Cord compression: a rare complication of chronic lymphocytic leukaemia

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Abstract
Two years after the diagnosis, while the disease was under good control, a patient with chronic lymphocytic leukaemia (CLL) developed spinal cord compression from an extradural solid tumour composed of leukaemic cells. He was treated successfully with resection of the tumour followed by local radiotherapy and systemic chemotherapy.

Central nervous system complications, though common in acute leukaemias and in the blastic phase of chronic granulocytic leukaemia, are extremely rare in chronic lymphocytic leukaemia (CLL). We report a patient who developed acute cord compression as a result of extradural leukaemic deposit two years after CLL had been diagnosed.

Case report
A 66 year old man presented with cervical lumps and general malaise of three months' duration. Clinical examination showed generalised superficial lymphadenopathy and an enlarged spleen palpable 3 cm below the costal margin. There were no other abnormal physical signs. A blood count showed the following: haemoglobin 12·8 g/dl; a white cell count of 33·3 x 10⁹/l, with neutrophils 4·1 x 10⁹/l and lymphocytes 28·0 x 10⁹/l; and platelets 197 x 10⁹/l. Liver and renal functions and serum immunoglobulins were normal. A chest x-ray picture showed hilar adenopathy but no parenchymal disease. Surface marker analysis of the peripheral blood lymphocytes showed HLA-DR 80%, B-7 70%, but no surface immunoglobulins, and T cell markers in <10%. A bone marrow aspirate and trephine biopsy specimen showed dense infiltration by well differentiated small lymphocytes. B cell CLL was diagnosed.

The patient was treated for four months with chlorambucil, at a dose of 0·4 mg/kg/day for four days each month. He showed a satisfactory response, with disappearance of splenomegaly, regression of superficial lymphadenopathy, and a fall in his peripheral blood lymphocyte count to 6·0 x 10⁹/l. A repeat chest x-ray picture showed regression of hilar adenopathy. He was completely asymptomatic and no further treatment was given. One year after the diagnosis a transient IgM-κ paraprotein (2·4 g/l) was noted on three occasions.

Twenty two months after the diagnosis of CLL he developed low back pain which rapidly got worse. Three days later he complained of weakness of his legs and on the next day developed paraplegia and paraesthesia of both legs with a clinically determined concentration of L₃. He had minimal superficial lymphadenopathy and no hepatosplenomegaly. Blood counts showed that the haemoglobin concentration was 13·4 g/dl, the white cell count 19·6 x 10⁹/l, with lymphocytes 15·7 x 10⁹/l, and a platelet count of 169 x 10⁹/l. A chest x-ray picture showed no progression of hilar adenopathy. A myelogram showed cord compression at the level of T₆. A specimen of cerebrospinal fluid showed raised protein (1·2 g/l) concentration but no normal cell count. No abnormal cells were seen.

Emergency laminectomy and excision of the lesion, an extradural solid tumour (6 cm x 4 cm), was carried out. Histological examination showed a homogeneous mass of well differentiated small lymphocytes with no suggestion of nodular configuration and fibrosis. The tumour cells had marker expression similar to that of peripheral blood lymphocytes. After surgery he received local radiotherapy and systemic chemotherapy with cyclophosphamide, vincristine, and prednisolone (CVP) for six months. His neurological deficits improved slowly and he was left with minimal residual paraparesis. His blood count at the end of chemotherapy was as follows: haemoglobin 12·9 g/dl; white cell count 6·5 x 10⁹/l, with lymphocytes 3·4 x 10⁹/l, and a platelet count of 141 x 10⁹/l. Residual superficial and hilar adenopathy persisted and he developed hypogammaglobulinaemia with a reduction in all three major classes of Ig. Over the next 12 months he had several bouts of chest infection. Blood counts and immuneparesis remained virtually unchanged. Three and a half years after the diagnosis of CLL he died of severe chest infection. Permission for necropsy was unfortunately refused.

Discussion
This patient had typical CLL at the time of diagnosis and responded well to a short course of chemotherapy. He remained asymptomatic and the disease remained under very good control without any further treatment until cord compression developed. Though the symptoms developed extremely rapidly, there was no evidence of progression or transforma-
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Thorotrast granuloma: an unexpected diagnosis

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Abstract

An example of a Thorotrast granuloma (thorotrastoma) occurred in the neck of a patient 44 years after a carotid angiogram in which Thorotrast was used as radiological contrast medium. The lesion had produced a "cold" abscess and the patient was undergoing treatment for retropharyngeal tuberculosis. Thorotrast leakage can produce unusual clinical symptoms and signs which are frequently misdiagnosed.

Thorotrast is, however, highly radioactive with predominant $\alpha$ emission and a half-life of $1.39 \times 10^9$ years. Excretion of $^{232}$thorium is negligible, and thus the body becomes the permanent site of deposition of this radioactive and potentially hazardous material. In spite of an increasing realisation of its carcinogenic potential $^{1}$ Thorotrast continued to be used widely in many countries, including the United Kingdom until the early 1950s.

Case report

A 66 year old man presented with a five week history of non-productive cough, sore throat, and severe pain in the right side of the neck and right occiput, with restricted neck movements and some dysphagia. He was treated, initially by his general practitioner, with antibiotics to no avail.

On subsequent examination in the ear, nose, and throat outpatients clinic the posterior pharyngeal wall on the right side was noted to be swollen with adherent inflammatory slough, and there was some induration of the right side of the neck. The palate was mobile and no abnormalities were detected in the nasal passages. Some fasciculation of the right side of the tongue was noted. Small lymph nodes were