Bone Chondroblastoma in Children: About Case and Review the Literature
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Abstract
Chondroblastoma is a primary bone tumor in children, adolescents and young adults, which accounts for 1% of all bone tumors. Epiphyseal or epiphysometaphyseal localization, this lesion usually develops from secondary ossification centers close to the knee, shoulder and hip. In our work we will report an original of a knee chondroblastoma in a 13-year-old girl with difficulties in surgical decision-making.

Keywords: Chondroblastoma, adolescents, bone tumors, Epiphyseal.

INTRODUCTION
Chondroblastoma is a rare, benign cartilaginous tumor, developed in an epiphysis or an apophysis, near the growth cartilage (young subject); sometimes it is calcified. A few cases are described in adolescents. In our work we will report an original of a knee chondroblastoma in a 13-year-old girl with difficulties in surgical decision-making.

PATIENT AND OBSERVATION
We report the clinical case of a 13-year-old girl who presented with a chronic joint blockage in the right knee. The revealing clinical symptomatology was an intermittent and painless joint blockage in the right knee, evolving for 2 months. The clinical examination revealed a meniscal syndrome with a positive meniscal test (Mc Murray, Thesaly ...). Standard radiography: Postero-external lesion of the proximal tibia (image of osteolysis) (Fig-1). CT: Epiphyso-metaphyseal tumor with probable articular rupture (Fig-2). The biopsy: bone chondroblastoma (Fig-3). The therapeutic procedure consisted in broad intra lesional curettage by antero-external route with a massive spongy graft. The postoperative follow-ups were simple with consolidation in 4 months (Fig-4) and complete disappearance of symptoms at 6 months.

After a 24-month follow-up, no recurrences were noted (Fig-5).

Fig-1: Postero-external lesion of the proximal tibia (image of osteolysis)
Fig-2: Epiphyso-metaphyseal tumor with probable articular rupture

Fig-3: Histologic appearance of chondroblastoma

Fig-4: X-Ray after 4 months showing consolidation.

Fig-5: No recurrence after 6 months
DISCUSSION

Chondroblastoma (CB) is a very rare benign cartilage tumor in adolescents, the first symptom is often pain, lameness; sometimes it is revealed by local swelling; It mainly affects the pure epiphyses, sometimes it is of epiphyso-metaphyseal or pure metaphyseal site [1].

Radiologically the lesion is described as a geographic osteolysis with a dense peripheral border at the level of the epiphysis (standard radiography); namely multiple gaps with intra-lesional calcifications in rings (computed tomography) [2].

There are three surgical approaches: anterolateral, posterior popliteal or arthroscopic; the definitive treatment consists of curettage of the tumor with filling by autograft or with cement, sometimes a mosaicoplasty [3].

Histological analysis may reveal an associated aneurysmal component. The S100 protein is often positive; except that it never reveals signs of malignancy or cellular atypia [4].

The evolutionary profile of chondroblastoma is variable: joint stiffness or growth disorder; recurrences after surgical treatment are possible [5].

CONCLUSION

Chondroblastoma remains a benign, metaphysoepiphyseal or epiphyseal tumor of long bones, infrequent (1% of primary bone tumors), occurring between 5-25 years with a male prevalence and a high rate of recurrences.

REFERENCE