Case report

A 10-year-old girl initially presented at an outside institution with right hip and knee pain. At that time, the patient and her family were told that her hip and knee x-rays were normal. The pain continued, and MRI of the hip without contrast was performed at the outside institution approximately nine months after initial presentation. This study revealed a focal, well-defined lesion that was hypointense on both the T1 and STIR sequences, with edema in the adjacent bone and soft tissues (Fig. 1). The patient was then referred to our institution for evaluation and treatment.

At the time of presentation to our institution, the patient was a well-developed girl with a benign past medical history. Physical exam revealed limited range of motion and weakness of the right hip musculature secondary to pain. The patient also had tenderness overlying the greater trochanter. Re-interpretation of hip radiographs from the outside institution revealed a well-defined lytic lesion within the greater trochanter (Fig. 2). The patient subsequently underwent CT-guided biopsy of the greater trochanteric lesion. Biopsy revealed chondroblastoma. The patient then underwent curettage and bone-grafting of the lesion. The patient’s postoperative recovery and clinical course were uneventful for the next eleven months.

At that time, the patient again presented with increasing right hip pain. MRI of the hip showed a small joint effusion, with edema in the femoral head and surrounding soft tissues (Fig. 3). A small mass in the soft tissues just medial to the posterior greater trochanter was not recognized. The study was interpreted as possible septic arthritis. The patient then underwent ultrasound-guided aspiration of the joint effusion. Laboratory evaluation of the fluid was negative for infection. At that time, it was thought the patient may have injured herself. The patient continued to have increased pain, to the point that she was not able to bear weight. At that time, a CT-guided biopsy of the greater trochanter was performed. A soft-tissue mass surrounded...
Subperiostial recurrence of chondroblastoma

Figure 2. 10-year-old girl with recurrence of chondroblastoma. Right hip radiograph. Frontal view of the right hip reveals a well-circumscribed lytic lesion (arrow) in the greater trochanter.

Figure 3. 10-year-old girl with recurrence of chondroblastoma. MRI of the right hip. (A) Axial fat-saturated T2 (TR 4000/TE 81), (B) axial T1 (TR 550/TE 513), and (C) axial postgadolinium fat-saturated T1 (TR 500/TE 13) MR images of the right hip. Relatively hypointense T2 and T1 lesion (arrows) in the soft tissues posteromedial to the greater trochanter, with an associated joint effusion and inflammatory changes in the soft tissues and bone.
by a thin rim of interrupted periostium posteromedial to the greater trochanter (Fig. 4) went undetected. The biopsy of the greater trochanter was nondiagnostic.

Figure 4. 10-year-old girl with recurrence of chondroblastoma. CT of the right hip obtained in conjunction with biopsy, depicted on the bone window. Soft-tissue mass posteromedial to the greater trochanter surrounded by a thin interrupted rim of calcification (arrows).

Figure 5. 10-year-old girl with recurrence of chondroblastoma. MRI of the right hip. (A) Axial fat-saturated T2 (TR 4000/TE 89), (B) coronal T1 (TR 650/TE 13), and (C) axial postgadolinium fat-saturated T1 (TR 566/TE 13) MR images of the right hip. Enlarging relatively hypointense T2 and T1 lesion (arrows) in the soft tissues posteromedial to the greater trochanter, with an associated joint effusion and inflammatory changes in the soft tissues and bone. Postgadolinium, the lesion demonstrates enhancement that is less intense than the surrounding inflammatory changes in soft tissues and bone.
The patient continued with pain and inability to bear weight, despite aggressive physical therapy. A repeat MRI was performed and revealed intense increased signal on fat-saturated T2-weighted and on postcontrast fat-saturated T1-weighted images throughout the soft tissues around the right hip. Embedded in this inflammatory edematous reaction was an oval, 3-cm mass posteromedial to the greater trochanter that enhanced less than the surrounding tissue and therefore was visible as an island of relatively low signal (Fig. 5). The patient was taken to the operating room for an open biopsy. The tumor was identified in a subperiostal location outside of the cortical bone. The frozen section was consistent with recurrent chondroblastoma. Therefore, the surgeon proceeded to excise the soft-tissue mass and perform a curettage and bone-grafting of the surface of the bone. Final pathology confirmed recurrent chondroblastoma. The histological appearance was similar to the original tumor. It was composed of a diffuse proliferation of mononuclear cells with intermixed multinuclear giant cells and scattered foci of chondroid matrix, some of which had undergone calcification (Fig. 6). At followup eight months after surgery, the patient was pain-free and fully ambulatory.

Figure 6. Blue to pink chondroid matrix (arrows) on a background of diffuse proliferation of mononuclear cells and focal giant cells (curved arrows) (100x).

Discussion

Chondroblastoma is a rare benign bone tumor accounting for approximately 1% of all benign bone tumors (1). The lesion typically presents as a well-demarcated lytic lesion in the epiphysis on radiographs. They are also well-known to occur in the apophyses, including the greater trochanter. Treatment consists of curettage and filling of the defect with either bone graft or methylmethacrylate. Recurrence rates range from 8.3% to 21.4% in the more recent literature (2–5). The single most important factor predictive of recurrence is retention of tumor material at surgery (2, 3). Lesions occurring in the region of the proximal femur and pelvis also demonstrate an increased rate of recurrence (2, 5). The reason for this higher rate of recurrence may be secondary to difficulty of curettage in this location. Given the location of our patient’s lesion, our patient was at an increased risk for recurrence. Biological aggressiveness of the tumor may also play a role in recurrence (2). In our patient, there was no evidence for aggressive features either in the primary tumor or recurrent tumor.

Soft-tissue recurrence of chondroblastoma has been reported, although it is rare. The first case was reported in 1966 (6). To our knowledge, a total of eight soft-tissue recurrences have been reported in the literature (6–11). Our patient presented with subperiostal recurrence of tumor along the posteromedial aspect of the greater trochanter. Therefore, the plain films of the hip were not particularly helpful, as there was no osteolysis to suggest tumor recurrence. On the CT performed at the time of the second biopsy, the soft-tissue mass surrounded by a thin perioskeletal shell remained unrecognized but was similar in appearance to a reported case of primary cortical chondroblastoma (12). The MRI in this case clearly depicted the recurrence and surrounding inflammatory change. The low-to-intermediate T2-signal intensity and lobular margin used to describe primary chondroblastoma was also present in our case of subperiostal recurrence (13).

Our patient presented with significant edema within the adjacent soft tissues and bone marrow, a finding that is also common in primary chondroblastoma (13,14). In fact, the surrounding inflammatory reaction can be used as sign of recurrence, as the inflammatory change resolves after removal of a lesion (14). In our case, the initial MRI after the return of the patient’s symptoms was thought to represent septic arthritis, given the extensive marrow and soft-tissue edema, and joint effusion. Although infection was a reasonable consideration, had the recurrent mass in the soft tissues been recognized, this might have allowed more timely initiation of definitive treatment. To our knowledge, this is the first case report of an MRI demonstrating subperiostal recurrence of chondroblastoma.

In summary, persistent or increasing inflammatory changes on MRI in a patient with previous removal of a chondroblastoma should lead to a careful search for recurrent tumor. Although this typically occurs in bone, it may on occasion also develop in soft tissue, which can be subtler and harder to recognize.

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