Orbital lymphoma in a patient with Felty’s syndrome

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Abstract
A 72-year-old woman presented with a diffuse infiltrating soft tissue mass involving the entire right orbit. She had a pre-existing phthisis of the right eye secondary to retinal detachment, and had developed painless proptosis on the same side. Her medical history included rheumatoid arthritis, leucopenia, and an enlarged spleen, characteristics consistent with the diagnosis of Felty’s syndrome. Risk of lymphoma is thought to be higher in patients with established connective tissue disorders. The case described herein, to our knowledge, represents the first report of an association between Felty’s syndrome and orbital lymphoma. The symptoms of Felty’s syndrome in our patient were initially attributed to spread of a malignant process involving the orbit.

Case report
A 72-year-old woman with pre-existing phthisis of the right eye secondary to retinal detachment developed painless proptosis on the same side. The proptosis was unresponsive to antibiotics or anti-inflammatory agents. Vision in the right eye was no light perception and a 4+ afferent pupillary defect was present. We obtained exophthalmometry readings of 29 mm on the right and 17 mm on the left, with a base of 110 mm. The left eye was normal. Magnetic resonance imaging (MRI) and computed tomography (CT) revealed a diffuse infiltrating soft tissue mass involving the entire right orbit. No bony changes were present and the sinuses were clear (Fig 1).

Systemic work-up demonstrated hepatosplenomegaly, a white blood cell count of 1.8×10⁹/l, haematocrit of 37.1%, and platelet count of 96×10⁹/l with giant forms present. Serum antinuclear antibody (ANA) titre was positive at 1:80, rheumatoid factor was negative, and the angiotensin-converting enzyme level was normal. The sedimentation rate was 20 mm/h (Westergren). Tests for anti-DNA double strand antibody, anti-scleroderma antibody, and antibodies associated with Sjögren’s syndrome were negative. We interpreted these findings as consistent with metastatic carcinoma or with disseminated lymphoma.

The physical examination prompted us to study further the patient’s medical history. She then volunteered a history of rheumatoid arthritis, leucopenia, and an enlarged spleen. Medical records from 9 years prior to presentation showed strongly positive rheumatoid factor and a positive ANA titre. A white blood cell count of 1.4×10⁹/l, a platelet count of 181×10⁹/l, and a broad-based polyclonal gammopathy were found. A bone marrow aspiration demonstrated myeloid and erythroid hyperplasia. Based upon these findings the patient had been diagnosed as having Felty’s syndrome.

The orbital mass biopsy and subsequent work-up were diagnostic for a stage I large cleaved cell lymphoma of the B type, localised to the orbit (Fig 2). A 5 week course of 5000 cGy of local radiation therapy to the right orbit produced complete regression of the tumour by radioimaging studies.

Discussion
In 1924 Felty1 reported five adults with chronic rheumatoid arthritis (RA) who had associated splenomegaly, leucopenia, and hypochromic anaemia, with or without a reduced platelet count. This syndrome occurs in less than 1% of patients with RA and is twice as common in women as in men.2 Patients with long-standing connective tissue

Figure 1 Diffuse infiltration of the right orbit, proptosis, and shrunken globe are demonstrated by axial CT scan (left) and by sagittal MRI (right).
disorders are thought to be at higher risk of developing lymphoma than is the general population. For instance, patients with Sjögren’s syndrome are about 44 times more likely to develop lymphoma than is the general population. However the lymphomas associated with Sjögren’s syndrome have been of the diffuse histiocytic type rather than of the large cleaved cell variety found in our patient; furthermore antigen associated with Sjögren’s syndrome was absent in our patient.

The development of lymphoma in the setting of long-standing connective tissue disease may be explained either by chronic immunostimulation leading to lymphoid hyperplasia and, eventually, to malignancy, or by a disturbance in immune surveillance.

As orbital lymphoma associated with Felty’s syndrome has not been previously reported, ophthalmologists should be alert to this diagnosis when evaluating patients with suggestive findings. In particular, the hepatosplenomegaly, neutropenia, and thrombocytopenia which often indicate diffuse spread of a malignancy may occasionally signify an underlying benign inflammatory process in association with localised lymphoma.