A rare case of bilateral mandibular simple bone cyst and the importance of differential diagnosis

Saulo Gabriel Moreira Falci a, Késsia Nara Andrade Sales b, Lorenna Vieira da Silva b, Cássio Roberto Rocha dos Santos c

ABSTRACT
Objective: To present a rare case of a mandibular bilateral simple bone cyst (SBC) with different radiological features and argue about the importance of the differential diagnosis.
Case description: A 16-year-old boy was referred by his dentist to evaluate two radiolucent mandibular lesions. The diagnostic hypotheses were ameloblastoma or keratocystic odontogenic tumor to the left side and odontogenic cyst on the right side due radiologic features. In the left side, needle aspiration and biopsy, were performed. During surgical approach there was an empty space in the surgical cavity compatible with SBC. At this moment the diagnostic was made and the cavity was submitted to rigorous curettage. The right side was assessed after 4 months. To our surprise, it was found an empty cavity, as the left side. Six months later the first biopsy the panoramic radiograph showed completely bone neoformation on both sides. The patient is in radiographic follow-up for three years, without relapse.
Conclusions: Bilateral SBC is a rare condition that can show different radiographic features. The diagnostic is defined at the surgery time. The radiographic features helps in diagnosis, however, should not be decisive. Surgical exploration should be done soon after radiographic diagnoses.

Key words: Bone cyst; Mandible; Curettage; Oral surgery
INTRODUCTION

Simple bone cyst (SBC) is a non-odontogenic intraosseous lesion that presents unknown etiology. The most common sites affected are the long bones, with only 10% of the cases in the jaw bones. Mandible is usually affected in the premolar and chin regions [1]. Despite the name, the SBC is not true cystic lesion, because the bone cavity is not usually coated by epithelium [2]. Other names for SBC are described in the literature as, idiopathic bone cavity, traumatic bone cyst and hemorrhagic bone cyst [3].

There are some theories to explain the etiology of the SBC as abnormal bone growth or tumor degeneration and intramedullary hemorrhage. Nowadays, the intramedullary hemorrhage theory is the most widely accepted to explain the etiology. The bone trauma, insufficient to fracture a bone healthy, may induce an intramedullary hemorrhage. The pressure of the hematoma causes venous stasis that promotes necrosis followed by osteoclastic resorption [1]. Some authors still states that this intramedullary hemorrhage may be caused by other factor as previous tooth extraction [4].

In the most of cases, the SBC is asymptomatic and found on routine radiographs [5]. Radiographic examination shows radiolucent lesions, with thin and irregular, but well defined, sclerotic margins [3]. In some cases SBC may have multilocular pattern, usually presenting appearance of “glove fingers” when it extends among the teeth roots [4]. Computed tomography may be helpful in diagnosis, because it allows assessing the intralesional density [6]. However, in most cases, the diagnosis is confirmed at the surgical moment by the presence of the empty cavity. The prognosis is excellent, but it takes a period of monitoring to ensure that the lesion will not relapse [1].

The differential diagnosis between SBC and other maxillary lesions is extremely important because the SBC does not show exclusive clinical and radiographic features. Understanding of clinical and radiographic features is extremely important to the oral and maxillofacial surgeon. Thus, this paper presents a rare case of bilateral mandibular SBC with different radiographic features, suggesting two different lesions.

CASE DESCRIPTION

A 16-year-old boy was referred to the School of Dentistry at the Federal University of Vales do Jequitinhonha e Mucuri, Brazil by his dentist, to evaluate two radiolucent lesions in the lower jaw, viewed on routine panoramic radiograph.

The interview does not reveal systemic symptoms, medication use, and history of trauma to the affected region. There were no reports of pain or swelling and maxillofacial exams showed no changes. The panoramic radiograph showed two lesions, one on the left and other on the right side. The right (between tooth 48 and 47) and left (from tooth 33 to tooth 38) lesions showed unilocular and multilocular characteristics, respectively. Both of them showed well-defined limits (Fig. 1). The teeth spanning 37 to 33 were vital. The diagnostic hypotheses were ameloblastoma or keratocystic odontogenic tumor to the left side and odontogenic cyst on the right side. Needle aspiration and biopsy were performed under local anesthesia, in the left lesion and, produced a yellowish liquid. During the surgical approach there was an empty space in the surgical cavity compatible with SBC (Fig. 2). At this moment the diagnostic of SBC was made and the cavity was submitted to rigorous curettage. To access the other lesion an arrangement with the patient was made to the following week. However, he did not show for the consultation. After four months the patient came back to the office. At this moment the right third molar was removed and, to our surprise, the approach showed an empty space below the third molar also compatible with SBC (Fig. 3). Six months later first biopsy, the patient has undergone to a new panoramic radiograph that showed completely bone neoformation in both sides (Fig. 4). The patient is in radiographic follow-up for three years, with no signs of clinical and radiographic relapse.
aspiration reinforced the diagnostic hypothesis for the multilocular lesion. Intra and extra-oral examination were normal, reflecting the potential of the lesion to develop among the marrow spaces. It is normal to see this lesion involving the interdental spaces, but, on the other hand root resorption, mobility, or losses of vitality are not seen [4], as we see in this case.

Treatment of simple bone cyst is usually a surgical exploration with vigorous bone curettage [3]. This procedure reestablishes the bleed inside of lesion, followed by clot reorganization and a bone neoformation due the presence of osteogenic cells [7]. The choice of treatment protocol depends on the radiographic characteristic of the lesion. In this case, the distinct radiographic features resulted in two different treatment plans, inciosional biopsy for the left side and excisional biopsy for the right side. This difference was based on the different diagnostic hypothesis. Even with different sizes and radiologic features both lesions were SBC, confirmed by intraoperative finding of an empty space. The occurrence of multiple SBC is rare in the English-language literature, and there are few reported cases, since 1929, when the first case was reported [5].

The most accepted theory for the development of SBC is through of intramedullary hemorrhage with a defect on healing or a clot lysis [1]. However, the etiology remains unclear, because in most of cases there is no clear history of trauma, as in this case. The lack of lymphatic synovial drainage has also proposed as a possible cause, once the interstitial fluid would accumulate in bone trabeculae, leading to hemodynamic disorder and consequently cyst formation [8]. In this case, the lesion development probably occurred due this disorder. This may explain the bilateral occurrence, since the theory of trauma would be more difficult to happen bilaterally. Histologically, bone covered by a thin layer of loose fibrous connective tissue may be seen [1]. However, in this case it was not necessary the histological evaluation, because the definitive diagnosis was made during surgical procedure, due a presence of empty cavity.

The treatment prognosis for SBC is excellent, however is necessary at least three years of radiographic follow up, to notice early evidence of treatment failure or relapse [9,10]. In this case, six months after curettage, new bone formation was observed in both lesions. Thus, after this period the chances of relapse are smaller once that it occurs after three months normally. However, this case is under three years of follow up, as preconized by Suei et al., [9] once this lesion can present slow growth.

SBC is a slow-growing lesion, asymptomatic, found in routine radiograph evaluation in young patients. Bilateral SBC is a rare condition that can show different radiographic features. Diagnostic is defined at the surgery time. The period of follow up should be long enough to prove the success of treatment and treat early any relapse. The radiographic features helps in diagnosis, however, should not be decisive as shown in this case. Thus, surgical exploration and biopsy should be done immediately after radiographic diagnoses.

Figure 3. Surgical area after third molar extraction and osteotomy of the left side, where we can see an empty cavity compatible with simple bone cyst

Figure 4. Panoramic radiograph six months after the first surgery showing a new bone formation in both sides

DISCUSSION

The differential diagnosis of SBC may be any radiolucent lesion as, keratocystic odontogenic tumor, ameloblastoma, dentigerous cyst, central giant cell lesion among other [2]. In this case, the distinct radiographic features have induced two different diagnostic hypotheses, ameloblastoma or keratocystic odontogenic tumor to the multilocular lesion and odontogenic cyst to the unilocular lesion. Needle aspiration reinforced the diagnostic hypothesis for the multilocular lesion. Intra and extra-oral examination were normal, reflecting the potential of the lesion to develop among the marrow spaces. It is normal to see this lesion involving the interdental spaces, but, on the other hand root resorption, mobility, or losses of vitality are not seen [4], as we see in this case.

Treatment of simple bone cyst is usually a surgical exploration with vigorous bone curettage [3]. This procedure reestablishes the bleed inside of lesion, followed by clot reorganization and a bone neoformation due the presence of osteogenic cells [7]. The choice of treatment protocol depends on the radiographic characteristic of the lesion. In this case, the distinct radiographic features resulted in two different treatment plans, inciosional biopsy for the left side and excisional biopsy for the right side. This difference was based on the different diagnostic hypothesis. Even with different sizes and radiologic features both lesions were SBC, confirmed by intraoperative finding of an empty space. The occurrence of multiple SBC is rare in the English-language literature, and there are few reported cases, since 1929, when the first case was reported [5].

The most accepted theory for the development of SBC is through of intramedullary hemorrhage with a defect on healing or a clot lysis [1]. However, the etiology remains unclear, because in most of cases there is no clear history of trauma, as in this case. The lack of lymphatic synovial drainage has also proposed as a possible cause, once the interstitial fluid would accumulate in bone trabeculae, leading to hemodynamic disorder and consequently cyst formation [8]. In this case, the lesion development probably occurred due this disorder. This may explain the bilateral occurrence, since the theory of trauma would be more difficult to happen bilaterally. Histologically, bone covered by a thin layer of loose fibrous connective tissue may be seen [1]. However, in this case it was not necessary the histological evaluation, because the definitive diagnosis was made during surgical procedure, due a presence of empty cavity.

The treatment prognosis for SBC is excellent, however is necessary at least three years of radiographic follow up, to notice early evidence of treatment failure or relapse [9,10]. In this case, six months after curettage, new bone formation was observed in both lesions. Thus, after this period the chances of relapse are smaller once that it occurs after three months normally. However, this case is under three years of follow up, as preconized by Suei et al., [9] once this lesion can present slow growth.

SBC is a slow-growing lesion, asymptomatic, found in routine radiograph evaluation in young patients. Bilateral SBC is a rare condition that can show different radiographic features. Diagnostic is defined at the surgery time. The period of follow up should be long enough to prove the success of treatment and treat early any relapse. The radiographic features helps in diagnosis, however, should not be decisive as shown in this case. Thus, surgical exploration and biopsy should be done immediately after radiographic diagnoses.
REFERENCES


