirus  
C  Mycobacterium tuberculosis  
D  Staphylococcus aureus  
E  Toxoplasma gondii

A 77-year-old man has been irritable for the past 2 days. He is otherwise healthy. On physical examination, he has a temperature of 39.1°C. Laboratory examination of the CSF from a lumbar puncture shows numerous neutrophils, slightly increased protein level, and decreased glucose concentration. On Gram staining of the CSF, which of the following is most likely to be seen microscopically?

A  Gram-negative bacilli  
B  Gram-negative diplococci  
C  Gram-positive cocci  
D  Gram-positive short rods  
E  No organisms

A 43-year-old woman has had a headache and fever for the past 2 weeks following a severe respiratory tract infection accompanying bronchiectasis. On physical examination, her temperature is 38.8°C. There is no papilledema. She has no loss of sensation or motor function, but there is decreased vision in the left half of her visual fields. CT scan of the head shows a sharply demarcated, 3-cm, ring-enhancing lesion in the right occipital region. A lumbar puncture is done, and laboratory analysis of the CSF shows numerous leukocytes, increased protein, and normal glucose levels. What is the most likely diagnosis?

A  Cerebral abscess  
B  Glioblastoma  
C  Metastatic carcinoma  
D  Multiple sclerosis  
E  Subacute infarction

An 11-year-old boy has had pain in his right ear for 1 week and a severe headache for 1 day. On physical examination his temperature is 37.5°C. He has marked tenderness on palpation posterior to the right ear. Pus exudes from the right tympanic membrane, and Streptococcus pneumoniae is cultured. Which of the following intracranial complications is he most likely to develop if untreated?

A  Encephalitis  
B  Epidural abscess  
C  Multicystic encephalopathy  
D  Subdural hematoma  
E  Trigeminal neuralgia
A 4-year-old girl residing near Cape Town, South Africa, has had worsening headache and irritability for the past week, and now exhibits nausea, vomiting, and diminished responsiveness to verbal commands. On examination she has a temperature of 37.2°C. A tremor is observed in her extremities. Her eyes do not move laterally. A lumbar puncture is performed and examination of the CSF shows 100 leukocytes/mm³, and 75% of them mononuclears. The CSF glucose is decreased, but there are no neutrophils. A Gram stain and India ink preparation of the CSF are negative. Her condition gradually improves over the next 6 months. Serum serologic tests are done, and the CSF shows only a few mononuclear cells and normal glucose and protein levels. Infection with which of the following organisms is the most likely cause of her disease?

A  Aspergillus niger  
B  Cytomegalovirus  
C  Herpes simplex virus  
D  Eastern equine encephalitis virus  
E  Neisseria meningitidis

A 26-year-old woman has headaches for 4 weeks along with increasing malaise. Physical examination yields no remarkable findings. CT scan of the head shows no abnormalities. A lumbar puncture yields clear, colorless CSF with a normal opening pressure. Laboratory analysis of the CSF shows a normal glucose concentration and a minimally increased protein level. A few lymphocytes are present, but there are no neutrophils. A Gram stain and India ink preparation of the CSF are negative. Her condition gradually improves over the next 6 months. Serum serologic tests are most likely to show an elevated titer of antibodies to which of the following infectious agents?

A  Listeria monocytogenes  
B  Rabies virus  
C  Rubeola (measles) virus  
D  Streptococcus pneumoniae  
E  Toxoplasma gondii  
F  Treponema pallidum

A 12-year-old boy develops fever, accompanied by occasional headaches, malaise, fatigue, and nausea a month after being bitten by a dog. One day later, he experiences episodes of rigidity, hallucinations, breath holding, and difficulty swallowing because of uncontrollable oral secretions. Dr. Louis Pasteur is consulted. He writes: “The death of this child appearing to be inevitable, I decided, not without lively and sore anxiety, as may well be believed, to try … the method which I had found constantly successful with dogs. Consequently, 60 hours after the bites [the child] was inoculated under a fold of skin with half a syringeful of the spinal cord of a rabbit. In the following days, fresh inoculations were made. I thus made 13 inoculations.” The boy survives. Which of the following histologic findings in the brain of the dog is most likely to be present?

A  Spongiform change  
B  Pseudocysts with bradyzoites  
C  Multinucleate giant cells  
D  Perivascular lymphocytes  
E  Neocortex
45 A 37-year-old man who is HIV-1-positive has had increasing memory problems for the past year. He is depressed. During the past 3 months, he has had increasing problems with motor function and is now unable to stand or walk. For the past 3 days, he has had fever, cough, and dyspnea. A bronchoalveolar lavage shows cysts of *Pneumocystis jiroveci*. MRI of the brain shows diffuse cerebral atrophy; no focal lesions are identified. On microscopic examination of his brain, which of the following findings is most likely to be present?

A. Cerebellar spongiform changes  
B. Cortical microglial nodules  
C. Lentiform nuclear lacunar infarcts  
D. Neocortical neuritic plaques  
E. White matter plaques of demyelination

46 A 52-year-old woman with leukemia undergoes chemotherapy. Two months later, she develops neurologic deficits with ataxia, motor weakness in the right arm, difficulty swallowing, and sensory changes in the left leg. MRI of the brain shows irregular areas of increased attenuation in white matter of the cerebral hemispheres and the cerebellum. A stereotaxic biopsy specimen shows perivascular chronic inflammation, marked gliosis, large reactive astrocytes with bizarre nuclei, and intranuclear inclusions within oligodendroglia. What virus most likely caused these findings?

A. Cytomegalovirus  
B. Herpes simplex virus  
C. JC polyomavirus  
D. Rabies virus  
E. Rubeola virus  
F. West Nile virus

47 A 38-year-old man with chronic renal failure received a kidney transplant. While being treated with cyclosporine, azathioprine, and high doses of corticosteroids, he began to experience headaches and became lethargic. On physical examination, he now has a fever and nuchal rigidity. A lumbar puncture is performed and the opening pressure is increased. A CSF cell count shows increased leukocytes. An India ink preparation shows the findings in the figure. Which of the following organisms is most likely infecting this man?

A. Aspergillus fumigatus  
B. Cryptococcus neoformans  
C. Plasmodium falciparum  
D. Taenia solium  
E. Toxoplasma gondii  
F. Trypanosoma gambiense

48 A previously healthy 21-year-old man with a severe headache for 5 days now has a new-onset seizure. Papilledema is noted on fundoscopic examination. An MRI of the brain shows multiple 0.5- to 1.5-cm cystic periventricular and meningeal lesions. Which of the following infectious organisms is most likely to produce these findings?

A. *Aspergillus* fumigatus  
B. *Cryptococcus* neoformans  
C. *Plasmodium* falciparum  
D. *Taenia* solium  
E. *Toxoplasma* gondii  
F. *Trypanosoma* gambiense

49 A 20-year-old HIV-positive man has had a decreased level of consciousness for the past week. He now experiences a generalized tonic-clonic seizure. On physical examination, his temperature is 37.6°C. MRI of the brain shows several 1- to 3-cm, ring-enhancing lesions in the cerebral gray matter bilaterally. A stereotaxic biopsy is performed. What pathologic finding is most likely to be present on microscopic examination of the biopsy specimen?

A. Budding cells with pseudohyphae  
B. Large atypical lymphocytes  
C. Metastatic squamous cell carcinoma  
D. Spongiform encephalopathy  
E. *Toxoplasma* pseudocysts

50 A 63-year-old previously healthy woman has become more forgetful over a period of 6 weeks. One month later, she has difficulty ambulating and is unable to care for herself. On physical examination, she has myoclonus. She is afebrile. CT scan of the head shows minimal cerebral atrophy. An EEG shows low-amplitude, slow background activity with periodic complexes and occasional repetitive sharp waves with intervals of 0.5 to 1 second. Which of the following histologic abnormalities is most likely to be found in her cerebral cortex?

A. Lewy bodies  
B. Microglial nodules  
C. Numerous neuritic plaques  
D. Plaques of demyelination  
E. Spongiform encephalopathy

51 A 27-year-old woman had an episode of weakness 3 months ago, which she attributed to job stress and fatigue. The neurologic examination shows mild residual weakness, with 4/5 motor strength in the right lower extremity. A lumbar puncture is done, and laboratory examination of the CSF shows increased IgG levels with prominent oligoclonal bands. MRI of the brain shows small, scattered, 0.5-cm areas consistent with demyelination, most of which are located in periventricular white matter. Which of the following complications is she most likely to develop?

A. Non-Hodgkin lymphoma  
B. Progressive dementia  
C. Quadriplegia  
D. Seizure disorder  
E. Tremor at rest  
F. Visual impairment
A 28-year-old man states that 3 years ago he experienced paresthesias of his left arm and had difficulty walking, but these problems resolved. During the past year, he developed difficulty seeing from his left eye. Six months ago, he had difficulty writing with his right hand. On physical examination, there is decreased visual acuity on the left, no papilledema, and no retinal lesions. There is decreased motor strength and decreased sensation in the right hand and forearm. MRI of the brain shows focal areas of increased signal intensity in periventricular white matter and in the left optic nerve. A lumbar puncture is performed. What finding is most likely to be present on examination of the CSF?

A Antitreponemal antibodies
B Cryptococcal antigen
C Malignant cells
D Oligoclonal bands
E Xanthochromia

A 74-year-old woman sustains blunt head trauma in a motor vehicle accident. On admission to the hospital, she is conscious but disoriented. CT scan of the head shows a right temporal bone fracture and mild cerebral edema. Her blood alcohol level is 0.24 gm%. Two days later, laboratory studies show serum Na⁺, 109 mmol/L; K⁺, 3.9 mmol/L; Cl⁻, 82 mmol/L; CO₂, 23 mmol/L; glucose, 73 mg/dL; and creatinine, 1 mg/dL. The hyponatremia is corrected over the next 2 hours with intravenous fluid and electrolyte therapy and diuretics. She then rapidly becomes confused and exhibits limb weakness. No papilledema is seen on funduscopic examination. An MRI shows the finding in the figure. What complication has most likely occurred in this woman?

A Central pontine myelinolysis
B Cerebellar tonsillar herniation
C Intraventricular hemorrhage
D Subacute combined degeneration of the cord
E Wernicke-Korsakoff syndrome

An 8-year-old boy recovered uneventfully from a viral upper respiratory infection 2 weeks ago, but now has the abrupt onset of lethargy and irritability. On neurologic examination he has diminished pupillary reflexes bilaterally along with ataxia of his extremities. CSF obtained by lumbar puncture microscopically shows small numbers of lymphocytes and erythrocytes. MRI shows multiple hyperintense lesions at the gray-white junction. Which of the following pathogenic mechanisms is most likely causing this child’s brain lesions?

A Abscess formation
B Demyelination
C Embolization
D Metastases
E Thiamine deficiency
F Vasculitis

A study is conducted of patients who had increased phosphorylated tau and decreased Aβ peptide in their CSF 5 to 10 years prior to death at ages ranging from 55 to 80 years. At autopsy their brain weights are less than normal for age and body size. On gross examination, these brains show hydrocephalus ex vacuo and cortical atrophy but no focal lesions. The figure shows the high power microscopic appearance of cerebral neocortex with Bielschowsky silver stain. Which of the following symptoms is most likely to be recorded in the medical histories of these patients?

A Choreiform movements
B Gait disturbances
C Grand mal seizures
D Progressive memory loss
E Symmetric muscular weakness
A 68-year-old woman with a 7-year history of progressive dementia dies of bronchopneumonia. At autopsy, there is cerebral atrophy in a predominantly frontal and parietal lobe distribution. Microscopic examination of the brain shows numerous neuritic plaques in the hippocampus, amygdala, and neocortex. Neurofibrillary tangles in the hippocampus contain tau protein. Congo red staining shows amyloid in the media of the small peripheral cerebral arteries. Which of the following genetic abnormalities is the most important factor in the development of her disease?

A  Expansion of CAG repeats on chromosome 4p16
B  HLA-DR3/DR4 alleles
C  Increased tandem repeats in the FMRI gene
D  Mutation of a prion protein gene
E  Presence of the e4 allele at the ApoE gene

A 63-year-old man had increasing irritability over 3 years. He wandered about his neighborhood, complaining to the neighbors about everything. He had no memory loss and was always able to find his way home. The neighbors were pleased when he developed aphasia. On physical examination, there were no motor or sensory deficits and no gait disturbances or tremor. MRI of the brain showed bilateral marked temporal and frontal lobe gyral atrophy. He died of pneumonia 1 year later. At autopsy, the frontal cortex microscopically shows extensive neuronal loss, and some remaining neurons show intracytoplasmic, faintly eosinophilic, rounded inclusions that stain immunohistochemically for tau protein.

What is the most likely diagnosis?

A  Alzheimer disease
B  Huntington disease
C  Leigh disease
D  Multiple system atrophy
E  Parkinson disease
F  Pick disease
G  Vascular dementia

A 60-year-old woman had problems related to movement for 5 years. Physical examination showed cogwheel rigidity of limbs and a festinating gait, which she had difficulty initiating. Her face was expressionless. She was given levodopa/carbidopa, and her condition improved. Two years later, she had difficulty performing activities of daily living and showed marked cognitive decline. She died of aspiration pneumonia. Autopsy findings include mild cerebral atrophy and loss of substantia nigra pigmentation. Microscopically, cortical neurons show spheroidal, intraneuronal, cytoplasmic, and eosinophilic inclusions. Immunohistochemical staining for which of the following proteins is most likely to be positive in these inclusions?

A  α-Synuclein
B  Amyloid precursor protein
C  Apolipoprotein E
D  Huntingtin
E  Presenilin
F  Tau protein

A 55-year-old man has had increasing difficulty with initiation of voluntary movements and increasing inability to perform activities of daily living for 1 year. On physical examination, he has difficulty initiating movement, but he can keep moving if he follows someone walking ahead of him. He has an expressionless facies. The left side of the figure shows the gross appearance of the midbrain of this patient; on the right is a section through normal midbrain.

What additional clinical feature is most closely associated with this abnormality?

A  Ataxia with ambulation
B  Choreiform movements
C  Loss of short-term memory
D  Symmetric weakness in the extremities
E  Tremor at rest

A 47-year-old woman from Venezuela has had difficulty performing activities of daily living for the past year. She is emotionally labile and often cries. She is disturbed and depressed by these developments because her mother and brother died 5 years after experiencing the same symptoms. On physical examination, she has choreiform movements of her extremities. Cranial nerves are intact. She has no motor weakness and no sensory deficits. Her memory remains intact. Which of the following genetic abnormalities is most likely to be present in this woman?

A  Abnormal prion protein
B  Decreased level of hexosaminidase A enzyme
C  Extra chromosome 21
D  Expansion of CAG repeats
E  Mutation in presenilin genes

A 4-year-old girl developed clumsiness and difficulty ambulating over 6 months. On physical examination, she showed difficulty with balance while walking, dysarthria, poor hand coordination, absent deep tendon reflexes, and a bilateral Babinski sign. Light touch and vibratory sensation were greatly diminished. There was no muscular weakness. Over the next 5 years, she developed congestive heart failure from hypertrophic cardiomyopathy. She also had hyperglycemia. At autopsy, there was increased perinuclear iron deposition within cardiac myocytes. Which of the following genetic abnormalities with trinucleotide repeat expansions was most likely present in this patient?

A  CAG repeats in the huntingtin gene
B  CGG repeats in the FMRI gene
C  CTG repeats in the dystrophila myotonia-protein kinase gene
D  GAA repeats in the frataxin gene
62 A 36-year-old man who had been healthy all his life now has progressive, symmetric muscular weakness. A year ago, he noted weakness in the area of the head and neck, which caused difficulty with speech, eye movements, and swallowing. In the past year, the weakness in the upper and lower extremities has increased, and he can no longer stand, walk, or feed himself. His mental function remains intact. Which of the following is most likely being destroyed in this man?

A Ependymal cell  
B Lower motor neuron  
C Microglial cell  
D Oligodendrocyte  
E Pigmented neuron  
F Spiny neuron

63 A 12-year-old girl has had progressively diminishing neurologic function over 3 years. She has difficulty with ambulation, decreased mental ability, seizures, and loss of control over bladder and bowel functions. An MRI of her brain shows atrophy, and the centrum semiovale and central white matter are shrunken. These findings correlate with widespread microscopic myelin loss, but subcortical myelin is spared. Which of the following degenerative CNS diseases best explains her illness?

A Acute disseminated encephalomyelitis  
B Metachromatic leukodystrophy  
C Multiple sclerosis  
D Progressive multifocal leukoencephalopathy  
E Tay-Sachs disease

64 A 49-year-old man develops an acute psychosis. He has a lengthy history of chronic alcoholism. He has difficulty performing a finger-to-nose test, and there is paralysis of the lateral rectus muscles. A deficiency of which of the following nutrients is most likely to produce these findings?

A Cobalamin  
B Folate  
C Niacin  
D Pyridoxine  
E Thiamine

65 A 53-year-old man with a lengthy history of chronic alcohol abuse has had an increasingly clouded sensorium over the past 2 days. On physical examination, he has a flapping tremor of his outstretched hands. MRI of the brain shows no abnormalities. Microscopic examination of his brain would show increased numbers of neocortical and basal ganglia astrocytes with pale, swollen nuclei (Alzheimer type II cells). Which of the following laboratory findings in his blood is most likely to be associated with these findings?

A Ammonia level of 100 µmol/L  
B Carboxyhemoglobin level of 5%  
C Glucose of 30 mg/dL  
D Hemoglobin A1c level of 10%  
E Sodium of 111 mmol/L

66 A 55-year-old man has experienced headaches for the first time in his life beginning 2 months ago. He comes to the emergency department following a generalized tonic-clonic seizure. On physical examination, he has weakness on the left side. An MRI of his brain shows a large, irregular, 6-cm mass in the centrum semiovale of the right cerebral hemisphere that extends across the corpus callosum. A stereotactic biopsy of the mass is done and microscopically shows pleomorphic cells positive for glial fibrillary acidic protein (GFAP). Molecular analysis shows abnormalities of TP53 and platelet-derived growth factor-alpha (PDGF-α). Which of the following neoplasms is he most likely to have?

A Diffuse large B-cell lymphoma  
B Glioblastoma  
C Hemangioblastoma  
D Medulloblastoma  
E Pilocytic astrocytoma

67 A 10-year-old boy has had persistent headaches for the past 3 months. On physical examination, he is afebrile. He has an ataxic gait and dysdiadochokinesia. CT scan of the head shows a 4-cm cystic mass in the right cerebellar hemisphere. Cerebral lateral ventricles are enlarged. A lumbar puncture is done. The CSF protein concentration is elevated, but the glucose level is normal. Neurosurgery is performed, and the mass is removed and sectioned. On gross examination, the mass is a cyst filled with gelatinous material. The cyst has a thin wall and a 1-cm mural nodule. Microscopically, the mass is composed of cells that stain positive for glial fibrillary acidic protein (GFAP) and have long, hairlike processes. What is the most likely diagnosis?

A Astrocytoma  
B Ependymoma  
C Hemangioblastoma  
D Medulloblastoma  
E Meningioma  
F Schwannoma

68 A 40-year-old man has been experiencing headaches for the past 6 months. He had a seizure 1 day ago. On physical examination, there are no remarkable findings. MRI of the brain shows a solitary, circumscribed 3-cm mass in the right parietal centrum semiovale. The mass has small cysts and areas of calcification and hemorrhage. Neurosurgery is performed, and the mass is removed. Microscopically, the mass consists of sheets of cells with round nuclei that have granular chromatin. The cells have a moderate amount of clear cytoplasm, and they mark with GFAP by immunohistochemical staining. The patient receives adjuvant radiation and chemotherapy, and there is no recurrence. Which of the following molecular markers is most likely to be found in the cells of this mass?

A BRAF mutation  
B CD20 expression  
C c-MYC amplification  
D EGFR amplification  
E 1p and 19q co-deletions
A 46-year-old woman has had increasing weakness and loss of sensation in the lower extremities for the past 5 months. She has been unable to walk without assistance for the past week. On physical examination, there is 4/5 motor strength in the right lower extremity and 3/5 motor strength in the left lower extremity. There is bilateral loss of sensation to light touch from the lateral mid thigh distally. MRI of the spine shows a 1 × 4 cm lesion in the filum terminale. The mass is removed. Microscopically, the mass is composed of cuboidal cells around papillary cores in a myxoid background. Which of the following lesions was most likely present in this patient?

A  Choroid plexus papilloma  
B  Ependymoma  
C  Meningioma  
D  Neurofibroma  
E  Pilocytic astrocytoma  
F  Schwannoma

An 11-year-old girl has had increasing headaches upon awakening for the past month. On examination, papilledema is present bilaterally. An MRI of her brain reveals a 3-cm solid circumscribed mass within the fourth ventricle. There is third and lateral cerebral ventricular dilation. The mass is excised and microscopically shows perivascular pseudorosettes with round, regular tumor cells arranged around vessels. Which of the following neoplasms is she most likely to have?

A  Astrocytoma  
B  Ependymoma  
C  Glioblastoma  
D  Medulloblastoma  
E  Schwannoma

A 5-year-old boy has complained of headaches for the past week. His gait has become ataxic. After sudden onset of vomiting, he is brought to the emergency department, where he becomes comatose. A lumbar puncture is done and cytologic examination of the CSF shows anaplastic cells with dark blue nuclei and scant cytoplasm. An MRI is most likely to show a mass in which of the following locations?

A  Centrum semiovale  
B  Cerebellar vermis  
C  Cranial nerve VIII  
D  Fourth ventricle  
E  Parietal lobe gray-white junction

A 39-year-old HIV-positive man has received no anti-retroviral therapy. He has had left-sided weakness for the past month and experienced a generalized seizure a day ago. On physical examination, he is afebrile. CT scan of the head shows no intracranial hemorrhage, but there is a midline shift. MRI of the brain shows a 4-cm mass in the region of the putamen near the right internal capsule, a 3-cm mass in the right centrum semiovale, and a 1-cm mass near the splenium of the corpus callosum. These masses are circumscribed and solid. CSF from a lumbar puncture shows an elevated protein concentration and a normal glucose level. Cytologic examination shows large cells with large nuclei and scant cytoplasm that mark with CD19, but not with GFAP or cytokeratin. What is the most likely diagnosis?

A  Cytomegalovirus encephalitis  
B  Glioblastoma  
C  Kaposi sarcoma  
D  Large B-cell lymphoma  
E  Progressive multifocal leukoencephalopathy  
F  Toxoplasmosis

A 45-year-old woman has had unilateral headaches on the right for the past 5 months. Physical examination yields no remarkable findings. The representative gross appearance of the lesion seen on CT scan of the head is shown in the figure. The mass is surgically removed and microscopic examination shows elongated cells with pale, oblong nuclei and pink cytoplasm with occasional psammoma bodies. Cytogenetic analysis shows 22q-. What is the most likely diagnosis?

A  Astrocytoma  
B  Ependymoma  
C  Meningioma  
D  Metastasis  
E  Tuberculoma

A 76-year-old man has a single episode of grand mal seizure. On physical examination, he is afebrile and normotensive. Motor strength is intact, and there is no loss of sensation. Cranial nerves are intact. His mental function is not diminished. There is a 1-cm, darkly pigmented skin lesion on the upper back. Brain MRI shows three solid, 1- to 3-cm mass lesions, without ring enhancement or surrounding edema, located at the gray-white junction in the right and left frontal lobes. The cerebral ventricles appear normal in size. What is the most likely diagnosis?

A  Glioblastoma  
B  Hemangioblastoma  
C  Meningioma  
D  Metastatic carcinoma  
E  Non-Hodgkin lymphoma  
F  Oligodendroglioma
**75** An 18-year-old student has had decreased vision in her right eye for 6 months. On physical examination, there is papilledema on the right. She has 14 scattered, 2- to 5-cm flat, hyperpigmented skin lesions with irregular borders on the extremities and torso. CT scan of the head shows no intracranial hemorrhage and no edema or midline shift, but there is a mass in the region of the right optic nerve. An optic nerve glioma is excised. Eight months later, she returns for a follow-up examination, and a mass is palpated on the right wrist. Histologic examination of the mass is most likely to show which of the following neoplasms?

A. Fibrosarcoma  
B. Lipoma  
C. Hemangioma  
D. Meningioma  
E. Schwannoma

**76** A 41-year-old woman has diminished hearing for the past 4 months. On physical examination, she has decreased hearing on the left. Sound lateralizes to the right ear on the Weber tuning fork test. A head MRI shows a sharply circumscribed, 4-cm mass adjacent to the left pons that extends toward the left inferior cerebellar hemisphere. A smaller 1-cm lesion is in a similar location on the right. Family screening reveals a similarly affected 38-year-old sibling. An inherited mutation involving which of the following genes is most likely to be present in this patient?

A. **NF2**  
B. **TP53**  
C. **PTCH**  
D. **TSC1**  
E. **VHL**

**77** A 20-year-old woman with learning difficulties had flank pain for 1 week. Physical examination showed right costovertebral angle tenderness. Patches of leathery-appearing (shagreen patches) and hypopigmented (ash-leaf patches) skin were scattered over her body. There was a subungual nodule on her right index finger. Abdominal CT scan showed bilateral renal cysts and tumor masses. MRI of the brain showed subependymal nodules and 1- to 4-cm cortical foci with loss of the gray-white distinction. CT scan of the chest showed a 3-cm mass involving the interventricular septum. Two years later, she now has sudden, severe headache. MRI now shows a nodule obstructing the cerebral aqueduct. Neurosurgery is performed, and a subependymal giant cell astrocytoma is removed. What is the most likely diagnosis?

A. Down syndrome  
B. Krabbe disease  
C. Neurofibromatosis type 1  
D. Neurofibromatosis type 2  
E. Tuberous sclerosis  
F. Von Hippel–Lindau disease

**78** A study of adults with cerebellar neoplasms reveals that some of them have an autosomal dominant inheritance pattern with von Hippel–Lindau disease. Their cerebellar tumors are cystic with a mural nodule. Molecular analysis of tumor cells shows increased amounts of hypoxia-inducible factor (HIF). The incidence of renal cell carcinomas is increased in these persons. Which of the following paraneoplastic manifestations is most likely to be found in these adults?

A. Cushing syndrome  
B. Hypercalcemia  
C. Polycythemia  
D. Syndrome of inappropriate ADH  
E. Trousseau syndrome

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**ANSWERS**

1. **D** Hypoxic and hypoglycemic injury leads to “red” neurons as the initial reaction. The larger pyramidal neurons are the most sensitive, particularly in the hippocampus with hypoglycemia, as well as neocortical Betz cells and cerebellar Purkinje cells. Astrocytes proliferate in reaction to brain injury in the process called gliosis. Endothelial cells are most likely to proliferate in neoplasms. Microglia respond to brain injury with a macrophage-like function. Oligodendroglial cells provide myelination for brain neuronal axons.

2. **D** Microglial cells are part of the body’s fixed macrophage system, derived originally from mesoderm. Microglia can respond to cerebral injuries by taking on a macrophage-like function. The arachnoid layer is part of the meninges covering the brain. Astrocytes can proliferate in response to brain injuries, a process called gliosis, but there is no phagocytosis. Ependymal cells line the ventricular system containing CSF. Oligodendroglial cells provide the myelin sheaths for neuronal axons in the CNS.

3. **C** The papilledema and the herniation are a consequence of brain swelling, typically the vasogenic form of edema from blood-brain barrier disruption adjacent to the neoplasm. A large aggressive neoplasm, such as a glioblastoma, may produce a mass effect via enlargement from rapid growth, hemorrhage, and surrounding edema. The mass effect with herniation of the medial temporal lobe results in a third cranial nerve palsy as the nerve is compressed. A chronic subdural hemorrhage accumulates slowly enough that herniation may not occur. An abscess may cause a mass effect with some associated brain swelling, but this patient is afebrile. There is no pressure effect with hydrocephalus ex vacuo, which is a consequence of cerebro atrophy. An infarct is not likely to produce pronounced associated brain swelling, and a mass effect in occipital lobe is unlikely to affect the third nerve. Rupture of a berry aneurysm produces subarachnoid hemorrhage at the base of the brain, which is less likely to cause a mass effect.

4. **D** This is noncommunicating hydrocephalus with obstruction below the level of the third ventricle. If hydrocephalus had been present at birth, there would be increasing...
head size because the sutures are not yet closed, and congenital aqueductal stenosis would be suspected. At his age, a neoplasm should be suspected, and ependymomas arise in the ventricular system, often in the fourth ventricle, to cause obstruction of CSF flow. The increased CSF protein comes from this tumor, but shedding of cells from the mass into the CSF is unlikely. Except for vascular malformation, the other options are uncommon at his age. An abscess is typically accompanied by fever, and most would be located in the cerebral hemispheres away from the ventricular system. Cryptococcal meningitis is accompanied by fever, and exudate can be found within the ventricular system and subarachnoid space, but there is more likely to be cerebral edema, not hydrocephalus. The demyelinating plaques of multiple sclerosis are small and do not usually act as mass lesions. Vascular malformations usually arise in the cerebral hemispheres.

PBD9 1254–1255  BP9 812–813  PBD8 1283  BP8 861–862, 884

5 F He has communicating hydrocephalus, and pneumococcal meningitis in adults often involves the vertex, where the arachnoid granulations that reabsorb CSF are found. Overproduction of CSF from a choroid plexus papilloma may also produce hydrocephalus, but is rare. Non-communicating hydrocephalus occurs when there is obstruction within the ventricular system. Both aqueductal stenosis and Dandy-Walker malformation are congenital conditions causing ventricular system obstruction. Cerebral infarcts and HIV infection produce lesions within the brain parenchyma, and are unlikely to obstruct the CSF flow unless there is a mass effect (from edema or hemorrhage with the infarction, or a CNS lymphoma with AIDS), but no mass was noted in this man’s CT scan.

PBD9 1254–1255  BP9 813  PBD8 1283  BP8 861–862

6 A Her contralateral right leg weakness is a consequence of subfalcine herniation with compression of the ipsilateral anterior cerebral artery caused by a lesion in the adjacent frontal lobe. A rapidly expanding mass lesion can lead to herniation. The other listed options, above the tentorium, are much less likely to explain anterior cerebral artery compression. Frontal lobe lesions might account for transtentorial herniation. A very large mass effect above the tentorium could also produce tonsorial herniation.

PBD9 1255  BP9 813–814  PBD8 1283  BP8 862

7 B Cerebral edema may be cytotoxic or vasogenic. In cytotoxic edema the blood-brain barrier remains intact, and edema is due to failure of ATP-dependent ion transport with cellular retention of sodium and water. Global processes such as ischemia (from heart failure in this case) or metabolic derangements are implicated, and consistent with diffuse swelling with bilateral papilledema. Inflammation and neoplasms are more likely to produce vasogenic edema. An acute meninitis might produce diffuse swelling, but there are no signs of infection here. Metastases are unlikely to produce rapidly expanding mass lesions. Obstruction of CSF flow leads to hydrocephalus with ventricular enlargement, not narrowing. Subarachnoid hemorrhage is most likely a consequence of a ruptured berry aneurysm, without a mass effect.

PBD9 1254  BP9 813  PBD8 1282  BP8 861

8 B The figure shows linear midline hemorrhages, called Duret hemorhages, in the pons. The acute bacterial meningitis led to brain swelling with edema and subsequent herniation of medial temporal lobe with Duret hemorrhages in the pons. Although not seen in this figure, an infection could organize with scarring of foramina to produce a noncommunicating hydrocephalus, or it could scar the vertex and impair reabsorption of CSF at the arachnoid granulations to produce a communicating hydrocephalus. An abscess infrequently complicates meningitis; conversely, an abscess in a paranasal sinus or mastoid air cell may extend into the cranial cavity to cause meningitis. Laminar necrosis could occur after brain death, but this finding is not specific for meningitis. The small meningeal vessels do not often bleed because of inflammation caused by meningitis.

PBD9 1255  BP9 814  PBD8 1284  BP8 862–863

9 B Anencephaly is a form of severe neural tube defect that results from failure of formation of the fetal cranial vault. This is one of the most common CNS malformations seen at birth. The defect allows fetal α-fetoprotein to enter amniotic fluid and reach the maternal circulation. The karyotype listed is that of a Robertsonian Down syndrome carrier; Down syndrome (trisomy 21) may be associated with brachycephaly, but rarely with anencephaly. Congenital cytomegalovirus infection can produce extensive fetal brain parenchymal necrosis, but not loss of the fetal cranial vault. Neural tube defects are not associated with maternal or neonatal jaundice. Diabetes mellitus, suggested by an elevated hemoglobin A1c concentration, can increase the risk of malformations (e.g., holoprosencephaly in the CNS), but not neural tube defects.

PBD9 1256  BP9 822–823  PBD8 1284  BP8 872

10 D Holoprosencephaly is a midline defect in which there is absent (alobar) or partial (semilobar) cerebral hemispheric development. It can occur in trisomy 13, as in this case, with other midline defects. It also may be seen in cases of maternal diabetes mellitus. Anencephaly is the absence of a fetal cranial vault, which leads to absence of most of the brain. Arnold-Chiari II malformation results in a small posterior fossa, a misshapen midline cerebellum with downward displacement of the vermis, and tenting of the tectal plate. Dandy-Walker malformation is characterized by aplasia or hypoplasia of the cerebellar vermis, cystic enlargement of the fourth ventricle, and hydrocephalus. Periventricular leukomalacia is a form of perinatal injury that is caused by hypoxic-ischemic events or infections.

PBD9 1257  BP9 823  PBD8 1285  BP8 872

11 E The syrinx is a tubular defect that extends laterally, disrupting the spinothalamic tracts and anterior horns, producing sensory and motor deficits. When the cavity extends
into the medulla, it is called syringobulbia. There is no treatment, but the lesion tends not to progress. Dandy-Walker malformation is a congenital condition with an enlarged posterior fossa. Diffuse axonal injury involves white matter tracts in the cerebrum. Rachischisis is a large open neural tube defect. Spina bifida is a posterior neural tube defect.

12 A Arnold-Chiari II malformation has features including in a small posterior fossa, misshapen midline cerebellum with downward displacement of the vermis, and tenting of the tectal plate, leading to hydrocephalus. MRI in this case is characteristic of hydromyelia. Cerebral palsy is a general term describing nonprogressive motor deficits that are present from birth. The corpus callosum is seen in the figure; agenesis may be associated with other anomalies, or may be found in normal persons. Holoprosencephaly is a severe malformation with total (alobar) or incomplete (semilobar) separation of the cerebral hemispheres in brain development. Polymicrogyria is characterized by numerous small, irregularly formed gyral contours.

13 C Germinal matrix hemorrhage is the most common cause of intraventricular hemorrhage in premature infants. The germinal matrix, composed of highly vascularized tissue with primitive cells, is most prominent between 22 and 30 weeks’ gestation. Hemorrhages within this area readily occur with common neonatal problems such as hypoxemia, hypercarbia, acidosis, and changes in blood pressure. Hemorrhages in the germinal matrix can extend into the cerebral ventricles and from there into the subarachnoid space. Smaller hemorrhages can resolve without sequelae. With larger hemorrhages, organization of the blood in the aqueduct of Sylvius or the fourth ventricle or foramina of Luschka and Magendie may obstruct the flow of CSF, producing hydrocephalus. Cytomegalovirus infection can cause considerable necrosis of the brain parenchyma, particularly in a periventricular location, but not hemorrhage with obstruction of CSF flow. Infants with Down syndrome may have vascular malformations that can bleed into the parenchyma. The bilirubin staining of kernicterus does not result in meningal or aqueductal scarring. Medulloblastomas are aggressive posterior fossa tumors that occur in children (not typically in infants) that could cause obstruction of CSF flow.

14 B Concussion leads to altered consciousness, starting with instantaneous onset of transient neurologic dysfunction with head trauma, amnesia for the event, then neurologic recovery or post-concussion syndrome with disabling neuro-psychiatric manifestations, worsening with repeated concusions. There are no radiologic or pathologic findings. Arteriolar sclerosis is typically seen with hypertension. Hemorrhage following the traumatic event could organize to obstruct CSF flow or absorption, but this would have been identified with the original CT scan. The term leukoencephalopathy applies to degenerative white matter changes, typically with metabolic and infectious insults. Post-traumatic dementia from repeated blows to the head often has pathologic findings of neurofibrillary tangles and neuritic plaques similar to Alzheimer disease.

15 E This patient has the classic “contrecoup” type of injury, in which the moving head strikes an object, and the force is transmitted to the opposite side of the head. A fall backward is most likely to produce contusions to the inferior frontal lobes, temporal tips, and inferior temporal lobes. A blow to a stationary head is more likely to produce a “coup” injury directly adjacent to the site of the blow. Basal ganglia putaminal hemorrhage is most likely to occur in hypertension. Hemorrhage into the pons is typical of a Duret hemorrhage, seen in medial temporal lobe herniation. Hemorrhage into the cerebral ventricles may occur in premature infants from germinal matrix hemorrhage; in adults, it is uncommon, but may occur with dissection of blood from an intraparenchymal lesion. A blow to the side of the head is more likely to lacerate the middle meningeal artery and produce an epidural hematoma. Subarachnoid hemorrhage at the base of the brain may dissect into the sella.

16 C The “lucid” interval is a classic feature of an epidural hematoma with rapid accumulation of blood from the injured middle meningeal artery. The dura is pushed against the brain, producing the lens-shaped blood collection seen with radiologic imaging, particularly CT imaging. Leptomeningitis is not associated with trauma. Contusions do not progressively worsen. In an acute subdural hematoma or a ruptured aneurysm, there typically is no lucid interval, but instead a sudden and progressive worsening of symptoms.

17 E A subdural hematoma results from tearing of the bridging veins beneath the dura. These veins are at risk of tearing with head trauma, particularly in elderly individuals, in whom some degree of cerebral atrophy may be present. Bleeding from low-pressure veins produces a variable time course for appearance of signs and symptoms, from hours to days to weeks. Basal ganglia hemorrhages are most often associated with hypertension. Epidural hemorrhages are most often preceded by a blow to the head that tears the middle meningeal artery; there is commonly a “lucid” interval between an initial loss of consciousness occurring with trauma and the later accumulation of blood. Pontine hemorrhages are likely to be Duret hemorrhages. Subarachnoid hemorrhage could occur in contusions with trauma.

18 D Tearing of bridging veins leads to an acute subdural hematoma; this almost always results from head trauma in which there has been a fall, which may have been minor and
gone unnoticed. The risk of hemorrhage is greater in elderly individuals because of cerebral atrophy, which leaves the bridging veins beneath the dura at the vertex more vulnerable to traumatic tearing. Arteriovenous (vascular) malformations are often located within the parenchyma of a hemisphere, and bleeding from them occurs more often in young adults. When saccular (berry) aneurysms rupture, the bleeding is typically subarachnoid and at the base of the brain. A tear of the middle meningeal artery can occur in head trauma (e.g., a blow to the head), but it results in an acute epidural hematoma. Thrombosis of an intracranial artery may result in an infarction, which can be hemorrhagic, but the hemorrhage typically does not extend into subarachnoid or subdural locations.

19 C Global cerebral ischemia has occurred in this man, leading to brain death. There is ongoing liquefactive necrosis, and in 2 to 3 weeks there will be numerous macrophages that phagocytize the cellular debris. These macrophages can persist for months. The neocortex has more vulnerable pyramidal cell layers with a pattern of pseudolaminar necrosis. Fibroblastic proliferation with collagen deposition is uncommon in the CNS, except around organizing abscesses. An abscess is where neutrophils would be found, as well as meningitis with acute meningitis. Inflammation of brain parenchyma is most often lymphocytic. Oligodendrocytes provide myelination to axons. Dying red neurons are seen after 12 hours following ischemic injury and may persist for days, but not weeks.

20 B This cerebral infarction is a large hemorrhagic and softened area of beginning liquefactive necrosis after vascular injury in the distribution of the middle cerebral artery. Emboli likely emanate from the heart. It takes weeks to months for macrophages to clear the debris of liquefactive necrosis and leave a cystic space. A vascular malformation has irregular vascular channels that give a mass effect with a dark red-to-bluish appearance. Metastases may be solitary masses but are most often multiple, and may have central hemorrhage or necrosis, but are often multiple. An organizing subdural hematoma would leave a uniform area of compression with flattening of the underlying hemisphere. A contusion could produce some minimal focal loss of cortex with brown hemosiderin staining.

21 B The brief episodes of neurologic dysfunction represent transient ischemic attacks (TIAs) and are a prodrome to stroke in many cases. Atherosclerotic cerebrovascular disease is a common antecedent to cerebral infarction, and metabolic syndrome with dyslipidemia and hyperglycemia is a risk factor. A vascular malformation most often produces symptoms caused by bleeding in young adults. A neoplasm is unlikely to produce such sudden, episodic symptoms and signs. Meningoencephalitis may produce general features such as fever, headache, confusion, and seizures, but not sudden localizing signs. A subdural hematoma, which most frequently results from head trauma sustained in a fall, is unlikely to develop in a few minutes and would not explain the TIAs.

22 D The neurons that are most sensitive to anoxia reside in the hippocampus, along with the cerebellar Purkinje cells and the larger neocortical neurons. In addition, the first areas in the neocortex to be affected are the “watershed” areas between the three major cerebral circulations, including the watershed located superiorly between the anterior cerebral and middle cerebral circulations, as in this study. “Red” shrunken neurons, especially in the areas mentioned, are typically seen in the early stages of global hypoxia, as may occur in a severe hypotensive episode. Focal hypoxia leads to infarcts. Autoimmunity can lead to vasculitis and subsequent hypoxia. Chemotherapy tends to damage actively dividing cells more severely. Diabetes mellitus causes atherosclerosis, which can lead to hypoxia. Lead poisoning leads to encephalopathy, not ischemia. Poor nutrition with a deficiency of thiamine (vitamin B1) can lead to Wernicke disease, which affects the mammillary bodies and periaqueuductal gray matter most severely.

23 E An acute cerebral infarction results from obstruction of blood flow causing focal cerebral ischemia. After 2 days, there would be some cerebral softening and edema with ischemia of neurons, but little else. Hypercholesterolemia is a risk factor for atherosclerosis, which is the major cause of thrombotic cerebral arterial occlusions. The antiphospholipid syndrome can produce thrombotic and embolic disease, but an embolic “stroke” is typically hemorrhagic, and antiphospholipid syndrome is uncommon at this age. A positive blood culture suggests sepsis with the possibility of meningitis or cerebral abscess formation, but abscesses typically have ring enhancement on CT scans. Elevated serum levels of very long chain fatty acids are present in patients with adrenoleukodystrophy, a rare disorder that leads to myelin loss at an early age. Hyperammonemia occurs in hepatic encephalopathy with liver failure; it produces Alzheimer type II gliosis, but no focal or gross lesions. Neurosyphilis is now rare; it does not produce focal ischemic lesions.

24 E A remote cerebral infarction is present. Thromboembolic disease with cerebral infarction most often results from a cardiac disease (e.g., endocarditis, mural thrombosis, prosthetic valvular thrombosis). Hemorrhagic infarcts are more likely to occur when emboli partially occlude a vessel or undergo dissolution. Thrombosis over atherosclerotic lesions is more likely to cause nonhemorrhagic infarcts. AIDS is not often associated with significant cardiovascular or cerebrovascular diseases. Head trauma with hemorrhage is more common in individuals with chronic alcoholism. Hypertension is most often associated with basal ganglia, pontine, and cerebellar hemorrhages and with small lacunar...
infarcts. Malignant neoplasms may be associated with para-neoplastic syndromes, including hypercoagulability and thrombosis, but this is much less common than heart disease as a cause for cerebrovascular disease and stroke.

25 D Primary angiitis of the CNS is uncommon. It involves small peripheral vessels. The cause is unknown, but it appears to be immunologically mediated, so immunosuppressive therapy is of benefit, such as cyclophosphamide and methylprednisolone. Infections are unlikely to produce such focal involvement of small vessels. Hypertensive encephalopathy produces fibrinoid necrosis of arterioles. NSAIDs or aspirin as anti-inflammatory or anti-platelet agents are not strong enough to address the inflammatory process in this case. Mannitol infusion is used to treat marked cerebral edema, but that is not the major finding in this case.

26 D Hypertensive hemorrhages are most likely to arise in the basal ganglia (shown in the figure), thalamus, cerebral white matter, pons, or cerebellum. Multiple hemorrhages are uncommon. The small vessels weakened by hyaline arteriolosclerosis are prone to rupture. Chronic alcoholism predisposes to falls with subdural hematomas or contusions. Metastases are typically multiple and peripheral at the gray-white junction. The plaques of multiple sclerosis occur in white matter and do not bleed. Thromboemboli can produce ischemic cerebral infarctions, sometimes with hemorrhage, and they usually involve the cerebral cortex.

27 A He developed acute hypertensive encephalopathy from the rapid and marked increase in blood pressure. There is extensive cerebral edema with increased intracranial pressure, but no localizing signs. The arterioles are involved, and like malignant hypertension involving the kidney, there is fibrinoid necrosis and petechial hemorrhages. Telangiectasias may be one form of vascular malformation, typically a localized lesion not related to hypertension. Lacunar infarcts develop with hypertension, but they tend to be small and are often silent, but may cause significant neurologic impairment, though more likely a focal deficit due to focal involvement. Hypertensive hemorrhages may be large, and the basal ganglia region is a common location, but such a lesion could produce a unilateral mass effect with papilledema on one side. Subarachnoid hemorrhages are more likely to occur from ruptured saccular aneurysms.

28 A This patient’s stepwise loss of cognitive function is consistent with vascular dementia, and his history of diabetes mellitus makes underlying heart disease with embolic events, or vascular disease involving the brain, more likely. Some cases have diffuse cortical laminar necrosis from global hypoxic events, whereas others have multiple lacunar infarcts, and still others have embolic infarcts. Lewy body dementia mimics Alzheimer disease (AD), with progressive cognitive decline, and a parkinsonian movement disorder is often present. Neuritic plaques and neurofibrillary tangles are characteristic for AD, which has progressive cognitive decline. Pick bodies are seen with Pick disease, which is similar to AD. Plaques of white matter demyelination are seen with multiple sclerosis, and the gray matter is relatively spared, so that severe cognitive impairment is not a usual feature.

29 A Alzheimer disease (AD) can be complicated by cerebral amyloid angiopathy and terminal hemorrhagic stroke. Formation and aggregation of the Aβ peptide is now considered central to the pathogenesis of AD. Aβ peptide is derived from abnormal processing of amyloid precursor protein (APP). When APP, a transmembrane protein, is cleaved by a-secretase within the Aβ sequence, followed by γ-secretases, a soluble nontoxic fragment is formed. Cleavage of the Aβ sequence by β-secretase and then γ-secretase gives rise to Aβ peptides that aggregate and form the amyloid cores that elicit a microglial and astrocytic response to form neuritic plaques. Conformational change in prion protein leads to Creutzfeldt-Jakob disease, a rapidly progressive dementia with spongiform encephalopathy but not neuritic plaques or amyloid deposition. Loss of dopaminergic neurons with deficiency of dopamine is central to the pathogenesis of Parkinson disease. Expansion of polyglutamine repeats owing to CAG trinucleotide repeat-expansion underlies Huntington disease. Although abnormally phosphorylated forms of tau protein are found in neurofibrillary tangles seen in AD, there is no mutation of the tau gene, and the tangles are not considered primary in the pathogenesis of AD. Mutation of the tau gene can be found in frontotemporal lobe dementias.

30 D About 1 in 50 individuals has a saccular (berry) aneurysm. Although this lesion is present at birth as a congenital defect in the arterial media at intracerebral arterial branch points, it can manifest later in life with aneurysmal dilation and possible rupture. These aneurysms are the most common cause of spontaneous subarachnoid hemorrhage in adults. The bleeding from amyloid angiopathy is in peripheral cortex, and most likely to occur in association with Alzheimer disease. A hypertensive hemorrhage from arteriolosclerosis tends to remain within the brain parenchyma. Thromboemboli can cause infarctions, most often in the distribution of the middle cerebral artery in cortex, and embolic infarcts can be hemorrhagic, but the blood typically does not reach the CSF. A subdural hematoma over the brain surface results from a tear of bridging veins.

31 D Intracranial aneurysms are typically saccular and enlarge slowly over time. Aneurysms that grow to 4 to 7 mm are at the greatest risk of rupture. Rupture occurs into the subarachnoid space at the base of the brain, where the cerebral arterial distribution originates around the circle of Willis, and where saccular aneurysms are most likely to arise. Neither a
berry aneurysm nor the bleeding that results is likely to cause a mass effect and herniation. In some cases of survival after rupture of a berry aneurysm, a noncommunicating hydrocephalus results from organization of the subarachnoid hemorrhage occluding foramina of Luschka and Magendie. Epidural hematomas arise from a tear of the middle meningeal artery, typically as a result of head trauma. Trauma also can cause a tear of bridging veins that produces a subdural hematoma.

**32 B** Arteriovenous (vascular) malformations most often occur in the cerebral hemisphere of a young adult and can often be completely resected without complication. There may be slow leakage of blood from the lesion over time, resulting in the clinical symptoms and the gliosis seen on biopsy in this case. An angiosarcoma with malignant-appearing cells is not a primary lesion in the brain parenchyma. A plaque of demyelination in multiple sclerosis can appear as a mass, but its features include loss of myelin with gliosis and macrophages, not vascular abnormalities. An abscess would have an organizing wall with collagen and gliosis, but no prominent larger vessels. Head trauma generally produces contusions and hematomas on the surface, but not hemorrhages in the brain parenchyma. A ruptured aneurysm can extend upward into the parenchyma in some cases; the outcome is always fatal with this complication.

**33 C** Acute meningitis, with a purulent exudate on the cerebral convexities shown in the figure, is indicative of bacterial infection. At his age, a common etiologic agent is *Neisseria meningitidis*, initially as pharyngitis, and untreated proceeding to Waterhouse-Friderichsen syndrome with disseminated intravascular coagulopathy. Cryptococcosis should be considered in immunocompromised patients, but some cases occur in immunocompetent patients; a meningoencephalitis can occur. Tuberculous meningitis does not manifest so acutely, and the exudate is typically on the base of the brain. Poliomyelitis could occur after pharyngitis, but onset is insidious, with increasing paralysis from loss of motor neurons. It does not cause meningitis. Cerebral toxoplasmosis may occur in immunocompromised patients, but the lesions are parenchymal abscesses, not meningitis.

**34 D** Headache, fever, pronounced neutrophilia, a high CSF protein level, and a low glucose concentration all point to bacterial meningitis. *Staphylococcus aureus* is a common infection among injection drug users. Herpes simplex virus produces encephalitis, not meningitis. Tuberculous meningitis has a more insidious onset. This patient does not have a mass lesion of toxoplasmosis or focal lesions of progressive multifocal leukoencephalopathy (PML) that may be associated with AIDS. PML is associated with the JC polyomavirus. Toxoplasmosis, which may occur in immunocompromised patients, produces parenchymal abscesses, not meningitis.

**35 C** The most common etiologic organism for acute bacterial meningitis in his age group, because of vaccinations, is now *Streptococcus pneumoniae*, a gram-positive coccus. Pneumococci also are likely to be seen in an adult. The number of CNS infections with *Haemophilus influenzae*, a gram-negative bacillus, in this age group has decreased because of widespread immunization. The gram-negative diplococci of *Neisseria meningitidis* are seen in young adults. The gram-negative bacilli of *Escherichia coli* are most often seen in neonates. The short, gram-positive rods of *Listeria monocytogenes* appear sporadically or in epidemics caused by food contamination. With these clinical and CSF features, it would be unusual if bacteria were not present in the CSF.

**36 A** A cerebral abscess is most often a complication of an infection, such as pneumonia or endocarditis, with onset days to weeks earlier. The bacteria spread hematogenously. As the abscess organizes, it is ringed by fibroblasts that deposit collagen; this feature is characteristic of an abscess in the CNS. A neoplasm occasionally may be ring enhancing, but a glioblastoma is an aggressive malignancy that is not well demarcated. Metastases are mass lesions that are typically multifocal. A multiple sclerosis plaque is generally not large, is found in white matter, and does not typically have ring enhancement. An infant would produce sudden signs and symptoms that improve over time, and the CSF protein would not be increased.

**37 B** Acute otitis media may be complicated by spread of the infection to the mastoid air cells, and then to skull bone, and then to the epidural space. This also may be termed *pachymeningitis* or *epidural empyema*. Spread to the underlying meninges may next occur. Encephalitis involves the substance of the brain. Multicystic encephalopathy is a severe complication of perinatal brain injury with infarction. Subdural hematomas are most often caused by trauma. The trigeminal nerve is located at a distance from the mastoid air cells so that it is unlikely to be involved; but the facial nerve adjacent to the ear may be involved.

**38 C** Tuberculous meningitis is a complication of disseminated tuberculosis, and young children are at increased risk in locations where tuberculosis is prevalent. It tends to produce a more chronic course. Cranial nerves, such as the abducens in this case, can be involved. A large granulomatous mass, a tuberculoma, may complicate some cases. Viral “aseptic” meningitis is marked by mononuclear cells, but the protein is not usually markedly elevated, and no mass develops. Bacterial meningitis is often associated with a low glucose, but not a mass, and the inflammatory response is predominantly polymorphonuclear. *T. pallidum* is the causative agent for neurosyphilis with a long, insidious course and no mass effect. *T. solium* can lead to cysticercosis, with cystic masses in the brain.
39 F He has neurosyphilis from infection by *Treponema pallidum*. This form of tertiary syphilis occurs years following primary infection. His eye finding is consistent with Argyll Robertson pupil; his positive Romberg test is the result of tabes dorsalis. His mentation has changed from general paresis. Listeriosis and streptococcal infections are most likely to produce an acute meningitis. Rabies may have an incubation period of 1 to 3 months, but then manifests with an acute severe encephalitis. Rubeola (measles) infection rarely leads to subacute sclerosing panencephalitis years later, marked by spasticity and seizures. Toxoplasmosis is an opportunistic infection with encephalitis and abscess developing over weeks.

PBD 1274–1275, BP 826, PBD 831–832, BP 875

40 B An acute lymphocytic meningitis is most typically caused by a virus, such as West Nile virus, an equine encephalitis virus, or an echovirus. It is sometimes referred to as *aseptic meningitis* because routine Gram staining and bacterial cultures are negative. Most cases are self-limited, occur in immunocompetent individuals, and resolve without significant sequelae. Cryptococcal and toxoplasmal infections can occur in immunocompromised patients, but the India ink test would be positive in the former, and CT scan would show focal ring-enhancing lesions in the latter. Listerial and meningococcal infections are seen sporadically and have a neutrophilic response.

PBD 356, 1275, BP 826–827, PBD 351, 1302, BP 874

41 C Hemorrhagic lesions of the temporal lobes are characteristic of herpes simplex virus encephalitis; cases are few and usually sporadic, occurring in apparently healthy individuals. The lesions of aspergillosis can be hemorrhagic, but they are typically seen in immunocompromised individuals. Cytomegalovirus infection occurs in neonates and in immunocompromised adults, but it does not produce hemorrhagic lesions. Arboviral infections produce focal lesions that may have an associated vasculitis with hemorrhage, but the CSF protein is usually elevated, and there is a neutrophilic pleocytosis. Meningococcal infections produce meningitis.

PBD 1275–1276, BP 827–828, PBD 831–832, BP 876–877

42 A Cytomegalovirus (CMV) infection is one of the congenital TORCH infections, and it can become widely disseminated to affect the CNS. Periventricular leukomalacia is characteristic of CMV infection. Group B *Streptococcus* infections cause premature rupture of membranes and sepsis without significant CNS findings. Heart failure in utero causes hydrops fetalis. Herpes simplex virus infection of neonates typically occurs during passage through the infected birth canal, not in utero. HIV infection produces no significant CNS findings in the perinatal period. Listeriosis can produce focal microabscesses in various organs, but usually there is minimal necrosis.

PBD 359–360, 1277, BP 828, PBD 833–835, 1304, BP 877

43 A Poliomyelitis is described, starting with the minor gastrointestinal illness, followed by the acute major illness, then recovery, then the late post-polio syndrome. This enterovirus attacks lower motor neurons, leading to neurogenic muscle weakness and atrophy. The anterior horns of the spinal cord include lower motor neurons. Cranial nerve nuclei with lower motor neurons may also be involved. The basal ganglia modulate muscle movement, and involvement of these structures may result in movement disorders, but not paralysis. The corticospinal tracts are axons from upper motor neurons found in the neocortex, and neither is involved with polio. The dorsal root ganglia are involved with sensory pathways. Myoneural junctions may be involved with myasthenia gravis or clostridial infections (botulism, tetanus).

PBD 356, 1277, BP 828, PBD 830–831, 1304, BP 878

44 B The child had rabies, an infectious disease in which the virus travels from the site of an infected animal bite up nerves to the CNS. Pathognomonic Negri bodies are cytoplasmic inclusions found in the hippocampal pyramidal cells and cerebellar Purkinje cells. Worldwide, unvaccinated dogs remain the most common vector for transmission of this bullet-shaped virus to man. The suspected animal must be sacrificed to identify the virus in the brain. Infected neurons generally do not undergo cell death; instead the virus induces neurotoxin production that affects surface receptor interaction with neurotransmitters. After symptoms develop, survival is rare. Multinucleated cells may be seen in microglial nodules associated with HIV infection. Perivascular lymphocytic infiltrates are often seen with a variety of viruses, such as arboviruses or echoviruses, causing encephalitis. *Toxoplasma gondii* infection is characterized by finding enlarged, infected cells filled with bradyzoites. Spongiform change is characteristic for prion disease.

PBD 1277–1278, BP 828, PBD 830–831, BP 878

45 B AIDS dementia complex occurs late in the course of HIV infection. HIV-1 produces an encephalitis characterized by a collection of reactive microglial cells (microglial nodules). HIV-1-infected mononuclear cells, particularly macrophages, can fuse to form multinucleate cells, which are seen within microglial nodules. Spongiform change suggests Creutzfeldt-Jakob disease, a rapidly progressive dementia unrelated to HIV infection. Lacunar infarcts are small, cavitary infarcts that result from arteriolosclerosis of the deep penetrating arteries and arterioles. Such arteriolar lesions occur in individuals with long-standing hypertension and are unlikely to be found in a 37-year-old man. Neocortical neuritic plaques and neurofibrillary tangles are typical of Alzheimer disease and are unlikely to manifest at this patient’s age. Plaques of demyelination are typical of multiple sclerosis.

PBD 1278, BP 828, PBD 830, BP 877–878

46 C Progressive multifocal leukoencephalopathy is caused by the JC polyomavirus and occurs in immunocompromised individuals, including individuals with AIDS. The patient was treated for chronic myelogenous leukemia. Cytomegalovirus infection also complicates the course of immunocompromised patients, but it causes large intranuclear
inclusions, most often in endothelial cells. Herpes simplex virus is uncommon, even in immunocompromised patients, and it most often produces hemorrhagic encephalitis in temporal lobes. Rabies virus produces CNS excitability with convulsions, meningismus, and hydrophobia. Subacute sclerosing panencephalitis is a rare complication of measles (rubeola) virus infection and leads to progressive mental decline, spasticity, and seizures. West Nile virus, similar to many arboviruses, can cause a meningoencephalitis.

**47** D Cryptococcal meningoencephalitis is a complication of his immunocompromised state. *Cryptococcus neoformans* typically has a thick capsule, making it easily visible with the India ink preparation, a procedure that can be performed within a few minutes on a CSF sample. A cryptococcal antigen test on the CSF would also be useful for this patient. Brain biopsies are not commonly performed, and other, less invasive methods should be pursued first. Bacterial meningitis is possible, and pneumococcus would be a common bacterial cause, but this description is consistent with cryptococcosis.

Of the remaining fungal organisms listed, none are readily identified with India ink. *Aspergillus* spp. appear as branching septate hyphae, not as yeasts. *Blastoscyces dermatitidis* is characterized by broad-based budding yeasts. *Coccidioides immitis* yeast forms appear as large spherules containing endospores. *Histoplasma capsulatum* organisms are small and tend to be found within macrophages.

**48** D Cysticercosis from eating uncooked pork can result in the release of larvae that penetrate the gut wall and disseminate hematogenously, often settling in gray and white cerebral tissue, where they develop into cysts. The cysts may cause obstructive hydrocephalus. Neurocysticercosis is a major cause for seizures in parts of the world where pork tapeworms are ingested. Aspergillosis is a fungal disease in which the foci of inflammation grossly resemble granulomas, but there is often minimal inflammatory response, and the propensity for vascular invasion often produces a hemorrhagic border to the lesions. Cryptococcosis most often involves the lungs and meninges. Malaria caused by *Plasmodium falciparum* produces hemolytic anemia, splenomegaly, and cerebral thrombosis. Toxoplasmosis can be a congenital infection. In immunocompromised adults, it can produce inflammation in multiple tissues, but most often it causes chronic abscessing inflammation in the brain. African trypanosomiasis produces sleeping sickness.

**49** E Toxoplasmosis is a common opportunistic infection that affects the CNS in immunocompromised patients, including those with AIDS, and produces abscesses that organize on the periphery to produce a bright ring on CT and MRI. *Candida* fungal infections of the CNS are rare, and disseminated candidiasis in AIDS is uncommon. Malignant lymphomas also can produce this picture, but without fever, and they generally occur as fewer, larger masses or as a solitary mass. Metastatic carcinoma, which typically presents with multiple lesions, is uncommon at this age. A spongiform encephalopathy, such as Creutzfeldt-Jakob disease, typically has no grossly visible or radiographic findings.

**50** E A rapidly progressive dementia is most consistent with Creutzfeldt-Jakob disease (CJD). CJD is one of a group of diseases that are called spongiform encephalopathies because they produce a microscopic vacuolated appearance of the neocortex. CJD occurs in sporadic or familial forms and is caused by prion protein. The normal prion protein of the brain, designated PrPc, can undergo conformational change to PrPsc, which then induces further change in PrPsc to PrPsc, which patient’s age and clinical findings are characteristic of the sporadic type of CJD, which may arise from spontaneous mutation or from exposure to PrPsc. Variant CJD, which may be linked to exposure to bovine spongiform encephalopathy, occurs in much younger patients and does not produce the characteristic EEG findings. Microglial nodules may be seen in patients with AIDS. Neuritic plaques are seen in Alzheimer disease, which occurs over many years. The demyelinating plaques of multiple sclerosis develop over years, as do the Lewy bodies of Parkinson disease and diffuse Lewy body disease.

**51** F Multiple sclerosis (MS) produces white matter plaques of demyelination, and most patients develop optic neuritis with visual difficulties, often unilateral. The course of MS varies, with many relapses and remissions, and some patients are affected more severely than others. Some patients have minimal problems, but most can be expected to have further neurologic problems. Severe neurologic impairment and death are unlikely, and most patients live for decades. There is no defined inheritance pattern. Although MS is an immunologically mediated disease, immune dysregulation leading to development of lymphoid neoplasms is unlikely to occur. The focality of white matter lesions and the sparing of gray matter make dementia and quadriplegia unlikely complications of MS. Seizure disorders in MS are uncommon. Tremor at rest is a feature of Parkinson disease.

**52** D This shifting spectrum of clinical findings over time in a young adult suggests the diagnosis of multiple sclerosis. The plaques of demyelination that give rise to the differing symptoms can be found in various locations, but they most often occur in periventricular white matter. CSF immunoglobulins are increased, and most patients show oligoclonal bands of IgG. Neurosyphilis, a form of tertiary syphilis, is rare at this age and probably would not cause localizing signs that change. Cryptococcal meningitis would manifest more acutely with meningeal signs. Primary or malignant brain tumors are uncommon at this age and would not have such a long course without serious sequelae. Xanthochromia from hemorrhage would suggest a more acute problem.
A The rapid correction of hyponatremia is a common antecedent to central pontine myelinolysis, with demyelination in the basis pontis, with the round area of decreased signal intensity shown in the figure. Extrapontine myelinolysis may also occur. A history of alcohol abuse increases the risk. Cerebral edema sufficient to produce herniation would cause papilledema, not noted in this case. Intracranial hemorrhages would not result from electrolyte and fluid disturbances. A subacute degeneration of the spinal cord occurs slowly as a consequence of vitamin $B_2$ (cobalamin) deficiency. Wernicke-Korsakoff syndrome is now a rare accompaniment to chronic alcoholism that affects mammillary bodies and periaqueductal gray matter.

B Acute disseminated encephalomyelitis is an immunologically mediated demyelinating disease that has some features similar to multiple sclerosis, but may occur in younger patients, have an abrupt onset, and be rapidly fatal. Most cases are preceded by an infection. The remaining choices are unlikely to have bilateral optic neuritis. Abscesses are likely to be bacterial in origin. Emboli are uncommon in children, and likely to produce infarctions. Metastases are unlikely in children. Thiamine deficiency may lead to Wernicke disease, which has hemorrhagic lesions and is unre- lated to infection. Vasculitis may produce focal lesions but is uncommon in children.

D The figure shows a neuritic plaque with a rim of dys- trophic neurites surrounding an amyloid core consistent with Alzheimer disease (AD), the most common form of progressive dementia. AD is marked by increased numbers of microscopic neuritic plaques and neurofibrillary tangles compared to controls for age. Choreiform movements suggest Huntington disease. Gait disturbances occur in Parkinson disease. Seizures are associated with many lesions, but often a pathologic finding is not discernible. Symmetric muscular weakness suggests amyotrophic lateral sclerosis.

E The clinical history of dementia and the presence of numerous neuritic plaques and amyloid deposition in blood vessel walls are characteristic of Alzheimer disease (AD). The e4 allele of the $ApoE4$ gene increases the risk of developing AD by unknown mechanisms. Expansion of CAG repeats on chromosome 4p16 causes Huntington disease. There is no association between HLA genes and AD. Increased repeats in the $FMR1$ gene occur with fragile X syndrome. Mutant prion genes give rise to spongiform encephalopathies, such as Creutzfeldt-Jakob disease.

F Pick disease is the best known form of frontotemporal lobar degeneration with tau-containing inclusions (FTLD-tau) and has clinical features similar to the features of Alzheimer disease (AD), but initially FTLD-tau causes less memory loss and more behavioral changes. The “knifelike” gyral atrophy of frontal and temporal lobes and relative sparing of parietal and occipital lobes are characteristic of Pick disease. Pick bodies with tau protein are seen in remaining neurons, but the neuritic plaques and neurofibrillary tangles seen in AD are not increased. Huntington disease affects mainly the caudate nuclei and basal ganglia; onset occurs in middle age, and choreiform movements are common. Leigh disease is a mitochondrial encephalomyopathy that can cause muscular weakness and neurologic deterioration beginning at a young age. Multiple system atrophy (MSA) has features that overlap those of striatonigral degeneration, olivopontocerebel- lar atrophy, and Shy-Drager syndrome; most patients with MSA exhibit symptoms similar to those of Parkinson disease. MSA is characterized microscopically by the appearance of glial cytoplasmic inclusions. In Parkinson disease, loss of pig- mented neurons in the substantia nigra leads to movement problems. Vascular dementia, or multi-infarct dementia, can have clinical features that mimic those of AD, but there are multiple small infarcts that collectively produce dementia, and the neurologic decline occurs in a stepwise fashion.

G Dementia with Lewy bodies (DLB) combines clinical features of Alzheimer disease (AD) and idiopathic Parkinson disease. Mutations in the gene for $\alpha$-synuclein have been linked to idiopathic Parkinson disease, and Lewy bodies can be found in the substantia nigra neurons, but the clinical dementia and cortical Lewy bodies point to DLB. Amyloid precursor protein (APP) is encoded by a gene on chromosome 21 (perhaps explaining early AD in trisomy 21) and is processed to form the $A\beta$ amyloid of neuritic plaques in AD. The e4 allele of apolipoprotein E can bind $A\beta$ and increase the risk of AD. Huntington in the protein product of the $HD$ gene in Huntington disease. Presenilin 1 and 2 mutations can increase production of $A\beta$ and increase the risk of early-onset AD. Tau protein is found in neurofibrillary tangles of AD and in Pick bodies of Pick disease.

H Loss of pigmented dopaminergic neurons in the substantia nigra of the midbrain is most characteristic of Parkinson disease. Pill-rolling tremors at rest are typical of this disorder. A variety of genetic abnormalities have been associated with forms of Parkinson disease, including $\alpha$-synuclein, parkin, DJ-1, and $PINK1$ gene mutations and mitochondrial dysfunction. Axatia suggests a disruption in the motor control pathways, such as the cerebellum, or proprioception, from dorsal spinal cord columns. Choreiform movements suggest Huntington disease, which affects the caudate, not the substantia nigra. Short-term memory problems suggest hippocampal lesions. Symmetric weakness suggests a motor neuron disease.

I Huntington disease is a progressive degenerative disorder that affects basal ganglia, including the putamen and caudate nucleus through loss of spiny striatal neurons that normally dampen motor activity. This disease, inherited
in an autosomal dominant pattern, is caused by an abnormal expansion of the trinucleotide CAG in the huntingtin gene on chromosome 4, which normally has 6 to 35 CAG copies. There may be 40 to 55 copies in patients with typical Huntington disease and 70 repeats in patients with earlier onset of the disease. A “premutation” with 36 to 39 repeats may have reduced penetrance. These patients have atrophy, with loss of neurons and gliosis starting in the caudate, as well as putamen and globus pallidus. About 10% of cases of Creutzfeldt-Jakob disease are genetically determined, with the inheritance of an abnormal prion protein that leads to spongiform encephalopathy in later adult life. Tay-Sachs disease of infancy and childhood is caused by a deficiency of hexosaminidase A. Trisomy 21 results in mental retardation present at birth. Mutations in presenilin genes cause familial Alzheimer disease (AD). The risk of AD increases with increased levels of apolipoprotein E.

61 D She had Friedreich ataxia, an autosomal recessive progressive illness that most often has an onset in the first decade of life. The frataxin gene encodes for a protein involved in iron regulation in cells, and a GAA trinucleotide repeat expansion results in decreased protein and decreased mitochondrial oxidative phosphorylation. The other options listed have no cardiac involvement. Mutations of the huntingtin gene are seen with Huntington disease marked by choreoathetosis beginning in young to middle-aged adults. Increased tandem repeats in the FMR1 gene account for cases of fragile X syndrome characterized by mental retardation. The dystrophila myotonia–protein kinase gene is abnormal in cases of myotonic dystrophy with muscular weakness and dementia.

62 B The progressive and symmetric nature of this patient’s disease is a classic feature of amyotrophic lateral sclerosis (ALS). The muscles show a denervation type of grouped atrophy from loss of lower motor neurons in anterior horns of the spinal cord. The “bulbar” form of ALS affects mainly cranial nerve nuclei and has a more aggressive course. Cortical upper motor neurons may also be lost, but mental function is preserved in ALS. Ependymal cells line ventricles, which are normal in ALS. Microglial cells have a macrophage-like function and may be involved in demyelinating plaques of multiple sclerosis that can produce various motor signs and symptoms over time, but symmetry is not a feature of this disease. Oligodendrocytes provide myelin to axons in the CNS. Parkinson disease with loss of substantia nigra pigmented neurons is characterized by rigidity and involuntary movements, not by muscular weakness. Huntington disease with loss of spiny neurons in the caudate nucleus causes abnormal movements, not weakness, and there can be associated dementia over time.

63 B In children, an inherited form of CNS disease that accounts for a progressively worsening course should be suspected. The leukodystrophies are various inborn errors of metabolism involving lysosomal (arylsulfatase A in this case) or peroxisomal enzymes that affect white matter extensively and cause myelin loss and abnormal accumulations of myelin from failure of generation, maintenance, or catabolism of myelin. There are no discrete plaques of demyelination, however, in contrast to multiple sclerosis. Sparing of subcortical myelin (U fiber) is often seen in leukodystrophies. Acute disseminated encephalomyelitis is a postinfectious process with abrupt onset. Progressive multifocal leukoencephalopathy is an infectious lesion that occurs in immunocompromised adults. Tay-Sachs disease affects infants.

64 E Wernicke disease results from a deficiency of vitamin B1 (thiamine). Wernicke disease is uncommon in individuals who have a varied diet, but individuals with a history of chronic alcoholism may not. Capillary proliferation, hemorrhage, necrosis, and hemosiderin deposition are often found in the mammillary bodies and the periaqueductal gray matter, resulting in paralysis of the extraocular muscles. If memory problems with confabulation are observed, the thalamus is involved and the diagnosis is Wernicke-Korsakoff syndrome. Though deficiency of either B12 or folate may produce macrocytic anemia, subacute combined degeneration of the spinal cord is seen with vitamin B12 (cobalamin) deficiency, but folate deficiency does not produce CNS signs. Dementia may be present in individuals with niacin deficiency. A deficiency of pyridoxine may result in a peripheral neuropathy.

65 A Liver failure with hepatic encephalopathy can occur in severe liver disease from various causes, including commonly chronic alcoholism. Hyperammonemia is a feature of liver failure. Carbon monoxide poisoning can produce obtundation and coma. Severe hypoglycemia can damage neurons in the hippocampus and neocortex. An elevated hemoglobin A1c level suggests a diagnosis of diabetes mellitus, and diabetic patients are most prone to peripheral neuropathies and autonomic neuropathies. Hyponatremia from diabetics insipidus may result in obtundation.

66 B He may initially have had an infiltrating astrocytoma, which is the most common primary brain neoplasm in adults, typically arising in a cerebral hemisphere. Lower grade astrocytomas in adults may have a more indolent course. Some of these patients go on to develop a high-grade glioma known as secondary glioblastoma. Most glioblastomas arise de novo (primary glioblastoma) and have MDM2 mutations. Regardless of origin, glioblastomas are aggressive and have a poor prognosis. Diffuse large B-cell lymphoma is the most common type of primary CNS lymphoma (without evidence for disease elsewhere); some arise in immunocompromised patients. Hemangioblastomas are uncommon neoplasms arising in the cerebellum, often with von Hippel-Lindau disease, and associated with polycythemia. Medulloblastomas and pilocytic astrocytomas are usually childhood brain tumors arising in the posterior fossa.
Primary malignant neoplasms of the brain in children most often occur in the posterior fossa. The two most common neoplasms at this site are pilocytic (cystic cerebellar) astrocytoma and medulloblastoma. Pilocytic astrocytoma is slow growing and has a better overall prognosis than glial neoplasms in adults. Both may enlarge and block CSF flow, causing hydrocephalus. Ependymomas can occur in childhood, but are most likely to arise in the fourth ventricle. A hemangioblastoma is a rare cystic mass in adults, typically arising in the cerebellum, and may be associated with polycythemia. Medulloblastomas often occur in the cerebellar midline, are composed of primitive round blue cells, and have a poor prognosis. Meningiomas occur in adults; they are circumscribed, solid mass lesions adjacent to the dura, and may be multiple in neurofibromatosis. A schwannoma typically arises in cranial nerve VIII in adults.

Oligodendrogiomas tend to have a better prognosis than most other glial neoplasms. Pilocytic astrocytomas with *BRAF* mutations tend to be less circumscribed. Diffuse large B-cell lymphomas can occur in association with AIDS; they are negative for glial fibillary acidic protein (GFAP), but positive for CD19 and CD20. Medulloblastomas are posterior fossa tumors that occur in children, and those with poor prognosis often have c-MYC amplification. Classic glioblastomas are highly aggressive, infiltrative gliomas, often with *EGFR* alterations.

The myxopapillary variant of ependymoma is more common in adults than in children. Ependymomas that arise in the ventricles (usually the fourth ventricle) are more common in the first 2 decades of life. Choroid plexus papillomas are rare tumors that arise in the cerebral ventricles, consistent with medulloblastoma. Pilocytic astrocytomas, are most often found within a cerebral hemisphere. Schwannomas most often occur at the cerebellopontine angle in the eighth cranial nerve.

The MRI shows a 4-cm mass in the cerebellar vermis, along with dilation of the third and lateral cerebral ventricles, consistent with medulloblastoma. Most intracranial neoplasms in children are located in the posterior fossa. The medulloblastoma, one of the “blue cell tumors” of childhood, arises in the midline, and the cells can seed into the CSF. A glioblastoma is also a high-grade malignancy that could seed the CSF, but this neoplasm occurs in adults in a cerebral hemisphere. Schwannomas most often involve the eighth cranial nerve, are benign neoplasms, and do not seed the CSF. Ependymomas can occur in children, but they typically arise in the ventricles. Metastatic lesions may involve the gray-white junction, but are uncommon in children.

The myxopapillary variant of ependymoma is more common in adults than in children. Ependymomas that arise in the ventricles (usually the fourth ventricle) are more common in the first 2 decades of life. Choroid plexus papillomas are rare tumors that arise in the cerebral ventricles. Meningiomas most often arise in the cranial cavity and are rare complications of disseminated tuberculosis and aspergillosis. Tuberculomas are granulomas large enough to produce a mass effect and are rare complications of disseminated tuberculosis and often appear at the base of the brain.

Multiple discrete neoplasms in the CNS found at the gray-white junction are more likely to be metastases than a...
primary brain tumor. Tumor cells may reach the brain in the
form of emboli through the cerebral arterial circulation. Most
embolic events occur at the gray-white junction, where nar-
orring and acute branching of the vessels tend to trap emboli.
The distribution of the middle cerebral artery, which receives
the most blood, is the most likely location. Metastases from
malignant melanomas are often widely disseminated, with
multiple mass lesions in organ sites of involvement. Glioblas-
tomas are large, malignant primary glial neoplasms; they are
invasive, but do not typically appear as multiple small lesions.
A hemangioblastoma is a rare cystic mass in adults, typically
arising in the cerebellum, and may be associated with poly-
cythemia. Meningiomas are circumscribed, solid mass lesions
adjacent to the dura; they may be multiple in patients with
neurofibromatosis. Cerebral malignant lymphomas are rare
except in HIV-positive patients. Oligodendrogliomas are soli-
dary, circumscribed mass lesions that occur in the cerebral
hemispheres of adults.

75 E The multiple pigmented skin lesions and optic nerve
glioma strongly suggest the diagnosis of neurofibromatosis
type 1. Patients with this condition, which has an autosomal
dominant inheritance, have multiple large café-au-lait spots
on the skin, and there is a propensity for development of
multiple nerve sheath tumors (schwannomas or neurofibro-
mas). CNS gliomas also may occur. The nerve sheath tumors
may become malignant and metastasize, most commonly to
the lungs. Of the other neoplasms listed, only meningioma is
associated with neurofibromatosis type 1, but it occurs
intracranially.

76 A A schwannoma in this location also is known as a cer-
ebellopontine angle tumor. Schwannomas in this location arise
from cranial nerve VIII; they are also called acoustic neuro-
mas. Most schwannomas act as benign, slow-growing tumors
that can be completely resected. Having bilateral acoustic
schwannomas is virtually pathognomonic for neurofibro-
matosis type 2. Other familial tumor syndromes involving
brain include Li Fraumeni syndrome with p53 mutations
and medulloblastomas, Gorlin syndrome with PTCH muta-
tions and medulloblastomas, tuberous sclerosis with TSCI
mutations and cortical hamartomas, and von Hippel–Lindau
disease with VHL mutations and hemangioblastomas.

77 E Tuberous sclerosis is one of the phakomatoses, a
group of rare inherited disorders in which hamartomas and
neoplasms develop throughout the body, along with cuta-
nceous abnormalities. Patients with tuberous sclerosis have
cortical tubers, which are hamartomas of neuronal and glial
tissue; other characteristic findings include renal angio-
myolipomas, renal cysts, subungual fibromas, and cardiac
rhabdomyomas. In Down syndrome (trisomy 21), patients
may develop acute leukemia, but not brain neoplasms, and
individuals who survive to middle age develop Alzheimer
disease. Krabbe disease is a leukodystrophy that results in
deficiency of galactocerebroside β-galactosidase and an onset
of neurologic deterioration in infancy. Neurofibromatosis
type 1 is characterized by deforming cutaneous and visceral
neurofibromas, cutaneous café-au-lait spots, and neurofibro-
sarcomas. In neurofibromatosis type 2, acoustic schwanno-
mas, meningiomas, gliomas, and ependymomas are present.
Von Hippel–Lindau disease is characterized by hemangiob-
lastomas in the cerebellum, retina, and spinal cord, and by
pheochromocytomas.

78 C Hemangioblastomas are part of the spectrum of
neoplasms seen in association with von Hippel–Lindau dis-
ease. The VHL gene encodes for a ubiquitin-ligase compo-
nent important in the degradation pathway of HIF, and this
increases vascular endothelial growth factor (VEGF) expres-
sion to drive vascular proliferation. Cushing syndrome
with increased corticosteroid production can be related to
ectopic ACTH production by a carcinoma. Cushing dis-
ease is caused by an ACTH-producing adenohypophyseal
adenoma. Hypercalcemia of malignancy often results from
parathormone-related peptide production by a carcinoma.
Syndrome of inappropriate antidiuretic hormone (SIADH)
may be caused by neuroendocrine tumors such as small cell
lung carcinomas. Carcinomas, particularly adenocarcinomas,
may produce hypercoagulability (Trousseau syndrome).
1. A 39-year-old man has increasing pain with left eye movement and swelling for the past week. On examination he has proptosis and ophthalmoplegia. There is edema and erythema of the eyelid with marked tenderness on palpation. A purulent nasal discharge is present. Which of the following orbital conditions is he most likely to have?
   - A. Cellulitis
   - B. Graves disease
   - C. Hemangioma
   - D. Non-Hodgkin lymphoma
   - E. Sarcoidosis

2. A 7-year-old child has had worsening performance in school for the past 4 months from decreased vision. Examination of the right eye shows diffuse punctate inflammation of the cornea and pannus extending as a growth of fibrovascular tissue from conjunctiva onto the cornea. Microscopic examination of a corneal scraping shows lymphocytes, plasma cells, neutrophils, and scattered corneal epithelial cells that have cytoplasmic inclusion bodies. Which of the following infectious agents is most likely to produce these findings?
   - A. Chlamydia trachomatis
   - B. Cytomegalovirus
   - C. Herpes simplex virus
   - D. Neisseria gonorrhoeae
   - E. Staphylococcus aureus

3. An 82-year-old man still surfs every week at Waikiki beach. He has noted clouding of vision on the right for the past year. On physical examination of the right eye there is a whitish irregular lesion on the conjunctiva extending onto the cornea. Which of the following is the most likely risk factor for his ocular lesion?
   - A. Chlamydia trachomatis infection
   - B. Hyperglycemia
   - C. Hypertension
   - D. Sunlight exposure
   - E. Vitamin A deficiency

4. A 27-year-old woman has had pain with cloudiness of vision in the right eye for the past 2 days. A similar episode occurred a year ago. On physical examination, there is no conjunctival erythema or vascular injection. Funduscopic examination shows no retinal lesions. A slit lamp examination with fluorescein dye shows a dendritic ulcer on the right cornea. Which of the following infectious agents has most likely produced these findings?
   - A. Chlamydia trachomatis
   - B. Cytomegalovirus
   - C. Herpes simplex virus
   - D. Neisseria gonorrhoeae
   - E. Staphylococcus aureus

5. A 26-year-old man has had severe visual impairment since birth. He is legally blind without his glasses, but his astigmatism in both eyes is so severe that he cannot get a correction better than 20/100 with eyeglasses. He tries rigid contact lenses and gets a correction of 20/40. Which of the following ocular conditions is he most likely to have?
   - A. Keratoconus
   - B. Pterygium
   - C. Stromal dystrophy
   - D. Trachoma
   - E. Vitamin A deficiency

6. A 34-year-old man has had decreasing vision for the past 3 years and now has severely impaired vision in both eyes. His brother is similarly affected. Both parents have normal vision. Ocular examination shows diffuse cloudiness of the anterior stroma with aggregates of gray-white opacities in the axial region of the corneal stroma. He undergoes bilateral corneal transplantation. The diseased corneas show basophilic deposits in the stroma that stain positively for keratan sulfate. What is the most likely diagnosis?
   - A. Cataract formation
   - B. Keratomalacia
   - C. Macular dystrophy
   - D. Pterygium
   - E. Trachoma
A 29-year-old woman has developed malaise with nausea over the past month. On physical examination, she has an erythematous rash on the cheeks of her face. Laboratory studies show a serum creatinine level of 3.3 mg/dL, urea nitrogen of 33 mg/dL, positive ANA of 1:2048, and positive anti-double-stranded DNA of 1:512. A renal biopsy specimen shows a proliferative glomerulonephritis. She receives long-term high-dose glucocorticoid therapy. What ocular complication is this patient most likely to develop?

A  Background retinopathy  
B  Cataracts  
C  Corneal stromal dystrophy  
D  Granulomatous uveitis  
E  Macular degeneration

An 82-year-old woman has had increasing visual problems that are worse in the right eye over the past 2 years. She is unable to see clearly when looking straight ahead because of cloudiness and opacification and has great difficulty reading printed material. Her peripheral vision is better. Her left eye is similarly affected and is shown in the figure. What pathologic process has most likely occurred in this woman?

A  Keratomalacia of the cornea  
B  Nuclear sclerosis of the lens  
C  Open-angle glaucoma  
D  Retinal macular degeneration  
E  Sympathetic ophthalmitis

A 77-year-old woman has experienced increasing pain accompanied by clouded vision in the right eye for the past 36 hours. She has worn corrective lenses for hyperopia for the past 70 years. On physical examination, there are no lesions of the cornea or crystalline lens. On funduscopic examination, there is excavation of the optic cup on the right. Which of the following pathologic processes is most likely to produce these findings?

A  Amyloid deposition within posterior chamber vitreous  
B  Crystalline lens dislocation into the anterior chamber  
C  Increased aqueous humor production by the ciliary body  
D  Resistance of aqueous humor outflow into Schlemm canal  
E  Shallow anterior chamber obstructing aqueous humor outflow  
F  Thromboembolism to the central retinal artery

A 48-year-old man was not using protective goggles while ripping plywood on his table saw, and he sustained a penetrating injury to the left eye. A wood splinter is removed. On funduscopic examination, there is a partial uveal prolapse, but he still has vision in the left eye. Three weeks later, he has loss of accommodation, photophobia, and blurred vision in the right eye. Choroidal infiltrates are now seen on funduscopic examination. What is the most likely diagnosis?

A  Aspergillus fumigatus infection  
B  Fuchs dystrophy  
C  Sarcoïdosis  
D  Sympathetic ophthalmitis  
E  Undiagnosed trauma

A 61-year-old woman has had decreasing visual acuity in the right eye for 6 months. She then experienced sudden loss of part of the vision in the left eye, which occurred “as though a shade had been pulled across” her field of view. On funduscopic examination, there is a dark uveal mass. The enucleated eye is shown in transverse section in the figure. What is the most likely diagnosis?

A  Granulomatous uveitis  
B  Malignant hypertension  
C  Melanoma  
D  Ocular trauma with hematoma  
E  Retinoblastoma  
F  Toxoplasmosis
12 A 68-year-old woman with a history of left ventricular congestive heart failure has had decreased visual acuity for the past 5 years. She has no ocular pain. Her intraocular pressure is normal. Findings on funduscopic examination include arteriolar narrowing, flame-shaped hemorrhages, cotton-wool spots, and hard, waxy exudates. Which of the following underlying diseases is she most likely to have?

A Advanced atherosclerosis  
B Cerebral edema  
C Diabetes mellitus  
D Hypertension  
E Retinitis pigmentosa

13 A 68-year-old woman has had chronic renal failure for 10 years and has been on hemodialysis. She had a myocardial infarction last year. She now has worsening vision bilaterally. The figure shows findings representative for her retina. Which of the following laboratory test findings in her blood is she most likely to have?

A Antinuclear antibody titer of 1:1024  
B Ferritin of 555 ng/mL  
C Hgb A1c of 9.6%  
D Renin activity of 9 ng/mL/hr  
E Total cholesterol 265 mg/dL

14 A 74-year-old man suddenly lost the upper half of the visual field in the right eye. Before this event, he had decreasing visual acuity in both eyes for the past 6 years. On physical examination, his height is 170 cm (5 feet 8 inches), and weight is 92.5 kg (body mass index 32). Laboratory studies show a fasting serum glucose level of 165 mg/dL. What underlying pathologic process is most likely to account for the sudden loss of vision in his right eye?

A Dendritic corneal ulcer  
B Macular degeneration  
C Retinitis pigmentosa  
D Traction retinal detachment  
E Uveal malignant melanoma

15 A 23-year-old primigravida with preeclampsia gives birth prematurely at 32 weeks’ gestation. The infant’s Apgar scores are 4 and 6 at 1 minute and 5 minutes, respectively. The infant has hyaline membrane disease and is intubated and administered positive pressure ventilation with 100% inspired oxygen. The infant survives and is discharged on the twenty-third day of life. Two months later, the mother notices that the infant does not always respond visually to her presence. The infant is examined and is found to be blind in nasal visual fields. What is the most likely diagnosis?

A Cataracts  
B Keratomalacia  
C Macular degeneration  
D Retinitis pigmentosa  
E Retrolental fibroplasia

16 A 15-year-old boy from Eastern Arabia has experienced episodes of chest, abdominal, and back pain as well as fatigue for the past 10 years. Physical examination is not remarkable. Laboratory investigations show hemoglobin, 8 g/dL; hematocrit, 24.7%; MCV, 95 μm³; and total serum bilirubin, 2.1 mg/dL. The serum AST, ALT, albumin, and total protein are normal. Prothrombin time and partial thromboplastin time are normal. Examination of a peripheral blood smear shows reticulocytosis and polychromasia with occasional deformed RBCs shaped like crescents. This patient is at increased risk for developing which of the following ocular complications?

A Cataract  
B Intraretinal hemorrhage  
C Keratoconus  
D Secondary angle-closure glaucoma  
E Uveitis

17 A 75-year-old woman with diabetes mellitus and congestive heart failure has sudden loss of vision in her left eye. Funduscopic examination shows a cherry-red appearance of the foveola, whereas the remaining retina appears pale. No abnormalities of the right eye are noted. Which of the following is the most likely cause for her sudden unilateral visual loss?

A Central retinal artery occlusion  
B Cytomegalovirus retinitis  
C Primary angle-closure glaucoma  
D Proliferative retinopathy  
E Tay-Sachs disease

18 A study is conducted of children 1 to 2 years of age who appeared healthy when born at term, but developed blindness and failure to meet developmental milestones for neural development. On funduscopic examination they have pale retinae with prominent red macular regions. A deficiency of which of the following enzymes is most likely to be found in these children?

A α-L-Iduronidase  
B Galactosylceramidase  
C Glucocerebrosidase  
D Hexosaminidase A  
E Lysosomal glucosidase
A pharmaceutical company is developing a product that would be useful to prevent age-related visual loss. A cohort of individuals 60 to 80 years old is followed for 5 years with periodic examinations of visual acuity, funduscopy, and fluorescein angiography. Some of these individuals develop progressive loss of vision characterized initially by diffuse deposits in Bruch membrane and by atrophy of retinal pigment epithelium. Later, a subset of these patients has a further decline in vision because of development of choroidal neovascularization. An antagonist to which of the following molecules is most likely to be useful in reducing vision loss in this subset of patients?

A. Epidermal growth factor (EGF)
B. Insulin-like growth factor (IGF)
C. Platelet-derived growth factor (PDGF)
D. Transforming growth factor beta (TGF-β)
E. Vascular endothelial growth factor (VEGF)

A 78-year-old man with a 30 pack-year history of smoking has had decreasing vision, mainly in a central pattern, for the past 3 years. He has no ocular pain. Intraocular pressures are normal. There are no abnormalities of the cornea or crystalline lens. On funduscopic examination, the retinal pigment epithelium appears atrophic, and deposits are seen in the Bruch membrane. What is the most likely diagnosis?

A. Macular degeneration
B. Proliferative retinopathy
C. Retinal detachment
D. Retinitis pigmentosa
E. Retrolental fibroplasia

A 33-year-old man has had increasing difficulty seeing at night, but has no problems with his vision during the day. Three years later, his daytime visual acuity also is decreasing. Funduscopic examination now shows a branching reticulated pattern to the retina. The optic disc appears pale and waxy, and there is attenuation of retinal blood vessels. These findings are most characteristic of what condition?

A. Arteriosclerotic retinopathy
B. Hypertensive retinopathy
C. Macular degeneration
D. Proliferative retinopathy
E. Retinitis pigmentosa

A 22-year-old woman gives birth at term to a boy after an uncomplicated pregnancy. No abnormalities are noted on a newborn physical examination. During infancy, a well-infant checkup shows leukocoria in the right eye. The eye is enucleated; the microscopic appearance of an intraocular mass is shown in the figure. What is the most likely diagnosis?

A. Glioma
B. Melanoma
C. Medulloblastoma
D. Retinoblastoma
E. Squamous cell carcinoma

A 70-year-old man with atrial fibrillation and poorly controlled hypertension is recovering from right hip replacement surgery. He has had a worsening headache for the past day. He is receiving anticoagulant therapy. On physical examination, he is afebrile with blood pressure 170/110 mm Hg. On funduscopic examination, there is papilledema of the left eye. What is the most likely cause of the papilledema?

A. Alzheimer disease
B. Glaucoma
C. Macular degeneration
D. Optic neuritis
E. Intracranial hemorrhage
F. Schwannoma

A 72-year-old woman has had decreasing vision for the past year. She now has increasing headaches. She has worn glasses since childhood because of myopia. Funduscopic examination shows deepening of the optic cup with excavation. The surrounding retina appears normal. Screening of which of the following would most likely have detected the disease that led to these findings?

A. Blood pressure
B. Homocystinuria
C. Intraocular pressure
D. Serum glucose
E. Visual acuity
ANSWERS

1 A Orbital cellulitis can be due to spread of infection from adjacent paranasal sinuses, such as ethmoid air cells. Organisms may include *Staphylococcus aureus*, *Streptococcus pyogenes*, *Streptococcus pneumoniae*, and *Haemophilus influenzae*. In diabetic ketoacidosis, mucormycosis should be considered. Idiopathic orbital inflammation (inflammatory pseudotumor) may present similarly, but is noninfectious. Graves disease is likely to present with bilateral proptosis, but without inflammatory changes, because accumulation of extracellular matrix proteins is the etiology. Hemangioma in children and lymphoma in adults are mass lesions of the orbit, likely without inflammatory changes. Sarcoidosis produces granulomatous inflammation without acute changes.  

PBD9 1320–1321 PBD8 1347

2 A Trachoma is a major cause of blindness worldwide. The initial inflammation from infection is followed by progressive conjunctival scarring with eyelid involvement, so that the eyelashes turn inward (trichiasis) to produce scarring of the conjunctiva and cornea. In children, cytomegalovirus (CMV), one of the herpesviruses, is a rare cause of ocular infection. CMV produces prominent intranuclear inclusions. Herpetic keratitis can result in ulceration and scarring; herpesviruses have intranuclear inclusions. Congenital rubella, which is now a rare disease because of immunization, produces a retinopathy. Congenital infections with *Treponema pallidum* result in an interstitial keratitis.  

PBD9 1323 BP9 311 PBD8 1349 BP8 323

3 D A pterygium can extend onto the cornea but does not cross the pupillary axis; it may induce mild astigmatism to affect vision. A pinguecula does not impinge upon the cornea, but it does affect distribution of the tear film to produce focal dehydration and corneal depression. Both of these lesions are composed of fibrovascular connective tissue, and ultraviolet light is the driving force. Trachoma leads to conjunctival scarring that is not so focal. Diabetes mellitus with hyperglycemia is more likely to affect the crystalline lens or the retina. Hypertensive retinopathy may occur. Vitamin A deficiency can lead to keratomalacia and not a localized scar.  

PBD9 1323 PBD8 1349

4 C The most common cause of corneal dendritic ulcers is herpes simplex virus infection. Such ulcers can perforate through to the globe, which is a medical emergency. Some chronic herpetic corneal infections cause localized opacity. Lymphocytes and plasma cells, and viral inclusions in the corneal epithelial cells, are present. Trachoma, an infection with *Chlamydia trachomatis*, is seen most often in children and produces inflammation leading to extensive corneal and conjunctival scarring. Cytomegalovirus infection rarely causes corneal lesions. It can cause retinitis in congenital infections and in immunocompromised adults. *Neisseria gonorrhoeae* infection can occur in sexually active persons, and it can be transmitted to neonates at birth, so prophylactic silver nitrate eye drops at birth are employed to prevent ophthalmia neonatorum. *Staphylococcus aureus* is a common infection at many tissue sites, and can produce conjunctivitis, which was not evident in this case.  

PBD8 1351

5 A Keratoconus patients are typically not candidates for laser in situ keratomileusis (LASIK) because the marked corneal thinning with breaks in the Bowman layer gives their corneas a conical shape. Penetrating keratoplasty may be performed. Corneal transplantation can also be considered as treatment for keratoconus. A pterygium is a localized area of conjunctival opacification from degenerative changes; unlike a pinguecula, it extends onto the cornea and may interfere with vision. Stromal dystrophies are uncommon inherited conditions with corneal clouding from deposition of mucopolysaccharides. Trachoma occurs from infection with *Chlamydia trachomatis* and may lead to corneal scarring with blindness. Vitamin A deficiency can lead to keratomalacia and eventual blindness if the deficiency is not treated.  

PBD8 1352

6 C Inherited corneal stromal dystrophy has several forms, and most are autosomal dominant. However, the most severe form is macular dystrophy, which has an autosomal recessive form of inheritance. It is essentially a form of mucopolysaccharidosis confined to the cornea in which keratan sulfate is deposited. Cataracts are seen most often in elderly individuals and result from opacifications of the crystalline lens. Keratomalacia can be a consequence of vitamin A deficiency. A pterygium is a localized area of basophilic degeneration of conjunctival epithelium that extends onto the cornea. Trachoma is caused by infection with *Chlamydia trachomatis*.  

PBD9 1326–1327 PBD8 1352–1353

7 B Cataracts of the crystalline lens are an important complication of systemic therapy with glucocorticoids. This patient has systemic lupus erythematous with lupus nephritis. Cataracts can be caused by aging, diabetes mellitus, glaucoma, ultraviolet light, or irradiation. Retinopathy is most often a feature of diabetes mellitus or hypertension. Stromal dystrophies are inherited conditions that affect the cornea. Granulomatous uveitis occurs with sarcoidosis. Macular degeneration is most often a disease of central vision in the elderly.  

PBD9 1327 PBD8 1353

8 B Nuclear sclerosis of the lens leads to cataracts in elderly individuals, typified by the lens opacification shown in the figure. This change causes opacification owing to compression of the lens fibers in the central (nuclear) portion of the lens. Keratomalacia can produce corneal scarring with opacification, but not in a central distribution pattern.
Glaucoma results from increased intraocular pressure and damages the optic nerve, and it is not characterized by loss of central vision, but more likely peripheral vision. Macular degeneration also occurs in elderly individuals and results in loss of central vision, but it does not cause cloudiness. Sympathetic ophthalmia in one eye occurs after trauma to the other eye.

9 E In some older individuals with hyperopia, the iris is displaced forward to narrow the angle at the anterior chamber, obstructing flow of aqueous humor, so-called primary angle-closure glaucoma, which manifests with acute pain. Increased pressure on the optic nerve causes excavation and produces progressive visual loss. Mutations in the myocilin (MYOC) and the optineurin (OPTN) genes may account for some cases of primary open-angle glaucoma. Amyloid deposition is quite rare and does not increase intraocular pressure. Crystalline lens dislocation can occur with trauma and with Marfan syndrome. Increased production of aqueous humor is a rare cause of glaucoma. Increased resistance to outflow of aqueous into Schlemm canal is typical of primary open-angle glaucoma. Increased resistance to outflow of aqueous humor, obstructing flow of aqueous, so-called primary angle-closure glaucoma, which occurs in individuals with myopia. Thrombosis or embolism to the central retinal artery can lead to occlusion with edema, pallor, and a cherry-red spot in the macula. Ischemia of the choroid, the iris, or the ciliary body. They are often pigmented. Dendritic corneal ulceration suggests infection with herpes simplex virus.

10 D Sympathetic ophthalmia is an unusual, but devastating, form of uveitis that can complicate penetrating ocular trauma. It results from the release of a sequestered antigen from one eye that causes an immune response with inflammatory reaction in the opposite eye. To prevent this complication, the traumatized eye must be removed before inflammation begins in the opposite eye. Trauma alone cannot produce this spectrum of findings. Louis Braille, the inventor of the Braille alphabet for tactile reading of text, had an eye injury that led to sympathetic ophthalmia. Infection with Aspergillus is unlikely to become disseminated in an immunocompetent individual, and is not typically associated with traumatic eye lesions. Sarcoidosis can affect the eye, but lesions in the choroid are uncommon. Fuchs dystrophy is an inherited condition that affects the corneal endothelium.

11 C The pigmented uveal mass is a melanoma causing retinal detachment, because the mass lifted off the overlying retina. After skin, the eye is the most common site for primary malignant melanoma. This is the most common intraocular malignancy in adults. Uveal melanomas can involve the choroid, the iris, or the ciliary body. They are often pigmented. In addition to causing retinal detachment, as in this case, they may cause choroidal hemorrhage or macular edema. Granulomatous uveitis can occur from sarcoidosis, but the inflammation does not produce a large mass lesion. Hypertensive retinopathy can produce small hemorrhages.

12 D Hypertensive retinopathy results from long-standing hypertension, with progressive changes that begin with generalized narrowing of the arterioles and proceed to the changes seen in this case. The pressure load from systemic hypertension causes hypertrophy and failure of the left side of the heart. Arteriosclerotic retinopathy causes vascular changes, including arteriovenous nicking and hyaline arteriolar sclerosis with "copper wire" and "silver wire" arterioles. Cerebral edema may result in papilledema. Various findings are associated with diabetic retinopathy, including capillary microaneurysms, cotton-wool spots, arteriolar hyalinization, and more severe changes of proliferative retinopathy with neovascularization. Retinitis pigmentosa describes a variety of abnormalities that arise as an inherited condition that may begin later in life (but usually earlier) and produce a waxy pallor of the optic disc.

13 C Proliferative retinopathy with neovascularization from long-standing diabetes mellitus is shown in the figure. Several other ocular changes also can occur with diabetes mellitus, including hemorrhages, arteriolar hyalinization, cotton-wool spots, fibroplasia. This patient's renal failure and cardiovascular disease are typical of diabetes mellitus. This funduscopic finding is not seen with autoimmune diseases, hemochromatosis, hypertension, or hyperlipidemia.

14 D The clinical features described suggest retinal detachment, which occurs in a late stage of proliferative retinopathy associated with diabetes mellitus. The neovascularization results in a membrane with fibrosis that increases traction on the retina, leading to sudden detachment. Dendritic corneal ulceration suggests infection with herpes simplex virus. Macular degeneration is a common cause of decreased vision in elderly individuals, but not of retinal detachment. Retinitis pigmentosa is an inherited, degenerative condition that is not related to diabetes mellitus. Uveal melanomas may cause retinal detachment, but they are not a feature of diabetes mellitus.

15 E Retrolental fibroplasia is a complication of prematurity birth that results from oxygen toxicity to the immature retinal vasculature, leading to neovascularization of the retina with growth into the vitreous. The incompletely vascularized lateral aspects of the retina are most severely affected by ischemia and subsequent up-regulation of vascular
endothelial growth factor (VEGF). In some cases, scarring continues and causes retinal detachment. Cataracts also are seen in older individuals. Keratomalacia is a feature of vitamin A deficiency that develops over a longer period. Macular degeneration is a disease of elderly individuals. Retinitis pigmentosa can be inherited in various patterns and has a variable onset from childhood to through older age.

16 B In sickle cell anemia the sickled RBCs can occlude the retinal microvasculature as oxygen tension decreases and the RBCs assume a sickle shape. These vascular occlusions can cause preretinal, intraretinal, and subretinal hemorrhages. Organization of preretinal hemorrhages can cause retinal traction and detachment. Cataracts are most commonly age-related, but may be secondary to systemic diseases such as galactosemia, diabetes mellitus, and Wilson disease. Keratoconus is characterized by progressive thinning of the cornea without any inflammation, which leads to an abnormal shape that is more conical than spherical, giving rise to severe astigmatism. This form of corneal degeneration can occur sporadically or in association with a systemic disease, such as Marfan syndrome. Secondary angle-closure glaucoma is caused by inflammation of the uvea and consequent formation of a neovascular membrane that blocks the trabecular meshwork. Uveitis may occur locally or be part of systemic diseases, such as sarcoidosis.

17 A Thromboembolization from the diseased heart to the central retinal artery causes a diffuse retinal infarct that obscures the underlying vascular choroid, except where the retina is thinner in the fovea, so that it appears red compared to the surrounding pale retina. The other listed choices are typically bilateral processes. Cytomegalovirus retinitis has edematous and hemorrhagic lesions; it is most often seen in immunocompromised patients. Primary angle-closure glaucoma typically occurs in hyperopic eyes, and some cases may be acute, with eye pain from elevated intraocular pressure. Proliferative retinopathy can occur with diabetes mellitus, but the major change is retinal neovascularization. Tay-Sachs disease seen in infancy and early childhood produces a cherry-red spot, because the fovea in the center of the macula is relatively spared; it contains few ganglion cells that contain the storage product.

18 D Tay-Sachs disease is an autosomal recessive inborn error of metabolism due to deficiency of hexosaminidase A, one of the GM_1 gangliosidoses, leading to lysosomal storage of intermediate metabolites in retinal ganglion cells and neocortical neurons. The cherry-red spot is the least affected area where the thinner foveal area allows the vascularized choroid to appear more prominent in the pale surrounding affected retina. The remaining choices represent autosomal recessive conditions that are unlikely to affect the eye. Mucopolysaccharidosis type I (Hurler syndrome) with deficiency of α-1-iduronidase affects mainly connective tissues. Galactosylceramidase deficiency leads to Krabbe disease, a leukodystrophy affecting cerebral white matter. Gaucher disease from glucocerebrosidase deficiency is most often nonneuropathic. Lysosomal glucosidase deficiency leads to Pompe disease.

19 E Age-related macular degeneration (AMD) is the leading cause of visual loss in the Western world. Its advanced stages (exudative AMD, or the “wet” form with choroidal neovascularization) are characterized by extensive choroidal neovascularization that is driven by the local production of VEGF. Clinical trials indicate that anti-VEGF agents reduce neovascularization and visual loss. None of the other listed factors significantly affects angiogenesis.

20 A Macular degeneration is most often an age-related condition (AMD) and is the most common cause of decreased vision in the elderly. An absence of retinal vessels in the center of the macula may contribute to this disease because the retina has high metabolic demands. The disease may result in fibrous metaplasia and scarring of the macular region, causing permanent loss of central vision. A majority of AMD cases may be related to inherited mutations in the complement factor H (CFH) gene, made worse through environmental exposures, such as cigarette smoke. Proliferative retinopathy is characterized by neovascularization of the retina. Retinal detachment may be a complication of diabetic proliferative retinopathy, but not macular degeneration. Retinitis pigmentosa is an inherited disorder that produces a characteristic waxy pallor of the optic disc. Retrolental fibroplasia is a complication of high-dose oxygen therapy for neonates (often premature).

21 E Retinitis pigmentosa describes a range of retinal abnormalities that can be inherited in various patterns, and progression of the disease is variable. Night blindness caused by loss of rod photoreceptors is an early symptom. Later, the cone photoreceptors also begin to degenerate, producing blindness. Vascular changes are seen with hypertensive and arteriosclerotic retinopathies, but these tend to occur in older adults, with no difference in effects on day or night vision. Macular degeneration is seen in elderly individuals and affects central vision first. Neovascularization is a feature of diabetic proliferative retinopathy.

22 B This infant has a sporadic form of retinoblastoma. Both mutations probably arose in the retinoblasts. The infant did not inherit susceptibility to develop retinoblastoma because both copies of the RB gene are normal in unaffected somatic cells (fibroblasts). If he had inherited one copy of the mutant (or deleted) RB gene, all the cells in the body would
have only one normal copy of the RB gene. His siblings are at no increased risk of developing retinoblastoma, and he is at no increased risk of developing osteosarcoma. For similar reasons, risk of developing a retinoblastoma in the left eye also is no greater than that of the general population.

23 D Retinoblastoma is the most common malignant ocular neoplasm in children. Histologically, there is clustering of small blue cuboidal or short columnar cells around a central lumen. These clusters are sometimes called Flexner-Wintersteiner rosettes, as shown in the figure. This tumor can spread to the orbit or along the optic nerve. Most cases occurring in children are familial, with an inherited mutated RB gene. Gliomas may affect the optic nerve in a child, but the microscopic pattern does not include the rosettes shown. Melanomas of the eye are seen in adults and have spindle or polygonal cell patterns, and are often pigmented. Medulloblastomas are cerebellar malignancies in children. Retinoblastomas, medulloblastomas, and neuroblastomas all are forms of small round blue cell tumors seen in children. Adenocarcinomas and squamous cell carcinomas are uncommon neoplasms in children and unlikely to be intraocular.

24 E Papilledema results from increased intracranial pressure produced by cerebral edema, intracranial hemorrhage, or rapidly expanding masses. This patient is most likely to have a basal ganglia hemorrhage based upon the risk factor of hypertension. A hemorrhagic stroke from embolization is also a possibility, with the history of atrial fibrillation. Patients receiving anticoagulation, which could be given following major surgery, are also at risk for intracranial hemorrhages. The blood collecting in the subdural space leads to increased intracranial pressure. Cerebral amyloid angiopathy can occur in association with Alzheimer disease and can lead to intracranial hemorrhage, but that is not the risk in this vignette. Glaucoma, with increased intraocular pressure, produces optic cup excavation (the opposite of papilledema). Macular degeneration affects the fovea more severely. Optic neuritis leads to decreased visual acuity and is most often a complication of multiple sclerosis. Schwannomas arise in peripheral nerves, including cranial nerves, but the optic nerve is really a tract, an extension of the forebrain diencephalon. Thus a glioma could arise in the optic tract, but not a schwannoma.

25 C The increased intraocular pressure with glaucoma is believed to cause the loss of nerve fibers, resulting in a characteristic cupped excavation of the optic disc. Because there are no obvious early signs or symptoms, screening for increased pressure is important for detection. Glaucoma has several causes; various medications are used to treat the disease. Hypertension can produce a retinopathy, but not increased intraocular pressure. Homocystinuria is a rare condition that also increases the risk of atherosclerosis. Hyperglycemia suggests a diagnosis of diabetes mellitus, which can increase the risk of glaucoma. Visual acuity may be maintained while glaucoma is progressing, and loss of acuity alone does not identify the cause.
A 33-year-old man has experienced onset of chest pain, diaphoresis, and dyspnea over the past 6 hours. In the emergency department, he has a serum troponin I level of 6 ng/mL. Additional laboratory findings include hematocrit of 41%, hemoglobin A1c of 4.2%, total serum cholesterol of 482 mg/dL, and serum triglyceride of 160 mg/dL. Emergent coronary angiography shows 65% stenosis of the left circumflex artery and 70% stenosis of the left anterior descending artery. He undergoes angioplasty with stent placement. He experiences a series of transient ischemic attacks 1 year later. He also has pain in the lower extremities when walking more than 300 m. He is given a drug that inhibits hepatic HMG-CoA reductase. The pathogenesis of his underlying disease is most likely related to a reduction in which of the following cellular receptors?

A  Acetylcholine

B  Glucose

C  Hepcidin

D  Insulin

E  LDL

An 18-year-old woman has had recurrent acute attacks of dyspnea for the past 10 years. Between these attacks, she has no medical problems. She is brought to the emergency department within an hour of onset of the latest episode. On physical examination, her temperature is 37.1°C, pulse is 110/min, respiratory rate is 28/min, and blood pressure is 110/70 mm Hg. Expiratory wheezes are auscultated over the chest bilaterally. Pulmonary function studies show severe limitation of airflow, which is relieved on injection of epinephrine. Sputum cytologic examination shows abundant mucus with an inflammatory infiltrate dominated by eosinophils, but mixed with neutrophils and macrophages. Which of the following immunologic mechanisms is of primary importance in the pathogenesis of her disease?

A  Activation of neutrophils and macrophages by IL-8

B  Chemoattraction of eosinophils by exotoxin

C  Proliferation of the T_{H}2 subset of CD4+ T cells

D  Recruitment of monocytes by interferon-γ

E  Stimulation of bronchial smooth muscle cells by ADAM-33

A 64-year-old man has had a low-volume mucoid diarrhea with five bowel movements per day, accompanied by cramping abdominal pain, for the past 2 months. The stool is occasionally blood-streaked. On physical examination, he appears pale. A colonoscopy is performed; the figure shows a representative image of the mucosa from the rectum to the lower portion of the sigmoid. The remaining colonic mucosa appears normal. Biopsy specimens of the affected colon show mucosal crypt distortion, focal crypt abscesses, and mixed inflammatory infiltrates extending to the lamina propria. Over the next 2 years, he develops polyarthritis with no joint deformity, and uveitis. Which of the following additional diseases is he most likely to develop?

A  Atrophic gastritis

B  Dermatitis herpetiformis

C  Orchitis

D  Primary sclerosing cholangitis

E  Rheumatoid arthritis

F  Thyroiditis
CHAPTER 30  Final Review and Assessment

4 A 26-year-old, previously healthy woman has developed fever and generalized diffuse erythematous macular rash resembling sunburn in the past day. Her menstrual cycles are regular. She has nausea, vomiting, abdominal pain, diarrhea, myalgias, sore throat, headache, and dizziness. On physical examination, her temperature is 39.4°C, pulse is 101/min, respirations are 23/min, and blood pressure is 90/40 mm Hg. She has oropharyngeal and conjunctival hyperemia. The vaginal mucosa is erythematous. A tampon is present in the vaginal vault. She is disoriented, but there are no focal neurologic deficits. Laboratory findings show hemoglobin, 13.5 g/dL; hematocrit, 41.4%; platelet count, 100,000/mm³; WBC count, 13,200/mm³; glucose, 70 mg/dL; creatinine, 2.5 mg/dL; total bilirubin, 2.4 mg/dL; AST, 82 U/L; and ALT, 29 U/L. She receives nafcillin with clindamycin.

5 A 19-year-old man is found unconscious and taken to the emergency department. On physical examination, his temperature is 41.2°C, pulse is 103/min, respirations are 27/min and shallow, and blood pressure is 145/100 mm Hg. He develops an intractable cardiac dysrhythmia and dies. At autopsy, the heart is slightly enlarged; microscopically, the distal coronary arteries are thickened. Sections of the brain show a 2-cm area of hemorrhage in the right superior parietal lobe and a 0.5-cm hemorrhage in the medulla. There is a partially cystic, 1-cm area with brown discoloration in the left anterior frontal lobe. This clinical picture is most likely to have developed by usage of which of the following substances?

- A Barbiturate
- B Cocaine
- C Ethanol
- D Heroin
- E Marijuana
- F Methamphetamine
- G Phencyclidine

6 A 49-year-old man has had increasing knee and hip pain for the past 10 years. The pain is worse at the end of the day. During the past month, he has experienced bouts of sharp, colicky, right upper abdominal pain. On physical examination, his temperature is 37°C, pulse is 82/min, respirations are 10/min, and blood pressure is 140/85 mm Hg. He is 175 cm (5 feet 9 inches) tall and weighs 156 kg (body mass index 51). Laboratory findings show glucose of 139 mg/dL, Hb A1c of 10, total cholesterol of 229 mg/dL, and HDL cholesterol of 33 mg/dL. An arterial blood gas measurement shows pH of 7.35; PaO₂, 50 mm Hg; and PaCO₂, 75 mm Hg. Which of the following additional conditions is most likely present in this man?

- A Hashimoto thyroiditis
- B Hypertrophic cardiomyopathy
- C Laryngeal papillomatosis
- D Nonalcoholic fatty liver disease
- E Panlobular emphysema
- F Rheumatoid arthritis

7 A 70-year-old man has had memory loss and decreased ability to perform activities of daily living for the past 2 years. He has increasing exercise intolerance and difficulty breathing for the past year. On physical examination, with auscultation of the chest, rales are audible in the lung bases, and there is a diastolic murmur. He has a marked decrease in sensation to light touch and pinprick over the lower extremities. His gait is ataxic, with the feet widely spaced. He cannot name any of three objects after 3 minutes. He thinks he is an astronaut returned from Mars. An echocardiogram shows aortic regurgitation with a widened aortic root and arch. MRI of the brain shows mild diffuse cortical atrophy and meningeal thickening. Infection with what organism would most likely produce these findings?

- A Borrelia burgdorferi
- B Coxsackievirus B
- C HIV
- D Mycobacterium leprae
- E Mycobacterium tuberculosis
- F Treponema pallidum
- G West Nile virus

8 A 19-year-old woman has sudden onset of severe abdominal and back pain and dyspnea. She has had similar episodes over a 12-year period. She had osteomyelitis of the left hip 1 year ago; the bone culture was positive for Salmonella enteritidis. On physical examination, she has tachycardia. Palpation of the abdomen reveals diffuse tenderness with rigidity of abdominal musculature, but no apparent masses. CT scan of the chest shows prominent pulmonary veins, but no infiltrates. Abdominal CT scan shows the presence of multiple 0.5- to 1-cm stones in the gallbladder, a very small spleen, and prominent hepatic veins. CBC shows hemoglobin of 10.2 g/dL, hematocrit of 30.9%, MCV of 99 μm³, RDW of 22, platelet count of 189,300/mm³, and WBC count of 6320/mm³. What additional laboratory test finding is most likely in this case?

- A Amylase, 694 U/L
- B Anticardiolipin antibody
- C Calcium, 12.3 mg/dL
- D Cholesterol, 250 mg/dL
- E Haptoglobin, 1 mg/dL
- F Triglyceride, 1140 mg/dL

9 A 49-year-old man seeks a prescription for erectile dysfunction. He is 168 cm (5 feet 6 inches) tall and weighs 93 kg (body mass index 33). On physical examination, there are bilateral carotid bruits and a midline palpable abdominal pulsatile mass. Decreased hair is noted over the lower extremities, and a 1-cm shallow ulceration is present in the skin over the right first metatarsal head. He has decreased sensation to light touch and pinprick in the lower extremities. Laboratory findings include hemoglobin, 12.9 g/dL; hematocrit, 42%; WBC count, 8950/mm³; and creatinine, 1.7 mg/dL. Which of the following laboratory test findings is most likely to be present in this man?

- A Oligoclonal IgG bands in CSF
- B Hemoglobin A1c, 8.8%
- C Plasma ACTH, 119 pg/mL
- D Plasma homocysteine, 23 μmol/L
- E Serum anti-parietal cell antibodies

10 A 19-year-old woman has sudden onset of severe abdominal and back pain and dyspnea. She has had similar episodes over a 12-year period. She had osteomyelitis of the left hip 1 year ago; the bone culture was positive for Salmonella enteritidis. On physical examination, she has tachycardia. Palpation of the abdomen reveals diffuse tenderness with rigidity of abdominal musculature, but no apparent masses. CT scan of the chest shows prominent pulmonary veins, but no infiltrates. Abdominal CT scan shows the presence of multiple 0.5- to 1-cm stones in the gallbladder, a very small spleen, and prominent hepatic veins. CBC shows hemoglobin of 10.2 g/dL, hematocrit of 30.9%, MCV of 99 μm³, RDW of 22, platelet count of 189,300/mm³, and WBC count of 6320/mm³. What additional laboratory test finding is most likely in this case?

- A Amylase, 694 U/L
- B Anticardiolipin antibody
- C Calcium, 12.3 mg/dL
- D Cholesterol, 250 mg/dL
- E Haptoglobin, 1 mg/dL
- F Triglyceride, 1140 mg/dL
10. A 10-month-old infant is not meeting developmental milestones and is below ideal weight and height. The parents noted an episode of convulsions 1 week ago. On physical examination, the infant has hepatosplenomegaly and generalized nontender lymphadenopathy. There is tenderness on palpation of the right upper extremity. No focal neurologic deficits are present, but the infant’s attention and movement are diminished. A radiograph of the right arm shows a healing fracture. Laboratory findings show hemoglobin of 9.7 g/dL, hematocrit of 28.4%, platelet count of 76,700/mm³, WBC count of 4200/mm³, glucose of 78 mg/dL, and creatinine of 0.4 mg/dL. A bone marrow biopsy is done and the microscopic appearance is shown in the figure. The infant is most likely to have the near absence of which of the following enzymes?

A. α-l-Iduronidase  
B. α-1,4-Glucosidase  
C. Arylsulfatase A  
D. Glucocerebrosidase  
E. Hexosaminidase A  
F. Sphingomyelinase

11. A 54-year-old man has had nausea for the past 6 months, but he does not report hematemesis. He has increasing malaise. On physical examination, he has decreased sensation to pinprick and light touch over the lower extremities bilaterally. He exhibits mild ataxia when walking. An upper gastrointestinal endoscopy study shows the absence of gastric rugal folds, but no ulceration or mass. Which of the following findings is most likely to be detected on further work up?

A. Positive anti-Smith antibody  
B. Deficiency of factor V  
C. Positive Helicobacter pylori antibody  
D. MCV 125 μm³  
E. Urine glucose 4+

12. A 46-year-old Welsh man has had worsening arthritis involving his hands, knees, hips, and elbows for the past 3 years. He has had increasing orthopnea and worsening pedal edema for the past year. On physical examination, he has decreased range of motion of the lower legs, but no apparent joint deformities, warmth, or swelling. There is a brownish hue to his skin, although he rarely goes outdoors. Laboratory findings show hemoglobin, 13.7 g/dL; hematocrit, 40.8%; MCV, 90 μm³; platelet count, 213,500/mm³; WBC count, 6690/mm³; glucose, 201 mg/dL; creatinine, 1.2 mg/dL; and calcium, 8.2 mg/dL. A mutation in a gene encoding for which of the following most likely explains these findings?

A. B-globin  
B. Glucokinase  
C. LDL receptor  
D. HFE  
E. TNF

13. A 33-year-old woman has had increasing lethargy and sensitivity to sunlight for the past 8 months. She has pain in her hands, elbows, knees, and feet, and muscle aches in her arms and legs. She has had increasing dyspnea for the past week. Physical examination shows no joint deformities, swelling, or redness. On auscultation of the chest, a friction rub is audible. A chest radiograph shows bilateral pleural effusions. Laboratory findings show hemoglobin, 11.6 mg/dL; hematocrit, 34.3%; MCV, 84 μm³; platelet count, 133,400/mm³; WBC count, 4610/mm³; glucose, 80 mg/dL; creatinine, 2.4 mg/dL; and calcium, 7.9 mg/dL. Which of the following additional laboratory tests would be most helpful to diagnose her underlying condition?

A. Anti–acetylcholine receptor antibody  
B. Anti–DNA topoisomerase antibody  
C. Anti–glomerular basement membrane antibody  
D. Antimicrosomal antibody  
E. Antimitochondrial antibody  
F. Antinuclear antibody

14. A 31-year-old woman has had a persistent fever for the past 2 months. Her temperature has ranged from 38.3° C to 38.6° C on multiple occasions. On physical examination, she has diffuse abdominal pain and mild splenomegaly, but no hepatomegaly or lymphadenopathy. Laboratory studies show Hgb, 13.2 g/dL; Hct, 39.8%; MCV, 930 μm³; platelet count, 242,000/mm³; and WBC count, 12,290/mm³ with 71% segmented neutrophils, 19% lymphocytes, and 10% monocytes. CT imaging of her abdomen shows an ill-defined pelvic soft-tissue density mass with a mottled lucent center and a small, square, radiopaque area. Review of her medical record reveals that a salpingo-oophorectomy for ectopic pregnancy was performed on the left side 3 months ago. Which of the following is the most likely cause of her persistent fever?

A. Abscess  
B. Non-Hodgkin lymphoma  
C. Ovarian cystadenocarcinoma  
D. Sarcoidosis  
E. Pelvic inflammatory disease
A 52-year-old woman has a 3-month history of fatigue on exertion, dizziness, and syncopal episodes. She has trouble keeping her eyes open toward the end of the day and has double vision. On physical examination, she is afebrile and exhibits marked pallor, but no hepatosplenomegaly or lymphadenopathy. There is decreased motor strength with repetitive motion, but no apparent muscle atrophy, joint deformity, pain, or redness. Laboratory findings show hemoglobin of 6.6 g/dL; hematocrit of 19.9%; platelet count of 199,800/mm³; WBC count, 7385/mm³; creatinine, 4.7 mg/dL; glucose, 238 mg/dL; total protein, 5.7 g/dL; albumin, 2.7 g/dL; AST, 167 U/L; ALT, 69 U/L; alkaline phosphatase, 48 U/L; total bilirubin, 1.5 mg/dL; and prothrombin time, 23 seconds. What is the most likely diagnosis?

A Anti-acetylcholine receptor antibody
B Anti-DNA topoisomerase
C Anti-glomerular basement membrane antibody
D Antimitochondrial antibody
E Antinuclear antibody

A 24-year-old, previously healthy man has developed a cough with bloody sputum along with decreased output of dark urine over the past 2 days. He smokes one pack of cigarettes per day and works as a histotechnologist. A chest radiograph shows diffuse infiltrates most pronounced in the lower lobes. Laboratory findings show hemoglobin of 6.6 g/dL, hematocrit of 19.9%; platelet count of 199,800/mm³; WBC count of 4780/mm³; and reticulocyte count of 0.1%. A bone marrow biopsy specimen shows markedly reduced erythropoiesis. The figure shows a chest CT scan. Which of the following serologic laboratory tests is most likely to be found in this woman?

A Anti-acetylcholine receptor antibody
B Anti-DNA topoisomerase
C Anti-glomerular basement membrane antibody
D Antimitochondrial antibody
E Antinuclear antibody

A 24-year-old woman has developed right-sided facial pain over the past 24 hours. She has experienced a 5-kg weight loss over the past 6 months, despite increasing caloric intake. On physical examination, there is swelling with marked tenderness over the right maxilla, exophthalmos on the right side, diffuse abdominal pain, poor skin turgor, and dry mucous membranes. Her temperature is 37.7°C. She has fever, tachycardia, but no murmurs, and tachypnea; the lung fields are clear. Laboratory findings show Na+, 131 mmol/L; K+, 4.6 mmol/L; Cl−, 92 mmol/L; CO₂, 36 mg/dL; glucose, 361 mg/dL; and urea nitrogen, 36 mg/dL. An abdominal ultrasound scan shows normal-sized organs. A renal biopsy specimen shows a crescentic glomerulonephritis. Which of the following pathologic mechanisms most likely produced this patient’s pulmonary disease?

A Antibody directed against basement membrane collagen
B Apoptosis induced by CD8+ lymphocytes
C Complement activation by circulating antigen-antibody complexes
D Macrophage activation by CD4+ lymphocytes
E Release of inflammatory mediators from mast cells

A 58-year-old man has noticed increasing abdominal girth and decreased libido for the past 7 months. Physical examination shows an enlarged abdomen with a fluid wave, but no tenderness or masses; the spleen tip is palpable. Bismarckian, and hyperpigmentation over the cheeks and neck are present. What is the most likely diagnosis?

A Adrenal atrophy
B Aortic valvular stenosis
C Autoimmune gastritis
D Chronic glomerulonephritis
E Hypertrophic cardiomyopathy
F Micronodular cirrhosis

A 72-year-old woman with chronic bronchitis from cigarette smoking has been bedridden for the past 2 weeks. She experiences sudden, severe dyspnea with chest pain. On examination, her temperature is 37°C, pulse is 104/min, respiratory rate is 31/min, and blood pressure is 100/60 mm Hg. Her left leg is swollen and painful on raising. She appears cyanotic. A systolic ejection sound and diastolic murmur are auscultated over the pulmonic region. A chest radiograph shows a prominent right border of the heart. Laboratory studies show an elevated D-dimer level. She then develops right leg weakness. MRI of the brain shows early infarction within the left hemisphere. Which of the following cardiac conditions best explains these findings?

A Constrictive pericarditis
B Dilated cardiomyopathy
C Ventricular septal defect
D Infective endocarditis
E Rheumatic heart disease

A 24-year-old woman has developed right-sided facial pain over the past 24 hours. She has experienced a 5-kg weight loss over the past 6 months, despite increasing caloric intake. On physical examination, there is swelling with marked tenderness over the right maxilla, exophthalmos on the right side, diffuse abdominal pain, poor skin turgor, and dry mucous membranes. Her temperature is 37.7°C. She has fever, tachycardia, but no murmurs, and tachypnea; the lung fields are clear. Laboratory findings show Na+, 131 mmol/L; K+, 4.6 mmol/L; Cl−, 92 mmol/L; CO₂, 36 mg/dL; glucose, 361 mg/dL; and urea nitrogen, 36 mg/dL. An arterial blood gas measurement shows pH, 7.2; P0₂, 98 mm Hg; PCO₂, 28 mm Hg; and HCO₃⁻, 10 mmol/L. Fine-needle aspiration of the right maxillary region is performed. What organism is most likely to be present in this aspirate?

A Actinomyces israelii
B Bacillus anthracis
C Cytomegalovirus
D Clostridium perfringens
E Cryptococcus neoformans
F Mucor circinelloides
UNIT II  Diseases of Organ Systems

20. A 31-year-old man has an infertility work-up. He has aspermia. He also has chronic diarrhea with elevated quantitative stool fat. He has had recurrent, severe respiratory tract infections since early childhood. As a neonate, he had bowel obstruction from meconium ileus. He is most likely to have an abnormality involving mutation in which of the following genes?

A. CFTR  
B. FGFR  
C. G6PD  
D. HFE  
E. NF1  
F. p53

21. Children 6 to 10 years old in the same community are observed by the local physician to be doing poorly in school, which has been attributed to behavioral problems. Their parents state that these children have poor appetites, complain of nausea, and have frequent headaches. On physical examination, they have decreased sensation to touch over the lower extremities. They exhibit loss of fine motor control of movement and have a slightly ataxic gait. A representative CBC shows hemoglobin of 11.8 g/dL, hematocrit of 35.2%, MCV of 82 μm³, platelet count of 282,300/mm³, and WBC count of 4745/mm³. Examination of the peripheral blood smear shows basophilic stippling of the RBCs. Excessive chronic ingestion of which of the following substances is most likely to explain these findings?

A. Cadmium  
B. Copper  
C. Iron  
D. Lead  
E. Nickel

22. A 45-year-old, previously healthy woman has had a chronic nonproductive cough for the past 2 months. One week ago, her cough was productive of blood-streaked sputum. She does not smoke. Physical examination shows temperature of 37.5°C and blood pressure of 140/90 mm Hg. On auscultation, bilateral crackles are audible in the lungs. A chest radiograph shows bilateral nodular and cavity infiltrates, but there are no masses. Laboratory findings show hemoglobin, 11.7 g/dL; hematocrit, 35.2%; platelet count, 217,000/mm³; WBC count, 6330/mm³; serum glucose, 72 mg/dL; creatinine, 2.6 mg/dL; and urea nitrogen, 25 mg/dL. Urinalysis shows 1+ proteinuria, 2+ hematuria, and no glucose or ketones. A transbronchial biopsy specimen shows necrotizing granulomatous vasculitis of the alveolar capillaries and small peripheral pulmonary arteries. A renal biopsy specimen shows a crescentic glomerulonephritis. Which of the following serologic tests is most likely to be positive in this woman?

A. Anti-DNA topoisomerase I antibody  
B. Anti-glomerular basement membrane antibody  
C. Anti-Jo-1 antibody  
D. Antimitochondrial antibody  
E. Anti-neutrophil cytoplasmic autoantibody  
F. Anti-double stranded DNA antibody  
G. Antiribonucleoprotein antibody

23. A 52-year-old man has a 20-year history of Crohn disease. Over the past 14 months, he has had increasing fatigue with worsening peripheral edema. On physical examination, he has pitting edema to his knees. Laboratory studies show serum urea nitrogen is 35 mg/dL and creatinine is 3.8 mg/dL. Urinalysis shows proteinuria without hematuria, glucosuria, or ketonuria. A renal biopsy specimen shows deposits of amorphous pink material in glomeruli, arterioles, and peritubular interstitium. By electron microscopy, these pink deposits are composed of nonbranching 7.5- to 10-nm fibrils. Which of the following proteins is most likely to form these fibrils?

A. Amyloid-associated  
B. Atrial natriuretic peptide  
C. β2-Microglobulin  
D. Calcitonin  
E. Lambda light chain  
F. Transthyretin

24. A 40-year-old man has been bothered by oral candidiasis, fever, and diarrhea for the past year. On physical examination, he has muscle wasting. His weight is 70% of normal for his height and age. He has generalized nontender lymphadenopathy, but no hepatosplenomegaly. He developed three irregular, 1- to 2-cm, reddish-purple, nodular skin lesions on his forearm in the past 3 months. Laboratory findings show hemoglobin, 12.2 g/dL; hematocrit, 36.5%; MCV, 85 μm³; platelet count, 188,000/mm³; and WBC count, 2460/mm³ with 82% segmented neutrophils, 4% bands, 6% lymphocytes, 6% monocytes, and 2% eosinophils. Infection with which of the following organisms is most likely to produce these findings?

A. Hepatitis C virus  
B. Herpes simplex virus  
C. HIV  
D. Mycobacterium leprae  
E. Staphylococcus aureus  
F. Streptococcus pyogenes

25. A 30-year-old man has noted joint pain in the right hip and left elbow and a headache for the past week. One month ago, he had similar pain in the left hip and knee, which slowly resolved. He remembers having a ringlike skin rash on his left thigh that developed 4 months ago after a camping trip in the woods in Connecticut. On physical examination, there is joint tenderness, but no swelling or deformity of the right hip and left elbow. His heart rate is slightly irregular. What infectious agent is most likely to produce these findings?

A. Borrelia burgdorferi  
B. Mycobacterium tuberculosis  
C. Streptococcus pyogenes  
D. Staphylococcus aureus  
E. Yersinia enterocolitica
A 63-year-old man has become more withdrawn, less talkative, and less active over the past 3 years. He now spends most of his day in bed, although he has minimal difficulty with movement. On physical examination, he has 5/5 motor strength in all extremities; there is no apparent tremor or ataxia. There are no focal neurologic deficits. He can remember only one of three objects after 3 minutes. His mood is depressed. His condition improves with use of an acetylcholinesterase inhibitor. One year later, he has an episode of aspiration while eating and dies 1 week later of pneumonia. At autopsy, the brain weighs 1000 g. The gross appearance is shown in the figure. Which of the following microscopic findings is most likely to be seen in the frontal cortex?

- A β amyloid deposits
- B Absence of Betz cells
- C Alzheimer type II cells
- D Arteriolosclerosis
- E Red neurons
- F Spongiform change

A 51-year-old man has had increasing lethargy over the past year. On physical examination, his blood pressure is 165/100 mm Hg. He has deformity and decreased range of motion of the first three metacarpophalangeal (MCP) joints on the right and the second two MCP joints on the left. There is a 2-cm firm, painless nodule over the left olecranon bursa. A similar 1-cm nodule is palpated in the helix of the right ear, and another 1.5-cm nodule is palpable over the right Achilles tendon. Urinalysis shows specific gravity of 1.012, pH 5.5, 1+ hematuria, 1+ proteinuria, and no glucose. The serum urea nitrogen level is 31 mg/dL, and the creatinine is 3.2 mg/dL. Aspiration of material from the nodule at the left elbow is performed. Which of the following types of crystals is most likely to be seen microscopically in this aspirate?

- A Calcium pyrophosphate dihydrate
- B Cholesterol
- C Cystine
- D Hydroxyapatite
- E Sodium urate

A 22-year-old woman incurs multiple blunt trauma with bilateral femoral and right humeral fractures from a fall. The fractures are treated with open reduction and internal fixation. She is in stable condition until 3 days later, when she becomes progressively delirious and then comatose. On physical examination, she is afebrile. Head CT scan shows generalized brain edema. The representative gross appearance of her brain is shown in the figure. Lumbar puncture yields clear CSF with no RBCs, one mononuclear cell, and normal protein and glucose. Her serum glucose is 102 mg/dL, and creatinine is 0.9 mg/dL. What is the most likely diagnosis?

- A Central pontine myelinolysis
- B Diffuse axonal injury
- C Fat embolism
- D Ruptured berry aneurysm
- E Staphylococcus aureus abscesses
- F Viral meningitis

A translational research project is focused on development of a pharmacologic agent that would affect molecular signaling pathways within cells. This agent binds to type 1 TNF receptor (TNFR1), which triggers activation of intracellular caspases. For what condition is this agent most likely to be useful?

- A Adenocarcinoma
- B Atherosclerosis
- C Bronchiectasis
- D Cirrhosis
- E Dementia
- F Osteoporosis

phosphatase, 80 U/L; total bilirubin, 4 mg/dL; and direct bilirubin, 3.1 mg/dL. Serologic test results are positive for anti-HCV and negative for anti-HBs and IgM anti-HAV. Urinalysis shows 4+ proteinuria and 1+ hematuria. CT scan of the abdomen shows a small amount of ascites, mild hepatomegaly, and no splenomegaly or lymphadenopathy. A biopsy specimen of an ulcerated skin lesion shows leukocytoclastic vasculitis involving the upper dermis. What is the most likely diagnosis?
31. A 32-year-old woman has noticed a decline in dexterity and strength of her right hand in her work as an auto mechanic for the past 2 years. She experienced painful burning sensations in the left upper extremity 1 year ago. She had an episode of decreased visual acuity in the left eye lasting 3 days. She is insensitive to heat. On physical examination, she is afebrile and her blood pressure is normal. Motor strength in the right extremity is 4/5, but 5/5 elsewhere. Vision is 20/100 in the left eye and 20/40 in the right eye. One year later, she reports chronic constipation and incontinence. A magnetic resonance image of her brain is shown in the figure. What is the most likely diagnosis?

A. Diabetes mellitus  
B. Graves disease  
C. HIV infection  
D. Multiple sclerosis  
E. Myasthenia gravis  
F. Systemic lupus erythematosus

32. A 32-year-old woman whose pregnancy was uncomplicated gives birth at term. A newborn physical examination shows a small lower lumbar skin dimple with a protruding tuft of hair. A radiograph shows that the underlying L4 vertebra has lack of closure of the posterior arches. What is the most likely diagnosis?

A. Arnold-Chiari malformation  
B. Dandy-Walker malformation  
C. Meningomyelocele  
D. Spina bifida occulta  
E. Tuberous sclerosis

33. A 22-year-old woman is diagnosed with bipolar disorder. Over the next year, she develops neurologic manifestations that include resting and intention tremors, rigidity, chorea, dysphagia, and dysarthria. On physical examination, she has bilateral Babinski signs. There are ringlike deposits of green material involving the cornea bilaterally, but her vision is not decreased. One year later, she has an illness that lasts 3 weeks, with nausea, vomiting, and malaise and scleral icterus. Laboratory findings include serum AST, 100 U/L; ALT, 122 U/L; alkaline phosphatase, 105 U/L; total bilirubin, 4.5 mg/dL; glucose, 77 mg/dL; and creatinine, 0.9 mg/dL. Serologic test results for hepatitis A, B, and C are negative. This episode subsides without treatment, but she eventually develops cirrhosis. A mutation in a gene encoding for what substance is most likely to be present in this woman?

A. α1-Antitrypsin  
B. CFTR  
C. Copper-transporting ATPase  
D. Galactose-1-phosphate uridyltransferase  
E. Glucocerebrosidase  
F. Glucose-6-phosphatase

34. A 57-year-old woman experiences a sudden loss of consciousness. She has a history of untreated hypertension, and has smoked one pack of cigarettes per day for the past 40 years. On physical examination, her temperature is 37.1° C, pulse is 70/min and irregular, respirations are 18/min, and blood pressure is 90/40 mm Hg. Carotid and radial pulses are diminished compared with femoral and posterior tibial pulses. Auscultation of the chest reveals faint heart sounds; lung fields are clear. A chest radiograph shows a widened mediastinum. The chest CT scan with contrast is shown in the figure. Pericardiocentesis is performed, and there is blood in the aspirate. What condition is most likely to produce these findings?

A. Aortic dissection  
B. Bicuspid aortic valve  
C. Small cell anaplastic carcinoma  
D. Takayasu arteritis  
E. Tertiary syphilis  
F. Thromboangiitis obliterans
35 A 62-year-old woman has had a chronic cough and increasing dyspnea for 10 years. A chest radiograph performed 1 year ago showed increased lucency of upper lung fields and bilateral flattening of the diaphragmatic leaves. She has had nausea and vague abdominal discomfort for 6 months. Biopsy specimens from an upper gastrointestinal endoscopic study show a chronic nonspecific gastritis with no detectable *Helicobacter pylori* organisms. During the past month, she has passed red urine on several occasions. Cystoscopic examination shows a 3-cm exophytic mass in the dome of the bladder, and biopsy specimens show a urothelial carcinoma. What is her most likely risk factor for this spectrum of findings?

A. α1-Antitrypsin deficiency
B. Chronic alcoholism
C. Cigarette smoking
D. Exposure to aniline dye
E. Vitamin C deficiency

36 A 42-year-old woman has had increasing weakness, nausea, vomiting, watery diarrhea, and a 5-kg weight loss over the past 7 months. She has generalized muscle weakness, muscle wasting, and increased skin pigmentation on physical examination. After an upper respiratory tract infection lasting 1 week, she develops abdominal pain and faintness and lapses into a coma. Her temperature is 36.9°C, pulse is 83/min, respiration is 17/min and shallow, and blood pressure is 80/40 mm Hg. Laboratory findings show Na+, 129 mmol/L; K+, 3.5 mmol/L; Cl−, 95 mmol/L; CO2, 23 mmol/L; glucose, 48 mg/dL; and creatinine, 0.6 mg/dL. Atrophy of which of the following tissues is most likely to be present?

A. Adrenal cortex
B. Hypothalamus
C. Islets of Langerhans
D. Parafollicular C cells in thyroid
E. Pineal gland
F. Thyroid epithelium

37 A 39-year-old man has experienced diminished libido for the past 4 months. Review of systems indicates that he has had frequent headaches over the past 2 months. On physical examination, he is normotensive, has gynecomastia bilaterally, has normal-sized testes in the scrotum, and exhibits difficulty with peripheral vision. His visual acuity is 20/20 bilaterally. Laboratory findings show Na+, 141 mmol/L; K+, 4.1 mmol/L; Cl−, 102 mmol/L; CO2, 25 mmol/L; glucose, 75 mg/dL; and creatinine, 0.6 mg/dL. Which of the following neoplasms is he most likely to have?

A. Prolactinoma
B. Carcinoid tumor
C. Medullary carcinoma
D. Pheochromocytoma
E. Renal cell carcinoma
F. Small cell anaplastic carcinoma

38 A boy infant is born to a 41-year-old woman following an uncomplicated pregnancy. He is noted at birth to be at the 70th percentile for height and weight. On physical examination, the infant has bilateral palmar transverse creases and absent distal flexion creases on the fifth digits. The palpebral fissures are oblique. He has brachycephaly. On auscultation of the chest, a holosystolic murmur is audible. During childhood, intellectual disability is exhibited, but the child is able to perform activities of daily living. At age 17 years, the boy has a series of severe upper respiratory tract infections. CBC shows hemoglobin, 10.2 g/dL; hematocrit, 30.5%; MCV, 89 μm3; platelet count, 103,000/mm3; and WBC count, 19,200/mm3 with 14% segmented neutrophils, 6% bands, 22% lymphocytes, 13% monocytes, and 45% blasts. What karyotype is most likely to be present in this boy?

A. 45,X
B. 46,XY
C. 47,XY,13
D. 47,XY,18
E. 47,XY,21
F. 47,XXY

39 A 52-year-old woman has noticed increasing thirst and urine output for the past 6 months. She has had flank pain on the right during the past month. On physical examination, her temperature is 37°C, pulse is 77/min, respiration is 14/min, and blood pressure is 150/95 mm Hg. There are bilateral palpable masses in the abdomen. The figure shows her abdominal CT scan. Urinalysis shows specific gravity of 1.010, pH 6.5, 2+ proteinuria, 2+ hematuria, and no glucose or ketones. Laboratory findings show hemoglobin of 10.4 g/dL, hematocrit of 31.3%, glucose of 102 mg/dL, creatinine of 5.5 mg/dL, and urea nitrogen of 53 mg/dL. One year later, she develops a sudden, severe headache. CT scan of the head shows a subarachnoid hemorrhage at the base of the brain. What is the most likely diagnosis?

A. Adult-onset medullary cystic disease
B. ANCA-associated granulomatous vasculitis
C. Autosomal dominant polycystic kidney disease
D. Cystinosis
E. Type 2 diabetes mellitus
F. Polyarteritis nodosa, classic type
40. A 13-year-old girl has been in foster care with 10 caregivers for the past 11 years. On physical examination, there are ecchymoses of the trunk, extremities, and gingivae. A hyperkeratotic, papular rash, with 0.4-cm lesions ringed by hemorrhage, is present in a similar distribution. The child has pain on movement of the arms and legs. There is abnormal depression of the sternum with prominence of the ribs and the costochondral junctions. Radiographs of the arms and legs show bowing of the long bones and widening of the metaphyses, with normal calcification. There is a right femoral subperiosteal hematoma. No fractures are noted. CBC shows hemoglobin of 10.8 g/dL, hematocrit of 32.4%, MCV of 77 μm³, platelet count of 201,300/mm³, and WBC count of 5730/mm³. She is most likely to have a metabolic defect involving which of the following?
   A. Carbonic anhydrase levels
   B. Factor VIII activity
   C. Fibroblast growth factor receptor
   D. Hydroxylation of collagen
   E. Vitamin D synthesis
   F. Scurvy

41. An 11-year-old girl has a respiratory tract infection and is treated with trimethoprim-sulfamethoxazole. Three days later, she develops a sore throat, malaise, fever, and a macular skin rash on the trunk and extremities. Some of the skin lesions have a central raised area of more pronounced erythema. Within 4 days, there are erosions of the oral mucosa and small blisters developing on purpuric skin macules. The blisters enlarge slightly and then show epidermal detachment. The total body surface area involved with blistering and detachment is less than 10%. The occurrence of cutaneous lesions is most likely mediated by which of the following cell types?
   A. CD8+ lymphocytes
   B. Eosinophils
   C. Langerhans cells
   D. Macrophages
   E. Neutrophils
   F. Natural killer cells

42. A 21-month-old child has had recurrent otitis media complicated by mastoiditis for the past 3 months. On physical examination, there is a seborrheic eruption on the skin of the trunk and scalp. Hepatosplenomegaly and generalized nontender lymphadenopathy are present. A chest radiograph shows bilateral 0.5- to 2-cm pulmonary nodules, and there is a 1-cm lesion on the right clavicle and a 1.5-cm lesion on the left seventh rib, both osteolytic. Laboratory findings show pancytopenia. A bone marrow biopsy specimen shows reduced hematopoiesis with an increased number of large cells having oval vesicular nuclei and vacuolated cytoplasm that mark immunocytologically for CD1a. What is the most likely diagnosis?
   A. Acute lymphoblastic leukemia
   B. Gaucher disease
   C. Langerhans cell histiocytosis
   D. Leishmaniasis
   E. Multiple myeloma
   F. Myelodysplastic syndrome

43. A 26-year-old, previously healthy man sustains blunt force trauma to the left upper arm. On physical examination, there is focal swelling and redness. Three weeks later, the superficial contusion has resolved, but now a slightly tender mass is palpated in the outer aspect of the upper left arm. A radiograph of the left arm shows a 5-cm mass in the soft tissue. There is a radiolucent center and surrounding irregular bone formation. One month later, the mass is now 3 cm and painless. CT scan of the arm shows a well-circumscribed mass within muscle with areas of bright calcification throughout. What is the most likely diagnosis?
   A. Gouty tophus
   B. Hemarthrosis
   C. Myositis ossificans
   D. Osteochondroma
   E. Osteosarcoma
   F. Polymyositis

44. A 35-year-old man has had bouts of severe, diffuse abdominal pain accompanied by fever, malaise, and myalgias over the past 4 months. On physical examination, his temperature is 37.7° C, pulse is 81/min, respirations are 20/min, and blood pressure is 145/90 mm Hg. There is diffuse abdominal tenderness, but no masses, and bowel sounds are present. A stool sample is positive for occult blood. Laboratory findings show serum glucose of 73 mg/dL, amylase of 44 U/L, AST of 54 U/L, ALT of 23 U/L, creatinine of 2.4 mg/dL, and urea nitrogen of 22 mg/dL. A renal biopsy specimen shows acute transmural vasculitis of medium-sized arteries; the glomeruli and tubules are unremarkable. Mesenteric artery angiography is performed with the appearance shown in the figure. Which of the following serologic tests is most likely to be positive in this man?
   A. Antimitochondrial antibody
   B. ANA
   C. C-ANCA
   D. Cryptococcus neoformans antigen
   E. HbsAg
   F. Histoplasma capsulatum antibody
45. A 4-year-old girl has become increasingly listless over the past year. She is at the 25th percentile for height and weight. On physical examination, there is pubic hair and clitoral and breast enlargement. There is no hepatomegaly, splenomegaly, or lymphadenopathy. The neurologic examination is unremarkable. Laboratory findings show hemoglobin, 13.7 g/dL; hematocrit, 41.8%; WBC count, 7120/mm³; Na⁺, 128 mmol/L; K⁺, 4.8 mmol/L; Cl⁻, 99 mmol/L; CO₂, 21 mmol/L; glucose, 69 mg/dL; creatinine, 0.5 mg/dL, and ACTH of 95 pg/mL with loss of diurnal rhythm of secretion. What disease process is most likely associated with these findings?

A. Adrenal cortical hyperplasia  
B. Islet cell adenoma  
C. Pituitary microadenoma  
D. Retroperitoneal neuroblastoma  
E. Suprasellar craniopharyngioma  
F. Thyroid medullary carcinoma

46. A 39-year-old man has had a 4-kg weight loss with watery diarrhea and flatulence over the past 8 months. He has had urticarial plaques on extensor surfaces of the elbows and knees and on the upper back for the past month. Some of the plaques have small, grouped vesicles. A biopsy specimen of one of the skin lesions shows neutrophils at the tips of dermal papillae with overlying basal cell vacuolization. Under immunofluorescence microscopy, granular IgA deposits appear at the tips of dermal papillae. The figure shows the microscopic appearance of a jejunal biopsy specimen. More than 20 years later, he develops a T-cell lymphoma of the jejunum. Serologic studies are most likely to reveal antibodies against which of the following?

A. Desmoglein 3  
B. Double-stranded DNA  
C. Cyclic citrullinated peptide  
D. Histone  
E. Ribonucleoprotein  
F. Tissue transglutaminase

47. For the past month, a 33-year-old woman has had burning epigastric pain and nausea and vomiting. An upper gastrointestinal endoscopic study shows multiple 1-cm shallow gastric antral and proximal duodenal ulcerations. She is treated with omeprazole and improves. One year later, she has an episode of severe, colicky lower abdominal pain and hematuria and passes a calcium oxalate calculus. She notes galactorrhea 1 month later, and over the next 2 months ceases to menstruate. She is given a dopamine agonist and improves. Laboratory findings show calcium, 11.1 mg/dL; phosphorus, 2.4 mg/dL; and creatinine, 1.1 mg/dL. Which of the following gene mutations and associated neoplasm is characteristic of this disorder?

A. MEN1 — Islet cell adenoma  
B. RET — Medullary carcinoma  
C. BCL6 — Non-Hodgkin lymphoma  
D. APC — Osteoma  
E. RET — Pheochromocytoma  
F. VHL — Renal cell carcinoma

48. A 60-year-old man has had worsening dyspnea and nonproductive cough over the past 2 years. On physical examination, his temperature is 37.4° C, pulse is 74/min, respirations are 20/min, and blood pressure is 110/70 mm Hg. A chest radiograph shows extensive interstitial lung disease and a prominent right-sided heart border. Spirometry reveals decreased FEV₁ and FVC. His pulmonary disease is most likely caused by exposure to which of the following?

A. Carbon monoxide  
B. Fungal hyphae  
C. Plant pollen  
D. Silica crystals  
E. Sulfur dioxide  
F. Wood dusts

49. A 42-year-old man has noticed worsening myalgias and increasing difficulty swallowing over the past 2 years. When exposed to cold, the skin of his hands turns white. On physical examination, he has an erythematous rash extending across the bridge of his nose. There is swelling and warmth in the joints of his hands. Laboratory findings show hemoglobin, 12.2 g/dL; hematocrit, 36.5%; platelet count, 180,000/mm³; WBC count, 4510/mm³; serum glucose, 72 mg/dL; total bilirubin, 1 mg/dL; AST, 41 U/L; ALT, 19 U/L; alkaline phosphatase, 69 U/L; creatine kinase, 483 U/L; and creatinine, 1.3 mg/dL. The presence of antibodies to which of the following is most characteristic of his condition?

A. ANCA  
B. Cyclic citrullinated peptide  
C. Histone  
D. Smith  
E. Thyroid peroxidase  
F. U1-RNP

50. A family pedigree reveals first- and second-generation female relatives with premature menopause and male relatives with a progressive neurodegenerative disorder starting by their sixth decade. There are more males than females exhibiting mental retardation from childhood by the fourth generation. Genetic analysis of affected persons reveals CGG repeat expansions in a gene encoding for a protein that binds mRNA transcripts in neurons and shuttles them to the synapses. An abnormality involving which of the following organs is most likely to be present in affected males?

A. Adrenal  
B. Pancreas  
C. Pituitary  
D. Testis  
E. Thyroid
A 44-year-old woman has noted dull right upper quadrant pain for the past year. On physical examination, there is right upper quadrant tenderness on palpation. Abdominal CT scan shows a 5-cm circumscribed mass in the superior right lobe of the liver. The figure shows the representative gross appearance of a similar mass. She experiences acute chest pain with rapid dyspnea with diaphoresis 1 month later. Multiple peripheral perfusion defects are seen on a pulmonary ventilation-perfusion scan. Which of the following combinations of pharmacologic agents taken by this patient regularly is most likely to be associated with these findings?

A. Allopurinol and sulfamethoxazole  
B. Ethynyl estradiol and norethindrone  
C. Ibuprofen and acetylsalicylic acid  
D. Isoniazid and rifampicin  
E. Phenacetin and acetaminophen

A 37-year-old primigravida at 30 weeks’ gestation has noted increasing pedal edema, headaches, confusion, and decreased urine output for the past 2 weeks. She now exhibits seizure activity and then lapses into a coma. On physical examination, her temperature is 36.8°C, pulse is 82/min, respirations are 24/min, and blood pressure is 145/95 mm Hg. Her heart rate is regular, and lung fields are clear. The abdomen is soft, and bowel sounds are present. There is pitting edema to the thighs. No vaginal bleeding is noted, and the cervix is not effaced. Laboratory findings show hemoglobin, 11.9 g/dL; hematocrit, 39.7%; WBC count, 12,700/mm³; glucose, 151 mg/dL; total protein, 6.1 g/dL; albumin, 3.2 g/dL; total bilirubin, 2.3 mg/dL; AST, 101 U/L; and ALT, 28 U/L. Urinalysis shows 1+ hematuria, and WBCs and WBC casts. A chest radiograph shows a 3-cm nodule with an air-fluid level in the right upper lobe. What organism is most likely to be cultured from his blood?

A. Candida albicans  
B. Cryptococcus neoformans  
C. Escherichia coli  
D. Listeria monocytogenes  
E. Staphylococcus aureus  
F. Streptococcus pyogenes  
G. Yersinia enterocolitica

A 41-year-old woman has had headaches with blurred vision for the past 3 days and increasing mental confusion in the past day. On examination, her temperature is 37.9°C, pulse is 104/min, respirations are 25/min, and blood pressure is 70/40 mm Hg. She has petechial hemorrhages over her arms and trunk. A stool sample is positive for occult blood. Laboratory findings show hemoglobin of 9.1 g/dL, hematocrit of 27.2%, MCV of 92 μm³, RDW of 19%, platelet count of 8900/mm³, and WBC count of 8950/mm³. The peripheral blood smear shows schistocytes. A serum panel shows creatinine, 3.3 mg/dL; urea nitrogen, 32 mg/dL; and glucose, 80 mg/dL. Ultralarge multimers of von Willebrand factor are present in plasma. What therapy should she receive emergently?

A. Two units of packed RBCs  
B. Six-pack of platelets  
C. Dobutamine  
D. Exploratory laparotomy  
E. Plasmapheresis  
F. Prednisone

A 30-year-old woman has noted a 5-kg weight gain over the past 3 months; she has not had a menstrual period during that time. She has experienced upper abdominal pain for the past month. Physical examination shows abdominal enlargement with apparent ascites. There is no peripheral edema. She has a positive pregnancy test. Additional laboratory findings show hemoglobin, 13.2 g/dL; hematocrit, 39.7%; WBC count, 12,300/mm³; glucose, 80 mg/dL; AST, 581 U/L; ALT, 611 U/L; total bilirubin, 1.3 mg/dL; total protein, 6.2 g/dL; and albumin, 3.5 g/dL. Because she has family history of pulmonary embolism, she is tested for and found to have Factor V Leiden mutation. An abdominal ultrasound scan shows hepatomegaly with heterogeneous echogenicity, and there is an intrauterine gestation with a fetus estimated at 12 weeks’ size. What pathologic finding is most likely to be present in her liver?

A. Choledocholithiasis  
B. Chronic passive congestion  
C. Hepatic venous thrombosis  
D. Hepatocellular adenoma  
E. Metastatic choriocarcinoma  
F. Microvesicular steatosis
56 A 26-year-old woman has had increasingly frequent infections over the past 5 years. Her most recent respiratory infection was due to *Streptococcus pneumoniae*. She now has watery diarrhea. On physical examination, she is below ideal weight. There is a vesicular rash in the T10 dermatomal distribution on the left. Laboratory findings include WBC count, 7200/mm³ with 55% segmented neutrophils, 2% bands, 35% lymphocytes, 6% monocytes, and 2% eosinophils. Quantitative immunoglobulins include IgA of 22 mg/dL, IgG of 175 mg/dL, and IgM of 40 mg/dL. Lymphocyte subsets by flow cytometry show CD4+ cells (absolute) of 630/μL, CD8+ cells (absolute) of 785/μL, B cells of 280/μL, and T cells of 2010/μL. A stool sample examined for ova and parasites shows *Giardia lamblia* cysts. What is the most likely immunodeficiency disorder in this woman?

A Chronic granulomatous disease  
B Common variable immunodeficiency  
C Hyper-IgM syndrome  
D Leukocyte adhesion deficiency  
E Severe combined immunodeficiency

57 A 44-year-old man has had worsening exercise tolerance and peripheral edema during the past 5 years. He has noted increasing central opacifications that interfere with vision. He has frontal baldness. During the past 2 years, he has had progressive memory loss with decreasing ability to perform activities of daily living. On physical examination, there is significant atrophy of masseter, temporalis, scalene, deltoid, trapezius, and sternocleidomastoid muscles. There is bilateral testicular atrophy. A 2-hour glucose tolerance test shows serum glucose of 156 mg/dL. There is hypogammaglobulinemia. His condition worsens over the next 3 years, with increasing muscular weakness. An abnormality in which of the following gene products is most likely to be present in this man?

A α-1,4-Glucosidase  
B Dystrophin  
C Fibroblast growth factor receptor 3  
D Mitochondrial oxidative phosphorylase  
E Myophosphorylase  
F Myotonic dystrophy protein kinase

58 A 53-year-old man is found comatose on the floor of his bathroom. His neighbors say there was a party at his residence and they saw many acutely intoxicated people staggering outside. Vital signs on arrival in the emergency department are temperature, 36° C; pulse, 88/min; respirations, 16/min; and blood pressure, 95/60 mm Hg. There are no signs of trauma. Laboratory studies show blood ethanol level of 0.20 gm%. Which of the following substances found in his urine are most likely to cause acute kidney injury?

A Cystine crystals  
B Glucose  
C Hyaline casts  
D Ketones  
E Myoglobin

59 A 22-year-old woman has sudden onset of severe lower abdominal pain. Her medical history includes *Chlamydia trachomatis* cervicitis. On physical examination, her temperature is 36.9° C, pulse is 90/min, respirations are 17/min, and blood pressure is 90/50 mm Hg. There is lower abdominal tenderness, but no palpable masses. No vaginal bleeding is present. The rectal examination is unremarkable, and a stool sample is negative for occult blood. Bowel sounds are reduced. An abdominal ultrasound scan is performed, and the uterus appears normal in size with no masses visualized, but there is a right adnexal mass. Culdocentesis is performed, and there is blood in the aspirate. Laboratory findings show hemoglobin of 9.5 g/dL, hematocrit of 28.6%, platelet count of 269,300/mm³, and WBC count of 9110/μL. Which of the following laboratory findings is most likely to be present in this woman?

A Decreased coagulation factor XIII  
B Decreased follicle-stimulating hormone  
C Increased carcinoembryonic antigen  
D Increased human chorionic gonadotropin  
E *Entamoeba histolytica* cysts in stool  
F *Schistosoma haematobium* eggs in urine

60 A 43-year-old woman has become increasingly tired and listless over the past 5 months. She has had menometrorrhagia for the past 3 months. On physical examination, there are no remarkable findings except for a positive result on stool guaiac testing. Laboratory studies show hemoglobin, 9.2 g/dL; hematocrit, 27.3%; and MCV, 75 μm³. Pelvic ultrasound reveals an enlarged uterus. A Pap smear shows abnormal cells of probable endometrial origin. Colonoscopy is performed, followed by partial colectomy; the gross appearance of the lesion is shown in the figure. Which of the following molecular abnormalities has most likely led to these findings?

A Germline inheritance of *APC* gene mutation  
B Homozygous loss of *PTEN* gene  
C Inactivation of the Rb protein by HPV-16  
D Mutation in a DNA mismatch-repair gene  
E Tyrosine kinase activation caused by c-KIT mutation
A 51-year-old man living on the island of St. Helena has had a downturn in his political fortunes. Over the past 3 years, and particularly over the past year, he has had increasing bouts of abdominal pain, anorexia, nausea, vomiting, dysuria, lethargy, spiking fevers, diarrhea, constipation, excessive weakness, heavy perspiration, and weight loss. He is given a large dose of calomel (a mercury-containing compound) a few days before his death on May 5, 1821, a treatment that has since vanished for good reason. An autopsy shows hepatomegaly (with steatosis?) and ulceration with thickening of the stomach. The autopsy report does not record skin and nail changes, such as hyperkeratosis and hyperpigmentation. If those changes had been present, and squamous cell carcinoma of the skin, the findings would have been most suggestive of chronic poisoning with which of the following metals?

A  Arsenic  
B  Beryllium  
C  Chromium  
D  Cobalt  
E  Lead  
F  Nickel

A 90-year-old woman died suddenly. Autopsy shows a 5-cm skin ulceration extending to the sacrum. She had diffuse muscle wasting; a microscopic section shows decreased size of muscle fibers without inflammation or fibrosis. Her bones show marked osteoporosis, and there is vertebral column kyphosis. A finding on examination of the lungs is shown in the figure. There is pneumonia in the lower lobe of the right lung. Which of the following conditions most likely predisposed this patient to the pathologic findings seen at autopsy?

A  Antiphospholipid syndrome  
B  Aplastic anemia  
C  Chronic alcoholism  
D  Elder abuse with blunt trauma  
E  Immobilization  
F  Malnutrition

A 29-year-old man notes burning pain on urination with a urethral discharge that has persisted for 3 days. A sample of the exudate is positive by ELISA for Chlamydia trachomatis. The man has increasing stiffness of the knees and ankles and lower back pain 3 weeks later. A radiograph of the lumbar spine shows narrowing with sclerosis of the sacroiliac joints. One month later, he develops painful erythema of the glans penis, and the conjunctivae are red. A follow-up examination shows a slightly irregular heart rate and a murmur suggestive of aortic regurgitation. The back pain continues off and on for 5 more months. Which of the following test results is most likely to be positive in this man?

A  ANCA  
B  ANA  
C  HLA-B27 genotype  
D  Anti-Borrelia antibodies  
E  Rapid plasma reagin  
F  Rheumatoid factor  
G  U1-RNP

A 19-year-old man has been having headaches for the past month. On physical examination, his blood pressure is 160/95 mm Hg. On funduscopic examination, there are bilateral retinal angiomas. Abdominal CT scan shows a 3-cm mass involving the right adrenal gland. Laboratory testing shows increased urinary catecholamines. The mass is removed surgically. He develops a movement disorder 5 years later with incoordination and ataxia. MRI of the brain shows a 2-cm mass in the left cerebellar hemisphere and a 1-cm mass in the vermis. These are removed surgically. Six years later, he has right flank pain with hematuria; his abdominal CT scan is shown in the figure. His hemoglobin concentration is 20.3 g/dL, and hematocrit is 60.9%. Which of the following gene mutations and associated syndrome does he most likely have?

A  APC—Gardner syndrome  
B  MET—Denys-Drash syndrome  
C  NF2—Neurofibromatosis type 2  
D  TSC1—Tuberous sclerosis  
E  VHL—Von Hippel-Lindau disease  
F  WT1—Beckwith-Wiedemann syndrome
65. A 43-year-old woman has had increasing difficulty swallowing over the past year. She notices that her hands turn white and are painful on exposure to cold. She remarks, “I may be getting older, but at least I don’t have any wrinkles on my face or hands yet.” On physical examination, her blood pressure is 115/75 mm Hg. The skin of her face and hands appears taut and shiny. A punch biopsy specimen of the skin of the hand shows dermal collagenous fibrosis and focal calcification. She receives yearly esophageal dilation for the next 20 years, during which time she develops no serious illnesses. Which of the following serologic test results is most likely to be positive in this woman?

A. Anticentromere antibody
B. Anti-DNA topoisomerase antibody
C. Antimicrosomal antibody
D. Antimitochondrial antibody
E. Anti-neutrophil cytoplasmic autoantibody
F. Antitransglutaminase antibody

66. A 32-year-old woman has had increasing malaise and a 10-kg weight loss over the past 6 months. Physical examination shows muscle wasting, and there is a tan-yellow, plaquelike coating on her tongue. A scraping of the material from her tongue microscopically shows budding cells with pseudohyphae. She develops watery diarrhea 6 months later; a stool specimen contains cysts of Cryptosporidium parvum. She then develops a fever, cough, and severe dyspnea. Bronchoalveolar lavage is done; the figure shows the microscopic findings with GMS staining. Which of the following laboratory findings is most likely to be present in this woman?

A. ANA titer of 1:1024
B. CD4+ lymphocyte count of 111/μL
C. Complement C2 undetectable
D. IgG of 88 mg/dL
E. Neutrophil oxidative burst assay less than 5%
F. Positive rapid plasma reagin

67. A 73-year-old man has had bilateral knee and hip pain for the past 25 years and has taken a medication for this pain for the past 5 years. During the past year, he has noticed increasing frequency of headaches, dizziness, tinnitus, confusion, and nausea. One week ago, he experienced an episode of hematemesis. On physical examination, there are scattered petechiae on his arms and legs. His heart rate is regular and the lungs are clear. No neurologic deficits are noted. Laboratory findings show hemoglobin, 11.1 g/dL; hematocrit, 33.1%; MCV, 72 μm³; platelet count, 317,200/mm³; and WBC count, 5915/mm³. The partial thromboplastin time and the prothrombin time are normal. Platelet function analysis shows decreased aggregation in response to ADP and collagen stimulation. An upper gastrointestinal endoscopy shows gastric mucosal erythema and a 1.8-cm, sharply demarcated, shallow, antral ulceration. Long-term use of which of the following pharmacologic agents is most likely to produce these findings?

A. Acetylsalicylic acid
B. Acetaminophen
C. Adalimumab
D. Methotrexate
E. Oxycodone
F. Propoxyphene

68. A 16-year-old girl has had irregular menstrual cycles since menarche 2 years ago and has not menstruated for 3 months. She has not used contraceptives. One week ago, she noticed a small amount of vaginal bleeding and now has sudden onset of severe abdominal pain. On physical examination, there is marked right upper quadrant abdominal tenderness, and bowel sounds are reduced. A stool sample is negative for occult blood. Brownish fluid is noted emanating from a reddish brown, 2-cm mass in the vaginal vault. Her serum hCG level is high. An abdominal ultrasound examination shows multiple 3- to 6-cm masses in the liver, and the uterus appears enlarged. A chest radiograph shows 1- to 3-cm nodules in the lungs. Paracentesis is done, and there is blood in the aspirate. What neoplasm would most likely produce these findings?

A. Adenocarcinoma
B. Choriocarcinoma
C. Clear cell carcinoma
D. Leiomyosarcoma
E. Malignant mixed müllerian tumor
F. Sarcoma botryoides

69. A 44-year-old woman has had easy fatigability along with pain and stiffness of both wrist joints and other small joints of the hands for the past 8 years. The stiffness is marked in the morning and abates as the day goes by. Radiographs of her hands reveal narrowing of the proximal interphalangeal and metacarpophalangeal joint spaces from synovitis and erosion of the cartilage. Laboratory tests show Hgb, 8.4 g/dL; Hct, 23.5%; MCV, 65 fl; and MCH, 23 pg. Her peripheral blood smear shows hypochromic, microcytic RBCs. Her serum iron and iron binding capacity levels are low and the ferritin level elevated. A high level of which of the following is most likely related to the causation of anemia in this woman?

A. C-reactive protein
B. GM-CSF
C. Hepcidin
D. Rheumatoid factor
E. TNF
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70  A study of children with bone deformities assesses risk factors for their disease. One cohort of these children is found to have bone with the microscopic appearance shown in the figure panel at the left, compared to that of bone in normal children, shown in the panel at the right. Symmetric deformities include frontal bossing, long bone bowing, pronounced lumbar lordosis, and costochondral junction widening. No skin or soft tissue abnormalities are noted. What is the most likely underlying risk for this spectrum of bone deformities?
   A  Dietary deficiency of fresh fruit
   B  Diminished sunlight exposure
   C  Hypopituitarism
   D  Inherited mutation in vitamin D receptor
   E  Trauma from child abuse

71  A 4-month-old boy was born at term to an 18-year-old woman, G1, P0, after a normal pregnancy. The woman returned home from work one evening and was told by her boyfriend, who is staying at her home, that the infant died suddenly. An autopsy shows no external anomalies. The infant’s height and weight are at the 45th percentile. Internal examination reveals subarachnoid hemorrhage at the vertex and subdural hemorrhage over the right parietal lobe. The right eye shows petechial hemorrhages at the ora serrata. There is a soft-tissue hemorrhage in the right upper arm. There is a recent fracture of the occipital bone. What is the most likely diagnosis?
   A  Congenital syphilis
   B  Hemophilia A
   C  Osteogenesis imperfecta
   D  Child abuse injuries
   E  Sudden infant death syndrome
   F  Thanatophoric dysplasia

72  A 35-year-old man presents with increasing breathlessness on exertion. On physical examination, his height is 208 cm (82 inches) with long arms and tapering fingers. His joints are exceptionally flexible. He also has bilateral subluxation of crystalline lenses. Echocardiography reveals mitral valve prolapse and an aneurysm in his ascending aorta. Which of the following experimental therapies is likely to be beneficial in treating this man?
   A  Anti–TGF-β antibody
   B  Bone marrow transplantation
   C  Low-calorie, low-cholesterol diet
   D  Lysosomal stabilizing agents
   E  Transduction of fibrillin-1 gene into T cells

73  A 5-year-old child who has received no medical care since birth has had gradual onset of markedly decreased vision bilaterally. The child also has a history of increased respiratory tract infections caused by Haemophilus influenzae, Streptococcus pneumoniae, Klebsiella pneumoniae, and rubeola. The figure shows the representative microscopic appearance of the bronchial mucosa. The child also has passed urinary tract calculi. On physical examination, generalized papular dermatosis is noted. The child has xerophthalmia, and there is marked keratomalacia with corneal clouding. Bilateral cracks are audible in the lungs on auscultation. Which of the following diseases would most likely lead to these findings?
   A  Cystic fibrosis
   B  Congenital syphilis
   C  HIV infection
   D  Kartagener syndrome
   E  Vitamin A deficiency

74  A 47-year-old woman has had increasing abdominal enlargement, with no significant pain, and diarrhea for the past 3 months. There is a fluid wave on examination of the abdomen. Abdominal CT scan shows massive ascites and scattered 0.5- to 1.5-cm cystic to solid nodules on the surfaces of the bowel and abdominal wall. Paracentesis yields a yellow, slightly cloudy fluid with a high protein content. Cytologic examination of the fluid shows clusters of malignant cells. Laboratory studies show a positive CA 125, no elevation of HCG level, and a negative carcinoembryonic antigen test result. What is the most likely neoplasm?
   A  Adenocarcinoma of the ileum
   B  Carcinoid tumor
   C  Choriocarcinoma
   D  Malignant mesothelioma
   E  Serous cystadenocarcinoma
   F  Mucinous cystadenoma

75  What is the ultimate answer in health care, as in life?
   A  A sound diet with exercise prevails
   B  Diagnose early and treat appropriately
   C  Environmental controls aid prevention
   D  Love is the healer for all our ills
   E  Research and develop new therapies
As a student in the health sciences, you have just finished this review book, learning to apply your knowledge base in pathology to clinical and experimental scenarios of human disease states. Which of the following represents your best application of this knowledge?

A  Collegial consultation  
B  Compassionate care  
C  Differential diagnosis  
D  Lifelong learning  
E  Patient education  
F  Research projects

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**ANSWERS**

**1 E** Acute myocardial infarction, cerebrovascular disease, and peripheral vascular disease at a young age suggest heterozygous familial hypercholesterolemia with a mutation in the gene encoding for the LDL receptor. Such individuals have early, accelerated atherosclerosis. The statin drugs are HMG-CoA reductase inhibitors that suppress endogenous cholesterol synthesis and increase LDL receptor synthesis. Acetylcholine receptors at the neuromuscular junction may be targeted by antibodies in myasthenia gravis. The patient does not have findings consistent with diabetes mellitus, another disease that promotes atherosclerosis, because his hemoglobin A1c level is normal, indicating that hyperglycemia has not been present. In type 1 diabetes mellitus, there is reduced beta cell mass; in type 2 diabetes mellitus, there can be a reduction in glucose transport, but insulin receptors are generally not involved. Hepcidin is involved in control of iron absorption.

PBD 9 147, 495  BP 9 222, 336  PBD 8 497  BP 8 232–233

**2 C** Atopic bronchial asthma usually begins in childhood. Although many of the symptoms of bronchial asthma can be related to type I hypersensitivity, this disease is fundamentally T cell–mediated chronic inflammation of the bronchial wall. The Th2 type CD4+ T cells drive type I hypersensitivity by favoring IgE production and eosinophil recruitment. The dominance of Th2 helper cells is accompanied by down-regulation of the Th1 helper cells because the CD4+ T-cell differentiation is skewed toward Th2 cells. Neutrophils, eosinophils, and macrophages also can be present in the inflammatory exudates, but these cells all are recruited secondarily to the Th2 response. ADAM-33, a metalloproteinase, has been linked to the airway remodeling seen in bronchial asthma. Specifically, certain polymorphisms of the ADAM-33 gene are associated with proliferation of bronchial smooth muscle cells and fibroblasts. There is, however, no evidence that ADAM-33 contributes to the immunologic alterations seen in bronchial asthma.


**3 D** Ulcerative colitis has a peak age of onset between 15 and 30 years. A second peak occurs between 60 and 80 years. The colonoscopic appearance is of a mucosal erythematous and finely granular surface that looks like sandpaper. Extra-intestinal manifestations of ulcerative colitis can occur, including primary sclerosing cholangitis (PSC), uveitis, migratory polyarthritis, and erythema nodosum. Sixty percent to 70% of cases of PSC have concurrent inflammatory bowel disease. Atrophic gastritis causes pernicious anemia and may coexist with other autoimmune disorders, such as Hashimoto thyroiditis. Dermatitis herpetiformis occurs in patients with celiac disease, both caused by gluten sensitivity. Orchitis is most often a complication of mumps virus infection, particularly in adults. Rheumatoid arthritis is typically associated with Sjögren syndrome. Thyroiditis may occur in association with other autoimmune endocrinopathies, such as Addison disease.


**4 E** Toxic shock syndrome can be caused by some strains of *Staphylococcus aureus* that produce exotoxins acting as superantigens, such as TSST-1, which are T-cell mitogens. These superantigens are able to stimulate more than 10% of the body’s T cells, provoking an exuberant and dysregulated immune response characterized by release of the cytokines that mediate cell injury. *Bacillus anthracis* can cause anthrax, a serious illness with pneumonia and meningitis as the predominant findings. *Clostridium perfringens* is known to cause wound infections with gas gangrene. *Enterococcus* generally does not cause such a rapidly progressing illness. Listeriosis is associated with food poisoning and can cause life-threatening disease, but typically without desquamation. *Vibrio vulnificus* organisms may cause a blistering dermatitis, but this infection follows ingestion of contaminated shellfish.

PBD 9 363  BP 9 321  PBD 8 357  BP 8 334

**5 B** Features of acute cocaine toxicity, such as hyperthermia, and chronic cocaine use, with coronary arteriopathy and cerebral hemorrhagic strokes, are present. The brown discoloration from hemosiderin deposition in the left anterior frontal lobe suggests a subacute to remote hemorrhage. Cocaine is a powerful vasoconstrictor. Similar to cocaine, amphetamine and methamphetamine are uppers, or stimulants; however, they do not typically produce arterial changes.
Barbiturates are downers, or depressants. Ethanol is a depressant that can produce liver disease, but the cardiac effect is obscure and results in dilated cardiomyopathy after years of abuse. Heroin is an opiate narcotic that has minimal effects itself, but the typical route of administration is intravenous, and this can lead to numerous infectious complications, such as hepatitis, endocarditis, meningitis, and AIDS. Marijuana is a mild tranquilizer. Phencyclidine is a schizophrenomimetic that causes erratic behavior, but not obvious tissue effects.

6 D Morbid obesity can be associated with complications that include obesity hypoventilation syndrome, probable sleep apnea, glucose intolerance, cholelithiasis, and osteoarthrosis. Macrovesicular steatosis with hepatomegaly is seen in obesity and may even progress to cirrhosis. Weight gain owing to hypothyroidism, which could occur in Hashimoto thyroiditis, is modest and does not lead to morbid obesity. An “obesity cardiomyopathy” resembles dilated cardiomyopathy, but not hypertrophic cardiomyopathy, which typically involves the interventricular septum with myofiber disarray. Laryngeal papillomatosis, which produces airway obstruction (without snoring), occurs more often in children and is not associated with obesity. The blood gas findings in this case could be seen in emphysema, which is not a complication of obesity; panlobular emphysema is much less common than the centrilobular emphysema associated with smoking. Rheumatoid arthritis tends to involve small joints first, and there is no relationship to obesity.

7 F Tertiary syphilis may manifest with syphilitic aortitis from endarteritis obliterans of the vasa vasorum, which is most marked in the proximal aorta. In addition, the patient has findings of neurosyphilis, with tabs dorsalis and general paresis. A positive VDRL test result on CSF aids in diagnosis. None of the remaining organisms listed produces aortic disease. Neuroborreliosis from Lyme disease can produce aseptic meningoencephalitis, encephalopathy, and polyneuropathy. Coxsackievirus B can produce meningoencephalitis and myocarditis. HIV can produce encephalitis characterized by dementia and motor and sensory deficits. Hansen disease, caused by Mycobacterium leprae infection, mainly affects the peripheral nerves. Tuberculosis can produce meningoencephalitis or a mass known as a tuberculoma; obstructive hydrocephalus may occur in chronic meningitis. West Nile virus is most likely to produce severe meningoencephalitis in elderly individuals.

8 E Sickle cell anemia (hemoglobin SS) may present with abdominal crisis and vertebral bone marrow infarction. The haptoglobin is low as a result of sickle cell crisis with hemolysis. The continued hemolysis leads to formation of pigmented gallstones. Autosplenectomy is a consistent finding. There is a predisposition to aseptic necrosis and osteomyelitis, particularly with Salmonella. The MCV is high because of reticulocytosis, and the RDW is quite high because of sickling and hemolysis. An elevated amylase level could be seen in acute pancreatitis, which could occur with gallstones, but this would not explain the anemia or the undetectable spleen. Antiphospholipid syndrome (APS) with anticardiolipin antibodies leads to thrombosis, but the abdomen is not the typical location, and the anemia is not explained by APS. Pancreatitis can complicate hypercalcemia and hypertriglyceridemia, but this does not explain the anemia or undetectable spleen. An increased cholesterol is a risk for atherosclerotic diseases.

9 B Hemoglobin A1c elevation occurs in diabetes mellitus. The patient has complications of type 2 diabetes mellitus that are typical of obesity: a “diabetic foot” as a consequence of peripheral neuropathy and decreased sensation, advanced atherosclerosis with carotid arterial occlusion, and an atherosclerotic aortic aneurysm. Autonomic dysfunction with neuropathy can lead to erectile dysfunction. The oligoclonal bands of IgG are characteristic for multiple sclerosis, which usually has more focal CNS involvement. Cushing disease owing to an ACTH-secreting pituitary adenoma leads to truncal obesity, skin that is easily bruised, and possibly to secondary diabetes mellitus, but not to advanced atherosclerosis. Hyperchromocysteinemia is a risk factor for atherosclerosis, but not for neuropathy. Anti-parietal cell antibodies occur in pernicious anemia with vitamin B12 deficiency, which can lead to decreased sensation in the lower extremities, but not to atherosclerotic complications.

10 D The acute neuronopathic form of Gaucher disease (type 2) is uniformly fatal in children and thankfully is much less common than the milder type 1, which has prolonged survival into adulthood. The marrow is infiltrated by large cells with abundant pale cytoplasm with the appearance of crumpled tissue paper. α-1,4-Glucosidase deficiency is seen in Hurler syndrome, characterized by progressive features of corneal clouding, coarse facial features, and mental retardation. α-1,4-Glucosidase deficiency is present in Pompe disease, a glycogen storage disease that causes marked cardiomegaly. Arylsulfatase A deficiency is present in metachromatic leukodystrophy and causes CNS degeneration without visceral organ involvement. Tay-Sachs disease with cherry-red maculae and progressive neurologic deterioration occurs as a result of diminished hexosaminidase A. Spinogomyelinase deficiency leads to Niemann-Pick disease, with foamy-appearing macrophages filling tissues of the mononuclear phagocyte system.

11 D Pernicious anemia can be due to atrophic gastritis, with lack of intrinsic factor to bind dietary vitamin B12 for absorption. This condition has led to megaloblastic anemia and subacute combined spinal cord degeneration. The anti-Smith antibody is a feature of systemic lupus erythematosus, which has many manifestations, but not typically gastritis or spinal cord degeneration. Factor V (Leiden) deficiency is a risk factor for recurrent thrombosis. Helicobacter pylori infection leads to...
gastrectomy (not typically atrophic), but not to neurologic problems. An elevated urine glucose level suggests diabetes mellitus, which may lead to peripheral neuropathy, but not to gastritis.


12 D The C282Y mutation in the HFE gene can be found in one in nine individuals of Celtic heritage and causes increased absorption of dietary iron. In men beginning around age 40 years, the increased iron stores lead to organ dysfunction, typically involving the heart (cardiomyopathy with congestive heart failure), pancreas (diabetes mellitus), skin (increased pigmentation), and joints (arthritis). In women, increased iron loss through menses delays the onset of this disease for 20 more years. A patient with β-thalassemia would have anemia, although the ineffective erythropoiesis leads to excessive iron absorption. Although this patient has diabetes mellitus and an increased glucose level, diabetes mellitus does not explain all the findings, such as the arthritis. Familial hypercholesterolemia could lead to coronary artery disease and heart failure, but it does not explain the patient’s diabetes or arthritis. Rheumatoid arthritis with joint inflammation mediated in part by TNF typically leads to joint deformities and mostly involves small joints.

PBD9 847–849  BP9 629–630  PBD8 861–863  BP8 654–656

13 F Systemic lupus erythematosus (SLE) manifestations may include photosensitivity, renal failure, body cavity effusions, pericarditis, arthralgias, myalgias, and cytopenias. The antinuclear antibody test is the most sensitive screening test for SLE, and if positive can be followed by the more specific anti–double-stranded DNA antibody test. Acetylcholine receptor antibody may be seen in myasthenia gravis, which would explain muscle weakness but not pain. Anti–DNA topoisomerase is seen in scleroderma, in which there is renal failure and skin thickening, but not muscle weakness after use. Anti–glomerular basement membrane antibody can be seen in Goodpasture syndrome, a form of rapidly progressive glomerulonephritis, often with pulmonary hemorrhage. Antimitochondrial antibody may be seen in primary biliary cirrhosis. ANA is characteristic of many systemic autoimmune diseases, most often systemic lupus erythematosus, which can be accompanied by myalgias, but not by muscle weakness with repetitive movement.

PBD9 206, 627, 655  BP9 115, 457  PBD8 636–637  BP8 476, 830

14 A She has fever of unknown origin (FUO), at least until a record review correlated with the CT imaging suggests that the mass is a residual hemostatic sponge placed at the time of surgery. Radiopaque markers or radio-frequency identification (RFID) chips can be incorporated into such objects for identification. The mild splenomegaly is consistent with intra-abdominal abscess. There is a long differential diagnosis list for FUO. A mass could represent a neoplasm, but lymphomas do not tend to have significant necrosis and a lucent center. Sarcoidosis tends to involve multiple organs, but lymph node enlargement is likely, although central cleftation is not. At the prior surgery, an ovarian neoplasm should have been identified, as would pelvic inflammatory disease that may produce a tubo-ovarian abscess.

PBD9 624  BP9 456  BP8 633  BP8 475

15 A The chest CT scan shows a thymoma posterior to the sternum just below the clavicles. The findings listed describe myasthenia gravis with complications of thymoma, including the rare finding of pure RBC aplasia, which is characterized by selective suppression of the erythroid lineage in the bone marrow. In about half of such cases, removal of the thymic tumor relieves the RBC aplasia, suggesting some autoimmune mechanism as the cause of the aplasia. Thymic disorders are common in myasthenia gravis, either thymic hyperplasia or thymoma (as in this case). Antibodies against the acetylcholine receptor disrupt the function of the myoneural junction. Anti-DNA topoisomerase is seen in scleroderma, in which there is renal failure and skin thickening, but not muscle weakness after use. Anti–glomerular basement membrane antibody can be seen in Goodpasture syndrome, a form of rapidly progressive glomerulonephritis, often with pulmonary hemorrhage. Antimitochondrial antibody may be seen in primary biliary cirrhosis. ANA is characteristic of many systemic autoimmune diseases, most often systemic lupus erythematosus, which can be accompanied by myalgias, but not by muscle weakness with repetitive movement.

PBD9 206, 627, 655  BP9 115, 457  PBD8 636–637  BP8 476, 830

16 A Goodpasture syndrome has antibody directed against the glomerular basement membrane, which also acts on basement membrane in the lung to produce pulmonary hemorrhage and hemoptysis. Cytotoxic CD8+ lymphocytes are part of a cell-mediated immune response effective in eliminating intracellular infections such as influenza. Circulating immune complexes are more likely to be seen in autoimmune diseases, such as systemic lupus erythematosus. Macrophage activation is more typical of chronic inflammation and granulomatous inflammation with type IV hypersensitivity. Release of mediators such as histamine from mast cells is typical of anaphylaxis with type I hypersensitivity.


17 F The ascites, edema, and splenomegaly together with laboratory evidence of hepatic dysfunction suggest a hepatic disorder with portal hypertension, and a common cause is hepatic cirrhosis from chronic alcohol abuse. Decreased estrogen metabolism results in testicular atrophy. The findings of right-sided and left-sided congestive heart failure are associated with alcoholic dilated cardiomyopathy. Macrocytic anemia is common, and the AST is slightly higher than the ALT, features typical of chronic alcoholism. Addison disease resulting from adrenal atrophy would not produce hypoalbuminemia or liver enzyme elevations, and the glucose level is often lower. Aortic stenosis may explain the pulmonary edema. Autoimmune gastritis may lead to gastric mucosal atrophy, loss of parietal cells, and megaloblastic anemia, but not to hepatic abnormalities. In chronic glomerulonephritis
severe enough to produce a hepatorenal syndrome, the renal failure would be much worse and would be indicated by a high serum creatinine. Dilated cardiomyopathy is a type of cardiomyopathy typical of chronic alcoholism.

18 C Elusive as the yeti, the paradoxical embolism explains her findings. She has thrombophlebitis with pulmonary embolism as a result of prolonged bed rest. Her chronic obstructive pulmonary disease has led to cor pulmonale with an enlarged right side of the heart, reversing the shunt across a ventricular septal defect. Shunt reversal allows a thromboembolus arising in the venous circulation to reach the systemic arterial circulation in the brain. A constrictive pericarditis yields a paradoxical pulse from impaired cardiac filling and greater than normal decline in systolic arterial pressure on inspiration; this is not associated with thromboembolism. A dilated cardiomyopathy should lead to global cardiac enlargement, not just enlargement of the right side of the heart, and although mural thrombosis with embolism can occur, there is usually no association with thrombophlebitis. Infective endocarditis can lead to embolization, but in her case there is no fever or infection. Rheumatic heart disease can affect one or more valves, and often leads to left atrial enlargement with mural thrombosis and embolism, but the left atrial border would be prominent, and the pulmonic valve is almost never involved.

19 F Type 1 diabetes mellitus with ketoacidosis is the setting for infection by *Mycor* in the paranasal sinuses, an otherwise unusual infection. T-cell and B-cell function is generally maintained in diabetes mellitus, although neutrophilic function may be depressed, so bacterial infections (staphylococcal, streptococcal, and coliform organisms) most often complicate diabetes mellitus. Actinomycosis can produce chronic subcutaneous abscesses, usually in the neck, lung, or abdomen, and usually following trauma or tissue devitalization. Cutaneous anthrax is rare and produces a localized eschar or ulcerated region. Cytomegalovirus and cryptococcal infections are typically seen in immunocompromised individuals with diminished cell-mediated immunity. *Clostridium perfringens* appears in the setting of soft-tissue infections with gas gangrene.

20 A Cystic fibrosis can be accompanied by agenesis of the vas deferens, a common finding that leads to infertility. With good medical care, patients with cystic fibrosis are living longer, and childbearing becomes an issue. Disorders of fibroblast growth factor receptor (FGFR) can include dwarfism. Glucose-6-phosphate dehydrogenase (G6PD) deficiency results in hemolysis on exposure to oxidants such as antimarial drugs (e.g., primaquine). The HFE gene is abnormal in hereditary hemochromatosis; however, the patient is young for the onset of this disease. *NFI* (neurofibromatosis) is associated with the appearance of various neoplasms, including neurofibromas, pheochromocytomas, and gliomas. *p53* is a tumor suppressor gene, and loss of both alleles can promote the appearance of various malignancies, mainly carcinomas.

21 D Lead poisoning is mainly manifested by neurologic disorders, particularly in children. Lead absorption is enhanced by zinc deficiency; zinc is a trace metal. Lead inhibits heme incorporation into hemoglobin, leading to increased amounts of zinc protoporphyrin with anemia. Cadmium is a heavy metal associated with toxicity to the gastrointestinal tract, kidneys, and lungs. Copper is a trace metal that is unlikely to cause toxicity from environmental sources, although copper accumulation can occur with Wilson disease. Acute iron poisoning is associated with gastrointestinal, renal, and CNS toxicities. Nickel jewelry may cause skin rash; inhaled nickel produces respiratory problems.

22 E Granulomatosis with polyangiitis (ANCA-associated vasculitis) is a multisystem vasculitis that most often involves the lungs and kidneys. C-ANCA is positive in 95% of patients. Anti–DNA topoisomerase I antibody can be seen in diffuse scleroderma, which produces pulmonary interstitial fibrosis and renal hyperplastic arteriolosclerosis. Anti–glomerular basement membrane antibody is seen in Goodpasture syndrome, which produces crescentic glomerulonephritis and pulmonary hemorrhage, but not necrotizing vasculitis. Anti–Jo-1 antibody accompanies polymyositis/dermatomyositis. Antimitochondrial antibody accompanies primary biliary cirrhosis. Anti-dsDNA is positive principally seen in systemic lupus erythematosus (SLE), which can produce a vasculitis, but not a necrotizing granulomatous vasculitis. Anti–RNP antibody is positive in cases of mixed connective tissue disease, which overlaps SLE, scleroderma, rheumatoid arthritis, and polymyositis, but it usually does not entail significant renal or pulmonary involvement.

23 A Reactive systemic amyloidosis has developed from her underlying granulomatous ileitis, and serum amyloid-associated protein is generated by chronic inflammation. Atrophic natriuretic peptide can be seen with isolated atrial amyloidosis. β2-Microglobulin is found with hemodialysis-associated amyloidosis. Calcitonin is a precursor to local amyloid deposition in medullary thyroid carcinomas. Excessive light chains predispose to amyloid deposition with multiple myeloma. Transthyretin can be associated with systemic senile amyloidosis.

24 C The reddish-purple lesions are typical of Kaposi sarcoma in a patient with wasting syndrome, oral thrush, and lymphopenia characteristic of HIV infection with AIDS.
Hepatitis C virus is unlikely to produce skin lesions or lymphopenia of this degree. Herpes simplex virus infections may be seen more frequently in HIV infection, but the lesions are typically vesicular and are located in the perioral or peri-anal regions. Hansen disease, caused by Mycobacterium leprae infection, may produce a faint reddish rash that fades, followed by hypopigmentation or anesthesia of affected skin and sometimes nodular deforming lesions developing over years. Staphylococcal skin infections tend to produce localized abscesses, such as furuncles and boils. Streptococcal skin infections may manifest as abscesses or as cellulitis.

PBD9 253–254 BP9 152 PBD8 246–247 BP8 164–165, 374–376

25 A Lyme arthritis is suggested by meningitis, carditis, and a past history of erythema chronicum migrans. The arthritis may appear weeks to a couple of years after a bite from the vector, the deer tick (Ixodes), and can be migratory and involve the large joints. Tuberculosis rarely can produce a chronic arthritis associated with osteomyelitis of the large joints, leading to ankylosis and deformity. Streptococci may cause an acute arthritis, although a polyarthritis with rheumatic fever owing to group A streptococcal infection can occur along with carditis. Staphylococcus aureus causes an acute suppurative arthritis. Yersinia enterocolitica can cause enteropathic arthritis with an abrupt onset; a yearlong course is possible in HLA-B27–positive individuals.

PBD9 381–382, 1214 BP9 789–790 PBD8 377–378 BP8 824

26 A Alzheimer disease is progressive over years, and the brain becomes decreased in size, with narrower gyri and widened sulci, usually in all lobes except the occipital lobes. The atrophy leads to ex vacuo ventricular dilation. Microscopically, there are neocortical neuron plaques with Aβ amyloid cores. Absence of Betz cells may be seen in amyotrophic lateral sclerosis, in which there is progressive muscular weakness. Alzheimer type II cells, despite the eponym, are not part of Alzheimer disease, but are seen with increased blood ammonia levels as a consequence of hepatic failure—hepatic encephalopathy. Arteriosclerosis can occur in association with hypertension. Red neurons are seen with acute infarction. Spongiform change is most likely to occur in a rapidly progressive dementia (over weeks to months), such as Creutzfeldt-Jakob disease.

PBD9 1287–1292 BP9 836–838 PBD8 1313–1317 BP8 891–893

27 E The findings are most typical of gouty arthritis. Gout can lead to renal failure and to tophaceous deposits in soft tissues and joints, and it is often accompanied by hyperlipidemia. Calcium pyrophosphate dihydrate deposition disease is more common in elderly individuals and usually is asymptomatic; the knees are most often affected, and soft-tissue deposition of the crystals away from joints is unlikely. Cholesterol crystals form in joint cavities after trauma with hemorrhage. Cystine crystals can be seen in the urine in cystinosis, a rare inborn error of metabolism. Hydroxypatite crystals may be found in joints affected by osteoarthritis, with either acute or chronic presentation.

PBD9 1214–1217 BP9 786–789 PBD8 1243–1246 BP8 820–823

28 D Patients with hepatitis C infection can develop chronic hepatitis with persistently elevated liver enzymes. Some patients with hepatitis C develop a mixed cryoglobulinemia with a polyclonal increase in IgG. Renal involvement is common, with either nephrotic or nephritic features. Cryoglobulinemic vasculitis leads to skin hemorrhages and ulceration. Autoimmune hemolytic anemia can lead to a predominantly indirect hyperbilirubinemia. Although hepatocellular carcinoma may complicate hepatitis C infection, a mass lesion would be seen on CT scan, and the alkaline phosphatase level usually is elevated when an intrahepatic mass is present. Chronic liver disease may complicate hereditary hemochromatosis, but these patients usually do not have hepatitis C, and the skin has a slate color, not purpura. Multiple myeloma can increase the serum globulin and produce renal disease, but hepatitis is usually not part of myeloma; vasculitis likewise is not part of myeloma.

PBD9 833–835, 927 BP9 617–618 PBD8 847–848 BP8 643–644

29 C The delay in onset after trauma is consistent with fat embolism to brain, and the figure shows the predominantly white matter petechiae of brain purpura. Fat embolism causing respiratory difficulty or neurologic findings typically has an onset of 1 to 3 days after trauma. The exact mechanism for development of these vascular lesions is unknown, but vascular occlusion with free fatty acid release and platelet activation play a role. Central pontine myelinolysis occurs when hyponatremia is rapidly corrected, leading to white matter edema most marked in the tightly packed crossing fibers of the pons. Diffuse axonal injury can occur as a result of trauma, but symptoms and signs should appear soon after the injury, and the white matter lesions are often microscopic. A ruptured aneurysm is usually not related to trauma, and most of the hemorrhage is at the base of the brain. Because she is afebrile, an abscess is unlikely, and the CT scan did not show a focal lesion. Viral meningitis can produce brain edema, and herpes simplex viral infection can produce hemorrhage, but there is no relationship to trauma, and there should be leukocytosis in the CSF with infection.

PBD9 128 BP9 91 PBD8 126–127 BP8 99

30 A Translational research takes an idea from bench to bedside. The mechanism described is that of apoptosis, or individual cell death. Turning on apoptosis within cancer cells would be a useful pharmacologic effect to cause the tumor to self-destruct. The other listed conditions would not benefit from induction of apoptosis to a similar degree.

PBD9 52–58 BP9 19–22 PBD8 25–29 BP8 19–22

31 D The MR image shows multiple areas of signal intensity representing white matter plaques of demyelination typical of multiple sclerosis, a disease that has various neurologic manifestations. The mean age at onset is 30 years, and it rarely occurs after age 60 years. Optic neuritis, weakness, and sensory changes are frequent manifestations. Diabetes mellitus can produce peripheral neuropathy and retinopathy, but typically in a more symmetric fashion, and white matter plaques are not present. Graves disease can lead to
hyperthyroidism with heat intolerance, but neurologic manifestations are not frequent, and diarrhea is typically present. HIV infection can lead to HIV encephalitis, which can involve white matter, although defined plaques are unlikely, and peripheral neuropathy is infrequent. Myasthenia gravis can produce generalized weakness, which becomes worse with repetitive movement and is not focal; there are no white matter changes. Systemic lupus erythematosus can cause cerebritis but without plaques of demyelination, and focal neurologic deficits are an infrequent finding.

**PBD9 1283–1285** BP9 832–834 PBD8 1310–1312 BP8 887–889

32 D Spina bifida occulta is the mildest form of neural tube defect. It is characterized by defective closure of the vertebral arches, with intact meninges and spinal cord, and “closed” by skin and connective tissue. A radiograph may show its presence in 20% of the population. Despite the lack of an open defect, there is an increased risk for meningitis. In an Arnold-Chiari malformation, there is a small posterior fossa and extension of the cerebellum into the foramen magnum and a lumbar meningomyelocele. Dandy-Walker malformation is detected by ultrasound evidence of a cyst in the fourth ventricle and agenesis of the cerebellar vermis. Meningomyeloceles are open neural tube defects. Tuberculous sclerosis is a rare disease that causes firm hamartomatous “tubers” in the cortex, and is not associated with neural tube defects.

**PBD9 1256** BP9 822–823 PBD8 1284 BP8 872

33 C Wilson disease (hepatolenticular degeneration) with Kayser-Fleischer corneal rings results from a mutation in the gene encoding for a copper-transporting ATPase (ATP7B). Excessive copper deposition occurs, particularly in the liver, putamen, and cornea. Psychiatric and neurologic disturbances are common in Wilson disease, and patients often develop chronic liver disease ranging from acute hepatitis to chronic hepatitis to cirrhosis. In α1-antitrypsin deficiency, there can be chronic hepatitis and pulmonary emphysema, but there are no neurologic changes. Cystic fibrosis can lead to pulmonary disease and pancreatic insufficiency. Galactosemia can lead to liver disease and cirrhosis in early childhood. In its most severe form, Gaucher disease from glucocerebrosidase deficiency can lead to neurologic deterioration, but it does not lead to chronic liver disease. Von Gierke disease is a form of glycogen storage disease that does not commonly progress to cirrhosis.

**PBD9 849–850** BP9 630–631 PBD8 863–864 BP8 656–657

34 A An aortic dissection can extend proximally to envelope and partially occlude the great vessels. Proximal dissections may result in minimal or no chest pain. Blood has dissected into the mediastinum, causing widening, and into the pericardial cavity, causing tamponade. Risk factors include hypertension and atherosclerosis. In addition, cystic medial necrosis makes aortic dissection a serious risk in individuals with Marfan syndrome. A bicuspid aortic valve leads to aortic valvular stenosis, but not to aortic rupture. Neoplasms involving the mediastinum, typically hematologic malignancies such as lymphomas, may produce a superior vena cava syndrome, but neoplasms virtually never invade the arterial media, and bleeding from neoplasms is produced from much smaller vessels. Takayasu arteritis can involve the aortic arch and lead to dissection, but this condition is rare and is most likely seen in women younger than 30 years old. Syphilitic aortitis may produce aortic root dilation and possible rupture, but this is much less common than aortic dissection resulting from hypertension and atherosclerosis. Thromboangitis obliterans (Buerger disease) is an uncommon disorder that affects small to medium-sized arteries in the arms and legs of middle-aged men who smoke.

**PBD9 501–505** BP9 344–348 PBD8 508–510 BP8 359–362

35 C Pulmonary emphysema and gastritis are related to smoking. Multiple malignancies are related to smoking, including urinary tract urothelial carcinoma and renal cell carcinoma. α1-Antitrypsin (AAT) deficiency can explain emphysema, but it would be panlobular, and AAT deficiency is not associated with urinary tract neoplasia. Chronic alcoholism can explain gastritis, but not emphysema or carcinoma. Aniline dyes increase the risk of urothelial carcinoma, but not of emphysema or gastritis. Vitamin C deficiency can lead to soft-tissue hemorrhages and bone pain, but not to carcinoma or emphysema.


36 A Addison disease with bilateral atrophy of the adrenal cortex is most often idiopathic and leads to electrolyte changes owing to loss of mineralocorticoid secretion, mainly aldosterone; when atrophy is marked, glucocorticoid secretion is diminished. Increased ACTH precursor hormones, resulting from loss of feedback from cortisol production, stimulate skin melanocytes. Stress, including infections, may precipitate an addisonian crisis. Loss of islets of Langerhans is a feature of type 1 diabetes mellitus, with hyperglycemia and possible ketoacidosis as complications. Loss of releasing or inhibiting hormones from the hypothalamus affects the pituitary and leads to multiple endocrinopathies, but not specifically to loss of ACTH. Parafollicular C cells of the thyroid produce calcitonin, which plays a minor role in calcium homeostasis. The pineal gland produces melatonin, which is involved in circadian rhythms, but is not significant in disease states. Loss of thyroid hormone from follicular epithelium leads to hypothyroidism typified by modest weight gain, coarse and dry skin, and constipation, but without significant electrolyte disturbances.

**PBD9 1130–1132** BP9 757–759 PBD8 1154–1157 BP8 793–795

37 A Prolactinoma is the most common tumor of the anterior pituitary. Macroadenomas produce homonymous hemianopsia and can secrete prolactin to cause gynecomastia. A carcinoid tumor can produce various hormones, but not prolactin. Medullary thyroid carcinomas can produce calcitonin, which has a minimal effect on calcium homeostasis. Pheochromocytomas can produce excess catecholamines, most often manifested by hypertension. Renal cell carcinomas
may produce various paraneoplastic syndromes, most often polycythemia, hypercalcemia, and Cushing syndrome, but not hyperprolactinemia. Likewise, small cell anaplastic (oat cell) lung cancers of neuroendocrine origin can produce paraneoplastic syndromes, most often Cushing syndrome and the syndrome of inappropriate antidiuretic hormone secretion.


38 E The child has trisomy 21 (Down syndrome) and has developed acute leukemia as a complication. The 45,X karyotype is seen in Turner syndrome only in females. The normal 46,XY karyotype is unlikely with the constellation of anomalies present in this case. Trisomies 13 and 18 are far less likely than trisomy 21 to be associated with long-term survival; affected children are more likely to have severe anomalies. The 47,XXY karyotype of Klinefelter syndrome is associated with nearly normal–appearing males of normal intelligence.

PBD9 161–163 BP9 237–238 PBD8 161–163 BP8 244

39 C About 10% of individuals with autosomal dominant polycystic kidney disease (ADPKD) have a berry aneurysm of the circle of Willis, which may be complicated by rupture and hemorrhage into the subarachnoid space. The cysts of ADPKD may appear in the liver, and rarely in the pancreas. The cysts in adult-onset medullary cystic disease are centrally located in the kidney. Although renal failure does occur in middle age, similar to ADPKD, the kidneys are small and shrunken, and there are no cysts in other organs. Abnormal renal resorption of amino acids, including cysteine, may lead to formation of cystine crystals and stones in the urine; maple syrup urine disease and severe liver disease may cause such a finding. Granulomatosis with polyangitis (ANCA-associated vasculitis) can affect multiple organs, principally the kidneys and lungs, but it does not produce cystic disease. Diabetic nephropathy includes nephrosclerosis, glomerulosclerosis, pyelonephritis, and papillary necrosis, but not cystic disease. Polyarteritis nodosa may produce small microaneurysms of arteries, typically in the kidneys, and may affect multiple organs, but cystic disease is not seen.


40 D Scurvy from vitamin C deficiency is manifested by a decrease in synthesis of collagen peptides from inadequate hydroxylation of procollagen. Diminished collagen synthesis affects bone matrix formation, vascular integrity, and epithelial function. There is bleeding into joints and soft tissues with minimal trauma. Diminished carbonic anhydrase levels are seen in some forms of osteopetrosis with risk for fracture, but not the other findings. Hemophilia A, which in some cases can be due to an acquired inhibitor of factor VIII, leads to hemorrhage into soft tissues with hemarthroses and joint deformities, but this condition is typically X-linked and unlikely to affect girls, and skin and bone are not primarily involved. Mutations in fibroblast growth factor receptor underlie some forms of dwarfism. Vitamin D deficiency could produce the bone deformities, but there is reduced calcification, and anemia and hemorrhage are not found.

PBD9 442–443 BP9 301–302 PBD8 437–438 BP8 312–313

41 A Stevens-Johnson syndrome (SJS) is a severe form of erythema multiforme that can complicate infections and drug therapy. Sulfonamides, allopurinol, phenytoin, and carbamazepine are the most likely drugs to be associated with SJS. Cytotoxic (CD8+) lymphocytes mediate SJS through epidermal cell necrosis. Eosinophils are common in allergic reactions, including drug allergies, but most of these reactions are accompanied by urticaria and erythema of short duration, without blistering or desquamation. Langerhans cells and macrophages are antigen-presenting cells in the epidermis and dermis that do not directly cause toxic damage to surrounding cells. Neutrophilic exudates are not a feature of SJS, although a leukocytoclastic vasculitis with purpura is a form of drug reaction. Natural killer cells are part of innate immunity and do not participate directly in drug reactions.

PBD9 1164–1165 BP9 853–854 PBD8 1189 BP8 839–840

42 C Letterer-Siwe disease is an acute disseminated form of Langerhans cell histiocytosis. Acute lymphoblastic leukemia generally produces an elevated WBC count, and cutaneous and skeletal manifestations are rare. Gaucher disease, an autosomal recessive condition resulting from diminished glucocerebrosidase activity, has cells with cytoplasm that resembles crinkled tissue paper, and the course is not as aggressive. Leishmania donovani can cause visceral leishmaniasis, but the amastigotes infiltrating the marrow are not positive for CD1a, and the course is usually not as aggressive. Multiple myeloma is seen in older adults, and although lytic bone lesions are common, they are caused by infiltrates of plasma cells. Forms of myelodysplastic syndrome are seen in older adults, with myeloid cells that do not mark for CD1a but are accompanied by numerous ringed sideroblasts in the marrow; some cases progress to acute myelocytic leukemia.

PBD9 621–622 BP9 449 PBD8 631–632 BP8 467–468

43 C Myositis ossificans is an uncommon, exuberant repair reaction following soft-tissue trauma to muscle in which there is metaplastic bone formation. The keys to diagnosis are the location within soft tissue, calcification beginning at the periphery, and decrease in size over time. Gouty tophi can form in soft tissues, but there is typically a history of gouty arthritis first, and the lesions do not calcify. A hemarthrosis forms with joint trauma and hemorrhage in and around the joint capsule, but does not involve calcification. An osteochondroma is a bony exostosis projecting from bone into soft tissue. An osteosarcoma that rarely arises in soft tissue must be distinguished from myositis ossificans; the latter is characterized by the mature shell of bone, lack of enlargement, and lack of bone or soft-tissue destruction. Polymyositis involves inflammation with degeneration and regeneration of muscle fibers, but there is no mass effect and no calcification.

PBD9 38 BP9 793 PBD8 1251 BP8 833 AP3 Figs. 17-88, 17-91

44 E In “classic” polyarteritis nodosa, HBsAg is positive in about one third of cases. The mesenteric artery angiogram reveals focal distal occlusions and microaneurysms of branches of the superior mesenteric artery. Antimitochondrial antibody is seen in primary biliary cirrhosis. ANCA
is associated with “microscopic” polyarteritis, but not with classic polyarteritis. The ANA test result is positive in a wide variety of autoimmune diseases, principally systemic lupus erythematosus, which can affect the kidney with glomerulonephritis, not typically vasculitis. Cryptococcal antigen can be detected in CSF of patients with meningitis, usually immunocompromised patients. Histoplasma capsulatum antibody may be detected in individuals with prior exposure to this agent, which mainly causes pulmonary disease.

48 D His pneumoconiosis (with progressive massive fibrosis) is producing restrictive lung disease. Silicosis occurs from prolonged and extensive exposure to inorganic dusts in occupations such as mining. Carbon monoxide poisoning produces hypoxemia without pathologic changes to lung tissues. Fungal hyphae may produce either an allergic response or a hypersensitivity pneumonitis. Plant pollens are most often associated with episodic atopic asthma. Sulfur dioxide contributes to chronic bronchitis. Wood dusts tend to produce bronchoconstriction.

49 F Mixed connective tissue disease has overlapping features of systemic lupus erythematosus (SLE), scleroderma, polymyositis, and rheumatoid arthritis. ANCA is most likely to be seen in vasculitides such as ANCA-associated granulomatous vasculitis or microscopic polyarteritis. Antibodies to cyclic citrullinated peptide (anti-CCP) have greater than 99% specificity for rheumatoid arthritis, and these patients are more likely to have severe disease. Antihistone antibodies are most characteristic of drug-induced SLE. Anti-Smith antibodies are very specific for SLE. Thyroid peroxidase antibodies are seen in autoimmune thyroid disorders, such as Hashimoto thyroiditis and Graves disease.

50 D This family has fragile X syndrome. Anticipation occurred, with premutations of limited triple repeat expansions present in the first two generations, whereas later generations had full mutations with larger CGG expansions. The FMR1 gene encodes for familial mental retardation protein (FMRP) expressed most abundantly in brain and testis, accounting for macro-orchidism in the latter. Absence of this protein in brain leads to increased mRNA translation that affects synaptic junctions and their function. The FMRP protein is widely expressed in other tissues such as those listed, but their function is not as markedly affected as brain and testis.

51 B Thromboembolic events in women of later reproductive years, and the hepatic adenoma shown here, are complications of oral contraceptive use. Note the green color (biliverdin) indicating that this neoplasm, derived from hepatocytes, is producing bile. Allopurinol and sulfonamides are associated with hepatic granuloma formation. Aspirin use in children has been associated with Reye syndrome with microvesicular steatosis. Isoniazid has been associated with acute or chronic hepatitis. Phenacetin and acetaminophen
use have been associated with analgesic nephropathy, and excessive acetaminophen use may produce acute massive hepatic necrosis.

**PBD9 867–869**  **BP9 636**  **PBD8 415**  **BP8 293–294**

**52 D** Risk factors for HELLP (hemolysis with elevated liver enzymes and low platelets) syndrome, a variant of severe preeclampsia, include nulliparity, advanced maternal age, diabetes mellitus, preexisting hypertension, a prior history of preeclampsia, and renal disease. Patients with HELLP syndrome may progress to disseminated intravascular coagulation. Emergent delivery is indicated. Abruptio placentae is an acute event marked by severe abdominal pain and vaginal bleeding. Hepatic vein thrombosis in Budd-Chiari syndrome can produce liver necrosis with elevated enzymes; pregnancy is a risk factor, but this does not explain the neurologic and renal findings. Likewise, a dilated cardiomyopathy that can occur in pregnancy does not explain these findings. In hydatidiform mole, preeclampsia is more likely, but 30 weeks is a long time to have it, and a fetus would not be present. Acute fatty liver of pregnancy may resemble Reye syndrome (a disease that occurs in children) and may be preceded by preeclampsia. Sheehan syndrome is postpartum pituitary necrosis leading to hypopituitarism.

**PBD9 1037–1039**  **BP9 703–704**  **PBD8 685**  **BP8 737–738**

**53 E** The findings are those of infectious endocarditis, and the needle track in the left arm suggests injection drug use as the risk factor. These individuals can have right-sided and/or left-sided valvular lesions. The vegetations are likely to embolize, and the septic emboli can lead to infection or infarction of multiple organs from left-sided lesions and of the lungs from right-sided lesions. The pulmonary nodule is likely a lung abscess. The patient probably has acute pyelonephritis from hematogenous infection. *Candida albicans* and *Cryptococcus neoformans* may cause endocarditis in immunocompromised individuals. *Escherichia coli* is a likely cause of ascending urinary tract infections with pyelonephritis, but it is an uncommon cause of endocarditis. Listeriosis most often results from food or water contamination and can lead to sepsis with meningitis, but rarely endocarditis. Streptococcal infections are more likely to cause endocarditis in individuals with preexisting valvular heart disease, and viridans streptococci are most often implicated. *Yersinia enterocolitica* can produce enterocolitis, not endocarditis (this organism can persist in stored blood and cause transfusion-related sepsis).

**PBD9 559–561**  **BP9 392–394**  **PBD8 567–568**  **BP8 406–407**

**54 E** Thrombotic thrombocytopenic purpura (TTP) can present with the classic pentad of neurologic changes, fever, thrombocytopenia, microangiopathic hemolytic anemia, and decreased renal function. The pathogenesis of TTP is related to von Willebrand factor (vWF). Monomers of vWF are linked by disulfide bonds to form multimers with various molecular masses that range to millions of daltons. A vWF-cleaving metalloproteinase in plasma normally prevents the entrance into the circulation (or persistence) of unusually large multimers of vWF. The metalloproteinase is referred to as ADAMTS 13 (a disintegrin and metalloproteinase, with thrombospondin 1-like domains). In most patients with TTP, plasma ADAMTS 13 activity is less than 5% of normal. The patient’s hemoglobin concentration is not low enough to justify transfusion of RBCs. Giving platelets to a patient with TTP would “add fuel to the fire” because the platelets would cause more thrombi to form, resulting in further organ damage. Simple pressor agents, such as dobutamine, are not primary therapy. Surgery is not indicated because the injury is occurring in the small vasculature of many organs. Prednisone may be given to a subset of patients who do not respond to plasmapheresis.

**PBD9 659–660**  **BP9 453–454**  **PBD8 669–670**  **BP8 472–473**

**55 C** Budd-Chiari syndrome is a rare condition that can complicate pregnancy or the postpartum state. Hepatic venous occlusion leads to hepatomegaly with severe centrilobular congestion and necrosis, much more pronounced than the typical nutmeg liver of chronic passive congestion with right-sided heart failure. Biliary tract obstruction with choledocholithiasis would increase the serum bilirubin to a greater degree than seen in this patient, and hepatomegaly is unlikely. Hepatic adenomas, which can be associated with use of oral contraceptives, are mass lesions, usually several centimeters in size. At 12 weeks’ gestation, with a fetus present, choriocarcinoma is very unlikely, and a marked increase in liver enzymes is unlikely. Acute fatty liver of pregnancy with microvesicular steatosis produces a more uniform density with hepatomegaly, a rare condition that is usually seen in the third trimester of pregnancy.

**PBD9 863–864**  **BP9 634**  **PBD8 872–873**  **BP8 662**

**56 B** In common variable immunodeficiency (CVID), there are normal numbers of T cells with normal to low numbers of B cells, and there is hypogammaglobulinemia with decreased IgG and possibly other immunoglobulin types. CVID occurs in young adults of both sexes, causing increased bacterial infections and giardiasis and recurrent herpes simplex (and herpes zoster) infections. The mechanisms are diverse and include failure of B-cell maturation to plasma cells, excessive T-cell suppression, and defective T helper cell function. Mutations in genes encoding NADPH oxidase proteins produce chronic granulomatous disease and recurrent infections with *Aspergillus*, *Staphylococcus*, *Serratia*, *Nocardia*, and *Pseudomonas*. In hyper-IgM syndrome, mutations in the CD40 ligand induce failure in B cells, with low IgG, IgA, and IgE levels, but increased IgM; in infancy and childhood, there is increased risk of severe infections with bacterial and viral agents and opportunistic agents, such as *Pneumocystis jiroveci*. Mutations in CD18, the common β chain of integrins, which aid in binding of leukocytes, lead to leukocyte adhesion deficiency with leukocytosis, but with absence of supplicative inflammation in areas of tissue necrosis and ulceration caused by *Staphylococcus aureus* and gram-negative enteric bacteria. In severe combined immunodeficiency (SCID), half of cases result from an X-linked mutation for the common γ chain of IL-2, a receptor for many cytokines needed for T-cell development, and the other half of cases
result from autosomal recessive mutation in adenosine deaminase with accumulation of purine metabolites toxic to T cells. In SCID, although T cells are primarily involved, there is secondary impairment of B-cell function, so that affected individuals have diminished IgG levels and no IgA or IgM, and decreased T-cell function, resulting in increased susceptibility to virtually all infectious organisms.

**PBD9 241–242** BP9 141  BPDB 233  BP8 152–153

**57 F** Myotonic dystrophy is a form of muscular dystrophy in which there are increased CTG trinucleotide repeat sequences in the gene. Weakness in skeletal, cardiac, and smooth muscle develops, along with cata racts, dementia, gonadal atrophy, and hypogammaglobulinemia. Deficiency of α-1,4-glucosidase leads to Pompe disease in infancy. Absence of dystrophin characterizes Duchenne muscular dystrophy, which affects young boys. The FGFR3 mutation is seen in achondroplasia, a form of short-limbed dwarfism. Mutations in the mitochondrial oxidative phosphorylase genes (mitochondrial myopathies) can produce neurodegenerative disorders and hearing loss, in addition to myopathy. Myophosphorylase is diminished in McArdle disease, which causes muscle pain on strenuous exercise.

**PBD9 1244** BP9 804  BPDB 1269–1270  BP8 828–829

**58 E** Both the ethanol and the immobilization with reduced local blood flow predispose to rhabdomyolysis from skeletal muscle injury. Myoglobin is released from the injured muscle cells, and it is toxic to renal tubules and can lead to toxic acute tubular necrosis. Cystine crystals are rare and may represent cystinuria or severe liver disease. Glucosuria is indicative of diabetes mellitus. Hyaline casts can be found in healthy persons, and large numbers suggest reduced urine output. Ketonuria occurs when intake or use of carbohydrates is reduced.

**PBD9 927–929** BP9 806  BPDB 936–938  BP8 564–566

**59 D** These findings strongly suggest a ruptured ectopic pregnancy—hence human chorionic gonadotropin levels would be increased. Gonococcal and chlamydial infections are risk factors for pelvic inflammatory disease, which increases the risk of ectopic pregnancy. A ruptured ectopic pregnancy may lead to disseminated intravascular coagulation with increased partial thromboplastin time and prothrombin time, but the normal platelet count in this patient suggests that this has not yet occurred. Factor XIII, which stabilizes fibrin clots, can be deficient, but this is extremely rare. Bleeding is first observed in the neonatal period; older patients may have poor wound healing, intracerebral hemorrhage, infertility (men), and abortion (women). Decreased follicle-stimulating hormone is typical of pituitary failure, which does not explain the bleeding. An increase in carcinoembryonic antigen is seen in some gastrointestinal tract neoplasms, but this patient is young to have such malignancies. Amebiasis would produce a bloody diarrhea, and perforation of the colon is uncommon. Schistosomiasis resulting from *Schistosoma haematobium* can produce hematuria, but the bladder is not perforated.

**PBD9 368, 383, 1036** BP9 701  BPDB 1053–1054  BP8 734–735

**60 D** The figure shows a mass that is typical of adenocarcinoma of the ascending colon. Such cancers are unlikely to obstruct, but they can bleed a small amount over months to years, causing iron deficiency anemia. This relatively young woman has evidence for an additional cancer, an endometrial cancer, and this combination is most likely due to an inherited mutation in one of the DNA mismatch-repair genes, such as *MSH2* and *MLH1*. Homozygous loss of these genes can give rise to right-sided colon cancer and endometrial cancer. Such a mutation is typically associated with microsatellite instability. In contrast the *APC* gene, a negative regulator of β-catenin in the WNT signaling pathway, is associated with familial adenomatous polyposis syndrome and most sporadic colon cancers. This latter pathway also is known as the adenoma-carcinoma sequence, because the carcinomas develop through an identifiable series of molecular and morphologic steps. Loss of the *PTEN* tumor-suppressor gene is seen in endometrial carcinomas not associated with colon carcinoma and with some hamartomatous polyps of the colon. Infection with some strains of human papillomavirus leads to chronic protein inactivation and development of cervical carcinoma. Mutation with activation of c-KIT tyrosine kinase activity occurs in gastrointestinal stromal tumors, which respond well to treatment with imatinib mesylate, a tyrosine kinase inhibitor also used to treat chronic myelogenous leukemia.

**PBD9 810–814** BP9 597–600  BPDB 822–825  BP8 622–624

**61 A** Was the death of Napoleon Bonaparte the result of arsenic poisoning perpetrated by former enemies? The FBI analyzed a hair sample in 1995 and found increased arsenic levels, but the amount was equivocal. Another hypothesis is that the wallpaper at Longwood House, where he lived, contained copper arsenate and became moldy, releasing arsenic vapor. A third hypothesis suggests that he died of the effects of gastric cancer, not related to arsenic ingestion, substantiated by an observation at autopsy of lymphadenopathy adjacent to the stomach. Chronic arsenic exposure has been associated with a greatly elevated risk of skin cancer and possibly of cancers of the lung, liver (angiosarcoma), bladder, kidney, and colon. Beryllium acutely produces a pneumonitis, and long-term exposure leads to nonnecrotizing granulomas, similar to sarcoidosis. Chromium or nickel exposure may lead to respiratory tract cancers. Cobalt poisoning can produce pulmonary interstitial fibrosis. Lead poisoning can produce abdominal pain, anemia, neuropathy, and decreased mental sharpness, but it is not associated with malignancies.

**PBD9 412–413, 1155** BP9 275–276, 863  BPDB 408  BP8 285

**62 E** This woman has features seen in individuals who are bedridden for extended periods. There is a large saddle thromboembolus in the pulmonary arterial trunk. Antiphospholipid syndrome can include thromboembolic events, but osteoporosis, decubitus ulcers, and muscle wasting are not features. Aplastic anemia leads to high-output congestive heart failure, bleeding diathesis from thrombocytopenia, and risk of infections. Chronic alcoholism produces chronic liver disease that predisposes to bleeding problems, not thrombosis. Blunt trauma is marked by contusions and fractures. Malnutrition could account for decreased bone density from osteomalacia, poor wound healing with skin ulceration, and...
muscle wasting, but coagulopathy with bleeding is more likely to occur than thromboembolism.

63 C The combination of nongonococcal urethritis, arthritis, and conjunctivitis suggests reactive arthritis, one of the spondyloarthropathies; the changes in the spine can resemble ankylosing spondylitis and can be equally debilitating. ANCA is indicative of various forms of vasculitis, such as granulomatous vasculitis and microscopic polyangiitis. The ANA test result is positive in many autoimmune diseases, such as systemic lupus erythematosus (SLE), but it is not a feature of spondyloarthropathies. Lyme disease can include large joint arthritis, but not urethritis or conjunctivitis. Rapid plasma reagin (RPR) is a screening test for syphilis, which can include arthritis of large joints (Charcot joint) in the tertiary form, but it takes decades to develop.

Rheumatoid arthritis is a feature of rheumatoid arthritis, which initially manifests more commonly in small joints of the hands and feet. U1-RNP is a marker for mixed connective tissue disease, which has features of rheumatoid arthritis, scleroderma, polymyositis, and SLE; arthralgias are not accompanied by joint destruction or deformity.

64 E Von Hippel–Lindau disease is rare and results from acquired or inherited mutation in a tumor suppressor gene. The neoplasms in this case include, in order, retinal angiomas, adrenal phaeochromocytoma, cerebellar hemangioblastomas, and renal cell carcinomas producing erythropoietin. Gardner syndrome has the same mutation in the adenomatous polyposis coli (APC) gene as familial polyposis, but also has osteomas, epidermal cysts, and fibromatoses. The MET oncogene is mutated in papillary renal carcinomas (not associated with other tumors elsewhere) and in Denys-Drash syndrome, which also includes Wilms tumor plus gonadal dysgenesis and nephropathy. Neurofibromatosis type 2 includes schwannomas, meningiomas, gliomas, and ependymomas. Tuberculous sclerosis is one of the phakomatoses associated with cortical hamartomas called tubers, renal angiomyolipomas, cardiac rhabdomyomas, and subungual fibromas. Beckwith-Wiedemann syndrome includes Wilms tumor, hemihypertrophy, and adrenal cytomegaly.

65 A Limited scleroderma (former CREST syndrome), a form of systemic sclerosis, does not progress to include serious pulmonary fibrosis or renal disease. Therefore she is less likely to have diffuse scleroderma, which is associated with the anti–DNA topoisomerase antibody. Antimicrobial (anti–thyroid peroxidase) antibodies are seen in autoimmune thyroid diseases, such as Hashimoto thyroiditis and Graves disease. Antimitochondrial antibody appears most frequently in primary biliary cirrhosis. ANCA can be a marker for various forms of vasculitis. Anti-transglutaminase antibodies are seen in celiac disease, which is marked by malabsorption, not esophageal dysmotility.

66 B Pneumocystis jiroveci pneumonia is shown with Gomori methenamine silver stain. The spectrum of opportunistic infections, wasting syndrome, and lymphopenia suggest AIDS complicating HIV infection. This spectrum of findings is not typical of patients with systemic lupus erythematosus (SLE), who have a positive ANA test result. When SLE is treated with immunosuppressive therapy, opportunistic infections are common. SLE-like disease is seen in C2 deficiency. B-cell function tends to be preserved in HIV infection, so marked hypogammaglobulinemia is unlikely to be present. The neutrophil oxidative burst assay is used to test for chronic granulomatous disease, an immunodeficiency condition in which bacterial infections are more likely to appear in children. Rapid plasma reagin (RPR) is a screening test for syphilis, which is a sexually transmitted disease that is not accompanied by immunodeficiency.

67 A Metabolic acidosis, tinnitus, platelet function abnormalities, and gastritis with ulceration are typical abnormalities of excessive aspirin ingestion. Acetaminophen in large quantities may produce hepatotoxicity. Adalimumab, a monoclonal antibody directed against tumor necrosis factor, and methotrexate are drugs used to treat rheumatoid arthritis and do not have antplatelet effects. Oxycodone is an opiate, and propoxyphene is a nonnarcotic analgesic; these drugs do not have significant effects on the gastrointestinal tract or on platelets.

68 B Choriocarcinoma is the most aggressive, and the least common, form of gestational trophoblastic disease. It can metastasize widely, particularly to the lungs and vagina, and also to the brain, liver, and kidney. The neoplasm is composed of a malignant-appearing syncytiotrophoblast and forms a soft, hemorrhagic mass that can rupture and bleed. In addition, the amount of human chorionic gonadotropin (hCG) produced by a choriocarcinoma is marked; hCG shares the same α-subunit as other glycoprotein hormones, such as thyroid-stimulating hormone (TSH), and may enhance the effect of TSH, leading to features of hyperthyroidism. Many choriocarcinomas respond to chemotherapy. Endometrial adenocarcinoma is most often seen in postmenopausal women. Clear cell carcinomas of the vagina most often appear in young women whose mothers were given diethylstilbestrol (DES) during pregnancy. A leiomyosarcoma is an uncommon lesion in women and usually produces a large uterine mass. Likewise, a malignant mixed müllerian tumor is a rare neoplasm seen in women. Sarcoma botryoides is a neoplasm found in girls younger than 5 years.

69 C She has rheumatoid arthritis with anemia of chronic disease. Despite abundant stored iron stores, such anemias are caused by impaired transfer of iron from macrophages to developing erythroid cells. Hepcidin is synthesized in the liver and normally released in response to increased intrahepatic iron levels. Hepcidin inhibits ferroportin function in macrophages and thus prevents transfer of iron. Hepcidin...
production in the liver is increased in chronic inflammatory states by the action of inflammatory mediators such as IL-6. Rheumatoid factor is likely present but does not affect hepcidin levels. TNF plays a role in the joint inflammation. GM-CSF can cause increased production of normal macrophages.

70 B Rickets in children (and osteomalacia in adults) results from vitamin D deficiency. The left figure panel shows a costochondral junction in which the palisade of cartilage is lost, with widened irregular trabeculae with uncalcified osteoid. Compare with a normal costochondral junction in the right panel with orderly transition from cartilage to new bone. Exposure to sunlight is essential for synthesis of vitamin D from endogenous sources. Sufficient vitamin D reduces the risk for cancer, inflammatory conditions, and atherosclerosis. Thus the sage advice for children: go outside and be active. Lack of dietary fresh fruits and vegetables can lead to scurvy from vitamin C deficiency, with poor osteoid formation, but other connective tissues would also be affected. Hypopituitarism could lead to reduced growth hormone with reduced stature but no deformities. Mutations affecting collagen genes could lead to osteogenesis imperfecta with risk for fractures. Trauma leads to fractures, and the callus of healing fractures has an orderly process of ossification. Polymorphisms of vitamin D receptor affect can be involved in the pathogenesis of osteoporosis.

71 D The sudden nature of the death requires investigation by the medical examiner (coroner), who would notify the local law enforcement agency on discovery of the findings in this case. The pattern of injuries is consistent with vigorous shaking of the infant. Because an infant’s head is larger in proportion to its body compared with an adult, it cannot counter the shaking with neck musculature. Grasping the arms of the infant strongly and pressing or hitting the head against a hard surface increases the risk of internal injuries, including fractures. Another feature of trauma with shaking of the infant’s head is diffuse axonal injury with axonal retraction balls from marked stretching and tearing with acceleration-deceleration forces. Congenital syphilis produces osteochondritis with skeletal deformities, not fractures, and without a bleeding tendency. Hemophilia A could account for hemorrhages in a child or adult, but not fractures. Osteogenesis imperfecta can account for fractures, but not hemorrhages. In sudden infant death syndrome, there would be no signs of trauma. Thalassemic dysplasia is a lethal form of short-limbed dwarfism.

72 A He has features of Marfan syndrome, and although the primary defect is mutation in the fibrillin-1 gene and hence poor formation of microfibrils, there is an important secondary effect on transforming growth factor beta (TGF-β) bioavailability. Normally microfibrils sequester TGF-β, but in Marfan syndrome, abnormal microfibrils allow excessive TGF-β signaling, which is responsible for the cardiac symptoms. Bone marrow transplants are useful in those diseases where the primary defect is in the hematopoietic cells, or the marrow cells can deliver normal genes to the affected tissues. Even if one could transduce the fibrillin-1 gene into T cells, the protein cannot be delivered to the extracellular matrix. Marfan syndrome does not have abnormal lysosomes, and the vascular disorders are not the result of atherosclerosis accelerated by dietary factors.

73 E Vitamin A deficiency leads to epithelial disorders affecting the cornea, skin, respiratory tract, and urinary tract. It is the leading cause for preventable blindness worldwide. Squamous metaplasia (shown) in the respiratory tract increases the risk of infection; desquamation of keratin debris forms the nidus of urinary tract calculi. Hyperkeratosis and follicular plugging affect the epidermis. Cystic fibrosis leads to an increased risk of respiratory tract infections, particularly infections caused by Pseudomonas, from widespread bronchiectasis; the skin and eye are not affected. Congenital syphilis can produce bone deformities and gummas. HIV infection can be complicated by opportunistic infections, including infections of the respiratory tract, but keratolacia is not a feature of HIV infection. Kartagener syndrome can lead to bronchiectasis from an altered respiratory tract epithelium in which the ciliary dynein arms are absent; situs inversus is present, but not eye and skin changes.

74 E The pattern of metastases with seeding of the peritoneal cavity along with the microscopic and serum tumor markers are most characteristic for an ovarian serous cystadenocarcinoma, the most common malignant neoplasm arising in the ovary. Ileal adenocarcinomas are rare and would probably lead to bowel obstruction. Carcinoid tumors are unlikely to be widely metastatic. Choriocarcinoma is associated with high hCG levels and hemorrhagic metastases. A malignant mesothelioma is a rare complication of asbestos. It is unlikely that a cystadenocarcinoma would recur 10 years later. Cystadenomas are benign.

75 D We go back to the future in the thirteenth century for this answer from Rumi of Balkh, and it is applicable for any time and place. The art of medicine is more than science, and relies upon human interaction. The concept of empathy is part of this. The authors have enough medical conditions to know its importance and are passing that understanding on to you. Do not let it go. You need it for your family, your friends, and your patients.

76 (All are correct) We hope that you have advanced your knowledge and are now better able to help others in your health science career. Go make the world a better place for everyone!
Chapter 5


FIGURE 5-18: Courtesy Department of Pathology, University of Pittsburgh Medical Center, Pittsburgh.

FIGURE 5-26: Courtesy Dr. Vijay Tonk, Department of Pathology, The University of Texas Southwestern Medical Center, Dallas.

FIGURE 5-28: Courtesy Dr. Nancy R. Schneider and Jeff Doolittle, Cytogenetics Laboratory, The University of Texas Southwestern Medical Center, Dallas.

Chapter 6

FIGURE 6-21: Courtesy Dr. Richard Sontheimer, Department of Dermatology, The University of Texas Southwestern Medical School, Dallas.

FIGURE 6-31: Courtesy Dr. Trace Worrell, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 8

FIGURE 8-16: Courtesy Dr. Arlene Sharpe, Brigham and Women’s Hospital, Boston.

FIGURE 8-60: Courtesy Dr. Willy Pressens, Harvard School of Public Health, Boston.

Chapter 9

FIGURE 9-25: Courtesy George Katsas, MD, forensic pathologist, Boston.

FIGURE 9-27: From the teaching collection of the Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 11

FIGURE 11-20: From the teaching collection of the Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

FIGURE 11-22: Courtesy Tom Rogers, MD, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

FIGURE 11-47: Courtesy Tom Rogers, MD, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 12

FIGURE 12-10: Courtesy Arthur Weinberg, MD, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 13

FIGURE 13-53: Courtesy Dr. George Murphy, University of Pennsylvania Perelman School of Medicine, Philadelphia.

Chapter 15

FIGURE 15-41: From the teaching collection of the Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 16


Chapter 17

FIGURE 17-64: Courtesy Dr. Tad Wieczorek, Brigham and Women’s Hospital, Boston.

Chapter 20

FIGURE 20-7: Courtesy Dr. M.A. Ventkatachalam, Department of Pathology, The University of Texas Health Sciences Center, San Antonio.

FIGURE 20-15: Courtesy Dr. Helmut Rennke, Department of Pathology, Brigham and Women’s Hospital, Boston.

FIGURE 20-40: Courtesy Dr. M.A. Ventkatachalam, Department of Pathology, The University of Texas Health Sciences Center, San Antonio.

Chapter 21

FIGURE 21-8: Courtesy Dr. Christopher Corless, University of Oregon, Eugene.

Chapter 22

FIGURE 22-7: Courtesy Dr. Jag Bhawan, Boston University School of Medicine, Boston.

FIGURE 22-41: Courtesy Dr. Christopher Crum, Brigham and Women’s Hospital, Boston.

Chapter 23

FIGURE 23-26: Courtesy Dr. Jack G. Meyer, Brigham and Women’s Hospital, Boston.
Chapter 28

FIGURE 28-55: Courtesy Eileen Bigio, MD, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

FIGURE 28-59: Courtesy Eileen Bigio, MD, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.

Chapter 30

FIGURE 30-62: Courtesy Dr. Linda Margraf, Department of Pathology, The University of Texas Southwestern Medical School, Dallas.
<table>
<thead>
<tr>
<th>NORMAL VALUES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BLOOD: PLASMA, SERUM</strong></td>
</tr>
<tr>
<td>Alanine aminotransferase (ALT), serum</td>
</tr>
<tr>
<td>Albumin</td>
</tr>
<tr>
<td>Alkaline phosphatase, serum (adult)</td>
</tr>
<tr>
<td>Amylase, serum (adult)</td>
</tr>
<tr>
<td>Aspartate aminotransferase (AST), serum</td>
</tr>
<tr>
<td>Bilirubin, serum (adult) total // direct</td>
</tr>
<tr>
<td>Calcium, serum (Ca++)</td>
</tr>
<tr>
<td>Cholesterol, total // HDL</td>
</tr>
<tr>
<td>Creatinine, serum</td>
</tr>
<tr>
<td>D-dimer</td>
</tr>
<tr>
<td>Electrolytes, serum</td>
</tr>
<tr>
<td>Sodium (Na+)</td>
</tr>
<tr>
<td>Potassium (K+)</td>
</tr>
<tr>
<td>Chloride (Cl-)</td>
</tr>
<tr>
<td>HCO3</td>
</tr>
<tr>
<td>Anion gap</td>
</tr>
<tr>
<td>Gases, arterial blood (room air)</td>
</tr>
<tr>
<td>pH</td>
</tr>
<tr>
<td>PCO2</td>
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<tr>
<td>PO2</td>
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<tr>
<td>HCO3</td>
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<tr>
<td>Globulin</td>
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<tr>
<td>Haptoglobin</td>
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<tr>
<td>Hemoglobin A1c</td>
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<tr>
<td>Iron, serum</td>
</tr>
<tr>
<td>Iron binding capacity, total (TIBC)</td>
</tr>
<tr>
<td>Lactate dehydrogenase (LDH), serum</td>
</tr>
<tr>
<td>Lipase, serum</td>
</tr>
<tr>
<td>Phosphorus (inorganic), serum</td>
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<tr>
<td>Proteins, serum, total</td>
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<tr>
<td>Thyroid stimulating hormone (TSH)</td>
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<tr>
<td>Thyroxine (T4), serum</td>
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<tr>
<td>Triglycerides, serum</td>
</tr>
<tr>
<td>Urea nitrogen, serum</td>
</tr>
<tr>
<td>Uric acid, serum</td>
</tr>
<tr>
<td><strong>HEMATOLOGIC</strong></td>
</tr>
<tr>
<td>Erythrocyte (RBC) count</td>
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<tr>
<td></td>
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<tr>
<td>Erythrocyte sedimentation rate (ESR)</td>
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<tr>
<td></td>
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<tr>
<td>Hematocrit</td>
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<td></td>
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<tr>
<td>Hemoglobin, blood</td>
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<tr>
<td></td>
</tr>
<tr>
<td>Leukocyte (WBC) count, total</td>
</tr>
<tr>
<td>Segmented neutrophils</td>
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<tr>
<td>Bands</td>
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<tr>
<td>Eosinophils</td>
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<tr>
<td>Basophils</td>
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<tr>
<td>Lymphocytes</td>
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<tr>
<td>Monocytes</td>
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<tr>
<td>Mean corpuscular hemoglobin (MCH)</td>
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<tr>
<td>Mean corpuscular volume (MCV)</td>
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<tr>
<td>Partial thromboplastin time, activated (aPTT)</td>
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<tr>
<td>Platelet count</td>
</tr>
<tr>
<td>Prothrombin time (PT)</td>
</tr>
<tr>
<td><strong>BODY MASS INDEX (BMI)</strong></td>
</tr>
</tbody>
</table>