Attention Drives Synchronization of Alpha and Beta Rhythms between Right Inferior Frontal and Primary Sensory Neocortex

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The right inferior frontal cortex (rIFC) is specifically associated with attentional control via the inhibition of behaviorally irrelevant stimuli and motor responses. Similarly, recent evidence has shown that alpha (7–14 Hz) and beta (15–29 Hz) oscillations in primary sensory neocortical areas are enhanced in the representation of non-attended stimuli, leading to the hypothesis that allocation of these rhythms plays an active role in optimal inattention. Here, we tested the hypothesis that selective synchronization between rIFC and primary sensory neocortex occurs in these frequency bands during inattention. We used magnetoencephalography to investigate phase synchrony between primary somatosensory (SI) and rIFC regions during a cued-attention tactile detection task that required suppression of response to uncertain distractor stimuli. Attentional modulation of synchrony between SI and rIFC was found in both the alpha and beta frequency bands. This synchrony manifested as an increase in the alpha-band early after cue between non-attended SI representations and rIFC, and as a subsequent increase in beta-band synchrony closer to stimulus processing. Differences in phase synchrony were not found in several proximal control regions. These results are the first to reveal distinct interactions between primary sensory cortex and rIFC in humans and suggest that synchrony between rIFC and primary sensory representations plays a role in the inhibition of irrelevant sensory stimuli and motor responses.

Key words: attention; inferior frontal cortex; magnetoencephalography; primary somatosensory cortex; somatosensation; synchrony

Introduction

Synchronization of oscillatory activity between cortical regions is hypothesized to regulate information flow and to be integrally involved in the attentional modulation of intracortical communication (Fries, 2005, 2009; Jensen and Mazaheri, 2010; Schroeder and Lakatos, 2009; Kveraga et al., 2011; Siegel et al., 2012; Buschman et al., 2012; Womelsdorf et al., 2014). The right inferior frontal cortex (rIFC) is specifically associated with attentional control via the inhibition of behaviorally irrelevant stimuli and motor responses. Similarly, recent evidence has shown that alpha (7–14 Hz) and beta (15–29 Hz) oscillations in primary sensory neocortical areas are enhanced in the representation of non-attended stimuli, leading to the hypothesis that allocation of these rhythms plays an active role in optimal inattention. Here, we tested the hypothesis that selective synchronization between rIFC and primary sensory neocortex occurs in these frequency bands during inattention. We used magnetoencephalography to investigate phase synchrony between primary somatosensory (SI) and rIFC regions during a cued-attention tactile detection task that required suppression of response to uncertain distractor stimuli. Attentional modulation of synchrony between SI and rIFC was found in both the alpha and beta frequency bands. This synchrony manifested as an increase in the alpha-band early after cue between non-attended SI representations and rIFC, and as a subsequent increase in beta-band synchrony closer to stimulus processing. Differences in phase synchrony were not found in several proximal control regions. These results are the first to reveal distinct interactions between primary sensory cortex and rIFC in humans and suggest that synchrony between rIFC and primary sensory representations plays a role in the inhibition of irrelevant sensory stimuli and motor responses.

Key words: attention; inferior frontal cortex; magnetoencephalography; primary somatosensory cortex; somatosensation; synchrony
Evidence also suggests that rFIC subregions are differentially engaged in attentional monitoring and motor response inhibition. Anterior dorsal rFIC subregions, including right inferior frontal gyrus (rIFG) and inferior frontal sulcus (rIFS), are involved in attentional monitoring and reflexive reorienting, whereas more posterior ventral subregions, including the right inferior frontal junction (rIFF), are additionally involved in stopping action (Sharp et al., 2010; Levy and Wagner, 2011; Baldauf and Desimone, 2014).

Considerable evidence suggests that primary sensory representations gate the flow of relevant sensory information to cortex and that attentional modulation of low-frequency (alpha and beta) power underlies this gating (Worden et al., 2000; Siegel et al., 2008; e.g., Jones et al., 2010). To date, there is no evidence of attentional modulation of synchrony between prefrontal cortex and topographically localized primary sensory representations in humans. This absence may be due to phase synchrony not acting as a mechanism for attentional control, or it may be that phase synchrony is limited to more proximal regions in the neocortical hierarchy.

Here, we used magnetoencephalography (MEG) to test the hypothesis that a cued-attention tactile detection task, which explicitly required suppression of distractor stimuli in non-attended representations, would selectively engage low-frequency synchrony between a topographically localized primary somatosensory cortex (SI) representation and rFIC. We additionally hypothesized that anatomically identified rFIC subregions (i.e., rIFS and rIFG) may differentially synchronize with SI. This study extends prior work in which we showed that attention modulates alpha and beta components of the somatosensory mu-rhythm during the same cued-attention tactile detection task (Jones et al., 2010).

Materials and Methods

The data analyzed in this study were used in prior investigations of attentional modulation of alpha and beta power in SI. Subject recruitment, experimental protocol, and data acquisition have been explained in detail previously (Jones et al., 2010; Kerr et al., 2011; Wan et al., 2011) and are reviewed below. The novel source estimation and data analysis performed in this study are detailed.

Subjects and experimental design. Data were collected from 12 healthy, right-handed adults (mean ± SD: age, 31.6 ± 7 years; 11 female). During MEG data acquisition, subjects’ hands and feet rested on plastic frames through which tactile stimuli (10 ms duration) were delivered to the left hand (third digit) and left foot (first digit) using a piezoelectric device (Noliac). We performed localization trials to functionally identify subject-specific SI hand regions (hSI), in addition to cued-attention trials in which subjects completed a cued-attention tactile discrimination task. During localization trials, suprathereshold tactile stimuli were administered to the third digit of the left hand. These data were used to localize the left hand representation in SI. Localization runs were 3 min in duration, with 3 s trials. Each subject completed at least three localization runs.

During cued-attention trials, a visual cue instructed subjects to allocate attention in anticipation of a brief, threshold-level tactile stimulus to the “hand,” “foot,” or “either” (i.e., hand or foot). Tactile stimulation was counterbalanced by area and jittered, occurring at least 1000 ms after the cue on the distal pad of the left hand or foot. On an individual subject basis, initial stimulus thresholds were identified using a Parameter Estimation Sequential Testing convergence algorithm (Leek, 2001). Three minutes of data was used for this procedure. The Parameter Estimation Sequential Testing procedure consists of series of three-trial sets, including a strong, moderate, and minimum stimulus that change strengths dynamically depending on subject detection. The first three-trial set is comprised of a strong stimulus (100% amplitude, 350 μm), then a weak stimulus (50% amplitude), and last a blank stimulus. When the subject reports detection of both the strong and weak stimulus, the strong stimulus magnitude is halved, the moderate stimulus magnitude is replaced with the mean of moderate and weak stimuli magnitudes, and the weak stimulus magnitude stays the same. When the subject fails to detect the moderate stimulus, the strong stimulus does not change, and the minimum stimulus is replaced by the mean of the strong and the moderate stimulus. This three-trial set procedure was iterated until the difference in the middle stimulus between the last two trials was <5 μm.

During MEG data collection in cued-attention runs, intensity was maintained at a 66% detection level across all attend condition (hand, foot or either) using a dynamic algorithm. If two correct responses were made, the threshold-level voltage sent to the piezoelectric was decreased by 0.005 V (a change of ~4.5 μm in piezoelectric movement), and the correct response counts were reset to zero. In contrast, if three incorrect responses were made, the voltage was increased by 0.005 V. Additionally, suprathereshold (%) and null-threshold (0% detection) stimuli were randomly interleaved for a 10% and 20% of trials, respectively.

At least 400 ms after the stimulus, when the visual cue changed to a fixation cross, subjects reported via right-hand button press (see Fig. 1A) detection or non-detection of tactile stimulation in the attended location. Here we report on data from the pretactile stimulus period, from (~200 to 1000 ms relative to the visual cue). Subjects completed at least five cued-attention runs with at least 40 trials per run (mean ± SD: total trials, 562.8 ± 69.0).

We were unable to compare phase-locking values (PLVs) across detected and not detected trials as subjects only reported detection in attended trials, and the majority of attended hand trials were detected. The total number of trials in which attention was cued to the hand and subjects reported stimulus detection was 82.5 ± 11.3 (mean ± SD), whereas the number of attended and non-detected trials was 9.9 ± 5.8 (mean ± SD); thus, this analysis was underpowered. Additionally, our study was not designed to assess relations between PLV and reaction time, as subjects were cued to respond to detection, limiting reaction time variability.

MEG and MRI data acquisition. Data were recorded using a 306 channel MEG with 102 magnetometers and 204 planar gradiometers located at 102 locations around the scalp (dc-SQUID VectorView system, Elekta-Neuromag) inside a magnetically shielded room (IMEDICO). Data were collected at a 600 Hz sampling rate and filtered to a 0.1–200 Hz bandpass. Both horizontal and vertical bipolar EEG were collected for eye movement-related artifact rejection. High-resolution T1-weighted anatomical MRIs (3.0T Trio, Siemens) were collected for coregistration using standard procedures. The FreeSurfer software suite (http://surfer.nmr.mgh.harvard.edu) was used for surface reconstruction and cortical labeling (Fischl et al., 2002; Dale et al., 1999; Mosher et al., 1999; Liu et al., 2002; Fischl et al., 2004; Yovinec et al., 2006). MEG-coordinate frames were manually aligned to anatomical MRI data using fiducial points and subsequently automatically.

MEG and source signal estimation. In contrast to our prior studies, which used an equivalent current dipole fitting procedure (Jones et al., 2010; Kerr et al., 2011; Wan et al., 2011), in this study minimum norm estimates (MNEs) were calculated on a subject-by-subject basis using the MNE software suite (Gramfort et al., 2014). MNE calculates distributed dipoles by minimizing the norm of estimated currents within the brain, allowing for the projection of signals from sensor space to the cortical surface (i.e., source space). The MNE software suite facilitated MEG-MRI coregistration, construction of cortically constrained source, conductivity, and forward and inverse models (Hämäläinen and Sarvas, 1989; Dale and Sereno, 1993; Hämäläinen and Ilmoniemi, 1994; Dale et al., 1999; Mosher et al., 1999; Liu et al., 2002; Fischl et al., 2004; Yovinec et al., 2006). MEG-coordinate frames were aligned using standard procedures. The FreeSurfer software suite (http://surfer.nmr.mgh.harvard.edu) was used for surface reconstruction and cortical labeling (Fischl et al., 2002; Dale et al., 1999; Mosher et al., 1999; Liu et al., 2002; Fischl et al., 2004; Yovinec et al., 2006). MEG-coordinate frames were manually aligned to anatomical MRI data using fiducial points and subsequently automatically.
calibrated using an Iterative Closest Point algorithm (15 iterations). Forward solutions were estimated using single-layer homogeneous boundary-element models (the inner skull surface was segmented to 5120 triangles). Inverse operators were then estimated. For synchrony analysis, inverse operators were calculated using empty room noise covariance matrices in conjunction with depth weighting and loose orientation constraints (Lin et al., 2006). For functional localization, a similar procedure was used to estimate MNEs, with several key differences: a 200 ms pretactile stimulus period was used for the noise covariance matrix and activity was estimated using a normalized MNE (dynamic statistical parametric mapping [dSPM]). An empty room noise covariance matrix was used for the noise covariance matrix and functional localization, a similar procedure was then used for synchrony analysis to reduce attenuation of preattentional cue rhythmic activity (i.e., spontaneous rhythmic activities). In contrast, the functional localizer task is explicitly postattentional cue, and thus attenuation of preattentional cue activity was important to specifically reveal stimulus-evoked activities.

**Defining ROIs.** For the localization of hSI, peak tactile-evoked responses were extracted from the Desikan–Killiany postcentral cortical label of localization trials. To identify an approximate window in which the peak evoked response occurred, these trial-averaged current distributions were manually inspected from a 40–70 ms post-tactile stimulus window, which was previously shown to contain a prominent peak response to this tactile stimulus (Jones et al., 2007). Responses were visualized on rendered inflated cortical surfaces (manually identified approximate time interval start/end, mean ± SD: 54.6/71.3 ± 4.8/5.1 ms), and dSPM estimates were extracted from each vertex of the label, for each time point in the manually identified time window. This resulted in millisecond-by-millisecond poststimulus dSPM estimates from the postcentral label in which the hand-related evoked response occurred. The precise time point of maximum dSPM value across all postcentral vertices was then algorithmically identified. The top 7.5% most active dipoles at this precisely identified time point were defined as hSI. If a vertex in the ROI was ≥5 vertices from the largest contiguous mass of vertices, it was replaced by the next most active vertex (for an example hSI, see Figure 1B).

As in our prior studies (Jones et al., 2010; Kerr et al., 2011), we had difficulty localizing the foot area of SI (fSI) using a similar procedure. Difficulty in localizing fSI using MEG is thought to be the result of low signal amplitudes from the foot (and lower limbs more generally) as detected by MEG (Nakamura et al., 1998). Here, in an attempt to analyze fSI, we used average peak response to suprathreshold stimulation to the left first digit toe and defined fSI ROIs as ≥14 vertices clustered near the midline in the Desikan paracentral lobule cortical label. We were only able to identify fSI in 8 of the 12 participants. Analysis of alpha and beta PLV between fSI and rIFC in these 8 subjects revealed no statistical significance. We view these results as underpowered and inconclusive.

Anatomically defined cortical labels based on the Destrieux cortical atlas (Destrieux et al., 2010) were used to define rIFC subregions. These labels were not selected based on source-localized activity patterns but rather selected based on predefined atlas parcellations with regions chosen a priori. As MEG predominately detects signals tangential to the cortical surface and MNE preferentially attributes signals to the cortical surface, ROIs were restricted to labels anatomically defined within cortical sulci; however, this does not completely rule out possible contribution from gyri (Hillebrand and Barnes, 2002). Specifically, based on prior literature and a priori hypotheses, we examined synchrony between hSI and two subregions of rIFC (i.e., right IFS and IFJ, rIFS and rIFJ, respectively) (Sharp et al., 2010; Levy and Wagner, 2011; Baldauf and Destirome, 2014). These regions were anatomically identified using Destrieux parcellation labels referred to as IFS, and as the anterior vertical ramus of the Sylvian fissure at the intersection of the posterior end of the IFS and the precentral sulcus, also known as IFJ (Baldauf and Desimone, 2014; Levy and Wagner, 2011) (see Fig. 1B). Because of individual differences in cortical morphometry, in two subjects the rIFJ could not be well defined. These subjects were omitted from analyses related to the rIFJ.

**Signal extraction.** For a given ROI, vertex-wise signals were averaged to compute synchrony and spectral power analyses. To address the possibility of signals partially cancelling each other due to variation in dipole orientations, before signal averaging, signals from different vertices were aligned within each subject’s ROI during the computation of inverse solutions from the raw data. Specifically, first the surface normals within the label were organized in an $n \times 3$ matrix. The preferred orientation was then calculated from the $n \times 3$ matrix by taking the first right singular vector, which is its largest singular value. If, after taking the dot product of the surface normal of a vertex, the result was negative, then the signal estimates at this vertex were inverted. This procedure was
implemented via the align_z option of the mne_compute_raw_inverse function from the MNE software suite. Thus, align_z ensures that signals do not cancel due to the interaction of geometry and the arbitrary assignment of positive and negative activities in source space. We acknowledge that such alignment could bias toward homogenous PLVs (as defined below), even if their sign is heterogeneous, and hence overestimate the PLV. Given the complementary nature of our results with prior literature on the role of IFC, we do not think this has significantly confounded our results.

Phase synchrony analysis. Synchrony was estimated on a trial-by-trial manner by calculating PLVs between signals from hSI and rIFC subregions. PLV quantifies synchrony by assessing variance in phase relations between signals at a given frequency (Lachaux et al., 1999). After aligning and averaging vertex-wise signals for each ROI, PLVs were calculated millisecond by millisecond from −733 to 1333 ms relative to visual attention cue onset, from 1 to 50 Hz (1 Hz step). The beginning and ending periods were discarded to prevent signal smearing effects at the edge of the wavelet; only values from −200 to 1000 ms relative to the cue were considered for further analysis (see Fig. 2). Both signals (s_i and s_j) were first convolved with a Morlet wavelet as follows:

\[ w(t, f_0) = A \exp(-r^2/2\sigma_i^2) \exp(2i\pi f_0 t) \]

with normalization factor \( A \) equal to:

\[ (\sigma_i \sqrt{\pi})^{-1/2} \]

This results in a complex representation of phase for trial \( i \), time \( t \) and frequency \( f_0 \) for each signal (here \( s_i \)):

\[ \Phi_i(t, f_0) = \frac{w(t, f_0) \ast s_i(t)}{|w(t, f_0) \ast s_i(t)|} \]

The width of the wavelet (\( m = f_0/\sigma_i \)) was selected as 7; with \( \sigma_i = 1/2\pi f_0 \). Finally, PLVs were calculated across \( N \) trials between signals \( s_i \) and \( s_j \):

\[ PLV (t, f_0) = \frac{1}{N} \sum_{i=1}^{N} (\Phi_i \ast \Phi_j) \]

The modulus of which is PLV. PLV is thus a measure of intertrial variability of phase between two signals at time \( t \), ranging from 0 to 1 (1 indicating perfect phase synchrony).

Statistical analysis of PLVs. Consistent with previous work indicating dominance of 7–14 Hz alpha and 15–29 Hz beta frequency bands in human SI MEG measured signals (Tiihonen et al., 1989; e.g., Jones et al., 2009, 2010), PLVs between SI and the frontal regions considered were dominant in these bands (see Fig. 2). Therefore, for all analyzes, PLV was averaged across the 7–14 Hz alpha band and 15–29 Hz beta band to test \( a \ priori \) hypotheses that synchrony would occur in these bands.

Statistically significant differences in PLVs between the attend-out (foot) versus attend-in (hand) condition were assessed using a binned analysis. Data were binned and averaged in 200 ms intervals in a 0–1000 ms postcue period, as in our prior study (Kerr et al., 2011). These data were analyzed as percentage change from baseline, with baseline defined as −200 to 0 ms prevical cue across both conditions. As components of the data were non-normally distributed (Lilliefors test, \( p < 0.01 \), non-parametric statistics (paired Wilcoxon sign-rank test) were used. Bonferroni correction was implemented for correction of multiple comparisons (five bins: 0.05/5 = 0.01). Time-binned data meeting criteria at \( p < 0.01 \) were considered significant, and \( p < 0.05 \) was considered a statistical trend.

Analysis of power. To understand whether changes in power might drive synchrony-related effects, we additionally investigated attentional power modulation. As with PLV, we calculated power by convolving signal time courses with a complex Morlet wavelet with width 7. Spectral power time-frequency representations were computed from the squared magnitude of the complex wavelet-transformed data, as in our prior work (Jones et al., 2010). Power was statistically analyzed in the same manner as PLV using five 200 ms bins postattentional cue, with Bonferroni multiple comparison correction across time bins.

**Assessment of laterality.** As a control to assess the laterality of observed PLV effects, we additionally analyzed PLV from left lateralized IFC subregions, left IFS (lIFS) and left (lIFJ). As with the right hemisphere, the left-lateralized ROIs were identified using anatomical parcellation according to the Destrieux atlas. PLV and power were analyzed using the same binned procedure as the right hemisphere.

**Volume conduction analysis.** We tested the possible contribution of volume conduction to our results in three ways. First, in addition to PLV, we examined the imaginary component of PLV (iPLV), a metric robust to volume conduction artifacts that additionally accounts for deep sources that are difficult to measure directly (Nolte et al., 2004). Second, we calculated source-space signal crosstalk between hSI and all other vertices on the cortical surface grid using a simulated signal. Crosstalk is defined as the contribution from multiple source signals to the signal localized at an individual source. The simulated signal (using a normal orientation to the surface) is generated from the lead field matrix. High crosstalk in rIFC would suggest volume-conduction-related confounding of PLV effects. Crosstalk was assessed for each individual in native space, with subsequent warping to the FreeSurfer group template. A group aggregate map was produced by averaging crosstalk across individuals in template space. Last, we assessed PLV in three control regions proximal to rIFC. These ROIs were defined using the Destrieux cortical parcellation and included the right middle frontal sulcus (rMFS), right inferior premotor cortex (rIPMC), and right lateral orbital sulcus (rLOS; see Fig. 1B).

**Results**

Attention induces increases in alpha and beta synchrony between rIFC and non-attended SI representations. We examined phase synchrony in the alpha and beta bands between hSI and rIFC subregions during a task in which subjects were cued to report detection of a threshold level tactile stimulation to either the hand or foot (Fig. 1A). Subjects were instructed to respond only to detection of stimulation in the attended region, despite the equal possibility of stimulation in the nonattended area. Thus, the task required suppression of stimulus processing and motor response to stimulation in the nonattended area.

Based on \( a \ priori \) hypotheses, we examined two parcellated subregions of rIFC defined anatomically using structural MRI.
One of these regions constituted the anterior dorsal portion of rIFC, referred to as rIFS, and the other region constituted the posterior ventral portion of rIFC, referred to as rIFJ (Fig. 1B). These regions are known to have distinct cytoarchitectonic profiles (Amunts and Cramon, 2006) and to exhibit functional and anatomical specificity from neighboring regions (Chikazoe et al., 2009; Sharp et al., 2010; Levy and Wagner, 2011; Derrfuss et al., 2012). PLVs were compared between hSI and each region across attentional conditions (attend-out = attention to foot; or attend-in = attention to hand) in the postcue period before tactile stimulation. This period is marked in Figure 1A (dashed line).

Figure 2 depicts average time-frequency representations of PLVs between hSI and rIFC subregions for attend-in (left panel) and attend-out (right panel) conditions across subjects, from 5 to 45 Hz and −200 to 1000 ms around the cue during. The highest PLV values were restricted to the alpha and beta bands, with greater values in the attend-out condition. All further analyses were thus restricted to these bands.

Statistical significance of differences in PLVs across conditions (attend-out and attend-in) was assessed between hSI and each rIFC subregion in 200 ms bins (Table 1). This analysis revealed significant differences in alpha band PLVs between attend-out and attend-in conditions in the early 200–400 ms postcue period between hSI and rIFJ (p < 0.001), with a subsequent trend in the 400–600 ms period (p < 0.05; Fig. 3A). Specifically, greater alpha synchrony occurred during the attend-out compared with the attend-in condition. Trend differences in alpha synchrony were also observed between hSI and rIFJ 0–200 ms (p < 0.05).

In the beta band, significantly greater synchrony was observed in the attend-out compared with the attend-in condition between hSI and rIFJ in the later 600–800 ms bin (p < 0.01), with a continued trend in the 800–1000 ms bin (p < 0.02; Fig. 3B). Statistical trends were also observed in the beta band between hSI and rIFS in both of these bins (600–1000 ms, p < 0.05).

### Table 1. Statistical comparison of PLV across attentional conditions

<table>
<thead>
<tr>
<th>Time Bin (ms)</th>
<th>Alpha PLV</th>
<th>Beta PLV</th>
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<tbody>
<tr>
<td></td>
<td>rIFS</td>
<td>rIFJ</td>
</tr>
<tr>
<td>0–200</td>
<td>0.038**</td>
<td>0.196</td>
</tr>
<tr>
<td>200–400</td>
<td>0.383</td>
<td>0.047**</td>
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<td>400–600</td>
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<td>800–1000</td>
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<td>0.204</td>
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<tr>
<td></td>
<td>0.016**</td>
<td>0.204</td>
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</tbody>
</table>

*Statistical comparison of PLVs across attentional conditions. Attend-out — attend-in alpha and beta PLVs between signals from functionally defined hSI and rIFC, including rIFS and rIFJ. PLVs were averaged in 200 ms time windows after cue and compared across attentional conditions using two-tailed sign-rank tests.

**Significant difference (p < 0.05), using Bonferroni correction for multiple comparisons across five time bins.

***Statistical trend (p < 0.05).

### Synchrony and attentional modulation of spectral power

To assess whether the attentional PLV modulation might be driven by changes in spectral power, we additionally investigated attentional power modulation in all three regions (Table 2). Differences in power with attention were observed only in hSI in the beta band 600–800 ms after cue (p < 0.01): a time window that overlapped with the PLV differences. As power differences were not found in any of the IFC regions, we conclude that power effects are not the primary driver of the PLV results. We note that the current results differ from our prior work (Jones et al., 2010). Specifically, here we did not find attentional effects in hSI alpha power, and the hSI beta power results vary slightly in the window of significance. This is likely due to differences in the inverse solution techniques used. In the current work, we use MNE, whereas our prior work used single dipole modeling. As such, the ROIs in the current study represent a smaller cortical volume than in our prior work.

### Laterality of synchrony effects

To test whether synchronization was lateralized to rIFC, we also examined synchrony between hSI and IIFC subregions (lIFS and lIFJ). Significant differences across attentional conditions were observed in alpha band PLVs between hSI and lIFS, in the 200–400 ms postcue period (p < 0.01). No other significant differences or trends were found. These effects appeared across attentional conditions despite that the mean raw PLVs were an order of magnitude larger in rIFS than hIFS (mean ± SD: baseline rIFS 0.18 ± 0.0074; baseline hIFS 0.075 ± 0.0027). No differences in power across attentional conditions were found in IIFS.

### Influence of volume conduction

Based on three different analyses, we concluded that the observed synchrony effects between hSI and rIFC were not influenced by volume conduction. First, we examined the imaginary component of PLV, a metric robust to volume conduction artifacts that additionally accounts for deep sources that may be difficult to
alpha synchrony was found early after the cue (200–600 ms) between hSI and rIFS. In contrast, modulation of beta synchrony was found closer to stimulus processing and motor response (600–1000 ms after cue) between hSI and rIFJ, with a trend in rIFS. As discussed below, we conjecture that the observed increase in interareal alpha synchrony with non-attended representations early after the cue may reflect anticipatory "inhibition" of irrelevant sensory information, whereas the subsequent beta synchrony may relate to mechanisms involved in facilitating accurate somatosensory detection and motor response. The unexpected finding of alpha synchrony between hSI and rIFC early after the cue (200–400 ms) may be related to retrieval of the appropriate stimulus–response rule associated with the attentional cue. These results are the first to show differential engagement of alpha and beta synchrony between primary sensory cortex and IFC in humans.

The role of rIFC in inhibition of processing

The rIFC has been identified as having an important role in mediating top-down attention via inhibition of non-relevant sensory information and motor actions, and has been explicitly implicated as part of the stop-signaling pathway in motor control (Hamshire et al., 2010; Neubert et al., 2010; Levy and Wagner, 2011; Swann et al., 2012; Weissman and Prado, 2012; Hwang et al., 2014). However, the precise role and temporal precision of the rIFC in mediating inhibitory control of perception and/or action are unclear. Our results indicate that low-frequency synchronization with primary sensory cortex may be involved in this processing.

A recent review suggests that rIFC subregions may be differentially engaged in attentional processes and motor control, with more anterior regions encoding higher-order goals (e.g., reflexive reorienting) whereas more posterior regions additionally encode more specific subgoals (e.g., action plans) (Levy and Wagner, 2011). Similarly, anterior-to-posterior medial frontal segregations have been shown to differentiate attentional capture and action inhibition (Sharp et al., 2010). Our results can be interpreted within this context, as we found that relatively anterior rIFC (i.e., rIFS) showed early effects in the alpha band specific to the attentional cues (greater alpha synchrony with non-attended representations in SI), which may act to inhibit distraction in irrelevant areas. In contrast, the relatively posterior ventral region (i.e., rIFJ) was differentially engaged in the beta band closer to the time of stimulus processing (greater beta synchrony with nonattended SI representation). This may be related to maintaining the inhibition of irrelevant representations and to inhibiting motor responses to stimulation in the irrelevant area. A similar statistical trend in beta was observed between hSI and rIFS.

Studies using fMRI suggest that rIFG is recruited when important cues are detected, regardless of whether the detection is followed by inhibition of a motor response (Hamshire et al., 2010). However, causal manipulation of presumed rIFG with transcranial magnetic stimulation has directly impacted "stop signal"
motor responses (Chambers et al., 2007; Neubert et al., 2010), suggesting at least part of rIFG is involved in motor control. Our findings may help to disambiguate the role of rIFG in cue detection versus motor inhibition, such that cue detection processes involve alpha synchronization whereas motor inhibition involves beta synchronization (Zanto et al., 2011; Gogulski et al., 2015).

Alpha synchrony: rIFG and the attentional modulation to “inhibit” distraction
Attentional modulation of alpha activity acts in a topographically specific manner to gate the flow of sensory information in cortex such that increased alpha power correlates with inhibition of sensory information across modalities (Worden et al., 2000; Klimesch et al., 2007; Jones et al., 2010; Klimesch, 2012; Kerr et al., 2013). We conjecture that, similar to commonalities in modulation of alpha power, the observed increase in primary sensory to right frontal alpha synchrony in non-attend conditions is a generalizable feature of cortical dynamics that facilitates perception during distraction.

Other studies have investigated attentional modulation of interareal alpha synchrony with parietal association areas. Consensus has not been reached regarding the direction of modulation, likely due to differences in the spatial localization and imaging modalities studied (Sauseng et al., 2005; Siegel et al., 2008). Similar to our investigation, Doesburg et al. (2009) studied alpha synchrony between source-localized low-level sensory areas, in this case visual cortex (inferior occipital gyrus), and rIFC, using EEG. They found no effect on synchronization between visual cortex and rIFC, and report an increase in alpha synchrony with attention between inferior occipital gyrus and superior parietal lobe. This contrasts with our findings of synchrony between low-level somato-sensory representations and rIFC. There are a number of possible explanations for this discrepancy, including the increased spatial specificity in the current MEG study, MEG biases to sulcal rather than gyral activity (Hillebrand and Barnes, 2002), and/or differences in the probability of distracting stimuli in each paradigm, as distraction occurred more often in our paradigm (Doesburg et al., 2009, 50% compared with 20%). This difference suggests that the blocking of distraction may be a key factor in engaging rIFC alpha synchrony with primary sensory cortex. Concordant with our results, an intracranial LFP study in behaving ferrets reports greater low frequency (10–20 Hz) synchrony between primary auditory cortex and prefrontal cortical sites during passive presentations of tone stimuli, compared with behaviorally relevant task conditions in a tone detection paradigm (Fritz et al., 2010).

Alpha synchrony: IIFs and the selection of stimulus–response contingencies
The majority of studies of IFC suggest that attention–driven inhibitory control is restricted to the right hemisphere (Aron et al., 2004, 2014; Levy and Wagner, 2011), although some EEG evidence shows bilateral engagement of IFG in top–down attentional control (Green and McDonald, 2008). The IIFC has more often been associated with the cognitive control of memory, facilitating the retrieval of rules and selection of the proper response given contingencies associated with the rule (Badre and Wagner, 2007; Bunge, 2004; Donohue et al., 2005; Sakai and Passingham, 2003). The middle ventrolateral and dorsolateral regions of the left PFC in particular have been related to rule based response selection (Bunge, 2004) and postretrieval resolution of competition among active representations (Badre and Wagner, 2007). In this context, our observation of early and brief synchrony between hSI and IIFs alpha synchrony (200–400 ms after cue) may be related to the rapid selection of the appropriate stimulus–response rule associated with the contingency to attend-in versus attend-out, rather than a longer lasting top–down inhibitory signal that is mediated by the rIFC.

Beta synchrony: attentional modulation to mediate “stop signaling”
Similar to alpha, topographically precise attentional modulation of beta power in sensory cortex has been observed, with higher power inversely correlated with perception (Bauer et al., 2006; Dockstader et al., 2010; Jones et al., 2010; van Ede et al., 2012). Beta activity is also explicitly tied to motor function, as beta rhythms have been observed in many nodes of the motor pathway in contraction or hold conditions (Leventhal et al., 2012; Miller et al., 2012). This beta activity has been hypothesized to maintain the continuation of currently active rules or motor plans (Engel and Fries, 2010) or to reflect an increased “offline” processing mode during motor decision making (Haegens et al., 2011). The latter conceptualization is consistent with our findings of increased beta synchronization when inhibition of a non-relevant area was necessary for both the detection decision and motor response.

The rIFC has been explicitly implicated in “stop signaling,” probably mediated by a top-down signal to subthalamic nucleus and basal ganglia structures, and/or premotor cortex, which all exhibit beta rhythmity during these conditions (Kühn et al., 2004; Swann et al., 2011; Wessel et al., 2013). rIFC beta synchrony with primary motor and premotor cortex has been explicitly associated with successful execution of stop compared with unsuccessful stop trials (Swann et al., 2009, 2011, 2012). The current finding of increases in beta synchrony between hSI and rIFC during non-attention appears to act as an inhibitory signal to take the non-attended representation offline from the detection process. This beta synchrony may also be related to inhibiting the motor response program, as subjects had to “stop” button press responses to felt stimuli in the non-attended representation. We did not find a significant difference in beta synchrony with attention between hSI and the anatomically defined rIPMC control region, suggesting that the observed synchrony effects between hSI and rIFC may be mediated by subcortical structures rather than premotor cortex (Swann et al., 2009).

Computational neural modeling has shown that beta frequency oscillations can emerge from the integration of two alpha frequency excitatory drives to granular and supragranular layers of neocortex (Jones et al., 2009; Ziegler et al., 2010). These results suggest that beta frequency input to SI is not required to observe beta synchrony between hSI and IFC, but rather a third source may drive both areas (Schmiedt et al., 2014). Further studies are necessary to explicitly delineate the mechanisms of coordination.

References
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