Cemento-ossifying fibroma juvenile of the oral cavity

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SUMMARY
Cemento-ossifying fibroma juvenile of the oral cavity. Objectives. Fibro-osseous neoplasm remains somewhat controversial, and differing concept have been advanced regarding their nature and the proper terminology for them. Cemento-ossifying fibroma juvenile (JOF) is a rare type of fibro-osseous tumor as also been included under the “umbrella” of cemento-ossifying fibroma. The JOF is most often seen in patients who are between 5 and 15 years of age. With this work we emphasize the importance of a correct diagnostic approach.

Material and methods. The case describes a form of cemento-ossifying fibroma hight active and aggressive like JOF. The patient thirteen older showed from 2004 to 2008 three times the palatal lesion, it was performed with a incisional biopsy and excisional biopsy. The tumor were fixed in 10% buffered formalin embedded in paraffin cut into thick sections and stained with ematoxylin-eosin.

Results. The incisional biopsy was inadequate to formulate a correct diagnosis. The histological exams have shown for three times different aspects.

Conclusion. Some authors in the past have suggested different classification. The COFs show different clinical, histological and radiographical patterns.

Key words: escissional biopsy, cemento-ossifying fibroma, fibro-osseous neoplasm.
Introduction

The cemento-ossifying fibroma (COF) is the most common benign fibro-osseous lesion of the jaw, with a slow-growing process (1). The COF is a mesodermal type of non-odontogenic aggressive tumour which is highly destructive and shows a marked tendency to recur. It arises from ectopic multipotential blast cells of the periodontal membrane of the jaws (2). Although it has principally been found in the jaws, it also been reported in the frontal, ethmoid, sphenoïd (3), temporal bones and in the orbit, in the anterior cranial fossa (4).

The COF exhibits a variable behaviour ranging from slow growth to aggressive local destruction; some cases recur after surgery. This variable behaviour cannot be predicted on the basis of the histopathology which is itself variable.

Material and methods

A 13-years-old boy was referred to the dental hospital “S. Giovanni Calibita” Fatebenefratelli of Rome in April 2004 for examination of a swelling of the right maxillary region between 1.5-1.6 dental elements. The surgeon decided to make an incisional biopsy. It was incomplete because there was partial removal.

In 2006, 2 years later, the patient was referred in the same hospital, but there was another surgeon group, he complained of swelling in the right maxillary with no signs of inflammation. There was nothing significant in his medical and dental history. Laboratory tests were all within normal limits. There was no history of trauma. The vitality test of the teeth was negative. Clinical examination showed a marked swelling of the right maxillary, the overlying mucosa was normal in colour and intact (Fig. 1).

At palpation the lesion appeared hard, immotile and asymptomatic. Radiographic examination, orthopantomography (OPT), revealed a large calcified mass, well circumscribed and demarcated, with an oval shaped and without root resorption (Fig. 2). An excisional biopsy was performed on the right maxillary lesion to know the real nature of lesion. The lesion was excised under local anesthesia, with 2% mepivacaine and epinephrine as vasoconstrictive 1/100.000. Subsequently the lesion was incisioned with a blade n° 12 and was performed a total palatal flap (Fig. 3).

During the dissection, the lesion was easily separable from adjacent palatal mucosa but it was firmly attached to the bone (Picture 4, 5); the cavity was controlled and it was washed with saline solution. The flap was sutured with silk 3-0 (Ethicon Johnson & Johnson) and it was protected by periodontal pack for ten days (Fig. 6, 7, 8). After an uneventful post-operative course the patient was seen every six months.
In April 2009, 5 years later, the patient had a recurrence of the lesion in the right maxillary (Fig. 9). Radiographic examination revealed a large radiopaque mass surrounded by a radiolucent zone. He had another excisional biopsy (Figs. 10, 11, 12, 13).
Results

In our case report the first approach to the lesion was conservative. An incisional biopsy was done. The histological features was “hyperplastic formation with edematous overlining and angiectatic stroma”. Two years later at the onset of the recurrence of the same lesion an excisional biopsy was done. The histological result was not diagnostic: “collection constituted by fibrous tissue and bone spicules with necrosis and regressive aspects; there aren’t atypical cells, there isn’t inflammation”. In this case a watch and wait approach was used. After three years during a follow up visit an initial relapse was found and a new complete removal of the lesion was
case report

The istological features were “the material in examination shows fragments of mucosa and Malpighian epithelium in continuity with a neoformation that it’s constituted by fibrous tissue with cementicles and bone trabeculae. The histopathological diagnosis was: “Cemento-Ossyfying Fibroma” (Fig. 10, 11, 12, 13).

Discussion

Cemento-ossifying fibroma (COF) is a neoplasm forming part of the spectrum of fibro-osseous lesions of the jaws (5). The object clinician case of the present work describes varying it youthful, introduced from Jonhanson in 1952 (JOF) (2, 9).

COF is a well demarcated, occasionally encapsulated neoplasm, contains fibrous tissue and varying amounts of calcified tissue resembling bone, cementum, or both. The histological appearance of cementoma of the jaws was first described in 1872 (Menzel). The term “ossifying fibroma” has been used since 1927 (Montgomery) and since 1968 cement-containing tumours were grouped together (2, 6, 7). In 1971 the WHO classified four types of lesions cement containing: fibrous dysplasia, ossifying fibroma, cementing fibroma and cemento ossifying fibroma (8). The WHO revised in 1992 its nomenclature and divided benign fibro-osseous lesions of the oral and maxillo-facial regions into two categories: osteogenic neoplasm (including COF) and non neoplastic bone lesions, including fibrous dysplasia and cemento-osseous dysplasia (9). COF of the jaw shows a predilection for females (10). Many authors have confirmed that COF of the jaw tends to occur in middle-aged patients (6, 10). The peak incidence is in the third and fourth decade with a wide age range (11). JOF is used to describe an actively growing lesion that mainly affects individuals aged <15 years, whereas COF seems to occur in people aged >20 years, and mostly occurs in patients in their 20s and 30s (10).

The WHO classification of odontogenic tumors in the second edition (10), defines juvenile (aggressive) ossifying fibroma as a lesion showing a particular histological aspects: cell, rich fibrous tissue containing bands of cellular osteoid without osteoblastic rimming, with trabeculae of more typical bone, giant cells may also be present. Juvenile ossifying fibroma, however, has also been employed as diagnostic label for lesion occurring predominantly in the paranasal sinuses and histologically characterized by the presence of a fibroblastic stroma containing small ossicles resembling psammoma bodies (17). The stroma varied from loose and fibroblastic to intensely cellular without intervening collagen. The mineralized material consisted of spherical ossicles that were acellular or included sparsely distributed cells. Accounts of the biological behaviour of JOF have varied with reports of recurrence ranging from 30-50%, including the form with multiple recurrences (18). JOF of the jaw bone is generally asymptomatic, it’s depend by the size, it may cause pain, swelling and paresthesia. When it involves the paranasal sinus and orbit regions, it may cause nasal obstruction, sinusitis, headache and proptosis (17). Radiological features depend on the stage of the lesion; early stages present as radiolucensis, late lesions as well-demarcated, dense radiopacities and a minority as mixed. Osteoblastic rimming is present in cemento-ossifying fibroma but according to the WHO classification of JOF is absent in the latter. This is an important point for differential diagnosis between two lesions. A well-defined border has been reported in 85% of cases of COF (17). COF exhibits a locally aggressive growth pattern, and requires radical surgery for tendency of recurrence and the possibility of malignant transformation (10, 13). Takeda and Fujioka (14) reported a case of multiple cemento-ossifying fibroma in three jaw quadrants, and recently, Yih (15) et al. reported multiple familial ossifying fibroma. The same authors demonstrate microscopically capsule presence. Su et al. reported that 88% of COF tumours consisted of a large enucleated pieces and that 44% of cases had capsules (16).

Conclusion

This case report describes a recurrence of a JOF and illustrates the diagnostic difficulties that may be encountered when treating fibro-osseous lesions. It is almost inevitable that diagnosis of JOF is com-
complicated, because pathologic features are common with other fibro-osseous lesions, such as cemento-osseous dysplasia or Juvenile cemento ossifying fibroma.

Radiographical exams, excisional biopsy and histological aspects, are the common instruments for a correct diagnosis. Histology’s aspect of the lesion, psammoma bodies, rich fibrous tissue containing bands of cellular osteoid and the absence of capsule, suggest diagnosis of JOF.

Different diagnosis instruments: radiographical exams, histological exams and pathologic findings is required to ensure an accurate diagnosis. Treatment depends on the individual clinical features and biological behaviour. Postoperative follow-up is essential, to supervising JOF recurred. It’s important that dentist and pathologist communicate between.

Deep biopsy is very important to know the real nature of lesion and avoid the recurrence. The JOF lesions should be excised with conservative enucleation.

References


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