Differential Immune System DNA Methylation and Cytokine Regulation in Post-Traumatic Stress Disorder

Alicia K. Smith¹, Karen N. Conneely², Varun Kilaru¹, Kristina B. Mercer³, Tamara E. Weiss¹, Bekh Bradley⁴, Yilang Tang², Charles F. Gillespie¹, Joseph F. Cubells¹,², and Kerry J. Ressler¹,³,⁵,*

¹Department of Psychiatry & Behavioral Sciences, Emory University School of Medicine, Atlanta, Georgia ²Department of Human Genetics, Emory University School of Medicine, Atlanta, Georgia ³Howard Hughes Medical Institute, Maryland ⁴Atlanta VA Medical Center, Decatur, Georgia ⁵Yerkes National Primate Research Center, Atlanta, Georgia

Abstract

DNA methylation may mediate persistent changes in gene function following chronic stress. To examine this hypothesis, we evaluated African American subjects matched by age and sex, and stratified into four groups by post-traumatic stress disorder (PTSD) diagnosis and history of child abuse. Total Life Stress (TLS) was also assessed in all subjects. We evaluated DNA extracted from peripheral blood using the HumanMethylation27 BeadChip and analyzed both global and site-specific methylation. Methylation levels were examined for association with PTSD, child abuse history, and TLS using a linear mixed model adjusted for age, sex, and chip effects. Global methylation was increased in subjects with PTSD. CpG sites in five genes (TPR, CLEC9A, APC5, ANXA2, and TLR8) were differentially methylated in subjects with PTSD. Additionally, a CpG site in NPFFR2 was associated with TLS after adjustment for multiple testing. Notably, many of these genes have been previously associated with inflammation. Given these results and reports of immune dysregulation associated with trauma history, we compared plasma cytokine levels in these subjects and found IL4, IL2, and TNFα levels associated with PTSD, child abuse, and TLS. Together, these results suggest that psychosocial stress may alter global and gene-specific DNA methylation patterns potentially associated with peripheral immune dysregulation. Our results suggest the need for further research on the role of DNA methylation in stress-related illnesses.

Keywords

PTSD; epigenetic; total life stress; TPR; APC5; TLR8; NPFFR2

INTRODUCTION

Post-traumatic stress disorder (PTSD) is a debilitating, stress-related psychiatric disorder, with US prevalence rates of 7–8% [Kessler et al., 1994; Kessler, 2000]. Cumulative exposure to traumatic or adverse events across the lifespan is an identified risk factor for developing PTSD, and studies suggest a dose–response relationship between the level of lifetime trauma exposure and development of PTSD [Breslau et al., 1999; Brewin et al.,...
2000; Ozer et al., 2003; Binder et al., 2004]. This relationship is especially relevant if the trauma was experienced during childhood since child abuse increases the risk for developing PTSD in adulthood [Duncan et al., 1996; Kessler et al., 1997].

Exposure to child abuse has been associated with increased physiological reactivity to stress [Weiss et al., 1999; Heim et al., 2001], which results in dysregulation of the hypothalamic–pituitary–adrenal (HPA) axis that persists into adulthood. These observations are consistent with both human and animal studies that highlight the mechanisms by which early maternal care experience can permanently alter HPA axis activity through DNA methylation of the glucocorticoid receptor promoter [Francis et al., 1999; Weaver et al., 2004]. Additionally, early maltreatment of rat pups increases methylation and decreases expression of BDNF [Roth et al., 2009]. Similar to these preclinical studies, the glucocorticoid receptor promoter in DNA extracted from umbilical cord blood from children born to depressed mothers was reported to be differentially methylated in children, depending on their cortisol levels and cortisol reactivity to stress measured at 3 months of age [Oberlander et al., 2008].

In adults, Uddin et al. [2010] reported associations between PTSD and methylation patterns from DNA extracted from peripheral blood, noting that differentially methylated CpG sites were over-represented in genes related to immune function and inflammation. They then observed in the same subjects that PTSD associates with a greater immune response to cytomegalovirus infection. Because the HPA axis interacts with the immune system to maintain stress-related allostasis, dysregulation of the HPA axis may result in excessive inflammation [Gill et al., 2009]. Subjects with PTSD have been reported to have increased levels of circulating inflammatory cytokines including IL-6 [Maes et al., 1999], TNFα [von Kanel et al., 2007], IL1B [Spivak et al., 1997], and lower anti-inflammatory cytokines such as IL4 [Kawamura et al., 2001; von Kanel et al., 2007] than individuals without PTSD. Further, case–control studies of PTSD have been shown to have distinct expression patterns in genes involved in immune activation in peripheral blood cells [Segman et al., 2005; Zieker et al., 2007].

This convergent data suggest that epigenetic alterations may contribute to the inflammatory and immune dysregulation observed in subjects with PTSD. The current study examines genome-wide methylation in order to confirm and extend previous studies. We hypothesize that DNA methylation will differ in subjects with PTSD. We further hypothesize that childhood trauma or high levels of cumulative stress over the lifetime will also result in independent changes in DNA methylation. Finally, we hypothesize that plasma cytokine levels will associate with PTSD as well as history of childhood trauma and lifetime stress in these subjects.

**MATERIALS AND METHODS**

**Subject Recruitment**

We evaluated African American subjects recruited as part of a larger study investigating the influence of genetic and environmental factors on response to stressful life events in a predominantly African-American, urban population of low socioeconomic status (SES) [Binder et al., 2008; Bradley et al., 2008; Gillespie et al., 2009]. Briefly, research participants were approached in the waiting rooms of the primary care clinic or obstetrical-gynecological clinic of a large urban, public hospital while either waiting for their medical appointments or while waiting with others who were scheduled for medical appointments. Subjects willing to participate provided written informed consent and participated in a verbal interview.
In this study, we assessed methylation in 110 subjects selected from the cohort described above such that they could be stratified into four categories: PTSD diagnosis (Clinician-Administered Post-traumatic Stress Disorder Scale; CAPS) [Blake et al., 1995] with (N = 25) and without (N = 25) a history of childhood trauma (Childhood Trauma Questionnaire; CTQ) and controls with (N = 26) and without (N = 34) childhood trauma. All subjects were matched for age and sex. All procedures in this study were approved by the Institutional Review Boards of Emory University School of Medicine and Grady Memorial Hospital.

**Subject Assessments**

Demographic information including subject age, sex, and race was provided on a self-administered form. The CAPS [Blake et al., 1995] is an interviewer-administered diagnostic instrument measuring PTSD and has been demonstrated to have excellent psychometric properties [Weathers et al., 2001] The CAPS provides a diagnostic measure of PTSD and assesses lifetime and current PTSD. Subjects were scored as having PTSD if they met DSM-IV PTSD criteria for current PTSD from the CAPS interview.

The CTQ [Bernstein et al., 1994; Bernstein 1998; Bernstein et al., 2003] is a 28-item, self-report inventory assessing three types of child abuse: sexual, physical, and emotional. Cutoff scores for each category have shown excellent sensitivity and specificity in correctly classifying cases of abuse in psychiatric patients. The CTQ yields a total score and subscale scores for each of the three types of child abuse. Bernstein and Fink [Bernstein et al., 1994; Bernstein 1998] established scores for mild, moderate, and severe for each type of abuse. In this study, the data from the CTQ were used to classify participants with physical, sexual, and/or emotional abuse into two categories: (1) Those with CTQ scale scores in the none to mild range, and (2) those with scores in the moderate to severe range.

The Stressful Events Questionnaire (SEQ) is a 39-item instrument assessing recent and lifetime Total Life Stress (TLS) via exposure to stressful life events. The SEQ was developed for the purposes of the current study and covers stressful events ranging from divorce, unemployment, crime, financial, and interpersonal stressors to knowing someone who was murdered. The SEQ collects data on whether participants have experienced these events in the last year or ever (coded as no = 0 to yes = 1). These events are summed to yield a total score reflecting the number of types of stressful life events experienced over the last year or prior to the last year (lifetime), revealing a continuous score of 0 (none) to 39 (most severe). In our larger study of over 3,000 participants, lifetime stressful events are normally distributed (mean = 14.96 ± 6.64). Supplementary Table 1 lists the 39 items making up the SEQ, and the clinical and demographic characteristics of subjects whose DNA samples passed QC (as described below) are detailed in Supplementary Table 2.

**DNA Methylation**

DNA was extracted from whole blood at the Emory University Biomarker Service Core with a KingFisher Flex Magnetic Particle Processor (ThermoFisher Scientifics) and the Mag-Bind SQ Blood DNA kit (Omega Biotek, Norcross, GA) or with a Qiagen M48 BioRobot Workstation using the MagAttract DNA Blood M48 kit (Qiagen, Valencia, CA). DNA concentration was determined by PicoGreen quantitation using the Quant-iT dsDNA Assay Kit (Invitrogen, Carlsbad, CA) on a SpectraMax Gemini XPS plate reader (Molecular Devices, Sunnyvale, CA). DNA concentration was determined by PicoGreen quantitation using the Quant-iT dsDNA Assay Kit (Invitrogen, Carlsbad, CA) on a SpectraMax Gemini XPS plate reader (Molecular Devices, Sunnyvale, CA). Samples were resolved on a 1% agarose gel to verify that the DNA was of high molecular weight (at least 2 kb). One microgram DNA was bisulfite-treated for cytosine to thymine conversion using the EZ DNA Methylation-Gold kit (Zymo Research, Orange, CA). The DNA was then whole-genome amplified, fragmented, and hybridized to the HumanMethylation27 BeadChip (Illumina, San Diego, CA). The XStain was performed on a Tecan Evo 150 liquid handling robot. The BeadChips were scanned...
using a BeadStation 500GX, and the methylation level (beta value) calculated for each queried CpG locus using the Methylation Module of BeadStudio software.

Samples with probe detection call rates <90% were excluded from further analysis as were those with an average intensity value of either <50% of the experiment-wide sample mean or <2,000 arbitrary units (AU). Hierarchical clustering was then performed to identify extreme outliers and global trends in methylation. One sample of pooled female DNA was included on each BeadChip as a technical control throughout the experiment and assessed for reproducibility using a Pearson R$^2$ coefficient. For each individual sample and CpG site, the signals from methylated (M) and unmethylated (U) bead types are used to calculate a β value: β = M/U + M + 100, which can be treated as an approximation of the proportion of CpG dinucleotides methylated at a particular site. Prior to computing β values, we performed quantile normalization on the signal data to adjust for any technical variability between samples.

**Statistical analysis**

We worked with the log ratio of β-values log ($\beta/1-\beta$), which is the same as the log signal ratio log(M/U) for quantile-normalized values. To identify CpG sites for which methylation varied significantly with each outcome (PTSD, child abuse, or TLS), we fit a separate linear mixed effects model for each CpG site. For each CpG site, we regressed log ($\beta/1-\beta$) on the variable of interest (PTSD diagnosis, child abuse, or TLS), as well as sex and age. We included chip-specific random effects to allow for potential batch effects due to chip in measurement of the proportion of DNA methylated [Leek et al., 2010]. We fit the above model for all 27,578 CpG sites. To define an initial set of CpG sites for which methylation varied significantly with each outcome, we applied a false discovery rate (FDR) cutoff of 0.05 using the method described by Storey [2002]. To examine global methylation, we also fit the above model with a measure of average methylation across all 27,578 CpG sites as the dependent variable log ($\beta\overline{\beta}/1-\beta$), where $\beta$ represents the average β value over all 27,578 sites for an individual.

**Cytokine Measures**

Fasting blood was drawn into EDTA tubes between 8 and 9 AM by trained phlebotomists and then placed immediately on ice. Plasma was separated from blood cells via centrifugation, aliquoted into 1 ml aliquots and frozen at −80°C until thawed for cytokine analysis. Both pro-inflammatory (IL6, IFNa, IL1b, IL2, and TNF-α) and anti-inflammatory (IL4 and IL10) cytokines were measured in plasma using a multiplex ELISA (R&D Systems) according to the manufactures’ instructions. Each sample was measured in duplicate and the results were averaged and log transformed prior to analysis.

Because of sample availability, 51 subjects had samples available for both DNA methylation and plasma cytokines. Because this may limit power to detect associations, plasma cytokine levels were also evaluated in an additional 126 subjects that did not have methylation data. The potential demographic and clinical confounding factors (age, sex, years of education, employment, disability, past history of substance abuse, current substance abuse, and past suicide attempt(s)) were not different between the groups in which cytokines were measured or the 110 subjects in methylation dataset. We then examined the association between these potential confounders and each outcome (PTSD, child abuse, or TLS) by applying a general linear model. To avoid adjusting for collinear variables, the associated factors were then entered into a forward and reverse stepwise regression, and only those factors that remained associated with the outcome at $P < .05$ were considered as covariates in the final analysis model. Because of independent associations ($P < 0.05$), we adjusted the analysis of PTSD for history of suicide attempts and the analysis of TLS for history of substance abuse.
RESULTS

Global Changes in DNA Methylation

We noted an increase in global methylation in subjects with PTSD compared to controls ($T = 3.95; P = 1.5 \times 10^{-4}$; Fig. 1). There was no change in global methylation levels in subjects with a history of childhood trauma ($T = -0.17; P = 0.87$) or with increased total life stress ($T = -0.29; P = 0.77$).

Gene-Specific Changes in DNA Methylation

We examined the association between each CpG site and PTSD diagnosis (Supplementary Table 3), child abuse (Supplementary Table 4), and TLS (Supplementary Table 5). CpG sites in five genes met experiment-wide criteria (FDR < 0.05) for association with PTSD diagnosis. CpG sites in translocated promoter region (TPR; $T = -5.08; P = 1.9 \times 10^{-6}$) and annexin A2 (ANXA2; $T = -4.69; P = 9.3 \times 10^{-6}$) demonstrated decreased methylation while CpG sites in C-type lectin domain family 9, member A (CLEC9A; $T = 4.89; P = 4.3 \times 10^{-6}$), acid phosphatase 5, tartrate resistant (ACP5; $T = 4.73; P = 8.0 \times 10^{-6}$), and toll-like receptor 8 (TLR8; $T = 4.65; P = 1.1 \times 10^{-5}$) showed increased methylation in PTSD subjects compared to controls (Supplementary Table 1; Fig. 2). Additionally, one CpG site met criteria for experiment-wide significant association with TLS (Supplementary Table 3; Fig. 2). This site, cg03017946, is located near neuropeptide FF receptor 2 (NPFFR2), and its DNA methylation was inversely associated with TLS scores ($T = -5.33, P = 6.6 \times 10^{-7}$). No CpG site reached experiment-wide criteria for significance for association with child abuse.

Replication of Differentially Methylated Genes from the Literature

To complement our hypothesis-neutral approach, we evaluated the association between CpG sites in genes that had been previously associated with PTSD, early life stress, or child abuse in the literature [McGowan et al., 2009; Roth et al., 2009; Beach et al., 2010; Uddin et al., 2010] and PTSD diagnosis, child abuse, and TLS in our population. In examining previously reported CpG sites in genes related to immune function and inflammation, we focused on the CpG sites identified by a gene ontology analysis of PTSD subjects [Uddin et al., 2010]. The results of the literature replication are presented in Table I. Of the 60 genes examined, 19 were differentially methylated with respect to PTSD diagnosis in these subjects. Interestingly, while none were associated with both PTSD and child abuse, CpG sites in both BDNF and CXCL1 were associated with both PTSD diagnosis and TLS. We did not observe differential methylation of CpG sites in NR3C1 or SLC6A4 with any outcome.

Plasma Cytokine Measures

Because a number of genes related to inflammation were differentially methylated with numerous stress-related outcomes, and because of numerous reports of immune system dysregulation in subjects with trauma history, we measured pro-inflammatory (IL6, IFNa, IL1b, IL2, and TNF$\alpha$) and anti-inflammatory (IL4 and IL10) cytokine levels in a subset of these subjects for which plasma was available (N = 51) using a multiplex ELISA. The association between these immune response markers and PTSD, child abuse, and TLS were evaluated (Table II). Interestingly, IL4 was decreased in subjects with PTSD. Plasma IL2 and TNF$\alpha$ levels were increased in subjects with a history of child abuse. Consistently, subjects with higher TLS scores also have higher TNF$\alpha$ levels.

We then investigated the association of these three cytokines (IL4, IL2, and TNF$\alpha$) in a larger sample of subjects that included the original 51 subjects and an additional 126 independent subjects (total N = 177; Table II). Decreased plasma IL4 levels remained associated with PTSD in this larger group of subjects when adjusted for previous history of suicide attempts. Also in these subjects, increased plasma TNF$\alpha$ levels are associated with a
history of child abuse and with higher TLS scores when adjusted for past history of substance abuse. However, the association between history of child abuse and IL2 levels did not replicate in the larger sample.

**DISCUSSION**

When examining individual CpG sites, thousands were nominally associated with PTSD diagnosis and symptoms, child abuse, and TLS (Supplementary Tables 3–5). Many of these associations were consistent with previous reports from the literature. For example, Roth et al. [2009] reported that early maltreatment of rat pups produced increased methylation of BDNF that resulted in persistent decreases in gene expression in the adult prefrontal cortex. In this study we observed a nominal association between increased methylation at a CpG site (cg27351358) in BDNF and current PTSD. A recent study [Uddin et al., 2010] identified a number of immune-related genes that were differentially methylated in subjects with PTSD; we also observed evidence of association between CpG sites in 19 of these genes with PTSD, 6 with child abuse, and an additional 5 with TLS. However, CpG sites in 34 genes were not associated with any outcome examined. The failure of more of these CpG sites to replicate may be due to a number of differences in how this study was conducted. For example, we controlled for factors that have been previously shown to alter DNA methylation including age, race, and early life stress [McGowan et al., 2009; Rakyan et al., 2010; Teschendorff et al., 2010; Adkins et al., 2011], and accounted for potential batch effects due to chip [Leek et al., 2010]. Another key difference is that we examined CpG sites in which the % methylation varied from 21–79%. As such, we evaluated a greater number of CpG sites but also set a more stringent threshold for statistical significance.

We further identified five additional genes that may be differentially methylated in PTSD. For example, TPR, which has lower levels of methylation in subjects with PTSD, interacts with the stress-induced transcription factor heat shock transcription factor 1 (HSF1) to upregulate HSP70 [Skaggs et al., 2007], which facilitates folding of the glucocorticoid receptor (NR3C1) [Dittmar et al., 1997]. Additionally, genes related to immune function were also identified. Specifically, TLR8 is expressed in the white matter of periventricular, subcortical, and cerebellar regions of the brain and has been implicated in neurogenesis and neurite outgrowth in the developing brain [Mishra et al., 2006; Mallard et al., 2009]. It remains expressed in adults and participates in inflammatory processes along with suppression of neurite outgrowth and induction of neuronal apoptosis [Ma et al., 2007]. Also, ACP5, also known as TRAP, is expressed in the brain [Sun et al., 2008] and is stimulated by monocyte chemotactic protein 1 (MCP1), which is produced during neuroinflammation and associated with neuronal death [Kim et al., 2006]. MCP1 increases the excitability of nociceptive neurons and is increased in rodent models of neuropathic pain [White et al., 2009].

A CpG site in NPFFR2 met criteria for experiment-wide significance. NPFF is a neurotransmitter that modulates the opioid system, which has been heavily implicated in PTSD [Holbrook et al., 2010]. Additionally, NPFFR2 is activated in response to inflammatory pain [Yang and Iadarola 2003; Yang et al., 2008; Lameh et al., 2010]. PTSD and other stress-related disorders are associated with increased reports of pain and pain-related disability, as well as increased healthcare utilization and lower quality of life [Bryant et al., 1999; Shipherd et al., 2007] in veteran populations [Asmundson et al., 2004]. We have reported a similar relationship between PTSD and chronic pain in civilian populations [Schwartz et al., 2006]. Also in civilians, PTSD has been shown to be a risk factor for headache chronicity [Peterlin et al., 2008]. Thus, PTSD appears to associate with abnormal regulation of pain responses. We hypothesize that stress-related methylation of NPFFR2
contributes to dysregulation of pain, and to the abnormal activation of inflammatory pathways in chronic stress conditions [Bauer et al., 2010].

While we observed an overall increase in global methylation, CpG sites in TPR and ANXA2 demonstrated decreased methylation. Since DNA methylation is often inversely correlated with gene expression, these data suggest that while transcription is generally reduced overall in peripheral blood, these two genes as well as NPFFR2 may be activated in subjects with PTSD or significant TLS and warrant further investigation.

To determine whether changes in DNA methylation could also be associated with immune dysregulation, we evaluated both pro-inflammatory and anti-inflammatory cytokines in plasma. In two groups of subjects, we observed an association between PTSD and lower IL4 levels, consistent with previous reports [Kawamura et al., 2001; von Kanel et al., 2007]. We also observed increased plasma TNFα levels associated with both child abuse and higher TLS scores. TNFα crosses the blood–brain barrier to stimulate the HPA axis, resulting in increased production of cortisol and symptoms including malaise, fatigue, and changes in sleep and appetite [Raison and Miller 2003; Silverman et al., 2005; Dunn et al., 2006; Sternberg 2006].

Our study only examined DNA extracted from peripheral blood. While this is appropriate for assessing differential methylation of immune-related genes, it is not clear to what degree methylation changes in the blood correlate with the brain. Stress-related outcomes such as PTSD, child abuse, or TLS should also be assessed in brain tissue through animal or post-mortem studies. Interestingly, NPFFR2, which is expressed in hypothalamus, amygdala, forebrain, and brainstem [Goncharuk and Jhamandas, 2008], has also been identified in primary diffuse large B-cell lymphoma [Sung et al., 2011] suggesting that the peripheral blood may be a reasonable proxy for some genes expressed in the brain. Finally, in order to present a complete picture of the factors that may affect DNA methylation in this population, three phenotypes were examined, potentially increasing the possibility of type 1 error.

In conclusion, this study provides evidence suggesting that psychosocial stress may alter global and gene-specific DNA methylation patterns associated with peripheral immune dysregulation and is consistent with previous reports. Further studies are necessary to evaluate the potential role of DNA methylation in chronic activation of the immune system in response to stress-related illness.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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References


FIG. 1.
Global methylation changes are associated with PTSD diagnosis. Box plots of the average methylation level ($\beta$-values averaged across all 27,578 CpG sites) for patients with PTSD (N = 50) and control subjects without PTSD (N = 60).
FIG. 2.
CpG sites in TPR, CLEC9A, ACP5, ANXA2, and TLR8 are differentially methylated in subjects with PTSD, and NPFFR2 methylation is associated with Total Life Stress. The methylation level (β-value, box plot of average methylation value) for each subject is plotted on the vertical axis while PTSD diagnosis is plotted on the horizontal axis. Box plots of methylation values in PTSD cases (N = 50) and controls (N = 60) are shown for TPR (A), CLEC9A (B), ACP5 (C), ANXA2 (D), and TLR8 (E). F: Total life stress (TLS) scores for all subjects (N = 110) are plotted on the horizontal axis and NPFFR2 methylation β values are presented as a scatter plot on the vertical axis. All six CpG sites met experiment-wide criteria for significance (FDR < 0.05).
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<td><strong>P</strong>-value</td>
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<td>Uddin et al. [2010]</td>
<td>2.24</td>
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<td>Uddin et al. [2010]</td>
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<td>SLAMF7</td>
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<td>Uddin et al. [2010]</td>
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NS indicates genes that were not significant with respect to the outcome at $P < 0.05$. Additionally, CpG sites in ATP1A1, BBS7, CD1D, CSNK1D, CYP2J2, DGAT2L6, GALR2, HOXD10, IL8, ISLR2, LST1, LTA4H, MRH1, MTPN, NALP2, NR3C1, NUMB, PTGIS, PTPN22, PYDC1, RIF1, SCN5A, SLC6A4, SOSTDC1, SRXN1, STAP1, STC1, TREM1, TFPI, TLR3, TNC, and TRAP1 were not associated at $P < 0.05$ with any outcome examined.
### TABLE II

Association of Plasma Cytokine Levels With PTSD, Child Abuse, and TLS.

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