Adenoid Cystic Carcinoma of Parotid Gland with Infratemporal Extension: A Rare Case Report

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Abstract
Adenoid cystic carcinoma (ACC) is a slowly developing malignant tumor of the salivary glands, occurring more commonly in minor salivary glands and rarely in parotid glands. It has the potential of retrograde perineural spread to the adjacent structures and spaces. Here we report a rare case of ACC of the parotid gland extended to the infratemporal fossa with perineural and vascular invasion, reflecting the advanced stage of the disease. After consulting with ENT specialists and radiation oncologists, palliative surgery was performed followed by adjuvant radio/chemo therapy.

Keywords: Adenoid cystic carcinoma, Infratemporal fossa, Parotid gland

Case Presentation
A 32-year-old man with chief complaints of limited mouth opening, right preauricular swelling, and pain over the past one year was referred to our Oral and Maxillofacial Surgery department of Beasat hospital. He did not have any history of cancer in the head and neck region or drug use.

The examination of the eyes and ears was normal with no evidence of intra-nasal mass or history of epistaxis. The tympanic membranes were intact bilaterally. Cranial nerves I to XII were unimpaired, with no deficiency in the muscles of mastication (VII). There was mild sensory loss in the V3 distribution. Facial mimetic muscles were intact and symmetrical. No evidence of crepitus, pain, or tenderness in temporomandibular Joint (TMJ) was found. Maximum mouth opening was limited (9 mm).

In clinical examination, there was an obvious swelling in the right preauricular area that distorted the facial contour. Palpation revealed a well-circumscribed mildly tender mass without any ulceration, pulse, or induration of the overlying soft tissue. Intraoral examination was normal and there was no intraoral extension of the lesion. The parotid papillae were noninflamed bilaterally, with clear saliva secretion from Stensen’s duct. No palpable lymphadenopathy in the submandibular or cervical areas was found.

Sonography of the right parotid and the angle of the mandible revealed a 14 × 30 mm hypoechoic area attached to the medial side of the parotid with extension to the depth of the mandible. Panoramic radiograph confirmed rarefaction in the right mandibular angle (Figure 1). In PNS CT scan with contrast, PNS and nasal mucosa...
were normal with no lesion. In the neck CT scan with contrast, there was a $26 \times 47$ mm soft tissue mass in the right infratemporal space with heterogeneous contrast enhancement and right parotid invasion. There was no evidence of bone erosion, airway obstruction, or vessel invasion (Figure 2). Skull base cerebellopontine angle (CPA) MRI with contrast demonstrated an ill-defined heterogeneously enhancing mass lesion with a cystic component measuring $43 \times 60$ mm in the deep portion of the right parotid gland. The lesion extended to the right masticator space and showed close contact with the skull base at the level of the foramen ovale. Internal acoustic meatus had normal width. The cranial nerves CN 6 to CN 9 were intact. No evidence of intracranial or lymph nodes involvement was observed. Brain MRI was normal (Figure 3).

Routine laboratory test results were within normal limits. Differential diagnoses included pleomorphic adenoma, ACC, schwannoma, lymphoma, juvenile nasopharyngeal angiofibroma, chondrosarcoma, fibrosarcoma, hemangioma, rhabdomyosarcoma, liposarcoma, meningioma, peripheral nerve sheath tumor, neurofibroma, and mucoepidermoid carcinoma.

Considering the advanced stage of the disease and possible malignant differential diagnoses, after consulting with ENT specialists and radiation oncologists, we decide to perform an incisional biopsy and tumor bulk reduction to alleviate trismus and pain and improve future radio/chemo therapy outcomes.

For surgical approach, we used Al-Kayat and Bramley incision to expose the maxillary arch. After zygomatic arch osteotomy, condylotomy with downward repositioning of masseter muscle was performed. Then, coronoidotomy and upward repositioning of the temporalis muscle provided access to the infratemporal lesion. The infratemporal part of the lesion was removed and sent for histopathological examination. Then, condylar reduction and fixation with L plate were performed. For maxillary arch reduction and fixation, we used the coronoid process as free bone graft (Figure 4).

The pathology report indicated low-grade cribriform and tubular variants of ACC with neural and vascular invasion.

**Discussion**
The term “adenoid cystic carcinoma” was introduced by...
Ewing in 1954 (4). The parotid gland ACCs occasionally cause a cranial nerve, particularly the facial nerve, to become paralyzed and manifest as a bulge or solid mass (9). Perineural invasion (22% to 46%) and numerous local recurrences are the major features of this neoplasm (10).

ACC has three histopathologic forms: cribriform, tubular, and solid. The most common pattern is cribriform (11). Any type may have noticeable perineural invasion (12).

Preoperative imaging is necessary for the accurate diagnosis. It mainly aids in determining the anatomical extension of the tumor. The modality of choice for bone invasion is CT scan, while MRI is useful in determining the type of the lesion and assessing its locoregional extent and the cervical lymph nodes and bone marrow infiltration. Considering the potential of retrograde peri-neuronal pathway and a caudal extension of the cervicothoracic passage, it is essential to take skull base imaging (13).

Surgery is the main form of treatment, but adjuvant radiotherapy plays a significant role in more advanced stages and in cases where there are positive margins. A total conservative or a radical parotidectomy is advocated for parotid ACCs though the main intent is to obtain a tumor-free area of at least 1 cm (14). Regional lymph
Adenoid cystic carcinoma in the infratemporal fossa


