Myoepithelioma of minor salivary glands – A diagnostic challenge: Report of three cases with varied histomorphology

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ABSTRACT
Myoepitheliomas are rare, benign neoplasms affecting predominantly parotid gland and to a lesser extent minor salivary glands. In this article we have reported three cases with different histomorphological patterns. Two cases are from oral cavity and one from sinonasal tract, a very rare location. We have discussed different histomorphological patterns of myoepitheliomas, which at times possesses a real diagnostic dilemma to a pathologist. Along with its morphology, immunohistochemical profile and clinical behavior are discussed in detail with relevant review of literature.

Key words: Myoepithelioma, minor salivary glands, sinonasal

CASE REPORTS

A detailed clinical profile of all the three cases is shown in Table 1.

Table 1: Detailed clinical profile of three cases

| Age (years)/sex | Anatomic location       | Size (cm)  | Clinical diagnosis       |
|----------------|------------------------|------------|--------------------------|
| 70/F           | Lateral wall of Rt. nasal cavity | 3.5 × 2 × 2 | Nasal Polyp             |
| 62/F           | Ulcerated cystic mass, soft palate | 1.4 × 1.3  | Haemangioma              |
| 30/F           | Hard palate             | 1.5 × 0.8  | Pleomorphic adenoma      |

INTRODUCTION

Myoepithelioma of salivary glands are very rare, accounting for only 1%-1.5% of all salivary gland neoplasms. The most common site is parotid gland with a few arising in minor salivary glands of the oral cavity and palate.[1] Since 1943 when Sheldon identified them and characterized them as myoepithelioma, less than 100 cases have been reported in the literature.[2] Reports of the tumor from sinonasal tract are scarce in most reviews.[3]

These tumors exhibit complex and variable histomorphological pattern, immune antigenic expression, and clinical behavior leaving many gray areas needing further exploration and study. This has prompted us to report three cases we came across in the recent past having different histomorphology and rare sites of origin. Two arose in the oral cavity and one in the sinonasal tract, an extremely rare site. The clinicopathological profile and histopathological diagnosis are highlighted with review of relevant literature.

CASE REPORTS

Case 1

Gross: Received two grayish white masses measuring 2.5 cm × 2 cm × 2 cm and 1 cm × 0.2 cm each. The histopathological examination showed a tumor tissue containing loose sheets of fairly uniform, plump, oval to spindle cells. These cells were having hyaline eosinophilic cytoplasm and eccentric nuclei giving them plasmacytoid appearance [Figure 1a and b]. Occasional ductal elements were seen entrapped within the tumor. There was no evidence of necrosis or abnormal mitoses.

The differential diagnoses considered were plasmacytoma and myoepithelioma. Immunohistochemistry: S100-Uniform strong positivity [Figure 1c] and smooth muscle actin (SMA)-Focal strong positivity [Figure 1d].

Final diagnosis: Plasmacytoid (hyaline) myoepithelioma of sinonasal tract.

Case 2

Gross: Received grayish brown ulcerated oval cystic mass, 1.5 cm in diameter. Cut surface was grayish white soft,
mucoid, partially cystic. The histopathological examination showed tumor tissue separated from normal salivary gland by fibrocollagenous tissue. The tumor was made up of solid sheets of round to oval cells having moderate amount of clear cytoplasm and regular round to vesicular nuclei [Figure 2a and b]. The diagnosis made was clear cell myoepithelioma.

Immunohistochemistry: S100 and SMA-positive [Figure 2c and d].

Final diagnosis: Clear cell myoepithelioma of soft palate.

Case 3

Gross: The specimen consisted of two grayish white soft tissue pieces, together measuring 1.5 cm × 1 cm.

The histopathological examination showed a well-circumscribed tumor made up of anastomosing cords of epithelial cells separated by abundant intercellular hyaline material. These cells were having moderate amount of eosinophilic cytoplasm and round to oval monomorphic nuclei [Figure 3a and b]. The differential diagnoses considered were monomorphic adenoma and reticular (epithelioid)-type myoepithelioma.

Immunohistochemistry: S100 and SMA: strong positivity [Figure 3c and d].

Final diagnosis: Reticular (epithelioid) myoepithelioma of hard palate.

DISCUSSION

Parotid gland and to a lesser extent glands in the palate are the preferred locations for myoepitheliomas. There are no specific clinical features that distinguish them and they usually present as slow-growing painless masses. None of our cases were in parotid gland and one was in a very rare site, namely, nasal cavity. Only three to four tumors reported in the literature earlier were at this site. Although no predilection with regard to age and gender was noted in the literature, which is similar to that of pleomorphic adenoma, all our cases were females in the age group of 30-70 years.

Myoepitheliomas of the parotid are usually encapsulated lesions without ulceration. Palatal lesions may be unencapsulated as in our case 3 and ulcerated as was in our case 2. Histomorphologically, standard text books describe four types mainly spindle cell, hyaline/plasmacytoid, clear cell, and oncocytic (a variant of spindle cell). Others identify the four types as fusiform, epithelioid, plasmacytoid, and clear cell types. There may be mixtures. Pilch et al., recognize four types as spindle cell, hyaline/plasmacytoid, epithelioid, and clear cell type. The epithelioid variant shows interconnected cords of cells with abundant mucoid stroma, also labeled as reticular type by Dardick. Morphology of one of our cases (Case 3) was consistent with that of this type [Figure 3a and b].

All of the few sinonasal myoepitheliomas reported in the literature were spindle cell type, whereas the one case we encountered (Case 1) was plasmacytoid/hyaline type [Figure 1a and b].

Cystic change seen in our case 2 is peculiar to the clear cell type of myoepithelioma as suggested by previous reports of cystic changes in two such tumors from the parotid and one from the hard palate.

To consider a histomorphological diagnosis of pure myoepithelioma, the epithelial component should be less

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Figure 1: (a) A fairly uniform plump spindle shaped tumor cells arranged in sheets, without pleomorphism and mitosis [H&E stain, ×100]. (b) Polygonal cells showing distinct border, eccentric nuclei, and hyaline eosinophilic cytoplasm (H&E stain, ×400). (c) Photomicrograph showing uniform strong positivity for S100 (IHC stain, ×400). (d) Photomicrograph showing focal strong positivity for smooth muscle actin (IHC stain, ×400)

Figure 2: (a) Normal salivary gland (arrow), separated from tumor by fibrocollagenous tissue. (H&E stain, ×40). (b) Tumor shows round to oval cells with clear cytoplasm and regular vesicular nuclei (H&E stain, ×400). (c) Photomicrograph showing diffuse strong positivity for S100 (IHC stain, ×100) (d) Photomicrograph showing diffuse positivity for smooth muscle actin (IHC stain, ×400)
Conflicting results have been obtained on immunostains of myoepitheliomas. The most consistent were the positive staining for cytokeratin, S100, and SMA, whereas vimentin and GFAP expressions vary. All our cases showed strong positive staining for S100 and SMA, except that the stain for SMA in the plasmacytoid variant was positive only focally. Ogawa et al. argue that the plasmacytoid variant lacks evidence of myoepithelial differentiation and hence should be classified as adenoma or plasmacytoid adenocarcinoma. However, others do not agree with this view and suggest that SMA expression may vary and may be negative at certain stage of differentiation of myoepithelial cells due to certain inhibitory effects of the extracellular matrix. The focal strong expression of SMA in our case of plasmacytoid myoepithelioma vindicate this stand.

The recommended management of myoepithelioma is surgical excision with a margin of uninvolved tissue around. A recurrence rate of 15%-18% is observed. Recurrence is possible with incomplete resection.

We conclude that rare type of salivary gland tumors, especially those arising in unusual locations with unfamiliar morphological patterns can cause difficulty in diagnosis. Awareness about these with their varied morphological patterns and diligent use of immunohistochemistry will help in their identification. This is important as most of them can be cured by adequate excision.

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