Case Report

Adenoid Cystic Carcinoma of the Lacrimal Gland in a 45 Years Old Male: Case Report and Review of Literature

Khan AH1, Ahmed N2, Shalike N3, Matin ABMA4, Amin FAA5, Barua S6, Kamal M7

Abstract:
Adenoid cystic carcinoma (ACC) is a rare malignant tumor that can manifests as proptosis in adult population. They account for 1.6% of all orbital tumors. Despite their rarity, they are the second most frequent epithelial neoplasms occurring in the lacrimal gland after pleomorphic adenomas. This kind of tumors are commonly occur in the salivary glands but can metastasize to lung, breast, brain and sinuses in hematogenous route. We describe a patient who presented with protrusion of right eyeball, developing over 8 years with history of intermittent watery discharge for 4 years. His magnetic resonance imaging showed a retrobulbarextraconal soft tissue lesion around the lacrimal fossa with invasion and erosion of the adjacent bone. The patient underwent right sided orbito-pterional craniotomy and gross total removal of tumor. Pathologic analysis showed neoplastic cells in a predominantly cribriform pattern with features of perineural invasion and diagnosed as a case adenoid cystic carcinoma of the lacrimal gland. We review the incidence, clinical features, radiographic and histopathologic features of these rare, aggressive malignancies along with current treatment options with reference to the relevant literatures.

Keywords: Adenoid cystic carcinoma, lacrimal gland, orbital tumor.

Introduction:
Adenoid cystic carcinomas of the lacrimal gland are rare, but most common malignant tumor of lacrimal gland origin with an estimated incidence of 0.073 per 100,000 individuals annually1. Patients may present with asymmetric facial pain or swelling, proptosis, ptosis, diplopia, visual disturbance in the form of dimness of vision and/or double vision2,3.
Magnetic resonance imaging (MRI) of brain and orbit is the preferred imaging modality for both the diagnosis and surgical planning. Adenoid cystic carcinomas are aggressive tumors having poor prognosis. Surgery is the primary modality of treatment followed by adjuvant chemo-radiation.

In our case report, we describe a patient who presented with right sided extra-axial painless non pulsatile proptosis with MRI findings of an irregular well circumscribed extra-conal soft tissue mass in the lacrimal fossa with globe deformation and surrounding bone erosion. Gross total removal of the tumor was achieved. Histopathology report confirmed the diagnosis of adenoid cystic carcinoma of the lacrimal gland. We review the incidence, clinical features, radiographic and histopathological features of these rare, aggressive malignancies along with current treatment options with reference to the relevant literatures.

**Case Report**

A 45-years-old man with no past medical history presented with gradual protrusion of his right eyeball for 8 years with intermittent excessive watering from the same eye for 4 years. He denied localized pain or numbness around his eye for preceding 7.5 years. But for last 6 months, he noticed orbital pain and a small mass at the upper eyelid. On examination, the patient had extra-axial proptosis of the right eyeball with mild congestion (Figure 1). There was a palpable, mildly tender soft tissue mass, having attachment with orbital rim. On examination, visual acuity was 6/18 in right eye and 6/6 in the left eye; visual fields of both eyes were intact. Ocular movements were full in all gazes. His pupils were equally round & reactive, and the fundoscopic examination was unremarkable. He had no palpable lymph nodes.

Computed tomography (CT) scan of the orbit showed an irregular, soft tissue mass in the area of lacrimal fossa, having calcification and erosion of orbital plate of frontal bone (Figure 2: A, B). MRI of brain and orbit revealed an heterogeneous enhancing, T1 iso- to hypo-intense and T2-hyperintense irregular soft tissue mass which was retrobulbar, extra-conal, associated with forward displacement of right eyeball, invasion of right lateral rectus muscle, and erosion of the adjacent orbital walls. The mass had extradural extension towards the anterior cranial fossa. It had also a positive dural tail sign (Figure 3: A, B).

The patient underwent right orbito-pterional craniotomy and gross total removal of tumor (Figure 4: A, B, C). Per-operatively the tumor was irregular, firm, greyish red in color with moderate vascularity. There was no clear demarcation between tumor and normal orbital content. Peri-orbita was eroded and tumor had extension into anterior cranial fossa. Based on the

![Fig.-1: Patient presented with right sided extra-axial proptosis.](image1)

![Fig.-2: CT scan of brain with contrast - sagittal (A) and coronal (B) sections showing a retrobulbar extra-conal lesion, causing displacement of optic nerve and erosion of orbital plate of frontal bone.](image2)
Fig.-3: MRI of brain with contrast- axial (A) and sagittal (B) sections showing T1WI iso- to hypo-intense retrobulbarextraconal lesion, having attachment with sclera and extension towards anterior cranial fossa. After giving contrast, there is presence ofdural tail sign.

Fig.-4 : Per operative photograph showing- single piece orbito-pterional bone flap (A), greyish red irregular tumor (B), attachment of the tumor with the eyeball (C).

Fig.-5: Adenoid cystic carcinoma showing small tumor cells and cystic spaces (H&E x400) (A), Adenoid cystic carcinoma showing perineural invasion (H&E x400) (B).
nature and extension, the tumor was classified as Stage - T4 according to American Joint Committee on Cancer (AJCC) classification. On histologic analysis there was small tumor cells and cystic spaces with perineural invasion (Figure 5: A, B). Post operatively, proptosis was aggravated with huge congestion and chemosis. However, at 8th POD these features were almost corrected. Visual acuity and status of extraocular muscle movements were same as pre-operative one. The patient was referred for radiation therapy.

Discussion:
Adenoid cystic carcinoma is slow growing tumor with aggressive clinical behavior. Though rare in overall incidence, it is the most common malignant lacrimal gland tumor; has a peak incidence in the fourth decade. Most common clinical presentation is facial pain or numbness following invasion to nerve and extraocular muscle. Distant metastasis through hematogenous route can be found in the lungs, liver, bone, brain, and kidney. Our patient presented at fifth decade with the features of painless, non-pulsatile proptosis with intermittent excessive watering (Figure 1). He had no feature of facial pain. The clinical course is 8 years before the diagnosis, which is longer in comparison to previously reported cases.

CT scan and MRI of brain and orbit with contrast are very useful to differentiate between benign and malignant lacrimal gland tumors to determine the extent of tumor invasion and surgical planning. On post-contrast sequence of CT scan of the orbits, benign lacrimal gland tumors are usually round, well-defined with no features of invasion, whereas adenoid cystic carcinomas may appear nodular, irregular, with bone erosion and calcification. However, MRI is the preferred modality to evaluate the nature of the lesion and relationship with surrounding neurovascular structure, especially for perineural spread and bone invasion. Adenoid cystic carcinomas typically appear isointense on T1WI and hyperintense on T2WI. After giving contrast, it takes heterogenous enhancement. In our patient, we did both CT scan and MRI for better delineation of tumor extent and surgical planning. On imaging, patient had a retrobulbar extraconal lesion at the area of right lacrimal fossa. Mass had attachment with posterior sclera with medial displacement of optic nerve. There was erosion of orbital plate of frontal bone and extradural extension to anterior cranial fossa through the bony gap (Figure 2: A, B; Figure 3: A, B). Based upon the imaging findings, we planned for orbito-pterional craniotomy for removal of tumor.

Histopathological analysis shows four patterns: cribriform, basaloid/solid, sclerosing, and tubular. Among them, cribriform growth is the most common pattern which is characterized by cystlike structures containing accumulations of basophilic, amorphous glycosaminoglycans or eosinophilic, hyalinized basal lamina. On the other hand, basaloid/solid growth is characterized by a predominance of basaloidmyoepithelial cells. But multiple patterns can be present in the same sample. Different growth patterns have significant impact on prognosis. For example, 5 years survival rate of patients with basaloid/solid patterns is 21% compared with that of 71% in patients with non-basaloid patterns. In our reported case, the growth pattern is cribriform pattern with an expectant better prognosis.

There are several treatment options for the management of this kind of aggressive tumor but the initial preferred modality should be surgical removal of tumor with or without orbitectomy. If gross total removal could not be achieved due to lack of cleavage plane between tumor and surrounding structures or if there is evidence of bone invasion, then radiotherapy is recommended for better outcome. If gross total removal could not be achieved, then patient may also benefit from neoadjuvant chemotherapy. We achieved gross total removal of tumor with removal of invaded roof and lateral wall of bony orbit. After that, we referred him for adjuvant radiotherapy as there was intracranial extension and bone erosion.

The American Joint Committee on Cancer (AJCC) classification defines T4 stage, if there is involvement of periosteum or bone. Patients with tumors ≥T3 at presentation exhibit shorter overall survival and time to metastasis. The recurrence rate of adenoid cystic carcinomas is up to 70% and worse prognosis is associated with neural invasion, basaloid growth pattern and tumor size >4 cm at the time of diagnosis. According to this, our reported case was staged as T4, which had perineural invasion. However, overall prognosis is poor in this aggressive malignant tumor.

Reference:


