Cerebellar learning distinguishes inflammatory neuropathy with and without tremor

ABSTRACT

Objectives: This study aims to investigate if patients with inflammatory neuropathies and tremor have evidence of dysfunction in the cerebellum and interactions in sensorimotor cortex compared to nontremulous patients and healthy controls.

Methods: A prospective data collection study investigating patients with inflammatory neuropathy and tremor, patients with inflammatory neuropathy without tremor, and healthy controls on a test of cerebellar associative learning (eyeblink classical conditioning), a test of sensorimotor integration (short afferent inhibition), and a test of associative plasticity (paired associative stimulation). We also recorded tremor in the arms using accelerometry and surface EMG.

Results: We found impaired responses to eyeblink classical conditioning and paired associative stimulation in patients with neuropathy and tremor compared with neuropathy patients without tremor and healthy controls. Short afferent inhibition was normal in all groups.

Conclusions: Our data strongly suggest impairment of cerebellar function is linked to the production of tremor in patients with inflammatory neuropathy.

GLOSSARY

ADM = abductor digiti minimi; ANOVA = analysis of variance; APB = abductor pollicis brevis; CIDP = chronic inflammatory demyelinating polyradiculoneuropathy; CR = conditioned blink responses; EBCC = eyeblink classical conditioning; FDI = first dorsal interossei; IgMPN = immunoglobulin M paraproteinaemic neuropathy; MEP = motor evoked potential; MMNCB = multifocal motor neuropathy with conduction block; MRC = Medical Research Council; ONLS = Overall Neuropathy Limitation Scale; PAS = paired associative stimulation; PF = peak tremor frequency; SAI = short afferent inhibition; TMS = transcranial magnetic stimulation; TP = total power of the spectra between 1 and 30 Hz used as surrogate measure of tremor amplitude; US = unconditioned stimulus; WE = wrist extensor muscles.

Inflammatory mediated neuropathies are common and potentially treatable.1 Tremor occurs with immunoglobulin M paraproteinaemic neuropathy (IgMPN)2-4 and less commonly in other inflammatory neuropathies.5 It has been suggested that temporally distorted peripheral inputs reach a normally functioning central processor, such as the cerebellum, which is misled into producing a delayed second agonist burst and tremor.6-8 The involvement of the cerebellum in neuropathic tremor is supported by functional imaging abnormalities.9 There does not seem to be a straightforward relationship between the development of tremor and conduction velocity.10 Further, no relationship seems to exist between tremor and the severity of neuropathy as assessed by proprioceptive loss, weakness, or fatigue.11,12 However, we have shown that although conduction velocity does not predict the presence of tremor, it is correlated with its severity for those in whom tremor is present.5 This indicates a second mechanism may be necessary to produce tremor.

Here we set out to explore aspects of CNS physiology in tremulous and nontremulous patients with inflammatory neuropathies compared to healthy controls. We hypothesized that the central compensation needed to account for delays caused by the peripheral neuropathy would most likely depend on plastic changes within the cerebellum and connections that mediate interaction between sensory and motor systems and therefore that patients with tremor would have evidence...
of dysfunction in the cerebellum and interactions in sensorimotor cortex compared to non-tremulous patients and controls.

**METHODS Subjects.** Eighteen out of 43 consecutive patients published recently with a diagnosis of inflammatory neuropathy (chronic inflammatory demyelinating polyradiculoneuropathy [CIDP], multifocal motor neuropathy with conduction block [MMNCB], or IgMPN) agreed to take part in all or just parts of the study. The latter depended on contraindications to electrical/magnetic stimulation and on the cumulative length of study sessions.

Patients were divided into tremulous and non-tremulous depending on whether arm tremor was clinically detectable. The Fahn-Tolosa-Marin scale, a summed Medical Research Council score (MRC score; maximum 70), a sensory score (maximum 56), and the Overall Neuropathy Limitation Scale (ONLS; maximum 12) were performed.

Ten tremulous patients (mean age 60.0 [7.7] years; mean disease duration 12.5 [8.2] years; total sensory score 41.7 [13.8]; total MRC score 65.3 [4.2]; ONLS score 3.6 [1.3]) were studied. They were compared with 8 non-tremulous patients who did not differ in these characteristics (mean age 63.3 [8.1] years; p = 0.46; mean disease duration 14.1 [10.6] years; p = 0.72; total sensory score 42.0 [16.1]; p = 0.97; total MRC score 63.2 [9.0]; p = 0.59; ONLS score 4.2 [1.2]; p = 0.38) (table 1). We also recruited 9 healthy age-matched controls (mean age 59.0 [7.7] years; p = 0.54).

**Table 1 Demographics and clinical characteristics**

<table>
<thead>
<tr>
<th>Age, y</th>
<th>Sex</th>
<th>Disease</th>
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<th>FTM score</th>
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<td>-</td>
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Abbreviations: FTM = Fahn-Tolosa-Marin total score (0 [minimum] to 4 points [maximum severity]) are assigned for tremor amplitude under a variety of conditions and 0–4 points for severity in daily activities; group NT = nontremulous; group T = tremulous; study E = eyeblink classical conditioning; study P = paired associative stimulation; study S = short afferent inhibition; study T = tremor analysis.

Standard protocol approvals, registrations, and patient consents. Before inclusion in the study, written informed consent was obtained from all participants. This study was approved by the local Research Ethics Committee.

**Electrophysiologic evaluation.** Surface EMG recordings were made with Ag-AgCl surface electrodes using a belly-tendon montage. Data were stored in a computer for display and off-line analysis using Signal version 4.00 (and Spike version 2 for tremor analyses).

**Accelerometry and EMG for tremor.** Nine patients with tremor (5 CIDP, 2 MMNCB, 2 IgMPN) took part in this evaluation. A triaxial accelerometer transducer (sensitivity ≥ 100 mV/G) was attached to the dorsal surface of the middle phalanges of the index fingers. EMG recordings were made of wrist extensor muscles (WE), wrist flexors, abductor pollicis brevis (APB), and biceps brachii bilaterally. Recordings were performed 1) with arms relaxed (rest), 2) with arms/wrists outstretched at shoulder level (posture), and 3) with a 500-g mass attached to the wrists (loading), and 4) while performing a goal-directed task (action). Accelerometry and EMG were recorded and analyzed for 30 seconds in each condition.

**Blink reflex and eyeblink classical conditioning.** Three age-matched groups were examined: 9 healthy controls, 7 non-tremulous patients (4 CIDP, 1 MMNCB, 2 IgMPN), 9 tremulous patients (6 CIDP, 2 MMNCB, 1 IgMPN). Tremulous and nontremulous patients did not differ in age (p = 0.97), disease duration (p = 0.59), total sensory score (p = 0.72), MRC score (p = 0.63), or ONLS score (p = 0.26).

Blink reflex and R2 blink reflex recovery cycle were assessed in all subjects according to a previously described protocol. The conditioning stimulus (CS) was a loud (50 dB above auditory threshold) 2,000 Hz tone lasting 400 ms played via binaural headphones. The CS consistently produced an acoustic startle response (“alpha blink”) occurring within 200 ms after the CS. An electrical stimulus (unconditioned stimulus [US]; 200 μs pulse width at 5× sensory threshold) was given to the left supraorbital nerve 400 ms after the CS, eliciting a blink reflex (unconditioned response).

Repeated pairs of CS and US at 400 ms intervals yield conditioned blink responses (CR) occurring within 200 ms before the US (see figure e-1 on the Neurology® Web site at www.neurology.org). EMG was recorded bilaterally from orbicularis oculi. Conditioning consisted of 7 acquisition blocks (such consisting of 9 CS-US pairs; 1 US only; 1 CS only trial). An eighth and ninth block consisted of 11 CS-only trials to measure extinction.

**Short afferent inhibition and paired associative stimulation.** Both short afferent inhibition (SAI) and paired associative stimulation (PAS) rely on precisely timed interactions between sensory afferents and motor cortical stimulation. In healthy subjects, these interactions occur at specific times related to the N20 response. We expected N20 responses to be delayed in our patients and therefore we evaluated N20 latency in each subject. One patient had to be excluded because N20 could not be identified. N20 could be measured in all other subjects studied (expressed as mean [SD]; healthy controls 20.3 [1.5]; neuropathic tremor 33.8 [11.5]; no tremor 32.6 [6.6]).

EMG recordings were made from the abductor APB, first dorsal interossei (FDI), and abductor digitii minimi (ADM) muscles of the right side. Test responses in the target muscles were evoked by transcranial magnetic stimulation (TMS) of the left primary motor cortex applied through Magstim 200 magnetic stimulators with a monophasic current waveform, connected to a figure of 8 coil. Standard techniques were used to identify the
motor “hot spot” and resting motor thresholds.13 Electrical stimulation was applied to the median nerve at the wrist at 500% of perceptual threshold using a constant current generator. The stimulus duration was 0.2 ms.

**Short afferent inhibition.** Three age-matched groups were examined, 6 healthy controls, 5 nontremulous patients (4 CIDP, 1 IgMPN), and 6 tremulous patients (4 CIDP, 2 IgMPN). Tremulous and nontremulous patients did not differ regarding age (\(p = 0.84\)), disease duration (\(p = 0.82\)), total sensory score (\(p = 0.63\)), MRC score (\(p = 0.90\)), or ONLS score (\(p = 0.44\)).

SAL was assessed as previously described.20 We assessed the response to a cortical stimulus alone and when preceded by conditioning stimuli at 10 interstimulus intervals in reference to subjects’ N20: –18 ms, –4 ms, –2 ms, 0 ms, +2 ms, +4 ms, +6 ms, +8 ms, +10 ms, +18 ms. Comparison of responses between groups was based on motor evoked potential (MEP) area.

**Paired associative stimulation.** Three age-matched groups were studied, including 6 healthy controls, 5 nontremulous patients (5 CIDP), and 8 tremulous patients (6 CIDP, 2 IgMPN). Tremulous and nontremulous patients did not differ regarding age (\(p = 0.61\)), disease duration (\(p = 0.72\)), total sensory score (\(p = 0.62\)), MRC score (\(p = 0.90\)), or ONLS score (\(p = 0.83\)).

A conditioning median nerve electrical stimulus was given 5 ms plus individual N20 (i.e., 25 ms if N20 latency was 20 ms) before a TMS pulse over the APB muscle “hot spot” at an intensity predetermined to yield a –1 mV resting MEP. Two hundred paired stimuli were delivered at a rate of 0.25 Hz.21 Thirty MEPs were recorded before, immediately after, 15 minutes and 30 minutes after PAS. Comparison of PAS response was based on MEP area.

**Data analysis and statistics.** A Fourier analysis of signals derived from accelerometer was performed to define peak tremor frequency (PF). Total power of the spectra between 1 and 30 Hz was used as surrogate measure of tremor amplitude (TP). All parameters were calculated for each accelerometer axis, and then averaged. For EMG, the signal was full-wave rectified and smoothed and Fourier analysis was performed to derive PF.

For measurement of eyeblink conditioning, CRs were counted manually. EMG bursts were regarded as “alpha blinks” if their amplitude exceeded 50 \(\mu\)V and if latency was <200 ms after the CS. EMG bursts were regarded as CRs if latency was >200 ms after the CS but before the US. For the CS only trials, EMG bursts occurring 200–600 ms after the CS were considered CRs. Statistical analysis was performed using PASW Statistics 18 (SPSS; Quarry Bay, Hong Kong). All post hoc comparisons were corrected by the Bonferroni method. The level of statistical significance was pre-set at \(p < 0.05\).

**RESULTS Tremor recordings.** In all 9 patients, a bilateral tremor was recorded during posture and action. Five patients had additional bilateral rest tremor. The power spectra of accelerometry and WE EMG showed corresponding peaks. Since there was no side-to-side difference in PF or TP in any position \((p > 0.3)\), we used the mean of both sides for PF and TP for further analyses. Mean PF and TP in the 4 recorded conditions are shown in table 2.

To compare PF and TP measured by accelerometry at rest, posture, and action, we computed 2 repeated-measures analyses of variance (ANOVA). For PF, there was no effect of condition \((F_{2,16} = 3.47; p = 0.06)\). For TP, there was an effect for condition \((F_{1,8} = 6.76; p = 0.03)\); however, post hoc comparisons showed no differences \((p > 0.09)\).

A \(t\) test for pairwise comparisons showed no difference in PF (accelerometry, WE EMG) before and after loading \((p > 0.2)\), indicating that loading did not decrease tremor frequency. Five out of 9 patients had an increase of tremor amplitude after loading by at least 100%. However, there was no difference regarding TP before and after loading \((p = 0.33)\) on group level.

In 3 out of 9 patients \((2\) with IgMPN, 1 with MMN), PF in the APB was more than 1 Hz lower compared to the biceps. However, a paired \(t\) test comparing PF during posture in biceps and APB in the whole group of patients showed no difference \((p = 0.17)\).

**Blink reflex and eyeblink classical conditioning.** R2 blink reflex recovery curves, R1 and R2 latencies, and latency variability did not differ among the 3 groups.

Repeated-measures ANOVA with block (7) as within-subject factor and group (3) as between-subject factor revealed an interaction of block \(\times\) group \((F_{12,132} = 3.34; p < 0.001)\). There were also effects of block \((F_{6,132} = 12.2; p < 0.001)\) and group \((F_{2,22} = 16.6; p < 0.001)\) (figure 1). Post hoc tests showed that tremulous patients had a lower rate of CRs as the blocks progressed compared to healthy controls and nontremulous patients \((p < 0.001)\). This difference was significant in conditioning blocks 3–7 (figure 1). Latencies of CRs, spontaneous blink rates, and “alpha blinks” were not different between the groups.

**SAL.** Repeated-measures ANOVA with state (11) as within-subjects factor and group (3) as between-subjects factor revealed an effect of state \((F_{3,48} = 6.64; p < 0.001)\). There was no effect of group or the group \(\times\) state interaction. Post hoc tests showed a reduction in MEP size occurring at interstimulus intervals of N20 \((p < 0.001)\) and N20–2 \((p = 0.007)\) (figure 2).

### Table 2 Mean (SD) PF and TP (derived from accelerometer) in the 4 recorded conditions

<table>
<thead>
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<th>Condition</th>
<th>Accelerometry</th>
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</thead>
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<td></td>
<td>PF (Hz)</td>
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<tr>
<td>Rest</td>
<td>7.1 (1.6)</td>
</tr>
<tr>
<td>Posture</td>
<td>6.1 (1.6)</td>
</tr>
<tr>
<td>Weight</td>
<td>6.4 (1.4)</td>
</tr>
<tr>
<td>Action</td>
<td>5.5 (1.4)</td>
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</tbody>
</table>

Abbreviations: action = repetitive finger-to-nose movements; PF = peak frequency; posture = arms outstretched; rest = rest position; TP = total power; weight = arms outstretched with weight loading.

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Mean intensity to produce 1 mV MEPs was not different between patients (56%) and controls (61%). Repeated-measures ANOVA with time (4) and muscle (3) as within-subject factors and group (3) as between-subject factor revealed that PAS produced a lasting increase in mean MEP area demonstrated by an effect of time ($F_{2,33} = 4.762, p = 0.014$). The size of MEP facilitation differed among groups and muscles, indicated by an effect of group ($F_{2,16} = 9.890, p = 0.002$) and an interaction of time $\times$ group ($F_{6,33} = 5.166, p = 0.002$). The interaction between time $\times$ muscle $\times$ group ($F_{6,62} = 3.436, p = 0.003$) demonstrates that the effect of PAS on the homotopically (APB) and heterotopically (FDI, ADM) conditioned muscles differed time-dependently between groups (figure 3).

To further explore the conditioning effects of PAS on MEP areas in each group, we computed separate repeated-measures ANOVAs with time and muscle as within-subject factors. In controls, an effect of time ($F_{3,15} = 3.212; p = 0.047$) was found. The facilitatory effect was stronger in the APB compared to the FDI/ADM, reflected by a strong time $\times$ muscle interaction ($F_{6,30} = 7.257; p < 0.001$). In patients without tremor, a different pattern of PAS-induced changes occurred. MEP facilitation was higher and spatial specificity was compromised as indicated by a main effect of time ($F_{3,12} = 6.570; p = 0.007$) without time $\times$ muscle interaction. Patients with tremor had an effect of time ($F_{3,21} = 3.479; p = 0.034$) due to overall MEP depression without time $\times$ muscle interaction.

Post hoc comparisons revealed that PAS induced an increase in MEP areas compared to baseline in the APB in controls (T15: $p = 0.023$), but not in neuropathy with and without tremor. A facilitation of the MEP area in the ADM and FDI was only observed in patients without tremor (T15 [FDI]: $p = 0.003$; T15 [ADM]: $p = 0.036$).

**DISCUSSION** We demonstrate that patients with inflammatory neuropathy and tremor differ from patients without tremor with regard to cerebellar function and sensorimotor plasticity. We found very low rates of EBCC in patients with inflammatory neuropathy and tremor compared to nontremulous patients and healthy controls, suggesting abnormal associative learning in the cerebellum that segregates with tremor. We also describe an absence of normal facilitation in TMS-evoked EMG potentials after PAS in patients with tremor, suggesting abnormal sensorimotor cortex plasticity. In nontremulous patients, sensorimotor plasticity, demonstrated by facilitation of TMS-evoked EMG potentials after PAS, occurred in neighboring muscles but without a normal facilitatory response in the target muscle, suggesting a lack of topographic specificity of sensorimotor plasticity.

Tremor in our patients with inflammatory neuropathies was invariably present during posture and action. Five patients had additional rest tremor. When present in all 3 conditions, tremor was worst
EBCC is a form of simple associative learning that is well-studied and for which the cerebellum is both necessary and sufficient. Structural or functional impairments of the cerebellum lead to abnormalities in acquisition of this conditioned response.8,17,18,23,24 We demonstrate abnormal EBCC in tremulous neuropathy patients that clearly differentiates them from the normal rates of conditioning in nontremulous neuropathy patients and controls. Mean R1 and R2 latencies and latency variability did not differ between groups, making it unlikely that desynchronization of the afferent volley alone may be a factor in the lack of conditioned responses in the tremulous patients. The degree of impairment of acquisition of conditioned responses reported here is in line with the degree of impairment reported in patients with cerebellar degeneration or cerebellar lesions. A previous study showed a delayed second agonist burst25 in patients with IgM neuropathy and tremor, suggesting that the cerebellum, although intact, would be a likely candidate for a central processor “tricked” into generating tremor in the context of distorted mistimed peripheral signals.8 Our data instead provide evidence that the cerebellum is not functioning normally in those patients who develop tremor.

We were able to record somatosensory evoked potentials, albeit delayed, in all CIDP or IgM patients with tremor. This is in line with the assertion that tremor occurs in the presence of distorted rather than absent sensory input.8 All patients, tremulous and nontremulous, had normal SAI as compared with normal controls. This suggests that despite the peripheral sensory-motor delay due to the demyelinating neuropathies, central processes have, remarkably, been able to adapt to such delays to reset to the new latency of the N20.

In healthy subjects, PAS causes a facilitation of motor evoked potentials in the “target muscle” only, lasting for 15–30 minutes. This response shares a number of features with long-term potentiation.19 Patients with tremor showed no response to PAS. The normal SAI in patients with tremor argues against afferent dysfunction and associated changes in the sensory motor cortex as sole explanation for the abnormal PAS response. This is supported by the findings in one tremulous CIDP patient with normal N20 and absent PAS response. In recent work, we have demonstrated that cerebellar suppression in healthy subjects by transcranial direct current stimulation impairs subsequent motor cortical facilitation by PAS.26 We therefore speculate that the absent PAS response in tremulous neuropathy cortical stimulation may reflect cerebellar dysfunction that is also responsible for their impaired EBCC.

In patients without tremor, PAS response was also abnormal. Facilitatory changes were seen but these occurred in neighboring ulnar-innervated muscles.
but not in the APB. This latter finding has not, to our knowledge, previously been described in any other group of subjects. It is conceivable that altered topographic representation triggered by the neuropathy may affect sensory-motor integration required to mediate changes associated with PAS.\textsuperscript{27,28} An additional speculation is that this unusual response to PAS may be explained by a peripheral phenomenon such as ephaptic transmission between peripheral nerve fibers.

We present evidence that tremor in patients with inflammatory neuropathy is associated with cerebellar dysfunction. We acknowledge that generalizability is limited by our relatively small sample size. Also, this study does not answer the question whether the cerebellar abnormalities in tremulous patients are secondary to the presence of tremor or primary. Regarding the latter, one possibility is that in those with tremor, the specific antibody involved in causing the peripheral neuropathy is capable of crossing the blood–brain barrier and binding to the cerebellum. There is indirect evidence for this in IgMPN, in which tremor is typical. It would be of interest to look for evidence of antibodies that bind to cerebellum in tremulous patients with CIDP: they may share a common causative antibody for their neuropathy and the cerebellar dysfunction that drives the development of tremor.

**AUTHOR CONTRIBUTIONS**


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**DISCLOSURE**

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