Modulation of primary orthostatic tremor by magnetic stimulation over the motor cortex

C H Tsai, J G Semmler, T E Kimber, G Thickbroom, R Stell, F L Mastaglia, P D Thompson

Abstract

Objectives—To study the role of cortico-motor neuronal pathways in primary orthostatic tremor.

Methods—Transcranial magnetic stimuli at an intensity 10% above the resting motor threshold were delivered over the leg motor cortex in two patients with primary orthostatic tremor while standing still. Electromyographic responses in both tibialis anterior muscles were recorded after 20 stimuli given randomly at intervals of 120 to 180 seconds. Differences between predicted and actual times of occurrence of tremor bursts after the stimuli were used to calculate a resetting index, with a value of 0 representing no resetting and a value of 1 representing complete resetting.

Results—Transcranial magnetic stimulation evoked EMG responses in both tibialis anterior muscles, followed by transient suppression of tremor before reappearance of rhythmic EMG activity. Analysis of the timing of tremor bursts from EMG recordings before and after the magnetic stimuli disclosed that the phase of orthostatic tremor could be reset by brain stimulation (mean resetting indices 0.93 and 0.82).

Conclusion—The results suggest that a central oscillator, involving the motor cortex, has a crucial role in either the generation or modulation of orthostatic tremor.

Keywords: orthostatic tremor; tremor resetting; transcranial magnetic stimulation

Patients and methods

Patients

Two female patients aged 58 and 68 years with a typical history and physical signs of orthostatic tremor of six and 30 years duration respectively were studied. Both patients gave informed consent before the study. Electromyographic recordings of tremor in leg muscles disclosed 16 Hz tremor during standing. Tremor was associated with unsteadiness. No other neurological abnormalities were evident.

Methods

Surface EMG recordings from both tibialis anterior muscles were made using silver/silver chloride electrodes taped 3 cm apart over the muscles. The EMG signals were amplified (MacLab/8 (AD Instruments, NSW, Australia)) bandpass filtered (10 Hz to 2.5 kHz), digitised (sampling rate 1 kHz per channel), and stored on computer. EMG activity was recorded for one second before and 1.5 seconds after the transcranial magnetic stimulus. Each trial was stored on computer and later retrieved, full wave rectified, and digitally smoothed for off line measurement.

The motor cortex was stimulated using a MAGSTIM 200 (Whitland, Dyfed UK). A double cone coil (inner diameter 9.6 cm; outer diameter 12.5 cm), designed for stimulation of the leg area of the motor cortex, was placed over the vertex. The stimulus intensity was set at 10% of stimulator output above the threshold for eliciting EMG responses in relaxed tibialis anterior muscles. Cortical stimuli were given randomly at intervals of 120 to 180 seconds. Patients were stimulated while standing still, in a position resulting in typical tremor of the legs. Between stimuli, they were seated comfortably.

The rectified tibialis anterior EMG of each trial was averaged by computer (fig 1). Because magnetic stimuli were given at random times within the tremor cycle, this had the effect of “averaging out” the EMG bursts preceding the stimulus, so producing a relatively flat average rectified EMG trace of prestimulus muscle activity. If magnetic stimuli had no effect on the tremor, there would be a level trace in the period after stimulation as well. However, if the timing of EMG bursts in the tibialis anterior muscle was modulated in a consistent manner (for example, phase reset) by magnetic brain stimuli, then the average rectified EMG trace after stimulation will show such modulation.

To quantify the effect of magnetic brain stimuli on tremor, we employed the “resetting index” (RI), which was calculated in the following manner. From the single raw EMG
traces, the average period between tremor bursts (average cycle length) was calculated for the 10 EMG bursts preceding the magnetic stimulus. The timing of the magnetic stimulus relative to the timing of the last EMG burst before the magnetic stimulus was then expressed as a proportion of the average cycle length (% cycle length). Predicted timings for the subsequent 10 tremor bursts (had there not been a magnetic stimulus) were then calculated based on the timing of the last burst before the stimulus and the average cycle length. The actual timing of 10 tremor bursts after stimulation was measured. A graph was then plotted of timing of the magnetic stimulus in the ongoing tremor cycle against the actual—expected timing of the subsequent 10 EMG bursts. For each EMG burst after the magnetic stimulus, a linear regression line was derived. The slope of the lines gives an indication of the resetting that has occurred; a slope of 0 implies no resetting, whereas a slope of 1 implies complete resetting. The RI was calculated by taking the average slope of the regression lines for the 10 EMG bursts after the magnetic stimulation.

In many trials it was often difficult to decide whether the initial EMG activity after magnetic stimulation was due to rebound during the silent period or resumption of orthostatic tremor. To account for this, the first tremor burst after the magnetic stimulus in each raw trace was defined by the timing of the first EMG burst of tremor in the averaged record. Results were obtained from a total of 20 trials in each patient.

Statistical analysis of group data was performed with repeated measure analysis of variance (ANOVA) and paired Student’s t tests as appropriate. Correlation was performed by Pearson’s correlation test. For all analysis, significance was reported for P<0.05.

Results
In all records, each magnetic stimulus resulted in a short latency motor response, followed by a transient suppression of the EMG activity (silent period), before rhythmic EMG activity resumed at about 160 ms after the stimulus. Figure 1 illustrates the effect of averaging all individual sweeps, aligned according to the timing of occurrence of the magnetic stimulus. The occurrence of oscillatory EMG activity in the averaged EMG signal after the stimulus indicates that the tremor phase has been reset and become time locked to the stimulus. The calculated RI in case 1 was 0.87 for the right leg and 0.99 for the left leg (fig 1, upper two traces), and in case 2 it was 0.86 for the right leg and 0.79 for the left leg (fig 1, lower two traces).

Figure 2 shows the time course of tremor behaviour after magnetic stimulation. In all 20
trials, the average cycle length of the initial 10 tremor bursts after the stimulus tended to be shorter than those of the prestimulus tremor. On resumption of tremor, the average cycle length was about 70%–80% of the prestimulus levels, and gradually returned to the prestimulus tremor period. Poststimulus cycle lengths were shorter by an average of 6.4 ms in case 1 and 4.8 ms in case 2 (t tests all P<0.01). Repeated measures ANOVA between prestimulus and poststimulus average cycle lengths in both cases were significantly different (right, P<0.01; left, P<0.01).

To investigate whether the tremor bursts on both legs were synchronised before stimulation, the timing of the magnetic stimulus within a tremor cycle was measured and expressed as a percentage of the total average cycle length (% cycle length; fig 3) on both sides. The degree of synchrony was estimated by a comparison of the % cycle length for right and left legs. The results showed a high correlation in % cycle length between left and right legs in both patients (case 1: correlation coefficient 0.91, P<0.001; case 2: correlation coefficient 0.97, P<0.001) and suggested a synchronisation of tremor bursts in both legs before magnetic stimulation (fig 3).

Discussion
This study has shown that transcranial magnetic stimulation can reset and modulate orthostatic tremor of the legs. As the volleys evoked by magnetic stimulation were conveyed from the motor cortex to the spinal cord, where spinal motor neurons were discharged, resetting could occur at each of these levels. Electrical stimulation of the peroneal nerve is ineffective in modulating or resetting orthostatic tremor bursts in the tibialis anterior, making it unlikely that a peripheral feedback mechanism accounts for the resetting of this tremor after brain stimulation (unpublished observations). Furthermore, the 16 Hz tremor found in primary orthostatic tremor greatly exceeds that expected from oscillations in an overactive spinal reflex arc. Accordingly, central circuits must have a crucial role in the tremor resetting. Several features suggest that central factors might be important. Firstly, the
phase of tremor was highly synchronised in both legs (fig 3). This finding was compatible with a previous single motor unit cross correlation study in primary orthostatic tremor, in which close linkage of motor unit discharges was detected between both tibialis anterior muscles, strengthening the central hypothesis. Secondly, in most patients with orthostatic tremor, including the present cases, a tremor of similar frequency in proximal upper limb muscles is usually present when patients stand with their arms outstretched, either in front of or behind their bodies. If central circuits are involved in the genesis of the primary orthostatic tremor, how does this fit with the current understanding of pathophysiological mechanisms of tremor? Recently, a "two loop" hypothesis has been proposed as a model for parkinsonian tremor. In this model, the intrinsic loop, involving striatothalamiccorticostriatal circuits, interacts with an extrinsic loop in which short latency reafferent input to the ventralis intermedius nucleus of the thalamus (via cerebellar pathways) projects to the motor cortex and may serve to stabilise oscillation in the intrinsic loop. In the case of parkinsonian tremor, it was postulated that the extrinsic loop was modified by magnetic cortical stimulation or mechanical perturbation, culminating in tremor resetting or modulation. Interruption of a cortical projection is likely to account for resetting of orthostatic tremor by magnetic brain stimulation. Subcortical circuits also seem to be involved in orthostatic tremor. A recent PET study of primary orthostatic tremor showed that, when patients maintained a posture with their right upper limb outstretched, abnormal bilateral cerebellar and contralateral lentiform and thalamic activation were detected.

The period of the tremor after stimulation, particularly in the early stages of tremor reappearance (fig 2). It is of interest that a shortening of the tremor period was also reported in parkinsonian postural tremor but not in essential or mimicked tremors after transcranial magnetic stimulation. The phenomena suggest the pathophysiological mechanisms underlying primary orthostatic tremor and parkinsonian tremor are different from those responsible for essential and mimicked tremors and indicate a prominent role of the motor cortex in their generation or modulation.

Figure 3  Relation between the % cycle length of tremor bursts in the left and right legs of two patients with primary orthostatic tremor. The % cycle length indicates the timing of the tremor bursts as a percentage of the total tremor cycle. The linear regression correlation coefficient for case 1 was 0.91 and case 2 was 0.97 (both P<0.001). The strong correlation between the % cycle length in left and right legs indicates a high degree of synchronisation of the tremor bursts in left and right legs.

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