Cerebellar degeneration with Hodgkin's disease

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The association of neurological symptoms with Hodgkin's disease is uncommon when there is no direct involvement of the central nervous system by tumour tissue. This paper describes a patient in whom the first symptoms of neurological disorder appeared four months before the clinical features of Hodgkin's disease developed.

CASE REPORT

A male glassworker, aged 19, who had previously been in good health developed severe occipital headache, made worse by movements of the head and by stooping. A fortnight later the headache had become bitemporal and gradually less severe so that he was able to return to work. During this initial period no abnormal signs appeared in the nervous system or elsewhere although the blood pressure was somewhat labile and was variously recorded as 150/90, 135/75, and 140/100 mm. Hg.

Two months later he was referred again because of brief dizzy spells for some of which he had to be taken home from work. During these he would have to hold on to objects to avoid falling and there would at times be nausea and vomiting. A week later he improved sufficiently to attempt to drive a car to visit relatives but on the way he developed double vision and was forced to return. Some days later he drove a hundred miles without trouble. Whilst on holiday, however, double vision recurred, his gait became unsteady, and his speech indistinct.

Neurological examination 10 weeks from the onset revealed a pure cerebellar disorder with staccato speech, and a horizontal nystagmus to the left and right. There was such marked ataxia of the limbs and trunk that he was unable to stand unsupported. There was no weakness or sensory loss and the deep and superficial reflexes were normal with flexor plantar responses. The optic fundi were normal and the visual fields full. The remainder of the cranial nerves and the rest of the nervous system showed no abnormality apart from the cerebellar signs.

He was admitted to hospital for detailed investigation. The peripheral blood showed a haemoglobin level of 116% (16-9 g.) white blood cells 8,400 per c.mm. (neutrophils 80%, lymphocytes 18%, monocytes 2%). The E.S.R. was 14 mm. in one hour. Urine examination was normal. The total serum protein concentration was 6-5 g.% (albumin 3-5, globulin 2-7, fibrinogen 0-3).

FIG. 1. Electroencephalogram showing excessive slow activity in all areas.
The cerebrospinal fluid was clear and colourless, under a pressure of 135 mm. with less than 1 cell per c.mm. The total protein content of the fluid was 60 mg. per 100 ml., the globulin content was not increased, and the gold curve showed 0000000000. The Wassermann reaction in both blood and cerebrospinal fluid was negative.

A chest radiograph on admission was normal. Bilateral carotid and vertebral angiograms were also normal.

The E.E.G. showed gross abnormalities. Very little alpha activity was present, the basic rhythms being predominantly delta and theta which were equal over both hemispheres (Fig. 1). The slow activity was appreciably increased by hyperventilation.

On completing the preliminary investigation, A.C.T.H. (20 units twice daily) was started as a symptomatic measure and was continued for eight weeks. During this period the patient became slightly less unsteady and was able to walk three paces between hospital beds unaided.

Two weeks after completing the course of A.C.T.H. he developed a fever of 100° to 104°F. Routine blood cultures were sterile and there was no response to tetracycline. The white cell count at this time rose steadily to 20,000/c.mm., the differential count being polymorphs 87%, lymphocytes 3%, and eosinophils 10%. For the first time the lower cervical lymph nodes became enlarged and a second chest radiograph showed the superior mediastinal shadow to be widened by enlarged lymph nodes. The sternal marrow was extremely cellular due to marked myeloid activity but the myeloid series was morphologically normal and erythropoiesis was normoblastic. A Paul-Bunnell test was positive to a titre of only 1 in 32 and the dye dilution and complement-fixation tests for toxoplasmosis were negative.

A cervical node was removed and showed Hodgkin's disease. Cyclophosphamide, 100 mg. twice daily, was then administered for four and a half weeks during which time the white cell count was reduced to 5,000/c.mm., but there was no improvement in the patient's general condition. The lymph nodes continued to enlarge and fever persisted. As the head hair began to fall out cyclophosphamide was discontinued.

During this time there was no improvement in the cerebellar dysfunction; speech remained slurred and there was some weakness of the palate and disturbance of the gag reflex. In the next four months the general condition gradually deteriorated although the neurological disorder remained essentially unchanged and at no time did consciousness or intellect become impaired.

Ten months after the onset the patient developed respiratory distress and high fever once more. The white cell count rose to 42,000/c.mm. (polymorphs 96%, lymphocytes 4%). The chest radiograph now showed a large mass in the superior mediastinum on the right side extending far out into the right lung field.

On examination, nystagmus and other cerebellar signs persisted as did the weakness of the gag reflex. The triceps tendon reflexes were present, but the biceps, knee, and ankle reflexes could not now be obtained. There were still no long tract abnormalities.

The patient died a week after the onset of this final episode.

POST-MORTEM FINDINGS A fluctuant subcutaneous swelling present over the manubrium was continuous with a necrotic mediastinal mass, 5 in. in diameter, which had infiltrated the wall of the trachea and the sternum, and which had indented the upper lobe of the right lung. There was bilateral bronchopneumonia but no extensive consolidation, and no macroscopical direct involvement of lung by tumour.

The pericardium was thickened beneath the mediasti- nal mass, but the heart appeared normal. The liver showed congestion with necrotic nodules up to 2 cm. in diameter surrounded by capillary proliferation. A necrotic mass was present in the region of the porta hepatitis, but no biliary obstruction. The spleen (400 g.) was slightly enlarged, of a homogeneous fleshy, pink colour, and showed loss of normal structure. The kidneys had a streaky appearance due to pyelonephritis. Lymph nodes in the neck and mesentery were enlarged and firm.

Externally and on macroscopical section the brain showed slight atrophy of the cerebellum and olives, but no gross abnormality. No tumour tissue was seen within the skull.

MICROSCOPY Sections of mediastinal mass, liver nodules, and spleen all showed infiltration by reticulum cell multinucleate giant cells, and large numbers of eosino- phils. Bands of fibrous tissue were prominent and there was much necrosis (Figs. 2 and 3). The appearances were typical of Hodgkin's disease.

Central nervous system The stains used on paraaffin embedded material were haematoxylin and eosin, Nissl, cresyl violet, Heidenhein stain for myelin, phosphotungstic acid haematoxylin, and Holzer's crystal violet and for glial fibres; on frozen sections, oil red O, periodic acid-Schiff, buffered toluidine blue, Bielschowsky's silver hydroxide method for axis cylinders, and Cajal's gold sublimate method for astrocytes were employed.

The most striking changes were present in the cere- bellum. There was extensive loss of Purkinje cells throughout, but particularly in the vermis (Fig. 4); the few remaining cells, seen in small groups in the lateral part of the hemispheres, showed shrinkage, hyperchromasia, and homogenizing change. In Bielschowsky's silver preparations basket fibres surrounded some of the Purkinje cell remnants and occasional axonal swellings were seen (Fig. 6). Some tangential fibres remained intact but many were short and stunted. The Bergmann astrocytes were much increased, even in the areas where Purkinje cells persisted, and there was a mild, diffuse gliosis increase in the molecular cell layer (Fig. 5). Granule cells were generally reduced in number but abnormal forms were not seen.

The nerve cells of the dentate nuclei showed shrinkage and homogenizing change. There was diffuse loss of myelin in the surrounding area, which contained many macrophages filled with sudanophilic droplets (Fig. 7).

Similar diffuse myelin loss was found in some of the cerebellar folia, particularly those more centrally placed. In these demylinated areas, and also in some of the folia where myelin loss was slight, marked fibrillary gliosis was seen (Fig. 8). Some of these astrocytes were large but bizarre, multinucleate, tumour-like forms were not seen.
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FIG. 2. Section of tumour nodule in liver. Haematoxylin and eosin $\times$ 160.

FIG. 3. Section of mediastinal tumour showing reticulum cells and tumour giant cells. Haematoxylin and eosin $\times$ 350.

FIG. 4. Cerebellar folium showing absence of Purkinje cells and increase in Bergman astrocytes. Nissl $\times$ 65.

FIG. 5. Same cerebellar folium as in Fig. 4. Nissl $\times$ 350.
FIG. 6. Cerebellar cortex showing degenerate Purkinje cell remnants and one swollen axon. Tangential fibres are short and irregular. Bielchowsky $\times 650$.

FIG. 7. Dentate nucleus showing shrinkage of nerve cell and lipid macrophages in adjacent white matter. Heidenhain's myelin $\times 350$.

FIG. 8. Cerebellum and medulla, showing gliosis around dentate and olivary nuclei and in white matter of cerebellar folia. Holzer $\times 14$.

FIG. 9. Edge of olive showing large fibrillary astrocytes. Holzer $\times 350$. 
Abnormal enlarged oligodendrocytes and eosinophilic
inclusions were not found.
The inferior cerebellar peduncles showed mild myelin
loss and gliosis but no definite changes were seen in the
superior and middle peduncles and in the nuclei pontis.
There was glial increase, including a few large astrocytes
in the inferior olives, particularly in the hilar region, and
although some apparently viable nerve cells persisted,
many were shrunken, distorted, and hyperchromatic
(Fig. 9). Occasional degenerate nerve cells were seen
also in the vagal and hypoglossal nuclei.
No gross changes were found in the red nuclei, sub-
stantia nigra, or subthalamic nuclei; nor in the thalamus,
mammillary bodies, putamen, or globus pallidus.
Sections of frontal, temporal, parietal, and occipital
cortex were examined and no significant lesions were
found, apart from mild nerve cell loss and moderate
gliosis in Ammon’s horn, which may well have been due
to terminal hypoxia associated with pulmonary in-
fec tion.
Perivascular lymphocytic cuffing was noted around
some brain-stem and cerebellar capillaries and arterioles,
and occasionally in the cerebral hemispheres where it was
most apparent in the subcortical white matter of the
temporal lobes. A few clusters of lymphocytes were seen
in relation to the meninges, but there was no significant
infiltration.
The whole spinal cord was not available but in the
upper cervical region no spino-cerebellar or other tract
degeneration could be shown by oil red O or myelin
stains.

DISCUSSION

Attention was drawn to the relation between
subacute cortical cerebellar degeneration and carci-
noma by Brain, Daniel, and Greenfield (1951).
They reviewed previously reported cases of subacute
cortical cerebellar degeneration and added four
cases of their own. In these cases cerebellar symp-
toms developed fairly rapidly, progressing from un-
steadiness of gait to include clumsiness of the hands
and dysarthric speech, till in a few weeks or months
the patient was no longer able to sit up unsupported.
In the whole group of 16 cases, 11 were associated
with carcinoma. This association in the subacute
cases contrasted with chronic forms of cerebellar
degeneration where no such association exists.
Henson, Russell, and Wilkinson (1954) reported 19
cases of carcinomatous neuropathy and myopathy;
five of them had cerebellar symptoms, but only one
showed marked cerebellar cortical degeneration,
though degeneration of the dentate nucleus, olives,
subthalamic nuclei, and motor neurons of the spinal
cord and medulla oblongata was found in another.
Brain and Henson (1958) reported further cases and
pointed out that neurological symptoms could
antedate the clinical manifestations or diagnosis of
the carcinoma by at least three years. These reports
of non-metastatic neurological complications of
malignant disease have all been in association with
carcinoma.
In 1958, Aström, Mancall, and Richardson re-
ported three cases of unusual demyelinating lesions;
in one the patient had died of Hodgkin’s disease, in
the other two cases of leukaemia. Subsequent similar
reports have appeared by Cavanagh, Greenbaum,
Marshall, and Rubinstein (1959) and Lloyd and
Urich (1959). A comprehensive report by Richardson
(1961) summarized 22 previously recorded cases of
progressive multifocal leucoencephalopathy and
gave a detailed description of the clinical features and
pathology. This condition usually occurs against
a background of chronic disease affecting the reticulo-
endothelial system, including the reticuloses and
leukaemias, carcinomatosis, sarcoidosis, and miliary
tuberculosis. The clinical neurological manifestations
are diffuse, eventually bilateral but asymmetrical,
and whilst the cerebral hemispheres are always
affected, lesions may occur in the brain-stem,
cerebellum, and basal ganglia.
Brain (1963) reviewed the neurological complica-
tions of neoplasms and discussed the non-metastatic
complications in three main groups: encephalo-
myeloneuropathy, including subacute cerebellar
cortical degeneration associated with carcinoma;
myopathic-myiasthenic syndrome; and progressive
multifocal leucoencephalopathy associated with
chronic disease of the reticulo-endothelial system,
most often Hodgkin’s disease, leukaemia, or lymp-
sarcoma.
The only previous report of a case of subacute
cerebellar degeneration associated with Hodgkin’s
disease is that of Rewcastle (1963) and he could only
find one similar case recorded, that included by
Malamud in his atlas of neuropathology (1957).
Rewcastle’s case occurred in a 62-year-old man, but,
apart from the age of the patient, the clinical and
pathological features are strikingly similar to the
present case. In both cases initial symptoms of
headache, dizziness, and diplopia were followed
after a short remission by progressive cerebellar
symptoms. Constituents of the cerebrospinal fluid
were normal in Rewcastle’s case and an E.E.G.
showed only a minimal irregularity over the left
frontal area. Another finding common to both cases
was the leucocytosis with eosinophilia (18,300 per
c.mm. with 7% eosinophilia in Rewcastle’s case,
and 20,000 per c.mm. with 10% eosinophilia in our
case). In both cases these high eosinophil counts
were made about five months after the initial symp-
toms, at the time the Hodgkin’s disease became
apparent; in our case the white cell count had been
normal on the first admission. Rewcastle found a
reduction in size of enlarged lymph nodes and a
partial remission in the cerebellar symptoms after a course of mechloretamine, whereas the present case failed to respond to cyclophosphamide, a similar cytotoxic agent.

Histologically also both cases were very similar and showed much in common with the cases of subacute cerebellar degeneration associated with carcinoma, such as depletion of Purkinje cells, axonal swellings, moderate loss of granule cells, demyelination, and gliosis of dentate nuclei and similar less intense changes in the cerebellar folia.

Focal demyelination was not seen in the cerebral hemispheres in the present case. Other typical features of multifocal leucoencephalopathy as mentioned by Richardson (1961), such as abnormal oligodendrocytes, eosinophilic inclusions, abnormal granule cells, bizarre or tumorose astrocytes, were not found.

Hutchinson, Leonard, Maudsley, and Yates (1958) reviewed 229 patients with reticulosis and found 45 with neurological disorders. These included an unusual form of myelomalacia, peripheral neuropathy, and papilloedema without raised intracranial pressure. Haynal and Regli (1964) reported 27 patients who presented with a lesion of the nervous system out of an unspecified total of cases of Hodgkin’s disease treated in the university neurological clinic at Zurich. In these cases the cerebral lesions were the result of focal demyelination and included cerebellar symptoms and also a granulomatous angiitis going on to meningoencephalitis. Cranial and peripheral nerve lesions as well as paraplegia were discussed and one case of polymyositis in Hodgkin’s disease was described. Deep, Fraumeni, Tashima, and McDivitt (1964) report leucoencephalopathy and dermato-myositis in a case of Hodgkin’s disease. In none of these papers were there any case reports of subacute cerebellar cortical degeneration associated with reticulosis.

The aetiology of non-metastatic neurological disorders complicating malignant disease remains obscure. If there is a general aetiological explanation, it must account for the rarity of these conditions, and the fact that more than one type of neurological disorder may complicate the same type of malignancy, possibly even in one individual. Furthermore the neurological disorder may precede or follow the appearance of malignancy, undergo a spontaneous remission, be arrested by surgery or chemotherapy directed at the malignant disease, and may even appear for the first time after such treatment. The same type of neurological disease, particularly multifocal leucoencephalopathy, may complicate benign conditions or even appear in elderly but otherwise normal subjects.

In the present case there was no known industrial or hereditary predisposition to neurological or malignant disease. The subacute stuttering onset suggested the start of an inflammatory or vascular process, but at this time there was no pyrexia, and the blood count was normal. Carotid and vertebral angiograms were also normal. The cerebrospinal fluid was not examined till two months after the clinical onset but then showed only a slight rise in protein content. At necropsy no inclusions were found and perivascular cuffing, though nowhere intense, was more evident in the cerebellum and brain-stem where it was thought to be reactive to neuronal degeneration rather than the result of infection.

The possibility of a circulating toxin was considered, particularly in view of the high eosinophil count noted also in Rewcastle’s case, which occurred at the time of the enlargement of glands due to Hodgkin’s disease, and the known relationship between Purkinje cell loss and experimental injection of eosinophils. However, considerable damage to the cerebellum clearly took place four months before the enlargement of lymph glands at a time when there was no increase in circulating eosinophils. The E.E.G. abnormalities were an unexpected finding in cerebellar degeneration, although Liversedge and Emery (1961) have described similar changes where there was no increase in intracranial pressure. It was thought that the abnormalities in the present case arose in the cerebral hemispheres although they were not the changes usually found in association with Ammon’s horn sclerosis. Possibly they were evidence of some toxin to which the cerebral cortical nerve cells were more resistant than the cerebellar neurones.

An alternative possibility that the agent producing the neurological damage also promoted the Hodgkin’s disease seems unlikely, in view of similar cases where cortical cerebellar degeneration antedates carcinoma arising in various organs.

In general, neoplastic-neuropathic associations follow patterns as described by Brain (1963), but less common associations may be important in providing aetiological clues although so far they only appear to have eliminated a type-specific concept.

SUMMARY

A case of subacute cerebellar degeneration with Hodgkin’s disease is described. The subject was a 19-year-old man who presented with intermittent headache and vertigo. A severe cerebellar disorder followed, becoming only slowly progressive in the later stages. Five months after the onset of neurological symptoms enlarged supraclavicular glands appeared. A biopsy of a cervical lymph node showed
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Hodgkin’s tissue. The condition was treated first with A.C.T.H. and later with cyclophosphamide without material benefit. Death occurred 10 months after the onset.

At necropsy Hodgkin’s disease of the mediastinal lymph nodes with tumour nodules in the porta hepatis and liver was confirmed. The brain showed cerebellar cortical degeneration with patchy demyelination and gliosis. Degeneration and gliosis were also prominent in the dentate nuclei and olives. Less severe nerve cell loss and gliosis occurred in Ammon’s horn.

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REFERENCES


