Failure of Anterior Cingulate Activation and Connectivity With the Amygdala During Implicit Regulation of Emotional Processing in Generalized Anxiety Disorder

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Objective: Clinical data suggest that abnormalities in the regulation of emotional processing contribute to the pathophysiology of generalized anxiety disorder, yet these abnormalities remain poorly understood at the neurobiological level. The authors recently reported that in healthy volunteers the pregenual anterior cingulate regulates emotional conflict on a trial-by-trial basis by dampening activity in the amygdala. The authors also showed that this process is specific to the regulation of emotional, compared to nonemotional, conflict. Here the authors examined whether this form of noninstructed emotion regulation is perturbed in generalized anxiety disorder.

Method: Seventeen patients with generalized anxiety disorder and 24 healthy comparison subjects underwent functional MRI while performing an emotional conflict task that involved categorizing facial affect while ignoring overlaid affect label words. Behavioral and neural measures were used to compare trial-by-trial changes in conflict regulation.

Results: Comparison subjects effectively regulated emotional conflict from trial to trial, even though they were unaware of having done so. By contrast, patients with generalized anxiety disorder were completely unable to regulate emotional conflict and failed to engage the pregenual anterior cingulate in ways that would dampen amygdalar activity. Moreover, performance and brain activation were correlated with symptoms and could be used to accurately classify the two groups.

Conclusions: These data demonstrate that patients with generalized anxiety disorder show significant deficits in the noninstructed and spontaneous regulation of emotional processing. Conceptualization of anxiety as importantly involving abnormalities in emotion regulation, particularly a type occurring outside of awareness, may open up avenues for novel treatments, such as by targeting the medial prefrontal cortex.
while abnormalities appear to exist within both limbic and prefrontal regions in generalized anxiety disorder, the nature of these abnormalities remains poorly understood.

Use of an experimental paradigm in which emotion regulation can be tracked from trial to trial may be a fruitful approach to understanding generalized anxiety disorder. We recently reported (15, 16) on a facial affect identification emotional conflict task in which healthy volunteers were asked to identify the expression of a face (fearful or happy) while ignoring an overlying emotion word (“fear” or “happy”) that either matched (congruent) or conflicted (incongruent) with the facial expression. Reaction time interference by emotionally incongruent stimuli was seen in nearly every participant (15, 16). Interestingly, there is less conflict, indexed by faster reaction times, for incongruent trials if they are preceded by an incongruent trial than if they are preceded by a congruent trial (15–20), which suggests that the emotional conflict generated by incongruency on the previous trial activates a regulatory mechanism that leads to improved emotional conflict regulation on the current incongruent trial (16, 21–23), thus optimizing task performance. We termed this across-trial effect “emotional conflict adaptation” (15, 16), in reference to the label previously applied to similar congruency sequence effects observed in nonemotional conflict tasks (21). Likewise, performance on postcongruent congruent trials is often superior to that on postincongruent congruent trials (21).

To date, the cognitive model that best accounts for the conflict adaptation effect, after eliminating potential confounders (24), is the “conflict monitoring hypothesis” (17–23, 25, 26). According to this model, conflict is continuously evaluated, such that greater conflict regulation can be flexibly recruited as required by the amount of conflict. Thus, the conflict-monitoring hypothesis distinguishes between two important functions—conflict evaluation and conflict regulation. Many studies have examined the regions associated with these functions during adaptation to nonemotional conflict (e.g., color-word Stroop or flanker tasks) by comparing activity during incongruent trials that differ only with respect to whether they were preceded by a congruent or an incongruent trial (17, 18, 20–23, 25, 26). Regions whose activity tracks the amount of conflict (i.e., postcongruent incongruent trials > postcongruent congruent trials) have been interpreted as conflict evaluation regions (17, 18, 20–23, 25, 26). Regions showing the opposite effect (postincongruent incongruent trials > postcongruent incongruent trials) have been interpreted as conflict regulation regions (17, 18, 20–23, 25, 26), as activity in these regions is greatest when conflict is minimized through regulation. Because these contrasts compare physically identical incongruent trials, the behavioral and neural effects differ only by virtue of expectation created by conflict on the previous trial (17–23, 25, 26).

In previous studies (15, 16) we applied this logic to the analysis of adaptation in a novel emotional conflict task. Greater activity during postincongruent incongruent trials (i.e., regulation-related) was seen in the pregenual anterior cingulate, and this was accompanied by strong negative coupling between the pregenual cingulate and the amygdala. These findings are consistent with other contexts in which emotion regulation is observed (27–29). By contrast, greater activity during postcongruent incongruent trials (i.e., evaluation-related) was seen in the amygdala and the dorsal anterior cingulate/dorsomedial prefrontal cortex.

We also compared activations during emotional conflict adaptation with those during nonemotional conflict adaptation (gender identification with the same emotional faces, while ignoring gender words overlaid on the faces) to determine the specificity of activations for emotion. Pregenual cingulate activation and coupling with the amygdala was specific to emotional conflict adaptation, whereas dorsal anterior cingulate/dorsomedial prefrontal cortex activation was shared by emotional and nonemotional conflict (15, 16), consistent with the role of

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### TABLE 1. Demographic and Clinical Characteristics of Healthy Comparison Subjects and Patients With Generalized Anxiety Disorder in an fMRI Study

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Comparison Subjects (N=24)</th>
<th>Patients With Generalized Anxiety Disorder (N=17)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>%</td>
<td>N</td>
</tr>
<tr>
<td>Female</td>
<td>18</td>
<td>75</td>
<td>11</td>
</tr>
<tr>
<td>Right-handed</td>
<td>24</td>
<td>100</td>
<td>17</td>
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<tr>
<td>Age (years)</td>
<td>36.5</td>
<td>11.8</td>
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<tr>
<td>Age (years)</td>
<td>17.3</td>
<td>2.0</td>
<td>16.6</td>
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<tr>
<td>Spielberger Trait Anxiety score</td>
<td>30.6</td>
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<td>Penn State Worry Questionnaire score</td>
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<td>8.3</td>
<td>61.9</td>
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<td>Beck Anxiety Inventory score</td>
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<td>3.4</td>
<td>21.6</td>
</tr>
<tr>
<td>Beck Depression Inventory score</td>
<td>3.6</td>
<td>3.4</td>
<td>14.6</td>
</tr>
<tr>
<td>Mood and Anxiety Symptom Questionnaire</td>
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<td></td>
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<tr>
<td>Anxious arousal subscale score</td>
<td>18.2</td>
<td>1.5</td>
<td>25.4</td>
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<tr>
<td>Anhedonic depression subscale score</td>
<td>48.6</td>
<td>10.3</td>
<td>69.2</td>
</tr>
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</table>
regulation of emotional processing is at the core of this disorder, we hypothesized that patients would show abnormalities in adapting to emotional conflict in our task. Additionally, to better understand emotional conflict adaptation more generally, and thus enhance interpretation of these latter regions in the evaluation of conflict in many other studies of nonemotional conflict adaptation (17, 20, 25, 26).

Considering that the clinical phenomenology of generalized anxiety disorder suggests that a deficit in the

FIGURE 1. Response to Emotional Conflict in Patients With Generalized Anxiety Disorder and Healthy Comparison Subjects

Panel A shows a sample task time course illustrating the contrasts made to examine adaptation during congruent or incongruent trials. Panel B shows reaction time difference scores reflecting the overall effect of emotional conflict (incongruent minus congruent trials), the facilitation in reaction times during emotional conflict adaptation (postincongruent incongruent trials [iI] faster than postcongruent incongruent trials [cI], resulting in a negative reaction time difference score), and similar adaptation on congruent trials (postcongruent congruent trials [cC] faster than postincongruent congruent trials [iC]). A group difference was observed only during adaptation on incongruent trials. The inset in panel B shows reaction times for each condition (a detailed table is presented in the online data supplement).

\* Panel A shows a sample task time course illustrating the contrasts made to examine adaptation during congruent or incongruent trials. Panel B shows reaction time difference scores reflecting the overall effect of emotional conflict (incongruent minus congruent trials), the facilitation in reaction times during emotional conflict adaptation (postincongruent incongruent trials [iI] faster than postcongruent incongruent trials [cI], resulting in a negative reaction time difference score), and similar adaptation on congruent trials (postcongruent congruent trials [cC] faster than postincongruent congruent trials [iC]). A group difference was observed only during adaptation on incongruent trials. The inset in panel B shows reaction times for each condition (a detailed table is presented in the online data supplement).

\* One-sample t test, p<0.001.
\* Two-sample t test, p<0.05.
\* One-sample t test, p<0.05.
\* One-sample t test, p<0.01.
of abnormalities in patients, we investigated in a separate cohort of healthy volunteers whether they were aware of these trial-to-trial adaptation effects and thus whether conscious attention is required for this process. We hypothesized that participants would not be aware of the adaptation effect and thus that this process is carried out at an implicit level.

**Method**

**Participants**

A total of 41 individuals, recruited locally through online advertisements, participated in the functional MRI (fMRI) component of this study; all provided informed consent. DSM-IV-based psychiatric diagnoses were determined through both an informal clinical interview with a psychiatrist and the Mini-International Neuropsychiatric Interview, a structured diagnostic interview (30, 31). Generalized anxiety disorder was the primary diagnosis for all patients, in terms of both onset and severity. Exclusion criteria were major depressive disorder, bipolar disorders, psychotic disorders, substance abuse, and posttraumatic stress disorder; a history of a neurological disorder, head trauma, or loss of consciousness; claustrophobia; or regular use of benzodiazepines, opioids, or thyroid medications. No patient was taking regular psychiatric medications or had used a benzodiazepine within 48 hours of the scan. No patient had ever received an evidence-based structured psychotherapy, and only five patients had ever received antidepressant medication. Nine patients had no comorbid disorders, five had one comorbid disorder (two with dysthymia and three with social anxiety), three had two comorbid disorders (two with social anxiety and panic disorder, and one with social anxiety and obsessive-compulsive disorder), and none had more than two comorbid disorders. All comparison subjects were free of any current or past axis I conditions or psychiatric medications. All participants completed the Spielberger State-Trait Anxiety Inventory (32), the Penn State Worry Questionnaire (33), the Beck Anxiety Inventory (34), the Beck Depression Inventory (35), and the Mood and Anxiety Symptoms Questionnaire (36, 37), from which the anxious arousal and anhedonic depression subscales were used. Resting state data from nine of the healthy comparison subjects and 10 of the patients were included in a previous study (12). The behavior-only study was conducted on a group of 19 healthy volunteers (mean age=25.2 years [SD=1.0]; 13 of them women) that did not overlap with the healthy comparison group involved in the fMRI study.

**Experimental Paradigm**

The emotional conflict task was performed as previously described (15, 16). Stimuli were presented with the Presentation software package (Neurobehavioral Systems, http://nbs.neuro-bs.com) during fMRI scanning and displayed through a custom-built MRI-compatible projection system. The task consisted of 148 presentations of happy or fearful facial expression photographs drawn from the set of Ekman and Friesen (38), overlaid with the words “FEAR” or “HAPPY.” Stimuli were presented for 1,000 msec, with a varying interstimulus interval of 3,000–5,000 msec (mean=4,000 msec), in a pseudorandom order, counterbalanced across trial types for expression, word, response button, and gender. Participants indicated facial affect with a button press response. Behavioral data were analyzed in SPSS (SPSS, Inc., Chicago). For the behavior-only task, a questionnaire was administered after the task to assess participants’ awareness of the conflict adaptation effect.

**fMRI Data Acquisition**

Images were acquired on a 3-T GE Signa scanner using a custom-built head coil. Twenty-nine axial slices (4.0 mm thickness with a 0.5 mm gap) were acquired across the whole brain using a T1*-weighted gradient echo spiral pulse sequence (repetition time=2,000 msec, echo time=30 msec, flip angle=80°, interleave=1, field of view=22 cm, matrix=64×64) (39). To reduce blurring and signal loss arising from field inhomogeneities, an automated high-order shimming method based on spiral acquisitions was used before acquisition of functional MRI scans (40). A high-resolution T1*-weighted three-dimensional inversion recovery spoiled gradient-recalled acquisition in the steady state MRI sequence was used with the following parameters: inversion time=300 msec, repetition time=8 msec; echo time=3.6 msec; flip angle=15°; field of view=22 cm; 124 slices in coronal plane; matrix=256×192; number of excitations=2; acquired resolution=1.5×0.9×1.1 mm. The images were reconstructed as a 124×256×256 matrix.

**fMRI Data Analysis**

The first five volumes were not analyzed to allow for signal equilibration effects. A linear shim correction was applied separately for each slice during reconstruction using a magnetic field map acquired automatically by the pulse sequence at the beginning of the scan (39). Functional MRI data were then preprocessed using the SPM5 software package (http://www.fil.ion.ucl.ac.uk/
FIGURE 3. Activation of the Pregenual Cingulate and Modulation of the Dorsomedial Prefrontal Activity During Emotional Conflict Adaptation in Healthy Comparison Subjects and Patients With Generalized Anxiety Disorder

Panel A shows the healthy comparison subject > patient contrast for the postincongruent incongruent trial (iI) minus postcongruent incongruent trial (cI) difference within the pregenual cingulate region of interest; panel B shows each group’s data extracted for the cluster, for both difference scores and individual trial types (inset). The pregenual cingulate was activated only in healthy comparison subjects. Panel C shows the healthy comparison subject > patient contrast for the postcongruent incongruent trial (cI) minus postincongruent incongruent trial (iI) difference within the dorsomedial prefrontal region of interest; panel D shows each group’s data extracted for the cluster, for both difference scores and individual trial types (inset). Healthy comparison subjects were found to exhibit less dorsomedial prefrontal activity in postincongruent incongruent trials (hence a positive difference score). By contrast, in patients, there was inappropriately greater activity in the dorsomedial prefrontal cortex in response to postincongruent incongruent trials (i.e., negative difference scores).

1 Two-sample t test, p<0.01.
2 One-sample t test, p<0.01.
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Finally, we asked a separate group of healthy volunteers whether they were aware of any pattern across trials that might help or hinder their performance. No participant mentioned previous trial conflict. In addition, discrimination in a forced-choice question of whether performance on a current incongruent trial was improved by a previous incongruent trial compared to a previous congruent trial did not differ from chance (p>0.25), which suggests that conscious awareness of the adaptation phenomenon is not required for successful adaptation.

Abnormal Medial Prefrontal Responses to Emotional Conflict in Patients

We first examined overall responses to emotional conflict (i.e., incongruent > congruent). As shown in Figure 2A, healthy comparison subjects exhibited greater activation to emotional conflict than did patients with generalized anxiety disorder in the dorsomedial prefrontal cortex (x=0, y=36, z=38; z=3.96; d=1.22; 2,832 mm³; x=6, y=44, z=34; z=3.33; d=1.14). This difference resulted from activation by emotional conflict within this cluster in comparison subjects (t=3.9, df=23, p=0.001; d=0.8) but not in patients (t=1.84, df=16, p=0.05; see Figure 2B). No group differences were observed in the pregenual cingulate or the amygdala.

Next, we explored the neural correlates of group differences in emotional conflict adaptation, guided by our behavioral results. Based on our previous findings with the emotional conflict task in healthy volunteers (15, 16), we examined the contrast of postincongruent incongruent trials minus postcongruent incongruent trials in the pregenual cingulate and the amygdala in patients with generalized anxiety disorder.
jects and found a significant cluster (x=−12, y=32, z=−4; z=3.49; 376 mm³; d=1.2; see Figure 3A). Average signal within this cluster was extracted for each group to further describe the effect. As predicted, in this cluster, healthy comparison subjects had greater activity during postincongruent incongruent trials (t=3.34, df=23, p<0.005; d=0.68), whereas in patients no difference was observed (t=1.6, df=16, p>0.1; see Figure 3B).

Next, we examined the contrast of postcongruent incongruent trials minus postincongruent incongruent trials in the dorsomedial prefrontal cortex and amygdala in both groups and found a significant cluster in the dorsomedial prefrontal cortex (x=−1, y=36, z=38; z=3.26; 568 mm³; d=1.15; see Figure 3C) but not in the amygdala. Extraction of average signal within this cluster revealed that the group difference was driven by the expected greater activity in postcongruent incongruent trials in healthy comparison subjects (t=2.36, df=23, p<0.05; d=0.48) and by the opposite effect in patients (t=2.66, df=16, p<0.05; d=0.64; see Figure 3D). Note that the inability of patients to decrease dorsomedial prefrontal activity in postincongruent incongruent trials paralleled patients’ inability to improve reaction times during these trials compared to healthy comparison subjects. No group differences were observed in any of the regions of interest for the contrast of postcongruent congruent trials with postincongruent congruent trials. Finally, comparing across all trial types, we found significantly greater activation in patients than in comparison subjects in the left amygdala (x=−22, y=−2, z=−18; z=3.04; 232 mm³; d=1.04). Using cytoarchitectonic probability maps of the basolateral, centromedial, and superficial amygdalar subregions (47, 48), we found that 78% of this cluster corresponded to the superficial amygdala and 21.1% to the basolateral amygdala.

**Absent Pregenual Cingulate-Amygdala Connectivity in Patients**

We next examined differential functional connectivity between the pregenual cingulate and the amygdala during postincongruent incongruent trials compared with postcongruent incongruent trials using psychophysiolgic interaction analyses, with the pregenual cingulate as the seed and the amygdala as the target, while controlling for task-related activations in both regions and task-nonspecific connectivity (44). As shown in Figure 4A, we found a significant group difference in both the left (x=−20, y=−4, z=−22; z=3.54; 536 mm³; d=1.15) and right amygdala (x=30, y=−4, z=−22; z=3.4; 168 mm³; d=1.15). Extraction of average connectivity strength within these clusters revealed that the group effect resulted from the predicted significant negative pregenual cingulate-amygdala connectivity in healthy comparison subjects during postincongruent incongruent trials, compared with postincongruent incongruent trials (left side: t=4.14, df=23, p<0.001; d=0.85; right side: t=3.08, df=23, p=0.005; d=0.63), but not in patients (see Figure 4B). We did not pursue further characterization of differential group cingulate-amygdala connectivity using effective connectivity methods such as dynamic causal modeling, as we had in a previous study of healthy volunteers (16), since we did not think it would add significant new information beyond the result from the functional connectivity analysis above. Finally, we found that the majority of the left amygdala differential connectivity cluster was in the basolateral amygdala (55%), with 44.3% in the superficial amygdala and only 0.4% in the centromedial amygdala. One hundred percent of the right amygdala cluster was in the basolateral amygdala.

**Additional Findings**

We conducted several additional analyses to better understand the group differences reported above. First, for the patients, we correlated symptom scale scores with behavior and brain activity within the group difference clusters. We found that the impairment in emotional conflict adaptation was greatest, in terms of both reaction times and dorsomedial prefrontal modulation, for the most anxious patients (see the online data supplement). Second, we conducted multivariate pattern classification to determine whether behavior and brain activation could be used to determine participants’ diagnostic group. Significant classification of patients and healthy comparison subjects could be achieved with both behavior and brain activation data, reaching 95% when whole-brain data were used (see the online data supplement).

**Discussion**

In this study, we investigated emotional conflict adaptation using a paradigm in which emotional processing is regulated spontaneously and in the absence of explicit instruction. We found that patients with generalized anxiety disorder were unable to adapt to emotional conflict through engagement of this regulatory process. By contrast, adaptation during congruent trials was similar in both groups, as was the overall reaction time interference due to emotional conflict, demonstrating the specificity of the deficit.

At the neural level, patients with generalized anxiety disorder failed to activate the pregenual cingulate and demonstrate negative top-down (16) pregenual cingulate-amygdala connectivity during the regulation of emotional conflict. As in previous studies of emotion regulation (27, 28), regulation-related changes in activity were seen in the context of overall task-independent medial prefrontal deactivation from an implicitly modeled baseline, and this deactivation did not differ between our groups (data not shown). Moreover, since the critical contrast involves only incongruent trials, the many processes that differ between incongruent and congruent stimuli are controlled for, as are nonspecific responses to task demands, leaving only the effect of previous trial conflict on processing of emotional conflict on the current trial.
We suggest that patients’ failure to show the neural effects related to previous trial conflict accounts for their behavioral regulatory deficit during emotional conflict adaptation, in accordance with predictions made by the conflict monitoring hypothesis about brain activity during conflict adaptation—a cognitive model supported by an extensive neuroimaging literature (17, 18, 20–23, 25, 26). These conclusions are also consistent with emotion regulatory roles attributed to ventromedial prefrontal regions through connectivity with the amygdala in other studies (27–29, 49). Moreover, we recently found functional connectivity and structural evidence for an intra-amygdalar abnormality at a subregional level in generalized anxiety disorder (12). Thus, it appears that patients have deficits both in activating relevant control regions (pregenual cingulate) and in the connectivity required for such regions to exert control over limbic structures. It is therefore interesting to speculate that the lateral prefrontal hyperactivation (7, 9, 14) or increased connectivity with the amygdala (12) previously reported in patients with generalized anxiety disorder may reflect the compensatory engagement of worry, an attention-demanding cognitive process, to regulate the effects of emotional stimuli in the absence of patients being otherwise able to recruit pregenual cingulate-based regulatory mechanisms.

Finally, there is controversy regarding overall emotional responsiveness in generalized anxiety disorder, indexed largely through activation in the amygdala. Pediatric studies have shown hyperactivity to negative emotional expression faces (7, 8), and adult studies have shown either no difference (10) or hypoactivation (9) to similar stimuli, although one adult study showed nonspecifically exaggerated amygdalar reactivity to both negative emotional and neutral cues (11). Consistent with the latter study, we found greater amygdalar activity in patients during both congruent and incongruent trials.

Our data also highlight an emerging theme in affective neuroscience, namely, that there are many ways by which emotional processing is regulated and that deficits in these functions contribute importantly to psychopathology. To date, the neurobiology underlying the regulation of emotional processing has been primarily studied by asking participants to deliberately alter their emotional responses to defined stimuli (i.e., “explicit” regulation) (50). Much of the normal regulation of emotional processing, however, probably occurs in the absence of explicit effort (51, 52). Far less is known about these “implicit” forms of regulation.

Based on the fact that behavioral performance in our task indicates the engagement of a mechanism for regulating emotional processing that occurs in the absence of specific regulation instructions, we and others have argued that emotional conflict adaptation is a type of implicit regulation of emotional processing (51, 52). In this study, we provide direct behavioral evidence for this in healthy participants. The striking deficit in emotional conflict regulation in patients, in the context of otherwise intact task performance, provides the strongest evidence to date linking abnormalities in a defined form of implicit regulation and a type of psychopathology whose clinical presentation suggests emotion regulatory abnormalities.

Several limitations are important to note. First, we are unable to report on subjective ratings of emotion during emotional conflict adaptation, as asking participants to report on subjective emotional states might itself lead to emotion regulation (53–57). Thus, we inferred the effects of emotion from behavioral indices, such as reaction times, and patterns of brain activation. Second, although we focused in this study primarily on the neural effects we previously found to be specific to emotional conflict (15), it would be useful in future experiments also to examine adaptation to nonemotional conflict. Finally, it is unknown whether medial prefrontal dysfunction during emotional conflict adaptation reflects a disorder-specific abnormality or a more general endophenotype of affective disorders, such as major depression. Nonetheless, the robust group differences seen at both the behavioral and neural levels suggest that the inability of patients to adapt to emotional conflict is an important aspect of the pathophysiology of generalized anxiety disorder—and potentially of other psychiatric disorders—and thus merits continued, deeper, study.

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Dr. Etkin has served as a consultant for Neostim. Dr. Schatzberg has served as a consultant to BrainCells, CeNeRx, CNS Response, Corcept, Eli Lilly, Forest Labs, GlaxoSmithKline, Inntapharma, Lundbeck, Merck, Neuronetics, Novartis, Pathway Diagnostics, Pfizer, PharmaNeuroBoost, Quintiles, Sanofi-Aventis, Synoasis, Takeda, Xytis, and Wyeth and has received speaking fees from GlaxoSmithKline and Roche; he has equity holdings in BrainCells, CeNeRx, Corcept (co-founder), Forest, Merck, Neurocrine, Pfizer, PharmaNeuroBoost, Somaxon, and Synoasis; and he was named an inventor on pharmacogenetic use patents on prediction of antidepressant response. The other authors report no financial relationships with commercial interests. Supported by funds for a residency research program of the Veterans Affairs-Palo Alto Health Care System to Dr. Etkin, and NIH grants HD047520 and NS058899 to Dr. Menon and 5K23HD054720 to Dr. Hoef.

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Supplemental Methods and Table S1

Supplemental Methods

Multivariate pattern classification

For the multivariate pattern classification analysis (1, 2), we converted the relevant individual-level contrast images, masked by the a priori regions of interest described in the main text, or a whole brain gray matter mask, into a matrix of vectors, using the set of 24 controls and 17 patients. Classification dimensions were reduced by recursive feature elimination (1, 2), using in-house tools based on Matlab (3). To do so, we constructed a classifier using all relevant voxels and rank ordered each voxel’s contribution to the discrimination. We iteratively removed the 40% worst-discriminating voxels, stopping at the point at which performance of the classifier began to deteriorate. This procedure determined the minimum number of required features (voxels). We then performed linear support vector machine-based classification analysis (regularization parameter C=1), with leave-one-out cross-validation. Significance was determined by randomly reassigning class labels in 2000 permutations of the leave-one-out analysis using the voxels identified by the recursive feature elimination process (alpha = p<0.05, two-sided).

Results

Correlations with symptom scales

For the patients, we correlated symptom scale scores with the reaction time difference between postincongruent incongruent trials minus postcongruent incongruent trials. The only
correlation that survived a Bonferroni correction for the six comparisons made was between greater scores on the anxious arousal subscale of the mood and anxiety symptom questionnaire and progressively greater reaction time difference scores (i.e. greater sensitization; \( r=0.68, p<0.005 \)). This correlation is particularly interesting, because this subscale was designed to differentiate anxiety from depression (4, 5). Indeed, after controlling for the same questionnaire’s depression-specific measure, the anhedonic depression subscale, which was not significantly correlated with either the reaction time difference scores \( (r=0.32, p>0.2) \) or the anxious arousal subscale \( (r=-0.1, p>0.7) \), the correlation between the anxious arousal subscale and the reaction time difference scores rose to \( r=0.75 \) \( (p=0.001) \); see Figure S1A). Independence from outliers in this correlation was confirmed with robust regression (data not shown) as well as by removing the potential outlier value at the top-right of the plot in supplemental Figure S1A \( (r=0.54, p<0.05) \).

Correlations were also made between symptom scale scores with the average difference between postincongruent incongruent trials minus postcongruent incongruent trials for the group contrast clusters in the pregenual cingulate and dorsomedial prefrontal cortices. After Bonferroni correction, the anxious arousal subscale and Penn State Worry Questionnaire were found to correlate with greater inappropriate dorsomedial prefrontal activation in postincongruent incongruent trials (anxious arousal: \( r=0.64, p=0.005 \); worry: \( r=0.68, p<0.005 \); see supplemental Figure S1B and S1C). After removing variance associated with the anhedonic depression subscale, which did not correlate with dorsomedial prefrontal activity \( (r=0.04, p>0.8) \), as well as worry scores, the correlation between dorsomedial prefrontal activity and the anxious arousal subscale remained strong \( (r=0.6, p<0.05) \); see supplemental Figure S1B). Likewise, dorsomedial prefrontal activity correlated with worry scores after removing variance associated with anxious
arousal and anhedonic depression scores ($r=0.63$, $p=0.01$, see supplemental Figure S1C). Worry scores and anxious arousal scores were not significantly correlated ($r=0.35$, $p>0.15$).

**FIGURE S1. Correlations Between Anxiety Symptoms and Reaction Times**

(A) A significant positive correlation in patients between the anxious arousal subscale of the Mood and Anxiety Symptom Questionnaire and reaction time difference scores for the contrast of postincongruent incongruent trials minus postcongruent incongruent trials (iI-cI), indicating worse performance, rather than facilitation, to repeated incongruent trials in more anxious individuals. Psychometric and behavioral data are expressed as Z-scores after accounting for variance related to the anhedonic depression subscale of the same questionnaire. (B, C) Greater inappropriate dorsomedial prefrontal activation is correlated with higher levels of anxiety. Correlations between iI-cI signal for patients in the group difference dorsomedial prefrontal cluster from Figure 3C and anxious arousal subscale scores adjusted for anhedonic depression subscale and worry scores (B), or the correlation with scores on the Penn State Worry Questionnaire (C), after adjusting for anxious arousal and anhedonic depression scores, all expressed as Z-scores.
Multivariate pattern classification

We implemented a linear support vector machine pattern classification approach, using recursive feature elimination for feature reduction and leave-one-out cross-validation, to determine whether fMRI signal in the emotional conflict task can discriminate between patients and controls. Within our *a priori* regions of interest, significant classification using the contrast of incongruent trials preceded by incongruent trials minus incongruent trials preceded by congruent trials was achieved in the pregenual cingulate (accuracy 88%, p<0.0005; sensitivity 88%, specificity 88%), and the dorsomedial prefrontal cortex (accuracy 76%, p<0.01; sensitivity 59%, specificity 88%), but not in the amygdala (accuracy 66%, p>0.1; sensitivity 24%, specificity 96%). Using whole brain data, we were able to achieve 95% accuracy (p<0.0005, sensitivity 94%, specificity 96%). Successful classification could also be achieved using reaction time difference scores (accuracy 73%, p<0.0005; sensitivity 53%, specificity 88%).

REFERENCES

3. Hoeft F, Lightbody AA, Hazlett HC, Patnaik S, Piven J, Reiss AL: Morphometric spatial patterns differentiating boys with fragile X syndrome, typically developing boys, and developmentally delayed boys aged 1 to 3 years. Arch Gen Psychiatry 2008; 65:1087–1097
### TABLE S1: Reaction Times (standard deviations in parenthesis)

<table>
<thead>
<tr>
<th></th>
<th>Healthy subjects</th>
<th>Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>Current trial</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>congruent</td>
<td>incongruent</td>
</tr>
<tr>
<td><strong>Previous trial</strong></td>
<td>congruent</td>
<td>736.6 (95.6)</td>
</tr>
<tr>
<td></td>
<td>incongruent</td>
<td>762.9 (84.8)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>850.2 (152.4)</td>
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<tr>
<td></td>
<td></td>
<td>821.4 (117.1)</td>
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