Extraosseous calcifying odontogenic cyst: a case report and a literature review

Cisto odontogênico calcificante: relato de caso e revisão da literatura

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Introduction

The calcifying odontogenic cyst (COC) is a rare odontogenic lesion first described as a distinct entity by Gorlin et al., in 1962(8). Since then, literature shows a large controversy regarding terminology and classification in spite of the currently acceptance of Gorlin’s original designation by the World Health Organization (WHO) in 1971(17).

Subsequently, WHO(11), who first recognized and defined the COC as a non-neoplastic cystic lesion(17), classified the entity and its variants as an odontogenic tumor rather than an odontogenic cyst.

The neoplastic lesions were subdivided in three subgroups based on location (intraosseous and extraosseous) and histological features. Despite its variable clinical characteristics, COC is often referred as an asymptomatic slow growing swelling of the jaws or gingival tissues, depending upon whether the lesion is intra- or extraosseous. The extraosseous COC is less common, comprising 12% to 20% of the

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reported cases. Clinically it appears as a localized or pedunculated gingival mass with no distinctive features, and radiographically it shows no or only superficial bone involvement with surface erosion.

The histological features of peripheral COC are similar to the intraosseous COC characteristics, and include an epithelial lining of varied thickness, composed of a distinct columnar (and occasionally cuboidal) basal layer, with cells that contain darkly staining nuclei, polarized away from the basement membrane and palisading, similar to the ameloblasts. Within the epithelial lining there is an irregular collection of cells, including sheets of stellate reticulum and eosinophilic cells so called ghost cells. Next to the basal layer, irregular or dysplastic dentin and osteoid can also be found.

By the year 1991 only 54 cases of extraosseous calcifying odontogenic cysts had been reported. Forty-five cases of COC were discussed by Buchner et al., in 1991, this including seven new cases of extraosseous COC. Eleven years after that, a MEDLINE (1993-2005) research was performed and exhibited only 13 new cases of peripheral COC published in the English literature (Table 1).

Treatment involves enucleation of the lesion and long-term follow-up. Recurrences are uncommon and probably related to incomplete cyst removal.

The present paper reports a documented case of an uncommon extraosseous calcifying odontogenic cyst (ECOC) with a review of the literature.

Case report

A 57-year-old black woman was referred to the Oral Diagnosis Department of Universidade de Pernambuco, for evaluation and treatment of an asymptomatic swelling in the retromolar region with a three-month evolution.

Clinically a well-defined mass on the hind right side of the mandible, with 0.5cm of diameter, firm consistency and a reddish overlying mucosa was observed. The orthopantomograph showed no osseous involvement, but a well-circumscribed extraosseous image with an osteogenic reaction of the subjacent bone. Amounts of calcified material scattered throughout the lesion were visible (Figure 1). Clinical diagnosis suggested ossifying fibroma, Gorlin cyst or a residual inflammatory cyst.

Excisional biopsy was performed and the surgical specimen was submitted to histopathologic evaluation at the Oral Pathology Department of the same center. The lesion was paraffin embedded and stained with hematoxylin and eosin (H&E), and after that, evaluated with optical microscopy.

Microscopically, sections showed a hemisected soft tissue specimen superficially composed of a stratified squamous epithelium lining and a subjacent fibrous connective tissue. Within this connective tissue odontogenic epithelium islands resembling ameloblastoma features could be observed (Figure 2). Those neoplastic islands exhibited basal cells with a cuboidal or columnar aspect similar to ameloblasts (Figure 3). Sheets of loose myxomatous tissue were also observed and interpreted as stellate reticulum-like tissue.

![Figure 1 – Orthopantomograph showing an osteogenic reaction of the subjacent bone. A relationship with the bone is not present](image)

Table 1: Cases of extraosseous calcifying odontogenic cyst reported in the literature until 2005

<table>
<thead>
<tr>
<th>Authors</th>
<th>Year of publication</th>
<th>Number of cases</th>
<th>Age</th>
<th>Gender</th>
<th>Localization</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mascrè et al.</td>
<td>1990</td>
<td>1</td>
<td>56</td>
<td>1 M</td>
<td>1 mandible</td>
</tr>
<tr>
<td>Buchner et al.</td>
<td>1991</td>
<td>7</td>
<td>10-92</td>
<td>3 M, 4 F</td>
<td>7 mandible</td>
</tr>
<tr>
<td>Hong et al.</td>
<td>1991</td>
<td>8</td>
<td>37-79</td>
<td>4 M, 4 F</td>
<td>7 mandible, 1 nonspecified</td>
</tr>
<tr>
<td>Moleri et al.</td>
<td>2002</td>
<td>1</td>
<td>66</td>
<td>1 F</td>
<td>1 mandible</td>
</tr>
<tr>
<td>Orsini et al.</td>
<td>2002</td>
<td>1</td>
<td>39</td>
<td>1 M</td>
<td>1 mandible</td>
</tr>
<tr>
<td>Fregnani et al.</td>
<td>2003</td>
<td>2</td>
<td>10-38</td>
<td>2 F</td>
<td>2 maxila</td>
</tr>
</tbody>
</table>
resembling the enamel organ and a variable number of ghost
cells (Figure 4). Irregular foci of calcified material resembling
dentin were visualized throughout the specimen.

Based on these features the proposed histopathologic
diagnosis was extraosseous calcifying cyst or Gorlin cyst.
Healing was uneventful, and the lesion had not recurred
after a four-year follow-up.

Discussion

The COC is also known as Gorlin cyst since its initial
description, in 1962, by that author(8). It is considered a
unique entity with both cystic and neoplastic behavior.
Peripheral varieties are usually located on the gingiva and
alveolar mucosa. They generally appear as circumscribed
elevated masses, with smooth surface and firm or soft
consistency. Since they have no characteristic clinical
appearance they can resemble granulomas, fibrous
hyperplastic or fibromas(12, 18, 19).

The presented case, similar to other reports in the
literature was an asymptomatic localized gingival swelling.
Clinical presentations of the ECOC are often described as
variable or non-specific(2, 5).

Our patient was a 57-year-old woman, but contradicting
the literature, her lesion was located in the retromolar
region. Review of the epidemiological characteristics of the
ECOC has shown that the tumor has predilection neither
for gender(6) nor for age(13, 15, 20). However, some authors
have described two peaks of occurrence in the second and
sixth decades(12, 12). Also, no overall predilection was shown
for either maxilla or mandible(12), although a tendency to
occur in the anterior region is described(2).

As an extraosseous lesion, COC exhibits no or minimal
radiographic changes(12). It may appear as a radiolucent area
with scattered amounts of opacities and no relationship
with the subjacent bone. We observed a slightly higher
condensation on the subjacent bone that was probably
caued by compressive pressure of the lesion.

Some attention concerning intraosseous lesions must
be taken to exclude cases that perforated the cortex and
exteriorized.

Although histological criteria have been established
for the diagnosis of the COC(17), its pathogenesis is still
speculative. Freedman et al.(6) proposed that the neoplastic
cell is originated from a well-differentiated ameloblast, and
its neural crest origin confirms to this cell a pleuripotential
capacity to undergo terminal differentiation. Starting from
the postulate that ameloblasts are stem cells, terminal
differentiation is not necessarily required to originate the
COC neoplastic cell. This fact might justify the ameloblastic-
like morphology observed in those neoplasms.

Praetorius et al.(18) and Buchner et al.(3) believe COC
cystic epithelium is originated from the reduced enamel
organ, from islands of odontogenic epithelium within
the tooth follicle or from the remnants of odontogenic
epithelium in the bone or gingival tissue. In the reported
case, we believe the neoplastic epithelium arised from the odontogenic remnants of the overlying mucosa due to the lesion intimacy with the oral surface and absence of tooth or bone involvement. But whether the cyst develops as central or peripheral lesion probably depends on the location of the odontogenic epithelium, which constitutes the source of the lesion. Nevertheless, the location does not seem to have any relation to either behavior or histological features of the cyst(3, 18).

According to some reports(11, 19), neoplastic or solid lesions constitute 2% to 10% of all cases, and are more often associated with peripheral location.

Treatment of the extrasosseous COC involves surgical excision(3-6, 12, 20) and recurrences are unexpected(1, 4, 20). Generally, cystic COC have good prognosis, but the neoplastic cases are uncertain. When a COC is associated with other odontogenic tumors, treatment and prognosis must be based on the associated lesion(9, 15, 21).

References