A Color Handbook

Pediatric Rheumatology

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Pediatric rheumatology is an evolving subspecialty of pediatrics. In 1992, pediatric rheumatology subspecialty certification was offered by the American Board of Pediatrics. With advances in basic science and clinical investigation and collaboration, newer illnesses such as the autoinflammatory syndromes have been added to the pediatric rheumatology repertoire. Advances in the treatments of adult rheumatic diseases are being translated to children with comparable conditions. Pediatric rheumatology is an ever-expanding field.

Children with rheumatic diseases frequently present with primarily musculoskeletal issues but may also present with multiple organ involvement. This diversity requires that providers caring for children with rheumatic disease have a well-developed set of clinical skills. These include general pediatrics as well as familiarity with concepts about immunology, musculoskeletal medicine, and the management of chronic diseases.

This book is formatted primarily based on the most frequently recognized symptom complexes in children with rheumatic diseases. The initial chapters focus on primarily musculoskeletal issues and subsequent chapters focus on chronic autoimmune diseases that may have multiorgan involvement. We hope that you find this handbook useful to your clinical practice and a tool with which better to assess children with potential rheumatic disease.
ACKNOWLEDGMENTS

We would like to thank the chapter authors for all their time and hard work that will allow improvement of the identification of pediatric rheumatic conditions. Further, we thank all the patients who have allowed us to use their images and understand the need for early identification of these conditions. We give special thanks to Laurie Metag for her assistance in editing the text.
Introduction

- Assessment of the child with rheumatic disease
- Approach to children with rheumatic disease

ASSESSMENT OF THE CHILD WITH RHEUMATIC DISEASE

Rheumatologic diseases in general, and particularly in children, are relatively uncommon. The most common rheumatologic diagnosis for children is probably juvenile arthritis, which has a prevalence of close to 1–2:1000 children. This is similar to the prevalence of childhood diabetes and epilepsy but significantly less common than that of childhood asthma.

Frequently the differential diagnosis in these cases is quite broad. Not surprisingly, with the rarity of rheumatic disease in children, the diagnosis is often arrived at after an extensive workup for other causes. Making the diagnosis of a rheumatic disease sometimes is challenging to the clinician. Other times it is quite obvious when there is a very large swollen joint, antalgic gait, and so on.

The timeliness of arriving at a diagnosis for rheumatic disease in a child is critical. Recognition of patterns of clinical signs and symptoms is the key to making an accurate assessment as well as having a foundation to guide therapy. In the following chapters you will find descriptions of the various childhood rheumatic conditions.

The impact of the sequelae of these conditions can be profound. Certainly life- and organ-threatening consequences are occasionally seen in patients with vasculitis or connective tissue diseases. Pain, suffering, and loss of function are frequently seen in children with inflammatory arthritic conditions such as juvenile arthritis and the spondyloarthropathies. Also the consequences of treatment must be carefully considered when embarking on therapy.
Approach to Children with Rheumatic Disease

Inflammatory versus non-inflammatory

In sorting out symptoms, it is important to take a careful medical history although such diseases may be intermittent, with periods of inactivity. Because conditions that have an inflammatory etiology have the potential for the highest risk for damage and disability, clinical priority is given to these conditions. Inflammatory features of conditions affecting the musculoskeletal system include joint swelling, joint redness, and joint warmth in addition to joint pain and tenderness. Typically the pain is worse in the morning and gets better as the day goes on. It is helped by activity and frequently associated with stiffness.

Non-articular signs and symptoms include fevers, weight loss, chills, or night sweats. Signs and symptoms of organ dysfunction such as weakness, dyspnea, edema, and rash are clinical clues to an inflammatory extra-articular process.

Conversely, the absence of these findings as well as weight gain, and difficulty with sleep, point to a potentially non-inflammatory etiology. Chronic pain syndromes and fibromyalgia are examples of conditions that may appear to be rheumatic in nature but are not.

Based on whether the primary symptom(s) (joint pain, rash, hematuria, and so on) is (are) chronic and the history and physical examination demonstrate an inflammatory/non-inflammatory pattern, an algorithmic approach to the diagnosis of rheumatic diseases in children can be utilized.

Acute, inflammatory, articular

Almost all of the pediatric rheumatology conditions are chronic. However, clinicians involved in the evaluation of children need to be competent in evaluating acute musculoskeletal problems as well. One of these is acute monoarthritis. While the differential diagnosis includes traumatic arthropathies, the strongest consideration needs to be given to infectious causes. Septic arthritis or adjacent osteomyelitis needs to be strongly considered in any child with an acute monoarthritis with or without fever. This is a medical emergency and needs to be evaluated appropriately in a timely fashion. Infection-related arthropathies are not reviewed in this atlas.

Acute organ dysfunction such as renal insufficiency, and pulmonary dysfunction is rarely attributable to rheumatic conditions in an undiagnosed child. But, a number of the pediatric rheumatic conditions can have organ dysfunction as part of the clinical course of their illness.

Chronic, inflammatory, articular

If a child presents with chronic symptoms and the symptoms are primarily referable to the musculoskeletal system such as pain, swelling, morning stiffness, limp then the differential diagnosis includes two categories of conditions. One of these is juvenile idiopathic arthritis (JIA) and the other is spondyloarthropathies. JIA is discussed in Chapter 2 and it is contrasted with spondyloarthropathies, which are well described in Chapter 3. Subtypes of JIA have characteristic patterns of joint involvement and extra-articular manifestations including autoantibodies and inflammatory eye disease. Spondyloarthropathies have a characteristic pattern of joint involvement which includes enthesitis (also seen in some forms of JIA) and involvement in the spine as well as association with HLA-B27. An example of joint swelling associated with juvenile rheumatoid arthritis (JRA) is shown (1). Evidence of early radiographic change in the sacroiliac joints in a patient with spondyloarthritis is also shown (2).

The most important part of the evaluation of patients with inflammatory joint disease is a careful medical history. Features of the onset, course, and duration of the symptoms is very important. Exclusion of other causes of joint pain and swelling are important as well such as those related to infectious causes. On physical examination, careful inspection and palpation of the joints are important in addition to a static and dynamic exam. Watching the child move the joint as well as the examiner moving the joint and moving the joint against resistance are important to discern the specific nature of the symptoms referable to a joint. In addition to these techniques for peripheral joints, techniques to examine the spine are outlined in the Chapter 3.
Approach to children with rheumatic disease

These patients may have evidence of inflammatory response in their blood with anemia, thrombocytosis, or leukocytosis. Acute-phase reactants such as CRP and sedimentation rate may be elevated. Some may also have autoantibodies such as RF.

In some conditions with chronic inflammatory joint symptoms, uveitis may also be seen. The uveitis may asymptomatic (posterior) or symptomatic (anterior) as shown (3).

To summarize, patients who have chronic inflammatory symptoms referable primarily to the musculoskeletal system and confirmatory findings on X-ray (4) are likely to have JRA or spondyloarthropathy. Laboratory studies can help verify the diagnosis and give prognostic information.

1 Wrist synovitis with cyst in polyarticular arthritis.

2 Scalloping of the sacroiliac joints in a 6 year old with spondyloarthropathy, severe back pain, and antalgic gait.

3 Red painful eye from anterior uveitis.

4 Hip arthritis with joint space loss in polyarticular arthritis.
**Chronic, inflammatory, organ involvement**

The next group of children with rheumatic diseases and inflammatory symptoms may have evidence of organ involvement apart from the musculoskeletal system. Examples of these include skin involvement with rashes from vasculitis (5) or vasospasm (6), or evidence of other organ involvement such as pleurisy (7), evidence of glomerulonephritis (8).

Since rheumatic diseases can affect any organ system, a very detailed history and a comprehensive physical exam are required to assess these patients. Unlike other subspecialties, patients may present with symptoms of any organ system so there are no characteristic historical or physical findings that would exclude a rheumatic condition. As with children with inflammatory joint symptoms, assessment of the duration and onset of the symptoms, things that make the symptoms worse or better, and interventions that have been done are important to note. A complete review of systems is also required.

On physical exam, characteristic findings such as a malar rash, painless mucositis, diminished breath sounds, edema, evidence of neurologic compromise, vasculitic rashes, and musculoskeletal weakness may be present and noted.

In these children, laboratory assessment is very important. They are necessary to assess the degree of organ damage. For example, a chest radiograph and perhaps pulmonary function studies would be required to assess chest pain or shortness of breath. Urinalysis, serum creatinine, and potentially renal biopsy for evidence to assess the cause of renal insufficiency might be needed as well. The degree of inflammation on blood tests is assessed as it is in inflammatory arthritis. The presence of autoantibodies is important as well.

To summarize, pediatric rheumatic diseases are uncommon and it often takes an astute clinician to make the diagnosis. Patterns of presentation include chronic illness that has inflammatory features. If the predominant clinical pattern is that of musculoskeletal involvement, the likely diagnostic categories include JIA and spondyloarthropathy. If organ involvement predominates, with or without musculoskeletal symptoms, then consideration of a vasculitis or connective tissue disease would be likely. My algorithm for connective tissue diseases is as shown (9).
Pleural effusion in systemic lupus erythematosus and acute pericarditis.

Glomerulonephritis in systemic lupus erythematosus.

A rheumatologic approach to categorizing chronic inflammatory symptoms.

SpA, spondyloarthritis; JIA, juvenile idiopathic arthritis; CTD, connective tissue disease; IIM, idiopathic inflammatory myopathy.
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Juvenile idiopathic arthritis
Juvenile idiopathic arthritis (JIA) is one of the most common pediatric rheumatic diseases. It is estimated that 300,000 children in the United States have arthritis, and 100,000 likely have some form of JIA (Sacks et al. 2007). Worldwide, the reported incidence has varied from 2 per 100,000 to 20 per 100,000 (Cassidy et al. 2005). True incidence is difficult to measure given the inherent difficulties in case ascertainment. Inflammatory arthritis has been described in all races and geographic areas. JIA comprises of different disease subtypes with heterogeneous phenotypic characteristics. Classification has been achieved by recognizing common clinical characteristics with the exclusion of potential mimics, and as such has been a source of debate through the years. For the purposes of this handbook, we will use the International League of Associations for Rheumatology (ILAR) criteria of classification. In general, juvenile arthritis is defined by intra-articular swelling, or the presence of two or more of the following signs: limitation in range of motion (ROM), tenderness or pain on motion, and increased heat that persists for longer than 6 weeks (Cassidy et al. 2005).

An international consensus conference was convened in 1997 to agree on nomenclature. The American College of Rheumatology (ACR) classification criteria are currently the only ones with corresponding ICD-9 codes in the United States. The three most widely used criteria, the ILAR classification, the ACR, and the European League Against Rheumatism (EULAR), are compared (Table 1) (Brewer et al. 1972, European League Against Rheumatism 1977, Petty et al. 1997). The ILAR classification includes the HLA-B27-associated spondyloarthropathies and psoriatic arthritis, subtypes categorized separately by the ACR criteria and covered in a different chapter in this handbook.

**Systemic arthritis**

**DEFINITION**

Systemic arthritis is a multisystemic inflammatory disease of which arthritis is only one of the multiple manifestations. It is characterized by the presence of arthritis and documented quotidian fever for at least 2 weeks plus at least one of the following: typical rash, generalized lymphadenopathy, hepatosplenomegaly, or serositis. Patients who fit the criteria for psoriatic arthritis, enthesitis-related arthritis, or other forms of arthritis are excluded from this category.

**EPIDEMIOLOGY AND ETIOLOGY**

Systemic juvenile arthritis was first described in 12 children from England by George Frederic Still in 1897. In North America and Europe systemic arthritis is one of the less-common forms of chronic arthritis, accounting for 10–20% of children seen in rheumatology clinics (Schneider & Laxer 1998, Woo 2006, Woo & Wedderburn 1998). There is no clear peak of onset as it can manifest in early childhood, adolescence, or during the adult years (then called adult-onset Still disease). Children of both genders are equally affected. It is uncertain as to whether there is seasonal variation in incidence. Recent research has indicated that these patients have high circulating levels of IL-1 and IL-6. IL-1 and IL-6 antagonists have been used clinically with improvement in some children with systemic arthritis (Pascual et al. 2008, Woo et al. 2005).

**CLINICAL HISTORY**

These children are usually ill with signs and symptoms of systemic inflammation (weight loss, fatigue, anemia, thrombocytosis, and elevated inflammatory markers) which tends to predominate early in the course of systemic arthritis. Temperature can rise to 39°C on a daily or twice-daily basis (quotidian pattern) and will subside to subnormal levels (10). Fever is characteristically elevated in the late afternoon or evening and tends to occur with the rash. Children will appear more systemically ill during fevers. Fever must be present for at least 2 weeks to fulfill diagnostic criteria. The rash that is associated with systemic arthritis is classically discrete, with erythematous macules...
<table>
<thead>
<tr>
<th>Classification</th>
<th>ACR (1977)</th>
<th>EULAR (1977)</th>
<th>ILAR (1997)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>JRA: juvenile rheumatoid arthritis</td>
<td>JCA: juvenile chronic arthritis</td>
<td>JIA: juvenile idiopathic arthritis</td>
</tr>
<tr>
<td>Minimum duration</td>
<td>6 weeks</td>
<td>3 months</td>
<td>6 weeks</td>
</tr>
<tr>
<td>≤4 joints at presentation</td>
<td>Pauciarticular</td>
<td>Pauciarticular</td>
<td>Oligoarthritis</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>A. Persistent</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>B. Extended</td>
</tr>
<tr>
<td>&gt;4 joints at presentation</td>
<td>Polyarticular</td>
<td>Polyarticular RF negative</td>
<td>Polyarticular RF negative</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Polyarticular RF positive</td>
</tr>
<tr>
<td>Fever, rash, arthritis</td>
<td>Systemic</td>
<td>Systemic</td>
<td>Systemic</td>
</tr>
<tr>
<td>Other categories included</td>
<td>Exclusion of other forms</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Juvenile ankylosing spondylitis</td>
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<td></td>
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<td></td>
<td>Juvenile psoriatic arthritis</td>
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<td></td>
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<td></td>
<td>Psoriatic arthritis</td>
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<td></td>
<td></td>
<td></td>
<td>Enthesitis-related arthritis</td>
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<td></td>
<td></td>
<td></td>
<td>Undifferentiated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>A. Fits no other category</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>B. Fits more than one category</td>
</tr>
</tbody>
</table>

ACR, American College of Rheumatology; EULAR, European League Against Rheumatism; ILAR, International League of Associations for Rheumatology (see text for references).

Fever pattern in a child diagnosed with systemic JIA.
2–5 mm in size (11). It is typically described as salmon-pink in color, migratory, and evanescent. It is most often found on the trunk and proximal extremities, but may develop on acral surfaces. Köbner phenomenon is often present (12), and it may be intensely pruritic in up to 10% of patients. Arthritis is often present initially, but may take weeks or months to manifest. Systemic JIA cannot be diagnosed without the presence of arthritis.

PHYSICAL EXAMINATION
Physical examination findings may include arthritis, tenosynovitis, fever, rash, lymphadenopathy and hepatosplenomegaly. The child may exhibit signs of cardiac or pleuropulmonary disease. Arthritis varies in manifestation, but is typically symmetric, starting in a few joints and progressing to a polyarticular course. It may present in any number of joints, and typically includes knees, wrists, and ankles. Cervical spine, hip, and TMJ disease may be present. In a minority of children, the arthritis is extremely resistant to treatment and results in an aggressively destructive course (13, 14). Tenosynovitis, present in 10% of children, may be seen in the extensor tendon sheaths on the dorsum of the hand (15), foot, and around the peroneus longus and brevis tendons of the ankle. Splenomegaly and lymphadenopathy are commonly seen, with hepatomegaly seen less commonly.
13 Radiographs of destructive hand arthritis in a child with unremitting systemic arthritis.

14 Cervical spine in a child with severe systemic JIA with fusion of multiple posterior facets allowing motion only at C5–6. Radiograph on left is at full extension, and on right at full flexion.

15 Tenosynovitis in the extensor tendons of the fingers, most marked in the right second, third, and fifth, and left second, third, and fourth digits.
The lymphadenopathy may be marked and symmetric. Serositis may be prominent, including pericardial and pleural effusions (16). Cardiac manifestations may include myocarditis, pericarditis, and rarely endocarditis and coronary arteryitis.

LABORATORY
There is no specific laboratory test for the diagnosis of systemic JIA. Laboratory markers of inflammation are typically markedly elevated (Table 2). White blood cell count may be in excess of 30,000 cells/mm\(^3\) with a predominance of neutrophils. Platelets are usually elevated as well, although rarely thrombocytopenia may be present in the early stages, raising questions about concurrent or alternative conditions. A normocytic hypochromic anemia, typical for chronic disease, is seen. Sedimentation rates are characteristically high except when associated with macrophage activation syndrome, where the ESR falls to normal and is typically accompanied by cytopenias. CRP and ferritin levels are also elevated. Elevated IgG may be found, but the presence of autoantibodies such as RF and ANA is atypical. Synovial fluid analysis usually reflects inflammation with white cells in the range 10,000–40,000/mm\(^3\). Biopsy of the rash will reveal minimal perivascular infiltration of mononuclear cells around capillaries and venules in the subdermal tissues. If there is any concern for macrophage activation syndrome or malignancy, bone marrow biopsy should be performed to evaluate for lymphoblasts or hemophagocytosis.

IMAGING
Serial radiographs may show joint space narrowing, erosions, growth arrest lines, and joint destruction. MRI and bone scan (17) may show ongoing inflammation in joints. Echocardiogram should be performed if there is any concern for pericarditis, coronary artery abnormalities, or myocarditis. If malignancy is suspected, a CT scan of the chest, abdomen, and pelvis should be considered.

DIFFERENTIAL DIAGNOSIS
At presentation systemic JIA can resemble infection, malignancy, inflammatory bowel disease (IBD), vasculitis, other connective tissue diseases, or one of the periodic fever syndromes. If corticosteroids are to be used for treatment, consideration of a bone marrow examination should be entertained before initiation of corticosteroids. Infections such as sepsis, acute rheumatic fever, Epstein–Barr virus, Bartonella sp., bacterial endocarditis, brucellosis, and Lyme disease should be considered. Malignancies such as leukemia and lymphoma may present like systemic arthritis with rash, fever, arthralgias or arthritis, and anemia. Unlike JIA, the fevers of infection and malignancy do not follow the quotidian pattern. Macrophage activation syndrome may be on a spectrum with hemophagocytic lymphohistiocytosis (HLH).

PROGNOSIS
Acute manifestations such as fever and rash are variable in duration, lasting from months to years, and may recur with disease flares. Approximately 50% of children with systemic arthritis recover completely, while the other half may show progressive arthritis with eventual chronic disability. In a series of patients from Cincinnati, 48% still had active arthritis 10 years later (Wallace & Levinson 1991). In a multicenter trial of 47 patients with systemic-onset JRA, there was a 37% probability of remission in 10 years (Oen et al. 2003). Predictors for prolonged disease and erosive arthritis include the presence of rash and fever 6 months into the disease course associated with a platelet count over 600 × 10^9 per liter. (Spiegel et al. 2000). Early studies showed a death rate of up to 14% (Stoeber 1981), but this is now thought to be much lower. Deaths in Europe were commonly due to amyloidosis; in the United States, deaths were more commonly associated with infections after corticosteroids. Macrophage activation remains a cause of mortality in these children. Unlike other forms of JIA, uveitis is generally not associated with systemic arthritis.

MANAGEMENT
Many of these children are hospitalized at diagnosis for systemic symptoms of illness and evaluation for possible sepsis or malignancy. These children rarely respond to NSAIDs alone, but these are used for fever and pain relief. Children with systemic JIA may have increased sensitivity to the hepatotoxicity of NSAIDs, especially aspirin. If the diagnosis is certain with no concern for possible malignancy, glucocorticoids may help with systemic manifestations and arthritis. Pulse methylprednisolone may be used.
Bone scan of systemic JIA showing abnormal bone tracer uptake in the left knee, right ankle, right midfoot, and possibly the left radial wrist and upper cervical spine.

<table>
<thead>
<tr>
<th>Laboratory test</th>
<th>Typical result</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC</td>
<td>Elevated (20,000–60,000)</td>
</tr>
<tr>
<td>Platelets</td>
<td>Elevated</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>Decreased (normocytic, hypochromic)</td>
</tr>
<tr>
<td>ESR</td>
<td>Markedly elevated</td>
</tr>
<tr>
<td>CRP</td>
<td>Elevated</td>
</tr>
<tr>
<td>Ferritin</td>
<td>Markedly elevated</td>
</tr>
<tr>
<td>Serum IgG</td>
<td>Elevated</td>
</tr>
<tr>
<td>RF, ANA</td>
<td>Usually negative</td>
</tr>
</tbody>
</table>

Echocardiogram showing a large pericardial effusion (white line) in a child with systemic JIA (16a); chest radiograph showing pleural effusion and mild mediastinal shift to the right (16b).
initially (30 mg/kg daily intravenously), with a lower daily dose of prednisone. Methotrexate has been used as a DMARD with some improvement. Anti-TNF therapy may be effective; however, there seems to be a higher proportion of non-responders to anti-TNF therapy compared with other subtypes of JIA (Carrasco et al. 2004, Quartier et al. 2003). New trials are under way for the evaluation of agents that disrupt IL-1 and IL-6/IL-6R dependent signaling pathways for therapy of systemic arthritis as anti-IL-1β, IL-1 TRAP, IL-1RA, and IL-6R have been reported to dramatically improve the systemic symptoms of the disease (18). Initial studies have showed marked response in children who were steroid dependent or resistant to other therapies (Pascual et al. 2008, Schneider & Laxer 1998).

COMPLICATIONS
Complications of systemic arthritis include macrophage activation syndrome (MAS), pulmonary hypertension, cardiac anomalies, chronic deforming arthritis, and side effects due to medication toxicity. Of these MAS is most noteworthy. MAS is a macrophage-related disorder that involves dysregulation of mature macrophages, which results in fever, pancytopenia, liver inflammation, disseminated intravascular coagulation, and neurologic involvement. Muscosal bleeding, pulmonary failure, and cardiac and renal involvement may occur as well. It is closely related to secondary HLH and is associated with multiple rheumatic diseases as well as infections and malignancies. Behrens et al. found that 53% of 15 patients with systemic JIA had evidence of occult MAS on bone marrow aspirate, while only 13% carried a clinical diagnosis of MAS (Behrens et al. 2007). Laboratory findings include prolonged PT and PTT, decreased sedimentation rate from hypofibrinogenemia, markedly elevated ferritin, elevated liver enzymes, elevated triglycerides, elevated lactate dehydrogenase, and hypoalbuminemia. Abnormalities in perforin expression, granzyme B expression, and low NK cell activity have been associated with this disorder (Tristano 2008). A bone marrow biopsy can be diagnostic (19). Treatment is not standardized, but commonly involves corticosteroids, cyclosporine A, etoposide, cyclophosphamide, and other chemotherapeutic agents. Etoposide is traditionally used in treatment of HLH, but cyclosporine A has

18 Fever curve in a hospitalized child with systemic JIA showing dramatic resolution of fevers with initiation of anakinra (given at 18:00 on 5/19).

19 Bone marrow smear from a child with systemic JIA showing hemophagocytosis.
been more frequently reported for treatment of MAS associated with systemic JIA. There has now been at least one case report describing the successful use of the IL-1R antagonist anakinra for the treatment of MAS.

Oligoarthritis

DEFINITION
Oligoarthritis is defined as chronic arthritis of four or fewer joints during the first 6 months of disease in a patient younger than 16 years of age. Patients who develop this in more than four joints after the first 6 months are then classified as having extended oligoarthritis.

EPIDEMIOLOGY AND ETIOLOGY
Oligoarthritis accounts for 50–60% of children with chronic arthritis in North American and European populations (Adib et al. 2005). Estimates of incidence are in the range 1–18/100,000 per year (Cassidy et al. 2005). Prevalence is estimated at 8–400 per 100,000 children (Arguedas et al. 1998, Danner et al. 2006, Manners & Bower 2002, Martinez Mengual et al. 2007, Oen & Cheang 1996, Pruunsild et al. 2007). There is a peak incidence between 2 and 4 years of age and a female:male preponderance of 3:1. Oligoarthritis developing in older boys may raise the possibility of other diagnoses such as spondyloarthropathy. It seems likely that oligoarthritis is a multigenic disease – it is seldom familial although there may be a preponderance of family members with autoimmune disease in children with oligoarthritis. HLA-A2 has been shown to be increased in children with early onset oligoarticular JRA (Murray et al. 1999). Oligoarthritis is strongly associated with chronic non-granulomatous uveitis.

CLINICAL HISTORY
These patients present with inflammation of four or fewer joints in the first 6 months. Oligoarthritis typically presents in the lower extremities, with knee arthritis being the most common finding (20). Ankles are also commonly involved (21). Involvement of small joints of fingers or toes is rare (presence of a finger or toe with tenosynovitis is more characteristic of psoriatic arthritis or enthesitis-related arthritis). Morning stiffness in the affected joints is common, but young children may not be able to verbalize this symptom; parents may notice a limp in the morning that resolves by afternoon. There may be a history of the child no longer standing in the crib in the morning. Children with oligoarthritis do not usually present with extra-articular manifestations other than anterior uveitis. Uveitis may present either before or after the diagnosis of arthritis and is often asymptomatic until irreversible scarring is present. Uveitis carries a cumulative incidence of 10–15% in patients with oligoarthritis, and is more common in children with a positive ANA (Carvounis et al. 2006). Children may develop limb-length discrepancies or joint contractures as complications of disease. Early during the course of the disease, increased blood flow to the inflamed joint may cause increased growth of the affected limb or digit; however, continued inflammation may
less commonly cause the growth plate to fuse earlier than on the opposite side, causing a foreshortened limb or digit on the affected side (22).

**PHYSICAL EXAMINATION**
The affected joint may be warm to the touch, with a ballotable fluid accumulation and restriction of range of movement. These patients do not usually complain of joint tenderness but may display joint guarding on passive range or an abnormal gait. Measurement of the length of the affected limb and examination of the back for a pelvic tilt may uncover a limb-length discrepancy (23). A Baker cyst may be present on posterior examination of the knee (24). Uveitis must be diagnosed by an ophthalmologist with a slit-lamp looking for inflammation of the anterior chamber of the eye as the uveitis is asymptomatic until scarring is present (25).

**LABORATORY**
Inflammatory markers will usually be normal. RF and anti-CCP are usually negative. ANA, present in about 50% of patients, is most helpful in predicting risk for chronic uveitis. Younger children who are ANA positive in the first few years of disease are at the most risk of uveitis and need to be screened frequently (Table 3). Analysis of synovial fluid should be culture negative with an increase of inflammatory cells typical of inflammatory arthritis in the range 10,000–40,000 cells/mm³. Early in the disease course, a culture of synovial fluid may be necessary to rule out infection-related arthritis.

**IMAGING**
Radiographs may show joint space narrowing, bony overgrowth, joint narrowing, or periartricular osteopenia (26), although radiographs may be normal.

---

**Table 3**
Recommended frequency of ophthalmologic screening examination in patients with juvenile rheumatoid arthritis (Cassidy et al. 2005)

<table>
<thead>
<tr>
<th>Type</th>
<th>ANA</th>
<th>Age at onset, years</th>
<th>Duration of disease, years</th>
<th>Risk category</th>
<th>Eye examination frequency, months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oligoarthritis or polyarthritis</td>
<td>+</td>
<td>≤6</td>
<td>≤4</td>
<td>High</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>≤6</td>
<td>&gt;4</td>
<td>Moderate</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>≤6</td>
<td>&gt;7</td>
<td>Low</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>&gt;6</td>
<td>≤4</td>
<td>Moderate</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>&gt;6</td>
<td>&gt;4</td>
<td>Low</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>−</td>
<td>≤6</td>
<td>≤4</td>
<td>Moderate</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>−</td>
<td>≤6</td>
<td>&gt;4</td>
<td>Low</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>−</td>
<td>&gt;6</td>
<td>NA</td>
<td>Low</td>
<td>12</td>
</tr>
<tr>
<td>Systemic disease</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Low</td>
<td>12</td>
</tr>
</tbody>
</table>
Same twelve-year-old with history of left second toe arthritis at age 2 years. Leg-length discrepancy due to JIA causing overgrowth of the right leg with pelvic tilt. A lift under the left knee corrects the pelvic tilt (courtesy of Lisa Mangino PT, DPT) (23); Baker cyst seen in the right posterior knee fossa (24); band keratopathy and iridocyclitis in a child with JIA (courtesy of Glenn Jaffe MD) (25); knee radiographs of a child with oligoarticular JIA with right knee involvement demonstrating epiphyseal bony overgrowth and joint effusion (26).
MRI with gadolinium can confirm presence of synovitis, increased intra-articular fluid and/or bone marrow edema (27). This may be helpful in joints that are difficult to examine clinically such as the TMJ or hips. Contrast is necessary in small joints, but not necessarily in larger joints such as the knee and hip.

DIFFERENTIAL DIAGNOSIS
Differential diagnosis includes trauma or infection, either an acute infectious arthritis due to *Staphylococci*, *Streptococci*, or *Kingella kingae*, or an indolent infection such as tuberculosis (28). Also included in the differential is malignancy (namely bone-infiltrative cancers such as leukemia or neuroblastoma), hemophilia, or other forms of early rheumatic disease (enthesitis-related arthritis, psoriatic arthritis, sarcoidosis, systemic lupus erythematosus (SLE), or periodic fever syndrome). Pain out of proportion to physical findings, night awakenings, and abnormalities with cytopenias on the blood smear are clues to an underlying malignancy.

PROGNOSIS
Prognosis is variable. Some children continue to have fewer than five joints affected and are more likely to go into remission, although flares may occur several years later. Children with extended oligoarthritis are more likely to have progressive disease. However, serious functional disability is uncommon, and mortality is extremely rare. Uveitis activity does not correlate with severity or timing of arthritis, and may be more difficult to treat than the arthritis. JRA-associated uveitis results in decreased visual acuity in 9% of patients, with cataracts or glaucoma in 20%, and band keratopathy in 16% (29) (Arguedas et al. 1998).

MANAGEMENT
Initial management consists of NSAID therapy and consideration of an intra-articular joint injection to the involved joint(s) with corticosteroids (Table 4 over page). Side effects of NSAIDs including gastritis and pseudoporphyria should be monitored. A physical therapy referral for passive and active stretching, along with gait training, may also be helpful. Knees and wrists that have contractures should be placed in an extension splint when sleeping (30). Intra-articular joint injection with triamcinolone hexacetonide is a well-tolerated, long-lasting preparation. Joint injection may be repeated up to three times per year without significant side effects, but those who continue to have ongoing disease may need methotrexate, or rarely biologic therapy. A shoe lift can be efficacious in children with significant limb-length discrepancy (>1 cm).
28 Child with tuberculous arthritis of the knee with corresponding radiograph demonstrating joint destruction.

29 Severe iridocyclitis with shallow anterior chamber, posterior synechiae, and secondary cataract, seen with a slit beam from a slit-lamp (courtesy of Glenn Jaffe MD).

30 Leg extension splint in a child with a knee contracture that is to be worn at night to leave the leg in maximal comfortable extension (courtesy of Lisa Mangino PT, DPT).
Uveitis is managed by ophthalmologists initially with topical steroid eye drops and mydriatic agents to prevent synechiae. For uveitis unresponsive to topical drops, systemic corticosteroids, local sub-Tenon corticosteroid injections, or systemic use of methotrexate is considered. There have been multiple reports of excellent efficacy of the TNF-α-blocking antibodies (adalimumab or infliximab) for the treatment of resistant uveitis.

**Table 4**

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Typical medications</th>
<th>Typical dosage</th>
<th>Indication</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>NSAIDs</td>
<td>Naproxen (Naprosyn)</td>
<td>15 mg/kg orally divided twice daily</td>
<td>Pain relief, mild anti-inflammatory effect</td>
<td>Gastritis, pseudo-porphyrin, renal toxicity</td>
</tr>
<tr>
<td></td>
<td>Ibuprofen</td>
<td>40 mg/kg orally divided three times daily</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Meloxicam</td>
<td>0.125 mg/kg orally daily</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corticosteroid joint injection</td>
<td>Triamcinolone hexacetonide</td>
<td>Large joints 1–2 mg/kg Small joints 0.25–0.5 mg/kg</td>
<td>First-line agent for oligoarticular disease, bridging therapy</td>
<td>Joint infection, skin atrophy</td>
</tr>
<tr>
<td></td>
<td>Betamethasone</td>
<td>Finger joints 0.3 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Classic DMARDs</td>
<td>Methotrexate</td>
<td>0.5–1 mg/kg orally or subcutaneously weekly (10–15 mg/m²) Give with folic acid 1 mg orally daily</td>
<td>Mild-moderate polyarticular arthritis</td>
<td>Oral ulceration, nausea, vomiting, teratogenicity, immunosuppression, hepatitis, blood count dyscrasias, concern for malignancy, GI upset, allergic rash, pancytopenia, and renal and hepatic toxicity</td>
</tr>
<tr>
<td></td>
<td>Sulfasalazine</td>
<td>Initial: 12.5 mg/kg orally daily Maintenance: 40–50 mg/kg orally divided twice daily</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**COMPLICATIONS**

Chronic complications of oligoarthritis include limb-length discrepancy, joint contractures, periarticular osteopenia, and bony overgrowth of the affected joint. Medications may cause side effects, especially NSAID-induced pseudoporphyria. Complications related to uveitis include blindness, visual loss, cataracts, and/or glaucoma.
<table>
<thead>
<tr>
<th>Drug class</th>
<th>Typical medications</th>
<th>Typical dosage</th>
<th>Indication</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leflunomide</td>
<td>10–20 mg orally daily</td>
<td>GI upset, hepatic toxicity, allergic rash, reversible alopecia, teratogenicity – needs washout with cholestyramine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biologics</td>
<td>Anti-TNF-α</td>
<td>Etanercept 0.8 mg/kg subcutaneously weekly</td>
<td>Moderate–severe polyarticular arthritis</td>
<td>Immunosuppression, concern for possible malignancy</td>
</tr>
<tr>
<td></td>
<td>Infliximab 3–10 mg/kg intravenously every 4–8 weeks</td>
<td></td>
<td>arthritis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Adalimumab &lt;30 kg: 20 mg subcutaneously every other week &gt;30 kg: 40 mg subcutaneously every other week</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-CTLA-4 Ig</td>
<td>Abatacept 10 mg/kg intravenously every 4 weeks max 1000 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-CD20</td>
<td>Rituximab 750 mg/m² intravenously * 2, 2 weeks apart, max 1000 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-IL-1</td>
<td>Anakinra 1–2 mg/kg</td>
<td>Systemic subcutaneously daily</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Immunosuppression arthritis</td>
</tr>
</tbody>
</table>
Polyarthritis

DEFINITION
Polyarthritis, or polyarticular JIA, is defined as chronic arthritis involving more than four joints during the first 6 months of disease in a patient younger than 16 years of age (see Table 1). Polyarticular JIA is further classified as RF-negative or RF positive. The RF must remain positive on two separate assays at least 3 months apart. Presence of enthesitis-related arthritis, psoriatic arthritis, and systemic arthritis is part of the exclusion criteria.

EPIDEMIOLOGY AND ETIOLOGY
Polyarthritis accounts for 30% of children with chronic arthritis. Estimates of incidence are in the range 0.29–8.9/100,000 per year (Danner et al. 2006, Malleson et al. 1996, Martinez Mengual et al. 2007, Moe & Rygg 1998, Pruunsild et al. 2007). Estimates of prevalence arc in a wide range of 4.4–54.2 per 100,000 (Danner et al. 2006, Gäre & Fasth 1992, Martinez Mengual et al. 2007, Moe & Rygg 1998). Age at onset has a bimodal distribution, with an early peak between the ages of 1 and 4 years and a later peak between 6 and 12 years. RF-positive children usually present in the teenage years. Girls are affected with polyarticular arthritis more often than boys, with ratios reported of up to 4.5:1 (Moe & Rygg 1998). As with the other subtypes of JIA, the etiology is likely multifactorial and unknown. Some children may have a genetic predisposition that likely involves interaction between multiple genes. Genetic polymorphisms may be more important in RF-positive disease. Anti-CCP antibodies are less prevalent in JIA than in adult arthritis but are detectable in a significant proportion of RF-positive polyarticular JIA patients. The disease involves dysregulation of both the humoral and cell-mediated immune responses, with a pro-inflammatory cytokine profile.

CLINICAL HISTORY
Polyarthritis commonly involves the large joints and tends to be symmetric. The cervical spine and TMJ are also commonly involved in this subtype (31a, 31b, 32a, 32b). The onset of arthritis is often subtle as a child may not complain of pain but rather may quit performing their usual functional activities. The onset is often more aggressive in those who are RF positive. A history of morning stiffness, sometimes lasting hours, or gelling after inactivity is typical. Extra-articular manifestations may include low-grade fever, anorexia with poor weight gain, and fatigue. Pericardial and pleural effusions, adenopathy, and hepatosplenomegaly may be seen, but not as
Coronal CT of the mandible of a child with erosive disease of the TMJ. Note the asymmetry and destruction of the condyle on the right (32a); moderate-to-marked degenerative changes involving bilateral TMJ joints including erosive changes involving the mandibular condyles (32b).

Rheumatoid nodules in a RF+ female with erosive arthritis and interstitial lung disease (33a, b); pseudorheumatoid nodule (33c).

Prominently as in systemic arthritis. Rheumatoid nodules occur almost exclusively in individuals with RF-positive disease (33a–c). A small-to-medium vessel vasculitis of the skin or internal organs may rarely occur in RF-positive polyarthritis. Other extra-articular manifestations seen in RF-positive polyarthritis, such as Felty syndrome, cardiac disease, and pulmonary involvement are rare in children compared with their adult counterparts. As in children with oligoarthritis, age and the presence of ANA are predictors for the development of uveitis and determine the frequency of necessary screening with a slit-lamp exam (see Table 3 page 22).
PHYSICAL EXAMINATION
Inspection of the joints is helpful to assess any asymmetry. Arthritic joints have palpable fluid effusions and/or pain with restricted ROM. The involved joints may emit heat or warmth. They may be tender to palpation and sometimes have a purplish discoloration overlying, especially in the small finger joints of RF-positive patients. Arthritis is often symmetrical and can involve small joints in the hand and wrist (34a–c). Children may not report pain as much as adults in the clinic setting; however, on examination they may guard due to pain with ROM (Anthony & Schanberg 2007). Cervical spine involvement is often evident and is seen clinically as restricted neck extension. Auscultation of the TMJ may reveal crepitance with jaw opening and closing. Measurement of the maximal interincisal distance helps to assess and follow ROM of the TMJ. Tenosynovitis, inflammation of tendon sheaths, and synovial cysts may be present, especially in the hands (35). Hip involvement is not uncommon and should be evaluated by performing a complete hip ROM exam. Assessment for a leg-length discrepancy is important in children with lower extremity involvement. Patients may have an antalgic gait with a limp from involvement of the hip, knee, ankle, or feet.

LABORATORY
The diagnosis of polyarthritis is a clinical one, thus laboratory tests are only supportive. Hematologic abnormalities may be present, including leukocytosis, thrombocytosis, and anemia of chronic disease. Evidence of inflammation with elevations in ESR and CRP, along with elevated immunoglobulins, are common in polyarticular JIA. Fewer than 10% of children with JIA will have positive RF, but the presence of high titer

34a–c, 35 Girl with polyarthritis with involvement of wrist and fingers (34a); wrist swelling in a girl with polyarticular arthritis (34b); wrist swelling in a girl with polyarticular arthritis (34c); hand of girl with polyarthritis. Note the tendon sheath effusion of the second digit with fluid extending beyond the joint line, and the flexion contracture of the fifth digit (35).
portends a poorer prognosis (Gilliam et al. 2008). The presence of antibodies to CCP also suggests more aggressive disease (Gilliam et al. 2008). The presence of ANA indicates an elevated risk of developing uveitis (Table 3).

**IMAGING**
Plain radiographs are often normal in part because of the reduced ability to detect changes in incompletely calcified structures. Changes on radiographs may be seen early, revealing periarticular osteopenia, or decreased joint space (36a–e).

36a–e Hand radiograph demonstrating multiple erosions (arrows) in a child with recalcitrant polyarthritis (36a); hand radiographs at diagnosis of a 7-year-old girl with polyarthritis demonstrating periarticular osteopenia around the carpal bones (36b, c); shoulder joint space loss, erosions, and osteopenia in a 16-year-old girl with severe RF+ arthritis (36d); soft tissue swelling over wrist arthritis (36e).
Later bony overgrowth, bone erosions or ankylosis may occur (37, 38). Accelerated ossification of carpal bones may be seen in younger children (39). As with any form of arthritis, MRI with gadolinium can confirm the presence of synovitis, bone erosions, increased intra-articular fluid, and/or bone marrow edema (40a, b, 41a, b, 42). Radiographs of the hip may show structural abnormalities (43), but are not sensitive for demonstrating hip effusions; however, they are used to demonstrate disease pre- and post-hip replacement (44a, 44b)

37 Ankylosis of the wrist bones in a patient with polyarticular arthritis.

38 Hand radiograph demonstrating carpal and metacarpal–phalangeal ankylosis in a child with recalcitrant polyarthritis.

39 Hand radiographs demonstrating asymmetric and accelerated carpal maturation on involved left hand compared with non-involved right hand. There is early ossification of the ulnar styloid and larger trapezoid and scaphoid bones on the left.

40a, b MRI of the TMJ: pre-contrast infusion showing flattening of the condyle (40a); post-contrast image with gadolinium showing a joint effusion of the TMJ (40b).
Eight year old with wrist arthritis seen on plain radiograph demonstrating erosions and crowding of the carpal bones (41a); 8 year old with wrist arthritis: contrast-enhanced MRI of the same wrist demonstrates erosions and synovial enhancement and fluid within the joint (41b); hip MRI demonstrating hip effusion on the right (42).

Hip radiograph revealing flattened femoral heads from long-standing polyarthritis (43); 16 year old girl with severe RA pre hip replacement (44a); 16 year old girl with severe RA post-hip replacement (44b).
Ultrasound can be particularly helpful in evaluation of possible hip effusion (45). Chest imaging is warranted with concerns for arthritis-related lung disease; CT scan demonstrates ground glass throughout the lung fields and nodules (46a, 46b).

DIFFERENTIAL DIAGNOSIS
Differential diagnosis includes polyarthritis associated with other rheumatologic diseases, including SLE, enthesitis-related arthritis, sarcoidosis, dermatomyositis, and scleroderma. The arthritis associated with these other autoimmune diseases may be difficult to differentiate from polyarticular JIA without looking for characteristic clinical or laboratory findings (e.g. presence of double-stranded DNA antibodies indicative of lupus or rash of dermatomyositis). Infections most often cause oligoarticular arthritis, but rarely present with a polyarticular picture, such as with infections caused by Neisseria gonorrhoeae, acute Lyme disease, or acute rheumatic fever. Polyarthritis may be found in children with serum sickness, IBD, and sickle cell disease. Whipple disease is another rare cause of polyarticular arthritis. Malignancies may cause periarticular pain and swelling or actual joint effusions, and are included in the differential diagnosis.

PROGNOSIS
Poor predictors of outcome include delay in diagnosis, longer disease duration, persistent active disease, early development of bony erosions, and presence of subcutaneous nodules. RF seropositivity and presence of anti-CCP are also poor predictors, and these patients tend to have more aggressive disease unless aggressively treated (Gilliam et al. 2008, Habib et al. 2008, Selvaag et al. 2005). Overall prognosis for a life without disability is good, however, especially with the increase in number of new available medications.

MANAGEMENT
NSAIDs are used for pain relief and decrease in inflammation but alone rarely prevent joint damage (see Table 4). Second-line medications, including DMARDs, are often required to treat polyarthritis. Methotrexate has long been the standard therapy for polyarthritis. It is given in low-dose, weekly administration either by mouth or by subcutaneous injection (for improved absorption). Both leflunomide and sulfasalazine have been shown to be beneficial (Silverman et al. 2005, van Rossum et al. 1998). Biologic agents, specifically TNF-α inhibitors, are becoming more important in the management of polyarticular disease (Lovell et al. 2000, Lovell et al. 2008, Ruperto et al. 2007). Anti-TNF-α agents include etanercept, adalimumab, and infliximab. Abatacept, which blocks T-cell costimulation, has recently been approved for use in polyarticular JIA (Ruperto et al. 2008). Rituximab, an antibody directed toward B cells, has been shown to be effective in adult rheumatoid arthritis (Cohen et al. 2006). Intra-articular injections of corticosteroids can be used to treat a few recalcitrant joints, or as a bridging therapy while awaiting the effects of DMARD treatment. Systemic corticosteroids should be used sparingly, with prolonged use discouraged due to the unique side effects on the growing skeleton such as growth retardation and osteopenia. Physical therapy and occupational therapy are also important elements in the treatment of children with polyarthritis as attention needs to be paid to not only current functional status but also expected changes with growth and maturity. Children may need special school accommodation provided.
CT chest scan of 11-year-old girl with RF+ arthritis, rheumatoid nodules, and interstitial lung disease with ground-glass appearance and pulmonary nodule.

Ultrasound scan of bilateral hips revealing large effusion on the right.
COMPLICATIONS
Chronic arthritis can lead to disturbances in growth of the involved joint, causing bony overgrowth, lengthening of an affected limb, or premature closure of growth plates and shortening of an affected limb. With chronically active disease, overall linear growth of a child may also be slowed. Chronic complications also include joint contractures with restriction in ROM, erosive changes in bone, and periarticular osteopenia (47, 48). Disease of the TMJ in polyarthritis can lead to micrognathia and retrognathia. Disease of the cervical spine in polyarthritis can lead to degeneration of the vertebral bodies and cervical spine fusion, and may lead to impingement of the spinal cord (see 31a, 31b). Medications used to treat arthritis may also have side effects.

47 Hand in a 19 year old with polyarthritis. He was initially diagnosed with polyarthritis at age 3 years and now has contractures at his MCPs with subluxation at the base of his thumb. His pip contractures have been surgically corrected.

48 The same young man (as shown in 47) demonstrating contractures of toes.
Spondyloarthropathies and reactive arthropathies

- Spondyloarthropathies
- Reactive arthropathies
- Post-streptococcal arthropathies
The term ‘spondyloarthropathy’ refers to a collection of inflammatory conditions in which both axial (spine) and appendicular (peripheral) joints are involved. Inflammation at the sites of ligamentous insertion, also known as enthesitis, is also a hallmark of these diseases. The joint involvement is primarily in the lower extremities and develops at age 16 years or younger (Gensler & Davis 2006).

Reactive arthropathies can be defined as a sterile inflammatory arthropathy temporally associated with a remote antigenic exposure. While this antigenic exposure is usually infectious, in conditions like serum sickness, a similar reactive inflammatory arthropathy can occur to non-infectious agents such as a medication. One of the challenges in writing about these conditions is the diversity in terminology that has been used in the past to describe these conditions. Several of these conditions have been labeled in alternative ways and presently could be better classified as forms of juvenile idiopathic arthritis (JIA). Examples of this include juvenile psoriatic arthritis and enthesitis-related arthritis which are subtypes of JIA in the ILAR classification system (see below). Unfortunately, much of the literature describing these conditions predates these criteria, and so the previous framework for describing these conditions is used in this chapter. An example of this approach is shown in the next section on the broad category of juvenile spondyloarthropathy.

**SPONDYLOARTHROPATHIES**

### Juvenile spondyloarthropathy

**DEFINITION**

In 1991 criteria were established by the European Spondyloarthropathy Study Group (ESSG) to diagnose spondyloarthropathies in which one major criterion and one minor one are required (Dougados et al. 1991). These criteria are summarized (Table 5). The ESSG criteria have been validated in children (Prieur et al. 1990).

**EPIDEMIOLOGY AND ETIOLOGY**

Determining epidemiologic data for juvenile spondyloarthropathy (JSpA) is challenging because of plural criteria for diagnosis, lack of key clinical features at diagnosis, and the lack of confirmatory radiographic data, particularly

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Criteria for the diagnosis of spondyloarthropathy (Dougados et al. 1991)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Major criteria</strong></td>
<td><strong>Minor criteria</strong></td>
</tr>
<tr>
<td>Synovitis that is asymmetric or predominantly of the lower limbs</td>
<td>Positive family history, which is defined as presence in first-degree or second-of any of the following: AS, degree relatives psoriasis, acute uveitis, reactive arthritis, or IBD</td>
</tr>
<tr>
<td>Inflammatory spinal pain</td>
<td>Psoriasis diagnosed by a physician; IBD (either Crohn disease or ulcerative colitis) diagnosed by a physician and confirmed by radiographic examination or endoscopy</td>
</tr>
<tr>
<td>Non-gonococcal urethritis or cervicitis, or acute diarrhea within 1 month before arthritis</td>
<td>Buttock pain alternating between right and left gluteal areas</td>
</tr>
<tr>
<td>Enthesopathy, defined as spontaneous pain or tenderness at the site of insertion of the Achilles tendon or plantar fascia</td>
<td>Radiographic evidence of sacroiliitis by plain films</td>
</tr>
</tbody>
</table>

At least one major criterion and one minor criterion are needed to make diagnosis of spondyloarthropathy. Inflammatory spinal pain is defined as three of the following: at least 3 months’ duration, onset before 45 years of age, insidious onset, improved by exercise, or associated with morning stiffness.

**Box 1 Keys to recognizing JSpA**

- Chronic, inflammatory arthritis usually oligoarticular, lower extremities
- Enthesitis
- IBP
- Anterior uveitis
- Associated disease features: nail pits, psoriatic rash, mucositis, dysuria, diarrhea, and so on.
of the axial skeleton. Estimates of the prevalence of JSpA based on inventories of patients seen at pediatric rheumatology centers suggest that from 1% to 20% of children with a specific rheumatic condition may have JSpA (Rosenberg 2000). The etiology of JSpA is largely unknown, but seems to be associated with HLA-B27 (see below).

**CLINICAL HISTORY**

Key clinical features of JSpA are shown (box 1). Since less than one-quarter of children with JSpA have axial spine pain, stiffness, or restricted motion or signs or symptoms of sacroiliitis at presentation (Rosenberg 2000), the diagnosis is often delayed. The diagnosis often rests on finding inflammatory arthritis in the lower extremities along with at least one of the minor ESSG criteria, for example knee involvement as shown (49).

Another prominent clinic feature of JSpA is uveitis. Anterior uveitis (50), which is usually associated with pain, photophobia and redness, is more associated with ankylosing spondylitis (AS) and reactive arthritis (ReA) (Paiva et al. 2000). Posterior uveitis (51) is more subtle, often painless, and even asymptomatic. In JSpA, posterior uveitis is seen more with psoriatic arthritis (PsA) and the arthropathy associated with IBD (Paiva et al. 2000).

49 Right knee effusion. Notice the swelling in lateral suprapatellar pouch of the right knee. This 14-year-old boy had a bulge sign and pain on extreme knee flexion, but no antalgic gait. Oligoarthritis of the lower extremities is a frequent presentation for spondyloarthopathies.

50 Acute anterior uveitis with hypopyon. Note the marked conjunctival injection and the cloudiness of the anterior chamber. These patients usually present with red eye and eye pain with photophobia (courtesy of D. C. Herman, MD, Mayo Clinic).

51 Severe posterior uveitis. In this funduscopic image the vitreous is so cloudy with inflammatory cells that it is difficult to make out other normal structures such as the disc, vessels, or retinal detail (courtesy of D. C. Herman, MD, Mayo Clinic).
PHYSICAL EXAMINATION
Enthesitis is a term that refers to inflammation at the sites of ligamentous insertion on to bone near a joint. Frequent areas of enthesitis in spondyloarthropathies include the insertion of the Achilles tendon (52) and that of the plantar fascia on the heel, and the patellar tendon insertion on the tibia. These areas are tender on exam, and may be red, warm or swollen. While joint pain, swelling, tenderness, limited ROM, and enthesitis are hallmark features of all forms of JSpA, other common distinguishing clinical features (including axial findings) will be discussed later.

LABORATORY
Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP and evidence of the anemia of chronic disease. The presence of HLA-B27 increases the odds ratio of a diagnosis of a JSpA, particularly if there is axial involvement. The prevalence of HLA-B27 is not clear in JSpA although estimates have ranged from 40% to 90% (Burgos-Vargas 2002, Rosenberg 2000). Autoantibodies are not commonly seen in JSpA.

IMAGING
Imaging of the peripheral joints may show soft tissue swelling that is apparent on clinical exam. Rarely destructive changes such as joint space narrowing or erosions may be present.

Plain radiographs of the spine are frequently normal, even in those with axial involvement of JSpA. With more sensitive imaging techniques, such as CT and MRI, the presence of inflammatory changes around the sacroiliac (SI) joints may be more apparent, for example early subtle changes in JSpA (53), and more dramatic SI disease (54a, 54b).

DIFFERENTIAL DIAGNOSIS
Since axial involvement is uncommon at the outset of JSpA, other conditions that share the clinical features of inflammatory arthritis, particularly an oligoarthritis of the lower extremities in children should be considered. One of these is enthesitis-related JIA. Children with enthesitis-related JIA may later develop a more classic spondyloarthropathy as described above. These criteria were developed in 1998 and updated in 2001 (Petty et al. 2004) (Table 6).
Early sacroiliitis. In this T2-weighted MRI scan, subtle findings of bone marrow edema (lighter areas) adjacent to the SI joints are highlighted by arrows (53); CT scan of SI joints of a 6-year-old boy with bilateral gluteal pain and waddling gait who was HLA-B27 negative. There are erosions (arrows) on the sacral side anteriorly of both SI joints, more impressively on the right. The SI joints were aspirated and all cultures were negative (54a); CT scan of SI joints of the same boy (54a) 26 months later. Notice the marked increase in number of erosions (arrows) especially on the iliac side of the SI joints and the demineralization of the sacrum in spite of treatment with sulfasalazine and oral methotrexate. The child responded dramatically to etanercept therapy (54b).

Table 6
Criteria for the diagnosis of the enthesitis-related arthritis subgroup of juvenile idiopathic arthritis (Petty et al. 2004)

<table>
<thead>
<tr>
<th>Diagnosis confirmed</th>
<th>Diagnosis confirmed</th>
<th>Exclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arthritis and enthesitis</td>
<td>Arthritis or enthesitis with at least two of the following:</td>
<td>Dermatologist confirmed psoriasis in at least one first-degree relative</td>
</tr>
<tr>
<td></td>
<td>• (+) HLA-B27</td>
<td>Systemic onset arthritis</td>
</tr>
<tr>
<td></td>
<td>• Sacroiliac tenderness and/or inflammatory spinal pain</td>
<td>Positive RF</td>
</tr>
<tr>
<td></td>
<td>• Family history in first-degree relative with uveitis, AS, ReA, IBD-related sacroiliitis, or enthesitis-related arthritis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Anterior uveitis, acute</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Arthritis in a boy after age 6 years</td>
<td></td>
</tr>
</tbody>
</table>
The other common condition with similar features is pauciarticular juvenile rheumatoid arthritis (pauci-JRA). In the US, many of these children will be initially diagnosed with pauci-JRA, reflecting the slight difference in nomenclature between JIA and JRA. There are similarities and differences between the spondyloarthropathies and pauci-JRA (Table 7).

**PROGNOSIS**

In population-based studies from Norway, children with spondyloarthropathies were a subset of children with chronic arthritis who had a poorer outcome (Selvaag et al. 2005). In these studies of nearly 200 children with JRA, 7 with JSpA, and 5 with juvenile PsA (JPsA), those with JSpA and JPsA had lower levels of physical function and wellbeing, and higher levels of pain compared with the rest of the cohort (Selvaag et al. 2005).

**MANAGEMENT**

Within the management principles for chronic rheumatic diseases in general, there are particular treatment principles for JSpA (box 2). More specific aspects of medical management of the axial and peripheral arthritis of JSpA will be highlighted in the following sections.

**COMPLICATIONS**

The potential complications of the various forms of JSpA will be discussed below.

---

**Box 2 Treatment principles for management of JSpA**

- Establish correct diagnosis
- Assess pain, functional status and risk for progression and damage
- Patient/family education
- PT/OT to optimize functional status
- Medications administered after risk/benefit ratio assessment

---

**Juvenile ankylosing spondylitis**

**DEFINITION**

Juvenile ankylosing spondylitis (JAS) is defined as the diagnosis of AS before age 17 years; it may also be called juvenile-onset ankylosing spondylitis (JoAS) to contrast with adult-onset ankylosing spondylitis (AoAS) (Burgos-Vargas et al. 1997). The diagnosis of AS is made based on a combination of clinical and radiographic features (Table 8). These criteria (van der Linden et al. 1984) are weighted heavily to radiographic findings.

**EPIDEMIOLOGY AND ETIOLOGY**

The prevalence of AS is in the range of around 70–210 cases per 100,000 adults and has an incidence of between seven and nine cases per 100,000 person-years depending on the population studied (van der Linden et al. 2008). In patients diagnosed with AS, about 10–20% will be before age 17 years (JAS), although there are populations with higher juvenile prevalence in Mexico and Korea (Burgos-Vargas et al. 1996). The etiology of JAS is not known, but is also associated with HLA-B27 (see below).

**CLINICAL HISTORY**

Patients with JAS frequently present with non-axial symptoms and a high index of suspicion is needed to make this diagnosis. Confirmatory radiographic findings are frequently absent early in the illness. The key feature of axial involvement is inflammatory back pain (IBP). Features of IBP include stiffness, particularly in the mornings, that is helped by activity and chronicity (usually months of symptoms). Sometimes the pain will be perceived in the buttocks.

In a Korean study of nearly 100 consecutive AS cases from 1997 to 1998, patients with JAS were more likely to present with peripheral joint symptoms and less likely to present with spinal symptoms compared with adults with AS (Baek et al. 2002). In this study radiographic findings of ankylosis and syndesmophytes were much less in the JAS group as was limitation of spinal ROM (Baek et al. 2002). Similar findings were described in 150 patients with AS and 60 with JAS from India (Aggarwal et al. 2005).
Table 7
Comparison of clinical features of juvenile spondyloarthropathies and juvenile rheumatoid arthritis

<table>
<thead>
<tr>
<th></th>
<th>Pauci-JRA</th>
<th>JSpA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at onset</td>
<td>Preschoolers</td>
<td>Older</td>
</tr>
<tr>
<td>Gender</td>
<td>Girls</td>
<td>Boys</td>
</tr>
<tr>
<td>Spine involvement</td>
<td>No</td>
<td>Characteristic, but often not at diagnosis</td>
</tr>
<tr>
<td>SI involvement</td>
<td>No</td>
<td>Characteristic, but often not at diagnosis</td>
</tr>
<tr>
<td>Enthesitis</td>
<td>No</td>
<td>Characteristic</td>
</tr>
<tr>
<td>Uveitis</td>
<td>Chronic posterior (increased with positive ANA)</td>
<td>Acute anterior</td>
</tr>
<tr>
<td>Joint involvement</td>
<td>Oligoarthritis</td>
<td>Oligoarthritis</td>
</tr>
<tr>
<td>Autoantibodies</td>
<td>ANA</td>
<td>None</td>
</tr>
<tr>
<td>HLA associations</td>
<td>Mostly class II, Class I: B27 multiple</td>
<td></td>
</tr>
<tr>
<td>Other associated conditions</td>
<td>–</td>
<td>Psoriasis, IBD, Gi/GU infections, oral mucositis</td>
</tr>
</tbody>
</table>

Table 8
Criteria for the diagnosis of ankylosing spondylitis (van der Linden et al. 1984)

<table>
<thead>
<tr>
<th>Radiographic findings</th>
<th>Clinical findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>At least moderate bilateral SI changes (at least 2 on a 4-point scale)</td>
<td>IBP</td>
</tr>
<tr>
<td>At least severe unilateral SI changes (at least 3 on a 4-point scale)</td>
<td>Restricted ROM in lumbar spine in at least planes of movement</td>
</tr>
<tr>
<td>Decreased chest expansion</td>
<td></td>
</tr>
</tbody>
</table>

At least one radiographic feature and one clinical feature are needed to make the diagnosis of AS.
PHYSICAL EXAMINATION

Inflammatory oligoarthritis, particularly in the lower extremities, and enthesitis are common features of JAS, and were discussed previously. An assessment of the ROM of the spine and the presence of SI tenderness is critical to evaluate children with JAS.

To assess SI tenderness, direct palpation is important, but may be difficult to reproduce. Compressive maneuvers such as the Patrick test may be more specific. This test is also known as the FABER maneuver (flexion, abduction and external rotation).

To assess spine ROM, the spine is examined in three regions: cervical, thoracic, and lumbar. There are several techniques for examining each region, and an example for each follows. To assess cervical spine ROM, the wall-to-tragus measurement is helpful. To assess thoracic spine mobility, measurement of chest expansion is done. Measurement of lumbar spine ROM may be assessed by the modified Schober test.

LABORATORY

Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP and evidence of the anemia of chronic disease. Almost 90% of JAS patients are HLA-B27 positive. Autoantibodies currently measured are most often absent in JAS.

55 Patrick test. The patient, while supine on an examination table (A), is asked to flex and externally rotate a hip (B). Then pressure from the examiner is placed on the contralateral anterior iliac crest and the ipsilateral femur towards the lumbar spine (C). SI pain is usually perceived in the ipsilateral SI area. This is repeated on the opposite side to assess each SI joint.
Wall-to-tragus distance. A patient stands against a wall or stable vertical planar surface, keeping knees straight, eyes looking straight ahead, and chin parallel to floor. The patient is then asked to extend his or her neck and the distance from the posterior tragus to the wall is measured and recorded. This measurement helps assess cervical spine ROM (56). Chest expansion. From behind a standing (or seated) patient, a tape measure is placed around the chest at T4. The patient is asked to take a deep breath (B), and then exhale. The chest circumference is measurement at the end of exhalation and noted (C). The patient is then asked to take another deep breath (D), and the measurement of chest circumference at maximum inhalation is noted. Chest expansion is the difference in measurements at points C and D (57).

Modified Schober test. From behind a standing patient, the surface marker of the L2 vertebra is noted (approximately at the level of the iliac crests). On either side of the spinous processes, this point is marked, and a second point 10 cm superior is also marked while the patient is still standing. The patient is then asked to bend forward, with knees extended. At the point of maximal flexion, the distance between the two points is re-measured. The difference between the first and second measurement is recorded.
IMAGING
If JAS is suspected, plain radiographs of the SI joints are required. Often a 30° tilt view is required to image the SI joint en face. Also AP and lateral views of the lumbosacral spine are helpful, which may include views in extremes of flexion and extension.

Early radiographic findings include changes of the SI joints (59). Later changes in the lumbar spine include ‘squaring’ and ‘shiny corners,’ which are radiographic findings that represent a proliferative process at areas of contact on the anterior longitudinal ligament and the anterior surface of the vertebral body (60).

These radiographic findings usually precede ‘bamboo spine’, which represents calcification of both the anterior and the posterior longitudinal spinal ligaments and is a very late finding (61).

Given that most JAS patients may not have axial symptoms, the frequency of radiographic changes at diagnosis of spine and SI joints is not clear. But even in those with axial symptoms who get conventional radiography, the frequency of characteristic changes of sacroilitis of JAS is low. The ability of MRI to detect potential inflammatory changes around the SI joints early in the course of adults with AS that predict diagnostic radiographic changes years later has been recently demonstrated (Bennett et al. 2008). In this study, gradation of bone marrow edema adjacent to the SI joints was the best predictor of subsequent development of AS in a cohort of patients with IBP of less than 2 years’ duration (better than B27 status) (Bennett et al. 2008). This important study may have significant implications for earlier diagnosis of JAS.

A study of nearly 80 patients with JAS and over 300 patients with AS who at least 20 years of disease duration confirmed earlier studies that axial radiographic changes are less in JAS compared with AS even with longer disease duration in the JAS cohort (Gensler et al. 2007).

DIFFERENTIAL DIAGNOSIS
Other conditions that present with inflammatory arthritis such as JRA/JIA share features with JAS, particularly if axial symptoms are absent. If enthesitis is present, other forms of JSpA should also be considered. Since peripheral inflammatory oligoarthritis ± enthesitis is a common presentation for JSpA in general and JAS in particular, specific questions about axial symptoms and assessment of spinal ROM are critical to avoid overlooking JAS in its early stages.

From an axial point of view, it is important to identify IBP from other forms of back pain. IBP is chronic as opposed to the acute back pain from injury or referred pain from urinary tract infection (UTI). Also IBP is better with activity as opposed to mechanical back pain, which is not. Back pain in a very young child, or that wakes the child from sleep is not likely to be from a spondyloarthropathy. Finally, there is stiffness in the back in the morning (or after prolonged rest, ‘gelling’) in IBP, which is not usually found in other forms of back pain.

PROGNOSIS
A cross-section of over 300 North American patients with JAS and 2000 patients with AS were assessed for functional status using the Bath Ankylosing Spondylitis Functional Index (BASFI) (Stone et al. 2005). In this study, those with JAS had longer duration of disease, greater delay in diagnosis, and higher BASFI scores (less function) than the AS cohort (Stone et al. 2005).

Also, JAS patients are at increased risk of developing severe hip involvement, as measured by an increased rate of total hip arthroplasty (Gensler et al. 2007). An example of the destructive hip arthropathy that can accompany JAS is shown (62).

MANAGEMENT
The important management principles for the axial features of JSpA, specifically JAS, are highlighted (box 3).

COMPLICATIONS
As with AS in adults, the primary potential complication of JAS is fusion of the spine and SI joints. The arthropathy in JAS is usually not destructive. Patients may also develop sequelae of uveitis, such as cataracts. In addition, complications of medical therapies such as gastritis associated with NSAIDs, infections associated with immunosuppressive medications, and bone disease related to corticosteroids may occur.
Unilateral sacroiliitis. ‘Bamboo spine.’ In this AP view of the lumbar spine, note the fusion of the SI joints, the preserved disc height, and the bilateral flowing calcifications (‘dripping candle wax’) from the proliferative enthesitis in this fused lumbar spine. Earlier, more subtle changes of this process are also noted on the superior aspect of the anterior surface of the L2 vertebral body.

61–62 Unilateral sacroiliitis. ‘Bamboo spine.’ In this AP view of the lumbar spine, note the fusion of the SI joints, the preserved disc height, and the proliferative enthesitis, which has calcified, is apparent on the anterior edge of the L4 vertebral body. Earlier, more subtle changes of this process are also noted on the superior aspect of the anterior surface of the L2 vertebral body.

Box 3 Overview of the treatment of spine disease in JSpA

- Good posture
- Maintain ROM, encourage cervical extension
- Longitudinal assessments
- NSAIDs are treatment mainstay
- TNF blockers
- Intra-articular injections (SI)
Juvenile psoriatic arthritis

DEFINITION
In 2004 a revised set of criteria were published for the diagnosis of JPsA (Petty et al. 2004), allowing a diagnosis of JPsA to be made if the child has psoriasis and arthritis or if the child has arthritis without psoriasis but fulfilled the criteria (Table 9).

EPIDEMIOLOGY AND ETIOLOGY
The overall prevalence of psoriasis is 1–3% and as many as one-third will develop it before age 16 years (Lewkowicz & Gottlieb 2004). The incidence of JPsA is about 3 cases per 100,000 children per year and has a prevalence of about 1/10,000 and accounts for up to 15% of children with chronic arthritis (Petty & Southwood 2005). The etiology of JPsA is unknown. It is not generally associated with HLA-B27 unless there is axial involvement (see below).

CLINICAL HISTORY
While most of the time psoriasis precedes the arthritis in adults, only 10–15% of children with JPsA have skin lesions at the time of diagnosis (Hafner & Michels 1996). The most common form of psoriasis in children is plaque psoriasis (Lewkowicz & Gottlieb 2004) (63). Another form is a predominantly guttate pattern of psoriasis, as shown in a 7-year-old boy at diagnosis (64).

Since the rash of psoriasis is frequently absent in children with JPsA, a careful inspection of the nails is important, as nail changes, particularly pitting (65, 66), may precede the rash of psoriasis in these patients.

There seem to be two groups of children who get JPsA. One is younger, preschool-aged girls who generally have a polyarthritis and dactylitis; the other group is older boys who may have axial involvement or enthesitis and are more likely to have evidence of skin disease (Stoll & Nigrovic 2006).

PHYSICAL EXAMINATION
In those with peripheral arthritis, frequently joints involved will have a ‘ray’ distribution. This refers to joints in the same ‘ray’ of an extremity, such as the metacarpophalangeal (MCP), proximal interphalangeal (PIP), and distal interphalangeal (DIP) joints (67). When the joints are closer together, such as the toe, this is called dactylitis and may be referred to as a ‘sausage digit’, which is a characteristic feature of JPsA (68). Another feature of the arthritis of JPsA is the frequency of DIP involvement, more so than in other forms of arthritis. This is noted frequently in a DIP adjacent to a fingernail with active psoriatic changes.

LABORATORY
Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP along with evidence of the anemia of chronic disease. The presence of HLA-B27 increases the odds ratio for a diagnosis of a JSpA such as JPsA, particularly if there is axial involvement. As with the other forms of JSpA, autoantibodies are not frequently found in patients with JPsA; however, when they are found they are most likely to be low-titer ANA.

Table 9
Criteria for the diagnosis of juvenile psoriatic arthritis subgroup of juvenile idiopathic arthritis (Petty et al. 2004)

<table>
<thead>
<tr>
<th>Diagnosis confirmed</th>
<th>Diagnosis confirmed</th>
<th>Exclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arthritis and psoriasis</td>
<td>Arthritis without psoriasis, but with at least two of the following:</td>
<td>Family history in first-degree relative</td>
</tr>
<tr>
<td></td>
<td>Dactylitis</td>
<td>uveitis, AS, ReA, IBD-related</td>
</tr>
<tr>
<td></td>
<td>Nail changes</td>
<td>sacroiliitis or enthesitis-related arthritis</td>
</tr>
<tr>
<td>Dermatologist confirmed</td>
<td>Positive RF B27 (+) boy older than 6 years</td>
<td>Systemic-onset arthritis</td>
</tr>
<tr>
<td>psoriasis in at least one first-degree relative</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

nasis (Hafner & Michels 1996). The most common form of psoriasis in children is plaque psoriasis (Lewkowicz & Gottlieb 2004) (63). Another form is a predominantly guttate pattern of psoriasis, as shown in a 7-year-old boy at diagnosis (64).
Psoriatic plaque. Oval-shaped, raised, scaly, hyperkeratotic lesion on the entensor surface of the leg of a young man with psoriasis (courtesy of Dawn M. R. Davis, MD, Mayo Clinic) (63); guttate distribution of psoriasis (courtesy of Dawn M. R. Davis, MD, Mayo Clinic) (64); nail pitting in a child with psoriasis. In the third and fourth fingernails, characteristic pitting is noted which may precede the development of arthritis (courtesy of Dawn M. R. Davis, MD, Mayo Clinic) (65); toenail changes in the child shown in 65. Note the asymmetry (courtesy of Dawn M. R. Davis, MD, Mayo Clinic) (66).

Inflammatory arthritis in right forefoot. Note the erythema and swelling over the medial aspects of the first metatarsophalangeal joint and interphalangeal joint of the great toe in this 14-year-old boy who was seronegative.

‘Sausage’ toe. Note the diffuse swelling of the second toe in this boy with severe nail changes from PsA (courtesy of C. J. Michet, MD, MPH, Mayo Clinic).
IMAGING
Since most patients with JPsA have peripheral joint involvement, they share imaging features with the other forms of JSpA.

DIFFERENTIAL DIAGNOSIS
With oligoarticular or axial presentation, the primary differential diagnosis is between the other forms of JSpA, enthesitis-related JIA and pauciarticular JRA (as discussed previously). The differential diagnosis of those with JPsA that present with polyarthritis includes polyarticular forms of JRA/JIA.

PROGNOSIS
Like most of the other forms of JSpA, the risk for erosive destructive arthritis in JPsA is low compared with JRA or JIA. As noted previously, however, the functional outcomes of children with JPsA seem to be worse than for children with other forms of chronic inflammatory arthritis (Selvaag et al. 2005).

MANAGEMENT
The principles of medical management of the peripheral musculoskeletal manifestations of JSpA such as JPsA are highlighted (box 4).

COMPLICATIONS
The arthropathy from JPsA is occasionally destructive. As with JAS, complications of medical therapies such as gastritis associated with NSAIDs, infections associated with immunosuppressive medications, and bone disease related to corticosteroids may occur.

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Inflammatory bowel disease-associated arthropathy

DEFINITION
The peripheral or axial arthropathy associated with IBD such as ulcerative colitis or Crohn disease may be called IBD-associated arthropathy.

EPIDEMIOLOGY AND ETIOLOGY
Up to one-quarter of patients with IBD will have onset of disease before the age of 20 years (Burgos-Vargas et al. 1997). Joint disease, primarily arthralgias, has been reported in up to 20% of children with IBD (Lindsley & Laxer 2005). The etiology of IBD-associated arthropathy is unknown. It is not generally associated with HLA-B27 unless there is axial involvement.

CLINICAL HISTORY
While there are significant clinical differences between the common forms of IBD, similar features include abdominal pain, diarrhea, weight loss, and fever. Endoscopic examination may show a range of findings from shallow ulceration to ‘cobblestoning’ and obstruction, as seen in these endoscopic images from three children with Crohn disease of varying severity (69–71).

---

Box 4 Overview of the treatment of peripheral arthritis in JSpA

- Maintain and improve ROM
- Longitudinal assessments
- NSAIDs are treatment mainstay
- Intra-articular injections
- DMARDS (sulfasalazine, methotrexate) as with JRA/JIA
- TNF blockers as with JRA/JIA
Mild Crohn disease. Note the aphthous changes of the intestinal mucosa of the small intestine of this 14-year-old adolescent referred for intermittent loose stools and infrequent, mild abdominal pain (courtesy of W. A. Faubion, MD, Mayo Clinic) (69). Moderate Crohn disease. Note the erythematous borders and large areas of mucosal ulceration of the small bowel in this 12-year-old girl referred for diarrhea, abdominal pain, weight loss, and an abnormal small bowel radiograph (courtesy of W. A. Faubion, MD, Mayo Clinic) (70a & b). Severe Crohn disease. Cobblestoning and stricture formation is apparent in the small bowel of this 12-year-old girl referred for diarrhea, abdominal pain, growth retardation with anemia, and hypoalbuminemia (courtesy of W. A. Faubion, MD, Mayo Clinic) (71).
Painful skin lesions may be seen with IBD. One of these is erythema nodosa (EN). It is usually painful, subcutaneous, and frequently on the extensor surfaces of the legs (72). Another painful ulcerative lesion occasionally seen with IBD is pyoderma gangrenosum (73). Oral mucositis, which is painful, may precede the GI manifestations of IBD; a severe case of mucositis associated with Crohn disease is shown (74). Mucositis may be seen with other forms of JSpA.

**PHYSICAL EXAMINATION**

IBD-related arthropathy is primarily in the lower extremities (75), and is generally not as destructive as JRA (Burgos-Vargas et al. 1997). The axial disease seen in adults is very uncommon in children (Burgos-Vargas et al. 1997).

**LABORATORY**

Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflamma-
Spondyloarthropathies

Inflammatory process with elevations in the WBC, platelet count, ESR, and CRP along with evidence of the anemia of chronic disease. Evidence of occult blood is frequently found in stool samples. Autoantibodies are uncommon.

**IMAGING**

Although clinical features and endoscopic evaluations are the mainstay of diagnosing IBD, imaging can assist the diagnostic process. In addition to excluding other reasons for the symptoms of enteritis, abdominal imaging such as CT scanning can show inflammatory findings such as bowel wall thickening (76a, b).

Since most patients with IBD-associated arthropathy have peripheral joint involvement, they share imaging features with the other forms of JSpA.

**DIFFERENTIAL DIAGNOSIS**

Since IBD-associated arthropathy occurs with or after the diagnosis of IBD, the primary differential diagnosis of a peripheral arthritis would include infectious (septic) arthritis or post-infectious (reactive) arthritis. The differential diagnosis of axial IBD-associated arthropathy would include the other forms of JSpA.

**PROGNOSIS**

Like most of the other forms of JSpA, the risk for erosive, destructive arthritis in IBD-associated arthropathy is low compared with JRA or JIA. Few data are available on the long-term functional outcomes of children with IBD-associated arthropathy.

**MANAGEMENT**

The principles of medical management of the peripheral musculoskeletal manifestations of JPsA such as the arthropathy associated with IBD are highlighted (see box 4). One of the challenges of managing these patients is that many are not candidates for NSAIDs because of their potential for adverse actions to the GI system.

**COMPLICATIONS**

The arthropathy in IBD-associated arthropathy is usually not destructive. Patients may rarely develop sequelae of uveitis, such as cataracts. Complications of medical therapies such as infections associated with immunosuppressive medications may occur. Bone disease related to corticosteroids may be enhanced by vitamin and mineral deficiencies related to enteritis. Poorly controlled IBD can increase the risk of bowel perforation, fistulae formation, and growth retardation.
**REACTIVE ARTHROPATHIES**

The hallmarks of reactive arthropathies are its post-infectious nature and its absence of infection in the joints affected (box 5).

**Reactive arthritis**

**DEFINITION**
ReA is a sterile inflammatory arthritis that occurs after an identifiable non-streptococcal infectious exposure.

**EPIDEMIOLOGY AND ETIOLOGY**
Estimates of the relative frequency of ReA in several pediatric rheumatology series ranges from around 10% to 40%, based on the criteria used (Burgos-Varga & Vazquez-Mellado 2005). The precise mechanisms of how the infectious agents in ReA initiate a remote, inflammatory response are not known, but depend on both host and pathogen characteristics.

**CLINICAL HISTORY**
ReA usually follows a pathogenic enteric bacterial infection or a lower UTI with *Chlamydia*. Post-enteric ReA tends to occur in younger children and the post-UTI form is seen more in adolescents and young adults. The pattern of joint involvement (lower extremity, oligoarthritis, asymmetric, and so on) is similar to the other forms of JSpA. Enthesitis may also be present. The involved joints are very painful.

**PHYSICAL EXAMINATION**
An oligoarthritis with acutely swollen, tender lower extremity joints (see 75) would be characteristic of ReA. Lower extremity enthesitis or dactylitis may also be seen. Findings of inflammatory eye disease, such as uveitis, may also be seen. The classic form of ReA is characterized by urethritis, conjunctivitis, and a painful oligoarthritis following a lower UTI.

Two characteristic skin findings may be seen in older children with ReA. In males, circinate balanitis may be seen in a few cases of ReA (77). Keratoderma blenorrhagicum is a pustular rash, on the soles of feet and may also be seen in an occasional patient with ReA (78).

**LABORATORY**
Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP along with evidence of the anemia of chronic disease. The presence of HLA-B27 increases the odds ratio of a diagnosis of ReA, particularly if there is axial involvement.

Since the time between the exposure and the development of arthritis could be several weeks, or the exposure may have generated minimal symptoms, a very high degree of suspicion is needed to look for potential GI or GU pathogens that could be provocative. Stool culture and urine polymerase chain reaction for evidence of previous chlamydia infection may be needed.

**IMAGING**
Since most patients with ReA have peripheral joint involvement, they share imaging features with the other forms of JSpA.

**DIFFERENTIAL DIAGNOSIS**
The primary concern is to be sure that the child has a post-infectious ReA and not an infectious form of arthritis, such as septic arthritis. Additionally the arthropathy of ReA shares many features of the other forms of JSpA.

**PROGNOSIS**
Like most of the other forms of JSpA, the risk for erosive destructive arthritis in ReA is low compared with JRA or JIA. Few data are available on the functional outcomes of children with ReA.

**MANAGEMENT**
The management principles and frequently used medications for ReA are shown (box 6). The use of antibiotics (beyond treating the initial, precipitating infection) to treat the arthritis of ReA is not felt to be helpful (Kvien et al. 2004).

**COMPLICATIONS**
The arthropathy in ReA is usually not destructive. Patients may also develop sequelae of uveitis, such as cataracts. Like JPsA, complications of medical therapies such as gastritis associated with NSAIDs, infections associated with immunosuppressive medications, and bone disease related to corticosteroids may occur.
Box 5 Keys to recognizing juvenile reactive arthropathies including ReA, ARF, and PSRA

- Acute, very painful, sterile arthritis
- After an identifiable exposure
- Typical exposures include enteropathic bacteria and Chlamydia
- Post-streptococcal conditions like rheumatic fever and PSRA are also reactive arthropathies

Box 6 Overview of the treatment of ReA

- Identify the exposure, treat as indicated
- Exclude septic arthritis
- NSAIDs are the treatment mainstay
- Longitudinal assessments
- Intra-articular injections
- DMARDs (sulfasalazine, methotrexate) rarely

77, 78  Circinate balanitis. This rash, which is usually painless, may be seen in a subset of males with ReA (courtesy of Dawn M. R. Davis, MD, Mayo Clinic). Keratoderma blenorhagicum. This painless, pustular rash on the soles of the feet may be seen in Reiter syndrome, a form of ReA (courtesy of Dawn M. R. Davis, MD, Mayo Clinic).
**POST-STREPTOCOCCAL ARTHROPATHIES**

**Acute rheumatic fever**

**DEFINITION**
Acute rheumatic fever (ARF) is defined by the Jones criteria established in 1944 (Jones 1944), which are characterized by major and minor criteria (Table 10) (Dajani et al. 1992).

**EPIDEMIOLOGY AND ETIOLOGY**
Although estimates of the incidence of ARF in developing countries may be as high as 100 cases per 100,000 per year, the incidence rate in the US is 0.5–3 cases per 100,000 children per year (Ayoub & Alsaeid 2005). The precise mechanisms of how the infectious agents in ARF initiate a remote, inflammatory response are not known, but depend on both host and pathogen characteristics.

**CLINICAL HISTORY**
ARF follows a group A β-hemolytic streptococcal (GABHS) upper respiratory infection (URI). This form of reactive arthropathy tends to have a migratory pattern, with primarily large joint involvement. In both ARF and ReA, the joint pain is generally out of proportion to physical findings.

**PHYSICAL EXAMINATION**
In adults the joint disease may be the primary clinical feature; in children cardiac involvement is more common. Insufficiency murmurs may be noted. The rash of erythema marginatum (79) is uncommon as are subcutaneous nodules. A late finding of rheumatic fever is chorea, which is also very uncommon. It may present months after the initial GABHS URI, and be a challenge to link to ARF.

**LABORATORY**
Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP and evidence of the anemia of chronic disease.

Documenting the recent URI with GABHS can be done with microbiologic techniques such as ‘rapid-strep’ screens or throat cultures. Alternatively, serologic studies may be preferred. Elevated serum titers of anti-streptococcal antibodies such as ASO or DNase B document a recent GABHS exposure. Unlike JSpA and ReA, there is no association with HLA-B27.

An ECG is done to document cardiac conduction abnormalities or other markers of structural cardiac damage.

**IMAGING**
Like the other forms of reactive arthropathies, the risk of joint damage is very low, and imaging affected joints may show evidence of soft tissue swelling, but no joint space narrowing or erosion.

Imaging of the heart with echocardiography is very important in assessing potential cardiac disease in ARF. While myocardial and pericardial disease may be present, the primary purpose of echocardiography is to assess for potential valvular abnormalities.

---

**Table 10**
Modified Jones criteria for the diagnosis of acute rheumatic fever (Dajani et al. 1992)
The diagnosis of ARF is made when at least two major or two minor and one major criteria are present following a group A, β-hemolytic streptococcal URI.

<table>
<thead>
<tr>
<th>Major criteria</th>
<th>Minor criteria</th>
<th>Previous strep URI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carditis</td>
<td>Fever</td>
<td>Increased Strep Ab titers</td>
</tr>
<tr>
<td>Arthritis culture</td>
<td>Arthralgia</td>
<td>(+) throat</td>
</tr>
<tr>
<td>Sydenham chorea</td>
<td>Increased ESR</td>
<td>(+) rapid strep screen</td>
</tr>
<tr>
<td>Erythema marginatum</td>
<td>Increased CRP</td>
<td></td>
</tr>
<tr>
<td>Subcutaneous skin nodules</td>
<td>Prolonged PR interval</td>
<td></td>
</tr>
<tr>
<td>Strep Ab, antistreptococcal antibodies such as ASO and anti-deoxynuclease B (DNase B).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
DIFFERENTIAL DIAGNOSIS
The primary diagnosis in the differential is post-streptococcal reactive arthritis (PSRA) if the arthropathy is post-streptococcal. Higher levels of acute-phase reactants and more treatment-resistant arthropathy favor the diagnosis of ARF (Barash et al. 2008). Also, the arthropathy of PSRA is not migratory like ARF. Other forms of ReA may have similar clinical features.

PROGNOSIS
Like most of the other forms of reactive arthropathies, the risk for erosive destructive arthritis in ARF is extremely low compared with JRA or JIA. The presence of valvular heart disease is the primary prognostic marker. Because the valvular disease can progress with repeated GABHS exposures, antibiotic prophylaxis is indicated (see below).

MANAGEMENT
Penicillin is the primary treatment for the acute URI and skin infections with GABHS responsible for ARF and PSRA. The American Heart Association (AHA) has guidelines for antibiotic prophylaxis for the prevention of progression of valvular heart disease in patients with rheumatic fever (Dajani et al. 1995).

The management of the arthropathy of post-streptococcal arthropathies, including ARF, is highlighted (box 7).

COMPLICATIONS
The arthropathy in ARF is not destructive. Complications of NSAID use include gastritis and renal insufficiency. The primary potential extra-articular complication is valvular heart disease.

Box 7 Overview of the treatment of ARF and PSRA
- Identify GABHS URI and treat if indicated
- Assess for extra-articular involvement
- NSAIDs are the treatment mainstay
- Extensive antibiotic prophylaxis if ARF, some antibiotic prophylaxis if PSRA

79 Erythema marginatum. This 7-year-old boy had a widespread rash, including involvement of his face, following a recent streptococcal URI (courtesy of Lisa Drage, MD, Mayo Clinic).
Post-streptococcal reactive arthritis

DEFINITION
Post-streptococcal reactive arthritis is a reactive arthropathy that follows a streptococcal URI, but doesn’t meet the criteria for ARF. Criteria for the diagnosis for PSRA (Ayoub & Ahmed 1997) are shown (Table I).

EPIDEMIOLOGY AND ETIOLOGY
The incidence of PSRA may be 1–2 cases per 100,000 children per year, and was about twice as frequent as ARF in a US study (Ayoub & Alsaeid 2005). The precise mechanisms of how the infectious agents in PSRA initiate a remote, inflammatory response are not known, but are similar to ARF.

CLINICAL HISTORY
While similar to ARF as a post-streptococcal reactive arthropathy, PSRA differs from ARF in that the joint involvement is not migratory, and is more persistent in PSRA. Since patients with PSRA cannot have other features of ARF, heart disease, chorea, and nodules are not seen at presentation.

PHYSICAL EXAMINATION
Some will have fever and most have arthritis in larger, lower extremity joints.

LABORATORY
Routine laboratory studies such as CBC, ESR, and CRP may show evidence of an inflammatory process with elevations in the WBC, platelet count, ESR, and CRP, and evidence of the anemia of chronic disease.

IMAGING
Like the other forms of reactive arthropathies, the risk of joint damage is very low, and imaging affected joints may show evidence of soft tissue swelling, but no joint space narrowing or erosion.

DIFFERENTIAL DIAGNOSIS
The primary diagnosis in the differential is rheumatic fever if the arthropathy is post-streptococcal. As noted previously, higher levels of acute phase reactants and more treatment-resistant arthropathy favor the diagnosis of ARF (Barash et al. 2008). Since the arthropathy of PSRA is not migratory like ARF, it could be confused with polyarticular forms of JRA/JIA.

PROGNOSIS
Like rheumatic fever, the risk for erosive destructive arthritis in PSRA is extremely low compared with JRA or JIA. Cases of carditis have been described following the diagnosis of PSRA, thus antibiotic prophylaxis is indicated (see below).

MANAGEMENT
Penicillin is also the primary treatment for the acute URI with GABHS responsible for PSRA.

Documenting the recent URI with GABHS can be done with microbiologic techniques such as ‘rapid-strep’ screens or throat cultures. Alternatively, serologic studies may be preferred. Elevated serum titers of anti-streptococcal antibodies such as ASO or DNase B document a recent GABHS exposure. Unlike JSpA and ReA, there is no association with HLA-B27.
The duration of antibiotic prophylaxis in patients who have had PSRA is unclear, but one reference suggests prophylaxis for 1 year after diagnosis (Dajani et al., 1995).

The management of the arthropathy of post-streptococcal arthropathies, including PSRA, is highlighted (box 7).

**COMPLICATIONS**
The arthropathy in PSRA is not destructive. Complications of NSAID use include gastritis and renal insufficiency. PSRA is not associated with cardiac involvement.

### Table 11

**Proposed criteria for the diagnosis of post-streptococcal reactive arthritis (Ayoub & Ahmed 1997)**

The diagnosis of PSRA may be made when an arthritis following a group A β-hemolytic streptococcal (GABHS) URI occurs with these features and doesn’t meet the criteria for acute rheumatic fever (ARF).

<table>
<thead>
<tr>
<th>Characteristics of arthritis</th>
<th>Evidence of GABHS URI</th>
<th>Exclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute-onset, non-migratory</td>
<td>Rapid strep screen</td>
<td>Any major criteria for ARF</td>
</tr>
<tr>
<td>Persistent or recurrent</td>
<td>Throat culture</td>
<td>Meets criteria for ARF</td>
</tr>
<tr>
<td>Suboptimal response to aspirin or NSAIDs</td>
<td>Strep Ab</td>
<td></td>
</tr>
<tr>
<td>Strep Ab, antistreptococcal antibodies such as ASO and antideoxynuclease B (DNase B).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Post-streptococcal arthropathies
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Lupus erythematosus
Systemic lupus erythematosus

DEFINITION
Systemic lupus erythematosus (SLE) is a heterogeneous, inflammatory, autoimmune disorder affecting multiple organ systems, which causes significant morbidity and mortality. Although its etiology, pathogenesis, clinical findings, and laboratory abnormalities are similar to adult-onset SLE, children affected with lupus have special considerations regarding physical and emotional growth and development, treatment toxicities, and lifelong burden of chronic disease. Childhood-onset SLE (onset <16 years of age) makes up approximately 15–20% of all SLE cases, and pediatric lupus is more severe than adult-onset SLE, having a more aggressive presentation and course, including more frequent renal involvement (Barron et al. 1993, Carreno et al. 1999, Tucker et al. 1995).

EPIDEMIOLOGY AND ETIOLOGY
The incidence and prevalence of SLE vary with geographic location and race/ethnicity. The estimated incidence of pediatric SLE is 0.6 per 100,000 (Fessel 1988). Lupus occurs throughout the world, and may be more frequent in non-white populations. For example, Asians, African-Americans, and Hispanics have a higher incidence of SLE compared with whites (Rus & Hochberg 2002). These racial/ethnic groups may also have a more severe disease course. SLE, like many autoimmune diseases, is more common in females than males.

The etiology of SLE is complex and incompletely understood. In a genetically susceptible individual, there is loss of normal homeostatic control of immunologic reactivity to antigens. SLE is an autoimmune disease characterized by production of antibodies against the patient’s own nuclear and cellular components. The triggers for the formation of these antibodies are still under investigation, but may include environmental factors, such as infections, UV light, and drug exposure. There may be decreased clearance of apoptotic cellular debris, causing self-antigens to be abnormally presented, triggering autoantibody formation and immune complex deposition (80). The immune complexes may be ineffectively cleared, resulting in cytokine release, inappropriate inflammation, and tissue destruction (81). Hormonal factors are also implicated, given the female-to-male predominance.

There is some evidence for a genetic susceptibility, with increased concordance of SLE among identical twins (Deapen et al. 1992). In approximately 10% of families with a lupus diagnosis, another connective tissue disease can be found. This illustrates the heritability of lupus, but the actual susceptibility to SLE is multifactorial and polygenic. Lupus predisposition is associated with selective IgA deficiency, the C4A null allele, and inherited complement deficiencies, such as C2, C1q, C1r, and C1 esterase inhibitor. HLA-DR3 and HLA-DR2 increase the risk of SLE in whites, and HLA-DR2 is associated in African-Americans with SLE (Cassidy & Petty 2005).

Drugs thought to classically induce a lupus-like syndrome include hydralazine, minocycline, and some anticonvulsants. However, many drugs have been implicated with this phenomenon. Often, the clinical symptoms of drug-induced lupus will resolve with discontinuation of the offending trigger. Drug-induced lupus is associated with anti-histone antibodies.
Pathogenesis of SLE.

**APOPTOSIS**
- Death signal
- Receptor ligation
  - ex: TNF, Fas
- Protease (caspase) cascade
- DNA fragmentation
- Chromatin condensation
- Cytoplasmic blebbing
- Apoptotic bodies

**AUTOREACTIVITY**
- Receptor ligation
  - ex: TNF, Fas
- Immune complex formation
- Complement fixation
- Release of inflammatory, vasoactive, and chemotactic mediators
- Disruption of endothelium
- Thickening of BM
- Infiltration of inflammatory cells
- Tissue damage

RBC, red blood cell; BM, basement membrane.
CLINICAL HISTORY
The American College of Rheumatology (ACR) revised diagnostic criteria for SLE in 1982 and again in 1997 (Table 12) (Hochberg 1997, Tan et al. 1982, Weening et al. 2004). In adults, SLE is diagnosed when 4 of the 11 criteria are met. The 1982 criteria have been validated in children with a sensitivity and specificity of 96% (Cassidy & Petty 2005). Patients who do not meet the full diagnostic criteria for SLE are often followed in a pediatric rheumatology clinic because as many as 50% of them will eventually meet the diagnostic criteria (Hallengran et al. 2004).

Constitutional symptoms such as fevers, malaise, and weight loss are common at disease presentation and with flares of disease activity. Musculoskeletal involvement may include arthralgias or arthritis that is usually non-erosive and non-deforming. Osteoporosis and avascular necrosis (82, 83) can also be seen and may be a complication of treatment with chronic corticosteroids. Raynaud phenomenon is a common vascular manifestation and is characterized by vasospasm of the extremities leading to red, white, and/or blue color changes, often triggered by cold or stress.

Table 12
American College of Rheumatology classification criteria for diagnosis of systemic lupus erythematosus

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Malar rash</td>
<td>Fixed erythema, flat or raised, over the malar eminences, tending to spare the nasolabial folds</td>
</tr>
<tr>
<td>2. Discoid rash</td>
<td>Erythematous raised patches with adherent keratotic scaling and follicular plugging; atrophic scarring may occur in older lesions</td>
</tr>
<tr>
<td>3. Photosensitivity</td>
<td>Skin rash as a result of unusual reaction to sunlight, by patient history or physician observation</td>
</tr>
<tr>
<td>4. Oral ulcers</td>
<td>Oral or nasopharyngeal ulceration, usually painless, observed by physician</td>
</tr>
<tr>
<td>5. Arthritis</td>
<td>Non-erosive arthritis involving two or more peripheral joints, characterized by tenderness, swelling, or effusion</td>
</tr>
<tr>
<td>6. Serositis</td>
<td>(a) Pleuritis – convincing history of pleuritic pain or rubbing heard by a physician or evidence of pleural effusion OR (b) Pericarditis – documented by ECG or rub or evidence of pericardial effusion</td>
</tr>
<tr>
<td>7. Renal disorder</td>
<td>(a) Persistent proteinuria greater than 0.5 grams per day or greater than 3+ if quantitation not performed OR (b) Cellular casts – may be red cell, hemoglobin, granular, tubular, or mixed</td>
</tr>
<tr>
<td>8. Neurologic disorder</td>
<td>(a) Seizures – in the absence of offending drugs or known metabolic derangements, e.g. uremia, ketoacidosis, or electrolyte imbalance OR (b) Psychosis – in the absence of offending drugs or known metabolic derangements, e.g. uremia, ketoacidosis, or electrolyte imbalance</td>
</tr>
</tbody>
</table>
9. Hematologic disorder  
(a) Hemolytic anemia – with reticulocytosis OR 
(b) Leukopenia – less than 4,000/mm³ total on two or more occasions OR 
(c) Lymphopenia – less than 1,500/mm³ on two or more occasions OR 
(d) Thrombocytopenia – less than 100,000/mm³ in the absence of offending drugs

10. Immunologic disorder  
(a) Anti-DNA – antibody to native DNA in abnormal titer OR 
(b) Anti-Sm – presence of antibody to Sm nuclear antigen OR 
(c) Positive finding of antiphospholipid antibodies based on: 
   (1) an abnormal serum level of IgG or IgM anticardiolipin antibodies 
   (2) a positive test result for lupus anticoagulant using a standard method, or 
   (3) a false-positive serologic test for syphilis known to be positive for at least 6 months and confirmed by Treponema pallidum immobilization or fluorescent treponemal antibody absorption test

11. ANA  
An abnormal titer of ANA by immunofluorescence or an equivalent assay at any point in time and in the absence of drugs known to be associated with ‘drug-induced lupus’ syndrome

* The proposed classification is based on 11 criteria. For the purpose of identifying patients in clinical studies, a person shall be said to have SLE if any 4 or more of the 11 criteria are present, serially or simultaneously, during any interval of observation.
Cardiac abnormalities in patients with lupus range from pericarditis (84, 85) and myocarditis to Libman–Sacks endocarditis and coronary artery disease. Pericarditis may manifest as recurrent chest pain and shortness of breath. Lupus is an additional independent risk factor for atherosclerosis, and adult lupus patients can have myocardial infarctions earlier than the general population (Deapen et al. 1992, Manzi et al. 1997).

The most common pulmonary lesions in SLE are pleural effusions which may present with pleuritic chest pain or shortness of breath (86). Pulmonary function testing is followed and lupus patients most commonly develop a restrictive pattern. When pulmonary hemorrhage occurs, it can be life threatening. Pulmonary emboli from antibodies against cardiolipin or a lupus anticoagulant can also cause shortness of breath and chest pain (87, 88).

Liver dysfunction in the form of acute autoimmune hepatitis is the most frequent liver pathology related to SLE; however, steatohepatitis is common secondary to steroid use and obesity (89).
SLE with slight enlargement of the cardiac silhouette felt to be related to a pleural effusion.

Diffuse pulmonary emboli in a patient with anticardiolipin antibody (87); CT scan demonstrating pulmonary artery thrombosis (88); CT scan of the liver showing diffuse fatty infiltrate in a patient with lupus on high-dose steroids (89).
Renal involvement is a common complication in both adults and children, although it is more prevalent in pediatric-onset SLE. When present, nephritis in children is often more severe than in adults with SLE. It can show at presentation or develop many years later. Patients with nephritis present with hematuria and proteinuria. Hypertension or edema may be present. Children commonly have a renal biopsy to evaluate the extent of lupus involvement if they present with or develop abnormalities in their urinalysis. Lupus nephritis has been most recently reclassified in 2003 by the International Society of Nephrology/Renal Pathology Society (ISN/RPS) into classes I–VI (Table 13) (Weening et al. 2004). Of these, class IV lupus nephritis, which is diffuse, proliferative glomerulonephritis, is the most common and severe (90–94) (Emre et al. 2001).

<table>
<thead>
<tr>
<th>Class</th>
<th>Description</th>
</tr>
</thead>
</table>
| Class I | Minimal mesangial lupus nephritis  
Normal glomeruli by light microscopy, but mesangial immune deposits by immunofluorescence |
| Class II | Mesangial proliferative lupus nephritis  
Purely mesangial hypercellularity of any degree or mesangial matrix expansion by light microscopy, with mesangial immune deposits. There may be a few isolated subepithelial or subendothelial deposits visible by immunofluorescence or electron microscopy, but not by light microscopy |
| Class III | Focal lupus nephritis  
Active or inactive focal, segmental or global endo- or extracapillary glomerulonephritis involving <50% of all glomeruli, typically with focal subendothelial immune deposits, with or without mesangial alterations. A, active lesions; A/C, active and chronic lesions; C, chronic lesions |
| Class IV | Diffuse lupus nephritis  
Active or inactive diffuse, segmental, or global endo- or extracapillary glomerulonephritis involving ≥50% of all glomeruli, typically with diffuse subendothelial immune deposits, with or without mesangial alterations. This class is divided into diffuse segmental (IV-S) lupus nephritis when ≥50% of the involved glomeruli have segmental lesions, and diffuse global (IV-G) lupus nephritis when ≥50% of the involved glomeruli have global lesions. Segmental is defined as a glomerular lesion that involves less than half of the glomerular tuft. This class includes cases with diffuse wire loop deposits but with little or no glomerular proliferation. A, active lesions; A/C, active and chronic lesions; C, chronic lesions |
| Class V | Membranous lupus nephritis  
Global or segmental subepithelial immune deposits or their morphologic sequelae by light microscopy and immunofluorescence or electron microscopy, with or without mesangial alterations. Class V lupus nephritis may occur in combination with class III or IV in which case both will be diagnosed. Class V lupus nephritis may show advanced sclerosis |
| Class VI | Advanced sclerosing lupus nephritis  
≥90% of glomeruli globally sclerosed without residual activity |
Class IV lupus glomerulonephritis (light microscopy).

Class IV lupus glomerulonephritis (immunofluorescence) (courtesy of Sanjeev Sethi, MD, PhD, Mayo Clinic).

Class II mild mesangial proliferative lupus glomerulonephritis (courtesy of Sanjeev Sethi, MD, PhD, Mayo Clinic).

Lupus crescentic glomerulonephritis (courtesy of Sanjeev Sethi, MD, PhD, Mayo Clinic).

Class IV lupus glomerulonephritis subendothelial deposits (electron microscopy) (courtesy of Sanjeev Sethi, MD, PhD, Mayo Clinic).
Lupus involvement of the central nervous system (CNS) is sometimes difficult to diagnose and the symptoms are very non-specific. Symptoms such as poor concentration, poor school performance, memory loss, depression, headache, and visual or auditory hallucinations should be evaluated in a patient with SLE. More serious symptoms can include seizures, strokes, bleeds, or coma (95–99). Neuropsychiatric testing can be helpful to make the diagnosis and to follow progress. Evaluation, especially for school, must be tailored to the child’s age and

95–99 CNS lupus (courtesy of Ann M. Reed, MD, Mayo Clinic) (95); stroke in a teenager with CNS lupus (96); CNS infarcts with lupus patient and antifactor antibodies to factor 2 (97); CNS infarcts: a child with new-onset SLE and antifactor antibodies (98); subdural bleed in an SLE patient with a midline shift (99).
developmental level. Patients with a chronic disease, such as SLE, are at high risk for depression, which can sometimes be difficult to distinguish from neuropsychiatric lupus.

**PHYSICAL EXAMINATION**

Cutaneous manifestations include malar (butterfly) rash, discoid rash, periungual erythema, mucosal ulcerations, photosensitivity, cutaneous erythema (100), and alopecia. The malar rash (101, 101b) is the most common cutaneous manifestation and is present over the nasal bridge, extends onto the cheeks, and spares the nasolabial folds. It is often symmetric and well demarcated, slightly raised, and with texture. Petechiae (102) and purpura (103) can be related to vasculitis or underlying thrombocytopenia.

![Cutaneous erythema in a patient with new-onset lupus](100)

**100** Cutaneous erythema in a patient with new-onset lupus (courtesy of Ann M. Reed, MD, Mayo Clinic).

![Malar rash](101, 101b)

**101, 101b** Malar rash (courtesy of Ann M. Reed, MD, Mayo Clinic) (101); malar rash (101b).

![Petechiae and purpura](102, 103)

**102, 103** Petechiae from a patient with SLE and vasculitis (courtesy of Dawn Davis, MD, Mayo Clinic) (102); purpura in a child with SLE and vasculitis (courtesy of Dawn Davis, MD, Mayo Clinic) (103).
The classic mucosal involvement of patients with SLE is a shallow, painless ulceration of the hard palate with surrounding erythema. Ulcerations present in the oral or nasal mucosa can be painful, especially when present on the tongue. These ulcerations are not as specific for SLE. Edema may be a sign of renal involvement. Joint swelling, tenderness, and limited ROM are indicators of arthritis in children with SLE. Serositis may be detected on exam by the presence of pleural or pericardial rubs.

LABORATORY
ANA is the most common autoantibody present in SLE patients and is present in 98% of children with SLE (Worrall et al. 1990). ANAs are present in 10–15% of healthy children and can be increased non-specifically with infections, but usually to a low titer. More specific ANAs common in SLE include anti-dsDNA, anti-SSA (anti-Ro), anti-SSB (anti-La), anti-Smith, and anti-histone antibodies. Only the ANA, dsDNA, and Smith antibodies are part of the ACR diagnostic criteria. The dsDNA antibodies can be followed over time, because they correlate with increased active systemic inflammation and lupus nephritis in particular. Ribonuclear protein (RNP) antibodies can be present in patients with SLE in low titers and with mixed connective tissue disease (MCTD) in high titers.

Hematologic abnormalities are frequently seen in SLE patients at presentation and with flares of disease. Anemia in SLE patients can be either hemolytic or anemia of chronic disease. A Coombs test will likely be positive in patients with hemolytic anemia due to IgG complement fixing antibodies against red blood cells (Cassidy & Petty 2005). Pancytopenia can also be present. Patients who develop thrombotic thrombocytopenic purpura require hospitalization and aggressive treatment because this can be fatal.

Coagulation abnormalities in SLE patients can be due to either the presence of anti-phospholipid (APL) antibodies, such as a lupus anticoagulant, or anti-cardiolipin antibodies. APL antibodies and the lupus anticoagulant are measured in a number of ways and must be present on at least two occasions 12 weeks apart for the ACR diagnostic criteria. SLE patients can have thrombosis and bleeding at the same time, making them difficult to treat.

SLE patients with active systemic inflammation often have increased non-specific markers of inflammation such as an elevated ESR. C3 and C4 complements are consumed by the circulating immune complexes and are often low in SLE patients with active disease. C3 and C4 are followed clinically, because decreases in these components of the complement cascade correlate with systemic disease activity.

Urinalysis is done regularly with a random urine protein-to-creatinine ratio to monitor for development of nephritis. Active renal disease is indicated by an increase in the first morning urine protein:creatinine ratio, hematuria, pyuria, or the presence of urinary casts. Renal biopsies are commonly done in children with SLE to characterize the ISN/RPS classification, activity and chronicity of renal involvement (Table 14), which helps guide treatment options.

IMAGING
There is no specific imaging test pathognomonic for lupus; instead, radiological studies can only support or challenge the suspicion of this diagnosis. For example, chest radiographs may show pleural effusion or an enlarged heart consistent with pericardial effusions characteristic of the serositis seen in lupus. Joint radiographs and MR images may show soft-tissue swelling and joint fluid, but the arthritis of lupus is usually non-erosive and easily treated. A kidney ultrasound may be required if there is nephritis, or an abdominal ultrasound if there are liver function test abnormalities. A head MR scan and/or MR angiogram is recommended for concerns of lupus cerebritis. For treatment-related toxicities, bone density scans help to evaluate the bone health of a patient on chronic corticosteroids.

DIFFERENTIAL DIAGNOSIS
The differential diagnosis includes the following: other autoimmune diseases (MCTD, drug-induced lupus, autoimmune hepatitis, Evan syndrome, primary antiphospholipid antibody syndrome, primary Sjögren syndrome, and so on); malignancy; systemic vasculitis; juvenile idiopathic arthritis (systemic); or systemic viral infection (Table 15) (Woo et al. 2007).
Oral ulceration (courtesy of Dawn Davis, MD, Mayo Clinic).

Table 14
Active and chronic glomerular lesions (adapted from Weening *et al.* 2004)

<table>
<thead>
<tr>
<th>Active lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endocapillary hypercellularity with or without leukocyte infiltration and with substantial luminal reduction</td>
</tr>
<tr>
<td>Karyorrhexis</td>
</tr>
<tr>
<td>Fibrinoid necrosis</td>
</tr>
<tr>
<td>Rupture of glomerular basement membrane</td>
</tr>
<tr>
<td>Crescents, cellular or fibrocellular</td>
</tr>
<tr>
<td>Subendothelial deposits identifiable by light microscopy (wireloops)</td>
</tr>
<tr>
<td>Intraluminal immune aggregates (hyaline thrombi)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chronic lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glomerular sclerosis (segmental, global)</td>
</tr>
<tr>
<td>Fibrous adhesions</td>
</tr>
<tr>
<td>Fibrous crescents</td>
</tr>
</tbody>
</table>

Table 15
Differential diagnosis for systemic lupus erythematosus (adapted from Woo *et al.* 2007)

<table>
<thead>
<tr>
<th>Disease category</th>
<th>Common features</th>
<th>Differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other autoimmune diseases (MCTD, drug-induced lupus, autoimmune hepatitis, Evan syndrome, primary antiphospholipid antibody syndrome, primary Sjögren syndrome, and so on)</td>
<td>Fever, cytopenia, fatigue, rash</td>
<td>Lack of specific autoantibodies (dsDNA, Smith), specific features unique to specific diseases (e.g. Gottron papules in juvenile dermatomyositis)</td>
</tr>
<tr>
<td>Malignancy</td>
<td>Fever, cytopenia, fatigue, pain, lymphadenopathy, hepatosplenomegaly</td>
<td>Night pain, bone tenderness, normal complement, no urinary changes</td>
</tr>
<tr>
<td>Systemic vasculitis</td>
<td>Fever, fatigue, rash</td>
<td>Nodules, calf pain, positive ANCA, bruits</td>
</tr>
<tr>
<td>Juvenile idiopathic arthritis (systemic)</td>
<td>Arthritis, fatigue, fever, rash, lymphadenopathy, marked anemia</td>
<td>Lack of specific autoantibodies, normal complements, no major organ dysfunction</td>
</tr>
<tr>
<td>Systemic viral infection</td>
<td>Fever, lymphadenopathy, hepatosplenomegaly, cytopenias</td>
<td>Lack of specific autoantibodies, normal complements</td>
</tr>
</tbody>
</table>
PROGNOSIS
The natural history of lupus is characterized by a widely variable and unpredictable clinical course with periods of flare and quiescence. Left untreated, SLE often results in progressive deterioration with a significant fatality rate. However, survival in pediatric SLE has dramatically improved over the last 20 years, the result of earlier and more accurate diagnosis, recognition of mild forms of disease, and better approaches to treatments (Ravelli et al. 2005).
In terms of long-term survival for all lupus patients in the 2000s, the 5-year survival rate approaches 100% and the 10-year survival rate is close to 90%. However, these numbers reflect all lupus patients and are not specific to lupus patients with a particular organ system disease.
Other important lupus outcome measures include disease activity, cumulative organ damage, and functional health status. Several studies have shown that cumulative organ damage is common in patients with pediatric-onset SLE, with a frequency of damage of 50.5–61% (Ravelli et al. 2005). Most of the damage occurs in the musculoskeletal, renal, and neuropsychiatric systems, although the current damage index does not take into account pediatric-specific issues such as growth retardation, pubertal development, and the regenerative ability of children to recover and reverse some forms of damage. The outcome is worse in patients with proliferative renal disease and/or CNS disease, and both are the leading causes of death in SLE after infectious causes.
Unfortunately, studies have also shown that patients with pediatric-onset SLE have poorer health-related quality of life and lower socio–economic status compared with their healthy peers (Ravelli et al. 2005). Patients with childhood-onset SLE are living longer, but are subsequently dealing with the burden of chronic disease and treatment-associated morbidities, which greatly impact their physical, psychological, and financial wellbeing. Patients with pediatric SLE are entering into adult life challenged with the sequelae of disease activity, permanent organ damage, medication toxicity, and other comorbidities such as recurrent infections, accelerated atherosclerosis, osteoporosis and/or avascular necrosis, hypertension, infertility, and increased risk of malignancies.
MANAGEMENT
There is a wide range of disease severity and it is crucial to tailor specific treatment to the degree of organ system involvement to prevent treatment-related toxicities. Most of the lupus treatments used in children have been extrapolated from adult clinical trial data, which may not be completely applicable. Special considerations need to be made regarding the growth and development of children on chronic, often aggressive, medications. Compliance issues in the adolescent period can lead to severe consequences and poor outcomes. Treatment for pediatric SLE can be roughly divided into mild, moderate, and severe categories. Patients with childhood-onset SLE require treatment with high-dose corticosteroid (Tucker et al. 1995) and other immunosuppressive agents more often than adults (Brunner et al. 2002). Patients with APL antibody syndrome may require prophylaxis with aspirin or other anticoagulation. The management of SLE is outlined in the table (Table 16).
COMPLICATIONS
Since SLE is a multisystem disease, many organs have the potential to be damaged. Decreased cognitive function, seizures, and paralysis are examples of possible neurologic complications. Renal failure and hypertension may be the result of renal involvement. The arthritis is not destructive. Bleeding can occur from severe thrombocytopenia. Opportunistic infections may occur from leukopenia. Complications of medical therapies such as gastritis associated with NSAIDs, infections associated with immunosuppressive medications, and bone disease related to corticosteroids may occur.

Table 16
Management principles for systemic lupus erythematosus

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild lupus</td>
<td>NSAIDs; low-dose corticosteroids (prednisone 0.5 mg/kg per day); antimalarials (hydroxychloroquine); low-dose methotrexate (0.5 mg/kg per week)</td>
</tr>
<tr>
<td>Moderate lupus</td>
<td>NSAIDs; medium-dose corticosteroids (prednisone 1–2 mg/kg per day); antimalarials; low-dose; methotrexate; azathioprine; mycophenolate mofetil</td>
</tr>
<tr>
<td>Severe lupus</td>
<td>High, often pulse-dose corticosteroids (2–3 mg/kg per day or pulse dose 30 mg/kg per day i.v.) in combination with immunosuppressives: cyclophosphamide, azathioprine, methotrexate, cyclosporine, mycophenolate mofetil, or rituximab. In certain situations: plasmapheresis, IVIG, and anticoagulation. Dialysis and renal transplantation in patients with end-stage renal disease</td>
</tr>
</tbody>
</table>
**Mixed connective tissue disease**

**DEFINITION**
Some children present with signs and symptoms characteristic of two or more major rheumatic disorders, including JIA, SLE, juvenile dermatomyositis, and/or systemic sclerosis. These children may be classified as having mixed connective tissue disease (MCTD), one of the least common disorders in pediatric rheumatology. Children with MCTD often have very high titers of antibodies to RNP. Three sets of classification criteria for MCTD are widely used that include a variety of clinical features. Kasukawa criteria are used most frequently in the pediatric setting (Table 17) (Kasukawa 1987).

**EPIDEMIOLOGY AND ETIOLOGY**
MCTD is one of the rarest pediatric rheumatology diseases, with a frequency of 0.3% in the US rheumatology database (Black & Isenberg 1992). Although pediatric presentations make up 23% of all cases of MCTD, one study estimated 0.6% prevalence of all pediatric rheumatologic patients (Mier et al. 2005). Median age at onset is 11 years, ranging from 4 years to 16 years. MCTD is three times more common in girls than in boys (Black & Isenberg 1992). The presence of haplotypes HLA-DR4 and HLA-DR2 is associated with MCTD. However, given that MCTD is a disease with overlapping features of other autoimmune diseases, the following associations have also been reported: HLA-DR2 and HLA-DR3 (associated with SLE); HLA-DR5 (associated with systemic sclerosis); and HLA-DR3 (associated with dermatomyositis) (Smolen & Steiner 1998).

**CLINICAL HISTORY**
Because of the wide range of symptoms encompassed by this disease, there are several variations reported for clinical manifestations: (1) SLE-like features; (2) scleroderma-like features; and (3) myositis-like features (Table 18). Clinical features of MCTD have included the following (Black & Isenberg 1992, Cassidy & Petty 2005, Michels 1997, Mier et al. 2005, Mukerji & Hardin 1993, Smolen & Steiner 1998, Tiddens et al. 1993, Yokota 1993):

- Constitutional – fatigue, weight loss.
- Musculoskeletal – arthralgias are common; myalgias are present most at disease onset.
- Cardiovascular – Raynaud phenomenon is one of the most common initial signs. Pericarditis, vasculitis, and hypertension have all been described.
- Gastrointestinal – dysphagia can occur secondary to abnormal esophageal and bowel motility.
- Neurologic – trigeminal neuropathy is the most frequent neurologic finding (25% in adults). Transverse myelopathy, seizures, and psychosis have also been reported.
- Pulmonary – signs of interstitial disease, pulmonary hypertension and serositis have also been reported.

<table>
<thead>
<tr>
<th>Kasukawa’s criteria for mixed connective tissue disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients must meet all three of the following criteria to be diagnosed with MCTD:</td>
</tr>
<tr>
<td>Raynaud phenomenon or swollen fingers or hands or both</td>
</tr>
<tr>
<td>Anti-RNP antibody positivity</td>
</tr>
<tr>
<td>At least one abnormal finding from two or more of the following categories:</td>
</tr>
<tr>
<td>Signs or symptoms of SLE (polyarthritis, facial rash, serositis, lymphadenopathy, leukopenia, or thrombocytopenia)</td>
</tr>
<tr>
<td>Signs or symptoms of scleroderma (sclerodactyly, pulmonary fibrosis, vital capacity &lt; 80% of normal, carbon monoxide diffusion &lt; 70% of normal, decreased esophageal motility)</td>
</tr>
<tr>
<td>Signs or symptoms of dermatomyositis (muscle weakness, elevated creatinine kinase, EMG abnormalities)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 18 Frequencies of mixed connective tissue disease manifestations at presentation and course</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease manifestation</td>
</tr>
<tr>
<td>-----------------------</td>
</tr>
<tr>
<td>SLE like</td>
</tr>
<tr>
<td>Scleroderma like</td>
</tr>
<tr>
<td>Myositis like</td>
</tr>
<tr>
<td>n = 34 patients</td>
</tr>
</tbody>
</table>
• Renal – glomerulonephritis and renal insufficiency can occur.
• Glandular – Sjögren syndrome reported in three-quarters of children but xerostomia, parotid swelling, and parotitis are present in only 15–20%.

**PHYSICAL EXAMINATION**


• Constitutional – weight loss.
• Musculoskeletal – non-erosive polyarthritis is possible. Inflammatory muscle disease with muscle weakness, mostly at disease onset.
• Dermatologic – sclerodermatous skin changes, hand edema (105), and nailfold capillary abnormalities (106). Digital ulcerations, subcutaneous nodules (107), calcinosis (108), telangiectasias, and photosensitivity can occur. One-third of patients complain of malar rash similar to SLE, and another third of rash similar to juvenile dermatomyositis (JDM), including heliotrope rash and/or Gottron papules.
Cardiovascular – Raynaud phenomenon is one of the most common initial signs. Pericarditis, vasculitis, and hypertension have all been described.

Gastrointestinal – hepatomegaly has also been described.

Neurologic – trigeminal neuropathy is the most frequent neurologic finding (25% in adults). Transverse myelopathy, seizures, and psychosis have also been reported.

Pulmonary – findings of interstitial lung disease and pulmonary hypertension are the two most common and dangerous manifestations of pulmonary involvement in MCTD. Serositis has also been reported.

Hematologic – splenomegaly can occur.

Renal – glomerulonephritis and renal insufficiency can occur.

Glandular – parotid swelling, and parotitis are present in only 15–20%.

LABORATORY

Most children with MCTD have ANAs and antibodies to RNP. Additional serologies that could be obtained include those more specific for particular rheumatologic diseases, such as:

1. dsDNA and anti-Smith antibodies for SLE;
2. ACA and anti-Scl-70 antibodies for scleroderma; and
3. RF for JIA. The arthritis of MCTD is often associated with a positive RF and can be present in two-thirds of children with MCTD. Children with MCTD may have more frequent and severe renal disease than adults, warranting periodic urine analyses. Children with MCTD also may have more hematologic complications such as thrombocytopenia than adults, and screening CBCs should be obtained regularly (Mier et al. 2005).

IMAGING

Plain films may be useful to evaluate the extent of arthritis, although erosive bone disease in MCTD is rare. A chest radiograph may be helpful in evaluating for possible cardiopulmonary disease. Children with MCTD have less pulmonary hypertension than adults, but a baseline cardiac echocardiogram and/or pulmonary function test with DLco may be indicated in a child with concerning symptoms. A high-resolution chest CT scan may be indicated if there is concern for pulmonary fibrosis. An upper GI series with small bowel follow-through or swallow study may be useful in evaluating for esophageal or GI dysmotility or strictures.

DIFFERENTIAL DIAGNOSIS

Given the amount of overlap MCTD has with other rheumatologic diseases, it is important to ensure that the patient does not actually have an unusual presentation of a particular connective tissue disease, such as SLE, juvenile idiopathic arthritis, dermatomyositis, polymyositis, systemic sclerosis, localized scleroderma, or Sjögren disease.

PROGNOSIS

There exists the possibility of clinical evolution of MCTD into a disease that better satisfies another connective tissue disorder, such as lupus, scleroderma, or dermatomyositis; therefore, heightened awareness is needed when treating a patient with MCTD. The serologic profile of patients with MCTD may also be somewhat prognostic. Some studies have noted associations between anti-Smith and renal disease, while other studies have noted associations between anti-RNP and Raynaud phenomenon, myositis, esophageal dysmotility, and pulmonary fibrosis. A meta-analysis reported that remissions occurred in 3–27% of children. Raynaud phenomenon and scleroderma-like skin changes were the most common finding, reported in up to 86% of children (Michels 1997). Another study showed that more than 50% of patients initially diagnosed with MCTD showed features more consistent with progressive systemic sclerosis after 5 years (Mukerji & Hardin 1993). Renal disease was uncommon but was the cause of the only two disease-related deaths in the study. Pulmonary hypertension and CNS disease have been shown to be persistent, possibly relaying worse prognosis (Kasukawa 1987). The annual disease-specific mortality in pediatric MCTD is reported to be 3–4 per 1000 patients (Mier et al. 2005).

MANAGEMENT

MCTD was initially characterized as a disorder with an excellent initial response to glucocorticoid therapy; however, therapy is directed at the specific symptoms of the child, such as arthritis, cutaneous disease, or other organ involvement. Tiddens et al. reported the use of NSAIDs in 78%, corticosteroids in 71%, and hydroxy-
chloroquine in 50% in their series of 14 patients (Tiddens et al. 1993). Other immunosuppressive medications used to treat children with MCTD include methotrexate, mycophenolate mofetil, azathioprine, cyclosporine, cyclophosphamide, etanercept, and infliximab (Cimaz et al. 2003). Calcium channel blockers such as amlodipine and nifedipine and topical nitropaste can be used for complaints of Raynaud phenomenon.

**COMPLICATIONS**

Since MCTD is an ‘overlap’ condition with features of several other connective tissue diseases, many organs have the potential to be damaged. Myocardial involvement can lead to cardiac dysfunction. Pulmonary fibrosis may occur after interstitial lung disease. Digital ischemia may also occur. Opportunistic infections can occur from severe skin changes. The arthritis is usually not destructive. Complications of medical therapies such as gastritis associated with NSAIDs, infections associated with immunosuppressive medications, and bone disease related to corticosteroids may occur.
Neonatal lupus syndrome

DEFINITION
Neonatal lupus syndrome (NLS) is a passively acquired autoimmune disease that occurs in neonates born to mothers who have specific autoantibodies. These autoantibodies pass through the placenta and can cause neonatal organ damage including to the skin, liver, brain, heart, blood vessels, and blood elements.

EPIDEMIOLOGY AND ETIOLOGY
In women with SSA/Ro antibodies, 27% of infants develop hematologic abnormalities, 26% develop liver enzyme elevation, 16% develop cutaneous lesions, and 1.6% develop complete congenital heart block (Cimaz et al. 2003). Half of infants with NLS have mothers who are clinically well.

NLS results from transplacental passage of autoantibodies which target fetal and neonatal tissues for immune destruction. Implicated autoantibodies include anti-SSA, anti-SSB, and anti-RNP antibodies (Buyon et al. 1994). The characteristic pathologic findings reveal inflammatory cell infiltrates and deposition of immunoglobulin, complement, and fibrin in the various organs and tissues.

CLINICAL HISTORY
Since NLS presents in the newborn, most of the relevant history is maternal, such as the presence of a connective tissue disorder such as SLE or Sjögren syndrome. These connective tissue disorders are characterized by the presence of autoantibodies that could cross the placenta perinatally. The presence of bradycardia, cytopenias, or characteristic skin lesions in a newborn also prompts an evaluation for NLS. The fetal bradycardia could be detected in utero prior to delivery.

PHYSICAL EXAMINATION
Each of the following manifestations can occur in isolation or in combination:
- Skin – the classic NLS rash is characterized by round or elliptical, erythematous, papulosquamous lesions with a fine scale and central clearing (111). Annular erythema (112), bullous lesions on the soles and palms, and facial telangiectasias have also been described. Lesions typically involve the face and scalp, but are also found on the neck, trunk, extremities, and intertriginous areas. A raccoon-eye rash is common, whereas the malar rash is rarely seen. The rash may be be present at birth but more commonly develops within the first few months of life. Lesions can develop at any time up to 6 months of age, and can be exacerbated by sunlight or phototherapy. The rash spontaneously resolves by 6–8 months as maternal autoantibodies disappear from the infant’s circulation. Residual dyspigmentation and atrophy are rare.
• Cardiac – the most severe and life-threatening complication is congenital heart block (CHB). CHB occurs in 1 in 14,000 live births and at least 90% of these cases are due to transplacental passage of maternal autoantibodies (Buyon et al. 1998). The incidence of heart block in infants born to mothers who have anti-SSA or anti-SSB antibodies is 1–2%, and the recurrence rate in subsequent pregnancies is 16–18% (Brucato et al. 2002, Buyon et al. 1998, Cimaz et al. 2003). Complete CHB causes significant morbidity and mortality, with 65% of surviving neonates ultimately requiring pacemakers (Buyon et al. 1998). Incomplete heart block can progress to higher degrees of block years after the postnatal period (Askanase et al. 2002). Additional ECG abnormalities have been reported including transient sinus bradycardia, QT interval prolongations, and Wolff–Parkinson–White syndrome (Askanase et al. 2002). Ventricular endocardial fibroelastosis can present prenatally or postnatally in the first year of life.

• Hematologic manifestations – thrombocytopenia and anemia are the most common hematologic manifestations and typically worsen in the first days after birth and resolve after disappearance of maternal autoantibodies. Cases of neutropenia, transient pancytopenia, aplastic anemia, hemolytic anemia, and thrombosis have also been reported.

• Liver – the primary manifestations are transaminitis, cholestasis, and hepatosplenomegaly. The two most common presentations of NLS liver disease are: (1) direct hyperbilirubinemia with mild or no transaminitis occurring in the first few weeks after birth, and (2) mild transaminitis occurring at 2–3 months of age (Lee et al. 2002). Most infants have spontaneous resolution of liver abnormalities by 6 months after clearance of maternal autoantibodies. Severe perinatal hepatic dysfunction, often with the phenotype of neonatal iron storage disease, suggests a poor prognosis and warrants treatment with corticosteroids (Frankovich et al. 2008).

• Central nervous system – cerebral hypomyelination, hydrocephalus, and macrocephaly are the most common reported findings (Boros et al. 2007, Prendiville et al. 2003). Cerebral dysmaturity and vasculopathy affecting the blood vessels that supply the basal ganglia and germinal matrix are also reported (Frankovich et al. 2008). Rarely, infants with NLS can have hypotonia, seizures, developmental delay, and feeding difficulty (Frankovich et al. 2008).

• Other manifestations – rhizomelic chondrodysplasia has been reported (Shanske et al. 2007). Pneumonitis, pulmonary capillaritis, glomerulonephritis, and nephrotic syndrome are rare manifestations of NLS and may be suggestive of infantile primary SLE.

LABORATORY
Evaluation for maternal autoantibodies should be undertaken if any of the previously described manifestations are present in the neonate without a clear cause. The initial screening can be done using maternal serum or cord blood. A complete evaluation for congenital infection should be pursued simultaneously since congenital infections mimic NLS and require prompt antibiotic therapy. The initial maternal antibody screen should include ANA, anti-SSA antibody, and anti-SSB antibody. Blood counts, liver function tests, and thyroid hormone levels should be evaluated in all infants born to mothers with autoimmune diseases and asymptomatic mothers with implicated autoantibodies.

IMAGING
If maternal autoantibodies are found, fetal cardiac echocardiography should be done to evaluate for evidence of heart block.

DIFFERENTIAL DIAGNOSIS
Congenital infections can cause hepatitis, basal ganglia calcifications, and hematologic abnormalities which are similar in presentation to NLS. Neonatal iron storage disease and giant cell hepatitis have similar liver pathology and presentation. Peroxisomal disorders can also present with rhizomelic chondrodysplasia.
PROGNOSIS
Most manifestations resolve spontaneously by 6 months of age after clearance of maternal antibodies. Permanent sequelae can occur when inflammation is extensive (skin, liver, or bone marrow) or when vulnerable tissues are involved (cardiac conduction system and small end-organ blood vessels). Maternal disease activity during pregnancy predicts worse outcomes. High titers of maternal SSA antibodies and the presence of maternal thyroid disease increase the risk of NLS complications. Future development of an autoimmune disease in early childhood warrants long-term follow-up of infants with NLS (Martin et al. 2002).

MANAGEMENT
All pregnant women with autoimmune symptoms should undergo early first-trimester screening of implicated antibodies. Women found to have positive anti-SSA and anti-SSB should be considered at high risk for development of CHB in the fetus. Current recommendations are to monitor the fetal PR interval by echocardiogram weekly between 16 and 26 weeks’ gestation, and biweekly between 26 and 32 weeks. For cardiac involvement, maternal oral dexamethasone therapy and IVIG in utero have been used but efficacy and risks are still questionable. Sympathomimetics are used for severe intrauterine and postnatal bradycardia until a pacemaker can be placed. Corticosteroids are recommended for active carditis postnatally. In cases of severe symptomatic fetal thrombocytopenia and anemia, IVIG and corticosteroids have been used. Corticosteroids are also used for severe or persistent liver involvement.

In most cases, affected neonates present with an isolated manifestation of NLS, although multiorgan dysfunction can occur. The most common multiorgan presentation of NLS is a sick preterm infant who has anemia, thrombocytopenia, liver dysfunction, and respiratory failure. In addition to the standard evaluation for congenital infection, a timely evaluation for NLS, including ECG and maternal autoantibody profiles, may facilitate diagnosis. In addition, maternal thyroid and APL antibodies may be measured and a careful neurologic examination undertaken.

COMPLICATIONS
The primary complication from NLS is permanent damage to the neonatal cardiac conduction system. There is risk for bleeding and infection with the associated cytopenias.

Cutaneous lupus erythematosus

DEFINITION
Cutaneous lupus erythematosus (CLE) is a heterogeneous group of connective tissue diseases represented by skin manifestations. The skin lesions are characterized by the presence or absence of interface dermatitis and the presence or absence of scarring. There are three categories based on Gilliam’s classification system (Gilliam & Sontheimer 1981).

EPIDEMIOLOGY AND ETIOLOGY
CLE is thought to occur two to three times more frequently than SLE with the most common age between 20 and 40 years. There exists a female:male predominance of 3–6:1 depending on the subtype of CLE (Tebbe & Orfanos 1997). In multiple studies of CLE patients, age of onset varied from 8 years to 84 years of age with only around 3% presenting younger than 20 years of age. African–Americans tended to have a higher prevalence of chronic CLE than whites (Tebbe & Orfanos 1997).

The etiology for CLE is thought to be multifactoral, with genetic predisposition, environmental factors, and host immune factors all contributing. Genetic associations have been reported of subacute SCL (SCLE) with C2 and C4 deficiencies, HLA-B8, HLA-DR3, HLA-DRw52, HLA-DQ1, and HLA-DQ2; associations of acute CLE (ACLE) with HLA-DR2 and HLA-DR3; and associations of chronic CLE (CCLE) with C1q, C3, C5 deficiencies (Patel & Werth 2002). Complement deficiencies are thought to cause failure of the clearance of apoptotic cells, which in turn causes increased immunologic stimulation. The cytokines that have been implicated in CCLE include IL-2 and interferon-γ, with CD45RA+ T cells being the prevailing inflammatory cells. It has been reported that 65–70% of CLE patients will have exacerbating or aggravating skin reactions when exposed to UVA and UVB light. This reaction can be delayed for up to months after the exposure. It is hypothesized
that keratinocyte-induced apoptosis by antibody-mediated cellular cytotoxicity reactions causes autoantibody production. Drugs such as hydrochlorothiazide, calcium channel blockers, angiotensin-converting enzyme inhibitors, and anti-TNF antagonists, certain viruses such as Epstein–Barr virus, cytomegalovirus, and human immunodeficiency virus, long-term silica exposure, and smoking have been implicated in CLE induction (Patel & Werth 2002).

CLINICAL HISTORY

- Acute cutaneous lupus erythematosus – photosensitivity is a primary feature. Alopecia can also develop.
- Subacute cutaneous lupus erythematosus – photosensitivity is also prominent.
- Chronic cutaneous lupus erythematosus – as with ACLE, and SCLE, photosensitivity occurs. Unlike ACLE and SCLE, CCLE can result in scarring. CCLE is the most common subtype of CLE. CCLE can be associated with systemic symptoms such as arthritis and arthralgias (15–20%). CCLE can present in a variety of ways: discoid lupus erythematosus (DLE) (localized and generalized), hypertrophic LE, lupus panniculitis, and mucosal DLE. The most common type is DLE (113a, 113b), which has sharply demarcated scaly erythematous papules or plaques that can extend into the follicles. Patients with localized DLE (>50% of CCLE) have lesions that involve only the neck and head as well as fewer manifestations of systemic diseases. Patients with widespread DLE (5–10% of CCLE) called DLE-SLE may progress to systemic disease within 1–3 years. DLE lesions can also develop into squamous cell carcinoma.

PHYSICAL EXAMINATION

- Acute cutaneous lupus erythematosus – non-scarring lesions with an intermittent course that usually involves the face, chest, shoulders, and extensor surfaces of the arm and hand. There can be an associated superficial ulceration of nasal and oral mucosa. The most common localized ACLE rash is the classic malar or butterfly rash. The most generalized ACLE rash is photosensitive lupus dermatitis. Furthermore, diffuse hair thinning, lupus
• Hair, telangiectasias, and proximal nailfold erythema have been reported.

• Subacute cutaneous lupus erythematosus – usually non-scarring lesions with an intermittent course that commonly involves the face and neck (114). It usually presents as a papulosquamous or annular polycyclic eruption. The lesions from SCLE can have a psoriasis-type or pityriasis look to them. In addition, malar eruption, livedo reticularis, periungual telangiectasias, and generalized poikiloderma have been reported.

• Chronic cutaneous lupus erythematosus – more generalized skin lesions that can result in scarring, atrophy, and follicular plugging.

LABORATORY

All lab results associated with SLE can also occur with CLE (more with ACLE and less with CCLE). However, association with SLE serology is neither diagnostic nor exclusionary. The following laboratory results have been reported in CLE: anti-dsDNA antibody (SCLE 5–30%, ACLE 60–80%), anti-SSA antibodies (SCLE 90%), anti-Sm antibodies, elevated ESR, low complements (C3, C4, CH50), hematologic abnormalities (anemia, leukopenia, thrombocytopenia), and hypergammaglobulinemia. The lupus band test, now rarely used, may reveal immunoglobulin and/or complement deposition at the dermal–epidermal junction (SCLE 60%, DLE 90%) (Callen 2006).

IMAGING

None is helpful unless organ system involvement is present.

DIFFERENTIAL DIAGNOSIS

• ACLE – drug eruptions, drug-induced photosensitivity, dermatomyositis, bullous lesions of toxic epidural necrolysis.

• SCLE – urticarial vasculitis and other cutaneous vasculitic syndromes; annular SCLE often needs to be differentiated from erythema annulare centrifugum, and papulosquamous SCLE from lichen planus, psoriasis, and/or dermatomyositis.

• CCLE – papulosquamous diseases and chronic eczema in DLE; non-melanoma skin cancers, keratoacanthomas, and warts.

PROGNOSIS

Overall, it appears that patients with isolated CLE have a better prognosis than patients with SLE; however, severe life-threatening systemic complications can occur in CLE patients. Renal and CNS involvement have been reported (Tebbe & Orfanos 1997). The extent of skin involvement may predict disease severity, with less systemic involvement in patients with skin lesions limited to the face or scalp (Tebbe & Orfanos 1997). Disease activity of CLE can be variable but has been reported to persist up to 41 years (Tebbe & Orfanos 1997). One study showed that CLE patients with signs of nephropathy, arthralgias, and high-titer ANAs (>1:320) should be carefully monitored for transition into SLE (Tebbe & Orfanos 1997). Patients with ACLE have a high risk (up to 100%) of developing SLE whereas patients with SCLE have a much lower risk (about 10%) of developing SLE. With regard to CCLE, patients with localized DLE have a 5% risk of developing SLE while patients with diffuse DLE have about a 20% risk in developing SLE.
MANAGEMENT
The main goals of management are to prevent lesion progression, provide counseling about systemic disease surveillance, and improve physical appearance. Patient education is very important as many factors are known to flare and exacerbate the disease; it consists of avoidance of certain drugs associated with CLE, taking sun- and heat-protective measures, and refraining from smoking. Patients should be aware that there exists the possibility of transition to SLE and they should be appropriately screened with routine blood and urine tests.

Localized CLE can be treated with varying potency topical steroids depending on the location of the lesions, with lower potency for facial lesions and higher potency for the palms and soles. Intrallesional steroids and lasers have been reported to treat localized CLE. Other non-steroidal topical agents used include: retinoids (i.e. tretinoin); immunomodulators (i.e. tacrolimus 0.1%); calcipotriene; and imiquimod. Systemic treatment of CLE includes the use of antimalarials such as hydroxychloroquine as a first-line agent. In more diffuse cases, systemic corticosteroids, antibiotics (i.e. dapsone, cefuroxime axetil, sulfasalazine, and clofazimine), auranofin (oral gold effective in SCLE), thalidomide, and oral phenytoin have been used. Immunosuppressive agents such as azathioprine, methotrexate, mycophenolate mofetil, cyclosporine, and cyclophosphamide have been reported for refractory cases. The use of biologics is limited with the concern that such drugs can lead to the development of drug-induced SLE and SCLE (Callen 2006, Patel & Werth 2002).

COMPLICATIONS
One of the primary complications of the cutaneous forms of lupus is the conversion to SLE. Skin cancers may also develop.
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Idiopathic inflammatory myopathies
The juvenile inflammatory myopathies are systemic, autoimmune inflammatory muscle disorders that occur in children younger than 18 years of age. Three general types commonly seen in pediatric rheumatology are juvenile dermatomyositis, juvenile polymyositis, and postinfectious myositis.

**Juvenile dermatomyositis**

**DEFINITION**

Juvenile dermatomyositis (JDM) is an inflammatory vasculopathy that affects primarily the skin and the skeletal muscles. JDM represents 20% of all inflammatory myositis and is the most common inflammatory myopathy in children. Classic findings are Gottron rash and papules, heliotrope rash, calcinosis cutis, and symmetric, proximal muscle weakness. Diagnosis is made by Bohan and Peter (1975a): a characteristic rash, proximal muscle weakness, elevated muscle enzymes, EMG results of polyphasic, low-amplitude motor unit potentials with insertional irritability and fibrillations, and muscle biopsy with inflammation and/or vasculitis. Skin findings with three other features are necessary to make the diagnosis (Table 19).

**EPIDEMIOLOGY AND ETIOLOGY**

The estimated annual incidence in whites is 3.4 cases per million; there is a higher incidence in Hispanics and African–Americans. Females are affected more often than males. The median age of onset is approximately 7 years with a median delay to diagnosis of 3–4 months.

The etiology is not completely understood. The innate and adaptive immune systems and environmental factors may trigger JDM in a genetically susceptible host. Seasonal clustering of JDM in the spring months suggests that environmental agents play an important role in disease onset. An upper respiratory infection may be seen in 56% of patients with JDM 3 months before onset of symptoms as per data collected by the National Juvenile Dermatomyositis Registry. The relationships to infectious agents associated with development of JDM such as Coxsackie B virus, adenovirus, echovirus, hepatitis B, toxoplasmosis, influenza, parvovirus B19, enterovirus, *Mycobacteria pneumoniae*, and *Streptococcus* spp. are all indirect. Molecular mimicry may play a causative role. However, the evidence is inconclusive as there has not been viral or bacterial DNA found in muscle biopsy specimens. Non-infectious agents such as exposure to UV light and bone marrow transplantation as in a graft-versus-host paradigm are also hypothesized to be factors in JDM (Table 20).

The pathogenesis of JDM involves the innate and adaptive immune system with the humoral and cell-mediated pathways causing vascular and muscle damage. There is increased expression of major histocompatibility complex class I and II. Autoantibodies against endothelial antigens, currently unknown, may produce vascular injury leading to ischemia and muscle damage. An inflammatory lymphocytic infiltrate involving CD4+ T cells, B cells, plasmacytoid dendritic cells, and macrophages is present in the muscles and the skin. These cells are arranged in a perivascular and perifascicular distribution (115). There is perifascicular atrophy of type I and type II fibers. Swelling and occlusion of the capillary lumen with muscle degeneration and regeneration are characteristic findings. Infarcts and necrosis of muscle fibers can occur in the fascicle. Immune complex deposition with membrane attack complexes produces muscle inflammation.

<table>
<thead>
<tr>
<th>Table 19 Clinical criteria for juvenile dermatomyositis</th>
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<tr>
<td><strong>Characteristic skin findings</strong></td>
</tr>
<tr>
<td>Gottron papules</td>
</tr>
<tr>
<td>Erythematous rash</td>
</tr>
<tr>
<td>Periungual erythema and telangiectasis</td>
</tr>
<tr>
<td>Alopecia</td>
</tr>
<tr>
<td>Calcinosis</td>
</tr>
<tr>
<td><strong>Characteristic muscle findings</strong></td>
</tr>
<tr>
<td>Symmetric proximal weakness</td>
</tr>
<tr>
<td>Elevation of muscle enzymes</td>
</tr>
<tr>
<td>Typical EMG pattern</td>
</tr>
<tr>
<td>Vasculitis and/or inflammation on muscle biopsy</td>
</tr>
</tbody>
</table>
Table 20
Environmental causes of myositis

<table>
<thead>
<tr>
<th>Agents</th>
<th>Observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infectious</td>
<td></td>
</tr>
<tr>
<td>Coxsackie virus</td>
<td>Antibodies to Coxsackie B viruses noted at onset of JDM compared with controls few in studies</td>
</tr>
<tr>
<td>Influenza virus</td>
<td>Seen with acute viral myositis children who subsequently developed JDM</td>
</tr>
<tr>
<td>Hepatitis B and C</td>
<td>Myositis development associated after acute illness</td>
</tr>
<tr>
<td>Parvovirus</td>
<td>Myositis development associated after acute illness</td>
</tr>
<tr>
<td>Echovirus</td>
<td>Myositis seen mainly in children with X-linked hypogammaglobulinemia</td>
</tr>
<tr>
<td>Streptococcus spp.</td>
<td>Elevated titers seen in children before onset of myositis</td>
</tr>
<tr>
<td>Mycoplasma pneumoniae</td>
<td>Case reports of myositis after pneumonia</td>
</tr>
<tr>
<td>Toxoplasmosis</td>
<td>Elevated titers noted before disease onset</td>
</tr>
<tr>
<td>Non-infectious</td>
<td></td>
</tr>
<tr>
<td>Drugs</td>
<td>JDM and JPM noted after penicillamine</td>
</tr>
<tr>
<td>Vaccines</td>
<td>Agents include MMR, DPT, BCG, hepatitis B several months after vaccination</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>JDM and JPM reported after transplantation in a graft-versus-host disease</td>
</tr>
<tr>
<td>UV light</td>
<td>JDM seen after excessive sun exposure</td>
</tr>
</tbody>
</table>

Table 21
Clinical presentation of juvenile dermatomyositis

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weakness</td>
<td>&lt;100</td>
</tr>
<tr>
<td>Rash</td>
<td>100</td>
</tr>
<tr>
<td>Muscle pain</td>
<td>72</td>
</tr>
<tr>
<td>Fever</td>
<td>65</td>
</tr>
<tr>
<td>Swallowing difficulty</td>
<td>45</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>37</td>
</tr>
<tr>
<td>Arthritis</td>
<td>36</td>
</tr>
<tr>
<td>Calcinosis</td>
<td>22</td>
</tr>
</tbody>
</table>

CLINICAL HISTORY
Constitutional symptoms may occur 3 months prior to disease onset. Photosensitive rashes can occur coupled with muscle pain and weakness. Thirty percent of children demonstrate a rash at least 3 months prior to muscle weakness; also, 30% of children may experience muscle weakness prior to a rash. Sixty percent of children develop skin and muscle manifestations within 3 months of each other. The child may have difficulty getting out of bed or tire easily in physical education classes or sporting events.

Other symptoms include dysphonia, hoarseness, and melena as a consequence of vasculopathy involving the GI tract. Perforation may occur in up to 5% of patients by occlusion and infarction of the vessels; secondary infection may result as a complication (Table 21).

Perivascular and perifascicular infiltrates seen in JDM.
PHYSICAL EXAMINATION
Skin findings may be prominent and are classic in JDM. The eyelids may be swollen with a heliotrope rash and an overlying scale (116–118, 119). A malar rash in a photosensitive distribution and sparing of the nasolabial folds may occur (120, 121, 122, 123). Also, patches of erythema with poikilodermatous changes may occur in various areas of the face and the anterior and posterior neck (124–128). The ‘shawl sign’ may be present (129).

116–122 Eyelid vasculitis and heliotrope rash with periorbital edema in a child with JDM (116); a heliotrope rash with periorbital edema in a child with JDM (117); facial erythema, scaling and heliotrope rash in a child with JDM (118); patchy, erythematous macular rash of the cheeks and upper eyelid in JDM. The erythematosus rash spares the perioral region (119); patchy, erythematous macular rash of the cheeks and upper eyelid in JDM. The erythemasus rash spares the perioral region (120); patchy, erythematous macular rash of the cheeks and upper eyelid and a heliotrope rash in JDM. The erythematous rash spares the perioral region (121); patchy, erythematous macular rash with atrophic changes of the cheeks and perioral area in JDM. The erythematous rash spares the perioral region (122).
Juvenile dermatomyositis

Patchy, erythematous macular rash of the cheeks and mouth in JDM. The erythematous rash spares the perioral region (123); erythematous rash on cheek in JDM (124); erythematous rash and atrophic changes on the face in JDM (125); erythematous rash with atrophic changes on the neck in JDM (126); erythematous rash and calcinosis on posterior neck in JDM (127); erythematous rash and calcinosis on neck in JDM (128); erythematous rash in a shawl pattern and a V-shaped distribution on the anterior chest, upper arms, and anterior neck in JDM (129).
Gottron papules are typical; they are violaceous papules and plaques over bony prominences as in the MCP, PIP, and DIP joints, knees, and ankles (130, 131, 132a, 132b). Erythematous, violaceous, scaly plaques may occur over the extensor surfaces of the extremities (133–135). Nailfold telangiectasias, periungual erythema, and hypertrophic ragged cuticles may be found with end-row loop capillary loss and bushy loops of capillary dilation and branching (130, 131, 132a). Children with inadequately treated disease may demonstrate persistent nailfold changes that may be reflective of disease activity, but not muscle involvement. Finally, mechanic’s hands may occur with hyperkeratosis and peeling of the skin over the medial and palmar aspects of the digits. Mechanic’s hands can be seen in patients with Jo-1 autoantibody (histidyl-tRNA synthetase) and lung disease. Additional skin findings include panniculitis (136) and gum erythema with bleeding (137).

130–132a, b Gottron papules on the DIP joints with periungual capillary loop enlargement and telangiectasias in JDM (130); Gottron papules on the MCP and DIP joints with periungual capillary loop enlargement and telangiectasias in JDM (131); Gottron papules on the DIP joints, atrophic PIP lesions with periungual capillary loop enlargement and telangiectasias in JDM (132a); Gottron rash and extensor erythema on the knees and shins in JDM (132b).
Gottron papules, erythematous, violaceous papules overlying the elbow joints in a child with JDM.

Gottron papules, erythematous, violaceous papules overlying the elbows in a child with JDM (134); Gottron papules, erythematous, violaceous papules overlying the knees in a child with JDM (135); panniculitis presenting as firm, painful erythematous lesions (136); gum erythema and telangiectasia in JDM (137).
Cutaneous vasculitis (138, 139) is not uncommon in JDM and may be seen with Raynaud phenomenon as well as cutaneous ulcer, if present, which raises concerns for more severe disease (140–143).

Calcinosis has been well described in JDM. The deposits are firm, flesh-colored nodules present over bony prominences, long strips of calcium in the muscle bundles or in the subcutaneous tissue (144, 145, 145a, 146b). The most commonly affected areas are the elbows, knees, and extremities (147, 148). Calcinosis onset may occur within 3 years of diagnosis, and predictors for devel-
144 Cutaneous calcinosis (courtesy of Lisa Rider MD).

145 Cutaneous calcinosis.

146a A girl with erythema and calcinosis on the anterior chest (146a); cutaneous calcinosis (146b); radiograph of liquified and crystallized calcinosis in the foot of a patient with JDM (147); radiograph of the arm of a patient with JDM showing sheet-like calcinosis of the fascia (148).

147
Development include poor clinical response to treatment, insufficient steroid use, and evidence of a severe vasculitis on biopsy.

Lipodystrophy is seen in a subset of JDM, usually those with long-standing and difficult-to-treat disease. Loss of extremity fat (149) and an increase in truncal fat, with elevated tryglycerides, alterations of glucose, and insulin metabolism. Poikiloderma is commonly seen associated with lipodystrophy (150). Complications such as striae, most commonly from corticosteroid use, may develop (151).

Symmetric, proximal muscle weakness is seen in most children, although it may present insidiously. Muscle involvement usually occurs in the deltoids and the quadriceps muscles, or both muscle groups. The child may demonstrate an abnormal Gower maneuver with an inability to rise to a standing position without the use of the arms.

Other organ systems may be commonly involved in JDM – including interstitial lung disease (152–155), GI vasculitis, dysphonia, and dysphagia – and less commonly cardiac, ophthalmologic, and CNS disease.

LABORATORY
The most common laboratory abnormalities seen are an elevation of the levels of muscle enzymes, greater than 90%, such as aspartate aminotransferase, lactate dehydrogenase, creatine kinase, and aldolase. Muscle enzymes may return to normal within a few months of disease onset if the child is treated adequately at times when disease remains active. The ESR may be normal in 30–50% of children. Up to 75% of children may also have a positive ANA, but the extractable nuclear antigens may be negative. The p155/140 kDa doublet protein has been found in 30% of children with JDM and may

149–151 Lipodystrophy with increase in truncal fat (149); poikiloderma, a brown skin discoloration (150); striae and cutaneous erythema (151).
correlate with extensive cutaneous involvement. Finally, approximately 10% of children have myositis-specific antibodies and myositis-associated antibodies such as Jo-1, PL-7 (threonyl-tRNA synthetase), PL-12 (alanyl-tRNA synthetase), anti-signal recognition particle, and anti-MI-2, which define clinical subsets and predictors of disease severity. Autoantibodies to the tRNA synthetases have been associated with the clinical presentation of fever, inflammatory arthritis, interstitial lung disease, Raynaud phenomenon, and mechanic’s hands.

152–155 Interstitial lung disease in JDM including bronchiolitis obliterans organizing pneumonia (152); interstitial lung disease in JDM including ground-glass changes (153, 154, 155).
**IMAGING**
Plain radiography does not generally play a large role in the workup or therapeutic decision management of JDM. However, areas of calcinosis may be detected over the extremities or joints in plain radiographs (156) and cortical thickening of long bones can also be seen (157). MRI with T2-weighted fat suppression and STIR technique is useful in illustrating muscle edema, suggesting muscle inflammation (158). There may be cutaneous, subcutaneous, and fascial inflammation. MRI may help to localize an area for muscle biopsy. Finally, MRI may detect areas of calcinosis in JDM (159). Muscle ultrasound is generally not used in the diagnostic workup, but may reveal increased muscle echogenicity and attenuation. However, muscle ultrasound may be used to localize and monitor calcinosis (160).

**DIFFERENTIAL DIAGNOSIS**
Given the extensive skin findings seen in JDM, the diagnosis may initially be confused with SLE or scleroderma. Other connective tissue disease mimickers include juvenile idiopathic arthritis and juvenile polyarteritis nodosa.
Other childhood muscle diseases that resemble myositis include the muscular dystrophies, Duchenne and Becker, and the metabolic myopathies. Other disease entities to consider are sarcoid, viral myositis, celiac disease, and IBDs (Table 22).

**PROGNOSIS**

Prior to the use of corticosteroids, the prognosis of JDM was poor. Approximately one-third of children died from the disease and another third suffered from severe functional disability. With adequate and early treatment, however, children are now able to live normal lives. Calcinosis cutis has been a major cause of morbidity due to cosmetic disfigurement, contractures, and ulcerations resulting in secondary skin infections.

**MANAGEMENT**

A multidisciplinary approach involving consultations with a pediatric rheumatologist, dermatologist, and a physical and an occupational therapist are essential to reduce long-term morbidity. Assessment scales of strength and physical endurance such as the Childhood Ultrasound of liquid calcinosis.

### Table 22

<table>
<thead>
<tr>
<th>Differential diagnosis of juvenile dermatomyositis</th>
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<tbody>
<tr>
<td><strong>Connective tissue diseases</strong></td>
</tr>
<tr>
<td>Systemic lupus erythematosus</td>
</tr>
<tr>
<td>Scleroderma</td>
</tr>
<tr>
<td>Mixed connective tissue disease</td>
</tr>
<tr>
<td>Juvenile idiopathic arthritis</td>
</tr>
<tr>
<td>Juvenile polyarteritis nodosa</td>
</tr>
<tr>
<td>ANCA + vasculitis</td>
</tr>
<tr>
<td><strong>Muscular dystrophies</strong></td>
</tr>
<tr>
<td>Duchenne</td>
</tr>
<tr>
<td>Becker</td>
</tr>
<tr>
<td><strong>Metabolic myopathies</strong></td>
</tr>
<tr>
<td>McArdle disease</td>
</tr>
<tr>
<td>Pompe disease</td>
</tr>
<tr>
<td><strong>Other diseases</strong></td>
</tr>
<tr>
<td>Sarcoid myopathy</td>
</tr>
<tr>
<td>Postinfectious myositis</td>
</tr>
<tr>
<td>Celiac disease</td>
</tr>
<tr>
<td>Inflammatory bowel disease</td>
</tr>
<tr>
<td>Myasthenia gravis</td>
</tr>
<tr>
<td>Leukemia</td>
</tr>
<tr>
<td>Multicentric reticulohistiocytosis</td>
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</tbody>
</table>
Health Assessment Questionnaire (CHAQ) and the Childhood Myositis Assessment Scale (CMAS) (Table 23) determine the clinical status and predict functional outcomes. Rehabilitation with physical therapy using isometric and isotonic exercises should be first-line management to restore and maintain muscle strength. To treat skin manifestations, sun avoidance and sunscreen that protects against UVA and UVB rays are important. Medical therapy for calcinosis has been controversial due to lack of adequate response; generally diltiazem or bisphosphonates have been used with varying success. Calcium and vitamin D supplementation is thought to be key in bone health.

For treatment of active muscle disease, oral corticosteroids are the backbone of therapy with high doses (1–2 mg/kg per day) initially used to induce remission over 4–6 weeks with a slow taper. Weaning of corticosteroids may take 2 years and when additional immunosuppressive agents are used weaning may at times occur over 1 year. For severe or refractory disease, pulse therapy with intravenous methylprednisolone of 30 mg/kg per day for 3 days should be used. Other immunosuppressive agents are added, at the onset of disease, to the corticosteroid regimen to reduce steroid side effects, disease relapse, and progressive disease. A commonly used agent is methotrexate with folic acid supplementation. Other immunosuppressive agents used include IVIG, cyclosporine, and rituximab. Combination therapy is often used as two agents generally work better to induce a rapid response and this is associated with less steroid toxicity.

**COMPLICATIONS**

Calcinosis cutis, interstitial lung disease, and GI involvement (including perforation) may occur in severe cases. Complications from long-term corticosteroid treatment may occur such as obesity, growth failure, acne, cataracts, hypertension, menstrual irregularities, psychiatric disturbances, avascular necrosis (161), osteoporosis, steroid myopathy, lipodystrophy, and metabolic toxicities such as insulin resistance and hypertriglyceridemia. Opportunistic infections can occur from corticosteroids and other immunosuppressive therapies used to treat JDM.

| Table 23
<table>
<thead>
<tr>
<th>Childhood Myositis Assessment Scale (details in Lovell et al. 1999)</th>
</tr>
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<tbody>
<tr>
<td>1 Head elevation</td>
</tr>
<tr>
<td>2 Leg raise/touch object</td>
</tr>
<tr>
<td>3 Straight-leg lift</td>
</tr>
<tr>
<td>4 Supine to prone</td>
</tr>
<tr>
<td>5 Sit-ups</td>
</tr>
<tr>
<td>6 Supine to sit</td>
</tr>
<tr>
<td>7 Arm raise/straighten</td>
</tr>
<tr>
<td>8 Arm raise/duration</td>
</tr>
<tr>
<td>9 Floor sit</td>
</tr>
<tr>
<td>10 All-fours maneuver</td>
</tr>
<tr>
<td>11 Floor rise</td>
</tr>
<tr>
<td>12 Chair rise</td>
</tr>
<tr>
<td>13 Stool step</td>
</tr>
<tr>
<td>14 Pick up</td>
</tr>
</tbody>
</table>

161 T2-weighted fat-saturated MR scan of a JDM patient on intravenous steroids with avascular necrosis of the femur and tibia.
Juvenile polymyositis

DEFINITION
Juvenile polymyositis (JPM) is less common than JDM. It is a systemic autoimmune disease of the muscles without skin involvement. Diagnosis is based on the Bohan and Peter criteria which involve proximal muscle weakness, elevated muscle enzymes, myopathic findings on EMG, and muscle biopsy with inflammation.

EPIDEMIOLOGY AND ETIOLOGY
JPM is an extremely rare childhood myopathy. Because of its rarity, the incidence of JPM is unknown. As in JDM, girls are affected more than boys with a sex ratio of 2–5:1. The age of onset of JPM is similar to JDM with the highest incidence occurring in ages 5–10 years.

The etiology of JPM is largely unknown. As in JDM, similar environmental triggers and genetic risk factors have been associated with JPM. Infectious agents implicated in the development of JPM include viral such as Coxsackie B and respiratory syncytial virus, and bacterial infections such as Streptococcus spp. Non-infectious associations include drugs such as penicillamine, vaccines such as hepatitis B and BCG, as well as after bone marrow transplantations as in a graft-versus-host disease paradigm.

Pathogenesis of JPM likely involves the innate and adaptive immune system including the cell-mediated pathways causing muscle damage. Cytotoxic CD8+ T cells infiltrate muscle fibers and there is endomysial inflammation.

CLINICAL HISTORY
Children with JPM do not have the characteristic skin rash as seen in JDM. Muscle weakness is the hallmark feature, usually occurring in the deltoids and the quadriceps muscles; sometimes it is severe resulting in hypotonia. Similar to JDM, constitutional symptoms may occur prior to obvious muscle weakness. The child may have difficulty getting out of bed or tire easily in physical education classes or sporting events. A non-erosive arthritis, dysphonia, hoarseness, esophageal dysmotility, and muscle pain are other associated features.

PHYSICAL EXAMINATION
Symmetric, proximal muscle weakness is seen in all children, although distal muscles may be affected as well. Muscle involvement usually occurs in the deltoids and the quadriceps muscles, or both muscle groups. There are no cutaneous or nailfold capillary abnormalities. As with JDM, the child may demonstrate an abnormal Gower maneuver with an inability to rise to a standing position without the use of the arms.

LABORATORY
The most common laboratory abnormalities are an elevation in the levels of muscle enzymes such as aspartate aminotransferase, lactate dehydrogenase, creatine kinase, and aldolase. Muscle enzymes may return to normal within a few months of disease onset if the child is treated adequately.

Similar myositis-associated and myositis-specific autoantibodies have been seen in JPM as in JDM. These autoantibodies include Jo-1, PL-7 (threonyl-tRNA synthetase), PL-12 (alanyl-tRNA synthetase), anti-signal recognition particle, and anti-MI-2, which define clinical subsets and predictors of disease severity. Autoantibodies to the tRNA synthetases have been associated with the clinical presentation of fever, inflammatory arthritis, interstitial lung disease, Raynaud phenomenon, and mechanic’s hands. Autoantibodies to the signal recognition particle have been observed in African–American girls with more aggressive and refractory disease, or cardiac manifestations.

IMAGING
Plain films do not play a large role in the workup or management of JPM. As in JDM, MRI with T2-weighted fat suppression and STIR technique is useful in illustrating muscle edema, suggesting muscle inflammation, and the signal intensity from STIR may correlate with the level of disease activity. With chronic muscle disease, there may be fatty infiltration of the muscle and/or muscle atrophy. As in JDM, muscle ultrasound generally is not used in the diagnostic workup, but may reveal increased muscle echogenicity and attenuation.

DIFFERENTIAL DIAGNOSIS
As with JDM, the childhood muscle diseases that mimic myositis include the muscular dystrophies, Duchenne and Becker, and the metabolic myopathies. Evaluation with muscle biopsy is key to differentiate JPM from other causes of myopathy.
PROGNOSIS
The course may be protracted and refractory to steroids. Complications from steroid treatment such as obesity, growth failure, acne, cataracts, hypertension, menstrual irregularities, psychiatric disturbances, avascular necrosis, osteoporosis, steroid myopathy, lipodystrophy, and metabolic toxicities may occur.

MANAGEMENT
As in JDM, a multidisciplinary approach involving a pediatric rheumatologist and a physical and occupational therapist is needed to reduce long-term morbidity. Assessment scales of strength and physical endurance such as the CHAQ and the CMAS have been developed to determine and predict functional outcomes. Rehabilitation with isometric and isotonic exercises should be the first-line management of childhood myositis to restore and maintain muscle strength.

For treatment of active muscle disease, oral corticosteroids are the backbone of therapy with high doses (1–2 mg/kg per day) initially used to induce remission over 4–6 weeks with a slow taper to avoid relapse. However, JPM is often chronic and may be refractory to steroids. Weaning of corticosteroids may take 2 years or longer. For severe or refractory disease, it is recommended to use pulse therapy with intravenous methylprednisolone of 30 mg/kg per day for 3 days. Other immunosuppressive agents are added to reduce steroid side effects, disease relapse, and progressive disease. As in JDM, a commonly used immunosuppressant is methotrexate with folic acid supplementation. Other immunosuppressive agents used include IVIG, cyclosporine, anti-TNF agents such as etanercept and infliximab, and rituximab. Combination therapy is often used in the treatment of childhood myositis as two agents together generally work better to induce a rapid response.

COMPLICATIONS
Muscle atrophy and interstitial lung disease may occur in severe cases. Complications from long-term corticosteroid treatment may occur such as obesity, growth failure, acne, cataracts, hypertension, menstrual irregularities, psychiatric disturbances, avascular necrosis (see 161), osteoporosis, steroid myopathy, lipodystrophy, and metabolic toxicities such as insulin resistance and hypertriglyceridemia. Opportunistic infections can occur from corticosteroids and other immunosuppressive therapies used to treat JDM.
Postinfectious myositis

DEFINITION
Postinfectious myositis is a transient myositis following an acute infection.

EPIDEMIOLOGY AND ETIOLOGY
Children and adolescents are affected, usually boys more than girls. Viruses associated with myositis include influenza A and B and Coxsackie B viruses. Other infectious causes include toxoplasmosis, *Trichinella spiralis*, and staphylococcal and streptococcal bacteremia. Also, clostridia, *Salmonella* spp., and schistosomiasis are less common causes. Pyomyositis, as from *Staphylococcus* spp., is a skeletal muscle abscess occurring after a local injury.

CLINICAL HISTORY
Coxsackie B virus may cause a pleurodynia with chest wall pain and fever after a headache, nausea, vomiting, and sore throat. Illness lasts 3–5 days. Trichinosis may cause fever, diarrhea, and abdominal pain. Pyomyositis may be associated with a low-grade fever.

PHYSICAL EXAMINATION
Focal tenderness of involved muscles such as the thigh, buttock, and calf muscles is present. Trichinosis may cause periorbital edema and facial swelling. If pyomyositis is present, the affected muscle may be warm and tender.

LABORATORY
Elevated serum muscle enzymes are usually seen such as creatine kinase and aspartate aminotransferase. Myositis associated with schistosomiasis or trichinosis may be associated with marked peripheral blood eosinophilia.

IMAGING
Ultrasound may localize the muscle abscess with pyomyositis.

DIFFERENTIAL DIAGNOSIS
As with JDM and JPM, the childhood muscle diseases that mimic myositis include the muscular dystrophies and the metabolic myopathies. Postinfectious myopathies would more often be seen in the acute setting, associated with the above-mentioned infectious symptoms.

PROGNOSIS
The illness should be self-limited. Proper treatment of the bacterial, parasite, or fungal infection should result in prompt recovery.

MANAGEMENT
Postinfectious myositis is usually self-limited, but glucocorticoids may be used to treat the inflammation. Mebendazole or thiabendazole is used to treat trichinosis.

COMPLICATIONS
There are usually no long-term complications from the muscle involvement.
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Vasculitis involves inflammation of blood vessels. Beyond this straightforward description, however, lies a family of complex and protean disorders. As a result of the ubiquity of blood vessels, the variability of their structure and function, and the multitude of ways in which they may be affected, vasculitis can lead to anything from numbness to pain, thrombosis to bleeding, aneurysms to obstruction.

Large gaps in the understanding of nosology, etiology, and pathogenesis confound the study of vasculitis. This lack of information, in turn, complicates attempts at classification. Primary vasculitides may be classified according to their clinical manifestations, the size of blood vessels involved, the histology of vascular damage, or the presumed disease pathogenesis. This chapter uses a working scheme for classifying pediatric vasculitides, attempting to include current knowledge of disease pathogenesis but leaving open the possibility of reclassification as new data emerge (Table 24).

Kawasaki disease

**DEFINITION**

Kawasaki disease (KD), or mucocutaneous lymph node syndrome, is unusual among the vasculitides in being a self-limited condition, with fever and manifestations of acute inflammation lasting for an average of 12 days without therapy. It is diagnosed by clinical criteria, not histology or angiography.

**EPIDEMIOLOGY AND ETIOLOGY**

KD is almost entirely a disease of children. The average age of patients with KD is about 2 years, 80–90% of cases occur before the fifth birthday, and occurrence beyond late childhood is extremely rare. Boys are affected 50% more frequently than girls. Asians are affected 5–10 times as frequently as whites; blacks and Hispanics have an intermediate risk.

The etiology of KD is unclear, but vascular damage likely plays a role. Blood vessel damage in KD results from an aberrant immune response leading to endothelial cell injury and vessel wall damage. Genetic factors seem to account for the varying susceptibility of different ethnic groups to KD, with polymorphisms of chemokines and TNF receptors and variations in HLA haplotypes all possibly contributing. After many failed attempts to find a single etiologic agent, many researchers now believe that KD represents a final common pathway of immune-mediated vascular inflammation following a variety of inciting infections.

The fever and systemic inflammation of KD reflect elevated levels of proinflammatory cytokines such as TNF, IL-1, and IL-6 that are also thought to mediate the underlying vascular inflammation. Pathologically, the finding of macrophages and IgA-producing plasma cells in vessel walls of children with KD is unique.

**CLINICAL HISTORY**

Guidelines for the diagnosis of KD were established by Tomisaku Kawasaki in 1967 (Table 25). The diagnosis requires the presence of fever lasting 5 days or more without any other explanation, combined with at least four of five manifestations of mucocutaneous inflammation. Children who do not meet the criteria may none-the-less develop coronary artery changes. In an attempt to identify patients with this incomplete form of KD, an
AHA working group has recommended modifications of the criteria.

**PHYSICAL EXAMINATION**

Fever in KD is above 38.5°C and typically hectic and minimally responsive to antipyretic agents. Bilateral non-exudative bulbar conjunctivitis is present in as many as 90% of cases of KD. Children are also frequently photophobic, and many develop anterior uveitis. Mucositis, marked by cracked, red lips and a strawberry tongue (162), is characteristic; discrete oral lesions, such as vesicles or ulcers, as well as tonsillar exudate, suggest a viral or bacterial infection rather than KD. Cutaneous manifestations of KD are polymorphous. The rash typically begins as perineal erythema and desquamation, followed by macular, morbilliform, or targetoid lesions of the trunk and extremities. Vesicular or bullous lesions are rare. Late in the course of the disease, children develop extremity changes, including indurated edema of the dorsum of the hands and feet and diffuse erythema of the palms and soles. The convalescent phase of KD may be characterized by sheet-like desquamation that begins in the periungual region of the hands and feet (163) and by linear nail creases (Beau’s lines). In addition, one-third of children have a self-limited small-joint polyarthritis during the first week of illness. Cervical lymphadenopathy is the least consistent of the cardinal features of KD, absent in as many as 50% of children with the disease. When present, lymphadenopathy tends to involve primarily the anterior cervical nodes overlying the sternocleidomastoid muscle, while diffuse lymphadenopathy or splenomegaly should prompt a search for alternative diagnoses.

**LABORATORY**

Systemic inflammation manifested by elevation of acute-phase reactants (e.g. CRP, ESR, and \( \alpha_1 \)-antitrypsin), leukocytosis, and a left shift in the WBC count is characteristic. By the second week of illness, platelet counts may reach 1,000,000/mm\(^3\) in the most severe cases.
Children with KD often present with a normocytic, normochromic anemia, and urinalysis commonly reveals microscopic pyuria. Intrahepatic congestion often leads to elevated transaminase levels or mild hyperbilirubinemia. CSF typically displays a mononuclear pleocytosis with normal sugar and protein, and arthrocentesis of involved joints demonstrates 50,000–300,000 white cells/mm³, primarily neutrophils.

**IMAGING**

An echocardiogram should be obtained early in the acute phase of illness; up to 20–25% of children develop coronary artery aneurysms within 10 days of the onset of fever if not treated with IVIG. Abdominal ultrasound often shows hydrops of the gallbladder, especially in children with fullness and tenderness of the right upper quadrant. Coronary angiography is rarely necessary as echocardiograms adequately visualize involved vessels in most children.

**DIFFERENTIAL DIAGNOSIS**

KD is most commonly confused with infectious exanthems of childhood. Measles, echovirus, and adenovirus may share many of the signs of mucocutaneous inflammation, but they typically cause less inflammation and lack the extremity changes seen in KD. Toxin-mediated illnesses, especially β-hemolytic streptococcal infection and toxic shock syndrome, lack the ocular and articular involvement typical of KD. Drug reactions such as Stevens–Johnson syndrome or serum sickness may mimic KD but with subtle differences in the ocular and mucosal manifestations. Systemic-onset JRA is marked by prominent rash, fever, and systemic inflammation, and may be difficult to distinguish from KD until persistent polyarthritis becomes evident.

**PROGNOSIS**

Although most small aneurysms fully resolve by echocardiogram, there is a concern that vascular reactivity may not return to normal despite grossly normal appearance. Children should thus be followed indefinitely after KD, and additional cardiac risk factors should be monitored closely. Overall, with modern treatment and follow-up, the prognosis of children with KD is excellent: 25-year follow-up of Japanese children without persistent coronary artery abnormalities demonstrates no increase in morbidity or mortality.

**MANAGEMENT**

Patients who fulfill the criteria for KD are hospitalized and treated with IVIG and aspirin. IVIG therapy within the first 10 days of illness reduces the incidence of coronary artery aneurysms by more than 70%. Fever persists or returns within 48 hours after treatment in 10–15% of patients. These children are at increased risk of developing coronary artery aneurysms, so they are treated with an additional dose of IVIG, one to three daily doses of pulsed methylprednisolone (30 mg/kg), or a single dose of infliximab (5 mg/kg). When symptoms are prolonged beyond 3–4 weeks, consideration should be given to an alternative diagnosis, including chronic vasculitides such as polyarteritis nodosa (PAN).

Patients suspected of having KD but not meeting diagnostic criteria may require further testing such as slit-lamp examination and echocardiography to confirm the diagnosis. At times, children are treated for suspected KD when the diagnosis is uncertain but no clear alternative explanation for the clinical findings can be identified. Markers of increased risk of developing coronary artery aneurysms, including age under 1 year, severe systemic inflammation, or consumptive coagulopathy, may shift the balance toward empiric therapy.

**COMPLICATIONS**

The primary complication is the development of coronary artery disease as noted above. Rarely other cardiac involvement occurs, such as myocarditis. Complications of therapy include risk of bleeding with aspirin and hypersensitivity to IVIG.
Henoch–Schönlein purpura

DEFINITION
Henoch–Schönlein purpura (HSP) is a small-vessel leukocytoclastic vasculitis that presents as the triad of non-thrombocytopenic palpable purpura, arthritis, and colicky abdominal pain. HSP is diagnosed on the basis of clinical findings; vascular imaging and biopsy are generally neither necessary nor diagnostic.

EPIDEMIOLOGY AND ETIOLOGY
HSP is the most common vasculitis in children, occurring in up to 1 child in 5,000 per year. Similar to Kawasaki disease, most cases occur in children under 7 years of age, almost twice as many boys as girls contract the condition, and peak incidence tends to be during the colder winter months.

HSP appears to represent an aberrant immune response to an infectious or antigenic stimulus. Infections, especially upper respiratory tract infections (most commonly streptococcal), precede the onset of illness in at least 50% of children with HSP, but immunizations and insect bites may also trigger the condition. As yet unidentified host and pathogen factors determine a child’s likelihood of developing this vasculitis.

In HSP, the alternate pathway of complement activated by large IgA-containing immune complexes may be important, though precisely how this contributes to the HSP phenotype is poorly understood. Small vessels have a perivascular infiltrate consisting of neutrophils and mononuclear cells, and deposits of IgA, C3, and fibrin may be detected using immunofluorescence and electron microscopy. Renal biopsies typically show an endocapillary proliferative glomerulonephritis involving endothelial and mesangial cells. Proliferation of extracapillary cells also may occur, resulting in variable degrees of crescent formation.

CLINICAL HISTORY
Skin involvement in HSP may begin as urticaria; in most cases it progresses to palpable purpuric lesions primarily involving dependent areas (buttocks and lower legs in ambulatory children, the sacrum, ears and buttocks in infants) (164). Arthritis may affect any joint in an asymmetric and migratory pattern, though lower extremities are more commonly involved. GI involvement ranges from colicky abdominal pain to profuse bleeding, intussusception, and perforation. Renal disease usually is observed more than 10 days after the onset of illness.

PHYSICAL EXAMINATION
Children often appear well and are afebrile despite having prominent purpura. Edema may involve the scrotum (165), dorsa of the hands or feet, and even the scalp. The arthritis of HSP is transient, causing neither chronic joint changes nor permanent sequelae. Signs of the syndrome recur within 6 weeks in up to 40% of children, most commonly abdominal pain and rash. Exacerbations are usually shorter in duration and the symptoms are generally milder than those of the initial disease.
LABORATORY
HSP is a clinical diagnosis; no specific laboratory abnormalities are characteristic. Laboratory tests usually reflect the triggering infection rather than the HSP itself, so investigations are generally limited to ruling out other vasculitides, infections, and hematologic disorders. Children should have a CBC to exclude thrombocytopenia as a cause of palpable purpura, and in some cases measurement of PT, aPTT, and D-dimers to rule out coagulopathies. BUN and creatinine levels and urinalysis are monitored to evaluate renal involvement. In severe cases, renal biopsy is helpful for excluding other causes of acute nephritis, nephrotic syndrome, or renal failure, and for estimating prognosis.

IMAGING
Imaging is generally limited to assessing the cause of abdominal pain in children with known or suspected HSP. Plain radiographs reveal free air in cases of ruptured viscus, and abdominal ultrasound is the examination of choice for determining whether an ileo-ileal intussusception has occurred. Since primarily small vessels are involved, skin or renal biopsies are generally more helpful than vascular imaging if definitive evidence of vasculitis is needed.

DIFFERENTIAL DIAGNOSIS
Clinicians must consider infectious and non-infectious causes of purpura. Sepsis, malignant, and metabolic causes of disseminated intravascular coagulopathy or thrombocytopenia can generally be excluded clinically, based on the relatively slow tempo of disease progression and well appearance of the patient. Exclusion of other vasculitides causing palpable purpura, especially hypersensitivity vasculitis, Wegener granulomatosis, and Churg–Strauss syndrome, might require testing for ANCA, serum complement levels, or even a skin biopsy. Acute hemorrhagic edema of infancy (166), a benign febrile condition of infants, presents with ecchymotic-like macular purpura distinct from that of HSP.

PROGNOSIS
HSP is a self-limited illness that typically resolves within a month in two-thirds of children. Younger patients tend to have milder disease of shorter duration, while adults have a higher incidence of severe renal involvement. Generally there is a correlation between the severity of urinary abnormalities and the chances of developing chronic renal disease, with those demonstrating both nephritic and nephrotic changes at greatest risk. Overall about 1–5% of patients with HSP develop some degree of chronic renal disease.

MANAGEMENT
Therapy of HSP is primarily supportive, aiming for symptomatic relief of arthritis and abdominal pain. Acetaminophen or NSAIDs appear to be effective and safe in most cases. Use of steroids, whether in children who fail to respond to NSAIDs or in those thought to be at highest risk of developing renal compromise, is controversial. Prednisone at a dose of 1–2 mg/kg per day appears to rapidly relieve symptoms in the majority of cases, but caregivers must avoid excessively rapid tapering of the steroids, as this commonly triggers a flare of symptoms. More potent immunosuppressive agents are reserved for children with biopsy-proven crescentic glomerulonephritis or other life-threatening complications such as cerebral or pulmonary hemorrhage.

COMPLICATIONS
Renal insufficiency can occasionally occur. Rarely, intestinal perforation has been reported. Potential complications related to therapy include gastritis with NSAIDs and opportunistic infection with corticosteroids.
Polyarteritis nodosa

DEFINITION
Polyarteritis nodosa (PAN) is a systemic necrotizing vasculitis affecting medium and/or small arteries. The segmental panmural fibrinoid necrosis with nodule and aneurysm formation give this condition its name. PAN is important historically as the first non-infectious cause of vascular damage to be identified, but at least one-third of children have a more limited cutaneous form that is restricted largely to the skin and joints. Diagnosis of PAN usually requires tissue biopsy and/or radiological documentation of vasculitis.

EPIDEMIOLOGY AND ETIOLOGY
Worldwide, PAN is most commonly associated with hepatitis B or C infections. Perhaps as a result of the relative infrequency of these infections in children, pediatric PAN is quite rare, especially in North America. Childhood incidence of PAN peaks at 9–10 years of age, and it may be slightly more common in boys than in girls.

PAN is characterized by transmural necrosis of the walls of medium-sized arteries. Both the patchy, interrupted involvement and the association with hepatitis B suggest an immune-complex-mediated process. The resulting recruitment of polymorphonuclear neutrophils and monocytes, leading to focal areas of necrosis, is typical of an Arthus reaction.

The prevalence of systemic PAN in adults with chronic hepatitis suggests that the development of vasculitis is related to the disordered formation or clearance of immune complexes. Although no clear genetic association has been identified, the fact that up to 1% of patients with familial Mediterranean fever (FMF) develop PAN raises the possibility that inflammasomes play a role.

CLINICAL HISTORY
Cutaneous PAN is usually limited to the skin and musculoskeletal system. It commonly occurs after a sore throat or streptococcal pharyngitis. Livedo reticularis, maculopapular rash, painful skin nodules, edema, and arthritis mostly affect the knees and ankles. Constitutional symptoms are generally mild and systemic involvement is, by definition, rare. Symptoms are more troubling than disabling, and treatment generally consists of NSAIDs and steroids.

Systemic PAN may involve virtually any muscular artery. Consequently, in addition to constitutional symptoms, it may cause a vast array of organ dysfunction. Palpable purpura, livedo, acral blanching (167), necrotic dermal lesions and autoamputation (168), abdominal pain, arthritis/arthralgia, myositis/myalgia, renovascular hypertension, neurologic deficits, pulmonary disease, and coronary arteritis may all be seen at presentation or during the course of the disease.

167 Acral blanching due to PAN of the digital artery.

168 Autoamputation due to PAN of the fingers.
PHYSICAL EXAMINATION
Evidence of cutaneous PAN is generally limited to rashes such as livedo reticularis or nodules (‘nodosa’) overlying medium-sized vessels. Arthritis and fever frequently accompany these findings, but other organ systems are rarely involved. Systemic PAN, on the other hand, may involve any organ system, so evidence of this protein and widespread condition may lead to anything from cranial nerve palsies and myocardial infarction to testicular pain and an acute abdomen.

LABORATORY
Laboratory evaluation usually reflects the ongoing systemic inflammation including anemia, leukocytosis, thrombocytosis, and elevated ESR, CRP, and immunoglobulins. Evidence of myositis (elevated creatine kinase and aldolase) and renal dysfunction (increased BUN and creatinine) are also common findings. Complement levels are normal.

IMAGING
Vessels affected by PAN typically demonstrate the so-called ‘beading’ of arteries, representing alternating narrowing and dilation of high-pressure vessels. CT or MR angiography is generally preferred to conventional angiography because they are less invasive and have less potential for complications (169a–c). Echocardiography may be used to visualize coronary arteries when cardiac involvement is suspected. None-the-less, in some patients, conventional angiography may be necessary to determine the extent and severity of vascular involvement (170).

DIFFERENTIAL DIAGNOSIS
PAN should be considered in the differential diagnosis of any undiagnosed systemic inflammatory condition or in cases of severe, unexplained hypertension. As a result of its pleomorphism, PAN may be confused with other rheumatologic conditions including systemic-onset JRA, Kawasaki disease, and dermatomyositis. Appendicitis or hypercoagulability must be excluded when abdominal pain or vascular insufficiency is the presenting feature, and testicular torsion is a potential confounder when scrotal vessels are involved. Small vessels are spared in classic PAN, so glomerulonephritis typically is not a feature of this condition.

PROGNOSIS
Recent reviews of PAN in children suggest an excellent overall prognosis, with a 4-year mortality rate under 5%. Such an outcome is predicated, however, on close follow-up and aggressive management of active disease or its complications.

MANAGEMENT
In cutaneous PAN, treatment is primarily aimed at controlling symptoms. In systemic disease, on the other hand, vascular inflammation must be controlled in order to prevent compromise of vital vascular beds. In both cases the mainstay of therapy is high-dose steroids, but, since the disease tends to persist or relapse, many patients require steroid-sparing agents for long-term management. These may include methotrexate, IVIG, or TNF inhibitors. Penicillin prophylaxis may prevent disease flares due to recurrent streptococcal infections.

COMPLICATIONS
Renal insufficiency can occasionally occur. Organ ischemia may occur secondary to vascular damage. Potential complications related to therapy include gastritis with NSAIDs, avascular necrosis, and opportunistic infection with corticosteroids. Immunosuppressive medications are also associated with an increased risk of opportunistic infections.
CT angiography showing hepatic artery narrowing (courtesy of Ann M. Reed, MD, Mayo Clinic) (169a); CT enterography showing mesenteric narrowing (courtesy of Ann M. Reed, MD, Mayo Clinic) (169b); CT enterography showing renal infarcts (courtesy of Ann M. Reed, MD, Mayo Clinic) (169c).

Obstruction of the fourth digital artery in PAN.
ANCA-associated vasculitides

DEFINITION
ANCA-associated vasculitides (AAVs) are a group of conditions marked by pauci-immune inflammation of the vessels of the lungs, kidneys, and skin in the presence of anti-neutrophil cytoplasmic antibodies. They include:

- Churg–Strauss syndrome (CSS) – a cause of steroid-dependent asthma with pulmonary infiltrates and systemic inflammation accompanied by ANCA in either a cytoplasmic or a peripheral pattern.
- Wegener granulomatosis (WG) – a necrotizing granulomatous inflammation of small- to medium-sized vessels positive for cytoplasmic staining (c-ANCA) against the neutrophil protease PR-3.
- Microscopic polyangiitis (MPA) – a non-granulomatous necrotizing vasculitis of small vessels characterized by a positive p-ANCA with reactivity to MPO.

EPIDEMIOLOGY AND ETIOLOGY
The total incidence of pediatric AAVs appears to be on the order of one new case per million children per year. The first report of CSS occurring in a pre-teen has been published only in the past decade, so, while these conditions must be included in the differential diagnosis of inflammatory disorders of the kidneys and lungs, most practitioners will not see a case during their careers.

These conditions share antibodies to antigens within the neutrophil cytoplasm and a microscopic pauci-immune polyangiitis marked by little or no complement deposition. They primarily target the respiratory, renal, and cutaneous systems, and they often develop in the wake of a viral infection. The genetic and environmental factors responsible for their development, however, are not known.

171–173 Sinus CT in Wegener granulomatous (171); pseudotumor of the orbit in a PR-3+ child (courtesy of Ann M. Reed, MD, Mayo Clinic) (172); pulmonary hemorrhage in Wegener granulomatous (173).
The pathogenesis of AAVs appears to be related to ANCA. These antibodies most likely stabilize adherence of rolling neutrophils to endothelium, and activate neutrophils and monocytes to undergo an oxidative burst. Activation of phagocytic cells causes upregulation of proinflammatory cytokines (such as TNF-α and IL-8), with resultant localized endothelial cell cytotoxicity.

**CLINICAL HISTORY**

Clinical manifestations of AAVs in children are similar to those in adults. Children with WG almost universally have nasal and sinus involvement and respiratory disease including pulmonary hemorrhage (171–173).

Arthralgias, ocular findings, and skin (174) or renal involvement (175) occur in more than half, while GI disease and CNS involvement are less typical. Unique to pediatric series is a high frequency of subglottic stenosis, noted in almost 50% of children with WG (176). CSS presents initially as allergic rhinitis and asthma. After months or years, peripheral eosinophilia and pulmonary infiltrates develop (177). Only later do manifestations of systemic vasculitis become evident, with weight loss, fever, arthralgia, myalgia, nodular rash, and neuropathy. MPA is most commonly limited to the kidneys, though the presence of constitutional and inflammatory involvement demonstrate the systemic nature of the condition.
PHYSICAL EXAMINATION
Most children with WG present with upper respiratory symptoms such as epistaxis, sinusitis, otitis media, or hearing loss, or with cough, wheezing, or dyspnea indicative of lower respiratory tract involvement. Similarly, children with CSS are recognized to have asthma, but then their symptoms worsen and become increasingly refractory to routine management. Unusually persistent or severe manifestations, or development of hemoptysis or hypertension, suggest evolution beyond routine childhood illnesses.

LABORATORY
While specific ANCA targeted against PR-3 or MPO is positive in the majority of patients with AAVs, this autoantibody may also be found in other more common diseases such as cystic fibrosis and IBD. Accordingly, a positive ANCA should not replace a tissue biopsy in confirming the diagnosis of vasculitis, nor should ANCA screening substitute for a careful history and physical examination. Similarly, while peripheral eosinophilia is characteristic of active CSS, it is more frequently encountered in common allergic conditions, and therefore should be only one of many factors weighed during the diagnostic process.

IMAGING
Chest radiographs may be particularly helpful when a diagnosis of WG or CSS is suspected. Even in asymptomatic children with WG, up to one-third have radiographic abnormalities (173, 178, 179). Similarly, the majority of children with CSS demonstrate pulmonary infiltrates that would be atypical for uncomplicated asthma (180).

DIFFERENTIAL DIAGNOSIS
Biopsy or radiographic confirmation of vascular inflammation is necessary in order to confirm the diagnosis of AAVs, and to exclude a variety of other conditions that may have a non-specific ANCA. Necrotizing granulomatous vascular inflammation in a child with suggestive clinical features is strongly suggestive of WG. In CSS, biopsy of involved tissue shows significant perivascular eosinophilic infiltrates and occasional extravascular granulomas.

PROGNOSIS
Despite progress in the management of WG, the disease continues to cause significant morbidity and mortality due to relapses and treatment-related toxicity. Subglottic stenosis does not respond to systemic therapy, but rather requires surgical dilation and local steroid injections. In patients with limited upper respiratory disease trimethoprim/sulfamethoxazole has been shown to be beneficial, perhaps by suppressing URIs that might activate vascular inflammation.

MANAGEMENT
AAVs, especially WG, are rapidly progressive and may be fatal without effective immunosuppressive therapy. Consequently, potent combination therapies are the mainstay of therapy, including steroids, cyclophosphamide, azathioprine, methotrexate, and more recently mycophenolate mofetil and anti-TNF antibodies (but not the TNF-receptor antagonist etanercept). Most recently, rituximab has been proposed as a potential replacement for both cyclophosphamide and long-term steroids in AAVs.

COMPLICATIONS
Renal insufficiency can occasionally occur. Large airway disease may lead to subglottic stenosis. Organ ischemia may occur secondary to vascular damage. Nasal septal perforation has been reported. Potential complications related to therapy include gastritis with NSAIDs, avascular necrosis, and opportunistic infection with corticosteroids. Immunosuppressive medications are also associated with increased risk of opportunistic infections.
177–179 Eosinophilic pulmonary vasculitis in Churg–Strauss syndrome (courtesy of Sarah Vargas, MD and Debra Boyer, MD) (177); Wegener granulomatous cavitary lesion (courtesy of Ann M. Reed, MD, Mayo Clinic) (178); Wegener granulomatous cavitary lesion (courtesy of Ann M. Reed, MD, Mayo Clinic) (179).

180 Chest CT scan in Churg–Strauss syndrome.
Takayasu arteritis

DEFINITION
Takayasu arteritis (TA), or ‘pulseless disease,’ is a large-vessel vasculitis that involves the aorta and its major branches. Although reports of patients losing their peripheral pulses have appeared for hundreds of years, TA is named after a Japanese ophthalmologist who first described, in 1908, the changes in retinal vessels due to increased outflow into unobstructed cerebral vessels.

EPIDEMIOLOGY AND ETIOLOGY
TA is the third most common form of childhood vasculitis. It most commonly develops during the third decade of life, but the disease has been reported in infancy. TA is more common in the Far East and West Africa than in Europe and North America. In a recent review of childhood TA the mean age of onset was 11.4 years and two-thirds of patients were female.

The cause of TA remains unknown, though histopathology and immunohistochemistry of biopsy and autopsy samples from adults with TA suggest a primarily T-cell-mediated mechanism. Certain HLA associations have been found in Japan, but these have not been confirmed in other populations.

TA lesions consist of granulomatous changes progressing from the vascular adventitia to the media, indistinguishable from those seen in giant cell and temporal arteritis. This histopathology, and the correlations between levels of IL-6 and RANTES and disease activity, suggest that T cells and macrophages play a central role in the disease.

CLINICAL HISTORY
TA is diagnosed on the basis of the distribution of involvement – primarily the aorta and its branches – and the young age of patients – typically under 40 years of age. The distribution of vessel involvement in children parallels that of adults, with diffuse aortic involvement predominating. Signs and symptoms include hypertension, cardiomegaly, elevated ESR, fever, fatigue, palpitations, vomiting, nodules, abdominal pain, arthralgia, claudication, weight loss, and chest pain.

PHYSICAL EXAMINATION
This is particularly helpful in light of the need to completely suppress the vasculitis in order to prevent disease progression. Laboratory markers may be entirely normal despite ongoing inflammation, so MRI offers a potentially more sensitive test for residual disease.

LABORATORY
During the early phases of the disease, markers of systemic inflammation correspond with active vasculitis. Later in the process, vascular lesions may progress in the absence of elevated ESR, CRP, or platelets, significantly complicating treatment aimed at fully controlling disease activity.

IMAGING
Angiography has been the standard method used for diagnosing TA. The size of the vessels involved and the spotty nature of the vascular inflammation make biopsies impractical. CT and MR angiograms have supplanted traditional angiograms as the modalities of choice (181–185). They are less invasive, and MRI offers the opportunity to monitor disease activity by revealing evidence of ongoing vessel wall inflammation. PET scanning may similarly allow simultaneous visualization of disease anatomy and physiology.

DIFFERENTIAL DIAGNOSIS
Different conditions must be considered depending on the presenting complaints of a patient with TA. Fever and systemic inflammation or other non-specific presentations of TA have an enormous differential diagnosis. Loss of peripheral pulses, on the other hand, generates a more restricted list, especially in children. A thorough and thoughtful history and physical examination is the most important element in making the accurate diagnosis.

PROGNOSIS
As with all vasculitides, early diagnosis and aggressive therapy are important in TA in order to prevent irreversible vessel damage with resulting compromise of vital organs. The delay in diagnosis in children is notably longer than that reported in most adult patient series (19 months in a recent report), possibly explaining the 33% mortality rate, significantly higher than that in adult series.

MANAGEMENT
Steroids as well as the typical immunosuppressive agents used in other vasculitides (including cyclophosphamide, methotrexate, and azathioprine) have shown variable efficacy in TA. A recent report in adults with TA from the Cleveland Clinic documented a high response rate to use of TNF inhibitors. Before starting these, however, it is important to test patients for tuberculosis, since aortitis is associated with mycobacterial infections, especially in less developed countries.
COMPLICATIONS

Organ ischemia may occur secondary to vascular damage. Hypertension may develop for many reasons. Potential complications related to therapy include avascular necrosis and opportunistic infection with corticosteroids. Immunosuppressive medications are also associated with increased risk of opportunistic infections.

181 Great vessel MRI in Takayasu arteritis.

182 Loss of blood flow in the femoral arteries and development of collateral blood flow in Takayasu arteritis.

183 Left brachial artery narrowing in Takayasu arteritis (courtesy of Ann M. Reed MD, Mayo Clinic).

184, 185 Internal femoral artery narrowing in Takayasu arteritis (courtesy of Ann M. Reed, MD, Mayo Clinic) (184); Takayasu arteritis – thickened aorta (courtesy of Ann M. Reed, MD, Mayo Clinic) (185).
Primary angiitis of the central nervous system

**DEFINITION**
Primary angiitis of the central nervous system (PACNS) is an inflammatory vasculopathy that is limited to cerebral vessels. It was originally described in adults in 1959, and diagnostic criteria were proposed in 1992. These same criteria, stressing the development of neurologic deficits in someone without a systemic condition that can account for the findings, are used in children as well.

**EPIDEMIOLOGY AND ETIOLOGY**
The frequency with which isolated CNS vasculitis occurs in children is unknown because of its novelty; it may go unrecognized by practitioners who are not aware that such an entity exists. In adults, PACNS is estimated to have an incidence of 2–3 cases per 1,000,000 person-years. In the largest pediatric series, boys represented 61% of cases.

Rare cases of CNS vasculitis following varicella have been recognized for decades, and in many cases a preceding infection appears to trigger the condition. In most cases, however, children are diagnosed with PACNS without a convincing environmental, infectious, or genetic explanation.

Biopsies of children with PACNS typically demonstrate lymphocytic infiltration of the wall of cerebral vessels (186). Adults typically have necrotizing granulomas that are not commonly seen in children. These findings suggest an ongoing immunologic attack against an unidentified antigen, though whether this represents true autoimmunity or a response to an exogenous agent is not known.

**CLINICAL HISTORY**
Headache (80%) and focal neurologic deficits (78%) are the most common presenting complaints in children diagnosed with PACNS of the large- or medium-sized cerebral vessels, followed by hemiparesis in 62% (187). Though small-vessel disease may present with the same findings, such patients are more likely to have systemic complaints such as fever or malaise as well. Global CNS dysfunction is uncommon, though children may have seizures or gradual cognitive decline.

**PHYSICAL EXAMINATION**
A thorough neurologic exam is essential in any child suspected of having CNS vasculitis. Small-vessel disease is commonly multifocal, so patients often have subtle evidence of isolated abnormalities of strength or sensation of which they may not even be aware. A patient examiner may also detect cranial neuropathies, optic neuritis, or myelitis.

**LABORATORY**
In PACNS, acute-phase reactants are typically normal, and examination of CSF might be unrevealing as well. Even when a patient has abnormal laboratory studies, they tend to be non-specific, such as elevated serum acute-phase reactants, CSF pleocytosis, or increased intracranial pressure. Thus, diagnosis depends on demonstration of vascular abnormalities on imaging or biopsy.

**IMAGING**
Brain and cerebral vessel imaging are indicated in any child with CNS abnormalities that cannot be ascribed to a clearly defined infectious, toxic, or vascular abnormality. A normal MR scan together with normal CSF have a high negative predictive value for PACNS. None-the-less in 5–10% of cases, only a meningeal and brain biopsy, guided by clinical or MRI abnormalities or performed blindly, is diagnostic of CNS vasculitis.

**DIFFERENTIAL DIAGNOSIS**
The differential diagnosis includes the following:
- Infectious process
- Atherosclerosis
- Embolic disease
- Systemic vasculitits
- Toxin/drug
- Malignancy
- Systemic autoimmune disorder.

**PROGNOSIS**
The number of pediatric cases of isolated CNS vasculitis is insufficient to generate useful prognostic information. Further, an unknown but significant minority of children with PACNS, especially those with localized medium- or large-vessel disease, appear to recover without therapy. It is clear, however,
that early recognition and treatment are essential to ensure optimal outcomes.

**MANAGEMENT**

Brain biopsy is indicated in many children with suspected PACNS, both to confirm the diagnosis and to exclude mimics of CNS vasculitis, especially atypical infections that could worsen if immunosuppression is started empirically. PACNS may be rapidly progressive and neurologically devastating, so risks of diagnostic procedures must be weighed against the need for prompt diagnosis and initiation of therapy. Treatment includes corticosteroids as well as a potent immunosuppressive agent, usually cyclophosphamide. Once remission is achieved, patients are often switched to methotrexate or azathioprine for maintenance therapy.

**COMPLICATIONS**

CNS sequelae depend on the area of brain involved. Potential complications related to therapy include avascular necrosis and opportunistic infection with corticosteroids. Immunosuppressive medications are also associated with increased risk of opportunistic infections.
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CHAPTER 7

Scleroderma
in children
Scleroderma in children includes a group of multisystem autoimmune conditions with the unifying characteristic of the presence of hard skin and the onset before 16 years of age. They may be classified into two main categories: juvenile systemic sclerosis (JSSc) and juvenile localized scleroderma (JLS), known also as morphea.

**Juvenile systemic sclerosis**

**DEFINITION**

Juvenile systemic sclerosis (JSSc) is a chronic, multisystem, connective tissue disease characterized by symmetric thickening and hardening of the skin, associated with fibrous changes in internal organs, such as the esophagus, intestinal tract, heart, lungs, and kidneys, arthritis, and myositis. Very recently, a Committee on Classification Criteria for JSSc, including pediatricians, rheumatologists, and dermatologists, developed new classification criteria (Zulian et al. 2007b).

On the basis of these criteria, a patient, aged younger than 16 years, shall be classified as having JSSc if the one major criterion, presence of proximal skin sclerosis/induration, and at least 2 of 20 minor criteria, grouped in nine main categories, are present (Table 26).

**EPIDEMIOLOGY AND ETIOLOGY**

In general, systemic sclerosis (SSc) has an estimated annual incidence ranging from 0.45 per 100,000 to 1.9 per 100,000, and a prevalence of approximately 15–24 per 100,000 (Mayes et al. 2003). Onset in childhood is very uncommon: children under 16 years account for less than 5% of all cases (Scalapino et al. 2006) and fewer than 10% develop SSc before the age of 20 (Black 1999, Kornreich et al. 1977, Medsger 1994). The onset occurs at a mean age of 8.1 years and the peak age is between 10 and 16 years (Martini et al. 2006, Scalapino et al. 2006). The disease is almost four times more frequent in females and there is no racial predilection (Martini et al. 2006).

The cause of SSc is unknown despite significant advances in the understanding of the pathogenic mechanisms since its initial description (Jimenez & Derk 2004). The disease can be represented as a tripartite process in which dysfunction of the immune system, endothelium, and fibroblasts gives rise to a heterogeneous phenotype that is characterized prominently by fibrosis.

Autoimmunity is evident by the elaboration of circulating disease-specific autoantibodies. Fibroblast dysfunction is manifested as fibrosis due to increased synthesis and deposition of extracellular matrix proteins. Raynaud phenomenon, capillary dropout, endothelial injury, and abnormalities in vascular tone are manifestations of endothelial cell dysfunction. These three aspects, although apparently unrelated to each other, are, actually, closely linked by several immunologic alterations.

The excessive accumulation of collagen in affected skin has led to the hypothesis that there may be abnormalities of collagen type or metabolism (Jelaska et al. 1996). Many reports suggest that cellular immunity plays a major role in the initiation of scleroderma. Several cytokines are increased in scleroderma serum including IL-1, IL-2, IL-4, IL-6, and IL-8 (Atamas et al. 1999, Hasegawa et al. 1999) (Table 27). Endothelial cell injury is another important pathogenic event and often predates fibrotic changes in scleroderma.
Table 26
Classification criteria for juvenile systemic sclerosis

<table>
<thead>
<tr>
<th>Major criterion</th>
<th>Minor criteria</th>
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<tbody>
<tr>
<td>Proximal sclerosis/induration of the skin</td>
<td>Skin Sclerodactyly</td>
</tr>
<tr>
<td></td>
<td>Vascular Raynaud phenomenon</td>
</tr>
<tr>
<td></td>
<td>abnomalities Digital tip ulcers</td>
</tr>
<tr>
<td></td>
<td>Gastrointestinal Gastroesophageal reflux</td>
</tr>
<tr>
<td></td>
<td>Renal Renal crisis</td>
</tr>
<tr>
<td></td>
<td>hypertension New-onset arterial</td>
</tr>
<tr>
<td></td>
<td>Cardiac Arrhythmias</td>
</tr>
<tr>
<td></td>
<td>Respiratory Pulmonary fibrosis (HRCT/radiograph) D</td>
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<tr>
<td></td>
<td>Musculoskeletal Tendon friction rubs</td>
</tr>
<tr>
<td></td>
<td>Neurologic Neuropathy</td>
</tr>
<tr>
<td></td>
<td>Serology ANA</td>
</tr>
<tr>
<td></td>
<td>SSc selective autoantibodies (anti-centromere, anti-topoisomerase I, anti-fibrillarin, anti-PM-Scl, anti-fibrillin, or anti-RNA polymerase I or III)</td>
</tr>
</tbody>
</table>

A patient, aged younger than 16 years, shall be classified as having juvenile systemic sclerosis if the one major criterion and at least 2 of the 20 minor criteria are present. This set of classification criteria has a sensitivity of 90%, a specificity of 96%, and a kappa statistic value of 0.86.

Table 27
Cytokines in juvenile systemic sclerosis

<table>
<thead>
<tr>
<th>Cytokine/growth factorBiologic effect</th>
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</thead>
<tbody>
<tr>
<td>TGF-β, PDGF, IL-1, synthesis</td>
</tr>
<tr>
<td>IL-4, IL-6</td>
</tr>
<tr>
<td>Increased collagen</td>
</tr>
<tr>
<td>IFN-β, IFN-γ, TNF-α, β</td>
</tr>
<tr>
<td>Decreased collagen synthesis</td>
</tr>
<tr>
<td>IFN-β, IFN-γ, TGF-β, PDGF, TNF, IL-1, IL-4</td>
</tr>
<tr>
<td>Fibroblasts proliferation</td>
</tr>
<tr>
<td>IFN-γ, TGF-β, PDGF, TNF, IL-4</td>
</tr>
<tr>
<td>Chemoattraction</td>
</tr>
<tr>
<td>TGF-β, TNF, IL-1 synthesis</td>
</tr>
<tr>
<td>Glycosaminoglycan</td>
</tr>
<tr>
<td>TGF-β, IL-4</td>
</tr>
<tr>
<td>Fibronectin synthesis</td>
</tr>
<tr>
<td>IFN-γ, TNF-α, β, IL-2, NK cell, granzyme A</td>
</tr>
<tr>
<td>Endothelial cell injury</td>
</tr>
<tr>
<td>TGF-β</td>
</tr>
<tr>
<td>Reduction of synthesis</td>
</tr>
<tr>
<td>TNF-α</td>
</tr>
<tr>
<td>Collagenase gene induction</td>
</tr>
<tr>
<td>TGF, transforming growth factor; IFN, interferon; PDGF, platelet-derived growth factor.</td>
</tr>
</tbody>
</table>
CLINICAL HISTORY
The clinical features at the onset of the disease include mainly Raynaud phenomenon and skin induration (188). Raynaud phenomenon is the first sign of the disease in 70% of patients and in 10% it is complicated by digital infarcts. Raynaud phenomenon is more common in the fingers (189) but can be observed in the toes, ears, lips, tongue, and tip of the nose. Proximal skin induration develops later and is the second most frequent symptom, being present in around 40% of patients at onset (Martini et al. 2006).

Other presenting complaints include arthralgia, arthritis, and, although less frequently, muscle weakness, dyspnoea, and calcinosis. Unlike adults, telangiectasia is rarely present in children with JSSc.

PHYSICAL EXAMINATION
Cutaneous involvement in JSSc includes nailfold changes as well as skin induration. These nailfold capillary changes are reported in 40% of patients but their frequency could be much higher if nailfold evaluation was regularly done (Martini et al. 2006). Raynaud phenomenon, positive ANA, and the presence of capillaroscopy changes represent the early signs of the disease and precede by months or years the onset of the other clinical features. These unspecific findings identify the so-called ‘pre-scleroderma’ status (LeRoy & Medsger 2001). Cutaneous changes characteristically evolve in a sequence beginning with edema, followed by induration resulting in marked tightening and contracture (190–192). The skin becomes waxy in texture, tight, hard, and bound to subcutaneous structures. This is particularly noticeable in skin of the digits and face where the characteristic expressionless appearance of the skin may be the first clue to diagnosis (193). Distal tuft loss can be seen on radiographs with severe vasospasm (194).

In children with JSSc, visceral organ involvement can be widespread and is associated with significant morbidity. GI involvement may occur in 30–70% of children with JSSc. Most affected patients have esophageal dysfunction, resulting in gastroesophageal reflux (GER) and dysphagia. Manometry, esophageal scintigraphy, and intraesophageal 24-hour pH monitoring provide more sensitive indicators of diminished lower esophageal sphincter tone and GER (Weber et al. 2000). Large bowel involvement is less frequent and presents as alternating complaints of constipation and diarrhea, bloating, or abdominal discomfort. Lactulose breath test to evaluate bacterial overgrowth, endoscopy, or colon scintigraphy is a useful tool to evaluate this portion of intestinal tract.
190, 191  Tightness of the skin and early mottling in a new-onset patient. (courtesy of Ann M. Reed, MD, Mayo Clinic).

192  Tightness of the skin and early mottling in a new-onset patient. (courtesy of Ann M. Reed, MD, Mayo Clinic).

193  Classic expressionless appearance of a 10-year-old girl with systemic sclerosis. Note also the waxy, translucent appearance of the skin in the upper trunk (courtesy of Dr Ruben Cuttica, Buenos Aires).

194  Distal tuft loss of the first and second DIP joints in a 10-year-old girl with systemic sclerosis and severe sclerodactyly and Raynaud phenomenon (courtesy of Ann M. Reed, MD, Mayo Clinic).
Pulmonary involvement, although frequently asymptomatic, can be present as a dry, hacking cough to dyspnea on exertion. Unlike adults, interstitial pulmonary fibrosis is not frequently reported in children with JSSc at onset but may develop later on during the course of the disease (195). Other abnormalities can be pleuritis, abnormal DLCO and pulmonary arterial hypertension (Martini et al. 2006, Scalapino et al. 2006). The classic radiographic features of interstitial lung disease consist of symmetric, reticulonodular shadowing, most pronounced at the lung bases. High-resolution CT (HRCT) may reveal pulmonary disease even in the presence of a normal chest radiograph. In children, HRCT findings include ground-glass opacification, subpleural micronodules, linear opacities, and honeycombing (Koh & Hansell 2000, Seely et al. 1998). In addition, DLCO and spirometry are sensitive measures of involvement of the respiratory tract. Pulmonary vascular disease occurs very rarely, either as a primary event or as a result of pulmonary fibrosis. In this case, echocardiography is an important tool in detecting early pulmonary hypertension, which should be confirmed by right heart catheterization.

Cardiac involvement is present in around one-fifth of the patients and represents a primary cause of morbidity among children with JSSc (Foeldvari et al. 2000, Martini et al. 2009, Scalapino et al. 2006). Pericardial effusions are not common and are usually of no hemodynamic significance. In addition, pulmonary hypertension caused by pulmonary vascular disease can lead to myocardial damage and right heart failure. Cardiorespiratory complications are the leading cause of death in children with JSSc (Foeldvari et al. 2000, Martini et al. 2009, Scalapino et al. 2006).

There are limited data on the prevalence of renal involvement in children with JSSc. In children with SSc, about 10% had some kind of renal involvement including either increased urinary protein excretion or raised creatinine level (Martini et al. 2006, Scalapino et al. 2006). Although renal involvement in children appears to be not so severe and frequent as in adults, the abrupt onset of accelerated hypertension with acute renal failure (scleroderma renal crisis) remains one of the most severe and life-threatening complications of JSSc.

LABORATORY

About a quarter of JSSc patients have anaemia of chronic disease, or, less commonly, macrocytic anaemia due to malabsorption. Leukocytosis is not prominent but correlates with the degree of visceral or muscle disease. Patients with myositis have elevated levels of creatine kinase (Martini et al. 2006, Scalapino et al. 2006). High titers of ANAs are commonly found in 80–90% of children with JSSc (Martini et al. 2000, Martini et al. 2009, Scalapino et al. 2006). The prevalence of both anti-topoisomerase I (Scl-70) and ACA, SSc-specific antibodies, is lower in children compared with adults, being present in 34% and 7% of the patients, respectively (Martini et al. 2006).
et al. 2006). Positive RF is present in 17% of patients and correlates with the presence of articular involvement (Martini et al. 2006).

**IMAGING**

Imaging is not a prominent tool for managing the skin involvement of JSSc. Joint radiography may help in the assessment of arthritis. Chest imaging, including CT, may be needed to evaluate potential interstitial lung disease. Contrast-based GI studies may be utilized to assess potential esophageal disease.

**DIFFERENTIAL DIAGNOSIS**

Primary Raynaud phenomenon is not associated with any organ involvement. While few other conditions are associated with skin induration, when this occurs in the hands (as it frequently does), it can be confused with an evolving inflammatory arthropathy. Also, particularly when other organ involvement is noted, other connective tissue diseases, such as SLE, are in the differential diagnosis.

**PROGNOSIS**

Generally, the prognosis of JSSc is poor. Skin tightness and joint contractures inevitably lead to severe disability (Bottoni et al. 2000). It has been reported that the skin may eventually soften years after onset of the disease. Survivorship has not been determined in any large series of children because of the rarity of this disease and very few retrospective data are available (Foeldvari et al. 2000, Martini et al. 2009, Scalapino et al. 2006). The overall mortality rate at 5 years ranges between 6% and 15% and is better than in adults. The causes of death in JSSc include cardiac failure (67%), end-stage renal failure (13%), respiratory failure (10%), infections (7%), and hypertensive encephalopathy (3%) (Foeldvari et al. 2000, Martini et al. 2009).

**MANAGEMENT**

Due to the rarity of SSc in children, very little information is available on the treatment and no controlled studies have been performed so far. The non-pharmacologic measures include physiotherapy to help maintain functional ability, muscle strength, and joint movement while preventing flexion contractures, and the use of corrective splints to treat or prevent contractures. Patients and parents should be told to avoid cold and trauma since they can exacerbate symptoms.

The pharmacologic management of patients with JSSc is challenging since no drug has been shown to be of significant benefit in either children or adults with SSc. Calcium channel blockers, usually oral nifedipine or nicardipine, should be considered as first-line therapy for Raynaud phenomenon, and iloprost, or other available IV prostanoids, for severe SSc-related Raynaud phenomenon and digital ulcers (Pope et al. 1998, Zulian et al. 2004). In accordance with the data for adults, cyclophosphamide may be considered for the treatment of SSc-related interstitial lung disease in children (Hoyles et al. 2006, Tashkin et al. 2006).

Since steroids seem to be associated with a higher risk of scleroderma renal crisis (DeMarco et al. 2002, Steen & Medsger 1998), patients on steroids, for concomitant myositis or arthritis, should be carefully monitored for blood pressure and renal function.

Methotrexate, which is widely used for the treatment of many rheumatic conditions in children, has been shown to improve skin softening in adults (Pope et al. 2001) and could be the treatment of choice for skin manifestations also for children, especially in the early phase of the disease.

New drugs for the treatment of pulmonary arterial hypertension, such as bosentan, sitaxsentan, and sildenafil, have been recently introduced in adult patients with SSc (Barst et al. 2004, Galie et al. 2005, Rubin et al. 2002). There is limited experience, at present, to recommend their use in children.

**COMPLICATIONS**

The most common causes of death in children are related to involvement of the cardiac, renal, and pulmonary systems. Cardiomyopathy, although rare, can be one of the causes of early death, especially in children (Quartier et al. 2002). Interstitial lung disease and renal failure or acute hypertensive encephalopathy supervene as a potentially fatal outcome in few children and seems more likely to occur early in the disease course (Foeldvari et al. 2000, Martini et al. 2009). Complications of medical therapies such as infections associated with immunosuppressive medications and bone disease related to corticosteroids may occur.
Juvenile localized scleroderma

**DEFINITION**

Juvenile localized scleroderma (JLS), known as morphea, comprises a group of conditions that involve essentially the skin and subcutaneous tissues. They have various features and range from very small plaques to extensive fibrotic lesions which may cause significant functional and cosmetic deformity.

**EPIDEMIOLOGY AND ETIOLOGY**

Although JLS is relatively uncommon, it is far more common than SSc in childhood, by a ratio of at least 10:1 (Peterson et al. 1997). There is a mild female predilection, the F:M ratio being 2.4:1 (Zulian et al. 2006a). The disease onset is usually during late infancy although a few cases with onset at birth have been described (Zulian et al. 2006b).

The etiology and pathogenesis of localized scleroderma (LSc) remain uncertain. Two possible contributing pathogenic processes include abnormal fibroblast function and autoimmune dysfunction. One study demonstrated increased collagen production by fibroblasts in skin biopsy specimens from patients with morphea (Kähäri et al. 1988). In addition, raised levels of cytokines that increase collagen synthesis by fibroblasts have been reported in LSc lesions (Liu & Connolly 1998). Autoantibodies are a frequent finding in patients with LSc (Harrington & Dunsmore 1989, Liu & Connolly 1998, Zulian et al. 2006a). Relatives of affected patients also have an increase in organ-specific autoantibodies compared with control patients (Harrington & Dunsmore 1989). In a large multicenter retrospective review of 750 patients, 13.3% of the patients reported specific environmental events as a potential trigger for their disease (Zulian et al. 2006a). These included infections, drugs, mechanical events (e.g. accidental trauma, insect bite, and vaccination), and psychological stress. Trauma has been implicated in the initiation of morphea lesions in 3–13% of patients with LSc (Harrington & Dunsmore 1989, Liu & Connolly 1998, Zulian et al. 2006a). The underlying mechanisms on how trauma would contribute to the development of LSc is unclear, although cytokines and neuropeptides (e.g. endothelin-1), which are normally involved in wound healing, have been suggested to play a causative role (Liu & Connolly 1998).

**CLINICAL HISTORY**

The primary presentation of JLS is dermatologic. Skin lesion(s) may develop in any skin area. They are generally painless, and may be expanding. Other features depend on what area of skin is involved. If the lesion is on the trunk, it may be completely asymptomatic or a change in skin color or texture. If the lesion occurs on the face, it will likely be noticed earlier in the disease course. The family may notice different skin color and texture of the child’s face, or facial movements may seem altered. If the lesion occurs on an extremity, it could impact the range of motion (ROM) of the underlying joint. The classification of JLS is based on clinical and pathologic findings discussed below.

**PHYSICAL EXAMINATION**

JLS can be classified into five subtypes: circumscribed morphea, linear scleroderma, generalized morphea, pansclerotic morphea, and a mixed subtype where a combination of two or more of the previous subtypes is present (Table 28). Circumscribed morphea is characterized by oval or round circumscribed areas of induration surrounded by a violaceous halo. It is confined to the dermis with only occasional involvement of the superficial panniculus. Sometimes, as in deep morphea, the entire skin feels thickened, taut, and bound down (196a, 196b).
### Table 28
Classification criteria of juvenile localized scleroderma (Consensus conference, Padua (Italy) 2004)

<table>
<thead>
<tr>
<th>Main group</th>
<th>Subtype</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circumscribed morphea</td>
<td>Superficial</td>
<td>Oval or round circumscribed areas of induration limited to epidermis and dermis, often with altered pigmentation and violaceous, erythematous halo (lilac ring). They can be single or multiple</td>
</tr>
<tr>
<td></td>
<td>Deep</td>
<td>Oval or round circumscribed deep induration of the skin involving subcutaneous tissue extending to fascia and may involve underlying muscle. The lesions can be single or multiple. Sometimes the primary site of involvement is in the subcutaneous tissue without involvement of the skin</td>
</tr>
<tr>
<td>Linear scleroderma</td>
<td>Trunk/limbs</td>
<td>Linear induration involving dermis, subcutaneous tissue, and, sometimes, muscle and underlying bone and affecting the limbs and/or the trunk</td>
</tr>
<tr>
<td></td>
<td>Head</td>
<td><em>En coup de sabre</em> (ECDS). Linear induration that affects the face and/or the scalp and sometimes involves muscle and underlying bone. Parry–Romberg syndrome (or progressive hemifacial atrophy): loss of tissue on one side of the face that may involve dermis, subcutaneous tissue, muscle, and bone. The skin is mobile</td>
</tr>
<tr>
<td>Generalized morphea</td>
<td></td>
<td>Induration of the skin starting as individual plaques (four or more and larger than 3 cm) that become confluent and involve at least two anatomic sites</td>
</tr>
<tr>
<td>Pansclerotic morphea</td>
<td></td>
<td>Circumferential involvement of limb(s) affecting the skin, subcutaneous tissue, muscle, and bone. The lesion may also involve other areas of the body without internal organ involvement</td>
</tr>
<tr>
<td>Mixed morphea</td>
<td></td>
<td>Combination of two or more of the previous subtypes. The order of the concomitant subtypes, specified in brackets, will follow their predominant representation in the individual patient, i.e. mixed (linear-circumscribed)</td>
</tr>
</tbody>
</table>

**Figure 196a** Circumscribed morphea in a 12-year-old boy: single indurate plaque with evident hyperemic border ('lilac ring').

**Figure 196b** Circumscribed morphea with keloid formation after biopsy.
When individual plaques are four or more, larger than 3 cm and become confluent, we have the so-called generalized morphea (197–200). Linear scleroderma, the most common subtype in children and adolescents, is characterized by one or more linear streaks that can extend through the dermis, subcutaneous tissue, and muscle to the underlying bone, causing significant deformities (201–206).

197–200 Generalized morphea in a 17-year-old male: indurate plaques that become confluent and involve the trunk symmetrically (197); generalized morphea in a girl with new-onset disease, which resolved entirely on treatment (courtesy of Ann M. Reed, MD, Mayo Clinic) (198); generalized morphea in a girl with new-onset disease, which resolved entirely on treatment (courtesy of Ann M. Reed, MD, Mayo Clinic) (199); generalized morphea in a girl with new-onset disease, which resolved entirely on treatment (courtesy of Ann M. Reed, MD, Mayo Clinic) (200).
Linear scleroderma involving the whole right lower limb and resulting in undergrowth of the foot and hyperpigmented skin (201); linear scleroderma involving the whole left lower limb and resulting in undergrowth of the foot and hyperpigmented skin (202); linear scleroderma involving the whole left lower limb; close-up of the foot (same patient as 202) (203); linear scleroderma of the right upper limb: note the flexion contractures of the wrist and fingers (204); linear scleroderma of the left chest wall (205); linear scleroderma of the right neck causing severe limitation of motion (206).
The upper or lower extremities can be affected but also face or scalp, as in the *en coup de sabre* (ECDS) variety so called because the lesion looks like the depression caused by a dueling stroke from a sword (207–211). This latter form can be complicated by CNS abnormalities, such as calcifications, vasculitis or simply MRI-unspecific abnormalities, and dental and even ocular abnormalities (Zannin *et al.* 2007). In patients with ECDS, associated neurologic manifestations include seizures (particularly complex partial seizures), headaches, and facial paralysis. The neurologic findings may occur before, concurrent with, or after the skin manifestations. Parry–Romberg syndrome (or progressive hemifacial atrophy) loss of tissue on one side of the face that may involve dermis, subcutaneous tissue, muscle, and bone is also seen (210, 211). Pansclerotic morphea, an extremely rare but severe subtype, is characterized by generalized full-thickness involvement of the skin of the trunk, extremities, face, and scalp with sparing of the fingertips and toes (212). Eosinophilic fasciitis is a form that extends into the fascia with inflammation and eosinophils present with boggy, tight extremities (213, 214).

Extracutaneous manifestations in patients with LSc mainly include articular, neurologic, ocular, and GI involvement (Zulian *et al.* 2005). Articular involvement is the most frequent finding, being reported in 19% of the patients (Dehen *et al.* 1994, Zulian *et al.* 2005). It is mainly present in patients with linear scleroderma. These children tend to have an accelerated course with predominant musculoskeletal disease, including rapid development...
**210–214** Linear scleroderma and Parry–Romberg syndrome (or progressive hemifacial atrophy) loss of tissue on one side of the face that may involve dermis, subcutaneous tissue, muscle, and bone (courtesy of Ann M. Reed, MD, Mayo Clinic) (210, 211); disabling pansclerotic morphea (212); eosinophilic fasciitis in a young girl on presentation (courtesy of Ann M. Reed, MD, Mayo Clinic) (213, 214).
of contractures. Neurologic manifestations, reported in 4% of the patients, consist essentially in epilepsy and headaches, behavioral changes, and learning disabilities (Blaszczyk et al. 2003, Kister et al. 2008, Zulian et al. 2005). Abnormalities on MRI, such as calcifications, white matter changes, vascular malformations, and changes consistent with CNS vasculitis, have also been reported (Kister et al. 2008). Ocular changes are reported in 3.2% of the children affected by localized scleroderma (Zannin et al. 2007). As expected, most of them have ECDS but, interestingly, around one-third had no facial skin lesion. The most frequent lesions are on eyelids and eyelashes, one-third consisting of anterior segment inflammation, such as anterior uveitis or episcleritis, the remaining being CNS-related abnormalities, paralytic strabismus, pseudopapilledema, or refractive errors.

Esophageal motor function abnormalities with GER have been reported in 2–4% of children with JLS (Guariso et al. 2007, Zulian 2004). These findings might support the indication for an extensive GI evaluation in patients with LSc, especially in the presence of specific GI symptoms.

LABORATORY

In a large cohort of patients, ANAs were found in 42% of patients with no correlation between their presence and a particular subtype or disease course (Table 29) (Zulian et al. 2006a). Anti-topoisomerase I antibodies (anti-Scl-70) and ACAs, specific markers of systemic sclerosis, were found to be positive in 2–3% of children with JLS (Zulian et al. 2006a). Whether these antibodies reflect just the immunologic component of the disease process or can also have a prognostic significance is unclear. RF detected, at low titre in 16% of the patients, significantly correlates with the presence of arthritis (Zulian et al. 2006a). Children with LSc who develop arthritis are more likely to have a positive RF, and sometimes an elevated ESR and circulating autoantibodies (Zulian et al. 2005).

IMAGING

The management of JLS is challenging and the detection of disease activity and progression remains a fundamental problem. Non-invasive technology may be helpful in following the extent of disease in selected patients. Thermography, which measures the radiation of infrared heat from the body, is able to discriminate active from inactive lesions (Martini et al. 2002). This procedure is non-invasive and usually well tolerated. It allows for easy interpretation and rapid results that are helpful in decision-making. There are false-positive results in cases of atrophic or old lesions, particularly in the scalp region (Martini et al. 2002).

Ultrasoundography is a non-invasive tool, which may also be useful in the evaluation of LSc patients (Cosnes et al. 2003). Unfortunately, this evaluation is difficult in overweight patients, operator dependent, and not yet standardized for children. Ultrasonography can also be used to monitor disease activity (Li et al. 2007). A computerized skin score method allows the measurement of single target lesions and comparison over time. It consists in the demarcation of the indurate borders of the lesions on an adhesive transparent film and subsequent transfer, as a scanned image, in a computer (Zulian et al. 2007a). The calculation of the affected area is then performed by computer software. This new technique is applicable in day-to-day practice, is not time-consuming, and does not need specialized equipment.

Bone growth malformations may be monitored with radiographs to determine bone maturation and possible surgical intervention (215).

<table>
<thead>
<tr>
<th>Serum autoantibody</th>
<th>Patients %</th>
<th>Healthy controls %</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANA</td>
<td>42.3</td>
<td>0–3</td>
</tr>
<tr>
<td>Anti-topoisomerase I</td>
<td>3.2</td>
<td>0</td>
</tr>
<tr>
<td>ACA</td>
<td>1.7</td>
<td>0</td>
</tr>
<tr>
<td>Anti-dsDNA</td>
<td>4.2</td>
<td>0</td>
</tr>
<tr>
<td>RF</td>
<td>16</td>
<td>0–4</td>
</tr>
<tr>
<td>APL antibody</td>
<td>12.6</td>
<td>1.5–9.4</td>
</tr>
</tbody>
</table>

Table 29 Serum autoantibodies in juvenile localized scleroderma
DIFFERENTIAL DIAGNOSIS
Other dermatologic conditions, such as vitiligo, are in the differential diagnosis of JLS. When it occurs over a peripheral joint(s), it can be confused with an inflammatory arthropathy. When presentation includes facial involvement, particularly in a young child, consideration of congenital abnormalities is given.

PROGNOSIS
The prognosis of JLS depends primarily on what area of the skin is involved. Facial disease, and skin involvement over peripheral joints may have greater consequences if disease progresses. Also, if the JLS involves deeper skin layers, it is frequently more difficult to treat.

MANAGEMENT
Over the years, many treatments have been tried for localized scleroderma. Unfortunately, the rarity of the disease and the difficulty in assessing outcomes in an objective way have limited the interpretation of most of these studies.

Treatment should be mainly topical such as moisturizing agents, topical glucocorticoids, or calcipotriene (Cunningham et al. 1998). When there is a significant risk for disability, such as in linear and deep subtypes, systemic treatment with methotrexate in combination with corticosteroids should be considered (Uziel et al. 2000, Weibel et al. 2006). Methotrexate can be used at a weekly regimen of 10–15 mg/m² as a single oral or subcutaneous dose for at least 1 year. During the first 2–3 months of therapy, a course of glucocorticoid steroids (prednisone 0.5–1 mg/kg per day administered orally or methylprednisolone 20–30 mg/kg per day administered intravenously for three consecutive days) may be used as adjunctive bridge therapy for those with rapidly progressive disease. Most patients show a response within 2–4 months and the side effects are usually mild and associated more with corticosteroid than with methotrexate treatment. Unfortunately, no controlled trials are available and the series of treated patients are very small.

COMPLICATIONS
Circumscribed morphea generally is of cosmetic concern only, and frequently lesions spontaneously remit with residual pigmentation as the only abnormality. With ECDS, facial disfigurement can occur. Rarely seizure disorders can be associated with this form of JLS. Persistent flexion contractures of peripheral joints may be seen as well. Complications of medical therapies such as infections associated with immunosuppressive medications and bone disease related to corticosteroids may occur.
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Autoinflammatory diseases

• The monogenic autoinflammatory diseases
• Other monogenic autoinflammatory syndromes
The autoinflammatory diseases or the periodic fever syndromes have opened a new chapter in the field of rheumatology. The systemic autoinflammatory diseases are a group of rare disorders characterized by unprovoked inflammatory episodes without high-titer autoantibodies (Stojanov & Kastner 2005). These are mainly monogenic diseases where mutations in a single gene in the inflammatory pathway have been associated with a disease. The most typical example is familial Mediterranean fever (FMF). However, recently a number of other complex diseases have been included in this group (Ozen et al. 2006).

THE MONOGENIC AUTOINFLAMMATORY DISEASES

Familial Mediterranean fever

DEFINITION
Familial Mediterranean fever (FMF, MIM249100) is a disease caused by mutations in the gene coding for pyrin (Aksentijevich et al. 1999). Clinically it is characterized by recurrent attacks of fever and serositis along with increased acute-phase inflammation (Samuels & Ozen 2006).

EPIDEMIOLOGY AND ETIOLOGY
The disease is caused by mutations in the Mediterranean fever (MEFV) gene. Mutations in this gene are known to be very common in people of eastern Mediterranean origin with carrier rates as high as 1:3 to 1:5 (Yilmaz et al. 2001). The frequency of these mutations has led to investigations on the advantage introduced by the carrier state for these mutations. Today it is widely believed that these mutations offer a better and maybe exaggerated inflammatory response to pathogens (Lachmann et al. 2006, Ozen et al. 2002).

More than 50 mutations in the MEFV gene have been associated with FMF (Aksentijevich et al. 1999, Stojanov & Kastner 2005). The MEFV gene consists of 10 exons and most of the disease-causing mutations are located on exon 10.

The gene codes for pyrin, an important protein of the inflammasome and the inflammatory pathway. Pyrin regulates IL-1 production through its interaction with caspase-1; a direct effect of pyrin on IL-1β activation and the idea that pyrin interacts directly with caspase-1 to modulate IL-1β production has been suggested in recent studies (Chae et al. 2006). Thus mutations in pyrin are associated with increased IL-1β production. In fact the IL-1 receptor antagonist anakinra was shown to suppress acute-phase proteins in a patient with FMF and amyloidosis (Chae et al. 2006).

CLINICAL HISTORY
The hallmark of clinical presentation is the typical attacks in the form of fever and serositis along with elevation in acute-phase reactants (Samuels & Ozen 2006). The typical attack of FMF lasts half a day to 3 days, and is accompanied by fever (Livneh et al. 1996). Myalgia may occur (Tunca et al. 2005). In the majority of patients serositis manifests itself as abdominal pain. The pain is incapacitating. Arthritis and/or arthralgia is also very frequent and is present in half of the patients in the large series reported from Israel and Turkey (Livneh et al. 1996, Tunca et al. 2005). Pleurisy in the form of chest pain is common; it is unilateral and often spreads to the shoulder.

The disease is clinically diagnosed on the basis of typical attacks with increased inflammatory response. Livneh et al., from Israel, have suggested a set of criteria for diagnosis; however, these have not been validated in other ethnic groups (Livneh et al. 1997). According to these criteria there are major and minor criteria as well as supportive criteria. The four major criteria include the typical attacks (≥3 of the same type; rectal temperature ≥38°C; 12–72 hour) with peritonitis, pleuritis, monoarthritis (hip, knee, and ankle), and fever alone. Minor criteria are defined as incomplete attacks, exertional leg pain, and favorable response to colchicines (Livneh et al. 1997).

PHYSICAL EXAMINATION
The only skin manifestation of FMF – except when it is associated with vasculitis – is erysipelas-like erythema (ELE), which is provoked by pressure-bearing exercise (216). ELE may not necessarily accompany an attack. In fact some manifestations of FMF are not associated with an attack. These include protracted febrile myalgia, hematuria, arthralgia, and exertional myalgia. Another
interesting feature of FMF is the association with vasculitides and especially polyarteritis nodosa (Ozen et al. 2001). A large series from Turkey has shown that these patients have increased rates of Henoch–Schönlein purpura, polyarteritis nodosa, and probably Behçet disease (Tunca et al. 2005).

LABORATORY
Subclinical inflammation persists even in between attacks (Lachmann et al. 2006). Lachmann et al. have shown that there was also some inflammatory activity between attacks as well (median SAA 6 mg/l, CRP 4 mg/l). They have thus suggested that substantial subclinical inflammation occurs widely and over prolonged periods in patients with FMF, indicating that the relatively infrequent clinically overt attacks represent the ‘tip of the iceberg’ in this disorder.

IMAGING
Imaging is not compulsory in diagnosing FMF. An ultrasound and chest radiograph may show fluid in the peritoneum and pleura, respectively. An echocardiogram may demonstrate pericardial fluid in rare cases who have pericarditis.

DIFFERENTIAL DIAGNOSIS
A mutation analysis for the MEFV gene is indicated in a patient who has recurrent attacks lasting less than 3 days with elevated acute-phase reactants, and lacks any skin findings except for ELE and lacks the features of the other periodic fever diseases (Table 30). The ethnic background is important in the differential workup of these patients, though one must not forget that FMF patients have been identified in many ethnic groups including Japanese individuals. Differential diagnosis includes all other periodic fever syndromes and is very problematic in a patient who does not have an eastern Mediterranean ancestry (Table 30) (Samuels & Ozen 2006). Urticaria and other skin findings that occur in other periodic fever syndromes are not present in FMF. The time frame of the attack helps in differentiating from TRAPS. However, the most important differentiating characteristic of FMF is probably the response to colchicine. This is the rationale for using colchicine for a trial period in patients who are thought to have FMF but lack genetic confirmation.

PROGNOSIS
This is the only periodic fever where an excellent quality of life is possible in the majority of patients – with a cheap pill. In fact colchicine has been shown to prevent amyloidosis in compliant patients in series from Israel and Turkey (Kallinich et al. 2007, Saatci et al. 1997, Zemer et al. 1991). Colchicine is also indicated for quality of life since it prevents attacks or decreases the severity and/or frequency of attacks (Kallinich et al. 2007).

MANAGEMENT
Evidence for the effectiveness of colchicine in children suffering from FMF came from subsequently performed open-label studies (evidence level IIA) (Kallinich et al. 2007). In the large cohorts of children (n = 809) no manifestation of amyloidosis occurred in 809 treated children during an observation period of up to 13 years (Zemer et al. 1991). In the Turkish FMF registry 2.3% developed amyloidosis, all being non-compliant with respect to regular colchicine intake (Saatci et al. 1997).

A starting dose of colchicine 0.5 mg/day in young children and 1.0 mg/day in older children (>5 years) is sufficient for disease control in approximately half of patients (evidence level IIB) (Kallinich et al. 2007). The dose is increased to the adult dose of 1.5–2 mg/day once the child weighs more than 27 kg. High SAA levels might indicate insufficient disease control and thereby the need for
dose adjustment (evidence level III) (Kallinich et al. 2007).

There are many problems that as pediatricians we should warn the families about: the risk in any other children; to follow the siblings closely for any symptoms of unexplained fever or typical attacks. A routine mutation analysis of the MEFV gene for sibs is probably not indicated. However, if there is a case of renal amyloidosis in the family, a genetic workup may be required. SAA levels may also be checked in siblings.

The diagnosis in a patient without genetic confirmation is more problematic. Since this is an autosomal recessively inherited disease we require the presence of two mutations in the MEFV gene to confirm the diagnosis of FMF. However, there may be patients who resemble the FMF phenotype but have failed to show two or even any mutation in the MEFV gene. A trial of colchicine is warranted if the patient has a phenotype suggestive of FMF since this is the only autoinflammatory disease that responds to colchicine. A trial with and without colchicine should be compared in terms of attacks and acute-phase response, especially the SAA levels. Features of other autoinflammatory diseases should be carefully sought out, especially in patients who do not have an eastern Mediterranean ancestry.

**COMPLICATIONS**

The primary potential complication of FMF is the development of amyloidosis. Complications of colchicine treatment include diarrhea.

<table>
<thead>
<tr>
<th>Table 30</th>
<th>Differential diagnosis of periodic fevers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inheritance&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Recessive</td>
</tr>
<tr>
<td>Predominant ethnicities/ ancestries</td>
<td>Jewish, Armenian, Arab, Turkish, Italian</td>
</tr>
<tr>
<td>Gene</td>
<td>MEFV</td>
</tr>
<tr>
<td>Protein</td>
<td>Pyrin (marenostrin)</td>
</tr>
<tr>
<td>Duration of episodes</td>
<td>1–3 days</td>
</tr>
<tr>
<td>Cutaneous disease</td>
<td>Erysipeloid-like erythema</td>
</tr>
<tr>
<td>Musculoskeletal findings</td>
<td>Episodic monoarthritis, sacroilitis</td>
</tr>
<tr>
<td>Abdominal involvement</td>
<td>Sterile peritonitis 85%</td>
</tr>
<tr>
<td>Other key features</td>
<td>Pleurisy, asymptomatic pericardial effusions; scrotal pain</td>
</tr>
<tr>
<td>Treatment</td>
<td>Colchicine prophylaxis</td>
</tr>
</tbody>
</table>

<sup>a</sup>Inheritance patterns are all autosomal.
(with permission from Samuels & Ozen 2006)

<table>
<thead>
<tr>
<th>TRAPS</th>
<th>FCAS</th>
<th>MWS</th>
<th>NOMID/CINCA</th>
<th>PAPA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dominant</td>
<td>Dominant</td>
<td>Dominant</td>
<td>Dominant</td>
<td>Dominant</td>
</tr>
<tr>
<td>Irish and Scottish, but expanding</td>
<td>Mostly European</td>
<td>Northern European</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td><strong>TNFRSF1A</strong></td>
<td><strong>CIAS1/NALP3/PYPAF1</strong></td>
<td><strong>CIAS1/NALP3/PYPAF1</strong></td>
<td><strong>CIAS1/NALP3/PYPAF1</strong></td>
<td><strong>PSTPIP1/CD2BP1</strong></td>
</tr>
<tr>
<td>55 dDa TNF receptor</td>
<td>Cryopyrin</td>
<td>Cryopyrin</td>
<td>Cryopyrin</td>
<td>PSTPIP1/CD2BP1</td>
</tr>
<tr>
<td>Often &gt;1 week</td>
<td>Usually &lt;24 hours</td>
<td>24–48 hours</td>
<td>Continuous but with flares</td>
<td>Variable</td>
</tr>
<tr>
<td>Migratory erysipelas-like rash overlying myalgias</td>
<td>Urticarial rash induced by exposure to cold</td>
<td>Urticarial rash</td>
<td>Urticarial rash</td>
<td>Pyoderma gangrenosum, acne</td>
</tr>
<tr>
<td>Severe migratory myalgia, arthralgia, often non-erosive monoarthritis</td>
<td>Myalgias, debilitating polyarthralgias</td>
<td>Myalgias, intermittent arthralgias, large joint oligoarticular arthritis</td>
<td>Epiphyseal/patellar overgrowth, periosteal elevation, intermittent or chronic arthritis</td>
<td>Pyogenic, sterile arthritis</td>
</tr>
<tr>
<td>Pain, peritonitis, diarrhea, constipation</td>
<td>Nausea</td>
<td>Pain</td>
<td>Uncommon but hepatosplenomegaly with flares</td>
<td>None</td>
</tr>
<tr>
<td>Pleurisy</td>
<td>Headache</td>
<td>Sensorineural deafness</td>
<td>Learning disability, chronic aseptic meningitis, headache, sensorineural deafness</td>
<td>Destructive, recurrent inflammation in skin, joints, and muscles</td>
</tr>
<tr>
<td>Anti-TNF-α prophylaxis</td>
<td>Avoidance of cold, anti-IL-1, NSAIDs</td>
<td>Anti-IL-1, NSAIDs, prednisone?</td>
<td>Anti-IL-1, anecdotal anti-TNF-α</td>
<td>Steroids (anecdotal anti-IL-1 and anti-TNF)</td>
</tr>
</tbody>
</table>
Cold-induced autoinflammatory syndrome 1 (CIAS1-pathies) – the cryopyrin-associated periodic fever syndromes

DEFINITION
The cryopyrin-associated periodic fever syndromes (CAPS) include familial cold autoinflammatory syndrome (FCAS, MIM120100), Muckle–Wells syndrome (MWS, MIM191900), and neonatal-onset multisystem inflammatory disease (NOMID, also known as chronic infantile neurologic cutaneous articular (CINCA) syndrome, MIM607115).

Cold-induced autoinflammatory syndrome 1 (CIAS1), CIAS1-pathies or cryopyrinopathies, or CAPS, is a group of autoinflammatory diseases caused by mutations in the gene coding for cryopyrin (Samuels & Ozen 2006, Stojanov & Kastner 2005). In this group we have three major diseases, with FCAS representing the mildest and NOMID the most severe form of the disease complex (Hawkins et al. 2004, Hoffman et al. 2004, Johnstone et al. 2003, Neven et al. 2004).

EPIDEMIOLOGY AND ETIOLOGY
The reported patients with FCAS and MWS have been mainly confined to European ancestry whereas NOMID/CINCA has been reported in a variety of ethnic groups (Stojanov & Kastner 2005). In this group we have three major diseases, with FCAS representing the mildest and NOMID the most severe form of the disease complex (Hawkins et al. 2004, Hoffman et al. 2004, Johnstone et al. 2003, Neven et al. 2004).

Cryopyrin shares a similar pyrin domain. Cryopyrin forms a macromolecular complex with ASC, CARDINAL (CARD8), and caspase-1 called the inflammasome, which mediates caspase-1 and IL-1β activation (Ozen et al. 2006, Stojanov & Kastner 2005). Mutations in cryopyrin are believed to cause an increased inflammasome activity.

Although these three syndromes may be associated with different mutations in CIAS1 and there is some phenotype–genotype correlation, the same single missense mutation can produce variant phenotypes and/or overlapping phenotype with partial criteria from two or more of these syndromes (Hawkins et al. 2004, Hoffman et al. 2004, Johnstone et al. 2003, Neven et al. 2004). There is considerable overlap among some patients and there is evidence that other genes and/or environmental factors are effective in the pathogenesis of these diseases.

CLINICAL HISTORY
All of the CIAS1-pathies are characterized by fevers and a non-pruritic, urticarial rash (217–221) usually presenting in infancy (later in MWS), and they may have some overlapping features as described below. FCAS presents with attacks precipitated by cold as suggested in the name. The main differentiating feature of MWS is the hearing defect. The naming of the third disease in this group reflects the characteristics of the disease: NOMID for the onset and CINCA for the involved systems. NOMID and sometimes the other diseases tend to have more persistent features and may lack the typical attacks that may prove important in the differential diagnosis (Samuels & Ozen 2006).

PHYSICAL EXAMINATION
There are some overlapping signs such as conjunctivitis (FCAS, MWS) or sensorineural hearing loss and systemic amyloidosis (MWS, NOMID) among these diseases. Unique features of each disease include the precipitation of attacks by generalized cold exposure in FCAS, the hearing defect in MWS and more severe CNS involvement including optic nerve irritation/papilledema, chronic aseptic meningitis, learning disability, as well as facial dysmorphism in NOMID (Hawkins et al. 2004, Hoffman et al. 2004, Johnstone et al. 2003, Neven et al. 2004). The musculoskeletal involvement can be in the form of arthralgias (FCAS) to synovitis (MWS) to the characteristic arthropathy of NOMID/CINCA with premature ossification and overgrowth and bone inflammation (222). The arthropathy of NOMID/CINCA is very characteristic.

LABORATORY
There are no specific laboratory tests. Acute-phase reactants are expected to be increased.

IMAGING
Imaging may be useful to identify bony abnormalities including inflammation and overgrowth (223) and to exclude other causes of periodic fevers.
The monogenic autoinflammatory diseases

**Urticarial rash in a patient with CIAS1-pathy.**

**218–223** Urticarial rash in a newborn with CIAS1-pathy (courtesy of Ann M. Reed, MD, Mayo Clinic) (**218, 219**); urticarial rash as seen in a patient with FCAS and MWS (courtesy of Dawn Davis, MD, Mayo Clinic) (**220, 221**); bone lesions seen in CIAS1-pathy (courtesy of Ann M. Reed, MD, Mayo Clinic) (**222, 223**).
DEFINITION

Hyperimmunoglobulinemia D with periodic fever syndrome (HIDS, MIM260920) is an autosomal recessively inherited periodic fever syndrome caused by mutations in the gene MVK, which codes for the enzyme mevalonate kinase (Drenth et al. 1999).

EPIDEMIOLOGY AND ETIOLOGY

The majority of the reported patients are from the Netherlands and France. However, cases have been reported worldwide. The lack of mevalonate causes an elevation of mevalonic acid along with a lack of production of the downstream elements, which are the isoprenoids (Frenkel et al. 2001). The deficiency of isoprenoids has been shown to contribute to increased IL-1β secretion. Thus it is not the increased IgD but rather the metabolic pathway that leads to the inflammatory burst (Frenkel et al. 2001, Ozen et al. 2006).

HIDS is actually a milder form of the disease named mevalonic aciduria, which results in even higher levels of urinary mevalonate. Those children have mutations on the same gene as HIDS but suffer from severe neurologic deficits and dysmorphic features as well as the fevers (Prietsch et al. 2003).

DIFFERENTIAL DIAGNOSIS

A mutation analysis for the CIAS1 gene is indicated in a patient with inflammatory attacks provoked by cold – even air conditioning, a patient with hearing loss and recurring attacks with urticaria unresponsive to colchicine, or a young feverish child with recurrent meningitis or other neurologic and skin features. The differential diagnosis includes all other periodic fever syndromes. The unique features of each disease are helpful. It should be noted that NOMID is probably the only disease with more persistent symptoms rather than attacks. Systemic juvenile idiopathic arthritis and a number of infectious diseases may also be considered in the differential diagnosis of NOMID/CINCA.

PROGNOSIS

NOMID/CINCA represents the most severe condition and can be complicated with CNS involvement including learning disability, chronic aseptic meningitis, increased intracranial pressure, papilledema, cerebral atrophy, and sensorineural hearing loss as well as chronic inflammatory and destructive arthropathy, skeletal dysplasia, and retinal dystrophy.

MANAGEMENT

Case reports and open-label studies highlight the success of anti-IL-1 treatment currently with anakinra (Lovell et al. 2005). This fits in with the pathogenesis of the disease as well since there is a loss of control in IL-1 production. Anti-IL-1 treatment has even been effective in suppressing amyloidosis. Symptomatic treatment and avoidance of cold are also important for FCAS. There are case reports with etanercept (only versus the arthropathy) as well (Federico et al. 2003).

COMPLICATIONS

Secondary amyloidosis may occur as a complication of ongoing inflammation. Opportunistic infections may occur with anti-IL-1 therapy.

Hyperimmunoglobulinemia D with periodic fever syndrome

DEFINITION

Hyperimmunoglobulinemia D with periodic fever syndrome (HIDS, MIM260920) is an autosomal recessively inherited periodic fever syndrome caused by mutations in the gene MVK, which codes for the enzyme mevalonate kinase (Drenth et al. 1999).

EPIDEMIOLOGY AND ETIOLOGY

The majority of the reported patients are from the Netherlands and France. However, cases have been reported worldwide. The lack of mevalonate causes an elevation of mevalonic acid along with a lack of production of the downstream elements, which are the isoprenoids (Frenkel et al. 2001). The deficiency of isoprenoids has been shown to contribute to increased IL-1β secretion. Thus it is not the increased IgD but rather the metabolic pathway that leads to the inflammatory burst (Frenkel et al. 2001, Ozen et al. 2006). HIDS is actually a milder form of the disease named mevalonic aciduria, which results in even higher levels of urinary mevalonate. Those children have mutations on the same gene as HIDS but suffer from severe neurologic deficits and dysmorphic features as well as the fevers (Prietsch et al. 2003).

CLINICAL HISTORY

The attacks typically last for 2–7 days. They recur without obvious periodicity. Along with the fever, abdominal pain, diarrhea, and vomiting are common (Frenkel et al. 2001, Prietsch et al. 2003). Arthralgia is common. The attacks may be provoked by vaccinations as well. The severity of symptoms varies among patients and tends to be milder as the child grows (Drenth et al. 1999, Frenkel et al. 2001).

PHYSICAL EXAMINATION

Along with the fever hepatosplenomegaly and cervical lymphadenopathy are seen (Frenkel et al. 2001, Prietsch et al. 2003). Arthritis may be noted. Rashes are present in three-quarters of the patients (224).
LABORATORY
There is leukocytosis and increased acute-phase reactants during attacks. Serum IgD levels are usually elevated during attacks; however, this is not a prerequisite for the diagnosis and it may be elevated in other autoinflammatory diseases as well (Frenkel et al. 2001). On the other hand, as a result of the mevalonate kinase deficiency there is an increased excretion of urinary mevalonic acid and this may be a helpful differentiating feature.

IMAGING
Imaging may be useful to exclude other causes of periodic fevers.

DIFFERENTIAL DIAGNOSIS
A mutation analysis of the MVK gene is indicated in a patient (especially from Dutch or French ancestry) with recurring attacks of fever and pain not responding to colchicine, who also has cervical lymphadenopathy, and some rash that lasts less than a week. Diagnosis depends on characteristic features with high serum IgD and high urine mevalonate levels. For genetic confirmation mutations need to be shown on the MVK gene. All inherited periodic fevers and especially PFAPA syndrome should be considered in the differential diagnosis (Samuels & Ozen 2006). There is no response to colchicine in HIDS.

PROGNOSIS
The frequency of attacks decreases by age. Although the risk of amyloidosis seems to be low the patients suffer from a poor quality of life in the early years.

MANAGEMENT
We lack a high level of evidence for the treatment of HIDS. Anti-TNF and anti-IL-1 treatment has been reported to be effective in recent papers (Bodar et al. 2005). Statins have been tried; however, they did not have a significant effect and had to be stopped in some patients due to side effects (Simon et al. 2004b).

COMPLICATIONS
There has been only one case of amyloidosis.
Tumor necrosis factor receptor-associated periodic syndrome

DEFINITION
Mutations on chromosome 12 in TNFRSF1A, the gene encoding the 55-kDa receptor for TNF has been associated with the autosomal dominantly inherited disease, tumor necrosis factor receptor-associated periodic syndrome (TRAPS, MIM142680) (McDermott et al. 1999). Both the number of known mutations (more than 30) and the spectrum of affected ethnicities have considerably increased over the last years.

EPIDEMIOLOGY AND ETIOLOGY
Although first defined in Scottish and Irish individuals, patients have now been diagnosed all around the world and the syndrome can occur in any ethnic group (Ozen et al. 2006, Stojanov & Kastner 2005).

In healthy controls, TNF-α activates TNFRSF1A receptors and releases these receptors from the cell surface to be shed into the extracellular compartment. The unattached TNF receptors are free to bind TNF-α and reduce the cellular inflammatory response to it (McDermott et al. 1999). In most of the patients with TRAPS there is a shedding defect in the aforementioned receptor (Stojanov & McDermott 2005). However, recent findings suggest that the shedding hypothesis does not account for all cases. Thus a more complex mechanism may be involved in at least some of the patients (Ozen et al. 2006, Stojanov & Kastner 2005).

The pedigree is compatible with an autosomal dominant inheritance.

CLINICAL HISTORY
Patients have prolonged (greater than 1 week) attacks of fever associated with abdominal pain, and myalgia.

PHYSICAL EXAMINATION
Patients may have an erythematous skin rash (225a), conjunctivitis, and/or periorbital edema (Hull et al. 2002, McDermott 1999). Muscle involvement is a frequent feature in these patients, usually involving only one muscle group at a time. Myalgia has been shown to frequently result from monocytic fasciitis, but also from myositis or lymphocytic vasculitis.

LABORATORY
Acute-phase reactants are elevated as in other periodic fever syndromes.

IMAGING
Imaging may be useful to exclude other causes of periodic fevers.

DIFFERENTIAL DIAGNOSIS
A family history is almost always present. Mutation analysis for the TRAPS gene is indi-
cated in a patient who presents with attacks longer than a week with myalgia or myositis and migratory rashes and with affected cases in the pedigree.

**PROGNOSIS**
AA amyloidosis develops in 25–40% of patients with TRAPS, with the kidneys and liver being the most common complication.

**MANAGEMENT**
Understanding the etiology has brought up the logical use of TNF-α inhibition. Case reports and open-label studies confirmed anti-TNF treatment as a safer and more effective alternative to corticosteroid therapy with regard to the frequency, duration, and severity of attacks (Drewe et al. 2004). Etanercept may also reverse or slow the clinical progression of renal amyloidosis in TRAPS. However, this may not always be the case. For the rare cases when anti-TNF-α therapy does not work anti-IL-1 therapy may prove useful (Simon et al. 2004a).

**COMPLICATIONS**
Amyloidosis has been reported in these patients; 14% of those reported so far have developed amyloidosis (225b). Thus urinalysis needs to be checked at each visit. Amyloidosis has been associated with cysteine substitutions (Hull et al. 2002).
OTHER MONOGENIC AUTOINFLAMMATORY SYNDROMES

Pyogenic sterile arthritis, pyoderma gangrenosum, and acne syndrome

Pyogenic sterile arthritis, pyoderma gangrenosum, and acne (PAPA) syndrome is a very rare disease, characterized by early destruction of the joints (226–228) by recurrent effusions. The patients also experience infiltration of muscle and numerous dermatologic manifestations including cystic acne, expanding purulent ulcerating lesions (pyoderma gangrenosum), and sterile abscesses at injection sites (Lindor et al. 1997). It is mainly a white American disease (Stojanov & Kastner 2005). It is inherited in an autosomal dominant fashion.

The disease is caused by mutations in proline serine threonine phosphatase-interacting protein (PSTPIP1, or CD2-binding protein 1 (CD2BP1)), which is a tyrosine-phosphorylated protein involved in cytoskeletal organization, also interacting with pyrin (Shoham et al. 2003). Pyrin binds the PSTPIP1/CD2BP1 protein, defining FMF and PAPA syndrome as disorders in the same pathway. Shoham et al. have found increased IL-1β production by peripheral blood leukocytes from a patient with clinically active PAPA with the A230T PSTPIP1/CD2BP1 mutation, consistent with the hypothesis that these mutations exert a dominant-negative effect on the previously reported activity of pyrin (Shoham et al. 2003, Stojanov & Kastner 2005). The disease is mediated predominantly by granulocytes as in FMF.

While some patients benefit from corticosteroids many of them become resistant to therapy. Biologics (anti-TNF and anti-IL-1) have been tried with varying success in resistant patients (Samuels & Ozen 2006, Stojanov & Kastner 2005).

Early onset sarcoidosis (sporadic granulomatous arthritis), Blau syndrome (familial granulomatous arthritis), and Crohn disease

CARD15 is a caspase recruitment domain; CARD15 serves as an intracellular sensor of bacteria and is another member of the inflammatory pathway leading to the activation of IL-1β. It participates in an inflammatory signaling cascade triggering NF-κB (Rose & Martin 2005). CARD15 mutations have been shown in 40% of patients with Crohn disease, 50% of those with Blau syndrome, and 90% of patients early onset sarcoidosis (EOS). Thus mutations in this protein have been associated with the aforementioned granulomatous (auto-)inflammatory diseases (Stojanov & Kastner 2005).

Blau syndrome is an autosomal, dominantly inherited syndrome characterized by synovitis, uveitis, and rash (229–231). The histology of the synovium, conjunctiva, and dermis show the typical non-caseating granuloma (232). EOS has similar characteristics but lacks family history. It is now widely accepted that Blau syndrome and EOS are the same disease, with an inherited mutation in Blau syndrome and a new mutation EOS (Rose & Martin 2005). In fact, CARD15 mutations are present in 50% and 90% of Blau syndrome and EOS patients, respectively (Rose & Martin 2005).

CROHN DISEASE

Some patients with Crohn disease also share a mutated form of CARD15. It should be stressed that similar to Blau syndrome, non-caseating granulomas and arthritis may also be found in Crohn disease (Rose & Martin 2005). Arthritis is a feature of this IBD. Understanding more about the pathophysiology may enable us to have more specific therapeutic options in these diseases.
Other monogenic autoinflammatory syndromes

Cutaneous skin granulomas in Blau syndrome in adolescence (courtesy of Ann M. Reed, MD, Mayo Clinic). (230–232)

Destructive arthropathy in PAPA syndrome (courtesy of Ann M. Reed, MD, Mayo Clinic). (226–228)

Cutaneous skin granulomas in Blau syndrome at birth (courtesy of Ann M. Reed, MD, Mayo Clinic) (230, 231): bone granuloma in Blau syndrome (courtesy of Ann M. Reed, MD, Mayo Clinic) (232).
NON-BACTERIAL OSTEITIS

Chronic recurrent multifocal osteomyelitis (CRMO) and synovitis, acne, pustulosis, hyperostosis, and osteitis (SAPHO) syndrome are characterized by recurring attacks of bone pain and fever. The osteomyelitis may recur months or years later (233–236). There is an autosomal recessive syndromic form of the disease, called Majeed syndrome, in which the mutated gene is identified (Ferguson et al. 2006). A similar, autosomal, recessively inherited mouse model has been identified where there is bone, cartilage, and skin inflammation. In these mice the mutated gene is PSTPIP2, which shares significant sequence homology to the PSTPIP1 (the mutated gene of PAPA syndrome) (Ferguson et al. 2006).

233–236 Radiograph of patient with CRMO (courtesy of Ann M. Reed, MD, Mayo Clinic) (233); bone scan of patient with CRMO (courtesy of Ann M. Reed, MD, Mayo Clinic) (234); clavicular osteomyelitis in a patient with SAPHO (courtesy of Ann M. Reed, MD, Mayo Clinic) (235); skin findings in SAPHO (236).
DEFICIENCY OF THE INTERLEUKIN-1-RECEPTOR ANTAGONIST
Deficiency of the IL-1-receptor antagonist (DIRA) presents in the neonatal period and is characterized by pustulosis, periostitis, and sterile osteomyelitis (237–240). The illness arises from mutations in the \textit{IL1RN} gene, which results in a non-functional IL-1-receptor antagonist, causing cellular hypersensitivity to the proinflammatory cytokine interleukin-1 (Aksentijevich \textit{et al.} 2009, Reddy \textit{et al.} 2009).

\textbf{237–240} Pustulosis in an infant with DIRA (237); skin erythema in an infant with DIRA (238); osteitis in an infant with DIRA (239); osteitis of the sixth rib in an infant with DIRA (240).
Autoinflammatory syndromes that are not monogenic/autoinflammatory diseases with complex genetic traits

The spectrum of autoinflammatory syndromes has now been enlarged to include a number of complex genetic syndromes. For example, Behçet syndrome has been suggested to be an autoinflammatory disease since it is characterized by remissions and exacerbations and by an acute-phase response without autoantibodies (Ozen et al. 2006). Again, at least a portion of systemic juvenile idiopathic arthritis patients with the high IL-1 levels may be representative of this group. In both of these diseases secondary amyloidosis develops similar to the group of aforementioned monogenic diseases. The group of diseases to be included in this section may expand in future. These diseases do have an autoinflammatory element; however, they have their differentiating features and they require immunosuppressive treatment unlike the typical monogenic diseases.

In conclusion, the autoinflammatory diseases have enlightened our understanding of the pathway of inflammation and innate immunity. Further developments are expected to provide insight into other rheumatic diseases and enable us to manage and treat these patients in a more proper fashion.
2. JUVENILE IDIOPATHIC ARTHRITIS


### 3. SPONDYLOARTHROPATHIES AND REACTIVE ARTHROPATHIES


Lupus erythematosus

4. LUPUS ERYTHEMATOSUS


Buyon JP, Waltuck J, Caldwell K, et al. (1994). Relationship between maternal and neonatal levels of antibodies to 48 kDa SSB(La), 52 kDa SSA(Ro), and 60 kDa SSA(Ro) in pregnancies complicated by congenital heart block. *J Rheumatol*, 21: 1943–50.


5. IDIOPATHIC INFLAMMATORY MYOPATHIES


6. VASCULITIS


Cardiovascular Disease in the Young, American Heart Association. Pediatrics, 114: 1708–33.


7. SCLERODERMA IN CHILDREN


Galie N, Ghofrani HA, Torbicki A, et al.; for Sildenafil Use in Pulmonary Arterial Hypertension (SUPER) Study Group


8. AUTOINFLAMMATORY DISEASES


Abbreviations

AAV  ANCA-associated vasculitide
ACA  anti-centromere antibodies
ACLE acute cutaneous lupus erythematosus
ACR  American College of Rheumatology
AHA  American Heart Association
ANA  antinuclear antibody
ANCA  anti-neutrophil cytoplasmic antibodies
AoAS  adult-onset ankylosing spondylitis
AP  anterior–posterior (view)
APL  anti-phospholipid
aPTT  activated partial thromboplastin time
ARF  acute rheumatic fever
AS  ankylosing spondylitis
ASO  anti-streptolysin O
BASFI  Bath Ankylosing Spondylitis Functional Index
BCG  Bacillus Calmette–Guérin
BUN  blood urea nitrogen
CAPS  cryopyrin-associated periodic fever syndromes
CARD15  caspase recruitment domain-15
CBC  complete blood count
CCLE chronic cutaneous lupus erythematosus
CCP  cyclic citrullinated peptide
CHAQ  Childhood Health Assessment Questionnaire
CHB  congenital heart block
CINCA  chronic infantile neurologic cutaneous articular (syndrome)
CLE cutaneous lupus erythematosus
CMAS  Childhood Myositis Assessment Scale
CNS  central nervous system
CRMO  chronic recurrent multifocal osteomyelitis
CRP  C-reactive protein
CSF  cerebrospinal fluid
CSS  Churg–Strauss syndrome
CT  computed tomography
DIP  distal interphalangeal
DLCO  diffusing capacity of the lung for carbon monoxide
DLE  discoid lupus erythematosus
DMARD  disease-modifying antirheumatic drug
DNase B  deoxyribonuclease B
DPT  diphtheria, pertussis and tetanus
dsDNA  double-stranded DNA
ECDS  en coup de sabre
ECG  electrocardiogram
ELE  erysipelas-like erythema
EMG  electromyography
EN  erythema nodosum
EOS  early onset sarcoidosis
ESR  erythrocyte sedimentation rate
ESSG  European Spondyloarthropathy Study Group
FCAS  familial cold autoinflammatory syndrome
FMF  familial Mediterranean fever
GABHS  group A β-hemolytic streptococcal
gastroesophageal reflux
GI  gastrointestinal
GU  genitourinary
HIIDS  hyperimmunoglobulinemia D with periodic fever syndrome
HLA  human leukocyte antigen
HLH  hemophagocytic lymphohistiocytosis
HRCT  high-resolution computed tomography
IBD  inflammatory bowel disease
IBP  inflammatory back pain
Ig  immunoglobulin
IL  interleukin
ILAR  International League of Associations for Rheumatology
ISN/RPS  International Society of Nephrology/Renal Pathology Society
IV  intravenous
IVIG  intravenous immunoglobulin
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tbody>
<tr>
<td>JAS</td>
<td>juvenile ankylosing spondylitis</td>
</tr>
<tr>
<td>JDM</td>
<td>juvenile dermatomyositis</td>
</tr>
<tr>
<td>JIA</td>
<td>juvenile idiopathic arthritis</td>
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<tr>
<td>JLS</td>
<td>juvenile localized scleroderma</td>
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<tr>
<td>JoAS</td>
<td>juvenile-onset ankylosing spondylitis</td>
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<tr>
<td>JPM</td>
<td>juvenile polymyositis</td>
</tr>
<tr>
<td>JPsa</td>
<td>juvenile psoriatic arthritis</td>
</tr>
<tr>
<td>JRA</td>
<td>juvenile rheumatoid arthritis</td>
</tr>
<tr>
<td>JSpA</td>
<td>juvenile spondyloarthropathy</td>
</tr>
<tr>
<td>JSSc</td>
<td>juvenile systemic sclerosis</td>
</tr>
<tr>
<td>KD</td>
<td>Kawasaki disease</td>
</tr>
<tr>
<td>LE</td>
<td>lupus erythematosus</td>
</tr>
<tr>
<td>LSsc</td>
<td>localized scleroderma</td>
</tr>
<tr>
<td>MAS</td>
<td>macrophage activation syndrome</td>
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<tr>
<td>MCP</td>
<td>metacarpophalangeal</td>
</tr>
<tr>
<td>MCTD</td>
<td>mixed connective tissue disease</td>
</tr>
<tr>
<td>MEFV</td>
<td>Mediterranean fever (gene)</td>
</tr>
<tr>
<td>MMR</td>
<td>measles, mumps, and rubella</td>
</tr>
<tr>
<td>MPA</td>
<td>microscopic polyangiitis</td>
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<tr>
<td>MPO</td>
<td>myeloperoxidase</td>
</tr>
<tr>
<td>MR</td>
<td>magnetic resonance</td>
</tr>
<tr>
<td>MRI</td>
<td>magnetic resonance imaging</td>
</tr>
<tr>
<td>MTX</td>
<td>methotrexate</td>
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<tr>
<td>MWS</td>
<td>Muckle–Wells syndrome</td>
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<tr>
<td>NF-κB</td>
<td>nuclear factor kappa B</td>
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<tr>
<td>NK</td>
<td>natural killer</td>
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<tr>
<td>NLS</td>
<td>neonatal lupus syndrome</td>
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<tr>
<td>NOMID</td>
<td>neonatal-onset multisystem inflammatory disease</td>
</tr>
<tr>
<td>NSAID</td>
<td>non-steroidal anti-inflammatory drug</td>
</tr>
<tr>
<td>OT</td>
<td>occupational therapy (therapist)</td>
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<tr>
<td>PACNS</td>
<td>primary angiitis of the central nervous system</td>
</tr>
<tr>
<td>PAN</td>
<td>polyarteritis nodosa</td>
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<tr>
<td>PAPA</td>
<td>pyogenic sterile arthritis, pyoderma gangrenosum, and acne</td>
</tr>
<tr>
<td>PET</td>
<td>positron emission tomography</td>
</tr>
<tr>
<td>PFAPA</td>
<td>periodic fevers with aphthous stomatitis, pharyngitis, and adenitis</td>
</tr>
<tr>
<td>PIP</td>
<td>proximal interphalangeal</td>
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<tr>
<td>PsA</td>
<td>psoriatic arthritis</td>
</tr>
<tr>
<td>PSRA</td>
<td>post-streptococcal reactive arthritis</td>
</tr>
<tr>
<td>PT</td>
<td>prothrombin time</td>
</tr>
<tr>
<td>PTT</td>
<td>partial thromboplastin time</td>
</tr>
<tr>
<td>RANTES</td>
<td>regulated on activation, normal T-cell expressed and secreted</td>
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<tr>
<td>ReA</td>
<td>reactive arthritis</td>
</tr>
<tr>
<td>RF</td>
<td>rheumatoid factor</td>
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<tr>
<td>RNP</td>
<td>ribonuclear protein</td>
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<tr>
<td>ROM</td>
<td>range of motion</td>
</tr>
<tr>
<td>SAA</td>
<td>serum amyloid A</td>
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<tr>
<td>SCLE</td>
<td>subacute cutaneous lupus erythematosus</td>
</tr>
<tr>
<td>SI</td>
<td>sacroiliac</td>
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<tr>
<td>SLE</td>
<td>systemic lupus erythematosus</td>
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<tr>
<td>SSc</td>
<td>systemic sclerosis</td>
</tr>
<tr>
<td>STIR</td>
<td>short tau inversion recovery (MRI)</td>
</tr>
<tr>
<td>TMJ</td>
<td>temporomandibular joint</td>
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<tr>
<td>TNF-α</td>
<td>tumor necrosis factor-α</td>
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<tr>
<td>TRAPS</td>
<td>tumor necrosis factor receptor-associated periodic syndrome</td>
</tr>
<tr>
<td>URI</td>
<td>upper respiratory infection</td>
</tr>
<tr>
<td>UTI</td>
<td>urinary tract infection</td>
</tr>
<tr>
<td>UV</td>
<td>ultraviolet</td>
</tr>
<tr>
<td>WBC</td>
<td>white blood cell</td>
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<tr>
<td>WG</td>
<td>Wegener granulomatosis</td>
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