Pleomorphic Adenoma in Ectopic Salivary Gland Tissue in the Neck

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ABSTRACT

A case of pleomorphic adenoma originating from ectopic salivary gland tissue (ESGT) of the upper neck is reported. A 34-year-old male patient was referred to our Department for a painless swelling in the right submandibular region. Preoperative evaluation (clinical examination, fine-needle aspiration cytology (FNAC) and imaging studies) was performed and the finding was that of a pleomorphic adenoma in ESGT. A modified «S» incision with extension to the submandibular region was performed and the tumour was extirpated. The histopathological report confirmed our initial diagnosis. No recurrence was obtained during a four-year follow-up period. Isolated neck mass may be overlooked as ectopic salivary gland tissue neoplasm (ESGTN). Proper preoperative assessment and optimal surgical treatment are the keys for successful management of these rare tumours. The distinction between metastatic lesion from a head and neck tumour and ESGTN may present considerable diagnostic problem. A review of the literature on ESGT and associated tumours with emphasis on clinical features, diagnosis and treatment is also presented.

Key words: ectopic salivary gland tissue, choristoma, pleomorphic adenoma, upper neck

Introduction

Salivary tissue neoplasms may originate from normal, accessory and heterotopic (ectopic) sites of salivary gland tissue1. Regular salivary tissue is organized in three major (parotid, submandibular and sublingual) and multiple minor salivary glands. Accessory parotid gland is salivary tissue that is located anteriorly and anatomically separated from the main parotid by Stensen’s duct2,3. Salivary tissue found in unusual locations is termed ectopic or heterotopic salivary tissue, as well as salivary tissue choristoma. In the head and neck region ectopic salivary gland tissue (ESGT) has been found in the middle ear, hypophysis, thyroglossal duct, mandible, tongue, gingiva, lymph nodes of the neck, thyroid gland, parathyroid glands and in the sternoclavicular joint4-10. The presence of ESGT is rare and neoplasms of this tissue are even rarer. In the neck it usually manifests as an asymptomatic lump, as a cyst or a draining sinus11. According to Willis12, there are three main hypotheses for salivary ectopia: abnormal persistence and development of vestigial structures, dislocation of a portion of definitive organ rudiment mass and further development along with abnormal differentiation of local tissues. In 1999, Perito et al.1 reviewed the literature on ESGTN of the head and neck and identified 111 cases. Ninety-three percent (104 of 111) of these neoplasms were found in the neck region. Neoplasms arising in the ectopic salivary tissue are mostly benign with Warthin’s tumour being the commonest histologic type, but several other benign and malignant tumours have been documented including pleomorphic adenoma, papillary cystadenoma, sebaceous lymphadenoma, oncocytoma, mucoepidermoid carcinoma and malignant oncocytoma13. We present a case of pleomorphic adenoma (mixed tumour) arising within salivary gland tissue in the neck. To the best of our knowledge, only 19 cases of pleomorphic adenoma in ESGT have been reported so far.
**Case Report**

A 34-year-old man was referred to the Department of Maxillofacial Surgery, University of Zagreb School of Medicine with a history of a painless swelling in the right submandibular area, just below the angle of the mandible. The neck lump was present for about five years ago and during that period the patient was symptomless. Clinically there was a palpable node posteriorly to the right submandibular gland, just below the angle of the mandible (Figure 1). The mass was 2.5 x 1.5 cm in size, non attached to the overlying skin, painless, freely movable and separated from the submandibular gland and the tail of the parotid. There were no other palpable lymph nodes in the neck. Ultrasound guided fine-needle aspiration cytology (FNAC) was performed and the finding was that of a pleomorphic adenoma. Additionally, computed tomography (CT) was done and it revealed a lesion of 23 mm in diameter not adherent to the submandibular nor the parotid gland (Figure 2). Intraoperative finding was identical to clinical and radiographic examinations (Figure 3). The tumour was extirpated and the pathohistologic review confirmed the cytologic report of a pleomorphic adenoma in ESGT (Figure 4). Postoperative course was uneventful, and function of the facial nerve remains intact. No recurrence was detected during the four-year follow-up period.

**Pathologic findings**

The well circumscribed, encapsulated oval mass was histologically revealed to be a tumour composed of uniform, mostly cuboidal epithelial cells forming tubuloductal structures, embedded within the chondromyxoid stroma. Remnants of un conspicuous salivary tissue overlies the fibrous capsule of the tumour (Figure 5).
lymph nodes have been proposed for the ectopic salivary tissue in the upper neck and heteroplasia of the ectodermal lining of the cervical sinus of His for the lower neck\(^\text{30}\). Clinically in the upper neck ESGT manifests as a slowly enlarging, painless and mobile mass, while lower cervical EST usually presents as a draining sinus\(^\text{21}\).

ESGTN is probably often overlooked in the differential diagnosis of a neck mass. After clinical exam and FNAC of the tumour, radiographic imaging (CT and ultrasound) should follow, because ESGTN should be considered a metastatic lesion until no primary tumour is identified. Daniel and McGuirt\(^\text{30}\) proposed an algorithm for evaluation and management of neck/periparotid masses, suggesting excision or parotidectomy alone for benign lesions and isolated low-grade malignant tumours, whereas high-grade malignant lesions require more extensive surgical treatment with possible irradiation which is consistent with reports of other authors\(^\text{11,18}\).

In our case we have treated our patient similarly, having a radiological staging performed with an ultrasound of the neck plus a FNAC of the tumour and CT head/neck after clinical evaluation. Since no primary tumour in major or minor salivary glands or metastatic nodes were found, the neck mass was classified as an ESGTN. The surgical approach included a modified «S» incision extended to the submandibular's gland which provided excellent exposure and facilitated extirpation of tumour mass.

To the best of our knowledge, excision for benign lesions has been reported as sufficient and we have treated our patient according to previously described guidelines\(^\text{11,18,30}\).

**Conclusion**

The evaluation and management of neck mass represents a complex task. The differential diagnosis of an isolated neck lump includes cysts of the neck, primary tumours (i.e. lymphoma), metastases from head and neck malignancies and extremely rare ESGTN. Recommended management, after proper preoperative evaluation (physical examination, FNAC and imaging studies), includes excision of the tumour with healthy margins and extended treatment (including neck dissection) with possible postoperative therapy depending on location, size and histological features of the tumour.

**REFERENCES**

PLEOMORFNI ADENOM EKTOPIČNOG TKIVA ŽLIJEZDE SLINOVNICE U VRATU

SAŽETAK

Prikazan je slučaj pleomorfognog adenoma ektopičnog tkiva žlijezde slinovnice u gornjem dijelu vrata. Tridesetgodišnji muškarac upućen je u našu Kliniku zbog bezbolne otekline desne submandibularne regije. Preoperativnom obradom ... tumor ektopičnog tkiva žlijezde slinovnice. Detaljna preoperativna obrada i optimalni kirurški zahvat ključ su uspješnog liječenja ovih rijetkih tumora. Značajan dijagnostički problem može predstavljati diferencijalna dijagnoza između metastaza tumora glave i vrata i tumora ektopičnog tkiva žlijezde slinovnice. Dat je pregled literature o ektopičnom tkivu žlijezde slinovnice i pridruženim tumorima s naglaskom na kliničke značajke, dijagnozu i liječenje.