Angiotropic intravascular large-cell lymphoma with massive cerebral extension

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Abstract
Angiotropic intravascular large-cell lymphoma (AILL) is a rare, generally fatal disease characterised by a multifocal proliferation of neoplastic mononuclear cells within small blood vessels. The diagnosis of a patient was made at necropsy. The malignant cells had infiltrated the periventricular areas of the brain.

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Angiotropic intravascular large-cell lymphoma (AILL) is an uncommon entity characterised by a massive neoplastic proliferation of mononuclear cells within the lumina of small and intermediate sized vessels. This condition was first reported under the designation of “angioendotheliomatosis proliferans systemata”. Later various terms, such as “neoplastic angioendotheliomatosis”, “malignant angioendotheliomatosis”, “proliferating endotheliosis”, and “systemic angioendotheliomatosis” have been used to describe this rare clinicopathological entity. Initially this process was thought to be a neoplasm of endothelial origin, but immunopathological and morphological evidence now supports a lymphoid origin.

Vascular occlusions in a variety of organs can produce a bewildering array of clinical symptoms. Most commonly, patients with AILL present with bizarre neurological manifestations. Because the disease is rare and there are no specific diagnostic procedures apart from cerebral biopsy, it can be difficult to diagnose during life, and in most cases the diagnosis is made after necropsy.

Case report
A 62 year old woman was admitted because of weight loss, fatigue, nausea and vomiting. A few days after admission the patient complained of diplopia. She was bradyphrenic, and had bilateral internuclear ophthalmoplegia and horizontal nystagmus on upward gaze.

MRI showed bilateral periventricular white matter lesions in both frontal lobes. The white matter abnormalities consisted of large confluent lesions of high signal intensity on T2-weighted spin echo images. Transverse T2-weighted images showed involvement of the thalamus, globus pallidus and brainstem. After intravenous administration of Gado-

linium DTPA areas of abnormal increased signal intensity were seen around the lateral, third and fourth ventricles. There was no abnormal enhancement of the leptomeninges (fig 1).

CSF examination repeatedly revealed an elevated cell count (224/μl 3; 90% lymphocytes and 10% monocytes) and an elevated protein content (1263 mg/l). The glucose level was 2.4 mmol/l (serum 5.9 mmol/l). Antibody titres against neurotropic viruses, CSF cultures and Ziehl-Neelson stains were negative. She was treated as for tuberculous meningitis without benefit. She developed hyperventilation and coma. The MRI showed progression of the periventricular and brainstem lesions. CSF examinations showed pleocytosis, but no specific abnormalities, and especially no malignant cells. Mechanical ventilation was started and dexamethasone 12 mg per day added. Over a period of 24 hours the patient regained consciousness but remained dependent on mechanical ventilation. At this point the CSF contained tumour cells. Because of her very poor clinical condition no further treatment was started. The patient died of pneumonia after 28 days of mechanical ventilation.

Materials and methods
A complete necropsy was conducted. Formalin (5%) fixed, paraffin embedded
tissue obtained at necropsy was examined histologically with a variety of stains including hematoxylin and eosin, periodic-acid Schiff, Gomori's reticulin, Van Gieson's elastin and chloroacetate esterase.

Immunoperoxidase stains were performed using monoclonal antibodies against leucocyte common antigen (LCA) (leucocytes), MT1 (T lymphocytes monocytes, granulocytes), UCHL1 (Subset of T lymphocytes), MB2 (B lymphocytes, endothelial cells), L26 (majority of B lymphocytes), LeuM1 (neutrophilic granulocytes), Vimentin (cells of mesenchymal origin), polykeratin (cells of epithelial origin) and Factor VIII related antigen (endothelial cell differentiation).

Primary antibodies were then applied for one hour at room temperature (MT1 and MB2 24 hours at 4°C) in moisture chambers. Sections to be labelled with anti polykeratins, LCA, UCHL1 and MT1 were first digested with 0.5% trypsin in phosphate buffered saline (pH 7.4) for 30 minutes at room temperature. This step was not employed for other immunostains. After incubation with the specific antibody, the biotinylated Goat-anti-Mouse IgG was followed; after washing avidin-biotin-peroxidase complex was added. Sections were developed with diaminobenzidine for three minutes, counterstained with hematoxylin, and mounted with depex.

**Pathological findings**

At necropsy, no macroscopic abnormalities were found except for massive bilateral bronchopneumonia.

The brain was examined after fixation, and on cut surface showed multiple white lesions around the ventricles and periaqueductal grey substance. The lesions extended into the basal ganglia, the hypothalamus, and the nuclei in the tegmentum pontis (fig 2).

On histological examination, almost all organs (heart, lungs, thyroid, liver, pancreas, kidney, adrenal glands, para-aortic lymph nodes, uterus and bladder wall) contained numerous dilated vessels of various sizes which showed partial or complete occlusion by proliferating neoplastic mononuclear cells, without evidence of cohesion or syncytial aggregation (fig 3). The neoplastic cells were generally large and had scanty amphophilic cytoplasm. The hyperchromatic nuclei were variable in size, irregular in shape with wrinkled, irregular nuclear membranes. They had clumped chromatin and one or several relatively prominent nucleoli. Mitotic figures were found easily. The cells showed immunohistochemical reactivity for LCA, L26, MB2 and Vimentin. No expression of LeuM1, MT1, UCHL1, cytokeratin and Factor VIII related antigen was found. Endothelial cells showed no pleomorphism or atypia. In part of the brain, neoplastic cells were found within the lumina of vessels, adhering to the endothelial cells. In periventricular locations, along the CSF pathways, tumour cells were found outside the vessels, within the brain parenchyma. Here, a perivascular arrangement was sometimes seen. The proliferation destroyed pre-existing structures, among them the hypothalamic nuclei. The cells were histologically and immunohistochemically the same as seen in the vessels elsewhere in the body.

**Discussion**

Clinically, AILL is a difficult disease to diagnose. Bizarre neurological manifestations, such as global encephalopathy, dementia, Bradyphrenia, confusional states, or altered consciousness are often predominant, but review of the reported cases reveals additional dermatological, ophthalmological and isolated lung and adrenal gland involvement. Infrequently, CNS involvement is occult and only discovered at necropsy.

Thirty nine patients have been described in the English literature initially presenting with neurological abnormalities. There were 23...
males and 16 females and both the median and average ages of onset were 63 years with a range of 41–79 years. The disease was fatal in 36 cases, the median survival from onset of symptoms was seven months. The longest surviving patient lived nine years, but only seven patients survived longer than 16 months. Among those 39 patients 19 were diagnosed during life, and 12 received chemotherapy other than corticosteroid hormones. One received brain irradiation and another received whole body irradiation. The prognosis of angiotropic intravascular large-cell lymphoma can therefore be considered dismal, even in those cases that were diagnosed pretumour, and in which treatment was attempted. Possible survival is more related to the extent of dissemination. Because the disease is very rare and lacks specific diagnostic features, the diagnosis is infrequently established before necropsy. A remarkable discrepancy exists in many cases (ours included) between the isolated neurological symptoms, and the abundant presence of intravascular tumour cells plugging vascular lumina of almost every organ. Skin, muscle and cerebral biopsies have been taken and transbronchial biopsies have been suggested to reach the diagnosis, but since AILL is a lymphoma with the propensity to localise and proliferate throughout the body, a biopsy of any organ showing dysfunction or abnormalities may be diagnostic. Remarkably, neoplastic cells have not been reported in CSF.

Routine radiological investigation is generally not diagnostic. However, the contribution of MRI analysis may be more substantial, but has been reported only rarely. The macroscopic abnormalities of the brain were consistent with the MRI examination. The large, confluent periventricular and white matter lesions of high signal intensity on T2-weighted images and a high signal intensity after intravenous contrast administration corresponded to lymphomatous tissue and oedema.

The characteristic histological pattern of AILL shows distended vascular spaces with intraluminal cells, some appearing adherent to the endothelium, some “free floating”. The neoplastic cells are generally large and, in cases where immunohistochemistry was performed, have been proved to be of B-cell origin. Perivascular parenchymatous foci of neoplastic cells are frequently seen, but extensive extravascular spread was only once reported in the kidneys. Our case was unique in that there was a massive parenchymatous extension of tumour in the brain, especially in the periventricular region.