Introduction

Malignancies of the temporal bone are rare with an incidence of less than 0.2% amongst all head neck cancers [1]. Amongst them, most are squamous cell carcinoma of the external auditory canal. Despite advances in surgical and radiotherapeutic techniques, prognosis for patients with this disease has remained poor [2,3]. The median age at presentation of squamous cell carcinoma (SCC) of the temporal bone is in the seventh decade. Occasionally, SCC of the temporal bone can develop in younger adults and the possibility should not be ignored merely on the basis of age [4]. We report a case of bilateral squamous cell carcinoma of both temporal bones in a young adult male patient who presented very late.

Case Report

30 years old male patient presented to ENT OPD with complaints of bilateral foul smelling, painful, occasionally blood mixed ear discharge for the past three years, painful swelling on the right side of the face for the past six months, and deviation of the angle of the mouth on the right side for the past two weeks. There was a history of the patient undergoing some ear surgery about three years prior, but no documents pertaining to the surgery were found. He was a farmer by profession, and there was no prior history of any exposure to radiation. On examination, there was a large swelling on the right side of the face, about 8cm x 5cm in size, which was hard, tender, fixed to the lower structures and skin, with the overlying skin inflamed, and ulcerated at places. The ear lobule was displaced laterally by the swelling which involved the right parotid. There was also right sided Lower Motor Neurone type facial palsy, with incomplete closure of the eyes on maximum effort (Figure 1a & 1b). Mouth opening was restricted to one and a half fingers. He had no other neurodeficit. On otoscopy and otoendoscopy, both ear canals were occluded by reddish proliferative tissue that was extremely vascular and bled on touch. A dirty white discharge was also found to be smeared around the reddish masses (Figure 2). He complained of decreased hearing in both the ears, but pure tone audiometry was impossible as mere placement of the headphones over the mastoid resulted in severe pain. Rest of the systemic examination did not reveal any abnormality.
The patient was admitted and started on intravenous antibiotics and analgesics. A high resolution CT scan of both temporal bones was ordered, which revealed a large heterogeneous soft tissue density behind right pinna, measuring about 37mm x 23mm in the maximum cross sectional diameter, which extended to involve the right parotid. Soft tissue densities were also noted in bilateral ear canals. There was a large post operative cavity on the right mastoid region. Bilateral inner ear structures were normal (Figure 3). To know the extent of soft tissue involvement, a contrast MRI of both temporal bone region along with neck was ordered. T2 weighted FSE & STIR images showed large heterogenous lesion with complete bony destruction of right middle ear, involving right ear canal, pinna, right parotid and extending to middle & posterior cranial fossa, with involvement of the neurovascular planes. Focal enhancing lesion was also seen in left ear confined to petrous bone, about 28mm x 24mm x 26mm in size. These pointed towards a neoplastic lesion (Figure 4).

Biopsy was taken from both the ears under general anaesthesia and the ears were packed with gauze packing to control the significant amount of bleeding. Histopathological studies revealed well differentiated infiltrating keratinising squamous cell carcinoma of both the ears, with the tumour cells arranged in solid nests and sheets with peritumoral chronic inflammatory cell infiltration (Figure 5).

Since the tumour on both sides did not have any interconnection between them, both were diagnosed as primaries. Using the University of Pittsburgh staging system for temporal bone tumours, the lesions on both the sides were independently staged. The right side was staged to Stage IV (T4N0M0) while the left side was staged to Stage II (T2N0M0).

Since such a large tumour required multimodality treatment, a tumour board was constituted. The lesion was deemed unresectable, so the patient was put up for palliative radiotherapy in a dosage of 30 Gy over 10 fractions. During the course of his treatment, he developed intractable pain, and his analgesics were stepped up gradually and ultimately, he had to be started on opioid analgesics (morphine tablets). Eye care for prevention of exposure keratitis was prescribed. Since prognosis was grave, the patient’s relatives were counselled about providing the best supportive care.

At the last follow up, patient had completed his course of palliative chemotherapy but his general condition was extremely poor. He is alive as of January 2018.

Discussion

Squamous carcinoma of the temporal bone is a rare but
aggressive tumour, usually having a poor prognosis. Survival rate decreases with increasing stage of the disease [7]. The main staging system used today, despite being flawed, is the revised Pittsburgh classification system [8,9]. Surgical resection of the tumour, with adjuvant radiotherapy for advanced stage disease appears to be the treatment of choice with the extent of surgery being a subject of debate [4,10]. Prasad et al. [11], in their meta-analysis on temporal bone carcinoma treatment, found that survival rates in case of advanced stage disease stayed poor, despite the extent of primary surgery.

Pensak et al. [12] considered that invasion of any of the following structures: cavernous sinus, carotid artery, infratemporal fossa, paraspinal musculature make a tumour unresectable. In the present case, the tumour was deemed unresectable due to spread to the infratemporal fossa and also had extensive involvement of the neurovascular planes. Since the patient already had very advanced disease in the right ear, there was no obvious benefit for operating on the left ear, and so surgery was not contemplated for the left ear although it was Stage II.

References