Squamous Odontogenic Tumor: A Rare Case Report

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ABSTRACT

Squamous odontogenic tumor is a rare benign locally infiltrative neoplasm first described by Pullon et al. (1975). Since then, there have been less than 50 cases reported. The tumor is often asymptomatic, although it can present with symptoms of pain and tooth mobility. The characteristic radiographic appearance is that of a triangular-shaped unilocular radiolucency with divergent roots of the involved teeth. Histologically, the tumor is characterized by the proliferation of benign appearing, squamous epithelium in the connective tissue stroma with occasional vacuolization and keratinization. Here, we report a rare case of Squamous odontogenic tumor in anterior mandible of a 40 years old female patient.

Key words: odontogenic tumor, Squamous epithelium

INTRODUCTION

Squamous odontogenic tumor (SOT) is a rare benign neoplasm first described in 1975 by Pullon et al.¹ It arises from the neoplastic transformation of epithelial rests of Malassez. The World Health Organization (WHO) in 2005 classified it as an epithelial odontogenic tumor with around 50 cases reported in the literature.²

SOT is defined as a benign but locally infiltrative neoplasm. It affects a wide age range showing a slight male preponderance and occurs more frequently in the mandible with preference for posterior mandible and anterior maxilla.² Maxillary lesions appear to be more aggressive in biologic behavior than those occurring in mandible.³ Radiographically, SOT is seen as a unilocular radiolucent triangular shaped area whose base is localized at the apices of the diverging adjacent roots.⁴ Histologically the lesion consists of islands of squamous epithelium in a fibrous stroma. Cystic degeneration or calcification may occasionally be observed in the epithelial islands.⁵ Clinico-pathologically, three main types are identified: Intraosseous, Mural and Extraosseous forms.¹ Intraosseous or central type is more common. Mural type presents as SOT like proliferation in the wall of cyst. Extraosseous or peripheral type is rare and only one case has been reported in the literature.¹

CASE REPORTS

A 40 years old female patient reported with the complaint of an asymptomatic swelling on the right side of mandible since 4 months. On clinical examination hard swelling of about 1.5 x1.5 cm was seen on buccal aspect of lower anterior region wrt 42, 43 and 44. The overlying mucosa was normal. On radiographic examination (Fig.1), areas of mixed radiolucency and radiopacities were seen wrt 42 and 43 and the roots of the same were divergent. On the basis of clinical and radiographical findings, clinical differential diagnosis were made of Ossifying fibroma or Calcifying epithelial odontogenic tumor. The incisional biopsy of the lesion was taken (Fig.2 & Fig.3) and sent for histopathological examination. Histological examination revealed proliferation of mature stratified squamous epithelial islands in a dense fibrous connective tissue stroma (Fig.4) These islands showed peripheral flat or cuboidal shaped cells and central squamous cells with intercellular bridges (Fig.5).

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Areas of cystic degeneration and keratinization were seen within these islands (Fig.6). Thus depending on clinical and histological features the final diagnosis of SOT was confirmed.

DISCUSSION

Squamous odontogenic tumor is a rare benign neoplasm with very few cases reported in literature till date. Pullon et al in 1975 reported six cases and described SOT for the first time. They established the diagnostic criteria and surgical approaches that are still followed today. Before 1975, this lesion was believed to represent an atypical acanthomatous ameloblastoma or even a squamous cell carcinoma.

Most of the cases reported in literature have been found to be located within the bone (intraosseous), although a few peripheral cases have also been discussed. The central or intraosseous variety is thought to arise from the rests of Malassez or derived from the periodontal ligament whereas the peripheral variety is thought to arise from the gingival surface epithelium or the rests of Serres.

SOT has been found in wide age range, from first decade to eighth decade of life, with mean age of occurrence being 38.2 years. Badni et al reviewed 44 cases of SOT and found slightly more predilection among males compared to females (F: M=1:1.8). Considering the reported cases, the most common location for development of SOT in the maxilla is anterior region and in case of mandible is posterior region. In our case the lesion was located in the anterior region of mandible which is in consistence with the finding of Bansal et al.

It has been stated that SOTs occurring in maxilla were found to be more aggressive than in mandible which is mainly due to the anatomy and porous nature of bone.

Clinically, SOT appears as a slow growing lesion, which leads to an increase in the volume of the maxilla or mandible, tooth mobility, ulceration of the soft tissue, painful symptoms, and tooth displacement. However, lesion may sometimes be asymptomatic and detected only on routine radiographic investigations.

Radiographically, common central variant of SOT shows a well-defined unilocular, triangular radiolucency between the roots of adjacent teeth, but in our case areas of mixed radiolucency and radiopacity was seen. The peripheral variant may show saucerization of underlying bone which is likely to be a pressure phenomenon rather than the result of true tumor infiltration.

Histologically, the lesion usually presents as islands of benign squamous epithelium in mature connective tissue stroma without the evidence of peripheral columnar cells, palisading nuclei, or stellate reticulum. Cystic degeneration in the center of the islands is a frequent finding. In some epithelial cells prekeratin is present and laminated calcifications may be seen inside keratin pearls. Sometimes the morphologic
appearance of the islands of odontogenic epithelium in SOT is similar to the follicular pattern of ameloblastoma, which may lead to the misdiagnosis of the tumor. However, lack of polarization of peripheral cells in the epithelial islands, which is typical of ameloblastoma, favors the diagnosis of SOT.\textsuperscript{17} In our case, squamous epithelial islands in mature connective tissue stroma, presence of intercellular bridges and areas of microcystic degeneration and absence of ameloblasts like cells at periphery was of major importance in establishing the diagnosis. Newer diagnostic techniques such as immunohistochemistry has shown expression of keratin and Notch receptor (Notch 1, 3, 4) and its ligand (jagged 1 and delta 1) in SOT.\textsuperscript{1} However, routine eosin and hematoxylin technique remains the gold standard for the diagnosis of SOT.

SOT can be managed conservatively by local excision, curettage, enucleation and scaling of the teeth. In case of maxilla, since the lesion is more aggressive, the lesion has also been treated surgically by en bloc excision and hemimaxillectomy.\textsuperscript{8,18,19} Recurrence of the lesion is rare and only a single case of recurrence has been reported in the literature, which may be due to incomplete removal of the lesion. Local curettage or extraction of adjacent teeth was found to be effective in management of recurrences.\textsuperscript{19} Ide et al. in 1999 reported malignant transformation of SOT such as Intraosseous Squamous Cell Carcinoma.\textsuperscript{20} Kim et al. in 2007 reported two cases of SOT that led to erosion of lingual cortical plate of mandible, which may impact the therapeutic modalities and management of these lesions. This indicates that these lesions do not always behave similarly.\textsuperscript{21}

CONCLUSION

Rare as it may seem since its first description till date SOT has always been an enigma in the field of dentistry. Hence analysis of each case clinically, radiographically and histologically is important in diagnosing the disease and choosing the right treatment modality.

REFERENCES

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