Automatic processing of unattended object features by functional connectivity

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INTRODUCTION

Humans can selectively attend to different features of the same object, such as shape, color, or motion (see Maunsell and Treue, 2006, for a review). At the cortical level, areas that preferentially respond to these separate object features have been identified. For instance, areas in the lateral occipital cortex show preferential responses to intact images of objects as opposed to scrambled images (referred to as the lateral occipital complex, LOC; Malach et al., 1995). Similar feature-preferring areas have been found for color (V4 and V8; McKeefry and Zeki, 1997; Hadjikhani et al., 1998) and motion (area MT; Zeki, 1980; Tootell et al., 1995). It is well established that attention to specific object features can increase the hemodynamic responses in areas that preferentially respond to those features (Corbetta et al., 1990, 1991; Peuskens et al., 2004; Schultz and Lennert, 2009; see Kanwisher and Wojciulik, 2000 for a review).

Thus, selective attention can help observers interact with a complex and dynamic environment by recruiting appropriate neural resources to process object features that are relevant for the task at hand. However, it may still be beneficial to implicitly process or encode information about unattended features because they may become task-relevant under different circumstances. It is therefore important to have neural mechanisms that automatically process features belonging to the same object irrespective of whether they are or are not relevant for the task at hand. This automatic processing can be considered a form of feature integration. The results of several imaging studies suggest that unattended features can still be processed and may even be integrated with attended features (O’Craven et al., 1999; Paradis et al., 2008; Xu, 2010). One possibility is that specific areas may encode the conjunction of multiple features. For example, areas along the fusiform gyrus (FG), posterior parietal cortex, and superior temporal sulcus have been shown to respond to feature conjunctions (Kawasaki et al., 2008; Schultz et al., 2008; Cavina-Pratesi et al., 2010). Other studies further suggest that feature-prefering areas such as visual areas, LOC and MT, may also respond to feature conjunctions (Corbetta et al., 1990; Self and Zeki, 2005; Sarkheil et al., 2008; Seymour et al., 2009).

These regional responses are important but they do not necessarily provide a complete picture of feature integration at the neural level as there are also anatomical and functional connections between areas (Fellman and Van Essen, 1991; Hagmann et al., 2008). These connections may, for instance, allow areas to pool information from other areas. This form of feature integration can occur either between two anatomically connected areas or in a functional network that may not necessarily be anatomically connected.

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integration could therefore also occur through the functional connectivity between different brain areas, which can be conceptualized as the correlation in the activation time series between brain areas (Friston et al., 1997; Horwitz, 2003; Bingel et al., 2007; Nummenmaa et al., 2010). These connections can further depend on the attentional state of the observer (Friston et al., 1997). Thus in addition to modulating regional responses, selective attention can modulate the functional connectivity between areas.

To date, no study has been designed to test whether functional connectivity may provide a mechanism to integrate multiple object features (in addition to regional responses to the conjunction of features). We used the Garner paradigm to test this possibility. This paradigm has been a powerful method to test whether features are integral or separable at the behavioral level (Garner, 1974, 1988; Gottwald and Garner, 1975; see also Cant et al., 2008). With integral features, observers’ performance on a task with an attended feature is affected by changes to the unattended feature (i.e., Garner interference). That is, integral features are processed automatically irrespective of whether they are task-relevant or task-irrelevant. By comparison with separable features, observers’ performance on the task with the attended feature is unaffected by changes to the unattended feature. Here we applied the logic of the Garner paradigm to a feature-attention task to test whether different features are integral or separable at the neural level. Applying the logic of the Garner paradigm to neural responses does not mean that we necessarily expect differences between conditions at the behavioral level.

Observers in our study were presented with novel objects which had distinct three-dimensional shape, color, and non-rigid motion. They were instructed to attend to the shape, the color, or the motion and detect changes to the attended feature while ignoring the remaining features in two types of blocks. On change blocks, both unattended features varied from trial to trial. By comparison, on no-change blocks, both unattended features were held constant which leads to fMRI adaptation (i.e., a decrease in the BOLD response to the unattended object features; Grill-Spector and Malach, 2001). The two block types allowed us to measure whether and how changes to the unattended features affect regional responses and functional connectivity between areas while keeping the task constant (see Sakai, 2008, for a review of possible effects of task sets induced by the different instructions). The Garner interference is typically measured by the difference in speeded response times between change and no-change blocks (e.g., Cant et al., 2008). We have previously shown that observers’ performance was affected by changes to unattended features using the same stimuli as the ones used in the present study (Mayer and Vuong, 2012). However, behavioral differences may drive changes in regional responses or functional connectivity between areas. We therefore designed our fMRI experiment so that performance was equated across the two block types (e.g., Nummenmaa et al., 2010). Thus, in contrast to the standard (behavioral) Garner tasks, we do not expect observers’ performance to differ on change and no-change blocks. If shape, color, and motion are separable features at the neural level, we predict no differences in regional responses and functional connectivity for change blocks compared to no-change blocks. However, if these features are integral as suggested by previous studies (e.g., O’Craven et al., 1999; Peuskens et al., 2004; Self and Zeki, 2005; Sarkheil et al., 2008; Schultz et al., 2008; Seymour et al., 2009; Cavina-Pratesi et al., 2010), then there are two possible BOLD signatures that parallel the behavioral signature originally reported by Garner and colleagues (Garner, 1974, 1988; Gottwald and Garner, 1975). If feature integration is accomplished by regional responses, then areas that encode the conjunction of features would show larger activation on change blocks compared to no-change blocks (e.g., O’Craven et al., 1999; Paradis et al., 2008; Xu, 2010). If feature integration is accomplished by functional connectivity between areas, then connections between areas would show larger correlations in their activation time series on change blocks compared to no-change blocks.

MATERIALS AND METHODS

PARTICIPANTS

Twelve volunteers (4 males, 8 females; age in years: \( M = 31 \), \( SD = 6 \); age range: 24–42 years) participated in this study. All participants were naive to the purpose of the study and gave informed consent. They had never seen the stimuli before. Due to a technical failure, behavioral results were only available for 11 out of 12 participants. The study was performed in accordance with the Declaration of Helsinki and approved by the ethics committee of Newcastle University.

STIMULI AND APPARATUS

A set of 64 novel colored dynamic objects was used (for details see Mayer and Vuong, 2012). Briefly, each object was defined by a combination of one of four shapes (e.g., cylinder or brick), one of four colors (e.g., red or blue), and one of four non-rigid motions (e.g., bending or twisting). Each object subtended approximately 7.7° (height) \times 3.8° (width) of visual angle. Examples of the stimuli can be found at: http://www.staff.ncl.ac.uk/q.c.vuong/MayerVuong.html

We used a canon XEED LCD projector (1280 \times 1024 pixels) to backproject the visual stimuli onto a projection screen at the foot-end of the scanner. Participants viewed the projection through an angled mirror attached to the head coil approximately 10 cm above their eyes. The experiment was run on a Windows PC using the Psychophysics toolbox version 3 (http://www.psychtoolbox.org; Brainard, 1997; Pelli, 1997) to control the experiment, present the stimuli and record responses. Participants responded via a MR-compatible response pad (LumiTouch™).

DESIGN

The experiment was set up as a 2 \times 3 within-subjects design with block type (change, no-change) and attended feature (shape, color, motion) as repeated measures. A block design was used in which the six experimental conditions were run in separate blocks. The six blocks were presented twice in a single functional run. The 12 blocks were grouped into four sets. Within each set, each attended-feature condition was presented once and one of the two block-type conditions was presented twice. Both the order of the attended-feature conditions and the repeated block type were randomly determined for each set. In addition to the six experimental blocks, there were five 12 s fixation blocks. The
fixed. Fixation blocks were presented at the beginning of each functional run and after each set of three experimental blocks. There were two functional runs each lasting approximately 7 min in the scanning session.

PROCEDURE
Participants were instructed outside the scanner and ran through a set of practice blocks of the experiment ensuring that every participant practised each experimental condition. In one experimental block, participants watched a sequence of eight objects. They were instructed to attend to one feature (color, shape, or motion) and to perform a 1-back feature matching task on the attended feature (see also Schultz and Lennert, 2009). On change blocks, the objects were selected so that the unattended features were different on consecutive trials (except on target trials; see below). On no-change blocks, the objects were selected so that the unattended features were the same throughout the block. Participants pressed a button on the response pad with the left thumb whenever the attended feature was repeated on consecutive trials (target trials). No response was required on mismatch trials. Therefore, a response was only required on a small set of trials. We did not explicitly ask participants to respond as quickly as possible as we only implemented the behavioral task to ensure that participants attended to the instructed feature for that block. For each participant, the 64 objects were used as equally often as possible. Furthermore across participants, the 64 objects occurred in all six experimental as equally often as possible.

Each experimental block began with a 2 s instruction (the word: “color,” “shape,” or “motion”) presented at the center of the screen, which indicated the attended feature for that block. The instruction was followed by a 0.7 s blank screen. Within a block, eight objects were shown in total (two of them were target objects; see below). Each object was then shown for 2.5 s at the center of the screen, and was followed by a 0.7 s blank screen. The presentation duration was less than a full cycle of motion for the objects. Based on our previous behavioral data (Mayer and Vuong, 2012), this duration was a sufficient amount of time to judge all three features. In total, an experimental block lasted 27.6 s.

There were two target trials in each experimental block. On target trials, the repeated object matched the preceding object on all three features. This was necessary to keep target trials on change and no-change blocks equivalent as the unattended features on a no-changed block were identical throughout the block. Participants were asked to respond when they detected a target trial. A response was counted as correct if it was made during the 2.5 s presentation of the repeated object on a target trial. Misses (i.e., not responding on a target trial) and responses made on any other occasions were counted as errors. No feedback was provided.

FUNCTIONAL LOCALIZERS
We used standard functional localizers to identify brain areas that preferentially responded to shape, motion, or color. The shape and color localizers were combined into a single functional run lasting ∼7 min. For these localizers, participants viewed blocks of colored images of common objects, phase-scrambled versions of these images, colored Mondrian patches, and grayscale Mondrian patches. Each image was shown for 0.3 s with a 0.4 s blank between images. There were four blocks of each type containing 18 different images each. The block order was counterbalanced and intermixed with nine fixation blocks (12 s). Data for the motion localizer were collected in a separate functional run lasting ∼5 min. The localizer contained eight blocks in which a random-dot field expanded and contracted from the center of the screen (15 s) and eight blocks in which the random-dot field remained static (15 s). The dynamic and static blocks alternated (starting with a dynamic block) with fixation blocks (12 s) intermixed between the two main block types.

IMAGE ACQUISITION
All participants were scanned at the Newcastle Magnetic Resonance Centre. In the scanner, head motion was restrained with foam pads that were placed between the head and the head coil. Anatomical T1-weighted images and functional T2*-weighted echo planar images (EPIs) were acquired from a 3 T Philips Intera Achieva MR scanner using a Philips 8-channel receive-only head coil. The high resolution T1-weighted scan (MP-RAGE) consisted of 150 slices and took approximately 5 min to acquire. The field of view (FOV) was 240 3 240 3 180 mm3 with a matrix size of 208 3 208 pixels. Each voxel was 1.15 3 1.15 3 1.2 mm3 in size. The T2*-weighted EPIs consisted of 27 axial slices acquired from the bottom to the top of the head. The parameters of the EPIs were: TR = 1.92 s, TE = 40 ms, flip angle = 90◦. The FOV was 192 3 192 3 107 mm3 with a matrix size of 64 3 64 pixels. Each voxel was 3 3 3 mm3 in size, with a 1 mm gap between slices. We use sensitivity encoding (SENSE) with factor = 2 to increase the signal-to-noise ratio of the functional images. For each participant, a total of 414 functional images were acquired (207 in each run). Before each functional run, four “dummy” scans were acquired to allow for equilibration of the T1 signal. The functional localizers were acquired in the same scanning session using T2*-weighted EPIs with the same parameters as the experiment.

fMRI PREPROCESSING
Functional images of the experiment were realigned to the first image of each participant and resliced to correct for head motion. These images were normalized to a standard MINI EPI T2*-weighted template with a resampled voxel size of 3 3 3 mm3. They were then spatially smoothed with a 6 mm full-width-at-half-maximum Gaussian kernel to improve the signal-to-noise ratio and to allow for comparisons across participants. To remove low-frequency drifts in the signal, we applied a high-pass filter with a cutoff of 128 s. We also applied an autoregressive model [AR(1)] to estimate serial correlations in the data and adjust degrees of freedom accordingly. The same procedure was applied to the functional localizers.

fMRI ANALYSES
The preprocessed data were analysed using SPM5 (http://www.fil.ion.ucl.ac.uk/spm; Friston et al., 1994). We used the general linear model (GLM) with a two-step mixed-effects approach. First,
a fixed-effects model was used to analyse each participant’s data set. Second, a random-effects model was used to analyse the individual datasets at the group level. No additional smoothing of the images was used at the group level.

The design matrix for each participant was constructed as follows. The onset and duration for each of the six experimental conditions (2 block types × 3 attended features; each with a duration of 27.6 s) and the fixation block (12 s) were modeled as boxcar functions. These boxcar functions were then convolved with a canonical hemodynamic response function (HRF), implemented in SPM5 as the difference of two gamma functions. In addition to these regressors of interest, the six movement parameters (roll, yaw, pitch, and three translation terms), the instruction period (2.0 s), and a constant term for each session were included in the design matrix as regressors of no interest. A linear combination of the regressors was fitted to the BOLD signal to estimate the beta weight for each regressor.

For the first-level analysis, contrast images for the main effect of block type, the main effect of attended feature, and the interaction between block type and attended feature were computed from the beta-weight images. For the second-level group analysis, 1-sample $t$-tests of participants’ contrast images were conducted at each voxel. We used an initial (uncorrected) threshold of $p < 0.001$ and a minimum cluster size of $k = 10$ voxels (unless otherwise stated). We accepted clusters as significant if $p < 0.05$, FDR-corrected for multiple comparisons at the cluster level.

Similarly, for our functional localizers, we created a design matrix that included block type as our regressors of interest, and the movement parameters and a constant term as regressors of no interests. After fitting this matrix, we computed contrast images for the contrast of interest for each functional localizer (shape: object > scrambled, color: color > grayscale, and motion: dynamic > static). At the second-level group analysis, 1-sample $t$-tests of participants’ contrast images were conducted at each voxel. As for the experimental data, we used an initial (uncorrected) threshold of $p < 0.001$ and a minimum cluster size of $k = 10$ voxels.

**PPI ANALYSES**

We used psychophysiological interaction (PPI) analyses (Friston et al., 1997) to identify target areas which show changes in functional connectivity between seed and target areas as a function of block type. For each PPI analysis, a design matrix was constructed for each participant that included the activation time series from a seed area (i.e., the physiological variable), the boxcar time series for the two block types convolved with the canonical HRF (i.e., the psychological variable), and the product between the two time series (i.e., the PPI). The activation time series was computed by deconvolving the first eigenvariate of the BOLD time series from the participants’ beta-weight image for the PPI regressor to a 1-sample $t$-test against zero.

**RESULTS**

**BEHAVIORAL RESULTS**

For accuracy, there was no main effect of attended feature or block type, and no interactions between these two factors (all $ps > 0.11$). For response times, there was only a main effect of attended feature [$F(2, 20) = 35.86; \ p < 0.001; \ \eta^2_p = 0.78$; all remaining $ps > 0.14$]. Tukey’s post-hoc tests revealed that response times were quicker on attention to color trials compared to the other two attend-feature conditions, and that response times were quicker on attention to shape trials compared to attention to motion trials ($ps < 0.05$). We did not find Garner interference at the behavioral level (Garner, 1974, 1988; Gottwald and Garner, 1975): there were no behavioral differences between change and no-change blocks. However, the experiment was designed to equate performance on these two block types as it has been shown that response times can affect both regional responses and functional connectivity (e.g., Ganel et al., 2005; Prado and Weissman, 2011).

**fMRI RESULTS: FUNCTIONAL LOCALIZERS**

Table 1 shows the significant cortical areas for the different localizers. For the shape localizer, we focused on the two areas along the...
the left and right lateral occipital cortex and the FG (e.g., Grill-Spector et al., 2001) for the subsequent overlap calculations although we report all significant areas in Table 1. The peak voxel in the left hemisphere was in the cerebellum (as reported by the WFU PickAtlas; http://fmri.wfubmc.edu/software/PickAtlas; Maldjian et al., 2003). We therefore reported a peak in cortex which was a sub-cluster of the area based on the coordinate of the peak voxel in the right hemisphere.

Within each hemisphere, the areas activated by the localizers overlapped with each other. However, we needed masks of the localizers that did not overlap with each other to determine whether the areas identified by our whole-brain and PPI analyses in the main study overlapped uniquely with functionally localized shape-, motion- and color-selective areas. Thus, voxels that were activated by more than one localizer were excluded when computing overlap (see below). Figure 1 shows only voxels that were uniquely activated by one of the localizers.

fMRI RESULTS

Attention to shape (S), color (C), and motion (M)

Following previous work (e.g., Corbetta et al., 1990; Peuskens et al., 2004), we identified areas that preferentially responded to the attended object feature by contrasting blocks in which a feature was attended with blocks in which the other two features were attended. Activated areas were labeled with the WFU Pickatlas (Maldjian et al., 2003). Figure 2 and Table 2 present areas which showed larger activation for the attended features with respect to the other two features. When participants attended to shape \((S > C + M)\), only one area in the right MOG was significant.

To test whether the activated area was consistent with the areas that the localizer identified as shape-areas we divided the number of voxels contained in a localizer mask by the total number of voxels of the area. The cluster activated when participants attended to shape did not overlap with the LOC localizer and only 11% of the area overlapped with the MT localizer (Table 3).

Previous studies have reported activation in temporal, parietal, and frontal areas when observers processed an object’s shape (Murray et al., 2003; Peuskens et al., 2004; Paradis et al., 2008; Schultz et al., 2008). To further explore our results, we lowered the initial threshold to \(p < 0.01\) and accepted areas whose peak voxel had an uncorrected \(p < 0.001\). As shown in Table 4, there was a trend that attention to shape activated areas in the right occipital cortex (one at the posterior part and one more ventrally at the occipito-temporal junction), in the left ventral occipito-temporal and in parietal cortices (Peuskens et al., 2004; Schultz et al., 2008).

Several areas showed larger activation when participants attended to motion relative to when they attended to the other features \((M > C + S)\). An area in the posterior temporal cortex extending to lateral occipital areas was activated in each hemisphere. These areas overlapped with the shape localizer (27 and 38%) and one of them overlapped slightly with the motion localizer (8%). Consistent with previous work (e.g., Peuskens et al., 2004; Paradis et al., 2008; Schultz et al., 2008), areas were also

![Figure 1](https://example.com/figure1.png)

**FIGURE 1** | Results of the functional shape, motion, and color localizers for the whole-brain group-analysis \((N = 12)\). Areas are significant at \(p < 0.05\), FDR-corrected at the cluster level for multiple comparisons across the whole-brain. Only voxels that were uniquely activated by one localizer are shown. L, left; R, right; A, anterior; P, posterior. Coordinates are in MNI space.
found in the parietal and lateral frontal cortex. Additionally, we found significant activation in the postcentral gyrus, the superior temporal gyrus (e.g., Schultz et al., 2008) and the cerebellum.

Attention to color (C > M + S) also activated several areas across the brain. There were significant bilateral areas along the collateral sulcus (CoS) and the cuneus (Cun). These areas are consistent with previously reported areas that were activated when observers processed the color of static objects (e.g., Cant and Goodale, 2007; Cant et al., 2008; Cavina-Pratesi et al., 2010). However, these areas did not overlap with the color localizer but three of them overlapped slightly with the shape and motion localizers (8–25%). Activations were also found in the temporal and frontal cortices. Previous studies have reported activation in frontal areas when observers attended to color (e.g., Cavina-Pratesi et al., 2010). We also found large areas in the cingulate cortex (CC) when observers attended color that have not been previously reported.

**Change vs. no-change blocks**

We also tested for cortical areas with larger responses during change blocks compared to no-change blocks. We expected very little adaptation to occur when both unattended features varied from trial to trial because of the drastic visual changes that occurred between trials. By comparison, we expected fMRI adaptation to occur when the unattended features were held constant within a block (Grill-Spector and Malach, 2001). Surprisingly, there were no significant areas localized by this contrast. Furthermore, no areas showed a significant interaction between block type and attended features.

**PPI RESULTS**

We ran PPI analyses to test whether the functional connectivity between areas depended on block type. All 24 areas from the whole-brain analyses were used as seed areas in the PPI analyses (Table 2). To control for unequal cluster sizes, we used spheres with a 6 mm radius centered on the peak voxel of each area. Thus, each seed area consisted of 33 voxels except for the left cuneus (25 voxels) from the attend-color condition, and the right MOG (26 voxels) from the attend-shape condition. This was due to the lateral location of the peak voxel for these areas which led to an overlap between the sphere and extra-cortical space. Although we interpret all significant PPIs at our p < 0.05, FDR-corrected at the cluster-level threshold, we further indicate those PPIs that also remain significant after a Bonferroni correction (see note in Table 5).

As shown in Figure 3 and Table 5, the PPI analyses identified several target areas that showed a positive change in the functional connectivity between the seed and the target areas on change blocks relative to no-change blocks (i.e., a significant positive
Table 2 | Brain areas which showed larger responses for attention to a feature relative to attention to the remaining features.

| Brain area | MNI | k | z |
|------------|-----|---|---|
| **ATTENTION SHAPE > ATTENTION MOTION/COLOR** |     |   |   |
| MOG       | 33  | 96| 3 | 18| 4.18 |
| ITG       | 48  | 63| 9 | 198| 4.80 |
| STG       | 66  | 45| 15| 28| 4.06 |
| MFG       | 27  | 6 | 57| 32| 3.98 |
| MFG       | 30  | 91| 1 | 7 | 3.85 |
| IFG1      | 54  | 32| 3 | 33| 3.92 |
| IFG2      | 54  | 32| 3 | 33| 3.92 |
| IFG3      | 50  | 32| 3 | 33| 3.92 |
| IFG4      | 57  | 32| 3 | 33| 3.92 |
| IPL       | 39  | 32| 3 | 22| 3.84 |
| PCG       | 39  | 32| 3 | 40| 4.41 |
| Cerebellum| 33  | 72| 27| 123| 4.57 |
| Cerebellum| 36  | 60| 30| 167| 4.69 |

| **ATTENTION COLOR > ATTENTION MOTION/COLOR** |     |   |   |
| MOG       | 24  | 72| 6 | 39| 3.68 |
| LG*       | 18  | 81| 9 | 35| 4.34 |
| Cun       | 3   | 96| 6 | 21| 3.75 |
| Cun       | 15  | 90| 27| 21| 3.40 |
| MTG       | 48  | 75| 27| 82| 3.97 |
| ITG       | 33  | 30| 12| 16| 3.88 |
| PHG       | 33  | 15| 24| 17| 4.28 |
| MFG       | 24  | 51| 21| 29| 3.79 |
| MeFG      | 3   | 63| 9 | 307| 6.64 |
| PreCun    | 3   | 12| 45| 59| 4.23 |
| CC        | 6   | 12| 6 | 17| 4.26 |

*p < 0.05, FDR-corrected at the cluster level.

k, number of voxels in the cluster; MOG, middle occipital gyrus; ITG, inferior temporal gyrus; MFG, middle frontal gyrus; IFG, inferior frontal gyrus; IPL, inferior parietal lobule; STG, superior temporal gyrus; MeFG, medial frontal gyrus; Cun, cuneus; PreCun, precuneus; MTG, middle temporal gyrus; PHG, parahippocampal gyrus; CC, cingulate cortex; PCG, postcentral gyrus; LG, lingual gyrus.

In addition to these findings, we found that four seed areas (attend-motion: right MOG and left ITG; attend-shape: right MOG; attend-color: right CoS) showed significant PPIs with target areas that overlapped with the shape localizer (Sarkheil et al., 2008). Shape- and motion-prefering regions have previously been shown to integrate multiple object features (Self and Zeki, 2005). The left Cun seed area also showed a significant PPI with a target area that completely overlapped with the motion localizer (Sarkheil et al., 2008). Shape- and motion-prefering regions have previously been shown to integrate multiple object features (Self and Zeki, 2005; Sarkheil et al., 2008). Moreover, five of the nine seed areas showed significant PPIs with both posterior and anterior targets. Thus, although there were no changes in regional responses as a function of block type (no main effect of block type and no interaction between block type and attention condition), there were changes to the functional connectivity between seed and target areas.

**DISCUSSION**

In the present study, we investigated the neural signature underlying the automatic processing of object features. We adapted a
Garnet paradigm for functional imaging (Garner, 1974, 1988; Gottwald and Garner, 1975; see also Ganel et al., 2005) in which observers either attended to the shape, color or non-rigid motion of novel colored dynamic objects while the unattended features changed or remained constant. Surprisingly we found that changes to unattended features did not modulate regional responses, despite large trial-by-trial changes to the visual stimulus. Rather, we found that changes to unattended features modulated the functional connectivity between areas.

Consistent with a large body of work, we found that selective attention to features increased regional responses in areas involved in processing those features (e.g., Corbetta et al., 1990, 1991; Paradis et al., 2000, 2008; Murray et al., 2003; Peuskens et al., 2004; Catt and Zeki, 2005; Cant and Goodale, 2007; Cant et al., 2008; Sarkheil et al., 2008; Schultz et al., 2008; Seymour et al., 2009; Cavina-Pratesi et al., 2010). This important replication demonstrated that we could identify feature-prefering areas with our paradigm despite using a simple task. The areas activated by selective attention generally did not overlap with areas identified by the functional localizers but they were spatially close. The differences in spatial location may reflect the fact that more complex stimuli were used in the Garnet paradigm compared to the stimuli used in the localizer runs. Another possibility is that selective attention to an object feature may shift the spatial locus of activation. This is an interesting possibility that could be explored in future studies.

Additionally, we found larger activation in CC when observers attended to color compared to when they attended to shape or motion. Activations in these areas were not found in previous studies on color processing (Self and Zeki, 2005; Cant and Goodale, 2007; Cant et al., 2008, 2009; Seymour et al., 2009; Cavina-Pratesi et al., 2010) but that may be due to the nature of the task used in present study compared to those used in previous studies (see Mayer and Vuong, 2012). The area we found in CC can be divided into an anterior and posterior part (ACC and PCC, respectively). From a theoretical perspective, activations in the ACC and PCC may reflect different strategies for processing shape, color, and motion. These areas may also be involved in how attention is allocated to different features (Schultz and Lennert, 2009). The ACC has been shown to be involved in performance monitoring on a Stroop task, for instance (MacDonald et al., 2000; see Botvinick et al., 2004, for a review). This finding is consistent with the ACC’s role in monitoring performance more carefully on mismatched trials. Observers in our study responded more quickly when they...
attended to color compared to when they attended to shape or motion, suggesting that they may be monitoring their performance differently when they attended to color than when they attended to shape or motion. For example, they may have been able to focus on local “patches” to discriminate color but had to focus on global shape or motion to discriminate the other two features (see Peuskens et al., 2004). The PCC activation seems to further support the possibility that observers used different processing strategies for color than for shape or motion. The PCC is part of the default-mode network (Buckner et al., 2008) and has been found to show deactivations when observers attend to external events (i.e., during an explicit task) relative to a resting state. In particular, larger activation of the PCC has been observed for “easy” tasks compared to “hard” tasks, presumably because “hard” tasks require more external monitoring (e.g., Leech et al., 2011). That said, it may not be straightforward to interpret ACC and PCC activation as a consequence of task difficulty. In our study, participants showed quickest responses for color. Accuracy, however, did not differ across features. Furthermore, Gilbert et al. (2012) suggested that measures such as response times and accuracy may not necessarily be good predictors for deactivations of areas that are part of the default-mode network. Future work will be needed to understand the precise role of the ACC and PCC in selective attention and how these roles may relate to different processing strategies for different object features.

Selective attention had a robust effect on the regional responses of several brain areas in our study. Previous research indicated that unattended visual features also elicit distinct BOLD responses (O’Craven et al., 1999; Peuskens et al., 2004; Ganel et al., 2005; Paradis et al., 2008). We therefore expected differences in regional responses in some brain areas when unattended features varied or remain constant because of adaptation (Grill-Spector and Malach, 2001). However, we found that changes to unattended object features did not affect regional responses in any area relative to when these features remained constant. We also found no areas which showed differential activations between these two conditions when we focused our analyses on each attended feature separately.

Although we did not find overall changes in BOLD responses depending on whether unattended features changed or not, we found that unattended features affected neural responses between areas using PPI analyses (Friston et al., 1997). First and most importantly, our PPI analyses identified target areas that showed an increased functional connectivity between seed and target areas. We re-emphasize that we designed the experiment to ensure that performance was equated across change and no-change blocks (e.g., Nummenmaa et al., 2010). Observers were not instructed to respond quickly. They responded only on trials in which the attended feature changed but not on other trials leading to two responses per eight trials. We have previously shown that changes to unattended features can affect
performance using the same stimuli (Mayer and Vuong, 2012). Thus, the modulation of functional connectivity by changes to unattended features is unlikely due to differences in performance (cf., Ganel et al., 2005). Rather, the functional connectivity between areas suggests that observers continued to automatically process the unattended features at the neural level. Our finding is consistent with Frith et al.'s (1997) finding that attention to motion increased functional connectivity between motion-selective areas when observers attended to the motion of moving dots relative to when they passively viewed the same stimuli. Our finding is also consistent with recent work by Prado and Weissman (2011). They found that trial-by-trial variations in response times were correlated with variations in functional connectivity between middle occipital gyrus and FG. They also showed that observers' covert spatial attention influenced the relationship between response times and functional connectivity, but did not influence regional responses in these areas. These studies highlight the importance of functional connectivity even if there are no differences in regional responses across conditions.

Second, increases in functional connectivity were predominantly found between occipito-temporal seed areas and their target areas (i.e., 8 occipito-temporal areas out of 24 seed areas). Only one seed area in the right IFG showed a significant interaction with a target area in the right MFG. Furthermore, six of the occipito-temporal seed areas showed a significant increase in functional connectivity between seeds and occipito-temporal target areas which overlapped with the LOC and MT localizers. These six target areas (lingual gyrus: LG, ITG, and MOG) were located posteriorly in areas of visual cortex that are known to process basic visual features such as color (McKeefry and Zeki, 1997), motion (Tootell et al., 1995), or shape (Malach et al., 1995). Importantly, shape- and motion-preferring areas have previously been found to integrate color and motion (LOC; see Self and Zeki, 2005) and shape and motion (MT; see Sarkheil et al., 2008). The increase in functional connectivity found in our study may indicate that these areas also pool information about unattended features processed in other areas, such as LG, as a form of feature integration. Alternatively, the increase in functional connectivity may indicate that shape- and motion-preferring areas suppress attentional resources automatically recruited by the unattended features and/or strengthen responses in areas processing the attended feature in the face of distraction by the unattended features. Future work is needed to tease apart these alternatives.

Lastly, we found that several occipito-temporal seed areas (5 out of 7) showed significant interactions with both posterior and anterior cortical target areas. The anterior target areas included those in the ACC, medial frontal gyrus, and IFG. These frontal areas have been implicated in monitoring external events or gauging task difficulty (e.g., MacDonald et al., 2000; Weissman et al., 2005; Leech et al., 2011; Gilbert et al., 2012). For example, MacDonald et al. (2000) found that the ACC monitored mismatches between color labels and their printed color. Furthermore, Weissman et al. (2005) showed that the activity in the dorsal parts of the ACC predicted the amount of behavioral interference by distracting events on a task in which participants had to attend to local or global features of a stimulus. Frontal areas have also been implicated as part of an attentional network which provides top-down cognitive control of a task (e.g., Cabeza et al., 2003). The significant interactions between seed areas and the frontal target areas observed in our study suggest that there may be some top-down control of posterior areas to monitor the larger amount of changing visual information on change blocks compared to no-change blocks. However, this interpretation is speculative as we need to be cautious about interpreting the direction of influence from PPI analyses (Gitelman et al., 2003; Nummenmaa et al., 2010). Overall, our results highlight the importance of functional connections between anterior task-monitoring areas and posterior feature-preferring areas and between posterior areas for processing multi-feature objects. These connections are particularly important when observers are attending to some features while ignoring others.

In our study, observers were instructed to attend to different features. This instruction may have induced different task sets which may have modulated both regional responses and interactions between areas (see Sakai, 2008, for a review). Sakai and Passingham (2003) found that the correlation in activity between anterior prefrontal areas and posterior prefrontal areas (e.g., superior frontal sulcus and IFG) depended on whether observers were instructed to remember items in the order they were presented or in reversed order. Interestingly, these interactions between prefrontal areas were observed during a delay period before the task began and were sustained throughout the task. Task set rather than selective attention may have partly driven some of the BOLD responses we observed in our study. However, the modulation of functional connectivity between areas cannot be driven solely by task set as the key comparison was whether unattended features varied or remained constant for a given attended feature (observers had to attend to shape, motion, and color during both change and no-change blocks).

Previous studies have identified several areas that respond selectively to the conjunction of features. These feature-conjunction areas have been identified along the ventral and dorsal processing pathways. We did not find any areas that responded selectively to the conjunction of multiple object features in our study (though caution is warranted for null findings). However, our results complement the results of these previous studies rather than contradict them. It is most likely the case that there are complementary mechanisms to automatically process features that belong to the same object. These mechanisms can work alone or in combination. Importantly, they allow observers to adaptively select those object features that are the most effective for the task at hand, while at the same time allowing them to encode or process unattended features that may become task-relevant under different circumstances.

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REFERENCES

Bingel, U., Rose, M., Glascher, J., and Buchel, C. (2007). fMRI reveals how pain modulates visual object processing in the ventral visual stream. Neuron 55, 157–167.

Botvinick, M. M., Cohen, J. D., and Carter, C. S. (2004). Conflict monitoring and anterior cingulate cortex: an update. Trends Cogn. Sci. 8, 539–546.

Brainard, D. H. (1997). The psychophysics toolbox. Spat. Vis. 10, 433–456.

Buckner, R. L., Andrews-Hanna, J. R., and Schacter, D. L. (2008). The brain’s default network – anatomy, function, and relevance to disease. Ann. N.Y. Acad. Sci. 1124, 1–38.

Cabeza, R., Dolcos, F., Prince, S. E., Rice, H. I., Weissman, D. H., and Nyberg, L. (2003). Attention-related activity during episodic memory retrieval: a cross-function fMRI study. Neuropsychologia 41, 390–399.

Cant, J. S., Arnott, S. R., and Goodale, M. A. (2009). fMR-adaptation reveals separate processing regions for the perception of form and texture in the human ventral stream. Exp. Brain Res. 192, 339–345.

Cant, J. S., and Goodale, M. A. (2007). Attention to form or surface properties modulates different regions of human occipitotemporal cortex. Cereb. Cortex 17, 713–731.

Cant, J. S., Large, M.-E., McCall, L., and Goodale, M. A. (2008). Independent processing of form, colour, and texture in object perception. Perception 37, 1–20.

Cavina-Pratesi, C., Ktentrige, R. W., Heywood, C. A., and Milner, A. D. (2010). Separate channels for processing form, texture, and color: evidence from fMRI adaptation and visual object agnosia. Cereb. Cortex 20, 2319–2332.

Corbetta, M., Miezin, F. M., Dobmeier, S., Shulman, G. L., and Petersen, S. E. (1990). Attentional modulation of neural processing of shape, color, and velocity in humans. Science 248, 1556–1559.

Corbetta, M., Miezin, F. M., Dobmeier, S., Shulman, G. L., and Petersen, S. E. (1991). Selective and divided attention during visual discriminations of shape, color, and speed – functional-anatomy by positron emission tomography. J. Neurosci. 11, 2383–2402.

Fellemann, D. J., and Van Essen, D. C. (1991). Distributed hierarchical processing in the primate cerebral cortex. Cereb. Cortex 1, 1–47.

Friston, K. J., Buechel, C., Fink, G. R., Morris, J., Rolls, E., and Dolan, R. J. (1997). Psychophysiological and modulatory interactions in neuroimaging. Neuroimage 6, 218–229.

Friston, K. J., Holmes, A. P., Worsley, K. J., Poline, J. P., Frith, C. D., and Frackowiak, R. S. J. (1994). Statistical parametric maps in functional imaging: a general linear approach. Hum. Brain Mapp. 2, 189–210.

Ganel, T., Valvrey, K. F., Goshen-Gottstein, Y., and Goodale, M. A. (2005). The involvement of the “fusiform face area” in processing facial expression. Neuropsychologia 43, 1645–1654.

Garnier, W. R. (1974). The Processing of Information and Structure. Hillsdale, NJ: Lawrence Erlbaum Associates.

Garnier, W. R. (1988). Facilitation and interference with a separable redundant dimension in stimulus comparison. Percept. Psychophys. 41, 321–330.

Gilbert, S., Bird, G., Frith, C., and Burgess, P. (2012). Does “task difficulty” explain “task-induced deactivation”? Front. Psychol. 3:125. doi: 10.3389/fpsyg.2012.00125

Gittelman, D. R., Penny, W. D., Ashburner, J., and Friston, K. J. (2003). Modeling regional and psychophysiological interactions in fMRI: the importance of hemodynamic deconvolution. Neuroimage 19, 200–207.

Gottwald, R. L., and Garner, W. R. (1975). Filtering and condensation tasks with integral and separable dimensions. Percept. Psychophys. 18, 26–28.

Grill-Spector, K., Kourtzi, Z., and Kanwisher, N. (2001). fMR-adaptation: a tool for evidence from fMRI adaptation and psychophysiologic interactions in neuromaging. Neuroimage 1233–1239.

Hadjikhani, N., Liu, A. K., Dale, A. M., Corbetta, M., Miezin, F. M., Dobmeyer, R. J., and Buchel, C. (2007). fMRI study. Neuroimage 39, 6:e159. doi: 10.1016/j.neuroimage.2006.06.015

Holland, R. L., Reppas, J. B., Benson, R. R., Kwong, K. K., Jiang, H., Kennedy, W. A., et al. (1995). Object-related activity revealed by functional magnetic-resonance-imaging in human occipital cortex. Proc. Natl. Acad. Sci. U.S.A. 92, 8135–8139.

Maldjian, J. A., Laurienti, P. J., Kraft, R. A., and Burdette, J. H. (2003). An automated method for neuroanatomic and cytoarchitectonic atlas-based interrogation of fMRI data sets. Neuroimage 19, 1233–1239.

Mansell, J. H. R., and Treue, S. (2006). Feature-based attention in visual cortex. Trends Neurosci. 29, 317–322.

Mayer, K. M., and Vuong, Q. C. (2012). The influence of unattended features on object processing depends on task demand. Vis. Res. 56, 20–27.

McKeefry, D. J., and Zeki, S. (1997). The position and topography of the human colour centre as revealed by functional magnetic resonance imaging. Brain 120(PT 12), 2229–2242.

Murray, S. O., Ohlhausen, B. A., and Woods, D. L. (2003). Processing shape, motion and three-dimensional shape-motion in the human cortex. Cereb. Cortex 13, 508–516.

Nammenmara, L., Passamonti, L., Rowe, J., Engell, A. D., and Calder, A. J. (2010). Connectivity analysis reveals a cortical network for eye gaze perception. Cereb. Cortex 20, 1780–1787.

O’Craven, K. M., Downing, P. E., and Poline, J. B. (2008). Visual perception of motion and 3-D structure from motion: an fMRI study. Cereb. Cortex 10, 772–783.

Paradis, A. L., Droulez, J., Cornilleau-Peres, V., and Poline, J. B. (2008). Processing 3D form and 3D motion: respective contributions of attention-based and stimulus-driven activity. Neuroimage 43, 736–747.

Pelli, D. G. (1997). The videotoolbox software for visual psychophysics: transforming numbers into movies. Spat. Vis. 10, 437–442.

Peukens, H., Claes, K. G., Todd, J. T., Norman, J. F., Van Hecke, P., and Orban, G. A. (2004). Attention to 3-D shape, 3-D motion, and texture in 3-D structure from motion displays. J. Cogn. Neurosci. 16, 665–678.

Prado, J., and Weissman, D. H. (11). Spatial attention influences trial-by-trial relationships between response time and functional connectivity in the visual cortex. Neuroimage 54, 465–473.

Sakai, K. (2008). Task set and prefrontal cortex. Annu. Rev. Neurosci. 31, 219–245.

Sakai, K., and Passingham, R. E. (2003). Prefrontal interactions reflect future task operations. Nat. Neurosci. 6, 75–81.

Sarkheil, P., Vuong, Q. C., Balthoff, H. H., and Noppeney, U. (2008). The integration of higher order form and motion by the human brain. Neuroimage 42, 1529–1536.

Schultz, J., Chiang, L., and Vuong, Q. C. (2008). A dynamic object-processing network: metric shape discrimination of dynamic objects by activation of occipitotemporal, parietal, and frontal cortices. Cereb. Cortex 18, 1302–1313.

Schultz, J., and Lennert, T. (2009). BOLD signal in intraparietal sulcus covaries with magnitude of implicitly driven attention shifts. Neuroimage 45, 1314–1328.

Selt, M. W., and Zeki, S. (2005). The integration of colour and motion by the human visual brain. Cereb. Cortex 15, 1270–1279.

Seymour, K., Clifford, C. W. G., Logothetis, N. K., and Bartels, A. (2009). The coding of color, motion, and their conjunction in the human visual cortex. Curr. Biol. 19, 177–183.

Tootell, R. B., Reppas, J. B., Kwong, K. K., Malach, R., Born, R. T., Brady, T. J., et al. (1995). Functional analysis of human MT and related visual processing of shape, color, and texture in the human ventral stream. Neuron 55, 157–167.
cortical areas using magnetic resonance imaging. *J. Neurosci.* 15, 3215–3230.

Weisman, D. H., Gopalakrishnan, A., Hazlett, C. J., and Woldorff, M. G. (2005). Dorsal anterior cingulate cortex resolves conflict from distracting stimuli by boosting attention toward relevant events. *Cereb. Cortex* 15, 229–237.

Xu, Y. (2010). The neural fate of task-irrelevant features in object-based processing. *J. Neurosci.* 30, 14020–14028.

Zeki, S. (1980). The responses of cells in the anterior bank of the superior temporal sulcus in macaque monkeys. *J. Physiol. Lond.* 308, P85.

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