

研究成果の刊行に関する一覧表

書籍

著者氏名	論文タイトル名	書籍全体の 編集者氏名	書籍名	出版社名	出版地	出版年	ページ

雑誌

発表者氏名	論文タイトル名	発表誌名	巻号	ページ	出版年
Shirao, N. <u>Okamoto, Y.</u> Mantani, T. Okamoto, Y. <u>Yamawaki, S.</u>	Gender differences in brain activity toward unpleasant word stimuli concerning body image: an fMRI study.	British Journal of Psychiatry	186	48-53	2005
Yamashita, H. <u>Okamoto, Y.</u> <u>Morinobu, S.</u> <u>Yamawaki, S.</u> Kahkonen, S.	Visual emotional stimuli modulate auditory sensory gating studied by magnetic P50 suppression.	European Archives of Psychiatry and Clinical Neuroscience	255	99-103	2005
Mantani, T. <u>Okamoto, Y.</u> Okada, G. Shirao, N. <u>Yamawaki, S.</u>	Reduced activation of posterior cingulate cortex during imagery in subjects with high degrees of alexithymia: a functional magnetic resonance imaging study.	Biological Psychiatry	57	982-990	2005
Shirao, N. <u>Okamoto, Y.</u> Okada, G. Ueda, K. <u>Yamawaki, S.</u>	Gender differences in brain activity toward unpleasant linguistic stimuli concerning interpersonal relationships: an fMRI study.	European Archives of Psychiatry and Clinical Neuroscience	255	327-333	2005

Kurosaki, M. Shirao, N. Yamashita, H. <u>Okamoto, Y.</u> <u>Yamawaki, S.</u>	Distorted images of one's own body activates the prefrontal cortex and limbic/paralimbic system in young women: a functional magnetic resonance imaging study.	Biol Psychiatry	59	380-386	2006
Kudo, K. Wati, H. Qiao, C. Arita, J. <u>Kanba, S.</u>	Age-related disturbance of memory and CREB phosphorylation in CA1 area of hippocampus of rats.	Brain Research	1054	30-37	2005
Qiao, C-X. Den, R. Kudo, K. Yamada, K. Takemoto, K. Wati, H. <u>Kanba, S.</u>	Ginseng enhances contextual fear conditioning and neurogenesis in rats.	Neurosci Res	51	31-38	2005
Nonaka, M. Doi, T. Fujiyoshi, Y. Takemoto-Kimura, S. <u>Bito, H.</u>	Essential contribution of the ligand-binding bB-bC loop of PDZ1 and PDZ2 in the regulation of postsynaptic clustering, scaffolding and localization of PSD-95.	J. Neurosci.	26	763-774	2006
Iki, J. Inoue, A. <u>Bito, H.</u> Okabe, S.	Bidirectional regulation of postsynaptic cortactin distribution by BDNF and NMDA receptor activity.	Eur. J. Neurosci.	22	2985-2994	2005
Inoue, K. Fukazawa, Y. Ogura, A. <u>Inokuchi, K.</u>	Two-dimensional neural activity mapping of the entire population of hippocampal CA1 pyramidal cells responding to fear conditioning.	Neurosci. Res.	51	417-425	2005

Yamamoto, K. Sakagami, Y. Sugiura, S. <u>Inokuchi, K.</u> Shimohama, S. Kato, N.	Homer-1a enhances spike-induced calcium influx via l-type calcium channels in neocortex pyramidal cells.	Eur. J. Neurosci.	22	1338-1348	2005
Maekawa, M. Takashima, N. Arai, Y. Nomura, T. <u>Inokuchi, K.</u> Yuasa, S. Osumi, N.	Pax6 is required for maintenance and differentiation of progenitor cells in postnatal hippocampal neurogenesis.	Genes to Cell	10	1001-1014	2005
<u>岡本泰昌</u> 上田一貴	ストレス事象の予測に関する脳機能画像解析	医学のあゆみ	212	1115-1119	2005
<u>岡本泰昌</u> <u>山脇成人</u>	うつ病と前頭前野	Clin Neurosciences	23	679-681	2005
<u>田中康雄</u>	発達障害の支援の向こう側-発達障害支援論序説-	教育と医学	630	1137-1145	2005
<u>田中康雄</u>	発達障害と児童虐待 (maltreatment)	子どもの虐待とネグレクト	7	304-312	2005

IV. 研究成果の刊行物・別刷

Gender differences in brain activity generated by unpleasant word stimuli concerning body image: an fMRI study

NAOKO SHIRAO, YASUMASA OKAMOTO, TOMOYUKI MANTANI, YURI OKAMOTO and SHIGETO YAMAWAKI

Background We have previously reported that the temporomesial area, including the amygdala, is activated in women when processing unpleasant words concerning body image.

Aims To detect gender differences in brain activation during processing of these words.

Method Functional magnetic resonance imaging was used to investigate 13 men and 13 women during an emotional decision task consisting of unpleasant words concerning body image and neutral words.

Results The left medial prefrontal cortex and hippocampus were activated only among men, and the left amygdala was activated only among women during the task; activation in the apical prefrontal region was significantly greater in men than in women.

Conclusions Our data suggest that the prefrontal region is responsible for the gender differences in the processing of words concerning body image, and may also be responsible for gender differences in susceptibility to eating disorders.

Declaration of interest None. Funding detailed in Acknowledgement.

Eating disorders, which have been associated with concerns about body shape and size (American Psychiatric Association, 1994), are about 10 times more common in women than in men (Weissman & Olfson, 1995). A possible reason for this difference in susceptibility might be a gender difference in the neural processing of unpleasant information about body image. We previously reported that women showed amygdalar activation while processing unpleasant words concerning body image and perceived these words to be emotionally negative (Shirao *et al*, 2003a). The medial prefrontal cortex has connections to the amygdala, constituting an interaction zone between emotional and cognitive processing (Drevets & Raichle, 1998). In this study we compared the brain activation between men and women while processing these words. We predicted that the amygdala would be less activated and the medial prefrontal cortex more activated in men than in women during the emotional decision task.

METHOD

Study sample

An age-matched sample of 13 men (mean age 25.3 years, s.d.=2.8, range 21–30) and 13 women (mean age 25.2 years, s.d.=3.2, range 21–30) participated in this study ($P=0.949$ by two-tailed, two-sample Student's *t* test). Participants were recruited by community announcement and paid incentives equivalent to their transportation expenses. All of them were right-handed and were native Japanese speakers. Handedness was determined using the Edinburgh Handedness Inventory (Oldfield, 1971). According to self-report, participants had no history of psychiatric, neurological or other major medical illness, and had never been treated with a psychotropic medication. There was no significant difference in the average years of education between

men and women: men 15.2 (s.d.=1.6) *v.* women 14.9 (s.d.=2.5); $P=0.645$ by two-tailed, two-sample Student's *t*-test. The average body mass index of the men was 22.4 kg/m² (s.d.=3.2, range 18.0–31.3) and that of the women was 21.5 kg/m² (s.d.=3.7, range 18.8–28.4); $P=0.543$ by two-tailed two-sample Student's *t*-test. The average of the total Eating Disorder Inventory – 2 (EDI–2; Garner, 1991) scores of men was 45.5 (s.d.=28.4, range 9–103) and that of women was 37.9 (s.d.=23.5, range 7–85); $P=0.330$ by two-tailed Wilcoxon single-rank test. The average score for the item 'body dissatisfaction' for the men was 7.43 (s.d.=5.45, range 2–19) and for the women was 11.31 (s.d.=7.00, range 0–22); $P=0.330$ by two-tailed Wilcoxon single-rank test. The study was conducted using a protocol approved by the ethics committee of Hiroshima University School of Medicine. All individuals provided written informed consent for participation in the study.

Emotional decision task

We used the emotional decision task developed by Tabert *et al* (2001), with some modifications. The words used in the task were selected from the database of Toglia & Battig (1978), which includes 2854 words that have been rated on several items such as familiarity and pleasantness, on a scale of 1 (very unfamiliar; very unpleasant) to 7 (very familiar; very pleasant), with 4 as the mid-point. For our study, 30 neutral words were selected from the database and translated into Japanese. We also selected 30 highly unpleasant words concerning body image, chosen from Japanese-language dictionaries and thesauri. The two groups of words did not significantly differ with regard to word length (mean length in Japanese letters: body image words 3.2, neutral words 3.1; $P=0.575$ by two-tailed, two-sample Student's *t*-test). Our previous validation study comparing women who had eating disorders with a control group of healthy women showed that there was no significant difference in familiarity between the two categories of words (eating disorder group mean familiarity score: body image words 4.2; neutral words 4.1, $P=0.727$; control group mean familiarity score: body image words 3.9, neutral words 4.1, $P=0.218$, by two-tailed Wilcoxon single-rank test) and there was no significant difference in the familiarity ratings of words concerning

body image between women with eating disorders and the control group ($P=0.365$ by two-tailed Wilcoxon single-rank test), whereas there were significant differences in pleasantness between the two categories of words (mean pleasantness score in the eating disorder group: body image words 2.4, neutral words 3.9, $P=0.0002$; mean pleasantness score in the control group: body image words 3.0, neutral words 4.0, $P=0.0001$, by two-tailed Wilcoxon single-rank test) and there were significant differences in the ratings of pleasantness between the eating disorders group and the control group ($P=0.030$ by two-tailed Wilcoxon single-rank test) (Shirao *et al*, 2003b). Both lists of words contained nouns, verbs, adjectives and adverbs.

The selected words were used to generate sets of unpleasant words concerning body image and sets of neutral words. Each word set comprised a unique combination of three words. The word sets were presented in six alternating blocks of two conditions (the task condition and the control condition) in three cycles (Fig. 1). During the task condition unpleasant word sets were presented, and during the control condition neutral word sets were presented. Each block began with a 3 s cue identifying the condition by displaying the word 'task' or 'control'. Five word sets were presented

in each block. Each word set was shown for 4 s with a 1.4 s interstimulus interval (Fig. 1). The blood oxygen level-dependent (BOLD) response was recorded during three blocks of unpleasant words and three blocks of neutral words. During each inter-stimulus interval, a fixation cross placed centrally on the screen replaced the word set. Baseline functional magnetic resonance images were obtained during a 9 s period prior to the first block of trials, during which the individual viewed a centrally placed fixation cross. During each trial, the word set was projected to the centre of the person's field of view by a Super Video Graphics adapter computer-controlled projection system. The timing of presentation of word sets was controlled by Presentation Software Version 0.51 (Neurobehavioral Systems, Inc., San Francisco, CA, USA) and the word sets were presented in a randomised order. Immediately before functional magnetic resonance imaging (fMRI) scanning was begun, each participant was given ten practice trials (five unpleasant word sets and five neutral word sets). The words presented in the practice trials did not overlap with the experimental words.

Participants were instructed to select the most unpleasant word from each set of unpleasant words based on their personal knowledge and experience, and for each set

of neutral words, participants were instructed to select the word that they thought was the most neutral; they indicated their choice by pressing one of three buttons on a response pad in the MRI scanner.

Image acquisition and processing

The MRI scanner used was a Magnex Eclipse 1.5 T Power Drive 250 (Shimadzu Medical Systems, Kyoto, Japan). A time-course series of 63 volumes was acquired with T_2^* -weighted, gradient echo, echo planar imaging (EPI) sequences. Each volume consisted of 28 slices, each 4.0 mm thick with no gap, encompassing the entire brain. The interval between two successive acquisitions of the same image (time to repetition, TR) was 3000 ms, the time to echo (TE) was 55 ms and the flip angle was 90° . The field of view was 256 mm and the matrix size 64×64 , giving voxel dimensions of $4.0 \text{ mm} \times 4.0 \text{ mm} \times 4.0 \text{ mm}$. After fMRI scanning, structural scans were acquired using a T_1 -weighted gradient echo pulse sequence (TR 12 ms, TE 4.5 ms, flip angle 20° , field of view 256 mm, voxel dimensions $1.0 \text{ mm} \times 1.0 \text{ mm} \times 1.0 \text{ mm}$), to facilitate localisation and co-registration of the functional data.

Image processing and statistical analysis were performed using Statistical Parametric Mapping (SPM) 99 software (Wellcome Department of Cognitive Neurology, London, UK) implemented in Matlab (Mathworks, Inc., Natick, MA, USA). The first two volumes of the fMRI run (pre-task period) were discarded because the magnetisation was unsteady, and the remaining 61 volumes were used for the statistical analysis. Images were corrected for motion and realigned with the first scan of the session, which served as the reference. The T_1 anatomical images were co-registered to the first functional images in each individual and aligned to a standard stereotaxic space, using the Montreal Neurological Institute (MNI) T_1 template in SPM99. The calculated non-linear transformation was applied to all functional images for spatial normalisation. Finally, the fMRI images were smoothed with a 12 mm full-width, half-maximum Gaussian filter.

Using group analysis according to a random effect model that allowed inference to the general population (Friston *et al*, 1999), we first identified brain regions that showed a significantly greater response to unpleasant word sets in comparison with the response to neutral word sets among

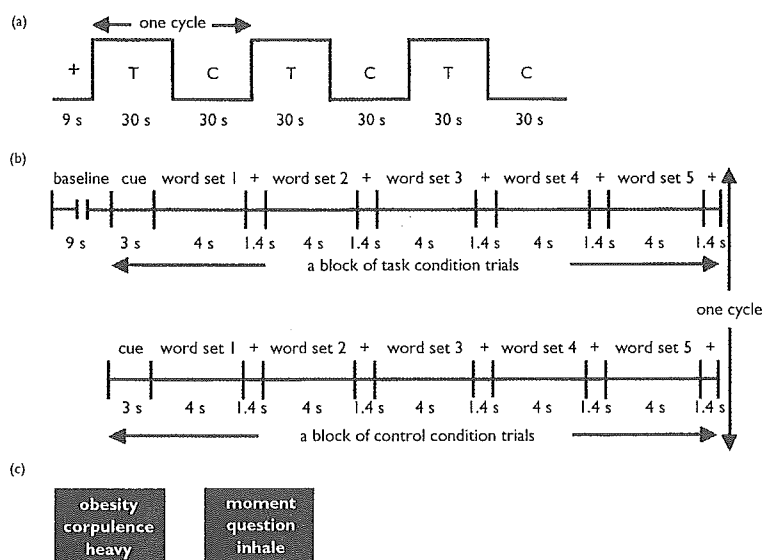


Fig. 1 Design of the study task. (a) Six alternating blocks of task condition (T) trials and control condition (C) trials were presented successively; the total scan time was 189 s (3 min and 9 s), yielding 63 images of 28 axial slices (1764 images). (b) Blocks of task condition and control condition trials were preceded by a baseline imaging period. Each block began with a cue ('task' or 'control'). The participant selected the word judged to be the most unpleasant or most neutral in each word set, by pressing one of three buttons. (c) Translations of typical word sets presented in this study (left-hand block, task condition; right-hand block, control condition).

male and among female participants, as brain areas related to the cognition of unpleasant word stimuli concerning body image in men and women, respectively. We then took the data of 13 of the 15 women who had participated in our previous study (Shirao *et al*, 2003a) and directly compared the activation of the entire brain in the male and female sub-samples using the two-sample Student's *t*-test. The resulting set of voxel values for each contrast constituted an SPM(t) map. The SPM(t) maps were then interpreted by referring to the probabilistic behaviour of Gaussian random fields. The data were given an initial threshold at an uncorrected $P < 0.001$ at the voxel level, and regions about which we had an *a priori* hypothesis were reported at this threshold (Elliott *et al*, 2000). For regions about which there was no clear hypothesis, a more stringent threshold of $P < 0.05$ corrected at the cluster level of multiple comparison was used. The *x*, *y* and *z* coordinates provided by SPM, which were in MNI brain space, were converted to the *x*, *y* and *z* coordinates in Talairach & Tournoux's (TT) brain space (Talairach & Tournoux, 1988) using the following formulae:

$$(a) x_{TT} = x_{MNI} \times 0.88 - 0.8;$$

$$(b) y_{TT} = y_{MNI} \times 0.97 - 3.32;$$

$$(c) z_{TT} = y_{MNI} \times 0.05 + z_{MNI} \times 0.88 - 0.44.$$

Labels for brain activation foci were obtained in Talairach coordinates using the Talairach Daemon software (Research Imaging Center, University of Texas, TX, USA), which provides accuracy similar to that of neuroanatomical experts (Lancaster *et al*, 2000). The labelling of areas given by this software was then confirmed by comparison with activation maps overlaid on MNI-normalised structural images.

Evaluation of pleasantness and familiarity of the word stimuli

Each participant was asked to rate the pleasantness and familiarity of all the words presented in the tasks on a scale from 1 (very unfamiliar; very unpleasant) to 7 (very familiar; very pleasant), immediately after scanning. For this rating procedure the list of words was presented in randomised order in a table format.

RESULTS

Rating of words

The ratings of familiarity with the two categories of words did not significantly

differ among men (mean familiarity score: unpleasant words 3.8, neutral words 4.4, $P=0.054$ by two-tailed Wilcoxon single-rank test) or women (mean familiarity score: unpleasant words 4.3, neutral words 4.3, $P=0.456$). However, all participants rated the unpleasant words concerning body image as significantly more unpleasant than the neutral words (mean pleasantness score: unpleasant words 3.1, neutral words 4.1, $P=0.007$ in men; unpleasant words 2.7, neutral words 4.1, $P=0.002$ in women). Neither the ratings of pleasantness nor the ratings of familiarity in each word category significantly differed between the male and female groups.

Brain activation

In men there was significantly greater activation of the left hippocampus, left superior temporal gyrus, left fusiform gyrus and left medial frontal gyrus when the emotional decision task involved unpleasant words compared with neutral words, whereas the women showed significantly greater activity of the left parahippocampal gyrus including amygdala, left thalamus and right caudate body in the same comparison (Table 1, Fig. 2).

The two-sample Student's *t*-test revealed that there was a significantly higher BOLD response in the left apical prefrontal region in men than in women during the

unpleasant word task compared with neutral word task (Table 1, Fig. 3). No brain area showed significantly higher activation in women than in men during any of the tasks.

Correlation between psychological data and brain activation

Among the 13 women participants, activation in the left apical prefrontal area, which was significantly lower than that in men during the unpleasant words task, was negatively correlated with the total EDI-2 score (Spearman's rank-order correlation analysis: correlational coefficient -0.699 , $P=0.008$). There was no correlation between any brain area showing significant BOLD response and the EDI-2 scores or the pleasantness rating of the unpleasant words.

DISCUSSION

We used the emotional decision task to examine the brain areas engaged in the perception of unpleasant words concerning body image and to compare the patterns of brain activation in men and women. Our results showed that the left medial part of the frontal gyrus, the left limbic area excluding the amygdala, the left superior temporal gyrus and the left fusiform gyrus play an important part in processing

Table 1 Relative increases in brain activity associated with unpleasant words concerning body image (task) and neutral words (control)

	Cluster	BA	t score	Coordinates ¹		
				x	y	z
Men (n=13)						
Left hippocampus	696*		9.59	-32	-13	-13
Left superior temporal gyrus		21	6.54	-50	-7	-8
Left fusiform gyrus		20	6.35	-43	-25	-17
Left medial frontal gyrus	359*	9	5.71	-4	53	9
Left superior frontal gyrus		10	5.34	-15	51	18
Women (n=13)						
Left parahippocampal gyrus	404*	37	7.08	-17	-13	-15
Left thalamus	485*		6.08	-3	-11	11
Right caudate body			4.65	10	1	5
Men > women						
Left apical prefrontal region	144	9	4.36	-15	49	20

BA, Brodmann area.

1. Stereotaxic coordinates were derived from Talairach & Tournoux (1988) and refer to the medial-lateral position (*x*) relative to the midline (positive=right), anterior-posterior position (*y*) relative to the anterior commissure (positive=anterior) and superior-inferior position (*z*) relative to the commissural line (positive=superior).

*Areas exceeding the extent threshold of $P < 0.05$ corrected at the cluster level, all other areas exceeding the height threshold of $P < 0.001$ uncorrected at the cluster level and belonging to a cluster of activation with an extent of at least 140 voxels are displayed.

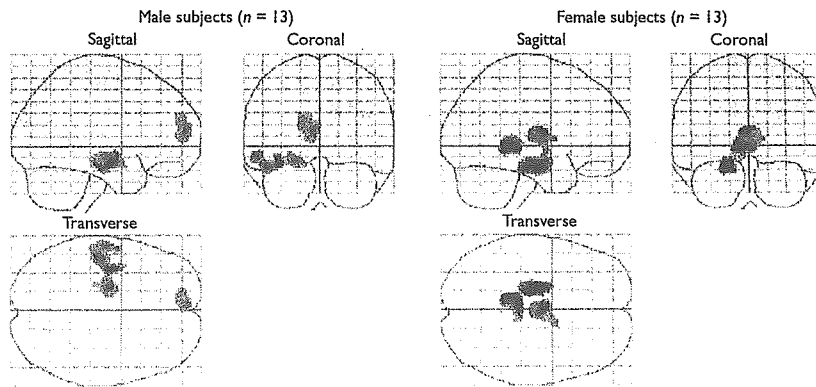


Fig. 2 Brain areas showing significantly greater activation during the task condition compared with the control condition. Three-dimensional 'look-through' projections of statistical parametric maps of the brain regions are shown (one-sample Student's *t*-test; corrected $P < 0.05$ at the cluster level; $n=13$; $d.f.=12$).

unpleasant words concerning body image in men.

Lack of amygdalar activation in men

Consistent with our hypothesis, the amygdala did not show significant activation among men; however, the gender difference of the BOLD response in the amygdala was not significant by two-sample Student's *t*-test.

The amygdala has been suggested by many studies to be strongly associated with stimuli signalling threat. Human lesion and imaging studies consistently indicate that the amygdala is concerned in fear conditioning (Morris *et al*, 1998), in the recognition of fearful facial expressions (Adolphs, 1999) and in the evocation of fearful emotional responses from direct

stimulation (Halgren *et al*, 1978). The amygdala is also considered to be important in the detection of environmental threat (Scott *et al*, 1997), including verbal stimuli (Isenberg *et al*, 1999). Therefore, the lack of significant activation in the amygdala among men suggests that men may not process unpleasant words concerning body image as fearful information, whereas women seem to do so.

Medial prefrontal cortex and emotional processing

The significant activation in the medial part of the frontal gyrus – Brodmann areas (BAs) 9 and 10; medial prefrontal cortex – was only detected in men, and there was a significantly higher BOLD response in men than in women in the left apical prefrontal region (BA 9) when performing the

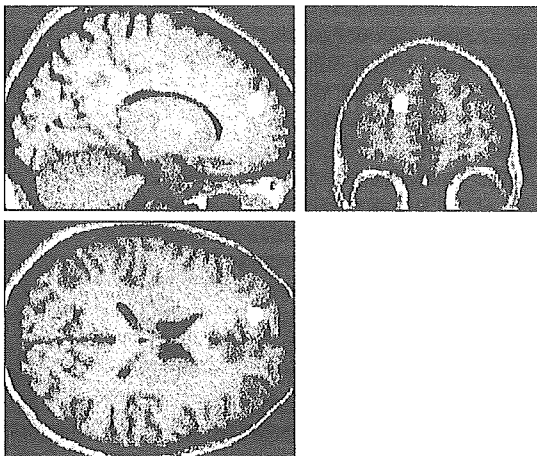


Fig. 3 Brain regions showing significantly greater activation in men than in women during the task condition of the emotional decision task compared with the control condition. Clusters of activation are overlaid onto a T_1 -weighted anatomical magnetic resonance image. The white spots show areas of high activation. Two-sample Student's *t*-test; uncorrected $P < 0.001$ in height; $n=26$ (13 men, 13 women); $d.f.=24$.

unpleasant word task compared with the neutral word task by two-sample Student's *t*-test. These results were consistent with our hypothesis. Many previous studies have suggested that the medial prefrontal cortex might have a role generally in emotional processing. It is reported that visual stimuli that evoke emotions, such as films or pictures, activated the medial prefrontal cortex, and that recall of various emotions such as happiness, sadness and disgust, and a mixture of these emotions, all separately engaged this brain region (Lane *et al*, 1997; Reiman *et al*, 1997). Several more recent studies suggest that when people turn their attention inwards to assess self-relevant attributes or emotional awareness, activity increases in the medial prefrontal cortex (Johnson *et al*, 2002; Zysset *et al*, 2002). The medial prefrontal cortex has connections to limbic structures, including the amygdala, constituting an interaction zone between emotional processing and cognitive processing (Drevets & Raichle, 1998), and this region may have a role in modulating the emotional response in the amygdala and other limbic structures. Limbic structures, including the amygdala, are likely to respond to emotional stimuli at a sensory or perceptual level (Reiman *et al*, 1997), whereas the medial prefrontal cortex may be involved in the cognitive aspects of emotional processing, such as attention to emotion, appraisal or identification of emotion (Drevets & Raichle, 1998). From this viewpoint, the gender differences detected in our study may demonstrate differences of cognitive pattern in men and women. Our results suggest the possibility that men processed the emotional decision task including words concerning body image more cognitively rather than emotionally, and activation in the medial prefrontal cortex was prominent; on the other hand, women processed this task more emotionally rather than cognitively, and the medial prefrontal cortex did not exhibit any significant activation. Both men and women perceived the unpleasantness of the words concerning body image to the same degree, according to their subjective ratings, but the fMRI data suggest that their processes are different: women are likely to use more intuitive processing whereas men use more rational processing. This discrepancy between the genders in cognitive style related to body image may contribute to the large gender difference in susceptibility to eating disorders.

Another possible explanation of the different patterns of activation in the medial prefrontal cortex between men and women may be the difference in men's familiarity with the unpleasant word set compared with the neutral words. Although the ratings of familiarity were not different between men and women ($P=0.133$ by Mann-Whitney U test), there was a trend for male participants to be less familiar with the unpleasant words concerning body image than with the neutral words ($P=0.054$ by two-tailed Wilcoxon single-rank test). When processing unfamiliar words concerning body image, men might turn more attention inwards, and subsequently the BOLD response in the medial prefrontal cortex was higher than while processing neutral words.

Among women, correlational analysis revealed that the BOLD response in the left apical prefrontal region (BA 9), which was significantly lower in women than in men, was negatively correlated with total EDI-2 scores; in other words women with higher EDI-2 scores exhibited lower activity in this brain area. These results suggest the possibility that the apical prefrontal region might be involved in the pathophysiology of eating disorders.

Comparison with other neuroimaging studies

To our knowledge, two fMRI studies concerning body image distortion have investigated the effects of pictorial body image stimuli in women with anorexia nervosa and healthy controls (Seeger *et al*, 2002; Wagner *et al*, 2003). One study reported that patients with anorexia nervosa showed activation in the right amygdala, right fusiform gyrus and brain-stem associated with stimulation with their own body image whereas healthy controls showed activation only in the fusiform gyrus (Seeger *et al*, 2002), and the other reported that patients with anorexia nervosa showed greater activation in the prefrontal cortex and the inferior parietal lobule than did controls (Wagner *et al*, 2003). The latter authors explain the discrepancy between these results as a consequence of the design of the task. Many differences in the experimental conditions between these studies and ours make it difficult to compare the brain activation data, but a possible explanation of the discrepancy between the study by Wagner *et al* (2003) and our study is the age of the participants: those in the former

CLINICAL IMPLICATIONS

- Gender differences in brain activation suggest differences between men and women in the style of cognition toward unpleasant stimuli concerning body image.
- This discrepancy in cognitive style may have relevance to the large gender difference in susceptibility to eating disorders.
- The medial prefrontal cortex may be the brain area linked to the pathophysiology of eating disorder.

LIMITATIONS

- We did not use a structured interview when selecting participants.
- We asked the participants to rate only pleasantness and familiarity of the word stimuli and we could find no clear relationship between brain activation and the subjective rating of the words concerning body image.
- It is unclear whether the patterns of activation in the prefrontal area were specific to the stimuli concerning body image.

NAOKO SHIRAO, MD, PhD, YASUMASA OKAMOTO, MD, PhD, TOMOYUKI MANTANI, MD, Department of Psychiatry and Neurosciences, Graduate School of Biomedical Sciences, Hiroshima University, Hiroshima, and Core Research for Evolutional Science and Technology (CREST), Japan Science and Technology Corporation, Seika; YURI OKAMOTO, MD, PhD, Hiroshima University Health Service Center, Hiroshima; SHIGETO YAMAWAKI, MD, PhD, Department of Psychiatry and Neurosciences, Graduate School of Biomedical Sciences, Hiroshima University, and CREST, Seika, Japan

Correspondence: Dr Shigeto Yamawaki, Department of Psychiatry and Neurosciences, Division of Frontier Medical Science, Programs for Biomedical Research, Graduate School of Biomedical Sciences, Hiroshima University, 1-2-3 Kasumi, Minami-ku, Hiroshima 734-8551, Japan. Tel: +81 82 257 5207; fax: +81 82 257 5209; e-mail: yamawaki@hiroshima-u.ac.jp

(First received 4 March 2004, final revision 10 August 2004, accepted 11 August 2004)

study were adolescents (approximately 15 years old), whereas we recruited young adults (approximately 25 years old). An fMRI study which investigated the brain activation of adult and adolescent men and women while processing emotional facial expressions reported that the adult men and adolescents (both boys and girls) showed significant activation in the bilateral orbitofrontal cortex and anterior cingulate cortex in response to an angry face, whereas the adult women showed significant activation in the left amygdala in addition to these brain areas (McClure *et al*, 2004). These results suggest that the patterns of neural responses to emotional stimuli may be different in adults and adolescents.

A positron emission tomography study of gender differences in brain activation patterns during recognition of emotional facial expressions revealed that greater amygdalar activation was observed in women and greater medial frontal

activation was observed in men (Hall *et al*, 2004); these authors suggest that men might take a more analytic approach and might regulate their emotional reaction to the stimuli more than women. Although the categories of stimuli are different, these results support our findings.

Study limitations

Our study has some limitations. First, we did not administer a structured interview when selecting the participants; however, they had no psychiatric or neurological illness at the time of their participation, although we cannot rule out its occurrence in the future. Second, participants were asked to rate only the unpleasantness and familiarity of the words used. If we had also asked about the fearfulness induced by the stimuli, we might have found gender differences in subjective rating and the results with brain image data would have been more clear-cut. Last, although our data

suggest that there is differential activation of the brains of men and women when processing unpleasant words concerning body image, we cannot conclude whether these results are specific to unpleasant stimuli concerning body image or would apply to a wide range of unpleasant stimuli. Among women, a lower BOLD response in the prefrontal region compared with men while processing unpleasant words concerning body image exhibited a negative correlation with the total EDI-2 score, but it is unclear whether this brain region is the focal area responsible for susceptibility to eating disorders.

In conclusion, our study revealed that the paralimbic area including the amygdala was activated only in women and that the left medial prefrontal cortex was activated only in men while performing the emotional decision task with unpleasant words concerning body image. These results suggest that gender differences in brain activation might explain the differences in the style of cognition towards unpleasant stimuli concerning body image. Further studies comparing people who have eating disorders with healthy controls and which include general unpleasant word stimuli to contrast with words specific to body image are needed to elucidate the neural substrate responsible for the onset of eating disorders.

ACKNOWLEDGEMENT

The study was supported by the Research on Psychiatric and Neurological Diseases and Mental Health, Ministry of Health, Labour and Welfare, Japan.

REFERENCES

- Adolphs, R. (1999)** Social cognition and the human brain. *Trends in Cognitive Science*, **3**, 469–479.
- American Psychiatric Association (1994)** *Diagnostic and Statistical Manual of Mental Disorders* (4th edn) (DSM-IV). Washington, DC: APA.
- Drevets, W. C. & Raichle, M. E. (1998)** Reciprocal suppression of regional cerebral blood flow during emotional versus higher cognitive process: implications for interaction between cognition and emotion. *Cognition and Emotion*, **12**, 353–385.
- Elliott, R., Friston, K. J. & Dolan, R. J. (2000)** Dissociable neural responses in human reward systems. *Journal of Neuroscience*, **20**, 6159–6165.
- Friston, K. J., Holmes, A. P. & Worsley, K. J. (1999)** How many subjects constitute a study? *Neuroimage*, **10**, 1–5.
- Garner, D. M. (1991)** *Eating Disorder Inventory – 2 (EDI-2)*. Odessa, FL: Psychological Assessment Resources.
- Halgren, E., Walter, R. D., Cherlow, D. G., et al (1978)** Mental phenomena evoked by electrical stimulation of the human hippocampal formation and amygdala. *Brain*, **101**, 83–117.
- Hall, G. B., Witelson, S. F., Szechtman, H., et al (2004)** Sex differences in functional activation patterns revealed by increased emotion processing demands. *Neuroreport*, **15**, 219–223.
- Isenberg, N., Silbersweig, D., Engelen, A., et al (1999)** Linguistic threat activates the human amygdala. *Proceedings of the National Academy of Science USA*, **96**, 10456–10459.
- Johnson, S. C., Baxter, L. C., Wilder, L. S., et al (2002)** Neural correlates of self-reflection. *Brain*, **125**, 1808–1814.
- Lancaster, J. L., Woldorff, M. G., Parsons, L. M., et al (2000)** Automated Talairach atlas labels for functional brain mapping. *Human Brain Mapping*, **10**, 120–131.
- Lane, R. D., Reiman, E. M., Bradley, M. M., et al (1997)** Neuroanatomical correlates of pleasant and unpleasant emotion. *Neuropsychologia*, **35**, 1437–1444.
- McClure, E. B., Monk, C. S., Nelson, E. E., et al (2004)** A developmental examination of gender differences in brain engagement during evaluation of threat. *Biological Psychiatry*, **55**, 1047–1055.
- Morris, J. S., Ohman, A. & Dolan, R. J. (1998)** Conscious and unconscious emotional learning in the human amygdala. *Nature*, **393**, 467–470.
- Oldfield, R. C. (1971)** The assessment and analysis of handedness: the Edinburgh inventory. *Neuropsychologia*, **9**, 97–113.
- Reiman, E. M., Lane, R. D., Ahern, G. L., et al (1997)** Neuroanatomical correlates of externally and internally generated human emotion. *American Journal of Psychiatry*, **154**, 918–925.
- Scott, S. K., Young, A. W., Calder, A. J., et al (1997)** Impaired auditory recognition of fear and anger following bilateral amygdala lesions. *Nature*, **385**, 254–257.
- Seeger, G., Braus, D. F., Ruf, M., et al (2002)** Body image distortion reveals amygdala activation in patients with anorexia nervosa – a functional magnetic resonance imaging study. *Neuroscience Letters*, **326**, 25–28.
- Shirao, N., Okamoto, Y., Okada, G., et al (2003a)** Temporomesial activation in young females associated with unpleasant words concerning body image. *Neuropsychobiology*, **48**, 136–142.
- Shirao, N., Okamoto, Y., Okamoto, Y., et al (2003b)** Ratings of negative body image words, negative emotion words and neutral words by eating disorder patients and healthy subjects. *Brain Sciences and Mental Disorders*, **15**, 141–147.
- Tabert, M. H., Borod, J. C., Tang, C. Y., et al (2001)** Differential amygdala activation during emotional decision and recognition memory tasks using unpleasant words: an fMRI study. *Neuropsychologia*, **39**, 556–573.
- Talairach, J. & Tournoux, P. (1988)** *Co-planar Stereotaxic Atlas of the Human Brain*. Stuttgart: Thieme.
- Toglia, M. P. & Battig, W. F. (1978)** *Handbook of Semantic Word Norms*. Hillsdale, NJ: Lawrence Erlbaum.
- Wagner, A., Ruf, M., Braus, D. F., et al (2003)** Neuronal activity changes and body image distortion in anorexia nervosa. *Neuroreport*, **14**, 2193–2197.
- Weissman, M. M. & Olfson, M. (1995)** Depression in women: implications for health care research. *Science*, **269**, 799–801.
- Zysset, S., Huber, O., Ferstl, E., et al (2002)** The anterior frontomedian cortex and evaluative judgment: an fMRI study. *Neuroimage*, **15**, 983–991.

Hidehisa Yamashita · Yasumasa Okamoto · Shigeru Morinobu · Shigeto Yamawaki · Seppo Kähkönen

Visual emotional stimuli modulation of auditory sensory gating studied by magnetic P50 suppression

Received: 18 November 2003 / Accepted: 11 June 2004 / Published online: 12 November 2004

Abstract The auditory sensory gating system modulates its sensitivity to incoming stimuli and prevents higher brain functions from sensory overload in the primary auditory cortex. We investigated whether visually evoked emotional stimuli affect auditory sensory gating. Magnetic P50 (P50m) suppression was evaluated by magnetoencephalography in fifteen healthy subjects while they viewed slides varying in emotional valence and arousal. The ratio of strength of dipole moments of the 2nd to the 1st P50m and the anatomical location of their sources were calculated. Negatively valenced slides significantly attenuated P50m suppression, as compared to neutral ones, while the effects of positive slides were insignificant. No effects on latencies or the location of P50m sources were observed. Thus, negative emotional stimuli may modulate sensory gating.

Key words emotion · auditory sensory gating · magnetoencephalography (MEG) · P50 suppression

H. Yamashita (✉)
1-2-3, Kasumi, Minami-ku
Hiroshima 734-8551, Japan
Tel.: +81-82/257-5208
Fax: +81-82/257-5209
E-Mail: hidehisa@hiroshima-u.ac.jp

H. Yamashita · Y. Okamoto · S. Morinobu · S. Yamawaki
Department of Psychiatry and Neurosciences
Graduate School of Biomedical Sciences
Hiroshima University, Japan

H. Yamashita · Y. Okamoto · S. Morinobu · S. Yamawaki
Core Research for Evolutional Science and Technology (CREST) of
Japan Science and Technology Corporation (JST)

S. Kähkönen
BioMag Laboratory
Medical Engineering Center
Helsinki University Central Hospital, Finland

S. Kähkönen
Cognitive Brain Research Unit
Department of Psychology
University of Helsinki, Finland

Introduction

Sensory gating is defined as the pre-attentive ability of the brain to modulate its sensitivity to an incoming stimulus, and is hypothesized to be a protective mechanism that prevents sensory overload of higher brain functions by filtering out the irrelevant sensory input (Braff and Geyer 1990). Deficit in sensory gating could result in an overload of irrelevant stimuli, which in turn may lead to perceptual and attentional impairments associated with psychiatric disorders such as schizophrenia (McGhie and Chapman 1961).

A paired click paradigm is used to evaluate the auditory sensory gating. In this paradigm, two identical stimuli (1st: conditioning stimulus and 2nd: test stimulus) are presented with a short inter-stimulus interval (ISI) of 500 ms and a longer inter-pair interval. The P50 appears as a positive peak in electroencephalography (EEG), at about 50 ms after the stimulus onset. Under normal conditions the amplitude of the P50 for test stimuli is smaller than that for conditioning stimuli, and this suppression of P50 is typically quantified as a ratio (S2/S1). This phenomenon is referred to as P50 suppression (Adler et al. 1982).

Magnetoencephalography (MEG) offers a non-invasive method for functional brain studies with high temporal resolution equal to that of EEG, but it enables more accurate source localization. MEG measures selectively the activity from tangential sources and is well suited for the measurement and localization of primary auditory cortical activity (Hämäläinen et al. 1993). MEG studies indicate that the magnetic P50 (P50m) counterpart is generated in the superior aspects of temporal lobes, near the primary auditory cortex (Hari et al. 1980; Reite et al. 1988; Mäkelä et al. 1994).

Cognitive and affective processing are two basic interacting modes of information processing (LeDoux 1993). Several studies have shown that emotional visual stimuli can have an effect on visual evoked potentials (VEPs), especially late component P300 in healthy sub-

jects (Lang et al. 1990; Laurian et al. 1991; Kayser et al. 1997; Cuthbert et al. 2000). Only a few studies have investigated the effect of visual emotional stimuli on auditory information processing and have shown that emotional stimuli may in fact affect auditory processing (Schupp et al. 1997; Surakka et al. 1998). However, the exact role of the interaction between emotion and cognition remains to be elucidated. In this study, we examined whether visually evoked emotional stimuli could affect auditory sensory gating, as measured by P50m suppression in healthy subjects.

Methods

Subjects

Fifteen healthy, right-handed volunteers (14 males and 1 female), aged 22–38 years (mean age 29.5 ± 5.1 y), participated in the experiment consisting of three sessions in a randomized order. The subjects reported having no history of neurological or psychiatric disorders or of any drug use for 2 weeks before the study. Because acute nicotine ingestion within 0.5 h of testing may alter P50 suppression, subjects were not permitted to smoke at least 1 h before until the end of the measurement (Adler et al. 1992). Informed consent was obtained from each subject according to institutional guidelines, and the study was approved by an institutional ethical committee.

Task procedure

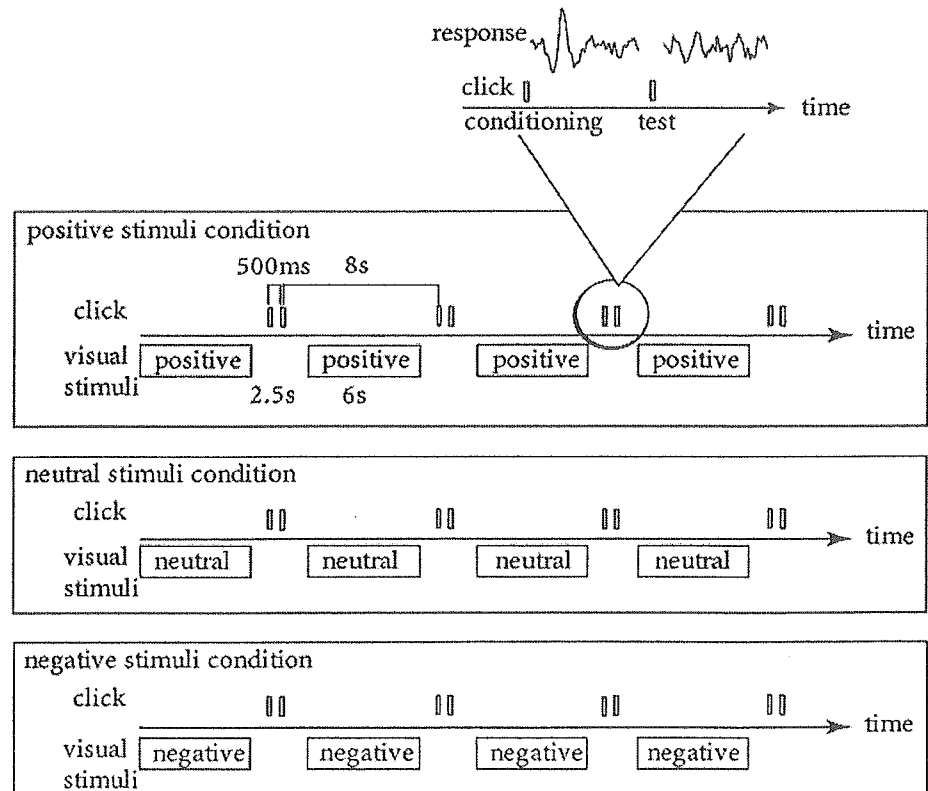
During the MEG recording, the subject sat in a comfortable chair in a magnetically shielded room. The auditory evoked magnetic fields

were recorded with a 204 channel MEG (Neuromag Ltd., Finland). The paired-click paradigm with an inter-click interval of 500 ms with click pairs (0.1 ms, 60 dB above the individually determined subjective hearing threshold) separated by 8 s inter-pair was used. The stimuli were delivered binaurally to subjects through a non-magnetic, echo-free plastic tube system, because all previous studies which investigated P50 suppression have used bilateral stimuli (Adler et al. 1982; Braff et al. 1990; Adler et al. 1992; Light et al. 1999; Patrick et al. 1999; Adler et al. 2001), and the ipsilateral P50 response to monoaural stimuli is so small that it is difficult to detect. Three hundred digitized pictures (one hundred per category) were chosen from the International Affective Picture System (IAPS) (Lang et al. 1997). The categories were negative, neutral, and positive (e.g., mutilations, buildings, and pleasant landscapes, respectively). The visual stimuli were presented by projection onto a screen during the interval of click pair presentation for 6 s, from 1 s after the second click to 1 s before the first click of the next pair (Fig. 1). The viewing distance was 3 m, and light was dimmed during the measurement. Three sessions were recorded for 15 minutes per session in a randomized order. The subjects rated their experiences evoked by stimuli, on two dimensions: valence and arousal (Lang et al. 1997). The ratings were made on a nine-point visual analog scale. The valence scale varied from unhappy to happy. The arousal scale varied from calm to highly arousing.

Neuroimaging data collection

The position of the subject's head relative to the recording instrument was determined by measuring the magnetic fields produced by marker coils in relation to cardinal points of the head (nasion, left and right pre-auricular points), which were determined before the experiment using an Isotrack 3D-digitizer (Polhemus, Colchester, VT, U.S.A.) (Ahlfors and Ilmoniemi 1989). The recording passband was 0.1–200 Hz for MEG and EOG, and the sampling rate was 600.025 Hz. Digital band-pass filtering was performed off-line at 5–50 Hz (Light et al. 1999). The first few responses and the entire epoch coinciding with EOG or MEG changes exceeding $150 \mu\text{V}$ or 3000 ft/cm, respec-

Fig. 1 Schematic diagram of the stimulus sequences in the three experimental conditions. Auditory stimuli (60 dBHL, click: depicted as bars) were presented as trains of pairs in a conditioning (1st)–testing (2nd) paradigm. The inter-pair interval was 500 ms and the intra-pair interval was 8 s. The rectangles below the sounds represent visual stimuli. The visual stimuli were presented during the interval of click pairs presentation for 6 s, from 1 s after the second click to 1 s before the first click of the next pairs. Three sessions (negative, neutral, and positive slide conditions) were recorded separately



tively, were omitted from averaging. An epoch lasted 550 ms, including a 100 ms prestimulus baseline. Electrodes were attached below the right eye and above the left eye to minimize potential electrooculogram artifacts. The electrode impedances were below 5 k Ω . Subjects were monitored visually for signs of sleep. At least, 70 responses were averaged in each condition.

MEG source localization

All analyses were conducted blind to the session condition. An individual sphere model of the head was constructed from the local radius of curvature on the basis of individual MRI images. MRI was performed using a 1.5-T apparatus (General Electric Co., Milwaukee, WI, USA). The P50m peaks were obtained from the latency ranges of 35–80 ms after the stimulus presentation. The latency, location, and strength of the P50m source were analyzed with single equivalent current dipole modeling, determined by a least-squares fit using a subset of 34 channels separately over each auditory cortex (Hämäläinen et al. 1993). Dipole fits with at most 40% residual variance and with at most 4186 mm³ confidence volume (10 mm radius sphere) were considered successful. The latency, location and dipole moments of P50m for conditioning stimuli (Qc) and test stimuli (Qt), and the t/c ratio (Qt/Qc) in the positive and negative slide conditions were compared to those in the neutral slide condition.

Data analysis

For statistical analysis, one or two-way analyses of variance (ANOVA) for repeated measures were used. Fisher's PLSD was used for post-hoc

tests. Stepwise forward multiple regression analysis was used to examine the Qc, Qt, and t/c ratio-related factors. Qc, Qt, and t/c ratio were entered into the analysis as a dependent variable. Sex, age, hemisphere, valence, and arousal during individual sessions (negative, neutral, and positive) in all subjects were entered as independent variables. In addition, the relationships between the three dependent variables and independent variables were investigated. The results are expressed as a mean \pm standard deviation.

Results

The self-ratings of experiences evoked by the positive, neutral, and negative slide sessions were, respectively, 7.3 ± 0.6 , 4.8 ± 0.5 , and 2.2 ± 0.6 for valence; and 4.4 ± 1.1 , 4.2 ± 1.0 , and 7.5 ± 0.9 for arousal. A one-way ANOVA revealed a significant main effect of slide category on valence ratings ($F[2,28] = 286.4$; $p < 0.01$). Post-hoc tests showed that all the pair-wise differences were significant ($p < 0.01$). For arousal ratings ANOVA also showed a significant main effect of slide category ($F[2,28] = 67.8$; $p < 0.01$). Post-hoc test showed that the negative slides were experienced as significantly more arousing, in comparison to neutral or positive slides ($p < 0.01$).

Fig. 2 shows the typical response waveform over the left primary auditory cortex and dipole location in one

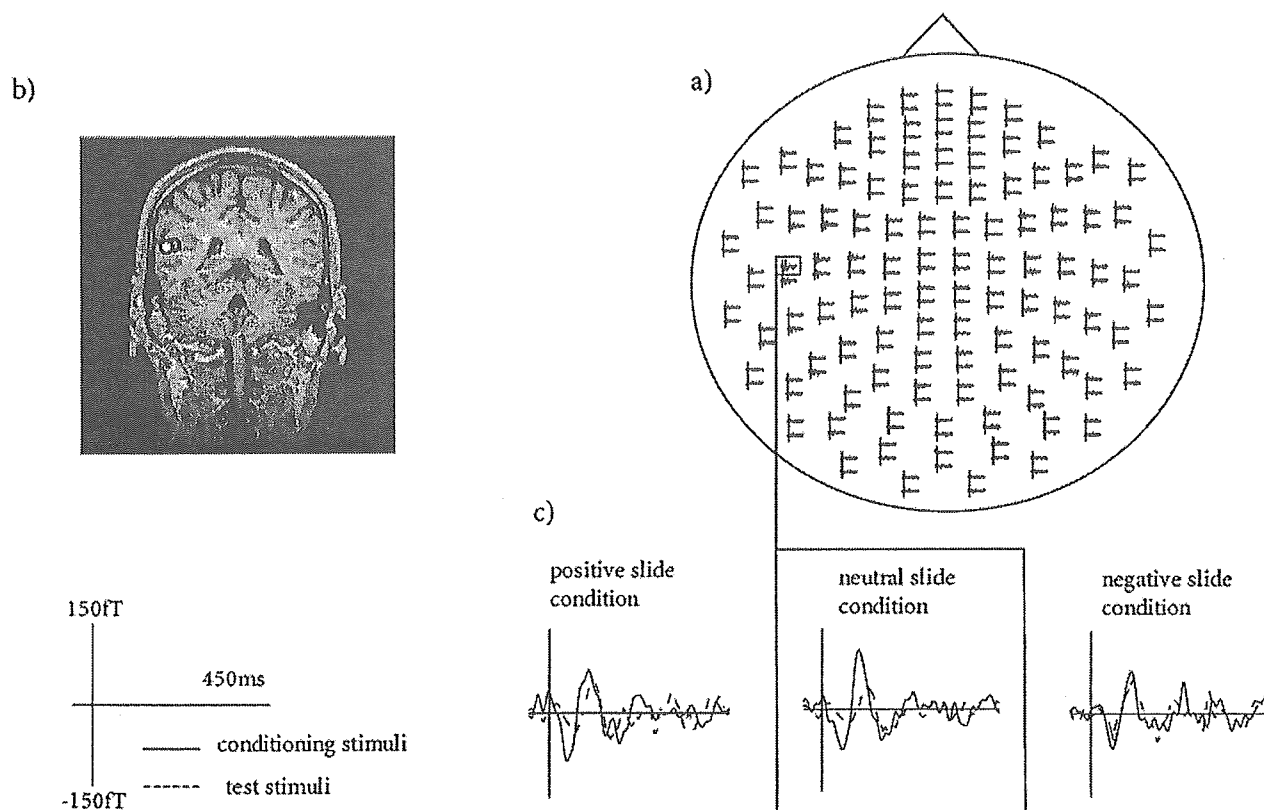


Fig. 2 a The average response waveform in the neutral slide condition of one representative subject. b Dipole locations over the left primary auditory cortex of one representative subject. The white dot represents the dipole to the conditioning click and the gray dot represents the dipole to the test click. c The response waveform at the channel showing maximal response in the left hemisphere in negative (left), neutral (middle), and positive slide conditions (right). The solid line represents the response to the conditioning click and the dashed line, the response to the test click. Note that the amplitude of response waveforms do not directly reflect the intensity of the underlying neural activation, which was estimated on the basis of magnetic field distribution at the sensors and the relative position of the head with respect to the sensors

subject. A significant difference between Qc and Qt in the neutral slide condition was observed in both hemispheres (left: $t=7.85$; $p<0.01$, right: $t=5.19$; $p<0.01$). The strength of Qc was not affected significantly by the slide category or hemisphere. A two-way ANOVA revealed a significant main effect of the slide category on Qt ($F[2,48]=4.27$; $p<0.05$), and t/c ratio ($F[2,48]=15.97$; $p<0.01$), and a significant main effect of hemisphere on t/c ratio ($F[1,24]=6.29$; $p<0.05$). Post-hoc tests showed that Qt and t/c ratio in the negative slide condition were larger than those in the positive and the neutral slide conditions, and t/c ratio in the right hemisphere was larger than that in the left hemisphere (Table 1). Slide category \times hemisphere interactions were not significant in any of the analyses. P50m latencies and the source location were not significantly influenced by emotional condition (data not shown).

Table 2 shows the predictors of Qc, Qt, and the t/c ra-

Table 1 Source activations in different emotional conditions

	Slide category		
	Negative	Neutral	Positive
Left hemisphere			
Qc (nAm)	15.0 \pm 3.7	18.3 \pm 7.4	18.4 \pm 6.7
Qt (nAm)	8.9 \pm 3.6*	6.0 \pm 3.4	6.5 \pm 4.2
t/c ratio	0.61 \pm 0.24**	0.33 \pm 0.17	0.33 \pm 0.17
Right hemisphere			
Qc (nAm)	14.0 \pm 7.3	16.7 \pm 7.6	16.7 \pm 7.0
Qt (nAm)	10.7 \pm 7.9*	8.0 \pm 3.6	7.2 \pm 3.9
t/c ratio	0.77 \pm 0.25** ^a	0.48 \pm 0.21 ^a	0.45 \pm 0.22 ^a

* $p < 0.05$; ** $p < 0.01$, compared to Neutral and Positive slide conditions

^a $p < 0.05$, compared to left hemisphere

Qc strength of the P50m response (dipole moment) to conditioning stimuli; Qt strength of the P50m response (dipole moment) to test stimuli

Table 2 Multiple regression analysis of predictors of Qc, Qt, and t/c ratio

Independent variables	Coefficient	Standardized coefficient	F	p
Qt				
Valence of stimuli	-0.508	-0.221	4.108	0.046
t/c ratio				
Valence of stimuli	-0.056	-0.454	15.338	< 0.0001
Right hemisphere	0.142	0.268		

There was no significant predictor of dipole moment of Qc response

Qt Multiple R = 0.221, Multiple R² = 0.049, Adjusted R² = 0.037

t/c ratio Multiple R = 0.529, Multiple R² = 0.280, Adjusted R² = 0.261

Table 3 Correlations between P50 responses and independent variables

	Qc		Qt		t/c ratio	
	r-value	p-value	r-value	p-value	r-value	p-value
age	-0.054	0.63	-0.063	0.57	-0.055	0.62
sex	0.084	0.45	0.028	0.81	-0.072	0.52
Arousal of stimuli	0.006	0.95	0.205	0.07	0.301	0.006
Valence of stimuli	0.189	0.09	-0.221	0.04	-0.456	< 0.0001

p-values were obtained with Pearson's correlation analysis

tio. There was no significant predictor of the dipole moment of the Qc response. A stepwise forward multiple regression analysis revealed that the self-rating valence of the emotional slides may have been predictive of dipole moment of Qt response, and t/c ratio and right hemisphere predicted a higher t/c ratio. The relationship between the separate independent variables and the dependent variables is shown in Table 3. The self-rating valence of the emotional slides was significantly correlated with the Qt response, and the t/c ratio. In addition, the t/c ratio was positively correlated with arousal by the emotional slides.

Discussion

The present study demonstrated that P50m suppression was attenuated by negative visual stimuli, and that this modulation effect predominated in the right, rather than the left hemisphere, and was related to emotional valence. Our study suggests that negative emotions could modulate sensory gating in the auditory cortex.

A deficit in sensory gating has been identified in a number of psychiatric disorders, most notably schizophrenia (Adler et al. 1982). Furthermore, drugs affecting emotional tone such as amphetamines (Light et al. 1999), marijuana (Patrick et al. 1999), and cocaine (Adler et al. 2001) have been reported to disrupt the P50 suppression. Although emotional stress contributes to the onset and exacerbation of illness in patients with psychiatric disorders, no study has been conducted which attempted to determine the effect of emotional stress on P50 suppression.

There have been a few studies which evaluate the effect of emotional stimuli on other components of event-related potentials (ERP). Schupp et al. (1997) reported that auditory P300 amplitudes were modulated by picture arousal, with smaller auditory P300 responses elicited when viewing highly arousing pictures regardless of their valence. They speculated that attentional resources are needed for the late information processing because they are very complicated and have more cognitive factors. On the other hand, Surakka et al. demonstrated that the mismatch negativity, an ERP component elicited by sound change that peaks about 150 ms after stimulus presentation, was attenuated by positively valenced and little arousing visual emotional stimuli (1998). Our results showed that emotional modulation of sensory processing could occur even at earlier (about

50 ms) cortical stages of auditory processing and that this modulation was related to emotional valence. Together, these results suggest that auditory processing can be affected by emotional stimuli, and in earlier stages, the emotional valence of the stimuli may predominate over the arousal effect. However, due to the small sample size, the results should be confirmed by further studies with a larger number of subjects.

The amygdala is thought to play an important role in the perception of emotionally meaningful information (Morris et al. 1996), and neurons in the amygdala of rats behave similarly to the human P50 in response to repeated auditory stimuli (Bordi and LeDoux 1992). In addition, it has been reported that fear conditioning enhances auditory evoked activity in the amygdala in response to repetitive auditory stimuli (Rogan et al. 1997). The amygdala is richly interconnected with the neocortex. Some amygdalofugal projections to auditory areas have been found, in addition to projections to visual regions (Amaral et al. 1992). Thus, we speculate that the effect of emotional visual stimuli on auditory P50m suppression, at least in part, might be mediated by the amygdala.

This P50m suppression study showed that auditory sensory gating at the primary auditory cortex was affected by emotional stimuli, and that this effect may be related more to the emotional valence than to the arousing effect of the stimuli. We suggest that a pre-attentive, automatic mechanism of the brain to gate out incoming irrelevant sensory input is affected by emotional valence.

■ **Acknowledgment** This study was supported in part by a Health Science Research Grant for Research on Brain Science from the Ministry of Health, Welfare, and Labor of Japan.

References

- Adler LE, Hoffer LJ, Griffith J, Waldo MC, Freedman R (1992) Normalization by nicotine of deficient auditory sensory gating in the relatives of schizophrenics. *Biol Psychiatry* 32:607–616
- Adler LE, Olincy A, Cawthra E, Hoffer M, Nagamoto HT, Amass L, Freedman R (2001) Reversal of diminished inhibitory sensory gating in cocaine addicts by a nicotinic cholinergic mechanism. *Neuropsychopharmacology* 24:671–679
- Adler LE, Pachtman E, Franks RD, Pecevich M, Waldo MC, Freedman R (1982) Neurophysiological evidence for a defect in neuronal mechanisms involved in sensory gating in schizophrenia. *Biol Psychiatry* 17:639–654
- Ahlfors S, Ilmoniemi RJ (1989) Magnetometer position indicator for multichannel MEG. In: Williamson SJ, Hoke M, Romani GL (eds) *Advance in Biomagnetism*, Plenum Press, New York, pp 673–676
- Amaral DG, Price JL, Pitkänen A, Carmichael ST (1992) Anatomical organization of the primate amygdaloid complex. In: Aggleton JP (eds) *The Amygdala: Neurological Aspects of Emotion, Memory, and Mental Dysfunction*, Wiley-Liss, New York, pp 1–66
- Bordi F, LeDoux J (1992) Sensory tuning beyond the sensory system: an initial analysis of auditory response properties of neurons in the lateral amygdaloid nucleus and overlying areas of the striatum. *J Neurosci* 12:2493–2503
- Braff DL, Geyer MA (1990) Sensorimotor gating and schizophrenia. Human and animal model studies. *Arch Gen Psychiatry* 47:181–188
- Cuthbert BN, Schupp HT, Bradley MM, Birbaumer N, Lang PJ (2000) Brain potentials in affective picture processing: covariation with autonomic arousal and affective report. *Biol Psychol* 52:95–111
- Hämäläinen M, Hari R, Ilmoniemi RJ, Knuutila J, Lounasmaa OV (1993) Magnetoencephalography – theory, instrumentation, application to noninvasive studies of working brain. *Rev Mod Phys* 65:413–498
- Hari R, Aittoniemi K, Jarvinen ML, Katila T, Varpula T (1980) Auditory evoked transient and sustained magnetic fields of the human brain. Localization of neural generators. *Exp Brain Res* 40:237–240
- Kayser J, Tenke C, Nordby H, Hammerborg D, Hugdahl K, Erdmann G (1997) Event-related potential (ERP) asymmetries to emotional stimuli in a visual half-field paradigm. *Psychophysiology* 34:414–426
- Lang PJ, Bradley MM, Cuthbert BN (1997) *International Affective Picture System (IAPS)*. NIMH Center for the Study of Emotion and Attention
- Lang SF, Nelson CA, Collins PF (1990) Event-related potentials to emotional and neutral stimuli. *J Clin Exp Neuropsychol* 12:946–958
- Laurian S, Bader M, Lanares J, Oros L (1991) Topography of event-related potentials elicited by visual emotional stimuli. *Int J Psychophysiol* 10:231–238
- LeDoux JE (1993) Emotional memory systems in the brain. *Behav Brain Res* 58:69–79
- Light GA, Malaspina D, Geyer MA, Luber BM, Coleman EA, Sackeim HA, Braff DL (1999) Amphetamine disrupts P50 suppression in normal subjects. *Biol Psychiatry* 46:990–996
- Mäkelä JP, Hämäläinen M, Hari R, McEvoy L (1994) Whole-head mapping of middle-latency auditory evoked magnetic fields. *Electroencephalogr Clin Neurophysiol* 92:414–421
- McGhie A, Chapman J (1961) Disorders of attention and perception in early schizophrenia. *Br J Med Psychol* 34:103–116
- Morris JS, Frith CD, Perrett DI, Rowland D, Young AW, Calder AJ, Dolan RJ (1996) A differential neural response in the human amygdala to fearful and happy facial expressions. *Nature* 383:812–815
- Patrick G, Straumanis JJ, Struve FA, Fitz-Gerald MJ, Leavitt J, Manno JE (1999) Reduced P50 auditory gating response in psychiatrically normal chronic marijuana users: a pilot study. *Biol Psychiatry* 45:1307–1312
- Reite M, Teale P, Zimmerman J, Davis K, Whalen J (1988) Source location of a 50 msec latency auditory evoked field component. *Electroencephalogr Clin Neurophysiol* 70:490–498
- Rogan MT, Staubli UV, LeDoux JE (1997) Fear conditioning induces associative long-term potentiation in the amygdala. *Nature* 390:604–607
- Schupp HT, Cuthbert BN, Bradley MM, Birbaumer N, Lang PJ (1997) Probe P3 and blinks: two measures of affective startle modulation. *Psychophysiology* 34:1–6
- Surakka V, Tenhunen-Eskelinen M, Hietanen JK, Sams M (1998) Modulation of human auditory information processing by emotional visual stimuli. *Brain Res Cogn Brain Res* 7:159–163

Reduced Activation of Posterior Cingulate Cortex During Imagery in Subjects with High Degrees of Alexithymia: A Functional Magnetic Resonance Imaging Study

Tomoyuki Mantani, Yasumasa Okamoto, Naoko Shirao, Go Okada, and Shigeto Yamawaki

Background: Although the brain areas involved in imagery have been reported, the neural bases of individual differences in imagery remain to be elucidated. People with high degrees of alexithymia (HDA) are known to have constricted imaginal capacities. The purpose of this study was to investigate neural correlates of imagery disturbance in subjects with HDA.

Methods: A functional magnetic resonance imaging (fMRI) study was undertaken in 10 subjects with HDA and 10 subjects with low degrees of alexithymia (LDA), who were selected according to their scores on the 20-item Toronto Alexithymia Scale (TAS-20). The two groups' regional cerebral activation was compared during various imagery conditions. In those conditions, the subjects imaged a past happy (PH) event, a past sad (PS) event, a past neutral (PN) event, a future happy (FH) event, a future sad (FS) event, and a future neutral (FN) event. The activation levels during these conditions were compared with those during a rest condition (REST).

Results: The *t* tests showed that the mean subjective ratings of both the vividness of the imagery and the intensity of emotion during the imagery were higher in the subjects with LDA than in those with HDA for the PS and FS imagery conditions. On the other hand, relative to the LDA group, the HDA group showed significantly less activation in the posterior cingulate cortex (PCC) during the PH and FH imagery conditions compared with REST and during the FH imagery condition compared with the FN imagery condition.

Conclusion: The present results suggest an association between an HDA and reduced activation of the PCC during happy imagery. Given the function of this brain region, these results might be related to a dysfunction of episodic memory retrieval during happy imagery in subjects with HDA.

Key Words: Alexithymia, imagery, posterior cingulate cortex, fMRI, future, happy

The brain areas involved in imagery have been investigated by neuroimaging studies in normal subjects. These studies revealed that, in addition to primary and secondary sensory areas (Cabeza and Nyberg 2000; Chen et al 1998; Halpern 2001; Le Bihan et al 1993; Shergill et al 2001; Yoo et al 2003), brain areas with other major cognitive functions, such as language, memory, and movement, are activated during imagery (Mellet et al 1998). The neural bases of individual differences in imagery, however, remain to be elucidated.

Alexithymia, a personality construct, was introduced by Nemiah and Sifneos in the early 1970s. The concept evolved initially from clinical observations of patients with psychosomatic disorders (Nemiah et al 1976). The salient features of this construct are as follows: 1) difficulty identifying and describing subjective feelings; 2) difficulty distinguishing between feelings and bodily sensations of emotional arousal; 3) constricted imaginal capacities; and 4) an externally oriented cognitive style (Nemiah et al 1976). Recently, high prevalences of alexithymia

have been reported in various psychiatric disorders, such as somatoform disorders (53%, Cox et al 1994), anxiety disorders (46.7%, Parker et al 1993; 41%, Shipko et al 1983), and substance dependence (51%, Taylor et al 1990). An approximately 10% prevalence has been reported even in normal populations (Honkalampi et al 2000; Salminen et al 1999). Taylor (2000) stated in his review that alexithymia is a deficit in emotional regulation that reflects three kinds of deficits: 1) deficits in the cognitive-experiential component of emotion response systems; 2) deficits at the level of interpersonal regulation of emotion; and 3) constricted imaginal capacities. Several studies have investigated imaginal capacity in people with alexithymia. Nemiah et al (1977) reported that whereas control subjects increased their oxygen consumption when instructed to think various emotional thoughts, subjects with alexithymia showed no such increase in these conditions. Whereas imaginal capacity correlates with hypnotic susceptibility (Varga 2001), Frankel et al (1977) reported a high prevalence of alexithymia in subjects with low hypnotic susceptibility. Hyer et al (1990) studied the responses of posttraumatic stress disorder patients who were listening to accounts of their own traumatic experiences and found that the more alexithymic the subject was, the less his heart rate differed between the stressor period and the baseline. Friedlander et al (1997) compared the responses of people with alexithymia with those of people without it to an autogenic relaxation exercise with guided imagery; the former reported less enjoyment and poorer (less vivid) imagery during relaxation than did the latter. These studies suggest that people with high degrees of alexithymia (HDA) might have low imaginal capacities and show less physical reactivity during imagery with emotional contents; however, these studies relied on indirect methods to characterize the brain function of people with alexithymia.

Imagery can be defined as the manipulation of sensory information that comes from memory without information from actual sensory input (Cabeza and Nyberg 2000). The memory is

From the Department of Psychiatry and Neurosciences, Division of Frontier Medical Science, Programs for Biomedical Research, Graduate School of Biomedical Sciences, Hiroshima University, Hiroshima; and the Core Research for Evolutional Science and Technology (CREST), Japan Science and Technology Corporation (JST), Seika, Japan.

Address reprint requests to Shigeto Yamawaki, M.D., Ph.D., Hiroshima University School of Medicine, Department of Psychiatry and Neurosciences, Division of Frontier Medical Science, Programs for Biomedical Research, Graduate School of Biomedical Sciences, 1-2-3 Kasumi, Minami-ku Hiroshima 734-8551, Japan; E-mail: yamawaki@hiroshima-u.ac.jp.

Received August 11, 2004; revised January 24, 2005; accepted January 28, 2005.

0006-3223/05/\$30.00
doi:10.1016/j.biopsych.2005.01.047

BIOL PSYCHIATRY 2005;57:982-990
© 2005 Society of Biological Psychiatry

subdivided into working memory, episodic memory, and semantic memory. Of these subcategories, the one most closely tied to the emotions of daily life is episodic memory, especially “autobiographical memory” (Rubin 1998). Because alexithymia is conceptualized as a multifaceted construct consisting of factors related logically to each other (Taylor et al 1997), the imagery disturbance of alexithymia is considered to be related to other factors, such as difficulty in identifying and describing feelings (i.e., in processing emotion). In fact, experimental studies suggest that imagery disturbance in subjects with alexithymia is related to emotion processing, as mentioned in the previous paragraph. Thus, we considered that subjects with HDA might have difficulty in retrieval of autobiographical memory. On the other hand, the restricted imaginal capacities of people with alexithymia have been considered to limit the extent to which individuals with HDA can modulate anxiety and other emotions by fantasy, dreams, interest, and play (Krystal 1988; Mightes and Cohen 1992). Thus, previous clinical reports have emphasized disturbances in imaging experiences that had not and were not going to occur, as well as past experiences. Furthermore, in examining the types of emotion that accompany imaging, the previous reports seem to have emphasized the difficulty that people with alexithymia have in imaging positive things to modulate negative emotions such as anxiety. So, we planned a functional magnetic resonance imaging (fMRI) study to directly investigate neural correlates of imagery disturbance in the subjects with HDA with the imagery task in which subjects recall affect-laden autobiographical memories or imagine affect-laden future episodes accompanied by positive (happy) or negative (sad) emotions.

Recent positron emission tomography (PET) and fMRI studies have suggested that episodic memory retrieval is associated with activation of the prefrontal, medial-temporal, medial parieto-occipital (posterior cingulate cortex [PCC], including the retrosplenial cortex and precuneus), lateral parietal, anterior cingulate, occipital, and cerebellar regions (Cabeza and Nyberg 2000). On the other hand, it has been reported that the retrieval of autobiographical memory, which is a kind of episodic memory, often activates the medial-frontal region and the left hippocampus and sometimes activates the medial parieto-occipital area (MPOA) (Maguire 2001). These studies suggested a relatively consistent activation of the prefrontal and medial-temporal cortices in episodic/autobiographical memory retrieval; however, there are also inconsistencies among studies of autobiographical memory retrieval, especially regarding the activation of the MPOA. Although several studies have reported the activation of this area (e.g., Maguire and Mummery 1999; Ryan et al 2001), a considerable number have reported no activation of this area (e.g., Conway et al 1999; Markowitsh et al 1997) during the retrieval of autobiographical memory. Meanwhile, emotional stimuli have been reported to consistently activate this area (Maddock 1999). Furthermore, whereas mental imagery is known to be a major component of episodic memory recall (Tulving 1983), MPOA in imagery domain studies seems to be specifically recruited whenever the generation of the mental image relies on the reactivation of a memorized percept (Ghaem et al 1997; Kosslyn et al 1993; Mellet et al 1995; Roland and Gulyas 1995). So, both episodic memory recall and imagery might share the MPOA as a common region. We considered that the inconsistencies concerning the activation of MPOA across studies might reflect not only differences among tasks but also individual differences in personality traits, in individual patterns of reactions to emotional stimuli, or in individual imaginal capacities such as alexithymia. We therefore studied brain activity during imagery in volunteers and investi-

gated the relationship between this brain activity and the level of alexithymia, while paying attention to MPOA (PCC and precuneus; Brodmann's areas 7, 23, 29, 30, 31). We hypothesized that the activation of the MPOA varies with the 20-item Toronto Alexithymia Scale (TAS-20) score during imagery, especially during emotional imagery conditions.

Meanwhile, although no study to date has used neuroimaging techniques to investigate the relationship between brain activation during imagery and alexithymic characteristics, two neuroimaging studies of alexithymia have recently been conducted (Berthoz et al 2002; Kano et al 2003). These studies suggested that there was no difference in the limbic structure between subjects with alexithymia and those without, and that the impairment of anterior cingulate cortex (ACC) functioning might be associated with alexithymia. Moreover, the medial prefrontal cortex (MPFC), adjacent to the ACC, also was activated in numerous neuroimaging studies of emotion (Phan et al 2002) and was reported to be impaired in subjects with alexithymia (Berthoz et al 2002). So, we were also interested in whether the ACC (Brodmann's areas 24, 32, 33) and MPFC (the medial regions of Brodmann's areas 8, 9, 10, 11) are associated with imagery disturbance in subjects with alexithymia.

Methods and Materials

Subjects

We recruited 14 men and six women aged 20–30 years, who were right-handed, nonanxious, and nondepressed (on the basis of Hospital Anxiety and Depression Scale scores; Zigmond and Snaith 1983; Zigmond et al 1993) from among 38 male and 22 female volunteers according to their alexithymia scores on the Japanese version of the TAS-20. The volunteers were recruited through community announcements and were paid incentives corresponding to their transportation expenses. The TAS-20 is the most psychometrically valid and commonly used measurement of alexithymia (Bagby 1994a, 1994b), and the Japanese version also has high construct validity and reliability (Fukunishi et al 1997). It is a self-report questionnaire containing 20 items rated on a 5-point scale. A high score indicates HDA. In accordance with methods used by Berthoz et al (2002), 10 subjects (7 men and 3 women) with high (≥ 56) TAS-20 scores were placed into an HDA group, and 10 subjects (7 men and 3 women) with lower (≤ 44) scores, who were age- and handedness-matched to the HDA group, were placed into an LDA group. The subjects in the HDA and LDA groups were aged 25.9 ± 3.3 years and 23.7 ± 3.0 years, respectively (mean \pm SD), and their TAS-20 scores were 61.9 ± 4.0 and 37.9 ± 3.9 , respectively.

All subjects were identified as right-handed according to the Edinburgh inventory (Oldfield 1971). According to the self-reported responses, the subjects had no history of psychiatric, neurologic, or other major medical illness and had never been treated with a psychotropic medication. After the study was described completely to the subjects, written informed consent was obtained from all of them. This study was approved by the Institutional Review Board and the Ethics Committee of Hiroshima University Hospital, Japan. The subjects received course credit for their participation.

Experimental Design

We developed our task by modifying that used in the study by George et al (1995). Before the scanning session, each subject was asked to name specific events that, when imaged, would make him or her happy (one past and one future event) or sad

(one past and one future event). Each subject was also asked to image a specific time when he or she was or would be emotionally neutral—that is, not experiencing any particular emotion (one past and one future event). As for the future events, the subjects were asked to imagine, in the greatest possible detail, specific events that they could realistically expect to occur. The researcher then reviewed each event to assess whether the emotion was appropriate (e.g., not a mixture of happiness and sadness or anger) to the type of event. Additional specific sensory stimuli were elicited that could possibly aid in imaging the event (e.g., the exact place where the subject was or would be at the peak moment of emotion or the clothing, time of year, sights, sounds, or smells associated with that moment). Finally, each subject was asked to supply key words that would simply represent each event (e.g., travel with friends, death of grandmother, brushing teeth).

We used a periodic design involving the presentation of an activation condition for 30 sec followed by a baseline condition for 18 sec. This cycle was repeated 18 times over the course of 864 sec. During the activation condition, subjects were cued by the visual presentation of Japanese words representing the six event conditions: “past happy” (PH), “past sad” (PS), “past neutral” (PN), “future happy” (FH), “future sad” (FS), and “future neutral” (FN); key words were also used to cue the subjects to generate imagery of each event. When the words were presented, the subjects were instructed to image each previously agreed-upon event to make themselves feel the emotions and senses that they would feel as if the past or future event was actually happening. Each word was presented for 30 sec. During the baseline condition (REST), the subjects were shown a cross symbol (“+”) and instructed to see only that symbol, with no imagery internally. Each presentation of the symbol lasted 18 sec. During each trial, the words were projected to the center of the subject’s field of view with a super video graphics array computer-controlled projection system. The order in which the activation conditions were presented was counterbalanced across the subjects. Each trial was started by presenting the cross symbol for 9 sec; this initial presentation was excluded from the analyses. For each event, each subject’s ratings of the vividness of his or her imagery and the intensity of the emotion were recorded immediately after the scanning session. Two nongraduated visual analogue scales were used: one assessed the vividness of the imagery (range, 0–10), from imaging nothing to imaging extremely vividly; and the other assessed the intensity of emotion during the imagery (range, 0–10), from feeling nothing to feeling extremely intensely.

Image Acquisition

Functional magnetic resonance imaging was performed with a Magnex Eclipse 1.5-T Power Drive 250 (Shimadzu Medical Systems, Kyoto, Japan). A time-course series of 291 volumes was acquired with T2-weighted, gradient echo, echo planar imaging sequences. Each volume consisted of 28 slices, with a slice thickness of 4 mm with no gap, and entirely covered the cerebral and cerebellar cortices. The interval between successive acquisitions of the same image (TR) was 3000 msec, the echo time (TE) was 55 msec, and the flip angle was 90°. The field of view (FOV) was 256 mm, and the matrix size was 64 × 64, giving voxel dimensions of 4 × 4 × 4 mm. Scan acquisition was synchronized to the onset of the trial. After functional scanning, structural scans were acquired with a T1-weighted gradient echo pulse sequence (TR = 12 msec; TE = 4.5 msec; flip angle = 20°; FOV = 256 mm; voxel dimensions of 1 × 1 × 1 mm), which facilitated localization.

Analysis

Image processing and statistical analyses were carried out with Statistical Parametric Mapping (SPM99) software (Wellcome Department of Cognitive Neurology, London, United Kingdom) implemented in Matlab (Mathworks, Natick, Massachusetts). The first three volumes of the fMRI run were discarded because the magnetic resonance signal was unsteady. The remaining 288 volumes were used for the analysis. Images were corrected for motion and realigned with the first scan of the session, which served as the reference. For each subject, the T1 anatomic images were coregistered to the first functional images and aligned to a standard stereotaxic space, with the Montreal Neurological Institute (MNI) T1 template in SPM99. The calculated nonlinear transformation was applied to all functional images for spatial normalization. Finally, the functional magnetic resonance images were smoothed with a 10-mm full-width at half-maximum Gaussian filter.

The group analysis was performed at two levels. At the first level, each subject’s signal time course was modeled with a delayed boxcar function convolved with a hemodynamic response function in the context of a general linear model. One contrast image per subject was created by contrasting each activation condition (PH, PS, PN, FH, FS, FN) with the baseline condition (REST) and by contrasting each emotional condition with the neutral condition. In the second step, with group analysis according to a random effect model that allows inference to the general source populations (Friston et al 1999), we first identified regions that showed significant responses during each activation condition (PH, PS, PN, FH, FS, FN) compared with the baseline condition (REST) and during each emotional condition compared with the neutral condition (PH > PN, PS > PN, FH > FN, FS > FN) in the HDA and LDA groups, with the one-sample *t* test. Next, the images were entered into a two-sample *t* test (to locate brain regions in which the two groups differed significantly) and into a regression analysis (to locate brain regions in which the magnitude of brain activation correlated significantly with the TAS-20 score). Although there were inequalities in the amount of data between the REST condition and the experimental conditions (6 vs. 10), SPM99 performed these analyses, correcting the inequalities of data size. The resulting set of voxel values for each contrast constituted an SPM (*t*) map. The SPM(*t*) maps were then interpreted by referring to the probabilistic behavior of Gaussian random fields. The data were thresholded at $p < .001$ uncorrected at the voxel level and at $p < .05$ corrected at the cluster level for regions about which there was no clear hypothesis. Moreover, for regions about which we had an a priori hypothesis (ACC, MPFC, and MPOA), the height and extent of thresholds were set to $p < .001$ uncorrected and $p < .05$ uncorrected, respectively (as justified by Friston 1997).

The x, y, and z coordinates provided by SPM, which were in the MNI brain space, were converted to the x, y, and z coordinates in Talairach and Tournoux’s (TT) brain space (Talairach and Tournoux 1988) with the following formula: (TT-x = MNI-x × .88 – .8; TT-y = MNI-y × .97 – 3.32; TT-z = MNI-y × .05 + MNI-z × .88 – .44). Labels for brain activation foci were obtained in Talairach coordinates with Talairach Daemon software (Research Imaging Center, University of Texas, San Antonio, Texas), the accuracy of which is similar to that of neuroanatomic experts (Lancaster et al 2000).

The accuracy of the labeling of the areas given by this software was then confirmed by comparison with activation maps overlaid on MNI-normalized structural magnetic resonance images.

Results

Subjective Ratings

The *t* tests showed that the mean subjective ratings of the vividness of the imagery were higher in the LDA group than in the HDA group for the PH, PS, and FS conditions. On the other hand, the mean subjective ratings of the intensity of emotion during the imagery were higher in the LDA group than in the HDA group for the PS and FS conditions (Table 1). The examples of the events that subjects imaged during the imagery conditions were sea bathing, party with friends, travel abroad; death of grandmother, death of dog, loss of bag; brushing teeth, cleaning, going to school; honeymoon, playing with child, entrance to ideal occupation; death of mother, loneliness at work, failing to pass on to the next grade; and washing face, changing clothes, driving for PH, PS, PN, FH, FS, FN conditions, respectively. There were no significant differences between the groups in the mean subjective remoteness of the time when the imaged event occurred or would occur in any of the imagery conditions.

fMRI Results (Brain Activation During Imagery Conditions Within Groups)

In the LDA group, at the higher level of significance (height and extent thresholds, respectively, set to $p < .001$ and $p < .05$ corrected), there was significantly greater activation of the PCC in PH and FH than in REST and of the left superior frontal gyrus in PS compared with REST. Moreover, at the lower level of significance (height and extent thresholds, respectively, set to $p < .001$ and $p < .05$ uncorrected), the ACC and right precuneus were significantly more active in PN than in REST and in FH than in FN, respectively. In the HDA group, at the higher level of significance (height and extent thresholds, respectively, set to $p < .001$ and $p < .05$ corrected), there was significantly greater activation of the fusiform gyrus and cerebellum in PH, PN, and FH than in REST (Table 2).

fMRI Results (Comparison of Brain Activation Between Groups)

With the threshold of significance at $p < .001$ uncorrected at the voxel level and at $p < .05$ corrected at the cluster level, FH (FH compared with REST) induced less activation in the HDA

subjects than in the LDA subjects in the bilateral PCC (Table 3, Figure 1). These were the only regions showing between-group differences at this level of significance.

Moreover, at the lower level of significance (height and extent thresholds, respectively, set to $p < .001$ and $p < .05$ uncorrected), PH compared with REST and FH compared with FN induced less PCC activation in the HDA group than in the LDA group (Table 3, Figures 2 and 3).

fMRI Results (Regression Analysis)

Regression analysis revealed a significant inverse correlation between the TAS-20 score and the magnitude of brain activation of the bilateral PCC in PH compared with REST ($x, y, z = -6, -54, 9$; area 30; *t* value 5.26; 1595 voxels; $r = -.846$; $p < .001$), in FH compared with REST ($x, y, z = -8, -50, 9$; area 30; *t* value 7.83; 1363 voxels; $r = -.810$; $p < .001$), and in FH compared with FN ($x, y, z = 6, -62, 21$; area 31; *t* value 6.52; 669 voxels; $r = -.812$; $p < .001$) at the higher level of significance (height and extent thresholds, respectively, set to $p < .001$ and $p < .05$ corrected). The Talairach coordinates presented in each parenthesis represents the coordinates in which the local maximum within the cluster was observed. No significant correlation was observed in any other investigated contrasts.

Discussion

As expected, the activation of the MPOA varied with the TAS-20 score (the degrees of alexithymia) during imagery. During FH imagery, the activation in the PCC was significantly lower in the HDA subjects than in the LDA subjects. Also in FH imagery, a significant inverse correlation was found between the TAS-20 score and the magnitude of PCC activation. These results support our hypothesis that the inconsistencies about the activation of MPOA across studies reflect individual differences in personality traits such as alexithymia. The variability of activation in this area according to alexithymic characteristics might explain why prior studies, which did not control for personality, reported various outcomes.

A qualitative comparison of the brain activation detected by one-sample *t* test in the two groups suggests other findings in addition to the intergroup difference in PCC activation. The LDA

Table 1. Subjective Ratings of Vividness of Imagery and Intensity of Emotion During Imagery

Imagery Condition	TAS-20 Group	Vividness of Imagery		<i>p</i>	Intensity of Emotion During Imagery		<i>p</i>
		Mean	SD		Mean	SD	
Past Happy	HDA	6.1	2.2	.007	6.0	1.8	.068
	LDA	8.3	.75		7.5	1.7	
Past Sad	HDA	4.6	2.7	.031	4.1	2.7	.009
	LDA	7.4	2.6		7.6	2.6	
Past Neutral	HDA	4.2	2.2	.052	2.3	1.9	.188
	LDA	6.3	2.3		3.8	2.8	
Future Happy	HDA	6.5	2.7	.133	5.9	2.7	.086
	LDA	8.1	1.9		7.8	1.8	
Future Sad	HDA	3.9	3.1	.003	3.7	2.8	.010
	LDA	7.7	1.8		7.0	2.4	
Future Neutral	HDA	4.5	2.2	.325	2.3	1.6	.321
	LDA	5.7	2.8		3.5	3.1	

Two nongraduated visual analogue scales were used (the vividness of imagery [imaging nothing to imaging extremely vividly] and the intensity of emotion during imagery [from feeling nothing to feeling extremely intense]). HDA, high degrees of alexithymia group: the subjects with a total 20-item Toronto Alexithymia Scale (TAS-20) score of ≥ 56 ; LDA, low degrees of alexithymia group: the subjects with a total TAS-20 score of ≤ 44 ; *p* = value of *t* test.

Table 2. Brain Regions Showing Significant Activation During Imagery Task

	k ^a	BA	t score	Talairach Coordinates ^b		
				x	y	z
Low Degrees of Alexithymia Group						
Past happy > REST						
Left posterior cingulate gyrus	325 ^c	30	6.93	-4	-54	8
Past sad > REST						
Left superior frontal gyrus	229 ^c	8	9.22	-4	22	48
Past neutral > REST						
Left anterior cingulate gyrus	128 ^d	24	6.85	-3	4	47
Future happy > REST						
Left posterior cingulate gyrus	528 ^c	30	9.85	-6	-54	9
Right posterior cingulate gyrus		30	6.25	6	-50	17
Right posterior cingulate gyrus		23	5.56	4	-58	14
Future happy > future neutral						
Right precuneus	97 ^d	31	8.35	6	-62	25
High Degrees of Alexithymia Group						
Past happy > REST						
Left cerebellum anterior lobe	369 ^c		10.54	-32	-48	-22
Left fusiform gyrus		37	7.81	-36	-60	-19
Left cerebellum posterior lobe			6.63	-31	-63	-25
Past neutral > REST						
Right cerebellum posterior lobe	398 ^c		10.48	38	-73	-20
Right fusiform gyrus		18	6.40	24	-85	-19
Right cerebellum posterior lobe			5.75	19	-75	-22
Left fusiform gyrus ^c	179 ^c	18	5.75	-20	-85	-20
Left cerebellum posterior lobe			4.76	-36	-73	-23
Future Happy > REST						
Left cerebellum anterior lobe ^c	566 ^c		9.25	-36	-40	-30
Left cerebellum posterior lobe			6.78	-29	-75	-24
Left fusiform gyrus		19	5.81	-41	-75	-11

BA, Brodmann's Area; REST, during a rest condition.

^aNumber of voxels in cluster.

^bCoordinates of the local points of maximal activation included in the cluster.

^cDifferences were significant at $p < .001$ (uncorrected) for voxel level and $p < .05$ (corrected) for cluster extent.

^dDifferences were significant at $p < .001$ (uncorrected) for voxel level and $p < .05$ (uncorrected) for cluster extent.

group showed significantly greater activity of the MPFC, which also extends to the ACC, during PS imagery, but the HDA group did not. This is partially in line with the ACC deficit model of alexithymia (Berthoz et al 2002; Lane et al 1997). There seems to be greater activation in the fusiform gyrus and cerebellum in the HDA group than in the LDA group, which might reflect visual attention (Allen et al 1997; Mangun et al 1998); however, such a

comparison does not allow us to measure voxel-by-voxel differences in the magnitude of activation between the groups. A more formal test of the null hypothesis of no between-group difference in activation was provided by a two-sample *t* test at each voxel. A direct comparison between the groups showed significantly lower brain activation in the PCC of the HDA group than in that of the LDA group during FH imagery.

Table 3. Brain Regions Showing Significant Activation in Nonalexithymic Group Compared with Alexithymic Group During Imagery Task

Area	k ^a	BA	t score	Talairach Coordinates ^b		
				x	y	z
Past Happy > REST						
Left posterior cingulate gyrus	220 ^c	30	4.53	-4	-56	7
Left posterior cingulate gyrus		31	4.23	-1	-60	25
Left posterior cingulate gyrus		23	3.93	-8	-58	18
Future Happy > REST						
Left posterior cingulate gyrus	1150 ^d	30	6.97	-4	22	48
Right posterior cingulate gyrus		31	5.61	8	-54	25
Future Happy > Future Neutral						
Right posterior cingulate gyrus	277 ^c	31	5.58	6	-59	23

BA, Brodmann's Area; REST, during a rest condition.

^aNumber of voxels in cluster.

^bCoordinates of the local points of maximal activation included in the cluster.

^cDifferences were significant at $p < .001$ (uncorrected) for voxel level and $p < .05$ (uncorrected) for cluster extent.

^dDifferences were significant at $p < .001$ (uncorrected) for voxel level and $p < .05$ (corrected) for cluster extent.