Covert Shifts of Spatial Attention in the Macaque Monkey

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In the awake state, shifts of spatial attention alternate with periods of sustained attention at a fixed location or object. Human fMRI experiments revealed the critical role of the superior parietal lobule (SPL) in shifting spatial attention, a finding not predicted by human lesion studies and monkey electrophysiology. To investigate whether a potential homolog of the human SPL shifting region exists in monkeys (Macaca mulatta), we adopted an event-related fMRI paradigm that closely resembled a human experiment (Molenberghs et al., 2007). In this paradigm, a pair of relevant and irrelevant shapes was continuously presented on the horizontal meridian. Subjects had to covertly detect a dimming of the relevant shape while ignoring the irrelevant dimmings. The events of interest consisted of the replacement of one stimulus pair by the next. During shift but not stay events, the relevant shape of the new pair appeared at the contralateral position relative to the previous one. Spatial shifting events activated parietal areas V6/V6A and medial intraparietal area, caudo-dorsal visual areas, the most posterior portion of the superior temporal sulcus, and several smaller frontal areas. These areas were not activated during passive stimulation with the same sensory stimuli. During stay events, strong direction-sensitive attention signals were observed in a distributed set of contralateral visual, temporal, parietal, and lateral prefrontal areas, the vast majority overlapping with the sensory stimulus representation. We suggest medial intraparietal area and V6/V6A as functional counterparts of human SPL because they contained the most widespread shift signals in the absence of contralateral stay activity, resembling the functional characteristics of the human SPL shifting area.

Key words: fMRI; human; monkey; shifting; spatial attention; superior parietal lobe

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ACC showed early onset single-cell activity in the context of attention shifts (Kaping et al., 2011). All aforementioned monkey areas, however, differ substantially in location or function from human SPL. Thus, it remains an unanswered question whether monkey cortex contains any region functionally homologous to human SPL as previously characterized by fMRI (Vandenberghhe et al., 2001; Yantis et al., 2002).

Our goal was to identify monkey brain regions involved in covert spatial attention shifts. Also, we sought to identify regions involved in sustained contralateral attention and to map the spatial relationship between these two attentional signals and the sensory stimulus representation. We trained monkeys to covertly shift attention whenever a relevant shape changed location. This task was similar to the human fMRI task previously used to isolate effects of transient spatial attention shifts against a sustained-attention baseline (Molenberghs et al., 2007). Cue events probing the monkey’s allocation of attention occurred independently of the events of interest (i.e., shift and stay events when one stimulus pair was replaced by the next). The cue events consisted of a dimming of relevant shapes while dimming of irrelevant shapes had to be ignored. Hence, we could dissociate attentional shifting operations from stimulus changes, sustained attention effects and processes engaged during target detection, response preparation, and execution.

Materials and Methods

Subjects. Three rhesus monkeys (female M13, female M24 and male M35; Macaca mulatta; 3–6 kg; 3–9 years of age) participated in the experiments. Animal care and experimental procedures were performed in accordance with the National Institute of Health’s Guide for the Care and Use of Laboratory Animals, the European legislation (Directive 2010/63/EU), and were approved by the Ethical Committee at the KU Leuven. The details of the general fMRI procedures and training of monkeys have been described previously (Vanduffel et al., 2001; Nelissen et al., 2005).

Stimuli and tasks: spatial shift experiment. The experimental paradigm was adapted from a human fMRI experiment (Molenberghs et al., 2007). Two stimuli were placed on the horizontal meridian at 9.25 degrees eccentricity, in the left and right hemifields (Fig. 1, I). All stimuli were white on a black background and of equal area (1.59 degrees²). There were two possible coupled stimulus pairs: a square with a triangle and a circle with a diamond. Each pair contained a relevant and an irrelevant stimulus. To obtain a reward, the monkey had to respond by manually interrupting a light beam when the relevant (Fig. 1, IVa), but not when

Figure 1. Stimuli and task. Top (schematic of display viewed by the monkey), Animals fixated during all trial types, also when responding to dimming events. One stimulus pair (consisting of a relevant and irrelevant shape) was presented at any given time that the monkey fixated. Each pair was replaced, with no temporal gap, by the succeeding pair after 2250 ms. Top left, Dimming of the relevant shapes had to be indicated by a manual response to receive a juice reward. Top right, Dimming of the irrelevant shapes had to be ignored. Examples of different event types: I, shift right, a feature change cues the monkey to make a covert spatial-attention shift to the right hemifield (the left relevant stimulus [square] is replaced by an irrelevant one [diamond]); II, stay right, a feature change cues the animal to maintain its covert attention to the right because the relevant stimulus [square] of the next pair appears at the same position as the preceding one (circle); III, shift left; IVa, a relevantdimming, the monkey responds and is rewarded; IVb, irrelevant dimming, the monkey continues fixating, no reward is given; V, stay left; and VI, null left, same shapes remain at the same position for 2 × 2250 ms.
the irrelevant-stimulus dimmed (Fig. 1, IVb). Monkeys were intensively trained (>4 months) to attend only the relevant stimulus of each pair and to fixate upon the central fixation point during the entire task. Fixation-only trials were not rewarded. Relevant stimuli were identical for M13 and M35 (square and circle) and swapped for M24 (triangle and diamond). Monkeys thus covertly maintained their attentional focus at the location of the relevant stimulus. Compliance was probed by the occasional randomly inserted dimming events (the relevant and irrelevant shapes dimmed for 100 ms in 14% of the trials each). Luminance contrast (gray level) was adapted on a run-wise basis to approximate a performance level of 1.0 for all conditions. None of the interactions of Repeated-measures ANOVAs was significant; therefore, data are not shown.

The sustained attention baseline was transiently interrupted by two types of events. These consisted of a replacement (without temporal gaps) of one stimulus pair by the second stimulus pair, occupying the same positions. In the first event type (Fig. 1, I, III), the relevant stimulus of the new pair appeared at the position previously occupied by the irrelevant stimulus of the preceding pair, and vice versa. This feature change, in combination with a positional change of the relevant stimulus, elicited a shift in spatial attention (shift event). A “shift-right” event (Fig. 1, I) corresponded to a spatial shift of the relevant stimulus from the left to the right hemifield and a “shift-left” (Fig. 1, III) to a spatial shift from right to left.

In event Type 2 (Fig. 1, I, V), the relevant stimulus of the new pair appeared at the same position as the relevant stimulus of the preceding pair. Hence, the relevant stimulus site remained unchanged, inducing no spatial shift in attention. This was called a stay event: “stay-right” (Fig. 1, II, V), the relevant stimulus of the new pair appeared at the position previously occupied by the same positions. In the first event type (Fig. 1, I, III), the relevant stimulus of the new pair appeared at the position previously occupied by the irrelevant stimulus of the preceding pair, and vice versa. This feature change, in combination with a positional change of the relevant stimulus, elicited a shift in spatial attention (shift event). A “shift-right” event (Fig. 1, I) corresponded to a spatial shift of the relevant stimulus from the left to the right hemifield and a “shift-left” (Fig. 1, III) to a spatial shift from right to left.

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within the response box was checked using infrared light beams. This procedure minimized unwanted (hand) movements or saccadic eye movements toward the peripheral stimuli. In cases where the monkeys made a hand movement (except during the dimming of the relevant stimulus) or an eye movement outside the fixation window, the trial was aborted and the stimulus display disappeared.

Stimulus localization experiment. To characterize the cortical representation of the visual stimuli, monkeys were scanned with a fixation-only task after the main experiment. The same stimuli (square, triangle, diamond, and circle) were shown unilaterally to the left, unilaterally to the right, and bilaterally at the same positions (9.25 degree eccentricity) as in the main experiment. All possible stimulus combinations were displayed and replaced every 200 ms (without gaps), resulting in blocks of 20 s. These three stimulation blocks were interleaved with a fourth block of 20 s without visual stimuli, during which only the red fixation dot was present. The monkeys fixated within a 2 × 3 window while their hands remained in the response box to receive a reward. The juice-reward interval was systematically decreased from 2500 to 500 ms to encourage long, uninterrupted fixation sequences.

fMRI acquisition: procedure. Monkeys were trained in a mockup of an MRI scanner for all tasks. They were scanned following several months (~4) of training, when fixation performance exceeded 90%–95%, and when false-alarm rates were close to 0%. The monkey sat in a sphinx position, head fixed to the plastic monkey chair, directly facing the screen. Eye position was monitored at 120 Hz, using pupil position and corneal reflection (Iscan).

Scanning. Functional images were acquired with a 3.0 tesla horizontal bore full-body scanner (TIM Trio; Siemens Healthcare), using a gradient-echo T2*-weighted echo-planar imaging-sequence: 40 horizontal slices; TR, 2 s; TE, 17 ms (M13, M24), 19 ms (M35); 1.25 × 1.25 × 1.25 mm3 isotropic voxels, image acceleration factor of 3 (M13, M24), or 2 (M35). M13 was scanned with a custom-built, 8-channel, phased-array receive coil, and a saddle-shaped, radial transmit-only surface coil (Kolster et al., 2009). M24 had been implanted with an 8-channel receive array coil on top of the skull, beneath the headpost, improving the sensitivity for MR-imaging (Janssens et al., 2012). M35 had been implanted with a 5-channel receive array coil (due to additional recording wells, only 5 coils were constructed). Scanning of M24 was performed with the same scanner, main parameters, and transmit coil as M13, but also using a 36 cm inner-diameter head gradient set (AC88, maximum strength: 80 mT/m; maximum slew rate: 800 T/m/s). Slices were oriented transversally, covering the entire brain. High-resolution anatomical images were acquired for each monkey during a separate session under anesthesia, using a single radial transmit-receive surface coil and an MPRAGE sequence (TR = 2200 ms; TE = 4.05 ms; 208 slices; 0.4 mm isotropic voxel size).

Before each functional scanning session, a contrast agent, monocrystalline iron oxide nanoparticle (MION) (Sinerem; Laboratoire Guerbet), was injected into the femoral/saphenous vein (~6–11 mg/talline iron oxide nanoparticle (MION) (Sinerem; Laboratoire Guerbet, voxel size).

Nonrigid slice-by-slice distortion correction (Kolster et al., 2009). Subsequent data analysis was performed using SPM5 software package (Wellcome Department of Cognitive Neurology, London) running under MATLAB (MathWorks). The preprocessing steps included the following: (1) skull-stripping the images, (2) coregistration of the anatomical and mean functional images, and (3) spatial normalization of all images to the 112-PM atlas (McLaren et al., 2009) aligned to the F99 surface-based atlas (Van Essen et al., 2001), with a voxel size of 1 × 1 × 1 mm. Images were smoothed with a 1.5 mm kernel.

Data analysis of main experiment (event related). As illustrated in Figure 1 (and described above), the following events entered the GLM: stay events, composed of a feature change (Fig. 1, I: stay-right, Fig. 1, V: stay-left); shift events, composed of a feature change and a spatial shift (Fig. 1, I: shift-right, Fig. 1, III: shift-left); and null events (Fig. 1, VI: null left, no feature change). An equal number of null-left and null-right events constituted the attention baseline. The number of correctly executed trials analyzed for each event type was equalized for left- and right-sided attention. Only trial sequences with at least 3 or more consecutive and correctly executed fixation trials were included in the analysis. The dimming and reward events within these series of correctly executed fixation trials were excluded from the analyses. The GLM included 5 regressors for the five conditions, 2 eye movement regressors, and 6 additional head motion regressors (translation and rotation in 3D), per run. Each condition was modeled by convolving a γ function (\( \gamma = 0, \tau = 8 \), and exponent = 0.3), modeling the MION hemodynamic response function, at the onset of the condition (transition of stimulus displays). The first two main contrasts of interest were stay-left versus stay-right (contrast 1) and vice versa (contrast 2). These delineated areas reflecting the current locus of attention and will be deemed “contralateral stay,” relative to the observed hemisphere.
Activations correlating with “transient” spatial attention shifts regardless of the direction of the shift were visualized by contrasting bilateral “shifts” versus “stay” (shift-left and shift-right vs. stay-left and stay-right, contrast 3).

Data were first analyzed per subject and scanning day (session) with a fixed-effects GLM. Runs with a fixation performance <85% were excluded. For M13, this resulted in 9 sessions with a total of 131 runs (on average 15 runs/session); for M24, 12 sessions with a total of 181 runs (on average 15 runs/session); and for M35, 10 sessions with a total of 171 runs (on average 17 runs/session) that were included in the analyses. Next, we computed a session-wise second-level mixed-effect analysis (MFX) (Friston et al., 2005), with the “con-images” of contrasts 1–3, obtained from the first level analyses as input, including a total of 31 images per contrast: 9 (M13) + 12 (M24) + 10 (M35) sessions. All statistical maps were based on this MFX analysis. Contralateral stay maps for contrasts 1 and 2 were thresholded at p = 0.001, whereas shift maps (contrast 3) were shown at a threshold of p = 0.01, both uncorrected. The relatively low threshold for the shifting contrast was motivated by the fact that we had a clear a priori hypothesis about superior parietal involvement.

Second, to compute raw time courses for the group, we performed a group fixed-effect analysis (FFX). This was done by selecting the best 69 runs for each animal according to behavior (percentage fixation and response performance), warped to group space. These 207 runs (reaching the upper limit of MATLAB memory for NIFTI files: 207 runs × 350 volumes) were pooled for the FFX analysis, and the final contrasts (same as used in the MFX analysis) were thresholded at family-wise error (FWE, corrected for multiple comparisons at p = 0.05). This procedure was performed to extract time courses and t-values for only those clusters reaching the above-mentioned threshold in the statistically more stringent MFX analysis. A cluster extent threshold of 10 voxels was applied to all volume maps.

**Analysis of stimuli level data (block design).** The data from the stimulus localization experiment were preprocessed and analyzed the same way as the event-related data of the main experiment, using a GLM. Data were first subjected to session-wise FFX analyses for each animal (two sessions per day for M13: 26 runs; M24: 35 runs; M35: 44 runs). The onset of each condition was modeled over the entire 20 s block extent: (1) red fixation dot only, “stimulus OFF” or “stimulus ON” conditions, (2) bilateral, (3) unilateral left, (4) unilateral right stimulus display. The FFX t-map was obtained by contrasting the bilateral (2) “stimulus ON condition” with the fixation-only condition, and thresholded at p = 0.01 uncorrected (as only 6 con-images entered the MFX analysis). The t-score maps were finally projected onto the macaque F99 surface to compare the stimulus representation with maps from the shift and stay contrasts of the main experiment. This was also done in the single subjects for the calculation of overlap between activated voxels obtained by the stimulus localization and the respective shift and stay contrasts. The single monkey stimulus localization and stay contrasts (contrasts 1 and 2) for M24 and M35 (scanned with implant coils) were thresholded at FWE corrected for multiple comparisons at p = 0.05, whereas the less statistically powerful shift map (see Results, contrast 3) was thresholded at p = 0.0001 (uncorrected). In M13 (scanned with external coils), a threshold of p = 0.0001 (uncorrected) was used for the stimulus localization and stay-contrasts (1 and 2), and of p = 0.001 (uncorrected) for the shift contrast (3).

### Table 3. Eye data

| Monkey | Null left | Null right | Stay left | Stay right | Shift left | Shift right |
|--------|-----------|------------|-----------|------------|------------|------------|
| M13    | -0.05     | 0.03       | -0.06     | 0.03       | -0.09      | -0.04      |
| SD     | 0.06      | 0.03       | 0.07      | 0.04       | 0.05       | 0.03       |
| M24    | -0.06     | 0.06       | -0.08     | 0.08       | -0.07      | 0.10       |
| SD     | 0.06      | 0.06       | 0.07      | 0.07       | 0.04       | 0.05       |
| M35    | 0.00      | 0.00       | -0.01     | 0.01       | 0.01       | -0.04      |
| SD     | 0.03      | 0.03       | 0.06      | 0.06       | 0.04       | 0.03       |

**Repeated-measures ANOVA 3 × 2**

| Condition (null, stay, shift) | Direction (left, right) |
|-------------------------------|-------------------------|
| F(2,16) = 56.97 p < 0.0001    | F(1,16) = 6.67 p = 0.03 |
| F(1,16) = 10.73 p = 0.0049    | F(1,16) = 20.23 p = 0.0009 |

**Surface-based projections.** For visualization on flat maps or inflated brains, the resulting t-maps from the MFX analyses were projected onto the macaque F99 brain (Van Essen, 2004). Flat maps were created using FreeSurfer (http://surfer.nmr.mgh.harvard.edu/). All surface maps where thresholded at a minimal surface area extent of 30 mm² (http://surfer.nmr.mgh.harvard.edu/fswiki/roi_surfcluster).

**Definition of anatomical labels.** The anatomical labels, used to designate the positions of our activations on inflated F99-brain surfaces, were derived from three separate sources. Areal labels for V1, V2, V3, V4, V4A, PITd, PITv, and area MT were derived from previous retinotopic experiments. These labels were defined based upon combined retinotopic labels over 3 monkeys (probability maps of 50% overlap, including M13 and M24) (Janssens et al., 2014).

The outlines of areas 24 d, 45, 46, and 5V were derived from the Lewis and Van Essen (2000) atlas in F99 space (Lewis and Van Essen, 2000a, b). Area 5V corresponded to area PE. Dorsal and ventral premotor labels F5, F6, and F7, as well as LIP (pooled LIPa and LIPi), FEF, areas 11 and 12, and V6A were based upon the atlas of M12 (Ekstrom et al., 2009) and on the template anatomy of M12 (Friesen et al., 2009), and were warped to the F99 surface using the FreeSurfer surface-to-surface registration algorithm. Activations not falling into any of those atlases were defined anatomically with the aid of the monkey atlas of Paxinos et al. (2008).

**Time courses.** The MFX analyses did not allow us to extract raw time courses. To display time courses for ROIs showing a significant attention effect (stay or shift) in the MFX and FFX analyses, all MFX contrasts (1–3) were inclusively masked with the corresponding FFX contrasts (at FWE correction, corrected for multiple comparisons, p = 0.05), applying a cluster extent threshold of 10 voxels. ROI names, atlas source, F99 coordinates centered on the anterior commissure, and respective t-values were listed in tables corresponding to each contrast. Next, raw time courses were extracted within a 2 mm sphere surrounding the closest FFX local maxima (mean ± SD vector distance of 1.6 ± 0.7 mm), using the FFX group data (207 runs, 69 runs per animal). The raw signal was then high-pass filtered (256 s). An independent baseline was determined for each data point by calculating the moving average of the MR signal inside a window of ±50 data points with a TR of 2 (±100 s) (Cui et al., 2009; Arsenault et al., 2013).

All conditions were aligned at time point zero, and a peristimulus average percent signal change (PSC) was computed. The baseline (null events) subtracted signal is displayed in all ROI time course plots. All conditions were defined as for the GLM, and the numbers of left and right attention events were equal.

**Shift-sensitive ROIs: degree of attentional direction selectivity?** To further characterize the activity within each shift-sensitive area, we calculated a two-way repeated-measures ANOVA on the PSC with factors (1) shift versus stay, and (2) left versus right (Yantis et al., 2002) using STATISTICA version 12 (StatSoft 2014; www.statsoft.com).

A shift ROI could show a main effect of shift versus stay (factor 1) with or without a directional effect (determined by significance of factor 2, left vs right attention).

A shift-selective ROI, according to our definition, should be transiently activated for bidirectional shifts, without coding the direction of attention (Yantis et al., 2002). In contrast, a shift-sensitive area should...
show a main effect of shifting attention, and a main directional effect. Finally, within each ANOVA, we also performed post hoc tests (Fisher’s least significant difference) to test for directional selectivity of the shift conditions only (shift left vs shift right).

Analysis of behavioral data. The behavioral data of each monkey were analyzed from all runs that were included in the group analysis. As for the GLM, behavioral data were obtained from at least 3 consecutively correctly executed trials.
Probing the allocation of attention: dimming of the relevant stimulus. Percentage hits (detected relevant dimmings) and misses (undetected relevant dimmings) were calculated with respect to the overall number of response trials (relevant dimmings). False alarms (responses to irrelevant dimmings) were calculated relative to the total number of irrelevant dimmings.

Reaction times were analyzed to assess whether (1) monkeys reacted equally quickly after dimmings of relevant stimuli in the left or right visual hemifield and whether (2) differences in reaction times existed between the dimmings occurring in the three trial types (null, stay, and shift). Reaction times were sampled for all correctly performed response trials during which a dimming of the relevant stimulus occurred no more than 600 ms after trial onset. Across-session repeated-measures ANOVAs were performed for each animal (M13, 9; M24, 12; M35, 10), with factors condition (null, stay, shift) and direction of attention (left, right), for reaction times, hits, and misses (Tables 1, 2).

Eye-movement data during fixation-only trials. Eye-position data during fixation-only trials were analyzed for the fixation trials (correct sequences of at least 3 consecutive trials) that were included in the GLM.

Across-session repeated-measures ANOVAs were performed for each animal (M13, 9; M24, 12; M35, 10), with factors condition (null, stay, shift) and direction of attention (left, right), for reaction times, hits, and misses (Tables 1, 2).
Contralateral modulation of attention, cortical areas. Group MFX data in 112-RM atlas (McLaren et al., 2009) aligned to the F99 group space (anterior/posterior levels in coronal slices indicated with respect to the anterior commissure), thresholded at $p = 0.001$ (uncorrected), cluster extent threshold of 10 voxels. Middle left panels (contrast 2), Hot colors represent activations for stay right versus stay left. Cold colors represent deactivations for stay right versus stay left. Middle right panels (contrast 1), Hot colors represent activations for stay left versus stay right. Cold colors represent deactivations for stay left versus stay right. PSC plotted from raw time courses against the baseline (null events), of the ROIs indicated with the white arrows. The polarity of all signal change values has been inverted, as increase in signal produces decrease in the MION CBV maps (see Materials and Methods). Numbers indicate the ROI locations on the flatmap in Figure 3. Middle/top, For illustrative purposes, we depict the “sustained model” (example for “right attention”) (adapted from Yantis et al., 2002).
provides additional evidence that monkeys effectively attended to the relevant stimuli.

M13 fixated without interruption for an average of 4 ± 1.4 consecutive trials (mean ± SD), whereas M24 averaged 15 ± 13.6 consecutive trials, and M35 14 ± 15.3. Only trial sequences during which monkeys fixated for at least 3 consecutive trials without dimming events and rewards were included in the statistical GLM analyses.

The average fixation performance of analyzed runs (within a fixation window of 2 × 3 degrees) for M13 ranged from 85% to 95% across runs, 93% to 100% for M24, and 85% to 99% for M35.

**Contralateral attention signals**

We first identified brain regions showing fMRI activity modulated by contralateral attention using the contrast stay-left versus stay-right (contrast 1, Fig. 3a–c, right hemisphere), and vice versa (contrast 2, Fig. 3d–f, left hemisphere). Modulation of fMRI activity during contralateral stay was observed, for example, in visual cortex (V1, V2, V3, V4, V4A, MT, FST, PITv/d, area TE, posterior superior temporal sulcus [STS]) with a clear tendency for more widespread and stronger attention-dependent modulations in higher-order areas (Kastner and Ungerleider, 2000). Also parietal (LIP spreading into PE) and frontal areas 45, FEF, 46, 11, 12, F5, ACC, and SEF/F7 showed modulation of activity by contralateral attention. This was also the case for subcortical regions in both hemispheres, such as the dorsal caudate nucleus, the ventral caudate, the dorsal lateral geniculate nucleus, and the superior colliculus (Tables 4, 5).

**Time course of contralateral attention signals**

A time course analysis was used to illustrate the properties of ROIs showing significant differences between stay right and stay left. Their raw time courses, plotted against a null trial baseline (see Materials and Methods), showed a pattern similar to the
sustained attention signal previously described and modeled by Yantis et al. (2002) (Fig. 4, top, example for a stay-right selective ROI). In addition to producing the strongest activation for the contralateral stay condition and deactivation for the ipsilateral irrelevant shapes in these stay ROIs, attention shifts to the ipsilateral visual hemifield (Fig. 4, left column, magenta plots, example for shift-left) resulted in a peak in fMRI activity ~2–4 s after the shift, followed by a deactivation ~6–10 s. These time periods correspond, respectively, to the initiation of the shift in the contralateral hemifield and the reengagement of attention ipsilaterally relative to the ROI. The opposite activation pattern was observed for a shift to the right (shift-right, Fig. 4, left column, cyan lines). As already shown in humans (Yantis et al., 2002), these stay ROIs revealed modulation of contralateral attention originating before time point zero, indicated by Figure 4 (top, black arrow, Sustained Model). This activation pattern was not unexpected because attention had already been deployed, for at least one trial, to the same target location, even before the exchange of the two relevant shapes.

Posterior parietal areas, such as LIP, areas in inferotemporal cortex, the STS (for example, area FST, shown in Fig. 4), and lateral prefrontal areas, such as areas 12 and 45, displayed the prototypical time course indicative of strong contralateral attentional signals illustrated in Figure 4, with the exception of ACC (Fig. 4, bottom).

Subcortical areas, such as the dorsal lateral geniculate nucleus, the superior colliculus, as well as ventral caudate and the dorsal caudate nucleus, also exhibited profound modulation of activity for contralateral attention, in accordance with the sustained model (Fig. 5). Visual areas V3 and V4 also showed this sort of contralateral attentional modulation, as did V1 to a lesser extent (Fig. 6). Counterintuitively, caudal occipital regions along the medial wall, close to peripheral dorsal V2 (Fig. 6, #8), showed the opposite activation pattern, indicating deactivations with respect to the contralateral attended stimulus, as described previously (Kagan et al., 2010).

**Shift-sensitive activations**

By contrasting shift with stay events, we could identify shift-sensitive activations (Fig. 7, hot colors). Table 6 lists the location and t values of the ROIs activated by shifts, as obtained from the MFX analysis across sessions (and also present in the FFX analysis). Posterior shift-sensitive activations were located in caudo-dorsal occipital cortex, the most posterior portion of the STS (termed TPOC), posterior medial parietal areas V6, V6A, medial intraparietal area (MIP), and left somatomotor cortex. Frontal activations for shifting were situated in areas F6, 11, 12, anterior portions of area 46,
Figure 7. Shift-related activation does not overlap with stimulus representation. FreeSurfer F99 inflated (c, b, f, e) and flattened (a, d) surfaces. MFX analysis, contrast 3 (shift left and right vs stay left and stay right) at \( p = 0.01 \). Green transparent overlay, Activation of the stimulus localizer (MFX: contrast, bilateral stimulus vs fixation, same conventions/contrast as in Fig. 3). Numbers indicate areas of which example time courses are plotted in Figures 9, 11, and 12. Sulci and areal boundaries: same conventions as in Figure 3.
part of the ACC, and the lateral bank of the anterior principal sulcus (aPCS). Comparing the maximal t values for contralateral attention (contrast 1, $t_{\text{max}} = 12.88$; contrast 2, $t_{\text{max}} = 16.63$) with those for shifts (contrast 3, $t_{\text{max}} = 5.06$), listed in Tables 4, 5, 6, revealed major differences in statistical power for contralateral stay and shifts. We therefore applied a more liberal threshold for localizing shift-sensitive regions. It has to be noted that computing a conjunction between the contrasts shift left versus stay (left and right) and shift right versus stay, as obtained from the MFX analysis, resulted in virtually the same activated areas as for main contrast 3 (Fig. 8, coronal slices).

Overlap of stimulus representation with shift and stay signals
First, we compared the direction-sensitive stay activations with the actual sensory representation of the stimulus as measured during the independent localizer experiment (Fig. 3, green outlines overlaying the contralateral stay activity). In most cortical sites, modulation of fMRI signals by contralateral attention was enhanced in areas where the stimuli are represented. Overall, activations overlapped in striate and extrastriate visual cortex, such as V1, V3, V4, and V4A, the MT cluster, PITv/d, but also in areas more anteriorly in the STS. Overlap was also observed in the lateral bank of the IPS (LIP) and (pre)frontal areas, including anterior F5, FEF, area 45, and posterior area 46. In single animals, 46% of the voxels (M13, 43%; M24, 52%; M35, 43%) activated during the stimulus localizer experiment showed a contralateral attention signal (both contrasts thresholded at $p = 0.0001$, uncorrected, for M13; and at FWE corrected for multiple comparisons at $p = 0.05$, for M24/M35, given the implant coils; see Materials and Methods). The latter numbers are threshold dependent. Therefore, we also showed virtually unthresholded activations of the stimulus localizer experiment (6 sessions entering the MFX analysis; Fig. 9, hot colors). This analysis confirmed that the majority of contralateral stay activity (white outlines, 31 sessions entering the MFX analysis, same as in Fig. 3, hot color) is indeed restricted to areas were the stimuli are represented. Next, we plotted the PSCs obtained during the stimulus localizer experiment within the stay-selective ROIs (Fig. 9, red and blue panels). This quantitative analysis also showed that most voxels showing a contralateral stay signal are indeed driven by passive viewing of the same contralateral stimulus during the localizer experiment. The only exceptions were rostral ACC, and area F7 (SEF) in frontal cortex, which showed a pronounced and sustained contralateral attention effect, although they were not activated during the stimulus localizer experiment (Figs. 3, 9, #6 and #11, blue/red panels), as well as dorsal caudate nucleus (Fig. 9, dCaud) bilaterally.

Second, unlike most regions selectively modulated by contralateral sustained attention, the shift-sensitive activations were distinct from the stimulus representations, except for area 46 (hot colors in Fig. 7 correspond to yellow outlines and black panel in Fig. 9). In single animals, the percentage of voxels activated during the stimulus localizer test that coincided with activations driven by attention shifts (M13, $p = 0.001$, uncorrected; and M24/M35, $p = 0.0001$, uncorrected, given the implant coils; see Materials and Methods) was very small (M13, 2%; M24, 6%; M35, 3%), whereas the stimulus localizer was thresholded even higher than the shift contrast ($p = 0.0001$, uncorrected, for M13; and at FWE corrected for multiple comparisons at $p = 0.05$, for M24/M35). Thus, the majority of shift-sensitive areas are not activated during the stimulus localizer test. This strongly suggests that attentional shift signals represent a distinct attention process compared with stay signals, executed by a separated neural substrate.

Do all shift-sensitive regions compute direction-independent shift signals?
Using the same contrast (shift vs stay) as the human study, where the major shift activation was confined to SPL (Molenberghs et al., 2007), we observed several shift-sensitive activations outside caudal-medial parietal (and occipital) cortex in the monkey. It has to be noted that this contrast does not differentiate between direction-dependent and -independent attention signals. First, to distinguish between these two, we computed a two-way repeated-measures ANOVA with factors condition (shift, stay) and direction of attention (left, right), exactly as in Yantis et al. (2002). We

### Table 6. Local maxima of contrast 3 (shift vs stay)*

| Area         | Hemisphere | Atlas     | F99 space, x, y, z (mm) (MFX) | Voxel, t-value | Shift versus stay: $F_{(1,206)}$ | Left versus right: $F_{(1,206)}$ | Interaction: $F_{(1,206)}$ | ROI (Figs. 7, 9) |
|--------------|------------|-----------|-------------------------------|---------------|----------------------------------|-------------------------------|-------------------------------|-----------------|
| cmoc         | Left       | R         | -2, -38, 9                   | 3.99          | 7.72*                            | 4.96*                         | 0.00                          |                 |
| cmoc         | Right      | R         | -6, -41, 9                   | 5.06          | 11.96**                         | 5.81*                         | 0.00                          |                 |
| V6A          | Left       | N         | -5, -35, 12                  | 4.35          | 10.92**                         | 1.29                          | 0.12                          | 11              |
| V6A          | Right      | N         | 7, -36, 12                   | 4.27          | 8.40**                          | 0.31                          | 0.47                          | 12              |
| V6           | Right      | N         | 5, -37, 7                    | 3.10          | 8.51*                           | 0.14                          | 0.52                          | 10              |
| MIP          | Right      | N         | -6, -31, 14                  | 4.49          | 7.28*                           | 0.60                          | 0.32                          | 10              |
| TPOC         | Left       | L         | -11, -31, 16                 | 3.46          | 9.45*                           | 0.55                          | 0.00                          | 9               |
| Somatomotor  | Left       | A         | -16, -9, 13                  | 3.60          | 9.02*                           | 0.63                          | 0.05                          |                 |
| F6           | Right      | N         | 2, 25, 11                    | 4.16          | 7.27*                           | 0.02                          | 0.24                          | 6               |
| Area 12      | Left       | N         | 22, 7, 3                     | 3.11          | 17.71**                         | 19.94**                       | 0.18                          | 4               |
| Area 11/12   | Right      | N         | 20, 11, 6                    | 3.54          | 16.56**                         | 68.84**                       | 26.39**                       | 3               |
| Area 46      | Left       | L         | -17, 12, 13                  | 4.01          | 9.65*                           | 30.13**                       | 2.32                          | 1               |
| Area 46      | Right      | L         | 15, 11, 11                   | 4.37          | 5.80*                           | 6.18*                         | 6.08*                         | 2               |
| ACC          | Left       | A         | -4, 13, 13                   | 4.91          | 15.02**                         | 0.61                          | 0.00                          | 7               |
| ACC          | Right      | A         | 3, 14, 16                    | 3.02          | 8.34*                           | 2.74                          | 0.68                          | 8               |
| aPCS         | Left       | A         | -12, 17, 11                  | 4.87          | 17.17**                         | 0.01                          | 0.83                          | 5               |

*Clusters are shown that are significant for contrast 3 in both the MFX analysis ($p < 0.01$; Fig. 7) and the FFX analysis (FWF corrected for multiple comparisons at $p = 0.05$ with a minimum cluster extent threshold of 10 voxels. Column 5 displays the t-values for the contrast shift versus stay. Same conventions as in Tables 4 and 5 for table columns 1–5 and 9. Columns 6–8 show results of the two-way repeated-measures ANOVA, with factors (1) shift versus stay and (2) left versus right attention, and their interaction, computed with PSCs obtained from the MFX analysis. The rightmost column shows results of the MFX analysis in the MIP coronal plane. 

For table columns 1–5 and 9, Columns 6–8 show results of the two-way repeated measures ANOVA, with factors (1) shift versus stay and (2) left versus right attention, and their interaction, computed with PSCs obtained from the MFX analysis. The rightmost column shows results of the MFX analysis in the MIP coronal plane. cmoc, caudo-medial-occipital cortex. Significance levels for FWE corrected for multiple comparisons at $p < 0.001$, uncorrected, for M13; and at FWE corrected for multiple comparisons at $p = 0.05$, for M24/M35, given the implant coils; see Materials and Methods.
tested those ROIs that were present in both the MFX \( (p = 0.01) \) and FFX analysis (FWE corrected for multiple comparisons at \( p = 0.05 \)), as listed in Table 6 (see Materials and Methods). From these shift-sensitive ROIs, all lateral prefrontal ROIs, such as left area 12, right area 11/12, and left/right areas 46, showed a main effect of left versus right attention \( (F_{(1,206)} = 19.94, F_{(1,206)} = 68.84, \text{both } p < 0.0001 \) and \( F_{(1,206)} = 6.18, \text{both } p < 0.02 \), respectively; Figure 10, orange box), in addition to a main effect of shifting. This was also the case for left and right caudo-medial-occipital cortex \( (F_{(1,206)} = 4.96 \text{ and } F_{(1,206)} = 5.81, \text{both } p < 0.05) \), but as opposed to the lateral prefrontal areas, this was driven by the ipsilateral direction of attention (Fig. 10, bar plots). The caudo-medial-occipital cortex was located in vicinity of peripheral dorsal V2, also activated for ipsilateral stay (Fig. 6, #8). Additionally, right areas 12 and 46 also showed a significant interaction \( (F_{(1,206)} = 26.39, p < 0.0001 \) and \( F_{(1,206)} = 6.08, p < 0.02 \). The other ROIs, as listed in Figure 10 (yellow box), showed a main effect of shifting only and were thus shift-selective. Second, to test for direction selectivity of the shift-signal only, we performed post hoc tests within all shift ROIs (Fisher’s least significant difference, comparing shift left vs shift right). This revealed that only left area 12 \( (p < 0.001) \), right area 12 \( (p < 0.01) \), and left area 46 \( (p < 0.05) \) showed a direction-selective shift signal, but none of the other ROIs. Example raw time courses of shift-sensitive ROIs that are not shift-selective are listed in Figure 11, whereas examples of shift-selective ROIs without a significant main effect of left versus right attention are shown in Figure 12. These included, for example, bilateral posterior medial parietal cortex, including areas V6/V6A, and MIP, but also the TPOC, ACC, the left aPCS corresponding to the rostrolateral border of area 46, and right area F6.

**Summary: shifting compared with contralateral stay**

Figure 13 summarizes the spatial relationships between areas showing contralateral stay activity (blue represents stay-right, left hemisphere; red represents stay-left, right hemisphere) and regions showing shift signals without (yellow represents shift only) and with a direction-selective attention component (orange represents shift together with left vs right attention). Because of the more stringent criteria applied here, a few small shift-sensitive ROIs that survived the \( p < 0.01 \) criteria in the MFX-based analysis (Fig. 7; e.g., LIP, PE, right TPOC, and bilateral insula) disap-
Figure 9. Segregation of shift signals compared with stay and stimulus representation signals. Top, FreeSurfer F99 inflated and flattened surfaces (LH, left hemisphere; RH, right hemisphere), displaying activation of the stimulus representation (from separate localizer experiment) obtained with the same contrast as in Figures 3 and 7 (green), but at virtually unthresholded level (hot color, \( t > 0.02 \)). The majority of areas activated by contralateral attention (white outlines, same as hot color in Fig. 3) overlaps with the stimulus representation, opposed to areas modulated by attention shifts (yellow outlines, same as hot color in Fig. 7), clustering outside the activated areas. Black outlines indicate the same as white outlines of retinotopic areas in Figures 3 and 7. Blue panel, PSC of localizer data within the stay-right selective ROIs in the left hemisphere. Red panel, PSC of localizer data within stay-left selective ROIs in the right hemisphere (same ROIs/numbers as in Figs. 3, 4, 5, 6). The stimulus is represented in all areas driven by contralateral attention, except in the ACC and SEF/F7 and the caudate nucleus, and much less so in subcortical areas. * \( p < 0.05 \) (t test, Bonferroni corrected for \( n = 14 \)). Black panel, Shift-selective areas (same ROIs/numbers as in Figs. 7, 11, 12) are not driven by the stimulus localizer, except for left and right areas 46. * \( p < 0.05 \) (t test, Bonferroni corrected for \( n = 12 \)). Bar plots represent conditions of the stimulus localizer experiment: (1) left unilateral, (2) right unilateral, and (3) bilateral stimulus display versus fixation (see Materials and Methods).
peared in Figure 13, which is a combined summary of the MFX map with the data of the two-way ANOVA based on the FFX analysis (Table 6). In general, a much larger cortical territory was modulated by stay compared with shift activity. The only position where shift signals were not adjacent to or overlapping with stay signals was in posterior medial parietal cortex. To estimate the regional distribution of cortical shift signals, we computed the number of voxels from contrast 3 that showed a main effect for shifting and no direction-selective attention signal (Fig. 13, yellow shift-selective voxels; see also Table 6). This analysis confirmed that the majority of these shift-selective voxels (63%) were found in posterior medial parietal cortex (Fig. 13, yellow within red half-circle).

Outside posterior medial parietal cortex, shift activations were typically adjacent to stay-selective voxels (with right F6 and left CS as the only exception). For example, the shift activations in left F6 neighbored stay selectivity in F7 of the left hemisphere. Areas TPOC, ACC, areas 12 and 11/12-border, and area 46 also showed adjacent or overlapping stay- and shift-sensitive activity (Fig. 13). This suggests functional heterogeneity of these areas. Furthermore, a gradual transition existed from stay to more shift-related activity along a caudorostral axis in left area 46, which is most likely a consequence of the heterogeneous connectivity of this region (Gerbella et al., 2013).

**Discussion**

We recorded fMRI activity while monkeys performed covert shifts of spatial attention compared with a sustained attention baseline. We found the most extensive shift-selective signals in posterior medial parietal cortex, including the posterior portion of the fundus of the IPS, extending medially into MIP and the parieto-occipital sulcus comprising V6 and V6A (Figs. 12, 13). The rostrolateral border of area 46, ACC, somatomotor cortex, F6, and the most posterior tip of the STS (TPOC) also contained regions showing shift signals, not coding direction-selective attention signals (Figs. 12, 13, yellow). Furthermore, functionally more heterogeneous areas 46, 11, 12, and ACC contained subregions sensitive to either shifts or contralateral attention. Unlike most areas modulated by contralateral attention (Fig. 3), regions showing shift signals were not activated by the visual stimulus representation, suggesting that they form a distinct functional network, most likely from a higher-order nature (Figs. 7, 9). Area TPOC is remote from monkey’s posterior medial parietal cortex and therefore considered an unlikely potential homolog of human SPL. It may correspond topologically to a human region near the temporoparietal junction (Corbetta et al., 2000).

Because the most extensive shift signal was located in posterior medial parietal cortical areas V6/V6A and MIP, where no stay activity was found, and because of its topographic location, we propose this region as the functional counterpart of human SPL, which is also activated by shifts of spatial attention.

**Comparison of monkey and human cortical areas showing attentional shift signals**

**Parietal lobe**

Several human fMRI studies, one using an experimental paradigm virtually identical to the present study, revealed signals coding spatial attention shifts in medial SPL regardless of the shift direction (Yantis et al., 2002; Molenberghs et al., 2007). Furthermore, the same region was also engaged when human subjects shifted attention between object (Serences et al., 2004), features, and task rules (Shomstein and Yantis, 2004; Chiu and Yantis, 2009; Shulman et al., 2009). Exactly as in the present monkey study, areas showing shift-related signals in humans were largely...
segregated from those maintaining the current locus of attention (Yantis et al., 2002; Vandenberghe and Gillebert, 2009). In addition, Simon et al. (2002) showed systematic activations of the SPL, using four different spatially specific visuomotor tasks, including covert shifts of attention, voluntary saccades, and spatially specific grasping and pointing. The computation of attention shifts is most likely the common feature of these tasks. Causal evidence for SPL playing a functional role in attentional shifting was provided by Ciavarro et al. (2013). They applied rTMS to posterior medial SPL, which they termed putative human V6A, and observed prolonged reaction times during reorienting trials in the context of a spatial attention and a spatially selective reaching task. Corroborating causal evidence comes from another recent TMS study showing that the medial SPL is involved in shifts of attention regardless of the current locus of attention (Capotosto et al., 2013). In sharp contrast to shift activity, which was confined to SPL, modulation of fMRI activity by contralateral spatial attention signals was observed in the posterior IPS and extrastriate human visual cortex (Yantis et al., 2002; Molenberghs et al., 2008; Shulman et al., 2009; Vandenberghe et al., 2012). This was also confirmed causally by rTMS of the IPS (Capotosto et al., 2013). This study showed impairment of target discriminations largely limited to the contralateral hemifield indicating that the IPS signals the current locus of attention. Thus, in human parietal cortex, SPL encodes attentional shifts, whereas activity in posterior IPS is modulated by sustained attention signals.

In the present monkey study, sustained contralateral attention also activated the IPS (LIP) while posterior medial parietal cortex was activated by shifts of attention. The former result is consistent with a lesion study showing impaired performance during the detection of contralateral targets in the context of a conjunction search task (Wardak et al., 2004). Thus, as in humans, different subregions of monkey parietal cortex are involved in different aspects of the control of spatial attention, such as sustained contralateral attention, versus spatial-attentional shifting. Monkey V6A receives most of its visual inputs via area V6 (Galletti et al., 2001; Passarelli et al., 2011), and it contains neurons showing attention, saccade, reach, and grasping-related activity (Galletti et al., 1996, 1999, 2010; Kutz et al., 2003; Fattori et al., 2004, 2005; Premereur et al., 2015). Areas V6A and MIP in macaque form a tightly linked complex where visual, somatic, and motor information is integrated (Passarelli et al., 2011). This matches the functional characteristics of human SPL and distinguishes V6A/MIP-complex from areas simply responsive to visual stimulation (Galletti et al., 2010) or modulated by sustained attention (Fig. 9). More generally, the present results suggest that the functional architecture of monkey and human SPL may be more similar than initially proposed (Grefkes and Fink, 2005; Margulies et al., 2009; Vanduffel et al., 2014). Indeed, several authors (Vandenbergh et al., 2001; Simon et al., 2002; Galletti et al., 2010) suggested tentatively that human SPL might be homologous to monkey area V6A. In humans, SPL has been hypothesized to be the source of attentional modulation, whereas extrastriate areas have been called the sites of attentional modulation (Yantis et al., 2002). Possibly, the monkey V6A/MIP complex similarly exerts higher-order control over extrastriate cortex,

Figure 11. Raw time courses of shift-sensitive ROIs (main effect shift and left vs right attention). MFX contrast 3 (shift left and shift right vs stay left and stay right, \( p = 0.01 \)), cluster extent threshold of 10 voxels, projected onto the freesurfer F99-macaque anatomical volume. Numbers within the circles indicate ROI locations on the flatmap in Figures 7 and 9. Error bars indicate the trial-wise SEM.
but this requires further study (e.g., by means of effective connectivity analysis) (Ekstrom et al., 2008; Friston, 2011).

**Frontal cortex**

Despite commonalities between human and monkey parietal cortex in present and previous studies focusing on attention shifts, differences were observed in frontal cortex. For example, shift-sensitive fMRI activations were observed in several frontal areas of monkeys but not humans (Molenberghs et al., 2007).

Specifically, shift signals were observed in frontal areas 46, 11/12, ACC, and pre-SMA/F6. A potential explanation for the apparent discrepancy between human and monkey studies may lie in the presence and absence of feedback signals, respectively. In monkeys, ACC is involved in computing context-dependent error likelihoods by integrating feedback signals (Kuwabara et al., 2014). Whereas reward contingencies have been shown to be represented independently of spatial attention shifts within human medial parietal cortex (Tosoni et al., 2013), no such evidence exists for monkey parietal cortex.

Another explanation may be that endogenously driven shifts in attention may require relatively more cognitive control for monkeys than for humans, explaining the stronger activation of monkey PFC. Corroborating evidence that task demand may be a critical factor regarding the contribution of frontal cortex comes from a human study directly comparing shifts between task rules and spatial-attention shifts within the same task (Chiu and Yantis, 2009). Unlike Molenberghs et al. (2007), these authors ob-
served, in addition to SPL, precentral and frontal activations when human subjects had to shift between different task sets in addition to spatial shifts in attention, thus requiring higher levels of cognitive control. In their comparative fMRI study, Nakahara et al. (2002) also showed activations in ventral orbitofrontal area 12 when humans and monkeys had to switch task sets during a cognitively demanding Wisconsin Card Sorting task. In our study, area 12 was activated during spatial attention shifts as well as stay-trials, corroborating its role in cognitive control. Finally, it is worth noting that a previous comparative fMRI study reported much stronger and widespread activations in monkey PFC compared with that of humans when subjects simply had to fixate to images of objects versus scrambled images (Denys et al., 2004).

**Shifts of attention or response preparation in posterior medial parietal cortex?**

Area MIP is part of the parietal “reaching” region (Cohen and Andersen, 2002), which also includes other areas, such as V6A and the intraparietal parts of area PE. Furthermore, it has been claimed that MIP neurons are involved in processing visuospatial transformations (Cohen and Andersen, 2002), even before a movement is initiated (Johnson et al., 1996). Hence, one could argue that the shift-selective activations observed in MIP and V6A are linked with preparatory motor activity in the monkey. We think this is an unlikely explanation for our results because dimming events were matched between stay and shift trials, between left and right hemifield, and across relevant and irrelevant shapes. Monkeys were trained to respond to dimming events occurring independently of shift events. Any response-preparatory activity should thus be equally distributed across event types; hence, this cannot explain the observed shift signals in medial parietal cortex.

Monkey V6A contains visual, somatic, oculomotor, and reaching neurons (Battaglia-Mayer et al., 2001; Gamberini et al., 2009). Consistent with our results, Galletti et al. (2010) showed that V6A neurons fire during covert shifts of attention, with equal numbers of cells tuned for contralateral and ipsilateral shifts. Compared with previous human and monkey studies, we attempted to match all sensory events as well as possible, measured eye and hand positions, and dissociated these potential confounds from the events of interest (shift/stay). Nonetheless, we observed shift-selective activations in this region of posterior medial parietal cortex, indicating that also in monkey, there may exist a region that is responsible for shifts of attention in different modalities.

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**Figure 13.** Spatial comparison between cortical stay and shift signals. FreeSurfer F99 inflated (top) and flattened (bottom) surfaces. Only those shift ROIs from Figure 7 are plotted that show a significant main effect (two-way ANOVA, shift/stay and left/right) according to Table 6 and Figure 10. Yellow represents shift-selective ROIs, without a main effect of left versus right attention. Orange represents shift-sensitive ROIs, displaying a main effect of left versus right attention. Contralateral stay (blue/red, same as in Fig. 3), left hemisphere (LH), contrast 2 (blue), and right hemisphere (RH), contrast 1 (red). Red dotted lines indicate medial occipitoparietal cortex: shift activations not overlapping with the stay activations. For all labels/ROIs, same conventions as in Figures 3, 7, and 9. cmoc, Caudo-medial-occipital cortex.
In conclusion, in monkeys as in humans (Molenberghs et al., 2007), posterior medial parietal cortex is involved in shifting spatial attention in the absence of overt behavior. Our results suggest that monkey posterior medial parietal cortex is functionally homologous to a topographically corresponding portion in the human (SPL). Hence, we posit that, in addition to the established role of this part of the cortex in directed hand and arm movements (and to a lesser extent also eye movements), it serves as a key area for shifting spatial attention. In the monkey, additional frontal areas are recruited during attention shifts, complying with the higher cognitive demands required for the computation of attention shifts, relative to humans. Also, in correspondence with human IPS, its monkey counterpart, and particularly LIP, may be instrumental in coding the location and significance of a stimulus, as others have suggested (Gottlieb and Goldberg, 1999; Wardak et al., 2002; Bisley and Goldberg, 2003b; Molenberghs et al., 2007).

The identification of a likely functional homology between human SPL and monkey posterior medial parietal cortex invites investigation into the neuronal mechanisms underlying attentional shifting using functional connectivity analyses, electrophysiological recordings, and the most refined causal methods in an animal model closely akin to humans (Vanduffel et al., 2014).

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