The nosological position of concentric lacunar leucoencephalopathy

S. CURRIE, A. H. ROBERTS, AND H. UРИCH
The London Hospital, Whitechapel, London

In 1960 Grcević reported an apparently unique case. The patient, a woman, had been well until the age of 30. In the following three years she became 'almost blind', her vision deteriorating in an illness characterized by exacerbations and partial remissions. She then became progressively demented, developing abruptly a hemiparesis and aphasia six months before her death in epileptic status at the age of 35. At necropsy extensive changes were found in the white matter of the rostral two-thirds of the cerebral hemispheres consisting of concentric lacunae separated by glial septa. In the affected areas there was total loss of axons and myelin sheaths. The lesions bore a superficial similarity to the concentric sclerosis of Baló (1928) from which they differed in being destructive and not demyelinating. The author discussed the nosological position of the condition which he named 'concentric lacunar leucoencephalopathy', and its relationship to demyelinating diseases, without reaching definite conclusions.

We report a case in which the association of similar lesions in the white matter with patchy demyelination in the optic pathways and an acute necrotizing myelopathy suggested a close relationship to the demyelinating diseases, particularly to neuromyelitis optica.

CASE REPORT

CLINICAL HISTORY Mr. G.R., a labourer, was admitted to his local hospital at the age of 64 in August 1966 with painless deterioration in the vision of both eyes. This appeared to have developed suddenly overnight. When he was seen two weeks later a left homonymous field defect was found which over the next two weeks progressed to complete blindness. His memory had become defective during the same period. Vision in the left eye was said to have been poor for years, but three years earlier severe myopia had apparently been corrected by an optician. Two weeks after the sudden deterioration in vision lumbar puncture was performed; the cerebrospinal fluid was found to contain 208 mg./100 ml. protein but no other abnormality, and manometriccs were normal. The Wassermann reaction was negative in blood and fluid. It was thought that the patient might have a suprasellar lesion and he was transferred to a neurosurgical centre. There he was found to be disoriented. The eyes were myopic and the optic discs were noted to be pale, but there appeared to be normal pupillary reactions despite imperfectness of light. There was a mild impairment of superficial sensory perception over the left side of the body. At ventriculography the fluid was xanthochromic and turbid, containing 375 red blood cells and 4 white cells/mm. and 140 mg/100 ml. of protein; manometrics and radiology were normal. A right vertebral angiogram was normal. Infarction of both occipital lobes was considered the most probable diagnosis.

The patient's condition then remained unchanged until five months later in January 1967 he developed a sudden left hemiplegia from which he recovered within 24 hours. Three months after this a grand mal epileptic attack occurred, followed by transient left-sided weakness. He was admitted to a local hospital again. There he was found to be orientated and co-operative, but blind without light perception, the pupils reacting only sluggishly to light. The optic discs were noted to be pale. No disturbance of left-sided motor or sensory function was apparent. He was normotensive. An ESR and blood count were normal. A right carotid angiogram demonstrated atheroma at the origin of the internal carotid artery but no abnormality of the intracerebral vessels. The patient was treated with phenobarbitone and had only one further major seizure during the following year.

He then remained well again until October 1967, five weeks before his death, when a sudden episode of weakness of both legs occurred lasting half an hour. This recurred within a fortnight, again transiently, on this occasion after a long walk. Two days later he awoke in the night with severe lumbar pain and on the following morning was incontinent of urine and unable to move his legs. On admission to hospital a dense flaccid paraplegia was found with cutaneous sensory loss from the fourth thoracic dermatome caudally; bladder drainage was required.

He was transferred to the London Hospital (no. 369575) the next day. On admission he was apyrexial, but drowsy, disoriented, and euphoric. He was blind without light perception in either eye. Pupillary reactions were present...
but not in response to light, and were thought to indicate
reflex attempts at convergence. The left eye was highly
myopic and both optic discs were atrophic. The retinas
and vessels were normal. Neck stiffness was present.
Initially a flaccid paraplegia with slight weakness of the
wrist extensors on the left was noted. The tendon
reflexes were sluggish in the arms, with the exception of
the triceps, which were brisk, and absent in the legs.
There was little response to plantar stimulation. Respiration
was largely diaphragmatic, only the upper inter-
costals contracting. There was anaesthesia from the
second thoracic dermatome caudally. Later the same day
the sensory level was found to have risen to the fourth
cervical dermatome on the left and to the eighth on the
right, and breathing had become entirely diaphragmatic.
Routine investigations including peripheral blood exam-
ination and radiographs of the chest and spine were
normal. The ESR was 20 mm in one hour. The cerebro-
spinal fluid contained 355 white cells/cu. mm, pre-
dominantly neutrophils, 40 mg/100 ml. protein, and 90
/100 ml. sugar. The Wassermann and Lange colloidal
gold reactions in the fluid were negative and no organisms
were seen or cultured. A myelogram was normal. A
presumptive diagnosis of infarction of the cord was made.
Within two days the sensory level had risen to the second
cervical dermatome, the arms had become asymmetrically
weak with slight power remaining only proximally in
deltoids and biceps, while the tendon reflexes could no
longer be elicited. He continued to deteriorate, the arms
becoming flaccid and totally paralysed the day before his
death. He died on 6 November 1967, 11 days after the
onset of persistent leg weakness.

Necropsy findings (PM 354/67) The following features
were observed: concentric lacunar leucoencephalopathy;
retrolublar neuritis; acute myelomalacia; deep femoral
vein thrombosis; pulmonary embolism; and acute
haemorrhagic cystitis.

Examination of the nervous system
Macroscopic The brain (1,176 g) was externally normal. Coronal sections
of the cerebral hemispheres revealed extensive lacunar
softening in the white matter, involving both occipital
lobes, parts of the right parietal and right temporal lobes,
and the splenium of the corpus callosum (Fig. 1). The
lacunae were arranged in approximately parallel, con-
centric rows, separated by lamellae of firm, white glial
tissue. In the lateral parts of the white matter the lacunae
were larger and more widely spaced out than those situ-
ated more medially. The cortex over the affected areas
appeared normal. Right Ammon’s horn was shrunken and
discoloured. Both lateral geniculate bodies were
atrophic. The intracranial portions of the optic nerves and
the optic chiasm showed no obvious lesions. The ventricles
were normal. No lesions were seen in the basal ganglia,
brain-stem or cerebellum.

The spinal cord externally appeared swollen and dis-
coloured from the lower cervical to the lower thoracic
segments. Transverse sections showed central softening
with partial peripheral preservation from C4 to C7.
From C8 to T11 the cord appeared completely necrotic
with patches of haemorrhagic discoloration. The lowest
thoracic and all lumbar segments were fairly well pre-
served and showed central small patches of brownish
discolouration. The anterior and posterior roots and
the posterior root ganglia appeared normal.

Microscopic The cerebral cortex showed good pre-
servation of its cyto-architecture with the exception of
right Ammon’s horn which showed a large defect in
Sommer’s sector of the pyramidal layer. Sections stained
for myelin showed partial demyelination of the cortex in
areas adjacent to the lesions in the white matter. This was
most striking in the calcineurine fissure where only the tips
of the cuneus and the lingual gyrus remained fully
myelinated, while in the deeper part of the fissure there
was total loss of myelin both in the radial fibres and in

\[\text{FIG. 1. Coronal section through both occipital lobes showing concentric arrangement of rows of lacunae, particularly on the right.}\]
the stria of Gennari (Fig. 2). There was a mild marginal gliosis throughout the occipital cortex and in addition a striking proliferation of large spider astrocytes in layer VI of the visual cortex in the floor of the calcarine fissure (Fig. 3).

The distributions of lesions in the white matter corresponded accurately with the macroscopic appearances. The central white matter of the occipital lobes was completely devoid both of myelin sheaths and of axons and consisted of rows of lacunae separated by glial septa of variable thickness (Figs. 4 and 5). Some of the septa separating the lacunae of the same rows were very slender and consisted of a blood vessel and a few glial fibres (Fig. 6). Others formed somewhat wider bridges of tissue, containing blood vessels, proliferated astrocytes, glial fibres, and phagocytic microglia. Similar appearances were seen in the broader septa separating the rows of lacunae from each other. This central area of lacunar rarefaction was surrounded by a zone of demyelination with partial preservation of axons, extending deeply into the subcortical white matter and encroaching in places upon the cortex as noted above. While axons were present in these demyelinated zones their density was less than normal and decreased rapidly towards the central area of cavitation. Astrocytic proliferation and fibrillary gliosis were prominent and phagocytic microglia abundant. Many blood vessels were surrounded by thick cuffs of mononuclear cells, predominantly of microglial origin. Similar appearances were observed in the splenium of the corpus callosum and the adjacent white matter of the cingulate gyrus.

The deep structures of the hemispheres were normal with the exception of the lateral geniculate bodies, which showed severe atrophy with almost total loss of neurones.

The optic pathway showed patchy demyelination in the optic nerves and chiasm. The intra-orbital part of the right optic nerve showed total loss of myelin and severe, but not complete, loss of axons (Fig. 7). The nerve appeared very cellular owing to astrocytic and microglial proliferation. A dense network of fine glial fibrils was present throughout the nerve. The left optic nerve appeared slender, the cords of myelinated fibres were thinner than normal and the fibrous septa crowded together (Fig. 8). Despite this appearance of mild atrophy the density of axons and their myelination seemed normal and fibrillary gliosis minimal. The chiasm showed massive demyelination of the decussation in continuation with the plaque in the right optic nerve (Fig. 9). There was also a small irregular plaque at the left lateral margin of the chiasm.

In the spinal cord the lesions became apparent at the level C4 where a central area of recent necrosis with breakdown of axons and myelin sheaths occupied mainly the deeper parts of the posterior columns (Fig. 10). Caudally, the lesion increased rapidly in size and at C8 occupied almost the entire cross-section of the cord with the exception of a thin peripheral rim of preserved myelinated fibres (Fig. 11). Amid this massive breakdown of the white matter the grey matter appeared relatively well preserved and the neurones of the anterior and posterior horns were apparently unaffected. The lesion reached its maximum intensity at T4 where destruction of the cord was total and apparently of longer duration than in the cervical segments (Fig. 12). There was diffuse proliferation of phagocytic microglia and patchy infiltration by polymorphonuclear leucocytes, but no

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**Fig. 2.** Right visual cortex: myelin is preserved only in the tips of the gyri while the deeper parts are totally demyelinated. Klüver and Barrera's luxol fast blue and cresyl violet, × 7.

**Fig. 3.** Right occipital cortex: proliferation of spider astrocytes in floor of calcarine fissure adjacent to glial wall of medial row of lacunae. Holzer's method for glial fibres, × 150.
FIG. 4. Section of right occipital lobe stained for myelin showing extent of demyelination and lacunar rarefaction. Klüver-Barrera, $\times 1.8$.

FIG. 5. Adjacent section stained for glial fibres: the concentric arrangement of glial septa is clearly seen. Holzer, $\times 1.8$.

FIG. 6. Right occipital lobe: groups of thin-walled lacunae separated by thick glial septum. Holzer, $\times 45$. 

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FIG. 7. Right optic nerve: total loss of myelin and increased cellularity due to proliferation of glial cells. Klüver-Barrera, × 30.

astrocytic reaction. Similar appearances were seen in lower segments of the thoracic cord down to T10, but with gradual reappearance of small islands of myelinated fibres. At T12 the necrosis was reduced to a central core involving the base of the posterior columns. This continued as a thin tail of circumscribed necrosis in the left posterior column down to the level L5 (Fig. 13).

DISCUSSION

The clinical diagnosis in this case presented considerable difficulty. In retrospect, when the full records were brought together from the five hospitals in which the patient had been investigated, the evolution of the disease and the distribution of the lesions might have suggested a demyelinating disease, possibly neuromyelitis optica. The unusually advanced age should not have precluded the diagnosis, as the incidence of this disease covers a span from 5 to over 60 years (McAlpine, 1938).

The clinicopathological concept of neuroopticomyelitis is not entirely clear. The clinical picture was fully reviewed by Stansbury (1949) on a material consisting of 200 reported cases. The syndrome consists of blindness, temporary or permanent, followed after a variable, though usually short, interval by paraplegia of rapid onset. The disease is progressive and often rapidly fatal, but cases running a remitting and relapsing course have been reported. Pathologically the lesions fall into three groups (Greenfield and Norman, 1963): some are examples of multiple sclerosis involving predominantly the optic pathways and the spinal cord, others resemble


FIG. 9. Optic chiasm: plaque of demyelination extending from right optic nerve into decussation; also small plaque at base of left optic nerve. Klüver-Barrera, × 7.

acute perivenous encephalomyelitis of similar distribution, others still consist of demyelinating lesions in the optic pathways associated with acute necrotic myelopathy. Hughes (1966) suggested that only the last group formed a separate entity for which the term neuromyelitis optica should be retained.

The association of necrotizing myelopathy, indistinguishable from the isolated myelomalacia described originally by Bassoe and Hassin (1921) and fully reviewed by Hughes (1961), with asymmetrical plaques of demyelination in the optic nerves clearly places our case in this last group. The unusual finding was the lesion affecting both occipital lobes and the splenium of the corpus callosum and closely resembling the concentric lacunar leukencephalopathy of Grcević (1960). Demyelinating lesions in the occipital lobes have been reported in several cases of Devic’s disease (Marinesco, Draganesco, Sager, and Grigoresco, 1930; Greenfield, 1950; Paarmann, 1952). The case of Marinesco et al. is of particular interest in that the extensive, though strictly unilateral, parieto-occipital lesion consisted of an outer demyelinated and inner lacunar zone. The lacunae, however, were not arranged concentrically, and the intervening tissue contained demyelinated axons as well as glial fibres. The significance of the concentric pattern and its possible relationship to the ‘encephalitis periaxialis concentrica’ of Baló (1928) was fully discussed by Grcević and our case throws no new light on this.


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problem. The essential difference between lacunar encephalopathy and Baló's sclerosis lies in the fact that in the latter zones of demyelination are separated by lamellae of preserved white matter, in the lacunar type zones of cavitation alternate with glial septa devoid of axons and myelin sheaths.

Our case differs from that of Grcevic mainly in the distribution of the lesions which in his affected the frontal lobes and the anterior third of the corpus callosum. The optic nerves and spinal cord were not examined. While there was no clinical suggestion of a myelopathy the history of recurrent episodes of loss of vision progressing towards almost total blindness suggested involvement of the optic pathways presumably by a demyelinating process. Nevertheless Grcevic considered that the available evidence did not warrant the inclusion of his case in any definite nosological group.

We feel that our case clarifies the nosological position of concentric lacunar leucoencephalopathy. The association with typical neuropticomyelitis places this lesion within the context of demyelinating diseases.

SUMMARY

A man aged 64 presented with left homonymous hemianopia rapidly progressing to total cortical blindness. This was followed by a transient left hemiparesis and grand mal seizures. Fourteen months later he developed a rapidly progressive ascending myelopathy to which he succumbed. Post-mortem examination of the nervous system revealed plaques of demyelination in the optic nerves and chiasm, an acute necrotizing myelopathy, and bilateral lesions in the occipital white matter identical with those described by Grcevic under the name of concentric lacunar leucoencephalopathy. The association of this condition with typical neuropticomyelitis suggests its close relationship to the demyelinating diseases.

We are indebted for the excellent documentation of this case, and for permission to publish it, to Dr. R. A. Henson and Mr. T. T. King, and also to Dr. F. Lees, Mr. B. Fairburn, and Lt. Col. A. V. Forage.

REFERENCES