Beta-Adrenergic Modulation of Tremor and Corticomuscular Coherence in Humans

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Abstract
Coherence between the bioelectric activity of sensorimotor cortex and contralateral muscles can be observed around 20 Hz. By contrast, physiological tremor has a dominant frequency around 10 Hz. Although tremor has multiple sources, it is partly central in origin, reflecting a component of motoneuron discharge at this frequency. The motoneuron response to ~20 Hz descending input could be altered by non-linear interactions with ~10 Hz motoneuron firing. We investigated this further in eight healthy human subjects by testing the effects of the beta-adrenergic agents propranolol (non-selective β-antagonist) and salbutamol (β2-agonist), which are known to alter the size of physiological tremor. Corticomuscular coherence was assessed during an auxotonic precision grip task; tremor was quantified using accelerometry during index finger extension. Experiments with propranolol used a double-blind, placebo-controlled crossover design. A single oral dose of propranolol (40 mg) significantly increased beta band (15.3–32.2 Hz) corticmuscular coherence compared with placebo, but reduced tremor in the 6.2–11.9 Hz range. Salbutamol (2.5 mg) was administered by inhalation. Whilst salbutamol significantly increased tremor amplitude as expected, it did not change corticomuscular coherence. The opposite direction of the effects of propranolol on corticomuscular coherence and tremor, and the fact that salbutamol enhances tremor but does not affect coherence, implies that the magnitude of corticomuscular coherence is little influenced by non-linear interactions with 10 Hz oscillations in motoneurons or the periphery. Instead, we suggest that propranolol and salbutamol may affect both tremor and corticomuscular coherence partly via a central site of action.

Introduction
Many studies have investigated oscillatory coupling between the sensorimotor cortex and the periphery using coherence analysis. During steady contractions, cortical recordings (electroencephalogram, EEG, or magnetoencephalography, MEG, in humans, local field potentials in animals) show coherence with contralateral rectified electromyogram (EMG) at frequencies of 15–30 Hz [1–6]. Both efferent and afferent (feedback) pathways contribute to this oscillatory coupling [7–11].

In the periphery, oscillations are greatest at ~10 Hz. Physiological tremor has a strong component around this frequency, probably as a result of multiple interacting factors. These include mechanical resonance at skeletal articulations [12], the stretch reflex loop [13], motor unit recruitment [14] and motor unit synchronisation [15]. A component of ~10 Hz physiological tremor is centrally generated [15–17]. Although both cortical and muscle recordings show oscillations around 10 Hz and 20 Hz, significant corticomuscular coherence is usually only seen for the higher frequency band, despite ~10 Hz oscillations being effectively carried from the motor cortex down the corticospinal tract [18]. We have recently provided evidence that neural circuits in the spinal cord phase invert signals around 10 Hz [19,20]. Convergence of in-phase and out-of-phase signals on motoneurons results in phase cancellation, reducing the amplitude of signals at frequencies related to physiological tremor. This may have functional consequences in improving movement precision.

The non-linear nature of spinal circuits means that interactions can occur across different frequency bands. For example, our previous computational modelling suggested that recurrent inhibition by Renshaw cells will boost signals around 30 Hz, at the same time as reducing those around 10 Hz, leading to a reciprocal modulation of power at these frequencies [19]. Conversely, the highly periodic nature of motoneuron firing at low forces can produce oscillations in motor output, even when the synaptic input to motoneurons is not oscillatory [19]. As the firing of the motoneuron pool becomes more or less periodic, this effect will mean that the amplitude of oscillations at 10 Hz and 20 Hz tends to co-modulate. We were interested to examine experimentally how motor systems oscillations at different frequencies might interact, since this could tease apart the different factors influencing oscillatory amplitude and coupling.

One approach to this issue is to manipulate the system pharmacologically. We have previously shown that the benzodiazepine diazepam modulates cortical oscillations in the beta band, without altering coupling to the periphery [21]. By contrast, the anti-epileptic drug carbamazepine increases beta band corticomuscular coherence, without affecting the amplitude of cortical oscillations [22]. However, neither of these agents affected oscillations around 10 Hz. Clinically, the most powerful effects...
on physiological tremor are produced by beta-adrenergic agents. The beta-adrenergic agonists enhance tremor (adrenaline [23]; isoprenaline [24]; salbutamol [25]), whereas the antagonist propranolol reduces it [24]. These drugs are generally believed to act by changing the gain of muscle receptors and spinal reflex loops, mediated via peripherally located β2-adrenergic receptors [23,24,26–28]. If the effects of beta-adrenergic agents are solely peripheral, they will change the gain of peripheral feedback, which would inject oscillations into motoneuron firing. This might co-modulate oscillations at 10 Hz and 20 Hz [19].

In this paper, we test this idea directly. Surprisingly, we show that propranolol increases ~20 Hz corticomuscular coherence but reduces tremor at ~10 Hz, and that salbutamol has no significant effect on ~20 Hz corticomuscular coherence, but markedly increases ~10 Hz tremor. These opposite effects on oscillations at 10 Hz and higher frequencies are similar to those seen in a computational model of recurrent feedback by Renshaw cells [29], and are not expected if these agents only act peripherally. We speculate that beta-adrenergic agents also have a central action, allowing opposite effects on physiological tremor and beta-band corticomuscular coherence.

**Methods**

Experiments were performed on 8 young healthy right-handed volunteer subjects (3 female; age range 21–32 years) without a family history of essential tremor. None of the subjects were taking prescription medication either regularly or as required. Subjects were not pre-selected based on their corticomuscular coherence. None of the subjects had received a prior diagnosis of enhanced physiological tremor or had evidence of enhanced physiological tremor when assessed by a clinical neurologist (MRB) at the time of experiments.

**Ethics statement**

Informed written consent was obtained in accordance with the Declaration of Helsinki and all procedures were approved by the Cambridge Local Research Ethics Committee.

**Electrophysiological Recordings**

Surface EMGs from the right upper limb were recorded with adhesive electrodes (Biotrace 0713C, MSB, Marlborough, UK). Recordings were made from first dorsal interosseous (1DI), abductor pollicis brevis (AbPB), abductor digitii minimi (AbDM), flexor digitorum superficialis (FDS), and extensor digitorum communis (EDC), with an inter-electrode distance of 1.5–2 cm. Differential EEG was recorded from the left sensorimotor cortex using two adhesive scalp electrodes (Neurilept 720 00-S, Medicotest, St Ives, UK) placed 30 mm lateral to the vertex and 20 mm anterior and posterior to the interaural line. The anterior EEG electrode was connected to the non-inverting input of the amplifier; this is the same montage as used in our previous work [7,11,21,22,30]. Signals were amplified (EMG gain 300–5000; EEG gain 50 k) and bandpass filtered (EMG 30 Hz–2 kHz; EEG 3 Hz–2 kHz) before being digitised at 4273.5 Hz by a Power1401 interface (Cambridge Electronic Design Ltd, Cambridge, UK) connected to a computer running Spike2 software (Cambridge Electronic Design Ltd).

**Measurement of Tremor**

Physiological tremor was quantified using accelerometry. A low mass splint was taped to the subject’s right index finger, and restricted movements to the metacarpophalangeal joint. Subjects gripped a fixed vertical pole with the thumb and digits 3–5, and extended the index finger. A miniature accelerometer (Isotron 25B, Endevco, San Juan Capistrano, CA, USA) was fixed to the end of the splint, with its sensitive axis aligned to detect finger flexion/extension movements. Recordings were made for 90 s.

**Precision Grip Task**

Measurement of corticomuscular coherence was carried out using an auxotonic precision grip task. Subjects held the two levers of a purpose-built manipulandum between thumb and index finger. The aluminium levers (20×80×1.5 mm) were attached to the shafts of two computer-controlled torque motors, which incorporated optical encoders for position measurement. At rest the levers were separated by 70 mm; a 1 N force was required to move the levers off their end-stops.

The task (see Figure 1) required subjects to maintain cursors representing each lever position within two moving target boxes on a computer screen. Movement of the levers was resisted by the torque motors, which simulated an auxotonic (spring-like) load. The target boxes produced a hold-ramp-hold pattern, with the first hold requiring a rapid lever displacement of 12 mm from rest followed by a hold period of 3 s at a force level of 1.3 N. The targets then produced a 2 s ramp movement to reach the second hold, with a displacement of 24 mm, 1.6 N force and duration of 3 s, before subjects released the levers. This is the COMP1 task of Kilner et al. [6], and has been used in several of our previous publications [7,11,21,22,30].

**Experimental Protocol and Drug Administration**

Drugs were prescribed and administered by one of the authors (MRB), who is a clinical neurologist. Potential adverse reactions were minimised by using the lowest therapeutic dose recommended by the Joint Formulary Committee [31].

Propranolol experiments were conducted as a double blind randomised placebo-controlled trial. For each subject, placebo and propranolol (40 mg of active compound mixed with vehicle compound), which were indistinguishable in appearance, were randomly assigned a label A or B by an independent randomizer. Each subject participated in experiments on two days, separated by a 2–3 week washout period. At the start of the morning session of each day, the subject was prepared for recording. A control experiment was then carried out, which consisted of 80 trials of the precision grip task and a measurement of tremor. After this, the electrode locations were marked on the skin with ink and the subject ingested experimental compound A (first day) or B (second day). Four hours later, when peak serum concentrations of the active agent should have been achieved [32], the subject returned to the laboratory. Fresh adhesive electrodes were applied at the marked muscle and scalp locations, and further recordings were made. The identities of compound A and B for each subject were only revealed by the randomizer once all experiments were complete and data were ready for analysis. By using a placebo, we were able to control for effects caused by the different time of day of the experimental and control recording sessions, which we have previously shown to be important [22].

The subjects who had participated in the propranolol experiments were also used in the salbutamol experiments, following a washout period of at least one week. After the control recordings, 2.5 mg of salbutamol was inhaled by the subject as an aerosolised solution produced by a nebuliser. Inhaled aerosolised salbutamol is absorbed rapidly via the lungs into the bloodstream. Maximum serum concentrations are achieved ~10 minutes after inhalation [33]. Nebulised salbutamol, at a dose of 40 μg/kg, results in an average serum concentration of 2.5 ng/ml after inhalation [34]. The rapid absorption meant that control and test experiments
were separated by only a short period, and we accordingly considered it unnecessary to carry out placebo experiments for salbutamol (see paragraph 12 of the Results section for further justification).

Electrocardiogram (ECG) was monitored throughout the salbutamol experiments. Salbutamol at an average plasma concentration of 2.5 ng/ml produces a 10% increase in heart rate above baseline [34]. The second recording session (after salbutamol) was therefore only commenced after the heart rate had increased by 10% relative to baseline.

Analysis

All data analysis was performed using custom written MATLAB (Mathworks Inc.) routines, and followed the procedures used in our prior publications [7,22]. Prior to analysis EMG recordings were full wave rectified.

Analysis of tremor power used 0.88 s long (4096 points for a sampling rate of 4630 Hz) sequential non-overlapping data sections, taken from the entire available duration of the accelerometer recording. These were processed with a Fast Fourier Transform, yielding a tremor power spectrum with frequency resolution of 1.13 Hz.

Analysis for the precision grip task focused upon the second hold phase, when β20 Hz corticomuscular coherence is greatest [6]. EEG and EMG power spectra, and coherence between EEG and EMG, were computed using three contiguous 0.88 s-long sections of data taken from this task phase of each trial, and a 4096-point-long Fast Fourier Transform [3].

Figure 1. Experimental tasks. A. Postural tremor task. B. Example of postural physiological tremor in a single subject, recorded via an accelerometer. C. Auxotonic precision grip task. D. Lever displacement and average rectified EMG recorded from extensor digitorum communis (EDC), flexor digitorum superficialis (FDS), abductor digiti minimi (AbDM), abductor pollicis brevis (AbPB) and first dorsal interosseous (1DI) muscles in a single subject during a single trial of the auxotonic task. doi:10.1371/journal.pone.0049088.g001
Accelerometer, EEG and EMG power spectra were first normalised to the total power in that signal across all frequencies. For acceleration, the normalisation used the total power in the first recording of that day, allowing overall changes in tremor power to be measured. For EEG and EMG, each spectrum was normalised to its own total power, revealing the magnitude of spectral peaks relative to the total. The spectra were then averaged across eight subjects for EEG or acceleration, and across eight subjects and five muscles for EMG. Coherence spectra were similarly averaged across subjects and muscles; significance limits for the averaged coherence were calculated according to the method described in Evans and Baker [35] and Baker et al [8].

Power was summed across a frequency band of interest (6.2–11.9 Hz for tremor; 15.3–32.2 Hz for beta-band oscillations) within a single subject. Changes of power before and after substance administration in these frequency bands were then assessed using paired t-tests (P<0.05). For propranolol, we needed to test not only whether propranolol produced a change, but whether this change was significantly different from that produced by placebo. Accordingly, the change in power in the relevant band was found for each subject; these changes were compared between placebo and propranolol (paired t-test, P<0.05).

For corticomuscular coherence, coherence was averaged across muscles and across bins within the 15.3–32.2 Hz band for each subject. The significance of the changes in coherence before and after substance administration was determined by finding

\[ Z = \sqrt{\frac{L}{5NsNf} \sum_{f=1}^{Nf} \sum_{n=1}^{Ns} \sum_{m=1}^{5} (\text{atanh}(\sqrt{C_{n,m}(f)}) - \text{atanh}(\sqrt{C_{\text{before},n,m}(f)}) \right) \]

\[ \text{atanh}(\sqrt{C_{\text{before},n,m}(f)}) \]

Where L is the number of disjoint sections used to calculate the coherence, and \( C_{n,m}(f) \text{ and } C_{\text{before},n,m}(f) \) are the coherence calculated at frequency bin f for subject n, muscle m before and after substance administration. \( Z \) is computed from a sum over all available \( Ns \) subjects, \( Nf \) frequencies (\( Nf = f_{1} - f_{f} + 1 \)) and 5 muscles. On the null hypothesis that coherence is the same before and after substance administration, \( Z \) will be normally distributed with zero mean and unit variance [21,30]. The significance of changes in coherence was thus determined with reference to values of the standard normal probability distribution.

In experiments investigating the effects of propranolol, it was necessary to compare the changes caused by propranolol with standard normal probability distribution. Coherence was thus determined with reference to values of the mean and unit variance [21,30]. The significance of changes in coherence before and after substance administration was determined by finding

\[ \Delta Z = \frac{1}{\sqrt{2}} (Z_{\text{propranolol}} - Z_{\text{placebo}}) \]

The normalisation by \( 1/\sqrt{2} \) once again ensured that, on the null hypothesis that the substances cause equal changes in coherence, \( \Delta Z \) will be normally distributed with zero mean and unit variance.

We determined the effect of a drug or placebo on the coherence phase as follows. Firstly, frequency bins in the beta-band region (15.3–32.2 Hz) for each muscle were identified which had significant coherence both before and after substance administration. The coherence phase was measured for those bins, and the difference in phase determined between the ‘before’ and ‘after’ conditions. The circular average of this difference was computed [36]. We then randomly shuffled the individual paired phase measurements, reallocating each member of a pair to the ‘before’ or ‘after’ categories; the circular average of the phase differences was recalculated. The shuffling procedure was repeated 10^5 times, allowing estimation of the distribution of the circular mean phase difference, on the null hypothesis that there was no change before and after substance administration. If the absolute value of the circular average phase difference from the actual experimental data was larger than n of the absolute values of the circular average phase differences determined from the shuffled data, this yielded an approximate Monte Carlo significance level of \( P<ns/10^5 \).

**Results**

Figure 2 illustrates the results obtained from a single subject in each experimental protocol tested. Figures 2A–C show the effect of the different substances administered on physiological tremor, measured from the acceleration power spectrum during finger extension. In this subject, two peaks were visible in the tremor spectra, around 8 Hz and around 20 Hz; the ~10 Hz region usually associated with physiological tremor is marked with yellow shading. Each panel of Fig. 2A–C relates to measurements made on a different day; it is notable that there was high day-to-day variability in the baseline tremor levels (compare Fig. 2C with 2A&B). There was little change in tremor following the placebo (Fig. 2A), but a substantial reduction after propranolol (Fig. 2B) in this subject. By contrast, salbutamol dramatically increased tremor (Fig. 2C).

The remaining panels of Fig. 2 show results from the auxotonic precision grip task, which was used to examine beta-band oscillations in the motor system. Accordingly, the yellow shading in these plots highlights the 15.3–32.2 Hz band, which encompassed the beta peaks. In this subject, there was a small increase in the size of the power spectral peak at these frequencies in the EMG following both placebo and propranolol (Fig. 2DE); this is probably attributable to the time of day of the recordings, as the two datasets were gathered four hours apart to allow time for absorption of the orally administered substance. By contrast, salbutamol appeared to have little effect on the beta band peak in the EMG (Fig. 2F). The placebo recordings in this subject showed a non-specific rise in EEG power across many frequencies (Fig. 2G), but neither propranolol nor salbutamol seemed to affect the beta-band EEG peak (Fig. 2HI).

In the initial recordings for placebo and propranolol administration (Fig. 2JK), this subject showed weak beta-band corticomuscular coherence which barely rose above the significance level (dashed lines). In both cases, the coherence peak was enhanced in the second recording of the day. The control recording for salbutamol showed clearer coherence (Fig. 2L), but this did not change consistently following salbutamol inhalation. The high day-to-day variability in baseline measures is again of note; we have previously reported fluctuations in corticomuscular coherence from single subjects both diurnally [22] and over longer timescales [11]. Diurnal changes in coherence probably explain the apparent effect of placebo in Fig. 2J: we have previously shown that coherence increases in measurements made later in the day [22].

Figure 3 shows population data on the effects of the two drugs tested on physiological tremor. Figure 3AB plots the tremor power spectra, averaged across all eight subjects. There appear to be some changes in the peak around 10 Hz, with an increase in the tremor following placebo, but a reduction following propranolol. Figure 3G shows how the average power over the tremor band (6.2–11.9 Hz) changed in individual subjects. Although there was a trend for increased tremor with placebo, and decreased tremor with propranolol, these changes were not significant. Figure 3D
Figure 2. Single Subject Data. Each column of this figure refers to the effects of a different substance: placebo (A,D,G,J), propranolol (B,E,H,K) and salbutamol (C,F,I,L). A–C, effect on postural tremor during an index finger extension task. Traces show the power spectrum of the finger acceleration, before and after substance ingestion. Yellow shaded region corresponds to the band of physiological tremor (6.2–11.9 Hz). D–F, effect on EMG power. G–I, effect on EEG power. J–L, effect on corticomuscular coherence. In (D–L), all measurements were made during the second hold phase of a hold-ramp-hold auxotonic precision grip task (see Methods). Yellow shaded region marks the 15.3–32.2 Hz band of beta oscillations. EMG power spectra (D–F) and corticomuscular coherence (J–L) have been averaged over all five muscles recorded. In (J–L), horizontal dashed lines indicate the significance level for the averaged coherence.

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placebo (P < 0.05, paired t-test and also binomial test). There may have been some weak differences between measures made from the first versus last half of the recorded data (P = 0.14; Fig. 6B); however, even if this effect were genuine, it would be in the opposite direction to that seen following salbutamol inhalation.

Figure 4 presents the effects of propranolol on oscillations during the second hold phase of the precision grip task. Figure 4AB shows the EMG power spectra, averaged over all eight subjects and five muscles. In both cases, the recordings made later in the day showed a more pronounced beta-band spectral peak. Figure 4C shows individual measures of beta-band power; there were significant increases following both propranolol and placebo administration. However, a comparison of the changes in power (Fig. 4D) revealed no significant difference between the effects of propranolol and placebo.

Using the same format, Fig. 4E–H presents the changes in EEG power following placebo and propranolol administration. In both cases there was a slight increase in the beta-band power peak, which became significant for propranolol (Fig. 4G). However, as for the EMG power, there were no significant differences between the effects of propranolol and placebo.

For both EEG and EMG power we normalised values in a given frequency bin relative to the total power, summed across all frequencies. We also verified that there were no significant changes in the total power, for any of the comparisons described by Fig. 4CDGH.

Finally, Fig. 4I–L shows how corticomuscular coherence altered following ingestion of these substances. There was a very small increase in coherence following placebo, but a more pronounced increase following propranolol (Fig. 4I); both of these effects reached significance (Fig. 4K). Propranolol increased beta-band corticomuscular coherence more than placebo (Fig. 4L) in 7/8 subjects, significantly so (using an individual difference of coherence test) in 3 subjects. At a population level, the increase following propranolol was significantly greater than after placebo using the difference of coherence test described in Methods (P < 0.001), and also on a binomial test (P < 0.05).

The results from the precision grip task following salbutamol inhalation are shown in Fig. 5. There were small or negligible changes in the averaged beta band power spectral peaks in both EMG and EEG signals (Fig. 5AC); these were not consistent across subjects (Fig. 5BD) and not significant. Similarly, the averaged corticomuscular coherence decreased slightly (Fig. 5E), but this was not significant (Fig. 5F).

Experiments with propranolol used a double-blind placebo controlled design, as the long absorption time consequent on oral administration made it necessary to control for an effect of the time of day when recordings were made. By contrast, the rapid action of the salbutamol delivered by inhalation meant that control and drug recordings were made with only brief temporal separation. We thus chose not to make the comparison with placebo in this case. Additionally, the effects of salbutamol on heart rate and tremor were clearly apparent to the subjects, so that a double-blinded study would have been impossible. Several pieces of evidence indicated that the changes we saw following salbutamol were genuine effects of the drug, rather than reflecting progressive changes in the successive recordings. For corticomuscular coherence, Pohja et al [37] previously showed that successive measurements were highly repeatable. We also tested directly for reproducibility, by dividing each dataset recorded before salbutamol administration into two. The results from this analysis are shown in Fig. 6, for both tremor (Fig. 6A) and corticomuscular coherence (Fig. 6D). In neither case were there significant differences between measures made from the first versus last half of the recorded data (P > 0.05). There may have been some weak trend towards reduced tremor in the later part of the recording (P = 0.14; Fig. 6B); however, even if this effect were genuine, it would be in the opposite direction to that seen following salbutamol administration (Fig. 6F).

Figure 3. Effects on Resting Tremor. All measurements in this figure were taken during index finger extension. A, B, changes in finger acceleration power spectra, averaged across all eight subjects, before and after administration of placebo (A) and propranolol (B). C, total acceleration power over the 6.2–11.9 Hz range (yellow shading in A,B) before and after substance administration. Each point shows data from one subject; lines link data from the same subject. Data points are colour coded to match the colours used in (A,B). There was no significant change in tremor power after either placebo or propranolol. D, changes in tremor power after administration of placebo or propranolol. Each point shows the difference between the ‘after’ and ‘before’ points from the corresponding subject shown in (C). Propranolol reduced tremor power significantly compared to placebo (P < 0.05), E, changes in finger acceleration power spectra, averaged across all eight subjects, before and after administration of salbutamol. F, total acceleration power over the 6.2–11.9 Hz range (yellow shading in E) before and after salbutamol administration. Tremor power was significantly increased after salbutamol (P < 0.05).

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These results confirm that – at the doses used here – propranolol reduces, but salbutamol increases, physiological tremor, which is in accord with previous work [24] and also common clinical experience.
Phase analysis provides important clues as to the underlying network and the likely explanation for changes in corticomuscular coherence. In order to investigate phase changes, in each EMG and each subject we found the frequency bins in the 15.3–32.2 Hz range which showed significant coherence both before and after substance administration. The phase of the coherence following administration of a substance was then plotted against the phase measured in the control experiment. Figure 7 shows the results of such an analysis, with points overlain from all EMGs and subjects. Results from placebo, propranolol, and salbutamol respectively are shown in Fig. 7A, B and C. Significant changes in phase were seen in the salbutamol dataset (circular mean phases before and after substance administration: 1.63 vs 1.51 rad, P = 0.026, Monte Carlo test). Similarly significant differences were seen when the two outlying points with negative phase before drug administration were excluded from the analysis (P = 0.017, Monte Carlo test). There were no significant changes in phase for either propranolol or placebo (both P > 0.05).

**Discussion**

**Differential Effects on Tremor and Corticomuscular Coherence**

Surprisingly, the drugs tested in this study had different effects on ~20 Hz corticomuscular coherence and ~10 Hz tremor (Fig. 3 versus Figs. 4&5). For propranolol, the effects were reciprocal. Tremor showed a weak but non-significant trend to increase at the later time of day of the second recordings with placebo (Fig. 3AC); by contrast, when propranolol was given there was a non-significant trend in the reverse direction (Fig. 3BD). Comparison of
these two effects revealed a significant difference, indicating that propranolol reduced tremor compared with placebo (Fig. 3D). By contrast, propranolol elevated corticomuscular coherence (Fig. 4L). Such results are reminiscent of findings from our computational model of the effect of recurrent inhibition [29]. In this model, 10 Hz frequency components of the motor cortical input to motoneurones are removed by Renshaw cells, behaving as phase inverting filters, and thus 10 Hz tremor is reduced. By manipulating the strength of Renshaw cell recurrent inhibition, this model showed a reciprocal relationship between 10 Hz tremor and beta-band corticomuscular coherence (see Fig. 4 in [29]). Although the largest effect on corticomuscular coherence in the model was at 30 Hz, coherence in the entire band from 20–40 Hz was raised by increasing Renshaw cell feedback.

Whereas salbutamol clearly increased ~10 Hz tremor (Fig. 3EF), there was no significant change in corticomuscular coherence across the subject population (Fig. 5F). It is not clear why salbutamol and propranolol failed to exert reciprocal effects on the corticomuscular coherence. This may simply be due to the low doses used, which were chosen to minimise the chances of side effects in our healthy volunteer subjects, or to β1-receptor or β3-receptor mediated effects (salbutamol is a β2-agonist, whereas propranolol is a non-specific β-receptor antagonist).

### Location of peripheral and central β-Adrenergic Receptors Mediating Effects

Whilst part of the action of the non-specific beta-blocker propranolol on physiological tremor could be via a peripheral site [23,24,28], it seems likely, in the light of our results, that there is also a central action [38]. In support of this hypothesis, propranolol, which is lipophilic, is significantly more effective at both reducing tremor [24], and penetrating the blood brain barrier [39] than the hydrophilic atenolol. Whether the central effects of propranolol are mediated via one specific class of beta-adrenergic receptor or a combination is unclear. All subtypes of beta-adrenergic receptors are widely distributed throughout the central nervous system. Immunohistochemistry has identified β1-adrenergic receptors in the amygdala, hippocampus, hypothalamus, cerebellum, midbrain reticular nuclei, inferior olive and striatum [40] and β2-adrenergic receptors in the locus coeruleus [42] and soma and proximal dendrites of thalamo-cortical neurones [41]. In situ hybridisation has further revealed β3-adrenergic receptors in the thalamic intralaminar nuclei, cerebellar cortex [42] and superficial dorsal horn of spinal cord [43]. β3-adrenergic receptor mRNA has been detected in homogenates of frontal, temporal and parietal cortex, hippocampus, striatum and midbrain by reverse transcription/polymerase chain reaction methods [44,45].
Of the likely candidate motor areas implicated, it is unlikely that \( \beta \)-adrenergic effects on corticomuscular coherence are explained by actions on the cerebellum. It is known that, 20 Hz oscillations are propagated around a loop from motor cortex to cerebellum and back [43]. However, changing the properties of this loop should change only the nature of oscillations observed in the cortex, and not alter the coupling between cortex and periphery.

Given our previous results from computational modelling, we speculate that propranolol acts centrally via Renshaw cells, producing a reciprocal modulation of ~10 Hz tremor and 20-30 Hz corticomuscular coherence. Because Renshaw cells themselves are not known to have \( \beta \)-receptors, these effects would have to be mediated indirectly via inputs from neurons expressing \( \beta \)-adrenergic receptors, including those within the locus coeruleus (LC) and spinal dorsal horn. Interestingly, in the cat, Renshaw cells are in receipt of direct adrenergic reticulospinal inputs from LC, which when activated reduce recurrent inhibition [43]. Moreover, the \( \beta_2 \)-agonist (SR58611A) increases the firing rate of noradrenergic neurons in the locus coeruleus [44].

In contrast to propranolol, the striking lack of any effect of the \( \beta_2 \)-agonist salbutamol upon coherence suggests that its predominant mode of action is via peripheral receptors. \( \beta_2 \)-adrenergic receptors are certainly found in abundance in extrafusal muscle where they mediate diverse functions including: metabolic regulation [46]; myocyte hypertrophy [47]; and repair [48]. These actions occur over a protracted time course and are therefore unlikely to be responsible for salbutamol’s rapid effects on tremor. Faster processes are also mediated via \( \beta_2 \)-receptors. They decrease muscle excitability by closing inward rectifying potassium channels, opening calcium-activated potassium channels [49] and blocking sodium currents [50]. In addition, because sympathetic activity produces rapid changes in blood flow to skeletal muscle, it has been argued that ischaemia (or its consequences e.g. hyperkalaemia) might affect tremor by changing the biophysical properties of muscle [28].

**Network re-organization**

Although the literature is conflicting [4,51–54], the phase of corticomuscular coherence often does not agree with simple efferent propagation of oscillations from motor cortex to the periphery [7,11,19]. Rather, coherence phase probably reflects a complex interplay of feedforward and feedback pathways. In the present work, we showed that salbutamol produced a small but significant reduction in the coherence phase (Fig. 6). Although salbutamol did not change the magnitude of coherence, the reduction in phase might reflect a slight shift in the balance between feedforward and feedback pathways contributing to the coherence [11]. This could occur either by central actions of the drug, or via its previously reported peripheral actions. Beta-agonists are likely to increase the sensitivity of muscle spindles [24], whose afferents are known to encode beta-band oscillations [8].

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**Author Contributions**

Conceived and designed the experiments: MRB SNB. Performed the experiments: MRB SNB. Analyzed the data: MRB SNB. Contributed reagents/materials/analysis tools: MRB SNB. Wrote the paper: MRB SNB.

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