PERIPHERAL AMELOBLASTOMA- A CASE REPORT WITH EMPHASIS ON ETIOPATHOGENESIS

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ABSTRACT:
Peripheral ameloblastoma, a rare and unusual variant of odontogenic tumour, comprises about 1% of all ameloblastomas. The extraosseous location is the peculiar feature of this type of tumour, which is otherwise similar to the classical ameloblastoma. It appears in the gingiva and oral mucosa and it usually does not show any bone involvement on radiographs, except for saucer shaped erosion of underlying alveolar bone. Recurrence is considered uncommon. We report a case of peripheral ameloblastoma of maxillary gingiva.

KEYWORDS: peripheral ameloblastoma, odontogenic tumours, alveolar mucosa, maxilla

INTRODUCTION
Peripheral ameloblastoma (PA) is a rare, benign epithelial odontogenic tumor, occurring in the soft tissues overlying the tooth bearing areas of maxilla and mandible. Clinically it presents as a firm, slow growing, and dome shaped swelling with a smooth surface covered with normal mucosa. Ideally they do not show any bony changes on radiographs. Conservative surgical excision with a small margin of healthy tissue is the recommended treatment. We present and analyse a case of peripheral ameloblastoma in the anterior region of maxilla, with special emphasis on etiopathogenesis.

Case report
A 40 years old male patient presented to our department with a swelling in the gingiva of right maxillary region. He had noticed the swelling a month back. The swelling was gradually increasing in size and was totally asymptomatic. Past dental and medical histories were non-contributory. On intraoral examination a solitary, well defined dome shaped swelling was present in the interdental area of 13 and 14 measuring about 1.5 cm in size. Swelling was non tender and firm in consistency with normal overlying mucosa (Fig.1). The teeth in the vicinity were non tender, undisplaced and vital. Based on history and clinical examination a provisional diagnosis of peripheral ameloblastoma in the anterior region of maxilla, with special emphasis on etiopathogenesis.

which time, bony erosion was noticed beneath the lesion (Fig.2). The specimen was fixed in 10% formalin and sent for histopathological examination. Histologically it comprised of deeper tissue islands, strands of odontogenic epithelium with peripheral tall columnar cells with centrally stellate reticulum like cells suggestive of peripheral ameloblastoma (Fig.3). The healing was uneventful and the patient has been followed up at regular intervals for past nine months and he is disease free (Fig.4).

Discussion
Ameloblastoma is a benign epithelial odontogenic tumor with three main clinical variants- solid/multicystic (86%), single cyst (13%) and peripheral ameloblastoma (1%). So peripheral ameloblastoma (extraosseous ameloblastoma) is a rare tumor occurring in the soft tissues of the tooth bearing areas of maxilla and mandible. Several authors refer to Kuru as having reported on the peripheral ameloblastoma for the first time. But what Kuru described was not a PA, infact it was intra-osseous ameloblastoma, which penetrated through the alveolar bone and got fused with oral epithelium giving an impression of a peripheral lesion. The first completely documented case of a PA is attributed to Stanley and Krogh.

Ameloblastoma is an epithelial odontogenic tumour of the jaw bones which is thought to arise from rests of the dental lamina or from basal cells of the surface epithelium. A recent investigation demonstrated that alterations of the ameloblastin gene form the genetic basis for ameloblastoma.
The origin of PA at cellular level, continues to be a challenge to academicians and researches alike. However two sources of origin should be considered. One, it arises from the remains of the dental lamina located in soft tissues overlying the tooth bearing areas of the jaw bones. In this case, the lesion is totally separated from the surface epithelium by a band of collagenous tissue. Stanley and Krough had mentioned that their case probably arose in this manner from odontogenic epithelial cell rests. Two, it arises directly from the basal cell layer of the overlying epithelium. But however, even the demonstration of fusion between overlying epithelium and tumor tissue will not clearly prove the origin, as the tumor can grow subepithelially and may get fused with the surface epithelium. Even electron microscopic studies have not been able to resolve this challenge emphatically.1,7

Peripheral Ameloblastoma is noticed during 4th to 6th decades of life with an average age of 50 years. It is interesting to note that the average age of occurrence of PA is much higher than its central counterpart (37.4 years). A slight male predominance has been reported with male/female ratio being 1.9:1. Mandible is most favoured site than maxilla. In mandible it is more often located on lingual aspect on premolar region, while in maxilla its common location being tuberosity area.8 In contrary in our case swelling was located in maxilla in canine premolar region on facial aspect.

Clinically it presents as a sessile, firm growth of gingiva measuring around 0.5 to 2 cm in size. The surface is usually smooth and pink/red in colour. Clinical appearance was similar in the present case, with swelling measuring about 1 X 1.5 cm, with normal overlying mucosa and smooth surface. Usually they are asymptomatic. In majority of cases there are no radiographic bony changes. In few cases, a superficial erosion of the bony or bony
depression can be noticed, which might be caused due to pressure resorption rather than by the invasion of the tumor. Clinically it needs to be differentiated from other lesions which are common in gingiva.

Pyogenic granuloma: it is sessile/pedunculated lesion, common in maxillary gingiva. The colour ranges from pink to red to purple. It bleeds profusely with little manipulation. Peripheral giant cell granuloma: common in mandibular gingiva. The colour ranges from red to bluish purple, but usually more bluish than pyogenic granuloma. It can also cause destruction of underlying bone (sauерization).

Peripheral ossifying fibroma: gingiva of maxilla is more often involved than mandible. It is common in younger individuals. The surface can be smooth or lobulated. Early lesions are red in colour but mature lesion appear pale pink. Scattered irregular radiopacities can be seen on the radiograph.

PA can be successfully managed with conservative supraperiosteal surgical excision with adequate distance from margins. Although there were no bony changes on radiograph in our case, but on surgical exploration resorption of cortical plate (sauерisation) was evident beneath the lesion.

Local recurrence rate is much lower in comparison with central ameloblastoma. There are reports of aggressive behaviour of PA invading intracranial structures and malignant transformation. Keeping that in mind, a long term follow up is prudent. Our patient is examined periodically every month, he is disease free since nine months. So PA should be considered in differential diagnosis of swellings located in the gingiva. But, what is more important is the way it needs to be managed; the word “ameloblastoma” sends a strong signal for the surgeons for aggressive treatment plan which is not at all mandated. A simple conservative excision is all that one needs to do in peripheral ameloblastoma.

References

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