The Aftermath of 9/11: Effect of Intensity and Recency of Trauma on Outcome

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Does trauma exposure have a long-term impact on the brain and behavior of healthy individuals? The authors used functional magnetic resonance imaging to assess the impact of proximity to the disaster of September 11, 2001, on amygdala function in 22 healthy adults. More than three years after the terrorist attacks, bilateral amygdala activity in response to viewing fearful faces compared to calm ones was higher in people who were within 1.5 miles of the World Trade Center on 9/11, relative to those who were living more than 200 miles away (all were living in the New York metropolitan area at time of scan). This activity mediated the relationship between group status and current symptoms of posttraumatic stress disorder. In turn, the effect of group status on both amygdala activation (fearful vs. calm faces) and current symptoms was statistically explained by time since worst trauma in lifetime and intensity of worst trauma, as indicated by reported symptoms at time of the trauma. These data are consistent with a model of heightened amygdala reactivity following high-intensity trauma exposure, with relatively slow recovery.

Keywords: amygdala, trauma, stress, neuroplasticity, 9/11

Psychological traumas have been defined as events that threaten death or injury to self or others and that engender intense feelings of fear, helplessness, or horror (e.g., rape, combat, witnessing violence or disaster, or the sudden death of a loved one; American Psychiatric Association, 2000). Trauma exposure is a potent environmental risk factor that predicts immediate and lifetime increases in a diverse array of mental health disorders (Breslau et al., 1998; Kessler, Sonnega, Bromet, Hughes, & Nelson, 1995), including anxiety and depression (Brown, 1993; Kendler, Hettema, Butera, Gardner, & Prescott, 2003; McCauley, Kern, Kolodner, Dill, & Schroeder, 1997). Trauma exposure is also a definitional prerequisite for posttraumatic stress disorder (PTSD) and a predisposing factor for further incidence of PTSD if there is a subsequent trauma (Bremner, Southwick, Johnson, Yehuda, & Charney, 1993). In cases where trauma exposure is severe and protracted, rates of mortality and chronic illness far exceed the norm later in life, and mean life expectancy is sharply decreased (McFarlane, 1997).

Trauma exposure is not uncommon. Results from an epidemiological study of 5,877 people within the United States found that more than 50% of women and 60% of men experienced at least one trauma in their lifetime, and more than a quarter of the sample experienced two or more traumas (Kessler et al., 1995). Although only a small percentage of the trauma-exposed population develops PTSD (6% to 9%, Breslau et al., 1998; 8% to 20%, Kessler et al., 1995), this disorder has been the focus of studies of the effects of trauma on humans. The response to trauma in people without a specific clinical disorder has been less well characterized (Bunce, Larsen, & Peterson, 1995; McFarlane, 1997). Thus, although trauma exposure is an established environmental risk factor for a wide range of disorders, the neural mechanisms underlying this overall increase in risk are unclear.

A small number of epidemiological and community-based studies have examined the effects of trauma exposure in individuals without PTSD (Otte et al., 2005; Yehuda, Golier, & Kaufman, 2005; Young & Breslau, 2004a, 2004b; Young, Tolman, Witkowski, & Kaplan, 2004) by using peripheral biological indicators of the stress response (e.g., urinary cortisol, dopamine, epinephrine). From these efforts, the physiological correlates of trauma exposure in healthy (nonclinical) populations are beginning to emerge. For example, trauma-exposed young adults without PTSD were found to have significantly lower urinary catecholamine levels (dopamine and epinephrine) than nontrauma-exposed individuals, whereas a group with lifetime PTSD
had higher levels (Young & Breslau, 2004b). Another study (Otte et al., 2005) found significantly higher basal levels of MHPG (3-methoxy-4-hydroxy-phenylglycol, a major metabolite of noradrenaline) and greater MHPG reactivity in individuals with childhood trauma as compared to those with no childhood trauma. Although these findings are not conclusive, they are suggestive of long-term effects of trauma exposure on the central nervous system that may be more directly investigated with noninvasive neuroimaging techniques.

There have been a number of neuroimaging studies examining brain function and structure in individuals with PTSD. This research has provided substantial evidence for long-term central nervous system effects of trauma exposure that is accompanied by PTSD. Some of these studies have found enhanced amygdala activation in response to a variety of negatively valenced stimuli (e.g., Rauch et al., 2000; Shin et al., 2004; although, see Lanius et al., 2003; Phan, Britton, Taylor, Fig, & Liberzon, 2006; Sakamoto et al., 2005). The majority of this work has involved reexperiencing paradigms (e.g., Britton, Phan, Fig, Taylor, & Liberzon, 2005; Lanius et al., 2005), which include reinstatement of memories of the traumatic event. The format of these paradigms makes it difficult to assess whether reported results are specific to memory reinstatement or are a reflection of the disorder itself. It has also been argued (Williams et al., 2006) that the use of standardized probes of limbic activity allows for generalization of findings to other populations. As a consequence, a growing number of neuroimaging studies of PTSD have used standardized probes of amygdala activity (e.g., emotional faces, Armony, Corbo, Clement, & Brunet, 2005; Rauch et al., 2000; Shin et al., 2005; Williams et al., 2006; emotional scenes, Hendler at al., 2003; Phan et al., 2006; emotional Stroop task, Shin et al., 2001; emotionally valenced words, Bremner et al., 2003; Protopopescu et al., 2005).

The most frequently used standardized probe of amygdala activity in samples with PTSD has been presentation of emotional faces (Armony et al., 2005; Rauch et al., 2000; Shin et al., 2005; Williams et al., 2006). In general, these studies have reported increased amygdala activity to negative emotional expressions in those with PTSD. Shin and colleagues (2005) found increased activity in the right amygdala and decreased activity in medial prefrontal regions in men with chronic PTSD of long duration. Rauch et al. (2000) and Armony et al. (2005) found increased amygdala activation in samples with PTSD, but did not report significant differences in prefrontal activation. Williams et al. (2006) reported increased activity in ventral amygdala and decreased activity in dorsal amygdala in a mixed sample of men and women with PTSD of relatively short duration. Thus, increased activity in the amygdala is the most consistently observed neural correlate of PTSD as probed with emotional faces, with some evidence that this activation is localized to subregions of the amygdala and is associated with alterations in medial prefrontal activity.

In comparison to the rich body of neuroimaging research on PTSD, there has been very little work examining the neural correlates of trauma exposure in people without a clinical disorder; this is surprising, considering the known impact of negative life events on psychological distress and disorder in the overall population (e.g., Dohrenwend, 2006). One recent structural MRI paper has addressed this question from a developmental perspective (Cohen et al., 2006), showing that retrospective report of accumulated adverse childhood events predicted smaller anterior cingulate and caudate volumes in adulthood. Adverse childhood events with the most impact on adult brain morphology were those more likely to meet criteria as traumas (e.g., death of a parent, witnessing domestic violence, sexual assault). Also, a recent functional neuroimaging study (Sharot, Martorella, Delgado, & Phelps, in press) used a flashbulb memory paradigm that focused on events of September 11th. These results showed significant increases in left amygdala activation in response to evoked memories of the events of September 11th in adults who were in downtown Manhattan that day, relative to those who were in midtown Manhattan, five miles away.

Animal models of stress have highlighted the effects of severe stressors on the amygdala and related structures. Exposure to acute uncontrollable stressors produces extended hypere excitability of the amygdala in laboratory animals (Adamec, Blundell, & Burton, 2005; Maier & Watkins, 1998), which renders the amygdala and related structures more readily activated independently of the triggering stimulus (Rosen & Schuckin, 1998). This effect has been associated with increased vigilance and fearful responses (e.g., freezing) to ambiguous or mild standardized stressors (Adamec et al., 2005; Rosen & Schuckin, 1998). Exposure to acute stressors increases spine synapese formation in the basolateral amygdala (BLA), which may underlie the associated increases in anxiety-like behavior (Mitra, Jadhav, McEwen, & Chattarji, 2005; Vyaz, Mitra, Shankaranarayana Rao, & Chattarji, 2002). Exposure to repeated/chronic stressors produces anxiety-like behavior in response to standardized stressors (e.g., an open field), along with dendritic growth in the BLA, greater increases spine density than seen with acute stressors, and dendritic retraction in the hippocampus (McEwen, 2005; Mitra et al., 2005).

Taken together, these data suggest the hypothesis that the amygdala and closely related structures are persistently more reactive after trauma exposure in healthy adults (i.e., in individuals without a clinical disorder) and that these effects will be observable using mild, standardized stressors (i.e., that they do not require a reexperiencing paradigm). Epidemiological studies indicate that it is the nature of the worst trauma (i.e., the event that precipitates diagnosis and/or the most symptoms) that best predicts long-term negative psychological consequences of trauma exposure. Worst traumas that are more recent (Kessler et al., 1995) and more severe (e.g., violent assault: Breslau et al., 1998; Kessler et al., 1995) are more likely to predict distress and disorder. Selection of a sample with relatively recent, high-intensity, worst-trauma exposure would maximize the possibility of observing the hypothesized increase in amygdala reactivity.

The events of September 11, 2001, and the experiences of the men and women who were in close proximity to that disaster, provide a unique window into the neural correlates of environmental stressor exposure. Notably for this study, proximity to this disaster have been shown to predict psychological morbidity (Blanchard et al., 2004; Galea et al., 2002), making it possible to prospectively titer the effects of the disaster across study groups by studying participants who were near the World Trade Center (WTC) in New York City on September 11, 2001, relative to those who lived far away at that time. We hypothesize that relative increases in amygdala reactivity would be more apparent in the group with closer proximity to the WTC on September 11th. Building on previous assays of amygdala reactivity (Brieter et al., 1996; Thomas et al., 2001), participants
passively viewed blocks of fearful and calm faces while undergoing functional MRI (fMRI).

**Methods**

**Participants**

Twenty-two right-handed adults participated in this study; all were living in the New York metropolitan area at time of scan. Recruitment material requested participants who were at different distances from the WTC on September 11th. Eleven participants were within 1.5 miles of the WTC on September 11, 2001 (5 women, aged 30.3 ± 2.5 years [mean ± SE]; range = 19 to 41 years). Eleven lived at least 200 miles from the New York City area on September 11, 2001 (5 women, aged 29.3 ± 1.4 years; range = 23 to 37 years). People who lived in Washington, DC, on September 11th and those who had friends or relatives on aircraft involved in the disaster were excluded. Data were collected between 41 and 48 months after September 11th.

Before imaging, all participants were screened for current or past psychiatric, neurological, or medical illness and trauma exposure, as well as for any contraindications for fMRI. Approximately 30 minutes elapsed between time of interview and time of scan. This investigation was conducted within institutional guidelines established for protection of human subjects. All participants provided informed written consent.

**Behavioral Measures**

Three different clinical and standardized assessments were used to assess psychiatric symptoms and diagnosis on the day of the MRI visit: (1) the PTSD module of the University of Michigan Composite International Diagnostic Interview (UM-CIDI: Kessler et al., 1994) was used in conjunction with the Life History Calendar methodology developed by Caspi et al. (1996) to assess lifetime incidence of trauma exposure, current PTSD, and PTSD symptoms in lifetime. The UM-CIDI is a fully structured diagnostic interview that allows the assessment of current and lifetime mental disorders in the form of the third revised Diagnostic and Statistical Manual for Mental Disorders (American Psychiatric Association, 1986); (2) the Impact of Events Scale (Horowitz, Wilner, & Alvarez, 1979) is a 15-item scale that was used to assess PTSD stress reactions in the seven days before scanning. This investigation was conducted within institutional guidelines established for protection of human subjects. All participants provided informed written consent.

**Stimuli and Procedure**

Participants viewed gray-scaled pictures of faces of eight different actors (four female) demonstrating fearful and calm facial expressions (see Figure 1). Images were selected from a standardized picture set (Tottenham et al., in press). Face stimuli were presented on an overhead liquid crystal panel in a pseudorandom sequence of nine blocks of fixation, fearful faces, or calm faces. Participants were instructed, "Please look at the faces and the +s. You don’t have to press any buttons." There was no reference to the events of September 11th during imaging. Order of presentation was counterbalanced across subjects and across runs using the two following sequences: +FC + FC + FC or + CF + CF + CF (where F indicates a block of fearful faces, C indicates a block of calm faces, and + indicates fixation). In each block of faces, 10 images were presented for 4 seconds each. Fixation blocks were 30 seconds. Total block duration was 330 seconds.

**Imaging Protocols**

Subjects were scanned with a General Electric Signa 3-Tesla fMRI scanner (General Electric Medical Systems, Milwaukee, WI) using a quadrature head coil. After a three-plane localizer and a whole-head coronal localizer, a T2-weighted two-dimensional anatomic image with a fast spin-echo (FSE) sequence was acquired: time for repetition (TR) = 4000, time for echo (TE) = 68 ms, flip = 90°, field of view (FOV) = 20, 29 slices, 5-mm slice thickness, 0-mm gap, matrix = 256 × 192, axial-oblique. A three-dimensional spoiled gradient recalled (SPGR) T1-weighted anatomic scan was also acquired (124 axial slices, TR = 25 ms, TE = 5 ms, flip = 20°, FOV = 24 cm, 1.5-mm thickness, 0 mm gap, matrix = 256 × 256 × 160). Functional data was acquired using a spiral in-out sequence (Preston, Thomason, Oechsner, Cooper, & Glover, 2004) and the same spatial prescription as the FSE: TR = 2000 ms, TE = 30 ms, matrix = 64 × 64 mm, 29 slices per volume.

**Data Analysis**

Preprocessing and statistical analysis of fMRI data was performed by using Statistical Parametric Mapping (SPM2: Wellcome Department of Neurology, London, United Kingdom) implemented on MatLab 7.0. During preprocessing, the first four acquisitions were discarded and functional scans were realigned to the initial image, generating a set of realignment parameters for each run and a mean functional image. The mean functional image was used to coregister functional scans to the FSE anatomic images, which were then coregistered to the SPGR. The resulting parameters were used to realign the functional scans. The SPGR was then transformed to Montreal Neurological Institute (MNI) space, and these parameters were applied to the functional scans. The normalized functional data was smoothed by using a 6-mm full-width/half-maximum kernel.

Individual level analysis was performed by using the general linear model (Friston, Holmes, Price, Buchel, & Worsley, 1999).
with a fixed effects model. Contrast images comparing each of the block types were generated for each individual. These individual-level contrasts served as the basis for group-level random effects analyses (Friston et al., 1999). Data were first analyzed by using whole-brain analysis to identify significant areas activated to fearful versus calm faces in all participants. Then, a one-way analysis of variance was performed to compare brain activation across groups (9/11-exposed and control) for fearful versus calm faces. For all analyses, results were reported as significant if they met a voxel-wise threshold of \( p < .001 \), with clusters of 20 or more contiguous voxels (Forman, Cohen, Fitzgerald, Eddy, Mintun, & Noll, 1995). Given our strong, directional hypotheses regarding the amygdala, results were reported as significant if \( p < .01 \), with clusters of 3 or more voxels (this provides a conservative estimate of statistical significance: Foreman et al., 1995). In addition, small volume corrections from a 5-mm sphere around a priori locations of activation in the amygdala are reported (Worsley, 1996). All activations are reported using MNI coordinates.

In order to test a priori hypotheses regarding activation in the 9/11-exposed group relative to the comparison group, a region-of-interest (ROI) analysis was conducted. ROIs were defined functionally as the voxels found to be reliably activated in the whole-brain analysis of the fearful versus calm contrast at thresholds of \( p < .01 \), with cluster extents of three or more contiguous voxels. Because there were no medial prefrontal areas reliably activated in the whole brain analysis of variance, post hoc ROI analyses were limited to the amygdala. The association between the behavioral measures and blood oxygen-level dependent (BOLD) signal change in the amygdala for the fearful-calm contrast in each ROI was examined by using multiple regression analyses.

**Results**

**Behavioral Results**

**Demographics and trauma.** Comparison of demographic and trauma variables across groups revealed no significant differences in age, gender, age at first trauma, number of traumas in lifetime, number of traumas in lifetime with associated shock and/or horror, or history of PTSD (see Table 1). The control group had a significantly longer mean time since their worst trauma in lifetime than the 9/11-exposed group. The 9/11-exposed group also retrospectively reported more symptoms of avoidance and arousal at time of worst trauma in response to probes from the PTSD module of the UM-CIDI. Most, but not all, 9/11-exposed individuals rated the 9/11 disaster as the worst trauma that they had experienced in their lifetime.

**Current symptoms and anxiety.** Although the 9/11-exposed group did not meet diagnostic criteria for any disorder, they had a higher mean level of current symptoms, as assessed by both subscales of the IES (see Table 1). There were no significant differences between groups on the state measure of the STAI-S.

**Imaging Results**

BOLD signal in left and right amygdala was elevated in the group that was closer to the WTC on September 11th in the contrast of fearful versus calm faces (see Figure 2). Voxel-wise analysis of variance results indicated that the 9/11-exposed group had significantly greater activity than controls for fearful versus calm faces in the left amygdala \((-25, -9, -20; z = 3.13, p = .001, p_{svc} = .01, 19 \text{ contiguous voxels})\) and in the right amygdala \((-22, -9, -20; z = 2.97, p = .001, p_{svc} = .01, 5 \text{ contiguous voxels})\).

The ROI analysis also showed that the mean signal change in the left, \( t (22) = -3.3, p = .003 \) and right amygdala, \( t (22) = -2.7, p = .01 \) was higher in the 9/11-exposed group than in the comparison group for the fearful versus calm contrast. Signal change in these regions was not correlated with gender, age, number of traumas in lifetime, age at first trauma, or years since most recent trauma.

Table 2 shows correlations between BOLD signal change in right and left amygdala (fearful-calm contrast) with behavioral variables across the whole group. Signal change in the left amygdala was negatively correlated with IES intrusion and avoidance subscales and positively correlated with intrusion symptoms at worst trauma and number of avoidance symptoms at worst trauma. The relationship between signal change in the right amygdala and PTSD variables was weaker, with some correlations just reaching significance. This suggests that the left amygdala may be more strongly associated with PTSD-related symptoms than the right amygdala.
Figure 2. Amygdala activity and proximity to the WTC on September 11th, 2001. (A) Fearful emotional faces elicited greater amygdala activity than calm faces in the whole group. Voxel-wise analysis of variance for this contrast indicated that the 9/11-exposed group showed significantly increased bilateral amygdala activation relative to the comparison group. Region of interest analysis showed mean signal change in the (B) left amygdala ($p < 0.01$) and (C) right amygdala ($p < 0.05$) (fearful vs. calm contrast) was higher in the group that was exposed to the disaster on September 11, 2001 than in the comparison group.
was correlated robustly with current symptoms as reported on the IES, as well as with retrospectively reported symptoms of avoidance on the UM-CIDI. Signal change in the right amygdala was less strongly correlated with overall current symptoms on the IES and with retrospective report of avoidance symptoms at time of worst trauma. One outlier was removed for these analyses (Mahalanobis distance = 12.37; Darlington, 1990). These relationships were not driven by data from individuals with a past history of PTSD; control for history of PTSD strengthened the significance of the relationship between BOLD signal and IES score (left amygdala: $\Delta R^2 = .17, p = .007$; right amygdala: $\Delta R^2 = .14, p = .05$) and had no effect on the association between signal and symptoms at worst trauma ($\Delta R^2 = .009$, not significant). With control of 9/11 group status, there was an association between STAI-S and percent signal change in the left amygdala that was significant ($\beta = .35; p = .05$) in this sample, which is in keeping with previous findings (Bishop, Duncan, & Lawrence, 2004).

**Mediation by amygdala activation.** If increased amygdala reactivity drives the association between trauma exposure and increased vulnerability, then it should statistically mediate the relationship between trauma exposure and amygdala activation (fear vs. calm faces). In support of this model, explained nearly half of the variance in current symptoms in this sample; $R^2_{adj} = 0.46$. Appropriate testing (MacKinnon & Dwyer, 1993; Sobel, 1986) found this mediated effect to be significant, $z = 2.80, p < .01$. Thus, amygdala reactivity is a reasonable candidate for a potentially causal mechanism associating 9/11 exposure and symptom presentation (Baron & Kenny, 1986). The next analysis identifies the key parameters of exposure to the 9/11 disaster that were associated with persistent amygdala reactivity and increased symptoms in this sample.

**Years since worst trauma.** Trauma exposure was prevalent in this sample, which provided the opportunity to examine temporal effects in the psychobiological correlates of trauma exposure, regardless of type of trauma. We previously noted that 9/11-exposed participants had significantly fewer years since worst trauma exposure than those in the comparison group (see Table 1). Of the 21 participants with trauma exposure, 11 identified the 9/11 disaster as their worst trauma in lifetime (eight in the group that was closest to the WTC on 9/11/01 and three in the group that was farther away). The remaining 10 identified other lifetime traumas as their worst trauma (three in the group that was closest to the WTC, seven in the group that was farther away). Overall, years since worst trauma ranged from less than one year to nearly 20 years, with the exception of one person who experienced very early abuse (nearly 40 years before scanning). In the whole group, years since worst trauma predicted reductions in current symptoms ($\beta = -0.51, p = .02$), as well as signal in the left ($\beta = -0.46, p = .05$) and right ($\beta = -0.45, p = .06$) amygdala to fearful versus calm faces (outlier excluded; see Figure 3). Slopes of these relationships are similar for the whole group and for the group that identified non-9/11-related traumas as their worst traumas in lifetime (left amygdala: $\beta = -0.49$; right amygdala: $\beta = -0.36$), suggesting that this finding may be generalizable to non-9/11 traumas. These analyses included control for age at scan, which was found to serve as a classical suppressor variable in these models (age was not significantly correlated to years since worst trauma or to amygdala activation but the presence of age increased the multiple correlation of the above regression models, suggesting that age was suppressing some of what would otherwise be error variance in the relationship between years since worst trauma and amygdala activation: Darlington, 1990). The associations between years since worst trauma and all elements of the mediation model discussed above suggest that 9/11 status may be secondary to this factor in the prediction of current symptoms and amygdala activation (fear vs. calm). In support of this, statistical control of years since worst trauma in the relationship between 9/11 group status and current symptoms rendered the contribution of 9/11 status nonsignificant, suggesting that the contribution of 9/11 status in this model was accounted for by variation in years since worst trauma. Similarly, control of this factor in the relationship between 9/11 status and percent signal

### Table 2

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<tr>
<th>Correlation of Percent Signal Change in Right and Left Amygdala (Fearful Versus Calm Contrast) With Behavioral Variables (Pearson r), in the Whole Group</th>
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<tr>
<td><strong>Signal change in left amygdala (r)</strong></td>
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<td>Total number of symptoms at worst trauma</td>
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<td>Number of intrusion symptoms at worst trauma</td>
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<td>Number of avoidance symptoms at worst trauma</td>
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<td>Impact of Events Scale (IES)</td>
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* $p < .05$, ** $p < .01$, † $p < .10$. 
change in left amygdala (fear vs. calm) rendered the contribution
of 9/11 group status less significant ($\beta = 0.64, p = .003$ to $\beta = 0.57, p = .02$), indicating that years since worst trauma accounts
for part of the explained variance in this relationship.

**Trauma intensity.** Trauma intensity is difficult to define and
quantify, except for its immediate impact on the individual.
This relative level of arousal and distress at the time of trauma
is an important predictor of long-term vulnerability to mental
health disorders (Ozer, Best, Lipsey, & Weiss, 2003). For the
purposes of this analysis, we operationally defined intensity by
the participant’s own report of symptoms at worst trauma. In
this sample, the 9/11-exposed participants reported more symp-
toms of avoidance and arousal at time of worst trauma (see
Table 1). Trauma intensity was highly related to current symp-
toms ($\beta = 0.60; p = .005$). Most of the variance in this
relationship was explained by years since worst trauma (its
inclusion with trauma intensity in a model predicting current
symptoms reduced the contribution of trauma intensity to a
trend: $\beta = 0.47; p = .07$). Together, years since worst trauma
and trauma intensity explained a substantial portion of the
variance in the relationship between 9/11 group status and
current symptoms ($R^2 = 0.40$).

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**Figure 3.** Amygdala activity and time since worst trauma. Partial plots for the whole group controlling for age
at scan of (A) mean signal change in left amygdala (fearful vs. calm) versus years since worst trauma in lifetime:
$\beta = -0.46, p = .05$. (B) mean signal change in right amygdala (fearful vs. calm) versus years since worst trauma
in lifetime: $\beta = -0.45, p = .06$. Slopes of these relationships for the group with non-9/11 traumas as worst
traumas are shown for comparison (see p. 232).
Increased symptoms of avoidance at time of worst trauma also predicted increases in percent signal change in the left and right amygdala to fearful versus calm faces (Table 2; Figure 4). While trauma intensity was correlated with years since worst trauma ($r = -0.48; p = .05$), it also contributed significantly to the explained variance of percent signal in the left amygdala (fear vs. calm) beyond that already accounted for by years since worst trauma ($\beta = 0.60, p = .02; \Delta R^2 = 0.23, p = .02$). When intensity of worst trauma was entered with years since worst trauma into a model predicting signal change in the left amygdala from 9/11 status, these two variables fully controlled the contribution of 9/11 group status. This suggests that the amount of time since an individual’s worst trauma and the intensity of that worst trauma accounted for the observed increases in amygdala reactivity (fearful vs. calm) in participants who were close to the WTC on September 11th, relative to those who were farther away.

**Figure 4.** Amygdala activity and trauma intensity. (A) Correlation between the number of retrospectively reported symptoms of avoidance at worst trauma and signal change in left amygdala (fearful vs. calm contrast): $r = 0.58, p < .01$. (B) Correlation between the number of retrospectively reported symptoms of avoidance at worst trauma and signal change in right amygdala (fearful vs. calm contrast): $r = 0.42, p = .06$. 
Discussion

This fMRI study examined the neural correlates of trauma exposure in a sample of healthy adults. We found that participants who were within 1.5 miles of the WTC on September 11th, 2001, had significantly higher mean levels of activation in bilateral amygdala to fearful versus calm faces relative to those who were living more than 200 miles away, despite having no current diagnoses of PTSD, depression, or anxiety disorder. The 9/11-exposed group also had higher levels of current symptoms and reported more symptoms at time of worst trauma. Signal change in the amygdala (fearful vs. calm contrast) fully mediated the relationship between 9/11-exposure and number of current symptoms. In turn, the effect of group status on both amygdala activation (fearful vs. calm faces) and current symptoms was statistically explained by time since worst trauma and intensity of worst trauma, as indicated by reported symptoms at worst trauma. These data are consistent with a model of heightened amygdala reactivity following high-intensity trauma exposure, with relatively slow recovery. Notably, similar results have recently been identified in a similar population (Sharot et al., in press) using a paradigm that evoked specific memories of the disaster of September 11th. The present study supports and extends this research by suggesting that long-term trauma-related modulation of the amygdala is observable using mild, standardized emotional stimuli (fearful vs. calm faces), indicating that these effects may extend further into everyday life than previously thought.

Animal models of exposure to stress may provide insight into the processes of brain plasticity that underlie these results. Traumatic stressors encountered in the natural environment are likely to involve a mixture of conditioned and unconditioned response (Adamec, Blundell, & Burton, 2006). The most analogous animal models are likely to be those involving relatively severe uncontrollable stressors (for reviews, see Adamec et al., 2006; unprotected predator stress; Maier & Watkins, 1998; inescapable shock; McEwen, 2005; acute and chronic restraint stress). Controllable stressors do not activate the amygdala enough to produce even temporally increased reactivity of the amygdala; uncontrollable stressors have, however, produced moderately persistent hyperexcitability of the amygdala and related structures in laboratory animals (Adamec et al., 2005; Maier & Watkins, 1998). This hyperexcitability has been associated with increased fearful response to ambiguous or mild stressors (Adamec et al., 2006; Maier & Watkins, 1998).

At the neuronal level, one type of uncontrollable stressor (chronic restraint stress or CRS) has been shown to produce hypertrophy of the dendritic arborization in the BLA and extended amygdala, accompanied by dendritic atrophy and decreases in spine density in medial prefrontal areas and the hippocampus (Mitra et al., 2005; Vyas et al., 2002). In rats, these changes are associated with standard behavioral indicators of anxiety in rodents, including decreased time spent in the open (exposed) arms of an elevated plus maze and reduction in open arm entries (Mitra et al., 2005; Vyas et al., 2002). Acute (single-event) restraint stress produces similar increases in spine density in the BLA and behavioral indicators of anxiety, with no associated changes in prefrontal areas (Mitra et al., 2005). Consistent with these findings, single-event unprotected predator stress produces persistent increases in evoked potentials from neurons in the BLA, which are also associated with increased open arm entry and decreased in exploration in the elevated plus maze (Adamec, Blundell, & Burton, 2005). Thus, a single noxious event of various types can drive alterations in the amygdala that are associated with persistent increases in behavioral reactivity in situations of uncertainty and potential threat. Notably, although CRS-induced changes to prefrontal areas and hippocampus are reversible with an extended stress-free period, hypertrophy of the amygdala and increases in anxiety-related behaviors are more persistent (Vyas, Pillai, & Chattarji, 2004). The effects of predator stress are also reported to be slow to return to baseline (Ademac et al., 2005). This highlights the role of the amygdala in maintaining stress-induced anxiety-like behaviors, as well as the slow recovery of this system over time after exposure to intensive stressors.

The findings of the present study point to the potential durability of human stress-related neural plasticity, even in adulthood, and suggest that such changes may be driven by the recency and intensity of worst adverse event (worst trauma in lifetime). Although accumulated trauma exposure did not add to the statistical prediction any of the outcomes in this analysis, it is a point for future research whether a more general measure of cumulative environment risk would do so. Research has established the relationship between accumulated environmental risk and negative socioemotional outcomes (e.g., Rutter, 1979; Sameroff, Seifer, Baldwin, & Baldwin, 1993). However, increases in cumulative environmental risk (e.g., low socioeconomic status, minority status, compromised social support, mental illness in family) may be confounded with increased likelihood for high-intensity trauma exposure in most community-based samples, and cumulative risk may also exacerbate trauma outcomes (Galea et al., 2002). More basic research is needed on the neural correlates of stressful life events of all types in healthy individuals; this is an exciting new area for investigation in cognitive neuroscience—one that in the past year has begun to show forward movement (e.g., Cohen et al., 2006; Sharot et al., in press; this study: also see recent work on the effects of in-scanner stressor exposure, Dedovic et al., 2005; Wang et al., 2005).

Finally, amygdala hyperexcitability after trauma may provide a tool for examining the neural processes that modulate stress-related neural plasticity. Chief among these potential modulatory processes is emotion regulation. Regulation of emotions is important for social adaptation, and neuroimaging evidence suggests that regulation of negative emotions, in particular, depends on modulation of amygdala response using prefrontal and anterior cingulate control systems (Ochsner & Gross, 2005). For example, amygdala activation is attenuated in response to fearful or angry emotional faces during cognitive evaluation of the type of emotion (Hariri, Bookheimer, & Mazzotta, 2000). The same occurs during cognitive evaluation of threatening emotional scenes and is accompanied by decline in skin conductance (Hariri, Mattay, Tessitore, Fera, & Weinberger, 2003). Different strategies of emotion regulation (e.g., reappraisal vs. suppression: Gross & John, 2003; John & Gross, 2004) are likely to modulate amygdala hyperexcitability and associated behavioral outcomes following trauma exposure. The latter would be a point of interest for intervention.

Limitations

This study uses a retrospective report of trauma exposure. The difficulties with retrospective reporting include revisionist recall, bias because an ensuing disorder is known to have occurred, bias by respondent’s depression or cognitive impairment, and normal
forgetting (Hardt & Rutter, 2004; Moffitt et al., 2006). Prospective studies of the effects of trauma exposure on the brain would be preferable in many ways. However, this is made difficult because trauma exposure typically occurs unexpectedly. In the absence of a prospective study, it has been argued (Moffitt, Caspi, & Rutter, 2006) that the use of the life history calendar (Caspi et al., 1998) is a highly valid and reliable method for collecting retrospective data on adverse life events. We used this method in the present study to reduce retrospective reporting bias to a minimum.

An additional limitation of this study is that the comparison group in the current sample exhibited a high rate of trauma exposure. This may be due in part to participant self-selection, although high rates of trauma exposure have also been observed in large community-based studies (e.g., Breslau et al., 1998; Young & Breslau, 2004a and 2004b; Young et al., 2004) that used the UM-CIDI measure (Kessler et al., 1994) that was employed in the present study. Because there were few subjects without trauma exposure in this sample, we were unable to test differences between participants with and without trauma exposure. We note, too, that because most of the sample consisted of trauma-exposed individuals without PTSD, this may have inadvertently become a study of resilience; if so, these findings suggest that resilience under stress does not imply lack of physiological consequences following stressor exposure. In addition, specific examination of gender differences was not possible due to the small size of the study, although we found no significant statistical contributions of gender to the results reported here.

A further limitation to this study is that we used a passive viewing paradigm in this “first look” for neural correlates of trauma exposure in adult humans, because passive viewing of emotional faces is one of the most powerful standardized methods for evoking amygdala response (e.g., Whalen et al., 2001) and because tasks that require an active response to emotional stimuli can lead to top-down suppression of the amygdala response (Lange et al., 2003). The limitation of this method is the lack of accompanying behavioral data to determine level of attention to the stimuli. Having established with Sharot et al. that there are robust neural correlates of trauma exposure, the next set of research questions would address the source of the observed differences between groups using more sophisticated paradigms, for example, examination of differences in attention or emotion regulation using more active paradigms.

Finally, in the present research, participants were aware that the study was related to the effects of the 9/11 disaster, and they were interviewed for trauma exposure before scanning. This knowledge, and the screening process, may have had a priming effect on amygdala response. This is an unexplored area and one that requires more research. Because there was trauma exposure in both groups (with the exception of one individual in the comparison group), the trauma inventory might have been expected to have a similar impact across all participants. However, it may be that specific variables related to prescreening interviews (e.g., length and timing of the trauma interview) had significant effects on the BOLD signal response, even years after the trauma. If so, it would suggest even more powerful and subtle long-term effects of previous trauma exposure on the brain than are indicated in the current study.

Conclusions

Close proximity to the WTC on September 11th, 2001, was associated with increased amygdala activation and higher levels of symptoms in healthy adults. Amygdala activation mediated the association between trauma exposure and current symptoms, suggesting a role for amygdala hyperexcitability in the association between trauma exposure and subsequent vulnerability to mental health disorder. Overall, amygdala activation was found to decrease with time since worst trauma, and to increase with the intensity of each individual’s worst-ever trauma, as measured by symptoms at time of trauma. Together, these two variables statistically explained the effect of proximity to the 9/11 disaster on both amygdala activation and current symptoms. This finding suggests that there is heightened amygdala activity from high intensity traumas, and that recovery occurs over many years, even in those without a current clinical disorder.

References


Appendix

Statistical mediation (Baron & Kenny, 1986, p. 1176)

A variable functions as a mediator of the significant relationship between an independent and dependent variable when (a) variation in the levels of the independent variable significantly accounts for variations in the presumed mediator, (b) variations in the presumed mediator significantly account for variations in the dependent variable, and (c) both the independent variable and the presumed mediator are entered simultaneously into a regression model predicting the dependent variable, then the previously significant relationship between the independent variable and the dependent variable is reduced in significance or (in the strongest condition) is no longer significant. MacKinnon & Dwyer (1993) provide a statistical test for significance of this overall process, which was used here.

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