Bilateral warthins tumour—it runs in the family

Abstract

Background: A Warthins tumour or adenolymphoma is a benign cystic tumour of the salivary gland. They commonly occur in smokers, in the 6th or 7th decade of life and are treated by surgical excision. To date, no reports exist in the literature of inherited bilateral Warthins tumour in the literature.

Case report: We describe the case of a 67-year-old man who presented with bilateral Warthins tumour of the parotid gland. The family history showed that the patient’s brother had also been diagnosed with this and his father’s medical history was also suggestive of Warthins tumour. The patient underwent bilateral superficial parotidectomy and suffered no major long term complications.

Conclusion: Familial bilateral Warthins tumour is an extremely rare clinical presentation in ENT practice. To our knowledge, this is the first case report which describes coexisting bilateral and familial disease.

Keywords: Warthins tumour; Bilateral; Familial; Superficial parotidectomy

Introduction

Warthins tumours represent approximately 2-15% of all parotid tumours. Most tumours are unilateral, though bilateral disease is well described in the literature (approximately 10% cases) and is often exclusive to the parotid gland. Risk factors for the development of Warthins tumour include; cigarette smoking, genetic predisposition and environmental factors such as radiation, diet and viruses. The role of cigarette smoking in tumourigenesis is thought to be secondary to irritation of the ductal epithelium within the gland. Some evidence has also shown that several genetic alterations (point mutation, translocations of chromosomal units and allelic loss) have been linked to tumour initiation and progression. Recent studies have highlighted the specific role of the t (11:19) translocation and its associated oncoogene (CRTC1-MAML2) in mucoepidermoid carcinoma and Warthins tumour.

Despite Warthins tumour being more traditionally associated with men, the increasing incidence of females smoking may somewhat alter these findings in future studies. As with most head and neck pathology which presents with a lump, fine needle aspiration is the investigation of choice, showing a sensitivity of around 90% and a positive predictive value of 98%. Though malignant transformation is rare (approximately 0.3%), treatment mainly includes surgical excision of the superficial lobe of the parotid (assuming there no disease spread elsewhere within the parotid). Well documented complications of superficial parotidectomy include damage to the facial and great auricular nerve, hematoma, Frey’s syndrome and salivary fistula. Familial Warthins tumour is far less documented and rarer than bilateral disease. Within the literature only 3 sets of cases have described familial disease. To date, there has been no documentation of familial bilateral Warthins tumour of the parotid gland. Our case report is therefore the first description of this condition amongst the literature.

Case presentation

A 67-year-old male smoker presented to clinic with few month history of a smooth, well-defined 3cm swelling overlying the right parotid gland. There were no other clinical features on examination. It was revealed from the family history, that the patient’s brother was diagnosed and underwent bilateral surgical excision for bilateral Warthins tumours of the parotid a few years previous. The patient’s father also reported to have had a parotid lump, though due to lack of clinical information, no definite diagnosis could be commented on. Ultrasound guided fine needle aspiration revealed a diagnosis of Warthins tumour. The patient underwent a right superficial parotidectomy using a modified Blair incision and was successfully discharged from clinic after a few months with no complications (Figure 1).

Figure 1 Modified Blair incision overlying right parotid gland.

Five years later, the patient presented with a metachronous lesion overlying the left parotid gland. Ultrasound guided fine needle aspiration again, was suggestive of Warthins tumour. A T2-weighted MRI scan showed evidence of a 2.5cm well defined lesion within the left superficial parotid tissue (Figure 2). The patient underwent another superficial parotidectomy using the same surgical approach (Figure 3). Histopathological findings of the specimen were classical for Warthins tumour, showing papillary finger-like processes of columnar epithelium, eosinophilic uptake of dye and lymphoid aggregates with germinal centre formation (Figure 4). Following surgery, the patient suffered with transient Grade 2 left sided facial weakness (as per House Brakeman Classification) lasting approximately 3weeks and permanent numbness of the left ear lobe. The patient was subsequently discharged from clinic after 3months.
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In general, superficial parotidectomy is recommended for the treatment of Warthins tumour. Alternative treatment options include local surgical excision (e.g. lumpectomy), total or deep lobe parotidectomy (usually carried out if there is evidence of deep lobe involvement) or non-surgical conservative management. As mentioned previously, postoperative complications often include ear lobe numbness (especially when performing surgery with a modified Blair incision, as in our case), facial nerve weakness, Frey's syndrome, infection, haematoma, salivary fistula and recurrent disease. Both of the patients we describe developed transient facial nerve weakness secondary to neuropaxia, which has been documented in up to 19% of cases.

Cases of familial unilateral Warthins tumour in the literature include 3 brothers (between 44 and 48 years old) and 2 sets of mother and son (both of whom were in their 70’s and 50’s respectively). All of these cases, (including our case series), describes patient presentation within the 4th and 7th decade of life which is consistent with the literature. All patients underwent uncomplicated surgical excision confirming the diagnosis. Interestingly, one of the sons presented with a swelling of the left submandibular gland and subsequently underwent excision, confirming a diagnosis of Warthins tumour. All cases presented with salivary gland swellings, without any other associated features, as in our case series. The 3 brothers were known smokers, which supports evidence that cigarette smoke has a likely role in tumourigenesis. The potential genetic predisposition of Warthins tumour, via point mutation and genetic translocation (as mentioned previously), is also supported in all of the cases described, as each case series were direct relatives. Perhaps the limited literature describing familial cases of Warthins tumour highlights the need for further scientific research to help substantiate the evidence linking specific genes with the tumour and indeed our own findings.

Key messages

i. Inherited bilateral Warthins tumour of the parotid gland is extremely rare.

ii. Treatment of Warthins tumour of the parotid is commonly by way of superficial parotidectomy.

iii. Imaging is not a necessary pre-operative investigation in the UK, but may help in identifying synchronous or bilateral disease that is undetectable on clinical examination.

iv. Our case report shows evidence of inheritance for bilateral Warthins tumour, which may prompt a clinician to include family history as part of the initial clinical consultation.

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Conflict of interest

The author declares no conflict of interest.

References


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