Progressive spinocerebellar degeneration “plus” associated with Langerhans cell histiocytosis: a new paraneoplastic syndrome?

H Goldberg-Stern, R Weitz, R Zaizov, M Gornish, N Gadoth

Abstract
Langerhans cell histiocytosis (LCH), formerly known as histiocytosis-X, manifests by granulomatous lesions consisting of mixed histiocytic and eosinophilic cells. The hallmark of LCH invasion into the CNS is diabetes insipidus, reflecting local infiltration of Langerhans cells into the posterior pituitary or hypothalamus.

In five patients who had early onset LCH with no evidence of direct invasion into the CNS, slowly progressive spinocerebellar degeneration accompanied in some by pseudobulbar palsy and intellectual decline was seen. Neurological impairment started 2-5 years after the detection of LCH. No correlation was found between the clinical syndrome and location of LCH or its mode of treatment.

An extensive search for metabolic, toxic, neoplastic, and hereditary aetiologies for progressive cerebellar degeneration was negative. It seems that the clinical entity described here may be considered a new paraneoplastic syndrome related to LCH. It may be induced by the eosinophil derived neurotoxin, which was shown to cause damage to Purkinje cells and pyramidal neurons.

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Keywords: Langerhans cell histiocytosis; central nervous system; cerebellum.

It is almost 40 years since Lichtenstein linked eosinophilic granuloma of bone, Letterer-Siwe disease, and Hand-Schuller Christian syndrome under the term “histiocytosis-X”. It seems that the presently used term Langerhans cell histiocytosis (LCH) is more appropriate as the pathological proliferation of Langerhans cells, with the typical Birbeck granules, is the uniform microscopic finding for all forms of histiocytosis-X.

Complications of the CNS as a result of LCH, manifested mainly by diabetes insipidus or cerebellar dysfunction, are traditionally believed to result from (a) CNS extension from neighbouring bones; (b) dural and leptomeningeal spread; (c) primary intraparenchymatous lesions; (d) side effects of chemotherapy and radiotherapy; (e) a combination of these.

In recent years, improved diagnosis and treatment has resulted in prolonged survival, enabling the study of long term sequelae of LCH.

Ranson et al. reported that about half of his patients with generalised LCH had “neuropsychiatric disability”, and the Southwest Oncology Group reported on 17 out of 56 long term survivors who had a variety of neurological disabilities including cerebellar ataxia in two of them; details of neurological state or neuroimaging studies were not noted.

The present report describes five patients who had extraneural LCH, in whom progressive spinocerebellar syndrome appeared several years after the initial diagnosis.

In all cases the family history was negative and the initial neurological examination, CSF content, and brain imaging at the time of LCH detection were normal. Diabetes insipidus was looked for but never found, neither initially nor during the long term follow up. A meticulous search for known causes of progressive spinocerebellar impairment of young onset was negative in all cases.

Case reports
PATIENT NO 1
In a previously healthy child, LCH, in the form of eosinophilic granuloma of the left mastoid, was excised at the age of 18 months. There were no symptoms or signs of invasion of the CNS. After operation he was irradiated locally and subsequently developed postradiation necrosis of the left side of his face and sternocleidomastoid muscle resulting in fixed left torticollis and left nerve deafness. His condition remained stable until the age of 4 years, when gradual gait instability appeared. Subsequently, progressive dysarthria developed. Severe motor dysfunction confined him to a wheelchair at the age of 30 years. Recurrent grand mal seizures occurred at the age of 37 years and have been successfully controlled with phenytoin. At the last clinic visit after 36 years of continuous follow up, the cardinal neurological abnormalities included emotional lability with uncontrolled crying, severe dysarthria, gaze directed horizontal nystagmus, abnormal otular smooth pursuit, slow saccades, generalised hypotonia, symmetric hyperreflexia, mild ankle clonus, bilateral extensor plantar response, severe disequilibrium and limb ataxia, and prominent dysmetria.

PATIENT NO 2
A girl was healthy until 3 months of age when
Patient No 3, a 27 year old man. (A) T1 weighted (TR/TE 440/27) sagittal; and (B) coronal (TR/TE 350/12) 0-5 T1 MR images. Note pronounced atrophy of vermis, cerebellar hemispheres, and pont with mildly prominent cerebral sulci for age.

erythematous seborrhoeic skin rash and purulent otitis were noted. An osteolytic lesion of the left mastoid was seen on skull radiographs. Skin biopsy disclosed typical LCH. The neurodevelopmental state, routine laboratory tests, spinal fluid, skeletal radiological survey, and cerebral CT were normal. The family history was negative. She was treated with vinblastine sulphate, cyclophosphamide, and prednisone divided into 17 single doses during the first year of life. At the age of 3 years and 10 months she started dragging her right leg. A year later she was seen for frequent falls. During the subsequent months, mental deterioration, severe progressive ataxia, limb dysmetria, muscle hypotonia, brisk deep tendon reflexes, and bilateral extensor plantar response were noted.

ANCILLARY TESTS
In all five cases routine laboratory tests, including a screen for metabolic errors of childhood, heavy metal storage, CSF composition, EEG, nerve conduction studies, and EMG were normal. In all cases brain CT disclosed mild to moderate cerebellar and pontine atrophy, symmetric dilatation of the ventricular system, and mild cerebral cortical atrophy. Similar findings were obtained in
patients 1–4 by MRI. Neither masses, nor hyperintense signals on T2 weighted images were detected. No gadolinium enhanced images were obtained. During the long follow up, additional CT and MRI were almost identical to the initial studies in case 1. In patients 2–4, slow progression of imaging abnormalities was evident. The most striking changes were seen in patient No 2. Initial studies were almost normal, but the typical radiological picture evolved during the next five to six years. The serum samples and CSF of patients Nos 1, 3, and 4 did not contain antibodies to Purkinje cells.

**Discussion**

In 1985 we first reported four cases with late neurological deterioration associated with LCH with no evidence of disease invasion into the nervous system. In the present report we have completed a follow up period ranging from 10 to 36 years and have added an additional patient.

Of the five patients described, four had the localised form of LCH, eosinophilic granuloma type and one had the diffuse type with mastoid lesions and skin involvement. The diagnosis in each case was confirmed histologically. The bony lesions were excised in four patients (with addition of local irradiation in two). Patient No 2, with the generalised form of the disease, was the only one who received chemotherapy. All patients developed signs of progressive spinocerebellar impairment 2–5 to seven years after the initial diagnosis. Neurological findings consisted of severe cerebellar ataxia, brisk deep tendon reflexes, and bilateral extensor plantar responses. In three patients intellectual deterioration and in two pseudobulbar signs subsequently appeared (table). The neuroradiological findings in all cases were similar and showed cerebellar and pontine atrophy with no signs of demyelination on MRI and no evidence of disease invasion.

An extensive search for hereditary, metabolic, neoplastic, immunological, and toxic aetologies was negative. No correlation was found between the clinical syndrome and the location of the lesion or its mode of treatment. The normal CSF protein, metabolic screen, EMG, and nerve conduction studies together with the negative family history helped in excluding the hereditary progressive cerebellar atrophies of young age.

Diabetes insipidus and other endocrine abnormalities resulting from hypothalamic or pituitary invasion of tumour cells are the most common neurological manifestations of LCH. Among the extradiencephalic sites of this disease, the cerebellum seems to be the most vulnerable. Recently, a diffuse infiltration of the CNS by Langerhans cells was reported.

Grois et al summarised 72 cases with LCH and CNS complications reported since 1924. In 72% those symptoms started a few years after the onset of LCH. The most frequent presentation was spinocerebellar impairment, as in our cases. In all patients there was evidence of invasion of the CNS manifested by diabetes insipidus or intraparenchymatous infiltration into the cerebrum, cerebellum, etc, detected by neuroimaging or necropsy.

Our cases are unique in that they lacked evidence of cranial or brain parenchymatous spread at the time of initial diagnosis or during the long term follow up. Rosenfield et al described the neuroimaging features of eight patients with LCH who had neurological manifestations. Their case No 4 showed clinical and radiological signs of progressive cerebellar degeneration without evidence of invasion of the CNS. We believe that this case resembles our patients. Kepes et al stated that a possible mechanism for neurological dysfunction without direct spread of Histiocytosis-X into the CNS may be secondary demyelination. The absence of hyperintense signals on the T2 weighted MRI in our patients, however, excludes this assumption. As patients with skull lesions have occasionally received radiotherapy, the possibility that cerebellar degeneration is a delayed effect of radiation was considered. The presence of clinical and radiological cerebellar degeneration in cases that were never irradiated exclude this possibility.

Considering that progressive spinocerebellar dysfunction in our patients could not be attributed to currently known aetologies and that it is not caused by tumour invasion, the possibility that a remote effect of LCH caused

**Clinical manifestations of patients**

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Sex</th>
<th>Age at LCH diagnosis (y)</th>
<th>Location of granulomas</th>
<th>Treatment</th>
<th>Age at onset of neurological impairment (y)</th>
<th>Duration of neurological disease (y)</th>
<th>Neurological signs</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>18/12</td>
<td>Left mastoid</td>
<td>Excision and local irradiation</td>
<td>4</td>
<td>36</td>
<td>SCD, Seizures</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>3/12</td>
<td>Skin, left mastoid</td>
<td>Chemotherapy</td>
<td>3 10/12</td>
<td>10</td>
<td>MD, Seizures, MD</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>2</td>
<td>Right mastoid</td>
<td>Excision and local irradiation</td>
<td>4 8/12</td>
<td>22</td>
<td>MD, Seizures, MD</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>18</td>
<td>Right 9th rib</td>
<td>Excision</td>
<td>21</td>
<td>20</td>
<td>MD</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>2</td>
<td>Biparietal Right proximal femur Right iliac bone</td>
<td>Excision of skull lesion</td>
<td>9</td>
<td>10</td>
<td>SCD, MD</td>
</tr>
</tbody>
</table>

SCD = Spinocerebellar degeneration; MD = mental deterioration; PBP = pseudobulbar palsy.
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The lesions of LCH consist of mixed histiocytic and eosinophilic infiltrates with varying multinucleated giant cells. It has been suggested that LCH may represent an uncontrolled immunological reaction to an unknown antigen. In 1932, Gordon reported that injecting rabbits intracerebrally with lymph node suspension obtained from patients with Hodgkin's disease caused ataxia and muscle paralysis after two to 20 days. The histopathological findings consisted mainly of damage to Purkinje cells, with a lesser hippocampal and pyramidal neuronal involvement. Further information on the potential tissue damaging effects of eosinophils has emerged from precise delineation of the unique intracytoplasmic content of the eosinophilic granules.

It was shown that intracerebral injection of eosinophil derived neurotoxin, a glycosylated protein with ribonuclease like activity can cause ataxia and limb paralysis in experimental animals probably by inactivation of protein synthesis. Necropsy studies show that the neurotoxin damages mainly Purkinje cells, pyramidal neurons, and myelin.

The fact that the median time of onset of neurological findings after the detection of LCH in our patients was 3-5 (range 2-5-7) years is compatible with the concept of para-neoplastic syndromes according to Posner, who stated that a remote effect of cancer implies any nervous system dysfunction that is not caused by direct invasion or metastases. There is no limitation of time between the appearance of the neurological deficit and the neoplastic process or vice versa. Although no antibodies against Purkinje cells or other neuronal elements were found in our cases and others the possibility that the damage is mediated by an autoimmune mechanism was not excluded.

We believe that in patients with cured extraneural LCH, progressive neurological impairment manifested mainly by spinocerebellar degeneration may be considered a remote effect on the nervous system, by a mechanism as yet unknown.
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