Fibrous Dysplasia of Maxillary Sinus

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SUMMARY

Introduction: The Fibrous Dysplasia is a benign bone disease, of slow growth and unknown etiology. The involvement of the craniofacial skeleton is not uncommon and, generally, produces facial asymmetries.

Case Report: In this article we report the case of a patient with fibrous dysplasia occupying the entire left maxillary sinus with orbital extension confirmed in the anatomopathological exam.

Final Comments: The surgical treatment remains as the main therapeutic approach and the postoperative follow-up is necessary due to this condition recurrent nature.

Keywords: dysplasia, monostotic, polyostotic, maxillary sinus.
INTRODUCTION

The Fibrous Dysplasia is a fibro-osseous lesion characterized by the replacement of normal elements of the bone for a disorganized fibrous tissue. It represents 2% of the osseous tumors (1). The first author to describe the characteristic osseous lesion, today known as fibrous dysplasia, was VON REICHINGHAUSEN in 1891, but it was LICHTENSTEIN, in 1938 who introduced the term fibrous osseous dysplasia into the worldwide literature (2).

There are two primary categories of the disease: monostotic fibrous dysplasia, that involves only one bone and represents 70% of the cases; and polyostotic fibrous dysplasia, that presents the involvement of several bones. It is generally developed at the end of the childhood with prevalence in the male sex (2:1) and has a major preference for the white race (3). The craniofacial skeleton is a frequent region of the disease, which may cause organic, aesthetic and psychological disorders.

The diagnosis suspicion is made by clinical and radiological methods which need anatomopathological confirmation (4). The typical radiological exam shows a characteristic aspect of “opaque glass” involved by a dense cortical tissue (5). The definitive treatment is the lesion surgical exeresis. The clinical follow-up of the patient is essential for the early diagnosis of a recurrence. We describe one case of maxillary bone fibrous dysplasia that, in spite of its benign nature, caused facial deformity.

CASE REPORT

The male sex, 11-year-old, white patient from Itaperuna, was forwarded to the Otorhinolaryngology Service of HSJA with his responsible parent with a clinical picture of a progressive increase in the left hemiface. He denied headache, deficit of visual accuracy, otalgia, nasal obstruction and toothache. He mentioned one case of sinusitis 1 year before.

The clinical exam, good general state, the evaluation of the organic systems didn’t reveal alterations. Absence of lymphonodomegaly and cutaneous stains. He presented with a painless tumoration in the left jaw and zygoma with normal subjacent oral skin and mucosa (Picture 1).

The Computed Tomography of the face sinuses showed a hyperdense, heterogeneous, expansive mass of uneven contour, with aspect of an “opaque glass” involving the maxilla, zygomatic bone and left maxillary sinus (Pictures 2 and 3).
The patient was forwarded to the surgical center for the lesion biopsy. The access was intra-oral through Caldwell-Luc surgery and the lesion fragment was removed for histopathological study, which revealed a compatible picture with fibrous dysplasia. He was not submitted to the systemic search of other osseous lesions.

**Discussion**

The Fibrous Dysplasia (FD) is defined as a benign osseous disease characterized by a process of normal bone reabsorption, followed by an abnormal proliferation of a disorganized fibro-osseous tissue (1). It represents about 7% of all benign osseous tumors and may affect any bone of the skeleton (6).

It's classified as for the number of affected bones and the presence or not of extra-skeleton abnormalities. The monostotic form affects only one bone and corresponds to 70-80% of the FD cases. The polyostotic form, in which several bones are affected, may be divided into three subtypes: craniofacial, in which only the craniofacial complex are involved including the jaw and the maxilla; Albright's Syndrome, characterized by the affection of several bones, coffee-with-milk pigmentations in the skin and endocrine affection with a remark for the early adolescence in girls. The polyostotic form corresponds to 20-30% of the cases (7).

The craniofacial bones are more affected in the polyostotic form (50-100%) than in the monostotic form (20%) (4).

The FD etiology is controversial. Lee e col. proposed that the abnormalities in the AMPc or kinase protein intracellular regulation etiologic factors are possible in the development of the FD. Other researchers have identified mutations in the GNAS gen, which result in the GTPase activity alterations, with a consequent increase of the AMPc intracellular concentrations (8).

It manifests more frequently in the childhood and, however, is not exclusive of this age range (4,9). It has an usually slow evolution, a tendency to stabilize after adolescence and a high recurrence rate (2, 9, 10). Such characteristics have a strong implication in the therapeutic approach. As for the distribution of the disease by sex, there is no uniformity between the studies (6, 9, 10).

The disease is initially asymptomatic. The FD signals and symptoms depend on the location of the lesion(s) and the compressive effect in the adjacent structure as the tumor progresses slowly: facial asymmetry and deformity; pathological fractures; obstruction of the paranasal sinuses which generate recurrence infections, cysts and mucoceles; anosmia; headache; loss of visual accuracy for compression of the optic nerve; alteration of the ocular movements; descent; exophthalmia, squint; conductive hearing loss (4, 9, 11, 12).

The CT is the choice exam for the study of the lesion(s), analysis of their extension and surgical preparation (6,11). Basically, three radiographic standards in the cranium fibrous dysplasia and facial bones are described: pategoid, that alternates the radiodense and radiotransparent areas; sclerotic, homogeneously dense; cystic standard, with spherical or ovoid radiolucent area surrounded by dense limits (12). In the case reported, the lesion tomographic images presented a hyperdense standard intermixed by imprecise limits hypodense areas, which resulted in the classical aspect of “opaque glass”. The FD definitive diagnosis is made by the correlation of clinical, radiological and anatomopathological findings (13).

The FD macroscopic exam reveals an expansion of the osseous trabeculate into a thin cortical; as there is no definitive capsule, there occurs an abrupt transition into the healthy bone. The microscopic evaluation shows a collagen matrix stroma with fibroblasts in a entangled standard with osseous trabeculate similar to the “Chinese writing” (12).

The FD differential diagnosis includes malignant (sarcoma, metastatic osteoblastic lesions) and benign lesions (ossifying fibroma, Paget’s disease, aneurismatic osseous cyst, cystic Cristeller Syndrome, ameloblastoma, osteochondroma, hyperthyroidism etc.) (1, 6, 8, 10, 12).

The main factors that guide the FD approach are the presence and the intensity of the symptoms, the tumor location and the patient’s age. The simple presence of the lesion does not justify surgical intervention. The main indications for surgical treatment of FD are the presence of significant clinical symptoms and the control of large aesthetic deformities (2, 9). Because of the lesion benign nature and its recurrence character (10-20%), the surgery must be relatively conservative with the main objective of preserving the function (9). In this case, we chose an expectant procedure, not only for the absence of organic and functional alterations, in spite of the aesthetic involvement, but also because of the age of the patient. He keeps on quarterly service follow-up until his growth and adolescent development is complete.

Radiotherapy is counter-indicated not only because the tumor is radioresistant but also because of the probable increase of the capacity for the dysplasia sarcomatous transformation, whose estimates range from less than 0.5%
of its natural evolution to about 44% after radiotherapy (2,9).

The isolated clinical treatment or associated to surgery has been reported in the literature and there are disagreements as for their benefits in FD. Lustig reports that, for inhibiting the osteoclasts, the biphosphonates are employed in the treatment of patients with osteodystrophies, but that their effects in DF have been limited (9). Moreno reports the following benefits in patients treated with biphosphonates: improvement of pain and inflammatory symptoms, stabilization and reduction of bone destruction, increase of the osseous density, recalcification of osteolytic lesion in 50% of the patients, improvement of the radiologic aspects and the osseous metabolism (6).

Some authors defend the need for a clinical, endocrinologic study and cintilography with Tc 99 in the patients with diagnosis of Monostotic Fibrous Dysplasia in search of other lesions, other regions and extra-skeleton involvement (6, 11).

The clinical and radiological follow-up by CT is essential in the patients with FD because of the moderate recurrence rate of the lesion(s) which may reach 37% according to some authors (9).

**Final Comments**

The Fibrous Dysplasia is significant for the otorhinolaryngology because it may affect facial and cranial bones and may cause deformities and dysfunctions. In spite of its benign nature, the signals and symptoms resulting from the compression of noble structures of the cranial base and orbit may generate diagnostic doubts as for the possibility of a malignant lesion.

The surgical treatment must take into account the disease’s harmful and recurrent potential, by choosing a more conservative approach and removing as much tissue as possible to prevent mutilations and functional deficits.

**Bibliographical References**


