Prospective memory (PM) denotes the function to realize intentions after a delay while being immersed in distracting ongoing (OG) activity. Here, we scrutinize the often-reported involvement of rostral prefrontal cortex (rPFC; approximating Brodmann area 10) in such situations: This region might mediate attention between external stimuli and the internally maintained intention, that is, between stimulus-oriented (SO) and stimulus-independent (SI) processing. Using functional magnetic resonance imaging (fMRI) we orthogonally crossed 1) PM versus OG activity only, with 2) SO versus SI attention. In support of the hypothesis, common regions of medial rPFC exhibited greater blood oxygen level–dependent (BOLD) signal for the contrasts of both OG task only versus PM and SO versus SI attending. However, activation related to the former contrast extended more superiorly, suggesting a functional gradient along a dorsal–ventral axis within this region. Moreover, region-of-interest analyses revealed that PM versus OG task only was associated with greater BOLD signal in left lateral rPFC, reflecting the requirement to maintain delayed intentions. Distinct aspects of this region were also transiently engaged at transitions between SO and SI conditions. These results are consistent with the hypothesis that some of the rostral prefrontal signal changes associated with PM performance reflect relative differences in SO versus SI processing.

Keywords: anterior prefrontal cortex, delayed intentions, frontopolar cortex, fMRI, task switching

Introduction

Prospective memory (PM) denotes the capacity to remember to carry out an intention after a delay (e.g., posting a letter), while being immersed in distracting ongoing activity (OG; e.g., commuting to work) (Ellis 1996; McDaniel and Einstein 2007). Typically, the implementation of the intention has to be self-initiated upon occurrence of a particular event (i.e., the PM target; e.g., presence of a postbox). Thus, PM requires a fine attentional balance between information that is externally derived (e.g., monitoring the traffic) versus internally maintained (e.g., the intention).

Realizing such delayed intentions critically depends on rostral prefrontal cortex functioning (rPFC; approximating Brodmann area [BA] 10): Lesions to this region lead to impairments in PM, typically in the context of spared episodic memory abilities (Burgess et al. 2000, 2009; Roca et al. 2010; Urczek and Gilboa 2010; Volle et al. 2011). Neuroimaging studies have further specified the temporal involvement of this region. rPFC appears to be engaged during the delay period between intention formation and execution (Okuda et al. 1998, 2007; Burgess et al. 2001, 2003; den Ouden et al. 2005; Simons et al. 2006; Gilbert et al. 2009, 2011; Reynolds et al. 2009). These studies employed a great variety of different OG tasks, intentions, stimuli, and response modalities, often within single experiments (i.e., conjunction approach). Furthermore, a recent study by Gilbert (2011) found that although lateral rPFC (lrPFC) exhibited robust activation during maintenance of delayed intentions, the content of those intentions (i.e., the nature of the target stimuli and appropriate PM responses) could not be decoded from this region. These data suggest that rPFC subserves central aspects of PM, that is, those that are not specific to individual stimuli, responses, or tasks. Importantly, this region is recruited when participants are instructed to carry out delayed intentions, but no actual PM targets are embedded in the OG task (Burgess et al. 2001; Simons et al. 2006). Similarly, lrPFC exhibits increased blood oxygen level–dependent (BOLD) signal during PM performance, even when statistically controlling for transient signal changes at the moments of target detection (Reynolds et al. 2009). Therefore, Burgess et al. (2001) concluded that this region supports the maintenance of the delayed intention in the context of OG task activity rather than target detection or actual realization of the intention (see also Okuda et al. 1998). This account is consistent with functions attributed to rPFC, such as the preparation for upcoming task demands (Sakai and Passingham 2003; Rowe et al. 2007), the coordination of multiple tasks (Koechlin et al. 1999; Braver and Bongiolatti 2002; Benoit 2008), or the integration of independent cognitive operations (Ramnani and Owen 2004).

However, PM performance is not just associated with increased rPFC activation. Instead, BOLD signal typically exhibits a double dissociation within this region, when contrasting conditions that require either sole engagement in OG activity (OG blocks) or additional performance of delayed intentions (PM blocks) (Burgess et al. 2008). Activation in lateral rPFC is commonly greater for PM compared with OG blocks, whereas the reverse contrast is associated with medial rPFC (mrPFC) recruitment. A complete account of rPFC involvement in PM thus
needs to explain this full pattern, and the gateway hypothesis of rPFC function (Burgess et al. 2007, 2008) has recently provided such a comprehensive account.

It posits that mrPFC and lrPFC comprise a gateway mechanism that mediates competition between stimulus-oriented (SO; based on the current environment) versus stimulus-independent (SI; decoupled from the environment) processing. This mechanism is thought to be engaged when either processing mode needs to be biased to an unusual degree or in situations that require frequent switches between the 2 modes. Accordingly, engagement for sole OG task performance versus additional PM performance has been attributed to relative differences in such attentional demands (Burgess et al. 2003, 2008): Whereas OG task activity on its own primarily requires attention toward external stimuli (i.e., SO attending; e.g., monitoring the traffic), the additional PM component necessitates a relative disengagement from the external environment. That is, this condition also demands SI maintenance of the intention (e.g., thinking about posting the letter) and frequent matching of the external world and the internally represented PM target (e.g., watching out for a postbox).

The gateway hypothesis is supported by a number of neuroimaging studies, all of which provided evidence for a consistent functional dissociation between mrPFC and lrPFC. Gilbert et al. (2005), for instance, instructed participants to perform 3 tasks, alternately based on externally presented (SO phase) or internally generated (SI phase) stimuli (see also Gilbert et al. 2007; Dumontheil, Gilbert, et al. 2010). Consistent across all tasks, SO contrasted with SI phases were associated with sustained BOLD signal increases in mrPFC. In contrast, lrPFC was transiently recruited when participants switched between SO and SI phases. These findings have been extended to a variety of different forms of SO versus SI attending, and lrPFC has been observed to also exhibit sustained activation during phases of SI processing (Gilbert, Simons, et al. 2006; Dumontheil, Gilbert, et al. 2010; Dumontheil, Hassan, et al. 2010; Henseler et al. 2011). Taken together, this pattern supports the account that mrPFC is involved in SO attending, whereas lrPFC supports SI attending (cf. Christoff and Gabrieli 2000).

Thus, maintaining an intention may activate lrPFC due to increased reliance on SI attending, while mrPFC deactivation may reflect concurrent attenuation of SO processing of the OG task (Burgess et al. 2003; Simons et al. 2006). In this case, at least some of the neuronal populations involved in PM might also be sensitive to a manipulation of SO versus SI attending. However, to date, no study has examined if both functions in fact recruit overlapping regions. Indeed, rPFC exhibits a functional specialization on a fine spatial scale (on the order of a few millimeters) (Gilbert, Spengler, et al. 2006; Gilbert et al. 2007; Gilbert et al. 2010; see also Krueger et al. 2007; Benoit et al. 2010), suggesting that areas implicated in PM might be distinct from those associated with the attentional gateway mechanism.

To test the gateway account of rPFC involvement in PM, we employed functional magnetic resonance imaging (fMRI) in a factorial design. Specifically, we crossed the requirements to engage in 1) PM versus OG only and 2) SO versus SI attending. This allowed us to assess whether at least some aspects of rPFC involved in PM performance are also involved in mediating between SO versus SI attending. Moreover, the factorial design enabled us to explore the general functional properties of rPFC in more detail, in addition to examining functional overlap versus segregation. If, for example, both the SI (vs. SO) condition and the PM (vs. OG only) condition recruit overlapping aspects of lrPFC, they might yield additive or multiplicative effects on BOLD signal. This would suggest that increasing the demand on SI processing increases the engagement of this region. On the other hand, an interaction effect between the 2 factors (i.e., the PM condition has an effect during SO but not SI phases) may suggest that this region does not need to be further engaged once the system is already in a state of SI processing.

Materials and Methods

Participants

Nineteen volunteers participated in this experiment. They were all right handed, had normal or corrected-to-normal vision and reported good health with no known history of neurological or psychiatric illness. Prior to the experimental session, they gave written informed consent. All participants received £ 30 reimbursement as approved by the local research ethics committee. Of the 19 participants, 3 had to be excluded from further analysis either due to technical problems, miscomprehension of task instructions, or chance performance. Thus, 16 participants (9 females; mean age = 22.51 years, age range = 19–28 years) remained for further analyses.

Tasks and Procedure

Participants performed 2 tasks (Fig. 1 and description below) that required responses either based on visually presented information (i.e., SO condition) or internally generated information (i.e., SI condition). These stimulus conditions changed randomly within experimental blocks. Thus, both tasks alternated between phases of SO and SI processing. Apart from this critical commonality, they were designed to differ in other aspects of required cognitive operations, that is, spatial navigation versus line discrimination (see below; Gilbert et al. 2005). Engagement in the basic tasks constituted the OG condition, whereas the PM condition additionally required carrying out delayed Intentions. Participants only learned about the PM condition in the MRI scanner, after they had performed the OG task alone.

In the "shape task," participants continuously navigated around the edges of a shape in clockwise direction. For each corner, they indicated whether they would have to take a left or right turn. In SO phases, the actual shape was presented, which resembled the outlines of the letters H and F attached to each other (Fig. 1). In contrast, during SI phases, the HF shape was replaced by a distractor, that is, a mirrored version of the joined letters T and E. Participants were instructed to continue the task by picturing the HF shape in their head, while keeping their eyes open and fixated on the center of the screen. At transitions between stimulus phases, they continued the sequence from where they had left off. The outlines of both shapes were white and covered approximately 6° of the visual field. The PM targets were 2 junctions on the HF shape, which participants memorized at the beginning of each PM block. Whenever they got to these corners, they had to indicate their detection rather than making the right/left decision.

In the "alphabet task," participants classified capital letters based on perceptual features. Specifically, they indicated whether any given letter was composed entirely of straight lines (e.g., A, H) or included any curved lines (e.g., C, P). In the SO condition, letters were presented in alphabetical order one at a time. As soon as a response was made, the letter was replaced by the next. When the end of the alphabet was reached, the sequence continued at "A." Letters were presented either in red or blue during SO phases. In contrast, during SI phases, letters were presented in random order and in the other color (i.e., blue or red) (Fig. 1). In this condition, participants had to ignore the distracting random letters and mentally continue the correct alphabetical sequence whilst carrying on with the classification based on the imagined letters. At transitions to the SO condition, the letter was
presented that participants would have imagined if they had correctly continued the alphabetical sequence throughout the SI condition. Letters always covered approximately 1° of the visual field. The PM targets were 3 letters, which were presented as a word (e.g., CUP) at the beginning of each PM block (see Supplementary Material). Presenting the targets as a word made them more memorable, which served to reduce the episodic memory component of the PM condition. Whenever participants reached one of the target letters, they had to perform an alternative response instead of the classification judgment.

Each block started at a unique position or letter but always with an SO phase for 2 s (Fig. 1). The starting point for the shape task was indicated by a green arrow, which disappeared after the first response. Blocks lasted for 7.5 s and were comprised of miniblocks. Their duration varied between 3 and 7 s to allow for an efficient estimation of BOLD signal changes and capped at 7 s to ensure a sufficient number of transitions between stimulus phases (Gilbert et al. 2005). Stimuli were presented at 1 of 2 possible screen locations (separated diagonally by ca. 6°) throughout a miniblock. At the beginning of each miniblock, the stimulus changed position to the other location, and the current stimulus phase (SO, SI) was randomly selected (Fig. 1). A ‘stay’ trial is the first one of a new miniblock, if the stimulus phase has not changed relative to the last miniblock. In contrast, a ‘switch’ trial marks the beginning of a miniblock after a change of stimulus phase. (All other OG trials are ‘non-switch’ trials.) Changing the stimulus location on stay trials provided a baseline for switch events, controlling for visual transients at the transition between stimulus phases. Furthermore, PM targets, that is, either junctions marked on the shape or the 3-letter word, were presented for 10 s directly before the blocks started.

The frequency of the PM targets was similar in both tasks (alphabet: 11.54%; spatial: 11.11%). They were chosen so that the delayed intention could not easily be incorporated in the OG task (see Burgess et al. 2003). Specifically, at least 4 OG trials preceded the appearance of the first target, and successive targets were separated by at least 4 intervening OG trials. The minimum gap between subsequent targets ensured that detection of one PM target could not act as a strong retrieval cue for the imminent next target. Thus, processes involved in target detection and intention realization had to be engaged for each individual target. Moreover, to avoid automatized processing of PM targets, each was used for a single block only, and the gap between targets varied pseudorandomly across blocks. This rendered the target sequence unpredictable and served to minimize the awareness for any rule underlying the sequence. In addition, half of the targets were from either half of the shape or alphabet, and they were equally often associated with either OG response (e.g., left or right key). Finally, the 3-letter words were chosen to lack strong emotional connotations. Participants responded with the index, middle, or ring finger of their right hand, each of which was assigned to 1 of 3 keys. A left key press indicated left turns or ‘just-straight’ letters, whereas a right one was used for right turns or letters including curved lines. A middle button was associated with PM target detection.

In addition, participants performed a simple reaction time task. The target was a row of 5 white ‘X’ (XXXXX), which were oriented alternately vertically or horizontally. They remained on the screen until a response was made and were replaced by a black screen until onset of the next target after a random ISI of 300–700 ms. Participants indicated target detection by pressing the left button with their index finger. A block lasted for 20 s. This task served as a baseline to allow comparison of contrasts across fMRI runs (see below).

The experiment was divided in 4 sessions, one for each combination of PM condition (OG, PM) and task (shape, alphabet). The OG session of a particular task was directly followed by its PM session. A session consisted of 6 blocks of the task, each succeeded by a block of the simple reaction time task. Thus, including presentations of short reminders of the instructions before each block (10 s for the experimental, 2 s for the reaction time task), a session lasted for approximately 12 min. Participants were allowed to rest between sessions. Order of tasks and assignment of color to stimulus condition in the alphabet task were counterbalanced. Before entering the scanner, participants were familiarized with the HF shape and practiced both OG tasks.

fMRI Recordings

A 3-T Siemens Allegra MRI scanner was used to acquire *T*1-weighted echoplanar images (64 × 64, 3 × 3 mm pixels; time echo: 30 ms; time repetition: 2.54 s) sensitive to BOLD contrast. The whole brain was covered by volumes that comprised 36 oblique axial slices (2 mm thick, separated by 1.7 mm) oriented at approximately 10° to the AC-PC plane to diminish the susceptibility artifact from the sinuses. For each of the 4 sessions, a separate functional scan of 305 volumes was acquired, of which the first 6 volumes were discarded to allow for *T*1 equilibration effects. Stimuli were projected on a mirror in direct view of the participant.

Data Analysis

Behavioral Analysis

Response times (RTs) for correct responses and accuracy of the ongoing trials (i.e., all trials of the OG condition and those of PM blocks that were not associated with a PM response) were analyzed separately for stay, switch, and nonswitch trials. The initial trial of an experimental block was excluded from analysis.
Due to the continuous nature of the shape task and the lack of an external marker for the subjects’ position on the shape, accuracy of single responses cannot be assessed. Thus, for this task, variance was estimated by examining successive overlapping sequences of 4 responses. It was then coded if a given sequence made up a valid response sequence given the shape (for details, see Gilbert et al. 2005 and Supplementary Material). Similarly, a PM response was taken for a hit, if it was embedded in a valid sequence of 5 responses (see Supplementary Material).

fMRI Analysis

fMRI data were analyzed using SPM5 (http://www.fil.ion.ucl.ac.uk/spm/software/spm5/). The volumes were first realigned, corrected for individual slice acquisition times, and smoothed with an isotropic 8-mm full-width at half-maximum Gaussian kernel.

The 4 sessions were treated as separate time series, and variance in BOLD signal was decomposed in a general linear model (GLM) (Friston et al. 1995). Regressors coded for sustained activation in the 4 main conditions of interest (OG_SO, OG_SI, PM_SO, and PM_SI) and the baseline condition (i.e., the RT task). Transient activation associated with switch and stay events was modeled by delta functions (coding for stimulus onsets). Additional regressors modeled the first miniblock of each epoch, the respective instruction periods for the baseline and the task blocks, and PM responses in either stimulus phase. These regressors were all convolved with a canonical hemodynamic response function (HRF) and comprised the full model for each session, in addition to regressors representing residual movement artifacts and the mean over scans. A 1/128-Hz high-pass filter was applied to the data and the model.

Parameter estimates for each regressor were calculated from the least-mean-squares fit of the model to the data. Effects of interest were assessed in a random effects analysis as follows: Eight contrasts were performed, each individually assessing the variance explained by a regressor representing 1 of the 4 main conditions of interest in the 2 tasks (i.e., shape OG_SO, alphabet OG_SO, etc.). These were then relative to the baseline condition of the relevant functional run. Assuming that cognitive processes associated with the RT task and their neural correlates are invariant over time, this allows comparison of conditions scanned in different functional runs (Simons et al. 2006; see also Supplementary Material). The 8 contrasts of each subject were entered into a repeated-measures analysis of variance (ANOVA) using nonparametric correction (Friston et al. 2002). Appropriate contrasts for effects of interest were conducted at this second level. Here, the data were analyzed averaged across tasks (e.g., Burgess et al. 2001, 2003; Gilbert et al. 2007). Hence, reported activations are unlikely to result from task-specific effects. A further second-level analysis was performed to assess transient BOLD signal changes associated with changes in stimulus condition (i.e., switch vs. stay trials). This analysis included contrast estimates for switch and stay events as a function of stimulus and PM condition. Contrasts were averaged across tasks and also corrected for session differences by the RT baseline task. Finally, to analyze brain-behavior relationships in the RT task, a separate GLM was estimated with 3 additional regressors created by a series of delta functions coding for switch and stay trials on their own. Following GLM analysis, a random effects analysis as described above was performed, each individually assessing the variance explained by the interaction of the stimulus and PM condition. All contrasts were corrected for false discovery rate (FDR) and family-wise error (FWE) at the cluster level using a software package (http://www.nitrc.org/projects/fdrcorrection/).

Results

Behavioral Results

Nonswitch Trials

Inspection of the nonswitch trials indicated slowed responses for SI compared with SO phases and for PM compared with OG blocks (Fig. 2). Moreover, the shape task was associated with greater RT than the alphabet task. This pattern was reliable, as confirmed by an ANOVA with the factors task (shape, alphabet), PM condition (OG, PM), and stimulus phase (SO, SI), which revealed all main effects (task: $F_{1,15} = 12.29, P < 0.006$; PM: $F_{1,15} = 46.22, P < 0.001$; stimulus: $F_{1,15} = 47.92, P < 0.001$). In addition, all interactions including the task factor were significant (task $\times$ PM: $F_{1,15} = 19.85, P < 0.001$, task $\times$ stimulus: $F_{1,15} = 21.74, P < 0.001$; task $\times$ PM $\times$ stimulus: $F_{1,15} = 46.22, P < 0.005$). However, both main effects (i.e., PM condition and stimulus phase) were significant for both tasks (see Supplementary Material).

Accuracy was generally high for both tasks (>84%). While it was greater for PM than OG blocks of the alphabet task, the reverse pattern was associated with the shape task. This was statistically confirmed by an ANOVA with the factors task, PM, and stimulus phase. The analysis revealed a main effect of task ($F_{1,15} = 20.02, P < 0.001$), indexing higher accuracy for the alphabet task as well as interactions between task and PM ($F_{1,15} = 13.63, P < 0.005$) and PM and stimulus phase ($F_{1,15} = 4.71, P < 0.05$). The former interaction reflected a crossover pattern, as corroborated by follow-up analyses establishing significant PM condition effects for both shape ($F_{1,15} = 8.54, P < 0.05$) and alphabet task ($F_{1,15} = 7.68, P < 0.05$). Note, however, that performance estimates for the shape task were more conservative for PM than OG data (Supplementary Material). The latter interaction between PM condition and stimulus phase resulted from a significant effect of the stimulus-phase factor (i.e., lower accuracy for SI than SO) for PM blocks only ($F_{1,15} = 6.37, P < 0.05$).

Switch versus Stay Trials

For each task a separate ANOVA was performed on RT with the factors trial type (switch, stay), PM condition, and stimulus phase (Fig. 2). For the alphabet task, all main effects were significant (i.e., slower responses for switch trials, the PM condition, and SI phases), and so were all interactions (all $F > 9.17, P < 0.01$). The interactions indicate that the trial type effect varied with PM condition and stimulus phase. Bonferroni-corrected comparisons of stay and switch trials were significant for all combinations of these factors (all $P < 2.97$; all $P < 0.005$). Except for OG_SO, switch trials were always associated with slower responses. For the shape task, the interactions between trial type and PM ($F_{1,15} = 5.03, P < 0.05$) and trial type and stimulus phase ($F_{1,15} = 11.92, P < 0.005$) were significant. Follow-up analyses revealed a trial type effect for SO phases ($F_{1,15} = 9.02, P < 0.01$) and a trend for PM blocks ($F_{1,15} = 4.34, P = 0.06$) (i.e., switch slower than stay trials in both cases). Thus, whereas RT were generally influenced by trial type in the alphabet task, this was only the case for SO phases of the shape task.

Analysis of the accuracy data revealed main effects of PM condition ($F_{1,15} = 4.83, P < 0.05$) and stimulus phase ($F_{1,15} = 5.85, P < 0.05$) for the alphabet task, indexing higher performance in OG blocks and SO phases. The shape task was associated with a main effect of PM condition ($F_{1,15} = 8.93, P < 0.01$) and an
interaction between PM condition, stimulus phase, and trial type ($F_{1,15} = 6.47, P < 0.05$). However, Bonferroni-corrected paired samples t-tests did not reveal a trial type effect on accuracy for any combination of stimulus phase and PM condition (all $|t| < 1.9$, all $P > 0.08$).

PM Targets
An ANOVA of RT data with the factors task and stimulus phase revealed significant main effects (task: $F_{1,15} = 10.23, P < 0.01$; stimulus: $F_{1,15} = 7.48, P < 0.05$), reflecting slower responses for the shape task and the SI phases, respectively (Table 1).

Analysis of hit rates yielded an effect of stimulus condition ($F_{1,15} = 11.56, P < 0.005$), indicating a greater hit rate for SO phases (Table 1). Thus, participants were faster and more reliable in detecting PM targets during SO phases.

fMRI Results
Sustained Engagement of mrPFC: Functional Overlap
Significant BOLD signal changes for the contrasts of PM condition (OG vs. PM) and stimulus phase (SO vs. SI) are summarized in Table 2. Contrasting OG with PM blocks (OG > PM) revealed increased BOLD signal within mrPFC, including aspects of the medial frontal gyrus, extending caudally into the cingulate gyrus, and more rostral parts of the superior frontal gyrus (Fig. 3a). SO compared with SI phases (SO > SI) elicited more widespread activation, covering bilateral occipital cortex, right superior parietal lobe, temporal lobe, and precuneus as well as right parahippocampal gyrus and left dorsal striatum. Importantly, this comparison was also associated with mrPFC activation (Fig. 3b).

Particularly, the areas associated with this contrast appeared to overlap with those observed for OG > PM. To formally test for this congruency, the contrast OG > PM was inclusively masked with SO > SI (thresholded at $P < 0.001$, uncorrected). Two clusters were identified in this analysis, one of which reflected overlap in an inferior caudal region (peak voxel: $x = 4, y = 42, z = -4$) and the other one indexing common recruitment of a more superior rostral area ($x = -12, y = 62, z = 12$). However, analysis of the interaction between PM condition and stimulus phase revealed no significant effect, suggesting that these 2 factors had additive effects on mrPFC BOLD signal. This
was further assessed by analyzing baseline-corrected estimates of the 4 regressor coefficients (i.e., OG_SO, OG_SI, PM_SO, and PM_SI) that were extracted from the peak voxel of the more caudal cluster (Fig. 3c). (Virtually identical results were obtained for the other cluster.) Consistent with the individual contrasts, the ANOVA revealed main effects of PM condition ($F_{1,15} = 18.07, P < 0.001$), reflecting greater recruitment for OG blocks and SO phases. The interaction was not significant ($F_{1,15} < 1, P > 0.5$), also indicating that the effects of stimulus and PM condition were additive. This pattern also emerged for analyzing the data separately for each task (Supplementary Material).

**Sustained Engagement of mrPFC: Relation to Performance**

One might suggest that activation of this mrPFC region merely reflected task-unrelated processes during the easier conditions (i.e., SO and OG) that are less likely to occur during the more difficult conditions (i.e., SI and PM). Therefore, we examined if within-subject signal changes as a function of PM condition (OG > PM) or stimulus condition (SO > SI) can be accounted for by associated differences in performance (as indices of “task difficulty”). Neither RT nor accuracy differences were significantly correlated with the parameter estimates ($-0.25 < r < 0.08$; all $P > 0.36$). Furthermore, signal changes within the peak voxel during the RT task were analyzed. This task primarily requires SO processing, that is, attending toward the externally presented stimuli. If mrPFC supports task-related processes during such low-demand situations, we expected to observe greater activation on faster trials. On a trial-by-trial basis, RTIs were negatively associated with BOLD signal, that is, greater activation was associated with better performance ($z = 1.87, P < 0.05$). (Note that RT and response–stimulus interval were uncorrelated [Fisher’s $z$: $-0.007 \pm 0.008$ (mean $\pm$ standard error of mean); $t_{15} = -0.82, P > 0.4$], implying that participants did not “rest” longer on trials associated with fast responses.) Thus, BOLD signal in mrPFC did not seem to merely decrease as a function of task difficulty (i.e., when task-unrelated processes have to be suspended). Instead, this region seems to support task-related processes during low-demand situations such as the RT task that require SO processing.

**Sustained Engagement of mrPFC: Functional Segregation**

In addition to recruiting overlapping aspects of mrPFC, the contrasts SO > SI and OG > PM were also associated with unique activations within this region. Particularly, segregation along a dorsal-ventral axis was observed, where the contrast of PM conditions was associated with more superior activation (Fig. 4a). To formally test for systematic spatial differences, $y$- and $z$-coordinates of each contrasts’ peak voxels were extracted. This was done for each sagittal slice within medial BA 10 (defined as $-8 \leq x \leq 8; y \geq 40; -12 \leq z \leq 30$), separately for each task and each subject (Fig. 4b for a similar approach, see Gilbert et al. 2007). Consistent across tasks, the PM contrast was associated with more superior peak coordinates ($z = 8.4$ vs. $z = 3.65; F_{1,15} = 4.83, P < 0.05$), whereas no significant difference on the caudal-rostral axis was obtained.

**Sustained Engagement of lPFC**

The reverse contrast PM > OG was associated with activations of the left parietal lobe, bilateral precuneus, and posterior medial and middle frontal gyrus (Table 2), while no supra-threshold activation was observed for SI > SO (for activations at a lower threshold of $P < 0.001$, uncorrected, and at least 10 contiguous voxels, see Supplementary Material). Thus, neither contrast yielded activation of lPFC. Since this area had previously been implicated in PM, ROI analyses were conducted at reported peak voxels in the left ($x = -30, y = 64, z = -4$) and right ($x = 40, y = 52, z = 4$) hemispheres (Burgess et al. 2001). Specifically, SVC were applied for 3 mm spheres, to test for the main effect PM > OG as well as for interactions between PM condition and stimulus phase. Both PM > OG and the interaction contrast [(PM_SO > OG_SO) > (PM_SI > OG_SI)] revealed significant BOLD signal changes in the left ROI ($z = 1.96$ and $z = 1.88; x = -30, y = 62, z = -2$). The right ROI ($z = 2.8; x = 40, y = 52, z = 6$) was associated with the interaction only.

To further assess the nature of these effects, baseline-corrected estimates of the 4 conditions were extracted from the peak voxels revealed by the SVC analysis (Fig. 5). For the left hemisphere data, the effect of PM condition (PM > OG) was only significant for SO ($t_{15} = -4.1, P < 0.005$) but not SI phases ($t_{15} = -0.95, P > 0.36$). In contrast, right hemisphere activation was associated with greater activation for OG than PM blocks during SI phases ($t_{15} = 2.54, P < 0.05$) (for parameter estimates by task, see Supplementary Material).

Thus, whereas PM compared with OG blocks were associated with left rostrolateral activation during SO phases only, activation of the right ROI was actually greater for OG than PM blocks during SI periods. To assess the reliability of this lateralization effect, an ANOVA was computed with the factors ROI (left, right), PM condition, and stimulus phase. In addition to revealing an interaction of ROI and PM condition ($F_{1,15} = 6.73, P < 0.05$), the analysis also showed a trend for the interaction between PM condition and stimulus phase ($F_{1,15} = 4.48, P < 0.06$). The 3-way interaction, however, was not significant ($F_{1,15} < 0.01, P > 0.96$). Hence, only the effect of PM condition differed significantly between the ROIs.
Figure 4. (a) BOLD signal changes for the contrasts 1) ongoing only versus prospective memory (OG > PM) and 2) stimulus-oriented versus stimulus-independent (SO > SI), and 3) for the overlap between both contrasts (thresholded at $P < 0.001$, uncorrected; averaged across both tasks). (b) Mean $z$-coordinates of the peak voxels for the individual contrast within each sagittal plane of rostromedial prefrontal cortex. Error bars indicate standard error of means.

Table 3

Significant BOLD signal changes within BA 10 for the contrast of switch versus stay trials, averaged across PM conditions, stimulus phases, and tasks

| Region               | Side | MNI coordinates | $Z_{\text{max}}$ | Voxels |
|----------------------|------|-----------------|------------------|--------|
| Rostromedial PFC l   | l    | $-32$  68 12    | 4.08             | 3      |
| Rostrolateral PFC r  | r    | 36  62 12      | 4.04             | 4      |
| Rostrolateral PFC l  | l    | $-32$  60 12    | 4.46             | 67     |
| Rostrolateral PFC r  | r    | 22  56 6      | 4.19             | 13     |

Note: small-volume corrected for BA 10. l, left; r, right; MNI, Montreal Neurological Institute.

Transient Engagement of rPFC

Regions within BA 10 exhibiting BOLD signal changes for switch versus stay trials are listed in Table 3. Whereas no area was more strongly activated for stay compared with switch trials, the reverse contrast was associated with several foci. These were primarily located laterally in both hemispheres but also a medial cluster was identified (Fig. 6). Activation patterns within these regions were further examined by analyzing baseline-corrected estimates of the regressor coefficients for switch and stay trials of the 4 main conditions (Fig. 6). Site-specific ANOVAs with the factors PM condition, stimulus phase, and trial type (switch, stay) revealed greater BOLD signal for switch versus stay trials at both the left lateral ($x = -32$, $y = 60$, $z = 12$; $F_{1,15} = 14.72$, $P < 0.005$) and the medial site ($x = -6$, $y = 68$, $z = 14$; $F_{1,15} = 6.73$, $P < 0.05$). The right rostrolateral site, in contrast, yielded a main effect of trial type ($x = 22$, $y = 56$, $z = 6$; $F_{1,15} = 14.45$, $P < 0.005$) and the interaction between trial type and PM condition ($F_{1,15} = 5.61$, $P < 0.05$). Follow-up analyses revealed that the trial type effect was restricted to OG blocks ($F_{1,15} = 22.14$, $P < 0.001$). Thus, switch compared with nonswitch trials were associated with greater recruitment of left rPFC and mrPFC, whereas such an effect was only present for OG blocks in right rPFC. This pattern was largely corroborated by an ANOVA with the additional factor ROI (left, middle, right), which revealed a strong trend for the 4-way interaction ($F_{2,30} = 3.2$, $P < 0.06$) in addition to the trial type main effect ($F_{1,15} = 39.34$, $P < 0.001$).
required SO processing of the externally presented targets. This suggests that processes supported by this area are functionally relevant in such low-demand situations (Gilbert, Simons, et al. 2006). Moreover, activation differences between stimulus or PM conditions were not related to behavioral differences between these conditions. Thus, the observed pattern cannot readily be explained by simple differences in task difficulty.

**Discussion**

This study scrutinized the involvement of rPFC in PM. In particular, it tested the hypothesis that maintaining an intention activates lrPFC due to increased reliance on SI attending, while mrPFC deactivation reflects concurrent attenuation of SO processing of the OG task (Burgess et al. 2003; Simons et al. 2006).

**Medial rPFC**

**Default Mode Suspension?**

Consistently across 2 different tasks, overlapping parts of mrPFC were more strongly recruited during both 1) sole engagement in the OG task and 2) SO processing. This congruency suggests that mrPFC activation for OG compared with PM blocks might indeed reflect relative differences in SO versus SI attending. However, SO phases and OG blocks were associated with faster responses than SI phases and PM blocks, indicating that the former conditions were easier. It has been proposed that increased activity during low-demand tasks actually reflects a greater degree of task-unrelated thought (i.e., “mind-wandering”; Mason et al. 2007). Accordingly, less activation for the more demanding tasks may reflect suspension of such proposed “default activation” (e.g., Gusnard and Raichle 2001). However, in these data, there is little support for this explanation: mrPFC BOLD signal was negatively associated with RT during the simple RT task. Thus, greater activation was associated with better performance in a situation that primarily requires SO processing of the externally presented targets. This suggests that processes supported by this area are functionally relevant in such low-demand situations (Gilbert, Simons, et al. 2006). Moreover, activation differences between stimulus or PM conditions were not related to behavioral differences between these conditions. Thus, the observed pattern cannot readily be explained by simple differences in task difficulty.

**Functional Overlap**

If the observed pattern of additive effects on BOLD signal does not simply reflect degrees of “default network” suspension, does it further elucidate rPFC functioning? Both stimulus phases in the current study involved some degree of engagement with the environment (e.g., by pressing buttons) and attending to internally represented information (e.g., retrieval of task instructions). The stimulus phases thus vary in relative rather than absolute terms in required attention toward the external world, that is, the SO condition is not a benchmark of pure SO attending. OG compared with PM performance could well be associated with separable and hence additive aspects of SO processing (see next section). This would imply that mrPFC does not categorically bias attention toward SO processing. Instead, the data indicate that increasing the relative demand on SO processing increases the engagement of this region. Thus, the SO condition of the OG task might be considered a dual demand situation, in which both task requirements (i.e., SO vs. SI and OG only vs. PM) involve SO attending.

Such an interpretation can also account for the observation of greater mrPFC activation for OG than PM blocks during SI phases. During these phases, neither PM condition (i.e., OG and PM) is likely to require greater SO processing (both involve interactions with the external world primarily via button presses). However, PM blocks additionally require carrying out delayed intentions. Slowed responses for this condition indicate that intention execution was not directly prompted by the environment (Einstein et al. 2005). Thus, processes subserving the PM demand were SI, which prolonged the time participants spent disengaged from the external world. Hence, mrPFC activation might vary with the relative amount of time devoted to SO processing, where different aspects of attending to the external world independently recruit overlapping neuronal populations. This view is consistent with the proposal that medial PFC mediates SO and SI attending (i.e., “surveillance of the internal and external environments,” Gusnard et al. 2001, p. 4259).

**Functional Specialization along a Dorsal–Ventral Gradient?**

The observed spatial segregation within mrPFC supports the idea that the contrasts of PM and stimulus conditions reflect distinct aspects of SO attending. Although there was considerable overlap within this brain region, the peak activation for the PM manipulation was more superior than the peak for the stimulus effect. Since this effect was unexpected, any interpretation of its functional significance is merely provisional. However, some evidence suggests that dorsal versus ventral regions within medial PFC differ in their functional properties and relative connectivity patterns. For example, a recent study associated activation in more ventral parts with the imagination of future episodes per se, whereas activation in a more dorsal subregion could be linked to the influence of imagination on subsequent decisions (Benoit et al. 2011). Regarding the anatomical connections, it has been proposed...
that a gradient of connectivity runs along the genu of the corpus callosum (Amodio and Frith 2006). In the rhesus monkey, most superior aspects (BA 9) have strong connections with motor control regions (i.e., lateral premotor cortex, supplementary motor cortex, and cingulate motor area) but, at most, few connections with the rhinal cortex. In contrast, the latter brain region is highly connected with the most inferior and caudal medial PFC regions (BAs 25, 24, and 32; Barbas et al. 1999). Additionally, the inferiorly adjacent orbitofrontal cortex is primarily connected with sensory association areas (Ongur and Price 2002).

Thus, dorsal rPFC might primarily be connected with areas involved in the control of actions. Accordingly, the superior peak for the PM contrast may reflect differences in SO versus SI control of actions, that is, whether to perform the OG activity that is triggered by the stimulus or the internally represented intention that is less strongly prompted. Consistent with this idea, dorsal mrPFC has recently been found to be more strongly engaged when PM intentions were more directly cued by the environment (Gilbert et al. 2009). This hypothesis could be tested by contrasting PM conditions that vary in association strength between PM target and intention (cf. McDaniel et al. 2004). If, in contrast, ventral parts of rPFC are more strongly linked to sensory association cortices, they might be involved in mediating competition between perceived and imagined stimuli, which are processed in partly the same perceptual areas (Kosslyn et al. 2001).

Taken together, mrPFC is jointly associated with 1) OG task activity compared with additional maintenance of delayed intentions and 2) SO compared with SI processing. Thus, reduced recruitment of this region during PM performance might indeed reflect attenuation of SO attending. Concurrently, however, spatial segregation of peak activation suggests a functional gradient along a dorsal–ventral axis, which might reflect the regulation of different aspects of SO versus SI attending. This study thus contributes to the increasing knowledge about functional variations within mrPFC (Gilbert, Spengler, et al. 2006; Krueger et al. 2007; Gilbert et al. 2010; Volle et al. 2010).

**Lateral rPFC**

**Sustained Engagement of Left lrPFC for Maintaining PM Intentions**

Consistent with previous evidence (Okuda et al. 1998, 2007; Burgess et al. 2001, 2003; den Ouden et al. 2005; Simons et al. 2006; Gilbert et al. 2009; Reynolds et al. 2009), BOLD signal in the left lrPFC ROI was greater when participants performed a PM task in addition to OG activity. Burgess et al. (2001) demonstrated recruitment of this region when participants were prepared to carry out an intention in the absence of any actual PM targets (see also Simons et al. 2006; Reynolds et al. 2009). Thus, lrPFC appears to support the maintenance of an intention in the context of OG activity rather than actual target detection or task execution.

However, what processes supported by left lrPFC might be engaged while participants maintain a delayed intention? This region has been implicated in the adaptation to upcoming task demands (Sakai and Passingham 2003; Rowe et al. 2007). It thus might be involved in the preparation for intention execution. Alternatively, this region might subserve the interposition of the PM task in the OG activity (cf. Koechlin et al. 1999; Braver and Bongirolli 2002). Consistently, lesions of left lrPFC have been associated with frequent rule breaks during such multitasking (Burgess et al. 2000). Both accounts, however, associate this region with processes that are not directly contingent on externally presented stimuli (i.e., task preparation precedes PM target onset; task coordination is not externally guided). Therefore, left lrPFC activation can be characterized as being associated with SI attending (cf. Burgess et al. 2003, 2007).

If this region is recruited to bias attention toward internally maintained information (i.e., the PM intention), one might expect an interaction between PM condition and stimulus phase. Specifically, the difference between PM and OG blocks might be smaller for SI than SO phases, since the system would already be in a relative mode of SI processing in the OG condition of SI phases. Consequently, there would be less need to bias attention toward this processing mode for the PM task. Thus, one may expect an underadditive effect of PM and stimulus condition. In contrast, accounts that implicate lrPFC in the integration of the outcomes of 2 or more cognitive operations would predict the opposite pattern, that is, a “superadditive” effect of the 2 factors (e.g., Ramnani and Owen 2004). The data yielded an underadditive pattern: The effect of PM condition was significant for SO phases only.

However, a more caudally located region in the right hemisphere exhibited an unexpected activation profile, that is, BOLD signal was greater for OG than PM blocks during SI phases. The functions supported by this subregion need to be further elucidated in future studies.

**Transient Engagement of lrPFC for Shifting the Attentional Focus**

In addition to supporting SI versus SO processing (Dumontheil, Gilbert, et al. 2010; Dumontheil, Hassan, et al. 2010), lrPFC also seems to be involved in shifting between both modes more generally (Gilbert et al. 2005; Dumontheil, Gilbert, et al. 2010). This was the case for the present data, where subregions of both mrPFC and left lrPFC were invariably associated with greater BOLD signal for switch than for stay trials. The recruitment of lrPFC during switch events may accordingly reflect changes of task-relevant stimuli (i.e., externally presented vs. internally generated) (see also Pollmann et al. 2000; Braver et al. 2003).

In contrast, for right lrPFC, the trial type effect was restricted to OG blocks. If SO phases of PM blocks already require more SI processing than SO phases of OG blocks, there would be less need to shift between the 2 attentional modes in PM blocks. Hence, right lrPFC seems to be primarily engaged when the alternative processing mode needs to be strongly imposed (cf. Gilbert et al. 2005; Burgess et al. 2007).

**Summary and Conclusions**

Overlapping parts of mrPFC exhibited BOLD signal increases both during 1) mere OG task activity compared with additional PM performance and 2) SO compared with SI attending. This pattern supports the hypothesis that some of the rPFC activations associated with prospective memory reflect the demands that PM tasks make for the control of stimulus-oriented versus -independent attending (Burgess et al. 2009). Thereby, this study corroborates a major prediction derived from the gateway hypothesis of rPFC function (Burgess et al. 2007). At the same time, the PM contrast was consistently associated with more dorsal peak activation than the stimulus contrast, implying additional engagement of distinct processes.
This finding suggests that mrPFC might be functionally fractionated along a dorsal-ventral gradient. However, the nature of this putative gradient needs to be systematically examined in future studies. Moreover, left IrPFC recruitment for PM compared with OG blocks may reflect processes involved in maintaining and/or implementing delayed intentions in the context of distracting OG activity. The observed underadditive interaction between PM and stimulus condition, however, argues against accounts that implicate rPFC in the integration of the outcomes of multiple cognitive operations (e.g., Rammani and Owen 2004).

Supplementary Material

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/

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Notes

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References

Amodio DM, Frith CD. 2006. Meeting of minds: the medial frontal cortex and social cognition. Nat Rev Neurosci. 7:268–277.

Barbas H, Ghoshgai H, Dombrowski SM, Rempel-Clower NL. 1999. Medial prefrontal cortices are unified by common connections with superior temporal cortices and distinguished by input from memory-related areas in the rhesus monkey. J Comp Neurol. 410:343–367.

Benoit RG. 2008. The role of rostral prefrontal cortex in establishing cognitive sets preparation or coordination? J Neurosci. 28:3259–3261.

Benoit RG, Gilbert SJ, Burgess PW. 2011. A neural mechanism mediating the impact of episodic projection on farsighted decisions. J Neurosci. 31:6771–6779.

Benoit RG, Gilbert SJ, Volle E, Burgess PW. 2010. When I think about me and simulate you: medial rostral prefrontal cortex involvement in self-referential processes. Neuroimage. 50:1340–1349.

Braver TS, Bongiollati SR. 2002. The role of frontopolar cortex in subgoal processing during working memory. Neuroimage. 15:523–536.

Braver TS, Reynolds JR, Donaldson DL. 2003. Neural mechanisms of transient and sustained cognitive control during task switching. Neuron. 39:713–726.

Burgess PW, Alderman N, Volle E, Benoit RG, Gilbert SJ. 2009. Mesulam’s fronto-ial lobe mystery re-examined. Restor Neurol Neurosci. 27:493–506.

Burgess PW, Dumontheil I, Gilbert SJ. 2007. The gateway hypothesis of rostral prefrontal cortex (area 10) function. Trends Cogn Sci. 11:290–298.

Burgess PW, Dumontheil I, Gilbert SJ, Okuda J, Schölvinck ML, Simons JS. 2008. On the role of rostral prefrontal cortex (area 10) in prospective memory. In: Kliegl M, McDaniel MA, Einstein GO, editors. Prospective memory: cognitive, neuroscience, developmental, and applied perspectives. Mahwah (NJ): Erlbaum. p. 235–260.

Burgess PW, Quayle A, Frith CD. 2001. Brain regions involved in prospective memory as determined by positron emission tomography. Neuropsychologia. 39:545–555.

Burgess PW, Scott SK, Frith CD. 2003. The role of the rostral frontal cortex (area 10) in prospective memory: a lateral versus medial dissociation. Neuropsychologia. 41:439–453.

Burgess PW, Veitch E, Costello A, Shallice T. 2000. The cognitive and neuroanatomical correlates of multitasking. Neuropsychologia. 38:848–863.

Christoff K, Gabrieli JDE. 2000. The frontopolar cortex and human cognition: evidence for a rostrocaudal hierarchical organization within the human prefrontal cortex. Psychobiology. 28:168–186.

den Ouden HEM, Frith U, Frith C, Blakemore S-J. 2005. Thinking about intentions. Neuroimage. 28:787–796.

Dumontheil I, Gilbert SJ, Frith CD, Burgess PW. 2010. Recruitment of lateral rostral prefrontal cortex in spontaneous and task-related thoughts. Q J Exp Psychol. 63:1740–1756.

Dumontheil I, Hassan B, Gilbert SJ, Blakemore S-J. 2010. Development of the selection and manipulation of self-generated thoughts in adolescence. J Neurosci. 30:7664–7671.

Einstein GO, McDaniel MA, Thomas R, Mayfield S, Shank H, Morrisette N, Breneriser J. 2005. Multiple processes in prospective memory retrieval: factors determining monitoring versus spontaneous retrieval. J Exp Psychol Gen. 134:327–342.

Ellis J. 1996. Prospective memory or the realization of delayed intentions: a conceptual framework for research. In: Brandimonte M, Einstein GO, McDaniel MA, editors. Prospective memory: theory and applications. Mahwah (NJ): Lawrence Erlbaum Associates. p. 1–22.

Friston KJ, Glaser DE, Henson RNA, Kiebel S, Phillips C, Ashburner J. 2002. Classical and Bayesian inference in neuroimaging: applications. Neuroimage. 16:481–512.

Friston KJ, Holmes AP, Worsley KJ, Poline J-P, Frith CD, Frackowiak RSJ. 1995. Statistical parametric maps in functional imaging: a general linear approach. Hum Brain Mapp. 2:189–210.

Gilbert SJ. 2011. Decoding the content of delayed intentions. J Neurosci. 31:2888–2894.

Gilbert SJ, Frith CD, Burgess PW. 2005. Involvement of rostral prefrontal cortex in selection between stimulus-oriented and stimulus-independent thought. Eur J Neurosci. 21:1423–1431.

Gilbert SJ, Gollwitzer P, Cohen A, Oettingen G, Burgess PW. 2009. Separable brain systems supporting cue versus self-initiated re-alization of delayed intentions. J Exp Psychol Learn Mem Cogn. 35:905–915.

Gilbert SJ, Henson RNA, Simons JS. 2010. The scale of functional specialization within human prefrontal cortex. J Neurosci. 30:1233–1237.

Gilbert SJ, Simons JS, Frith CD, Burgess PW. 2006. Performance-related activity in medial rostral prefrontal cortex (area 10) during low-demand tasks. J Exp Psychol Hum Percept Perform. 32:45–58.

Gilbert SJ, Spengler S, Simons JS, Douglas Steele J, Lawrie SM, Frith CD, Burgess PW. 2006. Functional specialization within rostral prefrontal cortex (area 10): a meta-analysis. J Cogn Neurosci. 18:932–948.

Gilbert SJ, Williamson IDM, Dumontheil I, Simons JS, Frith CD, Burgess PW. 2007. Distinct regions of medial rostral prefrontal cortex supporting social and nonsocial functions. Soc Cogn Affect Neurosci. 2:217–226.

Gusnard DA, Akbodak E, Shulman GI, Raichle ME. 2001. Medial prefrontal cortex and self-referential mental activity: relation to a default mode of brain function. Proc Natl Acad Sci U S A. 98:4259–4264.

Gusnard DA, Raichle ME. 2001. Searching for a baseline: functional imaging and the resting human brain. Nat Rev Neurosci. 2:685–694.

Henseler I, Krüger S, Dechent P, Gruber O. 2011. A gateway system in rostral PFC? Evidence from biasing attention to perceptual in-formation and internal representations. Neuroimage. 56:1666–1676.

Koechlin E, Basso G, Pietrini P, Panzer S, Graffman J. 1999. The role of the anterior prefrontal cortex in human cognition. Nature. 399:148–151.

Kosslyn SM, Ganis G, Thompson WL. 2001. Neural foundations of imagery. Nat Rev Neurosci. 2:635–642.

Krueger F, Moll J, Zahn R, Heincke A, Graffman J. 2007. Event frequency modulates the processing of daily life activities in human medial prefrontal cortex. Cereb Cortex. 17:2346–2353.

Luce RD. 1986. Response times: their role in inferring elementary processes. Cambridge (MA): MIT Press.

Mason MF, Norton MI, Van Horn JD, Wegner DM, Grafton ST, Macrae CN. 2007. Wandering minds: the default network and stimulus-independent thought. Science. 315:393–395.

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McDaniel MA, Einstein GO. 2007. Prospective memory: an overview and synthesis of an emerging field. Thousand Oaks (CA): Sage Publications.

McDaniel MA, Einstein GO, Guynn MJ, Breneiser J. 2004. Cue-focused and reflexive-associated processes in prospective memory retrieval. J Exp Psychol Learn Mem Cogn. 30:605-614.

Okuda J, Fujii T, Ohtake H, Tsukiura T, Yamadori A, Frith CD, Burgess PW. 2007. Differential involvement of regions of rostral prefrontal cortex (Brodmann area 10) in time- and event-based prospective memory. Int J Psychophysiol. 64:233-246.

Okuda J, Fujii T, Yamadori A, Kawashima R, Tsukiura T, Fukatsu R, Suzuki K, Ito M, Fukuda H. 1998. Participation of the prefrontal cortices in prospective memory: evidence from a PET study in humans. Neurosci Lett. 253:127-130.

Öngür D, Price JL. 2002. The organization of networks within the orbital and medial prefrontal cortex of rats, monkeys and humans. Cereb Cortex. 10:206-219.

Pollmann S, Weidner R, Müller HJ, von Cramon DY. 2000. A fronto-posterior network involved in visual dimension changes. J Cogn Neurosci. 12:480-494.

Ramnani N, Owen AM. 2004. Anterior prefrontal cortex: insights into function from anatomy and neuroimaging. Nat Rev Neurosci. 5:181-194.

Reynolds JR, West R, Braver T. 2009. Distinct neural circuits support transient and sustained processes in prospective memory and working memory. Cereb Cortex. 19:1208-1221.

Roca M, Parr A, Thompson R, Woolgar A, Torralva T, Antoun N, Manes F, Duncan J. 2010. Executive function and fluid intelligence after frontal lobe lesions. Brain. 133:234-247.

Rorden C, Brett M. 2000. Stereotaxic display of brain lesions. Behav Neurol. 12:191-200.

Rowe JB, Sakai K, Lund TE, Ramsoy T, Christensen MS, Baare WFC, Paulson OB, Passingham RE. 2007. Is the prefrontal cortex necessary for establishing cognitive sets? J Neurosci. 27:13303-13310.

Sakai K, Passingham RE. 2003. Prefrontal interactions reflect future task operations. Nat Neurosci. 6:75-81.

Simons JS, Schölvinck M, Gilbert SJ, Frith CD, Burgess PW. 2006. Differential components of prospective memory? Evidence from fMRI. Neuropsychologia. 44:1388-1397.

Uretzky S, Gilboa A. 2010. Knowing your lines but missing your cue: rostral prefrontal lesions impair prospective memory cue detection, but not action-intention superiority. J Cogn Neurosci. 22:2745-2757.

Volle E, Gilbert SJ, Benoit RG, Burgess PW. 2010. Specialization of the rostral prefrontal cortex for distinct analogy processes. Cereb Cortex. 20:2647-2659.

Volle E, Gonen-Yaacovi G, de Lacy Costello A, Gilbert SJ, Burgess PW. 2011. The role of rostral prefrontal cortex in prospective memory: a voxel-based lesion study. Neuropsychologia. 49:2185-2198.