A Genome-Wide Identified Risk Variant for PTSD is a Methylation Quantitative Trait Locus and Confers Decreased Cortical Activation to Fearful Faces

Lynn M. Almli,1 Jennifer S. Stevens,1 Alicia K. Smith,1 Varun Kilaru,1 Qian Meng,2 Janine Flory,3 Duna Abu-Amara,4 Rasha Hammamieh,5 Ruoting Yang,6 Kristina B. Mercer,1 Elizabeth B. Binder,1,7 Bekh Bradley,8,1 Steven Hamilton,9 Marti Jett,5 Rachel Yehuda,3 Charles R. Marmar,4 and Kerry J. Ressler1,10*

1Department of Psychiatry and Behavioral Sciences, Emory University, Atlanta, Georgia
2Department of Psychiatry, University Medical Center, New York, New York
3Mental Health Care Center, James J. Peters Veterans Affairs Medical Center, Bronx, New York/Traumatic Stress Studies Division, New York, New York
4Department of Psychiatry, New York University, Steven and Alexandra Cohen Veterans Center for Posttraumatic Stress and Traumatic Brain Injury, New York, New York
5Integrative Systems Biology, US Army Center for Environmental Health Research, Fort Detrick, Maryland
6Advanced Biomedical Computing Center, Frederick National Laboratory for Cancer Research/SAIC-Frederick Inc., Frederick, Maryland
7Department of Translational Research in Psychiatry, Max Planck Institute of Psychiatry, Munich, Germany
8Department of Veterans Affairs Medical Center, Clinical Psychologist, Mental Health Service Line, Atlanta, Georgia
9Department of Psychiatry, University of California, San Francisco, California
10Howard Hughes Medical Institute, Chevy Chase, Maryland

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Genetic factors appear to be highly relevant to predicting differential risk for the development of post-traumatic stress disorder (PTSD). In a discovery sample, we conducted a genome-wide association study (GWAS) for PTSD using a small military cohort (Systems Biology PTSD Biomarkers Consortium; SBPBC, N = 147) that was designed as a case-controlled sample of highly exposed, recently returning veterans with and without combat-related PTSD. A genome-wide significant single nucleotide polymorphism (SNP), rs717947, at chromosome 4p15 (N = 147, β = 31.34, P = 1.28 × 10^-8) was found to associate with the gold-standard diagnostic measure for PTSD (the Clinician Administered PTSD Scale). We conducted replication and follow-up studies in an external sample, a larger urban community cohort (Grady Trauma Project, GTP, N = 2006), to determine the robustness and putative functionality of this risk variant. In the GTP replication sample, SNP rs717947 associated

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15. SUBJECT TERMS
GWAS; PTSD; fMRI; meQTL; epigenetic

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**Key words:** GWAS; PTSD; fMRI; meQTL; epigenetic

**INTRODUCTION**

Although post-traumatic stress disorder (PTSD) requires the experience of a life-threatening trauma to occur, twin and family studies have suggested that up to 30–40% of the risk for PTSD following significant trauma is genetically heritable (e.g., [True et al., 1993]). Genetic studies to date have been relatively small, but have yielded approximately 15–20 different genes that might associate with PTSD or PTSD risk. These candidate studies are well-discussed in several recent reviews [Pitchman et al., 2012; Almli et al., 2014b]. There have been three published GWAS, at the time of this writing, focused on PTSD [Guffanti et al., 2013; Logue et al., 2013; Xie et al., 2013]. The genome-wide significant findings have yielded variants within or near potentially interesting candidate genes, including RORA, TLL, and COBL, as well as within a long noncoding RNA, lincRNA AC068718.1. The advantage of GWAS is that it is hypothesis neutral, and thus allows the identification of variants not selected for candidate gene/single nucleotide polymorphism (SNP)-based analysis. However, there is a stringent multiple test burden with GWAS, which is necessary to avoid attributing the finding to false discovery.

Since it is difficult to get very large sample sizes within a single cohort, and individual heterogeneity is introduced by them, another approach is to conduct GWAS in highly homogenous, well-characterized samples where exposure levels are controlled in both cases and controls. For instance, a study using very well-matched trauma control cohorts may provide statistical power that is not found in epidemiological samples of patient cases compared to non-patient controls. Alternatively, an extreme phenotype design, where there is enrichment of extreme cases and controls, can lead to increased odds ratios, and improved power to detect associations, especially with quantitative traits [Van Gestel et al., 2000].

An ideal solution from these methods would be a discovery sample in which the cases differ in having extremely severe PTSD compared to those similarly exposed to trauma, but with no PTSD at all.

In this study, we aimed to increase our understanding of genetic risk for PTSD by conducting a GWAS for PTSD in a discovery sample with an extreme phenotype design and cases and controls having similar exposures. Our discovery sample consisted of a deeply phenotyped, well-controlled, similarly exposed, but relatively small military sample (Systems Biology PTSD Biomarkers Consortium; SBPBC) [Yan et al., 2013; Lindqvist et al., 2014; Yehuda et al., 2015]. To validate GWAS findings, we conducted a replication study in a sample consisting of subjects from an independent cohort, the civilian Grady Trauma Project (GTP) [Binder et al., 2008; Almli et al., 2013]. The GTP is a large, primarily African-American highly-traumatized community cohort recruited from the general medical clinics of Grady Memorial Hospital, a publicly funded hospital that serves economically-disadvantaged individuals in Atlanta, Georgia. Using the replication cohort (GTP), we are able to expand our understanding of the GWAS findings through functional genomic and neuroimaging association analyses. Identification and replication of PTSD association across such divergent cohorts, combined with examination of GWAS findings at epigenetic and neural intermediate phenotype levels, has led to a novel and interesting potential PTSD-associated genomic locus.

**METHODS**

**Participants in Discovery Cohort**

The SBPBC cohort utilized for initial GWAS discovery in this report was recruited as part of a larger study that is designed to identify biomarkers for PTSD diagnosis in cross-sectional and longitudinal studies in male and female Iraq and Afghanistan veterans [Yan et al., 2013; Lindqvist et al., 2014; Yehuda et al., 2015]. Participants were recruited at New York University Langone Medical Center and Icahn School of Medicine at Mount Sinai and from New York City Veteran Affairs medical centers at Manhattan and Bronx, as well as other veteran service organizations and the community. All procedures were approved by the Institute Review Boards of NYU School of Medicine and Mount Sinai School of Medicine. Participants gave written informed consent after receiving a complete description of the study. General inclusion criteria include being a US veteran who served in Operation Enduring Freedom in Afghanistan and/or Operation Iraqi Freedom, between the age of 20 and 60 years, being able to understand the protocol and willing to provide written informed consent. Consistent with the goal of having extreme phenotypes, with PTSD being the primary difference between groups, exclusion criteria included loss of consciousness for more than 10 min, neurological disorders, current alcohol and substance abuse disorders, suicidality, lifetime history of any psychiatric disorder with psychotic features, bipolar disorder or obsessive-compulsive disorder.

Doctoral level clinical psychologists conducted structured diagnostic interviews. Criteria for the PTSD group include warzone exposure and related PTSD symptoms of at least three months duration as indexed by the clinician administered PTSD scale (CAPS) [Blake et al., 1995], consisting of the sum of symptom frequency and intensity. The criteria for control group also included warzone exposure, but subjects did not meet CAPS criteria for lifetime or current combat or civilian PTSD. Warzone exposure was defined as exposure to an OEF/OIF (GWOT) Criterion A event and was confirmed by clinical interview. The Structured Clinical Interview for DSM-IV Diagnosis [Ventura et al., 1998] was used to diagnose comorbid disorders and to assess for exclusion criteria. The military version of the PTSD Checklist (PCL-M) [Weathers...
et al., 2013], and the Beck depression index (BDI) [Beck et al., 1961] were also used as self-report questionnaires to assess symptom severity. After clinical screening, subjects who met the diagnosis criteria for PTSD with CAPS score $\geq 40$ were included in the PTSD group and those with CAPS score $< 20$ were included in the control group. These cut-offs were chosen to ensure clinically significant levels of symptomatology in the cases, and to exclude sub-threshold cases in the control group. Because of the nature of warzone trauma and the recruitment feasibility of combat veterans, PTSD subjects in the present study were chronic cases, and both groups had undergone prolonged warzone exposure. Demographic and psychiatric characteristics of the discovery sample are shown in Table I. We used t-tests to determine statistical differences between cases/controls with the quantitative data and chi-squared tests for categorical data.

Genetic Data: Quality Control and Analyses

**Genotyping.** Using DNA extracted from blood samples, genome-wide SNP genotyping of 147 subjects (we note that one individual withdrew from the study) from the SBPBC discovery cohort was conducted using Illumina’s HumanOmniExpress BeadChip. After calls were made with GenomeStudio (Illumina Inc), there were 730,493 individual SNPs with genotype information. We performed quality control on the SNP data using PLINK [Purcell et al., 2007] (Supplementary Fig. 1). We first used crude quality control filters prior to performing principal-component analysis (PCA) to infer axes of ancestry (Supplementary Fig. 2A). We removed 3666 SNPs for low call rates (less than 95%) and 16,786 SNPs with a frequency of less than 0.01. All individuals had call rates of 98% or greater. Prior to PCA, we used PLINK to prune the autosomal only data in windows of 50 base pairs, removing one SNP from each pair of SNPs with $r^2 > 0.05$ to obtain a set of roughly independent SNPs. We further filtered SNPs for analysis with call rates of less than or equal to 98%, minor allele frequencies (MAF) of less than 0.1, and Hardy–Weinberg equilibrium (HWE) $P$-values of less than $1 \times 10^{-5}$ (removing 13,935, 137,997, and 686 more SNPs, respectively). After quality control, there were 147 individuals (130 males and 17 females) that were genotyped on 557,423 SNPs.

### TABLE I. Demographic and Clinical Characteristics of the Discovery Sample {SBPBC Cohort} Shown as Means (SD) or Counts (Percent)

<table>
<thead>
<tr>
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<th>Total (N = 147)</th>
<th>PTSD — (N = 63)</th>
<th>PTSD — (N = 84)</th>
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<tr>
<td><strong>Age</strong></td>
<td>32.8 (8.0)</td>
<td>33.6 (8.1)</td>
<td>32.1 (8.0)</td>
</tr>
<tr>
<td><strong>Gender (% male)</strong></td>
<td>130 88.4%</td>
<td>57 90.5%</td>
<td>73 86.9%</td>
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<tr>
<td><strong>Race/ethnicity</strong></td>
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<tr>
<td>Hispanic</td>
<td>57 (38.8%)</td>
<td>32 (50.8%)</td>
<td>25 (29.8%)</td>
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<tr>
<td>Non-hispanic Asian</td>
<td>6 (4.1%)</td>
<td>0</td>
<td>6 (7.1%)</td>
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<tr>
<td>Non-hispanic black</td>
<td>35 (23.8%)</td>
<td>16 (25.4%)</td>
<td>19 (22.6%)</td>
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<tr>
<td>Non-hispanic white</td>
<td>45 (30.6%)</td>
<td>14 (22.2%)</td>
<td>31 (36.9%)</td>
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<tr>
<td>Non-hispanic other</td>
<td>4 (2.7%)</td>
<td>1 (1.6%)</td>
<td>3 (3.6%)</td>
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<tr>
<td><strong>Education</strong></td>
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<tr>
<td>Less than 12th grade</td>
<td>3 (2.0%)</td>
<td>1 (1.6%)</td>
<td>2 (2.4%)</td>
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<tr>
<td>High school grad or GED</td>
<td>39 (26.5%)</td>
<td>21 (33.3%)</td>
<td>18 (21.4%)</td>
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<td>2 years college, AA degree</td>
<td>47 (32.0%)</td>
<td>22 (34.9%)</td>
<td>25 (29.8%)</td>
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<tr>
<td>4 years college, BA degree</td>
<td>43 (29.3%)</td>
<td>17 (27.0%)</td>
<td>26 (31.0%)</td>
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<td>Masters degree</td>
<td>14 (9.5%)</td>
<td>2 (3.2%)</td>
<td>12 (14.3%)</td>
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<td>Doctoral degree</td>
<td>1 (0.7%)</td>
<td>0 (0%)</td>
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<td><strong>Relationship status</strong></td>
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<td>Single</td>
<td>59 (40.1%)</td>
<td>22 (34.9%)</td>
<td>37 (44.0%)</td>
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<td>Steady relationship or living together</td>
<td>23 (15.6%)</td>
<td>11 (17.5%)</td>
<td>12 (14.3%)</td>
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<td>Married</td>
<td>39 (26.5%)</td>
<td>16 (25.4%)</td>
<td>23 (27.4%)</td>
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<td>Divorced</td>
<td>26 (17.1%)</td>
<td>14 (22.2%)</td>
<td>12 (14.3%)</td>
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<tr>
<td>Early trauma exposurea</td>
<td>5.8 (4.9)</td>
<td>6.9 (5.7)</td>
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<td>PTSD severity (clinician-rated)h</td>
<td>31.7 (34.7)</td>
<td>68.3 (17.0)</td>
<td>3.5 (5.2)</td>
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<td>PTSD severity (self-reported)c</td>
<td>41.0 (20.3)</td>
<td>60.9 (12.5)</td>
<td>26.0 (8.9)</td>
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<td>Depression severityd</td>
<td>13.8 (13.0)</td>
<td>24.7 (11.4)</td>
<td>5.6 (6.3)</td>
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<td>Military experiencee</td>
<td>36.1 (14.2)</td>
<td>52.2 (12.3)</td>
<td>30.2 (9.6)</td>
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<td>Negative life eventsf</td>
<td>10.4 (10.9)</td>
<td>17.8 (12.0)</td>
<td>5.1 (5.9)</td>
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*AA, associate’s degree; BA, bachelor’s degree; GED, general educational development; PTSD, post-traumatic stress disorder.

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*a*ETISR total [Bremner et al., 2000].

*b*CAPS [Blake et al., 1995].

*PCL-M [Weathers et al., 2013].

*BDI [Beck et al., 1961].

*DRRI_D [Vogt et al., 2013].

*Life events scale negative-past 12 months [Sarason et al., 1978].

*P < 0.05.
**Statistical analyses for GWAS.** Using the statistical package PLINK, we regressed the outcome of current CAPS score on allele count assuming an additive model (0, 1, or 2 copies of the risk allele), including sex and the top three principal components of genome-wide data as covariates. We note that we used the first three principal components within the discovery sample to prevent possible model overfitting and loss of power. To verify the robustness of the top SNP, we used logistic regression to test for the association between PTSD diagnosis (using CAPS) and allele count, including the same covariates as above. Given the mixed ancestry of the discovery samples, we also tested significant variants using self-reported race instead of principal components, as a sensitivity analysis.

**Replication Cohort**

**Participants.** The replication cohort consisted of study participants from Grady Memorial Hospital (Atlanta, Georgia) as part of the GTP. As previously shown [Ressler et al., 2011], these participants were adult, primarily female, highly traumatized and impoverished, with high rates of PTSD. To minimize genetic differences due to ancestry, individuals within three standard deviations of the medians of the first and second principal components of self-reported African Americans, were selected for analysis [Almli et al., 2014a].

**Genetic association analysis.** Given that the discovery sample was primarily male, and prior findings of genomic differences in PTSD across the sexes [Ressler et al., 2011; Gillespie et al., 2013], we conducted an analysis stratified by sex. Using the categorical measure of PTSD, we conducted SNP-based association analysis in the GTP samples for the genome-wide significant SNP. Analyses were run parallel to those described in the SBPBC cohort. In males and females separately, we regressed PTSD diagnosis, based on DSM A-D criteria using responses to the modified PTSD Symptom Scale (PSS) [Falsetti et al., 1993], on allele count (0, 1, or 2 risk alleles) as above. Due to the larger sample size, we used the top 10 principal components [Price et al., 2006; Lin and Zhoa, 2009] and chip type (Illumina's HumanOmniExpress or Omni-Quad BeadChip) as covariates.

**Follow-Up Analyses in Replication Cohort**

**Methylation quantitative trait locus analysis.** DNA methylation was assessed as previously described [Mehta et al., 2013]. Briefly, 1 microgram of DNA was bisulfite-treated, and methylation at >485,000 CpG sites was interrogated using the HumanMethylation450 BeadChip (Illumina). Beta values were generated with BeadStudio and were set to missing (no call) if detection p-values exceeded .001. CpGassoc [Barfield et al., 2012] was used to remove samples with probe detection call rates <95% and those with an average intensity value of either <50% of the experiment-wide sample mean or <2,000 arbitrary units (AU). In addition, CpG sites with missing data for >10% of samples were excluded from analysis. Beta Mixture Quantile dilution (BMIQ) was used to normalize each dataset [Teschendorff et al., 2013]. Individual level data from the 11 CpG sites within 1 MB of rs717947 were examined for this study. For each CpG site, methylation is represented as the average of methylated and unmethylated DNA across all cells that make up an individual DNA sample.

Given that rs717947 is located in an intronic region that appears to be epigenetically-regulated based on human postmortem prefrontal cortex methylation data [Maunakea et al., 2010], we identified methylation quantitative trait loci (meQTL) in the GTP cohort by applying the approach described previously [Smith et al., 2014] to the HumanMethylation450 data. Using R (www.R-project.org), the relationship between the proportion of methylation at the CpG site and SNP was examined via linear regression, where methylation was modeled as a linear function of the number of reference alleles (0, 1, or 2) with sex, age and the top three principal components from GWAS as covariates. Thus, an association, if found between genotype and methylation, represents genotype-dependent averages across the entire cohort. We also performed the same analysis in females only, consistent with the strategy described for SNP replication above. We examined 11 CpG sites and adjusted for each as an independent test; thus, $P < 0.0045$ ($0.05/11$ sites) was noted as statistically significant.

**Neuroimaging.** To determine the neural correlates of the PTSD risk allele, we examined fMRI data among a subset of traumatized women drawn from the GTP replication cohort. Participants completed a task designed to engage threat-processing networks, passively viewing static fearful and neutral face stimuli. Fearful and neutral face stimuli were presented in a block design. Trials included a face stimulus presented for 500 ms, followed by a 500 ms presentation of a fixation cross. Subjects were instructed to pay attention to the faces, and did not make any behavioral response, to minimize motion artifacts and neural activation unrelated to processing the visual stimulus. Detailed procedures and data processing methods are described elsewhere [Stevens et al., 2013].

**RESULTS**

The demographic characteristics of the SBPBC subjects are shown in Table I. The SBPBC cohort had an overall mean of 31.7 (SD = 34.7, range 0–136) for PTSD symptom scores; however, because the cohort was designed with extreme phenotypes, symptom severity was much higher in cases [PTSD+: mean = 69.4 (16.9)] than controls [PTSD-: 3.5 (5.2)]. We report that the demographic characteristics were similar between cases and controls ($P > 0.05$). The following psychiatric variables, Early Trauma Exposure, PTSD severity (clinician-rated), PTSD severity (self-reported), Depressive severity, and Negative Life Events, showed differences between cases and controls, with cases endorsing higher symptoms.

**GWAS**

Using the military SBPBC cohort, we conducted a GWAS using a quantitative measure of PTSD symptoms as the outcome with sex, and the top three principal components as covariates. There was no evidence to suggest inflation of test statistics in our GWAS given a lambda of 1.00 (Supplementary Fig. 2B). The GWAS yielded one genome-wide significant SNP, rs717947, at chromosome 4p15 ($N = 147$, $\beta$ (SE) = 31.34 (5.19), $P = 1.28 \times 10^{-8}$) that associated
with current CAPS score (Fig. 1A). SNP rs717947 had a MAF of 0.18, and a HWE p-value of 1 (4 TT/45 TC/98 CC) in the whole sample and 0.18 in controls. Although most of the top SNPs for the GWAS were intragenic, with the exception of Collagen, Type IV, Alpha 2 (COL4A2), several loci are represented 4p15.1, 12q15, 20p12.1, with SNPs in likely linkage disequilibrium (Table II). Sensitivity analysis covarying for self-reported race instead of the top three principal components showed similar results for the GWAS were intragenic, with the exception of Collagen, Type IV, Alpha 2 (COL4A2), several loci are represented 4p15.1, 12q15, 20p12.1, with SNPs in likely linkage disequilibrium (Table II). Sensitivity analysis covarying for self-reported race instead of the top three principal components showed similar results for the association between rs717947 and PTSD symptoms (data not shown). The risk allele (T) carriers consistently show higher CAPS scores compared to individuals with the CC genotype (mean (SD): TT/TC = 52.7 (34.7), CC = 21.5 (29.8)) (Supplementary Fig. 3A). Follow-up analysis of SNP rs717947 showed that this SNP is robust to different measures of PTSD, both categorical and self-reported measures. Using the categorical PTSD diagnosis measure (based on CAPS) as an outcome revealed an odds ratio (OR) of 8.6 (N = 147, 95% confidence interval (CI) = 3.5–21.6, P = 3.84 × 10^-6) (Fig. 2A). Given that the discovery sample was an extreme phenotype design based on CAPS, we verified that self-reported PTSD symptoms (not used for initial sample selection) produced similar results (N = 147, β (SE) = 17.37 (3.10), P = 1.203 × 10^-7). Plots of self-reported PTSD symptoms are similar to symptoms using CAPS (Supplementary Fig. 3B).

Given uncertainty on whether trauma exposure should be considered as a covariate in GWAS of PTSD, we next examined whether the main effect of rs717947 would be significant for CAPS scores after controlling for early childhood trauma exposure or overall negative life events. In both cases, the SNP remained genome-wide significant [Covariate: Early Trauma Inventory, N = 146, β (SE) = 31.5 (5.2), P = 1.13 × 10^-5; Negative life events, N = 146, β (SE) = 24.9 (4.3), P = 3.29 × 10^-8].

**Replication of rs717947**

Using the civilian GTP cohort, we attempted to replicate the association between SNP rs717947 and PTSD diagnosis. Due to our past findings of SNPs differentially associating with PTSD in males and females (e.g., Ressler et al., 2011) in addition to the male majority of the discovery sample, we stratified our analyses by sex. In females, we found that carriers of the risk allele (T) had increased odds of PTSD diagnosis (N=2006, OR=1.25, CI=1.1-1.5, p=0.005) (Fig. 2B); in contrast, no association was found in males (N=862, p=0.37). Follow-up analyses showed that 6 of the 9 SNPs in the regional peak on chromosome 4 identified from the discovery sample also associate with PTSD diagnosis in the same direction in the GTP replication cohort (nominal p<0.1; Supplementary Table I). We also note that in the GTP sample, this SNP, rs717947, was not associated with comorbid disorders, such as depression as measured by BDI (p=0.8), alcohol abuse by AUDIT (p=0.4), or drug abuse by DAST (p=0.6).

**rs717947 is a Methylation QTL**

Although the gene structure and function of this region of chromosome 4 is unclear, bioinformatics analyses suggest that this peak of association is within an intergenic region that appears to be epigenetically-regulated based on postmortem prefrontal cortex methylation data (Supplementary Fig 4). Therefore, we examined whether the GWAS-associated SNP is potentially a functional SNP, based on its association with DNA methylation in a subset of the full GTP cohort with methylation data. Of the 11 CpG sites within 1 MB of the SNP, one probe was found to be significant after multiple test correction (Supplementary Table II). The number of risk alleles of rs717947 predicted the proportion of methylation from probe cg09242288 [N = 157, β (SE) = -0.02 (0.007), P = 0.002] in the overall cohort (Fig. 2C), indicating that the SNP is an meQTL. Though power to detect associations was reduced when the analysis was restricted to females (N = 99), the effect sizes was equivalent β = −0.02. Methylation of cg09242288 did not predict PTSD independent of genotype.
been unclear at the genomic or neural levels. Our data add to this
Furthermore, functional status of the prior GWAS SNPs has
been published, and of those, replication has been limited.
These regions have previously been shown to play key roles
associated with altered medial and dorsolateral prefrontal activa-
tion to fearful faces.
Indeed, the peak of SNPs likely in linkage disequilibrium with the
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We next wished to examine whether this SNP may be associated
with an intermediate neural phenotype related to fear or threat,
using prior data from fMRI with a fearful faces task in a subset of
the civilian GTP cohort. The subset of females with fMRI
data did not appear to significantly differ on the clinical or
demographic variables from the full genetic sample of GTP
females (Supplementary Table III). To determine brain regions
whose response to threat stimuli was influenced by genotype,
we conducted a whole-brain analysis of the response to fearful
relative to neutral pictures (Fig. 3, Supplementary Table IV).
We observed a linear association between the number of risk
alleles and decreasing activation in bilateral dorsolateral prefron-
tal cortex (dlPFC) and dorsomedial prefrontal cortex (dmPFC).
These regions have previously been shown to play key roles
in emotion regulation (e.g., Ochsner et al., 2002), and
decreased dlPFC activation has been linked with PTSD symptoms
[Aupperle et al., 2012; Fani et al., 2012].

**DISCUSSION**

Identified in a small military cohort designed based on extreme
phenotypes, we found a genome-wide significant SNP that asso-
ciates with PTSD that replicates in a much larger civilian commu-
nity cohort. In the discovery sample (SBPBC), the genome-wide
significant SNP, rs717947, at chromosome 4p15 associated with
PTSD symptoms from the CAPS, as well as the PTSD diagnostic
measure from the CAPS. Subsequently, we replicated the top SNP,
as well as several others within the association peak, with PSS-
determined PTSD diagnosis from the much larger traumatized
civilian cohort. Using the GTP replication cohort, our data also
suggest that rs717947 may be functionally significant as the SNP
was found to be an meQTL, and the risk allele of rs717947 was
associated with altered medial and dorsolateral prefrontal activa-
tion to fearful faces.

These findings are notable as very few GWAS of PTSD have yet
been published, and of those, replication has been limited.
Furthermore, functional status of the prior GWAS SNPs has
been unclear at the genomic or neural levels. Our data add to this
literature in identifying a novel candidate region in chromosome
4 that appears to be a region of moderate epigenetic regulation.
Using human tissue derived from postmortem brain, a research
team from the Department of Neurosurgery, UC San Francisco,
investigated DNA methylation at 24 out of 28 million CpG sites
across the haploid genome and found that a majority of cytosine
methylation (5mC) occurs in intronic and intergenic regions
[Mauankea et al., 2010]. In fact, less than 3% of CpG islands
located in 5’ promoter regions are methylated. While a lot of
emphasis has historically been focused on methylation at the 5’
promoter region for its role in regulation of gene expression,
intergenic DNA methylation may also play a key role in gene
regulation, cell-type specificity, and epigenetic response to envi-
ronment. Li et al have recently shown that distinct populations of
neurons have differential methylation at intergenic loci [Li et al.,
2014]. Furthermore, they suggest that methylation differences
within these intergenic regions may be a result of experience-
induced methylation that contributes to the development of
neuropsychiatric disorders. Interestingly, in silico analysis of the
genomic region surrounding rs717947 (200 bp) reveals a
strong signal for DNA methylation (MeDIP) within the
brain, suggesting that this intergenic region can be methylated
(Supplementary Fig. 4). In fact, rs717947 may also be part of a
CpG dinucleotide. Given this scenario, it is reasonable to assume
that if this SNP is indeed a CpG, a T allele, in place of the
ancestral C allele would eliminate the potential for DNA meth-
ylation at this site. This is highly speculative but could be tested
further by methylation specific PCR or pyro sequencing of
bisulfite treated DNA at this particular SNP.

The region at 4p15 may be a distal regulatory region in cis of one
or more of the neighboring genes, *PCDH7, ARAP2,* and* DTHD1*,
or that this region of DNA may harbor yet unknown functions such
as noncoding RNA (ncRNA) or chromatin regulatory regions.
Indeed, the peak of SNPs likely in linkage disequilibrium with the
genome-wide significant SNP on chromosome 4 are in, or very
close to, the ncRNA BC036345 (chr4: 33,897,961–34,041,515) of
unknown function. However, further work needs to be performed
to identify the effect of the ncRNA within this region. Related to its
possible roles in these neighboring genes, of note *PCDH7* (proto-
(cadherin 7) is expressed within the brain [Hertel et al., 2012], and it has recently been associated with sleep regulation, epilepsy, and neurocognitive function (e.g., Ollila et al., 2014). ARAP2 is also expressed in cortex and hippocampus, and appears to be involved in cellular cytoskeletal dynamics and endosome regulation (e.g., Chen et al., 2013). Less is known about DTHD1 functioning, but it has been associated with retinal dystrophy [Abu-Safieh et al., 2013]. Thus all of the genes surrounding this SNP have potentially interesting characteristics as candidates expressed within the central nervous system (CNS) and putatively associated with other forms of CNS pathology.

To further examine whether there were any associations between rs717947 and intermediate phenotypes associated with PTSD, we examined brain activation during a fear and threat processing task—passive viewing of fearful faces, which we and others have previously associated with differential responses to PTSD and PTSD-related genetic pathways [Andero et al., 2013; Stevens et al., 2013; Stevens et al., 2014]. We found that when confronted with threat stimuli, individuals carrying the rs717947 risk allele showed decreased activation among prefrontal regions that regulate negative emotion (dLPFC and dmPFC). Dysregulation of negative emotion, whether related to inability to extinguish fear, hypervigilance to threatening cues, or generalization of fearful stimuli, are all important intermediate phenotypes related to PTSD. These data suggest that SNPs within this region of chromosome 4 associated with PTSD and differential genomic methylation, are also associated with differential intermediate fear-related neural phenotypes.

There are several limitations of this study. The sample size of the discovery cohort (SBPBC) was very small; however, we believe that the extreme phenotype design and well-controlled nature of the study enhance its power to determine genetic differences between cases and controls. While the extreme phenotype design may yield interesting information, traditional modeling of the phenotype is challenging. Follow-up studies using all continuous values of this phenotype are essential for confirmation of rs717947 as a risk variant for PTSD. We have used alternative PTSD phenotypes, not based on extreme phenotype thresholds, to assess the robustness of the association. The cohort is also of mixed race/ethnicity. Although we used principal components to control for ancestry differences in the samples, it is possible that the susceptibility for disease in risk allele carriers is actually driven by subtle differences in the genetic backgrounds of the different races. Additionally, the discovery and replication cohorts are quite different, particularly in sex and trauma type. Finally, almost half of the cases in the discovery cohort had comorbid depression, a level of comorbidity common across PTSD patients and seen within our civilian cohort as well. Further research is necessary to determine whether SNP rs717947 is specific to PTSD or to comorbid PTSD and depression.

In summary, we have identified a novel genomic locus associated with PTSD symptoms and diagnosis across two very different cohorts—one military, primarily of mixed race and male, and one civilian, African American, and primarily female. The observation will need further replication across other cohorts. However, the combination of significance across mixed samples, association with differential regional DNA methylation, and

**FIG. 2. Replication of Genome-wide PTSD-associated SNP and Evidence for an meQTL.** Replication results for associations of genome-wide significant SNP, rs717947, with PTSD diagnosis in the (A) discovery cohort (N = 147, OR = 8.6 CI = 3.5–21.6, \(P = 3.84 \times 10^{-5}\)) and (B) replication cohort (females only, N = 2006, OR = 1.25, CI = 1.1–1.5, \(P = 0.005\)). (C) Of the 11 CpG within 1 MB of the genome-wide SNP, rs717947 affects expression levels (probe cg09242288) with the risk genotype (TT) showing decreased proportion of methylation (N = 157, \(\beta\) (SE) = −0.02 (0.007), \(P = 0.002\)).
association with neural intermediate phenotypes related to PTSD are all compelling, and suggest this region as a potentially important genomic region for further study. Further genome wide association studies for PTSD, such as those currently planned within the Psychiatric Genomics Consortium PTSD workgroup [Koenen et al., 2013], will continue to dissect the genomic architecture underlying PTSD, with hope for new biomarkers and therapeutic targets.

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