Cued Spatial Attention Drives Functionally Relevant Modulation of the Mu Rhythm in Primary Somatosensory Cortex

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Cued Spatial Attention Drives Functionally Relevant Modulation of the Mu Rhythm in Primary Somatosensory Cortex

Stephanie R. Jones, Catherine E. Kerr, Qian Wan, Dominique L. Pritchett, Matti Hämäläinen, and Christopher I. Moore

Cued spatial attention modulates functionally relevant alpha rhythms in visual cortices in humans. Here, we present evidence for analogous phenomena in primary somatosensory neocortex (SI). Using magnetoencephalography, we measured changes in the SI mu rhythm containing mu-alpha (7–14 Hz) and mu-beta (15–29 Hz) components. We found that cued attention impacted mu-alpha in the somatopically localized hand representation in SI, showing decreased power after attention was cued to the hand and increased power after attention was cued to the foot, with significant differences observed 500–1100 ms after cue. Mu-beta showed differences in a time window 800–850 ms after cue. The visual cue also drove an early evoked response beginning ~70 ms after cue with distinct peaks modulated with cued attention. Distinct components of the tactile stimulus-evoked response were also modulated with cued attention. Analysis of a second dataset showed that, on a trial-by-trial basis, tactile detection probabilities decreased linearly with prestimulus alpha modulation. These results support the growing consensus that cue-induced alpha modulation is a functionally relevant sensory gating mechanism deployed by attention. Further, while cued attention had a weaker effect on the allocation of mu-beta, oscillations in this band also predicted tactile detection.

Introduction

The historical view of electroencephalography/magnetoencephalography (EEG/MEG)-measured alpha rhythms (7–14 Hz) as a “resting” brain state is being challenged by evidence that they are actively and topographically deployed to gate information processing. Cued spatial attention leads to decreased alpha amplitudes in parietal-occipital EEG sensors contralateral to the attended site in visual (Worden et al., 2000; Kelly et al., 2006, 2009; Thut et al., 2006) and intersensory visual–auditory (Foxe et al., 1998; Fu et al., 2001) tasks. Alpha decreases are accompanied by increases in opposing hemifields, and lateralized alpha amplitudes predict reaction times and visual discriminability (Worden et al., 2000; Kelly et al., 2006, 2009; Thut et al., 2006). Recent work has also shown that attentional biases are tied to changes in components of the broadband cue-induced evoked response (ER) in early visual cortices (Kelly et al., 2009).

In the somatosensory neocortex in humans, a spontaneous mu rhythm containing a complex of mu-alpha (7–14 Hz) and mu-beta (15–29 Hz) components is commonly observed above rolandic cortex (Tiihonen et al., 1989; Jones et al., 2009). These rhythms show an event-related desynchronization (ERD) with stimulation or movement, and in premovement periods, with a subsequent synchronization (ERS). There is an increase in alpha band ERS over the sensorimotor neocortex during visual processing, with a simultaneous alpha ERD over visual cortices, and vice versa during movement (Pfurtscheller, 1992) (see also Rouguel et al., 1979). Further, premovement alpha ERD is accentuated in elite athletes over the entire brain, suggesting that this is a functionally relevant mechanism that can be enhanced with practice (Del Percio et al., 2009).

Several studies have shown that selective somatic attention impacts movement- or sensory-induced ERD and ERS changes in sensorimotor mu-alpha and mu-beta activity (Bauer et al., 2006; Babiloni et al., 2008; Dockstader et al., 2010). However, allocation of these rhythms following an attentional cue, in anticipation of tactile sensory processing, has not been investigated.

In the present study, we used MEG imaging to investigate whether similar effects to those observed in the visual system are also present in somatosensation, using the well localized hand representation in SI as our substrate (Jones et al., 2007, 2009). We investigated whether spatial attention directed to or away from
the hand impacted allocation of mu-alpha and mu-beta, the predictive value of these rhythms for detection, and the impact of cued attention on evoked responses. We observed an impact in each of these dimensions, indicating that the decreased expression of localized alpha oscillations could be causally beneficial to attentional regulation employed across neocortical areas.

Materials and Methods

Stimuli
Subjects
MEG data were collected from 12 neurologically healthy (exclusion criteria included musculoskeletal diseases, arthritis, lupus, multiple sclerosis, scleroderma, and diagnosed current psychiatric disorder), right-handed, 18- to 50-year-old adults (mean age = 31.6 years, SD = 7 years, 1 man and 11 female). Subjects were medication free or on stable doses of selective serotonin reuptake inhibitor medication. The experimental protocol was approved by the Massachusetts General Hospital Internal Review Board, and each subject gave informed consent before data acquisition.

Stimuli
Subjects’ hand and foot rested on solid plastic frames through which tactile stimuli were delivered. The stimulus (single cycle of a 100 Hz sine wave, 10 ms duration) was generated by fused multilayer piezoelectric benders, which provide more favorable force and higher-frequency resonance characteristics than typical ceramic wafers (Noliac) (see Jones et al., 2007). Stimuli were applied to the distal pads of the third digit of the left hand and first digit of the left foot via a Delrin contactor affixed to the piezoelectric bender (7 mm diameter presented within a 1 cm circular rigid surround). The device was not glued to the skin. Instead, matched intensity of stimulation, relative to perceptual threshold, was maintained individually for each subject using a parameter estimation sequential testing (PEST) convergence procedure (Dai, 1995; Leek, 2001). During the cued detection runs described below, stimulus strength was maintained at 66% detection threshold with suprathreshold (100%) detected and null stimuli randomly interleaved for 10% and 20% of the trials, respectively. The PEST procedure and the hand stimulation device and protocol were as used by Jones et al. (2007, 2009).

Experimental procedure
Localization runs. To aid localization of primary equivalent current dipoles (ECDs) in contralateral SI, each experiment began with presentation of suprathreshold stimuli to the left hand third digit for 3 min with an interstimulus interval of 3 s (60 trials per subject). Separate localization runs were also performed on the first digit of the foot, as an original goal of our study was to look at the somatotopic precision of attention allocation by comparing hand and foot activity. However, consistent dipoles could not be reconstructed from the foot localization data using the standard ECD localization techniques described below. Thus, only hand area activity is presented, as its precise location in the SI map could be confirmed.

Cued detection runs. Subjects were instructed to fixate on a cross on a projection screen. PEST procedure was used for 3 min at the beginning to determine subjects’ initial detection thresholds. This run was followed by at least 5 cued detection runs, described in Figure 1, consisting of a 3.5 s trial that began with the fixation cross changing into a visual word cue on a projection screen directing the participant to attend to the “Hand” (the attend-in condition), the “Foot” (the attend-out condition), or “Either” location. The visual cue was accompanied by a 60 dB, 2 kHz tone delivered to both ears to mask audible clicks created by the tactile stimulator and remained constant for 2.5 s. At a randomized time between 1.1 and 2.1 s (fixed 100 ms intervals) after the visual cue, the piezoelectric stimulator delivered a brief tactile stimulus to either the finger or toe. At the end of the 2.5 s visual cue, and at least 400 ms after tactile stimulus, subjects reported detection or nondetection of the stimulus at the cued location with button presses using the second and third digits of the right hand, respectively. The next trial began 1 s after cessation of the visual cue. There were 120 trials per run, 40 of each attention condition, totaling at least 200 trials of 3 stimulus strengths in each condition.

MEG data acquisition and source analysis. The MEG signals were recorded using a 306-channel whole-head planar dc-SQUID Neuromag Vectorview system. Data were acquired at 601 Hz and filtered from 0.1 to 200 Hz. Four head position coils recorded head position in the Dewar for coregistration with structural MR images. Vertical and horizontal electro-oculogram (EOG) signals were recorded with electrodes placed close to the left eye. Epochs with EOG peak-to-peak amplitude exceeding 100 μV were excluded from analysis.

The contribution from the left third digit representation in SI to the measured fields was estimated using a least-squares fit with a dipole forward solution calculated using a spherical symmetric conductor model of the head (Hämäläinen and Sarvas, 1989). Averaged data from the localization runs described above were used to find an ECD (Elekta-Neuromag software) at the time of peak activity (mean peak activity = 66.8 ms, SD = 6.4 ms) in the mean signal from the suprathreshold stimuli (minimum n = 50 runs per subject). The goodness of fit of this single dipole model was larger than 70% in all fit data during peak responses. Coregistration of the SI source localization with the individual’s anatomical MRIs confirmed that the source emerged from the anterior band of the postcentral gyrus finger representation of area 3b in SI (Moore et al., 2000) in all subjects (see Fig. 2A). All analysis considered the forward solution from this SI source.

Analysis

Time evolution of spectral power. This metric was calculated using a complex wavelet analysis, from which time–frequency representations (TFRs) of near instantaneous changes were determined. The TFRs were calculated from 1 to 40 Hz on the SI ECD time courses by convolving signals with a complex Morlet wavelet of the form \( w(t, f_0) = A \exp\left(-\frac{t^2}{2\sigma^2}\right) \exp\left(2i\pi f_0 t\right) \), for each frequency of interest \( f_0 \), where \( \sigma = m/2\pi f_0 \), and \( m \) is the imaginary unit. The normalization factor was \( A = 1/(\sigma\sqrt{2\pi}) \), and the constant \( m \) defining the compromise between time and frequency resolution was 7, as in the Jones et al. (2009) study. Time–frequency representations of mu-alpha and mu-beta power were calculated as the squared magnitude of the complex wavelet-transformed data averaged across the range of interest. For the postcue analysis (see Fig. 2B), baseline was calculated as averaged power \([-200, 0]\) ms relative to the cue—averaged across attend-in and attend-out trials separately. For the prestimulus analysis (see Fig. 2C), baseline was calculated as average power \([-500, 0]\) ms relative to the stimulus.

Data were analyzed using the last 100 trials of each condition. This choice reflects preliminary analysis across multiple studies in our laboratory (S. R. Jones and C. I. Moore, unpublished data) indicating that the initial 100 trials provide less stable and consistent neurophysiological activity patterns across individuals.

Visual cue and tactile stimulus broadband evoked responses. SI evoked responses were calculated across an equal number of attend-in and attend-out trials per subject from threshold-level tactile stimulation to the finger (number of trials mean = 88 trials; SD = 10). Averages were baseline normalized by subtracting the mean over \([-100, 0]\) ms from the cue or stimulus, for each subject (see Fig. 3). A response artifact from 1.7 to 2.1 s in one subject was removed from Figure 3A.
Impact of mu-alpha and mu-beta on detection probabilities. We could not use the current data to assess the impact of mu-alpha and mu-beta on detection probabilities because the subset of relevant data was a statistically small sample. The relevant data consisted of the “hit” and “miss” trials in the attend-hand condition, and there were only a small number of miss trials per subject in this subcondition (number of miss trials: mean = 11, SD = 6; number of hit trials: mean = 83, SD = 11). Therefore, we conducted analysis using a second dataset where the statistics were tractable. As in the present study, subjects detected taps applied to the third digit fingertip (of the right hand) using the same stimulator, and signals were localized to the hand dipole using identical means. Because foot trials and “either” trials were not interleaved—subjects attended to the hand throughout—we had a more extensive trial base for comparing hit and miss trials (last 100 trials analyzed). Details of data collection were described in detail in prior reports by our group (Jones et al., 2007, 2009; Ziegler et al., 2010).

Analysis methods were as in the Linkenkaer-Hansen et al. (2004) study. In brief, for each subject mu-alpha and mu-beta power was averaged over a 1 s prestimulus time window for each of the last 100 threshold-level stimulus trials, and binned into 10 power percentile bins (10 averages per bin) sorted from low to high. The probability of detection in each bin was calculated as a percentage change in hit rate (number of detected trials/total number of trials) from the mean (see Fig. 4).

Statistical analyses. Nonparametric Wilcoxon sign-rank tests were used to assess the statistical significance of differences in attend-in and attend-out conditions across subjects at every time point (see Figs. 2B,C,3). The Wilcoxon test was preferred over ANOVA because the data were non-normally distributed (Shapiro-Wilkst test). Linear regression on the mean percentage change in hit rate across subjects was used to assess the impact of mu-alpha and mu-beta power on detection (see Fig. 4).

Results

We investigated the temporal evolution of changes in mu-alpha (7–14 Hz) and mu-beta (15–29 Hz) power in a localized dipole source in the right hand area of SI after a cue to attend to tactile finger stimulation to the contralateral left hand (attend-in condition), or to tactile toe-stimulation to the left foot (attend-out condition). Figure 2A shows example localizations in two subjects in the SI hand representation in the anterior bank of the postcentral gyrus (area 3b), confirmed by proximity to the Ω-shaped bend (marked in red in Fig. 2A, bottom panel) in the central sulcus (Moore et al., 2000). Figure 2B shows the corresponding average percentage change from baseline [#n = 12 subjects (Ss)] in SI mu-alpha and mu-beta power during the time period [−100, 1100] ms relative to the cue in attend-in and attend-out conditions. A significant difference across subjects was observed between the conditions in the mu-alpha band during the anticipatory postcue time period [500, 1100] ms relative to the cue (#p < 0.05 marked with asterisks, Wilcoxon sign-rank test). Significant differences in the mu-beta band were observed for a time window of [800, 850] ms.

Aligning trials to the tactile stimulus onset ([-1000, 200] ms), rather than visual cue, also showed a dominant effect of cued attention on prestimulus mu-alpha activity (Fig. 2C). Significant differences between attend-in and attend-out conditions are seen in the mu-alpha across the entire prestimulus time period, and in the [−200, 0] ms time window for mu-beta, with another period of significance around ≈ 800 ms.

Next, we investigated attentional modulation of the broadband SI evoked response to the visual cue and subsequent threshold-level tactile stimulus (Fig. 3). There was a rapid response in SI to the visual cue with an initial peak near 70 ms (labeled pcM70, for “postcue M70,” for reference in Fig. 3A) that was greater in the attend-in condition. Several other time points showed a significant difference between attend-in and attend-out conditions, and the timings of these differences was consistent with previously reported modulation of EEG measured event-related potentials (ERPs) during attention deployment in parietal, frontal, and visual cortices (Kelly et al., 2009). Most notable are the statistically significant differences at 200 ms, 400 ms, and
several intervals between 500 and 900 ms (see magenta asterisks in Fig. 3A). The 200 ms difference is consistent with that seen in parietal cortices and typically referred to as an “early directing attention negativity” (EDAN), labeled pcMEDAN (for “postcue MEG EDAN”) in Figure 3A. The difference near 400 ms is consistent with the anterior directing attention negativity (ADAN) observed in frontal cortices, and the later noncontinuous differences between 500 and 900 ms are in line with those seen over occipital cortex known as late directing attention positivity (LDAP), labeled pcMADAN and pcMLDAP, respectively.

Figure 3B shows the SI tactile ER at [−100, 200] ms. Peaks in the waveform were consistent with previous reports using similar stimuli (Jones et al., 2007, 2009). Four peaks occurring at −50 ms (M50), 70 ms (M70), 100 ms (M100), and 135 ms (M135), respectively, are labeled for visualization as in the Jones et al. (2007, 2009) studies. A previous study showed that high prestimulus mu-alpha and mu-beta was correlated with an increase in the magnitude of the M50 peak and a subsequent trend toward decreased M70 and later response elements (Jones et al., 2009). Here, we found that there was also a significant difference in the magnitude of the ER near the M50 peak between the attend-in and attend-out conditions, such that the magnitude of the ER was greater in the attend-out conditions, when prestimulus mu-alpha and mu-beta were higher (Fig. 3B, significant time points marked with asterisks, p < 0.05 Wilcoxon sign-rank test). There was also a significant difference in the ER near the M100 peak (p < 0.05 Wilcoxon sign-rank).

Although a slow cue-locked fluctuation is apparent visually in the averaged cue ER in attend-in and attend-out conditions (Fig. 3A), this did not bias the averaged tactile ER (Fig. 3B) since the tactile stimulus was jittered within the [1.1, 2.1] s postcue time window (marked in Fig. 3A). However, it is possible this early ER was impacted by postcue attentional modulation of the mu-alpha and mu-beta activity that is not phase locked to the cue (Fig. 2) (see Discussion).

We assessed trial-by-trial impact of prestimulus mu-alpha and mu-beta in our SI signal on tactile detection probabilities, using a second dataset that used analogous MEG and tactile detection methods, but with sustained attention to the finger (Jones et al., 2007, 2009; Ziegler et al., 2010). This previously collected dataset gave greater statistical power than the current data, where the relevant hit and miss trials represented a statistically small sample (see Materials and Methods). Following prior convention (Linkenkaer-Hansen et al., 2004), on individual trials average prestimulus mu-alpha and mu-beta power was calculated (1 s before stimulus) and sorted from high to low power into 10 equally sized percentile bins. Detection probabilities in each bin were calculated as the percentage change in hit rate from the mean, as a function of mu-alpha power sorted into 10 power percentile bins. Bold traces, Mean and SE across subjects (n = 105); thin traces, linear fit from linear regression analysis ($R^2 = 0.65, p < 0.05$). This analysis was performed on data from the Jones et al. (2007, 2009) studies (see Materials and Methods).

**Discussion**

Consistent with findings in visual cortices, we observed that cued attention modulates anticipatory postcue mu-alpha activity and early peaks in the broadband visual cue- and tactile stimulus-induced ERs in the SI hand representation. We found a significant difference in the postcue change from baseline of mu-alpha power between attend-in and attend-out conditions. Detection probabilities were greater during lower prestimulus mu-alpha and mu-beta power. These results are consistent with the theory that attentionally induced focal alpha changes are an active mechanism for modulation of sensory information processing (Foxe et al., 1998; Worden et al., 2000; Kelly et al., 2006, 2009; Thut et al., 2006; Klimesch et al., 2007). Further, our results show that although attentional cuing has a weaker impact on SI mu-beta rhythms, they also predict tactile detection.

**Attentional modulation of low-frequency rhythms and their impact on perception**

Prior research connecting low-frequency oscillations and cued attention in somatosensory cortex in humans has focused on sensory-stimulus- and movement-induced ERD and subsequent ERS in alpha and beta frequencies (Bauer et al., 2006; Babiloni et al., 2008; Dockstader et al., 2010) (see also Pfurtscheller, 1992). These studies also investigated attentional modulation of somatosensory gamma (35–80 Hz) activity. Gamma was not a focus of our study because it is not robust in our spontaneous SI signal (see Jones et al., 2009).

Our finding of a linear relationship between mu-alpha and mu-beta power and tactile detection agrees with Schubert et al. (2009), who found a linear relationship between alpha and beta amplitudes in EEG electrodes over SI and tactile detection probability. These results also are consistent with the relationship between smaller alpha amplitudes and increased perception and decreased reaction times in the visual system (Thut et al., 2006; Hanslmayr et al., 2007; van Dijk et al., 2008; Kelly et al., 2009). Other studies in the somatosensory (Linkenkaer-Hansen et al.,
2004; Zhang and Ding, 2010) and visual (Rajagovindan and Ding, 2010) systems have reported an inverted-U relationship between alpha power and detection probabilities. We also found that, on trials with the lowest values of mu-alpha power, detection probability diminished compared to slightly higher values (compare 10th and 20th percentiles in Fig. 4), suggesting that a minimal baseline of alpha activity is necessary for optimal signal propagation. These results imply that the reduction of mu-alpha and mu-beta activity in SI with cued attention is an active cortical gating mechanism that increases the perceptual salience of tactile signals.

Attentional modulation of visual cue and tactile stimulus

SI ERs Studies have shown consistent effects regarding the timing and spatial specificity of distinct components of postcue EEG measured ERPs in parietal, frontal, and low-level visual areas (e.g., Kelly et al., 2009). We have shown that attentional modulation of broadband visual cue-induced ERPs components exists at similar times in our high-resolution localized SI signal. In addition to changes near 200 ms (pcMEDAN), 400 ms (pcMADAN), and between 500 and 900 ms activity (pcMLDAP), we found an earlier significant difference at ~70 ms (pcM70), such that the magnitude of the peak was greater in attend-in conditions (Fig. 3A), suggesting rapid attentional modulation in SI.

While fitting data to a single dipole is a necessary simplification in our study, the pcM70 response difference is likely restricted to activity from SI. Rapid responses (<100 ms) to visual stimuli have been observed previously in SI. Zhou and Fuster (1997, 2000) showed a change in firing rate (predominantly increases) in single units in the anterior parietal cortex of trained monkeys, including the hand representation area in SI, during a visual cue indicating a subsequent tactile choice. In this visuo-haptic memory task, a change in firing rate was observed as early as ~50–100 ms after visual cue and lasted throughout a memory retention period (~20 s). Further, some units showed differential activity depending on haptic choice indicated by the cue—vertical or horizontal bars indicating corresponding rod with oriented ridges to pull (Zhou and Fuster, 1997, 2000).

Our findings of increased magnitude near the M50 and M100 SI tactile ER peaks (Fig. 3B) with attention are consistent with the theory that attention increases the gain of sensory-evoked responses, and are in line with previous studies that showed increased magnitudes of evoked SI activity in MEG/EEG sensor data at similar time points (Schubert et al., 2008; Dockstader et al., 2010). However, the decreased M50 peak in the attend-in condition contrasts with previous studies showing increases near 50 ms with attention (Iguchi et al., 2002; Schubert et al., 2008; Dockstader et al., 2010). These discrepancies are likely due to differences in experimental design and the fact that the previous studies used stimuli that were perceptually salient (electrical and Braille stimuli), in contrast to our stimulus amplitude that was maintained at a 50% perceptual threshold. The ~50 ms differences in Iguchi et al. (2002) appeared during a two-finger discrimination task, but not during a nondiscrimination task. Schubert et al. (2008) used an oddball detection paradigm, and Dockstader et al. (2010) engaged attention by counting stimuli. Further, the relative magnitude of evoked activity in SI can vary as a function of context, with identical stimuli causing enhanced or suppressed responses dependent on the amplitude of sensory drive (Moore et al., 1999). In our study, greater prestimulus mu activity may similarly prime the enhancement of this early response component under conditions of weak sensory drive.

Biophysical mechanisms of SI attentional modulation: connection to previous computational modeling predictions

SI mu-alpha rhythms

We have developed a model of SI that predicts the origin of the MEG mu rhythm and tactile ERs (Jones et al., 2007, 2009). The model predicts that the mu-alpha component of the mu rhythm is created by a 10 Hz lemniscal thalamic input to granular and infragranular layers, while the mu-beta component is dependent on a second 10 Hz input to the supragranular layers. Given our model, the observed attentional decrease of mu-alpha is predicted to occur through suppression of ongoing 10 Hz lemniscal thalamus activity. This is consistent with the notion proposed by Francis Crick that the inhibitory reticular thalamus controls an attentional “searchlight” in topographic sensory representations (Crick, 1984) and recent recordings by McAlonan et al. (2008) showing that attention modulates sensory-evoked thalamic responses before reaching the neocortex (see also Suffczynski et al., 2001).

SI ERs Our previous modeling further predicts that the decreased M50 tactile stimulus peak responses with attention may arise, at least in part, from excitatory neurons that are less depolarized during low prestimulus mu and hence have a smaller immediate post-stimulus (M50) response. This decreased excitatory response leads to decreased poststimulus inhibition in the network, which enables greater subsequent M70 and M100 responses (see Jones et al., 2009). Recent research suggests that alpha rhythms may be amplified by a recruitment of a class of low-threshold spiking inhibitory neurons (Fanselow et al., 2008; Vierling-Claassen et al., 2010). These predictions provide a direct neural correlate for the theory that mu-alpha exerts inhibitory control in the neocortex (Worden et al., 2000; Klimesch et al., 2007).

Further, the negative polarity of the postcue SI pcM70 suggests it was driven by excitatory input to the supragranular layers, from higher-order neocortical or nonspecific thalamic sources (Jones et al., 2007). A subsequent peak with negative polarity emerges ~100 ms later, reflective of a two-cycle cue-locked ~10 Hz drive, which is stronger with attention. The attentional differences between 500 and 900 ms (pcMLDAP) appear to be tied to the cue-locked slow fluctuation. Slow oscillations are known to modulate higher-frequency rhythms (Schroeder and Lakatos, 2009); however, an investigation of this is beyond the scope of this study.

In summary, our results support the idea that alpha deployment may be a general mechanism of cued attention for active sensory gating. Mu-beta rhythms are impacted less by attentional cuing, but are well correlated with the probability of tactile detection. This dissociation suggests that while both oscillations impact signal processing, mu-alpha deployment may be more readily recruited during attentional allocation. Future research is needed to identify whether attentional mu-alpha modulation is somatotopically precise on finer spatial scales, as has been explored in the visual system.

References


