Involvement of the central nervous system in chronic inflammatory demyelinating polyneuropathy: a clinical, electrophysiological and magnetic resonance imaging study

I E C Ormerod, H M Waddy, A G Kermode, N M F Murray, P K Thomas

Abstract

In a consecutive series of 30 patients with chronic inflammatory demyelinating polyneuropathy (CIDP) minor clinical evidence of CNS involvement was found in five. Cranial magnetic resonance imaging (MRI) was performed in 28 and revealed abnormalities consistent with demyelination in nine patients aged less than 50 years and abnormalities in five aged 50 years or over. Measurements of central motor conduction time (CMCT) were obtained in 18 and showed unilateral or bilateral abnormalities in six. It is concluded that subclinical evidence of central nervous system (CNS) involvement is common, at least in patients with CIDP in the United Kingdom, but that clinically evident signs of CNS disease are infrequent. The association of a multiple sclerosis-like syndrome with CIDP is rare.

Chronic inflammatory demyelinating polyneuropathy (CIDP) has close clinical and pathological similarities to the Guillain-Barré syndrome but pursues a chronic progressive or chronic relapsing course.1-4 In recent years a number of patients have been described with a multifocal central nervous system (CNS) disorder, the clinical features of which resemble multiple sclerosis (MS),5,6 in association with CIDP. The frequency of this association is not known. The occurrence of CNS lesions was not noted in any of the series of 53 cases of CIDP reported by Dyck et al3 or the series of 92 documented by McCombe et al.5 All six of the selected patients with combined peripheral nervous system (PNS)/CNS demyelination described by Thomas et al5 had CNS lesions demonstrated by magnetic resonance imaging (MRI). In a group of 16 patients with CIDP, Mendell et al6 found periventricular, subcortical or brainstem white matter lesions in six, of whom three had clinical or laboratory abnormalities considered to indicate MS. This series, however, was partially selected.8

In this study we have examined a consecutive unselected series of 30 patients, 28 of whom consented to MRI. Studies of central motor conduction time (CMCT) were also performed as CMCT was found to be abnormal in all four patients with CIDP and multifocal CNS demyelination reported by Thomas et al in which this was assessed.9

Patients and methods

Thirty consecutive patients with CIDP referred to one of us (PKT) formed the basis of this study (see table); of these, 28 consented to MRI studies. The diagnosis of CIDP was made according to the criteria of Dyck et al.3 This was based on the history, the clinical findings, nerve conduction studies and examination of the cerebrospinal fluid (CSF). Nerve biopsy was performed in 18. The age of the patients at the time of the study was 9-83 (mean 37-4) years; 14 were male and 16 female. The series will be biased against childhood cases as the majority were derived from the National Hospital for Neurology and Neurosurgery where paediatric cases are under-represented.

Patients with a chronic demyelinating polyneuropathy associated with a paraproteinaemia or leukodystrophy or who had other possible causes for neuropathy, such as diabetes mellitus, were excluded.

Cranial MRI was performed using a 0-5 Tesla Picker MR imaging system. Multislice T2-weighted spin echo images (SE 2000/60) were obtained with a slice thickness of 5 mm. The images were subjected to blind assessment by two of the authors (IECO and AGK) independently and only those images considered to be abnormal by both assessors were accepted as such. The images were evaluated for the number, pattern and distribution of abnormalities.

CMCT was measured with surface recording from the abductor digiti minimi muscle and magnetic stimulation of the motor cortex as previously described.9 The proximal motor roots were stimulated electrically in the region of the intervertebral foramina by high voltage (300-400 V) shocks applied transcutaneously over the cervical spine at the C7/T1 interspace and the ulnar nerve was stimulated supramaximally at the wrist. CMCT was obtained by subtraction of the latency of the response evoked by cervical stimulation from that following activation of the cortex. The upper limit of normal was taken as 8-3 ms (mean CMCT 6-2 ms, SD 0-86 ms in 46 sides of 32 subjects, aged 21-78 years). This value contains a small peripheral component estimated,
in the normal subject, to be about 0-4 ms from conduction in the roots and therefore correction of normative data is necessary where there is slowing of peripheral motor conduction. Proximal root conduction time was calculated as 1-0 ms for motor conduction velocity (neck-wrist) of 30 ms⁻¹ and 1-5 ms for 20 ms⁻¹ giving upper limits of CMCT of 8-9 ms and 9-4 ms respectively.

**Results**

**Clinical features**

The duration of illness ranged from 1 to 22 (mean ± SD: 10) years. The clinical course was relapsing and remitting in 17, and chronic progressive in 13. In three of the latter, spontaneous improvement occurred with a monophasic course. The mean age of onset in the relapsing cases was 25-1 years and 43-7 years in the chronic progressive cases (excluding the three patients with a monophasic course with spontaneous improvement).

Fourteen presented with weakness, four with combined weakness and sensory loss, nine with sensory disturbance and one with dysphonia. At some stage in the course of the illness, distal limb weakness was present in all and sensory disturbance was present in 20. Spontaneous pain was a complaint in 10 and upper limb postural tremor was observed in nine patients. Abnormalities of the cranial nerves were encountered in 16. Enlargement of peripheral nerves was noted only in three patients.

Evidence of involvement of the CNS was obtained in five (possibly six) cases: two had extensor plantar responses (Cases 1, 4; see table), one had an acute brainstem disturbance (Case 7; case 2, Waddy et al), one showed downbeat nystagmus (Case 27) and one had sensory loss in the legs and lower trunk with an upper sensory level (Case 20). A further patient had generally exaggerated tendon reflexes but no other evidence of corticospinal tract damage (Case 3).

A possible precipitating event in association with one or more clinical episodes was noted in nine patients. These were smallpox vaccinations in two, pregnancy in two and nonspecific infections in the other five.

Nerve conduction studies were performed in all 30 patients. Motor conduction velocities were reduced in all (table) with values of 25 ms⁻¹ or less in 11. The slowest value recorded was 7 ms⁻¹. The cerebrospinal fluid was examined in 25 patients. The CSF protein content was elevated (>0-5 g/l) in 17 (table). Oligoclonal IgG bands were present in four patients. Only one of these had clinical evidence of CNS involvement (Case 1, table).

Nerve biopsies were abnormal in the 18 cases in which they were obtained and showed a mixed pattern of demyelination and axonal loss. Minor hypertrophic changes were frequent but prominent "onion bulb" changes were seen in only two patients. Focal infiltrates of inflammatory cells were present in only two cases, but occasional mononuclear cells (monocytes and lymphocytes) were observed in the endoneurium in a number of the patients.

**Magnetic resonance imaging**

Fourteen out of the 28 patients had an abnormal MRI scan; of these, nine were aged 49 years or less. In the five older patients there were multiple discrete white matter lesions and irregular periventricular lesions in two (Cases 4, 11), smooth periventricular changes with multiple discrete cerebral white matter lesions in two (Cases 12, 27) and only discrete cerebral

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**Table Summary of clinical, MRI and CMCT data**

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*The numbering is not in the order of admission to the study.

**MNCV =** motor nerve conduction velocity, upper or lower limbs; P = chronic progressive; R = relapsing; M = monophasic; + = present (for oligoclonal CSF IgG, and clinical CNS involvement) or abnormal (for MRI and CMCT); ND = not done.
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Nine of the 14 patients with abnormal cranial MRI had cranial nerve lesions at some time during their illness, compared with eight out of 14 with normal MRI. Clinical evidence of CNS involvement was seen in four of the 14 with abnormal MRI but only in one of the 14 with normal MRI.

Oligoclonal IgG bands were detected in the CSF in three out of the 14 patients with abnormal MRI findings but only in one of 14 with normal MRI.

Central motor conduction time
An attempt was made to measure CMCT in 23 of the 30 patients. The technique employed requires the ability to record a reproducible evoked muscle action potential from abductor digitii minimi on ulnar nerve stimulation. In four patients the degree of conduction block or denervation, or both, was such that this was not possible and CMCT therefore could not be assessed. Recordings were accordingly obtained from 18 patients and of these, six were abnormal. The most common abnormality was a moderate prolongation of latency (fig 2) which was unilateral in Cases 3, 11, 16 and 21 and bilateral in Cases 9 and 23. The amplitude of the evoked muscle action potential following magnetic brain stimulation was not a helpful parameter. The potential was frequently dispersed and of low amplitude, but identical in morphology to that obtained from cervical root stimulation and therefore reflecting the peripheral conduction abnormality (fig 3).

When the CMCT results were compared with the MRI findings, seven of the 14 patients with abnormal MRI also had CMCT recordings; of these, three were abnormal. Of the 14 patients with normal MRI, 11 had CMCT recordings of which three were abnormal. In the five patients with clinical evidence of CNS disease, a normal result was obtained in both of the two cases in which CMCT was measured.

Discussion
The clinical features in this series of patients with CIDP are similar to those in other reported series. A relapsing course was seen in 57%, which is slightly higher than the figure of 34% found by Dyck and Arnason but close to the value of 65% given by McCombe et al. The age of onset in our patients was again earlier in...
the relapsing cases (25-1 years) than in those with a chronic progressive course (43-7 years). McCombe et al reported mean ages of onset of 26-8 and 51-4 years respectively. Possible precipitating factors were identified in 44% by McCombe et al, which is slightly higher than the present series (32%).

The main purpose of this study was to assess the frequency of CNS involvement in CIDP. No cases with florid evidence of this, similar to those described by Thomas et al and Rubin et al were found. Minor CNS signs were encountered in five and were possibly present in one further case. In our series, cranial nerve involvement was found in 60%, which is higher than in previous series. It was not always possible on clinical grounds to be certain whether this was the result of brainstem or extra-axial involvement. The occurrence of focal cranial nerve lesions in CIDP related both to CNS and peripheral lesions has been reported. MRI abnormalities were no commoner in the present patients who had cranial nerve involvement than in those who did not.

Mendell et al, Pakanlis et al and Barohn et al found that six out of 18 patients had abnormalities on MRI and that four of these also had clinical evidence of disease. However, four of the six were over 50 years of age and three over 70. In view of the high incidence of MRI abnormalities in older patients, these findings have to be interpreted with caution.

The nature and distribution of the MRI abnormalities found in the present series is of some importance. In six out of nine patients aged under 50 years there were irregular periventricular lucencies as well as other lesions within the brain. These appearances, although nonspecific, are indistinguishable from those seen in MS. In two out of these nine patients there were no periventricular lesions. This pattern of abnormality, which is even less specific, may also be encountered in MS. Of the older patients with abnormal MRI examinations, two out of five had smooth periventricular lesions of a type more frequently associated with vascular change than demyelination. This type of abnormality, sometimes combined with discrete lesions elsewhere in the brain, is also seen in some normal subjects and is of increasing frequency with age. Nevertheless, there was clinical evidence of CNS involvement in four of the patients with abnormal MRI findings but only in one of those without detectable abnormalities.

MRI and CMCT have both been available for some years, but this is the first time that they have been applied together systematically for the investigation of CNS involvement in CIDP. In our series, although some patients with MRI abnormalities also had delayed CMCT, there was not complete concordance between these techniques. This is not surprising as the two investigations demonstrate qualitatively different abnormalities within the nervous system. Furthermore, there is only a partial overlap in the anatomical territory encompassed by the two techniques. MRI demonstrates lesions where there are changes in the amount of tissue water and also alterations in the physical behaviour of water protons. Some of the lesions within the brain seen on MRI may be demyelinated mature plaques where myelin has been replaced by astrocytes, with relative preservation of the axons and a local increase in water content. Recent studies of serial MRI examinations in patients with established MS have shown that some lesions appear only briefly and may be undetectable on MRI within a matter of weeks. The pathological correlates of the evanescent lesions are not established but there is some evidence that the earliest change in the brain in the evolution of a lesion is a defect in the blood-brain barrier. Subsequent to the defect there is an increase in the tissue water content which may be partially extracellular and which in some instances appears to resolve without a detectable glial plaque. It is not known to what extent the various stages of evolution of such lesions will interfere with axonal function. Even in an established plaque there may be significant function retained by the residual axons. Abnormalities of CMCT demonstrate a functional disturbance in the central motor pathways. This technique has the additional advantage of including the cervical cord but of course will not demonstrate lesions within the brain which are outside the motor pathways. An abnormality of CMCT for corticospinal fibres to the T1 cord segment was detected in eight of the 19 patients from whom recordings were obtained.

Function in central pathways can also be examined by evoked potential studies. Visual (VEP) and brainstem auditory evoked potentials (BSAEP) were abnormal in all six of the cases of CIDP with multifocal CNS demyelination reported by Thomas et al. Somatosensory evoked potentials (SEP), when they could be recorded, were also abnormal, sometimes with very considerably prolonged latencies. VEP and BSAEP abnormalities in patients with CIDP have subsequently been reported by Pakanlis et al and Gigli et al. From the study and the previous reports employing MRI and evoked potential recordings, it can be concluded that although subclinical CNS abnormalities are common in CIDP and minor clinical abnormalities are observable in a small proportion, concurrent severe CNS disease is rare. Conversely, there have been reports dating back for many years of a demyelinating neuropathy affecting spinal roots or more widely in the PNS in patients with a primary diagnosis of MS. It is not yet clear whether the CNS lesions that may occur in patients with CIDP are morphologically similar to those of MS, but the severity of prolongation of SEP latencies indicates that they are demyelinating in nature. The temporal coincidence of the CNS and peripheral nerve disorder favours the operation of a single disease process or the existence of shared trigger mechanisms.
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8 Mendell JR. Personal communication. 1988.
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