Human Amygdala Responses During Presentation of Happy and Neutral Faces: Correlations with State Anxiety

Leah H. Somerville, Hackjin Kim, Tom Johnstone, Andrew L. Alexander, and Paul J. Whalen

**Background:** Previous functional imaging studies demonstrating amygdala response to happy facial expressions have all included the presentation of negatively valenced primary comparison expressions within the experimental context. This study assessed amygdala response to happy and neutral facial expressions in an experimental paradigm devoid of primary negatively valenced comparison expressions.

**Methods:** Sixteen human subjects (eight female) viewed 16-sec blocks of alternating happy and neutral faces interleaved with a baseline fixation condition during two functional magnetic resonance imaging scans.

**Results:** Within the ventral amygdala, a negative correlation between happy versus neutral signal changes and state anxiety was observed. The majority of the variability associated with this effect was explained by a positive relationship between state anxiety and signal change to neutral faces.

**Conclusions:** Interpretation of amygdala responses to facial expressions of emotion will be influenced by considering the contribution of each constituent condition within a greater subtractive finding, as well as 1) their spatial location within the amygdaloid complex; and 2) the experimental context in which they were observed. Here, an observed relationship between state anxiety and ventral amygdala response to happy versus neutral faces was explained by response to neutral faces.

**Key Words:** Amygdala, functional magnetic resonance imaging, human, state anxiety, happy faces, neutral faces

Numerous studies have demonstrated that the amygdala responds to facial expressions of emotion, with a particular emphasis on the study of fearful facial expressions (Breiter et al 1996; Morris et al 1996; Phillips et al 1997, 1998; Whalen et al 1998, 2001). The amygdala has also been shown to be responsive to other types of facial expressions (Anderson et al 2003; Yang et al 2002), including positive expressions (i.e., happy), when compared with either a neutral-face baseline (Breiter et al 1996; Canli et al 2002; Killgore and Yurgelun-Todd 2001; Yang et al 2002) or a low-level fixation baseline (Pessoa et al 2002; Whalen et al 1998).

To date, the experimental designs of studies reporting amygdala signal changes to happy facial expressions have included at least one negatively valenced facial expression. To determine whether the human amygdala is responsive to happy facial expressions in their own right (and not because of the presence of primary negative expressions in the experimental context), the present experimental design included only happy and neutral faces, as well as an interleaved fixation baseline condition. Previous behavioral research has demonstrated that the interpretation of neutral faces can vary depending on the presence of other primary expressions in the experimental context (e.g., Russell and Fehr 1987), so we were particularly interested in assessing amygdala response to neutral faces, given that they represent the most “potentially” negative stimulus in the current paradigm.

Previous neuroimaging research has demonstrated a relationship between amygdala activation and individual personality characteristics (Abercrombie et al 1998; Canli et al 2002; Fischer et al 2001; Johnson et al 1999; Schaefer et al 2002). Indeed, the results of one study (Canli et al 2002) suggest that amygdala responses are more variable to happy expressions (compared with fear) and that personality measures are particularly useful in explaining this variability. Therefore, we used such personality measures to assess their relationship with amygdala responses to happy versus neutral faces, as well as the unique contribution of each condition considered as a change from the fixation baseline. Consistent with our previous reports (Kim et al 2003; Whalen et al 1998, 2001), we assessed the current functional magnetic resonance imaging (fMRI) responses with a particular emphasis on spatial location within the amygdaloid complex.

**Methods and Materials**

**Subjects**

Subjects were 16 right-handed (Oldfield 1971) adults (eight female; aged 24.4 ± 3.67 years [mean ± SD]). All subjects underwent a brief clinical interview to ensure an absence of current or past psychiatric, neurologic, or medical illness in themselves and an absence of psychiatric illness in first-degree relatives. This investigation was conducted in accordance with the guidelines of the Human Subjects Committee of the University of Wisconsin-Madison; all subjects provided written informed consent for participation.

**Stimuli and Apparatus**

Face stimuli consisted of happy and neutral expressions of six individuals, three female (Ekman and Friesen 1976; identities used were C, EM, GS, MF, SW, and WF) normalized for size and luminance. For four subjects, stimuli were back-projected onto a screen within the imaging chamber and were viewable by a mirror (1.5 in × 3.5 in [3.8 cm × 8.9 cm]) approximately 6.5 in (16.5 cm) from the subject’s face. The remaining 12 subjects viewed stimuli through Avotec Silent Vision 4000 Enhanced, High Resolution Fiber Optic Video System (Avotec, Inc., Stuart, Florida) goggles centered and focused above the subjects’ eyes.

From the W.M. Keck Laboratory for Functional Brain Imaging and Behavior (LHS, HK, TJ, ALA, PJW) and the Departments of Psychology (HK, PJW) and Psychiatry (LHS, ALA, PJW), University of Wisconsin, Madison, Wisconsin.

Address reprint requests to Paul J. Whalen, Ph.D., University of Wisconsin, W.M. Keck Laboratory for Functional Brain Imaging and Behavior, 1500 Highland Avenue, Waisman Center, Room T227, Madison, WI 53705.

Received October 28, 2003; revised January 8, 2004; accepted January 9, 2004.

0006-3223/04/$30.00 doi:10.1016/j.biopsych.2004.01.007

Within each of the current effects, examination of individual subject data verified that subjects who viewed stimuli via back-projection did not represent extreme scores but rather were normally distributed among the responses for the group. Head stabilization was achieved by supporting the subject’s head with pillows.

Procedure

Subjects passively viewed blocked presentations of faces during two functional scans (see Figure 1). During each scan, 18-sec blocks alternated between presentations of happy (H) and neutral (N) faces. Within a scan, face presentation blocks were interleaved with 18-sec blocks during which a fixation point (+) was presented on an otherwise blank screen. An example of a typical scan consisted of the following: + N + H + N + H + N + H + + . The order of happy and neutral face blocks was counterbalanced within and across subjects. Each scan lasted 4 min 14 sec.

During each 18-sec block, subjects viewed 36 happy or neutral stimuli (six presentations of six individual faces, matched for presentation number and identity). Each face stimulus was presented for 200 msec at an interstimulus interval of 300 msec (i.e., two per second), for consistency with previous studies (Breiter et al 1996; Kim et al 2003; Whalen et al 2001). The background color of the screen was black across all blocks.

Upon exiting the scanner, subjects were again presented with the face stimuli in an 18-sec blocked format and were asked to label the expression and then provide a valence rating for the block (scale 1–9: 1 = very positive, 3 = positive, 5 = neither negative nor positive, 7 = negative, 9 = very negative). Subjects provided ratings of blocks of faces, rather than individual faces, to facilitate comparison of these data with their blocked fMRI data (see also Kim et al 2003). One block of each expression was presented (in the same order as their first functional scan), during which subjects provided one label and valence rating for each block of faces at the midpoint of the block.

Subjects also completed the following behavioral scales: Edinburgh Handedness Inventory (Oldfield 1971), Beck Depres- sion Inventory (BDI; Beck et al 1961), State- Trait Anxiety Inven- tory (STAI; Spielberger et al 1988), the NEO Five-Factor Inventory (NEO-FFI [Neuroticism and Extraitversion subscales only]; Costa and McCrae 1991), and the Positive and Negative Affect Scale (PANAS; Watson et al 1988).

Image Acquisition and Analysis

Subjects were scanned with a General Electric Signa 3.0 Tesla fMRI scanner (General Electric Medical Systems, Milwaukee, Wis- consin) with a quadrature head coil. For each subject, a whole- brain, high-resolution, T1-weighted anatomic scan (three-dimen- sional spoiled gradient; 250 × 256 in-plane resolution, 240-mm field of view [FOV]; 124 × 1.1-mm axial slices) was acquired for transformation and localization of functional data to Talairach space (Talairach and Tournoux 1988). An echo planar imaging se- quence (repetition time = 2000 msec, echo time = 33 msec, 60° flip angle) was used to collect functional data, with 18 contiguous 3-mm-thick coronal oblique slices (5-mm interslice gap; 64 × 64 in-plane resolution, 180-mm FOV). This functional acquis- ition scheme provided for slices with roughly isotropic voxels (2.812 × 2.812 × 3.0 mm t 5-mm skip), or 27 mm3) that were centered over the amygdala and tilted approximately 30° in an anterior direction. These slices covered the medial prefrontal cortex (mPFC) in an anterior direction, extending posteriorly to the posterior extent of the splenium of the corpus callosum. Thus, we did not image the frontal or occipital poles, posterior cingulate, or much of the parietal lobe. Standard software (AFNI; Cox 1996) was used to perform a random-effects analysis of functional data. Raw functional blood oxygen level–dependent (BOLD) images were motion-cor- rected, yielding estimated rotations and translations of less than 1° and 1.5 mm for every subject. Next, functional data were spatially smoothed with a Gaussian kernel with 6 mm full-width half maximum. A linear model with boxcar regressors, lagged by 2 repetition time (TR) intervals, for happy and neutral conditions was used to generate linear contrast maps of happy versus neutral, happy versus fixation, and neutral versus fixation. Each contrast map was then converted to units of percent signal change and spatially normalized into Talairach space. Percent signal change contrast maps were also averaged across runs to create one averaged percent signal change map per subject. Contrast maps were then analyzed across subjects with voxel-wise one-sample t tests. Voxel-wise bivariate correlations across subjects were calculated between happy versus neutral percent signal change (or the change of either from the fixation baseline) and raw scores of the behavioral measures. Data from one female subject was excluded from the analysis of state anxiety (Figure 3) because she was found to have signal changes that were more than 2 SDs above the group mean and twice that of the next highest subject’s signal change at this locus.

Given this study’s focus on responses within the amygdala, we first defined the anatomic boundaries of this region. The amygdala boundaries constituted a search volume of approximately 3500 mm3 bilaterally (Mai et al 1997). Because we discuss response distinctions made between dorsal versus ventral amygdalal, we note that the dorsal/ventral cut-off within Talairach space was z = −10 mm (see Kim et al 2003). The maximally activated voxels of all reported results were statistically significant at a threshold of p < .05, corrected for multiple compari- sions, as stipulated by Monte Carlo simulations (AlphaSim within AFNI), based on the search volume stated above. Susceptibility-related signal “dropout” attributable to B0 inho- mogeneity is of particular concern within the amygdala (Ojemann et al 1997). We set acquisition parameters and took precautions to avoid susceptibility-related signal dropout in the amygdaloid region as follows: 1) relatively small and roughly isotropic voxels were acquired to reduce intravoxel signal dephasing; 2) data were acquired in coronal slices to minimize through-plane signal dephasing; and 3) a relatively short echo time (TE) of 33 msec was used to minimize phase dispersion at the time of echo. Parameters similar to these, used in the current and in our previous studies (Kim et al 2003; Whalen et al 2001), have recently been reported to provide optimal coverage of the amygdaloid region (Chen et al 2003). In addition, we verified that
all subjects moved less than 1.5 mm (i.e., half a voxel) in all directions (anterior–posterior, right–left, inferior–superior), thereby ruling out movement-related susceptibility “edge” effects. Finally, because differences in signal quality across subjects might mimic a correlational finding, we verified that there was no significant relationship between state anxiety scores and baseline signal levels across subjects (i.e., fixation condition) at the reported left and right amygdala correlational loci (both p > .20).

Results

Behavioral Measures

Mean valence ratings of happy faces (1.63 ± .719; range = 1–3; mode = 1) were significantly more positive than those of neutral faces (5.75 ± .856; range = 4–7; mode = 6; t(15) = −11.716, p < .0001). The behavioral measures administered yielded the following mean scores: Edinburgh Handedness Inventory: 94.44% Right ± .05; Beck Depression Inventory: 2.26 ± 2.29; STAI’s: 29.5 ± 4.7; STAI-t: 29.38 ± 3.69; NEO-FFI Neuroticism subscale: 12.31 ± 4.3; NEO-FFI Extraversion subscale: 31.31 ± 5.13; PANAS Positive Affect subscale: 28.56 ± 6.84; PANAS Negative Affect subscale: 11.78 ± 2.33. There were no gender differences within any of these measures (all p > .10). These behavioral measures indicate that all subjects were right-handed, all subjects perceived happy faces as positive in nature, and all scores for depression and anxiety were within normal limits.

fMRI Data: Convergence with Previous Studies

When considered as a change from the low-level fixation condition, both happy and neutral faces produced significant signal increases across the dorsal amygdala/substantia innominata (SI) region (see Figure 2). The localization of these effects within the dorsal amygdala/SI is consistent with the dorsal nature of previously reported loci (Breiter et al 1996; Pessoa et al 2002; Whalen et al 1998). In addition, consistent with previous reports (Breiter et al 1996), these signal increases habituated significantly to repeated presentations within scans [i.e., block 1 > block 2 > block 3; run 1: F(2,14) = 15.50, p = .000012; run 2: F(2,14) = 7.28, p = .0027] and showed a trend toward decreases across scans [i.e., scan 1 > scan 2; F(1,15) = 5.63, p = .016; not significant after cluster threshold correction]. Given the similar responsivity seen to both conditions, we did not observe a significant signal difference to happy versus neutral faces within the dorsal amygdala/SI region.

There was a signal increase to happy versus neutral faces in the anterior and ventral amygdala/l/t(15) = 3.516, p = .0031, max voxel: x = 25, y = 0, z = −14, in a location consistent with a previous report (Yang et al 2002). This activation did not exceed corrected statistical thresholding, because it was rather spatially circumscribed (e.g., this locus comprised only 17 mm³ that exceeded p < .01, uncorrected). Furthermore, at this locus, a significant difference between happy versus neutral faces was observed only during the second scan [happy versus neutral, scan 1: t(15) = 1.094, p = .29; happy versus neutral, scan 2: t(15) = 3.574, p = .0028].

At the posterior border of the right ventral/amygdala hippocampus, there was a main effect of gender: female subjects were more responsive to all face stimuli compared with male subjects [F(1,14) = 2.34, p = .0032, x = 21, y = −8, z = −17]. This effect differs from that observed in the dorsal amygdala/SI, where magnitude of responsivity to these stimuli was similar for male and female subjects. This effect is more strongly supported by 1) a lack of a significant gender × expression interaction; and 2) its posterior location and contiguous throughout the hippocampus (reflecting perhaps a gender difference in greater medial temporal lobe responsivity rather than specifically in the amygdala).

fMRI Data: Correlations with State Anxiety

Figure 3A presents loci within the right (r = −.646, p = .009; x = 19, y = −5, z = −21) and left (r = −.685, p = .005; x = −22, y = −5, z = −18) ventral amygdala showing a significant negative correlation between STAI’s scores and response to happy versus neutral faces. Within Talairach space (Talairach and Tournoux 1988; Mai et al 1997), these loci were within the basolateral complex (BLC), which in the human is located within the ventral amygdala (see Methods). Figure 3B and C present scatterplots depicting the negative relationship between state anxiety and response to happy versus neutral faces for these left and right ventral amygdala loci. Greater levels of state anxiety were associated with smaller signal changes to happy versus neutral faces, indeed even reversing (i.e., neutral > happy) in the most highly state-anxious subjects.

To illustrate the relative contributions of response to happy and/or neutral faces to this subtractive finding, Figure 3D–G present separate state anxiety–amygdala response scatterplots depicting responses to happy faces (vs. fixation) and neutral faces (vs. fixation) for the left and right loci depicted in Figure 3A. Figure 3D and E suggest that the relationship between state anxiety and amygdala response to happy faces is not compelling within either the right (r = −.060, p = .83; Figure 3D) or left (r = −.174; p = .555; Figure 3E) amygdala. Figure 3F and G show a significant positive relationship between state anxiety and amygdala response to neutral faces within the right (r = .480, p = .035; Figure 3F) and left (r = .475, p = .038; Figure 3G) amygdala. Thus, the observation that signal “change” between happy and neutral faces is related to levels of state anxiety (Figure 3B and C) can be explained as a greater level of amygdala reactivity to neutral faces with greater levels of state anxiety (Figure 3F and G).

No other behavioral measures showed a significant relationship with amygdala responsivity to happy versus neutral faces (or either condition compared with fixation).
Discussion

This study assessed amygdala response to happy and neutral faces in an experimental context devoid of primary negatively valenced expressions. The fully interleaved fixation baseline condition allowed for the assessment of amygdala responsivity to happy and neutral faces separately. We observed a negative correlation between state anxiety scores and percent signal change to happy versus neutral faces. (B, C) Scatterplots depicting these negative correlations for the right and left amygdala, respectively. (D–G) Scatterplots D and E present the respective contribution of right and left amygdala response to happy faces (vs. fixation) at the locus depicted in A; scatterplots F and G present the respective contribution of right and left amygdala response to neutral faces (vs. fixation) at the locus depicted in A. These scatterplots suggest that the observed negative relationship between state anxiety and happy versus neutral faces is actually due to a positive relationship between amygdala response to neutral faces and levels of state anxiety. The x-axis of each scatterplot represents raw State-Trait Anxiety Inventory (STAI) scores, the y-axis represents percent signal change values. Image parameters as in Figure 2.
neural faces. Assessment of responsivity to each face type as a change from the fixation baseline showed that a positive relationship between state anxiety and amygdala response to neutral faces was the basis of this effect in the present experimental context.

Considering Human Amygdala Response to Happy Facial Expressions

First, fMRI responses to happy and neutral faces were not uniform across the human amygdaloid complex. Consistent with previous studies, the dorsal amygdala/SI demonstrated signal increases to both happy faces (Breiter et al 1996; Pessoa et al 2002; Whalen et al 1998) and neutral faces (Breiter et al 1996; Pessoa et al 2002). The human dorsal amygdala/SI region comprises the central nucleus of the amygdala, as well as basal forebrain cell groups, such as nucleus basalis of Meynert and sublenticular extended amygdala neurons. These neuronal groups might constitute a functional unit whose activity is related to the arousal properties of presented stimuli (Detari and Vanderwolf 1987; Whalen et al 1994; Kapp et al 1992). Thus, signal increases to faces in this region can be observed to a host of expressions (Breiter et al 1996; Pessoa et al 2002; Whalen et al 1998, 2001) and in the present experimental context did not discriminate between happy and neutral faces.

The human ventral amygdala comprises the BLC. The BLC has been shown to be critical to the detection and convergent processing of stimuli that predict biologically relevant outcomes (LeDoux et al 1998, 2001) and in the present experimental context did not discriminate between happy and neutral faces.

Human amygdala lesion patients are not impaired in their processing of happy faces (Adolphs et al 1994, 1995), which suggests that amygdala response to these stimuli might play more of a monitoring role rather than being causally related to behavioral outcomes. Of course, this point can only speak to the behavioral outcomes that have been measured to date. Like negative expressions, happy expressions offer predictive information concerning the ensuing probability of threat (or a positive outcome). Thus, human ventral amygdala response to happy faces converges with the demonstrated role of the BLC in appetitive conditioning in animal subjects (e.g., Schoenbaum et al 1999).

Given the human neuroimaging data garnered to date, amygdala response to fearful expressions seems to be more fixed, automatic, and perhaps even preattentive (Anderson et al 2003; Canli et al 2002; Whalen et al 1998), whereas response of this system to happy expressions is perhaps more variable and elaborative (Canli et al 2002). In the present study, amygdala response to happy faces was significantly different than that observed to neutral faces only for later stimulus presentations (i.e., scan 2 only). Therefore, discrimination of happy versus neutral faces within the ventral amygdala was qualitatively different than that observed to fearful expressions in previous studies, in which discrimination was observed within initial stimulus presentations (Breiter et al 1996; Whalen et al 2001). This distinction in temporal profile might offer a hint at potential behavioral outcomes (in response to happy faces) that might be attenuated by amygdala lesions.

Interpreting Amygdala Response as a Function of State Anxiety

A face constitutes a biologically relevant environmental “canvas” from which predictive information is routinely gleaned. Here, state anxiety levels were related to the level of ventral amygdala reactivity to this source of potential predictive information. Indeed, this effect was located within the BLC, the portion of the amygdaloid complex most notably involved in detection and convergent processing of stimuli that predict biologically relevant and often aversive outcomes (Davis and Whalen 2001; LeDoux et al 1990). This result might be consistent with the notion that elevated state anxiety affects the ability to disengage visual attention from threat-related expressions (e.g., see Fox et al 2001). Though neutral faces are not necessarily threat-related, it is possible that directly contrasting neutral faces with only happy faces in the present study design (an alternating-block design intended to “polarize” categorical discrimination between expression conditions) created an experimental context in which state anxiety correlated with response to the most “potentially” negative predictive stimulus in the environment. Such a hypothesis would be consistent with data showing that primary facial expressions can serve as “anchors” for neutral faces, the latter being rated more negatively when presented with positively valenced expressions than when presented with negatively valenced expressions (Russell and Fehr 1987).

An alternative hypothesis would contend that the observed relationship between amygdala response to neutral faces and state anxiety was observed, not because of the status of happy faces as positively valenced per se, but because of their clarity of valence or category. To elaborate, happy expressions are discriminated more quickly and more accurately than any other expression (e.g., Ekman et al 1987; Pollak et al 2000, 2001). In addition, subjects report having more prior experience with these expressions compared with other primary expression categories (Bond and Siddle 1996; Whalen 1998). Thus, within the present experimental context, amygdala responsivity might begin to track the “least clear” expression category, varying as a function of state anxiety. This hypothesis would predict that amygdala response to neutral faces should show a similar relationship with state anxiety, even when these faces alternate with a clearly negative expression (e.g., anger).

The two hypotheses considered thus far treat response to neutral faces as a contrast effect, contextually dependent on the comparison expression within a given experimental paradigm. Alternatively, amygdala response that varies as a function of state anxiety might be related to the inherent uncertainty of neutral faces. That is, this effect might be more absolute than relative. Such a hypothesis would predict that this relationship between state anxiety and amygdala response to neutral faces would be observed regardless of the experimental context.

Addressing Variability in a “Baseline” Condition

Identifying a source of variability influencing amygdala response to a “neutral” face baseline is both scientifically exciting and experimentally complicating. Documenting that such variability exists is just the first step. Upon knowing of its existence, one strategy is to try to counteract some of this variability. For example, some studies use 25% happy faces as a baseline condition when assessing responsivity to primary negative expression categories (e.g., Phillips et al 1998). Presumably, the use of such stimuli decreases variability in amygdala responsivity within the baseline, allowing for a clearer comparison of primary expression stimuli. To the extent that state anxiety contributes to this variability, one could test such an assumption within the present experimental design, hypothesizing that substitution of 25% happy faces as the “neutral” face baseline would attenuate (at least to some degree) the relationship with state anxiety observed here.
Alternatively, variability associated with presentation of a given neutral face condition can be measured and shown to be systematically related to variability in an additional dependent measure, here shown to vary as a function of state anxiety. Such a strategy has two advantages. First, upon knowing the contributions of state anxiety to variability within the neutral face baseline, one could statistically control for this variability and observe the residual effects related to other variables. Second, an investigator might seek to study the present effect in its own right. For example, ventral amygdala signal increases to neutral faces observed as a function of state anxiety could be theoretically consistent with the purported role of the BLC in potential threat detection (Davis and Whalen 2001; LeDoux 1996). To elaborate, greater levels of state anxiety have been hypothesized to lead to greater levels of vigilance in response to potential threat (see Bower 1981; Eysenck 1992). To the extent that neutral faces are usefully considered in terms of their ambiguous predictive value concerning potential threat, the observed relationship between state anxiety and amygdala response to neutral faces converges with data from studies utilizing ambiguous verbal information where greater anxiety levels have been shown to bias interpretation of these stimuli in a negative direction (Blanchette and Richards 2003; Mathews et al 1989; Eysenck et al 1991). Future studies could use the present experimental paradigm to assess the response of individuals with psychopathology related to exaggerated threat-assessment (e.g., anxiety disorders), hypothesizing that greater response to neutral faces might underlie between-group differences (see Birbaumer et al 1998; Thomas et al 2001).

Baseline issues in fmri are difficult to address, given the lack of an absolute baseline that is comparable across subjects (see Gusnard and Raichle 2001). In one sense, it could be argued that subtractions from the fixation condition (as opposed to the neutral face condition) merely substitute one arbitrary baseline for another. The current demonstration that the “neutral” baseline condition explained the majority of variability within a larger subtractive effect becomes more compelling if it can be shown that this will not always be the case. To elaborate, we recently observed a relationship between subjects’ valence ratings and ventral amygdala response to surprised versus neutral faces (Kim et al 2003). In that study, amygdala responses to surprised faces (vs. fixation) showed a greater degree of relationship to valence ratings ($r = .56$, $p = .013$) compared with neutral faces (vs. fixation; $r = -.06$, $p = .42$). Thus, in the present study, the demonstrated contributions of the neutral face condition to the greater subtractive finding are not an artifact inherent to the processing of BOLD signal changes within the amygdala, but rather a real phenomenon that perhaps provides a clue concerning the relationship between anxiety and amygdala function.

Methodologies such as positron emission tomography or perfusion fmri, which enable the measurement of an absolute baseline, could offer valuable additional information concerning the influence of resting (e.g., see Zald et al 2002) and dynamic (e.g., Gusnard and Raichle 2001) baselines concerning subtractive fmri effects. In the interim, given the infancy of fmri as a field, consideration of additional baseline conditions will lead to greater flexibility in the interpretation of a given finding, the stimulation of future research, and possibly a greater understanding of the nature of BOLD fmri signal changes. Indeed, the additional data considering responses to faces as a change from the fixation baseline (Figure 2D–G) prevent the possibility that the current correlational finding observed to happy versus neutral faces will be later “shorthanded” and oversimplified to suggest that more state anxious subjects are less sensitive to happy facial expressions. Figure 2D and E demonstrate that this is clearly not the case.

**Caveats and Limitations**

It is important to remember that anxiety scores measured in the current study represent variability within the normal range. Indeed, the present effects were observed despite the fact that our sample represented a lower extent within the normal range for state anxiety (mean percentile = 32.56 ± 17.64; range = 6th–64th percentile). Future studies could determine the extent to which this effect would be observed in normal subjects with higher levels of state anxiety.

It is also noteworthy that trait anxiety scores were relatively low across our subjects (mean percentile = 29.95 ± 13.97; range = 9th–69th percentile). Although this represents a weakness in terms of population estimation, this homogeneity in trait anxiety scores might have been fortuitous. Some models of anxiety posit that state anxiety will interact differentially with low versus high trait anxiety (MacLeod and Mathews 1988; see Egloff and Hock 2001; Eysenck 1992; Mathews and MacLeod 1994; Williams et al 1997; also see Beck 1976; Bower 1981). Thus, it is possible that state anxiety could show a different relationship with amygdala response in high trait-anxious subjects. Such studies would provide important data for predictions concerning application of the present paradigm to the study of pathologic anxiety.

Numerous neuroimaging studies have demonstrated that facial expressions of emotion represent convenient and well-tolerated stimuli that produce reliable activation of the human amygdala. These studies proceed against the historical backdrop of an animal literature that has focused on the role of the amygdala in aversive conditioning. Broadly consistent with this work, fearful facial expressions have been shown to produce reliable activation of the human amygdala. Comparison of amygdala response to other facial expressions represents a strategy for better understanding amygdala response to fearful expressions and human amygdala function more generally. The present study demonstrates amygdala responsiveness to both happy and neutral faces within the dorsal amygdala/SI region, consistent with previous studies. Discrimination of amygdala responses to these two facial conditions occurred within the ventral amygdala, although 1) the temporal aspects of these responses differed from that previously observed to fearful expressions; and 2) responses were more variable across subjects compared with those observed within the dorsal amygdala/SI. Indeed, here we documented that ventral amygdala response to happy versus neutral faces was related to individual state anxiety levels, and response to neutral faces explained the majority of the variability associated with this effect. Future studies will be needed to determine whether use of the current experimental context (devoid of primary negatively valenced expressions) is necessary to produce these effects.

This research was supported by a National Institute of Mental Health grant (MH-01866) and by the Howard Hughes Medical Institute.

We thank Ashly McLean, Sara Polis, Caroline Clemmons, Donald McLaren, Michael Anderle, and Ron Fisher for technical assistance, and Lisa Shin, Ned Kalin, Richard Davidson, Bill Kelley, and Scott Rauch for advice and comments.

L.H. Somerville et al


www.elsevier.com/locate/biopsych


www.elsevier.com/locate/biopsych