Primary biphasic synovial sarcoma of the orbit

N Ratnatunga, J R Goodlad, N Sankarakumaran, R Seimon, S Nagendran,
C D M Fletcher

Abstract
Synovial sarcoma is one of the most common soft tissue malignancies of adolescents and young adults. Despite its name, it is no longer thought to be histogenetically derived from the synovium. What seems to be the first case of synovial sarcoma to arise in the orbit presented in a 21 year old woman as a slowly enlarging subconjunctival mass. Although this tumour was typically biphasic, the monophasic spindle cell variant arising at this site could easily be confused with less aggressive orbital connective tissue neoplasms.

Soft tissue tumours of the orbit, particularly sarcomas, are rare, with the notable exception of rhabdomyosarcoma (most often of the embryonal type) in infants and young children. In adults, the most common lesions are benign fibrous histiocytoma and haemangioendothelioma. Synovial sarcoma represents one of the most common soft tissue malignancies in adolescents and young adults and predominates in the leg, most often arising in the thigh or around the knee. Less than 10% of cases arise in the head and neck region, occurring most often in a paravertebral location. But the existence of these tumours, which include lesions of the oral cavity and cheek, combined with comparable lesions arising in such sites as the anterior abdominal wall and retroperitoneum, provided one of the earliest pieces of evidence that synovial sarcoma is not histogenetically derived from synovium. In fact, histogenesis is no longer regarded as tenable in malignant mesenchymal tumours generally. To our knowledge, no such tumour has been reported as arising in the orbit.

Case history
A 21 year old Sri Lankan woman presented with a five year history of a progressively enlarging, painless mass in the left orbit which had resulted in lateral displacement of the eyeball. Examination showed the presence of a 2 cm subconjunctival tumour on the medial side, associated with limited adduction of the left eye. There was no proptosis and vision was normal. At surgery the tumour was adherent to the epimysium of the medial rectus muscle and extended posteriorly along the medial orbital wall into the retrobulbar region. In the absence of a specific diagnosis an incomplete excision was performed (leaving a 7 mm portion of tumour on the posterior aspect of the globe) and the eye was preserved. The patient has since refused further treatment. Following the definitive histological report a thorough clinical and radiological search has been made for a primary lesion elsewhere in the head and neck, trunk, and limbs with negative results.

Pathological findings
The specimen consisted of a lobulated mass of soft pale tissue measuring 3 x 1.5 x 1.5 cm with no normal surrounding tissue. Histologically it was composed of two elements. The predominant component consisted of interlacing fascicles of spindle cells with palely eosinophilic or amphophilic cytoplasm and tapering vesicular nuclei with an indistinct nucleolus (fig 1). The cells showed up to 12 mitoses per 10 high power fields (1 hpf = 0.159 mm²) and were set in a predominantly myxoid but focally hyaline stroma. Within spindle cell areas there was a branching, thin-walled vascular pattern, reminiscent of haemangioendothelioma. The other epithelioid component comprised two patterns, consisting either of solid circumcribed nests of plump epithelioid cells with

Figure 1 The spindle cell component is not easily distinguishable from a fibrosarcoma or malignant peripheral nerve sheath tumour.
vesicular nuclei and more eosinophilic cytoplasm, or of well formed glandular spaces arranged in small clusters (fig 2). Some of these spaces contained diastase or periodic acid Schiff positive material but most showed central hyalinisation. Both types of epithelioid area were clearly delineated by reticulin staining. In both components of the tumour, but most notably in glandular areas, there was a striking infiltrate of mast cells. At the periphery of the biopsy specimen, tumour irregularly infiltrated the medial rectus muscle.

Immunohistochemically, using the avidin-biotin complex method, both solid and glandular epithelioid elements expressed epithelial membrane antigen (EMA) (Dako) and pan-keratin (DPC Ltd) and, in fact, these antibodies showed more extensive epithelial areas than were apparent in haematoxylin and eosin stained sections (fig 3). A few spindle cells also expressed EMA but were keratin negative. Stains for desmin, smooth muscle actin, S-100 protein and carcinoembryonic antigen (CEA) were negative. The appearances were typical of biphasic synovial sarcoma.

Discussion
We believe that this is the first reported example of a primary synovial sarcoma of the orbit. If this is the case it therefore expands both the range of primary sites for this tumour and the range of connective tissue tumours found in the orbit. Although this tumour was biphasic and

Figure 2 Areas showing epithelial differentiation formed either solid aggregates (A) or glandular spaces (B), many of which showed central hyalinisation.

Figure 3 Glandular spaces show predominantly luminal positivity for EMA. Note also the scattered positive spindle cells (ABC method).
Chicken pox infection (varicella zoster virus) and acute monoarthritis: Evidence against a direct viral mechanism

C G Fink, S J Read, G Giddins, R P Eglin

Abstract
A 9 year old boy developed acute monoarthritis of the left knee concurrent with the appearance of a varicella zoster virus (VZV) rash. Repeated VZV DNA hybridisation of the cells within the synovial fluid and synovial membrane failed to show any evidence of intracellular virus. Virus was isolated from synovial fluid 24 hours after the start of clinical infection but not later. These findings suggest that the mechanism of the arthritis is not due to viral replication inside the swollen joint.

Acute arthritis is a rare complication of varicella (chicken pox). Sixteen cases have been reported,¹ and in an earlier review of eight cases five involved one large joint alone.²

Case report
A 9 year old boy presented with a 24 hour history of mild papular-vesicular eruption on the trunk. This eruption developed concurrently over 24 hours with a tender, hot and swollen left knee joint which was sufficiently uncomfortable after 12 hours to inhibit weight bearing. Further crops of vesicles developed over the trunk during the next five days and the knee joint remained acutely inflamed. No vesicles were seen near to the joint. There was no history of previous injury or arthritis and a clinical diagnosis of varicella infection (VZV) was made when the child was admitted for observation and bed rest. The vesicular rash resolved over seven days and the knee joint swelling resolved over three weeks, but complete recovery of the knee joint took two months.