MR Imaging of Atraumatic Muscle Disorders

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Atraumatic disorders of skeletal muscles include congenital variants; inherited myopathies; acquired inflammatory, infectious, or ischemic disorders; neoplastic diseases; and conditions leading to muscle atrophy. These have overlapping appearances at magnetic resonance (MR) imaging and are challenging for the radiologist to differentiate. The authors organize muscle disorders into four MR imaging patterns: (a) abnormal anatomy with normal signal intensity, (b) edema/inflammation, (c) mass, and (d) atrophy, highlighting each of their key clinical and imaging findings. Anatomic muscle variants, while common, do not produce signal intensity alterations and therefore are easily overlooked. Muscle edema is the most common pattern but is nonspecific, with a broad differential diagnosis. Autoimmune, paraneoplastic, and drug-induced myositis tend to be symmetric, whereas infection, radiation-induced injury, and myonecrosis are focal asymmetric processes. Architectural distortion in the setting of muscle edema suggests one of these latter processes. Intramuscular masses include primary neoplasms, metastases, and several benign masslike lesions that simulate malignancy. Some lesions, such as lipomas, low-flow vascular malformations, fibromatoses, and subacute hematomas, are distinctive, but many intramuscular masses ultimately require a biopsy for definitive diagnosis. Atrophy is the irreversible end result of any muscle disease of sufficient severity and is the dominant finding in disorders such as the muscular dystrophies, denervation myopathy, and sarcopenia. This imaging-based classification, in correlation with clinical and laboratory data, will aid the radiologist in interpreting MR imaging findings in patients with atraumatic muscle disorders.

Introduction

Atraumatic skeletal muscle disorders encompass a diverse group of conditions: anatomic variants; congenital, metabolic, acquired, inflammatory, autoimmune, infectious, and ischemic conditions; and benign and malignant masses. Conventional radiography and computed tomography (CT) are useful for detecting intramuscular mineralization, gas, and radiopaque foreign bodies, but they lack the sensitivity for detecting the majority of muscle disorders. Ultrasonography (US) affords high spatial resolution of muscle but is less sensitive than magnetic resonance (MR) imaging for mild edema and early myopathy. US plays an important role in the evaluation of masses, particularly for recognizing muscle variants, distinguishing cystic from solid neoplasms, assessing lesion vascularity, and guiding percutaneous interventions such as aspiration and biopsy. Muscle biopsy can be performed openly or percutaneously, with a greater than 95% success with both methods in obtaining...
Despite the sensitivity of MR imaging for detecting altered muscle signal intensity and architecture, arriving at a specific diagnosis is often challenging for radiologists, as there is a limited response of muscle to insult, resulting in substantial overlap in the imaging appearances of atraumatic muscle disorders. Correlation with clinical and laboratory findings, other imaging modalities, and, occasionally, tissue sampling is necessary before a diagnosis can be made.

Accessory muscles are usually asymptomatic and are detected incidentally at imaging, but they may manifest clinically as a painless palpable mass or cause symptoms related to compression of nearby neurovascular structures.

Edema is the most common pattern of muscle abnormality at MR imaging, appearing as increased signal intensity on T2-weighted or STIR images. T2 mapping affords quantitative assessment of muscle signal intensity and is more sensitive for mild degrees of muscle edema. Correlation with history is especially important in cases of atraumatic muscle edema. Dividing edema into symmetric or asymmetric patterns helps narrow the differential diagnosis: symmetric edema is typical of inflammatory and drug-related myopathies, whereas infection, radiation, myonecrosis, and compartment syndrome tend to be asymmetric and often focal. Additional findings such as architectural distortion of the tissues, intramuscular fluid collections, and extramuscular edema are also helpful.

Intramuscular masses include malignant or benign primary neoplasms, metastatic disease, and nonneoplastic mass-like lesions related to infection, trauma, inflammation, or other underlying disorders. The differential diagnosis of an intramuscular mass depends on the patient’s age, pertinent clinical history, location of the mass, and its imaging appearance. A few intramuscular masses demonstrate unique MR imaging characteristics, allowing a specific diagnosis.

Atrophy is characterized by loss of muscle tissue that manifests as decreased muscle size, often associated with fatty infiltration. It is the irreversible end stage of numerous muscle diseases, including focal disorders such as trauma or denervation, as well as generalized disorders that are initially inflammatory. Atrophy is the principal finding in hereditary degenerative myopathies that produce disseminated muscle tissue loss without a prodromal inflammatory phase. Atrophy related to aging is referred to as sarcopenia, which is increasingly recognized as a substantial predictor of overall health. Atrophy is an important finding predicting loss of muscle power and function.

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Patterns of Muscle Abnormality
Muscle disorders typically manifest as one of four MR imaging patterns: (a) abnormal anatomy with normal signal intensity, (b) edema/inflammation related to increases in water content, (c) intramuscular mass causing anatomic distortion, or (d) atrophy resulting in tissue loss, usually accompanied by fatty replacement.
matic fascial weakening, and surgical procedures altering myotendinous anatomy can also produce a similar pattern of architectural abnormality without signal intensity alterations (Fig 3). Recognition of anatomic variations can be challenging at MR imaging unless one is thoroughly familiar with normal regional anatomy. Fiducial markers are helpful for identifying this category of muscle abnormalities. Congenital absence of a small extremity muscle is often asymptomatic, whereas absence of a critical muscle involved in fine motor control (eg, ocular muscle) or a large muscle (eg, pectoralis) may result in dysfunction or deformity, respectively. In cases of accessory and anomalous muscles, T1-weighted images best depict the muscular architecture of the “lesion,” while T2-weighted images are useful to help confirm muscle signal intensity.

Pattern 1: Abnormal Anatomy with Normal Signal Intensity
Congenital muscle variants include absence of a muscle or muscle group; presence of an accessory (supernumerary) muscle; and deviations of size, course, insertion, or origin of a normally present (anomalous) muscle (11). Muscle herniation, which can be caused by congenital or posttraumatic fascial weakening, and surgical procedures
of individuals, typically involving the hands and feet (11,12). Accessory muscles are usually asymptomatic and are detected incidentally at imaging, but they may manifest clinically as a painless palpable mass or cause symptoms related to compression of nearby neurovascular structures. The extensor digitorum brevis manus muscle, located dorsally at the level of the carpometacarpal joint, typically arises from the dorsal wrist capsule and inserts on the index or middle finger (11). It commonly manifests as a painless mass overlying the carpus, often mistaken clinically for a ganglion cyst or carpal boss (13).

The accessory soleus muscle also commonly manifests as a mass, typically in adolescents or young adults, possibly related to increased muscle mass and physical activity at these ages (14). It arises from the fibula, soleal line of the tibia, or anterior soleus muscle and traverses through the pre-Achilles fat pad to insert on the superior or medial calcaneus (15) (Fig 4). Accessory muscles can lead to nerve compression when they overlie a nerve within a fibro-osseous tunnel. Compression of the ulnar nerve at the cubital tunnel by an accessory anconeus epitrochlearis muscle, the median or ulnar nerves at the wrist by palmar accessory muscles, and the tibial nerve by an accessory flexor muscle at the tarsal tunnel are all well recognized (11,15) (Fig 5).

**Anomalous Muscle**

Numerous anomalies of the course, insertion, or origin of muscles have been described. The majority are asymptomatic, but they can occasionally result in clinical symptoms, similar to those of accessory muscles. Anomalous muscles from
the forearm or palm extending into the carpal tunnel can cause median nerve compression. An anomalous plantaris insertion on the lateral patellar retinaculum can cause iliotibial band syndrome or result in patellar maltracking (16). The medial and lateral gastrocnemius muscles may have anomalous insertions that can cause popliteal artery entrapment, manifesting as diminished pulses with ankle movement or intermittent claudication with exertion (11) (Fig 6).

**Pattern 2: Edema/Inflammation**

Edema is the most common pattern of muscle abnormality at MR imaging, appearing as increased signal intensity on T2-weighted or STIR images. T2 mapping affords quantitative assessment of muscle signal intensity and is more sensitive for mild degrees of muscle edema (Fig 7). Correlation with history is especially important in cases of atraumatic muscle edema.

Dividing edema into symmetric or asymmetric patterns helps narrow the differential diagnosis: symmetric edema is typical of inflammatory and drug-related myopathies, whereas infection, radiation, myonecrosis, and compartment syndrome tend to be asymmetric and often focal (Table 2). Additional findings such as architectural distortion of the tissues, intramuscular fluid collections, and extramuscular edema are also helpful (1). While it is difficult to establish a specific diagnosis in many cases of muscle edema, MR imaging is still helpful for localizing active disease to guide biopsy (2,9,17).

**Idiopathic Inflammatory Myositis**

The idiopathic inflammatory myopathies are a group of chronic autoimmune disorders that result in muscle weakness and inflammation, primarily affecting proximal extremity muscles. These disorders have overlapping clinical and imaging features, and the diagnostic criteria and classification systems are continually evolving (18,19). The original classification proposed by Bohan et al (20) divided inflammatory myopathies into five types: (a) adult dermatomyositis, (b) adult polymyositis, (c) juvenile myositis, (d) myositis associated with cancer, and (e) myositis associated with connective tissue disease (overlap syndrome). Subsequently, other forms of autoimmune myositis, such as necrotizing autoimmune myopathy and inclusion body myositis, have been included (18). Current classification systems no longer distinguish between the adult and juvenile forms of these disorders (18).

**Polymyositis and Dermatomyositis.**—The best-known inflammatory myopathies are polymyositis and dermatomyositis. Both initially involve the proximal lower limb muscles, particularly the hamstrings, before ascending to involve the buttocks,
proximal upper limbs, neck flexors, and pharyngeal muscles (18,21). MR imaging shows symmetric edema with preserved muscle architecture, typically associated with considerable subcutaneous and perifascial thickening and fluid (21,22). Muscle edema can be patchy, inhomogeneous, and variegated, even within an individual muscle (1,21) (Fig 8).

In dermatomyositis, amorphous or sheetlike muscle and fascial calcifications may also manifest and are often difficult to appreciate at MR imaging. Edema correlates roughly with disease activity and may resolve with anti-inflammatory medications (22). At the chronic/burned-out stage, MR imaging shows nonspecific patchy symmetric muscle atrophy without inflammation (Fig 9). Whole-body MR imaging can potentially help assess disease burden and activity and reveal atypically involved sites, such as distal muscles (21).

**Overlap with Connective Tissue Disease.**—Autoimmune connective tissue disorders such as scleroderma, systemic lupus erythematosus (SLE), Sjögren syndrome, rheumatoid arthritis, mixed connective tissue disease, and overlap syndromes can be complicated by inflammatory myopathy (24,25). Muscle involvement is most common in scleroderma or in an overlap syndrome that includes scleroderma, affecting 5%–96% of such patients, portending a poor prognosis. Up to 28% of patients with rheumatoid arthritis or SLE demonstrate myositis with muscle necrosis and inflammation of connective tissues (25–27). At MR imaging, the myositis related to collagen vascular disorders is indistinguishable from polymyositis (25).

**Inclusion Body Myositis.**—Inclusion body myositis is a common myopathy in older patients that is diagnosed histologically by distinct inclusion bodies in the nuclei and cytoplasms of affected muscle cells. It develops insidiously, often without substantially elevated inflammatory markers and muscle enzymes. Unlike conventional inflammatory myositis, inclusion body myositis can be asymmetric and often affects the pharyngeal muscles, upper extremities, and distal musculature early in the course of the disease (18). The dominant finding is prominent patchy muscle atrophy without substantial inflammation or perifascial edema (2,9) (Fig 10).

**Drug-related Myopathy**

Muscle can be exposed to drugs systemically or directly through inoculation, resulting in diffuse...
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<td>Dermatomyositis</td>
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<td>Proximal lower limbs</td>
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<td>Dependent on the site of the radiation portal</td>
<td>Normal</td>
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<td>Normal (acute) or atrophy (subacute and chronic)</td>
<td>Mild uniform edema (acute and subacute)</td>
<td>Normal</td>
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<td>No enhancement</td>
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<td>Pyomyositis</td>
<td>Can be multifocal, if immunocompromised</td>
<td>Thighs</td>
<td>High-signal-intensity rim with abscesses</td>
<td>Edema</td>
<td>Fluid collections</td>
<td>Extensive edema</td>
<td>Rim enhancement of abscesses</td>
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<td>Acute compartment syndrome (ACS)</td>
<td>Surgical emergency</td>
<td>Lower leg</td>
<td>Mild hyperintensity</td>
<td>Edema</td>
<td>Normal</td>
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<tr>
<td>Chronic compartment syndrome</td>
<td>Can manifest as a mass</td>
<td>Lower leg</td>
<td>Low-signal-intensity calcifications</td>
<td>Central fluid collection</td>
<td>Peripherally calcified mass</td>
<td>Variable</td>
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<td>Diabetic myonecrosis</td>
<td>Poorly controlled diabetes</td>
<td>Anterior thigh or calf; can be multifocal</td>
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<td>Edema</td>
<td>Fluid collections</td>
<td>Perifascial edema</td>
<td>Nonenhancement of necrotic areas</td>
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or local tissue damage, respectively. Dysfunction related to systemic drug administration affects multiple muscles bilaterally, causing muscle weakness, myopathy, inflammation, and, in extreme cases, rhabdomyolysis.

Imaging findings are nonspecific, consisting of symmetric increased muscle size and edema, typically affecting the buttocks, quadriceps, adductors, and deep calf muscles (28). The most common drugs associated with myopathy are the HMG-CoA reductase inhibitors (statins), used for treating dyslipidemia. Statin-induced muscle pain and weakness develop in up to 5% of treated patients (28). This mild form, referred to as toxic statin myopathy, is dose dependent and resolves with cessation of the drug. A small subgroup, approximately one in 10,000 patients treated with statins, develops severe myositis that may persist or progress even after discontinuation of the drug. This form, known as autoimmune necrotizing myopathy, can progress to rhabdomyolysis and requires immunosuppressive therapy (29,30). Substantial creatine kinase level elevation and autoantibodies directed

Figure 8. Dermatomyositis in a 50-year-old man with bilateral lower extremity weakness. (a) Axial T1-weighted MR image shows no substantial distortion of the muscle architecture. (b) Axial fat-suppressed T2-weighted MR image shows relatively symmetric muscle edema, with subcutaneous and perifascial fluid (arrows) in both thighs. There is involvement of multiple muscle compartments, but it is patchy, with relative sparing of the rectus femoris and portions of the hamstring musculature. (c) Axial fluorine 18 fluorodeoxyglucose (FDG)-positron emission tomography (PET)/CT image shows matching increased activity in the inflamed musculature. The results of a muscle biopsy disclosed dermatomyositis.

Figure 9. Symmetric patchy atrophy in a 55-year-old woman with a long-standing history of inflammatory myositis, currently in remission. The patient had been variably diagnosed with either polymyositis or myositis associated with connective tissue disorder. Axial T1-weighted MR image of the proximal thighs shows bilateral symmetric atrophy of the pelvic and thigh musculature. The atrophy is somewhat patchy, most prominently at the gluteus maximus, vastus lateralis, and lateral portion of the rectus femoris muscles, with relative sparing of the adductor muscles. Symmetric patchy atrophy is a common sequela of all forms of long-standing inflammatory myositis.

Figure 10. Inclusion body myositis in a 71-year-old man who presented with slowly progressive bilateral leg weakness. Axial fat-suppressed T2-weighted MR image of both thighs shows symmetric increased signal intensity in the quadriceps muscles bilaterally. There is no substantial associated muscle atrophy or subcutaneous edema. The results of a subsequent muscle biopsy showed scattered subsarcolemmal vacuoles with blue granular rims, findings typical of inclusion body myositis.
against HMG-CoA reductase are identified in such patients. The exact mechanism of muscle damage may relate to this autoantibody, other soluble factors, aberrant drug processing, or infiltration by immune cells (29,30).

Antiretroviral drugs (eg, azidothymidine), corticosteroids, calcineurin inhibitors, vitamin metabolites, anabolic agents, and a host of other medications can also lead to myopathy (31). Azidothymidine myopathy develops in up to 6% of patients with HIV infection, related to drug interaction with mitochondrial DNA polymerase; it differs from pyomyositis by being symmetric, without intramuscular fluid collections (32). Drug-induced muscle disease is a diagnosis of exclusion. A temporal link between symptoms and initiation of drug therapy, or improvement following tapering/cessation of the drug, suggests this diagnosis (33). Illicit agents, such as cocaine, can result in extensive myonecrosis related to arterial vasoconstriction or direct muscle toxicity; such patients may not disclose their drug use (34) (Fig 11).

HIV-related Myopathy

The pathophysiology of HIV-related myopathies is multifactorial, related to the virus, its components, and the host’s altered immune response (35). Patients with HIV infection are at increased risk for infectious pyomyositis, drug-induced myopathy, and polymyositis (35–37). While the increased frequency of infection is well recognized in immunocompromised patients, noninfectious inflammatory polymyositis is actually more common than pyomyositis in patients with HIV infection, resulting in proximal muscle weakness and elevated creatine kinase levels. Inflammatory polymyositis may be the sole manifestation of HIV infection or appear years after clinically apparent immunodeficiency (36,37). Other less common HIV-related myopathies include rhabdomyolysis, necrotizing noninflammatory myopathy, nemaline rod myopathy, subclinical myopathy, and denervation caused by HIV-related peripheral neuropathy (36,38).

Muscle Infection/Pyomyositis

Pyomyositis, an acute bacterial infection of muscle, can occur by hematogenous dissemination, spread from adjacent osteomyelitis or septic arthritis, or direct inoculation through a soft-tissue defect (Fig 12). Infection risk is increased in muscle that is compromised by trauma, surgery, malnutrition, or ischemia (39–41). While originally considered a tropical disease of children and young adults, it is commonly associated with diabetes, intravenous drug use, and an immunocompromised state and can be multifocal or bilateral in up to 40% of such patients (2,41). *Staphylococcus aureus* is implicated in up to 90% of cases, but atypical and multiorganism infections do occur, particularly in immunocompromised patients (39,42). The thigh muscles are most commonly affected, particularly the quadriceps, followed by the gluteal and psoas muscles (40,41).

MR imaging demonstrates increased volume of the affected muscles on T1-weighted images; T2-weighted images show increased muscle signal intensity related to fluid and pus. Images obtained after the administration of contrast media often reveal small fluid collections related to myonecrosis or focal fluid accumulation. Large areas of
nonenhancement typically represent an abscess, the hallmark of pyomyositis (41) (Fig 13). Abscesses are usually hypointense or isointense on T1-weighted images and hyperintense on T2-weighted images. Intralosional gas may manifest. T1-weighted images may show a thin peripheral rim of increased signal intensity, referred to as the “penumbra sign” (35,42–44). There is typically extensive inflammation of the adjacent tissues, which helps distinguish an abscess from a soft-tissue tumor (41).

Cysticercosis
Cysticercosis is a parasitic infection that is caused by *Taenia solium* and is common in developing countries, where persons are infected by eating improperly cooked pork (45,46). The central nervous system is the most common site of symptomatic disease; when cysticercosis affects muscle, there is typically concomitant neural involvement (45,46). Muscle involvement by cysticercosis can be asymptomatic or manifest as myalgia, a solitary mass, or, most commonly, multiple intramuscular cysts (46) (Fig 14). The appearance of intramuscular lesions varies as the disease progresses through its vesicular, colloid, granular-nodular, and calcific stages (47). The early stages, before calcification, are nonspecific, particularly when there is a solitary lesion (45,46). The late calcific stage is distinctive, showing multiple small ovoid calcifications oriented parallel to the muscle fibers.

Radiation Therapy
Radiation therapy results in muscle edema caused by radiation-induced myositis, as well as intermuscular septal edema related to radiation-induced vasculitis (48). Signal intensity alterations at MR

Figure 13. Calf abscess in a 30-year-old man with endocarditis of a bicuspid aortic valve, with septic emboli related to *S aureus* vegetations. (a) Axial fat-suppressed PD-weighted MR image shows pyomyositis of the anterior and deep posterior compartments of the lower leg, with an intramuscular abscess of the soleus muscle. (b) Axial contrast-enhanced fat-suppressed T1-weighted MR image shows a small area of nonenhancing fluid, with a thin rim of enhancement, compatible with an abscess (arrows). The patient underwent incision and drainage.

Figure 14. Disseminated muscle involvement related to cysticercosis in an adult man. Coronal fat-suppressed T2-weighted MR image of the posterior chest and abdomen shows innumerable subcentimeter hyperintense lesions, oriented parallel to the muscle fibers in all visible muscles. (Courtesy of Travinder Buxi, MD, Ganga Ram Hospital, Delhi, India.)

Figure 15. Radiation-induced myositis in a 34-year-old man who underwent treatment for metastatic disease to the right femur, related to a primary malignant nerve sheath tumor of the ipsilateral ankle. Axial STIR MR image of the right midthigh shows a sharp bandlike demarcation (white line) between the abnormal irradiated muscle (predominantly of the anterior right thigh) and the normal muscle more posteriorly. Note that the demarcation is independent of any normal compartment boundaries.
imaging peak 6–12 months after therapy but may persist for years (9,48). Radiation-induced changes should be considered whenever there is a sharp bandlike demarcation between abnormal and normal muscle, particularly if this interface is within a single muscle (9,49) (Fig 15). Uncommonly, irradiated muscle can infarct, resulting in distorted enhancement and liquefaction. Rarely, irradiated muscle can exhibit a phenomenon known as radiation recall, whereby a drug, typically a chemotherapeutic agent, generates an acute inflammatory reaction at a prior site of radiation years after therapy (50). Over time, irradiated muscles can atrophy and undergo fibrosis.

Myonecrosis/Rhabdomyolysis

Myonecrosis refers to infarction of muscle, typically followed by tissue liquefaction. Rhabdomyolysis refers to myonecrosis complicated by myoglobinemia, which can lead to disseminated intravascular coagulation, acute renal failure, or even death (51,52). Muscle damage of sufficient severity from any cause can result in myonecrosis. Causes include compartment syndrome, diabetes, trauma, exercise, heatstroke, radiation, infection, metabolic disorders, seizures, envenomation, toxins, and medicinal or illicit drug use (51,53). Clinical findings can be nonspecific, and muscle pain may not be a prominent feature. T1-weighted images show mild hyperintensity owing to hemorrhage or protein concentration, while T2-weighted images demonstrate patchy hyperintense signal owing to a combination of edema, necrosis, and hemorrhage. There are two appearances seen following contrast medium administration: type 1 demonstrates nonspecific homogeneous edema and enhancement, reflecting the initial stages of myonecrosis before liquefaction; whereas type 2 (“stipple” sign) shows inhomogeneity with foci with low signal intensity within demarcated regions of nonenhancement, representing either residual viable muscle or inflammatory vessels (54,55). However, the use of a contrast medium is precluded owing to acute renal failure in at least 13% of patients with rhabdomyolysis (51).

Acute Compartment Syndrome.—Elevated compartmental pressure related to trauma, compression, overexertion, infection, or neoplasm can cause abnormal pressure gradients between the vasculature and the muscle, resulting in ACS (2,56) (Fig 16). ACS most commonly affects the
lower leg but also occurs in the thighs, forearms, and paraspinal musculature. ACS is diagnosed clinically, confirmed by intracompartmental pressure measurements, and requires prompt surgical decompression to prevent myonecrosis. Imaging plays a limited role in diagnosis when the clinical signs of arterial insufficiency (pain, paresthesia, pallor, and pulselessness) have already manifested (57). If MR imaging is performed because the diagnosis is not suspected, it demonstrates normal or increased signal intensity on T1-weighted images owing to hemorrhage and increased signal intensity on T2-weighted images owing to interstitial edema. Contrast-enhanced MR images best illustrate the perfusion failure within an affected compartment.

Chronic compartment syndrome may result in atrophy, fibrosis, calcification, or, rarely, masslike lesions referred to as calcific myonecrosis. Calcific myonecrosis is an unusual condition, typically affecting the calf, in which a chronic painless fusiform mass develops where liquefied necrotic muscle is surrounded by peripheral calcifications (58). MR imaging shows a heterogeneous mass on T2-weighted images, with a fluid-filled nonenhancing center following contrast medium administration (Fig 17). The peripheral calcifications are more apparent at radiography and CT.

**Diabetic Rhabdomyolysis.**—Myonecrosis/rhabdomyolysis is an uncommon complication of poorly controlled type 1 or 2 diabetes (52,59,60). Patients present with abrupt severe pain and swelling, typically at the anterior thigh or posterior calf muscles (59). Fever manifests in 10% of cases, making diabetic rhabdomyolysis difficult to distinguish clinically from infection. Diabetic

**Figure 17.** ACS of the lateral compartment of the left lower leg in a 48-year-old woman with hereditary angioneurotic edema, resulting in repetitive episodes of compartment syndrome. (a) Axial fat-suppressed PD-weighted MR image shows marked swelling and diffuse increased signal intensity involving the peroneus longus and brevis muscles (arrows). (b) Frontal radiograph of the left lower leg obtained 2 years following the last acute episode shows calcification corresponding to the lateral compartment, indicating that the tissues have undergone myonecrosis. (c) Corresponding axial fat-suppressed PD-weighted MR image shows regression of muscle edema, with formation of a masslike area with low signal intensity and a hypointense periphery (arrows), representing peripheral calcifications, enclosing muscle that has undergone liquefactive necrosis, consistent with calcific myonecrosis.

**Figure 18.** Rhabdomyolysis in a 52-year-old woman with a history of poorly controlled diabetes who presented with acute right thigh pain. Axial T1-weighted (a) and contrast-enhanced fat-suppressed T1-weighted (b) MR images of the right thigh show a well-margined area of nonenhancement related to muscle infarction (arrows) at the epicenter of the muscle inflammation. The slightly higher T1 signal intensity within the lesion is likely due to hemorrhage. This appearance is indistinguishable from a muscle abscess. The results of a needle aspiration confirmed rhabdomyolysis.
rhabdomyolysis is frequently multifocal and bilateral, with up to 40% of patients showing alterations of multiple muscle compartments. At T1-weighted imaging, diabetic myonecrosis results in muscle enlargement, effacement of intramuscular fatty septa, and occasionally hemorrhagic necrosis; perifascial fluid and increased muscle signal intensity are seen on T2-weighted images.

At imaging after the administration of contrast media (postcontrast), the ischemic muscle contains patchy nonenhancing regions of necrotic muscle that can simulate abscess. While myonecrosis tends to not show thick rim enhancement or as much mass effect as would be expected for the amount of excess fluid within the affected muscles, it is difficult to distinguish from an abscess (Fig 18). Because there is considerable overlap in the imaging appearances of pyomyositis and diabetic myonecrosis, fluid aspiration is typically necessary to exclude infection (59,60).

Pattern 3: Muscle Mass
Intramuscular masses include malignant or benign primary neoplasms, metastatic disease, and nonneoplastic masslike lesions related to infection, trauma, inflammation, or other underlying disorders. The differential diagnosis of an intramuscular mass depends on the patient’s age, pertinent clinical history, location of the mass, and its imaging appearance. A few intramuscular masses demonstrate unique MR imaging characteristics, allowing a specific diagnosis. It is beyond the scope of this article to discuss neoplasia in detail; only a few intramuscular masses are highlighted, emphasizing their characteristic imaging findings.

Malignant Primary Neoplasms of Muscle
The incidence of primary malignancy arising in muscle is surprisingly low given the large volume of this tissue in the body. The most common primary tumors arising in muscle are soft-tissue sarcomas of mesenchymal origin. While 12,000 new soft-tissue sarcomas are diagnosed annually in the United States (61), it should be noted that a substantial proportion arise in extramuscular tissues. In muscle, liposarcomas, neoplasms of fibrous origin, and lymphomas represent the most common primary malignancies. Less commonly, leiomyosarcomas, angiosarcomas, extraskeletal osteosarcomas (osseous), and synovial sarcomas develop in skeletal muscle (62). Intramuscular malignancies typically appear as heterogeneous infiltrative masses with variable degrees of enhancement and necrosis. Liposarcoma can be recognized if it contains macroscopic fat that is bright on T1-weighted images and suppresses with selective fat saturation. Tumors of fibrous origin may contain confluent regions of low signal intensity on T2-weighted images, but this finding is neither sensitive nor specific. In the majority of cases of skeletal malignancy, a biopsy is needed to establish a specific histologic diagnosis.

Malignant neoplasms derived from skeletal muscle cells and their precursors, known as rhabdomyosarcomas, are uncommon and account for less than 5% of all sarcomas (63,64). The majority of rhabdomyosarcomas are of the embryonal subtype and develop in children at extramuscular sites such as the head, neck, and genitourinary tract. The less common alveolar subtype affects all ages and typically arises in muscle, but overall, fewer than 25% of rhabdomyosarcomas arise within muscle (62). Adult rhabdomyosarcoma is rare, and the most common histologic subtype is anaplastic (also referred to as pleomorphic) (65). Rhabdomyosarcomas are nonspecific in appearance, resembling other soft-tissue sarcomas and often with central necrosis (Fig 19).

Skeletal muscle involvement is seen in 1%–2% of patients with lymphoma and typically results from muscle metastasis from systemic lymphoma or extension from the adjacent bone or lymph nodes (66). Primary extranodal lymphoma of
muscle is rare, typically affecting the thigh, trunk, upper arm, or calf in older patients, with an increased incidence in males and patients with HIV infection (67–69).

At MR imaging, muscle lymphoma most commonly manifests as an infiltrative mass, characteristically elongated along the course of the muscle fascicles, often with relative preservation of the intramuscular fat planes. It typically exhibits hyperintense signal on T1-weighted images and intermediate signal intensity on T2-weighted images, with diffuse enhancement or thick peripheral bandlike enhancement of the involved muscle (66,68,69). Involvement of multiple compartments is seen in up to 70% of patients. The presence of multiple masses, intratumoral vessels, extension of tumor along neurovascular bundles, thick fascial enhancement, and adjacent subcutaneous infiltration is suggestive of lymphoma (66–69) (Fig 20). Less commonly, lymphoma manifests as increased muscle signal intensity without a discrete mass, although the involved muscle tends to be enlarged and enhances in such cases.

**Benign Primary Neoplasms of Muscle**

The most common benign neoplasms arising within muscle are lipomas, hemangiomas/vascular malformations, myxomas, and fibromatoses (70). Lipomas are the most common soft-tissue neoplasm and can be classified as superficial or deep. Deep lipomas are commonly intramuscular but may also be intermuscular, typically affecting the large muscle groups in the extremities of adult men (71). Simple lipomas are isointense relative to fat on all pulse sequences. Large size, local symptoms, rapid growth, and nodular or thick (>2 mm) septal enhancement suggest an atypical lipomatous tumor or a well-differentiated liposarcoma (71).

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**Figure 20.** Lymphoma involving multiple muscle compartments in a 74-year-old woman. Axial fat-suppressed T2-weighted (a) and postcontrast fat-suppressed T1-weighted (b) MR images of the left calf show near-complete replacement of the normal musculature, with an intermediate-signal-intensity infiltrative tumor that enhances following contrast medium administration. A few areas of tumor necrosis are evident on the postcontrast image (arrowheads in b). Note the enlargement and infiltration of the tibial nerve (arrow in b), which also enhances.

**Figure 21.** Low-flow vascular malformation in a 35-year-old woman who presented with a mass along the ulnar aspect of the right elbow. (a) Coronal T1-weighted MR image shows a heterogeneous lesion (arrowheads) with low signal intensity and interspersed fat within. (b) Corresponding coronal fat-suppressed T2-weighted image shows nonuniform increased signal intensity, with low-signal-intensity bands corresponding to the fat striations within the mass (arrowheads). Note the small hypointense focus (arrow) at its deep aspect, which represents a phlebolith, typical of a low-flow vascular malformation.
Hemangiomas and vascular malformations are common benign and dysplastic vascular epithelial lesions, respectively, which typically demonstrate a lobulated and septated appearance, markedly hyperintense T2-weighted signal intensity, and enhancement on postcontrast images (72). Phleboliths are common in low-flow malformations (Fig 21), whereas enlarged serpentine flow voids suggest high-flow malformations.

Intramuscular myxoma is a benign tumor of uncertain differentiation that histologically resembles the mucinous substance of the umbilical cord (73). It is typically well marginated and homogeneous and, thus, is easily mistaken for a ganglion or benign cyst. MR imaging features that distinguish myxoma from a cyst include a peripheral rim of fat, edema in the surrounding musculature, and, most importantly, internal enhancement on postcontrast images (73) (Fig 22).

The fibromatoses are a group of benign fibroblastic proliferations that show infiltrative growth and can be locally aggressive. These lesions are classified by location as superficial or deep. Deep lesions are further categorized by age, location, and the specific tissue deposited. Desmoid-type fibromatosis is usually intramuscular or intermuscular and most frequently manifests in women during the 2nd to 4th decades of life. Lesions may be well defined or infiltrative with fascial plane extension (74). The MR imaging appearance reflects the amount of collagen and cellularity in the lesion, with more collagenous-containing lesions demonstrating low T1 and T2 signal intensities (74).

**Metastatic Disease**

Although muscle is highly vascularized and comprises 50% of total body mass, skeletal muscle metastases are rare, with autopsy studies citing a 0.03%–17.5% prevalence (75). Muscle metastases are most commonly located in the abdominal wall or paravertebral musculature, related to lung carcinoma, melanoma, or primary malignancies of the genitourinary and gastrointestinal systems; the latter types presumably disseminate to muscle by the paravertebral vascular plexus (75,76) (Fig 23). Muscle metastases are typically asymptomatic; the majority are diagnosed incidentally during routine staging. Their most common imaging appearance, seen in over half of cases, is that of a discrete intramuscular mass. Metastases can also appear abscess-like, related to central necrosis; ill-defined owing to infiltration; calcified; or hemorrhagic, simulating posttraumatic bleeding (75).
Hematomas

Hematomas are commonly encountered intramuscular masses, typically resulting from trauma or anticoagulant therapy. Hematomas can lead to elevated compartment pressure, serve as a nidus for infection, or cause compressive neuropathy (40,55,77). At MR imaging, hematomas demonstrate variable signal intensity, depending on their stage of blood product degradation. A peripheral high-signal-intensity ring related to methemoglobin on T1-weighted images in the subacute stage is the most distinctive finding indicating a hematoma (Fig 24).

Organizing hematomas often appear irregular and heterogeneous and may show patchy internal enhancement that can be mistaken for malignancy. Hematomas typically show rim enhancement at postcontrast imaging, a finding that may also be seen in necrotic neoplasms. When there is no clear history of injury or anticoagulation therapy, or when an intramuscular mass shows thick rim enhancement or substantial internal enhancement, an underlying neoplasm as the source of bleeding should be considered (8,9). It is prudent to follow such lesions with US to resolution to exclude an underlying neoplasm.

Nodular Myositis

Focal nodular myositis is a rare inflammatory pseudotumor of skeletal muscle that manifests as a rapidly enlarging and painful mass (78). Initial reports favored a traumatic or ischemic cause, but subsequent series describe an association with systemic polymyositis, suggesting an autoimmune basis (78,79). Other reports describe an association with underlying malignancy, suggesting that focal nodular myositis is a paraneoplastic phenomenon (80).

Focal nodular myositis is typically limited to a single muscle, but involvement of multiple muscles, either in the same or adjacent compartments, has been described (79). MR imaging findings are nonspecific, demonstrating a variably enhancing mass with surrounding edema (Fig 25). Tissue sampling is necessary to establish the diagnosis.

Sarcoidosis

Sarcoidosis is a chronic systemic noncaseating granulomatous disease that affects muscle in two forms: a generalized myopathy related to microscopic infiltration, and a nodular form with multiple discrete intramuscular masses (81,82). The diffuse form is not well depicted at MR imaging and is best diagnosed with gallium 67 ($^{67}$Ga) scintigraphy.

The nodular form, present in 1.4% of patients with sarcoidosis, typically affects the lower extremities and is distinctive. The masses are ovoid or cordlike, oriented longitudinally along muscle fibers (81), often at the musculotendinous junction (82). Axial images reveal central spiculated hypointense signal related to hyalinized connective tissue from chronic inflammation, surrounded by hypercellular granulomas peripherally (“dark star” sign). In the longitudinal plane, this arrangement produces cordlike signal intensity alterations, with a hypointense inner stripe encased by hyperintense outer stripes (“three stripes” sign) (81) (Fig 26). Sarcoid nodules can resolve after steroid therapy and may be substituted by regions of atrophy and fatty replacement (82,83).

Postinjection Masses

Adverse reactions to illicit, therapeutic, cosmetic, or preventative inoculations can result in localized muscle infection, inflammation, and fibrosis. Injection-related fibrosis after intramuscular vaccinations resulting in cordlike contractures of the deltoid muscle is well described, causing muscle atrophy and fibrosis, skin dimpling, and secondary scapular winging (84). MR images show an elongated mass of low-signal-intensity fibrotic tissue, while radiographs may show soft-tissue calcifications (85) (Fig 27).

Aesthetic intramuscular injections can result in peculiar appearances related to inoculation of autologous fat, hyaluronic acid, or, most commonly, free silicone. Free silicone injections are widely used for buttock cosmesis, producing a distinctive starry sky–like appearance related to the innumerable round heterogeneous deposits within the gluteal muscles and subcutaneous tissues (86) (Fig 28).

Intramuscular Extension of Calcium Hydroxyapatite

Calcium hydroxyapatite most commonly involves the shoulder tendons. Tendinous deposits can extrude into nearby tissues, inciting an intense
inflammatory response, typically affecting the adjacent bursa. Intramuscular extension is uncommon, occurring most commonly at the supraspinatus and infraspinatus muscles, producing a small intramuscular mass with low signal intensity surrounded by a larger inflammatory response (87) (Fig 29). Intramuscular hydroxyapatite migration may be along the central myotendinous junction or deep to the epimysial muscle covering. Radiographic correlation showing intramuscular extension of calcification confirms the diagnosis (87).

**Pattern 4: Atrophy**

Atrophy is characterized by loss of muscle tissue that manifests as decreased muscle size, often associated with fatty infiltration (88) (Fig 30). It is the irreversible end stage of numerous muscle...
diseases, including focal disorders such as trauma or denervation, as well as generalized disorders that are initially inflammatory. Atrophy is the principal finding in hereditary degenerative myopathies that produce disseminated muscle tissue loss without a prodromal inflammatory phase. Atrophy related to aging is referred to as sarcopenia, which is increasingly recognized as a substantial predictor of overall health. Atrophy is an important finding predicting loss of muscle power and function (89,90).

**Congenital Myopathies**

Muscular dystrophies are a heterogeneous group of progressive degenerative skeletal muscle disorders related to inherited gene defects (91). The best known are the Duchenne and Becker muscular dystrophies, inherited as X-linked recessive disorders, related to dystrophin gene dysfunction. Affected men present in childhood with pseudo-hypertrophy of the calf muscles and progressive lower extremity weakness. Maximal abnormalities are seen in the gluteal, thigh, and calf muscles bilaterally, with relative sparing of the adductor, gracilis, and sartorius muscles (91,92).

A host of other forms of muscular dystrophy have been characterized; almost all preferentially affect the lower extremities (91). A few forms, such as facioscapulohumeral muscular dystrophy, are characterized by preferential involvement of the face, chest wall, trunk, and upper extremity musculature (92,93) (Fig 31). The diagnosis of muscular dystrophies is based on clinical examination, muscle enzymes measurements, genetic sequencing, and tissue sampling. MR imaging has been used primarily to identify disease distribution, monitor disease severity, and select optimal sites for biopsy (91,93–95). Quantitative techniques such as MR spectroscopy and T2 mapping show promise for earlier and more accurate assessment of muscle degeneration and fatty replacement in patients with muscular dystrophies (96,97).

**Denervation Myopathy**

Denervation myopathy refers to the physiologic and anatomic changes that occur when muscle...
loses its innervation due to abnormalities of the central nervous system, spinal cord, peripheral motor nerve, or neuromuscular junctions at the muscle surface. Proposed mechanisms for denervation myopathy include capillary enlargement, increased vessel permeability, local metabolic effects, degeneration of muscle fibers, increased extracellular water, and proteolysis (98,99). MR imaging is sensitive for detecting muscle denervation and is a valuable adjunct to nerve conduc-
tion studies and electromyography for mapping the location and extent of disease, regardless of the underlying cause (100).

Denervation myopathy has variable appearances, depending on its duration. Acutely denervated muscle develops signal intensity alterations within 4 days of injury, preceding changes detected at electromyography (101). In acute denervation, T1-weighted images are normal, whereas fluid-sensitive images demonstrate uniformly increased muscle signal intensity (102,77). Unlike inflammatory disorders, there is no perimuscular fascial or subcutaneous fluid around the edematous muscle. In the subacute phase, signal intensity alterations persist on fluid-sensitive images, but T1-weighted images start to demonstrate atrophy (Fig 32). In chronic denervation, signal intensity alterations on fluid-sensitive images resolve, leaving a fat-replaced muscle that is irreversibly damaged (99).

The role of MR imaging in assessing denervation related to peripheral entrapment neuropathies has been emphasized in the musculoskeletal literature. Entrapment occurs in locations where a nerve courses through a fibro-osseous or fibromuscular tunnel, penetrates a muscle, or is impinged on by a mass (77,103). The key finding in peripheral nerve entrapment is that all affected muscles share a common innervation (Fig 33). The most common entrapment neuropathy relates to suprascapular nerve compression by a paralabral cyst arising from a glenoid labral tear. Cysts in the suprascapular fossa result in supraspinatus and infraspinatus denervation, whereas cysts located more distally in the spinoglenoid notch cause isolated infraspinatus denervation. Other common forms of entrapment neuropathy include compression of the peroneal nerve.

Figure 29. Intramuscular migration of calcium hydroxyapatite crystals in a 65-year-old woman with acute exacerbation of progressive left shoulder pain. Oblique sagittal fat-suppressed PD-weighted MR image of the left shoulder shows marked perifascial and intramuscular edema of the supraspinatus and infraspinatus muscles, related to intramuscular extension of the calcium hydroxyapatite (arrows). Calcifications were confirmed at radiography (not shown).

Figure 30. Fatty atrophy of supraspinatus muscle from a chronic rotator cuff tear in a patient undergoing reverse total shoulder arthroplasty. Photomicrograph of atrophic supraspinatus muscle specimen shows lobules of adipose tissue (arrows) interspersed between muscle fascicles (arrowheads), infiltrating and replacing the normal muscle tissue. Small foci of fat can also be seen within the degenerated fascicles. (Hematoxylin and eosin stain; original magnification, ×150.) (Courtesy of Samuel Ward, MD, University of California, San Diego, San Diego, Calif.)

Figure 31. Facioscapulohumeral muscular dystrophy in a 35-year-old man with progressive arm weakness and loss of ability to whistle. Coronal T1-weighted MR image of the upper chest shows severe fatty atrophy of the trapezius (arrows) and serratus anterior (arrowheads) muscles bilaterally, characteristic of the preferential chest wall and shoulder girdle muscle involvement seen in this form of muscular dystrophy.
near the fibular neck, the posterior interosseous branch of the radial nerve at the anterior elbow, and the median nerve at the carpal tunnel (103).

**Sarcopenia**

Sarcopenia, also known as *myopenia* or *muscle wasting*, is defined as diminished muscle mass and function. Aging is an important risk factor; up to 50% of an adult’s normal muscle mass can involute with advanced age, which has profound consequences on muscle function and physical health. Although aging is associated with muscle loss, the cause of sarcopenia is multifactorial and is also related to poor nutrition, inactivity, neurologic decline, hormonal alterations, and chronic disease (104). Sarcopenia has been associated with a number of adverse health outcomes such as prolonged disability, higher rates of hospital readmission and prolonged hospital stays, postoperative complications, and increased mortality (105). Sarcopenia appears to be an independent predictor of adverse outcomes beyond other comorbidities that may manifest (106).

Histologic analysis reveals diminished quantity and quality of muscle fibers, particularly of the type II fiber fraction; fatty infiltration; and quiescence of satellite cells involved in muscle regeneration (104). CT of the abdomen/pelvis is frequently used to opportunistically assess paraspinal and truncal muscle volume as it is widely available and has been shown to be a valid measure of sarcopenia (106). Routine MR imaging sequences demonstrate volume loss and fatty infiltration; more sophisticated analysis of muscle composition can be obtained by using quantitative techniques such as fat-water imaging, lipid-suppressed mapping, diffusion-tensor imaging, and lipid spectroscopy (105) (Fig 34).
Conclusion

The MR imaging appearance of atraumatic muscle conditions can be classified into four broad patterns: (a) abnormal anatomy with normal signal intensity, (b) edema/inflammation, (c) intramuscular mass, and (d) atrophy. These four patterns may be seen in isolation or in combination, depending on the underlying disease and its severity and duration. An understanding of the disorders leading to each of these imaging patterns, coupled with clinical information, will help radiologists formulate appropriate differential considerations, suggest a specific diagnosis, plan biopsy, and guide appropriate management.

References


