CASE REPORT

A primary tumour of the oesophagus with both melanocytic and schwannian differentiation. Melanocytic schwannoma or malignant melanoma?

R M Brown, S J Darnton, L Papadaki, G N Antonakopoulos, J Newman

A 76 year old white woman presented with a four month history of dysphagia and weight loss. Clinical, radiological, and endoscopic examination revealed a pigmented mass in the lower third of the oesophagus. The preoperative diagnosis, including biopsy examination, was that of malignant melanoma. Following oesophageal resection, the mass was found to be a localised, relatively superficial tumour with light, electron microscopic, and immunohistochemical features common to both Schwann cells and melanocytes. The patient survived 46 months after surgery and died of a stroke, with no evidence of tumour recurrence. The tumour is presented as a case of melanocytic schwannoma, with unique features when compared with oesophageal melanotic schwannomas and malignant melanomas described in the literature. The differential diagnosis is discussed and an origin from a common precursor cell of neural crest origin is postulated.

Malignant melanoma is a rare aggressive tumour, accounting for 0.2% of all oesophageal tumours. Less than 160 such tumours have been documented.1 Schwannomas of the oesophagus are even less common, with only 11 cases having been described previously,2-5 of which only three were melanotic.3 Melanotic schwannoma is most often associated with spinal nerve roots, but has been described at numerous other sites and as a part of Carney’s syndrome, which includes psammomatous melanotic schwannomas, myxomas, spotty pigmentation, and endocrine overactivity. It must be distinguished from classic schwannoma because its behaviour is unpredictable.3,4 Complete resection of oesophageal melanotic tumours is recommended to achieve a chance of survival.1

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We describe a tumour diagnosed preoperatively as a malignant melanoma. Ultrastructural examination of the resected specimen allowed revision of the diagnosis to that of a melanocytic schwannoma.

CASE REPORT

The patient, a 76 year old white woman, presented with a four month history of dysphagia and weight loss. Endoscopy showed a pigmented polyoid mass in the lower third of the oesophagus. A preoperative histological diagnosis of malignant melanoma was made. She had no cutaneous lesions and there was no radiological evidence of advanced local infiltration. Neither she, nor any family member, had Carney’s syndrome or neurofibromatosis. Subtotal oesophagectomy was performed. She had long standing cardiovascular disease and the postoperative period was complicated but the patient left hospital 27 days after surgery. She died 46 months after surgery, of stroke, with no evidence of tumour recurrence.

On resection the smooth, black, polyoid lesion measured 5 × 4 × 2 cm. Two centimetres proximal to this was a separate, flat, pigmented area measuring 2.5 × 2 cm. Excision was macroscopically complete.

Light microscopy showed that the superficial lesion was covered by attenuated squamous epithelium. It was nonencapsulated but had a circumscribed margin. The tumour was positive for CD34, epithelial membrane antigen, smooth muscle antigen, and desmin. Intratumoral macrophages were also melanin laden. Mucosa from the separate area showed nested junctional melanocytes without cytological atypia, and melanin in macrophages in the subepithelial stroma. Regional lymph nodes showed reactive changes only. Tumour staging was T1N0M0.
Electron microscopy showed that the tumour cells had deeply indented nuclei and long, elaborate, interlocking cytoplasmic processes, which also interdigitated laterally with neighbouring cells and formed intercellular junctions (fig 2). Melanosomes in varying stages of maturation, from the typical internal lamellar structure to dense pigmentation, occurred in most cells (fig 3A), a spectrum suggesting melanin synthesis.

An incomplete external lamina surrounded groups of closely apposed cells and long spacing collagen was sometimes seen in extracellular spaces (fig 3B).

**DISCUSSION**

The preoperative differential diagnosis between malignant melanoma and melanotic schwannoma is difficult because the appearances in small biopsies cannot be distinguished by light microscopy. Diagnosis is often not made until after examination of the resected specimen and may, as in this case, require ultrastructural confirmation.

Our patient fell within the age span expected for melanoma (fifth decade onwards). Melanotic schwannoma is a tumour of young adults, the mean age of those occurring outside Carney’s syndrome being 33 years. The mode of presentation and macroscopic appearance of our case would fit with a diagnosis of either melanoma or schwannoma.

Primary melanomas of the oesophagus are as aggressive as melanomas at other sites, with widespread metastasis and poor prognosis (five year survival is only 5.7%). Although exceptions exist, the long term survival of our patient argues against a diagnosis of melanoma. The prognosis for melanotic schwannoma is impossible to predict from pathological features, but outside the spinal cord it tends to follow an indolent course.

Immunohistochemistry did not permit distinction between the differential diagnoses but our case has the light microscopic features of a melanotic schwannoma: circumscript growth without encapsulation, high cellularity, a predominance of spindle cells, which are heavily pigmented and arranged in whorls or palisades, and a lack of cytological atypia and mitoses. However, there were no psammoma bodies or adipose tissue, as described by Assor and by Carney, the latter of whom puts great emphasis on the finding of psammomas. Melanoma, although renowned as a pathological mimic, usually displays obvious signs of malignancy, with cellular anaplasia and necrosis.

A confounding feature in our case is the presence of a junctional proliferation of melanocytes in nearby mucosa. Coexisting melanosis of the oesophagus is present in continuity with about 25% of cases of primary malignant melanoma and is used as a criterion in the differentiation between primary and secondary melanomas. However, in our case the junctional proliferation was clearly separate from the tumour and was not cytologically atypical.

“A confounding feature in our case is the presence of a junctional proliferation of melanocytes in nearby mucosa”

It seems reasonable to assume that this melanogenic primary oesophageal tumour has arisen from a cell of neural crest origin. The neural crest gives rise to a range of cells, including melanocytes and Schwann cells. During embryogenesis, melanocytes migrate from the neural crest to other sites, including the oesophagus. Cells of neural crest origin retain the capability in culture to differentiate along alternative lines, and it seems reasonable to assume that they may do so in vivo. Benign melanotic schwannoma and malignant melanoma may represent two ends of a spectrum of tumours arising from a common precursor cell of neural crest origin. In those cases of melanotic schwannoma that metastasise, it is interesting that the metastases have features more in common with melanoma than with the primary tumour.

Light microscopy indicated that the reported tumour has bland rather than aggressive features, favouring the diagnosis
Take home messages

- In this patient the preoperative diagnosis, including biopsy examination, was that of malignant melanoma.
- However, after surgery the tumour was seen to have bland rather than aggressive features, and ultrastructural examination confirmed that the neoplastic cells have characteristics of both Schwann cells and melanocytes, thus allowing a firm diagnosis of melanocytic schwannoma.
- This tumour is a unique case of primary melanocytic schwannoma of the oesophagus, occurring without psammoma bodies and outside Carney’s syndrome.
- An origin from a common precursor cell of neural crest origin is proposed.

of melanotic schwannoma over malignant melanoma. However, only ultrastructural examination proves that the neoplastic cells have characteristics of both Schwann cells and melanocytes, thus allowing the precise diagnosis. The tumour is a unique case of primary melanocytic schwannoma of the oesophagus, occurring without psammoma bodies and outside Carney’s syndrome.

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REFERENCES


PostScript

CALENDAR OF EVENTS

Full details of events to be included should be sent to Maggie Butler, Technical Editor JCP, The Cedars, 36 Queen Street, Castle Hedingham, Essex CO9 3HA, UK, email: maggie.butler2@btopenworld.com

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22–26 April 2002, Hammersmith Hospital (Imperial College Faculty of Medicine), London, UK
Further details: Wolfson Conference Centre, Hammersmith Hospital, Du Cane Road, London W12 0NN, UK. (Tel: +44 20 8383 3117/3227/3245; Fax: +44 10 8383 2428; email wcc@ic.ac.uk)

UK NEQAS for Blood Coagulation Annual Meeting

18, 19 June 2002, Conference Centre, Sheffield Hallam University, Sheffield, UK
Further details: TAL Woods, UK NEQAS for Blood Coagulation, Rutledge Mews, 3 Southbourne Road, Sheffield S10 2QJ, UK. (Tel: +44 114 267 3300; Fax: +44 114 267 3309; email neqas@coagusa.demon.co.uk)

Short Course on the Autopsy

25–28 June 2002, Department of Pathology, University of Sheffield, Sheffield, UK

Further details: Mrs S Clary, Department of Pathology, University of Sheffield Medical School, Beech Hill Road, Sheffield S10 2RX, UK. (Tel: +44 0114 271 2501; Fax: +44 0114 278 0059; email s.clary@shef.ac.uk)

Diagnostic Histopathology

8–19 July 2002, Department of Pathology, University of Sheffield, Sheffield, UK
Further details: Mrs S Clary, Department of Pathology, University of Sheffield Medical School, Beech Hill Road, Sheffield S10 2RX, UK. (Tel: +44 0114 271 2501; Fax: +44 0114 278 0059; email s.clary@shef.ac.uk)

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