Pharyngeal angiosarcoma following multimodal treatment for oropharyngeal squamous cell carcinoma

SL Ball, FNK Kwong, F Young, AK Robson

North Cumbria University Hospitals NHS Trust, UK

ABSTRACT

It is well established that angiosarcoma can develop following radiotherapy. We present an unusual case of angiosarcoma of the pharynx that developed three years after treatment with surgery and adjuvant chemoradiotherapy for a T2N2bM0 squamous cell carcinoma of the oropharynx. The patient was tumour free until developing dysphagia, which was found to be caused by an angiosarcoma. The patient underwent surgery of the pharyngeal angiosarcoma by laryngopharyngectomy, tongue base resection, selective neck dissection and radial forearm microvascular free flap reconstruction. Angiosarcoma following head and neck malignancy is rare but must be considered as part of the differential diagnosis in patients with new symptoms after radiotherapy.

KEYWORDS

Squamous cell carcinoma – Angiosarcoma – Pharynx – Radiotherapy – Chemotherapy

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CORRESPONDENCE TO

Stephen Ball, Department of ENT Surgery, Freeman Hospital, High Heaton, Newcastle upon Tyne NE7 7DN, UK
T: +44 (0)191 223 1086; E: s.l.ball@ncl.ac.uk

Angiosarcomas of the head and neck region are rare tumours. Radiotherapy and lymphoedema are well established risk factors for angiosarcoma.1 We present a case of angiosarcoma of the pharynx occurring approximately three years after curative multimodal treatment for a T2N2bM0 oropharyngeal squamous cell carcinoma with surgery and chemoradiotherapy. On reviewing the literature, we are not aware of any similar case of post-treatment pharyngeal angiosarcoma.

Case history

A 57-year-old man presented with a right neck lump in June 2008. Examination identified a 4cm cervical lymph node in level II and an enlarged right tonsil. Computed tomography (CT) confirmed this finding and excluded distant metastasis. Examination under anaesthesia revealed a mobile tumour confined clinically in the tonsil. Magnetic resonance imaging (MRI) suggested that the tumour did not extend beyond the tonsil but that level II neck nodes were involved. Histopathological analysis showed poorly differentiated squamous cell carcinoma (Fig 1).

The patient was treated by a laser excision of the tonsillar tumour with a right modified radical neck dissection. Histopathological analysis of the specimen confirmed a right tonsillar squamous cell carcinoma 24mm in maximum diameter, with 18mm depth of invasion and four positive level II neck nodes. There was a positive superior margin and close margins of 1mm anteriorly, posteriorly and laterally.

Following a multidisciplinary meeting, the patient received concurrent adjuvant chemoradiotherapy. A total dose of 60Gy in 30 fractions over 59 days was administered to the tonsillar region and right neck field together with docetaxel, cisplatin and fluorouracil chemotherapy. Following his multimodal treatment, he remained tumour free under combined clinical and radiological surveillance.

In March 2011 the patient presented with slowly progressive dysphagia for solids. Fibreoptic nasendoscopy identified a slightly oedematous supraglottis. CT and MRI did not reveal any signs of recurrence. Consequently, a panendoscopy and biopsy was performed (Fig 2). A striking feature at pharyngoscopy was that of very bland, ‘glistening’ mucosa involving multiple sites in the hypopharynx, supraglottic larynx, tongue base and posterior wall of the oropharynx. There was no bulky submucosal disease apparent. Histopathological analysis of multiple biopsy sites confirmed mucosal angiosarcoma (Fig 3).

Laser excision was unsuccessful in obtaining clearance of the tumour. Mapping biopsies demonstrated the tumour extending on to the anterior tongue, up the posterior pharyngeal wall to the level of the soft palate but not extending below the cricopharyngeus. Imaging showed no evidence of metastatic disease. Excision of the recurrent tumour was performed by total pharyngolaryngectomy, excision of
tongue base, left selective neck dissection and reconstruction with a tubed radial forearm microvascular free flap. Despite a wide resection, tumour clearance was not obtained and the patient was managed with best supportive care. He died 12 months following diagnosis of the pharyngeal angiosarcoma owing to complications of local disease progression and metastatic disease.

Discussion

We present an unusual post-treatment case of mucosal pharyngeal angiosarcoma. A particular feature was that the angiosarcoma appeared to involve all of the irradiated mucosa with a clear demarcation (at least inferiorly) between normal and abnormal mucosa. Gradual onset of dysphagia was the first presenting symptom, with initial examination and imaging consistent with post-chemoradiotherapy changes. Intractable dysphagia following chemoradiotherapy for head and neck cancer is a well recognised complication of treatment. Recurrent squamous cell carcinoma or treatment induced fibrosis. The implications for this in our practice may necessitate a lower threshold for examination under anaesthesia and biopsy following treatment.

Conclusions

Although this is a rare tumour, the lesson we have learnt is that dysphagia following treatment is not always due to recurrent squamous cell carcinoma or treatment induced fibrosis. The implications for this in our practice may necessitate a lower threshold for examination under anaesthesia and biopsy following treatment.

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