Histiocytic necrotising lymphadenitis in systemic lupus erythematosus

M D Litwin, B Kirkham, D R F Henderson, S C Milazzo

Abstract
Histiocytic necrotising lymphadenitis is the pathognomonic histological appearance of lymph nodes in Kikuchi’s disease, a condition characterised by a brief systemic illness and lymphadenopathy.

The case is described of a young man, originally diagnosed as having Kikuchi’s disease by lymph node histology, who subsequently developed systemic lupus erythematosus with symmetrical polyarthritis, Coombs’ positive haemolytic anaemia and haemorrhagic pneumonitis. The case emphasises that a range of diseases is associated with histiocytic necrotising lymphadenitis, belying the unitary impression given by the term Kikuchi’s disease.

Histiocytic necrotising lymphadenitis was first described in Japan in 1972,1 2 and is now increasingly recognised in Europe and the USA.3 Its diagnosis is based on a characteristic histological appearance of ‘lymphadenitis showing focal reticulum cell hyperplasia with nuclear debris and phagocytosis’.3 Recent case reports4-8 and an editorial9 have tended to reinforce the association of histiocytic necrotising lymphadenitis with a brief illness in previously healthy women which has a benign, uncomplicated prognosis. So confidently has this pathological lymphadenopathy been assumed to represent a discrete clinical entity that it has received an anonymous name, ‘Kikuchi’s disease’, after one of the first Japanese pathologists to describe it. Although it is important that doctors recognise the characteristic, self limiting presentations of histiocytic necrotising lymphadenitis with systemic illness and lymphadenopathy, they need also to appreciate that some patients with the same pathological diagnosis go on to develop progressive and severe disorders. One review has briefly described the development of systemic lupus erythematosus (SLE) after an initial diagnosis of histiocytic necrotising lymphadenitis.3 We report a florid case of SLE, presenting initially as histiocytic necrotising lymphadenitis.

Case report
A 19 year old man of Greek descent presented with leathery, anorexia, night sweats, and weight loss of 6 kg over a period of one month. Both his mother and grandmother had Hodgkin’s disease. His past history included chronic asthma and eczema, and repair of a duplex oesophagus. Physical examination showed marked bilateral cervical lymphadenopathy. The nodes were firm and not tender, with the largest measuring 6 x 6 cm. Smaller bilateral axillary and inguinal nodes were also noted.

Investigations showed a mild lymphopenia of 0.8 x 10^9/l (normal 1.3-3.4 x 10^9/l) with occasional atypical lymphocytes, a microcytosis of 69 fl (normal 81-98 fl) due to a β thalassaemia trait, with haemoglobin 135 g/l (normal 130-165 g/l) and an erythrocyte sedimentation rate of 18 mm/hour. Viral, toxoplasma, and HIV serology were all negative and a chest radiograph was normal.

Histology of a cervical node obtained by open biopsy showed scattered areas of necrosis, mainly in the interfollicular areas, surrounded by large numbers of proliferating, immature, mononuclear cells, some of which were atypical and mitotically active. There were also scattered foamy macrophages, lymphocytes, and a few polymorphs. Some lymphoid follicles could still be seen but the germinal centres were not prominent. The diagnosis made was histiocytic necrotising lymphadenitis, or Kikuchi’s disease.

The initial symptoms, other than lethargy, improved spontaneously and the lymphadenopathy became less prominent. Two months after presentation lymphopenia persisted and the erythrocyte sedimentation rate was 35 mm/hour. Total haemolytic complement was 20% (normal 90-96%) with a low C3 of 0.34 g/l (normal 0.48-1.35 g/l), and a low C4 of 0.09 g/l (normal 0.15-0.48).

Seven months after presentation he was less well and had developed a symmetrical, generalised polyarthritis with early morning stiffness lasting one hour and reported dyspnoea with small intermittent haemoptyses. Synovitis was present in most joints, bilateral basal pulmonary crepitations were audible, he was clinically anaemic, and generalised lymphadenopathy persisted.

Investigations were now consistent with an IgG/C3d Coombs’ positive haemolytic anaemia, with haemoglobin 65 g/l, reticulocytes 6% (normal 0.2-2%), erythrocyte sedimentation rate 96 mm/hour, and total haemolytic complement <20%. Antinuclear antibody was positive at a titre of 1/2560, homogeneous pattern, with antibody to double stranded DNA 140 IU/ml (normal 0-7 IU/ml). A chest radiograph showed cardiomegaly and peripheral bronchial thickening, but a computed tomography chest scan also showed patchy left upper lobe consolidation. Ventilation perfusion pulmonary scintigraphy was normal. Respiratory function tests showed a restrictive pattern with a forced vital capacity 67% of the predicted value. The carbon mon-
oxide transfer factor (K_{CO2}) was 120% of the predicted value. Bronchoscopy without biopsy or lavage was normal.

A diagnosis of SLE was made and treatment with prednisolone, 30 mg/day, was begun. His arthropathy, lymphadenopathy, dyspnoea, and malaise rapidly improved. The anaemia was resistant, despite an increase of the dose of prednisolone to 100 mg/day. Blood loss was suspected and an active duodenal ulcer was found at endoscopy. The haemoglobin increased steadily after ranitidine (H2 receptor antagonist) and iron replacement were added.

Twelve months later he remained well apart from mild, non-limiting dyspnoea during strenuous sporting activities, and small streaking haemoptyses on most days. His ongoing treatment was hydrochloroquine 400 mg/day and prednisolone 10 mg every other day. Chest radiographs at this time showed a transient right sided pulmonary infiltrate clearing partially within one week. The cardiac profile was normal. His forced vital capacity was 74% of the predicted value and the K_{CO2} 84% of the predicted value. Maximum inspiratory and expiratory airway pressures were normal, suggesting unimpaired respiratory muscle strength. The erythrocyte sedimentation rate was 9 mm/hour, the antinuclear antibody titre was 1/640 with antibodies to double stranded DNA 17 IU/ml; C3 was normal whereas C4 was marginally reduced at 0.13 g/l. A transbronchial lung biopsy showed pulmonary haemorrhage with widened septa which stained for IgG, consistent with lupus pneumonitis.

High dose prednisolone, 60 mg/day, was started again. After four months treatment his chest radiograph was clear and the dyspnoea and haemoptysis had improved, though they had not entirely abated. A second lung biopsy showed decreased but continuing fresh haemorrhage and decreased septal IgG. Azathioprine 100 mg/day was introduced. Four months later he still has haemoptyses of minimal volume approximately once a week.

Discussion

The patient had a triphasic illness beginning with a lymphoma-like cervical lymphadenopathy, followed by a florid multisystemic lupus which resolved to leave a chronic minimally symptomatic pneumonitis. The first two phases show an association between the histological diagnosis of histiocytic necrotising lymphadenitis and SLE, whereas the third phase questions how intensively we treat isolated, symptomatically mild, yet evident pulmonary abnormalities in chronic SLE. Published work on histiocytic necrotising lymphadenitis, which is written mainly by pathologists, not only confirms its distinction from malignancy, but emphasises that histiocytic necrotising lymphadenitis is a separate entity associated with the benign clinical features of Kikuchi’s disease. For example, Pileri et al state13: ‘...Clinically, the lesion often appears as lymphadenopathy in the neck; the enlarged nodes are painful. Many increase in size other than the cervical region and generalised lymphadenopathy are less common. Fever and leukopenia are observed frequently. The prognosis is always excellent, and many patients recover without treatment...’

The diagnosis of Kikuchi’s disease was reconsidered as multiple features of SLE became obvious. Support for our contention that the original illness was SLE came subsequently from a positive antinuclear antibody titre of 1/640, with antibodies to double stranded DNA 13 IU/ml, results available retrospectively from serum samples saved from the first presentation.

Generalised lymphadenopathy of the size seen in this patient is rare as a presenting feature of SLE. Nodal size is usually not more than three to four centimetres.11 Dubois found cervical lymphadenopathy to be the presenting feature of SLE in 1.7% of patients, and generalised adenopathy in 0.5–2.0%.11 Of established cases 50% had generalised lymphadenopathy. The histology of nodes in SLE is variable12 and can overlap those changes seen in histiocytic necrotising lymphadenitis.3

Both lymphocyte activation was assessed in our patient by assaying the number of peripheral blood cells secreting IgG, IgA, and IgM, at various times during his illness (see table). This paralleled other parameters of disease activity. Increased numbers of antibody secreting cells are a characteristic feature in active SLE but findings in histiocytic necrotising lymphadenitis have not previously been reported.

Despite his overall improvement, mild dyspnoea and haemoptysis persisted. Fleeting pulmonary infiltrates and increased K_{CO2} values suggested pulmonary haemorrhage, which was confirmed. Reports of alveolar haemorrhage in SLE describe acute, life threatening events usually in the context of glomerulonephritis and hypocomplementaemia, unlike the relatively asymptomatic nature of this problem.14–16 High doses of steroids and immunosuppressive drugs were given empirically because of the perceived threat to his lungs. There is, however, a lack of prognostic and management data about this aspect of SLE.

The association of SLE with histiocytic necrotising lymphadenitis has been suggested in one previous review, but the five patients included were neither as florid as this patient, nor were they fully documented clinically.1 Medeiros et al17 reported established SLE preceding the appearance of histiocytic necrotising lymphadenitis-type lymphadenopathy. Ohta et al18 and Lyberatos19 describe patients in whom the typical lymphadenopathy was among the presenting features of Still’s disease.

The clinical spectrum of associations of histiocytic necrotising lymphadenitis is wide and ranges from otherwise asymptomatic

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**Number of antibody secreting cells in peripheral blood (expressed per 10⁶ mononuclear cells for IgG, IgA, and IgM)**

<table>
<thead>
<tr>
<th>Time after presentation (months)</th>
<th>IgG (reference &lt;0.5)</th>
<th>IgA (reference &lt;1.6)</th>
<th>IgM (reference &lt;0.3)</th>
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</tr>
<tr>
<td>19</td>
<td>0.69</td>
<td>0.91</td>
<td>0.26</td>
</tr>
</tbody>
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adrenopathy, pyrexia of unknown origin, non-specific rashes, myalgias and arthralgias, mesenteric adenitis, Still’s disease, and SLE, through to death. This spectrum of associations of histiocytic necrotising lymphadenitis belies the unitary impression given by the term ‘Kikuchi’s disease’. We believe it would be best for doctors to avoid using the latter name as this suggests a discrete, benign, clinical entity, and hence reduces awareness of more serious diseases that may present with identical pathological and clinical features.

The aetiology of both histiocytic necrotising lymphadenitis and SLE is unclear. Reports of patients initially diagnosed as having histiocytic necrotising lymphadenitis who subsequently developed SLE and adult Still’s disease suggest that these conditions have similar causative factors. Although white patients are well represented in case series of histiocytic necrotising lymphadenitis, a large number of case reports have been of Asian patients. Our patient was of Greek descent and had lived in Australia most of his life, but the repair of his duplex oesophagus at the age of 13 years was performed in Hong Kong, from where 10 cases of histiocytic necrotising lymphadenitis have been reported. It is interesting to speculate if an operation in this country with a high occurrence of histiocytic necrotising lymphadenitis could have been the mode of transmission of an infectious agent, which has subsequently predisposed to both histiocytic necrotising lymphadenitis and SLE.

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