MICROBIOLOGY
VOL. II, 3E

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Notice
The authors of this volume have taken care that the information contained herein is accurate and compatible with the standards generally accepted at the time of publication. Nevertheless, it is difficult to ensure that all the information given is entirely accurate for all circumstances. The publisher and authors do not guarantee the contents of this book and disclaim any liability, loss, or damage incurred as a consequence, directly or indirectly, of the use and application of any of the contents of this volume.
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ACKNOWLEDGMENTS

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Please let us know if your name has been misspelled or mispelled and we will be happy to make the update in the next edition.
PREFACE TO THE 3RD EDITION

We were very pleased with the overwhelmingly positive student feedback for the 2nd edition of our *Underground Clinical Vignettes* series. Well over 100,000 copies of the UCV books are in print and have been used by students all over the world.

Over the last two years we have accumulated and incorporated over a thousand *updates* and improvements suggested by you, our readers, including:

- many additions of specific boards and wards testable content
- deletions of redundant and overlapping cases
- reordering and reorganization of all cases in both series
- a new master index by case name in each Atlas
- correction of a few factual errors
- diagnosis and treatment updates
- addition of 5–20 new cases in every book
- and the addition of clinical exam photographs within *UCV—Anatomy*

And most important of all, the third edition sets now include two brand new *COLOR ATLAS* supplements, one for each Clinical Vignette series.

- The *UCV-Basic Science Color Atlas (Step 1)* includes over 250 color plates, divided into gross pathology, microscopic pathology (histology), hematology, and microbiology (smears).
- The *UCV-Clinical Science Color Atlas (Step 2)* has over 125 color plates, including patient images, dermatology, and funduscopy.

Each atlas image is descriptively captioned and linked to its corresponding Step 1 case, Step 2 case, and/or Step 2 MiniCase.
How Atlas Links Work:

Step 1 Book Codes are:
- A = Anatomy
- BS = Behavioral Science
- BG = Biochemistry
- M1 = Microbiology, Vol. I
- M2 = Microbiology, Vol. II
- P1 = Pathophysiology, Vol. I
- P2 = Pathophysiology, Vol. II
- PH = Pharmacology

Step 2 Book Codes are:
- ER = Emergency Medicine
- IM1 = Internal Medicine, Vol. I
- IM2 = Internal Medicine, Vol. II
- NEU = Neurology
- OB = OB/GYN
- PED = Pediatrics
- SUR = Surgery
- PSY = Psychiatry
- MC = MiniCase

Case Number

Indicates Type of Images:
- H = Hematology
- M = Microbiology
- PG = Gross Pathology
- PM = Microscopic Pathology

Indicates UCV1 or UCV2 Series

- If the Case number (032, 035, etc.) is not followed by a letter, then there is only one image. Otherwise A, B, C, D indicate up to 4 images.

Bold Faced Links: In order to give you access to the largest number of images possible, we have chosen to crosslink the Step 1 and 2 series.

- If the link is bold-faced this indicates that the link is direct (i.e., Step 1 Case with the Basic Science Step 1 Atlas link).
- If the link is not bold-faced this indicates that the link is indirect (Step 1 case with Clinical Science Step 2 Atlas link or vice versa).

We have also implemented a few structural changes upon your request:

- Each current and future edition of our popular First Aid for the USMLE Step 1 (Appleton & Lange/McGraw-Hill) and First Aid for the USMLE Step 2 (Appleton & Lange/McGraw-Hill) book will be linked to the corresponding UCV case.
- We eliminated UCV → First Aid links as they frequently become out of date, as the First Aid books are revised yearly.
• The Color Atlas is also specially designed for quizzing—
captions are descriptive and do not give away the case name
directly.

We hope the updated UCV series will remain a unique and well-
integrated study tool that provides compact clinical correlations
to basic science information. They are designed to be easy and
fun (comparatively) to read, and helpful for both licensing
exams and the wards.

We invite your corrections and suggestions for the fourth edi-
tion of these books. For the first submission of each factual cor-
rection or new vignette that is selected for inclusion in the
fourth edition, you will receive a personal acknowledgment in
the revised book. If you submit over 20 high-quality corrections,
additions or new vignettes we will also consider inviting you to
become a "Contributor" on the book of your choice. If you are
interested in becoming a potential "Contributor" or "Author"
on a future UCV book, or working with our team in developing
additional books, please also e-mail us your CV/resume.

We prefer that you submit corrections or suggestions via
electronic mail to UCVteam@yahoo.com. Please include
"Underground Vignettes" as the subject of your message. If you
do not have access to e-mail, use the following mailing address:
Blackwell Publishing, Attn: UCV Editors, 350 Main Street,
Malden, MA 02148, USA.

Vikas Bhushan
Vishal Pall
Tien Le
October 2001
This series was originally developed to address the increasing number of clinical vignette questions on medical examinations, including the USMLE Step 1 and Step 2. It is also designed to supplement and complement the popular *First Aid for the USMLE Step 1* (Appleton & Lange/McGraw Hill) and *First Aid for the USMLE Step 2* (Appleton & Lange/McGraw Hill).

Each UCV 1 book uses a series of approximately 100 "supra-prototypical" cases as a way to condense testable facts and associations. The clinical vignettes in this series are designed to incorporate as many testable facts as possible into a cohesive and memorable clinical picture. The vignettes represent composites drawn from general and specialty textbooks, reference books, thousands of USMLE style questions and the personal experience of the authors and reviewers.

Although each case tends to present all the signs, symptoms, and diagnostic findings for a particular illness, patients generally will not present with such a "complete" picture either clinically or on a medical examination. Cases are not meant to simulate a potential real patient or an exam vignette. All the boldfaced "buzzwords" are for learning purposes and are not necessarily expected to be found in any one patient with the disease.

Definitions of selected important terms are placed within the vignettes in (small caps) in parentheses. Other parenthetical remarks often refer to the pathophysiology or mechanism of disease. The format should also help students learn to present cases succinctly during oral "bullet" presentations on clinical rotations. The cases are meant to serve as a condensed review, not as a primary reference. The information provided in this book has been prepared with a great deal of thought and careful research. This book should not, however, be considered as your sole source of information. Corrections, suggestions and submissions of new cases are encouraged and will be acknowledged and incorporated when appropriate in future editions.
### Abbreviations

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<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tr>
<td>5-ASA</td>
<td>5-aminosalicylic acid</td>
</tr>
<tr>
<td>ABGs</td>
<td>arterial blood gases</td>
</tr>
<tr>
<td>ABVD</td>
<td>Adriamycin/bleomycin/vincristine/dacarbazine</td>
</tr>
<tr>
<td>ACE</td>
<td>angiotensin-converting enzyme</td>
</tr>
<tr>
<td>ACTH</td>
<td>adrenocorticotropic hormone</td>
</tr>
<tr>
<td>ADH</td>
<td>antidiuretic hormone</td>
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<tr>
<td>AFP</td>
<td>alpha fetal protein</td>
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<tr>
<td>AI</td>
<td>aortic insufficiency</td>
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<td>AIDS</td>
<td>acquired immunodeficiency syndrome</td>
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<tr>
<td>ALL</td>
<td>acute lymphocytic leukemia</td>
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<tr>
<td>ALT</td>
<td>alanine transaminase</td>
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<tr>
<td>AML</td>
<td>acute myelogenous leukemia</td>
</tr>
<tr>
<td>ANA</td>
<td>antinuclear antibody</td>
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<tr>
<td>ARDS</td>
<td>adult respiratory distress syndrome</td>
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<tr>
<td>ASD</td>
<td>atrial septal defect</td>
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<tr>
<td>ASO</td>
<td>antistreptolysin O</td>
</tr>
<tr>
<td>AST</td>
<td>aspartate transaminase</td>
</tr>
<tr>
<td>AV</td>
<td>arteriovenous</td>
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<tr>
<td>BE</td>
<td>barium enema</td>
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<tr>
<td>BP</td>
<td>blood pressure</td>
</tr>
<tr>
<td>BUN</td>
<td>blood urea nitrogen</td>
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<tr>
<td>CAD</td>
<td>coronary artery disease</td>
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<tr>
<td>CALLA</td>
<td>common acute lymphoblastic leukemia antigen</td>
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<tr>
<td>CBC</td>
<td>complete blood count</td>
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<tr>
<td>CHF</td>
<td>congestive heart failure</td>
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<td>CK</td>
<td>creatine kinase</td>
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<td>CLL</td>
<td>chronic lymphocytic leukemia</td>
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<td>CML</td>
<td>chronic myelogenous leukemia</td>
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<td>cytomegalovirus</td>
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<td>central nervous system</td>
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<td>creatine phosphokinase</td>
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<td>cerebrospinal fluid</td>
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<td>computed tomography</td>
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<td>cerebrovascular accident</td>
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<td>disseminated intravascular coagulation</td>
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<td>distal interphalangeal</td>
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<td>Abbreviation</td>
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<td>electrocardiography</td>
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<td>echocardiography</td>
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<td>endoscopic retrograde cholangiopancreatography</td>
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<td>ESR</td>
<td>erythrocyte sedimentation rate</td>
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<td>forced expiratory volume</td>
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<td>FTA-ABS</td>
<td>fluorescent treponemal antibody absorption</td>
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<td>forced vital capacity</td>
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<td>glomerular filtration rate</td>
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<td>human chorionic gonadotrophin</td>
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<td>human leukocyte antigen</td>
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<td>history of present illness</td>
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<td>heart rate</td>
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<td>human rabies immune globulin</td>
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<td>ID/CC</td>
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<td>immunoglobulin</td>
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<td>insulin-like growth factor</td>
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<td>jugular venous pressure</td>
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<td>kidneys/ureter/bladder</td>
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<td>lower esophageal sphincter</td>
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<td>LFTs</td>
<td>liver function tests</td>
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<td>lumbar puncture</td>
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<td>LVH</td>
<td>left ventricular hypertrophy</td>
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<td>Lytes</td>
<td>electrolytes</td>
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<td>MCHC</td>
<td>mean corpuscular hemoglobin concentration</td>
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<td>MCV</td>
<td>mean corpuscular volume</td>
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<td>MEN</td>
<td>multiple endocrine neoplasia</td>
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<tr>
<td>Acronym</td>
<td>Definition</td>
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<tr>
<td>MGUS</td>
<td>monoclonal gammopathy of undetermined significance</td>
</tr>
<tr>
<td>MHC</td>
<td>major histocompatibility complex</td>
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<tr>
<td>MI</td>
<td>myocardial infarction</td>
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<tr>
<td>MOPP</td>
<td>mechlorethamine/vincristine (Oncovin)/procarbazine/prednisone</td>
</tr>
<tr>
<td>MR</td>
<td>magnetic resonance (imaging)</td>
</tr>
<tr>
<td>NHL</td>
<td>non-Hodgkin's lymphoma</td>
</tr>
<tr>
<td>NIDDM</td>
<td>non-insulin-dependent diabetes mellitus</td>
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<tr>
<td>NPO</td>
<td>nil per os (nothing by mouth)</td>
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<tr>
<td>NSAID</td>
<td>nonsteroidal anti-inflammatory drug</td>
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<td>PA</td>
<td>posterior/anterior</td>
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<td>PIP</td>
<td>proximal interphalangeal</td>
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<td>PBS</td>
<td>peripheral blood smear</td>
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<tr>
<td>PE</td>
<td>physical exam</td>
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<td>PFTs</td>
<td>pulmonary function tests</td>
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<td>PMI</td>
<td>point of maximal intensity</td>
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<td>PMN</td>
<td>polymorphonuclear leukocyte</td>
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<tr>
<td>PT</td>
<td>prothrombin time</td>
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<td>PTCA</td>
<td>percutaneous transluminal angioplasty</td>
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<td>parathyroid hormone</td>
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<td>partial thromboplastin time</td>
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<td>PUD</td>
<td>peptic ulcer disease</td>
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<td>RBC</td>
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<td>respiratory rate</td>
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<tr>
<td>RS</td>
<td>Reed-Sternberg (cell)</td>
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<tr>
<td>RV</td>
<td>right ventricular</td>
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<tr>
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<td>right ventricular hypertrophy</td>
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<tr>
<td>SBFT</td>
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<td>UGI</td>
<td>upper GI</td>
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<td>ultrasound</td>
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<td>Definition</td>
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<td>VDRL</td>
<td>Venereal Disease Research Laboratory</td>
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<tr>
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<td>Wolff–Parkinson–White (syndrome)</td>
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<tr>
<td>XR</td>
<td>x-ray</td>
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A 30-year-old female presents to the ER with severe, sudden-onset shortness of breath and an extensive pruritic skin rash.

**HPI**

She was prescribed cotrimoxazole by her general physician for a UTI; she took the first dose only a few minutes before developing symptoms.

**PE**

VS: hypotension. PE: severe respiratory distress; central cyanosis; extensive urticarial wheals noted all over body.

**Labs**

IgE antibody demonstrated to sulfonamides by RAST.

**Treatment**

Epinephrine (1:1000); antihistaminics; steroids; ventilatory support; adequate IV fluid administration or vasopressor agents to treat hypotension.

**Discussion**

Systemic anaphylaxis is the most serious and life-threatening IgE-mediated type 1 hypersensitivity reaction; its recognition and prompt treatment are critical to survival.
A 2-year-old male is admitted to the hospital for evaluation of a suspected immune disorder.

He has a history of recurrent fungal diaper rashes and staphylococcal cervical furunculosis requiring multiple incisions and drainage in addition to antibiotics. His mother also reports chronic diarrhea and a prior perianal fistula.

Cervical lymphadenopathy; mild hepatomegaly and splenomegaly; no pallor, purpuric patches, or sternal tenderness.

CBC/PBS: neutrophilic leukocytosis. Elevated ESR; normal serum immunoglobulins; absence of respiratory burst (negative nitroblue tetrazolium test and chemoluminescence assay); negative Mantoux test.

CXR: hilar lymphadenopathy. US, abdomen: hepatosplenomegaly; hepatic and splenic nodular lesions (due to granulomas).

Characteristic granuloma formation with phagocytes, giant cells, and occasional histiocytes in lymph nodes, liver, spleen, and lungs.

Long-term TMP-SMX prophylaxis, γ-interferon.

Chronic granulomatous disease is most commonly an X-linked disorder of neutrophil function (may have variable inheritance patterns) that is due to a deficiency of NADPH oxidase. Neutrophils of affected patients demonstrate normal chemotaxis, degranulation, and phagocytosis but cannot use the oxygen-dependent myeloperoxidase system for microbial killing, making patients susceptible to recurrent staphylococcal infections.
ID/CC  A 19-year-old male has recurrent attacks of bilateral periorbital and hand swelling coupled with respiratory difficulty that lasts up to 24 hours and often requires hospitalization.

HPI  He does not, however, complain of itching. His father and his aunt both suffer from a similar illness.

PE  Physical examination unremarkable.

Labs  Decreased C4 (best screening test); decreased C1 inhibitor (confirmatory test) and C2; normal C3; normal IgE.

Treatment  Synthetic androgens (e.g., danazol), fresh frozen plasma.

Discussion  C1 esterase inhibitor deficiency is inherited as an autosomal-dominant trait; death may result from laryngeal edema. Also known as hereditary angioedema.
A 25-year-old white female is referred to an internist by her family doctor for a workup of recurrent sinusitis, chronic otitis media, one episode of pneumonia that required hospitalization, and recurrent bouts of watery diarrhea.

HPI: She has seen an allergy specialist for several years and has received desensitization shots for multiple allergies, including pollen, dust, and cat hair.

PE: Normal except for hypopigmented spots on neck and arms (vitiligo).

Labs: Markedly decreased serum IgA; normal IgG and IgM.

Imaging: XR, sinus: opacification of paranasal sinuses (due to chronic sinusitis).

Treatment: Largely supportive; antibiotic therapy; try to avoid blood or plasma transfusion (anaphylaxis or serum sickness due to presence of antibodies to IgA).

Discussion: Selected IgA deficiency is the most common congenital immunodeficiency, especially in patients of European descent. Diarrhea is usually caused by Giardia lamblia; recurrent sinopulmonary infections are caused by Streptococcus pneumoniae, Haemophilus influenzae, or Staphylococcus aureus; associated with an increased incidence of allergies and autoimmune diseases such as SLE and rheumatoid arthritis. Selective IgA deficiency may be due to a specific defect in isotype switching.
ID/CC A 4-month-old male presents with chronic diarrhea and failure to thrive.

HPI The infant was diagnosed with extensive mucocutaneous candidiasis in the early neonatal period and shortly thereafter developed a fulminant Pseudomonas septicemia that required intravenous antibiotic therapy for an extended period of time. A paternal cousin had developed similar and equally devastating bacterial and fungal infections in the neonatal period and subsequently died.

PE Emaciated; mucocutaneous candidiasis noted; tonsils not seen; lymph nodes not palpable despite recurrent infections.

Labs CBC: severe lymphopenia. PBS: lack of mature lymphocytes. Tests for cutaneous delayed hypersensitivity and contact sensitization negative; serum immunoglobulin levels (IgG, IgA, and IgM) low; adenosine deaminase (ADA) deficiency demonstrated in red cells.

Imaging CXR: absent thymic shadow.

Gross Pathology Thymus fails to descend into the anterior mediastinum from the neck and resembles fetal thymus of 6 to 8 weeks.

Micro Pathology No lymphoid tissue in the lymph nodes, spleen, tonsils, and appendix.

Treatment Bone marrow transplant from an HLA-identical sibling; IV immunoglobulin; infusion of normal ADA-containing erythrocytes (ADA-PEG is also very successful); antibiotics; gene therapy for ADA; genetic counseling (SCID caused by ADA deficiency can be diagnosed prenatally by amniocentesis).

Discussion Severe combined immunodeficiency syndrome is characterized by marked depletion of the cells that mediate both humoral (B-cell) and cellular (T-cell) immunity. SCID may be transmitted as either an autosomal-recessive trait or an X-linked recessive trait, or it may be sporadic; half of the cases inherited in an autosomal-recessive manner are caused by a deficiency in ADA.
A 7-month-old male is admitted for a workup of recurrent upper respiratory tract and skin infections of several months' duration.

His parents state that he has had recurrent URIs, one episode of *Haemophilus influenzae pneumonia*, and severe otitis media.

Low weight and height for chronological age; chronic bilateral suppurative otitis media; asymmetric arthritis of knees; no tonsillar tissue seen; no lymphadenopathy or hepatosplenomegaly.

Panhypogammaglobulinemia: very low IgG; IgA and IgM undetectable.

Parenteral gamma globulin; monitor pulmonary function to guard against chronic lung disease.

An X-linked disease (manifests only in males) characterized by a selective B-cell defect with recurrent bacterial infections. Also known as Bruton's disease, X-linked hypogammaglobulinemia is due to a genetic defect in tyrosine kinase receptor found on antibody precursors, resulting in impaired maturation and development of antibodies. Male infants demonstrate infections when maternal antibodies have cleared from their system.
A 27-year-old white female complains of mouth ulcers, prolonged fever, flulike symptoms, and increasing fatigue and weight loss over the past 2 months.

She recently moved from a large metropolitan area to a farm in Ohio, where she spent 1 week cleaning a pigeons' loft.

VS: fever (38.5°C). PE: pallor; weight loss; enlarged liver and spleen; generalized lymphadenopathy; scattered sibilant rales over lung fields.

CBC/PBS: anemia; leukopenia. Small, budding fungus found intracellularly in reticuloendothelial cells (macrophages) on silver stain; elevated LDH; positive blood culture for dimorphic fungus.

CXR: nonsegmental shifting pneumonic infiltrates; mediastinal adenopathy with popcorn calcifications; bilateral hilar adenopathy. CT, abdomen: splenic calcifications.

Nodules with granuloma formation; central area of necrosis and caseation with sclerosis and calcification; any organ may be involved, mainly reticuloendothelial system (RES) and adrenals.

Granulomas with epithelioid cells, Langhans' giant cells, and organisms within macrophages; in disseminated disease, organisms present in RES throughout body with proliferation.

Itraconazole; amphotericin B.

Histoplasmosis is a systemic fungal infection sometimes resembling TB that is caused by *Histoplasma capsulatum*, a dimorphic fungus. The yeast form is found intracellularly; the mold form is found in soil associated with bird or bat feces. Transmitted by inhalation of mold spores, it varies in intensity from asymptomatic to fulminant (in immunocompromised patients). The disease is most prevalent in the southeastern, mid-Atlantic, and central regions of the United States.
A 57-year-old black male complains to his doctor of increasing weakness, swollen glands in the armpits and groin, and a feeling of heaviness in the abdomen (due to hepatosplenomegaly).

The patient is an immigrant from Trinidad and Tobago and has a history of nonresolving skin rashes and recurrent respiratory infections.

Marked pallor; extensive papular skin rash with few erythematous plaques over abdomen; generalized lymphadenopathy and hepatosplenomegaly.

CBC/PBS: marked leukocytosis (83,000) with relative lymphocytosis and atypical lymphocytes. Increased LDH; hypercalcemia.

CXR: normal.

Skin biopsy reveals infiltration by leukemic CD4+ T lymphocytes.

Aggressive combination chemotherapy.

Adult T-cell leukemia/lymphoma (ATLL) is associated with HTLV-I type C, a retrovirus that has a higher incidence in blacks from the Caribbean and southeastern United States as well as in people from southern Japan and sub-Saharan Africa. The infection is acquired via transmission from mother to child (breast milk), from sexual activity, from blood transfusion, or from IV drug use.
A 2-day-old neonate is evaluated for an eye discharge.

The baby's mother is a prostitute who did not receive any prenatal cervical cultures during pregnancy.

Normal full-term male neonate; mucoid eye discharge, conjunctival congestion, and chemosis noted in both eyes; nonfollicles seen on palpebral conjunctiva (due to absence of subconjunctival adenoid layer at this age); mild superficial keratitis also present.

Gram stain of swab reveals increased PMNs and no bacteria; characteristic intracellular inclusion bodies demonstrated by the DIF test; cell culture yields Chlamydia trachomatis serotypes D through K; chlamydia also grown from maternal cervical swab.

Erythromycin syrup; azithromycin suspension has also been shown to be beneficial; no topical therapy.

Chlamydia trachomatis is an important cause of preventable blindness; its strains can be further differentiated into 18 serotypes by microimmunofluorescence tests. Serotypes A, B, Ba, and C are principally associated with endemic trachoma in developing countries; serotypes D through K primarily cause sexually transmitted infections in adults and inclusion conjunctivitis and pneumonia in infants, transmitted through an infected birth canal; and serotypes L1, L2, and L3 cause lymphogranuloma venereum.
ID/CC
A 20-year-old male college student complains of sore throat, fatigue, fever, swollen lymph nodes on the back of his neck, anorexia, cough, and malaise of 10 days' duration.

HPI
He was initially given ampicillin by his school nurse, after which he developed an extensive skin rash.

PE
VS: fever. PE: enlargement of submaxillary and cervical lymph nodes; exudative tonsillitis; petechiae on soft palate; slightly enlarged spleen and liver.

Labs
CBC/PBS: anemia; thrombocytopenia; leukocytosis with absolute lymphocytosis (50%); atypical lymphocytes. Elevated ALT, AST, and bilirubin; positive heterophil antibody test (PAUL-BUNNELL TEST); IgM antibodies to viral capsid antigen/monospot positive.

Gross Pathology
Enlarged spleen, lymph nodes, and, to lesser extent, liver; hepatitis may be present along with brain involvement; splenic rupture rare complication.

Micro Pathology
Proliferation of reticuloendothelial system; infiltration of spleen by atypical lymphocytes.

Treatment
Supportive.

Discussion
Infectious mononucleosis is a systemic viral infection that is caused by Epstein–Barr virus (EBV), a herpesvirus, and is transmitted through respiratory droplets and saliva. In developed countries, it most commonly affects teenagers and young adults ("kissing disease"); in underdeveloped countries, it is seen as a subclinical infection of early childhood. EBV infection is associated with an increased risk of Burkitt's lymphoma, Hodgkin's disease, and nasopharyngeal carcinoma.
A 65-year-old male presents with a high fever, headache, extreme prostration, a nonproductive cough, and severe breathlessness.

He had been receiving chlorambucil for treatment of chronic lymphocytic leukemia (CLL) and was in an extremely debilitated state.

VS: fever; tachypnea; cyanosis. PE: conjunctival congestion; pharyngeal inflammation; rales and wheezes heard on auscultation over both lung fields; splenomegaly and lymphadenopathy (due to CLL).

No organisms seen or cultured from sputum; fluorescent antibody directed against influenza virus was positive; viral cultures of nasopharyngeal washings grew influenza virus; fourfold rise in hemagglutination inhibition antibody titer against influenza virus demonstrated.

CXR (PA view): bilateral, diffuse interstitial infiltrates suggestive of atypical pneumonia.

Amanitidine or rimantadine for influenza A (zanamivir or oseltamivir for influenza A and B); ventilatory support, antipyretics, and IV fluids. Secondary staphylococcal pneumonia should be treated with parenteral antibiotics; yearly vaccination prevents excessive morbidity and mortality, especially among the elderly.

Influenza viruses are medium-sized spherical RNA viruses termed orthomyxoviruses; influenza A and B viruses each contain 8 RNA segments and 10 viral proteins. Influenza infection is most common in winter, with the severity of a given outbreak depending on the status of immunity in the community.

Previous natural infection or immunization with viruses that are immunologically close to the current strain limits new infection, but if antigenic drift results in reduced cross-reactivity, the new strain will spread more rapidly. New strains produced by antigenic shift account for most major outbreaks. Influenza affects all segments of the population, but severe infections and major complications are most common in patients who are young, elderly, or debilitated.
A 30-year-old female presents with fever, chills, malaise, headaches, and myalgias.

She was diagnosed as suffering from secondary syphilis with an extensive nonpruritic skin rash, condylomata lata, and mucous patches in the mouth, for which she received a dose of intramuscular penicillin 6 hours ago.

VS: fever; tachycardia; mild hypotension.

No specific treatment; symptoms subside in 24 hours.

The Jarisch-Herxheimer reaction consists of fever, chills, mild hypotension, headache, and an increase in the intensity of mucocutaneous lesions 2 hours after initiating treatment of syphilis with penicillin or another effective antibiotic; symptoms usually subside in 12 to 24 hours. The reaction occurs in 50% of patients with primary syphilis and in 90% of those with secondary syphilis. The Jarisch-Herxheimer reaction also occurs after treatment of other spirochetal diseases (e.g., louse-borne relapsing fever caused by Borrelia recurrentis). It has been suggested that the release of treponemal lipopolysaccharides might produce this symptoms complex.
ID/CC  A 40-year-old male smoker complains of acute-onset high fever, chills, a nonproductive cough, tachypnea, and pleuritic chest pain.

HPI  A number of similar cases have been reported in his workplace in recent months. The patient admits to significant alcohol and tobacco consumption and uses a humidifier at night.

PE  VS: fever; dyspnea. PE: rales present bilaterally on auscultation.

Labs  Sputum exam with Gram stain reveals no pathogenic organisms. CBC: neutrophilic leukocytosis. Cold agglutinins absent; indirect fluorescent antibody technique reveals stable titer of > 1:256 (considered diagnostic); direct immunofluorescent staining of sputum confirms presence of Legionella.

Imaging  CXR, PA: bilateral diffuse, patchy infiltrates and ill-defined nodules.

Gross Pathology  Nodular areas of consolidation that may progress to involvement of one or more lobes of the lung.

Micro Pathology  Alveolar exudate with PMNs, macrophages, and fibrin; in more severe cases, destruction of alveolar septa.

Treatment  Erythromycin or an active fluoroquinolone.

Discussion  Legionnaire’s disease is caused by Legionella pneumophila, a filamentous, flagellated, aerobic gram-negative, motile bacillus, and is more common in immunocompromised patients. Epidemiologic studies have established drinking water and air conditioners as the sources of outbreak.
A 30-year-old Pakistani immigrant complains of chronic fever, weight loss, increased abdominal girth, a feeling of heaviness, and appetite loss.

Almost a year ago, the patient had a small, pruritic red papule on his left arm that was caused by an insect bite and disappeared spontaneously.

Skin darkening; trophic changes in hair; massive nontender, hard splenomegaly; hepatomegaly without jaundice; generalized lymphadenopathy; peripheral edema; ecchymosis.

CBC/PBS: anemia, leukopenia, thrombocytopenia (pancytopenia), and monocytosis; amastigotes in buffy coat. Hypergammaglobulinemia; decreased albumin; increased ALT and AST.

CT/US, abdomen: splenomegaly.

Massively enlarged spleen: also greatly increased in weight, dark colored, and congested with Leishman–Donovan bodies.

Proliferation of reticuloendothelial system cells; biopsy or aspiration reveals parasite-filled macrophages in infected locations.

Pentavalent antimony (e.g., sodium stibogluconate); amphotericin B or pentamidine isethionate.

Also known as kala azar, leishmaniasis is a zoonosis that is produced by Leishmania donovani and is transmitted through the bite of the Phlebotomus sandfly. It is associated with a high fatality rate when left untreated.
A 30-year-old male from India presents with slowly progressive hypopigmented skin patches and nodules together with a peculiar deformity of the nose.

The patient has a history of nasal stuffiness and bloody nasal discharge; he also complains of loss of libido.

Leonine facies (thickened facial and forehead skin); loss of eyebrows and eyelashes (MADAROSIS); depressed nasal bridge ("SADDLE-NOSE" DEFORMITY); gynecomastia; testicular atrophy; numerous symmetrical, hypopigmented macules with vague edges and erythematous, smooth, shiny surfaces; skin plaques and nodules; partial loss of pinprick and temperature sensation (HYPOESTHESIA); no anhidrotic changes; symmetrically enlarged ulnar and common peroneal nerves.

CBC: mild anemia. ESR elevated; slit skin smears reveal numerous acid-fast bacilli on modified ZN staining.

Dermis massively and diffusely infiltrated with foamy histiocytes with bacilli and globi (masses of acid-fast bacilli) containing Virchow giant cells; bacilli found only rarely in epidermis and in subepidermal "clear zone"; epidermis thinned out with flattening of rete ridges.

Multidrug therapy with rifampicin, dapsone, and clofazimine.

The discovery of one or more of the following is pathognomonic of leprosy: (1) anesthetic skin lesions (found in all tuberculoid and many lepromatous cases); (2) thickening of one or more nerves (found in many lepromatous and some tuberculoid cases); and (3) the presence of acid-fast bacilli in skin smears (found in all lepromatous and some tuberculoid cases). Mycobacterium leprae has not been cultured in vitro thus far. Frequent complications include hand crippling (secondary to nerve damage) and blindness. It is currently believed that in most instances, the mode of transmission is via person-to-person contact.

15 LEPROSY—LEPROMATOUS
A 26-year-old male from India presents with a hypopigmented, anesthetic skin patch over the left side of his face.

He also complains of an occasional "electric current"-like sensation radiating from his left elbow to his hand.

Dry, hypopigmented, anesthetic patch over left cheek; left ulnar nerve enlarged and palpable; eye, ear, nose, and throat exam normal; testes normal (vs. signs that are often demonstrable in lepromatous leprosy).

Glucose-6-phosphate dehydrogenase (G6PD) levels within normal range (done to prevent dapsone-associated hemolysis); slit skin smears reveal few acid-fast bacilli; skin biopsy from patch diagnostic of tuberculoid leprosy.

Single or small number of lesions with macular or raised edges.

Skin biopsy reveals many well-formed epithelioid granulomas with very few acid-fast bacilli.

Chemotherapy with rifampin and dapsone.

Caused by Mycobacterium leprae, an acid-fast bacillus. The organism has two unique properties: it is thermolabile, growing best at 27°C to 30°C, and it divides very slowly; generation time is 12 to 14 days. Consequently, leprosy in humans typically evolves very slowly. Tuberculoid leprosy predominantly affects the skin with limited nerve involvement (most commonly ulnar and peroneal); lepromatous leprosy has diffuse involvement of the skin, eyes, nerves, and upper airway with disfigurement of the hands and face (leonine facies).
A 35-year-old British dairy farmer complains of a high remittent fever with chills, severe muscle aches, decreased urine output, and dark-colored urine for the past 2 days.

He also complains of an extensive skin rash and nasal bleeding (epistaxis). A careful history reveals that the area in which he works is infested with rodents.

VS: fever; tachycardia; hypotension. PE: icterus; extensive hemorrhagic maculopapular skin eruption; conjunctival suffusion; lymphadenopathy.

CBC: leukocytosis with neutrophilia; thrombocytopenia. Mild hyperbilirubinemia, predominantly conjugated: increased alkaline phosphatase; elevated BUN and creatinine. UA: proteinuria, casts, and RBCs. Blood culture (positive during first 10 days of illness) and urine culture (positive after second week of infection) on Fletcher’s medium isolated Leptospira interrogans; serologic diagnosis (positive during second week of illness): microscopic slide agglutination demonstrated significant titer of antibody to L. interrogans.

CXR: patchy alveolar infiltrates consistent with alveolar hemorrhage.

Severe infection damages both the liver and kidneys.

Liver biopsy shows focal centrilobular necrosis with focal lymphocytic infiltration and disorganization of liver cell plates together with proliferation of Kupffer cells with cholestasis; kidney biopsy reveals mesangial proliferation with PMN infiltration.

Penicillin (dose modified due to presence of renal failure), doxycycline; hemodialysis.

Weil’s disease, a severe form of leptospirosis caused by Leptospira interrogans complex, is characterized by fever, jaundice, cutaneous and visceral hemorrhages, anemia, azotemia, and altered consciousness; major vectors to humans are rodents.

Transmission occurs through direct contact with the blood, tissue, or urine of infected animals. Person-to-person transmission is highly unlikely. Preventive measures include limiting the rodent population and vaccinating animals.
A 2-week-old female is brought to the emergency room because of high fever and convulsions.

She also has an extensive skin rash on her legs and trunk.

VS: fever. PE: generalized hypotonia; extensive maculopapular skin rash; nuchal rigidity; involuntary flexion of hips when flexing neck (BRUDZINSKI’S SIGN).

CBC: neutrophilic leukocytosis. LP: elevated CSF cell count (750 cells/mL), mostly neutrophils; elevated CSF protein; low CSF sugar. Gram-positive, facultative, intracellular, nonsporulating motile bacilli on Gram stain and culture.

Purulent meningitis.

Bacillus provokes both acute suppurative reaction with neutrophilic infiltration and chronic granuloma formation with focal necrosis.

IV antibiotics (high-dose ampicillin).

Listeriosis is caused by *Listeria monocytogenes*. Bacterial infection may occur early (acquired in utero) or later (drinking contaminated milk) in neonatal life. May be rapidly fatal if disseminated. Also occurs in adults immunocompromised by disease (e.g., renal disease or HIV). *Escherichia coli* and group B streptococcus are two other common causes of neonatal meningitis.
ID/CC  A 12-year-old male presents with fatigue, fever, headache, fleeting joint pain, and a reddish rash on his trunk and left leg of 1 week’s duration.

HPI  The patient is a native of Connecticut and attended a summer camp in the state’s national park 2 weeks ago. He recalls having noticed a tick bite on his leg about 2 weeks ago.

PE  VS: fever. PE: red macule on site of bite that has grown circumferentially; active border and central clearing (erythema chronicum migrans); femoral lymphadenopathy; mild neck stiffness; normal CNS exam.

Labs  Positive IgM ELISA for Borrelia burgdorferi; diagnosis confirmed by Western blot assay. ECG: normal. LP: lymphocytic pleocytosis; increased proteins. B. burgdorferi grown on Noguchi medium.

Gross Pathology  Erythema chronicum migrans (ECM) is characteristic of Lyme disease; must be minimum of 5 cm in diameter for diagnosis to be made; center may desquamate, ulcerate, or necrose; satellite lesions sometimes seen; may spontaneously disappear with time.

Treatment  Doxycycline; amoxicillin; ceftriaxone.

Discussion  The most common disease transmitted by vectors in the United States, Lyme disease is caused by Borrelia burgdorferi, a spirochete, and is transmitted through Ixodes species tick bites. Ticks acquire B. burgdorferi from deer mice, which are the natural reservoir. There are three recognized stages: stage 1 consists of ECM and constitutional symptoms; stage 2, cardiac or neurologic involvement; and stage 3, persistent migratory arthritides, synovitis, and atrophic patches on the distal extremities (acrodermatitis chronicum atrophicans).
A 57-year-old black female from Kenya complains of increasing weight and edema of the lower legs with difficulty walking.

HPI Over the years she has had episodes of fever with swelling of inguinal lymph nodes and itching. She has also had numerous attacks of malaria.

PE Inguinal lymph nodes indurated and slightly increased in size; marked deformity in both legs with thickening of skin and greatly increased diameter; rubbery consistency.

Labs PBS: several microfilariae; prominent eosinophilia.

Imaging Lymphangiogram: partial lymphatic obstruction at iliac level.

Gross Pathology Presence of adult worms in lymphatics; marked fibrosis surrounding obstructed vessels.

Micro Pathology Granulomatous reaction with plasma cell and lymphocytic infiltration; giant cell formation; intense fibroblastic hyperplasia.

Treatment Ivermectin; diethylcarbamazine; surgery in advanced cases.

Discussion Lymphatic filariasis is a chronic disease that is due to lymphatic obstruction and is caused by several types of filarial roundworms, mainly Wuchereria bancrofti and Brugia malayi; it is transmitted by female mosquito bites. Also known as elephantiasis.

Atlas Link U52M M-M2-020

LYMPHATIC FILARIASIS
A 25-year-old male complains of swollen, tender masses in his groin and very painful genital ulcers of 1 week's duration.

The patient admits to having had unprotected sex with multiple partners.

Swollen, erythematous, tender inguinal nodes, usually bilateral, with draining sinuses (inguinal adenitis, buboes); multiple small genital lesions.

Inguinal node biopsy diagnostic; positive complement fixation test; positive immunofluorescence test.

Primary lesion is ulcerated nodule; gives rise to inguinal bubo, an enlarged lymph node sometimes characterized by fistulous tract formation; balanitis, phimosis, and rectal involvement with stricture may also be present.

Neutrophilic infiltration of primary lesion with areas of necrosis; lymphoid hyperplasia of lymph nodes with foci of macrophage accumulation; abscess formation with fibrosis.

Doxycycline; tetracycline; azithromycin; erythromycin; TMP-SMX; ceftriaxone; ciprofloxacin.

Lymphogranuloma venereum is an STD that is due to Chlamydia trachomatis (L1, L2, L3). Counseling should be given about other STDs (e.g., AIDS, syphilis, gonorrhea).
A 30-year-old missionary comes to the emergency room complaining of high fever, chills, severe headache, and confusion.

Upon returning from Africa 2 weeks ago, he began to feel weak and experienced backaches, pain behind the eyes, and sleepiness.

VS: fever (39°C); tachycardia. PE: pallor; profuse sweating; mild splenomegaly without lymphadenopathy.

CBC/PBS: anemia; thrombocytopenia; plasmodia in erythrocytes on thick peripheral blood smear. Slight hyperbilirubinemia and hypoglycemia.

Liver and spleen moderately enlarged and soft in consistency, with sequestration and hemolysis of erythrocytes and macrophages; hyperplasia of Kupffer cells; malarial pigment in spleen and liver; brain capillaries may show thromboses.

Hypertrophy of phagocytic system; ischemic necrosis surrounding occluded blood vessels in brain.

Chloroquine; quinine for cerebral malaria; sulfadiazine-pyrimethamine, mefloquine, tetracycline for areas with chloroquine-resistant strains; primaquine for radical treatment.

Malaria is transmitted by female Anopheles mosquitoes. Plasmodium falciparum may be lethal, producing cerebral malaria. Other types include P. vivax, P. ovale, and P. malariae.
A 3-year-old female is brought to the emergency room with a high fever of 7 days' duration, accompanied by redness of the eyes, persistent dry cough, and coryza.

Her family doctor had treated her illness as a viral URI, but no improvement was seen. One day before her admission, her mother noticed a skin rash starting behind her ears and face that has now spread to her trunk and extremities.

Pallor; injected conjunctiva; hyperemic throat; erythematous maculopapular rash on face, neck, trunk, and extremities; retroauricular lymphadenopathy; bluish-gray spots surrounded by erythematous areola on buccal mucosa in region of first molar (Koplik's spots).

CBC: leukopenia.

Koplik's spots pathognomonic of measles; appearance presages rash by approximately 2 days; uniform lesions (vs. varicella).

Lymphocytic dermal infiltration; multinucleated giant cells in reticuloendothelial system (Warthin-Finkeldey cells).

No specific antiviral therapy available; treat complications.

Also called rubella; not to be confused with rubella. Measles is produced by a paramyxovirus and is transmitted by respiratory droplets; a live attenuated vaccine is available. Measles has an incubation period of 10 to 14 days. Sequelae include encephalitis, subacute sclerosing panencephalitis (SSPE), and giant cell pneumonia.

PED-030A, PED-030B
ID/CC A 12-year-old white female is brought to the emergency room because of sudden fever with chills, severe headache, pain in the extremities and back, stiff neck, and generalized rash; she also fainted while in school.

HPI She had been well until admission, with no relevant history. In the emergency room, she vomits bright red blood twice.

PE VS: tachycardia; hypotension (BP 70/50). PE: altered sensorium; pallor; moist, cold skin; nuchal rigidity and positive Kernig's sign; petechial rash all over body; minimal papilledema on funduscopic exam; no focal neurologic signs.

Labs Hypoglycemia. Labs: hyponatremia; hyperkalemia. CBC/PBS: thrombocytopenia; neutrophilic leukocytosis. LP: CSF cloudy and under increased pressure; increased proteins; low sugar. Gram-negative diplococci (Neisseria meningitidis) seen within and outside WBCs on Gram stain; negative India ink and ZN stain; growth of meningococci later revealed on blood culture.

Imaging CT, head: normal. CT, abdomen: bilateral adrenal hemorrhage.

Gross Pathology Bilateral adrenal hemorrhagic necrosis; skin necrosis; pyogenic meningitis.

Micro Pathology Meningeal hyperemia with abundant purulent exudate; diplococcus-containing PMNs; acute hemorrhagic necrosis of adrenal glands.

Treatment Steroid replacement; IV fluids; dopamine; IV penicillin G; prophylactic rifampin for close contacts.

Discussion Meningococcemia is a fulminant disease caused by several groups of Neisseria meningitidis; the cause of death is adrenal necrosis with vascular collapse. A meningococcal vaccine is available. Also known as Waterhouse-Friderichsen syndrome.
ID/CC A 24-year-old white female with insulin-dependent diabetes mellitus (IDDM) is hospitalized for ketoacidosis following a night out drinking; on the fifth day she develops right periorbital swelling and a mucopurulent postnasal discharge that fails to respond to antibiotics.

HPI She admits to irregular adherence to glucose control and insulin dosing.

PE Right periorbital and paranasal edema; swelling of conjunctiva (chemosis); exophthalmos; black ulceration of nasal mucosa; third cranial nerve (CN III) palsy.

Labs Large, irregular, nonseptate hyphae branching at wide (> 90°) angles on nasal culture.

Imaging XR, plain: opacification of paranasal sinuses.

Gross Pathology Necrotic destruction of paranasal sinuses and orbit with dissemination to lung and brain.

Micro Pathology Purulent arteritis with thrombi composed of hyphae; inflammation and necrosis with polymorphonuclear infiltrate.

Treatment Maintain tighter glucose control; amphotericin B; surgical drainage.

Discussion Mucormycosis is a phycomycosis produced by Mucor and Rhizopus molds; it should be suspected in cases of antibiotic-resistant sinusitis, especially in the presence of underlying diabetes, burns, lymphoma, or leukemia.


MUCORMYCOSIS
A 6-year-old white male presents with fever, nausea, vomiting, swelling, and tenderness of the mandibular angle; he finds it difficult to talk, eat, or swallow.

Two of his classmates were diagnosed with mumps 2 weeks ago. There is no vaccination record.

VS: fever. PE: outward and upward displacement of ear; obliterated mandibular hollow; orifice of Stensen’s duct swollen and hyperemic; right testicle enlarged and painful.

CBC: leukopenia with lymphocytosis. Hyperamylasemia; positive complement fixation antibodies.

Parotid glands enlarged with areas of necrosis and mononuclear infiltrate; encephalitis, orchitis, oophoritis, meningitis, and pancreatitis may also be present.

Examination of parotid glands reveals perivascular mononuclear, lymphocytic, and plasma cell infiltrate with necrosis; ductal obstruction and edema; testicular interstitial edema; perivascular cerebral lymphocytic cuffing.

Supportive; analgesics for pain; treat complications.

A systemic infection caused by the mumps virus, an RNA paramyxovirus, mumps is transmitted by droplets and direct contact. Bilateral testicular involvement may lead to sterility; one of the most common causes of pancreatitis in children. A vaccine is available with measles and rubella (MMR).
A 20-year-old male college student presents with a **productive cough**, headache, malaise, runny nose, and fever.

**HPI** He has a history of sore throat preceding the onset of the cough, which initially was nonproductive.

**PE** VS: fever. PE: mild respiratory distress; auscultation reveals fine to medium rales over right lower lobe.

**Labs** Gram stain of sputum negative; routine cultures of both blood and sputum negative. CBC: **leukocyte count normal**. Fourfold rise in complement fixation titer in paired sera; cold agglutinin titer > 1:128.

**Imaging** CXR: patchy alveolar infiltrates involving right lower lobe; appears worse than the clinical picture.

**Gross Pathology** Unilateral lower lobe pneumonia with firm, red pulmonary parenchyma in affected areas.

**Micro Pathology** Bronchial mucosa congested and edematous; inflammatory response consists of perivascular lymphocytes initially and PMNs later in infection. **Organism lacks cell wall** (thus penicillins and cephalosporins are ineffective).

**Treatment** Erythromycin.

**Discussion** *Mycoplasma pneumoniae* is the most common cause of primary atypical pneumonia. Transmission is by droplet spread; rapidly infects those living in close quarters.
ID/CC  A 50-year-old diabetic male presents with fever, pain, and a necrotizing swelling over his left leg.

HPI  His symptoms began about a week ago with redness and swelling of the left leg followed by bronze discoloration of the skin and the appearance of hemorrhagic bullae.

PE  Extensive cutaneous gangrene observed over left leg with many ruptured bullae; black necrotic eschar with surrounding erythema resembles a third-degree burn.

Labs  Swab staining reveals presence of chains of gram-positive cocci; culture isolated β-hemolytic group A streptococci (Streptococcus pyogenes).

Micro Pathology  Biopsy specimen reveals areas of necrosis in dermis and subcutaneous fat, infiltration with PMNs, and vasculitis and thrombosis in vessels in the superficial fascia.

Treatment  Treatment includes rapid surgical excision of necrotic tissue in combination with appropriate antibiotics.

Discussion  Streptococcal gangrene is a group A streptococcal cellulitis that rapidly progresses to gangrene of the subcutaneous tissue and necrosis of the overlying skin; the disease process usually involves an extremity. Necrotizing fasciitis is also recognized as a polymicrobial infection that is caused by aerobes and anaerobes ("SYNERGISTIC NECROTIZING CELLULITIS"). Infection spreads quickly through various fascial planes, the venous system, and lymphatics. Predisposing etiologies include surgery, trauma, and diabetes.
A 45-year-old white male undergoing chemotherapy for Hodgkin’s lymphoma is brought to the emergency room by his wife because of shortness of breath and cyanosis.

HPI For the past 3 months, he has been complaining of intermittent weakness, fever with chills, and foul-smelling, thick greenish sputum.

PE VS: fever (38°C); tachypnea; tachycardia. PE: pallor; mild cyanosis; localized dullness with bronchial breathing; diminished breath sounds over left lower lobe.

Labs CBC: leukocytosis with neutrophilia; anemia. Sputum culture reveals gram-positive, filamentous, partially acid-fast staining bacteria (due to Nocardia).

Imaging CXR: nodular infiltrate in left lower lobe with air-fluid level (abscess) and left pleural effusion.

Gross Pathology Lung lesions or disseminated lesions (brain, liver, kidney, subcutaneous tissue) consist of necrotic centers within regions of consolidation and abscess formation resembling pyogenic pneumonia.

Micro Pathology Consolidation of alveoli with pus formation (exudate of PMNs and fibrin) and surrounding granulomatous reaction.

Treatment Six-month course of TMP-SMX; surgery.

Discussion A chronic bacterial infection seen in diabetics, leukemia and lymphoma patients, and immunocompromised patients, nocardiosis usually involves the lungs with possible dissemination to the brain, subcutaneous tissue, and other organs. It is caused by Nocardia asteroides, a branching, aerobic, gram-positive organism that is weakly acid fast and is sometimes confused with Mycobacterium tuberculosis.

NOCARDIOSIS
A 60-year-old male who was hospitalized following a stroke presents with a high-grade fever with chills and obtundation.

He had been catheterized due to urinary incontinence and was receiving cephalosporin for treatment of aspiration pneumonitis.

VS: fever.

Blood culture grew Enterococcus fecalis (morphologically indistinguishable from streptococci and immunologically similar to members of group D streptococci, the enterococci are metabolically unique in their ability to resist heat, bile, and 0.5% NaCl); urine culture also isolated Streptococcus fecalis.

Ampicillin with gentamicin (vancomycin can be substituted for ampicillin in patients with penicillin allergies).

Enterococci constitute a relatively common cause of UTIs, wound infections, and peritonitis and intra-abdominal abscesses; they have also become an increasingly prominent cause of bacteremia, which usually originates from a focus in the urinary tract or abdomen. The incidence of nosocomial bacteremias caused by these organisms is also increasing, particularly in patients who have received cephalosporins or other broad-spectrum antibiotics. All clinically significant isolates should be subjected to testing for $\beta$-lactamase production, high-level aminoglycoside resistance, and vancomycin resistance to determine if an alternative therapy is necessary. Infections caused by enterococci that produce $\beta$-lactamase are treated with an antimicrobial agent that combines a penicillin with a $\beta$-lactamase inhibitor; infections caused by strains that are highly resistant to aminoglycosides are treated with vancomycin.
A 56-year-old white female is referred to an ophthalmologist for an evaluation of diminished visual acuity.

HPI She has spent most of her adult life as a missionary in rural Senegal and Mali. She admits to chronic generalized itching, mostly while showering.

PE Wrinkling and loss of elastic tissue in skin; marked hypopigmentation of shins; 2- to 3-cm, nonfixed, firm, nontender subcutaneous nodules on iliac bones, knees, and elbows; chronic conjunctivitis, sclerosing keratitis, and chorioretinal lesions on eye exam.

Labs CBC/PBS: eosinophilia. Fifty-milligram dose of diethylcarbamazine produces severe pruritus, rash, fever, and conjunctivitis (positive Mazzotti reaction).

Micro Pathology Skin biopsy at iliac crest shows microfilariae.

Treatment Ivermectin; suramin.

Discussion Onchocerciasis is caused by Onchocerca volvulus and is transmitted by the blackfly (Simulium), which breeds near rivers; hence it is also known as “river blindness.” Larvae migrate through subcutaneous tissue, producing painless soft tissue edema (CALABAR EDema); with time, subcutaneous nodules form and filariae obstruct dermal lymphatics, producing atrophy and hypopigmentation. Microfilariae concentrate in the eyes, leading to chorioretinitis and blindness.
A 9-year-old male is admitted for an evaluation of a suspected underlying immune deficiency.

He has been hospitalized and treated several times for recurrent life-threatening septicemia due to Streptococcus pneumoniae, meningococcus, and Haemophilus influenzae. Careful history reveals that a few years ago he underwent an emergency splenectomy following traumatic splenic rupture in a motor vehicle accident.

Left paramedian postsurgical scar seen on abdomen.

Reduced IgM levels; reduced antibody production when challenged with particulate antigens; PBS reveals Howell-Jolly bodies.

US, abdomen: spleen is absent.

Pneumococcal vaccine and prophylactic antibiotics (penicillin, amoxicillin, TMP-SMX).

Patients who have undergone splenectomy or who are functionally asplenic are at increased risk for overwhelming bacteremia: pathogens include organisms that possess a polysaccharide capsule, such as meningococcus, Staphylococcus, the DF2 bacillus, and, especially, Streptococcus pneumoniae and Haemophilus influenzae type B. Such functionally asplenic patients include individuals with sickle cell disease and those who have undergone splenic irradiation. Pneumococcal vaccine is indicated in all patients who have undergone splenectomy, particularly children and adolescents.
A 12-year-old girl arrives in the emergency room with pain, swelling, and limited motion of her left hand; she also complains of fever and chills.

The girl was bitten by a cat yesterday while playing at a friend's house.

Hand is erythematous, shiny, and markedly edematous; on palpation, hand is tender with fluctuation (cellulitis); limited passive and active motion; yellowish-green purulent fluid drains from wound; left epitrochlear and axillary lymphadenitis without lymphangitis.

Gram-negative rods with bipolar staining of abscess aspirate; catalase and oxidase positive (Pasteurella multocida).

XR, plain: soft tissue swelling; no periostitis or erosions (vs. osteomyelitis).

Incision and drainage, amoxicillin/clavulanate; tetracycline; penicillin.

Pasteurella multocida is the most common bacterium isolated from cat bite wounds and may progress to osteomyelitis. Human bite infections are most commonly caused by Eikenella corrodens and are treated with penicillin.
A 44-year-old male archaeologist presents with high fever, malaise, intense headache, severe myalgia, and painful swelling in the inguinal region.

HPI He recently returned from a trip to Arizona.

PE VS: tachycardia. PE: drowsy looking; no meningeal signs; pustule seen at site of an insect bite on left upper arm; inguinal lymph nodes enlarged, fluctuant, and tender (buboes); no lesion on external genitalia.

Labs CBC/PBS: normal; no malarial parasites. Gram-negative bacilli with “safety pin” appearance seen in aspirates from buboes; culture of aspirate reveals Yersinia pestis.

Gross Pathology Enlarged lymph nodes are necrotic and suppurative; pneumonic form shows lobar consolidation.

Micro Pathology Numerous organisms in suppurative and necrotic lymph tissue.

Treatment Streptomycin; gentamicin; doxycycline prophylaxis for close contacts; tetracycline.

Discussion Plague is usually acquired after contact with rodents and fleas in endemic areas (southwestern United States). Septic shock, pneumonia, DIC, and vascular collapse are life-threatening sequelae. Death rapidly ensues in the absence of treatment.
An 11-year-old white male presents with a high-grade fever, a productive, blood-tinged cough, mucoid sputum, and pleuritic left-sided chest pain of a few days' duration.

The child had previously been well and is fully immunized.

VS: fever; tachypnea. PE: use of accessory respiratory muscles; central trachea; decreased left respiratory excursion; increased vocal fremitus in left infrascapular area with dullness to percussion; bronchial breathing with coarse crackles heard over left lung area.

CBC: increased WBC count; preponderance of neutrophils. ABGs: hypoxemia without hypercapnia. Gram-positive diplococci in sputum; α-hemolytic colonies of gram-positive diplococci (Streptococcus pneumoniae) on blood agar culture.

CXR: homogenous opacification of left lower lobe (lobar consolidation) with small left pleural effusion.

Consolidation of lung parenchyma passes through four stages: congestion and edema, red hepatization, gray hepatization, and resolution.

Vascular dilatation with hyperemia and alveolar edema; PMNs rich in purulent exudate; fibrin deposition; hardening of lung parenchyma with fibrin clotting inside alveoli (consolidation).

Parenteral therapy with penicillin; monitor with radiologic imaging; supplemental oxygen for respiratory distress.

S. pneumoniae is the most common cause of community-acquired pneumonia and produces typical lobar pneumonia.

PNEUMOCOCCAL PNEUMONIA
A 32-year-old HIV-positive male presents with progressively increasing dyspnea over the past 3 weeks.

He also complains of a dry, painful cough, marked fatigue, and a continuous low-grade fever. He has been noncompliant with cotrimoxazole prophylaxis.

VS: fever; marked tachypnea. PE: pallor; generalized lymphadenopathy; respiratory distress; intercostal retraction; mild central cyanosis; nasal flaring; coarse, crepitant rales auscultated at both lung bases.

ABGs: hypoxemia out of proportion to clinical findings. Pneumocystis carinii on methenamine silver stain of induced sputum or bronchoalveolar lavage; ELISA/Western blot positive for HIV. CBC: leukopenia with depressed CD4+ cell count. Serum LDH typically elevated.

CXR: diffuse, bilaterally symmetrical interstitial and alveolar infiltration pattern, predominantly perihilar; no lymphadenopathy or effusion.

Congestion and consolidation of lungs with hypoaeration.

Eosinophilic exudate in alveoli with multiple 4- to 6-mm cysts containing oval bodies (merozoites) on lung biopsy or bronchial lavage; Pneumocystis abundant on Gomori methenamine silver stain.

TMP-SMX: pentamidine; steroids for severe disease.

Pneumocystis carinii pneumonia is an opportunistic infection that causes interstitial pneumonia in many immunocompromised patients. Traditionally it has been classified as a protozoan; however, P. carinii ribosomal RNA indicates that the organism is fungal. It is seen in the upper lobes in patients receiving inhaled pentamidine prophylaxis. Treat HIV patients prophylactically with TMP-SMX for P. carinii pneumonia if the CD4 count is < 200.

A 25-year-old HIV-negative homosexual male presents with rectal burning, itching in the anal region, diarrhea, tenesmus, and a bloody, mucopurulent discharge per rectum.

One month ago he was hospitalized with severe febrile proctocolitis that was diagnosed as lymphogranuloma venereum. He has also been treated several times in the past for amebiasis and shigellosis and admits to having receptive anal intercourse. Further history reveals that his most recent sexual partner has been suffering from urethral pain and discharge.

Condylomata acuminata noted in perianal distribution; remainder of physical exam normal.

Gram stain and culture of rectal swab reveals gram-negative diplococci identified as Neisseria gonorrhoeae on Thayer-Martin medium; urethral swab from partner also isolates N. gonorrhoeae.

Sigmoidoscopy: proctitis with bloody mucopurulent discharge noted.

Ceftriaxone and doxycycline (to treat likely concomitant chlamydial infection) for both patient and partner. Most apparent failures of correct antibiotic therapy are in fact due to reinfection; in resistant cases, spectinomycin, fluoroquinolones, or other cephalosporins can be used.

The term “gay bowel syndrome” is used in reference to enteric and perirectal infections that are commonly encountered in immune-competent homosexual men; in homosexuals with HIV, opportunistic organisms play a more important role. Common etiologic agents include Chlamydia trachomatis, lymphogranuloma venereum serovars, Neisseria gonorrhoeae, HSV, Treponema pallidum, human papillomavirus, Campylobacter species, Shigella, Entamoeba histolytica, and Giardia.

PROCTOCOLITIS
ID/CC  A 35-year-old male presents with high fever, malaise, headache, and a hacking cough productive of a small amount of mucoid sputum.

HPI  He has two pet parrots at home who have recently shown signs of illness.

PE  VS: fever; bradycardia. PE: auscultation of chest reveals crepitant rales over both lower lung fields; splenomegaly with mild hepatomegaly noted; multiple purpuric macules seen over abdomen ("HORDER'S SPOTS").

Labs  Greater than fourfold rise in complement-fixing antibody titer to a group antigen suggestive of infection with *Chlamydia psittaci*: definitive diagnosis of psittacosis was made from sputum by isolation of *C. psittaci* in pretreated tissue culture cells.

Imaging  CXR. PA: interstitial patchy, bilateral infiltrates.

Gross Pathology  Principal lesions found in lungs, liver, and spleen.

Micro Pathology  Pulmonary lesion is an interstitial pneumonitis; mononuclear cells with ballooned cytoplasm containing inclusion bodies are observed. In the liver, focal necrosis of hepatocyte occurs along with Kupffer cell hyperplasia.

Treatment  Azithromycin or clarithromycin.

Discussion  Psittacosis is an acute infection caused by *Chlamydia psittaci*: it is characterized primarily by pneumonitis and systemic manifestations and is transmitted to humans by a variety of avian species, principally psittacine birds (parrots, parakeets). A history of contact with birds, particularly sick birds, or of employment in a pet shop or in the poultry industry provides a clue to the diagnosis of psittacosis in a patient with pneumonia, especially if bradycardia and splenomegaly are also present.
A 54-year-old female being treated in the ER is noted to have developed progressively worsening abdominal pain and high-grade fever with chills.

HPI She presented to the ER a few hours ago with colicky abdominal pain and was diagnosed with choledocholithiasis.

PE VS: fever (39.5°C), hypotension (BP 80/60); tachycardia (HR 120). PE: toxic-looking; icteric; abdominal exam reveals extremely tender RUQ with hepatomegaly.

Labs CBC: leukocytosis with neutrophilia. LFTs: markedly elevated bilirubin, AST, ALT, alkaline phosphatase and GGT. Blood cultures grew Escherichia coli.

Imaging CT, abdomen: multiple hepatic abscesses; distended gallbladder with perihepatic and pericholecystic fluid collections.

Treatment Prolonged IV antibiotic therapy; emergent endoscopic (ERCP) or surgical biliary decompression; surgical drainage of the abscesses if no response to IV antibiotics.

Discussion A pyogenic liver abscess is a pus-filled cavity within the liver caused by a bacterial infection, typically polymicrobial. The causes of liver abscess include abdominal infection such as appendicitis, diverticulitis, or perforated bowel; sepsis; biliary tract infection; or liver trauma leading to secondary infection. The most common bacteria involved are E. coli, Klebsiella spp., Enterococcus, Staphylococcus spp., Streptococcus spp., and Bacteroides. Positive blood cultures are found in about half of patients with a pyogenic liver abscess and sepsis is a life-threatening complication. There is significant mortality even in treated patients and mortality is higher in those with multiple abscesses.
A 30-year-old dairy farm worker presents with complaints of fever, headache, cough, pleuritic chest pain, and malaise.

His work at the dairy involves milking cows and looking after parturient cattle.

VS: fever; tachypnea. PE: mild icterus; bilateral crackles on chest auscultation.

CBC: normal WBC count. Mild elevation of serum bilirubin and liver enzymes; greater than fourfold increase in complement-fixing antibody (against Coxiella burnetii) titer between acute and convalescent sera (IFA technique for early detection of specific IgM Ab is the serodiagnostic method of choice); negative Weil-Felix reaction; C. burnetii isolated from sputum by inoculation of cultured human fetal diploid fibroblasts.

CXR: right upper lobe rounded opacity that increased in size over a few days and cleared completely with treatment.

Doxycycline is the first-line agent of therapy (erythromycin can also be used).

Q fever is caused by the rickettsia-like organism Coxiella burnetii and produces the clinical picture of primary atypical pneumonia. Q fever differs from the other human rickettsioses in that rash is absent and transmission is usually by the airborne route. C. burnetii localizes in the mammary glands and uterus of pregnant cattle, sheep, and goats, in which infection is mild or inapparent; infected placentas, postpartum discharges, and the feces of these animals are the principal sources of contaminated material in the environment. Humans acquire Q fever by inhaling aerosolized particles from such substances; particularly at risk are dairy and slaughterhouse workers.
A 12-year-old white female is rushed to the emergency room because of numbness of the right foot and leg followed by fever and convulsions. The child refuses to drink any fluids (hydrophobia).

She had been camping 5 weeks ago. When questioned, her mother recalls that one night the child had apparently stepped on a bat that bit her in the right foot.

VS: no fever. PE: child is disoriented, hyperventilating, extremely agitated, and actively moving all four limbs; thus difficult to restrain; no meningeal signs; fundus normal; saliva viscous and foaming.

LP: lymphocytic pleocytosis with mildly elevated proteins and normal sugar in CSF. Positive rabies antigen in corneal scrapings.

Characteristic cytoplasmic inclusion bodies (Negri bodies) in corneal scrapings or Ammon's horn.

Supportive; almost always fatal; prevent with vaccine; postexposure prophylaxis with diploid cell vaccine and human rabies immune globulin (HRIG).

Rabies is a fatal viral disease that is transmitted to humans by the bites of bats, raccoons, skunks, foxes, coyotes, dogs, and cats. Rabies virus is an enveloped, single-stranded RNA virus. Rabies has a long incubation period (approximately 3 to 8 weeks); death usually results from respiratory failure.
A 27-year-old male researcher presents with sudden-onset fever, chills, headache, a skin rash, and painful swelling of multiple limb joints.

Careful history reveals that he was bitten by a rat in his laboratory a few days ago; the bite wound has now healed.

VS: fever. PE: morbilliform rash noted over extremities, particularly the hands and feet; painful swelling and restriction of movement noted over both wrist and knee joints.

CBC: leukocytosis. *Streptobacillus moniliformis* isolated from blood and synovial fluid of inflamed joints; agglutinins to *S. moniliformis* demonstrated in significant titers.

Amoxicillin/clavulonic acid (doxycycline can also be used).

*Rat bite fever*, which is caused by *Streptobacillus moniliformis*, is an acute febrile illness that is usually accompanied by a skin rash; most cases result from the bites of wild or lab rats, although mice, squirrels, weasels, dogs, and cats may also transmit the disease by bites or scratches. The disease is called Haverhill fever when *S. moniliformis* is transmitted by drinking rat-excrement-contaminated milk. Distribution is probably worldwide, with most cases occurring in crowded cities characterized by poor sanitation.
A 30-year-old male who lives in the western part of the United States presents with high fever, shaking chills, severe headache, myalgias, and diarrhea.

He reports having had similar symptoms 10 days ago that lasted for 4 to 5 days, followed by defervescence accompanied by drenching sweats and marked prostration. He had been hiking in a tick-infested forest until about a week before the development of symptoms.

VS: fever.

Spirochetes found on thick smears of peripheral blood obtained during febrile period and stained with Wright or Giemsa stain.

Doxycycline is the drug of choice (erythromycin may also be used).

Relapsing fever is an acute louse-borne or tick-borne infection that is caused by blood spirochetes of the genus Borrelia; it is characterized by recurrent febrile episodes separated by asymptomatic intervals. Unlike other spirochetes, the etiologic agent can readily be detected with Giemsa stain or Wright's stain. B. recurrentis is the cause of louse-borne relapsing fever, whereas a variety of different species produce the tick-borne disease. In the United States, the predominant species are B. hermsii and B. turicatae. Most patients experience the Jarisch-Herxheimer reaction within the first 2 hours of treatment.
A 6-year-old male presents with fever, intense headache, myalgia, dry cough, and a rash that began peripherally (on his wrists and ankles) but now involves the entire body, including the palms and soles.

The child lives in North Carolina and indicates that he was bitten by an insect a few weeks ago while playing in the woods near his home.

VS: fever; PE: lethargy; ill appearance; petechial rash all over body, including palms and soles.

Gross Pathology
Hemorrhagic necrosis in brain and kidneys; nodular formation in glia.

Micro Pathology
Inflammatory lymphocytic and plasma cell perivascular infiltration; endothelial edema with abundant rickettsiae; microthrombus formation with necrotic vasculitis.

Treatment
Doxycycline or chloramphenicol.

Discussion
*Rickettsia rickettsii* is the causative organism of Rocky Mountain spotted fever; *Dermacentor*, a wood tick, is the vector. The organism's tropism for endothelial cells results in vasculitis, edema, thrombosis, and ischemia. Ironically, Rocky Mountain spotted fever is endemic to the East Coast of the United States.
A 5-month-old male infant is brought to the pediatric clinic with wheezing and respiratory difficulty of 3 hours' duration.

He has had rhinorrhea, fever, and cough and had been sneezing for 2 days prior to his visit to the clinic.

VS: tachypnea, PE: nasal flaring; mild central cyanosis; accessory muscle use during respiration; hyperexpansion of chest; expiratory and inspiratory wheezes; rhonchi over both lung fields.

CBC/PBS: relative lymphocytosis. ABGs: hypoxemia with mild hypercapnia. Normal flora on bacterial culture of sputum; respiratory syncytial virus (RSV) demonstrated on viral culture of throat swab.

CXR: hyperinflation; segmental atelectasis; interstitial infiltrates. Humidified oxygen; bronchodilators; aerosolized ribavirin.

RSV is the most common cause of bronchiolitis in infants under 2 years of age; other viral causes include parainfluenza, influenza, and adenovirus. Infections typically occur during the fall and winter months. Transmission occurs via close contact with contaminated fomites but can also occur after coughing or sneezing. The majority of infections occur during an RSV epidemic.
A 4-month-old girl brought in for a well-child visit is found to be low in weight and height for her age and to have lens opacities (due to congenital cataracts).

HPI Her mother had a skin rash and fever during her first trimester. The mother states that when the child was born, she too had a rash like a “blueberry muffin” and was jaundiced.

PE Deaf and globally retarded; malnourished; microcephaly and bulging anterior fontanelle; microphthalmia with unilateral left cataract; discrete black, patchy pigmentation found in retina on funduscopic exam; hepatosplenomegaly; machinery murmur heard at second intercostal space on left sternal border (due to patent ductus arteriosus).

Labs CBC/PBS: leukopenia; thrombocytopenia. Increased serum bilirubin (both direct and indirect); rubella virus isolated from urine and saliva; markedly increased IgM specific antibody for rubella.

Imaging XR, plain: radiolucent (lytic) bone lesions (metaphyseal).

Treatment None.

Discussion Congenital rubella, transmitted in utero, is caused by rubella virus, a single-stranded RNA togavirus. In children and adults it is a transitory and unremarkable disease. If acquired in utero it has devastating consequences.
ID/CC A 10-year-old female Asian immigrant presents with a low-grade fever and coryza of 3 days' duration.

HPI She also complains of arthralgias and a skin rash that began on her face and spread to her trunk. Her mother says she cannot remember any details of her vaccination history.

PE VS: fever. PE: maculopapular rash over face and trunk; enlarged postauricular, posterior cervical, and occipital lymph nodes.

Labs CBC: leukopenia; thrombocytopenia. Rubella virus hemagglutination inhibition test demonstrates fourfold rise in titer to 1:32.

Gross Pathology Erythematous skin rash resembling rubeola measles but lighter in color and more discrete; similar distribution pattern in both.

Treatment Symptomatic treatment.

Discussion Rubella (German measles) is caused by a togavirus. Live attenuated rubella virus vaccine (part of MMR) should be given to all infants and to susceptible girls before menarche. The course of illness is self-limiting and mild; in females the major implication is the potential for congenital rubella syndrome. Females with rubella can get polyarthritis secondary to immune complex deposition.

Atlas Link IM2-026
ID/CC A 14-year-old male who is known to have sickle cell anemia presents with throbbing pain, redness, and swelling of the right thigh.

HPI The patient also complains of fever and chills of 1 week's duration. He has a few pet turtles at home.

PE VS: fever, tachycardia, PE: pallor; redness, swelling, and tenderness over right thigh; effusion demonstrated in right knee joint; limitation of range of motion of right knee.

Labs CBC: leukocytosis; elevated ESR. PBS: irreversible sickling; blood culture reveals *Salmonella typhimurium* (most common); organism also isolated from pus aspirated from right femur (diagnostic of osteomyelitis).

Imaging Nuc: increased uptake in metaphyseal region of right femur. XR (usually normal during the first 10 days of illness) may reveal changes of bone resorption, detached necrotic cortical bone (SEQUSTRUM), and laminated periosteal new-bone formation (INVOLUCRUM).

Gross Pathology Dense, pale, sclerotic-appearing area in shaft.

Micro Pathology Changes include suppurative and ischemic destructive necrosis, fibrosis, and ultimate bone repair.

Treatment Parenteral antibiotics, with fluoroquinolones being first-line agents (third-generation cephalosporins may also be used).

Discussion A striking association has been noted between diseases producing hemolysis (e.g., sickle cell anemia, malaria, and bartonellosis) and salmonella infections; elderly patients with impaired host defense mechanisms, those with hepatosplenic schistosomiasis, and AIDS patients are also at increased risk of severe and recurrent salmonella bacteremia. *Salmonella* osteomyelitis in sickle cell patients presents primarily in young individuals and typically affects long bones. It is believed that the functional asplenic state found in most sickle cell patients contributes to the increased prevalence of salmonella osteomyelitis.
**ID/CC** An Asian refugee family comprising a 30-year-old man, his wife, and two schoolchildren present with complaints of severe itching over their entire bodies except for their face; the itching increases during the night.

**HP** The male family members also report penile and scrotal skin lesions. The family is of low socioeconomic status and lives in a single room under crowded conditions.

**PE** Papulovesicular lesions; "burrows" seen in the dorsal interdigital web spaces and flexor aspects of both wrists; lesions also seen around elbows, anterior axillary folds, periumbilical area, lower buttocks, and thighs; face was spared; scrotal and penile lesions seen in male members were nodular and reddish.

**Labs** Female adult mite was seen with a hand lens when teased out of her burrow with a needle.

**Treatment** Apply lindane or permethrin (lindane is contraindicated in small children and in pregnant women). All family members must be treated; clothing, linen, and the like should be boiled and washed; fingernails should be trimmed. Use antihistamines or calamine lotion to help control itching.

**Discussion** Scabies is caused by infestation with *Sarcoptes scabiei*, a mite that bores into the cornal layer of the skin, forming burrows in which it deposits its eggs. The scabies organism does not survive for more than 48 hours away from the host; modes of transmission include close contact with infected individuals, unsanitary conditions, and sexual contact. In adults, certain areas of the body are generally spared, including the face, scalp, and neck.

**Atlas Links** [M-M2-049] [MC-185]
ID/CC  A 10-year-old white female complains of difficulty swallowing, pain in both ears, and fever of 1 week’s duration; she also complains of an extensive skin rash.

HPI  The child is fully immunized and has been well until now.

PE  VS: fever. PE: extensive erythematous rash ("GOOSE-PIMPLE SUNBURN") on neck, groin, and axillae; desquamation and peeling of fingertips; circumoral pallor; lines of hyperpigmentation with tiny petechiae (PASTIA'S SIGN) in antecubital fossae; bright red lingual papillae superimposed on white coat ("STRAWBERRY TONGUE"); pharyngitis with exudative tonsillitis; cervical lymphadenopathy; normal eardrums.

Labs  CBC: leukocytosis with neutrophilia. Group A β-hemolytic Streptococcus pyogenes on throat swab and culture; elevated ASO titer.

Gross Pathology  Toxin-induced vasodilation; complications include otitis media, pneumonia, glomerulonephritis, osteomyelitis, and rheumatic fever.

Micro Pathology  Inflammatory polymorphonuclear epidermal infiltrate; interstitial nephritis; lymph node hyperplasia.

Treatment  Penicillin; erythromycin.

Discussion  Scarlet fever is a streptococcal infection that is characterized by morbilliform rash due to hypersensitivity to erythrogenic toxin.

A 27-year-old Peace Corps volunteer working in the Congo is sent home after developing fever, sweats, and abdominal pain that have not responded to antimalarial treatment.

HPI
Five weeks ago, he developed severe itching and a macular rash (swimmer's itch) after swimming in a nearby pond.

PE
VS: fever. PE: moderate enlargement of liver and spleen; tender abdomen but no peritoneal irritation.

Labs
CBC/PBS: marked eosinophilia. Characteristic large parasite eggs with lateral spines may be found in stool specimen.

Imaging
Sigmoidoscopy: swollen and erythematous mucosa; many small ulcerations. CT/US, abdomen: hepatosplenomegaly; portal vein dilatation.

Gross Pathology
Skin and liver sites of principal lesions in acute stage; eggs may be found in liver, lungs, intestines, pancreas, spleen, urogenital organs, and brain; chronic stage characterized by granuloma formation in bladder and liver (periportal fibrosis).

Micro Pathology
Granulomatous reaction and fibrosis.

Treatment
Praziquantel.

Discussion
Schistosomiasis is among the most common parasitic diseases in the world; infection with Schistosoma mansoni or S. japonicum is acquired by swimming in snail-infested ponds and lakes. Long-standing infection may lead to noncirrhotic portal fibrosis and portal hypertension. Also known as bilharziasis.

Atlas Link [M2-051]
A 12-year-old immigrant from the Middle East presents with terminal hematuria, dysuria, and increased frequency of micturition.

He remembers having played and bathed in snail-infested streams while he was in his native country; on one occasion he had developed an intensely pruritic skin eruption after bathing in one such stream ("cercarial dermatitis").

Pallor noted.

UA: hematuria; mild proteinuria and sterile (abacterial) pyuria. Microscopic exam of urine and rectal biopsy reveals presence of ellipsoid eggs with a sharp terminal spine containing a miracidium surrounded by a thick, rigid shell.

XR: bladder wall calcification.

Praziquantel, metrifonate.

Three major species exist. Schistosoma mansoni, S. japonicum, and S. haematobium infect humans. S. mansoni is found in Africa, the Arabian Peninsula, South America, and parts of the Caribbean; S. japonicum is found in Japan, China, and the Philippines; and S. haematobium is found in Africa and the Middle East.

Transmission of schistosomiasis cannot occur in the United States because of the absence of the specific freshwater snail that is an intermediary host. In S. haematobium infection, the principal symptoms are terminal hematuria, dysuria, and frequent urination; hydronephrosis, pyelonephritis, and squamous cell carcinoma of the urinary bladder may develop as complications. In S. mansoni and S. japonicum infection, manifestations may include fever, malaise, abdominal pain, diarrhea, or hepatosplenomegaly. Presinusoidal hepatic trapping of eggs and the consequent granulomatous reaction induce portal hypertension.
A 36-year-old male executive comes to the emergency room because of the development of sudden nausea, vomiting, and diarrhea with blood and mucus (dysentery) as well as crampy abdominal pain for 2 days.

**HPI**
He had just returned from a business trip in South America.

**PE**
VS: low-grade fever. PE: mild dehydration; hyperactive bowel sounds; tender abdomen without definite peritoneal irritation.

**Labs**
Leukocytes on stool examination; *Shigella* isolated on stool culture; on microbiology, organism does not ferment lactose and is not motile.

**Micro Pathology**
*Shigella* enterotoxin acts by activating adenylate cyclase; organism invades intestinal mucosa.

**Treatment**
Rehydration with antibiotic therapy (ampicillin or TMP-SMX).

**Discussion**
Shigellosis outbreaks occur primarily in areas with overcrowding and poor hygiene (fecal-oral transmission); arthritis, conjunctivitis, and urethritis (Reiter’s syndrome) may be complications in HLA-B27-positive individuals. Like *Salmonella*, *Shigella* causes bloody diarrhea by invading the intestinal mucosa, causing intestinal ulceration and inflammation.
A 56-year-old hospitalized male is found to have an abrupt-onset high-grade fever with chills a few hours after he underwent nephrolithotomy.

**HPI**

He was diagnosed with chronic nephrolithiasis with recurrent UTIs; a surgery intern also noted poor urine output.

**PE**

VS: fever; tachycardia; hypotension; tachypnea. PE: confused and disoriented; hyperventilating; diaphoresis; hands warm and pink with rapid capillary refill; pulse bounding; on chest auscultation, air entry found to be bilaterally reduced.

**Labs**

CBC: leukocytosis with left shift; neutrophils contain toxic granulations, Döhle bodies, and cytoplasmic vacuoles; band forms > 10%; thrombocytopenia. Prolongation of thrombin time, decreased fibrinogen, and presence of D-dimers (suggesting DIC); raised BUN and creatinine. ABGs: metabolic acidosis (increased anion gap due to lactic acidosis) and hypoxemia (due to ARDS). Blood and urine culture yields *Escherichia coli*.

**Imaging**

CXR: evidence of noncardiogenic pulmonary edema (ARDS).

**Treatment**

IV antibiotics (with adequate gram-negative coverage); management of multiorgan failure (azotemia, ARDS, and DIC).

**Discussion**

Almost any bacterium can cause a bacteremia, including *E. coli* (most common), *Klebsiella*, *Proteus*, *Pseudomonas* (associated with antibiotic therapy and burn wounds), *Bacteroides fragilis* (causes of anaerobic septicemias), *Staphylococcus aureus*, *Streptococcus pneumoniae*, and pediatric septicemia due to *E. coli* and *Streptococcus agalactiae*. Gram-negative bacteria release endotoxins; the release of endotoxin into the circulation leads to the activation of macrophages and monocytes, which in turn release cytokines. These cytokines trigger cascade reactions that lead to the clinical and biochemical manifestations of the sepsis syndrome.

**Atlas Links**

[Link M-M2-054A, M-M2-054B]
A 37-year-old gardener complains of lumps with red streaks on his arm and swelling of the axillary lymph nodes.

Two weeks ago, he pricked his hand with a thorn while pruning roses. A nodule then formed which subsequently ulcerated and filled with pus.

Nonpainful nodular lesion on dorsum of hand with ulcer formation and suppurative; tender, palpable inflammation and hardening of lymph vessels (lymphangitis); swelling, inflammation, and suppurative of lymph nodes (lymphadenitis); nonulcerated satellite nodules along course of lymphatics.

Cigar-shaped budding cells (Sporothrix schenckii) visible in pus; diagnosis confirmed by culture of aspirate of nodule.

Nonpainful, soft, ulcerated nodule at inoculation site (sporotrichotic chancre); may extend to deep tissues and bone with osteitis and synovitis.

Usually area of suppuration with polymorphonuclear infiltrate surrounded by granulomatous reaction of varied intensity with epithelioid and giant cell formation; chlamydorescere asteroid bodies present.

Itraconazole; potassium iodide.

Also called “rose gardener’s disease,” sporotrichosis is a fungal infection caused by Sporothrix schenckii; a dimorphic fungus that lives on vegetation. It is typically transmitted by a thorn prick and causes localized infection with few systemic manifestations.

Atlas Link  [HHS] M-M2-055
A 7-year-old girl is seen by the embassy doctor in Nigeria for abdominal pain, diarrhea, fever, dry cough, and marked dyspnea of 2 weeks' duration.

She is the daughter of an American diplomat working in Nigeria. Despite her parent's admonitions, she frequently walks barefoot.

VS: fever. PE: moderate respiratory distress; no cyanosis; no clubbing; coarse, crepitant rales and wheezing heard over both lung fields; mild abdominal tenderness.

CBC/PBS: marked eosinophilia. Typical motile rhabditiform larvae on sputum exam as well as in freshly passed stool; positive filarial complement fixation test.

CXR: bilateral, transient migratory infiltrates.

Pneumonitis produced by migration of larvae through respiratory tract.

Ivermectin, thiabendazole.

Strongyloidiasis is seen in the presence of poor hygiene and in tropical countries. Larvae penetrate the skin, gaining entrance to the venous system and to the lungs, and then ascend to enter the GI tract.
A 54-year-old white female complains of spiking fever, chills, loss of appetite, several bouts of diarrhea, and right upper quadrant pain.

Ten days ago she underwent an apparently uncomplicated emergency surgery for suppurative cholecystitis and was subsequently discharged and sent home.

VS: fever; PE: pallor; slight icterus; pain on percussion of right costal region; well-healed surgical wound with no evidence of infection; liver not palpable; crepitant rales on right lung base.

CBC: elevated WBC count (17,000) with predominance of neutrophils.

CXR: elevated right hemidiaphragm; slight right pleural effusion. US/CT: complex fluid collection below diaphragm.

Percutaneous drainage under ultrasonic or fluoroscopic guidance followed by regular blood and radiologic exams; surgical exploration and drainage.

Subdiaphragmatic abscess most commonly occurs after abdominal surgery, mainly with septic, emergency procedures; it typically presents 1 week or more postoperatively.
A 6-week-old male, the son of a prostitute, is brought to the family doctor because of persistent, sometimes bloody mucopurulent nasal discharge, anal ulcers, and a generalized rash.

HPI
The child was delivered at home, and the mother did not receive any prenatal care.

PE
Weak-looking, icteric infant with hoarse cry; does not move right limb (pseudoparalysis); bloody purulent discharge evident at nares; generalized lymphadenopathy; hepatosplenomegaly; maculopapular rash with desquamation on back and buttocks; bullae on hands and feet.

Labs
CBC: anemia. VDRL in both mother and child positive; direct hyperbilirubinemia: negative Coombs' test; Treponema pallidum seen on nasal exudate and anal ulcers.

Imaging
XR. plain: periositis of long bones; bilateral moth-eaten lesions; focal defect in proximal tibial epiphysis with increased density of epiphyseal line (Wimberger's sign).

Gross Pathology
Pathologic features seen if neonatal disease is left untreated include syphilitic chondritis and rhinitis (causes saddle-nose deformity), pathologic fractures, bowing of the tibia (SABER SIGN), V-shaped incisors (HUTCHINSON'S TEETH), multicupid molars (MULBERRY MOLARS), interstitial keratitis, and deafness.

Treatment
Penicillin.

Discussion
Treponema pallidum is a spirochete; in utero vertical transmission occurs from an infected mother to the fetus. Congenital syphilis occurs maximally during 16 to 36 weeks of gestation and may be the cause of stillbirth. It is preventable if the mother has received adequate treatment.
An 18-year-old male presents with a painless ulcer on his penis.

He admits to having had unprotected intercourse with a prostitute 3 weeks ago.

Painless, single, rounded, firm papule with well-defined margins on dorsal aspect of glans penis that ulcerates ("hard chancre"); nontender, rubbery bilateral inguinal lymphadenopathy.

Treponemes on dark-field examination of exudate from chancre; VDRL positive; FTA-ABS positive; ELISA for HIV negative.

A 1.2-cm ulcerated papule with rolled edges and induration; regional lymphadenopathy.

Capillary dilatation with plasma cell, PMN, and macrophage infiltration; fibroblastic reaction.

Benzathine penicillin G 1M, 2.4 MU single dose.

An STD caused by Treponema pallidum, a spirochete, primary syphilis is characterized by the appearance of a painless chancre in the area of inoculation. If left untreated, secondary and tertiary syphilis may ensue. Other STDs, such as AIDS, are more prevalent in patients with syphilis.
A 23-year-old female presents with a **nonpruritic skin eruption**, hair loss, and generalized fatigue and weakness.

She admits to having had **multiple sexual partners** and unprotected sex. She has had two spontaneous abortions.

**Extensive raised, copper-colored, maculopapular, desquamative rash on palms and soles; generalized nontender lymphadenopathy with hepatosplenomegaly; large, pale, coalescent, flat-topped papules and plaques** in groin (**condylomata lata**); dull, erythematous mucous patches in mouth; hair loss (**alopecia**) in tail of eyebrows.

Skin lesions, mucous patches in mouth, and condylomata lata positive for **treponemes; positive VDRL; positive FTA-ABS; ELISA negative for HIV; CSF VDRL negative.**

**IM benzathine penicillin G.**

**Sexual partners must be treated.**
A 54-year-old man presents with ataxia, mental status changes, grossly deformed ankle joints, and shooting pains in his extremities.

He remembers having had a "boil" on his penis (PRIMARY SYPHILIS CHANCRAE) many years ago that went away by itself. He also recalls having had a scaling rash on the soles of his feet and the palms of his hands (due to secondary syphilis) some time ago.

Painless subcutaneous granulomatous nodules (GUMMAS); reduced joint position and vibration sense in both lower extremities (due to bilateral dorsal column destruction); loss of deep tendon reflexes in both lower limbs; loss of pain sensation and deformed ankle and knee joints with effusion (CHARCOT'S NEUROPATHIC ARthropathy); broad-based gait, positive Romberg's sign (due to sensory ataxia); pupillary light reflex lost but accommodation reflex retained (ARGYLL ROBERTSON PUPILS).

Positive VDRL and Treponema pallidum hemagglutination assay (TP-HA). LP: pleocytosis and increased proteins in CSF; VDRL positive. Normal blood glucose levels.

CXR: "tree-bark calcification" of ascending aorta.

Obliterative endarteritis and meningoencephalitis.

Proliferation of microglia; demyelination and axonal loss in dorsal roots and columns.

Penicillin.

Tabes dorsalis usually develops 15 to 20 years after initial infection. There may also be visceral involvement (can cause neurogenic bladder).

IM2-028
A 12-year-old white male presents with stiffness of the jaw and neck along with inability to swallow.

HPI Twelve days ago he stepped on a rusty nail, which produced a small puncture wound; the area is now red, hard, and swollen with pus. He has been experiencing tingling sensations and spasms in his calf muscles. He has not received any immunizations within the past 10 years.

PE Jaw muscle rigidity (trismus); facial muscle spasm (risus sardonicus); dysphagia; neck rigidity; normal deep tendon reflexes; profuse sweating; patient alert, apprehensive, restless, and hyperactive during PE; loud noise elicits painful spasms of face, neck, abdomen, and back, the latter producing opisthotonos.

Labs CBC, CSF, blood chemistries normal.

Gross Pathology There may be fractures of ribs or vertebrae with sustained spasms.

Treatment Surgical debridement of wound; tetanus immune globulin intramuscularly or intrathecally; diazepam; phenobarbital; tetanus toxoid; penicillin IV.

Discussion Tetanus is caused by tetanospasmin, a neurotoxin produced by Clostridium tetani, an obligate anaerobic, spore-forming, gram-positive rod; the toxin blocks the release of the inhibitory neurotransmitter glycine in the anterior horn cells. Tetanus often occurs in IV drug abusers; neonates of nonimmunized mothers may become infected through the umbilical cord stump. The disease may occur even years after injury or infection and may also involve the autonomic nervous system (arrhythmias, high/low blood pressure).

Atlas Link M-M2-062
A 15-day-old infant is brought by his mother to the pediatric emergency room in a state of marked muscle rigidity and spasm.

The mother is illiterate and did not receive any prenatal care; the delivery was conducted at home and, according to her, was uneventful and full term. The child did not receive any immunizations; directed questioning reveals that he has been crying excessively for the past 2 weeks and has not been feeding normally.

Extremely ill-looking infant in a state of generalized rigidity and opisthotonus; on slightest touch or noise, spasm worsens and he develops a stridor and becomes cyanosed.

Diagnosis is largely clinical; culture of umbilical stump yields Clostridium tetani.

Ventilatory assistance; supportive management; maintenance of nutritional, fluid, and electrolyte balance; tetanus antitoxin; control of tetanic spasms with diazepam.

Tetanus neonatorum accounts for a considerable proportion of infant deaths in developing countries, primarily owing to the lack of availability of good prenatal care (no tetanus immunization); untrained birth attendants in rural areas use contaminated material to cut or anoint the umbilical cord. Tetanus is caused by Clostridium tetani, a gram-positive, motile, nonencapsulated, anaerobic, spore-bearing bacillus; its effects are mediated through production of a powerful neurotoxin (tetanosasmin). The toxin acts principally on the spinal cord, altering normal control of the reflex arc by suppressing the inhibition regularly mediated by the internuncial neurons.
A 40-year-old male who recently went hiking in a forest in the 
western United States presents with symmetric weakness of the 
lower extremities that has now progressed over the past few days 
to involve the trunk and the upper arms.

The patient does not report any sensory symptoms.

Higher mental functions intact; symmetric flaccid paralysis with an ascending pattern of spread noted; no sensory loss demonstrated; on careful examination of hairy areas of the body, a tick was found embedded in the scalp.

LP: CSF normal. EMG: nerve conduction velocity and compound 
muscle action potentials decreased.

Tick was detached without being squeezed, and this led to 
resolution of symptoms over the next few days.

Feeding ticks may elaborate a neurotoxin that causes tick paralysis; symmetric weakness of the lower extremities progresses to an ascending flaccid paralysis over several hours to days, although the sensorium remains clear and sensory function is normal.
A 40-year-old male diagnosed with AIDS presents with a severe headache.

He suffered a grand mal seizure 2 hours before his arrival in the emergency room. He denies any history of seizures and adds that he has many pets, including cats.

Generalized lymphadenopathy; bilateral papilledema; left-sided hemiparesis with hyperactive deep tendon reflexes on left side; positive Babinski’s sign on left side.

Positive indirect fluorescent antibody test for toxoplasmosis; positive Sabin-Feldman dye test.

MR/CT, head; single or multiple rounded mass lesions with ring or nodular enhancement.

Large brain abscesses with concomitant focal neurologic abnormalities, seizures, or altered mental status.

Parasites appear in tissue as tachyzoites or encysted bradyzoites; aggregates of nonencapsulated organisms constitute pseudocysts.

Pyrimethamine; sulfadiazine.

The definitive host of Toxoplasma gondii is the domestic cat. The intermediate hosts are many and varied, including humans. Toxoplasmosis is also transmitted by ingestion of raw or undercooked meat.
A 50-year-old man presents with generalized myalgia and a persistent low-grade fever.

In addition, the patient recalled having severe abdominal pain and diarrhea several weeks ago. The patient worked in a pig slaughterhouse for many years.

VS: fever. PE: periorbital and facial edema; tenderness over calf, thigh, and shoulder muscles; conjunctival and splinter hemor- rhages; no neurologic deficit seen.

CBC: eosinophilia. Normal ESR; elevated serum CPK, LDH, and AST; latex agglutination test positive for *Trichinella*.

Facial, neck, biceps, lower back, and diaphragm most frequently affected muscles.

Biopsy of sternocleidomastoid muscle reveals cysts of *Trichinella spiralis*.

Albendazole; mebendazole; high-dose corticosteroids.

The organism causing trichinosis, *Trichinella spiralis*, can be transmitted when raw or undercooked pork is ingested. The larvae develop only in striated muscle cells.

TRICHINOSIS
A 6-year-old male is brought to the ER in a delirious state with fever and marked dyspnea that have rapidly progressed over the past 2 days.

His mother, an Asian immigrant, was diagnosed and treated for pulmonary tuberculosis a few months ago. He has had a low-grade fever, cough, malaise, and night sweats for the past 2 months. The child has not received prophylactic isoniazid or BCG vaccination.

VS: fever; tachycardia; marked tachypnea; hypotension. PE: toxic and stuporous; pallor; central cyanosis; extensive rales and rhonchi bilaterally; hepatosplenomegaly; lymphadenopathy; funduscopy reveals choroidal tubercles.

CBC: lymphocytosis; normochromic, normocytic anemia. Increased ESR. Mantoux skin test negative (false negative may occur during incubation and with severe disease); staining and culture of transbronchial and bone marrow biopsy specimens reveal presence of Mycobacterium tuberculosis; PCR for tuberculosis positive; ELISA for HIV negative.

CXR: soft, uniformly distributed fine nodules throughout both lung fields (miliary motting).

Myriad 1- to 2-mm granulomas demonstrable in lungs, liver, and bone marrow biopsy specimens.

Granulomas with central caseous necrosis surrounded by epithelial cells, Langerhan’s cells, lymphocytes, plasma cells, and fibroblasts in affected tissues.

Multidrug antitubercular therapy with isoniazid, rifampicin, pyrazinamide, and ethambutol or streptomycin; steroids may be indicated.

Miliary tuberculosis results from widespread hematogenous dissemination and often presents with a perplexing fever, dyspnea, anemia, and splenomegaly; the disease is more fulminant in children than in adults.
A 14-year-old male immigrant complains of malaise, weight loss, fever, and night sweats of 6 weeks' duration; he also has a mild cough that began to produce bloody sputum 3 days prior to his admission.

**HPI**
The patient's mother has been diagnosed with pulmonary tuberculosis and is currently receiving treatment for it.

**PE**
VS: mild fever. PE: malnourished; low height and weight for age; bronchial breath sounds with crepitant rales heard over right supramammary area.

**Labs**
CBC/PBS: normocytic, normochromic anemia; WBC count normal with relative lymphocytosis. Increased ESR; sputum stained with ZN stain positive for acid-fast bacilli; positive radiometric culture for *Mycobacterium tuberculosis*; positive ELISA for TB; positive intradermal tuberculin injection (MANTOUX TEST).

**Imaging**
CXR: small cavity with streaky infiltrates in right upper lobe; hilar lymphadenopathy; calcified lung lesion (GHON'S LESION); Ghon's lesion and calcified lymph node (RANKE COMPLEX).

**Gross Pathology**
Primary tuberculosis usually consists of lesions in lower lung lobes and in subpleural locations; cavitation rare; secondary TB or reinfection characterized by cavitary lesions usually located in apical regions.

**Micro Pathology**
Multinucleated epithelioid Langerhan's cells surround core of cascading necrosis in lung parenchyma, producing fibroblastic reaction at periphery with lymphocytic infiltration and proliferation (TUBERCLE).

**Treatment**
Multiple drug therapy with isoniazid (INH), rifampin, ethambutol, pyrazinamide, and/or streptomycin.

**Discussion**
Pulmonary tuberculosis is caused by *Mycobacterium tuberculosis*, an acid-fast, gram-positive aerobic bacillus. An increasing incidence in AIDS patients has been observed; drug resistance is becoming common.

**Atlas Links**
A 12-year-old white male is brought to his pediatrician because of an ulcer on his right wrist together with swelling of the lymph nodes in the right axillae with suppuration.

He had just returned from summer camp and, upon questioning, admits to having played with rabbits at the camp’s breeding grounds. He has been suffering from fever, headache, and muscle aches for almost a week.

VS: fever. PE: indurated erythematous nodule with ulcer formation on right wrist; right axillary adenopathy with pus formation; lymphangitis; mild splenomegaly; scattered rales in both lung bases.

CBC: normal WBC count. Increased ESR; elevated C-reactive protein; positive agglutination test; Francisella tularensis on direct fluorescent antibody staining of material from ulcer.

CXR: bilateral basilar interstitial infiltrates.

Enlarged, indurated lymph nodes with necrosis and suppuration; skin nodule at site of inoculation with ulcer formation.

Necrosis and suppuration of lymph nodes; pulmonary and disseminated lesions; granulomatous nodules with central caseating necrosis.

Streptomycin and gentamicin.

Tularemia is an acute zoonosis caused by Francisella tularensis, a nonmotile, aerobic, gram-negative bacillus; it is transmitted through contact with rabbits, squirrels, or other rodents or tick bites. It may be ulceroglandular, tonsillar, ocular, pneumonic, or typhoidal.
A 27-year-old male is admitted to the hospital for evaluation of increasing fever of unknown origin along with malaise, headache, sore throat, cough, and constipation.

He visited Southeast Asia 3 weeks ago but did not receive any prior vaccinations.

VS: bradycardia; fever; fever charting reveals “stepladder” pattern. PE: mild hepatosplenomegaly; faint erythematous macules seen over trunk (“rose spots”).

CBC: neutropenia with relative lymphocytosis. Widal’s test positive in significant titers; blood and stool cultures reveal Salmonella typhi.

Infection of Peyer’s patches in terminal ileum leads to necrosis of underlying mucosa, producing longitudinal oval ulcerations.

Ulcers bordered by mononuclear cells; typhoid nodules with lymphocytes and macrophages may be present in liver, spleen, and lymph nodes.

Ciprofloxacin is curative.

Because infection is acquired from contaminated food or water, typhoid vaccine is recommended for all those traveling to areas that have had typhoid epidemics. Three vaccines are available: the parenteral vaccine containing the capsular polysaccharide and the oral vaccine containing live attenuated organisms are more effective than the parenteral vaccine containing whole killed organisms. S. typhi is transmitted only by humans, whereas all other Salmonella species have an animal as well as a human reservoir.
A 19-year-old male goes to his health clinic complaining of painful urination and discharge.

The patient had casual sex with a classmate while at a party 2 weeks ago. He has had no previous STDs.

Watery yellowish-green discharge from meatus; no penile ulcerations or inguinal lymphadenopathy.

Numerous neutrophils but no bacteria on Gram stain of discharge; positive direct immunofluorescence using monoclonal antibody against Chlamydia; routine bacterial cultures, including Thayer-Martin, do not show growth.

Tetracycline; doxycycline; azithromycin; treat both patient and sexual partner.

The most common cause of nongonococcal urethritis is Chlamydia trachomatis; less frequently it is caused by Ureaplasma urealyticum. It is frequently coincident with gonococcal urethritis.
ID/CC A 25-year-old sexually active female complains of burning on urination.

HPI She also complains of pain in the lower abdomen and increased frequency of urination.

PE Mild suprapubic tenderness.

Labs UA: mild proteinuria; hematuria; WBCs but no casts seen. Urine culture reveals > 100,000 Escherichia coli organisms present.

Gross Pathology Infection ascends the urinary tract (urethritis, cystitis, pyelonephritis); mucosal hyperemia and edema.

Micro Pathology Urothelial hyperplasia and metaplasia.

Treatment Ciprofloxacin.

Discussion Eighty percent of UTIs are caused by E. coli; Staphylococcus saprophyticus is the second most common cause. Other causes, in order of frequency, are Proteus, Klebsiella, Enterobacter, Serratia, Pseudomonas, and Enterococcus; Chlamydia and Neisseria are also causes of urethritis. Risk factors include female gender, sexual activity, pregnancy, obstruction, bladder dysfunction, vesicoureteral reflux, and catheterization.
A 25-year-old sexually active woman presents with burning during micturition (dysuria), increased frequency and urgency of micturition, and low-grade fever.

She is otherwise in perfect health.

VS: fever.

UA: abundant WBCs; mild proteinuria but no casts; staining of sediment reveals presence of gram-positive cocci. Urine culture isolates coagulase-negative Staphylococcus saprophyticus.

Antibiotics (ampicillin, cotrimoxazole, or ciprofloxacin).

Enterobacteriaceae such as Escherichia coli, Klebsiella species, and Proteus and Pseudomonas species are the most common organisms causing UTI. After E. coli, Staphylococcus saprophyticus is the most common cause of primary nonobstructive UTI in sexually active young women.
ID/CC  A 5-year-old male presents with malaise, anorexia, fever, and a pruritic rash on his scalp, face, and trunk.

HPI  He also complains of a headache. Six of his classmates recently missed school because of similar symptoms.

PE  VS: fever (39°C). PE: skin lesions consist of macules, papules, vesicles, pustules, and scabs, all present at same time, predominantly over trunk, face, and scalp.

Labs  Multinucleated giant cells on scraping samples from vesicles. CBC: leukopenia.

Gross Pathology  Macular, papular, vesicular, and pustular rash with scab formation; characteristically, all lesions present at same time (vs. variola); lesions appear in crops every 3 to 5 days; myocarditis and pneumonitis may be present.

Micro Pathology  Intranuclear, acidophilic inclusion bodies (Lipschutz bodies) in epithelial cells with clear halo around them and multinucleated giant cells on histologic exam of skin lesions (on Tzanck smear).

Treatment  Acetaminophen; antihistamines and calamine lotion; hygienic measures, including isolation.

Discussion  A highly contagious dermatrophic viral disease caused by varicella-zoster virus, a DNA herpesvirus, chickenpox is transmitted by respiratory aerosol or by direct contact. Complications include secondary bacterial infection of the skin and pneumonia; high-risk individuals may be protected passively with immunoglobulin and/or acyclovir.

Atlas Links  VARICELLA (CHICKENPOX)
An 8-year-old male is brought to a physician with complaints of impairment of vision in the left eye, urticarial skin rashes, and ill-defined muscle aches.

The child’s mother has caught the child eating dirt or soil on many occasions (FICA). The family also has a pet dog at home.

Rounded swelling near the optic disk seen on fundus exam of left eye; urticarial wheals observed on extremities and trunk; mild hepatosplenomegaly noted.

Leukocytosis with marked eosinophilia; enzyme immunoassay using extracts of excretory-secretory products of *Toxocara canis* larvae positive.

Biopsy of liver reveals larvae with granuloma and eosinophilic infiltration.

Diethylcarbamazine; albendazole or mebendazole; steroids to control symptomatic inflammatory response; laser photocoagulation of visible ocular larvae.

When the nondefinitive human host is infected with parasites that normally infect animals, the parasites do not mature completely, but the larvae introduced persist and induce an inflammatory reaction. The syndrome of **visceral larva migrans** develops when nematode larvae of animal parasites (mostly cat or dog ascarids such as *Toxocara canis*) migrate in human tissues; the syndrome of **cutaneous larva migrans** (creeping eruption) develops when the larvae of various parasites (including the dog or cat hookworm *Ancylostoma braziliense*) penetrate human skin and form pruritic, serpiginous cutaneous lesions along the migratory tracts of the larvae.
A 2-year-old female is brought to the emergency room because of paroxysms and multiple coughs in a single expiration, followed by a high-pitched inspiratory whistle or whoop.

For the past 2 weeks she has had a runny nose, low-grade fever, muscle pains, and headache. Her immunization schedule is incomplete.

VS: fever. PE: child apprehensive and becomes cyanotic during cough paroxysm; thick green mucus expelled with cough; conjunctival injection.

CBC: marked leukocytosis with lymphocytosis. *Bordetella pertussis* on fluorescent antibody staining of nasopharyngeal secretions; diagnosis confirmed by culture on Bordet-Gengou medium.

Small conjunctival and brain hemorrhages may appear during paroxysms; bronchiectasis may also be a complication.

Signs of acute inflammation in upper respiratory tract mucosa, with erythema, petechiae, polymorphonuclear infiltrate, and necrosis.

Largely supportive; erythromycin.

A bacterial infection of the upper respiratory tract caused by *Bordetella pertussis*, a gram-negative coccobacillus, whooping cough is transmitted by droplets and comprises three stages: prodromal (catarrhal), paroxysmal (coughing), and convalescent. It is largely preventable with universally administered diphtheria toxoid, tetanus toxoid, and pertussis a cellular (DTP) vaccine. Pertussis toxin is a heat-labile exotoxin in which ADP ribosylates the inhibitory G protein, thus inactivating it and leading to constant activation of adenylate cyclase and increased cAMP. The remarkable lymphocytosis is due to pertussis toxin inhibiting chemokine receptors. As a result, lymphocytes are unable to leave the bloodstream.
A 10-year-old child who lives in tropical Africa presents with multiple papillomatous skin lesions and pain in both legs.

**HPI**
The first lesion had appeared on the leg as a small indurated papule that ulcerated into a granulomatous papilloma.

**PE**
Multiple papillomatous skin lesions seen, especially in intertriginous areas; lesions were painless and exuding a serous fluid; painful hyperkeratotic lesions seen on palms and soles; both tibia were tender to palpation.

**Labs**
Dark-field microscopic examination of exudate from lesions established the diagnosis by revealing organisms with the characteristic morphology and rotational motion of pathogenic treponemes; non-treponemal serologic tests (i.e., VDRL and RPR tests) and treponemal tests (i.e., FTA-ABS test) were positive.

**Imaging**
XR, legs: evidence of periostitis of the tibia.

**Treatment**
Long-acting intramuscular benzathine penicillin G is the treatment of choice.

**Discussion**
Yaws, the most common of the nonvenereal treponematoses, is a chronic infection of skin and bones caused by *Treponema pertenue*. Yaws occurs in tropical areas of Africa, Asia, and Central and South America; it is principally a disease of childhood, and initial infection occurs between 5 and 15 years of age. Transmission is by direct contact with infected skin lesions containing treponemes and is fostered by conditions of overcrowding and poor hygiene. The disease may occur in three stages: primary, secondary, and tertiary. Only lesions of primary and secondary yaws are infectious.
ID/CC  A 24-year-old white South American male develops sudden high fever, chills, generalized aches and pains, retro-orbital headache, nausea, and vomiting.

HPI  He gradually improves, but the fever returns 4 days later along with a yellowing of his skin and eyes and an episode of fainting and abundant coffee-ground emesis.

PE  VS: fever (39°C); hypotension (BP 90/60). PE: jaundice; petechiae on lower legs; swollen, bleeding gums; cardiomegaly; hepatomegaly.

Labs  CBC: leukopenia. UA: oliguria; albuminuria; hematuria.

Gross Pathology  Normal-sized liver with yellowish hue and petechiae: pale, swollen kidneys.

Micro Pathology  Characteristic midzonal lobular necrosis, fatty accumulation, and eosinophilic intracytoplasmic Councilman bodies on liver biopsy; hyperplasia of endothelial cells surrounding lymphoid follicles of spleen; severe renal tubular damage with epithelial fatty degeneration and necrosis.

Treatment  Symptomatic; prevention with mosquito control and live viral vaccination.

Discussion  Yellow fever is a viral hemorrhagic fever that is caused by a flavivirus transmitted by *Aedes* mosquitoes; it is preventable by a vaccine, which is required prior to travel to certain countries. It is associated with a mortality rate of 5% to 10%, but most cases are self-limiting and mild. It is similar to malaria but does not recur.
A neonate died shortly after birth.

Review of the medical record reveals history of refusal to feed, an extensive maculopapular skin rash on his legs and trunk, respiratory distress, diarrhea, and seizures shortly after birth.

Neonatal listeriosis may occur early or late in neonatal life. Infants may be acutely ill at birth and may die within hours as a result of disseminated listeriosis, which is also called granulomatosis infantiseptica. This condition is characterized by hepatosplenomegaly, thrombocytopenia, generalized skin papules, whitish pharyngeal patches, and pneumonia. Commonly, a stained smear of meconium will reveal gram-positive bacilli, suggesting the diagnosis.
A 25-year-old woman visits her family physician because of marked burning pain while urinating (dysuria), increased frequency of urination with small amounts of urine (pollakiuria), and passage of a few drops of blood-stained debris at the end of urination (hematuria).

HPI She got married 2 weeks ago and has just returned from her honeymoon.

PE VS: no fever; BP normal. PE: no edema; no costovertebral angle tenderness; moderate suprapubic tenderness with urgency.

Labs UA: urine collected in two glasses; second glass more turbid and blood-stained; urine sediment reveals RBCs and WBCs; no RBC or WBC casts; Gram stain of urine sediment reveals gram-negative bacilli; *Escherichia coli* in significant colony count (>100,000) on urine culture.

Treatment Oral antibiotics (Bactrim, fluoroquinolone); adequate hydration.

Discussion *E. coli* is the most common pathogen: *Proteus, Klebsiella, Staphylococcus saprophyticus*, and *Enterococcus* are other common bacteria causing cystitis. Hemorrhagic cystitis may result from adenoviral infection.
ID/CC A 28-year-old man comes to the ER with gradually worsening and now severe scrotal swelling and pain radiating to the inguinal area.

HPI The patient has no significant medical history. He reports pain on urination (due to concomitant urethritis) and notes that he is sexually active with multiple partners. He also notes that the pain is greater on standing and walking and is relieved by rest and elevation of the legs.

PE VS: normal. PE: scrotal edema and erythema; right epididymis enlarged and tender; induration present; elevation of scrotal contents relieves pain (PREHN’S SIGN).

Labs UA: pyuria. Culture negative; biopsy of epididymis inoculated into cell cultures grows *Chlamydia trachomatis*; immunofluorescence reveals subtype D.

Imaging US: hypoechoic, enlarged epididymis with hypervascularity.

Gross Pathology Nonspecific inflammation characterized by congestion and edema.

Micro Pathology Early stage of the infection is limited to the interstitial connective tissue with white cell infiltration.

Treatment Antibiotics like doxycycline, minocycline for chlamydia. Course of ofloxacin covers all possibilities of causative organisms.

Discussion Differentiate epididymitis from testicular torsion and tumor (scrotal ultrasonography or isotopic flow study may be needed for differentiating). Transmitted sexually in young adults and most often caused by *Chlamydia trachomatis* subtypes D through K and *Neisseria gonorrhoeae*. In those older than 40, *Escherichia coli* and *Pseudomonas* cause most infections. If associated with rectal intercourse, it may be due to *Enterobacteriaceae*. 

81 EPIDIDYMITIS
A 15-year-old male presents with **painful bilateral swelling of the parotid glands**, left-sided scrotal pain, and fever.

**HPI** Nothing in the patient's history suggests that he had childhood mumps. He has not received a measles-mumps-rubella (MMR) vaccination.

**PE** VS: fever. PE: bilateral parotid gland enlargement with obliteration of mandibular hollow; hyperemia and edema of Stensen's duct (parotid duct) orifice; retroauricular lymphadenopathy; left-sided scrotal and **testicular swelling with tenderness**.

**Labs** CBC: leukopenia with **lymphocytosis; hyperamylasemia**. Positive complement fixation antibodies; positive serologic enzyme immunoassay (EIA) for mumps antibody (repeat test after 1 week to demonstrate a fourfold rise).

**Imaging** US, scrotum: increased color flow and edema.

**Gross Pathology** Enlarged, edematous testicle.

**Micro Pathology** Parotid glands show perivascular mononuclear, lymphocytic, and plasma cell infiltrate with necrosis; ductal obstruction and edema; testicular interstitial edema; perivascular cerebral lymphocytic cuffing.

**Treatment** Scrotal support; analgesics, ice packs; corticosteroids.

**Discussion** Orchitis may be caused by bacterial infections such as *Escherichia coli* and other enterobacteria; viral infections such as **mumps**; STDs such as *Chlamydia* species or gonorrhea; or pathogens such as *Mycobacterium tuberculosis*. **Mumps** orchitis may give rise to sterility if bilateral.

**Atlas Link** [UOVA FG-MZ-082](#)
A 10-year-old child presents with complaints of acute-onset voiding of tea-colored urine and reduced urinary output.

The child was treated 1 week ago for streptococcal pyoderma that was confirmed by culture. He also complains of puffiness around the eyes and mild swelling of both feet.

VS: hypertension (BP 140/96); fever; tachycardia. PE: periorbital swelling; mild pitting pedal edema; no ascites or kidney mass palpable.

CBC: mild leukocytosis. Elevated BUN and creatinine; elevated ASO titer; serum cryoglobulins present. UA: RBC casts; proteinuria. C3 levels reduced in blood.

Smooth, reddish-brown cortical surface with numerous petechial hemmorhages.

Biopsy shows diffuse glomerulonephritis resulting from proliferation of endothelial, mesangial, and epithelial cells; granular, "starry-sky" pattern of IgG, IgM, and C3 on immunofluorescence; electron microscopy shows subepithelial "humplike" deposits (antigen-antibody complexes).

Penicillin if still infected with Streptococcus; diuretics, salt and water restriction and antihypertensives.

Poststreptococcal glomerulonephritis is a classic immune complex-mediated entity that is associated with acute nephritic syndrome, which develops following infection with nephritogenic group A β-hemolytic streptococci (e.g., types 1, 4, and 12, which are associated with pharyngitis, and types 49, 55, and 57, which are associated with impetigo).
A 25-year-old male presents with complaints of sudden-onset fever and chills, urgency and burning on micturition (dysuria), and perineal pain.

His symptoms developed a day after he underwent urethral dilatation for a stricture.

VS: fever. PE: suprapubic tenderness; rectal exam reveals asymmetrically swollen, firm, markedly tender, hot prostate; prostatic massage is avoided owing to risk of inducing bacteremia; epididymitis and extreme pain.

Examination and culture of urine and prostatic secretions reveal infection with *Escherichia coli*.

Edematous gland enlargement with suppuration of entire gland, possibly abscesses and focal areas of necrosis that have coalesced.

Initially minimal leukocytic infiltration of stroma. Later, necrosis of the gland may lead to gland fibrosis.

Antibiotic therapy as directed by urine and blood culture sensitivity tests.

*Escherichia coli* is the most common cause of acute prostatitis; many cases follow the use of instrumentation for the urethra and prostate (e.g., catheterization, cystoscopy, urethral dilatation, transurethral resection). Remaining infections are caused by *Klebsiella*, *Proteus*, *Pseudomonas*, and *Serratia*. Among the gram positives, enterococci and *Staphylococcus aureus* are frequent causative organisms.
A 65-year-old male complains of recurrent burning, urgency, and frequency of micturition together with vague lower abdominal, lumbar, and perineal pain.

He also complains of a mucoid urethral discharge. He was previously diagnosed via ultrasound with benign prostatic hypertrophy but does not report any severe symptoms of prostatism; his medical history reveals frequent UTIs due to Escherichia coli.

He also complains of a mucoid urethral discharge. He was previously diagnosed via ultrasound with benign prostatic hypertrophy but does not report any severe symptoms of prostatism; his medical history reveals frequent UTIs due to Escherichia coli.

VS: stable; no fever. PE: rectal exam reveals enlarged, nodular prostate; biopsy obtained to rule out carcinoma.

Examination and culture of expressed prostatic secretions reveal leukocytosis and E. coli.

IVP/voiding cystourethrogram (to rule out underlying anatomic cause): normal.

Enlarged prostate with nodularity and calculi.

Chronic inflammation and few PMNs around glands and ducts on biopsy; dilated ducts containing inspissated secretions (CORPORA AMMACEA).

Antibiotics (TMP-SMX, carbenicillin, quinolones). High fluid intake and abstinence from alcohol. Recurrences are common.

Bacterial prostatitis is usually caused by the same gram-negative bacilli that cause UTIs in females; 80% or more of such infections are caused by Escherichia coli. Chronic bacterial prostatitis is common in elderly males with prostatic hyperplasia and is a frequent cause of recurrent UTIs in males (most antibiotics poorly penetrate the prostate; hence the bacteria are not totally eradicated and continuously seed the urinary tract).
A 28-year-old black woman who is in her 27th week of pregnancy complains of right flank pain, high-grade fever, malaise, headache, and dysuria.

HPI

Thus far her pregnancy has been uneventful.

PE

VS: fever. PE: no peripheral edema; right costovertebral angle tenderness; acutely painful fist percussion on right lumbar area (positive Giordano's sign).

Labs

CBC: leukocytosis with neutrophilia. UA: proteinuria; hematuria; abundant WBCs and WBC casts; pyocytes on sediment; alkaline pH; urine culture > 100,000 colonies of Escherichia coli.

Imaging

US, renal: slightly enlarged kidney.

Gross Pathology

Kidney enlarged, edematous, and hyperemic with microabscesses in medulla.

Micro Pathology

Pyocytes in tubules; light blue neutrophils on supravital stain (GLITTER cells); PMN infiltration of interstitium.

Treatment

Antibiotics according to sensitivity: ampicillin; in nonpregnant patients, fluoroquinolone or ampicillin and an aminoglycoside constitute initial treatment.

Discussion

An acute bacterial kidney infection caused mainly by gram-negative bacteria such as E. coli, Klebsiella, Proteus, and Enterobacter; acute pyelonephritis usually results from upward dissemination of lower urinary tract bacteria.

Atlas Links

[PG-M2-086A, PG-M2-086B]
A 10-year-old female presents with a high fever, headache, vomiting, and impaired consciousness.

She suffered a generalized seizure about 45 minutes ago. She was previously diagnosed with cyanotic congenital heart disease (ventricular septal defect with Eisenmenger's syndrome).

VS: fever. PE: altered sensorium: papilledema; nuchal rigidity; clubbing; central cyanosis; cardiac auscultation suggestive of VSD with severe pulmonary arterial hypertension.

Blood culture reveals mixed infection with Bacteroides, microaerophilic streptococci, Staphylococcus aureus, and Klebsiella; staining and culture of pus aspirated from brain abscess confirm polymicrobial infection.

CT (with contrast): multiple ring-enhancing lesions with low attenuation centers (ABSCESS) surrounding cerebral edema and ventricular compression.

Cavity filled with thick, liquefied pus surrounded by fibrous capsule of variable thickness; pericapsular zone of gliosis and edema.

Central portion contains degenerated PMNs and cellular debris; capsule is composed of collagenous fibrous tissue with blood vessels and mixed inflammatory cells.

High-dose, extended parenteral broad-spectrum antibiotic coverage; CT-directed drainage of pus.

Brain abscesses arise secondary to hematogenous spread from another infection (bronchietasis, endocarditis), from contiguous spread from adjacent infection (chronic otitis media, mastoiditis, sinusitis), or following direct implantation from trauma. Patients with congenital heart disease with right-to-left shunt are particularly predisposed because the normal filtering action of the pulmonary vasculature is lost.
A 43-year-old male Mexican migrant worker visits his ophthalmologist because of pain and loss of vision in his right eye.

Recently he has also suffered from severe headaches and projectile vomiting.

Papilledema on left funduscopic exam; free-floating cyst in vitreous body of right eye; chorioretinitis and disk hemorrhage; multiple nontender subcutaneous nodules.

CBC: eosinophilia. LP: lymphocytic and eosinophilic pleocytosis in CSF with elevated protein and decreased glucose. Eggs of Taenia solium in stool sample.

XR, plain: small nodular calcifications. CT/MR, brain: characteristic ring-enhancing intracranial cysts or calcifications; can cause obstruction and hydrocephalus.

Fluid-filled cysts containing scolex surrounded by fibrous capsule in anterior chamber of eye; intraventricular and parenchymal invasion of brain, subcutaneous tissue, and striated muscle.

Inflammatory infiltration of cyst by PMNs; necrotic inflammation with calcification upon death of parasite.

Surgical removal of parasite from eye; albendazole, corticosteroids/praziquantel for brain disease.

Produced by Cysticerus colubriformis, the larval form of the pork tapeworm Taenia solium, neurocysticercosis is due to the ingestion of ova and spreads through fecal-oral transmission.
A 30-year-old male presents with a high fever and chills, headache, nausea, vomiting, and muscle aches.

Yesterday he had an episode involving abnormal movements of his right hand and face (FOCAL SEIZURE). He also has difficulty comprehending speech and has olfactory hallucinations. He has no history of psychiatric illness.

VS: fever; tachycardia; mild tachypnea; BP normal. PE: confused and disoriented; papilledema; mild nuchal rigidity; Kernig’s sign positive; paraphasic errors in speech; deep tendon reflexes normal and bilaterally symmetric.

L.P: cells 400/μL with mononuclear pleocytosis; mildly elevated protein; normal glucose; CSF PCR reveals herpes simplex virus type 1 (HSV-1); serum complement-fixing antibody titer > 1:1000. EEG: spiked and slow waves localized to temporal lobes.

CT: characteristic changes of encephalitis seen over temporal lobes.

Hemorrhagic, necrotizing encephalitis is most severe along inferior and medial regions of temporal lobes and orbitofrontal gyri.

Brain biopsy reveals Cowdry intranuclear viral inclusion bodies in both neurons and glial cells with perivascular inflammatory infiltrates.

Intravenous acyclovir.

Herpes simplex virus is the most common cause of acute sporadic encephalitis in the United States. In the newborn, HSV-2 is usually the cause; after the neonatal period, most cases result from HSV-1. Neonatal infection (usually HSV-2) occurs after exposure to maternal genital infection at the time of delivery. The precise pathogenesis of HSV-1 encephalitis in the older child or the adult is not clear, but viral spread into the temporal lobe by both olfactory and trigeminal routes has been postulated.

Atlas Link [HERPES SIMPLEX ENCEPHALITIS]
An 11-year-old girl is brought to the ER with high fever, chills, severe headache, vomiting, and obtundation.

Her parents report that she suffered a generalized seizure about an hour ago. A few days ago, the family had returned from a summer vacation in south India, where the child often played in irrigated rice farms. She did not receive any immunizations prior to her travel.

VS: fever. PE: patient is stuporous; neck stiffness and Kernig's sign positive (due to meningeal irritation); mild papilledema; tremors noted in hands.

CSF reveals pleocytosis with predominant lymphocytosis, mildly elevated proteins, and normal sugar (suggestive of aseptic meningitis); IgM enzyme immunoassay performed on acute and convalescent sera and CSF reveals significant titer of antibodies to Japanese encephalitis virus.

GT, head: areas of low density in the thalamus and basal ganglia.

Supportive; experimental intrathecal α-interferon therapy.

Japanese encephalitis virus is a flavivirus that causes disease in humans, horses and pigs. It is widely distributed in Asia from Japan and Eastern Siberia to Indonesia and westward to India; epidemics occur in summer months coincident with the abundance of the mosquito vector Culex tritaeniorhynchus. The vector breeds in irrigated rice fields and bites preferentially at sunset and sunrise; pigs are the amplifying hosts, whereas man is the incidental "dead-end" host. A vaccine is available for routine use for childhood immunization in Japan and in developed countries to protect travelers.
ID/CC A 30-year-old male laboratory researcher presents with a high fever, neck rigidity, retro-orbital pain, and severe myalgias of a few days’ duration.

HPI The patient also complains of a sore throat and photophobia. His work in the lab involves close contact with experimental animals such as hamsters, white mice, and nude mice. He was adequately vaccinated.

PE VS: fever. PE: neck stiffness and Kernig’s sign positive (due to meningeal irritation); pharyngeal inflammation but no exudate noted.

Labs CBC: mild leukopenia. LP: CSF suggestive of aseptic meningitis; LCM virus isolated from CSF. IgG and IgM antibodies detected in serum by immunofluorescent assay.

Treatment Supportive; ribavirin may play a role.

Discussion Lymphocytic choriomeningitis virus is an arenavirus. Sporadic cases occur after infection with feral mice, but the most common sources of human infection are pet/lab rodents. The virus is considered a major lab hazard, and care must be taken to avoid accidental infection. There is no commercially available vaccine.
A 50-year-old white male develops sudden fever with chills, pain in the back and extremities, and neck stiffness; he vomited six times and had a convulsion prior to admission.

The patient is a heavy smoker and is diabetic. Two weeks ago, he had a URI. He is also very sensitive to light (photophobia).

Markedly reduced mental status (obtunded); petechial rash over trunk and abdomen; nuchal and spinal rigidity; positive Kernig's and Brudzinski's signs; no focal neurologic deficits.

L.P: elevated pressure; cloudy CSF; elevated protein; markedly decreased glucose; high cell count with mostly WBCs. CSF Gram stain reveals gram-positive diplococci. Spinal fluid culture grows Streptococcus pneumoniae.

CT/MR, brain: meningeal thickening and enhancement.

Pia-arachnoid congestion results from inflammatory infiltrate; thin layer of pus forms and promotes adhesions while obstructing normal CSF flow (can cause hydrocephalus); brain covered with purulent exudate, most heavily on base.

Early empiric high-dose IV antibiotics; cefotaxime; vancomycin; high-dose steroids.

Bacterial meningitis is a pyogenic infection of the CNS that requires prompt treatment. Streptococcus pneumoniae is the most common cause of adult meningitis.

PG-M2-092A, PG-M2-092B, M-M2-092
NEU-027

MENINGITIS—BACTERIAL (ADULT)
A 4-year-old female presents with a 1-week history of fever, severe headache, irritability, and malaise; 2 days ago she developed neck stiffness, and her parents report projectile vomiting over the past 24 hours.

HPI The child is also very sensitive to light (photophobia). She is fully immunized and has no history of ear, nose, and throat infection, skin rashes, dog bites, or foreign travel.

PE VS: fever. PE: irritability; resistance to being touched or moved; minimal papilledema of fundus; no focal neurologic signs; no cranial nerve deficits; positive Kernig’s and Brudzinski’s signs.

Labs CBC: neutrophilic leukocytosis. LP: increased pressure; cloudy CSF; neutrophilic pleocytosis; decreased glucose; increased protein; gram-negative coccobacilli. Negative ZN and India ink staining; normal serum electrolytes; on chocolate agar, blood culture grew Haemophilus influenzae; negative Mantoux.

Imaging CT/MR, brain: meningeal thickening and enhancement.

Gross Pathology Abundant accumulation of purulent exudate between pia mater and arachnoid; meningeal thickening; cloudy to frankly purulent CSF.

Micro Pathology Intense neutrophilic infiltrate.

Treatment IV antibiotics (ampicillin, cefotaxime); consider steroids.

Discussion A pyogenic infection of the nervous system primarily affecting the meninges, bacterial meningitis is most commonly caused by pneumococcus (Streptococcus pneumoniae, associated with sickle cell anemia), meningococcus (Neisseria meningitidis, associated with a petechial skin rash), and H. influenzae (most commonly in children). It is less commonly caused by enterobacteria, Streptococcus species, Staphylococcus species (due to dental infections), and anaerobic organisms (due to trauma).

Atlas Link [URL] M-M2-093
A 33-year-old HIV-positive white male is brought into the emergency room by his mother because of a persistent headache.

The patient’s mother states that her son has been suffering for a long time from headaches and stiff neck as well as from fever and chills.

VS: fever (39°C). PE: severe nuchal rigidity; lack of responsiveness to any command; positive Kernig’s and Brudzinski’s signs; diminished patellar and Achilles reflexes; clear lung sounds.

LP: increased CSF pressure; variable pleocytosis (75 lymphocytes/mm³); elevated protein; decreased glucose. Heavily encapsulated, nondimorphic spherical fungal cells (Cryptococcus neoformans) revealed on India ink staining; polysaccharide capsular antigen detected on latex agglutination test; diagnosis confirmed by culture on Sabouraud’s medium.

CT/MR, brain: multiple ring-enhancing lesions.

Granuloma and abscess formation, mainly at base of brain; CNS primarily affected; lungs affected less commonly.

Abundant fungi in CSF and leptomeninges, with slight mononuclear inflammatory reaction; typical nodular granulomatous meningitis with exudate.

Amphotericin B and 5-flucytosine: fluconazole.

Once called torulosis, cryptococcosis is the most common cause of mycotic meningitis; it is acquired through the inhalation of dried pigeon droppings and is usually seen in immunocompromised patients.

ID/CC  A 6-year-old male being treated for primary pulmonary tuberculosis presents with diplopia, increasing drowsiness, irritability, and unexplained, recurrent vomiting.

HPI  The child has had a low-grade fever, loss of appetite, and a persistent headache over the past few weeks.

PE  VS: fever. PE: stuporous; signs of meningeal irritation noted (neck rigidity, Kernig's sign); CN III and IV palsy on right side; funduscopy reveals papilledema.

Labs  LP (guarded): CSF under increased pressure and turbid; on standing, a "cobweb" coagulum formed at center of tube; CSF studies reveal lymphocytic pleocytosis, greatly elevated protein, and low sugar; ZN staining of CSF coagulum reveals presence of acid-fast bacilli; radiometric culture yields Mycobacterium tuberculosis.

Imaging  CT: suggests basal exudates, inflammatory granulomas, and a communicating hydrocephalus; striking meningeal enhancement noted in post-contrast studies.

Gross Pathology  Meningeal surface covered with yellowish-gray exudates and tubercles that are most numerous at base of brain and along the course of the middle cerebral artery; subarachnoid space and arachnoid villi obliterated (leading to poor absorption of CSF and hence a communicating hydrocephalus).

Micro Pathology  Subarachnoid space contains gelatinous exudate of chronic inflammatory cells, obliterating cisterns, and encasing cranial nerves; well-formed granulomas occasionally seen, most often at base of brain; arteries running through subarachnoid space show "obliterative endarteritis."

Treatment  Antituberculous therapy with rifampin, isoniazid, ethambutol and pyrazinamide; steroids, ventriculoperitoneal shunt to relieve hydrocephalus.

Discussion  Tuberculous infection reaches the meninges through the hematogenous route, resulting in a clinically subacute form of meningitis; it is often complicated by cranial nerve palsies, a communicating hydrocephalus, decerebrate posturing, convulsions, coma, and death.
A 3-year-old male, the child of recent African immigrants, is brought to the local health center because of asymmetrical legs.

His parents give a history of incomplete immunization. They add that 5 months ago the boy had fever and diarrhea that subsided spontaneously; a few weeks later they noted that he could not use his right leg.

Right leg thin, short, wasted, weak, and flaccid; absent deep tendon reflexes in right leg; no sensory deficit; upper limbs normal; mental status and cranial nerves normal.

EMG: chronic partial denervation with abnormal spontaneous activity in resting muscle and reduction in number of motor units under voluntary control; normal sensory conduction studies.

Rehabilitation, supportive.

A symptomatic disease caused by poliovirus that is more common in infants and children, poliomyelitis can result in muscular atrophy and skeletal deformity. It attacks motor neurons in the anterior horns and may affect cranial nerves (bulbar polio); it is preventable by vaccine.
A 26-year-old nurse presented with headaches and recent-onset seizures; she also complained of increasing right-sided numbness and blurring of vision.

A clinical diagnosis of HSV encephalitis had previously been made, for which the patient was treated with two courses of acyclovir without any amelioration of symptoms; the disease continued to progress both radiologically and clinically. On serology she tested HIV positive.

Neurologic exam reveals cognitive mental impairment; visual field defects and sensory dysphasia seen; an ill-defined sensory loss on right side of body.

HIV positive by ELISA and Western blot.

MR (T2-weighted): patchy high-intensity lesions in the deep white matter of left cerebral hemisphere involving temporal, parietal, and occipital lobes.

Stereotactic biopsy sections show abnormal brain with rarefaction, numerous reactive astrocytes, foamy histiocytes, and inflammatory infiltrate around some vessels; JC virus in situ hybridization shows many positive nuclei; no herpesvirus inclusions seen; electron microscopy demonstrates cells with typical papovavirus structures in nucleus.

Disease was relentlessly progressive and resulted in death within 6 months.

Progressive multifocal leukoencephalopathy is a progressive demyelinating disease related to JC papovavirus infection; the largest number of cases occur in persons who are immunocompromised for any of a variety of reasons, including organ transplantation, hematologic and other malignant diseases, chronic immunosuppressive therapy, and AIDS.
A 30-year-old male is seen with complaints of a rash along with pain in his left ear and inability to move the muscles of his face with accompanying asymmetry.

He suffered an attack of chickenpox during childhood but has no history either of a similar rash over his face or of any visual symptoms (to rule out herpes zoster ophthalmicus).

Vesicular rash over left pinna (OTITIS EXTERNA); left-sided lower motor neuron-type facial nerve palsy (patient is unable to frown and unable to blink left eye; eyeballs roll up during attempt to close eye; patient is unable to whistle; taste sensation over anterior two-thirds of tongue lost on left side).

Although the diagnosis is predominantly clinical, a Tzanck test examining lesion scrapings (showing evidence of multinucleate acantholytic cells), direct culture, and immunohistochemical identification of infected cells allow identification of the virus.

Neuritis and vesicular skin lesions confined to distribution of geniculate ganglion of facial nerve.

Vesicular skin lesions with herpes viral inclusions, i.e., intranuclear, acidophil inclusions with a halo around them (COWDRY TYPE A INCLUSIONS); syncytial cells also seen.

Systemic steroids are mainstay of management.

Herpes zoster of the geniculate ganglion, or Ramsay Hunt syndrome, presents as a vesicular rash on the pinna followed by ipsilateral LMN facial nerve palsy.
ID/CC  A 30-year-old male presents with a **high fever, neck stiffness, and drowsiness**.

HPI  He also complains of nausea and vomiting. He **recently traveled along the Mississippi-Ohio River basin**.

PE  VS: fever. PE: neck stiffness and Kernig’s sign positive (due to meningeal irritation); right oculomotor nerve palsy noted; mild papilledema.

Labs  IgM enzyme immunoassay done on paired sera, and CSF confirms the diagnosis of **St. Louis virus infection**. LP: CSF exam reveals pleocytosis with predominant lymphocytosis suggestive of aseptic meningitis.

Micro Pathology  Inflammation and neuronal degeneration, principally in the thalamus, midbrain, and brainstem.

Treatment  Supportive treatment.

Discussion  St. Louis encephalitis virus is the **most common cause of epidemic encephalitis** in the United States; cases occur annually as isolated events or in summer and autumn encephalitis epidemics. **Most infections are asymptomatic**. The disease occurs throughout the United States, but outbreaks have also occurred in the Caribbean as well as in Central and South America.
100 SUBACUTE SCLEROSING PANENCEPHALITIS
A 25-year-old recently married woman is concerned about a scanty, offensively malodorous vaginal discharge.

She states that the discharge is thin, grayish-white, and foul-smelling. She does not complain of vulvar pruritus or soreness.

Pelvic exam confirms presence of a homogenous, grayish-white, watery discharge adherent to the vaginal walls that yields a "fishy" odor when mixed with KOH; no injection and excoriation of the vulva, vagina, or cervix.

Vaginal pH > 5; saline smear reveals presence of characteristic "clue cells" (squamous epithelial cells with smudged borders due to adherent bacteria).

Single dose of metronidazole (2 g) effective in treating the infection. Oral clindamycin is an alternative drug.

Although bacterial vaginitis was originally thought to be caused by Gardnerella vaginalis, this organism is now recognized to be part of the normal vaginal flora. Bacterial vaginosis is now known to result from a synergistic interaction of bacteria in which the normal Lactobacillus species in the vagina is ultimately replaced by high concentrations of anaerobic bacteria, including Bacteroides, Peptostreptococcus, Peptococcus, and Mobiluncus species along with a markedly greater number of G. vaginalis organisms than is encountered in normal vaginal secretions. Bacterial vaginosis is known to increase the risk of pelvic inflammatory disease, chorioamnionitis, and premature birth.
ID/CC A 25-year-old puerpera who was lactating her week-old infant presents with pain and swelling in her left breast.

HPI The symptoms commenced acutely, and she does not recall any previous breast lumps or swellings.

PE Skin overlying left breast is red, edematous, tender, and hot; area of tense induration felt underlying inflamed skin.

Labs Culture of pus drained from breast abscess and nasopharyngeal swab taken from the infant grew Staphylococcus aureus.

Imaging USG: nearly anechoic area with posterior enhancement.

Treatment Penicillinase-resistant antibiotic; incision (in a radial direction over the affected segment) and dependent drainage of intramammary abscess; breast feeding was temporarily discontinued.

Discussion Bacterial mastitis most commonly occurs in lactating women due to infection of a hematoma or secondary infection of plasma cell mastitis; the infecting organism is mostly penicillin-resistant Staphylococcus aureus.
A 38-year-old white female visits her gynecologist for a routine Pap smear.

She admits to early sexual activity, many sexual partners, and unprotected sex.

Pallor; cervical tenderness; a few small, raised, flat lesions on cervix; genital warts also seen on vulva (condylomata acuminata).

Presence of HPV in cervical cells revealed on DNA hybridization and immunofluorescent antibody assays for viral antigen.

Rounded basophilic cells on Pap smear with large nuclei occupying most of surface; nuclei show mitoses and coarse clumping of chromatin with perinuclear halo (severe koilocytic dysplasia).

Cryotherapy, conization, or local excision with follow-up.

Infection with HPV types 16, 18, and 31 is strongly associated with cervical cancer preceded by dysplasia. Spread of the infection to partners may be prevented by barrier contraception.
A 28-year-old sexually active woman presents with crampy lower abdominal pain, yellowish vaginal discharge, and general malaise.

She also complains of continuous low-grade fever and reveals that the pain is exacerbated during and immediately after menstruation (CONGESTIVE DYSMENORRHEA). She uses a copper intrauterine device for contraception.

VS: low-grade fever. PE: lower abdominal tenderness; bimanual pelvic exam demonstrates purulent vaginal discharge, bilateral adnexal tenderness, and pain on movement of cervix (MUCOPURULENT CERVICITIS).

CBC: leukocytosis with left shift. Increased ESR; endocervical swab sent for microscopic exam; staining and culture revealed combined infection with Neisseria gonorrhoeae (cultured on Thayer-Martin medium) and Chlamydia trachomatis (identified on cell culture, immunofluorescence, and antigen capture assay); laparoscopy ("gold standard" for diagnosis) confirmed diagnosis.

USG: free pelvic fluid, dilated tubular structure in adnexa.

Erythema and swelling of fallopian tubes on laparoscopy; sero-purulent exudate noted on surface of tubes from fimbriated end.

Endocervical swab reveals increased neutrophils and gram-negative diplococci seen both intra- and extracellularly; cervical biopsy reveals inclusions containing Chlamydia within columnar cells.

Antibiotic therapy with cefoxitin (for N. gonorrhoeae) and doxycycline (for chlamydial infection); male partners must be treated for STDs.

Pelvic inflammatory disease usually occurs as a primary infection that ascends from the lower genital tract due to STDs caused by Neisseria gonorrhoeae and Chlamydia trachomatis. Sequelae of PID include peritonitis; intestinal obstruction due to adhesions; dissemination leading to arthritis, meningitis, and endocarditis; chronic pelvic pain; infertility; ectopic pregnancy; and recurrent PID.
A 20-year-old Asian woman presents with complaints of infertility and heavy bleeding during menses (menorrhagia).

She was treated for pulmonary tuberculosis a few years ago. She has been unable to conceive despite unprotected intercourse for the past 2 years. Her husband's semen analysis is normal.

On pelvic exam, small, fixed adnexal masses are palpable that are matted and fixed to uterus ("frozen pelvis").

Culture of endometrial curettings yields Mycobacterium tuberculosis; histologic examination of curettings reveals presence of characteristic tubercles; Mantoux skin test strongly positive.

CXR: left apical fibrosis (evidence of old healed pulmonary tuberculosis). (Hysterosalpingography (HSG) is contraindicated in a proven case of tuberculosis. When done in asymptomatic cases, HSG yields certain typical findings, including a rigid, nonperistaltic, pipelike tube; beading and variation in filling density; calcification of the tube; cornual block; jagged fluffiness of the tubal outline; and vascular or lymphatic extravasation of the dye.)

Tubes are enlarged, thickened, and tortuous; examination of uterine reveals evidence of synechiae and adhesions (leading to Asherman's syndrome).

Microscopic exam of tubes, ovaries, and endometrium reveals evidence of granulomas with giant cells and caseation.

Four-drug therapy with isoniazid, pyrazinamide, ethambutol, and rifampicin; pyridoxine to prevent isoniazid-induced deficiency.

Genital tuberculosis is almost always secondary to a focus elsewhere in the body, with the bloodstream by far the most common method of spread. The fallopian tubes are the most frequently involved part of the genital tract, followed by the uterus. Ninety percent of patients are cured with chemotherapy, although only 10% regain fertility.
A 30-year-old woman presents to the ER with an abrupt-onset high fever, vomiting, profuse diarrhea, severe muscle aches, and disorientation.

One day ago she developed an extensive skin rash all over her body. Her husband says she used a vaginal sponge for contraception.

VS: fever; tachycardia; hypotension. PE: extremely toxic-looking; drowsy but responding to verbal commands; extensive scarlatiniform rash seen involving entire body; pharyngeal, conjunctival, and vaginal mucosa congested (frank hyperemia); no neck rigidity or Kernig’s sign demonstrable; funduscopic exam normal; no localizing neurologic deficits.

CBC: leukocytosis; thrombocytopenia. UA: mild pyuria (in absence of UTI). BUN and creatinine elevated; blood cultures sterile; culture of cervical secretions grows Staphylococcus aureus. LP: CSF normal. Serology for Rocky Mountain spotted fever, leptospirosis, and measles negative.

Vigorous IV fluids and parenteral penicillinase-resistant penicillin or first-generation cephalosporins; patient in this case recovered, and typical skin desquamation was seen over palms and soles during convalescence.

Toxic shock syndrome results from infection with Staphylococcus aureus. Its effects are mediated through the exotoxin TSST-1, which functions as a superantigen, stimulating the production of interleukin-1 and tumor necrosis factor. Staphylococcal TSS has been associated with the use of vaginal contraceptive sponges and with many types of localized staphylococcal soft tissue infections. Most cases of TSS occur in menstruating women.
ID/CC A 28-year-old primigravida at 36 weeks' gestation presents with a high fever.

HPI She was being monitored following a premature rupture of the membranes.

PE VS: fever; fetal tachycardia. PE: uterine tenderness.

Labs Elevated maternal total lymphocyte count; vaginal swab culture revealed colonization with group B streptococcus.

Treatment Presence of group B streptococcus in vagina after premature rupture of membranes was an indication for immediate delivery and treatment of the infant; mother was also treated with intravenous antibiotics.

Discussion A significant proportion of the population is colonized in the vagina and rectum with group B streptococcus, which is correlated with preterm labor, premature rupture of membranes (PROM), chorioamnionitis, and neonatal sepsis. Neonates with group B streptococcus sepsis have a 25% mortality rate. Among preterm neonates, this figure doubles to over 50%; therefore antibiotic prophylaxis is recommended in the setting of preterm delivery and PROM even without the diagnosis of frank chorioamnionitis. When chorioamnionitis is suspected, intravenous antibiotics are started and delivery is hastened.
ID/CC An 8-month-old male infant is brought to a pediatrician because of severe, intractable chronic diarrhea and failure to thrive.

HPI The mother died of AIDS shortly after the baby was delivered. The baby was asymptomatic at birth.

PE VS: fever; tachycardia. PE: emaciated, grossly malnourished; oral thrush; generalized lymphadenopathy; hepatosplenomegaly.

Labs Decreased CD4+ cell count; increased serum immunoglobulin level with impaired production of specific antibodies; ELISA and Western blot for HIV-1 positive (could be due to placental transfer of antibodies to HIV, but strongly supports diagnosis in presence of symptoms); PCR for HIV RNA positive (confirming HIV infection).

Treatment Nutritional support, Pneumocystis carinii prophylaxis, azidothymidine (zidovudine, or AZT) therapy (suppresses replication by inhibiting viral reverse transcriptase), and anti-infective agents for specific infections; IV serum immunoglobulin to reduce frequency of bacterial infections; oral polio vaccine and BCG contraindicated.

Discussion Vertical transmission of HIV-1 may occur in utero through transplacental passage of the virus, during delivery, or postnatally through breast feeding; however, it is believed that most infections are acquired at birth through contact with contaminated blood or secretions. Women who carry the virus should thus be discouraged from becoming pregnant or from breast feeding. The rate of transmission of HIV-1 from mother to infant has varied from 13% to 45%, with an average of 25%; however, when AZT is administered to HIV-1-infected pregnant women and to infants during the first 6 weeks of life, the risk of maternal-infant transmission is significantly reduced.
A 4-year-old white male presents with fever, chills, malaise, pain, and immobility of the right knee of 1 week's duration.

Two weeks ago the child fell while playing, but no abnormality was found by the school nurse.

Overlying skin warm and red; swelling of distal third of thigh and knee; tenderness on palpation.

CBC: leukocytosis. Elevated ESR. Gram stain and culture confirm diagnosis and isolate pathogen.

XR, plain: early findings include soft tissue edema and thin line running parallel to diaphysis (periosteal thickening); later findings include bone erosion, subperiosteal abscess with periostitis, and sequestrum formation (due to detached necrotic cortical bone); involucrum formation (laminated periostial reaction). MR: marrow edema; abscess. Indium-labeled WBC, scan: hot spot.

New osteoblastic periosteal bone formation (involucrum); trapping of detached necrotic bone by involucrum (sequestrum); isolated localized abscess (Brodie’s abscess); sinus tract formation, draining pus to skin.

Purulent exudate formation, usually metaphyseal, with ischemic necrosis of bone due to increased pressure of pus in rigid bone walls; vascular thrombosis.

IV antibiotics according to sensitivity; surgical debridement.

Osteomyelitis is an acute pyogenic bone infection which, if left untreated, produces functional incapacity and deformities. The most common pathogen is Staphylococcus aureus; less frequently Streptococci and enterobacteria are involved. In sickle cell anemia Escherichia coli and Salmonella species are seen; diabetics are at risk for Pseudomona infection. Immunocompromised patients may show Sporothrix schenckii osteomyelitis; human bites, anerobes; puncture wounds, Pseudomonas aeruginosa; and cutaneous wounds, Pasteurella multocida.
A 21-year-old female college student complains of low-grade fever along with pain and swelling in the left knee of 5 days' duration.

She had been to her family physician 2 weeks ago because of dysuria and a purulent vaginal discharge (due to gonococcal infection) and was given an “antibiotic shot”. She was asymptomatic until 4 days ago. She then developed fever, chills, and pain in both wrists and in her left ankle, which disappeared when the pain appeared in her left knee (migratory polyarthritis).

Swollen, tender, warm left knee with limited range of motion; white vaginal discharge.

Intracellular, bean-shaped gram-negative diplococci (gonococci) and markedly elevated WBC count on urethral smear and synovial fluid aspirate culture of synovial aspirate grows gonococci.

XR, knee: soft tissue swelling.

IV ceftriaxone.

Almost always accompanied by synovitis and effusion, gonococcal arthritis can rapidly destroy articular cartilage and is often associated with skin rash and C5, C6, C7, and C8 complement deficiencies. Single joints are usually affected, most often the wrists, fingers, knees, and ankles.