Case Report

An infrequent histopathological subtype of ameloblastoma: Adenoid granular cell ameloblastoma with dentinoid

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ABSTRACT

Adenoid ameloblastoma with dentinoid is a rare odontogenic tumor. Granular cell ameloblastoma also is a less common histological subtype of ameloblastoma. In this report, the patient was a 31-year-old male. The lesion was located in the right mandible and was unicystic with well-defined borders. The tumor tissue was showing a combination of follicular, plexiform, and desmoplastic patterns of ameloblastoma with wide areas of granular cells, fibrous stroma, glandular pattern, and dentinoid calcified. Very few cases of distinct forms of ameloblastoma that show the formation of dentinoid has been reported. However, there are no cases of adenoid granular cell ameloblastoma with dentinoid reported.

Key Words: Adenoids, ameloblastoma, cytoplasmic granules, odontogenic tumors

INTRODUCTION

Ameloblastomas are benign but locally invasive tumors of odontogenic epithelial origin and generally without induction in the connective tissue.[1] Rare cases of ameloblastoma associated with an odontome have been reported and are known as “odontooameloblastoma.”[2] In 1992, the first case of an ameloblastoma showing evidence of induction of dentinoid which diagnosed as a “dentinooameloblastoma” was reported by Slabbert et al.[3] The term “adenoid ameloblastoma with dentinoid” was first introduced by Evans et al. in 2004. They described a neoplasm showing histopathological features similar to ameloblastoma and adenomatoid odontogenic tumor (AOT) along with hard tissue formation.[4] Histopathologically, ameloblastoma exhibits follicular, plexiform, acanthomatous, granular cell, desmoplastic, and basal cell patterns. Granular cell ameloblastoma is rare to occur and accounts for only 5% of all ameloblastomas.[5] Based on current knowledge, there is no reported case of granular cell ameloblastoma with dentinoid in the literature. In this paper, we present a case of a rare histological subtype of ameloblastoma with large amounts of granular cells, glandular pattern, and dentinoid areas in the right mandible of a 31-year-old male. Final histopathological diagnosis of adenoid granular cell ameloblastoma with dentinoid was made for this entity.

CASE REPORT

A 31-year-old man was referred to us with a chief complaint of a swelling on the left side of the
mandible, which had been presented for nearly 1 month. Clinical examination revealed a localized expansion of the left mandible covered by normal oral mucosa.

The swelling was painful with palpation, soft in consistency, and fluctuant in nature. No history of numbness or other abnormal sensations was mentioned. The tumor caused mobility and displacement of left mandibular canine, first and second premolars. The involved teeth were vital. Cervical lymph nodes were not palpable. Patient’s medical and dental history was unremarkable.

Radiographically, panoramic view showed a unilocular radiolucency with well-defined borders extending from the left canine to the left mandibular second molar. There was root divergence, but no root resorption and the lamina dura of the affected teeth were destroyed [Figure 1a].

Cone beam computed tomography scan showed a radiolucent lesion in the left mandibular area with buccal and lingual cortices destruction [Figure 1b].

Radiographic differential diagnosis includes central giant cell granuloma, ameloblastoma, odontogenic myxoma, and keratocystic odontogenic tumor was given. Then, incisional biopsy was performed. Specimen was sent to the department of oral and maxillofacial pathology. The sections showed a combination of follicular, plexiform, and desmoplastic patterns of ameloblastoma [Figure 2a and b] with wide areas of granular cells and fibrous stroma [Figure 2 c and d]. Dentinoid calcified [Figure 3a and b] and occasionally glandular area were observed in some sections [Figure 3c and d].

Considering these features, a provisional histopathological diagnosis was considered.

The patient was submitted for left partial hemimandibulectomy from left central incisor to left third molar area.

Histopathological examination of the last specimen showed neoplastic proliferation of odontogenic epithelial cells in follicular and plexiform patterns and large amounts of granular cells. Occasionally, wide fibrous stroma, glandular pattern, and dentinoid areas were observed. Considering the histopathological features of present and previous tissue sections, the diagnosis of adenoid granular cell ameloblastoma with dentinoid was confirmed. The patient is under close follow-up and shows no evidence of recurrence 18 months after the operation.

DISCUSSION

Granular cell ameloblastoma is a rare subtype of ameloblastoma in which granular cells located in the center of the follicles. Its median age is 35-year-old with no distinct gender predilection in the literature.
Granular cell ameloblastoma should be differentiated from other odontogenic and nonodontogenic lesions along with granular cells, including granular cell tumor, congenital epulis, and granular cell odontogenic tumor.\(^9\)

The granular cell tumor is a soft tissue tumor and should be distinguished from ameloblastomas that cause cortical perforation and soft tissue extension.\(^9\) Congenital epulis is also a soft tissue tumor and with regards to its age outbreak, there is no need to distinguish it from a granular cell ameloblastoma.\(^9\) Granular cell odontogenic tumors exhibit a proliferation of large cells with eosinophilic granular cytoplasm and may show dystrophic calcifications.\(^10\) Granules of granular cells in granular cell ameloblastoma are positive for cytokeratin, CD68, lysozyme, and alpha-1-antichymotrypsin, and in granular cell odontogenic tumor are positive for vimentin, CD68, lysozyme, muscle-specific actin, alpha-smooth muscle actin, calponin, neuron-specific enolase, CD138, and bcl-2 in immunohistochemistry (IHC) staining.\(^11\) Granular cell lesions are easy to rule out with observing follicular, plexiform, and desmoplastic patterns of ameloblastoma in the specimens.

In this case, differential diagnosis between adenoid ameloblastoma with dentinoid and AOT is also important. This is due to the presence of glandular pattern and dentinoid areas in the histopathological assessment. Duct-like structures are the characteristic of the AOTs, and dystrophic calcifications are usually seen in these lesions.\(^10\) Considering that granular cell is not exist in AOTs, distinction between the present case and AOT was performed without IHC.

Few reports of ameloblastoma along with hard tissue formation were found in the literature. Bone formation was only reported in one neoplasm,\(^12\) two polycystic ameloblastomas showed diffuse interstitial ossification\(^13\) and enamel formation were described in few reports.\(^14\) In most other reported cases, the hard tissue had been construed as dentin or dentinoid.\(^15\) Consequently, hard tissue in these unusual cases of ameloblastoma has a wide variety of nature.

Finally, because of an existing neoplasic proliferation of odontogenic epithelial cells in follicular, plexiform and glandular patterns, dentinoid areas, and abundant granular cells, we concluded the present tumor to be a rare case of adenoid granular cell ameloblastoma with dentinoid. To make accurate diagnosis, it is essential that all the oral and maxillofacial pathologist to be aware of this entity.

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Conflicts of interest
The authors of this manuscript declare that they have no conflicts of interest, real or perceived, financial or non-financial in this article.

REFERENCES


