Desmoplastic ameloblastoma with osteoplasia: Review of literature with a case report

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ABSTRACT
Desmoplastic ameloblastoma (DA) is a relatively rare histological variant of ameloblastoma with specific clinical, radiological, and histological features. This tumor is more commonly seen in the anterior region of jaws as a mixed radiopaque-radiolucent lesion resembling benign fibro-osseous lesions. Histologically, DA is characterized by small nests and strands of “compressed” odontogenic epithelium supported by pronounced collagenized stroma. Metaplastic bone formation has been reported in few cases. This report describes a lesion in the left anterior maxilla of a young female patient.

Key words: Ameloblastoma, desmoplasia, osteoplasia

INTRODUCTION
Ameloblastoma is the second most common odontogenic tumor. It arises from odontogenic epithelium and is known for its distinct aggressive clinical behavior and characteristic histologic picture. Based on clinical and radiographic characteristics, histopathology, behavioral and prognostic aspects, the subtypes of ameloblastoma are classic solid/multicystic, unicystic, peripheral and desmoplastic types.[1] The first detailed report in English literature on desmoplastic ameloblastoma (DA) was given by Eversole in 1984, who called it an “ameloblastoma with pronounced desmoplasia.”[2]

DA is a rare variant of ameloblastoma accounting for approximately 4-13% of ameloblastomas.[3] It is characterized by an unusual histomorphology, including extensive stromal collagenization or desmoplasia. A characteristic feature is an almost equal distribution in location between the maxilla and mandible. Radiologically, the desmoplastic variant exhibits atypical and varied radiographic features such as: Localized irregular multilocular radiolucency with indistinct borders, or a mixed radiopaque-radiolucent appearance with ill-defined margins similar to fibro-osseous lesion, or a massive expansile osteolytic lesion with honeycomb, mottled or multilocular appearance.[4] Tooth displacement is a common feature. DA may be defined as a benign but locally invasive variant consisting of proliferating, irregular, often bizarrely shaped islands and cords of odontogenic epithelium of varying sizes embedded in a desmoplastic connective tissue stroma. Formation of metaplastic bone trabeculae (osteoplasia) rimmed by active osteoblasts has been described in a few cases.[5] A possible transitional form of DA, showing microscopic features of desmoplastic variant together with areas of classical follicular/plexiform ameloblastoma has been described as a “hybrid lesion.”[6]

CASE REPORT
A 26-year-old female reported with a swelling in relation to the upper front teeth. The swelling was slow in growth and present since 4 months. She had also noticed a slight downward movement of the affected teeth and had mild pain. On examination, a bony hard swelling was seen in relation to 21, 22 and 23 on the buccal aspect measuring about 2.5 cm × 1.5 cm and extending into the palate. It was non-tender. Grade I mobility of 21 and 22 was also noticed.

Radiographs revealed the presence of irregular radiolucencies with faint radio-opacities in the periapical region of 22 and 23. Lamina dura of the teeth was hazy and teeth were seen to be displaced [Figure 1]. A provisional diagnosis of benign fibro-osseous lesion was made and an incisional biopsy was done.

Both hard-tissue and soft-tissue were received. The specimens showed few irregular epithelial islands in a densely fibrous connective stroma. The islands consisted of peripheral cuboidal cells and central spindle cells. Some of the islands showed cystic degeneration [Figure 2]. Extensive desmoplasia...
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was seen throughout the stroma compressing the epithelial islands [Figure 3]. Areas of osseous metaplasia with active osteoblasts were also noted [Figure 4]. With the above picture, a diagnosis of DA with osteoplasia was made.

DISCUSSION

Clinically, DA may occur in all age groups with more cases in the 4th to 5th decade. There is no gender predilection and is seen equally in males and females. In this aspect, it is comparable with the conventional ameloblastoma. DA is reported to occur in the anterior or premolar region of either maxilla or mandible. This is in contrast to the conventional ameloblastoma which is found predominantly in the posterior region of mandible. Usually, the patient’s first symptom is a painless swelling. Involved teeth may be displaced with cortical plate expansion. The size of DA is mostly smaller than other types of ameloblastoma. This could be attributed to the fact that it is detected earlier because of its anterior location. A case of peripheral DA has also been reported recently. The clinical presentation in our case falls in the same category as generally accepted.

The radiographic features of DA also differ from the conventional ameloblastoma. A wide range of appearance from radiolucency to radioluency with flecks of opacity to mixed radiolucent/radio-opaque has been reported. Conventional ameloblastoma usually presents as a well-defined unilocular/multilocular radioluency. Again, the borders of DA are usually poorly defined. DA usually infiltrates into marrow spaces and surrounding non-neoplastic bone without a fibrous capsule. Also, osteoplastic bone may be produced by the lesion. The unique radiographic picture seen is thought to be due to its infiltrative nature. The presence of osteoplastic bone can also present similarly. The unconventional clinical site along with the mixed radiolucent/radio-opaque picture, in our patient, led us to the diagnosis of a benign fibro-osseous lesion.

Histological features consisted of proliferating, irregularly shaped islands and cords of odontogenic epithelium of varying sizes embedded in a desmoplastic connective tissue stroma. The tumor islands may show a pointed, stellate or “kite-like” appearance. The epithelial cells at the periphery of the islands are cuboidal and occasionally show hyperchromatic nuclei.
Columnar cells demonstrating reversed nuclear polarity are rarely conspicuous although some islands may show ameloblast-like cells peripherally. The center of the islands may contain spindle-shaped or squamoid epithelial cells. DA exhibiting granular cell transformation in some of the tumor cells has been reported. Furthermore, a case with associated follicular, plexiform, acanthomatous and basaloid changes has been documented. True glandular differentiation with mucus cells has been reported in the tumor islands. Cystic degeneration within the tumor has also been described. In our case, few islands with kite-like appearance with spindle cells in the interior of the island and stromal desmoplasia were noted.

Extensive stromal desmoplasia is a striking finding with a tendency to squeeze or compress the odontogenic islands at the periphery. The mechanism of desmoplasia is not understood even after two decades. Various immunohistochemical studies have reported DA tumor cells as showing variable expression of S-100 protein and desmin, high expression of caspase-3 and Fas, decreased expression of cytokeratin 19 and high expression of p63. Intense staining for collagen Type IV adjacent to tumor islands has been noted in DA. This indicates active synthesis of extracellular matrix proteins and that the stroma is not scar tissue. Marked expression of transforming growth factor-β has been demonstrated and it may play a part in the desmoplasia. It has also been suggested that it may indicate a maturation process and lesions in the anterior region may mature faster, hence the finding of desmoplasia. Oxytalan fibers have been identified in the stromal tissue. This finding indicates that this tumor may be periodontal in origin. The extracellular matrix has an important role in the behavior of neoplastic cells and immunohistochemical studies on the expression of tenasin and fibronectin proteins and Type I collagen in hybrid lesions have also been reported. Both tenasin and fibronectin showed positive staining around the conventional follicles rather than the desmoplastic areas. This indicates the greater aggressive potential of conventional follicular ameloblastoma as compared to DA as tenasin has been observed in unstable environments like neoplasia. Type I collagen showed intense staining in a fibrillar pattern in DA areas reflecting its high grade in the matrix.

Prominent osteoplasia within the tumor tissue seen in our case has been reported only in eight other cases of DA so far. The metaphasic bony trabeculae varied from woven to mature bone, which contained osteocytes and were lined by plump active osteoblasts. Remnants of non-neoplastic bone often seem to remain in the tumor tissue. Studies have been done to find whether the bone found in new bone is mature bone or osteoplasia. Fibrous bone can be recognized using Masson-trichrome staining. The presence of peripheral un-calcified fibrous bone gave the impression of newly forming bone rather than destroyed trabecular bone. Hence it can be suggested that if the desmoplasia is caused by tumor cell stimulation of stromal fibroblasts, then that stimuli could also affect another cell type, i.e., osteoblasts for formation of new bone (osteoplasia).

CONCLUSION

The biologic behavior of the lesion is still debatable. Conflicting reports are available regarding the recurrence because of paucity of adequate samples. The radiographic picture suggests an infiltrative process with the propensity to recur and the stromal reaction may be viewed as a defensive response of the host to the aggressive tumor. Furthermore, the lesions in maxilla are close to many vital structures and the thin cortical bone is a weak barrier and the tumor usually infiltrates into narrow spaces. Thus, a clear-cut border may be difficult to identify. The radiological and histological findings of poor encapsulation and ill-defined borders warrants a long-term follow-up. A radical approach to its treatment is recommended as in a case of conventional ameloblastoma.

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