Human proprioceptive adaptations during states of height-induced fear and anxiety

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Davis JR, Horslen BC, Nishikawa K, Fukushima K, Chua R, Inglis JT, Carpenter MG. Human proprioceptive adaptations during states of height-induced fear and anxiety. J Neurophysiol 106: 3082–3090, 2011. First published September 14, 2011; doi:10.1152/jn.01030.2010.—Clinical and experimental research has demonstrated that the emotional experience of fear and anxiety impairs postural stability in humans. The current study investigated whether changes in fear and anxiety can also modulate spinal stretch reflexes and the gain of afferent inputs to the primary somatosensory cortex. To do so, two separate experiments were performed on two separate groups of participants while they stood under conditions of low and high postural threat. In experiment 1, the proprioceptive system was probed using phasic mechanical stimulation of the Achilles tendon while simultaneously recording the ensuing tendon reflexes in the soleus muscle and cortical-evoked potentials over the somatosensory cortex during low and high threat conditions. In experiment 2, phasic electrical stimulation of the tibial nerve was used to examine the effect of postural threat on somatosensory evoked potentials. Results from experiment 1 demonstrated that soleus tendon reflex excitability was facilitated during states of height-induced fear and anxiety while the magnitude of the tendon-tap-evoked cortical potential was not significantly different between threat conditions. Results from experiment 2 demonstrated that the amplitudes of somatosensory-evoked potentials were also unchanged between threat conditions. The results support the hypothesis that muscle spindle sensitivity in the triceps surae muscles may be facilitated when humans stand under conditions of elevated postural threat, although the presumed increase in spindle sensitivity does not result in higher afferent feedback gain at the level of the somatosensory cortex.

posturography; somatosensory-evoked potential; balance; emotion

Recent evidence has highlighted the potential for emotions, such as fear and anxiety, to influence human balance control. For example, young and older adults adopt different postural control strategies depending on the level of fear they experience when standing on the edge of an elevated surface height (Carpenter et al. 1999, 2001, 2006; Adkin et al. 2000, 2002; Davis et al. 2009; Huffman et al. 2009). However, the exact neural mechanisms that contribute to the observed changes in postural control remain unclear.

There is some evidence to suggest that changes in muscle spindle sensitivity may provide one mechanism through which fear and anxiety may influence postural control. For example, significant decreases in soleus (SOL) H-reflex amplitude have been observed when individuals stand quietly on elevated surfaces (Sibley et al. 2007) or when they are engaged in a threatening task such as walking along a raised narrow beam (Llewellyn et al. 1990). This decrease in SOL H-reflex amplitude has been attributed to an increased tonic Ia discharge of muscle spindles that results in homosynaptic postactivation depression of the SOL lower motor neuron pool (Sibley et al. 2007). Other evidence suggests that muscle spindles become more sensitive to stretch under threatening conditions; subjects standing on high compared with low surface heights demonstrated a significant increased amplitude of soleus tendon reflex (STR), without any accompanying changes in H-reflex amplitude (Horslen 2010). Similar observations of increased STR have been observed in seated individuals when presented with other sources of arousing stimuli (Bonnet et al. 1995; Both et al. 2005; Kamibayashi et al. 2009; Hjorstkov et al. 2005).

One explanation that has been offered for why spindle sensitivity may be increased in threatening or novel conditions is that it may “provide supraspinal areas with increased feedback gain and resolution” (Ll Lewellyn et al. 1990). While there is current evidence indicating that later cortical potentials (i.e., N100) evoked by balance perturbations are increased under conditions of postural threat (Adkin et al. 2008; Sibley et al. 2010), the extent to which threat can influence the gain of afferent information initially received at the primary somatosensory cortex is currently unknown.

It is possible to determine whether an increased STR excitability also results in a larger magnitude afferent volley to the somatosensory region of the parietal cortex by simultaneously evoking STRs and cortical potentials (TEPs) via mechanical stimulations of SOL using tendon taps (Cohen et al. 1985; Frascharelli et al. 1993), although this technique has yet to be applied to standing individuals. Conversely, somatosensory-evoked potentials (SEPs) elicited by mild electrical stimulation of the tibial nerve allow for changes in the sensitivity of the somatosensory cortex to be measured independent of any changes in muscle spindle sensitivity (Gandevia and Burke 1984; Nelson et al. 2000; Staines et al. 2000; McIlroy et al. 2003).

Therefore, two experiments were performed to achieve the two aims of this study. Experiment 1 was designed to investigate whether elevated muscle spindle sensitivity during states of height-induced fear and anxiety results in facilitated spinal reflexes and a concomitant increase in the gain of afferent information delivered to the somatosensory cortex by comparing changes in STR excitability and TEP amplitude. Experiment 2 was designed to investigate the potential change in cortical sensitivity to incoming afferent information during states of height-induced fear and anxiety, independent of any changes in muscle spindle sensitivity, by comparing changes in SEP amplitude.

Based on previous observations (Horslen 2010; Bonnet et al. 1995; Hjorstkov et al. 2005; Kamibayashi et al. 2009; Both et al. 2005; Llewellyn et al. 1990), it was hypothesized that there...
would be a significant increase in both TEP and STR amplitude when standing under conditions of elevated postural threat compared with when standing under conditions of low postural threat. Furthermore, it was hypothesized that there would be a null effect of threat manipulation on SEP amplitude reflecting no change in the sensitivity at the level of the somatosensory cortex to incomingafferent information, independent of any change in spindle sensitivity.

METHODS

Participants

Two independent groups of participants volunteered for two separate experimental protocols in this study. In experiment 1, 35 young healthy adults (17 female, means ± SE: age 22.6 ± 0.6 yr, height 173.9 ± 1.7 cm, and weight 70.6 ± 1.8 kg) volunteered to participate. In experiment 2, 31 young healthy adults (16 male, means ± SE: age 24.9 ± 0.8 yr, height 174.5 ± 0.9 cm, and weight 68.9 ± 0.4 kg) volunteered to participate. All participants were recruited from the local undergraduate and graduate student community. Each participant completed a survey of their relevant medical history before testing. All participants were free of any relevant neurological, vestibular, and/or orthopedic conditions and were not taking any prescription medications that may have affected their balance performance during the study. Each participant provided written informed consent before testing. The University of British Columbia Clinical Research Ethics Board approved all experimental procedures.

Experiment 1: Mechanically Evoked Somatosensory Potentials (TEPs)

In experiment 1, participants were separated into one of two groups: Free Standing or Braced Standing. During testing, those in the Free Standing group stood unsupported during all experimental procedures, whereas those in the Braced Standing group stood in custom-made ankle braces designed to immobilize the ankle joint in the anterior-posterior (A-P) direction (Fig. 1). The ankle braces were used to prevent the characteristic posterior lean that occurs when participants stand under conditions of height-induced postural threat (Davis et al. 2009). By maintaining a constant ankle angle in the Braced Standing group throughout the experiment (during both testing and rest periods), we attempted to control for any lean-associated changes to prevent the characteristic posterior lean that occurs when participants are instructed to lean backward slightly against the back of a chair that provided some postural support.

A computer driven magnetic linear motor (E2000-AT; LIN Mot) was used as the tendon hammer to deliver mechanical taps to the Achilles tendon. The tendon hammer travelled a distance of 1 cm and took 13 ms to reach peak displacement. The distance between the tendon hammer and skin was measured at the beginning of each block of testing in each threat condition to ensure that a constant distance of 5 mm was maintained between the tendon hammer and the skin over the Achilles tendon throughout the experiment. Based on findings from initial pilot work, taps were applied to the Achilles tendon at an interstimulus interval of 1.2–1.7 s to ensure that any postural perturbation caused by a given tendon tap and/or tendon reflex response fully subsided before a subsequent tap was delivered.

The force of each tap was recorded with a dynamic force sensor (Isotron Dynamic Force Sensor and Conditioner; Endeveco) mounted on the contact surface of the tendon hammer. The force of each tap was amplified 10 times and sampled at 1,000 Hz (Spike 5; CED). Clinical investigations of STR sensitivity have demonstrated that a tendon tap force ranging from 21 to 50 N is adequate to elicit an STR in healthy young adults (Marshall and Little 2002). Therefore, the force of each tap was kept within this range for all participants included in the study. To evoke an optimal STR, the tendon hammer contact location was adjusted in the vertical plane to ensure that the tap force for each participant was kept above sensory perception threshold and evoked a clear STR. The force of each tap was monitored online during testing to ensure that a consistent tap force was delivered between threat conditions and was recorded for offline analysis.

Fig. 1. Experimental setup used to support stance in the braced standing condition. Participants in the Braced Standing condition stood in custom-made ankle braces designed to attenuate postural sway in the anterior-posterior direction.

While standing in each threat condition, participants in both the Free Standing and Braced Standing groups received 2 blocks of 120 mild mechanical taps delivered to their right Achilles tendon for a total of 240 taps per threat condition. A 30-s rest period was provided between blocks during which participants were instructed to lean backward slightly against the back of a chair that provided some postural support.

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Physiological and psychosocial estimates of height-induced fear and anxiety. Electrophysiological activity (EDA) of the nondominant hand was used to provide an estimate of the level of physiological arousal participants experienced during each threat condition. Participants were fitted with disposable recording electrodes on the thenar and hypothenar eminences to measure their skin conductance (2502 Skin Conductance Unit; CED) with a range of 0–100 μS. The skin conductance signal was collected at a sampling frequency of 1,000 Hz. EDA was averaged across the first 60 s of each trial and subsequently averaged across trials for each threat condition.

Before they were tested in each threat condition, participants were instructed to rate how confident they were that they would be able to maintain their balance and avoid a fall during the threat condition on a scale of 0% (no confidence) to 100% (complete confidence) (Adkin et al. 2002). Immediately following each standing trial, participants were instructed to rate how stable they felt during the trial on a scale of 0% (very unstable) to 100% (completely stable) and how fearful of falling they were during the trial again on a scale of 0% (completely unafraid) to 100% (extremely afraid) (Adkin et al. 2002). Participants also completed a 16-question survey of their perceived anxiety modified from Adkin et al. (2002). The scores from all 16 questions were summed to generate a total perceived anxiety score for each threat condition.

Statistical analysis. All dependent variables were compared between standing conditions (Low Threat and High Threat) and between groups (Free Standing and Braced Standing) using 2 × 2 (condition × group) mixed design ANOVAs (SPSS, IBM). Within-subjects effect sizes (Cohen’s d values) were calculated for each dependent variable as well. The criteria for statistical significance was set to $P < 0.05$.

Experiment 2: Electrically Evoked Somatosensory Potentials (SEPs)

In experiment 2, participants stood under the same threat conditions described in experiment 1: Low Threat and High Threat. While standing in each threat condition, participants received 2 blocks of 120 mild cutaneous electrical stimulations ($48$ with SIU5 stimulus isolation unit and CCU1C constant current unit; Grass Instruments) separated by a rest period of ~30 s for a total of 240 stimuli per threat condition (see details below). All participants in experiment 2 stood unsupported; a braced standing group was not included. The rationale for not including a braced standing group was based on the fact that the location of SEP stimulation, the popliteal fossa, is proximal to the lower limb musculature. Therefore, the ascending afferent volley evoked by the cutaneous electrical SEP stimulation would not have been subject to any changes in muscle physiology or lower motor neuron pool excitability associated with the height-induced posterior lean.

Before the experiment, all participants performed a practice trial with the hydraulic lift resting at its lowest height of 0.8 m to remove potential first trial effects. The order of presentation of Low and High Threat conditions was counterbalanced across participants to minimize potential order effects. At the end of the experiment, participants performed a posttest standing control trial in the Low Threat condition that was used to confirm the stability of the SEP signals over time.

COP measures. Ground reaction forces and moments were sampled (no. K00407; Bertec) at 100 Hz (Power 1401; CED) and low-pass filtered offline using a 5-Hz dual-pass Butterworth filter (MatLab; Mathworks) before calculating COP in the A-P direction. Mean position of the COP during each trial was calculated in the A-P direction and subtracted from the COP signal. From this unbiased signal, the root mean square and mean power frequency of COP displacement were calculated in the A-P direction. Note that COP was not measured in experiment 1. Pilot testing revealed that the braces influenced the ability to measure normal COP displacements in the Braced Standing group, and initial force from the hammer, and resulting STR, both represented dynamic postural perturbations that confounded the COP signal in the Free Standing group.
SEP stimulation procedure. SEPs were evoked by electrical stimulation of the right tibial nerve. The anode (10 × 3 cm, coal rubber pad; AMG Medical) was placed just superior to the patella, and the Ag-AgCl cathode (0.25 cm in diameter; Kendall) was placed in the popliteal fossa. To determine the optimal SEP stimulation intensity for each participant, a series of SOL H reflexes were elicited via a submaximal 0.5-ms square wave pulse to the tibial nerve. The SEP stimulus intensity was set to 40–60% of the intensity required to elicit a small M wave, as this range of intensities has been shown to provide an optimal range for the SEP to fluctuate within and not become saturated (Gandevia and Burke 1984). However, this intensity was just above sensory perception threshold for each participant and was not sufficient to elicit a reliable SOL H reflex or M wave in our participants. Therefore, to ensure that the intensity of the stimulation to the tibial nerve did not change between standing conditions, a M-wave test pulse equal to 50% of M-max (DS5; Digitimer UK) was evoked before and immediately following each 120-s standing trial in a subgroup of 10 participants. The peak to peak amplitude of each evoked M wave was monitored to confirm the consistency of the stimulation over the duration of the stimulus.

SEP recording and analysis procedure. A different recording system from the one used in experiment 1 was used to collect EEG in experiment 2. Sintered Ag/AgCl scalp electrodes (EASYCAP) were placed along the sagittal midline at the Cz and Fpz’ (2 cm caudal to Fpz) locations on the participant’s scalp according to the international 10/20 system for EEG recordings. A ground electrode was placed on the back of the participants’ neck (5 × 5 cm, coal rubber pad; AMG Medical). The impedance between all three electrodes was tested before and after testing (Grass EZM5 impedance meter; Grass Instruments) to ensure that the electrode impedance was kept at <10 KΩ during testing. The EEG signal was amplified 20,000 times, sampled at 10,000 Hz, and band-pass filtered between 1 and 1,000 Hz (Grass, P511 AC amplifier). Each recording was visually inspected offline to remove any stimuli that may have been contaminated by artifacts related to eye blinks or facial contractions. The remaining stimuli were used to create a spike-triggered average (Spike 5, CED, UK) of the SEP signal between 100 ms before and 150 ms after the stimulus onset. The ensemble spike-triggered average was baseline corrected using the mean of the signal 100 ms before stimulus onset and band-pass filtered between 1–100 Hz using customized software (LabVIEW; National Instruments). P1 and N1 peaks of the SEP waveform were identified and used to calculate time to peaks and the peak-peak amplitude of P1-N1 for each subject.

Two surface Ag/AgCl EMG electrodes were applied in a belly-tendon preparation on the right SOL muscle to record the SOL M-waves used for stimulus calibration. Two surface Ag/AgCl electrodes were placed ∼2 cm apart in a belly-belly preparation to collect background EMG activity in SOL. All EMG signals were amplified 2,000 times, sampled at a 1,000 Hz, and band-pass filtered between 30 and 300 Hz (P511 AC Amplifier; Grass Instruments).

Physiological and psychosocial estimates of height-induced fear and anxiety. Both EDA and psychosocial estimates of height-induced fear and anxiety were recorded in the same manner described for experiment 1.

Statistical analysis. All dependent variables were compared between Low and High Threat conditions using paired-samples t-tests (SPSS, IBM). Paired t-tests were also used to compare SEP amplitudes measured at the low height during the experiment and the posttest control trial. Within-subjects effect sizes (Cohen’s d values) were calculated for each dependent variable as well. Correlation analysis was performed using a Pearson correlation on the changes between the High Threat and Low Threat conditions for the physiological and posturographic variables. The criteria for statistical significance was set to $P < 0.05$.

**RESULTS**

**Experiment 1: Mechanically Evoked Somatosensory Potentials (TEPs)**

In the Low Threat condition, the TEP waveform was characterized by a distinct P1-N1 component (Fig. 2A). The mean time to peak for the P1 and N1 peaks occurred at 35.46 ± 0.92 and 48.12 ± 0.96 ms, respectively, following stimulus onset. The mean peak-peak amplitude of the P1-N1 was 4.80 ± 0.61 μV. A similar TEP waveform was observed in the High Threat condition. There were no observable differences in the grand average waveforms between threat conditions with respect to the timing or amplitude of the P1-N1 component of the TEP waveform (Fig. 2A).

The lack of observable differences in TEP timing and amplitude between Low and High Threat conditions was confirmed statistically. A mixed design ANOVA revealed no significant difference between threat conditions for the time to peak of the P1 [$F(1,22) = 2.311; P = 0.143; \delta = 0.13$] and N1 [$F(1,22) = 0.335; P = 0.568; \delta = 0.07$] when standing in the High Threat compared with Low Threat condition (Table 1). There was also no significant main effect of threat condition on
the amplitude of the P1-N1 component \(F(1,22) = 0.904; \ P = 0.352; \ \delta = 0.07\) of the TEP waveform (Table 1). However, there was a significant main effect of group on the peak-peak amplitude of the P1-N1 \(F(1,22) = 6.302; \ P = 0.020; \ \delta = 0.74\) whereby larger P1-N1 amplitudes were observed in the Braced Standing group compared with the Free Standing group. There were no significant interaction effects between threat condition and group.

Despite very little change in the observed TEP (Fig. 2B), there was a distinct increase in the STR peak-peak amplitude in the High Threat compared with the Low Threat condition (Fig. 2C). This observation was confirmed statistically across participants. There was a significant main effect of threat condition on both the peak-peak amplitude \(F(1,22) = 4.430; \ P = 0.047; \ \delta = 0.12\) and peak latency \(F(1,22) = 5.109; \ P = 0.034; \ \delta = 0.15\) of the STR. The peak-peak amplitude of the STR was significantly larger and occurred significantly earlier when standing in the High Threat condition compared with the Low Threat condition, independent of whether participants were in the Free Standing or Braced Standing group (Fig. 3, A and B). Furthermore, there was a significant main effect of threat condition on both the mean level of background SOL EMG activity \(F(1,22) = 9.547; \ P = 0.005; \ \delta = 0.30\) and mean level of TA EMG activity \(F(1,22) = 9.895; \ P = 0.005; \ \delta = 0.48\). In both the Free Standing and Braced Standing group, there was a significant decrease in the mean level of background SOL activity and a significant increase in mean level of background TA activity when participants stood in the High Threat compared with the Low Threat condition (Fig. 3C).

There was a significant main effect of threat condition on the level of confidence \(F(1,22) = 62.64; \ P < 0.001; \ \delta = 1.57\) and perceived anxiety \(F(1,22) = 24.06; \ P < 0.001; \ \delta = 0.085\) and fear \(F(1,22) = 33.51; \ P < 0.001; \ \delta = 1.17\). As shown in Table 1, there was a significant increase in both self-reported fear and perceived anxiety and a significant decrease in self-reported confidence when standing in the High Threat condition compared with the Low Threat condition.

There was also a significant main effect of height observed on EDA \(F(1,22) = 29.498; \ P < 0.001; \ \delta = 0.54\) whereby participants demonstrated higher EDA in the High Threat compared with the Low Threat condition (Table 1).

**Experiment 2: Electrically Evoked Somatosensory Potentials (SEPs)**

In the Low Threat condition, the mean SEP waveform was characterized by a P1 peak with a mean latency of 38.98 ± 1.44 ms following stimulus onset, followed by an N1 peak at 49.61 ± 1.78 ms (Fig. 4A). In the High Threat condition, the mean latency of the P1 peak was 38.98 ± 1.38 ms following stimulus onset, followed by an N1 peak at 50.55 ± 2.04 ms. There was no observable difference between threat conditions in the time to peak for any component of the SEP and there was

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**Table 1. Mean change of all neurophysiological, physiological, and psychosocial measures observed between the Low Threat and High Threat conditions**

<table>
<thead>
<tr>
<th>Dependent Measure</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neurophysiological measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P1-N1 amplitude, (\mu V)</td>
<td>4.80 ± 0.61</td>
<td>5.03 ± 0.63</td>
</tr>
<tr>
<td>P1 latency, ms</td>
<td>35.46 ± 0.92</td>
<td>34.87 ± 0.94</td>
</tr>
<tr>
<td>N1 latency, ms</td>
<td>48.12 ± 0.96</td>
<td>47.84 ± 0.80</td>
</tr>
<tr>
<td><strong>Physiological measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean EDA, (\mu S)</td>
<td>18.11 ± 1.91*</td>
<td>24.72 ± 2.50*</td>
</tr>
<tr>
<td>Mean SOL EMG, mV</td>
<td>0.012 ± 0.002*</td>
<td>0.009 ± 0.002*</td>
</tr>
<tr>
<td>Mean TA, mV</td>
<td>0.007 ± 0.002*</td>
<td>0.015 ± 0.002*</td>
</tr>
<tr>
<td>M-wave amplitude, mV</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><strong>Psychological measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perceived anxiety (sum score)</td>
<td>28.36 ± 1.75*</td>
<td>45.41 ± 4.01*</td>
</tr>
<tr>
<td>Fear, %</td>
<td>2.92 ± 1.27*</td>
<td>24.79 ± 3.32*</td>
</tr>
<tr>
<td>Confidence, %</td>
<td>96.87 ± 0.94*</td>
<td>77.71 ± 2.50*</td>
</tr>
<tr>
<td><strong>Posturographic measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean A-P position, mm</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>A-P MPF, Hz</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>A-P RMS, mm</td>
<td>–</td>
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</tr>
</tbody>
</table>

Data are means ± SE. P1 and N1, peaks of tendon tap-evoked potential waveform; EDA, electrodermal activity; SOL, soleus; EMG, electromyogram; TA, tibialis anterior; A-P, anterior-posterior; MPF, mean power frequency; RMS, root mean square. *\(P < 0.05\), significant main effects of threat condition.
The primary aims of this study were to determine whether muscle spindle sensitivity increases during states of height-induced fear and anxiety and whether this increase in spindle sensitivity results in higher afferent feedback gain at the level of the somatosensory cortex. The results from experiment 1 demonstrated that STR excitability was facilitated and STR latency was reduced when participants stood in the High Threat condition compared with the Low Threat condition. These results are consistent with the facilitation of STR observed in subjects that were seated under conditions of increased arousal (Bonnet et al. 1995; Both et al. 2005; Kamibayashi et al. 2009; Hjorstkov et al. 2005). We would argue that our findings cannot be explained by posterior leaning or consequent changes in length or tension of the muscle/tendon or intrafusal muscle fibres, since similar threat-related changes in STR were observed in both the Braced and Unbraced Standing group. Moreover, the observed increase in STR excitability in the High Threat condition cannot be explained by changes in background EMG activity because standing in the High Threat condition was associated with a significant decrease in SOL and a significant increase in TA EMG activity independent of bracing condition, and such a pattern of change in EMG background activity would lead to inhibition of the STR (Crone et al. 1987), as opposed to an increase in STR as currently observed. In this regard, the results suggest that not only is SOL STR excitability increased in High Threat conditions, but the increase is sufficiently large enough to overcome any inhibitory influence of the observed changes in background SOL and TA activity. Alternatively, it could be argued that the increase in STR amplitude under threatening conditions is caused by a decrease in presynaptic inhibition acting at the level of the spinal cord. However, evidence by Morita et al.
(1998) have demonstrated that the T reflex is less susceptible to influences of PSI, compared with the H-reflex amplitude, and there was no evidence for decreased H-reflex amplitudes in studies that have controlled for postural leaning (Horslen 2010). Therefore, based on the current evidence, the most likely explanation for the observed increase in amplitude, and decrease in latency, of STR is an increase in muscle spindle sensitivity during conditions of increased threat.

Supraspinal Ia Afferent Signal Attenuation

Contrary to our original hypothesis, facilitation of STR excitability in the High Threat condition was not accompanied by an increase in the afferent signal recorded at the somatosensory cortex; no changes in TEP amplitude were observed between Low and High Threat conditions. Furthermore, the results from experiment 2 clearly show that the sensitivity of the somatosensory cortex to incoming afferent information does not change during states of height-induced fear and anxiety as demonstrated by the null effect of threat condition on SEP amplitude. Instead, it appears that the heightened Ia volley associated with the increase in muscle spindle sensitivity and STR excitability is somehow gated or dampened at the somatosensory cortex- or along the posterior-column medial-lemniscus pathway.

Gating of afferent sensory afferent information has been previously observed during different phases of gait or voluntary ankle movement (Abbruzzese et al. 1981; Rushton et al. 1981; Dietz et al. 1985; Duyssens et al. 1995; Morita et al. 1998), perhaps as a means to facilitate voluntary motor control in such situations (Morita et al. 1998). Therefore, it is plausible that a similar gating mechanism may be activated to facilitate greater cortical control over posture during states of height-induced fear and anxiety (Huffman et al. 2009). However, if sensory gating or cortical inhibition was indeed responsible for the results of experiment 1, an attenuation of the electrically evoked SEP would have been expected to be observed in experiment 2, which was not the case. The lack of observable changes in SEP amplitudes with postural threat is consistent with previous reports that early auditory-evoked potentials are not influenced by postural threat (Adkin et al. 2008). This is contrary to later cortical potentials, such as the N100, that are significantly modulated by postural threat (Adkin et al. 2008; Sibley et al. 2010) or emotion (Bar-Haim et al. 2005; Eysenck et al. 1987).

One limitation of the current study is that TEPs and SEPs were evoked in different experimental populations. However, this protocol was deemed necessary to balance the needs of acquiring a sufficiently large number of TEP and SEP stimuli to generate reliable ensemble averages, while minimizing the time for subjects to habituate to the threat. Future studies could address this limitation by eliciting TEPs and SEPs from the same individuals. Likewise, further investigations of sensory gating and threat could incorporate recordings of evoked potentials from surface electrodes placed at different levels of the spine (Cameron et al. 2008; Murakami et al. 2008) or via indwelling thalamic electrodes in clinical populations (e.g., Katayama and Tsubokawa 1987; Klostermann et al. 2002) during states of height-induced fear and anxiety.

Utility of Studying Changes in TEP Amplitude during Stance

Only two studies have examined TEPs in humans. Cohen et al. (1985) was the first to descriptively compare SEPs and TEPs evoked in participants lying prone. Similarly, the magnitudes of TEP responses to mechanical stretch of the Achilles tendon have been compared between the paretic and functional limbs of hemiparetic patients while lying prone (Frascarelli et al. 1993).

Despite the observed null effect of height-induced fear and anxiety on TEP amplitude in the current study, the results provide the first evidence of TEPs recorded during stance. The mean peak-peak TEP P1-N1 amplitude (4.80 ± 0.61 μV) and latency (35.46 ± 0.92 ms) measured during stance in the Low Threat condition are comparable to measures recorded from prone individuals by Cohen et al. (1984) (3.2 ± 3.1 μV amplitude; 27–37 ms latency) and Frasarelli et al. (1993) (3.67 ± 0.44 μV; no latency reported).

Of note, however, was the observed main effect of bracing on TEP amplitude, with increased TEPs observed in the Braced compared with Unbraced Standing condition. One possible explanation for the difference may be the additional cutaneous inputs provided by the braces. However, this explanation is unlikely, since prior evidence has shown that decreasing cutaneous sensitivity through cooling had no effect on TEP amplitude (Cohen and Starr 1985). Alternatively, an increase in the sensitivity of the cortex may have been required to stand in the braces, as observed when subjects have been asked to control balance in other novel or challenging conditions (Staines et al. 2000). Since the SEP experiment did not involve a braced condition in this study, we cannot address this possibility; however, the potential for TEPs to be modulated by such factors indicates that further research needs to be applied to this technique, within a standing model in particular.

Potential Mechanisms and Significance of Increased Muscle Spindle Sensitivity

If the increase in muscle spindle sensitivity does not serve to increase the gain of afferent input received at the somatosensory cortex as originally hypothesized the question remains as to “how” and “why” the spindles are made more sensitive to stretch under conditions of increased threat.

As to how muscle spindle sensitivity may be increased during conditions of increased threat, there are at least two possible mechanisms. One possible mechanism involves direct connections between the sympathetic nervous system and muscle spindles. There is evidence that feline muscle spindles receive direct innervation from the autonomic nervous system (Barker and Saito 1981) that facilitates increases in muscle spindle firing rates when activated (Hunt 1960). Likewise, performing mental arithmetic or a static handgrip contraction to directly activate the human sympathetic nervous system has been shown to facilitate stretch reflex excitability (Hjorstovk et al. 2005; Kambayashi et al. 2009). Therefore, the increased sympathetic response (indexed by elevated EDA) observed during the High Threat condition may have contributed to increased SOL muscle spindle sensitivity and facilitation of STR excitability in the current study. Although some studies have demonstrated that sympathetic nervous system activation does not facilitate tonic muscle spindle firing rates in semi-intact animal preparations (Passatore et al. 1996; Hellström et
al. 2005) or in humans (Macefield et al. 2003; Birznieks et al. 2008), it should be noted that in these studies, the muscles that were tested were not posturally relevant or tested under a loaded condition.

Alternatively, the observed increase in spindle sensitivity and STR reflex facilitation observed may occur as a result of heightened gamma-motor drive to SOL muscle spindles (Prochazka et al. 1976, 1985, 1988; Ribot-Ciscar et al. 1986; Hospod et al. 2007). Evidence from cats has shown changes in Ia afferent firing patterns under difficult or threatening conditions (i.e., standing on a beam or ladder), which are consistent with patterns caused by activation of gamma static and dynamic neurons (Prochazka et al. 1985, 1988). While some studies have also provided evidence in support of alpha-gamma decoupling in humans (Ribot-Ciscar et al. 1986; Hospod et al. 2007), others have argued that selective modulation of gamma-motor neuron activity in humans is unlikely (Gandevia et al. 1997). Therefore, more detailed experiments are clearly required to establish the potential influence of sympathetic drive and or alpha-gamma decoupling on muscle spindle sensitivity during stance.

At this point we can also only speculate as to why the muscle spindle sensitivity may increase under conditions of increased threat. For example, it may be part of a more global fight/flight response, just as the sensitivity of other sensory receptors are known to increase with sympathetic activation (Hall 2011). Alternatively, increased muscle spindle sensitivity under conditions of increased postural threat may serve a functional role in maintaining postural control. For example, the strategy of stiffening the ankle joint to maintain a tighter control of the center of mass during quiet stance has been frequently observed in conditions of increased postural threat (Carpenter et al. 1999, 2001) and would be potentially facilitated by increased gain of spinal stretch reflexes in the soleus muscles. How effective this may be, is questionable, particularly in light of some arguments that the small ankle rotations during quiet stance may be taken up mostly by the tendon, creating a "dead-zone" in which the spindles are incapable of detecting the small changes involved in swaying (Loram et al. 2009). More likely, the increased muscle spindle sensitivity could be used to defend against unexpected postural perturbations to balance, as a stretch of the soleus muscle would be countered by a larger, more rapid, plantar-flexion torque to move the center of mass away from the edge of the high surface height. Although, observations from unexpected platform rotations failed to demonstrate increases in monosynaptic reflexes under conditions of increased threat, the observations of larger automatic postural responses initiated within 100 ms of an unexpected platform rotation (Carpenter et al. 2004), suggest that increased afferent gain may be used to modulate long-latency reflex pathways. If this is the case, the current observations suggest that this afferent input might be used to augment spinal-bulbar or spino-cerebellar reflex pathways, since no changes were observed in either TEPs or SEPs recorded at the cortical level.

Conclusion

The current study provides further evidence that muscle spindle sensitivity in the triceps surae muscles may be facilitated when humans stand under conditions of elevated postural threat (Horslen 2010; Sibley et al. 2007). The increase in spindle sensitivity observed in High Threat conditions did not produce a concomitant increase in the magnitude of theafferent volley received by the somatosensory cortex nor was it accompanied by an observable change in the cortical sensitivity to incoming afferent inputs generated by electrical stimulation of the nerve above the spindle. These findings challenge the notion that muscle spindle sensitivity is increased under difficult or threatening postural conditions to increase the gain of somatosensory information received by the cortex, instead, threat-induced facilitation of the ascending afferent volley appears to be gated or inhibited at the level of the brainstem, thalamus or cortex. Although the exact mechanism is currently unknown, the results of the study provide new insight into how proprioception may contribute to threat-induced changes in static and dynamic postural control and highlight the need for further study of the neurophysiological outcomes of fear and anxiety.

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AUTHOR CONTRIBUTIONS


DISCLOSURES

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