Simple Bone Cysts (Solitary Bone Cyst, Unicameral Bone Cyst)

Simple bone cysts are benign tumors of childhood and adolescence. They represent approximately 3 percent of all biopsied primary bone tumors and nearly always occur during the first two decades of life, most often between 4 and 10 years of age.\(^1\) There is a male predominance, with a 2:1 male-female ratio. The majority of cysts occur in the metaphyseal region of the proximal humerus or femur, with approximately 50 percent of cases involving the humerus and 18 to 27 percent affecting the femur. The next most common sites are the proximal and distal tibia. Occasionally, cysts may be found in the calcaneus, fibula, ulna, radius, pelvis, talus, or axial skeleton* (Fig. 37-1).

Rarely does more than one cyst occur in an individual, thus the term solitary bone cyst. The term unicameral bone cyst implies that one chamber exists. Although one large cavity is usually found, a cyst may become multiloculated following a fracture because of the formation of multiple bony septations, thus making the term unicameral technically incorrect.

Simple cysts are often categorized as “active” or “latent” based on their proximity to the growth plate.\(^2,3,5,6,10\) A cyst that is juxtaphyseal (less than 0.5 cm from the physis) is considered “active” and possesses greater potential for growth. A cyst that has grown away from the plate is considered “latent” and, theoretically, no longer has the capacity for growth (Fig. 37-2). In reality, though, latent cysts continue to have growth potential, as proved time and again by their unexpected recurrence following treatment in the young individual. After skeletal maturity, it is uncommon for the cysts to recur or progressively worsen.

**ETIOLOGY**

The cause of simple bone cysts remains uncertain. Any theory relating to the etiology of simple bone cysts should be able to explain the following factors: (1) more than 70 percent are discovered in childhood; (2) more than 95 percent arise from or involve the metaphysis; (3) most occur in the proximal humerus or femur; (4) a cyst wall and fluid high in protein content are common; and (5) simple bone cysts represent a benign process with a significant recurrence rate following treatment.\(^1,3,5,6,10\)

Mirra hypothesized an intraosseous synovial cyst in which a small amount of synovial tissue became entrapped in an intraosseous position during early infant development or secondary to trauma at birth.\(^1,3,5,6,10\) Over time, increased pressure secondary to secretions would lead to expansion within the bone. Jaffe and Lichtenstein postulated that cysts resulted from a localized failure of ossification in the metaphyseal area during periods of rapid growth.\(^1,3,5,6,10\) Cohen proposed that the cause of the cyst was blockage of the circulation (venous obstruction) and drainage of interstitial fluid in rapidly growing bone. He based this theory on the finding that the chemical constituents of the fluid in simple bone cysts are similar to those of serum.\(^1,3,5,6,10\) Current literature further substantiates this theory of a disturbance in or occlusion of the intramedullary venous circulation.\(^1,3,5,6,10\)

Drilling, trepanation, and reaming of the medullary cavity...
FIGURE 37–1  Imaging findings in a 14-year-old boy with pain in his heel. A, Radiograph demonstrating the lucent solitary bone cyst in the calcaneus. B, A CT scan showed the extent of the lesion, which is surrounded by a rim of cortical bone.

FIGURE 37–2  A and B, Active solitary bone cyst in the proximal left femur of a 4-year-old boy (D.J.). The cyst is juxaphyseal. C and D, A latent solitary bone cyst has grown away from the proximal physis in the right humerus of an 8-year-old boy (D.T.). See Figure 37–4 for posttreatment results.
to open vascular channels between cysts and the intramedullary venous system have all been found effective in healing unicameral cysts. The cyst fluid itself may be both a causative factor and an obstacle to healing. Bone-resorptive factors, such as prostaglandins, interleukin-1, and lysosomal enzymes, are found in cyst fluid. Oxygen free radicals, which are cytotoxic and are known to be generated under ischemic conditions, have also been found in cysts.

**PATHOLOGY**

Simple bone cysts tend to expand by eroding the cortex, resulting in a localized bulge of the bone. Despite this fact, reactive or periosteal bone formation is not present unless a pathologic fracture occurs. Where the cortical tissue is thinnest, the wall can actually be flaccid and a bluish tinge from the underlying fluid can be seen. Once the affected bone has fractured, the cortical wall is thicker and multiple bony septa may occur throughout the cyst.

The fluid found within simple bone cysts is straw-colored or serosanguineous, a feature distinguishing simple bone cysts from aneurysmal bone cysts. Often, significant pressure within the cyst (which can be greater than 30 cm H2O) is evident when a needle is introduced. After a fracture, though, the cyst may become filled with blood clot, granulation, or fibro-osseous tissues. The most characteristic histopathologic finding is the thin membranous lining of the cyst (Fig. 37–3). Composed primarily of flattened to plump epithelial-like cells, the lining may also possess osteoclast-type giant cells, cholesterol cells, and fat cells. Hemosiderin, fibrin, calcification, and reactive bone may be seen in focal areas of the cyst.

**CLINICAL FEATURES**

Clinically, cysts can be asymptomatic and may be discovered incidentally when radiographs, such as a chest film, are obtained for other reasons. More often, though, the cysts are diagnosed because of pain. The pain may be mild and reflective of a microscopic pathologic fracture. More abrupt discomfort occurs when a pathologic fracture occurs following relatively minor trauma, such as a fall. The fractures heal readily, but the cysts do not. Following these pathologic fractures, premature physeal closure has been reported in nearly 10 percent of patients.

**RADIOGRAPHIC FINDINGS**

There are several characteristic radiographic features of simple bone cysts. Approximately 50 percent occur in the proximal humerus and 18 to 27 percent in the proximal femur. The cyst is metaphyseal and usually extends to, but not across, the physis. On rare occasions, it crosses the physis into the epiphysis. Typically the cyst is symmetrically expansile and radiolucent, with a thin cortical rim surrounding it. Over time, the physis grows away from the cyst, changing from the active to the latent phase. In many newly diagnosed cases there is a pathologic fracture with or without displacement. The one pathognomonic manifestation of a simple bone cyst is the "fallen fragment" sign. This represents a portion of fractured cortex that settles to the most dependent part of the fluid-filled cyst. However, it is seen in less than 10 percent of cases, and it should not be expected if the cyst has become multiloculated following a previous pathologic fracture.

**DIFFERENTIAL DIAGNOSIS**

The diagnosis can usually be established based on the presence of typical radiographic findings. Other lesions to be considered in the differential diagnosis include aneurysmal bone cyst, monostotic fibrous dysplasia, and atypical eosinophilic granuloma. All of these lesions may be radiolucent. Aneurysmal bone cysts and fibrous dysplasia may be expansile and metaphyseal. However, features typically associated with these lesions usually help differentiate them from simple bone cysts.

**TREATMENT**

A common misconception in the treatment of simple bone cysts in children is that once the pathologic fracture heals, the cyst also has an excellent chance of spontaneously healing. However, most investigators examining this phenomenon have found that the likelihood of spontaneous healing of the cyst following pathologic fracture is very low, probably less than 5 percent. Thus, if treatment of the cyst is deemed necessary, it should be undertaken as soon as the fracture has healed. However, overtreatment in mature persons should be avoided. In these individuals, if the cyst has a sufficiently thick cortex and is located in the upper extremity, periodic observation may be all that is needed. If the patient is asymptomatic, it may not be necessary to restrict activities.

The treatment approach is more aggressive for all simple bone cysts in younger children and in mature individuals when the cyst is located in weightbearing bones of the lower extremities. In these cases, plans should be made for definitive treatment of the cyst in order to prevent future fractures and possible associated complications (e.g., shortening due to growth arrest and deformity).

The preoperative evaluation of patients with simple bone cysts rarely requires more than good-quality radiographs of the lesion. If the diagnosis is equivocal, a bone scan will verify the presence or absence of other abnormal areas.
Computed tomography (CT) may be helpful in differentiating simple bone cysts from other lesions, such as aneurysmal bone cysts or fibrous dysplasia. The diagnosis is usually confirmed at surgery, when straw-colored fluid is aspirated through a large-bore needle introduced into the cystic cavity.

Treatment modalities include injection of corticosteroids into the cyst, injection of autologous bone marrow, multiple drilling and drainage of the cavity, and curettage of the membranous wall followed by bone grafting. Older forms of treatment, such as subtotal resection with or without bone grafting and total resection, are rarely, if ever, used today.

Corticosteroid Injections. The successful healing of cysts following injections of methylprednisolone acetate was reported by Scaglietti and colleagues in 1979. They noted favorable results in 90 percent of lesions, and consequently concluded that treatment by curettage was seldom necessary. Healing was believed to have occurred if the cortex thickened and the cystic cavity became radiographically opaque.
of the cyst with "bone scar" was considered evidence of healing (Fig. 37–4). Actual remodeling, with complete disappearance of the cystic cavity, often took several years. Subsequent reports have continued to substantiate the effectiveness of injecting steroids into cysts, although the success rates have been lower, ranging from 40 to 80 percent. This method continues to be the treatment of choice for the initial management of simple bone cysts. The antiprosta-glandin action of steroids constitutes the rationale for their use in the treatment of cysts.

Operative Technique of Corticosteroid Injections. The patient is given a general anesthetic in the operating room and the procedure is performed using strict aseptic techniques. Fluoroscopy with image intensification is used to locate the margins of the cyst. Two large needles with stylets (≥14-gauge or Craig biopsy needles) are utilized. The first needle is introduced percutaneously, the stylet is withdrawn, and fluid is allowed to drip out. The presence of straw-colored (serosanguineous) fluid confirms the diagnosis of simple bone cyst. Vigorous aspiration must be avoided since blood may be returned, making it difficult to distinguish a simple bone cyst from an aneurysmal bone cyst. If straw-colored fluid is returned, contrast material (usually Renograin diluted 1:1 with normal saline) is injected to confirm the presence or absence of intracycst fibrous or osseous septa and loculation. If the cyst is not filled completely, the incidence of failure of healing increases. Needles must be introduced into each separate cystic cavity to ensure delivery of the steroid.

After the contrast material clarifies the structure of the cystic cavity, the second needle is introduced. The cavity is thoroughly flushed with normal saline solution. The operator should not aspirate when a second needle is in the cyst, as air can be aspirated into the cyst from the second needle, leading to an air embolus. When lavage with normal saline is completed, the second needle is withdrawn. Through the remaining needle, 40 to 120 mg (1 to 3 mL) of methylprednisolone acetate is introduced into the cyst and a simple compression dressing is applied.

This procedure is usually repeated every 2 months, requiring between two and five injections, with three being the usual minimal number to obtain healing. Radiographic changes usually are not noted in the first 2 to 3 months; thus radiographs are not needed before then. Subsequently, radiographs are obtained every 2 to 3 months to assess healing. Evidence of healing includes diminution in the size of the cyst, cortical thickening, remodeling of the surrounding bone, and increased internal density (such as "ground-glass" ossification).

Serial steroid injections is the most popular treatment mode because the procedure is simple, injury to the adjacent physis is avoided, the procedure causes a minimal operative scar, there is little morbidity, the patient is able to return promptly to a previous activity level, and the reported results are excellent. A potential disadvantage is a temporary systemic response to the steroid (Cushing's syndrome). It is best not to exceed a total of 120 mg of methylprednisolone during any one injection.

Autologous Bone Marrow Injections. Recent interest in the injection of other materials to stimulate healing has led to the successful use of autologous bone marrow. Collagen and demineralized bone matrix are other injectable materials that are under investigation.

Decompression of Cysts by Multiple Drilling. Multiple percutaneous drilling has been shown to be effective in the

*See references 36, 44, 50, 51, 89, 114, 193, 234, 239.
treatment of simple bone cysts. Following trepanation, the cyst is thoroughly lavaged with saline. Multiple holes are then created in the cyst wall. Fluid escapes through the drill holes, decreasing the internal pressure in the cyst. When the cysts are drilled with Kirschner wires, the wires are either left in place or removed. Leaving them in place theoretically keeps the holes open and allows for continuous drainage through the cyst wall. However, we have no personal experience with this technique.

Curettage of Cysts Followed by Bone Grafting. Once the common form of treatment for simple bone cysts, bone grafting was replaced in the late 1970s and early 1980s by steroid injections because of reports of better healing of cysts using methylprednisolone acetate (Fig. 37–5). Nearly 50 percent of cysts recur following curettage and bone grafting. However, there are some cysts for which curettage followed by bone grafting remains necessary. Patients with displaced pathologic fractures of the hip need open reduction and internal fixation. At the time of internal fixation, curettage of the cyst and bone grafting are also performed. Materials other than autogenous bone have recently been used with success, including cubes of high-porosity hydroxyapatite and tricalcium phosphate ceramic.

Aneurysmal Bone Cyst

An aneurysmal bone cyst is a solitary, expansile, radiolucent lesion of uncertain etiology. It is generally located in the metaphyseal region of the long bones. Seen much less often than simple bone cysts, aneurysmal cysts represent approximately 1 percent of all biopsied primary bone tumors. Nearly 70 percent of affected patients are between 5 and 20 years of age, but the lesion has been reported in infants. There is no sex predilection.

Aneurysmal bone cysts can be found throughout the skeleton. Nearly 50 percent of reported cases occur somewhere within the long bones of the extremities. The most common sites are the femur and tibia. Although they usually involve the metaphyseal region, aneurysmal cysts may on occasion cross the physis into the epiphysis or may extend into the diaphysis.

Approximately 20 percent of aneurysmal bone cysts involve the spine, and they can occur anywhere between the axis and the sacrum. Within the vertebra itself, the cyst may be found in the body, pedicles, lamina, and spinous process (Fig. 37–6). Involvement of two or more adjacent vertebrae is not uncommon. Aneurysmal bone cysts may also occur in the pelvis, maxilla, clavicle, hands, and feet.

FIGURE 37–5 A 10-year-old boy with a latent solitary bone cyst in the humerus sustained a nondisplaced fracture two separate times (A). Once the fracture had healed for the second time, curettage and bone grafting was performed. Three months postoperatively the cyst was healing (B, C). Five years postoperatively the cyst had healed (D, E).
FIGURE 37-6 Imaging findings in an 11-year-old girl with moderate back pain in the thoracolumbar spine for 6 months. A and B, AP and lateral spine radiographs demonstrated an aneurysmal bone cyst of L1. C and D, CT showed that the cyst extensively involved the posterior elements on one side. E, Two years after resection and bone grafting, the radiograph still was not normal.
ETIOLOGY

Aneurysmal bone cysts represent either a primary neoplastic condition or a secondary response (arteriovenous malformation) to the destructive effects of an underlying primary tumor. Development of an aneurysmal cyst as a secondary response is supported by the association of aneurysmal cysts with other primary lesions, such as nonossifying fibromas, fibromyxomas, fibrous dysplasia, chondroblastomas, giant cell tumors, single bone cysts, telangiectatic osteosarcomas, chondrosarcomas, and metastatic disease. 180,193 Sixty-five percent of aneurysmal bone cysts have been reported to be primary, with 35 percent believed to be secondary to other lesions. 133,139 Thus, once the diagnosis of aneurysmal bone cyst is considered, a thorough preoperative evaluation is necessary, adequate tissue must be obtained at the time of surgery, and careful pathologic studies are needed to ensure that the aneurysmal cyst is not secondary to a more serious primary neoplasm.

PATHOLOGY

Aneurysmal bone cysts vary considerably in size, with the potential of becoming quite large during the rapid destructive growth phase. On gross inspection, the cyst consists of an encapsulated mass of soft, friable, reddish brown tissue, usually contained within a thin subperiosteal shell of new bone. At the time of surgery, a large amount of blood may exude from a mesh of honeycomb spaces. In most cases the blood is dark red owing to a slow but continuous circulation. If the circulation to a portion of the aneurysmal cyst has been blocked, the cyst may be filled with serous or serosanguineous fluid or with focal organized blood clots.

Microscopy discloses a variable number of vascular spaces whose walls are lined with tissue composed of fibroblastic cells with collagen, giant cells, hemosiderin, and osteoid (secondary to microfractures) (Fig. 37-7). Extensive sampling should be performed to identify possible benign or malignant precursor lesions. The histologic diagnosis of primary aneurysmal bone cyst should be made only after other possible lesions have been excluded. Fibrous tissue, bone, and giant cells are the usual elements seen in most other benign precursor lesions associated with an aneurysmal bone cyst. Any solid area that is 1 cm or greater should raise the suspicion that it may represent another lesion.

There is another entity known as a solid aneurysmal bone cyst or giant-cell reparative granuloma. 27,40 This is a solid yet radiolucent lesion that appears grayish brown and often is friable. Histologic features include fibrous proliferation with giant cells, fibromyxoid areas, and bone production. Characteristically, the giant cells, which are clustered in areas of recent and old hemorrhage, are found throughout the lesion. The solid aneurysmal bone cyst lacks the normally large blood-filled channels (Fig. 37-8). In a recent review of a large series of aneurysmal bone cysts, the incidence of the solid entity was 7.5 percent. 27

CLINICAL FEATURES

The clinical presentation includes localized pain of several weeks’ or months’ duration, tenderness, and, if the aneurysmal bone cyst occurs in an extremity, swelling. When the cyst involves the spine, progressive enlargement may compress the spinal cord or nerve roots, resulting in neurologic deficits such as motor weakness, sensory disturbance, and loss of bowel or bladder control. Spinal involvement mandates urgent care.

RADIOGRAPHIC FINDINGS

The classic radiographic feature of aneurysmal bone cyst was described by Jaffe as a periosteal “blowout” or ballooned-out lesion that is outlined by a thin shell of subperiosteal new bone formation. 125 In about 80 percent of cases the cyst involves the metaphyseal region of the long bones and, unlike simple bone cysts, is eccentric in its location. In the spine, it more often involves the posterior elements (spinous process, transverse process, and pedicles) than the vertebral body. In the shorter tubular bones of the feet, the cysts are more central and extend into the diaphysis and subarticular region (this is explained by the smaller size of the bones).

Three phases of aneurysmal cysts have been described. 186 The incipient phase is characterized by either a small eccentric lucent lesion or by a pure lifting off of the periosteum
from the host bone without evidence of an intramedullary lesion. Most patients do not present with disease in this phase. Except for focal cortical thinning, the cortex may otherwise be preserved and the periosteum may show no reaction. In this phase, the lesion can be mistaken for a simple bone cyst, nonossifying fibroma, or possibly a lytic osteosarcoma. The midphase designates the period of rapid, destructive growth and is characterized by extreme lysis of the bone, focal cortical destruction, and the development of Codman’s triangles (periosteal ossification at the corner of the expanded cyst). It is during this phase that the “blowout” appearance is seen on radiographs, and aneurysmal bone cysts can easily be mistaken for an aggressive malignant lesion. In the late healing or stabilization phase, the lesion grows more slowly and the periosteum has sufficient time to lay down new bone. The cyst will exhibit eccentric (or possibly concentric) smooth-bordered expansion, a trabeculated or “bubbly” intramedullary appearance, and surrounding host bone sclerosis.

Capanna and colleagues proposed a radiographic classification system that is commonly used today. Inertive denotes a complete periostal shell, with the intraosseous margin defined by a sclerotic rim of reactive bone. Active indicates an incomplete periostal shell and a sharply defined intraosseous border. Aggressive cysts show no evidence of reparative osteogenesis, no periostal shell, and an ill-defined endosteal margin.

Once these lesions are identified on radiographs, the tumor can be better clarified with CT, particularly if it is located in the spine. The extent of involvement of the vertebral body and any encroachment of the spinal canal will be readily evident. CT will also demonstrate the characteristic fluid-fluid levels if the patient is able to lie still long enough for the serosanguineous fluid to separate from the blood within the chambers of the cyst that do not have active circulation. Magnetic resonance imaging (MRI) is indicated if there is evidence of spinal cord compression or if the edges of the rapidly expanding cyst cannot be defined with CT. Fluid-fluid levels are readily evident on MRI (Fig. 3.7–9).

**TREATMENT**

Although spontaneous healing of aneurysmal bone cysts has been reported, it is uncommon. Thus, expectant management should be considered only when the diagnosis has been made with confidence and the lesion is in a location and at a stage that do not entail any risk of fracture or further destruction. More often, when the diagnosis of aneurysmal bone cyst is made, active treatment is recommended.

Curettage followed by bone grafting of aneurysmal cysts has been the standard treatment for many years (Fig. 3.7–10). Unfortunately, this tumor has a high incidence of local recurrence (20 to 59 percent) after curettage. Thus, adjunctive therapy, such as cementation, cryotherapy, or embolization, should be considered along with curettage. Cementation of the lesion with polymethylmethacrylate, followed 4 to 6 months later with replacement bone grafting, has been reported to be more effective than curettage and bone grafting alone. Cryotherapy as an adjunct to curettage and bone grafting also increases the likelihood of healing. Embolization has been used as the sole treatment for aneurysmal bone cysts, but it is much more commonly used preoperatively to interrupt the vascularity to the lesion. Embolization is useful in treating aneurysmal cysts located in areas of limited access, such as the spine and pelvis. If the cyst is located in an expendable bone, such as a rib or fibula, the surgeon should consider performing a wide or en bloc excision.

Anecdotal reports of techniques useful in the treatment of aneurysmal bone cysts include oral dexamethasone (an angiotropic agent), and the use of multiple Kirschner pins inserted into the cyst. Aneurysmal cysts in the spine most commonly involve the elements of the posterior column, but the cysts may extend anteriorly into the body. More than one vertebra may be affected. On occasion, neurologic deficit due to compression of the spinal cord by the lesion requires emergency resection. More often, however, time is available to plan the necessary preoperative embolization, surgical approaches, and reconstruction of a surgically destabilized spine.

Radiation therapy has been used for some aneurysmal bone cysts, especially those located in areas that are difficult to gain access to, such as the spine. The dose should be minimized (between 3,000 and 5,000 cGy) to decrease the risk of radiation-induced sarcoma. Because of this concern, radiation therapy should be limited to cases of cysts that are inoperable or have become inoperable and cases in which embolization has failed.

**Fibrous Dysplasia**

The term fibrous dysplasia was originally proposed by Lichtenstein in 1938. He, along with Jaffe, McCune, and Albright, described this disorder of bone, as well as other extraskeletal abnormalities with which it is occasionally associated. Their descriptions remain among the best for fibrous dysplasia—a benign, nonfamilial disorder characterized by the presence of expanding intramedullary fibrous tissue in one or more bones. The incidence of fibrous dysplasia is not known, but it is not an uncommon primary bone tumor. It occurs more frequently in females than in males, particularly the polyostotic form. Although most lesions are probably present in early childhood, they usually do not become evident before late childhood to adolescence.

In general, fibrous dysplasia can be classified into one of three categories. Monostotic fibrous dysplasia involves only one bone, and many of these patients remain asymptomatic unless a fracture or swelling occurs. The polyostotic form is more severe, involving multiple bones. Nearly any bone in the body may be affected, including the long bones of the extremities, the skull, vertebrae, pelvis, scapula, ribs, hands, and feet. Often one side of the body (in particular, one of the lower extremities) is more severely affected, resulting in deformity and limb length discrepancy. Craniofacial involvement occurs in nearly 50 percent of patients with polyostotic disease. The third category, polyostotic form with endocrine abnormalities, is the least common form. Precocious puberty, premature skeletal maturation, hyperthyroidism, hyperparathyroidism, acromegaly, and Cushing’s syndrome can occur in these patients. The triad of precocious puberty (endocrinopathy), cafe-au-lait spots, and polyos-
FIGURE 37-9 Imaging findings in a 9-year-old girl with a swollen tender first ray of the left foot. A, AP radiograph demonstrating an expansile well-contained lesion. B to D, MRI images better defined the lesion and showed the layering effect seen with aneurysmal bone cysts. E, Three years after curettage and bone grafting, the metatarsal was normal.
totic bone involvement is commonly referred to as McCune-Albright (or Albright’s) syndrome.

ETIOLOGY

The exact cause of fibrous dysplasia is not known. The condition is not believed to be hereditary. Fibrous dysplasia probably results from a failure of maturation from woven to lamellar bone. Recent studies have reported an abnormality of a gene encoding a G protein, which is important in the development of bone. Somatic activating mutations of the signal transducer Gs of the alpha chain differentiate fibrous dysplasia (particularly McCune-Albright syndrome) from other lesions, and the mutations may be responsible for the loss of control of local proliferation and growth factor expression.

PATHOLOGY

The outer surface of expanded bone is usually smooth and covered by reactive periosteal bone. The underlying dysplastic tissue is firm and grayish white, and the proliferative tissue is fibrous. It may feel gritty when palpated, almost like sandpaper. There may be degenerative cystic changes due to cellular necrosis. Islands of hyaline cartilage may be seen.

Histologically, irregular foci of woven (nonlamellar) bone trabeculae are seen in a cellular fibrous stroma (Fig. 37–11). Under the microscope, the bony spicules are often described as looking like the letters C, J, or Y, or resembling Chinese characters. Osteoclastic resorption may be seen, but osteoblastic rimming of the bony spicules is uncommon. In unusual cases, as much as 95 percent of the lesional tissue may be fibrous. Cartilage islands, multinucleated giant cells, foamy histiocytes, and callus may all be seen if a fracture has occurred. The histologic features of polyostotic lesions are identical to those of monostotic lesions.

CLINICAL FEATURES

The clinical manifestations are usually mild in monostotic fibrous dysplasia. Pain and a limp may be evident when the neck of the femur is involved. Local swelling may be seen when the lesion is in a superficial bone, such as the mandible, skull, or tibia. The skeletal changes are usually more severe in the polyostotic form and may result in pain, swelling, deformity, and limb length discrepancies. The classic example of this is found in the proximal femur. Repetitive microfractures can lead to a “shepherd’s crook” deformity with pain, significant varus at the femoral neck, shortening of the femur, an obvious Trendelenburg gait, and limited mobility. Deformity can occur in all of the long bones, but usually not to the degree seen in the femur. When the facial bones are affected, progressive deformity may become very evident to the patient and family. Numerous reports of craniofacial abnormalities are found in the dental and maxillofacial literature. Spinal involvement is uncommon but can result in vertebral collapse, angular deformity, and possible spinal cord compression.

The most common nonskeletal manifestation associated with fibrous dysplasia is abnormal cutaneous pigmentation, or café-au-lait spots. These have irregular borders (“coast of Maine”), are not raised from the surrounding skin, and may be extensive, involving large areas of the trunk, face, or limbs. The café-au-lait spots can coexist with polyostotic fibrous dysplasia without endocrine changes or precocious puberty, or they may be absent when endocrinopathies accompany fibrous dysplasia. The pigmentation changes usually are not present in the monostotic fibrous dysplasia.

When sexual precocity occurs, it is most often seen in the female secondary to premature ovarian stimulation and may occur as early as 1 year of age. Most cases of McCune-Albright syndrome occur in females and are accompanied by accelerated maturation and advanced skeletal age. In such children, abnormally rapid growth may result in tall stature at a young age. Ultimately, however, their adult stature will usually be below average because of their early maturation.

RADIOGRAPHIC FINDINGS

Fibrous dysplasia can affect any bone. In the long bones, the lesions start in the metaphysis or diaphysis and rarely involve the epiphysis. The flat bones, ribs, jaw, and skull are commonly involved, but the spine is not.

Some of the smaller lesions of fibrous dysplasia remain confined to the intramedullary region, are often surrounded by sclerosis, and may appear “bubbly” or trabeculated. Normally, though, there is slow replacement of the cortex as expansion takes place. Larger lesions may result in an even or eccentric, smooth-bordered expansion of the bone, but they usually remain confined within a rim of periosteal bone. Angular deformity may occur in the long bones, such as the shepherd’s crook deformity in the proximal femur and occasionally in the humerus. This is usually a result of remodeling of the bone following repetitive microfractures or obvious fractures through the dysplastic bone (Fig. 37–13).

The radiographic density of the lesions depends on the amount of woven bone produced and on the amount of cortex replaced. If the lesion is small, has produced little woven bone, or has not replaced cortex, it will appear radiolucent in comparison to surrounding normal bone. If the cortex is thinned and if the fibrous dysplasia has expanded and replaced most of the normal bone, the characteristic “ground-glass” appearance is seen. When this feature is extensive, it is nearly pathognomonic of fibrous dysplasia. CT will clearly demonstrate this appearance.

A bone scan will demonstrate increased uptake throughout the lesion and is helpful in determining the extent of the disorder if radiographs are unable to do so.

DIFFERENTIAL DIAGNOSIS

With monostotic fibrous dysplasia, it may be difficult to differentiate small lesions from simple bone cysts on radiographs. Less often, small lesions may be confused with histiocytosis or enchondromas. In these cases a biopsy may be necessary. Larger lesions with cortical thinning and a ground-glass appearance usually do not require a biopsy to confirm the diagnosis. Polyostotic fibrous dysplasia is readily identified on radiographs.
FIGURE 37-10 Aneurysmal bone cyst. Imaging findings in a 3-year-old boy who presented with a painful swollen distal forearm. A and B, AP and lateral radiographs demonstrating the "blown-out" appearance of the distal radius. C, The edges of the lesion could not be seen well on CT. D and E, MRI demonstrated containment of the lesion.

TREATMENT

The mere presence of fibrous dysplasia in bone is not, in itself, an indication for surgery, and surgical overtreatment should be avoided. Surgery is indicated when the lesions (1) are complicated by fractures, (2) cause significant or progressive deformity that jeopardizes the integrity of the long bone or that results in unacceptable disfigurement, or (3) are symptomatic and cause the patient pain.

In children, it is nearly impossible to restore dysplastic bone to normal bone following surgery. Thus, if the rare biopsy is needed to confirm the diagnosis of a monostotic lesion, surgical intervention probably should not be undertaken unless there is a fracture or painful deformity. The reason for this conservative approach is that simple curettage of the lesion inevitably leads to local recurrence. In addition, the risk of pathologic fracture is increased during the months immediately following surgery. Bone graft used to replace part or even all of the tumorous bone is also predictably resorbed and replaced by the dysplastic bone. A proposed exception to this nonoperative approach in children (in the absence of fracture or deformity) is infantile fibrous dysplasia. In these children with polyostotic disease, early surgical treatment to prevent development of skeletal deformities that are difficult to correct later may provide long-term benefit. Prophylactic intramedullary nailing with nails of appropriate size was found to be most effective. In the adult, bone grafting in fibrous dysplasia has been reported to be more successful in healing the dysplastic bone.
Operative intervention is needed when repeated pathologic fractures have occurred, deformity increases, or associated pain becomes persistent. These pathologic fractures can occur following mild trauma, they are often minimally displaced, and they heal at a normal rate. Delayed union or nonunion is not a problem, but progressive deformity is. The primary goal of treatment is to realign the deformed bone, particularly in the weightbearing lower extremities. This objective, along with ready healing of the fracture, can often be achieved with cast immobilization in the young child. If repeated fractures occur in long bones or if a fracture involves the proximal femur, then surgical intervention is the preferred treatment approach. Internal fixation maintains proper alignment and can be achieved with intramedullary rods in the long bones or with compression screws with side plates in the proximal femur. Prior to the insertion of internal fixation, osteotomies may be required to achieve satisfactory alignment because bowing of the bone (malunions) is common following fractures. This is particularly true in the proximal femur, in which a shepherd’s crook deformity may be present. Intraoperative blood loss may be excessive because of increased vascularity in the bone. In addition, the dysplastic bone may not allow good cortical screw purchase, necessitating alternative plans for internal fixation. Despite the successful use of internal fixation and near-anatomic bone realignment, progressive deformity can still occur, leading to the need for additional surgery.

Some authors advocate attempts to augment bone strength in addition to or in place of internal fixation. Enneking and associates reported that cortical strut grafting was effective in strengthening the bone in the proximal femur (Fig. 37–14). In their opinion, the strength of the bone was greater if cortical rather than cancellous graft was used. Allograft cortical struts avoid the morbidity of harvesting autogenous graft and also appear to slow the resorption process by the dysplastic bone. Ultimately, though, the type of graft utilized probably does not affect the rate of recurrence.

Although malignancy in fibrous dysplasia is rare, when it does occur the prognosis is poor. Most cases occur in patients who have undergone radiation therapy. Thus, radiation treatments should be avoided because of the association with malignant transformation. In addition, no beneficial effects on the progression of polyostotic fibrous dysplasia have been demonstrated from medications or medical management. The hormonal abnormalities in McCune-Albright disease should be managed by an endocrinologist.
Osteofibrous Dysplasia of the Tibia and Fibula (Campanacci’s Disease)

Osteofibrous dysplasia of the tibia and fibula has been described as a variant of fibrous dysplasia. It may also have a histogenetic relationship to adamantinoma.* Cytokeratin-positive cells are found in the stroma of both osteofibrous dysplasia and adamantinomas, but not in fibrous dysplasia.

The disorder was first reported in the literature in 1921 by Frangenheim, who used the term congenital osteitis fibrosa. Other terms for this disorder include congenital fibrous dysplasia, congenital fibrous defect of the tibia, and ossifying fibroma. Osteofibrous dysplasia of the tibia and fibula was proposed by Campanacci in 1976.43,45

Osteofibrous dysplasia differs from the more common fibrous dysplasia with regard to age distribution, site, radio-

*See references 10, 25, 31, 37, 116, 124, 155, 265, 275, 285.
graphic features, and clinical course. The lesion is slightly more common in males. The symptoms almost always appear in the first decade of life. Nearly two-thirds of the lesions are noted before 5 years of age, and the disorder has been noted in infants. The tibia is almost always involved and the ipsilateral fibula may be affected. Bilateral involvement has been reported, as has involvement of the radius and ulna. The diaphysis is mainly affected, with localization to the middle third in the tibia. Extension into the proximal or distal metaphysis is seen sometimes. On rare occasions there is diffuse involvement of the entire shaft of the tibia. When the disorder is limited to the fibula, the distal third of the bone is affected.

ETIOLOGY

The pathogenesis of osteofibrous dysplasia remains unknown; however, several theories have been proposed: (1) it results from excessive resorption of bone with fibrous repair of the defect; (2) it is a congenital lesion or a variant of fibrous dysplasia; and (3) it results from abnormal blood circulation in the periosteum. Most recent literature has suggested either that the disorder is a reactive process secondary to adamantinoma or that it is a precursor of adamantinoma.*

*See references 25, 37, 116, 124, 155, 265, 285.

PATHOLOGY

On gross inspection the periosteum is intact. The affected cortex is thinned and may be perforated. The lesion has been described as either whitish yellow or reddish.

Histologically, the tissue is similar to fibrous dysplasia, with irregular spicules of trabecular bone and fibrous or collagenous stroma. In contrast to fibrous dysplasia, the spicules are usually lined with osteoblasts (Fig. 37–15). The finding of woven bone with juxtaposed lamellar bone (from osteoblasts) is thought to be characteristic of osteofibrous dysplasia. Recent immunohistochemical studies have demonstrated isolated cytokeratin-positive cells in the stroma of osteofibrous dysplasia. These cells are not seen in fibrous dysplasia but are found in adamantinomas. Based on this finding, it is believed that there is a relationship between osteofibrous dysplasia and differentiated adamantinoma; however, it is not known whether osteofibrous dysplasia represents a possible precursor or is a secondary reaction to adamantinoma. Nests of epithelial cells are a consistent histologic finding in adamantinomas, but they are not found in osteofibrous dysplasia.

CLINICAL FEATURES

The common presenting complaint is firm swelling localized over the tibia with associated mild to moderate anterior tibial bowing. Osteofibrous dysplasia is usually painless unless there is a coexisting pathologic fracture.
RADIOGRAPHIC FINDINGS

The lesion usually is extensive, involving the anterior cortex of either the diaphysis or the metaphysis of the tibia; the epiphysis usually is not affected. Characteristic eccentric, intracortical osteolysis is found, with moderate or marked expansion of the cortex (Fig. 37–16). In small areas, the cortex may actually appear absent and a "bubbled" appearance may be evident. In some areas, the osteolytic areas may have a ground-glass appearance. The tibia may be bowed anteriorly or anterolaterally. If the fibula is involved, it is usually evident in the distal third of the bone, involving nearly the entire circumference of the shaft.

DIFFERENTIAL DIAGNOSIS

The two entities that must be distinguished from osteofibrous dysplasia are monostotic fibrous dysplasia and
Solitary Osteochondroma

Osteochondroma is the most common benign bone tumor, reportedly accounting for 36 to 41 percent of all such tumors. It is characterized by a cartilage-capped osseous projection protruding from the surface of the affected bone. The exostosis is produced by progressive enchondral ossification of the hyaline cartilaginous cap, which essentially functions as a growth plate. More than 50 percent of solitary osteochondromas occur in the metaphyseal area of the distal femur, proximal tibia, and proximal humerus. Other areas in which solitary osteochondromas may be found include the distal radius, distal tibia, proximal and distal fibula, and occasionally the flat bones such as the scapulae, ilium, or ribs.

ETIOLOGY

A focal herniation of the medial or lateral component of the epiphyseal plate results in the formation of an aberrant cartilage-capped eccentric small bone. Several theories have been proposed to explain this phenomenon. Virchow in 1891 put forth the physis theory, according to which a portion of the physis cartilage becomes separated from the parent tissue, rotates 90 degrees, and grows in a direction transverse to the long axis of the bone. However, he did not provide an explanation for the separation and rotation of the detached physis cartilage. In 1920, Keith proposed that the cause was a defect in the perichondral ring surrounding the physis. Müller in 1913 theorized that the exostoses were produced by small nests of cartilage derived from the cambium layer of the periosteum. By producing osteochondroma using physis cartilage transplantation, D’Ambrosia and Ferguson provided support for the physis plate defect theory. Current thought is that the cause is misdirected growth of a portion of the physis plate, with lateral protrusions causing the development of the eccentric cartilage-capped bony prominence.

Unlike the more extensive hereditary (autosomal dominant) multiple exostoses, solitary osteochondromas do not appear to be genetically transmitted.

PATHOLOGY

Solitary osteochondromas may be sessile (broad-based) or pedunculated (narrow-based). The surface usually is lobular, with multiple bluish gray cartilaginous caps covering the irregular bony mass. The cartilaginous caps are covered by either thin or comparably thick perichondrium, which may be adherent to the underlying irregular surface and is continuous with that of the adjacent bony cortex. When this perichondrium is removed, the shiny cartilaginous cap is exposed. The cartilaginous cap is usually 1 to 3 mm thick, but in the younger patient it may be noticeably thicker. The thickness of the cartilaginous cap may be much greater if the tumor has undergone sarcomatous change. On cut section the cartilaginous cap will vary in thickness and often has an opaque yellow appearance owing to calcification within the cartilaginous matrix.

The tumor often resembles a cauliflower; however, it may also be flat, hemispheric, or tubular with a prominent end. Its base is contiguous with the normal cortical bone, and
the interior of the lesion (spongiosa bone) blends with that of the host bone.

A bursa may develop over the osteochondroma, particularly in larger lesions, where movement of the adjacent soft tissue leads to irritation. This bursal sac may contain mucinous fluid and fibrinous rice bodies.

Histologically, the cartilaginous cap is composed of bland hyaline cartilage. Variable degrees of cellularity are seen, but anaplastic cells are not characteristically evident. Normal enchondral ossification is seen at the cartilage-bone junction. In younger patients, cartilage cores may be present within the subchondral spongiosa near the physis, and these
cores may be responsible for recurrence of the lesion should an incomplete resection be performed. Aside from the cartilage cores, the cancellous bone underlying the cartilaginous cap resembles that of the host, although on occasion the marrow in the interior is predominantly fatty.

The appearance of the cartilaginous cap depends on the stage of growth, becoming thinner over time. Remnants of the quiescent cap may persist well into adult life. Should increased thickness of the cartilage become evident in an adult, malignant degeneration must be considered and the lesion should be carefully examined on histologic sections.

**CLINICAL FEATURES**

In the majority of affected individuals, the osteochondroma becomes evident between the ages of 10 and 20 years. There is a slight male preponderance. An osteochondroma may be discovered as an incidental radiographic finding or it may be detected on palpation of a protruding bump. Other factors that often draw attention to the osteochondroma include localized pain, growth disturbance of an extremity, compromised joint motion, abnormal cosmetic appearance, or secondary impingement of soft tissues (tendon, nerves, and vessels). On occasion, a fracture may occur through a stalk of a pedunculated (narrow-based) lesion following minor trauma.

**RADIOGRAPHIC FINDINGS**

There are several pathognomonic radiographic findings associated with an osteochondroma (Fig. 37–17).\(^{186}\) (1) The lesion protrudes from the host bone on either a sessile (broad-based) or pedunculated bony stalk. (2) It occurs either in the metaphysis or, as the main epiphyseal plate grows away from the lesion, in the diaphysis. It is never found in the epiphysis. (3) The cortex and cancellous bone of the osteochondroma blend with the cortex and cancellous bone of the host. This is the main radiographic finding, and any deviation from this feature should raise suspicion of a more serious lesion. (4) The lesion ranges in size from 2 to 12 cm.
radiographs. If there is evidence of cord compression, CT and MRI will clearly show the impingement.

Bursal osteochondromatosis overlying an osteochondroma of the rib has been described and, on occasion, ultrasound may delineate the extent of the swollen bursa. If there is concern over a progressively enlarging mass, ultrasound may also help determine the thickness of the cartilaginous cap. Steady growth of the cartilaginous cap is acceptable during childhood and early adolescence, but growth should cease when skeletal maturity is reached. If the cartilaginous cap continues to grow after skeletal maturity, malignant transformation should be considered and the appropriate follow-up studies undertaken.

DIFFERENTIAL DIAGNOSIS

Because of their typically distinct radiographic appearance, solitary osteochondromas are usually easily diagnosed. On occasion, they may be confused with a juxtaarticular chondroma or, less commonly, with myositis ossificans with a cartilaginous cap. Juxtaarticular chondromas usually have a scalloped cortical defect with a sclerotic margin. With myositis ossificans, the apparent tumor does not blend with the cortex and cancellous bone of the host bone, even though it may be attached to the peristeum. This is usually apparent radiographically, thus distinguishing the long-standing lesion of mature myositis ossificans from an osteochondroma. In the skeletally mature individual, enlargement of a solitary osteochondroma (particularly one that is associated with progressive discomfort) must alert the physician to the possibility of malignant degeneration into a chondrosarcoma.

TREATMENT

Because a solitary osteochondroma is a benign tumor, it does not need to be surgically excised if it is asymptomatic. Excision usually is reserved for those lesions that cause pain or symptomatic impingement on neurovascular structures or that interfere with joint function. Pain usually becomes an issue when an osteochondroma is repeatedly bumped on its prominence, if a pedunculated base fractures following trauma, or if a painful bursa develops. Neurovascular impingement may include peroneal nerve compression at the knee, median nerve compression at the wrist, or, rarely, spinal cord compression from a vertebral osteochondroma. Sometimes the osteochondroma is considered cosmetically unacceptable and the adolescent will ask to have it removed, preferring a scar to a bump. Finally, surgical excision is indicated any time there is a possibility of malignant transformation of an underlying osteochondroma, as demonstrated by an increase in the size of the lesion or in symptoms following skeletal maturity.

Excision of osteochondromas should, if possible, be postponed until late adolescence, for the following reasons. First, the growth potential of osteochondromas in younger children is unknown, and the full extent of the tumor cannot be appreciated until its growth potential is recognized. Second, because there are small pockets of cartilaginous cores within the spongiosa bone of osteochondromas in young children, the risk of local recurrence following excision is significant. In this situation, if the osteochondroma is removed, the perichondrium and the periosteum along the base of the lesion need to be excised. Finally, because of this dissection, the potential for growth arrest exists if the osteochondroma is very near a physis. In the maturing adolescent, the excision does not need to be quite as extensive, because the potential for recurrence is considerably less.

Occasionally a peripheral nerve (e.g., the peroneal nerve at the fibular head or the radial nerve along the humerus) may be in close proximity to the underlying lesion. Preliminary dissection of the nerve above the lesion can help avoid inadvertent injury to the nerve during excision of the osteochondroma. This anatomic situation, though, is more likely to occur with multiple hereditary exostoses rather than with a solitary osteochondroma.

SARCOMATOUS CHANGE

Malignant degeneration of a peripheral solitary osteochondroma leads to chondrosarcoma. However, malignant degeneration of solitary osteochondromas is rare, probably happening in less than 0.25 percent of lesions. Although Jaffe stated that 1 percent undergo malignant change and Dahlin reported an incidence of 4.1 percent in solitary osteochondromas treated surgically at the Mayo Clinic, these figures represent select cases referred to oncology centers. Malignant change evolves very slowly, usually occurring in adult life. When malignant changes occur, the lesions become painful and show evidence of growth.

MRI has been found useful in evaluating possible sarcomatous deterioration. The imaging modality delineates extension of the tumor mass into the adjacent soft tissues and allows proper planning of a wide resection of an underlying chondrosarcoma. Scintigraphic imaging with technetium-99m methylene diphosphonate has not been shown to qualitatively differentiate benign active exostoses from chondrosarcoma. Likewise, gallium scans cannot sufficiently distinguish between benign and sarcomatous lesions. Imaging criteria differentiating osteochondroma from chondrosarcoma are provided in Table 37–1.

Biopsy prior to surgical excision of a presumed chondrosarcoma may be of limited value, as there is a significant chance of a nonrepresentative biopsy and a potential risk of seeding the biopsy tract. The prognosis following excision of a chondrosarcoma is excellent.

Hereditary Multiple Exostoses

Hereditary multiple exostoses is an autosomal dominant condition affecting numerous areas of the skeleton that have been performed in cartilage. The overall prevalence has recently been reported to approach one in 50,000, which doubles the previously reported prevalence of one in 100,000. The disorder is known by a variety of terms: multiple hereditary osteochondromas, cartilaginous exostosis, diaphyseal aclasis (stressing abnormality of the modeling process), multiple osteochondromatosis, chondral osteoma, chondral osteogenic dysplasia of direction, deforming chondrodysplasia, hereditary deforming chondrodysplasia, and multiple cancellous exostoses. The term most commonly used today, hereditary multiple exostoses, was proposed by Jaffe in 1943.

The median age at the time of diagnosis in affected indi-
TABLE 37-1  **Criteria Differentiating Osteochondroma from Chondrosarcoma**

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Osteochondroma</th>
<th>Chondrosarcoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relation to parent bone</td>
<td>Continuity of cortex and medullary cavity with parent bone</td>
<td>Gradual loss of continuity of cortex</td>
</tr>
<tr>
<td>External surface of tumor</td>
<td>Distinct, well demarcated</td>
<td>Fuzzy and indistinct</td>
</tr>
<tr>
<td>Cartilaginous cap (best visualized on MRI)</td>
<td>Thin, &lt;1 cm</td>
<td>Thick, &gt;3 cm, lobulated, extending into soft tissues</td>
</tr>
<tr>
<td>Matrix pattern</td>
<td>Dense at periphery with solid cortex</td>
<td>Periphery granular in appearance with small areas of rarefaction and disorganized calcification</td>
</tr>
<tr>
<td></td>
<td>Normal cancellous bone centrally</td>
<td>Later, blotchy areas of calcification within center of tumor with streaky densities extending peripherally</td>
</tr>
<tr>
<td>Adjacent soft tissue</td>
<td>Normal</td>
<td>Large soft tissue mass containing disorganized areas of calcification</td>
</tr>
</tbody>
</table>


Individuals is approximately 3 years. Hereditary multiple exostoses has a penetrance of 50 percent by age 3½ years. By the end of the first decade of life, 80 percent of affected persons will have exostoses. By 12 years of age, nearly all affected individuals have evidence of exostoses, as penetrance of the disorder has been found to reach 96 to 100 percent (Fig. 37–18). Although some studies have reported that incomplete penetrance preferentially affects females, other investigations have shown no such reduction.

**ETIOLOGY**

The disorder is of autosomal dominant inheritance, with penetrance approaching 96 percent. If a person whose family is affected by hereditary multiple exostoses has not had an exostosis by 12 years of age, it is unlikely that exostoses will develop later. However, there remains a small risk that a particular individual will have affected children because the gene is nonpenetrant in approximately 4 percent of carriers. Approximately 10 percent of affected individuals have no family history of hereditary multiple exostoses.

Numerous genetic studies have found anomalies on chromosomes 8, 11, and 19, making this a genetically heterogeneous disorder. Specifically, the three loci include 8q24.1 (EXT1), 11p11–12 (EXT2), and 19p (EXT3). The EXT1 and EXT2 genes show extensive sequence homology to each other. These genes are deleted in exostoses-derived tumors, supporting the hypothesis that they encode tumor suppressors. Still another gene, EXT3 (or EXT-like), has been identified that shows a striking sequence similarity to both EXT1 and EXT2. All of this information suggests that a family of related genes may be responsible for the development of hereditary multiple exostoses.

**PATHOLOGY**

The gross pathologic and microscopic features of hereditary multiple exostoses are similar to those described for solitary osteochondromas.

*See references 4, 29, 38, 69, 157, 164, 217, 218, 277, 296, 301, 309, 310.

FIGURE 37-18 Hereditary multiple exostoses in a 12-year-old boy. Extensive involvement of the femurs, tibias, and fibulas was evident clinically and radiographically.
CLINICAL FEATURES

Multiple exostoses usually manifest during early childhood (although rarely before 2 years of age) with several knobby, hard, subcutaneous protuberances near the joints. Numerous sites can be involved. On presentation, five or six exostoses typically may be found, involving both the upper and lower extremities. The “knobby” appearance of the child is so characteristic that one can usually make the diagnosis by clinical inspection alone. Over time, the upper and lower extremities may appear short in relationship to the trunk. Shortening of the limbs is usually disproportionate. Approximately 10 percent of affected individuals will have a lower limb length inequality. Patients are not considered dwarfs, and the trunk-limb growth difference usually does not become obvious until the pubertal growth spurt. On occasion, concerned affected parents may bring in their normal-appearing child for screening.

For those affected with hereditary multiple exostoses, 70 percent will have involvement of the distal femur, 70 percent the proximal tibia, and 30 percent the proximal fibula (Fig. 37–19). The likelihood of involvement near the knee is at least one of these three locations is approximately 94 percent. The proximal humerus is affected in 50 percent of cases, the scapula and ribs in 40 percent, the distal radius and ulna in 30 percent, the proximal femur in 30 percent, the phalanges in 30 percent, the distal fibula in 25 percent, the distal tibia in 20 percent, and the bones of the foot in 10 to 25 percent. Tibia-fibular synostosis often develops from chronic apposition of osteochondromas proximally or distally but rarely causes symptoms or functional impairment. As the lesions enlarge, they may cause discomfort secondary to pressure on adjacent soft tissues, and they may hinder normal joint mobility.

Osteochondromas of the proximal humerus are often readily palpable but rarely cause neurologic dysfunction (Fig. 37–20). However, because of their proximity to major nerves, great care must be taken if resection is necessary. The scapula is involved in 40 percent of affected individuals, with osteochondromas located on the anterior or the posterior aspect of the scapula. The presence of osteochondromas on the anterior aspect of the scapula may lead to discomfort during scapulothoracic motion. Winging of the scapula due to the presence of osteochondromas has been described.70

Obvious deformity in the forearm is seen in 39 to 60 percent of patients.116,241,267 The ulna is shorter than the radius and the radius is bowed laterally, with its concavity toward the short ulna (Fig. 37–21). Often the distal end of the ulna is more severely affected than the distal end of the radius, leading to this discrepancy in length. A mild flexion deformity of the elbow is usually present. Loss of forearm pronation and supination occurs with increasing age.267 Dislocation of the radial head occurs and is usually associated with a negative ulnar variance. The resulting forearm deformities are usually asymmetric. Often, the patient’s main complaint is an undesirable cosmetic appearance. The natural history of these deformities has been described as progressive, with variable weakness, functional impairment, and worsening cosmetic deformity of the extremity. Some authors, however, report that the deformities are well tolerated and lead to little loss of function.267

Deformity of the hand is uncommon. The main area of involvement appears to be around the metacarpophalangeal
joint, but the proximal interphalangeal joint is the most common site of deformity. Metacarpal shortening usually does not cause functional problems and does not need to be treated. Angular deformity, although uncommon, does cause problems and requires surgical intervention. There is no evidence that deformity can be prevented by early excision of the osteochondromas.

In the lower extremity, valgus of the proximal tibia is frequently present and is nearly always found in the proximal metaphyseal region of the tibia. The valgus progressively increases during growth spurts. Occasionally there will be some angulation distally at the femur, and there may be recurrent dislocation of the patella.

Osteochondromas of the proximal femur may lead to progressive hip dysplasia which, on occasion, will require corrective varus osteotomy (Fig. 37–22). Adverse effects on femoral growth are less with proximal involvement than with distal involvement.

Although osteochondromas near the ankle are not uncommon, the reported prevalence of ankle deformities ranges widely, from 2 percent to 54 percent.

The characteristic ankle valgus deformity is often accompanied by decreased ankle motion. This deformity is caused either by retardation of the normal distal fibular growth or by deficient growth of the lateral half of the distal tibial epiphysis. The severity of the ankle valgus varies, as the distal fibular physis progressively rises in relation to the tibial-talar articular space.

There have been numerous reports of spinal cord impingement by vertebral osteochondromas. The cervical, thoracic, or lumbar region can be affected. Lower extremity discomfort associated with decreased balance, impaired coordination, or other central neurologic dysfunction should raise the consideration of a vertebral osteochondroma. The presence and extent of the lesion are best delineated with CT, while MRI of the spinal cord demonstrates the area of spinal cord impingement.

RADIOGRAPHIC FINDINGS

Unlike solitary osteochondromas, hereditary multiple exostoses involve a significantly greater portion of the metaphysis or diaphysis and are generally more irregular in shape. Over time, lesions that begin in the metaphyseal region migrate into the diaphysis of the long bones. The exostoses vary in number, size, and configuration. Like solitary osteochondromas, they may be sessile or pedunculated, cauliflower-like, or even narrow with a pointed end. They nearly always point away from the physis. In nearly 95 percent of cases, evidence of the osteochondroma will be found around the knee. Thus, the disorder can often be confirmed with radiographs of the knee. Irregular zones of calcification may be present, particularly in the cartilaginous cap. In older individuals, however, extensive calcification with changes in the shape of the cartilaginous caps suggests possible malignant degeneration.

DIFFERENTIAL DIAGNOSIS

The numerous lesions associated with hereditary multiple exostoses make the radiographic findings pathognomonic of the disorder. If painful growth of an osteochondroma becomes evident in a mature individual with hereditary
multiple exostoses, chondrosarcomatous transformation must be ruled out.

**TREATMENT**

The only treatment for hereditary multiple exostoses is surgery. However, because the exostoses are numerous and many are asymptomatic, a cautious approach is called for. The mere presence of an osteochondroma is not an indication for surgery. Reasonable indications include (1) pain from external trauma or irritation of surrounding soft tissues, (2) growth disturbance leading to angular deformity or limb length discrepancy, (3) joint motion compromised by juxta-articular lesions, (4) soft tissue (tendon, nerve, or vessel) impingement or tethering, (5) spinal cord compression, (6) false aneurysm produced by an osteochondroma, (7) painful bursa formation, (8) obvious cosmetic deformity, and (9) a rapid increase in the size of a lesion. During childhood, numerous operations may be necessary, and this prospect should be discussed in detail with the parents soon after the disorder is diagnosed. Life expectancy is normal unless malignant degeneration of an osteochondroma has occurred and metastases have developed.

Osteochondromas involving the forearm frequently lead to surgical intervention. Early excision of the lesions on the radius and ulna does not alter or correct existing deformity, but it may delay progression of the deformity. If ulnar shortening has occurred with bowing of the radius, lengthening of the ulna combined with distal radial hemiepiphyseal stapling has been found effective in correcting the deformity. Ulnar lengthening is best done gradually by distraction osteogenesis. Creation of a “one-bone forearm” has been successful as a salvage procedure for severely affected forearms. Deformities of the forearm should be treated early and aggressively in an effort to prevent further progression and to reduce disability. A prominent, symptomatic, dislocated radial head can be safely excised following skeletal maturity.

Marked shortening of the humerus may result from a proximal physeal growth disturbance. If the discrepancy is significant, distraction osteogenesis has been successful in increasing the length of the humerus.

For skeletally mature individuals with significant genu valgum, varus osteotomy of the proximal tibia can result in an improved appearance. However, if an osteochondroma is present in the proximal fibula, any osteotomy of the proximal tibia or fibula carries a significant risk of peroneal nerve palsy. In the skeletally immature patient, stapling along the medial side of the proximal tibial physis or distal femoral physis may be sufficient to correct the valgus. If correction is achieved by this method, the staples should be left in longer (slight overcorrection) to prevent recurrence of the deformity following staple removal.

Ankle valgus is treated either by medial distal tibial physeal arrest in the skeletally immature patient or by a varus corrective osteotomy in the more mature individual (Fig. 37–23). Tibia-fibular synostoses frequently occur distally, but there usually are no symptoms or functional impairment from this occurrence and the condition does not need to be surgically treated.

**SARCOMATOUS CHANGE**

Transformation of a lesion in hereditary multiple exostoses to chondrosarcoma during childhood is exceedingly rare. In general, transformation in adulthood remains uncommon, with current reports indicating the risk to be 0.9 to 5 percent. The apparent risk of malignant degeneration may become greater as the duration of follow-up increases. In Schmale and associates' 1994 study, two-thirds of the individuals with transformations were younger than 40 years when the authors reported an incidence of 0.9
FIGURE 37-23  A, Excessive ankle valgus accompanied by a distal tibiofibular synostosis led to discomfort. B to D, A varus closing wedge osteotomy in the distal tibial/fibular region resulted in improved alignment and resolution of the patient's discomfort.
percent. At this time, the lifetime risk of chondrosarcoma is estimated to be approximately 1 to 2 percent. The genetic abnormalities found on chromosomes 8, 11, and 19 (tumor suppressor genes) may play a role in the development of chondrosarcoma.

The most frequent sign of sarcomatous change is a painful, enlarging mass, usually one of long duration. The relative frequency of chondrosarcomas is reported to be highest with osteochondromas of the pelvis or shoulder girdle. The clinical course of the tumor is slow and metastasis occurs late, usually to the lungs.

If the cartilaginous cap of the exostoses exceeds 1 cm in thickness, malignancy should be suspected. Inadequate surgical removal almost always results in recurrence. Patient prognosis is good if metastasis has not occurred. Radiation therapy has no effect on the tumor.

**Solitary Enchondroma**

Intramedullary cartilaginous solitary enchondromas are relatively common, accounting for approximately one-fourth of all benign tumors. Unlike multiple enchondromatosis, which is frequently diagnosed during childhood, solitary enchondromas are usually diagnosed after the second decade of life. The peak age of presentation is approximately 35 years.

**PATHOLOGY**

Enchondromas appear as glistening white, grayish white, or pearly tissues that have a gritty feel on palpation, owing to the intrinsic calcification. The tumor is easily cut with a knife, as if it were soft chalk.

Histologically, enchondromas are proliferating nests of cartilage cells that lack obvious atypia. Foci of calcification are present. Plates of lamellar bone surround the lobules of cartilage in a partial to complete circumferential manner. Invasive infiltration of the bone marrow spaces is not characteristic of benign solitary enchondromas.

Although mitotic figures within the dysplastic cartilaginous lesions may be found in specimens from growing children, the likelihood of malignancy is low. In the adult, however, it may be difficult to differentiate an enchondroma from a low-grade sarcoma. In general, small peripheral cartilage tumors are usually benign, whereas the large axial tumors in the adult are more likely to be malignant. A malignant change of a solitary enchondroma in childhood or adolescent would be a rare event.

**CLINICAL FEATURES**

Nearly 50 percent of diagnosed solitary enchondromas occur in the hand, involving the phalanges in particular. The carpal bones are occasionally affected. Of the long tubular bones, the femur and humerus are frequent sites of localization. Occasionally the ribs, sternum, innominate bones, and vertebral columns may also be affected.

Clinically, enchondromas in the fingers are generally diagnosed following local trauma. Some patients, however, present with a firm, local swelling in the region of the affected phalanx or metacarpal without a fracture or local pain. Nearly 75 percent of patients with enchondromas involving the hands or feet have a solitary lesion. The remaining patients have multiple enchondromatosis. When the solitary enchondroma involves the femur or humerus, it is usually quiescent, with no clinical signs evident until adulthood.

**RADIOGRAPHIC FINDINGS**

Solitary enchondromas usually appear as well-delineated lucent defects in the metaphyseal region of long bones. In the phalanges of the hand or foot, the entire shaft may be involved. The cortical rim is generally intact unless a fracture has occurred through the weakened bone (Fig. 37–24). Calcification is usually present within the lesion and appears as fine punctate stippling. In the larger long bones, calcification may be more pronounced, which can make the differentiation between enchondroma and bone infarct difficult. Most lesions are 3 to 4 cm in size, with a range of 1 to 8 cm. Generally there is no evidence of focal cortical erosion, scalloping of the cortex, significant cortical thickening, or bone expansion. In the older symptomatic adolescent, cortical thinning and expansion in the larger long bones can be troubling because these radiographic findings may represent features of low-grade chondrosarcoma.

CT is useful for evaluating cartilaginous tumors, especially in the long bones, pelvis, and spine. Technetium bone

**FIGURE 37–24** A solitary enchondroma in the middle phalanx of the ring finger. Radiographs showed a well-delineated lucent lesion with an intact cortical rim. It was noticed because of mild painless swelling of the digit.
scanning generally is not necessary when evaluating a child with presumed enchondroma.

DIFFERENTIAL DIAGNOSIS

Confusion between an enchondroma and a bone infarct may occur when the radiolucent lesion has a significant amount of calcification. However, in general, the calcification seen with bone infarcts is more peripherally located. In the phalanges of the hand and foot, solitary enchondromas may be difficult to differentiate from epithelial inclusion cysts. In a metacarpal, it may be difficult to distinguish a solitary enchondroma from a small solitary bone cyst, a nonossifying fibroma, or a focus of fibrous dysplasia.

TREATMENT

For solitary enchondromas in the hand, complete curettage followed by autogenous bone grafting usually results in cure. Similar treatment is undertaken for solitary enchondromas in the long bones, where the recurrence rate remains low. If an en bloc wide excision were performed, the recurrence rate would be even lower. However, the possibly unacceptable postoperative functional deficit makes this more aggressive approach unnecessary.

The risk of malignant degeneration of an isolated enchondroma is rare in childhood. However, these lesions should be followed over time. Enchondromas in the pelvis, scapulae, sternum, vertebrae, and proximal area of the large long bones have a greater likelihood of malignant transformation in adulthood.

Multiple Enchondromatosis (Ollier’s Disease) and Maffucci’s Syndrome

Ollier in 1889 described a condition of multiple, typically unilateral enchondromas associated with deformity of the extremity. He referred to the condition as dysenchondroplasia, implying that it resulted from a developmental defect related to abnormal growth of cartilage.

PATHOLOGY

On gross inspection of the lesions, the bones show numerous islands of glistening cartilage, which are generally located in the diaphyseal and metaphyseal regions. Infractrarily, abnormal cartilage may also be seen in the epiphyseal region. This close proximity of the tumor to the physis can lead to profound inhibition of growth, resulting in severe limb length discrepancies or angular deformities. As longitudinal growth continues, the abnormal enchondroma cartilage derived near the epiphyseal plate will form long linear masses within the shaft. This phenomenon is seen only in Ollier’s disease and explains the pathognomonic “fanlike” metaphyseal septation seen on radiographs. The dense lines represent bone formed by normal enchondral ossification and the intervening columns of lucency represent the epiphyseally derived areas of abnormal cartilage.

The histologic features in multiple enchondromatosis generally resemble those of a solitary enchondroma. However, in multiple enchondromatosis the appearance might be that of a highly cellular lesion with ominous nuclei mimicking a low-grade chondrosarcoma. Eventual malignant transformation into chondrosarcomas has been reported to occur in 20 to 33 percent of those affected. Therefore, communication between the surgeon, pathologist, and radiologist is imperative when making the diagnosis in the older adolescent or adult. In general, aggressive histologic findings are still likely to represent a benign lesion if the biopsy site is the hand rather than the long bones or the pelvis. Conversely, if the biopsy site is the pelvis or larger bones, suspicion of low-grade sarcomas is higher. The atypical cellular features in Ollier’s disease may explain the relatively high incidence of chondrosarcomatous transformation in older individuals who are moderately or severely affected.

CLINICAL FEATURES

Multiple enchondromatosis is an uncommon, nonhereditary disorder. The physical signs related to this condition commonly begin in childhood and vary with the extent of the lesions and their effect on the weakened bones. The number of bones affected can vary greatly, with the phalanges, femur, and tibia most commonly affected (Fig. 37–25). Because of the tendency toward unilateral involvement, severe lower limb length discrepancies and angular deformities (most commonly genu valgum) result. Discrepancies may be in the range of 10 to 25 cm by maturity. Deformity and enlargement of fingers may impair normal function. Forearm abnormalities such as bowing, limited rotation, and ulnar deviation of the hand may be evident.

Maffucci’s syndrome is a condition of enchondromatosis associated with multiple hemangiomas involving soft tissue. They may be associated with superficial phleboliths, which can appear on radiographs as roundish opacities in the soft tissues. Hemangiomas have also been noted to occur within internal organs. Multiple pigmented nevi and vitiligo are other occasional nonskeletal manifestations. An abnormality in neuropeptides has been identified, and their presence appears to stimulate growth of abnormal blood vessels.

Another related entity, generalized enchondromatosis, represents a more severe expression of the disorder. Nearly all of the metaphyseal regions in all of the long and short tubular bones are affected.

RADIOGRAPHIC FINDINGS

Bone abnormalities are usually more extensive than the physical examination would suggest. In the long bones, enchondromatosis is recognized as radiolucent longitudinal streaks that involve the metaphysis and extend down into the diaphysis. Epiphyses are usually not affected but, as reported recently, may be involved. The cortex overlying the enchondroma is usually thin, and calcification within the lesion is common. Significant shortening and angular deformity are frequently noted in the involved long bones, whether in the hands, feet, or limbs.
CT is useful for evaluating multiple enchondromatosis, particularly in the long bones and pelvis. CT clarifies cortical and osteal scalloping better than plain radiography and allows precise comparisons to be made over time.

**TREATMENT**

Because of the extent of the disease, multiple enchondromatosis cannot be cured by curettage and bone grafting. The numerous deformities that often accompany multiple enchondromatosis require repeated operative interventions over several years to correct the angular deformities and achieve similar limb lengths at maturity. These procedures include osteotomies, limb lengthenings, and epiphysiodeses (Fig. 37–26). In the older child, use of the Ilizarov apparatus is the best method to achieve both angular correction and equalization of limb lengths. Use of multiple wires or half-pins allows sufficient purchase in the enchondromatous bone so that lengthenings can be successfully achieved.

**MALIGNANT POTENTIAL WITH ENCHONDROMATOSIS**

The incidence of secondary chondrosarcoma in patients with Ollier’s disease is approximately 25 to 30 percent by age 40 years. Those patients with Maffucci’s syndrome have a similar or higher likelihood of developing malignant de-
FIGURE 37–26 Findings in a 4-year-old boy with Ollier’s disease. A, Radiograph demonstrating unilateral left lower extremity involvement. B, A valgus ostectomy of the distal femur was performed to correct the notable genu varum. C, Two years later, the genu varum recurred, owing to the abnormal growth at the distal femoral physis. Another ostectomy was planned; future limb lengthening will be needed.

generation,7,8 with Schwartz and associates reporting nearly 100 percent expectation.25 Over the long term, periodic surveillance of the brain and abdomen for occult malignant lesions is indicated in patients who have enchondromatosis.122,123,124 Increased localized growth of a lesion in an extremity accompanied by pain is the hallmark of possible malignancy. In such a situation, biopsy of the lesion is indicated. Hematopoietic malignancies (acute lymphoid leukemia) have been described in association with Maffucci’s syndrome.226

Chondroblastoma

Chondroblastomas are uncommon benign cellular cartilage tumors that are most often located in the epiphyses of the long bone of the extremities.121,122 Chondroblastomas are twice as common in males as in females. The peak age of occurrence is in the second decade, the majority of patients presenting before age 30 years.

Chondroblastoma was first described in detail by Codman in 1931.19 He called the lesion an epiphysial chondromatous giant-cell tumor. Prior to his description, the tumors were often thought to represent chondrosarcomas. Today, chondroblastomas are still occasionally referred to as Codman’s tumor.

ETIOLOGY

Jaffe and Lichtenstein conjectured that the lesion arises from cartilage “germ cells” or cells of the epiphyseal cartilage.125 However, because of reports of the lesion involving the skull or rib (where cartilage “germ cells” would not likely be found), this hypothesis is difficult to confirm.

Abnormalities in chromosomes 5 and 8 have been reported in chondroblastoma; however, specific locations have not been clearly identified.223

PATHOLOGY

Gross specimens obtained at the time of curettage are often characterized by pieces of gray-pink or hemorrhagic tissues intermixed with gritty, calcified, cholesterol-laden tissues. There may be small islands of bluish to white chondroid. Because chondroblastomas are often found to contain cystic or degenerative areas, the amount of tissue removed may be less than expected based on the radiographic appearance.

Histologically, the tumor is characterized by polygonal cells (chondroblasts), giant cells, islands of chondroid or hyaline cartilage, “chicken wire” calcification, and nodules of calcification in the stroma (Fig. 37–27).309,313 The “chicken wire” calcification results when lacelike deposits of calcium are intermixed on the intercellular chondroid matrix.
A small percentage of chondroblastomas may be primarily cystic or hemorrhagic, making differentiation from aneurysmal bone cysts difficult histologically. Another more worrisome tumor, clear cell chondrosarcoma, may histologically resemble chondroblastoma. However, clear cell chondrosarcomas occur in adults with closed physes, and their radiographic appearance and clinical presentation would not be consistent with chondroblastomas.

**CLINICAL FEATURES**

The most common locations are the proximal humerus, distal femur, and proximal tibia. Chondroblastomas have also been found in the skull, maxilla, spine, ribs, pelvis, hands, patella, talus, and calcaneus. Multifocal, benign chondroblastomas have been reported. Nonepiphysial locations in the long bones have been described.

Symptoms usually are mild, consisting of pain and localized tenderness. The discomfort is often present for 6 months to several years prior to diagnosis. Because the lesion is epiphysial, the adjacent joint may be swollen and may have limited range of motion. If tumor is present in a lower extremity, an antalgic limp may be evident. Pathologic fractures are uncommon.

**RADIOGRAPHIC FINDINGS**

Chondroblastomas are usually located in the epiphyses, but they may extend into the metaphyseal region (Fig. 37–28). They are usually eccentric, involving less than one-half of the entire epiphysial. The lesion is rimmed by a border of host bone sclerosis, and small punctate calcifications are present within the tumor. Commonly, the physis adjacent to the lesion is present at the time of diagnosis. If all of these features are present, this radiographic appearance is pathognomonic for chondroblastoma.

CT will clearly demonstrate the extent of the lesion within the epiphysial region and its proximity to the physis or subarticular surface.

MRI findings associated with chondroblastoma have been reported. Adjacent bone marrow and soft tissue edema, as well as periosteal reactions, are more dramatically demonstrated on MRI than on plain radiographs. Bone marrow edema is common. Knowledge of the MRI findings of chondroblastoma may allow for more accurate diagnosis and help to avoid confusion with infection or aggressive neoplasms.

Fine-needle aspiration yields satisfactory material for interpretation and confirming the diagnosis.

**DIFFERENTIAL DIAGNOSIS**

The differential diagnosis includes giant-cell tumors, enchondromas, synovial lesions (e.g., pigmented villonodular synovitis, rheumatoid arthritis), and atypically located eosinophilic granuloma.

**TREATMENT**

Complete curettage and excision of the lesion is often successful. The surgeon should avoid interfering with the joint surface or disturbing the physes in the immature skeleton. The defect is filled with either autogenous or allograft bone. Although there is a definite risk of recurrence following intracapsular curettage, a recent study reported local control in approximately 80 percent of cases. Preservation of the physis should be considered a secondary concern compared to complete and thorough excision of the chondroblastoma.

If the tumor is beneath articular cartilage, adequate excision may occasionally require removal of some of the joint cartilage. Arthroscopy has been used as an adjunct in the excision of lesions in the proximal tibia and proximal femur.

Occasional chondroblastomas require wide marginal resection, especially those that have recurred locally. Reconstruction after marginal or wide en bloc resection usually requires a partial or full osteoarticular allograft.

Chondroblastomas have been known to undergo benign pulmonary metastasis. When found in the lung, these metastases are usually rimmed with bone. When identified, these lesions should be surgically removed to ensure proper diagnosis.

**Chondromyxoid Fibroma**

Chondromyxoid fibromas are rare, benign tumors representing less than 0.4 percent of biopsied primary bone tumors. They consist mainly of cartilaginous tissue intermixed with areas of myxomatous and fibrous elements. The myxomatous components are most probably due to degeneration of chondroid tissue, while the fibrous component may result from repair of the degenerated areas.

**ETIOLOGY**

Cytogenetic analysis of chondromyxoid fibromas has found an unbalanced reciprocal translocation between the short arm of chromosome 3 and the long arm of chromosome 6. Two known cartilage-related genes are located in the regions affected by this unbalanced rearrangement. These genes function to control growth and maturation of endochondral bone, the site of origin of cartilaginous tumors.
FIGURE 37-28 Chondroblastoma. A and B, AP and lateral radiographs of the left hip in a 16-year-old girl showing a subchondral radiolucent lesion in the femoral head (epiphysial region) that is rimmed by a border of host bone sclerosis. C, Bone scan shows significant uptake in the left femoral head. D, CT of the hips shows calcification within the lesion.
PATHOLOGY
Most chondromyxoid fibromas are less than 5 cm in size. The tissue is firm, grayish white, and often covered on the outer surface with a thin rim of bone or periosteum. Cysts or areas of hemorrhage may be found within the lesion (Fig. 37–29). Histologically, chondromyxoid fibromas have a lobulated pattern, with some of these lobules sparsely cellular and others more cellular. Those lobules that have few cells are composed of a myxoid or chondroid matrix. Microscopic areas of cystic degeneration may contribute to the myxoid appearance. Other features of a chondromyxoid fibroma include osteoclast-like giant cells, intermixed fibrous tissue, and occasionally cholesterol, hemosiderin, and lymphocytes. Distinct calcification is rare.

CLINICAL FEATURES
Chondromyxoid fibromas usually occur in older children and young adults, with individuals most commonly presenting for treatment in the second and third decades of life. Although most of the lesions are found in the tibia, other sites of predilection include the femur, fibula, metatar-
sals, and calcaneus. The upper extremities are rarely involved. Males and females are equally affected.

As with other benign bone tumors, it is not uncommon for a chondromyxoid fibroma to be discovered incidentally on a radiograph obtained for an unrelated reason. If symptoms are present from the lesion, the local discomfort usually is mild and intermittent. Swelling of the area and tenderness on palpation are occasionally noted.

**RADIOGRAPHIC FINDINGS**

Chondromyxoid fibromas are usually ovoid or round. They are slow-growing and usually evoke a border of reactive host bone sclerosis. This sclerotic border, most commonly seen in patients less than 20 years old, is a useful sign for determining the lesion's benign nature. Many of the tumors will have a trabeculated or bubbly appearance, but it is uncommon for calcification of the cartilage to be evident on radiographs. Chondromyxoid fibromas appear in the metaphyseal region of the long bones. They are usually eccentric and juxtacortical or even periosteal in location. It may be difficult to differentiate periosteal chondromyxoid fibromas from an aneurysmal bone cyst. In younger children, chondromyxoid fibromas may appear next to the physis, but with growth, the lesions tend to migrate away from the physis.

**DIFFERENTIAL DIAGNOSIS**

The radiographic appearance of a chondromyxoid fibroma can be very similar to that of a nonossifying fibroma. Both lesions are usually metaphyseal, eccentric, surrounded by a border of sclerosis, and trabeculated. Unlike nonossifying fibromas, chondromyxoid fibromas may bulge from the original bony contour. When the chondromyxoid fibroma is notably eccentric and associated with periosteal expansion, differentiation from aneurysmal bone cysts may be difficult. Other lesions to consider include solitary eosinophilic granuloma, enchondroma, simple bone cyst, and, in the older individual, worrisome entities such as chondrosarcoma or myeloma.

**TREATMENT**

Fine-needle aspiration cytology can be used to diagnose chondromyxoid fibroma. Many of these lesions have been effectively treated by simple curettage. In younger patients, however, incomplete removal may lead to recurrence, which is estimated to occur in as many as one-fourth of patients. Therefore, en bloc excision should be considered. Because of the benign nature of this tumor, the surgeon should avoid radical procedures. If the lesion is next to the physis, consideration should be given to delaying surgical intervention until the tumor has grown away from the physeal area. Radiation therapy and chemotherapy are not considered in the management of chondromyxoid fibromas. Chondrosarcomatous transformation is rare.

Osteoid Osteoma

Osteoid osteomas were described as a distinct entity by Jaffe in 1935. Earlier reports referred to this entity as "sclerosing nonsuppurative osteomyelitis," osteomyelitis of Garré, or "localized or cortical bone abscess."

Osteoid osteomas are solitary, benign, painful lesions of the bone. They have a nidus, 1.5 to 2.0 cm in size, that consists of osteoid, osteoblasts, and variable amounts of fibrovascular stroma. This nidus is surrounded by an area of reactive, dense bone. Osteoid osteomas are relatively common, benign bone lesions, exceeded in incidence only by osteochondromas and nonossifying fibromas. Osteoid osteomas account for approximately 10 to 11 percent of benign bone tumors and 2 to 3 percent of all primary biopsied bone neoplasms. They are characteristically seen in children and adolescents. The male-female ratio is approximately 2:1.

**ETIOLOGY**

The etiology of osteoid osteomas remains unknown.

**PATHOLOGY**

At the time of surgery, the cortical bone overlying the osteoid osteoma may be mildly pink compared with the surrounding cortex because of the increased local vascularity. The lesion itself is often a small, round or oval, cherry-red or reddish brown tumor 1.0 cm or less in diameter. Differentiation between osteoid osteoma and osteoblastoma depends on the size of the lesion: lesions less than 2.0 cm in diameter are technically classified as osteoid osteomas. The nidus may have a very dense, gritty texture if a significant amount of calcification is present, or it may be quite soft and granular if it is predominantly vascular with little calcification.

Histologically, osteoid osteomas are characterized by small spicules of immature trabeculae, most often lined by prominent osteoblasts and osteoclasts (Fig. 37-30). In mature lesions, the intervening stroma is sparsely cellular with readily apparent vascular spaces. Cartilage is not present. The demarcation between reactive surrounding bone and nidus is readily apparent microscopically. The pain associated with osteoid osteoma is thought to be caused by the numerous nonmyelinated axons that are present within the nidus.
CLINICAL FEATURES

The patient with osteoid osteoma typically presents with a history of dull, aching pain in the region overlying the affected long bone. The pain may have been present for several months before presentation, tends to be worse at night, and is relieved significantly by salicylates or nonsteroidal anti-inflammatory drugs (NSAIDs).

The most commonly involved site is the lower extremity, particularly the metaphyseal or diaphyseal region of the femur and tibia. Occasionally the tumor is periarticular in location. Less frequent sites of involvement include the humerus, spine, foot (talonavicular, and metatarsals), calvarium, maxilla, mandible, clavicle, scapula, ribs, pelvis, and patella.

A limp is often noted during evaluation of the patient’s gait. Muscle atrophy may be apparent if the lesion has been present for several months. However, direct tenderness, erythema, or swelling is uncommon. If the lesion is in the vertebral column (most commonly in the posterior elements), a secondary painful scoliosis may be evident. The concavity of the curvature is usually on the side of the lesion and is attributed to spasm of the paravertebral muscles. Excision of the nidus in the vertebral column will often result in complete resolution of the scoliosis.

RADIOGRAPHIC FINDINGS

The radiographic appearance depends on the location of the osteoid osteoma within the bone. Most of the tumors are intracortical, with the nidus appearing as a radiolucent lesion. This nidus rarely exceeds 1.0 cm in diameter but may be as large as 2.0 cm. The dense surrounding reactive sclerotic bone may extend for several centimeters away from the nidus. Less commonly, the osteoid osteoma may be intramedullary, subperiosteal, or periarticular in location. These atypical sites usually do not provoke reactive bone formation around the nidus. Calcification in the central portion of the nidus may be evident on radiographs.

Other radiologic studies may need to be made to correct diagnosis if the typical radiographic findings are not present or if the lesion is in an atypical location and lacks the associated reactive sclerosis. Technetium-99 bone scan is useful if osteoid osteoma is suspected but the lesion is not clearly demonstrated on plain radiographs. Technetium-99 bone scan nearly always demonstrates an intense focal increase in uptake of the technetium in the nidus and is of considerable value in evaluating the spine, pelvis, and long bones.

Once the general area of the lesion has been localized with bone scan, cross-sectional imaging with CT will best demonstrate the well-circumscribed area representing the nidus (Fig. 37–31). Thin sections (1.0 to 2.0 mm) may be needed for optimal detail.

MRI will demonstrate the soft tissue and bone marrow edema that accompanies osteoid osteomas. However, MRI is rarely needed because CT both better demonstrates the nidus and aids in differentiating between the nidus and reactive bone.

NATURAL HISTORY

Osteoid osteomas are described as self-limiting lesions that may mature spontaneously over the course of several years. The nidus will gradually calcify, then ossify, and finally blend into the sclerotic surrounding bone. During the maturation period, the local pain gradually diminishes. Knowing this, some clinicians advocate conservative management of osteoid osteomas, with NSAIDs or aspirin recommended for those patients choosing not to undergo operative intervention. In reality, though, very few patients are willing to continue with conservative management because of the intensity of the pain and the favorable outcomes likely with surgery.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis includes subacute osteomyelitis and osteoblastoma. On radiographs, a quiescent bone abscess may appear very similar to osteoid osteoma. Laboratory studies further assist in distinguishing between the two entities. Local aspiration of subacute osteomyelitis will confirm the diagnosis.

Osteoid osteomas are generally differentiated from osteoblastomas by size (osteoblastomas are larger, usually exceeding 2 cm in diameter), degree of sclerosis (osteoid osteomas, in general, have a greater degree of surrounding dense bone), and natural history (osteoblastomas can be more aggressive).

TREATMENT

Salicylate and NSAIDs are effective in relieving symptoms of pain associated with osteoid osteoma. If the symptoms are moderate and controlled by this treatment program, observation alone is sufficient. The possibility of spontaneous improvement over the course of several years may make medical management feasible for some patients. However, it is not possible on an individual basis to conclusively determine the ultimate outcome with medical management. Most families understand this and elect a surgical approach to this benign lesion.

The purpose of surgery is to eradicate the pain-producing nidus. Surgical excision has proved extremely effective in accomplishing this goal. More recently, less invasive maneuvers utilizing pinpoint CT-guided localization have become popular. With either approach, once the nidus is removed, the surrounding sclerotic bone will usually remodel. Relief from the pain is immediate, dramatic, and permanent unless the nidus has been incompletely excised. Patients often remark that the incisional pain is far different from the pain of the osteoid osteoma itself.

Accurate intraoperative localization of the nidus is crucial for the success of surgical intervention. Radiography, CT, tetracycline labeling, and bone scintigraphy have all been used for this purpose. Conventional intraoperative radiographs of the excised specimen may help confirm the presence of the nidus. CT-guided exploration, performed under anesthesia in the radiology suite, is helpful in localizing the nidus itself but will be inconclusive regarding the excision of the nidus. Tetracycline labeling can be used in children more than 8 years old. The risk of permanent staining of the dentin may preclude use of this technique in younger children. A dental consultation may be useful in determining the maturity of the teeth if the use of tetracycline is considered in younger children. Tetracycline, which is avidly taken up by the nidus, is administered orally 1 to 2 days
preoperatively. Tetracycline fluoresces under ultraviolet light, thus providing an intraoperative method of determining whether the nidus has been removed. With the operating room lights dimmed and a Wood's lamp emitting the ultraviolet light, the nidus can be readily identified in the resected portion.

In a similar fashion, radioactive isotope can be used intraoperatively to assist in identifying the osteoid osteoma. The isotope is administered preoperatively and a scintillation probe is used intraoperatively to detect the increased counts per minute in the area of the lesion. However, few centers use this method because of the expense involved and the sometimes equivocal results. We have no experience with this technique.

The two most common surgical methods for removing the nidus are en bloc resection and the bur-down technique. En bloc resection is performed by placing drill bits around the lesion and confirming their placement with fluoroscopy in the operating room. The lesion is then removed en bloc with the margin of reactive bone. This requires a larger resection of bone than the bur-down technique, and therefore either bone grafting or internal fixation may be necessary. With the bur-down technique, the sclerotic reactive bone is burred until the nidus is visible. The nidus is then curetted and the specimen is sent to pathology. The cavity of the lesion is then thoroughly burred. This technique has even been applied arthroscopically in the talus. The advantages of this procedure over en bloc resection include removal of less reactive bone (thus reducing the need for bone grafting) and a decrease in the risk of a postoperative pathologic fracture.

Newer techniques are showing favorable results. These include CT-guided percutaneous excision using a trephine and CT-guided percutaneous radiofrequency coagulation.*

*See references 80, 156, 192, 213, 231–233, 280.
A radiofrequency probe is placed in the center of the lesion and a 1-cm area around the bone is heated to destroy the nidus itself. Reports by those who have used this technique indicate its results are equivalent to those obtained with surgical excision. Its advantages include the fact that it is an outpatient procedure, there is a lower risk of pathologic fracture, and convalescence is rapid.

Osteoblastoma

Osteoblastomas have a histologic pattern very similar to that of osteoid osteomas, but they are usually much larger (2 to 10 cm). They are one-fifth as common as osteoid osteomas and represent approximately 0.5 percent of biopsied primary tumors.\(^{185}\) Most osteoblastomas occur in persons ages 10 to 25 years, with the peak incidence noted at around 20 years.\(^{186}\) More than 80 percent of patients are younger than 30 years at the time of diagnosis. Affected males outnumber females by a ratio of 2:1.

PATHOLOGY

On gross pathology, osteoblastomas vary in size from 2 to 10 cm. At surgery, an osteoblastoma is found to consist of hemorrhagic, granular, friable, and calcified tissue.\(^{40}\) The lesions are gritty on palpation, usually deep red to reddish brown or pink (reflecting their vascularity), and, if removed intact, often well circumscribed and surrounded by a shell of cortical bone or thickened periosteum.

Histologically, osteoblastomas are identical to osteoid osteomas, consisting of vascular spindle-cell stroma with abundant irregular spicules of mineralized bone and osteoid.\(^{73}\) Osteoblasts and osteoclasts are readily evident on the edges of the bone spicules (Fig. 37–32). Cartilage is distinctly absent. Because of their similar histologic pattern, osteoblastomas have sometimes been referred to as giant osteoid osteomas.

Occasional osteoblastomas appear aggressive on radiographs, with bone destruction and extension into soft tissues (Fig. 37–33).\(^{21,168}\) Microscopically, these infrequent lesions may reveal notable cellular atypia with large plump osteoblasts, making it difficult to differentiate an aggressive osteoblastoma from a low-grade osteosarcoma histologically. Osteoblastomas do not metastasize.\(^{40}\)

CLINICAL FEATURES

Unlike osteoid osteomas, approximately 30 to 40 percent of osteoblastomas are found in the spine, where they most often affect the posterior elements, including the spinous and transverse processes, lamina, and pedicles.\(^*\) Osteoblastomas exceed several centimeters in size, and spinal lesions may extend into the vertebral body. On occasion, the lesion appears to originate from within the vertebral body. All areas of the spine may be involved, from the upper cervical region to the sacrum.\(^{243}\) The clinical presentation may include myelopathic or radicular symptoms.\(^{267}\) Progressive painful scoliosis may develop. If the visceral spine is affected, torticollis may be evident.

Other common sites include the long bones, especially the femur and tibia. In the long bones, the osteoblastoma involves the metaphyseal or diaphyseal region. The lesions are centered in the medullary portions of the shaft, unlike osteoid osteomas, which tend to be located within the cortex or subperiosteally.\(^{153,186}\) Less often the mandible, foot, calvarium, pelvis, scapula, patella, ribs, clavicle, or hands may be affected. In these nonvertebral locations, pain is usually the prominent complaint. Symptoms may be present for a few months to a year. The pain is less localized than the pain of osteoid osteomas and much less likely to be relieved by salicylates.

Because osteoblastomas are several centimeters in size, physical examination may reveal a palpable mass. Tenderness over the area of the tumor is the most consistent physical finding. If the lesion is located near a joint, there may be some loss of joint motion.

RADIOGRAPHIC FINDINGS

Osteoblastomas usually result in a uniform fusiform expansion of the bone. The borders of the lesion are well delineated from the surrounding host bone, and often there is a thin rim of reactive intramedullary bone sclerosis.\(^{153}\) Most lesions are 3 to 6 cm in size, although the range is 2 to 10 cm. The reactive bone formation is noticeably less intense and the margins are less defined than those of osteoid osteomas. Although most lesions are metaphyseal or diaphyseal in location, epiphyseal lesions may be seen in the long bones of the hand or foot. The center of the lesion varies: it may be lucent, mixed lucent and blastic, or predominantly blastic.

In the spine, the posterior elements are predominantly affected. Cortical expansion is common and is similar to

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\(^*\)See references 35, 40, 95, 168, 187, 236, 253.
that seen with aneurysmal bone cysts. Osteoblastomas, however, are usually more radiodense than aneurysmal bone cysts.

Because of the size of the lesion, osteoblastomas can usually be seen on plain radiographs. CT better delineates the extent of involvement, particularly with vertebral lesions. Radionuclide bone scintigraphy may be helpful in localizing smaller osteoblastomas that are not readily apparent on plain radiographs.

**DIFFERENTIAL DIAGNOSIS**

Expansile osteoblastomas may be difficult to differentiate radiographically from aneurysmal bone cysts. Clarification,
however, is usually obtained with CT. Differences in the size and location of the lesions usually distinguish osteoblastomas from osteoid osteomas.

Up to 10 percent of low-grade osteosarcomas may have radiographic features that suggest osteoblastoma. As mentioned, an aggressive osteoblastoma may be difficult to differentiate histologically from a low-grade osteosarcoma. In a benign osteoblastoma, however, there is an absence of sarcomatous large, plump connective tissue stromal cells, sarcoma giant cells, and tumor cartilage and bone.

**TREATMENT**

Treatment consists of curettage or local excision. The risk of recurrence after such treatment is approximately 10 to 20 percent. If a spinal osteoblastoma impinges on the spinal cord or nerve roots, surgical decompression is required. Unlike osteid osteoma, soft tissue extension of an osteoblastoma into the epidural space may become adherent to the dura. Once the tumor is excised, internal fixation of the unstable spine and bone grafting may be necessary. Osteoblastomas located in sites inaccessible to surgical excision have been reported to respond to radiation therapy or chemotherapy.

**Histiocytosis X (Langerhans’ Cell Histiocytosis)**

Histiocytosis X, a term introduced by Lichtenstein in 1953, is a syndrome that consists of a group of clinical pathologic entities: eosinophilic granuloma of bone, Hand-Schüller-Christian disease, and Letterer-Siwe disease. Because these entities are the result of proliferation and dissemination of pathologic histiocye cells or Langerhans'-like cells, the term *Langerhans’ cell histiocytosis* was introduced as an alternative to histiocytosis X by Nezelof and associates in 1973.

The disseminated forms of the disease (Hand-Schüller-Christian disease and Letterer-Siwe disease) were reported prior to our understanding of the pathologic entity of eosinophilic granuloma of bone. Alfred Hand, in 1893, Arthur Schüller, in 1915, and Henry Christian, in 1920, independently described the complex of polyuria, exophthalmos, and defects found in membranous bones. Their descriptions were combined to form what is currently known as Hand-Schüller-Christian disease. Erich Letterer in 1924 and Sture Siwe in 1933 described a generalized disease process with multisystem involvement, including bone. Present in younger children, Letterer-Siwe disease has a poor prognosis. The term *eosinophilic granuloma* was introduced in 1940 and was used to describe solitary bone destruction by large histiocytic cells intermingled with eosinophilic leukocytes. Approximately 80 percent of cases of histiocytosis X are solitary eosinophilic granulomas, 6 percent are multiple eosinophilic granulomas, 9 percent are Hand-Schüller-Christian disease, and 1.2 percent are Letterer-Siwe disease.

The etiology of histiocytosis X is poorly understood. There is speculation that immunologic stimulation of a normal presenting cell, the Langerhans’ cell, continues in an uncontrolled manner, resulting in these cells’ proliferation and accumulation. This may not truly represent a neoplasm but, instead, a proliferative lesion that may be secondary to a defect in immunoregulation. In contrast to this theory, a recent study reported that histiocytosis X is probably a clonal neoplastic disorder with highly variable biologic behavior. Langerhans’ histiocyte is the cell of origin for this spectrum of the disease. There is no hereditary pattern described.

Langerhans’ cell histiocytosis can present at any age, from birth to old age. The incidence in children has been estimated at three to four per million, with a 2:1 male-female ratio.

**EOSINOPHILIC GRANULOMA: SOLITARY AND MULTIPLE WITHOUT EXTRASKELETAL INVOLVEMENT**

The mildest, most favorable form of histiocytosis X is an eosinophilic granuloma that is confined to a single bone, or occasionally to several bones, without extraskeletal involvement. The lesion is a benign process.

**Pathology.** Eosinophilic granulomas are usually soft, reddish brown material. They often show areas of hemorrhage and, occasionally, cysts.

Histologically the tissue is characterized by a mixture of eosinophils, plasma cells, histiocytes, and peculiar large mononuclear giant cells (Langerhans cells) with abundant pale-staining cytoplasm and indented or cleaved nuclei (Fig. 37–34). Necrosis, fibrosis, and reactive cells (foamy macrophages) may be evident. There is minimal mitotic activity. The lesions may consist primarily of the histiocytic infiltrates or there may be a mixture of histiocytes and eosinophils.

Electron microscopy can be used to confirm the diagnosis; the specific pathologic finding is the presence of Birbeck granules in the cell cytoplasm near the nucleus. These granules are rod-shaped structures characterized by central striation and a vesicular expansion resembling the strings of a tennis racket. The origin and function of Birbeck granules are still uncertain; however, when present, these structures are pathognomonic of Langerhans’ cell histiocytosis.

**Clinical Features.** About two-thirds of cases are diagnosed in individuals less than 20 years old, with most diagnoses made in the 5- to 10-year-old age group. The first symptom is localizing pain, occasionally accompanied by swelling and low-grade fever. The sedimentation rate may be elevated.

**FIGURE 37–34** Eosinophilic granuloma (×40). The histologic picture is characterized by a mixture of eosinophils, histiocytes, and Langerhans’ cells (large mononuclear giant cells with pale-staining cytoplasm).
The skull is the most common site of involvement, followed by the femur. About 40 percent of solitary eosinophilic granulomas are found at one of these two sites. In the case of multiple lesions, the skull and femur again are most commonly affected. Other sites of involvement include the pelvis, ribs, and spine. The tarsal and carpal bones are rarely affected. In the long bones, the lesions are usually intramedullary and most commonly located in the diaphysis.

**Radiographic Findings.** A rapidly destructive lytic process occurs in the bone, producing a “punched-out” appearance on radiographs. In the early phases the lesion may be poorly delineated, show a “moth-eaten” pattern of destruction, and exhibit erosions of the cortices. The periosteum may be stimulated, showing some periosteal elevation.\(^{5,18}\) It is in this phase that the condition most closely mimics osteomyelitis or Ewing’s sarcoma. Later the borders of the lesion become sharp and the contours become round or oval. During the early radiographic phase of the solitary lesion, a biopsy is often necessary to rule out a malignant process (Fig. 37–35). In the skull, the lesion is oval or round, with several satellite lesions sometimes present, making this particular radiographic appearance almost pathognomonic for eosinophilic granuloma. Periosteal new bone formation usually does not occur in the flat bones of the skull or pelvis. Marginal sclerosis during healing can be secondary to treatment or can occur spontaneously.

Another nearly pathognomonic sign of eosinophilic granuloma is the presence of *vertebra plana* in the spine (Fig. 37–36). This occurs with insidious collapse of the vertebral body, which is eventually compressed into a thin wafer.\(^{289}\) The patient’s neurologic status usually remains intact, although spinal cord or nerve root compression may occur on rare occasions as a result of severe vertebral body destruction.\(^{142,274}\) With healing, a variable degree of vertebral height is restored in these spinal lesions.

Approximately 10 percent of patients who initially present with a solitary eosinophilic granuloma develop multifocal lesions with extraskeletal involvement (Hand-Schüller-Christian disease). Nearly any bone other than those in the hands and feet may be affected. Chest radiographs should always be obtained to rule out pulmonary involvement.

CT is used to delineate the extent of the lytic lesions, particularly in the pelvis, spine, and skull. MRI is superior to both radiography and CT in delineating the medullary...
extent of eosinophilic granulomas and surrounding soft tissue changes (Fig. 37-37). The degree of peritumoral edema accompanying an eosinophilic granuloma is less extensive than that seen with Ewing’s sarcoma or osteomyelitis.

Radionuclide bone scintigraphy does not consistently demonstrate eosinophilic granulomas. The scans may be completely normal in patients with radiographic evidence of extensive bone involvement. A plain radiographic skeletal survey is superior to scintigraphy for the diagnosis of multiple lesions.

Differential Diagnosis. The differential diagnosis includes osteomyelitis, Ewing’s sarcoma, malignant lymphoma, metastatic disease, and, in the long bones, aneurysmal bone cyst and solitary bone cyst. Unless pathognomonic findings, such as multiple skull luencies or vertebra plana, are found on radiographic evaluation, biopsy is needed to differentiate the more serious lesions. Fine-needle aspiration yields sufficient material to confirm the diagnosis. In osteomyelitis, the fine-needle aspirate will contain pus, neutrophils, or organisms. Another benign musculoskeletal tumor, nonossifying
fibroma, may also resemble the late healing phase of eosinophilic granuloma. Unlike nonossifying fibromas, though, eosinophilic granulomas usually are diaphyseal and are not distinctly eccentric.

**Treatment.** Patients with solitary eosinophilic granulomas generally have a benign clinical course. They have a good chance of spontaneous remission and a favorable outcome over a period of months to years. The single bony lesion usually does not require treatment other than a biopsy to confirm the diagnosis. At that time, curettage may be performed. Curettage may require augmentation with bone grafting when performed on lesions in weightbearing bones of the lower extremities that are at risk for spontaneous fracture or on lesions where curettage alone could result in unacceptable deformity. If vertebral plana is identified but the associated back discomfort has resolved, observation alone is sufficient.

Intraleisional infiltration with steroids has been reported to be safe and effective. Although this is a minimally
invasive procedure, injections are not needed if the diagnosis is clear.

Lesions can occur in areas where they threaten neurologic function (e.g., the spinal cord or optic nerve) and where local steroid infiltration or surgical resection may not be possible. In these cases, treatment with low-dose radiation may be a good alternative.172,237 The use of radiation therapy to manage localized bone lesions has decreased considerably, however, because of the favorable natural history (spontaneous remission) and the risk (though low) of developing a secondary malignancy. Chemotherapy has been used with some success in cases of diffuse eosinophilic granuloma and in patients with systemic disease and multiple organ involvement (Letterer-Siwe disease).238

HAND-SCHÜLLER-CHRISTIAN DISEASE: MULTIFOCAL EOSINOPHILIC GRANULOMA WITH EXTRASKELETAL INVOLVEMENT (CHRONIC DISSEMINATED TYPE)

The classic description of Hand-Schüller-Christian disease includes multiple eosinophilic granulomas involving bone, diabetes insipidus (due to pituitary gland involvement), and exophthalmos (due to the presence of retro-orbital granulomas). This term is now used to include instances of more chronic evolution, even without the classic findings, that generally occur in children more than 3 years old with involvement of other systems. In fact, the triad of calvarial defects, exophthalmos, and diabetes insipidus is present in only 10 percent of cases.160,239 More than 70 percent of patients with Hand-Schüller-Christian disease are diagnosed before age 5 years. In addition to the features just mentioned, fever, hepatosplenomegaly, lymphadenopathy, anemia, and abnormal liver chemistries may be evident. In contrast to solitary eosinophilic granulomas, the bones of the hands and feet may also be affected in Hand-Schüller-Christian disease. Pathologic fractures may occur, particularly in the spine.

In the early phases of the disease, the histologic picture is similar to that of a solitary eosinophilic granuloma. The later phases are characterized by a greater proportion of lipid-laden macrophages and scarring. Significant morbidity is associated with this disorder.

Treatment consists of a combination of low-dose irradiation and corticosteroids. Surgical curettage is occasionally indicated. Chemotherapy, consisting of a combination of prednisolone and then vinblastine, has been used primarily in cases with evidence of fever, pain, severe involvement of the skin, failure to thrive, or dysfunction of vital organs.

LETTERER-SIWE DISEASE: MULTIFOCAL EOSINOPHILIC GRANULOMA (ACUTE DISSEMINATED OR INFANTILE FORM)

This acute, disseminated, progressive form of histiocytosis is rare. Characteristically it occurs during the first year of life. All patients are identified before age 2 years. Visceral involvement is diffuse and severe. The patient may present with fever and debilitating infection secondary to marrow failure. Hepatosplenomegaly, lymphadenopathy, papular rash, bleeding diathesis, anemia, and occasionally exophthal-

mos and diabetes insipidus may be present. The pulmonary parenchyma may have a granular appearance on chest radiographs. The destructive "punched-out" lesions of the bones, although not a major source of complaints, are identifiable on radiographs.

In the past, Letterer-Siwe disease was considered to be invariably progressive and fatal, with death caused by marrow failure, apathy, or septicemia. Today, appropriate treatment with chemotherapy, steroids, and high-dose antibi-otics may lead to survival.

Nonossifying Fibroma and Fibrous Cortical Defect

Fibrous defects in bone are common lesions in childhood. They are found in the metaphyseal regions of the long bones, particularly the femur and the tibia. Often they are cortical in location, but they can also be found in the cancellous area of bone. In 1942, Jaffe and Lichtenstein reported that when biopsied, these lesions contained fibrous tissue.130 They coined the terms fibrous cortical defect and nonosteoegenic (nonossifying) fibroma. Other terms used to describe these fibrous lesions include fibrous metaphyseal defect and fibrous endosteal defects.

The histologic appearance of all of these lesions is similar. They differ in size and in radiographic appearance, which reflects the varying phases of the development of the same lesion.

PATHOLOGY

Surgical curettage usually reveals soft, friable, yellow or brown tissue. Hemosiderin pigment contributes to the brownish color. The tumor is usually surrounded by ridges of bone septa, which gives it the trabeculated radiographic appearance.

Histologically, the two basic components are fibroblastic tissue and osteoclast-like giant cells (Fig. 37–38). Foamy pale histocytes, focal hemorrhage, and hemosiderin pigment may also be extensively present. These microscopic findings may cause some confusion with other lesions that contain giant cells, such as solid aneurysmal bone cysts.

FIGURE 37–38. Nonossifying fibroma (×10). The two basic components are fibroblastic tissue and osteoclast-like giant cells.
CHAPTER 37—Benign Musculoskeletal Tumors

CLINICAL FEATURES

The term fibrous cortical defect refers to the small fibrous lesions that occur in young children. These fibrous lesions appear to be developmental defects due to a localized disturbance of bone growth and may not be representative of true neoplasms. Most fibrous cortical defects are eventually obliterated by reparative ossification or by gradual extrusion from the cortex during remodeling at the metaphyseal (growing) end of the bone. In a small percentage of cases, these fibrous cortical defects not only persist but increase in size, penetrate into the medullary canal, and may become symptomatic, producing a pathologic fracture. Jaffe and Lichtenstein considered this to be an evolutionary process by which fibrous cortical defects matured into nonossifying fibromas.130

RADIOGRAPHIC FINDINGS

Both lesions are sharply delineated, radiolucent, multiloculated, eccentric, and outlined by a sclerotic border (Fig. 37–39). They are usually metaphyseal in location, but on rare occasions they are found in the epiphyseal region. Nonossifying fibromas have greater extension into the medullary cavity.

The radiographic findings are usually so characteristic of fibrous cortical defect or nonossifying fibroma that further radiologic studies are unnecessary. For those lesions that appear painful but lack evidence of pathologic fracture, better clarification will be obtained with CT. Bone scans may show mild uptake in this isolated lesion and help in differentiating it from other multifocal abnormalities, such as eosinophilic granuloma.

MRI is rarely needed. The MRI features of nonossifying fibroma include either hypo- or hyperintensity signals on T2-weighted images.17,133 Signal intensity on T1- and T2-weighted MR images and the patterns of contrast enhancement depend on the amounts of hypercellular fibrous tissue, hemosiderin, foamy histiocytes, and bone trabeculae.

DIFFERENTIAL DIAGNOSIS

Unicameral bone cysts radiographically resemble nonossifying fibromas more than any other lesions. Other similar benign bone tumors include aneurysmal bone cyst, chondromyxoid fibroma, and eosinophilic granuloma.

NATURAL HISTORY

The fibrous cortical defect usually appears near the physis and then migrates away during its growth. Usually the lesion regresses spontaneously, becoming smaller and less distinct and eventually disappearing. Occasionally the fibrous cortical defect proliferates and increases in size, extending into the endostium or medullary cavity and involving a greater portion of the width of the bone. At this stage the diagnosis of nonossifying fibroma is made.

TREATMENT

Fibrous cortical defects do not require treatment. They usually regress over time. Larger, nonossifying fibromas may
lead to some discomfort and possible pathologic fractures. Even so, the majority of patients with nonossifying fibromas can be monitored without surgical intervention, and if fractures do occur, they can be successfully managed nonoperatively. Occasionally, discomfort or repeated fractures may require intralesional excision by curettage down to normal bone, with the defect filled with autogenous or allograft bone (Fig. 37–40). Local recurrence is rare with this type of treatment, and there is little to no risk of malignant degeneration.

Primary Synovial Chondromatosis

Synovial chondromatosis is characterized by the formation of metaplastic and multiple foci of cartilage in the intimal layer of the synovial membrane of a joint. The lesion also occurs in bursae and tendon sheaths. The term synovial osteochondromatosis is used when the cartilage is ossified.

This benign neoplasm is very rare. It usually occurs in persons more than 40 years old but occasionally occurs in adolescents. It is twice as common in men as in women.

ETIOLOGY

The etiology of primary synovial chondromatosis is unknown, although cytogenetic findings strongly suggest that it is a clonal proliferation. Trauma has been postulated as a possible stimulus of metaplasia of the synovial cells into chondrocytes.

PATHOLOGY

Arthrotomy reveals the synovium to be thickened and studded with innumerable small, firm, flat or slightly raised, grayish white nodules. These cartilaginous or osteocartilaginous foci may become pedunculated and detached from the affected membrane, entering the joint cavity as loose bodies. Histologic studies disclose numerous foci of cartilaginous metaplasia of the synovium, which may be calcified or ossified (Fig. 37–41).

CLINICAL FEATURES

Clinical complaints consist of pain, swelling, and stiffness of the affected joint; joint locking may also be a symptom when there are loose bodies. Months or years may elapse before a patient seeks treatment. On examination the synovial membrane is noted to be thickened and the joint is limited in its range of motion. Other physical signs that can be elicited are crepitus and palpable loose bodies.

RADIOGRAPHIC FINDINGS

Radiographs reveal multiple areas of stippled calcification in and around the affected joint when the lesion is cartilaginous (Fig. 37–42). In such cases the findings are those of capsular distention and synovial thickening.

TREATMENT

Treatment consists of simple removal of the loose bodies and partial synovectomy, often performed arthroscopically. Extensive and complete synovectomy is impractical and usually not necessary. The condition has a definite tendency to resolve eventually. Malignant transformation into chondrosarcoma is unusual.

Pigmented Villonodular Synovitis and Giant-Cell Tumor of the Tendon Sheath

Pigmented villonodular synovitis (PVNS) is a benign lesion that develops in joint linings. Giant-cell tumor of the tendon sheath (histologically identical to PVNS) develops in the fibrous sheath of tendons.

PATHOLOGY

During arthroscopic or open synovectomy, the synovial membrane is found to be diffusely thickened and tan or brownish red in color. Sessile or pedunculated nodules may cover the surface of the synovium. The synovial texture may vary in firmness, depending on how much fibrous tissue is present. Extensive hemosiderin deposition may be evident. In the tendon sheaths of the fingers, the lesion is usually solitary and well circumscribed.

Histologically, the villous nodular appearance of the synovium is characteristic, with tightly packed histiocytes filling the subsynovial tissue (Fig. 37–43). Some of the histiocytes are laden with hemosiderin. Multinucleated giant cells and lipid-laden macrophages are seen in varying numbers. There are few mononuclear cells, lymphocytes, or plasma cells. Histologically, PVNS is similar to the hemosiderotic synovitis that results from multiple episodes of bleeding into a joint, such as seen in hemophilia. Abundant production of collagen may be evident in patients with long-standing disease. Occasionally, cellularity of the lesion may produce a pseudo-sarcomatous appearance.

CLINICAL FEATURES

PVNS is locally aggressive and almost always monarticular. Most patients are young to middle-aged adults. The most common sites of involvement are the knee and the tenosynovial region in the hand and wrist. Other areas that may be affected include the foot, ankle, hip, and shoulder. Multifocal involvement, although rare, has been reported in children. In these patients, genitourinary and other congenital anomalies may be noted. Multiple sites are involved in less than 1 percent in reported series of PVNS. The disorder has a slight male predilection.

Patient complaints consist of localized pain and swelling of the affected joint. The proliferated synovial membrane may become caught between the articulating ends of the bone, creating a locking within the joint. Range of motion may be limited. Joint aspiration yields a dark brown or serosanguineous fluid. In the absence of trauma to the joint, this finding is of diagnostic significance.

RADIOGRAPHIC FINDINGS

Radiographic findings that are highly suggestive of PVNS include soft tissue (synovial) swelling within the joint and...
FIGURE 37-40 Imaging findings in a 15-year-old boy with persistent discomfort. A and B, Radiographs showed a persistent distal tibial nonossifying fibroma. C and D, Two years after curettage and bone grafting, the lesion had healed.
lucent areas involving the epiphyseal (and/or metaphyseal) ends of two continuous bones across a joint. These radiolucencies may have a border of benign sclerosis. The lytic bone lesions may appear very aggressive, particularly in the femoral head and acetabulum. CT will further clarify involvement on both sides of the joint. A marked narrowing of the joint may be present (Fig. 37–44).

MRI findings include scattered areas of low signal intensity that represent hemosiderin deposits in hypertrophied synovium on T2-weighted images and dotted areas of low signal intensity, presumably due to fibrous components of the lesion, on T1-weighted images.16

Following aspiration of the joint (dark brown or serosanguineous fluid is usually obtained), contrast arthrography will reveal multiple filling defects due to the abundant hypertrophic synovium.

**FIGURE 37–42** Synovial chondromatosis of the shoulder. Note the multiple areas of stippled calcification in and around the joint.

**FIGURE 37–43** Pigmented villonodular synovitis (×10). The villous nodular appearance of the synovium is characteristic, with tightly packed histiocytes filling the subsynovial tissue.

**DIFFERENTIAL DIAGNOSIS**

Abnormalities that may affect both sides of a joint are most often considered in the differential diagnosis of PVNS. These abnormalities include chronic monarticular rheumatoid arthritis, synovial hemangiomatosis, low-grade infection, and, rarely, other inflammatory joint conditions such as tuberculosis. Hemophilia usually is readily diagnosed from its accompanying clinical symptoms, although the histologic findings are similar to those of PVNS.

**TREATMENT**

The treatment for PVNS consists of total synovectomy. In the knee, this is best performed arthroscopically. Recurrence of PVNS following synovectomy is common, and the patient and family should be made aware of this probability.24

Radiation therapy has been found useful in recurrent cases with extensive bone involvement and joint destruction in the adult patient.

**Dysplasia Epiphysealis Hemimelica (Trevor’s Disease)**

Dysplasia epiphysealis hemimelica is a rare developmental disorder of epiphyseal osteocartilaginous growth in children, usually in the lower limbs. The lesion consists of osteocartilaginous tissue arising from the epiphysis and usually is hemimelic (either the lateral or the medial part of the ossification centers is involved). Although the incidence has been reported as one per million, it is likely higher than that.

Mouchet and Belot in 1926 first described this as a tarsal bone disorder and used the term *tarsomegalie*.18 In 1950 Trevor used the term *tarsop-epiphyseal aclasis*. In 1956 Fairbank used the now common term *dysplasia epiphysealis hemimelica*.27

**ETIOLOGY**

The etiology of dysplasia epiphysealis hemimelica is unknown.11,15,26 There is no strong evidence to suggest a he-
editary component. It has been hypothesized that this condition represents a fundamental defect in the regulation of cartilage proliferation in the affected epiphyses, tarsal, or carpal bones.

**PATHOLOGY**

The findings are similar to those described for solitary osteochondromas. The lesion may be a pedunculated mass with a cartilaginous cap, or it may be seen only as an enlarged irregularity of the articular surface. Histologically, the lesion appears similar to benign osteochondromas.

**CLINICAL FEATURES**

The most common sites of involvement are the distal femur, proximal tibia, talus, and tarsal navicular (Fig. 37-45). Other affected areas have included the hip, first cuneiform, scapula, and, infrequently, the upper extremity. The presenting complaint usually is not discomfort but instead deformity and limited range of motion in the affected joint. Other symptoms include a limp, muscle wasting, and, if long standing, limb length discrepancy. Angular malalignment of the knee (valgum or varum), ankle, and hindfoot (valgus) may be evident. The affected portion of the epiphysis is enlarged and a mass may be palpable. Articular surface irregularity may lead to early secondary osteoarthritis.

The male-female ratio is reported as 3:1, with patients commonly diagnosed between 2 and 14 years of age.

**RADIOGRAPHIC FINDINGS**

The radiographic findings depend on the patient's age at presentation. With infants or toddlers, radiographs may be normal or may demonstrate minimal metaphyseal widening. As the affected bone matures, a multicentric radiodensity develops adjacent to the epiphysis or tarsal bone (Fig. 37-46). In adolescents or adults, the lesion appears as an irregu-
CT has been useful in accurately demonstrating the relationship between the normal bone and abnormal ossification, particularly at the articular surface. The use of MRI has allowed better imaging of the soft tissue component of the lesion. Most of the recent MRI literature reports a distinct plane of separation between the lesion and the normal epiphysial bone. However, this plane is much more difficult to define during surgery.

**NATURAL HISTORY**

Dysplasia epiphysialis hemimelica usually stops growing once maturity is reached. Incongruity that occurs in the joint will lead to subsequent osteoarthritis. This condition remains benign; malignant transformation has not been reported in dysplasia epiphysialis hemimelica.

**TREATMENT**

Observation is warranted if the condition is asymptomatic and has not led to angular deformity or caused significant limitation of joint range of motion. Surgical excision should be undertaken if the lesion is painful, deformity is occurring, or joint function is limited. Recurrence is common, and repeated local excision is often required. Any angular deformities can be treated with corrective osteotomy at the time of lesion excision. Generally, the results are very good following excision of lesions that are juxta-articular. Unfortunately, less successful outcomes are achieved with excision of intra-articular lesions.

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