Giant Synovial Chondromatosis of the Knee Mimicking a Parosteal Osteosarcoma: A Case Report

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Synovial chondromatosis is a benign nodular cartilaginous proliferation arising in the synovium of joints. The radiologic features of this condition are variable. Rarely, it would be confused with malignancy such as chondrosarcoma, osteosarcoma or synovial sarcoma. We report a case of primary synovial chondromatosis of the posterior aspect of the proximal tibia mimicking a parosteal osteosarcoma on the radiography, which showed a homogeneously radiopaque juxtacortical mass. However, subsequent computed tomography (CT) showed multiple intra-articular masses containing chondroid mineralization, suggesting synovial chondromatosis.

Key words: synovial chondromatosis, parosteal osteosarcoma, diagnostic imaging

Case Report

A 24-year-old man was referred to Orthopedic Clinic complaining of intermittent pain in the posterior aspect of the knee. During physical examination, the flexion of his left knee joint was limited to 110° and a fixed hard mass was palpated in the popliteal fossa. Neither tenderness nor local heating was detected. He did not give any trauma or previous specific medical history.

Plain radiographs showed a densely ossified mass which seemed to abut on the posterior aspect of the proximal tibia (Fig. 1). Lateral radiograph shows a soft tissue mass (arrow) containing dense mineralization extending into the popliteal fossa. Partial attachment to the cortex as well as radiolucent cleavage plane was evident. Two small radiodense soft tissue masses (arrowheads) were initially missed on the radiograph.
and two small radiodense masses in the suprapatellar pouch and the Hoffa’s fat pad were initially missed on the radiograph. A radiolucent cleft separating the part of the ossified large mass from the posterior cortex of the proximal tibia was detected radiographically. The combined radiological and clinical differential diagnosis at this point included a parosteal osteosarcoma as well as myositis ossificans and tumoral calcinosis, with parosteal osteosarcoma being preferentially considered in the differential diagnosis. CT scan was performed to further evaluate, which showed a large, well-circumscribed calcified mass extending into the popliteal fossa with another small mineralized masses (Fig. 2). According to the findings of CT scan, a radiologist reported that radiologic diagnosis was compatible with a synovial chondromatosis. We thought that limitation of the knee joint motion was occurred due to the mechanical block of the huge posterior mass. The operation for mass excision was performed. The large mass in the popliteal fossa, as well as another small sclerotic nodular masses were excised through direct posterior approach. The intra-articular mass showed only thin, tenuous attachments to the synovial membrane and could easily be removed. Because the synovium and the capsule of the joint appeared grossly unremarkable, a partial or complete synovectomy was not performed. Histologically, the mass was composed of hyaline cartilage with focal areas of acellular bone, diagnostic of primary synovial os- teochondromatosis. On microscopic examination, variably cellular hyaline cartilage covered by a fine fibrous layer and some myxoid change was reported (Fig. 3). After operation, the range of knee joint motion was markedly improved with normal range and no evidence of recurrence was observed postoperatively 14 months.
Discussion

Synovial chondromatosis is a benign arthropathy characterized by synovial tissue undergoing metaplastic transformation to produce foci of cartilage. It is most frequently encountered in the third to fifth decades, occurring in males approximately two times more commonly. Most frequent site of involvement is the knee joint. Clinical manifestations are generally nonspecific. They include mild and chronic, with intermittent swelling, low grade dull pain, and reduced motion. Intermittent acute exacerbations of pain, swelling, and joint locking can occur as sequel to the osteochondral fragment being entrapped between opposing joint surface.

Radiographic evaluation will be unrewarding if the loose bodies lack calcification or ossification and may be visible just as indistinct soft tissue masses. However, when mineralization occurs, radiographs reveal radiopaque, round loose bodies within the joint or periarticular area and the diagnosis can be established preoperatively. CT is useful in identifying the lesion in its early stages when there is insufficient mineralization to be documented in plain radiographs. MR imaging findings are more variable owing to the extent of mineralization and ossification of the chondral bodies. CT and MR imaging depict the extent of the synovial disease and lobular growth. The classical imaging appearance of synovial chondromatosis is multiple, oval, well-defined, intra-articular and homogeneous calcified bodies which are typically distributed evenly throughout the joint. These calcification frequently show a pathognomonic appearance of ring-and-arc, popcorn-like, or feathery pattern of mineralization. In addition, chondral bodies may also progress to further maturation and enchondral ossification with a peripheral rim of cortex and inner trabecular bone. In rare cases, the individual chondral bodies coalesce to form a larger, conglomerate, mineralized mass. Edeiken described 10 patients with a giant synovial chondromatosis showing osteocartilagenous bodies varying in size from 1 cm to even 20 cm, with the knee involved in five of the cases. They suggested that on the basis of radiologic findings, the giant solitary chondroma is formed by the fusion of multiple synovial chondromas or by continued growth of a solitary synovial chondroma into a large, well-circumscribed calcified mass, which was similar to our case. Occasionally, giant synovial chondromatosis and the classical type of synovial chondromatosis may occur concurrently in the same region as our case. However, our cases are different from theirs in that large calcified masses had broad base with adjacent bone, which was reminiscent of juxtacortical lesions and would be difficult to distinguish from parosteal osteosarcoma.

The radiologic appearance of parosteal osteosarcoma is often characterized by a large, lobulated (cauliflower-like), ossific (denser centrally), juxtacortical mass. Initially, only a narrow zone (stalk) of attachment to the cortex may be apparent, creating a partial radiolucent cleavage plane between the lobulated osseous mass and the remaining bone and has been referred to as the cleavage plane or string sign of parosteal sarcoma. Our case showed the combination of a large, densely mineralizing juxtacortical mass, partial attachment to the cortex as well as radiolucent cleavage plane on the plane radiograph, which could cause diagnostic problems to the unwary radiologist as this lesion tend to be mistaken for parosteal osteosarcoma. Indeed, our case was referred to orthopaedic oncology unit with presumptive diagnosis of a malignant process. However, identification of accompanying small nodular loose bodies, the rings-and-arcs morphology of the calcifications, and the intra-articular location on CT makes us discriminate between parosteal osteosarcoma and synovial chondromatosis.

References

방골성 골육종과 유사한 슬관절의 거대 활막 연골증식증

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활액막성 연골증증은 관절 활액막으로부터 기원하는 증식성의 결절성 연골성 악성 종양이다. 영상의학적 소견은 다양하게 나타나는데 드물게 연골육종, 골육종 그리고 활액 육종과 같은 악성 종양과의 감별을 요하는 수가 있다. 저자들은 컴퓨터 단층 활영에서는 관절 내 다발성의 연골성 무기질침착이 드러난 종괴로 인해 활액막성 연골증증을 예상할 수 있으며, 단순 방사선 소견상 골의 피질골에 근접하고 균질한 방사선이상과의 모습을 보여 방골성 골육종과 유사한 활막으로 보이는 근위 경골의 원위부에 발생한 원발성 활액막성 연골증증 1예를 보고한다.

색인단어: 활액막성 연골 증증, 방골성 골육종, 영상 검사
