Adenoid Cystic Carcinoma of the Submandibular Gland: A Case Report

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Abstract
Tumors of the salivary gland are comparatively rare, comprising less than 3% of all tumors. The distribution of tumors in the major salivary glands varies by site, with 85% to 90% occurring in the parotid gland, 10% to 15% occurring in the submandibular gland, and only 1% occurring in the sublingual gland. There is a higher incidence of malignancy in submandibular and sublingual tumors as compared with parotid tumors. Adenoid cystic carcinoma (ACC) is a malignancy of the seromucinous glands of the head and neck, accounting for 35% to 43% of malignant submandibular gland neoplasms. The majority of patients with ACC present with an asymptomatic mass that may have been present for months or even years prior to diagnosis. Adenoid cystic carcinoma is well known for its indolent pattern of recurrence and propensity for pulmonary metastases. Furthermore, it has a prolonged natural history and is slow growing even when there has been local recurrence and distant metastases. We report the case of a 46-year-old patient diagnosed with adenoid cystic carcinoma of the submaxillary gland, evolving for 5 years.

Keywords: Adenoid cystic carcinoma, submaxillary gland, surgery, radiotherapy, recurrence, metastases.

INTRODUCTION
Adenoid cystic carcinomas (ACC) or cylindromas were initially described by Robin and Laboublène in 1852 [1]. Adenoid cystic carcinomas are rare and usually develop at the expense of the salivary glands. They represent the main histologic type of accessory salivary gland tumors and account for approximately 10-20% of all salivary gland tumors [2]. ACC represent approximately 1% of ENT cancers, but can more rarely be found in all sites including secretory glands (breast, cervix, colon, prostate, etc.) [3]. Despite the non-aggressive histological character and the good short-term therapeutic results 4, the management of ENT cylindroma remains difficult because of their insidious growth, neurological tropism, and their metastatic potential. These associated elements delay the diagnosis which is often made when the tumor is locally advanced, and make the surgery difficult and late recurrence frequent. The treatment of these tumors has been long relied on exclusive surgery. Until the publications of Stewart et al., then of King et al. in the 1970s, these retrospective studies have shown the benefit of a treatment combining surgery and radiotherapy, in terms of local control and disease-free survival, compared to surgery or radiotherapy alone [5-8].

We report the case of a 46-year-old patient diagnosed with adenoid cystic carcinoma of the submaxillary gland, evolving for 5 years.

CASE REPORT
This is a 46-year-old patient, chronic tobacco user, who has presented for 5 years with right submandibular tumefaction, gradually increasing in size, painful, without local heat or redness. Clinical examination found a right submandibular tumefaction, approximately 7cm long, of firm consistency, painless on palpation. With an intraoral expression of the mass, without flow through the Wharton orifice, and without palpable cervical lymphadenopathies.

The cervical CT scan shows the presence in the right submaxillary gland, a hypodense tissue mass enhanced moderately after injection of contrast dye, with multiple tortuous vascular structures exceeding the...
mass and extending towards the parapharyngeal space and the right parotid region. This mass is well limited responsible for a thinning of the mandibular angle. Angio-MRI was also performed, shows a voluminous formation with regular contours, with hypointense signal in T1, and heterogeneous hyper signal in T2. Also the presence of bilateral cervical lymphadenopathies of which the most voluminous is in the left chain IIa measuring 11mm. A right submandibulectomy with homolateral angular mandibulectomy was realized. The aftermath of the surgery was simple. The final anatomic-pathologic study concluded adenoid cystic carcinoma of the submaxillary gland. A postoperative radiotherapy at a dose of 60 Gy was performed. The evolution was favorable with a follow-up of 24 months.

**DISCUSSION**

Three histological subtypes of ACC are known, as described above: the cribriform, tubular, and solid subtypes. Among the three subtypes, the solid subtype is the most aggressive [9, 14, 15]. Immunohistochemistry plays an important role in identifying the current case. CD117 was strongly expressed in this carcinoma. P63, S-100, and SMA are intensely stained in myoepithelial cells [16, 17]. In contrast, CK5 and CK7 are expressed in luminal cells [17].

The peak incidence of ACC among people is from 40 to 60 years of age [11, 15], which suggests that ACC is predominately a tumor of adulthood. Although a 1:1.3 male to female ratio was observed [11], another report demonstrated ACC with a slight female predominance, 3:2 male to female ratio [18]. The typical clinical symptom involves a slow-growing, firm, unilobular mass in the gland. An important associated symptom is painful swelling. Sometimes clinical symptoms are ignored so the tumor can grow rather large.

Surgical treatment of this lesion should therefore entail wide local excision of the submandibular space, including the submandibular gland, digastrics muscle, posterior mylohyoid muscle, lingual and marginal mandibular nerves, and the periosteum of the mandible. The hypoglossal nerve should be biopsied and resected if found to have perineural invasion on frozen section. Mandibular resection should be reserved for cases where the mandible is clinically or radiographically invaded. Neck dissections should be performed only in patients with known local lymph node involvement.

Most authors agree that ACC is radiosensitive but not radio-curative. However, several studies have indicated that combined therapy may offer improved local control [17, 20, 23, 24] and possibly prolonged survival [25]. Based on the study of Cohen and al [35], postoperative radiotherapy is recommended in the following situations: perineural invasion, positive surgical margins, T2 or larger tumors, and regional metastases. (Figure 3).

The recommendation of adjuvant radiotherapy often depends on putative prognostic indicators, which include stage, tumor site, microscopic or gross disease at the surgical margin, histological type, and perineural or perivascular invasion [20]. In previous reports, the dose of adjuvant radiotherapy ranged from 44 to 68.4 Gy, with a mean dose of 60 Gy [20, 21, 22]. Patients who received a dose greater than 60 Gy after surgery were reported to have significantly higher local control rates than those who received less than 60 Gy [23]. The total dose should be no less than 66 Gy. In cases with a positive margin, the dose should reach 66 to 70 Gy. It
seems that cisplatin, as well as high-dose melphalan, chemotherapy should cause a favorable result, but no statistically significant difference was confirmed in survival compared to placebo [24, 25].

DM is one of the most common features of ACC and usually occurs as a sign of the late course of disease26. Sung et al. [12] reported that the median DM interval was 48 months after the initial treatment in 46 ACC patients. The most common site of metastasis is lung15. However, ACC can also metastasize to other organs including bone, cerebrum, liver, thyroid, and spleen [27] Lung metastases usually progress slowly. On the contrary, bone metastases indicate a rapid progression28 [29].

Fig-3: TNM staging system

CONCLUSION

Adenoid cystic carcinoma of the submandibular gland is a slow-growing, insidious lesion with a prolonged natural history that includes a tendency toward local recurrence and distant metastasis. Early diagnosis is important, since advance tumor size is a poor prognostic factor. Wide surgical excision aimed at negative surgical margins offers the best chance for local control, and neck dissection should be reserved for those patients with clinically positive nodal metastases. Finally, postoperative radiation therapy should be strongly considered in those patients with perineural invasion, positive surgical margins, T2 or larger tumors, and/or regional metastases.

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