CASE REPORT

Synovial sarcoma of the hypopharynx – a case report and literature review

Sarcoma sinovial da hipofaringe – relato de caso e revisão da literatura

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Introduction

Synovial sarcoma of the head and neck is very rare, and sarcoma of the larynx and hypopharynx rarer still. As such, correct diagnosis and decision-making regarding surgery and adjuvant therapies pose clinical challenges to the otolaryngologist. We present a case of a synovial sarcoma arising from the hypopharynx, and a review of the literature.

Case report

A 77-year-old woman presented with intermittent hoarse voice and episodes of choking when swallowing. Panendoscopy showed a smooth, cystic mass arising from the right posterior aryepiglottic fold (Fig. 1). Initial biopsy showed only mild mucosal dysplasia. However, an urgent Computed Tomography (CT) scan was reported as showing a malignant-looking mass arising from the right superior thyroid pole. An ultrasound-guided biopsy showed spindle cells; differential diagnoses were a spindle cell variant of anaplastic thyroid carcinoma or Reidel’s thyroiditis, but the sample was too small for definitive diagnosis.

The case was reviewed at the head and neck oncology MDT, consisting of two Radiology consultants, two Oncology consultants and four head and neck surgeons, as well as...
speech therapy and cancer nurse specialists. For this case we also invited input from the thyroid surgeon. It was felt that the quality of the first CT made it difficult to establish the origin of the mass; therefore the patient underwent a repeat CT. In these images, a visible tissue plane separated the mass and the right thyroid lobe (Fig. 2), indicating that the mass was in fact hypopharyngeal in origin. A further incision biopsy was performed under local anaesthetic and sent to the regional sarcoma panel. Fluorescence In Situ Hybridisation (FISH) analysis signal pattern was consistent with SS-18 gene rearrangement. The diagnosis was a synovial sarcoma of the hypopharynx.

Our patient had a background of idiopathic pulmonary fibrosis, with progressive dyspnoea on minimal exertion. She was offered curative surgery; however she repeatedly declined any therapy. She explained that she wanted to spend what time she had with her family, preserving the quality of her life as long as possible. With support from the palliative care team, she died at home, nine months after the confirmation of her diagnosis.

**Table 1** Selected reports of synovial sarcoma in the pharynx or larynx.

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Age</th>
<th>Presentation</th>
<th>Site</th>
<th>Treatment</th>
<th>Follow up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jernstrom</td>
<td>1954</td>
<td>21/M</td>
<td>Dyspnoea, dysphagia, neck lump</td>
<td>Hypopharynx</td>
<td>Died post laryngoscopy</td>
<td>N/a</td>
</tr>
<tr>
<td>Miller</td>
<td>1974</td>
<td>23/F</td>
<td>Dyspnoea, dysphonia</td>
<td>Intra-arytenoid</td>
<td>Total laryngectomy</td>
<td>Disease-free 2 years</td>
</tr>
<tr>
<td>Gatti</td>
<td>1975</td>
<td>23/F</td>
<td>Acute stridor</td>
<td>Intra-arytenoid</td>
<td>Supraglottic laryngectomy</td>
<td>Disease-free 2 years</td>
</tr>
<tr>
<td>Gatti</td>
<td>1975</td>
<td>27/M</td>
<td>Dysphagia &amp; dysphonia</td>
<td>Laryngopharyngeal</td>
<td>Partial laryngopharyngectomy</td>
<td>1 year – pulmonary metastases, Ch-RTX, died after 2.5 years</td>
</tr>
<tr>
<td>Fernandez-Acenero</td>
<td>2009</td>
<td>12/M</td>
<td>Not documented</td>
<td>Supraglottic</td>
<td>WLE + ChTX</td>
<td>Disease-free 4 months</td>
</tr>
<tr>
<td>Bao</td>
<td>2013</td>
<td>37/M</td>
<td>Sore throat, haemoptysis</td>
<td>Aryepiglottic fold</td>
<td>Partial laryngectomy + Ch-RTX</td>
<td>Died distant metastases 41 months</td>
</tr>
<tr>
<td>Yang</td>
<td>2013</td>
<td>44/M</td>
<td>Dysphagia, dyspnoea</td>
<td>Hypopharynx</td>
<td>Total laryngectomy, pharyngoesophagectomy, neck dissection</td>
<td>Local recurrence 5 months – RTX, disease – free 2 years</td>
</tr>
<tr>
<td>Current case</td>
<td>2013</td>
<td>77/F</td>
<td>Dysphonia + dysphagia</td>
<td>Arytenoid</td>
<td>Refused therapy</td>
<td>Died at 22 months</td>
</tr>
</tbody>
</table>

RTX, radiotherapy; ChTX, chemotherapy; Ch-RTX, chemoradiotherapy; WLE, wide local excision.
Discussion

Synovial Sarcoma (SS) is a distinct malignant tumour of non-epithelial soft tissues. The name is a historical misnomer, as it is now recognised that they do not arise from joint synovia. It is most commonly seen as a slow-growing tumour in the extremities of adolescents and young adults; it is estimated that of approximately 200 presentations of SS per year in the U.K., 90% are in the extremities. They metastasise primarily haematogenously, to lung and bone. SS of the head and neck was first reported in the pharynx in 1954, but it remains a rare and poorly understood diagnosis.

On review of the English-language literature, we found 31 cases of laryngeal or hypopharyngeal SS (Table 1 shows a selection of reported cases). Median age was 21 years (range 11–79), with 72% male preponderance. The most common presenting symptoms were dysphagia (58%) followed by dysphonia (54%), similar presentations to other laryngopharyngeal malignancies.

In the reported cases, imaging did not identify any consistent radiological features. In a review of the MRIs of 6 Head & Neck SS cases, Hirsch et al. reported the lesions were most commonly isointense to grey matter on T1 MRI, and may also be heterogeneous, haemorrhagic and multi-loculated. However Sigal et al. (3 cases) and Rangheard et al. (8 cases) did not identify consistent radiological features. In the reviews and the case reports, the tumour was usually reported to be well-circumscribed, unlike other malignancies. It is worth noting that in several of the cases, including ours, the clinical appearance of the tumours was also well-circumscribed and smooth, suggesting benign pathology.

The diagnosis in SS cases therefore rests heavily on the histology. The core biopsy fragments showed spindle cells; thin, elongated cells with scant cytoplasm. They are a non-specific finding, present in inflammatory tissue, benign masses and various neoplasms. SS may be “monophasic”, consisting solely of these spindle cells, or “biphasic”, consisting of spindle cells and epithelial elements. Immunohistology shows epithelial markers including cytokeratin and epithelial membrane antigen among others, and gene analysis yields t(X;18;p11;q11) translocation in over 90% of SS.

Prognosis and management of these tumours has a limited evidence base. Of the 22 cases with a year or more of follow-up, 82% were disease-free at one year. The cases followed for 3 years, 67% were disease-free, 25% had died and one patient had a recurrence. 46% of the cases received primary surgery alone, 46% received surgery and adjuvant radiotherapy, and 7% received primary chemo-radiotherapy. Differing lengths of follow-up prevent meaningful comparisons of patient outcomes from the different therapeutic strategies.

Conclusion

Synovial sarcoma of the head and neck is a rare but significant pathology. The lack of specific clinical and radiological features, and the smooth well-circumscribed lesions, can delay diagnosis and even recognition of their malignant nature. This case highlighted the importance of multi-disciplinary collaboration, and of early suspicion of sarcoma. This case, with those previously reported; serve to further clinical recognition of these rare but devastating tumours.

Conflicts of interest

The authors declare no conflicts of interest.

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References