Cemento-ossifying fibroma of the mandible: Presentation of a case and review of the literature

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Abstract
Introduction: Cemento-ossifying fibroma is a benign fibro-osseous maxillary tumor belonging to the same category as fibrous dysplasia and cement-ossifying dysplasia. The aim of present study was report a case of cement-ossifying mandibular fibroma and offers a review of the literature on this disorder.
Case report: A 41-year-old woman presented for conventional dental treatment. A control panoramic X-ray study revealed a rounded, mixed-type image in the mandible, located in the periapical zone from tooth 4.3 to 3.3. The patient presented no symptoms and had no medical history of interest. A sample of the lesion was obtained for histopathological study, which yielded a definitive diagnosis of cemento-ossifying fibroma.
Discussion: The World Health Organization classifies cemento-ossifying fibroma as a fibro-osseous neoplasm included among the non-odontogenic tumors derived from the mesenchymal blast cells of the periodontal ligament, with a potential to for fibrous tissue, cement and bone, or a combination of such elements. These are slow-growing lesions, and are more frequent in women between the third and fourth decades of life. Although the underlying cause is not known, there have been reports of past trauma in the area of the lesion. Due to the good delimitation of the tumor, surgical removal and curettage is the treatment of choice.

Key words: Ossifying fibroma, cemento-ossifying fibroma, benign fibro-osseous lesions.
Introduction
Cemento-ossifying fibroma is a benign fibro-osseous maxillary tumor belonging to the same category as fibrous dysplasia and cement-ossifying dysplasia (1). It is most commonly seen between the third and fourth decades of life (2-6), and is more frequent in women than in men (4:1). The most common location is the mandible, with 70-90% of all cases (1,2). Clinically, these tumors manifest as a slow-growing intrabony mass that is normally well delimited and asymptomatic – though over time the lesion may become large enough to cause facial deformation (3).

Radiologically, cemento-ossifying fibroma shows a number of patterns depending on the degree of mineralization of the lesion. The latter manifests as a well delimited unilocular lesion containing variable amounts of radiopaque material (1-6).

Histologically, these tumors are composed of well vascularized fibrocellular tissue with the capacity to form immature bone trabeculae and cementoid formations, though these findings are not specific of the lesion and can also be seen in fibrous dysplasias (4). A definitive diagnosis therefore requires correlation of the clinical, radiological and histological findings (5).

Treatment comprises surgical resection of the lesion with enucleation and curettage of the bone bed (2,6). The aim of present study was report a case of cement-ossifying mandibular fibroma and offers a review of the literature on this disorder.

Case Report
A 41-year-old woman presented for conventional dental treatment. A control panoramic X-ray study (Fig. 1) revealed a rounded, mixed-type image in the mandible, located in the periapical zone from tooth 4.3 to 3.3. The patient reported having suffered trauma in that same area years ago. She presented no symptoms, had no medical history of interest, and reported no toxic habits or drugs use.

Oral exploration revealed the presence of a number of small caries and fillings in the posterior teeth, and some molars were missing (3.6, 4.6, 4.7). The oral mucosa was normal, and the teeth in the lesion zone (between both lower canines) showed no alterations. Pulp vitality of the teeth adjacent to the lesion proved positive.

The radiological study showed no dental displacement. In addition to the panoramic X-ray study, we requested a mandibular computed tomography scan, which revealed a rounded and elongated osteolytic lesion located between the lower canines. Axial imaging showed the lesion to cause thinning and bulging of the anterior vestibular cortical layer, which was ruptured at some point (Fig. 2). Likewise, the core region of the lesion showed a series of rounded radiopaque images.

Histological study, which revealed the presence of a fusocellular tumor containing calcifications with concentric and acellular mineralization at the center (cementoid type), and other areas presenting recently formed osteoid with peripheral osteoblasts and signs of progressive calcification (Fig. 3). There were no atypias or mitotic figures. The definitive diagnosis was cemento-ossifying fibroma.
Complete surgical removal of the lesion was carried out under infiltrating local anesthesia, raising a mucoperios-veal flap and performing ostectomy to gain access to the lesion. The bone bed was subjected to curettage. The postoperative course was favorable, and one year later the patient reported no discomfort in the zone. A control panoramic X-ray study confirmed good bone regeneration in the zone.

Discussion
Cemento-ossifying fibroma is a benign fibro-osseous maxillary tumor (6). It is a slow-growing lesion most often seen in women between the third and fourth decades of life. While one-half of all cases are asymptomatic, the growth of the tumor over time may lead to facial asymmetry, with the appearance of a mass causing discomfort or mandibular expansion, and the possible displacement of dental roots (2,7). In our case the lesion was casually identified on occasion of a panoramic X-ray study. Although the underlying cause is not known, there have been reports of past trauma in the area of the lesion (8). In coincidence with the data found in the literature, our patient reported having suffered trauma in the affected zone years ago. This points to trauma as a possible triggering factor in some presentations of the lesion, postulating the latter as representing a connective tissue reaction rather than a genuine neoplasm (9).

The World Health Organization classifies cemento-ossifying fibroma as a fibro-osseous neoplasm included among the non-odontogenic tumors derived from the periodontal ligament, with a potential for fibrous tissue, cement and bone, or a combination of such elements (1,7). However, there is controversy over such an origin, since tumors of similar histology have been reported in bone lacking periodontal ligament and not located in the maxillary region, such as ethmoid bone, frontal bone or even long bones of the body (cementiform fibrous dysplasia)(10-14).

Clinically, the tumor tends to present as a slow-growing intrabony mass most often located in the region of the mandibular premolars and molars and in the ascending ramus – in contrast to the anterior mandibular location of our case. The growth is usually asymptomatic, though there may be a degree of root reabsorption or displacement of neighboring teeth (2,6,8). These phenomena were not noted in our patient. There have been reports of more aggressive lesions characterized by rapid and extensive growth, capable of causing mandibular fractures, and multiple lesions have also been documented. Likewise, a more aggressive juvenile form has been described as juvenile cemento-ossifying fibroma (1,15).

Radiologically, these tumors may present a number of patterns depending on their degree of mineralization (6). Two basic patterns have been defined: one characterized by the presence of a unilocular or multilocular radiopaque areas are seen of multilocular appearance in the advanced stages presents extensive radiopaque areas surrounded by a radiotransparent halo (1,2,6). Since our patient had a history of trauma in the affected zone, consideration is likewise required of other lesions that can also be associated with antecedents of trauma, such as central giant cell granulomas. However, although there are radiotransparent zones due to such lesions, no radiopaque areas are seen of multilocular appearance in the course of their evolution.

Due to the good delimitation of the tumor, surgical removal and curettage is the treatment of choice (2,3). In the case of very large lesions with important tissue ablation, the challenge is to replace the affected tissue. The prognosis is usually good, since recurrences are not frequent. This was confirmed in our case, since repair of the affected area was seen to be correct one year after...
treatment – though longer follow-up is required.

References