The Role of the Parietal Lobe in Visual Extinction Studied with Transcranial Magnetic Stimulation

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Abstract

Interhemispheric competition between homologous areas in the human brain is believed to be involved in a wide variety of human behaviors from motor activity to visual perception and particularly attention. For example, patients with lesions in the posterior parietal cortex are unable to selectively track objects in the contralesional side of visual space when targets are simultaneously present in the ipsilesional visual field, a form of visual extinction. Visual extinction may arise due to an imbalance in the normal interhemispheric competition. To directly assess the issue of reciprocal inhibition, we used fMRI to localize those brain regions active during attention-based visual tracking and then applied low-frequency repetitive transcranial magnetic stimulation over identified areas in the left and right intraparietal sulcus to assess the behavioral effects on visual tracking. We induced a severe impairment in visual tracking that was selective for conditions of simultaneous tracking in both visual fields. Our data show that the parietal lobe is essential for visual tracking and that the two hemispheres compete for attentional resources during tracking. Our results provide a neuronal basis for visual extinction in patients with parietal lobe damage.

INTRODUCTION

Theories of visual attention have postulated the presence of two networks in the brain, located in the right and left parietal cortices, which drive attention to the corresponding contralateral hemispace. There has been a longstanding debate about the extent of interhemispheric competition between these two networks (Mesulam, 1999; Kinsbourne, 1977). In particular, an activation-orienting hypothesis has been suggested in which each hemisphere generates a contralateral attentional bias when stimulated by an external stimulus by inhibiting its contralateral counterpart (Reuter-Lorenz, Kinsbourne, & Moscovitch, 1990). In a seminal paper studying cats, Sprague (1966) introduced the concept of inhibition between homologous areas in the two hemispheres, but similar effects have rarely been reported in humans studies. However, evidence for interhemispheric competition has come from neuropsychological studies on parietal neglect patients, which suggest that visual extinction is likely caused by an imbalance between homologous areas in the two hemispheres (Corbetta, Kincade, Lewis, Snyder, & Sapi, 2005; Vandenbergh et al., 2005; Driver & Vuilleumier, 2001; Vuilleumier, Hester, Assal, & Regli, 1996). More specifically, a unilateral cerebral lesion may have the effect of disinhibiting the contralateral healthy hemisphere. Consequently, an exaggerated physiological contralateral bias becomes evident when two visual stimuli are simultaneously presented one in each hemifield. Under these conditions, the contralesional stimulus is extinguished by the ipsilesional stimulus (Duncan et al., 1999). Recent fMRI (Geng et al., 2006; Corbetta et al., 2005; Fink, Driver, Rorden, Baldeweg, & Dolan, 2000) and animal (Rushmore, Valero-Cabre, Lomber, Hilgetag, & Payne, 2006) studies support this notion. However, there is no direct evidence in healthy observers that homologous areas in both hemispheres interact in a mutually inhibitory manner under conditions of sustained attention. Understanding this mutual inhibition could have considerable implications for treatment and rehabilitative strategies for patients with parietal lobe lesions (Fecteau, Pascual-Leone, & Theoret, 2006; Oliveri & Caltagirone, 2006).

A particularly important aspect of sustained attention is the ability to keep track of multiple moving targets simultaneously. Tracking enables us to compute spatial relations between objects, to navigate in our environment while avoiding moving obstacles, and to monitor multiple moving objects, such as keeping track of our children at the playground. Studies investigating attentive tracking have suggested competing interactions between attentional resources in the two hemispheres. Patients with right parietal lesions are unable to track two targets when presented in the left and right visual fields simultaneously, while they perform normally when tracking one target either in the left or in the right visual field (Battelli et al., 2001). Although the deficit is more

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pronounced in the contralesional visual field, ipsilesional deficits in patients with right parietal lesion have also been reported (Battelli, Cavanagh, Martini, & Barton, 2003; Duncan et al., 1999). This finding is likely related to visual extinction, a common neuropsychological deficit after lesion to the parietal lobes, characterized by an inability to detect a contralesional target when another target is present in the ipsilesional field (Driver & Vuilleumier, 2001).

Within the parietal lobe, the intraparietal sulcus (IPS) may be particularly important for attentive tracking. fMRI studies have shown that the IPS is active during visual tracking (Culham et al., 1998), and the activity in the IPS (among other areas) varies with the change in attentional load (i.e., activity is higher as the number of targets to be tracked is increased). In contrast, the nonparietal area MT+, although strongly activated during visual tracking, does not change activity with attentional load, suggesting that it is activated by the motion per se but not by attentional allocation (Culham, Cavanagh, & Kanwisher, 2001). These results suggest that IPS activity is more directly involved in tracking performance.

The present study provides a direct measurement of the interaction between homologous brain areas using TMS as a “virtual lesion” technique to test the role of a brain area in a specific task. We demonstrate that a TMS-induced temporary inactivation over the IPS causes extinction-like behavior in normal subjects during attentional tracking. Experiment 1 shows that the IPS is directly involved in visual tracking, and that the two hemispheres are in competition when attention is split between the two hemifields. Experiment 2 rules out an alternative explanation based on the number of items tracked.

**EXPERIMENT 1: AN “EXTINCTION” EFFECT DURING FULL-FIELD ATTENTIONAL TRACKING**

**Methods**

**Observers**

We tested a total of 11 healthy participants (3 authors and 8 naïve observers) in Experiments 1 and 2. All participants were 26 to 38 years of age, and had normal or corrected-to-normal vision. All participants were checked for TMS exclusion criteria (Wassermann, 1998) and gave written informed consent to the study which had been approved by the Beth Israel Deaconess Medical Center’s Institutional Review Boards. The study was conducted in the Harvard–Thorndike General Clinical Research Center at BIDMC in order to provide the safest environment for the subjects. Eight individuals (4 men and 4 women) participated in Experiment 1, including two authors and six naïve observers.

**Stimuli**

Stimuli were displayed on a Macintosh G4 laptop using Matlab (MathWorks) in conjunction with the Psychophysics Toolbox (Brainard, 1997; Pelli, 1997). The displays consisted of a central fixation point (a black circle, radius = 0.15") and eight moving items (black circles, radius = 0.3") presented on a gray background. Four of the circles moved within a 9.4° × 9.4° region inset 2° to the left of fixation, and the other four moved within an equal size region inset 2° to the right of fixation. Items moved at a constant speed, repelled each other to maintain a minimum center to center spacing of 1.5° and “bounced” off of the invisible edges of the square region in which they moved.

**Procedure**

In the half-field condition, subjects tracked two target circles (in either the left or the right visual field), and in the full-field condition, subjects tracked four target circles (2 on each side simultaneously) while keeping their eyes fixated on the central fixation point throughout each trial in both conditions. These conditions were first equated for difficulty by adjusting the target speed in a preliminary threshold session.

**Threshold session.** On the first day of testing, we equated all conditions for difficulty by determining the speed at which observers could perform the task with 75% accuracy. At the beginning of each trial, the fixation point was presented for 1 sec, then eight circles appeared (4 on the left, 4 on the right), and a subset blinked off and on at 2 Hz for 2 sec to identify them as targets for tracking. Then all of the circles moved without crossing the midline for 3 sec. After the items stopped, one of them was highlighted in red (50/50 target or distractor). The observer then indicated by keypress whether the red item was a target or distractor, with a response time cutoff of 3 sec. After the response, the fixation point turned green for a correct response or red for an incorrect response for 1 sec. The next trial began immediately following this feedback.

Subjects first performed a practice block (16 trials) in which the circles moved at 5 deg/sec to learn the task, and then a test block in which the circles moved at one of eight different speeds (5–26 deg/sec) on each trial, with the speeds randomly interleaved over 7 blocks of 32 trials each. It took approximately 45 min to complete this session. This threshold procedure was used to identify the speed at which two targets or four targets could be tracked with 75% accuracy. Different threshold speeds were obtained for each individual subject, and different speeds were obtained for two half-field targets and for four full-field targets.

**Main experiment.** In the main experiment, the response cutoff time was reduced to 2.5 sec and the feedback duration was reduced to 0.5 sec, so that the duration of tracking could be increased to 6 sec. A 3-sec duration
was used in the threshold session to reduce the duration of that session.

There were two conditions presented randomly intermixed within each block of trials. In the **half-field** condition, two targets were presented either in the left or the right hemifield, whereas in the **full-field** condition, four targets were presented with two in each hemifield (Figure 1A). As described above, these conditions were matched for difficulty by setting the speed to the 75% accuracy threshold for each individual subject in each condition. In each block of trials, observers completed 48 trials (24 per condition) at their threshold speed. There were two experimental sessions (on separate days), each consisting of four blocks of trials, with a 20-min rest period in between each block. The first session (Day 1) had the following four blocks: (1) baseline, (2) TMS Site 1, (3) TMS Site 2, (4) baseline. The second session (Day 2) had the following four blocks: (1) baseline, (2) TMS Site 3, (3) TMS Site 4, (4) baseline. The four TMS sites were the left IPS, right IPS, left MT+, and right MT+. Within a session, subjects received TMS over one IPS site on one hemisphere, and one MT+ site on the opposite hemisphere, with the brain area (IPS/MT+) and hemisphere (left/right) orders counterbalanced across subjects.

### fMRI Protocol and Stimulation Site Reconstruction

Subjects were scanned on a 3-T Phillips Intera scanner at the Boston University Center for Biomedical Imaging. Functional imaging was conducted using an echo-planar imaging sequence (TR = 2000 msec, TE = 30 msec, flip angle = 90°, FOV = 192 mm, matrix size = 64 x 64). Scanned volumes were 30 oblique slices (3 x 3 mm in-plane resolution, 3 mm thick; interslice gap of 0.3 mm) covering the parietal and occipital lobes. The first four images of each 580 acquisition functional run were discarded to allow for T1 equalization. High-resolution T1-weighted structural scans (1 x 1 x 1 mm) were acquired for each subject to verify the anatomical locations of selected ROIs.

**IPS and MT+ localizer.** IPS and MT+ were localized using a block-design multiple-object tracking experiment similar to that used by Culham et al. (2001). Subjects were shown eight randomly moving dots presented in a fixed window approximately 15° x 15° of visual angle. Subjects were instructed to either passively view or actively track a varying number of dots (1, 2, or 4) cued at the beginning of the trial. Each of the four conditions...
was repeated four times in each run. Trials were 24 sec in length followed by a 12-sec rest period.

Acquired data were analyzed using the Brainvoyager QX (version 1.6). Functional data were motion corrected, slice scan time corrected, spatial smoothed (4 mm FWHM), and coregistered to high-resolution anatomical data. A model of the expected response of IPS and MT+ was generated using previously reported response properties for the two ROIs (Culham et al., 2001). ROIs were defined as a significant ($p < .001$, uncorrected) cluster of voxels in the parietal and medial temporal lobes. For all subjects, the most robust activation during multiple object tracking was observed in the parietal lobe bilaterally extending between the IPS and the postcentral sulcus.

High-resolution functional images were overlaid onto the anatomical images, obtained from the same fMRI experimental session using a frameless stereotaxy system (BrainSight, Rogue Research, Montreal, Canada). A three-dimensional anatomical reconstruction was used to visualize and measure the Talairach coordinates of the projected cortical target of the IPS and MT+ stimulation sites in all subjects (Figure 1C, example of one representative subject). The projected target of stimulation over the IPS corresponded to the posterior extent of the IPS, the same region implicated in visual tracking responsive in the fMRI study we conducted beforehand (Culham et al., 1998). The MT+ stimulation site projected onto the ascending extent of the lateral occipital sulcus, corresponding to reported location of MT+ as determined through ours and previous neuroimaging studies (Grossman, Battelli, & Pascual-Leone, 2005; Seiffert, Somers, Dale, & Tootell, 2003; Huk, Dougherty, & Heeger, 2002; Sunaert, Van Hecke, Marchal, & Orban, 1999).

Although fMRI revealed some individual variability in the functional activation during active tracking, for stimulation over the IPS, the average center of stimulation in stereotaxic space for all subjects corresponded to the following Talairach x, y, and z values: −24, −62, and 50 for the left and 25, −60, and 50 for the right hemisphere, respectively (Talairach & Tournoux, 1988). For MT+, the average coordinates were −36, −64, and 13 for the left and 46, −63, and 3 for the right hemisphere, respectively.

**TMS Protocol**

Subject performance was calibrated individually prior to any brain stimulation, as described above (Threshold Session). TMS was delivered using a MagStim stimulator (MagStim, Whitland, UK) and a commercially available 70-mm figure-of-eight Magstim stimulation coil. We applied a 10-min train of repetitive low-frequency (1 Hz) stimulation at 75% maximum stimulator output over one of the four brain sites, right IPS or MT+ and left IPS or MT+. Each subject underwent two testing sessions separated by at least 24 hr. To aid in brain site localization, subjects wore a Lycra swimmer’s cap on which the reference points for simulation were marked (see previous section for localization technique). The coil was held with the handle pointing backward toward the back of the head and positioned perpendicular to the stimulated region.

Immediately following the repetitive stimulation over the targeted brain site, subjects performed the visual tracking task (same task as the prestimulation baseline). The time required to perform the psychophysical task (approximately 10 min) is within that for which rTMS has been shown to have lasting effects in parietal regions (Hilgetag, Theoret, & Pascual-Leone, 2001). After completion of the task, observers rested for 15 min to allow complete recovery from the stimulation. Stimulation was then applied to the other brain site in the opposite hemisphere, and the subject again performed the task.

The order of brain site stimulation was counterbalanced across observers, which allowed us to control for any effects of “double-dose” of stimulation (there were none).

Following the second stimulation, task completion, and a 15-min recovery period, participants repeated the task to obtain a final poststimulation baseline. The same procedure was used in the second testing session.

**Results**

Only a slight reduction in speed (on average, 2 deg/sec) was necessary for subjects to track four targets in each field as well as two targets presented within a single half-field ($t < 1$ for targets tested in left and right visual fields, respectively; mean speed: full-field = 13.5 deg/sec; half-field = 15.5 deg/sec; Figure 1B). This is consistent with previous work demonstrating that it is possible to track four targets in separate hemifields with little or no cost over tracking two targets within a single hemifield, suggesting that tracking is functionally independent in the two visual fields (Alvarez & Cavanagh, 2005). The tracking task was then run at these threshold speeds throughout the entire TMS experiment.

Previous studies using off-line TMS with low-frequency stimulation have shown that the behavioral effect after TMS can last, on average, half as long as the time of the stimulation (Robertson, Theoret, & Pascual-Leone, 2003). Because we stimulated for 10 min, we divided our post-TMS data into two halves with the first and the last 5 min after stimulation analyzed separately. Of principal interest was whether performance in the contralateral visual field (opposite TMS) was different than in the ipsilateral visual field (same side as TMS), and in particular, whether there was a contralateral deficit (worse performance in the contralateral field).

TMS over the left IPS induced a significant contralateral impairment in the full-field tracking condition (track four targets, two in each hemifield), but only in the first 5 min post-TMS. Ipsilateral and contralateral performance in the full-field condition were significantly different from each other for the first 5 min after TMS ($t(7) = 3.72,$ Battelli et al. 1949)
EXPERIMENT 2: CONTROL FOR THE NUMBER OF OBJECTS TRACKED

Although we carefully matched the half-field and full-field conditions in terms of attentional load by equating task performance, the total number of targets tracked was greater in the full-field condition than in the half-field condition of Experiment 1. Thus, it remains possible that the number of targets tracked, independent of attentional load, accounts for the contralateral deficit observed in the full-field condition of Experiment 1. To address this possibility, we required observers to track either two or four targets all within the same half of the visual field. If increasing the number of targets from two to four results in a contralateral deficit during half-field tracking, then it would seem that it is the number of targets that is important, not whether tracking occurs in one hemifield or both hemisfields simultaneously. However, if we do not see a contralateral deficit for tracking four half-field targets, it would support the conclusion that the large contralateral deficit observed in the full-field condition of Experiment 1 was due to simultaneous tracking in both halves of the visual field.

Methods

All aspects of the method were identical to Experiment 1, except as noted here.

Observers

Eight individuals participated. Three were authors (two also participated in Experiment 1), and five were naïve subjects (three also participated in Experiment 1). We had functional localizers for all the subjects for the left IPS (the only site we stimulated in Experiment 2).

Stimuli and Procedure

The following modifications were made to the displays to allow for the presentation of eight items in each visual field so that up to four items could be tracked within a single half of the visual field. The displays consisted of a central fixation point (a black circle, radius = 0.15") and 16 moving items (black circles, radius = 0.3") presented on a gray background. Eight of the circles moved within a 17.1" × 17.1" region inset 2° to the left of fixation, and the other eight moved within an equal size region inset 2° to the right of fixation. Participants tracked either two or four targets in the right visual field or in the left visual field (i.e., always within half of the visual field). The same speed threshold procedure as in Experiment 1 was used to estimate the speed thresholds for tracking two or four circles within a hemifield. Within a trial, the items moved at a constant speed (eight speeds from 2 to 23 deg/sec for the threshold session, and at about 75% accuracy threshold speed for each individual participant in the main experimental session). Items repelled each other (minimum center to center spacing = 2.5") and “bounced” off of the invisible edges of the square region in which they moved.

We tested only the half-field condition, and the targets were always presented either in the left or in the right visual field, and separate 75% correct speed thresholds were measured for tracking two and tracking four targets to equate the two conditions for difficulty or attentional load.

TMS Protocol

Each subject underwent one testing session during which we applied TMS over the left IPS at the same intensity and frequency as in Experiment 1.

Results

A significant reduction in speed (9 deg/sec on average) was necessary for subjects to track four targets in a half-field as well as two targets within a half-field [2 half-field targets = 17.8 deg/sec; 4 half-field = 8.8 deg/sec; t(7) = 10.27, p < .001]. This is consistent with previous work, which suggests that targets within a hemifield compete for attentional resources (Alvarez & Cavanagh, 2005), and that more targets can be tracked when the target speed is reduced (Alvarez & Franconeri, 2007). The tracking task was then run at these threshold speeds throughout the entire TMS experiment.

With the speed adjustment, baseline performance in the main experiment was well matched for tracking two
Figure 2. Subjects’ performance following left (A) and right (B) IPS stimulation, and following left (C) and right (D) MT+ stimulation for the first 5 min post-TMS. Black bars indicate ipsilateral performance (same side as TMS) and white bars contralateral performance (opposite side of TMS). Error bars represent one within-subject standard error of the mean, calculated with Cousineau’s (2005) modification of Loftus and Masson’s (1994) method. Asterisks indicate where performance was significantly different between the contralateral and ipsilateral conditions (only during full-field tracking following left or right IPS stimulation). Insets show individual-subject difference scores (contralateral − ipsilateral), where positive values indicate a relative advantage in the contralateral field, and negative values indicate a relative deficit in the contralateral field. Authors are shown with dashed lines (A = 2), and naı ¨ ve observers with solid lines (N = 6).
or four targets within a hemifield (79% for 2 targets, 79%
for 4 targets, \( t < 1 \)). We applied TMS to the left IPS,
where we saw the largest, most robust contralateral
effect in Experiment 1, and the question was, do we see
a large contralateral deficit whenever four targets are
tracked, even if all of the targets are within a single half
of the visual field? The answer appears to be no (Fig-
ure 3). None of the ipsilateral versus contralateral differ-
ences were significant in the first 5 min or in the second
5 min after stimulation (all \( ts < 1 \), all \( ps > .59 \)). Part-
icularly notable is the lack of difference in the first 5 min
for the contralateral condition (Figure 3, compare white
bars for 2 and 4 targets): Performance was comparable
for two half-field targets and four half-field targets (77%
vs. 77%, \( t < 1 \)). This is in stark contrast to the results
of Experiment 1, where in the first 5 min the difference
in the contralateral condition between two half-field
targets and four full-field targets was large and significant
[91% vs. 51%, respectively; \( t(7) = 3.76, p < .01 \)]. Given
that a contralateral deficit does not arise under a single
hemifield load of four targets, the contralateral deficit
for four full-field targets in Experiment 1 does not ap-
pear to be due to the number of targets. Instead, it ap-
pears to be due to the full-field, bilateral nature of the
tracking task in that condition.

DISCUSSION

Our results show that low-frequency rTMS over the IPS
impairs performance on attention-based visual tracking
of moving stimuli, whereas TMS over the motion area
MT+ does not alter behavioral performance. These find-
ings are consistent with those of Culham et al. (1998),
who used fMRI to show that the IPS is selectively ac-
tivated during attentive tracking, whereas activity in
MT+ does not distinguish between active tracking and
passive viewing of the same moving stimuli. An alterna-
tive explanation of our results could be a disruption of
eye movements after TMS over the IPS. However, this is
very unlikely, as a disruption of eye movements should
have had a similar effect for both the full-field and half-
field tracking condition, but this was not the case in
our experiment. Several fMRI studies have also shown
that MT+ is robustly activated when subjects are asked
to perform visual pursuit tasks (Dukelow et al., 2001;
Barton et al., 1996), however, only the IPS is selectively
activated during active tracking (Culham et al., 1998),
and this ability was affected by TMS over the IPS in
Experiment 1.

A Neural Model for Attentive Tracking

Although we found that the IPS plays a selective role
during attentive tracking, TMS had an effect only during
full-field tracking, and not during half-field tracking. This
difference cannot be due to differences in task difficulty
between the full-field and half-field tasks because we
psychophysically measured the subjects’ speed sensitiv-
ity before the TMS experiment and equated the two
conditions for difficulty. Moreover, Experiment 2 ruled
out the number of targets tracked as a possible expla-
nation of the deficit we found during full-field tracking as
TMS had no effect when subjects were asked to track
four targets within a single hemifield. So what is the dif-
ference between the half-field and full-field conditions?

Previous cognitive research on attentive tracking has
found evidence for independent tracking of targets in
the left and right visual hemifields (Alvarez & Cavanagh,
2005), as if independent resources were operating in
each visual field. Similar findings using different exper-
imental paradigms have also been found in normal sub-
jects (Kingstone, 2004; Corballis & Gratton, 2003) as well
as in commissurotomy patients (Holtzman & Gazzaniga,
not make any claims about the underlying neural basis of
hemifield independence, the most obvious possibility
was that the left and right cerebral hemispheres operate
independently in this task. From this view, the left
parietal lobe would be solely dedicated to tracking tar-
gets in the right visual field, and the right parietal lobe
would only track targets in the left visual field. The cur-
rent study rules out this simple possibility and suggests
that the functional independence in tracking between

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Figure 3. Subjects’ performance for two and four targets in one-half
of the visual field following left IPS stimulation in Experiment 2, for the
first 5 min after stimulation. Black bars indicate ipsilateral performance
(same side as TMS) and white bars contralateral performance
(opposite side of TMS). Error bars represent one within-subject
standard error of the mean, calculated with Cousineau’s (2005)
modification of Loftus and Masson’s (1994) method. The
inset shows individual-subject difference scores (contralateral –
ipsilateral), where positive values indicate a relative advantage in
the contralateral field, and negative values indicate a relative deficit
in the contralateral field. Authors are shown with dashed lines (A = 3),
and naïve observers with solid lines (N = 5).
the hemifields must arise from a more complex network of competitive interactions between the left and right cerebral hemispheres.

The current results, and the functional independence observed in attentive tracking (Alvarez & Cavanagh, 2005), can both be accounted for by the following neural model of attentive tracking. Each IPS is able to control the selection of targets in both the ipsilateral and contralateral visual field. However, in normal conditions, inhibitory connections between the left and right IPS reduce the amount of ipsilateral processing performed by each hemisphere.

For example, when a single target was being tracked in the right visual field, the left IPS would strongly inhibit the right IPS. In the baseline condition of our experiment (without TMS), the left IPS would serve as the primary control center for tracking the target, and would actively inhibit the right IPS. There would be little or no change from this situation following TMS to the right IPS because the right IPS is already inhibited without TMS. However, following TMS to the left IPS, the right IPS would be released from inhibition and would then serve as the primary control center for tracking the target, whereas the left IPS would make a reduced contribution.

When tracking targets in both visual fields, each IPS would inhibit ipsilateral processing of the other IPS. Thus, in the baseline condition, the left IPS would be the control center for tracking the target in the right visual field, and the right IPS would be the control center for tracking the target in the left visual field, where neither IPS would contribute to tracking in the ipsilateral field. Following TMS to the left IPS, the amount of inhibition on the right IPS would be reduced. However, given that there are targets in both visual fields, the right IPS is dedicated to tracking the target in the left visual field, and therefore, cannot compensate for the reduced functioning of the left IPS. Behaviorally, this results in an inability to track the target in the right visual field. In the case of TMS to the right IPS, the pattern would be reversed.

This model also accounts for the functional independence observed in the Alvarez and Cavanagh (2005) study. Under normal conditions, due to inhibitory connections between the left and right IPS, each IPS contributes primarily to tracking targets in the contralateral visual field. Thus, when tracking a target in the left visual field (controlled primarily by left IPS), there is little to no cost to adding a second target to the right visual field (controlled by the right IPS), and vice versa.

This model of attentive tracking has two key components: (1) The IPS in each hemisphere can control the selection of targets in either the ipsilateral or the contralateral visual field, and (2) under normal conditions inhibitory processes reduce the amount of ipsilateral processing by each IPS. Future research will be required to determine exactly how these inhibitory effects are realized. There could be direct links between the left and right IPS via the corpus callosum, in which case the inhibition would be from IPS to IPS, and would result in reduced activity in the inhibited IPS. Alternatively, there could be projections from lower-level visual areas to the contralateral IPS, and the inhibition could occur by limiting the amount of communication over these connections. In this case, the amount of IPS activity would not be reduced by inhibition, but activity in areas which communicate to the contralateral IPS would be reduced (Geng et al., 2006).

**MT+ and Attentive Tracking**

Several recent studies on nonhuman primates have all shown that neurons in middle temporal area (MT) are modulated by attention (Recanzone & Wurtz, 2000; Treue & Martinez Trujillo, 1999; Treue & Maunsell, 1996). However, the attentional effects are always relatively small (Seidemann & Newsome, 1999; Ferrera & Lisberger, 1997; Newsome, Wurtz, & Komatsu, 1988) and the attentional modulation strength depends on stimulus attributes such as contrast and size (Pack, Hunter, & Born, 2005; Martinez-Trujillo & Treue, 2002). In particular, neurons in area MT fire more strongly to large low-contrast stimuli (Pack et al., 2005), whereas in our study, we used small high-contrast stimuli and this could have contributed to a moderate involvement of the MT+ area in our visual tracking task.

We are not claiming that MT+ is not involved in tracking motion, however, it might well be that its contribution is solely due to the presence of a strong motion signal, as has been argued previously (Culham et al., 2001).

**The Neuronal Basis of Visual Extinction**

The contralateral deficit we found in our study appears to manifest only under conditions in which there are competing tasks in the two hemifields, as in extinction with neglect patients (Mesulam, 1999).

In our experiment, there was a trend for TMS to exert a stronger effect over the left IPS than over the right. Indeed, for left TMS, subjects' performance dropped to chance level in the full-field condition during the first 5 min after TMS. Our results lend support to Kinsbourne's (1977) biased hemispheric-competition model of visuospatial attention, which postulated a stronger contralateral bias of the left hemisphere than the right hemisphere. In addition, many studies on visual neglect have emphasized that the rightward bias of the left attentional system is stronger than the leftward bias of the right attentional system (Bartolomeo & Chokron, 2002). Our results are consistent with this view because TMS over the left IPS caused a more severe temporary drop in contralateral performance than TMS over the right IPS. Finally, recent studies have suggested a left hemisphere superiority for selective attention (Chokron, Brickman,


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