Oscillatory Activity in Human Parietal and Occipital Cortex Shows Hemispheric Lateralization and Memory Effects in a Delayed Double-Step Saccade Task

We applied magnetoencephalography (MEG) to record oscillatory brain activity from human subjects engaged in planning a double-step saccade. In the experiments, subjects \( n = 8 \) remembered the locations of 2 sequentially flashed targets (each followed by a 2-s delay), presented in either the left or right visual hemifield, and then made saccades to the 2 locations in sequence. We examined changes in spectral power in relation to target location (left or right) and memory load (one or two targets), excluding error trials based on concurrent eye tracking. During the delay period following the first target, power in the alpha (8–12 Hz) and beta (13–25 Hz) bands was significantly suppressed in the hemisphere contralateral to the target. When the second target was presented, there was a further suppression in the alpha- and beta-band power over both hemispheres. In this period, the same sensors also showed contralateral power enhancements in the gamma band (60–90 Hz), most significantly prior to the initiation of the saccades. Adaptive spatial filtering techniques localized the neural sources of the directionally selective power changes in parieto-occipital areas. These results provide further support for a topographic organization for delayed saccades in human parietal and occipital cortex.

Keywords: human, MEG, oscillations, parietal cortex, saccade, sensorimotor, spatial memory

Introduction

The delayed-saccade task is an important tool for studying working memory processes in the brain. In this task, a subject fixes a central target while a light is flashed onto the retinal periphery; then, after a short time interval, he or she is cued to look at the remembered location of the flash. Several cortical and subcortical regions, in both human and nonhuman primates, have been shown to retain increased activity during the memory interval until the saccade is made (see Pierrot-Deseilligny et al. 2004 for review).

In the monkey, one of these regions is the lateral intraparietal area (LIP) (Barash et al. 1991; Colby et al. 1996; Mazzoni et al. 1996) located in the posterior parietal cortex, and recognized as an important interface between higher-order visual areas and the oculomotor system (Andersen and Buneo 2002). Increased firing rate in this region has not only been associated with the planning of saccades but also with sensory attention, decision making, and spatial updating. Some studies have reported that neurons in LIP are arranged in a topographical order, preferentially representing target locations from the contralateral visual field (Blatt et al. 1990; Ben Hamed et al. 2001, but see Platt and Glimcher 1998).

Recently, functional magnetic resonance imaging (fMRI) studies have identified a putative human equivalent of monkey area LIP by describing a bilateral region within the posterior superior parietal lobe that topographically represents remembered target locations for delayed saccades (Sereno et al. 2001; Medendorp et al. 2003, 2005; Koyama et al. 2004; Schluppeck et al. 2005). It was further shown that the directionally selective activity in human LIP is modulated with memory load in a double-saccade task, which implicates the region in the active maintenance of multiple target locations (Medendorp et al. 2006). Notably, most of these fMRI studies also reported other regions showing directional selectivity for delayed saccades (Schluppeck et al. 2005). One region of particular interest was identified in anterior-occipital cortex (Sereno et al. 2001; Medendorp et al. 2003, 2005) possibly corresponding to monkey extrastriate area V3A, which has been shown to carry memory and saccade-related signals as well (Nakamura and Colby 2000).

Yet, sustained blood oxygen level-dependent (BOLD) signals or increased firing rates are not the only signatures of ongoing neural activity in these regions. Neural processing is also characterized by both event-related and oscillatory signals, which can be recorded as electroencephalography (EEG) or MEG from the scalp. Given that the oscillatory signals are produced by large ensembles of neurons oscillating in synchrony, they are bound to play an important role in neuronal processing (see Hari and Salmelin 1997; Engel et al. 2001, for reviews). In both human and nonhuman primates various reports have linked the internal rhythms generated within distinct regions of the brain to memory maintenance in various types of working memory tasks. Oscillations in the theta (4–8 Hz) and gamma band (30–80 Hz) have been associated with neuronal activity responsible for active memory maintenance, whereas alpha-band (8–12 Hz) activity has been proposed to reflect inhibition of regions not required for the memory task (Tallon-Baudry et al. 1996, 2001; Gevins et al. 1997; Klimesch et al. 1999; Jensen et al. 2002; Lutzenberger et al. 2002; Pesaran et al. 2002; Howard et al. 2003; Kaiser et al. 2003; Lee et al. 2005; Scherberger et al. 2005).

The present study was carried out to characterize human oscillatory brain activity during the memory period in a delayed double-saccade task. More specifically, we addressed the question whether the neural mechanisms that give rise to the spatially selective and load-dependent BOLD activity observed in human extrastriate and parietal regions (Medendorp et al. 2006) would be reflected by similar modulations of oscillatory activity, originating from neural sources at corresponding locations.

To our knowledge, spatially tuned spectral activity in relation to memory-guided saccades has not been demonstrated in human subjects. In close connection though, using a spatial attention task, Worden et al. (2000) demonstrated a lateralized power distribution in the alpha band (8–12 Hz), in EEG sensors overlying occipital cortex. Although it is unclear whether the
same neural sources are involved in coding targets for delayed saccades, these data show that spatial-tuning effects on oscillatory activities can be measured noninvasively. Okada and Salenius (1998), who performed a single saccade task in MEG, observed strong alpha effects over occipital and parietal sensors but they did not address the issue of laterality or target load. Neither Okada and Salenius (1998) nor Worden et al. (2000) have identified the source locations of the alpha-band activity.

Here, we investigated oscillatory brain activity with magnetoencephalography (MEG), applying a slightly modified version of the delayed double-saccade paradigm used by Medendorp et al. (2006). This task has proven effective in dissociating target load and spatial selectivity for delayed saccades in fMRI. We examined spectral power while subjects saw 2 brief visual targets, separated by a short time interval, in either the left or right visual hemifield. Subjects were required to memorize these 2 target locations and, after another delay, make saccades to them in sequence. Our results show that the strength of spectral power at various posterior sensors was modulated by the number of target locations kept in memory as well as by the location of these targets within the visual hemifield. Using adaptive spatial filtering techniques, we localized the sources of the oscillatory activity in posterior parietal and occipital cortex.

Materials and Methods

Subjects
Eight healthy paid volunteers (5 males, 3 females, mean age of 23 years) participated in the experiments. All subjects gave their written informed consent in accordance with the institutional guidelines of the local ethics committee (CMO Committee on Research Involving Human Subjects, region Arnhem-Nijmegen, The Netherlands). Each subject practiced all tasks extensively before data acquisition to ensure that the tasks were performed correctly. Moreover, eye movement recordings and psychophysical measures were taken to confirm accurate behavior, as described below.

Experimental Setup
Subjects were seated upright in the MEG system that was placed in a magnetically shielded room. They were instructed to sit comfortably but still and to look at the stimulus screen, positioned at about 40 cm in front of them. Visual stimuli, generated with Presentation 9.10 software (Neurobehavioral Systems Inc., Albany), were presented using an LCD video projector (SANYO PROXtraX mutiverse, 60-Hz refresh rate) and back-projected onto the screen using 2 front-silvered mirrors. MEG data were recorded continuously using a whole-head system with 151 axial gradiometers (Omega 2000, CTF Systems Inc., Port Coquitlam, Canada). Head position with respect to the sensor array was measured using localization coils fixed at anatomical landmarks (the nasion and at the left and right ear canal). These measurements were made before and after the MEG recordings to assess head movements during the experiment. In addition, horizontal and vertical electro-oculograms were recorded using electrodes placed below and above the left eye and at the bilateral outer canthi. Electrode impedance was kept below 20 kOhm. During the experiment, these recordings were continuously inspected to check subjects’ task behavior and vigilance. MEG, electro-oculography, and electrocardiography signals were low-pass filtered at 300 Hz, sampled at 1200 Hz, and then saved to disk. Subjects’ psychophysiological performance was recorded by means of key presses using a button box (LUMI-Touch).

For each subject, a full-brain anatomical MR image was acquired using a high-resolution inversion prepared 3D T1-weighted scan sequence (flip angle = 15°; voxel size: 1.0 mm in-plane, 256 × 256, 164 slices, time repetition = 0.76 s; time echo = 5.3 ms). The anatomical MRIs were recorded using a 1.5-T whole-body scanner (Siemens, Erlangen, Germany), with anatomical reference markers at the same locations as the head position coils during the MEG recordings (see above). The reference markers allow alignment of the MEG and MRI coordinate systems, such that the MEG data can be related to the anatomical structures within the brain.

Experimental Paradigm
We investigated the temporal structure in human brain activity during delayed double-step saccades. Figure 1 illustrates the paradigm, which is a modified version of the fMRI paradigm used by Medendorp et al. (2006). Subjects were first asked to make an eye blink and then fixate centrally on a white cross. Subsequently, a red square (stim 1, size 0.7° × 0.7°) was flashed for 200 ms, either left or right of central fixation, at a random eccentricity between 9° and 18° and at a random angular elevation within a range of 6° and 13°, in either the lower or upper visual field. This was followed by a 2-s memory delay during which the subject maintained fixation. Then, another peripheral red square (stim 2) was flashed for 200 ms, in the same hemifield as the first stimulus. The location of the second stimulus was separated from the location of the first by at least 6° of visual angle. Subjects had to memorize the locations of both stimuli both of which were thus either right or left from the fixation direction. Next, 2 s after offset of the second stimulus, the fixation cross was turned off, prompting the subject to make saccades to the remembered locations of the stimuli, in the same sequence as they had appeared. To stimulate the subjects to memorize the spatial positions accurately, memory performance was probed in the following way: Small letters, either “c” or “o,” were shown from 300 to 400 ms after fixation cross offset at the original location of the first stimulus and from 600 to 700 ms after fixation cross offset at the original location of the second stimulus (not shown in Fig. 1). Due to their very small size and short appearance, the letters were distinguishable (“c” or “o”) only if the subject fixated at the correct location at the right time. This in turn could only be achieved if the subjects remembered accurately the
spatial positions of the 2 original targets. The letters were sufficiently small and short lived that subjects could not base their saccades on the presentation of the letters themselves. Subsequently, the central fixation cross (Fix) was turned on again, instructing the subject to make a saccade back to the centre of the screen and fixate till the end of the trial. During this period, subjects had to report which letters they had seen by pressing the corresponding buttons on a key pad using their right hand. The subject's actual performance was determined from the eye movement recordings, described below.

In addition, subjects performed trials in which they first blinked and then fixated centrally at all times, without presentation of any stimulus. These trials had the same duration as the test trials, described above, and served as baseline of the MEG recordings in further analyses. This baseline was not intended to match the memory trials in aspects like, for example, difficulty. Rather, it was meant solely to serve as low-level baseline for the normalization of power estimates obtained in the memory trials. The total duration of each trial was 9.35 s. Each subject performed 240 trials, consisting of 96 trials in which the 2 stimuli were flashed in the right visual field, 96 trials with stimuli in the left field and 48 baseline trials. The recording session was divided in 8 blocks of 30 trials, in which test and baseline trials were pseudorandomly interleaved. For each block of trials, subjects were given 1 min of rest, and their psychophysical performance hitherto was indicated on the screen. The total experiment lasted for 45 min.

Behavioral Analysis
As measured by the button presses, the subjects' performance on the letter detection task was over 80% correct. We did not exclude any trials based on this measure: the psychophysiology was merely incorporated to motivate subjects to perform optimally. Eye movement recordings in all subjects also confirmed that they generally followed the instructions correctly: on average in 6% of the trials did the subjects either break fixation (i.e., eye movements which exceeded the noise of our recording system) or make their saccades in the wrong direction. We excluded these error trials from further analysis.

Data Analysis
Data were analyzed using Fieldtrip software (http://www.mr.nl/fcdonders/fieldtrip), an open source Matlab toolbox for EEG and MEG data analysis developed at the Donders Centre for Cognitive Neuroimaging. As indicated above, we discarded trials in which subjects failed to keep eye fixation or made eye movements in the wrong directions. From the remaining trials, data segments that were contaminated with eye blinks, muscle activity or jump artifacts in the SQUIDs were excluded using semiautomatic artifact rejection routines. Power line noise was removed using a Fourier transformation of 10-s-long signal epochs and subtracting the 50 Hz component and its harmonics. This line noise was removed using a Fourier transformation of 10-s-long signal epochs and subtracting the 50 Hz component and its harmonics. This was done separately for all 10-s periods around all periods of interest from the continuous data record.

We analyzed the data in the frequency domain. For the sensor-level analysis, an estimate of the planar gradient was calculated for each sensor using the signals from the neighboring sensors. The horizontal and vertical components of the planar gradients approximate the signal measured by MEG systems with planar gradiometers. The planar field gradient simplifies the interpretation of the sensor-level data if sources are superficial and dipolar because then the maximal signal is located above the source (Hamalainen et al. 1993). A further advantage of planar gradiometers is that a superficial dipole activates a contiguous set of sensors (whereas for axial gradiometers, 2 separate sets are activated). This is important for cluster-based randomization statistics, as described below. Power spectra were computed separately for the horizontal and vertical planar gradients of the MEG field at each sensor and the resultant vector length of both was computed to obtain the power at each sensor location irrespective of the orientation of the gradient. For each subject, we visually inspected the averaged power spectral densities. Following the approach suggested by Klimesch (1999), we determined individual peak frequencies for the alpha (8-12 Hz) and beta bands (13-25 Hz), as shown by Table 1.

In the actual analysis of the data, we examined the changes in spectral power in the test trials relative to the baseline trials to determine the neural response related to the task. To investigate target load effects, we compared power differences between data epochs in which one target was memorized (first delay period 1, see Fig. 1) and data epochs in which 2 targets were kept in memory (second delay period 2). Likewise, we examined laterality effects by comparing the power in those trials in which the stimuli appeared in the left visual field with those in which stimuli were presented in the right visual field, for both the first and second memory period. We considered the power changes at various frequency bands, using the data at the individual peak frequencies when examining scalp topography across subjects (see Figs 2, 4, and 5). Using a jackknife procedure (Efron and Tibshirani 1991), we determined the variance of the power in the selected frequency bands separately for each time point across trials. Using these estimates, we expressed the difference in power between 2 conditions as a t-score separately for each subject. The resulting t-scores were transformed into z-scores and pooled across subjects to obtain a fixed-effects measure of the significance of the mean power change across subjects for each of the frequency bands. The critical value was Bonferroni corrected with

<table>
<thead>
<tr>
<th>Subject</th>
<th>Alpha band (Hz)</th>
<th>Beta band (Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>10</td>
<td>21.5</td>
</tr>
<tr>
<td>S2</td>
<td>9.5</td>
<td>18.5</td>
</tr>
<tr>
<td>S3</td>
<td>9</td>
<td>23.5</td>
</tr>
<tr>
<td>S4</td>
<td>10.5</td>
<td>20</td>
</tr>
<tr>
<td>S5</td>
<td>11.5</td>
<td>19</td>
</tr>
<tr>
<td>S6</td>
<td>9.5</td>
<td>17</td>
</tr>
<tr>
<td>S7</td>
<td>10.5</td>
<td>20</td>
</tr>
<tr>
<td>S8</td>
<td>10.5</td>
<td>23.5</td>
</tr>
<tr>
<td>Mean</td>
<td>10.1 ± 0.8 (standard deviation)</td>
<td>20.4 ± 2.3</td>
</tr>
</tbody>
</table>

Figure 2. Topographic distribution of power in the alpha band (8-12 Hz) averaged across subjects. (A, B) Suppression of the alpha-band power relative to baseline for stimuli presented in the left visual hemifield, for the first (delay 1) and second (delay 2) retention period, respectively. (C) Difference between delay 2 and delay 1 showing that alpha is further suppressed with target load. Middle panels, alpha-band suppression for stimuli in the right visual hemifield (D, E) and their difference (F). The degree of laterality does not increase with target load. Blue regions indicate a stronger suppression for remembered target locations in the left hemifield compared with the right. Red regions represent the opposite pattern. Open circles: positions of sensors selected in further analysis. Each map has been scaled equally between −9 < z < 9 (following the color range indicated) |z| > 3.77, P < 0.05 Bonferroni corrected. LH/RH: left/right hemisphere.
the number of sensors and time points, corresponding to $|z| > 3.77, P < 0.05$ for Figures 2 and 5, and $|z| > 4.07, P < 0.05$ for Fig. 4). We used repeated-measures analyses of variance for post hoc comparative power analysis of selected sensor groups across subjects, setting the type I error at the 0.05 level ($P < 0.05$).

Time-frequency representations (TFR), estimating the time course in power, were computed using a multitaper method. This approach seeks to optimize spectral concentration over the frequency of interest (Mitra and Pesaran 1999). We analyzed 2 frequency ranges separately: 1–30 Hz (alpha/beta) and 30–120 Hz (gamma). The lower band was analyzed

Figure 3. Regions within the posterior parietal and occipital cortex (transversal view) that show contralateral suppression effects in the alpha band during the first (A) and second (B) delay period. Population average of power difference between Target-Left and Target-Right conditions. Only significant clusters of voxels are shown ($P < 0.05$, randomization statistics with correction for multiple comparisons). The opacity of each voxel is scaled by its z-value. Blue regions indicate a stronger suppression for remembered target locations in the left hemifield compared with the right. Red regions represent the opposite pattern. LH/RH: left/right hemisphere.

Figure 4. Temporal development of the topography in the alpha band (8–12 Hz). (A, B) Alpha-band suppression relative to baseline for stimuli in the left and right visual hemifield, respectively. (C) Difference in alpha-band activity between leftward and rightward stimuli showing phasic and tonic components. Blue regions indicate a stronger suppression for remembered target locations in the left hemifield compared with the right, and vice versa for red regions. $|z| > 4.07, P < 0.05$ Bonferroni corrected. LH/RH: left/right hemisphere.
A spatial filter was constructed that passes activity originating from this volume was divided into a regular 8-mm 3D grid. For each grid point, we applied an adaptive spatial filtering or “beamforming” technique (Gross et al. 2001; Hoogenboom et al. 2006). Each subject’s brain was also spatially normalized to the International Consortium for Brain Mapping template (Montreal Neurological Institute, Montreal, Canada). The individual anatomical MRIs and the corresponding statistical maps were spatially normalized to a fixed effect statistic using the same procedure as for the sensor data. Multiple comparison corrections at the source level were made by applying a 3D cluster-randomization analysis (which operates along the lines described above for the TFR cluster-randomization analysis).

Results

We investigated the modulations of oscillatory activity with target location and target load in a double-delayed double-saccade task. In this task, subjects first saw the target of the first saccade, then, after a 2-s delay, the target for the second saccade and then, after another 2-s delay, they made saccades successively to the remembered stimulus locations (see Fig. 1). The paradigm had two different conditions of interest regarding the locations of the 2 targets: both were cued either in the right visual field (Target-Right condition) or in the left visual field (Target-Left condition).

As a first step in our analysis, we determined the scalp topography of the changes in power in the alpha band (8–12 Hz) (expressed as a z-score) for the 2 memory periods (always excluding the first 400 ms after target offset in order to reduce contributions from visually evoked fields) in the 2 task conditions, respectively, relative to the baseline trials (no targets). Figure 2, top row, presents the results of this analysis for the Target-Left condition. Panel A, illustrating the alpha topography for the first retention interval (one-target memory: delay 1), shows a clear decrease in alpha-band activity relative to baseline, strongest over the parietal areas in the right hemisphere.

What happens when the location of a second saccade target must be stored in memory? Panel B, showing alpha changes for the second memory period (2-target memory: delay 2), demonstrates a further but bilateral suppression of alpha relative to baseline over the posterior areas. Panel C then shows the effect of target load by plotting the difference in alpha power for the 2-target and the one-target load in the Target-Left condition (delay 2-delay 1). As this plot shows, alpha power decreases with increasing target load but almost equally in the posterior areas of both hemispheres.

Figure 2, middle row, shows the same analysis for the memory intervals in the Target-Right condition. Again, in this condition, alpha power decreases in the hemisphere contralateral to the remembered target location (panels D and E). The effects of target load, depicted in panel F, are consistent with the pattern in C showing that alpha power decreases with increasing memory demands. Although this effect can be seen in both hemispheres, the alpha decrease is most notable in the right (ipsilateral) hemisphere.
To determine the spatial selectivity of memory storage in more detail, we compared the alpha power for the Target-Left and Target-Right conditions. Thus, we subtracted the alpha activity for targets in the right visual field from the activity for targets in the left visual field. The results are depicted in Figure 2, bottom row, showing a clear contralateral suppression of alpha power for both the one-target memory interval (delay 1, panel G) and the 2-target memory period (delay 2, panel H). Blue regions indicate a stronger suppression for remembered target locations to the left than to the right of fixation, and vice versa for red regions. Note the hemispheric asymmetry in the pattern of alpha suppression. The suppression seems stronger and to cover a larger region in the left hemisphere than in the right. But clearly, alpha is suppressed in the hemisphere contralateral to the hemifield of stimulus presentation, most prominently over the sensors marked by the open circles. Consistent with these observations, a repeated-measures multivariate analysis of variance (MANOVA) over these sensors, with hemisphere (left/right) and target location (left/right hemifield) as factors, revealed a significant 2-way interaction for both the one-target memory period ($F_{1,7} = 11.7, P = 0.011$) and the 2-target memory period ($F_{1,7} = 9.6, P = 0.018$). Finally, Figure 2(I) demonstrates the interaction between the laterality effects and target load. As shown, the degree of laterality did virtually not depend on target load; if anything, it seems reduced during the 2-target memory period. Thus, in summary, 1) alpha is suppressed in the hemisphere contralateral to the hemifield of stimulus presentation and 2) alpha power decreased with the same amount in both hemispheres, when memory load is increased from one to two target locations.

Where in the brain can the oscillatory sources be identified that account for the contralateral suppression in the alpha band, as depicted in Figure 2? Spatial filtering based on the cross-spectral density matrix (“beamforming,” see Methods) was used to estimate the sources in the brain that gave rise to the laterality effects in the alpha band, as observed on the scalp (Fig. 2G,H). Figure 3 provides an overview of these results based on standardized averaged group results, for the one-target memory (Fig. 3A) and the 2-targets memory period (Fig. 3B), showing only significant clusters of voxels based on a randomization analysis correcting for multiple comparisons ($P < 0.05$). Again, as in Figure 2(G,H), blue regions show a stronger suppression to leftward than rightward targets, whereas red regions represent the opposite. As shown, in both Figure 3(A,B), widespread contralateral alpha suppression can be observed in extrastriate brain regions. More precisely, significant contralateral suppression extended from close to the intraparietal and parietal-occipital sulcus into anterior-occipital cortex. It should also be noted that no such bilateral topographic regions were identified in frontal cortex, in this frequency range.

Thus far, the tuning and target load effects in the alpha band were observed as averaged effects during the retention periods. To determine the temporal evolution of the alpha activity, Figure 4 demonstrates the alpha topography at various time intervals of the Target-Left and Target-Right condition (2 upper rows), as well as the difference between them (bottom row). As shown, alpha modulations with respect to the baseline trials were absent just before and during the onset of the first stimulus (~0.2 to 0.2 s). Then, at the first ~400 ms following the offset of the first stimulus, which characterizes the phasic response, there was a clear contralateral suppression in the alpha band at the posterior sensors, which was sustained at a slightly lower level throughout the first retention interval (0.6–2.0 s). Around the presentation of the second stimulus, alpha activity increased again but remained below baseline level. At the intervals of the second retention period, 2.4–2.8 s and 2.8–4.2 s, respectively, there were similar phasic and tonic responses. Finally, at the end of the trial, at the onset of the eye movements, alpha suppression vanished, as if alpha played no role in actually executing the response.

The effects described above were not restricted to the alpha band. Also the beta band (13–25 Hz) showed clear suppression effects, which became stronger after presentation of the second target, with a similar topography as the alpha band. This is shown in Figure 5, in the same format as in Figure 2. Compared with the alpha band, however, the degree of laterality was less robust. A repeated-measures MANOVA over the selected sensors (open circles), with hemisphere (left/right) and target location (left/right hemifield) as factors, revealed only a significant 2-way interaction for the one-target memory period ($F_{1,7} = 6.4, P = 0.039$). Laterality of the activity during the 2-target memory period remained below statistical significance ($F_{1,7} = 4.0, P = 0.087$).

Figure 6(A) (top panel) illustrates TFR of power difference between the Target-Left and Target-Right conditions for frequencies between 1 and 30 Hz, for the sensors marked in Figure 2. Data are presented in a pooled comparison across hemispheres computed as the power difference between contralateral and ipsilateral target conditions for each sensor, which was then averaged across all selected sensors and subjects. The bottom panel illustrates the statistically significant time-frequency clusters based on a cluster-randomization approach for multiple comparisons ($P < 0.05$, see Methods). The contralateral suppression effects are significant in the alpha and beta bands, as described above. In the beta band, the sustained, tonic component seems much weaker than the phasic component, suggesting that it may not be related to memory retention. Furthermore, with the frequency range 1–8 Hz there was a significant contralateral transient enhancement of activity in the theta range (at about 5 Hz) in response to each of the contralateral and ipsilateral target conditions for each sensor, which was again but remained below baseline level. At the intervals of the presentation of the second stimulus, alpha activity increased clear and significant ($P < 0.05$) lateralized activity, around 60–90 Hz, in a 1-s period prior to the initiation of the saccades.

To further examine this locus of activity, Figure 7(A) demonstrates the power modulations in the higher frequency bands (gamma band), in the same format as Figure 6(A). Enhancements can be discerned in response to both stimuli, with $P < 0.1$ for the first cue and $P < 0.05$ for the second cue but note that these enhancements are not retained into the memory periods. In fact, the first delay period does not show any significant power component within this frequency range. The second delay period, however, does demonstrate clear and significant ($P < 0.05$) lateralized activity, around 60–90 Hz, in a 1-s period prior to the initiation of the saccades.
They demonstrate enhancements at about 40–60 Hz, prior to the presentation of the first stimulus. Those enhancements likely reflect the anticipation of a test trial as opposed to a baseline trial. Also, following the go cue for the saccades, a substantial gamma-band component can be noticed. More importantly, however, power in the gamma band was reduced relative to baseline during the first delay period, irrespective of whether a contralateral or an ipsilateral target was kept in memory. During the second delay period, gamma power remained at a negative level when 2 ipsilateral targets were memorized but increased during the memory of 2 contralateral targets, most prominently in the frequency range 60–90 Hz and toward the end of the second delay period. As such, this would explain the findings of Figure 6(B): The gamma band shows a general desynchronization during the delay intervals, with less desynchronization for contralateral than ipsilateral targets during the second delay period. This observation is replotted in Figure 7(C), only for the second delay period and clusters that are significant at \( P < 0.05 \). It clearly shows that the gamma-band lateralization was not sustained during the entire duration of the period but arises at some point in time during the second delay period, perhaps when subjects start planning their saccades. In this respect, the contralateral gamma effects during the second delay period may reflect a neuronal correlate for preparatory set associated with saccade direction (Pesaran et al. 2002).

**Discussion**

We have investigated modulations of power in various frequency bands when subjects are processing and storing spatial information in a delayed double-step saccadic working memory task. To do so, we used a slightly modified version of the double-saccade experiment by Medendorp et al. (2006) performed in fMRI. Subjects were tested in a paradigm that employed 2-s delays between the occurrence of the first target, the second target, and the saccadic responses. In effect, this paradigm allowed us to dissociate the modulations in spectral power related to the first target (one-target memory load) and the first and second targets together (2-targets memory load). Our results showed a significant spatially selective suppression of power in the alpha and beta bands over posterior sensors during spatial working memory maintenance. This suppression was strongest over the hemisphere contralateral to the hemifield of the presented cue. We further found that oscillatory power in these bands decreased after presentation of the second cue but this memory effect itself did not exhibit any clear laterality. We localized the neural sources of these effects in parietal and occipital areas. Working memory maintenance of one target reduced also higher frequency (40–120 Hz) power over posterior sensors. However, in comparison with one-target memory, 2-target memory led to an increased gamma-band (60–90 Hz) power in the hemisphere contralateral to the targets, most notable shortly before the initiation of the saccades.

Let us now discuss the putative roles of the observed rhythms and their relation to previous interpretations. Recent studies, simultaneously recording electrophysiological and hemodynamic signals, have shown negative correlations between hemodynamic signals and spectral power in the alpha band (8–12 Hz) (Goldman et al. 2002; Laufs et al. 2003; Moosmann et al. 2003) but positive correlations in the gamma band (30–90 Hz) (Logothetis et al. 2001; Niessing et al. 2005). On this basis, our sources for lateralized alpha suppression in anterior-occipital and posterior parietal cortex are consistent with findings of recent fMRI studies that reported enhanced and directionally
selective BOLD activation at virtually the same locations during the retention of a working memory for delayed saccades (Sereno et al. 2001; Medendorp et al. 2003, 2005; Koyama et al. 2004; Schluppeck et al. 2005). Also several studies using macaque monkeys have described areas at similar locations showing sustained activity during the memory period in delayed-saccade tasks, including extrastriate area V3A (Nakamura and Colby 2000) and posterior parietal area LIP (Barash et al. 1991; Colby et al. 1996; Mazzoni et al. 1996; Pesaran et al. 2002). The lateraled enhancement in gamma-band power during the second delay period compared with the first delay period also corroborates previous fMRI findings (Medendorp et al. 2006) showing directionally-selective increases in BOLD activation in human posterior parietal cortex with memory load in a double-saccade task. Even the apparent gamma depression during the one-target memory (Fig. 7) does not necessarily disagree with earlier findings but can be explained by the difference in spatial resolution between BOLD and MEG imaging, as we will further argue below. All together, our findings are well in accord with the notion that a decrease in alpha activity and increase in gamma-band activity reflect the engagement of a cortical region whose neural activity can also be measured as an increased BOLD signal or discharge rate in a spatial working memory task.

A reduction in alpha power is generally interpreted as a marker of activated brain regions (Singh et al. 2002). Areas not processing sensory information or motor output have been shown to retain increased power in the alpha band. Basically, our results showing clear suppression of alpha compared with its level obtained in the separate baseline trials are in support of this view. Our results also show that alpha is selectively suppressed in specific posterior regions, with more suppression in the hemisphere contralateral to the target. Again, this is compatible with the idea that activated brain regions are characterized by a reduction of alpha-band activity.

Our alpha findings confirm the results of Okada and Salenius (1998), who found a sustained suppression of alpha-band activity during a spatial working memory task, most prominently over posterior areas. Which role should be attributed to the sustained component? When subjects in the Okada and Salenius study passively viewed the stimuli, the sustained suppression was no longer present. This suggests that the tonic alpha decrease is involved in a function that transcends simple visual analysis. Worden et al. (2000) found this marker of neural processing to be directionally selective in a spatial attention task, with more alpha suppression over posterior cortex contralateral to the direction of attention (see also Sauseng et al. 2005; Kelly et al. 2006). Our findings add the notion that alpha power suppression is also lateralized during the coding of working memory for saccades. This is compatible with the idea that coding attention and planning saccades involve the same neural network (Posner et al. 1980; Rizzolatti et al. 1987; Findlay and Walker 1999).

Is the reduction in alpha a specific prerequisite for maintaining a working memory, or just a general mechanism operative to put particular brain areas into function? Working memory studies have revealed variable results as to the activity in the alpha band. Jensen et al. (2002) reported increases in alpha with memory load in a Sternberg task with letters. The same was found by Krause et al. (1996) in an auditory memory task. Also Klimesch et al. (1999) found alpha activity to be enhanced with working memory demands. By contrast, Gevins et al. (1997) found this marker of neural processing to be directionally selective in a spatial attention task with letters. The same was found by Krause et al. (1996) in an auditory memory task. Also Klimesch et al. (1999) found alpha activity to be enhanced with working memory demands. By contrast, Gevins et al. (1997) reported decreased alpha in an n-back spatial working memory paradigm. Also the present study was performed in the spatial domain. The picture arising from these various results is that alpha is reduced when spatial functions are to be performed, whereas it is increased during nonspatial working memory tasks.

This basic result is incompatible with the notion of alpha activity as a general mechanism of working memory representations. In the same vein, it cannot be interpreted in support of the idling hypothesis (reviewed in Pfurtscheller et al. 1996). Rather, it argues in favor of a regulatory mechanism, put in place to allocate resources processing visuospatial information (Kelly et al. 2006; Thut et al. 2006). This puts forward a striking analogy with the classical idea that spatial and nonspatial functions are segregated in the brain, within a dorsal pathway and a ventral pathway, respectively (Ungerleider and Mishkin 1982). We propose that the alpha decrease during spatial working memory maintenance is due to an engagement of the dorsal stream.
Within this context, it can then be easily understood that the degree of alpha synchronization depends on the nature of the memory task. The increase in alpha power with working memory load during maintenance of letters or faces (Jensen et al. 2002) is due to an inhibition of the dorsal stream while the ventral stream is being engaged. More studies are required to test if this relationship holds up in general.

Interestingly, in the present study, we found that the degree of alpha lateralization did not further increase with target load, even though the alpha power was bilaterally reduced after presentation of the second target (Fig. 2). One possible reason for this effect is that the observed alpha laterality reflects anticipation of the second target (Worden et al. 2000). That is, after presentation of the first target, subjects anticipate the presentation of the second target which will be in the same hemifield. It could also mean that the laterality relates to the encoding of the first target, which is the goal for the first action, and remains this even after the second target has been presented. If so, this would suggest that the modulation in alpha power cannot be explained by visuospatial processing alone, but is also due to target selection for saccades. To distinguish between these and other possible explanations it would be useful to repeat the experiment with the second target presented at a location randomized across the 2 hemispheres (Medendorp et al. 2006).

Virtually the same sensors showing alpha laterality also exhibited contralateral suppression effects of power in the beta-band (Fig. 5). This finding is consistent with numerous earlier studies reporting beta-band desynchronization in activated brain regions. Specifically, beta-band power over sensorimotor cortex of humans was found reduced during hand movements (Salmelin et al. 1995), beta-band activity in visual cortex was reduced during visual stimulation (Hoogenboom et al. 2006), and beta-band activity over typical language areas was reduced during language tasks (Ressel et al. 2006).

It has often been found that a desynchronization in the lower frequency bands (alpha, beta) goes hand-in-hand with synchronization in the higher frequency bands (gamma) (Munk et al. 1996; Fries et al. 2001; Schoffelen et al. 2005; Bauer et al. 2006; Hoogenboom et al. 2006). Moreover, several reports have suggested that oscillatory firing in the gamma band may be significant for active memory maintenance in delayed response tasks (Tallon-Baudry et al. 1998, 2001; Pesaran et al. 2002; Howard et al. 2003). For example, Tallon-Baudry et al. (1998) have reported clear enhancements of gamma-band activity in occipitotemporal and frontal regions during the delay in the memory condition of a delayed-matching-to-sample task with visual shapes. Pesaran et al. (2002) found that, during delays when monkeys are planning a saccade, there are broadband gamma oscillations (25–90 Hz) in the local field potentials (LFP) of posterior parietal area LIP. They found the LFP tuned to the direction of planned movements, changing its strength with behavioral state. Therefore, at first, one might be surprised by the predominant decrease of power in the gamma band during the memory periods in the present oculomotor paradigm (Fig. 7A). One possible explanation comes from recent evidence that local gamma enhancements are surrounded by more widespread gamma reductions (Lachaux et al. 2001; Shmuel et al. 2006). Studies using intracranial recordings typically do not see this effect, because they usually tailor stimulation to the recording site to obtain focal activation. Also BOLD-fMRI may attain high enough spatial resolution to not be contaminated by these effects (Logothetis et al. 2001; Medendorp et al. 2006). However, MEG signals that average over a wider region of cortex might be dominated by the widespread gamma reduction in the surround such that the net effect at the MEG sensor is a gamma power reduction. This notion is fully compatible with our finding that contralateral gamma-band power did increase during the second delay period as compared with the first (see Figs 6B and 7C). The available evidence suggests that gamma-band synchronization in the absence of appropriate stimuli or tasks is weak. It is thus reasonable to assume that the gamma desynchronization (surrounding a focal synchronization) essentially abolishes any (nonspurious) gamma-band synchronization in the surround. As a consequence, the resulting level of minimal gamma-band synchronization would not be further reduced even if more focal activations would be added, as, for example, in our case by adding the second target. This model can explain the observations in the gamma-band: The first target induces one focus of enhanced gamma synchronization surrounded by widespread gamma desynchronization with the net effect at the MEG sensor of reduced gamma power. The second target adds a second focus of enhanced gamma synchronization, while the surround is already completely desynchronized and does not desynchronize further. The net effect at the MEG sensor is an increase in gamma power relative to the first target. It remains important to emphasize that the increased activity did not occur over the total duration of the second delay period, but only arose shortly before the initiation of the saccades. In a strict interpretation, the laterality effects that are observed during the second delay period do not so much reflect a target load effect but perhaps more so a motor planning effect, which would be consistent with the findings by Pesaran et al. (2002) for monkey parietal cortex.

In summary, the present study has provided evidence that changes in synchronized oscillatory neural activity are tuned to specific regions of space and modulated by target load during memory periods in an oculomotor task. Future research is required to determine more precisely how neural signals are processed in regions that are involved in storing and transforming spatial information into motor action.

Notes

We thank Markus Bauer for assistance with the data analysis and Stan van Pelt for comments on an earlier version of the manuscript.

Grants: This research was supported by grants from the Human Frontier Science Program (P.M., P.F.), the Netherlands Organization for Scientific Research (P.M., P.F., O.J.), and the Volkswagen Foundation (P.F.). Conflict of Interest: None declared.

Address correspondence to W. P. Medendorp, PhD, Nijmegen Institute for Cognition and Information, Radboud University Nijmegen, PO Box 9104, NL-6500 HE Nijmegen, The Netherlands. Email: p.medendorp@nici.ru.nl.

References


