Hemispheric Asymmetry in Visuotopic Posterior Parietal Cortex Emerges with Visual Short-Term Memory Load

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Visual short-term memory (VSTM) briefly maintains a limited sampling from the visual world. Activity in the intraparietal sulcus (IPS) tightly correlates with the number of items stored in VSTM. This activity may occur in or near to multiple distinct visuotopically mapped cortical areas that have been identified in IPS. To understand the topographic and spatial properties of VSTM, we investigated VSTM activity in visuotopic IPS regions using functional magnetic resonance imaging. VSTM drove areas IPS0 –2, but largely spared IPS3 –4. Under visual stimulation, these areas in both hemispheres code the contralateral visual hemifield. In contrast to the hemispheric symmetry observed with visual stimulation, an asymmetry emerged during VSTM with increasing memory load. The left hemisphere exhibited load-dependent activity only for contralateral memory items; right hemisphere activity reflected VSTM load regardless of visual-field location. Our findings demonstrate that VSTM induces a switch in spatial representation in right hemisphere IPS from contralateral to full-field coding. The load dependence of right hemisphere effects argues that memory-dependent and/or attention-dependent processes drive this change in spatial processing. This offers a novel means for investigating spatial-processing impairments in hemispatial neglect.

Introduction

Visual short-term memory (VSTM) refers to the encoding, maintenance, and retrieval of visually presented stimuli that are difficult to name and unlikely to be held in long-term memory (Phillips, 1974). Although our subjective visual experience suggests an effortless ability to encode all of the information before us, VSTM capacity is limited to only three or four objects (Luck and Vogel, 1997). Human neuroimaging studies have demonstrated that activity in the intraparietal sulcus (IPS) closely reflects the number of items held in VSTM (Todd and Marois, 2004; Vogel and Machizawa, 2004; Xu and Chun, 2006; Xu, 2007; Harrison et al., 2010); however, the nature of spatial processing during VSTM is unresolved.

The medial bank of human IPS contains at least five areas, as follows: IPS0 (also known as V7); IPS1; IPS2; IPS3; and IPS4 with visuotopic maps (Tootell et al., 1998; Sereno et al., 2001; Schluppeck et al., 2005; Silver et al., 2005; Swisher et al., 2007; Konen and Kastner, 2008a; Silver and Kastner, 2009). Research has only begun to delineate the specific functions of what may be close to a dozen specific cortical areas in IPS (Astafov et al., 2003; Greffkes et al., 2004; Orban et al., 2006; Konen and Kastner, 2008a,b).

Each hemisphere of IPS0 – 4 maps the contralateral visual field in a highly symmetric manner. The hemispheric symmetry of topographic maps in IPS sharply contrasts with findings from hemispatial neglect syndrome. Patients suffering from hemispatial neglect exhibit severe deficits in contralesional attentional processing predominantly after damage to right hemisphere parietotemporal and/or frontal lobes (Bisiach and Lazzati, 1978; Heilman and Van Den Abell, 1980; Mesulam, 1981; Vallar and Perani, 1986; Mesulam, 1999; Pouget and Driver, 2000; Dorichchi et al., 2008). Although acute, structural damage in hemispatial neglect is normally found ventral to topographic regions of the posterior parietal cortex (PPC), a seminal study (Corbetta et al., 2005) has demonstrated that activity in dorsal PPC regions diminishes during the acute phase of neglect and normalizes with the amelioration of neglect symptoms. These observations suggest that hemispatial neglect emerges from damage to a network including visuotopically mapped IPS (Doricchi et al., 2008). Several studies have investigated the contralateral bias during visual mapping and visual attention (Sereno et al., 2001; Silver et al., 2005; Swisher et al., 2007; Shulman et al., 2010; Szczepanski et al., 2010), but there has not yet been a systematic study of spatial processing in IPS during VSTM. Moreover, at present, we have few insights as to how to study the neglect network in normal healthy human subjects.

Here, we mapped the spatial and functional relationships between VSTM activation and the five visually driven visuotopic cortical areas IPS0 –4 (Swisher et al., 2007) using functional magnetic resonance imaging (fMRI). As visuotopic IPS areas are relatively small and reported Talairach coordinates of VSTM IPS activation show considerable variability, we first investigated the spatial overlap of visuotopic maps and areas subserving VSTM within individual subjects. Then, we investigated whether contralateral bias, characteristic of visuotopic maps, is also found for VSTM representations in the parietal cortex.
Materials and Methods

Subjects. Nine healthy subjects (4 female, 5 male) with normal or corrected-to-normal vision gave informed written consent to participate in the study, which was approved by the Institutional Review Boards of Boston University and Partners Healthcare. Subjects were recruited from the Boston University community. All subjects were right handed and were between the ages of 23 and 31 years. One author (S.L.S.) served as a subject.

Visual stimuli and experimental paradigm. Visual stimuli were driven by a Macintosh MacBookPro using the Vision Egg software package (Straw, 2008) and were presented using a liquid crystal display projector illuminating a screen within the scanner bore. Subjects viewed the rear-projection screen through a mirror angled at ~45°. The screen extended across a visual angle of ~12° radius horizontally and ~9° radius vertically. Subjects fixated a centrally located cross, while eight colored bars oriented vertically or horizontally were presented to the left visual field and eight colored bars were presented to the right visual field (see Fig. 1). The total number of bars per hemisphere was kept constant across all trials, but in a given block of trials one, three, or six of the bars were memory targets, whereas the others were distractors. Targets were distinguished from the distractors by color (blue/red, counterbalanced across subjects). Subjects were asked to remember the orientation (horizontal or vertical) of each target. Each bar subtended 0.7° × 0.23° of the visual angle. The memory sample was presented for 100 ms. After a memory delay of 900 ms, a memory probe was presented for 2000 ms. Trials were 6 s long, with a 3000 ms fixation period at the end of each trial. Subjects could respond during the memory probe or during the first 1000 ms of the post-probe period. In half of the trials, all the bars in the probe had the same orientation as in the memory sample. In the other half of the trials, one of the target bars changed from horizontal to vertical or from vertical to horizontal. Subjects were then asked to report, via button press, whether or not the orientation of the probe target bars matched those of the memory target bars. In the remember-left conditions, all targets were presented left of the fixation cross, whereas the other stimuli were distractors. In the remember-right conditions, all targets were presented to the right. In the remember-full-screen condition, the targets appeared either on the left or the right side of the screen with equal probability. The targets were restricted to a single hemifield in each trial to control for memory capacity across the conditions, as prior studies have revealed greater capacity when targets are spread across both hemifields (Alvarez and Cavanagh, 2005; Delvenne, 2005). Three of the subjects switched their response hand from run to run, whereas six subjects used their right hand exclusively. During passive-viewing trials, subjects were asked to simply press a key when the second set of stimuli appeared.

Subjects’ performance for VSTM was measured in terms of capacity. Although multiple estimates of K have been reported, we used Cowan’s K (Cowan, 2001). Cowan’s K is defined as follows:

\[ K = SS × (HR − FA) \]

where SS is the set size, HR is the hit rate, and FA is the false-alarm rate. K traditionally increases with set size until maximum capacity is reached and then plateaus at larger set sizes. An ANOVA was performed to measure the effects of set size and stimulus location.

Data acquisition. Data were acquired from a 3T Siemens Tim Trio magnetic resonance imager located at the Martinos Center for Biomedical Imaging at Massachusetts General Hospital, Charlestown, MA. Each subject participated in three critical types of scans across multiple sessions: high-resolution structural MRI, visuotopic mapping fMRI, and VSTM fMRI. In an initial session, high-resolution (1.0 × 1.0 × 1.3 mm) structural images were obtained for surface reconstruction, using a 12 channel birdcage head coil. Computer representations of each cortical hemispheric surface were unfolded and flattened using the Freesurfer software package (Dale et al., 1999; Fischl et al., 1999, 2001). Functional scans measured changes in the intrinsic blood oxygenation level-dependent (BOLD) contrast. Cortical representations of eccentricity and polar angle were mapped with flashing checker-board stimuli in separate runs and were combined using standard techniques (Engel et al., 1994; Sereno et al., 1995; Swisher et al., 2007) with parameters optimized to reveal maps in the parietal cortex. An F test was performed for each voxel to test for a difference between the power at the stimulus frequency and the power of the other frequencies in the time series. For the polar mapping, a wedge was created by contrasting a color-reversing checkerboard against a black background. Subjects were asked to maintain fixation while the wedge rotated around a central fixation point. The polar angle wedge had an arc of 72°, flashed at 4 Hz, had a sweep period of 42.67 s, and swept out 12 cycles per run (for a more detailed description of the visuotopic-mapping methods, see Swisher et al., 2007). Visual stimuli were projected into the patient chamber onto a rear-projection screen (Da-Plex, Da-Lite Screen) and viewed via an adjustable mirror.

In the VSTM study, stimulus location (remember-left, remember-right, and remember-full-screen) and VSTM load (set sizes 1, 3, and 6) varied by block. In addition, subjects performed a single block of passive viewing per run, and block order was randomized within runs and counter-balanced across runs. Each subject performed eight runs, each lasting 7 min and 12 s, which consisted of 10 40 s blocks as well as 16 s of blank fixation both before the first block and after the last block. Each block consisted of a 4 s cue, which indicated the set size and location of the target stimuli, followed by six trials each lasting 6 s. The 40 s blocks were used to allow subjects to maintain stable attentional and cognitive sets. A total of 32 slices oriented parallel to the calcarine sulcus were collected (TR = 2 s; voxels = 3.125 × 3.125 × 5 mm) using a bilateral quadrature-phase surface coil placed posterior to the occipital pole.

Data analysis. Intensity normalization was performed before signal averaging (Dale et al., 1999; Fischl et al., 1999). Single-subject VSTM fMRI data were analyzed voxel by voxel using a general linear model that included a predictor for each condition (e.g., each location-set size pairing and passive viewing). A t test was performed on each voxel to compare activation differences between conditions, and significance values were projected onto flattened cortical hemisphere representations. For the voxel-by-voxel analysis, the BOLD signal was modeled as a linear time-invariant system: a response function was assumed or each stimulus condition, with a delay of \( d = 2.25 \text{s}\) and a decay time constant of \( r = 1.25 \text{s} \). An estimated response was generated by convolving the response function with the stimulus time course (i.e., each block) and minimizing the residual error (FS-FAST, Cortech). Random-effects group analysis was performed using surface-based averaging techniques (Fischl et al., 1999). For the region of interest (ROI) analysis, the percentage signal change data were extracted (from all time points for a block) and averaged by condition, across runs, to construct time-course data for all voxels with a functionally defined ROI. The percentage signal change measure is relative to the average activation level during the fixation condition. Contralateral biases (see Fig. 6) were estimated at each set size for each ROI using the following formula:

\[ \text{Contralateral}_{\text{set size}} - \text{Ipsilateral}_{\text{set size}} \] / \( \text{Contralateral}_{\text{set size}} + \text{Ipsilateral}_{\text{set size}} \)

VSTM biases were compared across hemispheres using a paired t test.

Visuotopic IPS ROIs IPS0 (also known as V7), IPS1, IPS2, IPS3, and IPS4 were determined using polar angle retinotopy scan data and techniques reported previously (Swisher et al., 2007). For the phase angle analysis of Figure 6, we created binary masks for IPS0—2 using only those voxels significantly activated (\( p < 0.05 \)) during visuotopic mapping. For each region, we then extracted the preferred phase angle on a voxel-by-voxel basis. Data were summed across subjects and represented using a single polar histogram.

Eye position monitoring. Two subjects returned to perform VSTM during tracking of eye position. The stimuli and procedures for the eye-
ANOVA revealed a significant effect of set size (member-full-screen left size 1, to 1.94 at set size 3, and to 2.17 at set size 6 (remember-full-screen condition, the targets appeared either on the left, whereas all of the stimuli on the right were distractors. In the remember-left conditions, the targets were presented to the left visual field and eight colored bars were presented to the right visual field and eight colored bars oriented vertically or horizontally were presented to the left visual field and eight colored bars were presented to the right visual field (Fig. 1). Because visuotopic IPS areas are sensitive to visual drive (Swisher et al., 2007), the number of visual stimuli was kept constant for all conditions. Subjects were asked to remember the orientation (horizontal or vertical) of a subset of the bars that were distinguished from the distractors by color (red or blue). The number of the targets (1, 3, or 6) and their location (right visual field or left visual field) varied across blocks. In the remember-left conditions, the targets were presented to the left, whereas all of the stimuli on the right were distractors. In the remember-right conditions, the targets were presented to the right, whereas all of the stimuli on the left were distractors. In the remember-full-screen condition, the targets appeared either on the left or the right side of the screen with equal probability.

VSTM behavioral performance
VSTM behavioral capacity was assessed according to Cowan’s K (see Materials and Methods), a measure of the number of items that can be held simultaneously in short-term memory (Cowan, 2001). For all three target locations (full-screen, right, and left), K score increased and reached a plateau with increasing set size (Fig. 2). These results are consistent with prior behavioral reports (Cowan, 2001; Todd and Marois, 2004; Vogel and Machizawa, 2004; Xu and Chun, 2006). Overall, K increased from 0.91 at set size 1, to 1.94 at set size 3, and to 2.17 at set size 6 (member-left = 0.90, 1.85, 2.00; remember-right = 0.93, 1.87, 2.21; remember-full-screen = 0.92, 2.09, 2.29). A repeated-measures ANOVA revealed a significant effect of set size ($F_{(3,16)} = 44.07$, $p < 0.001$), which represented a significant difference between set sizes 1 and 3 ($t_{(26)} = 9.80$, $p < 0.001$) and set sizes 1 and 6 ($t_{(26)} = 8.64$, $p < 0.001$), but no significant difference between set sizes 3 and 6. There was neither a significant effect of target location nor an interaction between set size and target location, demonstrating that memory performance was not different across the visual hemifields.

Results
Each of nine subjects participated in a minimum of three sets of scans across multiple sessions. First, high-resolution (1.0 × 1.0 × 1.3 mm) structural scans were performed to support anatomical reconstruction of the cortical hemisphere surfaces. Retinotopic mapping fMRI scans were performed to identify visuotopic areas IPS0–4 within the PPC of each subject hemisphere (Swisher et al., 2007). Visual short-term memory fMRI scans were performed with varying degrees of memory load and varying visual field location of memory targets.

In VSTM scans, subjects fixated a centrally located cross while eight colored bars oriented vertically or horizontally were presented to the left visual field and eight colored bars were presented to the right visual field (Fig. 1). Because visuotopic IPS areas are sensitive to visual drive (Swisher et al., 2007), the number of visual stimuli was kept constant for all conditions. Subjects were asked to remember the orientation (horizontal or vertical) of a subset of the bars that were distinguished from the distractors by color (red or blue). The number of the targets (1, 3, or 6) and their location (right visual field or left visual field) varied across blocks. In the remember-left conditions, the targets were presented to the left, whereas all of the stimuli on the right were distractors. In the remember-right conditions, the targets were presented to the right, whereas all of the stimuli on the left were distractors. In the remember-full-screen condition, the targets appeared either on the left or the right side of the screen with equal probability.

Figure 1. VSTM trial structure. A, Subjects encoded a set of briefly presented targets among distractors, and after a delay were asked whether the orientation of any of the targets had changed. During the experiment, targets and distractors were presented in red and blue, respectively (colors randomized between subjects). Here, targets are depicted in black, and distractors in white. B, Target location conditions. In trial blocks, targets appeared always in the right visual hemifield (remember-right), always in the left visual hemifield (remember-left), or could appear in either hemifield (remember-full-screen). The number of targets also varied across blocks (set sizes 1, 3, and 6), but the total number of stimuli remained constant.

Figure 2. Average VSTM K score for each set size and location. K score plateaus for each memory location, with no significant differences between the locations.
Overlap between VSTM activation and visuotopic IPS

Visualization of fMRI activation patterns for the remember-full-screen VSTM data revealed significant overlap of the areas activated by VSTM and visuotopic mapping (Fig. 3A, B). Both visuotopic mapping and VSTM showed a band of activity along the medial bank of IPS. Along the posterior branch of IPS, VSTM activity and visuotopic areas IPS3–4 showed less overlap between visuotopic mapping and VSTM in IPS3–4 than in IPS0–2 (IPS3 vs IPS0: t(17) = 5.19, p < 0.001; IPS3 vs IPS1: t(17) = 5.172, p < 0.001; IPS3 vs IPS2: t(17) = 5.336, p < 0.001; IPS4 vs IPS0: t(17) = 6.310, p < 0.001; IPS4 vs IPS1: t(17) = 6.277, p < 0.001; and IPS4 vs IPS2: t(17) = 4.908, p < 0.001), but there were no significant differences among regions IPS0–2 or between IPS3 and IPS4. There was neither a significant effect of hemisphere nor an interaction between hemisphere and ROI demonstrating that in the remember-full-screen condition, the right and left hemispheres exhibited highly symmetric VSTM activation. These results also demonstrate a functional dissociation for VSTM in IPS. Areas recruited in VSTM are strongly coincident with areas IPS0–2, but largely spare areas IPS3–4, with the overlap primarily in the fundus of IPS. Although one must always be cautious when comparing effects across ROIs, we note that visuotopic areas IPS0–4 follow the medial bank of IPS, whereas VSTM activation follows the medial bank of posterior IPS and the lateral/inferior bank of anterior IPS. This anatomical distinction also indicates that VSTM activation is coincident with IPS0–2, but largely lateral/inferior to IPS3–4.

Effects of set size and visual hemifield on BOLD signal in IPS

We analyzed the remember-full-screen, remember-left, and remember-right conditions in both hemispheres for each VSTM set size. For each hemisphere, the conditions were treated as full-screen, contralateral, and ipsilateral locations. Because there was minimal overlap between VSTM and visuotopic mapping for regions IPS3–4, we restricted further analyses to regions IPS0–2. A repeated-measures ANOVA revealed a significant main effect of set size (F(2,34) = 4.52, p = 0.029). This can be seen qualitatively in Figure 4A–C (remember-full-screen) and quantitatively in Figure 4D (all 3 locations together). BOLD signal mirrored subjects’ memory capacity, with a significant difference between set sizes 1 and 3 (t(16) = 6.33, p < 0.001) and set sizes 1 and 6 (t(16) = 6.10, p < 0.001), but not between set sizes 3 and 6. These results indicate that visuotopic areas IPS0–2 are increasingly recruited as subjects remember a greater number of items, arguing for a functional role during VSTM. The location of activity is similar to prior reports of IPS that did not identify visuotopic areas (Todd and Marois, 2004; Xu and Chun, 2006; Xu, 2007).

To be activated during visuotopic mapping, a visually responsive voxel must show a spatial preference in terms of its population receptive field. If the VSTM BOLD signal in IPS0–2 reflects activity resulting from the same neural populations, these voxels would be expected to show a strong contralateral bias for VSTM representations. A significant main effect of memory stimulus location (i.e., contralateral > ipsilateral) would indicate that during VSTM there is a bias for remembered items presented in the contralateral visual hemifield. Consistent with this hypothesis, the repeated-measures ANOVA revealed a significant main effect of visuotopic location (F(2,32) = 3.70, p = 0.048). Paired comparisons showed a significant difference between the contralateral and ipsilateral (t(16) = 4.86, p < 0.001) and ipsilateral and full-screen (t(16) = 2.44, p < 0.029) conditions, but no significant difference between IPS0–2 and IPS3–4.
difference between contralateral and full-screen conditions. These results are largely expected, given the demonstrated overlap between VSTM activation and visuotopic maps.

Although there was not an overall effect of hemisphere, there was evidence of hemispheric asymmetry in VSTM processing. There were significant interactions between set size and hemisphere \( (F_{2,16} = 3.65, p = 0.049) \), and visual location and hemisphere \( (F_{2,16} = 5.00, p = 0.021) \). In addition, there was a marginally significant three-way interaction among visual location, hemisphere, and set size \( (F = 2.38, p = 0.07) \).

To better investigate the hemispheric asymmetry, we compared contralateral and ipsilateral responses for the two hemispheres. The group average for the high-load conditions (contralateral vs ipsilateral for set sizes 3 and 6) revealed a dramatic asymmetry (Fig. 5a). Under high-load conditions (set sizes 3 and 6), the left hemisphere exhibited a pattern of strong contralateral bias for VSTM that one would expect for visuotopic areas. In contrast, no contralateral bias was observed for the right hemisphere. Figure 5B displays the activation level for each hemisphere, for each set size and location. The left hemisphere exhibited increasing BOLD signal with increasing set size for contralateral stimuli only (set size 1 vs set size 3, \( t_{23} = 3.51, p < 0.01 \); set size 1 vs set size 6, \( t_{23} = 4.12, p < 0.001 \)); ipsilateral responses were flat across set sizes. In contrast, the right hemisphere exhibited increasing BOLD signal with increasing set size for both contralateral and ipsilateral memory targets (contralateral: set size 1 vs set size 3, \( t_{23} = 3.18, p < 0.01 \); set size 1 vs set size 6, \( t_{23} = 4.58, p < 0.001 \); ipsilateral: set size 1 vs set size 3, \( t_{23} = 7.23, p < 0.001 \); set size 1 vs set size 6, \( t_{23} = 4.94, p < 0.001 \)).

These results reveal a surprising, load-dependent, hemispheric asymmetry in VSTM processing in which the left hemisphere is strongly biased toward coding contralateral targets, but the right hemisphere robustly codes targets in both hemifields. Control studies ruled out eye position and response hand explanations of this hemispheric asymmetry (supplemental materials, available at www.jneurosci.org as supplemental material).

The combined results present a paradox. Visuotopic IPS shows a strong bias for stimuli in the contralateral visual field for both cortical hemispheres. This hemispheric symmetry contrasts with the asymmetry observed for VSTM activation. To confirm that both results hold for the same ROI voxels, we conducted an
analysis in which the preferred polar angle during retinotopic mapping scans was extracted for each voxel of each subjects’ ROIs. We created binary brain masks for IPS0–2 using only those voxels significantly activated (p < 0.05) during visuotopic mapping. The preferred angles were then summed across subjects and plotted on a polar histogram (Fig. 6A). The results demonstrate that each ROI showed a consistent and hemispherically symmetric contralateral bias during visuotopic mapping.

Conversely, analysis of these same voxels during VSTM revealed there was a hemispheric asymmetry in the degree of contralateral bias (Fig. 6B). In the left hemisphere, there was a significant contralateral bias for set sizes 3 and 6 for IPS0 (set size 3, t(26) = 3.564, p < 0.01; set size 6, t(26) = 6.20, p < 0.001), IPS1 (set size 3, t(26) = 2.449, p = 0.040; set size 6, t(26) = 3.029, p = 0.016), and IPS2 (set size 3, t(26) = 2.351, p = 0.047; set size 6, t(26) = 3.744, p < 0.01). There was no significant effect of stimulus location for the right hemisphere for any ROI at any set size. The contralateral bias between the hemispheres was not different at set size 1, but was significantly greater in the left hemisphere than the right for set sizes 3 (t(26) = 3.31, p < 0.01) and 6 (t(26) = 2.80, p < 0.01). These results confirm that the hemispheric asymmetry for VSTM emerges in IPS voxels that are strongly and symmetrically driven by contralateral visual representations.

In addition to areas IPS0–2, the memory task also activated a region lateral and anterior to areas of overlap between VSTM and visuotopic mapping. Although we did not observe visuotopic maps in this region, we cannot rule out the existence of visuotopic maps within this lateral and anterior IPS region (Konen and Kastner, 2008b). We performed an additional repeated-measures ANOVA for this region, referred to as lateral anterior IPS, to investigate effects of set size and stimulus location to characterize the involvement of this region in VSTM. A main effect of set size in this region approached significance (F(2,26) = 2.907, p = 0.08) but a significant interaction between set size and hemisphere was observed (F(2,26) = 7.077, p < 0.01). Consistent with VSTM activation in visuotopic IPS, paired comparisons demonstrated that this was the result of a significant effect of memory stimulus location for the left hemisphere (t(26) = 3.158, p < 0.01), but not for the right hemisphere. Thus, this lateral, anterior IPS activation also exhibited the same hemispheric asymmetry observed in IPS0–2, but we did not observe the symmetric visuotopic maps of the contralateral field.

Control measurements: eye position monitoring and response hand
It is difficult to imagine how eye position effects could account for the observed hemispheric asymmetry. Conceivably, subjects could shift their gaze rightward during remember-right trials, thus placing some targets contralateral to both right and left hemispheres, and yet hold central fixation during remember-left trials, resulting in only the right hemisphere receiving contralateral targets. Such an eye position asymmetry seems highly improbable; nevertheless, we examined this with two subjects. Eye position was monitored in the scanner while subjects performed remember-left, remember-right, and remember-full-screen conditions at set size 3. Subjects held fixation well across the conditions with no significant difference between remember-right and remember-left conditions (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). The subjects exhibited the same degree of parietal asymmetry that they did in the full experiments. Thus, eye position effects do not account for our observations.

It is also possible that the hand with which subjects responded influenced the spatial bias, resulting in a contralateral bias for the left but not the right hemisphere. Again, it is hard to conceive how right-handed button presses would influence only the ipsilateral hemisphere; nevertheless, we did investigate this issue. To determine whether subjects’ responses influenced the hemispheric spatial biases, we asked a set of our subjects (n = 3) to respond with the left hand for half of the runs. There was no interaction among the response hand, hemisphere, and location. Supplemental Figure 2 (available at www.jneurosci.org as supplemental material) displays the hemifield biases of each hemisphere for the subjects who only used their right hand to respond with those who switched hands. If anything, the hemispheric asymmetry is greater in those subjects who switched hands. At higher load, the switched-hand subjects tend toward an ipsilateral bias for the right hemisphere, compared with the strong contralateral bias for the left hemisphere. Thus, neither response hand nor eye position accounts for the hemispheric asymmetry observed in VSTM.

Discussion
We observed that activation in the human intraparietal sulcus reflected visual short-term memory performance across varying memory loads. As VSTM set size increased, BOLD signal and K score both increased and then reached a plateau. In contrast to previous studies (Todd and Marois, 2004; Xu and Chun, 2006; Xu, 2007), here the number of visual stimuli were constant across memory loads, confirming that set size effects in IPS are due to memory load, not to visual drive. IPS, the most prominent ana-
stimuli in either visual hemifield. Such hemispheric asymmetries, but exhibit mnemonic responses equally strongly for Right hemisphere IPS0–2 exhibit contralateral spatial maps for between the sensory and mnemonic spatial representations.

However, in the right hemisphere, we observe a major difference short-term memory representations appear to be consistent; hemisphere, the spatial representations of visual percepts and thus the activation reflects memory processes. In the left hemisphere, IPS0–2, but does not exhibit contralaterally biased visuospatial maps. This dual-population hypothesis implies that the right hemisphere responses should still be greater for contralateral than ipsilateral memory targets; however, our observations do not support this. Alternatively, right IPS could be driven by different sets of inputs that switch the effective spatial representation, depending on which are more strongly activated by the task. This dual-input hypothesis suggests that bottom-up visual stimulation arrives only from the contralateral field, whereas mnemonic inputs arrive from the entire visual field. This hypothesis is consistent with our observations and with single-unit primate studies that have shown that the same neurons can alter their firing patterns depending upon the task demands both in terms of representation of items (Toth and Assad, 2002) and the coordinate system in which objects are encoded (Battaglia-Mayer et al., 2003). These mnemonic inputs might come from ventral parietal and superior temporal regions, the primary loci of lesions underlying hemispatial neglect, or from lateral anterior IPS, which exhibits the same hemispheric asymmetry for VSTM as IPS0–2, but does not exhibit contralaterally biased visuospatial maps.

Hemispatial neglect is a complex syndrome that reflects a disruption in spatial attention in which contralesional neglect occurs almost exclusively after right hemisphere damage. To explain this hemispheric asymmetry, representational models of hemispatial neglect (Heilman and Van Den Abell, 1980; Mesulam, 1981, 1999; Pouget and Driver, 2000) have suggested that the right hemisphere should contain bilateral maps, whereas the left hemisphere should contain only contralateral maps. Prior neuroimaging studies have observed multiple spatial maps in the dorsal parietal attentional network (Sereno et al., 2001; Schluppeck et al., 2005). One possible explanation is that motor preparation, a confound in the delayed saccade task (Barash et al., 1991), may result in different patterns of topography compared with the memory component. Alternatively, the number of remembered stimuli may influence the topography, as we observed this asymmetry only when more than one target was remembered.

This hemispheric asymmetry could potentially be explained by either of two hypotheses regarding right hemisphere IPS. Right IPS, like many cortical areas, might contain two distinct neuronal populations: one population that performs mnemonic spatial indexing of the contralateral field and another population that performs nonspatial memory processes. In contrast, the left hemisphere might contain only contralateral spatial indexing neurons. This dual-population hypothesis implies that the right hemisphere responses should still be greater for contralateral than ipsilateral memory targets; however, our observations do not support this. Alternatively, right IPS could be driven by different sets of inputs that switch the effective spatial representation, depending on which are more strongly activated by the task. This dual-input hypothesis suggests that bottom-up visual stimulation arrives only from the contralateral field, whereas mnemonic inputs arrive from the entire visual field. This hypothesis is consistent with our observations and with single-unit primate studies that have shown that the same neurons can alter their firing patterns depending upon the task demands both in terms of representation of items (Toth and Assad, 2002) and the coordinate system in which objects are encoded (Battaglia-Mayer et al., 2003). These mnemonic inputs might come from ventral parietal and superior temporal regions, the primary loci of lesions underlying hemispatial neglect, or from lateral anterior IPS, which exhibits the same hemispheric asymmetry for VSTM as IPS0–2, but does not exhibit contralaterally biased visuospatial maps.

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As neglect symptoms recede, dorsal parietal activation increases. To explain this phenomenon, we suggest a network model of hemispatial neglect in which temporoparietal damage causes a disruption of inputs to the dorsal parietal regions that explicitly encode for spatial coordinates (Corbetta et al., 2005). Our findings are consistent with such a network model and provide fMRI evidence that hemispatial neglect mechanisms may be studied in normal human subjects. Our work suggests that one key is to use tasks that sufficiently tax subject capacities, such as requiring the coding of multiple stimuli within a visual hemifield.

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