Oral medicine case book 75: Mucoepidermoid carcinoma of the lower lip: review and a case report

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ABSTRACT

Mucoepidermoid carcinoma (MEC) is the most common malignancy of salivary gland origin, comprising 30% of all these neoplasms. Major salivary glands are more commonly involved, with most MECs presenting in the parotid gland. The palate is the most commonly involved site in minor salivary glands. MEC of the lower lip is extremely rare with only a few cases reported in the literature. We report a case of a low grade MEC of the lower lip in a 49 year old female patient. She presented with a mass in the lower lip of six months duration with a history of a recent rapid increase in size. Surgical resection was the treatment of choice and histopathological examination of the excised mass confirmed a diagnosis of low grade MEC. We present this case to highlight that although relatively rare, MEC should be included in the differential diagnoses of lower lip masses.

INTRODUCTION

MEC is a malignant epithelial salivary gland neoplasm characterised by proliferation of epidermoid, mucous and intermediate cells in varying proportions.1-3 The majority of tumours arising from the minor salivary glands are malignant.4 The MEC was first described by Stewart et al in 1945 as a mucoepidermoid tumour categorised into “favourable” (benign) and “unfavourable” (malignant) variants.5,6 It later, however, became apparent that all variants of the neoplasm are malignant and capable of loco-regional and distant metastasis, hence the adoption and endorsement of the term “MEC” by the Second 2005 WHO classification of salivary gland neoplasms.1,2 MEC is reported to represent 12–29% of all salivary gland neoplasms.1-3 MEC is a malignant epithelial salivary gland neoplasm with more than 50% presenting at the junction between the hard and soft palate where minor salivary glands are most numerous.1,2,5,7-9 Other sites that may be involved include buccal mucosa, upper and lower lips, and the retromolar region. Minor salivary gland neoplasms account for 22% of all salivary gland malignancies and 12-30% of all salivary gland neoplasms.10-12 Although tumours of minor salivary glands are not as common as those involving major salivary glands, a biopsy is crucial when there is suspicion as more than 80% of minor salivary gland neoplasms are malignant in nature.5,9,11 Smoking, viral infections, working in rubber manufacturing and genetic susceptibility are some of the risk factors that have been reported in association with MEC.12 Prior exposure to radiation, however, is the only well established and most commonly associated risk factor for the development of MEC.13 Hiroshima and Nagasaki atomic bomb survivors and patients treated with radiation for various childhood malignancies, including lymphoma, sarcoma, retinoblastoma, and thyroid carcinoma, have an increased risk of developing MEC, with atomic bomb survivors reported to have an increased relative risk of 9.3 times.1,2,12,13 A recurring chromosomal translocation (t 11; 19) (q21; p13) resulting in the formation of the MECT1-MAML2 fusion oncogene has been identified in MECs.14-16 The translocation is demonstrated in 55-75% of low grade MECs and up to 46% of high grade MECs.10,17 MECT1-MAML2 fusion positive MECs are associated with improved survival. MEC gender predilection reported in literature has been cited as either non-existent or showing a slight female predisposition with a ratio of 3:1.1,2,5,7,9 Although MEC can occur at any age, the peak incidence is between the third and sixth decades of life.1,2,18 The MEC is the most common malignant salivary gland tumour in children and in adolescents under the age of 20.17 Palatal salivary gland tumours tend to present in young patients, 5-7 years younger than those involving other intraoral minor salivary glands, whereas MEC of the tongue tends to present in older patients.1-3 The MEC demonstrates diverse histomorphological features and biological behaviour, with prognosis correlated with histopathological grade, early detection and management.5,6,19 Treatment options for MEC include complete local excision for low grade MEC and surgical resection with or without radiation for high grade MECs.1,3

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ACRONYMS

AFIP: Armed Forces Institute of Pathology
MEC: Mucoepidermoid carcinoma

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CASE REPORT

A 49-year-old African female patient presented with a six month history of a painless, slow growing swelling on the lower lip. The patient reported a 34-year history of snuff dipping on the lower labial sulcus, with a medical history that was non-contributory. Extraoral examination revealed palpable, mobile, non-tender bilateral submandibular lymph nodes. On intra-oral examination, a solitary dome-shaped, smooth surfaced, non-ulcerated intramuscular swelling of normal colour was visible on the lower left labial mucosa (Figure 1).

The clinical differential diagnoses included mucocele, fibro-epithelial polyp, salivary gland neoplasm, lipoma and haemangioma. The lack of an associated history of trauma favoured the latter three options. The mass was surgically excised (Figure 2A). The specimen was multilobular; round to oval in shape with a yellowish cut surface, and measured 12 x 8 x 6 mm (Figure 2B).

Histopathological examination of the specimen showed features consistent with those of a mucoepidermoid carcinoma characterized by proliferation of neoplastic epidermoid, mucous and intermediate cells lying within a fibrous connective tissue stroma (Figure 3).

Cystic spaces lined by mucous cells and a few scattered basoloid and cuboidal intermediate cells were observed (Figure 4 A). On higher magnification of the tumour section, a focal area showing epidermoid, mucous and intermediate cells is observed (Figure 4B).

Figure 1: A solitary dome-shaped, smooth surfaced, non-ulcerated intramuscular swelling on the lower left labial mucosa.

Figure 2: A. Biopsy of multi-lobular mass. B. Specimen sent for histological assessment. C. Post-operative surgical site.

Histochemical examination demonstrated the presence of intra-cytoplasmic mucin in mucous cells and pools of mucin as highlighted by diastase resistant Periodic Acid Schiff (PAS) and mucicarmine positivity. The neoplastic cells were negative for S100 and SMA but positive for the P63 stain. The tumour had a mostly solid composition, and showed no signs of neural invasion, necrosis, mitotic division nor cellular anaplasia; thus giving it a total score of Two and a classification of a grade One/low grade MEC according to the Armed Forces Institute of Pathology grading system (AFIP).3,5 Positive tumour margins were identified and the patient was subsequently recalled three weeks later for wider excision margins. More than a year after the second surgical procedure the patient remains tumour free.

The correlation between the tumour size and the histological grade is in line with the observation made by Triantafillidou et al. who suggested low grade MECs showed a disposition for a diameter less than 4cm, circumscription, lack of a capsule and a predominantly cystic composition.18 In our case the MEC was less than 4cm in diameter, lacked circumscription but had a predominantly solid composition. By demonstrating a cystic component of less than 20%, our low grade MEC underscores the significant role played by factors other than the architectural growth pattern in the grading of MECs.

GRADING CRITERIA

The most popular grading systems for MECs are the Modified Healey, the AFIP and the Brandwein systems.2,3,7,8,10–23 All three grading systems classify MECs into three grades: low, intermediate and high grade. The Healey system places great emphasis on the morphological features of the tumour whereas the other two systems are point based and assess various histopathological parameters (Tables 1, 2 & 3).

The Brandwein grading system is a modified AFIP grading system with three additional histopathological features (Tables 2 & 3).21,22 Currently no consensus exists with regard to which of the three systems is the best. The AFIP grading system is reproducible and demonstrates relatively good correlation between histological grade and clinical behaviour1,2,5 but tends to downgrade MECs and to cluster intermediate and high grade tumours as reported by Brandwein et al.22 In an attempt to correct this, Brandwein et al. modified the AFIP grading system.
Regardless of these challenges, MECs which are largely dependent on the histological implications on treatment modalities and prognosis for intermediate tumours. These differences have significant parameters examined and scored to eight. Of these parameters necrosis, mitotic activity, perineural, lymphovascular and bony invasion are considered to be the main determinants of tumour grade. In both grading systems the individual scores are added and a grand total score is then used to classify MEC into low, intermediate, and high grades. However, Nance et al. demonstrated that the Brandwein system is not without fault as it has a tendency to upgrade tumours, by clustering low grade and intermediate tumours. These differences have significant implications on treatment modalities and prognosis for MECs which are largely dependent on the histological grade of MECs. Regardless of these challenges, numerous studies have proven histological grading based on these systems to be the most significant independent prognostic factor for MECs. An exception to the rule is the MEC of the submandibular gland which is associated with increased risk for metastasis and poor prognosis irrespective of the histological grade. In addition to the site of involvement, gender, age, clinical stage, disease-free margins and lymph node metastasis have proven to be of prognostic significance.

**DISCUSSION**

The most common malignant tumours to arise from minor salivary gland tissue include mucoepidermoid carcinomas (21.8%), adenocarcinomas (7.1%), adenoid cystic carcinomas (6.3%), and acinic cell carcinomas (1.6%). In the past, prior to proper description, MECs, particularly the high grade variants have been misdiagnosed as squamous cell carcinoma or unspecified adenocarcinoma. This is not surprising considering that high-grade tumours are predominantly solid, and show a great degree of cellular atypia similar to that seen in squamous cell carcinoma. The misdiagnoses and changes in the diagnostic and grading criteria of these salivary gland neoplasms also explain the conflicting data regarding the frequency of MECs reported in literature over the years, which on average was reported as 27% of all malignant salivary gland tumours before 1990, and 45% thereafter. Although there are a few studies reporting on the MEC of the lower lip, found in their study MECs, particularly the high grade variants have been misdiagnosed as squamous cell carcinoma or unspecified adenocarcinoma. The misdiagnoses and changes in the diagnostic and grading criteria of these salivary gland neoplasms also explain the conflicting data regarding the frequency of MECs reported in literature over the years, which on average was reported as 27% of all malignant salivary gland tumours before 1990, and 45% thereafter. Although there are a few studies reporting on the MEC of the lower lip, found in their study MECs, particularly the high grade variants have been misdiagnosed as squamous cell carcinoma or unspecified adenocarcinoma. The misdiagnoses and changes in the diagnostic and grading criteria of these salivary gland neoplasms also explain the conflicting data regarding the frequency of MECs reported in literature over the years, which on average was reported as 27% of all malignant salivary gland tumours before 1990, and 45% thereafter. Although there are a few studies reporting on the MEC of the lower lip, found in their study MECs, particularly the high grade variants have been misdiagnosed as squamous cell carcinoma or unspecified adenocarcinoma. The misdiagnoses and changes in the diagnostic and grading criteria of these salivary gland neoplasms also explain the conflicting data regarding the frequency of MECs reported in literature over the years, which on average was reported as 27% of all malignant salivary gland tumours before 1990, and 45% thereafter. Although there are a few studies reporting on the MEC of the lower lip, found in their study MECs, particularly the high grade variants have been misdiagnosed as squamous cell carcinoma or unspecified adenocarcinoma.

**CONCLUSION**

In conclusion, we present this case to highlight that, MEC, although relatively rare, should be included in the differential diagnoses of lower lip masses to guide the treatment approach, avoid recurrences and possible repeat surgery in order to achieve curative margins. Whilst...
it is unlikely that the MEC in this case was associated with the reported history of the use of smokeless tobacco (snuff dipping), given the ‘coincidental’ existence of the MEC on the site where the smokeless tobacco was habitually placed over the years, future studies should evaluate the possibility of this association.

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