Malignant Degeneration into Chondrosarcoma of Multiple Exostosis Disease: The First Senegalese Case Report

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Introduction

Multiple exostosis disease (MED) is a hereditary benign bone tumor with autosomal dominant transmission1, 2, 3. It is the consequence of a mutation of tumor suppressor genes, EXT1 (chromosome 8q23-q24)4,5, EXT2 (chromosome 11p11-p13)6 and possibly EXT3 (chromosome 19)2. MED is a rare disease and his prevalence has been estimated at 1/50000 in the western population7, 8. Malignant degeneration is the most serious complication; however, it is rare and found in 2 % to 5% of cases9, 10. We report the first Senegalese observation of malignant degeneration into chondrosarcoma of multiple exostosis disease.

Observation

A 43-year-old man had been followed since March 2016 in our department for a family form of MED (MED was also found in his daughter). The disease has been evolved since the age of 2 years. Exostosis was found in metaphysis's long bones and in the left iliac bone (Figure 1). The patient had a Bessel Hagen deformity in his left forearm (Figure 1). He had consulted after 40 years of evolution

Figure 1: Before surgery: exostosis of the metaphysis of the long bones (A, B) and the left iliac bone (B). Bessel Hagen disease: arched deformation of the ulna and dysharmony of length between the ulna and the radius (A).
for pain of osteochondroma of the left iliac bone. The surgical resection of the mass was performed. Histological examination showed compact cartilage and bone tissue casings. The bone tissue consisted of regular spans around the Havers canals. The cartilage tissue contained medullary tissue with fibrous reworking. There was no suspicious sign of malignancy.

The evolution was marked after 2 years by a recurrence of the mass with signs of local compression (a large left lower limb without venous thrombosis on Doppler ultrasound with neuropathic pain in the form of electric shocks and burning, resistant to Paracetamol and Tramadol) (Figure 2).

Biological explorations showed an inflammatory syndrome with a sedimentation rate at 95 mm in the first hour, a C-reactive protein at 133.64 mg/l and a microcytic anemia at 11 g/dl. Serum calcium was 85.25 mg/l and serum creatinine was 7.9 mg/l.

The abdominal and pelvic CT scan followed by magnetic resonance imaging showed a large tumor formation centered on the left iliac bone with an abdominal development portion and a parietal development portion (figure 2). It was an infiltration of the gluteal muscles and the left psoas. The left kidney was pushed forward and bladder, rectum, sigmoid to the right.

Histological examination of the mass biopsy piece showed a malignant degeneration to grade 1 chondrosarcoma (Figure 3). The technical platform did not allow an immunohistochemical examination of the biopsy piece, and genetic tests.

The evolution was marked by the occurrence of respiratory failure death in a context of cachexia and pulmonary metastasis (Figure 4). The patient's cachectic state did not allow a biopsy of the metastatic lesions to be performed.

Discussion

We reported the case of malignant degeneration of multiple exostosis disease. This observation has an epidemiological also diagnostic and prognostic interest.

The prevalence of malignant transformation of MED has been subject to variations according to the literature. In a Western multicenter study including 742 patients followed for multiple exostosis disease, it was 2.7%. The most recent studies have estimated this prevalence between 2% to 5%. In Africa, this prevalence has not been estimated; however, in North Africa, Ahmed et al reported a series of 107 cases of chondrosarcomas secondary to multiple exostosis disease in Egypt. In Morocco, Shahdi et al described 01 case of this condition. In Sub-Sahara, some observations have been reported in South Africa by Solomon L (3 cases) and Vlok S.C.S et al (1 case). Malignant degeneration is thus an extremely rare complication of the
MED which is a rare disease\textsuperscript{12}. Our observation is the first Senegalese case report.

In more than 90\% of cases, it is the cartilage cap that degenerates, giving a chondrosarcoma\textsuperscript{8, 17} as in our patient. In the remaining cases, malignant transformation occurs in the stem of the exostosis as osteosarcoma or sarcoma of fusiform cells\textsuperscript{8, 17, 18}. It can occur at any age, however, according to our observation, it is more frequent around 30 years of age, with an increased risk with age\textsuperscript{8}. In the study by Czajka C M \textit{et al}, the age ranged from 19 to 48 years\textsuperscript{11}.

\textbf{Figure 3}. Histological image showing a malignant sarcomatous tumour proliferation consisting of many atypical chondrocytes with hyperchromatic nuclei and mitoses in small quantities. They are arranged in layers within a fibrous stroma.

\textbf{Figure 4}. Thoracic radiography showing pulmonary metastasis (diffuse images in the form of balloon releases dispersed to both pulmonary fields).
It is important to pay attention to the predictive factors of malignant degeneration in any patient followed for MED\textsuperscript{2} for a preventive approach. In a study including 529 patients in 2011, Pedrini, et al showed that the risk of malignant transformation of exostosis was correlated with their location in the pelvic girdle (pelvic region, proximal hip) and scapular girdle (scapula)\textsuperscript{11}. In the study by Czajka M.C et al, the site most frequently concerned was the pelvic region (8 cases out of 21 cases), followed by the scapula (4 cases 21 cases)\textsuperscript{11}. According to some authors, this association could be linked to delayed detection of osteochondromas in these zones as a result of the least compressive complications of exostosis\textsuperscript{8}. Other risk factors for malignant degeneration reported in the literature were: genetic predisposition (especially EXT1 mutation), male sex, number of exostosis, reactivation of exostosis activity after the stop of growth\textsuperscript{2} 19. All these factors were found in our patient, with the exception of genetic predisposition, which has not been explored. The reactivation of activity of exostosis was characterized in our patient by the occurrence of pain of iliac exostosis which increased in volume\textsuperscript{2} 19. The technical platform did not allow genetic tests to be carried out; which is a limitation in this study. However, the multiple forms of exostosis and the existence of the disease in a first-degree relative suggests a genetic predisposition in our patient\textsuperscript{9}. Thus, a family history of the disease would be described in 60% of cases\textsuperscript{9}.

In the study by Czajka C.M et al, the exostosis that recurred after surgical treatment were all localized to the pelvic girdle (pelvis and upper end of the femur) and the scapular girdle (scapula and sternum)\textsuperscript{11}. Surgical removal indicated for any active exostosis, as was the case with our patient, remains the primary treatment for osteochondromas\textsuperscript{2} 20, 21. However, it is a symptomatic treatment\textsuperscript{2} 20. However, it is a symptomatic treatment and is often repeated in patients with MED\textsuperscript{2} 11.

On the prognostic level, malignant degeneration represents the most serious complication of exostosis. It may be responsible for death as was the case with our patient\textsuperscript{2} 13. In the Egyptian study, the mortality rate was 20% in the group of solitary exostosis and 44% in the group of multiple exostosis disease\textsuperscript{13}.

**Conclusion**

The malignant degeneration of multiple exostosis is rarely described in African literature. Our observation is the first Senegalese case report. The occurrence of death in our patient suggests regular monitoring of MED. This allows early detection and management of risk factors for malignant degeneration.

**References**