Is postural tremor size controlled by interstitial potassium concentration in muscle?

M Lakie, N R Hayes, N Combes, N Langford

**Objectives:** To determine whether factors associated with postural tremor operate by altering muscle interstitial K⁺.

**Methods:** An experimental approach was used to investigate the effects of procedures designed to increase or decrease interstitial K⁺. Postural physiological tremor was measured by conventional means. Brief periods of ischaemic muscle activity were used to increase muscle interstitial K⁺. Infusion of the β₂ agonist terbutaline was used to decrease plasma (and interstitial) K⁺. Blood samples were taken for the determination of plasma K⁺.

**Results:** Ischaemia rapidly reduced tremor size, but only when the muscle was active. The β₂ agonist produced a slow and progressive rise in tremor size that was almost exactly mirrored by a slow and progressive decrease in plasma K⁺.

**Conclusions:** Ischaemic reduction of postural tremor has been attributed to effects on muscle spindles or an unexplained effect on muscle. This study showed that ischaemia did not reduce tremor size unless there was accompanying muscular activity. An accumulation of K⁺ in the interstitium of the ischaemic active muscle may blunt the response of the muscle and reduce its fusion frequency, so that the force output becomes less pulsatile and tremor size decreases. When a β₂ agonist is infused, the rise in tremor mirrors the resultant decrease in plasma K⁺. Decreased plasma K⁺ reduces interstitial K⁺ concentration and may produce greater muscular force fluctuation (more tremor). Many other factors that affect postural tremor size may exert their effect by altering plasma K⁺ concentration, thereby changing the concentration of K⁺ in the interstitial fluid.

**Definition of terms**

Postural tremor is a fine shaking of the limbs. It is classically studied by recording the inadvertent motion of a limb (usually the hand) when the subject attempts to hold it in a stationary posture. When hand tremor is recorded by conventional accelerometry it can be seen to possess a more or less rhythmic component, usually with a peak frequency between 7 and 11 Hz. There are several views concerning the reason for this predominant frequency of oscillation, but as far as we know there is only one way of altering the frequency of tremor and that is by changing the mechanical properties of the limb. In this paper we do not concern ourselves with the frequency of the oscillation. However, as will be seen, there are many ways of altering the size of the shake. In this paper we do not address the question of overt pathological tremors, such as those caused by Parkinsonism or cerebellar disease. Our findings may, however, have implications for one form of essential tremor and its pharmacological treatment.

**METHODS**

**Subjects**

We carried out β₂ agonist infusion experiments on 12 subjects (eight male, four female), mean (SD) age 21.4 (1.7) years. Ischaemia experiments were done on six subjects (five male, one female), mean age 37 (9.0) years. Permission was obtained from the local ethics committee, and written informed consent was obtained from the subjects. The experiments conformed with the declaration of Helsinki. All subjects declared themselves to be in good health and none had overt neurological symptoms or were taking β agonist or β blocker drugs.
Apparatus

Hand tremor was recorded using a miniature accelerometer (5.0 g; Eurosensor, Towcester, Northants, UK) attached by a tight Velcro® strap above the nail bed of the middle finger of the right hand. Subjects were seated in a comfortable chair which had a shaped polyether foam cradle which supported the right forearm as far as the wrist joint. The unsupported pronated hand was extended and the subject was instructed to maintain it in an approximately horizontal position with the fingers slightly separated. The method and tremor measurements made with it have been described previously. The acceleration signal was dc amplified so that 1.0 g = 3.75 V, and an offset was subtracted to null the effects of gravity on the accelerometer. The resulting signal was further amplified with a gain of 2.5 to 25 (to accommodate the subjects’ tremor size) and filtered with a passband of 2–40 Hz. The signal was converted by an analogue to digital interface (CED 1401, 12 bit resolution, sampling rate 1000 Hz) and stored on a personal computer for subsequent analysis. In these experiments an electromyogram (EMG) was routinely recorded from the active extensor digitorum communis muscle, using surface electrodes with an inbuilt 1000 gain preamplifier and a passband of 20–350 Hz.

Procedures

In separate experiments, tremor was recorded after limb ischaemia and following infusion of the β2 agonist drug terbutaline (Bricanyl sulphate). For the drug trials, cannulae were inserted into the overnight fasted subject, who was allowed to relax for a period of at least 45 minutes, during which three control measurements were made. The drug was infused at a rate of 8 μg kg⁻¹ h⁻¹ through a cannula inserted into the left antecubital vein, over a period of 45 minutes. Venous blood samples were periodically taken from a second cannula implanted in the right antecubital vein. Blood samples were processed and stored at −80°C and were subsequently analysed for plasma K⁺ (indirect ion specific electrode). In these experiments there were infusions of different drugs, dosages, and placebos, with each subject being infused on five occasions. The subject was blind to the agent being infused. We only report here the effects of the terbutaline infusion. In the ischaemia experiments, the recording arrangements were the same. Ischaemia was produced by inflating a cuff placed around the upper right arm to a pressure of 200 mm Hg. The duration of the ischaemic period was two minutes.

Analysis

Various procedures can be used to quantify the size of a shake. In these experiments, fast Fourier transform (FFT) analysis was used to calculate the amplitude spectrum of the tremor. The duration of the epoch that was analysed was variable. It was normally ~30 seconds (ischaemia control measurements) or ~60 seconds (terbutaline) but was shorter (~15 seconds) when tremor size was likely to be changing relatively rapidly (that is, in the ischaemia and muscle activity experiments). The total amplitude in the spectrum between 4 and 14 Hz was computed and its mean level was calculated and used as an index of tremor size. Most of the power in the tremor acceleration signal lies between 4 Hz and 14 Hz, and time domain measurements made using root mean square (RMS) values of the waveform, low pass filtered at 40 Hz, revealed a very similar pattern of change. Tremor size commonly reduces in the first few seconds after adopting a posture. To discount this effect we always allowed a period of at least five seconds between adoption of posture and the start of the analysis period. EMG was quantified by determining the area under the spectrum between dc and 45 Hz. Plasma concentration of K⁺ was expressed as mmol/litre.

RESULTS

Ischaemia

Figure 1 shows the change in the tremor (raw acceleration signal) in a typical subject. When a posture was maintained continuously during ischaemia, tremor size started to decline progressively after cuff inflation and was strikingly reduced after two minutes. However, when posture was not maintained continuously during ischaemia, there was only a small reduction in tremor size when posture was readopted by the arm after two minutes. When postural activity was restarted after ischaemia under resting conditions, the tremor size again showed a slow progressive decline as the posture was maintained. A substantial reduction in tremor was only apparent when muscular effort was combined with the ischaemia. In contrast to the mechanical record, the visual appearance of the raw EMG signal changed little, suggesting no change in the level of activation.

Figure 2 shows the power spectrum of the tremor signal recorded in the control period compared with the spectrum obtained two minutes after ischaemia, with and without concomitant postural activity. This figure also shows how the size of the tremor signal was quantified for use in the subsequent data analysis. This frequency domain analysis shows that all frequencies of tremor are substantially reduced...
by ischaemia, but only when there is accompanying postural activity. Ischaemia on its own only causes a small reduction in tremor size.

Figure 3 shows that for all our subjects the decline in tremor size over a two minute period was related not to ischaemia itself but to ischaemic muscle activity. The overall reduction in tremor size was large, consistent, and rapid when the ischaemic arm is active (mean (SD) reduction, 60 (10)% and much smaller and more variable (24 (18)%) when it was relaxed between measurements. The gross EMG necessary to generate the posture was not significantly different in the two conditions. The envelope of the EMG signal will contain components with similar frequencies to the tremor. The rectified and integrated (leaky integrator 20 ms time constant) EMG signal was subjected to FFT analysis. The resulting spectrum was divided into three bands, and the power in each was quantified (table 1). There was no significant change in the power in the band associated with tremor frequencies, or in the other two following ischaemia with or without postural effort. Peripheral ischaemia combined with postural effort reduced tremor size without a concomitant change in neural drive.

β₂ Agonist

Figure 4 shows how tremor size and plasma K⁺ changed in the infusion experiments. Terbutaline infusion began at time zero and lasted for 45 minutes. Tremor size increased steadily and reached a maximum around 60 minutes after the start of infusion (that is, about 15 minutes after the infusion ended). The change in tremor size is mirrored almost exactly by the change in K⁺, which falls from an initial mean value of 3.75 mmol l⁻¹ to 2.95 mmol l⁻¹ at time 60 minutes. At the end of the experiment (30 minutes after the infusion had ended) the tremor size was still increased and the K⁺ was still depressed.

Figure 5 shows how tremor size and plasma K⁺ changed in the infusion experiments. Terbutaline infusion began at time zero and lasted for 45 minutes. Tremor size increased steadily and reached a maximum around 60 minutes after the start of infusion (that is, about 15 minutes after the infusion ended). The change in tremor size is mirrored almost exactly by the change in K⁺, which falls from an initial mean value of 3.75 mmol l⁻¹ to 2.95 mmol l⁻¹ at time 60 minutes. At the end of the experiment (30 minutes after the infusion had ended) the tremor size was still increased and the K⁺ was still depressed.

Ischaemia

Ischaemic reduction of tremor size was originally attributed to changes in muscle spindle sensitivity. However, the role of muscle spindles in the genesis of this form of physiological tremor is debatable. For example, an artificially produced “tremor” (a small movement of the hand produced by repetitive ~10 Hz electrical stimulation at low intensity) was also rapidly attenuated by ischaemia. Thus the effect appeared less likely to be a reflex action and more likely to be a muscular phenomenon, but these investigators were unable to explain its basis, although they suggested it might be a form of “fatigue.” Our present observations provide an explanation. The reduction in tremor size is not produced by...
the ischaemia itself but rather by muscular activity in the presence of ischaemia. This has the hallmark of an effect produced by depletion of a substrate or accumulation of a metabolite. We suggest this substance may be K\(^+\) which accumulates in the ECF of the active muscle. There is no change in the EMG size or frequency spectrum. This further suggests that the reduction mechanism is muscular rather than neural. There was some reduction in tremor in the limb under resting conditions, but in general it was difficult for subjects to maintain the arm in a completely relaxed state.

**Infusion of terbutaline**

Terbutaline is a \(\beta_2\) adrenergic drug and is strongly tremorogenic. The classic experiments by Marsden and coworkers\(^4,7\) revealed a considerable increase in postural tremor and a slight decrease in the twitch time of slow muscles when adrenaline was infused. These workers concluded that it was the increase in the fusion frequency of the muscle that led to the great increase in tremor size observed. Thus a frequency of motor unit firing that would normally produce a fused force output would result in an output with much more ripple in the presence of an adrenergic drug. Adrenaline mediated changes in contractile characteristics had earlier been reported in animal studies.\(^4\) \(\beta_2\) Agonists are thought to activate membrane located \(\beta_2\) receptors (first described by Bowman and Nott\(^9\)), altering the twitch properties of muscle as a result of cyclic 3',5'AMP released into the muscle cytosol. However, as far as we are aware there is no direct evidence that this pathway is actually involved in the alteration of contractile characteristics. The increase in tremor is uncharacteristically slow. Abila \textit{et al} commented that “processes taking up to one hour to develop would be unusual for mechanisms expected to be mediated via adenylyl cyclase”.\(^10\)

The tremorogenic effect in the present experiments was slow (fig 4). The tremor that can occur when these drugs are used to treat respiratory problems develops some time after the therapeutic effect. The rate constant of the tremorogenic effects of isoprenaline and the tremolytic effects of the selective \(\beta_2\) blocker propanolol are very much slower than the heart rate responses.\(^10\)

The delayed increase in tremor after a frightening experience which causes an immediate release of adrenaline will be familiar to most people. Also, tremor is the final symptom to appear following the adrenergic response to an insulin induced hypoglycaemic crisis.\(^11\) We suggest (see below) that the contractile characteristics may be altered not by a specific direct adenylyl cyclase mediated action of the drug on muscle \(\beta_2\) receptors but by the more generalised reduction in K\(^+\) concentration that it produces by its well established effect on the muscle membrane located Na\(^+\)/K\(^+\) pump (see for example, Bengtsson\(^12\)). Two recent studies\(^1\) 2 and this one (fig 4) show that tremor size varies reciprocally and simultaneously with plasma K\(^+\). This relation might be even clearer were it not for the facts that tremor size is likely to be affected by the rate of change in K\(^+\) concentration as well as the absolute concentration, and that plasma K\(^+\) is not a precise indicator of muscle interstitial concentration, although it will reflect it.

**The proposed mechanism linking potassium concentration and tremor size**

As excitable cells are relatively permeable to K\(^+\) ions their membrane potential is close to the equilibrium potential for that ion. Consequently, alterations in extracellular K\(^+\) will have a profound effect on membrane potential and excitability. As far as the cells of the nervous system are concerned this effect is minimised by the presence of the blood–brain barrier, which is impermeable to K\(^+\), and by glial cells which “mop up” K\(^+\) and distribute it over a larger area (“spatial buffering”). Muscle cells have no such protective mechanism. Muscle fibres represent an enormous reserve of K\(^+\) which can
leak into the interstitium. Although the muscle cell membrane (sarcolemma) is relatively permeable to K⁺, accumulation in the interstitium does not normally occur at rest because, first, the resting membrane potential is close to the equilibrium potential for potassium; and second, the Na⁺/K⁺ pump (which is active at −5% of its maximum level in resting muscle) can prevent accumulation by returning K⁺ into the cell. In active muscle the situation is very different. There is a considerable efflux of K⁺ from the active muscle, largely as a result of repolarisation by the delayed rectifier K⁺ channels but also with a contribution from other K⁺ channels, with the result that K⁺ efflux exceeds Na⁺ influx. The K⁺ in the interstitium is diluted by an increased osmotic interstitial accumulation of water; however, very high local levels of K⁺ have been predicted and observed experimentally. The interstitial concentration of K⁺ is also in equilibrium with capillary blood. As the capillary membrane is permeable to K⁺ there is a flux of K⁺ from interstitium to blood, and the venous effluent from the muscle acts to reduce the local interstitial K⁺ concentration by raising that of the plasma. This equilibrium will be altered if there is any disturbance to muscle blood flow. Although plasma K⁺ will be influenced by the efflux from active muscle this is to a large extent cushioned by the uptake into other inactive muscle. Consequently, although plasma K⁺ does reflect muscular activity it is an imperfect indicator of local interstitial concentration. In severe exercise there is a generalised increase in activity of the Na⁺/K⁺ pump which ensures that plasma K⁺ does not usually exceed 5.5 mmol/l (although extreme values of 8.2 mmol/l have been recorded). This increased pumping is brought about by β adrenergic stimulation of the pump (for a fuller account see Sjøgaard and McComas).

Whereas increased Na⁺/K⁺ pump activity may be able to preserve a near normal transmembrane potential of the muscle cell it is very unlikely that this is the case in the T tubules. T tubule permeability to K⁺ is exceptionally high, and the very narrow lumen will restrict diffusion of the ion. Na⁺/K⁺ pumps are relatively scarce in the T tubule. Consequently, inward transmission of the action potential will become increasingly precarious. Also, some delay in activation of the centrally located myofibrils becomes possible. Thus interstitial K⁺ accumulation could cause the characteristic changes toward smaller and slower twitches. Only inward transmission of the impulse into the muscle fibre would be impaired; the EMG recorded from the muscle fibres, which is caused almost entirely by membrane depolarisation/repolarisation, would be little changed.

### Table 2 Factors associated with alterations in K⁺ and tremor size

<table>
<thead>
<tr>
<th>Condition</th>
<th>Potassium effect</th>
<th>Tremor effect</th>
<th>Treatment (if appropriate)</th>
<th>Explanatory note</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adrenaline and other β agonists</td>
<td>Reduces plasma K⁺ by pumping into muscle</td>
<td>Large increase</td>
<td>β₂ blockers</td>
<td>A</td>
<td>6,7,8,12,14,15,27</td>
</tr>
<tr>
<td>Hyperthyroidism</td>
<td>Plasma K⁺ is low</td>
<td>Large increase</td>
<td>β₂ blockers</td>
<td>B</td>
<td>14,15</td>
</tr>
<tr>
<td>Raised muscle temperature</td>
<td>Reduced interstitial K⁺ because of increased washout?</td>
<td>Large increase</td>
<td>NA</td>
<td>C</td>
<td>35</td>
</tr>
<tr>
<td>Age</td>
<td>Plasmas K⁺ generally lower in old age</td>
<td>Generalised increase in tremor in old age</td>
<td>NA</td>
<td>D</td>
<td>3,16</td>
</tr>
<tr>
<td>Ethanol withdrawal and delirium tremens (DT)</td>
<td>Low plasma K⁺ only in patients who develop DT</td>
<td>Large increase</td>
<td>β₂ blockers</td>
<td>E</td>
<td>17,18,19,20</td>
</tr>
<tr>
<td>Essential tremor</td>
<td>Not known</td>
<td>Moderate to large increase</td>
<td>Potassium G</td>
<td>G</td>
<td>23,24</td>
</tr>
<tr>
<td>Essential tremor</td>
<td>Not known</td>
<td>Decrease followed by increase</td>
<td>NA</td>
<td>H</td>
<td>25,26,27</td>
</tr>
<tr>
<td>Ischaemia and muscle activity</td>
<td>Interstitial K⁺ raised?</td>
<td>Large decrease</td>
<td>NA</td>
<td>I</td>
<td>This paper</td>
</tr>
<tr>
<td>Lowered muscle temperature</td>
<td>Interstitial K⁺ raised?</td>
<td>Large decrease</td>
<td>NA</td>
<td>J</td>
<td>4,35</td>
</tr>
<tr>
<td>Acute ethanol administration</td>
<td>Variable</td>
<td>Large decrease</td>
<td>NA</td>
<td>K</td>
<td>29,30</td>
</tr>
</tbody>
</table>

A: The peripheral β₂ receptor associated with tremor is one which controls sodium/potassium exchange (general stimulation of muscle Na⁺/K⁺ pumps), rather than one that has a direct effect on the contractile machinery. This hypothesis may explain the delayed tremor response. There are accounts of dramatic tremor and potassium changes with acute clonial poisoning.14,15

B: Thyrotoxic tremor has been attributed to a synergistic effect of thyroid hormones on adrenaline sensitivity because the tremor is symptomatically “cured” by β blockers. Our suggestion is that the tremor is a direct consequence of the low plasma K⁺ caused by the disease. β Blocks ameliorate the tremor by raising extracellular potassium concentration.

C: Sudden increases in muscle temperature greatly increase tremor size. It may be that the changes are partly consequent on temperature induced changes in blood flow. An increase in muscle blood flow may increase washout of interstitial K⁺ and keep the concentration low, thus enhancing tremor.

D: One study11 found “a large prevalence of … potassium deficiencies in the elderly, an observation we could not attribute to pathology or treatment”. The increased tremor of old age may be related to decreased interstitial K⁺.

E: The tremor of delirium tremens (DT) is indistinguishable from essential tremor. Three separate studies have shown that following alcohol withdrawal plasma or total body K⁺ was significantly reduced29 only in patients who developed DT. Resolution of the tremor was accompanied by return of the plasma K⁺ to normal values.

F: Two studies have reported a weak association between essential tremor size and circulating catecholamines.32–33 It would be worth investigating whether there is a systematic relation with plasma K⁺ in these patients.

G: Postural tremor is a common problem for patients taking lithium.34 Potassium has been employed successfully to treat these side effects.35

H: Vigorous exercise results in a subsequent prolonged increase in tremor.36-37 This may be attributable to the effects of associated sympathetic release of adrenaline on the Na⁺/K⁺ pump, tending to produce a lower interstitial K⁺ concentration, particularly but not exclusively in the muscles which have been active.

I: Ischaemia only reduces tremor when combined with muscular activity.

J: As in the case of heating (C) this has been attributed to a direct effect on the contractile apparatus. It is possible that a reduction in blood flow and reduced washout of K⁺ may also be a contributory factor.

K: Acute administration of alcohol causes a progressive and profound reduction of both essential tremor and interstitial potassium concentration. This may be attributable to the effects of associated sympathetic release of adrenaline on the Na⁺/K⁺ pump, tending to produce a lower interstitial K⁺ concentration. Even in the absence of tremor, this effect may be seen in patients who have been alcoholic and in whom tremor has been produced by alcohol withdrawal. The tremor of delirium tremens (DT) is indistinguishable from essential tremor.36-37 Three separate studies have shown that following alcohol withdrawal plasma or total body K⁺ was significantly reduced only in patients who developed DT. Resolution of the tremor was accompanied by return of the plasma K⁺ to normal values.
We suggest that under postural conditions the local concentration of $K^+$ may reach a level where it changes function in that part of the muscle. Postural conditions involve repetitive activation of a small population of small motor units, lack of movement of the muscle, and reduced or absent perfusion. The combination of increased $K^+$ efflux and diminished circulatory washout may produce a sufficiently high local interstitial $K^+$ to impair the force generation of the muscle fibres that are active. This impairment would act to reduce tremor as a posture is maintained, so in that respect it is not an undesirable feature, but rather an asset. Thus for the small forces associated with tonic postural maintenance the response of the muscle fibres involved will become very much less brisk and the response of the muscles will be blunted.

A depression of plasma $K^+$ will be associated with a generally decreased interstitial $K^+$ concentration. As a result, muscle function will be less easy to compromise and tremor size will be high. Therefore any factor that reduces plasma $K^+$ should be associated with large tremor size. Conversely, any factor that allows plasma $K^+$ to rise should blunt the muscle response and decrease tremor size. A search of published reports shows that many, if not all, conditions associated with changed tremor size have been found to involve altered plasma potassium (table 2). This provides considerable circumstantial evidence in support of our theory.

Conclusions

Many factors that affect tremor size are also known to affect plasma $K^+$ concentration, and by inference, interstitial $K^+$ concentration. This correlation has been reported previously but a cause and effect mechanism has not been proposed. A simple mechanism may link the two factors. By partial blockade of the T tubule, an increase in interstitial $K^+$ can cause muscle function to become changed in such a way that the fusion frequency is reduced. This will produce a much more pulsatile, predisposing to increased tremor. Our muscular output will be brisker and force generation will be more easy to compromise and tremor size will be high. Therefore any factor that reduces plasma $K^+$ concentration, and by inference, interstitial $K^+$ is not an undesirable feature, but rather an asset. Thus for the large peripheral compartment.

Acknowledgements

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