BOLD Repetition Decreases in Object-Responsive Ventral Visual Areas Depend on Spatial Attention

E. Eger,1,2 R.N.A. Henson,1,2 J. Driver,1,2 and R. J. Dolan1
1Wellcome Department of Imaging Neuroscience, London WC1N 3BG; and 2Institute of Cognitive Neuroscience, London WC1N 3AR, United Kingdom

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INTRODUCTION

Studies of visual object processing with functional magnetic resonance imaging (fMRI) have increasingly used blood oxygenation level–dependent (BOLD) repetition-suppression effects, during adaptation or “priming” paradigms, as a tool beyond basic subtraction designs for studying the neural representation of visual stimuli (e.g., see Grill-Spector et al. 1999; Henson 2003; Naccache and Dehaene 2001). Such repetition effects can provide information about the level of representation achieved in a given cortical region, by determining whether these effects generalize across specific transformations in the repeated stimulus, such as size and viewpoint in the case of object processing (Grill-Spector et al. 1999; James et al. 2002; Vuilleumier et al. 2002). fMRI studies using this approach have some parallels with the literature on behavioral priming effects [measured, e.g., with reaction times (RTs)] for objects repeated across various transformations (e.g., see Biederman and Kalocsai 1997; Bülthoff et al. 1995). It has been suggested that BOLD repetition decreases might provide a neural substrate for behavioral priming effects (Wiggs and Martin 1998), although caution should be exercised in assuming a direct relationship (Henson and Rugg 2003).

A question often addressed in behavioral priming studies, but rarely in fMRI work to date, is the extent to which specific repetition effects depend on attention. Some behavioral studies show that, although explicit awareness of object identity may depend on attention, some priming effects may nevertheless be found from unattended objects, at least in some circumstances (e.g., deSchepper and Treisman 1996; Tipper 1985). However, behavioral work addressing this issue has led to variable outcomes. In some studies, “positive” priming (i.e., shorter RTs) was observed from putatively unattended objects (e.g., Stankiewicz et al. 1998), whereas others reported “negative” priming (longer RTs) when a previously unattended object was subsequently attended (e.g., Tipper 1985; Tipper and Driver 1988). In other instances, no behavioral priming was observed in the absence of attention (e.g., Crabb and Dark 1999). Researchers using purely behavioral measures have begun to suspect that the variable behavioral priming observed might reflect the combined expression of several different types of neuronal “priming” effects.

There has been surprisingly little fMRI work to date on how BOLD repetition effects for visual objects depend on attention, although in principle this might disentangle different types of priming effects that could become superimposed in purely behavioral measures. Accordingly, we used event-related fMRI here to examine the influence of spatial attention on object priming as tested by BOLD repetition decreases. Our design included repetition of the identical object image, but also repetition across a mirror reversal that substantially changed the retinal image. Our main finding is that whereas fMRI repetition-suppression effects were reliably found from an attended object for a subsequent probe (in both original and mirror conditions), these effects were eliminated for unattended objects. This demonstrates a strong dependency of BOLD repetition suppression on spatial attention to the object at initial exposure.

METHODS

Subjects and imaging

Thirteen healthy right-handed volunteers (7 male/6 female, mean age 27.7 ± 5.8 yr) had normal or corrected vision. The study was approved by the Joint Ethics Committee of the National Hospital and Institute of Neurology, London. Functional images were acquired on a 4.0-Tesla whole-body magnetic resonance imaging scanner using a 9-cm-diameter circular array head coil. A whole brain volume was acquired using a T2*-weighted echo-planar imaging sequence (TR/TE = 4000/55 ms; slice thickness, 4 mm; in-plane resolution, 3 × 3 mm). Electrooculogram monitoring was performed using subdermal Ag-AgCl electrodes placed nasion, left and right outer canthus, and left and right lateral canthus. fMRI analyses were performed using SPM99 software (Welcome Department of Cognitive Neurosciences, London WC1N 3AR). Functional images were spatially realigned using a least-squares approach to remove head movements. The realigned images were then spatially normalized into the standard MNI (Montreal Neurological Institute) 152 template image as defined by a linear transformation. The images were then smoothed with an 8-mm half-width Gaussian kernel to increase the signal-to-noise ratio. Stimuli were presented on a rear-projection screen at a distance of 1 m. Staircase methods and a 600-ms interstimulus interval were used to minimize eye movements during the experiment. Subjects were instructed to make size judgments for probe objects as quickly and accurately as possible. Reaction times were measured from the onset of the probe image to the subject’s button press. The fMRI experiment consisted of four conditions: unattended primes (i.e., attended probes), attended primes (i.e., attended probes), unattended probes, and attended probes. Each condition was presented in blocks of 40 trials. Each condition included 50% of trials in which the probe was identical to the prime and 50% of trials in which the probe was a mirror image of the prime. Blocks consisted of 10 trials for each of the four conditions. Trials were presented in 20 successive blocks for a total of 800 trials. Subjects were instructed to press a Flannigan button (Model 3500, Superior Electric Corporation, Bridgeport, CT) to indicate whether the probe was the same object as the prime when the probe is the same object as the prime. Reaction times were measured from the onset of the probe image to the subject’s button press. The fMRI experiment consisted of four conditions: unattended primes (i.e., attended probes), attended primes (i.e., attended probes), unattended probes, and attended probes. Each condition was presented in blocks of 40 trials. Each condition included 50% of trials in which the probe was identical to the prime and 50% of trials in which the probe was a mirror image of the prime. Blocks consisted of 10 trials for each of the four conditions. Trials were presented in 20 successive blocks for a total of 800 trials. Subjects were instructed to press a Flannigan button (Model 3500, Superior Electric Corporation, Bridgeport, CT) to indicate whether the probe was the same object as the prime when the probe is the same object as the prime.
a 3-Tesla MR system with standard head coil (Siemens Allegra, Erlangen, Germany) as T2*-weighted echoplanar image (EPI) volumes every 2 s (TE 30 ms, 32 transversal slices with 30° anterior–posterior angulation, voxel size 3 × 3 × 2 mm, distance factor 75%, approximate slice position in Fig. 2).

**Stimuli**

A total of 120 digitized photographs or realistic shaded renderings of asymmetrical familiar objects served as stimuli. These came from different sources (Object Databank: http://www.cog.brown.edu/object_db; MasterClips image collection: http://www.imisoft.com; and Hemera Photo Clipart: http://www.hemera.com). Images were converted to grayscale on a white background. Object categories were household items (31), animals (18), tools (13), vehicles (12), musical instruments (6), food (6), and others (34). Stimuli were divided into 6 sets of 20 objects each, approximately equated for category content. They were rotated around the 6 experimental conditions across subjects for counterbalancing. An additional 120 objects of comparable categories served as “fillers” (see following text) in prime trials. During a “localizer” scan to determine areas responding visually to the objects, the objects from the priming study were presented in addition to “noise” versions of these stimuli (created by adding Gaussian white noise to the Fourier phase angles).

**Experimental protocol and task**

Stimuli were back-projected onto a translucent screen located approximately 60 cm above the subjects’ head and viewed by a mirror on the head coil. Objects subtended approximately 4.5°.

In 3 sessions of about 12 min each, trials started with a red cued square of 5° for 100 ms, centered at an eccentricity of 5.5°, unpredictably to left or right of central fixation (Fig. 1). This was followed by presentation of 2 concurrent objects for 100 ms, one inside the square, the other on the opposite side of fixation (cue plus display duration was thus 200 ms, which should be too brief for deliberate saccades to the cued object while it was still displayed1). Subjects were instructed to attend covertly to the object in the cued location and perform a speeded size-judgment task (press one of 2 buttons depending on whether in real life the object would fit into a shoebox). Subjects were instructed to ignore the object presented on the uncued side. After an interstimulus interval of 3 s, a single probe object appeared centrally for 200 ms and subjects again performed the size judgment. The probe was either a new object, the same as the attended (or unattended) object in the preceding prime display, or a mirror-reversed version of this. There were 6 experimental conditions: primed attended; primed unattended; primed mirror image attended; primed mirror image unattended; unprimed (baseline for attended); unprimed (baseline for unattended). The 2 baselines accounted for the fact that when the attended object primed the subsequent probe, there was a congruent size-judgment response to prime and target (as in the attended baseline also), whereas when the unattended image primed the subsequent probe, responses to prime and target were incongruent, as in the unattended baseline (see Fig. 1). All experimental conditions where randomly intermingled with an intertrial interval of 3 s. Filler objects (occupying the opposite side to the potentially probe-related object in prime displays) were randomly assigned to each trial.

Our analysis focused on immediate repetition effects (within each trial pair). Twenty different objects were used for each of the 6 experimental conditions. The trials for each object were presented twice within one session and therefore 6 times altogether. Any additional long-lag between-trial repetition effect that might in principle have occurred within a condition should apply equally to all experimental conditions and is thus unlikely to systematically confound our comparisons; session was nevertheless considered as a factor in some of our analyses.

After the main experiment, an 8-min “localizer” session mapped regions responding to the visual objects (vs. noise patterns). Subjects passively viewed 10 blocks of 12 object pictures, alternating with 10 blocks of 12 phase randomized “noise” patterns, all stimuli being presented for 500 ms with a stimulus onset asynchrony (SOA) of 1 s and blocks separated by baseline periods of 12 s. Stimulus presentation and recording of behavioral responses used in-house software (Cogent 2000v1.24: http://www.vislab.ucl.ac.uk/Cogent) within Matlab6.

**Image processing and data analysis**

Analysis of imaging data used SPM2 (http://www.fil.ion.ucl.ac.uk/spm2.html). Image preprocessing included realignment and unwarping, slice-time correction with middle slice as reference, spatial normalization (EPI-template), and spatial smoothing (10 mm full width at half-maximum Gaussian kernel). Responses to probe displays for the 6 experimental conditions, and separately to the 2 types of prime display for cue left or cue right, were modeled by delta functions convolved with a canonical hemodynamic response function (HRF) and its temporal derivative. Incorrect responses to probes constituted a separate regressor. Parameter estimates for all regressors were obtained by maximum-likelihood estimation, while using a temporal high-pass filter (cutoff 128 s), and modeling temporal autocorrelation as an AR(1) process. All statistical comparisons were performed as random-effects group analyses across the 13 subjects, using one-sample t-tests on images of the contrasts of HRF parameter estimates.

We report activations significant at P < 0.001, uncorrected, within the mask of object-responsive areas (see Fig. 2A), where object-responsive areas were defined by the group contrast of objects > phase-randomized noise at P < 0.001, uncorrected, from the localizer scans. In an additional region-of-interest (ROI) analysis, parameter estimates were extracted and averaged across voxels from spherical regions of interest of 10-mm radius, centered on left and right lateral occipital (LO) and fusiform response maxima, defined individually from each subject’s localizer contrast (objects > noise).
RESULTS

Behavior

Table 1 gives mean probe RTs and percentage correct in the size-judgment task for the 6 conditions during scanning, showing shorter reaction times for conditions with attended related primes. RTs for primed conditions were subtracted from their respective baseline and analyzed in a 2×2 ANOVA (original/mirror × attended/unattended). This yielded a significant main effect of attention on priming \[ F(1,12) = 35.5, P < 0.0001 \]. The main effect of view \[ F(1,12) = 0.003 \] and the interaction \[ F(1,12) = 3.7 \] were nonsignificant, indicating equivalent priming for repetition of the same image and for mirror images, but a critical dependency on attention. Mean priming was 98 ms from attended objects \[ t(12) = 7.2, P < 0.001 \] but only 5 ms from unattended objects \[ t(12) = 0.6, \text{n.s.} \]. Analysis of error rates yielded no significant effects, but confirmed that the priming effects in RTs for attended object were not due to speed/accuracy trade-offs. Figure 4A shows individual RT priming effects for all 13 subjects, with 95% confidence intervals for latter overlapping with zero. Thus the absence of priming from unattended objects was not caused by some subjects showing strong positive priming and others showing strong negative priming; instead, the group results are representative of most individuals.

Imaging

We were specifically interested in BOLD repetition-suppression effects in brain regions showing significant responses to objects (as defined by the localizer, which activated lateral occipital and ventral occipitotemporal regions bilaterally; see Fig. 2). To restrict further interrogation of the data to these regions, we masked and small-volume–corrected all further comparisons using the localizer contrast at \( P < 0.001 \), uncorrected.

Repetition decreases in the event-related BOLD response as a function of object repetition were first investigated by the simple effects for each of the primed conditions subtracted from its respective baseline condition. Analyses for primed attended original and mirror images each yielded a similar pattern of repetition decrease in lateral occipital and fusiform regions bilaterally (Fig. 2, Table 2). By contrast, the same comparison for primed unattended original images and for unattended mirror images showed no effects, not even at the lower threshold of \( P < 0.01 \), uncorrected. To directly compare repetition effects from attended and unat-

<table>
<thead>
<tr>
<th>Experimental Condition</th>
<th>Attended</th>
<th>Unattended</th>
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<tbody>
<tr>
<td>Original Image</td>
<td>609 ± 30</td>
<td>713 ± 33</td>
</tr>
<tr>
<td>Mirror Image</td>
<td>625 ± 31</td>
<td>699 ± 25</td>
</tr>
<tr>
<td>Unprimed (Baseline)</td>
<td>716 ± 26</td>
<td>712 ± 31</td>
</tr>
<tr>
<td>Original Image</td>
<td>713 ± 33</td>
<td>712 ± 31</td>
</tr>
<tr>
<td>Mirror Image</td>
<td>699 ± 25</td>
<td>859 ± 2.5</td>
</tr>
<tr>
<td>Unprimed (Baseline)</td>
<td>712 ± 31</td>
<td>842 ± 2.9</td>
</tr>
</tbody>
</table>

TABLE 1. Behavioral data (mean of 13 subjects ± SE)

FIG. 2. Imaging findings from group analyses. A: mask of object responsive regions used for small-volume correction, which was derived from the contrast of objects > phase-randomized noise, at \( P > 0.001 \), uncorrected. Tilted lines indicate the approximate slice position during scanning. B: regions expressing significant repetition decreases for attended original images. C: attended mirror images, in contrast with the unprimed baseline condition. For display purposes, findings were color-coded at \( P < 0.005 \), uncorrected (masked inclusively by localizer at \( P < 0.001 \)) and superimposed on the normalized structural scan of one participating subject. In both the original and mirror-image conditions, similar repetition decreases were found in lateral occipital and fusiform regions bilaterally.
Repetition decreases for priming from attended objects (separately for original and mirror images)

<table>
<thead>
<tr>
<th>Contrast</th>
<th>Region</th>
<th>MNI Coordinates</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Attended original image</strong></td>
<td>Lateral occipital and fusiform L</td>
<td>39 -87 -6</td>
<td>3.88</td>
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<tr>
<td></td>
<td>and fusiform R</td>
<td>45 -66 -21</td>
<td>3.58</td>
</tr>
<tr>
<td></td>
<td>Lateral occipital</td>
<td>-36 -90 12</td>
<td>4.30*</td>
</tr>
<tr>
<td></td>
<td>and fusiform L</td>
<td>-42 -72 -21</td>
<td>3.61</td>
</tr>
<tr>
<td></td>
<td>Lateral occipital R</td>
<td>-30 -45 -24</td>
<td>3.37</td>
</tr>
<tr>
<td></td>
<td>Posterior IPS L</td>
<td>-30 -90 27</td>
<td>3.04</td>
</tr>
<tr>
<td></td>
<td>IPS L</td>
<td>-24 -57 45</td>
<td>3.01</td>
</tr>
<tr>
<td><strong>Attended mirror image</strong></td>
<td>Posterior to midfusiform R</td>
<td>36 -60 -3</td>
<td>4.38*</td>
</tr>
<tr>
<td></td>
<td>Lateral occipital</td>
<td>48 -72 -24</td>
<td>3.37</td>
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<tr>
<td></td>
<td>and fusiform L</td>
<td>36 -54 -21</td>
<td>2.85</td>
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<td></td>
<td>Midfusiform R</td>
<td>-42 -54 -15</td>
<td>3.94</td>
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<td></td>
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<td>-36 -54 -27</td>
<td>3.52</td>
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<tr>
<td></td>
<td>and fusiform L</td>
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<td>3.44</td>
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<tr>
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<td>-30 -93 9</td>
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<td></td>
<td>IPS L</td>
<td>-45 -78 9</td>
<td>3.75</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-27 -78 30</td>
<td>3.01</td>
</tr>
</tbody>
</table>

Regions that survived correction for multiple comparisons at voxel level for the volume of areas defined by the localizer at $P < 0.05$ are indicated with *. Other regions are reported at a threshold of $P < 0.001$, uncorrected, in both priming contrast and the localizer mask.

For priming from attended objects, we further investigated these according to a $2 \times 2$ factorial design (original/mirror image × attended/unattended) after subtraction from the respective baseline conditions, analogous to the analysis of the behavioral data. The main effect of attention (repetition decrease for attended $> \text{unattended}$) gave significant results in lateral occipital and posterior to midfusiform regions bilaterally (Fig. 3, Table 3), and thus confirmed the pattern observed in the simple-effect comparisons. No voxels showed stronger repetition decreases for unattended than attended images.

To assess effects of view, we first subtracted the primed original from primed mirror-image conditions. This comparison yielded no effects at $P < 0.001$, uncorrected, nor did the reverse comparison (i.e., primed mirror image $> \text{primed original}$).

Finally, we performed ROI analyses on lateral occipital and fusiform maxima as defined individually by each subject’s localizer scan. Unlike the group SPM approach, this analysis can take into account any variability in the location of object-responsive regions across subjects, and could potentially be more sensitive to small levels of signal change that are coherent across these individually defined regions, as it aggregated over multiple voxels in those regions (see METHODS).

Data from the 4 ROIs for each subject are plotted in Fig. 4B in addition to the individual behavioral priming effects. The ROI fMRI analysis confirmed a main effect of attention on BOLD repetition suppression [$F(1,12) = 25.09, P = 0.0001$, whereas there was no difference across the regions tested, as indicated by nonsignificant interactions with region [$F(1,12) = 0.044$] or laterality [$F(1,12) = 0.005$]. Furthermore, the attentional influence on repetition effects did not interact with experimental session [$F(1,9,22.8) = 0.284$]. Tests for simple effects of repetition carried out separately for attended or unattended objects revealed significant BOLD repetition suppression for attended objects [$F(1,12) = 10.67, P = 0.007$] but not for unattended objects. If anything, there was a marginal tendency for the opposite pattern of BOLD repetition increases with unattended objects [$F(1,12) = 4.84, P = 0.048$]. We shall not base any strong conclusions on that outcome, which requires further replication. For now, the important point is that the results clearly show no evidence for BOLD repetition suppression.

The view × attention interaction testing for larger differences in priming between attended and unattended original than mirror images yielded nothing except an activation at $P < 0.001$, uncorrected, in the left intraparietal sulcus, but this did not survive correction for multiple comparisons across the volume of interest, and in any case fell outside the ventral visual areas which our hypotheses concerned. The reverse interaction contrast produced no effects at the same threshold.

Note that the vast majority of subjects all showed positive behavioral priming from attended objects, but no significant behavioral priming (confidence intervals overlap with zero) from unattended objects. Analogously, the majority of subjects showed fMRI repetition-suppression effects from attended objects, but not from unattended objects.

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**FIG. 3.** Imaging findings from group analyses. A: shows are regions expressing larger repetition decreases for priming from attended than from unattended images collapsed across view. After subtraction from the appropriate unprimed baseline conditions. For display purposes, findings were color-coded at $P < 0.001$, uncorrected (masked inclusively by objects $> \text{phase-randomized noise at } P < 0.001$) and superimposed on the normalized structural scan of one participating subject. B: plots of responses (contrasts of parameter estimates) for the 4 priming conditions from the maxima in left and right lateral occipital and fusiform cortices (OA, priming from original image attended; OU, priming from original image unattended; MA, priming from mirror image attended; MU, priming from mirror image unattended). Effects are displayed after subtraction from the unprimed baseline conditions; positive values thus denote repetition decreases. Error bars represent residual error from a one-way ANOVA on the 4 contrasts.
suppression from unattended objects, given that any residual tendency was actually in the opposite direction.

To summarize, attended objects produced reliable repetition effects in BOLD for both the same image and for repetition across a mirror transform, in lateral occipital and fusiform regions bilaterally. In striking contrast, no such repetition decreases were observed from unattended objects, demonstrating the dependency of these effects on attention, as confirmed by the main effect of our attentional factor.

**Discussion**

The degree to which priming-related repetition decreases in BOLD depend on attention has received surprisingly little research in previous imaging studies (although see Murray and Wojcikulik 2004). Some studies have found that behavioral priming accompanied by fMRI repetition suppression can occur with masked priming, and thus without explicit awareness of repetition (e.g., Dehaene et al. 2001). However, that situation is quite different from the spatial attention manipulation used here, where 2 objects were presented concurrently to “compete” for attentional resources. In the present situation, we demonstrated for the first time that BOLD repetition decreases for objects can be entirely dependent on spatial attention to the prime object.

One previously published study (Bentley et al. 2003) did assess fMRI repetition effects as a function of spatial attention. It reported that some repetition decreases in ventral occipito-temporal cortex may occur for unattended, as well as attended faces. Faces rather than objects were used, and the apparent contrast with our finding may also relate to a weaker attentional manipulation in that study (e.g., longer presentation duration, less spatial separation) that might conceivably allow attentional resources to be deployed even to task-irrelevant stimuli. That study compared only second with first presentation of a given face, without including the appropriate baseline of unprimed items for unconfounding repetition, time, and order effects as here.

Another recent fMRI study (Murray and Wojcikulik 2004) investigated adaptation in LO while directing attention either to schematic “objects” or to color changes at fixation. The effect of changes in object orientation on BOLD repetition suppression was reduced when attending to the fixed colors, but repetition decreases were not completely abolished. The difference between the “unattended” results for that study and ours might relate to the strength of the attentional manipulations (Lavie 1995), to the stimuli used, or to the involvement of spatially cued attention (here) versus object-based attention; these are now issues for future work.

Our own behavioral and fMRI repetition effects clearly both depended strongly on spatial attention, but not on view (original or mirror image). This contrasts with the proposal by Stankiewicz et al. (1998) that same-view priming should be found without attention, whereas only priming across mirror-image views should not. Our experiment differs from the purely behavioral work of Stankiewicz et al. in the format of the objects used (shaded grayscale instead of line drawings), and in the task used (size judgment instead of naming). Task factors can sometimes influence the outcome of behavioral priming, for instance with respect to the degree of image specificity (Bruce et al. 2000), although this does not undermine our demonstration that BOLD repetition-suppression effects can depend strongly on spatial attention.

The present fMRI repetition effects from attended primes generalized across view (for original and mirror images) in fusiform and lateral occipital cortex bilaterally (see Fig. 2), and did not interact with laterality in the ROI analysis. This differs from the proposal by Vuilleumier et al. (2002) that fMRI repetition effects may generalize across view changes more in the left than in the right hemispheres. However, these authors used more dissimilar and heterogeneous views (involving both plane and depth rotations), unlike the strict mirror imaging used here. Some models of object recognition that do not incorporate invariance to depth rotation can nevertheless generalize across mirror reversal (e.g., Riesenhuber and Poggio 1999). Moreover, invariance to mirror reversal in shape processing has been demonstrated in single-cell recordings in macaque inferotemporal cortex (Baylis and Driver 2001).

In conclusion, our results demonstrate that whereas repetition suppression for objects in ventral visual cortex generalizes across mirror reversal of the image, it is strongly dependent on spatial attention to the related prime, thus indicating that it requires attentional selection of relevant information in the ventral visual system.
visual input and is not an as automatic and intrinsic property of cortical cells (cf. Wiggs and Martin 1998), as often assumed.

GRANTS
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REFERENCES


