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Parenting stress and DNA methylation among African Americans in the InterGEN Study

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Introduction. General life stress has been associated with altered DNA methylation in individuals of African Ancestry, although the relationship between parenting stress and DNA methylation has not been described. The purpose of this study was to examine the relationship between maternal parenting stress and DNA methylation among African Ancestry mother-child dyads.

Methods. We evaluated epigenome-wide DNA methylation relative to parenting stress in 74 mother-child dyads using linear mixed models.

Results. Significant variation in maternal DNA methylation at 95 CpG sites was associated with level of parenting stress. Notably, we identified a change in DNA methylation associated with poly (ADP-ribose) polymerase-1, which plays a key role in stress signaling. We did not identify any significant variation in child DNA methylation related to maternal parenting stress.

Conclusions. However, DNA methylation patterns observed in children mirrored patterns observed in their mothers. The results suggest that differential maternal DNA methylation is associated with higher levels of parenting stress.

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socioeconomic status in childhood [7]. Stress related to finances has also been associated with increased DNA methylation at sites associated with accelerated epigenetic aging in AA women [8]. One recent study evaluated DNA methylation in adults relative to their perception of parenting and maternal stress during 3 developmental periods from their upbringing [9]. The authors did not find any significant correlation between maternal stress and DNA methylation in adult children, although they did identify statistically significant associations between offspring’s perception of parenting from middle childhood to adulthood and the DNA methylation in offspring’s adult genomes. The authors also reported stronger associations with indicators of the child’s perceived parenting than with maternal reports of maternal parenting stress, suggesting that changes in perceived parental rejection have a stronger association with the offspring’s adult epigenome than measures of maternal parenting stress.

There has been extensive research investigating DNA methylation in children exposed to maternal stress in utero [10], although there is a gap in the literature evaluating the relationships among parenting stress and the epigenome of the mother or child. Schechter et al. evaluated DNA methylation at the nuclear receptor subfamily 3 group C member 1 (NR3C1) gene, a gene that encodes glucocorticoid receptor, related to post-traumatic stress disorder (PTSD) and parenting stress in mothers of young children (12–42 months old) [11]. The authors suggest PTSD and parenting stress were negatively correlated with NR3C1 methylation, which could be associated with altered hypothalamic-pituitary-adrenal axis responsiveness. However, the authors noted that the study was not sufficiently powered to differentiate if the loss of DNA methylation at NR3C1 was associated with PTSD or higher levels of parenting stress independent of each other. Combined, the studies completed by Schechter and Naumova’s research groups suggest there may be differential DNA methylation in mothers and children associated specifically with parenting stress of the mother. Therefore, the purpose of this study was to examine the relationship among maternal parenting stress and epigenome-wide DNA methylation in AA mothers and their children.

Materials and Methods

Data for this study were collected from our ongoing study: The Intergenerational Impact of Genetic and Psychological Factors on Blood Pressure (InterGEN) [12, 13]. This sample contains 74 mother-child dyads (n=148) involved in our current prospective study examining the intergenerational impact of genetic and psychological factors on blood pressure. Extensive details on recruitment and statistical methods have been described elsewhere [12, 13]. In brief, after approval from the institutional review board was obtained, mothers were recruited from the community who had a child between the ages of 3 through 5 years of age and would be followed for 2 years to evaluate factors that may contribute to variations in blood pressure in mothers and children. Baseline demographic data and the Parenting Stress Index—Short Form (PSI-SF) [14] was obtained via audio self-assisted interviewing software. Saliva samples for analyses with the Illumina Infinium Methylation EPIC (850 K) BeadChip [15] were collected from the mother and child at the time of the interview. After 2 years of data collection, 81 mother-child dyads have completed baseline interviews and provided saliva for DNA analysis. Six mothers did not complete the PSI during the inter-Phase 1 and could not provide saliva at the time of the interview for DNA analysis; these dyads were excluded from these analyses (n=148 complete cases). Four mothers had 1 missing value from a PSI subscale. The mean for the subscale of the missing item for each individual was imputed and used for analyses per PSI protocol manual [14]. Quality control and preprocessing analytic pipeline of 850 K DNAm has been described elsewhere [12]. All individual samples passed the quality control procedures. In the epigenome-wide association study (EWAS), a total of 847,155 autosomal CpG sites were included. After merging with phenotypic data, 74 African American mothers and 74 children were retained so that only complete cases were analyzed. Epigenome-wide association analyses were conducted among mothers and children separately. Among children, the effects of maternal stress on children’s DNA methylation were examined. Linear regression models were applied with DNAm β-values as dependent variables, and total PSI-SF score as independent variables. Age and maternal smoking status were adjusted for potential confounding effects. Sex was also adjusted in the analyses among children. A reference-free EWAS method was used to account for potential heterogeneity in cell proportions from saliva samples [16]. Genomic control approach [17] adjustment was applied to address potential unmeasured confounding and inflation of type I error. Genomic control is a statistical method to measure the global deviation of a distribution of test statistics from a null distribution [17]. This method is not limited to a specific type of trait or study design, and does not require a case-control design. The method of genomic control has been widely applied to genome-wide association studies and EWAS of both quantitative and qualitative traits to measure and correct for the global inflation of low p-values to reduce false positive finding [18, 19]. False discovery rate was applied to account for multiple testing of epigenome-wide CpG sites. All analyses were conducted in the statistical computing environment [20] and a selection of packages from Bioconductor.

Results

The sample population is comprised of self-identified African American/Black women and one of their biological children who are between the ages of 3 through 5 years old. Demographic data are listed in Table 1 and in online Supplementary Table S1. The mean age of our maternal participants was 32.4 years old and 24% indicated that they currently smoke cigarettes. The mean age of our children was 3.7 years old, and 35% of children were male.

Significant variation in maternal DNA methylation at 95 CpG sites were associated with levels of parenting stress after genomic control (false discovery rate adjusted p < 0.05) (Fig. 1). Notably, the relationship between parenting stress and DNA methylation was inversely proportional at most sites (83 out of 95 sites, 87%) and had decreased DNA methylation with higher levels of parenting stress (online Supplementary Table S2). We did not identify any epigenome-wide significant differences in DNA methylation in children related to

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean</th>
<th>SD</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal age</td>
<td>32.351</td>
<td>6.042</td>
<td>21</td>
<td>46</td>
</tr>
<tr>
<td>Child age</td>
<td>3.730</td>
<td>0.746</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Parenting Stress Index total score</td>
<td>69.459</td>
<td>21.214</td>
<td>36</td>
<td>143</td>
</tr>
<tr>
<td>Parental distress subscale (12 items)</td>
<td>26.243</td>
<td>9.775</td>
<td>12</td>
<td>55</td>
</tr>
<tr>
<td>Parent-child dysfunctional interaction (12 items)</td>
<td>18.797</td>
<td>7.133</td>
<td>12</td>
<td>48</td>
</tr>
<tr>
<td>Difficult child (12 items)</td>
<td>24.419</td>
<td>8.153</td>
<td>12</td>
<td>46</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variable</th>
<th>Count</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal cigarette use</td>
<td>18</td>
<td>24.32%</td>
</tr>
<tr>
<td>Child gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>26</td>
<td>35.14%</td>
</tr>
<tr>
<td>Female</td>
<td>48</td>
<td>64.86%</td>
</tr>
</tbody>
</table>

Parenting Stress Index items were scored on a 5-point Likert scale, with larger scores corresponding to increased parenting stress.
the level of maternal parenting stress after genomic control. We compared the top 20 epigenetic associations with maternal parenting stress levels among children with the 95 significant CpG sites discovered among mothers. Summarized in Table 2, 4 CpG sites were found to meet epigenome-wide significance among mothers and were also marginally significant among children ($p < 5 \times 10^{-7}$).

We identified a change in DNA methylation associated with poly(ADP-ribose) polymerase-1 (PARP-1), which plays a key role in stress signaling. One of the CpG sites maps to 2 genes, kelch like family member 35 (KLHL35) and glycerophosphodiester phosphodiesterase domain containing 5 (GDPD5). KLHL35, is involved in the ubiquitination processes, although its specific role in the process

**Table 2.** Epigenome-wide associations with parenting stress identified among African Ancestry mothers and replicated among children

<table>
<thead>
<tr>
<th>CpG sites</th>
<th>Chr</th>
<th>bp</th>
<th>Gene</th>
<th>Analysis</th>
<th>$\beta$</th>
<th>SE</th>
<th>T-stat</th>
<th>$p$-value after GC</th>
<th>FDR adjusted $p$-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>cg10836755</td>
<td>1</td>
<td>226,546,944</td>
<td>PARP-1</td>
<td>Mother</td>
<td>$-8.61E-04$</td>
<td>1.47E-04</td>
<td>$-5.84$</td>
<td>1.07E-06</td>
<td>0.022</td>
</tr>
<tr>
<td>cg14574088</td>
<td>20</td>
<td>5,451,809</td>
<td>LOC643406</td>
<td>Mother</td>
<td>$-8.62E-04$</td>
<td>1.38E-04</td>
<td>$-6.26$</td>
<td>6.26E-07</td>
<td>0.090</td>
</tr>
<tr>
<td>cg15575851</td>
<td>5</td>
<td>68,168,648</td>
<td>KLHL35</td>
<td>Mother</td>
<td>$-8.51E-04$</td>
<td>1.44E-04</td>
<td>$-5.91$</td>
<td>8.55E-07</td>
<td>0.022</td>
</tr>
<tr>
<td>cg24260882</td>
<td>11</td>
<td>75,144,708</td>
<td>GDPD5</td>
<td>Mother</td>
<td>$-8.35E-04$</td>
<td>1.40E-04</td>
<td>$-5.97$</td>
<td>6.86E-07</td>
<td>0.022</td>
</tr>
</tbody>
</table>

bp, base pair; GC, genomic control; FDR, false discovery rate.

* FDR adjustment applied after GC approach.
Fig. 2. Regional plots of common significant epigenetic associations with maternal parenting stress in African Ancestry mothers and children (red circles: epigenetic associations in mother; blue circles: epigenetic associations in children). (a) through (d) correspond to the CpG sites listed in Table 2. CpG sites are listed at the top of each figure within the box annotating the location.
Discussion

Our study results suggest that levels of parenting stress are associated with differential DNA methylation in mothers, but not significantly in their young children. Of particular importance, we observed that higher levels of parenting stress was associated with hypomethylation in CpG sites located in the 3-UTR and gene body of PARP-1 (Fig. 2a). In this study, we conducted an EWAS of parenting stress, in which we interrogated individual CpG sites to identify significant epigenetic associations based on test for single CpG site, rather than test for multiple associated CpG sites within a local genomic region (ie, differential methylation regions). We presented regional plots of 4 regions harboring significant epigenetic associations in both mothers and children. Only the PARP-1 region has additional epigenetic associations, not the other 3 regions. Multiple associations from the same region may increase the confidence of true epigenetic associations. However, single associated CpG site within a region does not indicate a false positive given the stringent significant threshold and replication between mothers and children. The density of the EPIC array, ~3.7 Kb mean interval between CpG sites does not guarantee the coverage of co-methylated sites. In our experience, significant and replicated epigenetic associations can be either clustered or stand-alone. In the case of cigarette smoking, one of the most established epigenetic modifiers, AHRR region has multiple strongly smoking-associated CpG sites, and F2RL3 contains a single smoking-associated CpG site using 450 K chip [21]. To further explore the regional associations, we plotted all regions harboring significant association with parenting stress among mothers (online Supplementary Figure). Fifty out of 95 regions contain more than one suggestive associations (p < 0.001).

Altered DNA methylation associated with PARP-1 warrants further follow up in human population as the gene is known to play a key role in cellular function and response to stress. Alterations in the PARP-1 gene have been associated with genome instability, shorter lifespans, and spontaneous tumors in animal models [22]. Gene activity is often induced by DNA strand breaks and other conditions that signal the cell may come under stress, such as inflammation or changes in metabolic conditions. Up-regulation of PARP-1 can result in death of the cells by decreasing the amount of ATP and β-nicotinamide adenine dinucleotide (NAD+) available to carry out cellular processes [23]. Furthermore, PARP-1 functions in the regulation of DNA methylation modifications and is necessary for reprogramming somatic cells [24]. PARP-1 is widely expressed when DNA damage or other threats to the cell are present and deficiencies in PARP-1 have been associated with increased levels of DNA methylation [24]. Since high levels of PARP-1 are associated with cell death, and extreme deficiencies of PARP-1 leads to reduced transcription of genes that help repair DNA damage in response to cellular threats, the production of PARP-1 must be maintained in a specific range for optimal functioning. Alteration in DNA methylation at the promoter site of PARP-1 has resulted in altered gene expression in laboratory settings [25, 26]. Further studies are needed to determine if differences in DNA methylation at PARP-1 are associated with altered protein production or other measurable health effects in clinical populations. Since altered expression levels of PARP-1 are associated modification of DNA methylation and gene expression at other sites in the genome, differential methylation at this site in clinical populations may be precursor to other alterations that could have a lasting impact on future health outcomes.

Although the specific role of KLHL35 has not yet been identified in humans, others have observed differential methylation at CpG sites of the gene. Methylation at a cluster of CpGs near the second exon of KLHL35 was decreased in children that were born of older mothers [27]. The researchers also determined that the differential methylation persisted into adulthood and was replicable in another smaller sample. Since the role of KLHL35 is not yet known, the biological significance of the observed changes in DNA methylation are unclear. Similar to Markunas’ [27] results on maternal age at delivery, we identified a decrease in DNA methylation of KLHL35 with increased parenting stress. Conversely, hypermethylation of the KLHL35 sites has been associated with liver and renal cancers [28, 29]. It is unclear what health implications for hypomethylation associated with KLHL35 entail in our population or Markunas’ study. Although downstream protein production was not evaluated in our nor Markunas’ study, the observed DNA methylation mark may serve as a biomarker for parenting stress.

We chose to use saliva sampling in this study due to the ease of access and reduction of invasive procedures while working with young children. Our protocol produced high quality samples that passed QC protocols for the new EPIC array technology. One drawback to the use of saliva for DNA methylation analysis is the heterogeneity of DNA sources in saliva samples (epig, epithelial cells, leukocytes, and even foreign bacteria). However, a recent study comparing saliva and blood samples using the Illumina HumanMethylation450 BeadChip, found that DNA methylation levels in saliva may be more similar to brain tissue than blood, and the authors advocate for further exploration of the utility of saliva samples in psychiatric genetics [30]. Furthermore, our study detected differences in DNA methylation within the stress response pathway using saliva, a biofluid historically used to measure stress hormones such as cortisol. Future studies may choose to use saliva samples to concurrently measure DNA methylation and other biomarkers of stress with minimally invasive sample collection.

Because the study did not collect multiple samples for DNA methylation over time, it is unclear if the DNA methylation differences observed were a direct result of parenting stress or were preexisting differences that contribute to the mother’s response or perceived level of parenting stress. We also did not collect protein samples and are unable to evaluate if the differences in DNA methylation observed related to parenting stress are associated with differences in expression of PARP-1 or KLHL35. Since transcription levels PARP-1 that are too high and too low have been associated with negative outcomes in laboratory models, it will be important to evaluate implications in clinical settings related to health outcomes. Further research on the function of KLHL35 is necessary to determine if alterations contribute to changes in health or merely mark differences between groups.

Acknowledgments

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Disclosures

The authors have no conflicts of interest to declare.

Supplementary materials

To view supplementary material for this article, please visit https://doi.org/10.1017/cts.2018.3

References


