Hemispheric Asymmetry in Global/Local Processing: Effects of Stimulus Position and Spatial Frequency

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We examined the neural mechanisms of functional asymmetry between hemispheres in the processing of global and local information of hierarchical stimuli by measuring hemodynamic responses with functional magnetic resonance imaging (fMRI). In a selective attention task, subjects responded to targets at the global or local level of compound letters that were (1) broadband in spatial-frequency spectrum and presented at fixation; (2) broadband and presented randomly to the left or the right of fixation; or (3) contrast balanced (CB) to remove low spatial frequencies (SFs) and presented at fixation. Central broadband stimuli induced stronger activation in the right middle occipital cortex under global relative to local attention conditions but in the left inferior occipital cortex, stronger activation was induced under local relative to global attention conditions. The asymmetry over the occipital cortex was weakened by unilateral presentation and by contrast balancing. The results indicate that the lateralization of global and local processing is modulated by the position and SF spectrum of the compound stimuli. The global attention also produced stronger activation over the medial occipital cortex relative to the local attention under all the stimulus conditions. The nature of these effects is discussed. © 2002 Elsevier Science (USA)

Key Words: compound stimulus; contrast balancing; fMRI; global/local processing; hemispheric asymmetry; spatial frequency; occipital cortex.

INTRODUCTION

Visual perception of complex scenes requires the coordinated processing of images on both global and local levels. During global processing local elements are grouped into perceptual wholes, whereas during local

¹ To whom correspondence should be addressed. Fax: 8610-6276-1081. E-mail: shan@pku.edu.cn. processing scene elements are analyzed as individual objects. Global and local processing has been widely studied using compound letters similar to that shown in Fig. 1a, where the global letter is made up of many smaller letters on a local level (Navon, 1977).

Neuropsychological studies indicate that separate subsystems in the human brain are critical for the processing of global and local aspects of hierarchical objects (Robertson and Lamb, 1991). In particular, lesions of the temporal/parietal junction in the right hemisphere impair the processing of global structure in Navon-type compound stimuli, whereas corresponding left hemisphere lesions impair the processing of local elements (Lamb *et al.*, 1989, 1990; Robertson *et al.*, 1988). These results suggest that the right and left hemispheres are specialized for global and local processing, respectively.

Behavioral studies of the lateralization of global/ local processing in normal subjects have shown small and inconsistent asymmetries when stimuli are presented unilaterally in the left or right visual fields. Some researchers have found faster reaction times (RTs) to global targets presented in the left visual field (LVF) than in the right visual field (RVF) and a reverse pattern for local responses (Sergent, 1982; Kimchi and Merhav, 1991). Other researchers, however, failed to find such asymmetry (Boles and Karner, 1996; Martin, 1979; Polich and Aguilar, 1990), or have shown that the apparent asymmetries may result from differences in criterion rather than detection (Han *et al.*, 2002b).

Event-related potential (ERP) studies have shown conflicting results as well. Evans *et al.* (2000) found that an early positive component between 80 and 120 ms after sensory stimulation was enhanced over the left hemisphere in the local relative to the global condition and over the right hemisphere in the global relative to the local condition. Such a pattern of hemispheric asymmetry in global/local processing was also observed in a late occipito-temporal negativity (Heinze *et al.*, 1994) and global and local target-specific differ-



ence waves (Han *et al.,* 2000; Proverbio *et al.,* 1998). However, other ERP studies did not find such a pattern of hemispheric asymmetry in global/local processing (Han *et al.,* 1997, 1999a, 2002b).

The results of brain imaging studies are also conflicting. For example, positron emission tomography (PET) studies found that attention to global letters enhanced activation over the right lingual gyrus whereas locally directed attention enhanced activation over the left inferior occipital cortex (Fink *et al.*, 1996, 1997). In a task to identify the orientation of compound gratings, Fink et al. (1999) further showed that global attention increased occipital activity in the right hemisphere only when relatively high spatial frequency (SF) stimuli were used and local attention increased occipital activity in the left hemisphere only when relatively low SF stimuli were used. Because the global shapes appeared more prominent than the local shapes for the high SF stimuli whereas the reverse was true for the low SF stimuli, Fink *et al.* argued that the asymmetric occipital activities associated with global and local processing were modulated by perceptual salience of global and local shapes. A functional magnetic resonance imaging (fMRI) study found similar hemispheric asymmetry in global/local processing over the occipitotemporal cortical junctions (Martinez et al., 1997). However, other PET (Heinze et al., 1998) and fMRI (Sasaki et al., 2001) studies failed to replicate hemispheric asymmetry over the occipital cortices possibly because perceptual salience of the global shapes was impaired by composing global letters with few local letters (Heinze *et al.*, 1998) or nontypical Navon-type compound stimuli were used (Sasaki et al., 2001).

The current experiment further addressed the issue of hemispheric lateralization in global/local processing using fMRI. The first objective was to verify the hemispheric asymmetry in global/local processing, as observed in the previous PET and fMRI studies using compound letters presented in the center of the visual field. The experiment was designed to identify the relative difference between the patterns of activation associated with global and local processing. We were particularly interested in whether the left occipital cortex showed stronger activation under local rather than global conditions and the right occipital cortex showed stronger activation under global rather than local conditions.²

The second objective was to determine how the asymmetry is affected by stimulus position. Previous studies that showed hemispheric asymmetry in global/ local processing usually displayed compound stimuli in the center of the visual field (Fink et al., 1996; Han et al., 2000; Heinze et al., 1994; Martinez et al., 1997) or in the two visual fields simultaneously (Evans *et al.*, 2000), whereas those failed to replicate such asymmetry presented compound stimuli unilaterally in the LVF or RVF (Han et al., 1999a, 2002a). Han et al. (2002a) suggested that central presentation allows the two hemispheres equal access to stimulus information and leads to competition between the two hemispheres for processing the information. The competition during a global task results in stronger activation in the right than in the left hemisphere with a converse pattern observed in local tasks. When compound stimuli are presented unilaterally, however, the conditions conducive to competition are eliminated and the hemispheric asymmetry in global/local processing may be attenuated. Since there has been no research to study the effect of stimulus position on laterality associated with global/local processing systematically, the current work examined this hypothesis by recording hemodynamic responses associated with global and local processing of compound stimuli presented in the center of the visual field or randomly in the LVF or RVF. If the lateralized occipital activities elicited by global and local processing reflect the competition between the two hemispheres, according to Han *et al.* (2002a), the asymmetrical activities should be eliminated or weakened by presenting compound stimuli unilaterally.

The third objective of the current work was to examine the relationship between SF contents of compound stimuli and hemispheric lateralization in global/local processing. The SF content of compound stimuli is interesting because low SF channels can only transmit information about the global shape, while high SF channels can transmit information about both global and local shapes. The evidence that global and local shape perception is based on low and high SF channels, respectively (Shulman *et al.*, 1986; Shulman and Wil-

² To test stimulus-position-based and low-SF-based models of hemispheric asymmetry in global/local processing of compound stimuli, it would be nice to show first the activation associated with global and local processing, respectively, and then test how the activation is modulated by stimulus positions and SF contents of the compound stimuli. However, it is always difficult to establish a proper baseline condition for brain activation involved in global/local processing of compound stimuli. Most behavioral studies have shown an unbalanced processing of global and local properties, which suggests that the visual system intends to process global information earlier or

faster than local information. These findings imply that global properties may be mandatorily processed even when subjects are not asked to attend to the global level (see Rauschenberger and Yantis, 2001, for recent evidence). Thus it is possible that, for any stimuli used as a baseline condition in which subjects passively view the stimuli, the global structure is processed more extensively than local parts and thus results in stronger activation of the cortex associated with global processing relative to the cortex associated with local processing. Subtraction between global-attention condition and such a baseline condition may weaken activation associated with global processing and hemispheric asymmetry in global/local processing. Even using rest periods as a baseline condition is not good because activities in specific brain areas are substantially higher during rest than during task and show hemispheric asymmetry (Stark and Squire, 2001).

son, 1987), has led to the development of a model of hemispheric lateralization in global/local processing based on SF filtering (Ivry and Robertson, 1998), which assumes that the right hemisphere operates as a lowpass filter and carries on global processing based on low SFs, and the left hemisphere operates as a highpass filter and accomplishes local processing based on high SFs. The evidence supporting this model comes from patient studies that suggest right hemisphere dominance in global processing and left hemisphere dominance in local processing (Lamb et al., 1989, 1990; Robertson et al., 1988) and normal subjects studies that suggest that the left and right hemispheres are biased toward efficient use of higher and lower SF information, respectively (Christman et al., 1997; Kitterle et al., 1991). According to Ivry and Robertson's (1998) model, when the compound stimuli include only a limited range of high SFs, differentiation between global and local information cannot be based on low and high SFs and thus other features must be used. Under this condition, the hemispheric asymmetry associated with low and high SF processing during hierarchical analysis should be weakened or eliminated. We tested this hypothesis by comparing hemodynamic responses related to global/local processing of compound stimuli that were either broadband in SF spectrum or contrast balanced (CB) to eliminate low SFs³ that is critical for global processing. Although contrast balancing may also reduce medial SFs that may partially contribute to local processing there has been evidence that contrast balancing produced larger effects on global rather than local processing (Lamb and Yund, 1993, 1996).

MATERIALS AND METHODS

Subjects

Ten normal healthy adults (6 men, 4 women; aged between 19 and 43 years) participated in this study as paid volunteers. All had normal or corrected-to-normal vision. All participants were right-handed, without neurological disorders, and gave informed consent according to the guidelines of the Veterans Administration and the University of California, Davis.



FIG. 1. Illustrations of the compound stimuli used in the current study. (a) A broadband compound letter (i.e., a global E is made up of local Hs, which were composed of white lines on a gray background); (b) A CB compound letter (i.e., a global E is made up of local As, the white lines of which are surrounded by lines that were darker than the background).

Stimuli

Stimuli were presented with a PC running Presentation software (www.neurobehavioralsystems.com) through a Sharp LCD projector onto a rear-projection screen located at the subject's feet. The screen was viewed with an angled mirror positioned on the head coil. The stimuli were global letters made up of local letters in a 7 \times 7 matrix, as illustrated in Fig. 1. Letters "H" and "S" served as targets while letters "A" and "E" served as distractors. Stimulus patterns either contained one target at the global level (i.e., global/ local letters were H/A, S/A, H/E, or S/E) or the local level (i.e., global/local letters were A/H, E/H, A/S, or E/S), or contained no targets (i.e., A/E or E/A), resulting in 10 stimulus figures. From a viewing distance of 277 cm, global letters were 4.2° wide and 6.1° high. Local letters were 0.45° wide and 0.70° high.

Compound stimuli were presented on a gray (16.4 cd/m²) background. Each local letter of the broadband stimuli was composed of lines that were brighter (29.3 cd/m²) than the background. The CB stimuli were identical to the broadband stimuli except that the bright lines composing each local letter were surrounded by lines that were darker (10.1 cd/m²) than the background. The change in luminance from background was approximately twice as great for bright lines as for dark lines. Because dark lines occupied twice as much area as bright ones, the space-averaged luminance of the CB stimuli equaled that of the background. This arrangement of the luminance levels removed virtually all detectable spectral power below 3 cycles/degree from the CB stimuli (see Lamb et al., 1999, for the results of spectral power analysis). All lines (both bright and dark) were approximately 0.02° thick.

There were three stimulus conditions in the current experiment: (1) central broadband stimuli, (2) unilat-

³ If Ivry and Robertson's hypothesis (i.e., the hemispheric asymmetry in global/local processing of compound stimuli is based on lowand high-pass filtering operated in the right and left hemispheres, respectively) is correct, it may be predicted that the hemispheric asymmetry in global/local processing should be weakened if perception of compound stimuli is only conducted based on high SFs. Thus it is sufficient to test Ivry and Robertson's model by using high-pass compound stimuli. Low-pass filtering (i.e., blurring local elements of compound stimuli) impairs local perception seriously and makes the difficulty of global and local perception incompatible, and thus was not used in the current study.

eral broadband stimuli, (3) central CB stimuli. A fixation cross of $0.30 \times 0.45^\circ$ was continuously visible in the center of the screen under all the stimulus conditions. Subjects were instructed to maintain fixation at all times on the fixation cross.⁴ Under the central broadband and CB stimulus condition, the compound stimuli were displayed in the center of the visual field. The center of compound letters was 0.83° above the fixation so that compound stimuli and the fixation cross did not overlap. The compound stimuli were randomly displayed to the left or the right of the fixation in the unilateral broadband stimulus condition.⁵ The fixation and the inner edge of the compound stimulus were 1.7° apart. The stimulus displays were presented for 200 ms. Stimulus intervals were randomized between 500 and 1000 ms.

Experimental Design

Two scans of 308 s were obtained under each stimulus condition and were averaged together.⁶ Each scan

⁵ In our ERP study (Han, 2002b), the LVF and RVF stimuli were intermixed in the same blocks of trials. Such a unilateral-presentation condition was designed to eliminate the effect of eye movements. The current fMRI study used the same experimental design (unilateral presentation condition) as that in our ERP study so that we were able to compare the results from both studies. Such a design of the fMRI experiment made us unable to examine Hemisphere imes Visual Field interaction without using event-related analysis. However, this is not essential to the current work because what we were really interested in is the interaction of Attention (global vs local) \times Hemisphere (left vs right), which is useful for demonstrating hemispheric asymmetry in global/local processing (Fink et al., 1996, 1997). Because of limitation of tools for data analysis, we did not do eventrelated analysis of the fMRI data under the unilateral-presentation condition. However, the results of the block design did not affect the accuracy of localizing activation associated with global and local processing as the results of central broadband stimuli of the current work coincided very well with the results of previous work (Fink et al., 1996, 1997). Most important, such a block design does not affect the conclusion we made here.

⁶ We performed retinotopic mapping using grating-like stimuli to identify the boundaries between V1 and V2 in the current experiment. However, for unknown reasons, we failed to obtain clear results from most of the subjects. This made it difficult to localize the effects of global/local attention in V1 and V2 based on retinotopy. Thus we had to use coordinates in Talairach space to refer to the Brodmann areas that showed differential activation between global and local processing.

consisted of 11 epochs of 25 trials (28 s for each epoch), alternating between global and local attention conditions. The first epoch of each scan was excluded from statistical analysis to obtain a similar baseline for all the following epochs of trials. There were 10% target stimuli in each scan. At the beginning of each epoch, a large or a small letter made up of solid black lines was presented for 1000 ms to inform subjects of target shape (H or S) and target level (global vs local). The sizes of global and local cues were the same as those of global and local letters of the compound stimuli. Subjects pressed a hand-held fiber-optic response button to indicate detection of the designated global or local target. The order of stimulus conditions was counterbalanced across subjects. Subjects were given 6 to 10 epochs of trials for practice prior to brain imaging.

Image Acquisition

Brain imaging was performed using a Marconi 1.5-T scanner and a three-axis local gradient head coil. Twenty axial slices of functional images that covered the whole cerebral cortex were acquired using echoplanar imaging ($128 \times 128 \times 20$ matrix with $1.87 \times 1.87 \times 5$ -mm spatial resolution, TR = 2000 ms, TE = 40 ms, FOV = 240 mm, flip angle = 90°). Anatomical images were obtained with a standard 3D T1-weighted sequence (resulting in a $256 \times 212 \times 256$ matrix with 0.938 $\times 1.13 \times 0.938$ -mm spatial resolution, TR = 15 ms, TE = 4.47 ms, flip angle = 35°). Subjects' heads were immobilized during the scanning sessions using pieces of foam.

Analysis of Imaging Data

SPM99 (Welcome Department of Cognitive Neurology, London, UK) implemented in MATLAB (Math Works, Natick, MA) was used for imaging data processing and analysis. For each subject, functional images were realigned to the first scan to correct the head movement between scans. The structural image was coregistered with the mean image produced during the process of realignment. All images were normalized to a 2 \times 2 \times 2-mm³ Montreal Neurological Institute (MNI) template in Talairach space (Talairach and Tournoux, 1988) using bilinear interpolation. Functional images were spatially smoothed using a Gaussian filter with a full-width at half-maximum (FWHM) parameter set to 5 mm. Data were modeled using a boxcar function. Contrasts were used to compare the effect of attention (global vs local) for each stimulus condition. Regions preferentially engaged by global processing were defined as areas more activated by the global attention than by the local attention. Regions preferentially engaged by local processing were defined as areas more activated by the local attention than by the global attention. Statistical effects of attentional conditions for each type of stimulus were first assessed

⁴ We did not measure subjects' eye movements when subjects were in the scanner because of limitation of recording technique. However, the effects of eye movements on global/local processing were evaluated by the results of our ERP study (Han *et al.*, 2002b), which used the same experimental design (unilateral presentation) as that in the current fMRI study and recorded subjects' vertical and horizontal eye movements with electrodes below the left eye and lateral to the left and right external canthi. The electrophysiological results showed that eye movements were small (less than 2 μ v) and the patterns of eye movements under the global and local conditions were very similar. The results suggest that differential eye movements under the global/local conditions contributed trivially to the modulation of hemispheric asymmetry in global/local processing under the condition of the current experiment.

in individual subjects. Random effect analyses were then conducted across the group of 10 subjects based on

TABLE 2

Brain Areas Activated during Global and Local Tasks Under Each Stimulus Condition in the Group Analysis

Condition/region	Voxels, No.	BA	X	Y	Ζ	Z value
Central broadband						
Global attention						
Right middle occipital gyrus	213	19,37	52	-64	-10	3.83
Medial occipital gyrus	61	18	12	-96	22	3.58
Parieto-occipital sulcus	121	19	-10	-74	30	3.44
Local attention						
Left inferior occipital gyrus	52	18	-30	-92	-8	3.45
Unilateral broadband						
Global attention						
Cuneus and precuneus	281	19	2	-84	30	3.58
Right temporal-parietal junction	137	39	56	-66	20	3.74
Central contrast balanced						
Global attention						
Right middle occipital gyrus	79	19,37	54	-64	-10	3.50
Medial occipital gyrus	88	18	-8	-98	14	3.72
Local attention						
Precuneus	162	7	4	-64	42	3.57
Left inferior occipital gyrus	36	18	-26	-94	-6	3.55
Right inferior occipital gyrus	34	18	22	-96	-4	3.45
Left superior temporal gyrus	312	22,42	-42	-12	8	4.84
Left postcentral gyrus	191	1,2	-40	-26	60	4.53
Right temporal-parietal junction	99	39	50	-52	22	3.99

Note. Uncorrected P < 0.01 for all clusters listed in the table; Voxels, No., number of voxels in a cluster.

For unilateral broadband stimuli, global attention generated stronger activation over the medial occipital cortex (BA 19) relative to the local attention condition, as shown in Fig. 4. Unlike central broadband stimuli, there was no evident lateralized activation over the occipital cortex associated with global or local attention. However, globally directed attention produced stronger activation over the right temporal-parietal junction relative to locally directed attention. Because no significant activation was observed under the local attention condition, Fig. 4 shows only the results under the global attention condition.

For central CB stimuli, stronger activation was also observed over the medial occipital areas (BA 18) under the global relative to the local attention condition (Fig. 5). Global attention also activated the right middle occipital areas (BA 19, 37). Local attention activated bilateral inferior occipital gyrus (BA 18), left superior temporal gyrus (BA 22, 42), left postcentral gyrus (BA 1, 2), and the right parieto-temporal junction (BA 39). However, the interaction between Level of Attention and Hemisphere was not significant in the ROI analysis of the occipital activation (F(1,9) = 2.13, P > 0.1). To further confirm the difference in the occipital asymmetry associated with global/local processing between



FIG. 4. Brain areas activated by attention to the global level of unilateral broadband stimuli. The results of the group analysis from 10 subjects were rendered on a 3-D structural image of the MNI template provided by SPM99. Threshold for activation of all clusters was P < 0.01 (uncorrected).



FIG. 5. Brain areas activated by attention to the global or local level of central CB stimuli. The results of the group analysis from 10 subjects were rendered on a 3-D structural image of the MNI template provided by SPM99. Threshold for activation of all clusters was P < 0.01 (uncorrected).

central broadband and CB stimuli, an ANOVA was conducted on the mean values of the ROI over the occipital cortex with factors being Stimulus Set (broadband vs CB), Level of Attention, and Hemisphere. There was a significant interaction of Stimulus Set × Level of Attention × Hemisphere (F(1,9) = 5.53, P < 0.04), indicating that the pattern of hemispheric asymmetry over the occipital cortex associated with global/ local processing was different between central broadband and CB stimuli.

DISCUSSION

The present study used a target detection task to examine hemispheric asymmetry in global and local processing of compound stimuli. The same procedure and motor response were used for all the three conditions so that the difference in both behavioral and hemodynamic responses between the conditions reflects the effects of stimulus position and SF contents of stimuli. Response accuracies were high, indicating that subjects were successful in attending to the global and local levels of compound letters. Similar to Pomerantz (1983), we found that moving broadband stimuli from the fovea to the periphery slowed RTs particularly under the local condition. In contrast, removing low SFs from compound stimuli by contrast balancing slowed RTs particularly under the global condition and resulted in a local RT advantage, consistent with previous reports (Lamb and Yund, 1993, 1996).

There has been debate regarding whether activities of the left and right occipital cortices are differentially modulated by attention to the global or local level of compound stimuli (Fink et al., 2000; Mangun et al., 2000). In a task of selectively naming the global or local letters that were presented in the center of the visual field, Fink et al., (1996) found that the regional cerebral blood flow (rCBF) was increased over the left inferior occipital cortex and the right lingual gyrus by locally and globally directed attention, respectively. However, Heinze et al. (1998) failed to replicate these results in a similar selective attention task. We showed here that. when broadband compound letters were presented in the center of the visual field, locally directed attention was associated with stronger activation over the left inferior occipital cortex in an area close to that of Fink et al. (1996). The focus of the right lateral occipital activation in the global attention condition is more lateral and anterior compared to the results of Fink et al. (1996), but is consistent with the fMRI results of Martinez et al. (1997). The lateralized modulations of occipital activities by global/local attention revealed by our fMRI results is in agreement with the ERP studies (Evans et al., 2000; Han et al., 2000), which found that occipito-temporal activities show larger amplitudes over the right hemisphere when attending to the global shape and larger amplitudes over the left hemisphere

when attending to the local shape. Taken together, these results support the proposition that global and local properties of compound stimuli are differentially represented in the two hemispheres at an early stage of visual processing.

More interestingly, we found that the lateralized extrastriate activation associated with global/local processing was attenuated by unilateral presentation. These results are in agreement with the hypothesis that lateralization in global/local processing depends, to a certain degree, upon stimulus position in the visual field (Han Yund et al., 2002b). A possible account for these effects is that both hemispheres are capable of processing stimuli on either global or local levels, with the right hemisphere being superior in analyzing global patterns and the left hemisphere superior in analyzing local patterns. When stimulus information arrives in the two hemispheres simultaneously, the two hemispheres compete for processing the information at the proper level. Consequently, globally directed attention results in stronger activation in the right than in the left hemisphere and locally directed attention leads to a reverse pattern, as reflected in the asymmetric occipital activation induced by the central broadband stimuli. When compound stimuli are pree(the)-T*[(at)-573.8[(at)-co5.9(c2(unilat[(at)-305.9(occipi(a block of trials and asymmetric temporal-parietal activation when attention had to switch between global and local levels. Fink et al. suggest that temporalparietal regions may exert attentional control over global and local processing. If this is correct, it may be proposed that the findings of the lesion studies (Lamb et al., 1989, 1990; Robertson et al., 1988) may principally reflect impairment of high order processing in global and local perception (such as attentional control). We showed here that hemispheric asymmetry associated with global and local processing may take place at different levels of the brain structure depending on whether the compound letters were displayed in the fovea or periphery. More attentional control may be required under the unilateral than central presentation conditions and resulted in asymmetric activations at high levels of the brain structure.

We also found that contrast balancing modulated the lateralized extrastriate activation associated with global and local processing differentially. Attention to the global shape of the CB stimuli generated stronger activation relative to local attention only in the right lateral occipital cortex. Attention to the local shape of the CB stimuli, however, resulted in stronger activation relative to global attention over bilateral inferior occipital cortices. Apparently, the results suggest that contrast balancing did not influence the asymmetrical occipital activation associated with global processing but weakened the asymmetrical occipital activation associated with local processing by increasing the activity over the right occipital cortex. However, the results in Table 2 suggest that the spatial extent of the activation may be different between the central broadband and CB stimuli. For example, there were more activated voxels under the global condition for the central broadband than CB stimuli (213 vs 79). Similarly, there were more activated voxels under the local condition for the central broadband relative to CB stimuli (52 vs 36). The results suggest that even though both sets of stimuli activated the right and left occipital cortices under the global and local conditions, respectively, the spatial extent of the activation was smaller for the CB relative to the broadband stimuli. Accordingly, it may be proposed that the occipital activation in the right and left hemisphere observed for the central broadband stimuli may partially reflect SF filtering process involved in global and local processing respectively and was weakened by contrast balancing.

Alternatively, contrast balancing might increase the difficulty of both global and local processing by decreasing the perceptual salience of global and local shapes (as reflected by slower global and local RTs to CB relative to broadband stimuli). According to Fink *et al.*'s (1999) hypothesis that decreasing the perceptual salience of global and local shapes attenuated the occipital activation associated with global and local processing, CB would attenuate both the right hemisphere

activation during global processing and the left hemisphere activation during local processing. The right occipital activation observed under the local condition of central CB stimuli may stem from decreased activities under the global condition instead of increased activities under the local condition.

Note that neither the account based on SF filtering nor the account based on decrease of perceptual salience can explain our results alone. Other mechanisms may also contribute to the difference in the occipital activation between central broadband and CB stimuli. For example, the right occipital activation may partially be due to a process of local element grouping that is required for perceiving global shapes but not for local processing and is independent of SF contents of the compound stimuli (Han et al., 1999b; Han and Humphreys, 1999). The central presentation may require more grouping than unilateral presentation because single local elements would stand out more easily in the fovea and compete with the grouping process, and thus induced stronger occipital activation under global rather than local conditions. This proposal is consistent with our recent ERP findings that grouping of local elements elicited enhanced activation over the right occipito-parietal cortex between 180 and 280 ms after stimulus onset (Han et al., 2001b, 2002a).

Attention to the local shape of CB compound stimuli also activated the right temporal-parietal junction and the left superior temporal and postcentral lobes. Parietal areas (particularly in the right hemisphere) have been revealed to play a critical role in human visual selective attention (Corbetta *et al.*, 1993; Hopfinger *et al.*, 2000; Posner *et al.*, 1984). It is possible that, when local shapes could not be differentiated from global shapes by a high-pass filtering process, the perception of easJ/F10 1p975d a Heinze *et al.*) may weaken local element grouping (Han, 1999b; Han and Humphreys, 1999, 2002), and thus reduce the enhanced activity over the right occipital cortex associated with global attention. Similarly, when subjects were asked to discriminate X vs + compound shapes that were displayed in the center of the visual field (Sasaki *et al.*, 2001), subjects could use only a few local figures around fixation to perform the global task. Indeed, the location of any single local shape (other than the one at fixation) provides sufficient information to identify the global shape. This might diminish the difference in SF used in the global and local tasks, and thus weakens the asymmetric occipital activities based on SF filtering or on a grouping process.

Our current experiment also found stronger activation over the medial occipital cortex in the global relative to local attention conditions for the central broadband stimuli, which is similar to Sasaki *et al.*'s (2001) results. We showed further that this occipital activity associated with global processing was evident irrespective of retinal position and SF content of the compound stimuli. Because the medial occipital activation was observed when both global and local shapes were displayed in the peripheral visual field, it may not simply reflect enhanced peripheral representation of the stimulus displays in the global than in the local task. A possible explanation is that the enlargement of an attentional window in the global relative to the local task (Robertson et al., 1993) may enhance the neuronal responses associated with the processing of the stimuli inside the attentional spotlight. This top-down attentional modulation is not affected by where compound stimuli are presented and whether low SFs are available in the stimuli. Alternatively, it is possible that the medial occipital activation may reflect the process of perceptual grouping that is required for the perception of global shape (Han, 1999b; Han and Humphreys, 1999). The latter hypothesis is consistent with our recent ERP study which found that grouping of local elements produced enhanced activation over the medial occipital cortex between 100 and 120 ms after sensory stimulation (Han et al., 2001b, 2002a).

CONCLUSIONS

The present study provided fMRI evidence that the left and right extrastriate cortices are differentially activated by attention to the global or the local aspects of compound letters that were displayed in the center of the visual field. In addition, we showed that this asymmetry was eliminated by unilateral stimulus presentation, consistent with a model of lateralization of global/local processing based on competition between the two hemispheres. The asymmetric occipital activation associated with global and local processing was also diminished by removing low SFs from the stimuli, suggesting a role of SF filtering in that cortical area during the processing of compound letters. Mechanisms other than SF filtering (such as local element grouping) may also contribute to the lateralized occipital activity related to global processing.

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REFERENCES

- Boles, D. B., and Karner, T. A. 1996. Hemispheric differences in global versus local processing: Still unclear. *Brain Cogn.* **30**: 232– 243.
- Christman, S., Kitterle, F. L., and Niebauer, C. L. 1997. Hemispheric asymmetries in the identification of band-pass filtered letters. *Psychon. Bull. Rev.* **4:** 277–284.
- Corbetta, M., Miezin, F. M., Shulman, G. L., and Petersen, S. E. 1993. A PET study of visuospatial attention. *J. Neurosci.* 13: 1202–1226.
- Evans, M. A., Shedden, J. M., Hevenor, S. J., and Hahn, M. C. 2000. The effect of variability of unattended information on global and local processing: Evidence for lateralization at early stages of processing. *Neuropsychologia* **38**: 225–239.
- Fink, G. R., Halligan, P. W., Marshall, J. C., Frith, C. D., Frackowiak, R. S. J., and Dolan, R. J. 1996. Where in the brain does visual attention select the forest and the trees? *Nature* **382**: 626–628.
- Fink, G. R., Halligan, P. W., Marshall, J. C., Frith, C. D., Frackowiak, R. S. J., and Dolan, R. J. 1997. Neural mechanisms involved in the processing of global and local aspects of hierarchical organized visual stimuli. *Brain* 120: 1779–1791.
- Fink, G. R., Marshall, J. C., Halligan, P. W., and Dolan, R. J. 1999. Hemispheric asymmetries in global/local processing are modulated by perceptual salience. *Neuropsychologia* 37: 31–40.
- Fink, G. R., Marshall, J. C., Halligan, P. W., and Dolan, R. J. 2000. Neuronal activity in early visual areas during global and local processing: A comment on Heinze, Hinrichs, Scholz, Burchert, and Mangun. J. Cogn. Neurosci. 12: 355–356.
- Han, S., Fan, S., Chen, L., and Zhuo, Y. 1997. On the different processing of wholes and parts: A psychophysiological study. J. Cogn. Neurosci. 9: 686-697.
- Han, S., Fan, S., Chen, L., and Zhuo, Y. 1999a. Modulation of brain activities by hierarchical processing: A high-density ERP study. *Brain Topography* 11: 171–183.
- Han, S., and Humphreys, G. W. 1999. Interactions between perceptual organization based on Gestalt laws and those based on hierarchical processing. *Percept. Psychophys.* 6: 1287–1298.
- Han, S., Humphreys, G. W., and Chen, L. 1999b. Parallel and competitive processes in hierarchical analysis: Perceptual grouping and encoding of closure. *J. Exp. Psychol.: Hum. Percept. Perform.* 25: 1411–1432.
- Han, S., He, X., and Woods, D. L. 2000. Hierarchical processing and level-repetition effect as indexed by early brain potentials. *Psychophysiology* 37: 817–830.
- Han, S., He, X., Yund, E. W., and Woods, D. L. 2001a. Attentional selection in the processing of hierarchical patterns: An ERP study. *Biol. Psychol.* **5**: 31–48.
- Han, S., and Humphreys, G. W. 2002. Segmentation and selection contribute to local processing in hierarchical analysis. Q. J. Exp. Psychol. Sect. A 55: 5–21.

- Han, S., Yund, E. W., and Woods, D. L. 2002b. The Role of spatial frequency in the processing of hierarchical patterns: An event-related brain potential study. Under review.
- Han, S., Song, Y., Ding, Y., Yund, E. W., and Woods, D. L. 2001b. Neural substrates for visual perceptual grouping in humans. *Psychophysiology* **38**: 926–935.
- Han, S., Ding, Y., and Song, Y. 2002a. Neural mechanisms of perceptual grouping in humans as revealed by high density event related potentials. *Neurosci. Lett.* **319**: 29–32.
- Heinze, H.-J., Johannes, S., Münte, T. F., and Magun, G. R. 1994. The order of global- and local-level information processing: Electrophysiological evidence for parallel perception processes. In *Cognitive Electrophysiology* (H. Heinze, T. Muente, and G. R. Mangun, Eds.), pp. 1–25. Birkhaeuser, Boston.
- Heinze, H. J., Hinrichs, H., Scholz, M., Burchert, W., and Mangun, G. R. 1998. Neural mechanisms of global and local processing. A combined PET and ERP study. J. Cogn. Neurosci. 10: 485–498.
- Hopfinger, J. B., Buonocore, M. H., and Mangun, G. R. 2000. The neural mechanisms of top-down attentional control. *Nat. Neurosci.* 3: 284–291.
- Kitterle, F. L., Christman, S., and Hellige, J. B. 1990. Hemispheric differences are found in the identification, but not the detection, of low versus high spatial frequencies. *Percept. Psychophys.* 48: 297– 306.
- Mangun, G. R., Heinze, H. J., Scholz, M., and Hinrichs, H. 2000. Neural activity in early visual areas during global and local processing: A reply to Fink, Marshall, Halligan, and Dolan. *J. Cogn. Neurosci.* **12**: 357–359.
- Ivry, R. B., and Robertson, L. C. 1998. Two Sides of Perception. MIT Press, Cambridge, MA.
- Kimchi, R., and Merhav, I. 1991. Hemispheric processing of global form, local form, and texture. *Acta Psychol.* **76**: 133–147.
- Lamb, M. R., Robertson, L. C., and Knight, R. T. 1989. Attention and interference in the processing of global and local information: Effects of unilateral temporal-parietal junction lesions. *Neuropsychologia* 27: 471–483.
- Lamb, M. R., Robertson, L. C., and Knight, R. T. 1990. Component mechanisms underlying the processing of hierarchically organized patterns: Inferences from patients with unilateral cortical lesions. *J. Exp. Psychol. Learning, Memory, Cogn.* 16: 471–483.
- Lamb, M. R., and Yund, E. W. 1993. The role of spatial frequency in the processing hierarchically organized stimuli. *Percept. Psychophys.* **47**: 489–496.
- Lamb, M. R., and Yund, E. W. 1996. Spatial frequency and attention: Effect of level-, target-, and location-repetition on the processing of global and local forms. *Percept. Psychophys.* 58: 363–373.
- Lamb, M. R., Yund, E. W., and Pond, H. M. 1999. Is attentional selection to different levels of hierarchical structure based on spatial frequency? *J. Exp. Psychol. Gen.* **128**: 88–94.

- Martin, M. 1979. Hemispheric specialization for local and global processing. *Neuropsychologia* **17**: 33–40.
- Martinez, A., Moses, P., Frank, L., Buxton, R., Wong, E., and Stiles, J. 1997. Hemispheric asymmetries in global and local processing: Evidence from fMRI. *NeuroReport* 8: 1685–1689.
- Navon, D. 1977. Forest before trees: The precedence of global features in visual perception. *Cogn. Psychol.* **9**: 353–383.
- Polich, J., and Aguilar, V. 1990. Hemispheric local/global processing revisited. *Acta Psychol.* **74:** 47–60.
- Pomerantz, J. R. 1983. Global and local precedence: Selective attention in form and motion perception. J. Exp. Psychol. Gen. 112: 512–540.
- Posner, M. I., Walker, J. A., Friedrich, F. J., and Rafal, R. D. 1984. Effects of parietal injury on covert orienting of attention. *J. Neurosci.* 4: 1863–1874.
- Proverbio, A. M., Minniti. A., and Zani, A. 1998. Electrophysiological evidence of a perceptual precedence of global vs. local visual information. *Cogn. Brain Res.* 6: 321–34.
- Rauschenberger, R., and Yantis, S. 2001. Attentional capture by globally defined objects. *Percept. Psychophy.* **63**: 1250–1261.
- Robertson, L. C., Lamb, M. R., and Knight, R. T. 1988. Effects of lesions of temporal-parietal junction on perceptual and attentional processing in humans. J. Neurosci. 8: 3757–3769.
- Robertson, L. C., and Lamb, M. R. 1991. Neuropsychological contributions to theories of part/whole organization. *Cogn. Psychol.* 23: 299–330.
- Robertson, L. C., Egly, R., Lamb, M. R., and Kerth, L. 1993. Spatial attention and cuing to global and local levels of hierarchical structure. J. Exp. Psychol. Hum. Percept. Perform. 19: 471–487.
- Sasaki, Y., Hadjikhani, N., Fischl, B., Liu, A. K., Marret, S., Dale, A. M., and Tootell, R. B. 2001. Local and global attention are mapped retinotopically in human occipital cortex. *Proc. Natl. Acad. Sci. USA* 98: 2077–2082.
- Sergent, J. 1982. The cerebral balance of power: Confrontation or cooperation? J. Exp. Psychol. Hum. Percept. Perform. 8: 253–272.
- Shulman, G. L., Sullivan, M. A., Gish, K., and Sakoda, W. J. 1986. The role of spatial frequency channels in the perception of local and global structure. *Perception* 15: 259–279.
- Shulman, G. L., and Wilson, J. 1987. Spatial frequency and selective attention to local and global information. *Perception* 16: 89–101.
- Stark, C. L., and Squire, L. R. 2001. When zero is not zero: The problem of ambiguous baseline conditions in fMRI. *Proc. Natl. Acad. Sci. USA* 98: 12760–12766.
- Talairach, J., and Tournoux, P. 1988. *Co-Planar Stereotaxic Atlas of the Human Brain*. Thieme, New York.
- Van Kleeck, M. H. 1989. Hemispheric differences in global versus local processing of hierarchical visual stimuli by normal subjects: New data and a meta-analysis of previous studies. *Neuropsychologia* 27: 1165–1178.