Superficial epithelioid schwannoma presenting as a subcutaneous upper eyelid mass

Z Butt, J W Ironside

We describe a case of superficial epithelioid schwannoma presenting as a mass in the superotemporal quadrant of the orbit. This was treated by local excision with no evidence of recurrence or metastases on follow up. To the best of our knowledge, this is the first case report of the tumour in this particular area.

Case report
A 55-year-old woman presented to our casualty department in 1989 with a 2 year history of an asymptomatic enlarging mass along the temporal aspect of her left upper eyelid. There was no significant medical or ocular history and the patient displayed none of the stigmata of neurofibromatosis.

On examination, corrected visual acuity was 6/6,N5 right eye and 6/9,N5 left eye. Pupils were reactive to light, she had a full range of extraocular movements and there was no proptosis. Her discs were normal. She had a 1 cm diameter smooth firm mobile mass along the superotemporal quadrant of her orbit. Posterior
extension of the mass was suspected and a skull x-ray and computed tomography (CT) scan of her orbits were requested. The CT scan showed the features of a well defined soft tissue lesion with no associated extension into retroconal tissues or bony destruction. A chest x-ray, full blood count, urea and electrolytes, liver function tests, and erythrocyte sedimentation rate were all normal. She was referred to a consultant neurosurgeon for elective excision of her mass. This was excised completely and sent for histopathological analysis.

HISTOPATHOLOGY

The specimen was an apparently encapsulated smooth surfaced nodule measuring 8×5×4 mm. The lesion had been incised before receipt, demonstrating a solid grey/white cut surface. Histological examination showed a neoplasm with a composite pattern comprising spindle cells arranged in irregular bands and a population of larger epithelioid cells in solid sheets, with occasional large multinucleate cells (Fig 1). An extensive reticulin framework was present, with a predominantly pericellular distribution (Fig 1). The epithelioid cells contained abundant cytoplasm with well defined cell boundaries (Fig 2). Numerous thin walled capillaries were present throughout the tumour, but there was no sign of endothelial hyperplasia or haemorrhage. Occasional mitotic figures were identified (less than one mitotic figure per 10 high power fields). Although the tumour appeared encapsulated on naked eye inspection, on microscopy the tumour boundary was irregular with infiltration of the surrounding fibrous pseudocapsule.

Stains for neutral and acidic mucins and melanin were negative. Immunocytochemistry showed a positive reaction for S100 protein in polygonal and epithelioid cells (Fig 2B). Occasional large epithelioid cells also gave a positive reaction using an antibody to epithelial membrane antigen, but immunocytochemistry for cytokeratins was negative. Many of the tumour cells and vascular endothelial cells gave a positive staining reaction for vimentin, but no reaction was observed on immunocytochemistry for glial fibrillary acid protein, neurofilament protein, desmin, myoglobin, and the leucocyte common antigen. Electron microscopy showed irregular
polygonal cells surrounded by an abundant basement membrane. A few poorly formed intercellular junctions were noted, but no true desmosomes were identified. No premelano- somes, keratin filaments, muscle filaments, or neurosecretory granules were present in the tumour cell cytoplasm.

Comment
Epithelioid schwannoma is a rare variant of peripheral nerve sheath tumour and usually presents as an asymptomatic mass either in superficial (dermis or subcutaneous) tissue or deep soft tissue of the extremity. Superficial epithelioid schwannoma in the head and neck region therefore represents an uncommon variant of a rare tumour. The majority of superficial lesions are curable if wide local excision is accomplished expeditiously. Those in deep tissues, however, tend to be highly malignant and should be treated aggressively with possible adjuvant therapy.

The histological appearances of epithelioid schwannomas are variable, and in the past may have been confused with melanoma or metastatic carcinoma. The presence of epithelioid cells in a vague nodular pattern (as in this case) is characteristic. The immunocytochemical and ultrastructural features in this case are also characteristic and enable a distinction from metastatic carcinoma and melanoma. Epithelioid schwannomas are uncommon tumours; previous literature reports indicate that these are fully malignant neoplasms and warrant aggressive treatment. Mitotic activity and other histological features indicating malignancy are not uniformly present; the histological appearance cannot therefore be relied upon as a predictor of biological behaviour.

Our case has behaved in a benign fashion with complete local excision and no evidence of recurrence or metastasis on a 2 year follow up. Accurate histological diagnosis and early complete wide excision of superficial epithelioid schwannoma should be the objectives in managing this tumour.


Bilateral abducens nerve lesions in unilateral type 3 Duane’s retraction syndrome

Mark Mulhern, Catherine Keohane, Gerard O’Connor

Duane’s retraction syndrome (DRS) is a congeni- tal ocular motility disorder of neurogenic origin but of unknown aetiology. Clinically there is deficiency or loss of abduction and adduction and there may also be palpebral fissure diminution and globe retraction on attempted adduction.

Upshooting or downshooting of the globe may occur when adduction is attempted. DRS was first described in 1887 and early reports sug- gested a myogenic disorder; however, absence of the abducens nerve and nucleus with innervation of the affected lateral rectus muscle by the