Chondroblastoma of the Patella: A Case Report

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Abstract
Chondroblastoma is a rare benign cartilaginous bone tumor representing less than 1% of all primary bone tumors. This lesion usually develops in the epiphysis of long bones and occurs during the second decade of life. However, there are rare atypical localizations like the patella. We report an atypical case of chondroblastoma of the patella and discuss the clinical, radiological and therapeutic features of this rare location. A young 20-year-old patient consulted for right knee pain evolving for 2 months with a discreet knee effusion and pain on palpation of the patella.

The radiological and CT showed lytic lesion of the distal half of the patella, lobulated, with sclerotic outlines and blowing the internal cortex. The patient underwent surgery with curettage and filling with autologous iliac cancellous bone graft. Histological examination led to the diagnosis of chondroblastoma. At last follow-up, 18 months, the patient had a normal function of his knee with no recurrence for radiological control. The location of chondroblastoma in the patella is exceptional but its radiological features resemble those of usual epiphysseal locations. The most popular treatment is curettage associated with cancellous bone grafting and thus to avoid the functional consequences of a patellectomy.

Keywords: Chondroblastoma; Benign bone tumor; Patella; Knee

Introduction
Chondroblastoma is a rare benign cartilaginous bone tumor representing less than 1% of all primary bone tumors. This lesion usually develops in the epiphyses of long bones and occurs most often during the second decade of life. However, there are rare atypical localizations like the patella.

We report an atypical case of chondroblastoma of the patella and discuss through our observation and a review of the literature the clinical, radiological and therapeutic features of this rare localization.

Clinical Case
A 20-year-old has consulted for right gonalgia evolving for 2 months felt mainly when exercising. The examination revealed a discreet lameness. The right knee was swollen with minimal joint effusion and 10° flessum. The palpation of the patella awoke pain. There was also amyotrophy of the ipsilateral quadriceps. Active flexion was painful beyond 90°.

Standard X-rays showed a lytic, multi-lobed, sclerotic image extending to the entire distal half of the patella and blowing the internal cortex (Figure 1). The computed tomography revealed a multilobulated geode, finely circled by a line of condensation, blowing the anterior and internal cortices which are thinned. In some sections, we could note a break-in of the cortex on the articular surface (Figure 2).

The biological assessment was without abnormalities including infectious and inflammatory markers (CBC, ESR, and CRP).

The patient had a surgical treatment with anterior approach of the patella, collapse of the anterior wall of the cyst then resect of the cystic cavity whose content was firm, pink dotted with chalky areas (Figure 3A). Curettage of the cavity was performed (Figure 3B) followed by filling with an autologous cancellous bone graft taken from the ipsilateral iliac crest (Figure 3C) and closure of the anterior window with an iliac cortical fragment (Figures 3D and 4).

Microscopic examination revealed a sheet-like proliferation of small to intermediate-sized round polygonal cells within a chondroid matrix (Figure 5a). The cytoplasm is eosinophilic and the nucleus is centrally placed with longitudinal nuclear groove (Figure 5b). Some osteoclast-type giant cells are randomly distributed.

In the postoperative period, the knee was immobilized by a removable knee splint in extension for 2 weeks with authorization of the support. Rehabilitation was then started to strengthen the quadriceps and to foster the passive recovery of the articular amplitudes. Normal activity was resumed on the 45th postoperative day.

At the 18-month follow-up, the clinical results were rated excellent on functional criteria with total indolence and mobility at 0/140 of the right knee. Radiograph showed sub-total filling of the cavity and no signs of recurrence (Figure 6).

Figure 1: Initial plain radiograph displaying a lytic, multi-lobed, sclerotic lesion extending to the entire distal half of the patella and blowing the internal cortex.

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Figure 2: CT scan demonstrated a multilobulated geode, finely circled by a line of condensation, blowing the anterior and internal cortices which are thinned. In some sections, we could note a break-in of the cortex on the articular surface.

Figure 3: Intraoperative footage displaying: (A) Collapse of the anterior wall of the cyst then recess of the cystic cavity whose content was firm, pink dotted with chalky areas. (B) Resection of the lesion and curettage of the cavity. (C) Filling with an autologous cancellous bone graft taken from the ipsilateral iliac crest. (D) Closure of the anterior window with an iliac cortical fragment.

Figure 4: Postoperative radiograph: Cavity filled with cancellous bone taken from the iliac crest.

Figure 5a: A sheet-like proliferation of small to intermediate-sized round polygonal cells with scattered osteoclast-type giant cells (Hematoxylin-eosin, original magnification 100x).
Discussion

Chondroblastoma is a benign cartilaginous bone tumor consisting of chondroblastic cells with presence of chondroid substance. It is a rare lesion since it represents only 0.5% to 1% of biopsied primary bone tumors and 9% of benign bone tumors [1,2]. It has a preference for the male sex (2 to 3/1), usually between the ages of 10 and 20 years. It is therefore a tumor of the young subject, exceptional in early childhood and after the age of 30 years [3].

Chondroblastoma is typically located on an epiphysis or an apophysis always near a fertile phase. It develops in 80% of the cases at the end of a long bone with a predilection for the shoulder, knee and hip [4-7]. More rarely, it can occur on other skeletal segments and on flat bones. The localization of chondroblastoma at the patella is extremely rare estimated between 1 to 3% [3-5]. It represents 16% of benign bone patella tumors [4,5].

The revealing clinical signs are usually unobtrusive and nonspecific. The pain is most often moderate and can be revealed as a result of a trauma. Swelling may be associated with peripheral locations. When the tumor is located in the lower limbs, a lameness can be seen, as is the case in our patient. But this can sometimes be the only appealing clinical finding. Signs of joint irritation such as decreased mobility or hydarthrosis are common and have been observed in our patient. The occurrence of a pathological fracture remains exceptional [8,9].

In typical forms, standard radiographs are often sufficient to evoke the diagnosis [10] by showing an osteolysis zone of oval shape well bounded by a sclerous outline which testifies to its slow evolution, eccentric to the epiphysis and its size is often between 1 and 6 cm. Small calcified pits can be seen within this osteolytic image in half of the cases. CT allows a better analysis of the contours of the lesion. MRI may be useful in assessing the relationship of the tumor to the joint, soft tissue and growth cartilage that can be crossed by the lesion [11]. In our patient, CT, particularly on sagittal reconstructions, revealed a rupture of the subchondral lamina in some places with a joint break-in, explaining the signs of joint irritation present at the clinical examination. In typical epiphyseal locations, the differential diagnosis is essentially with the other “epiphyseal gaps” namely essentially chronic bone abscess and bone infarction and in adults the giant cell tumor and the clear cell chondrosarcoma [9].

Imaging is less contributory to the diagnosis when the lesion is located in flat bones, where also other lesions usually localized in the metaphyseal zone, such as the essential bone cyst, the aneurysmal cyst, but also the intraosseous lipoma and tuberculous osteitis are suspected [9]. The diagnosis in this case can only be pathological. In our patient, several diagnostic hypotheses were evoked on clinical and imaging findings without being able to decide. Only the histological study of the removed tissue allowed to make the diagnosis of chondroblastoma of the patella.

The progression of chondroblastoma is usually slow but may be rapid in some cases with extension to the metaphysis through the cartilage or more exceptionally to the joint [12]. The risk of malignant transformation is very controversial. Rather, it is an undifferentiated or clear cell chondrosarcoma initially confused with chondroblastoma or a sarcomatous transformation induced by radiotherapy on a benign lesion [13]. The occurrence of always histologically benign lung metastases has been reported [9]. In the latter case, excision of lung lesions leads to healing.

The treatment of chondroblastoma is always surgical curettage-bone grafting with the removal of the bone layer that corresponds to the wall of the tumor [9,14]. Some authors perform in addition a wash with phenol or alcohol, a thermal cauterity or cryotherapy, or even a filling with acrylic cement.

Segmental resection is only conceivable in certain localizations (fibulae, ribs, posterior vertebral arch) and in cases of very extensive lesions or recurrences.

In patella locations, patellectomy is indicated only in cases of major damage to the joint or resulting in a discontinuity of the extensor apparatus. The long-term functional results vary from one author to another.

The radio sensitivity of chondroblastoma is uncertain. Radiation therapy is therefore not recommended. It is formally contraindicated in children because of the risk of alteration of growth plates and the occurrence of radiation-induced sarcoma [9]. The recurrence rate of chondroblastoma varies according to the authors: 5.7% for Schajowicz [12], 15% for Meary [15] and up to 38% for Huvos [16]. The presence of a significant aneurysmal component would increase this risk. This recurrence seems to be related neither to the age of the patient nor to the non-union of the fracture [17].

In our patient, CT, particularly on sagittal reconstructions, revealed a rupture of the subchondral lamina in some places with a joint break-in, explaining the signs of joint irritation present at the clinical examination. In typical epiphyseal locations, the differential diagnosis is essentially with the other “epiphyseal gaps” namely essentially chronic bone abscess and bone infarction and in adults the giant cell tumor and the clear cell chondrosarcoma [9].
to the seat of the tumor and it is usually seen in the bone adjacent to the initial tumor, the surrounding soft tissues and even in the joint. Adequate clinical and radiological monitoring is required.

The functional prognosis of chondroblastoma depends on its location, its degree of aggressiveness that may be responsible for joint destruction or an extension to a nearby bone, and in children, the involvement of the growth plate may lead to growth defect of the bone segment and shortness of a limb [9]. But this prognosis also depends on the quality of the surgical procedure. Thus, segmental resections must be exceptional. In our patient, conservative surgical treatment with a curettage associated with autologous cancellous bone filling of the chondroblastoma of the limb led to an excellent functional result with less morbidity and no recurrence at the last follow-up.

Conclusion

The patellar localization of chondroblastomas is exceptional. Its radiological characteristics are comparable to those of the usual epiphyseal localizations, but the diagnosis is resolutely anatomopathological in front of the lack of specificity of the radiological findings. The most popular treatment is the curettage associated with cancellous bone grafting to avoid the functional consequences of patellectomies. The risk of recurrence of this lesion requires clinical and radiological monitoring.

References