Associations among child abuse, mental health, and epigenetic modifications in the proopiomelanocortin gene (POMC): A study with children in Tanzania

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Abstract

Child abuse is associated with a number of emotional and behavioral problems. Nevertheless, it has been argued that these adverse consequences may not hold for societies in which the specific acts of abuse are culturally normed. Epigenetic modifications in the genes of the hypothalamus–pituitary–adrenal axis may provide a potential mechanism translating abuse into altered gene expression, which subsequently results in behavioral changes. Our investigation took place in Tanzania, a society in which many forms of abuse are commonly employed as disciplinary methods. We included 35 children with high exposure and compared them to 25 children with low exposure. Extreme group comparisons revealed that children with high exposure reported more mental health problems. Child abuse was associated with differential methylation in the proopiomelanocortin gene (POMC), measured both in saliva and in blood. Hierarchical clustering based on the methylation of the POMC gene found two distinct clusters. These corresponded with children’s self-reported abuse, with two-thirds of the children allocated into their respective group. Our results emphasize the consequences of child abuse based on both molecular and behavioral grounds, providing further evidence that acts of abuse affect children, even when culturally acceptable. Furthermore, on a molecular level, our findings strengthen the credibility of children’s self-reports.

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ity in all six countries. However, the strength of the relation did vary by the perceived normativeness across countries. Many other studies demonstrated detrimental consequences for the psychological well-being and development of abused children, regardless of whether the surrounding society deems such practices acceptable (Ani & Grantham-McGregor, 1998; Hecker, Hermenau, Isele, & Elbert, 2014; Hermenau et al., 2011).

There are many countries in which many of the acts constituting child abuse are legal and socially accepted. In Tanzania, for example, a national survey with a representative sample of more than 3,700 youths revealed that the great majority (almost 75%) of both girls and boys had experienced physical abuse and more than one-quarter faced emotional abuse prior to the age of 18 (UNICEF, 2011). In concordance, we and others reported the use of harmful physical acts and psychological tactics on behalf of caregivers toward children to be highly prevalent in Tanzanian families and schools (Feinstein & Mwahombela, 2010; Hecker et al., 2014). In April 2013, the Tanzanian government reportedly confirmed that the use of corporal punishment in public schools persists (Tanzania Daily News, 2013). Given such high prevalence of child abuse, it is vital for both individuals and societies to have a better understanding of the potential effects of abuse. In particular, we must study whether the negative consequences of physical and emotional abuse of children are diminished in societies where such acts are legal and socially accepted.

Most studies on mental health problems have been conducted in Western samples. However, findings from the Democratic Republic of the Congo, Ethiopia, and Nigeria have shown that various mental health problems such as anxiety disorders, affective disorders, and hyperactivity are also common phenomena in sub-Saharan Africa (Adelekan, Ndom, Ekpo, & Oluboka, 1999; Kashala, Elgen, Sommerfelt, & Tylleskar, 2005). Adelekan et al. (1999) indicated a prevalence rate of internalizing problems of 7.3% and of externalizing problems of 8% in a representative sample from Nigeria. Kashala et al. (2005) compared their findings in a study with a representative sample in the Congo (Goodman, Meltzer, & Bailey, 1998) with prior findings from Great Britain. They found that the mean scores on all subscales of the Strength and Difficulties Questionnaire (SDQ) were significantly higher than the British mean scores of a comparable sample. Hence, Cortina, Sodha, Fazel, and Ramchandani (2012) concluded that child and adolescent mental health problems are also common in sub-Saharan Africa.

Child Abuse and the Hypothalamus–Pituitary–Adrenal (HPA) Axis

The HPA axis, when functioning properly, helps us to deal with crises. It describes a set of interactions between the hypothalamus, the pituitary gland, and the adrenal gland, which results in the release of its effector cortisol (Chrousos & Gold, 1992; de Kloet, Joëls, & Holsboer, 2005). Upon stress per-

ception, corticotropin-releasing hormone (CRH) and arginine vasopressin (AVP) are released from the hypothalamic paraventricular nucleus to activate the synthesis of proopiomelanocortin (POMC) in the anterior pituitary. POMC is processed into several peptides including adenocorticotropic hormone (ACTH). Finally, ACTH is released into the bloodstream and triggers the secretion of cortisol from the adrenal cortex. At each organizational level, the HPA axis is tightly regulated by negative feedback loops mediated by glucocorticoid receptors (GRs). After binding their ligand, cortisol, GRs dampen HPA axis activity.

Child abuse is translated into negative long-term mental health outcomes via the HPA axis. It plays a central role, because it is tuned to experiences occurring early in life, making it highly susceptible to early childhood adversities (Heim & Nemeroff, 2001). For example, adults with a history of childhood maltreatment displayed altered ACTH and cortisol responses following exposure to an acute stressor (Carpenter et al., 2007; Heim et al., 2000). HPA axis dysregulation is a key feature of a range of psychopathological symptoms (Chrousos & Gold, 1992; de Kloet et al., 2005). Both human and animal studies show that unremitting threat or stress weakens the immune response and increases abdominal fat, mental ill health, and depression via alterations of HPA functioning (McEwen & Lasley, 2002). HPA axis function, and with it behavioral changes, may be stably altered through aberrant epigenetic modifications, established as the result of child abuse.

Epigenetic Modifications of HPA Axis Genes

Of the various and complex mechanisms leading to epigenetic modification, DNA methylation is currently being studied most extensively. In humans, the relationship between early life adversities and the methylation of the GR has been extensively studied. GR promoter methylation is associated with both child abuse and psychopathology (Dammann et al., 2011; Hompes et al., 2013; Labonte, Azoulay, Yerko, Turecki, & Brunet, 2014; McGowan et al., 2009). Suicide victims with a history of childhood abuse displayed increased GR methylation in brain tissue (Labonte et al., 2012; McGowan et al., 2009). Higher GR methylation in peripheral blood mononuclear cells has been observed in patients suffering from borderline personality disorder (i.e., in individuals who have usually been exposed to severe forms of abuse during development). Disruption or lack of adequate nurturing, as measured by child maltreatment and inadequate parental care, was also associated with increased GR promoter methylation (Perroud et al., 2011; Tyrka, Price, Marsit, Walters, & Carpenter, 2012). In addition, epigenetic changes in the POMC gene may promote HPA axis dysfunction. Recent studies suggest epigenetic programming of POMC operates through nutritional cues, such as being underweight (Ehrlich et al., 2010), while other research suggests an association with alcohol abuse and dependence via increased craving (Muschler et al., 2010). Animal models
demonstrate epigenetic programming of additional HPA axis genes such as CRH (Mueller & Bale, 2008) and AVP (Mur-
gatroyd et al., 2009). Thus, current research has highlighted epigenetic modifications in genes associated with the HPA axis as being a possible driving force producing child abuse-induced disorders.

In the present study we investigated associations of child abuse with both the phenotype and the methylation status of genes related to the HPA axis in Tanzanian children. We limited our analyses of DNA methylation to the genes coding for the main components of the HPA axis, that is, the genes coding for AVP, CRH, and POMC, from which ACTH is cleaved. In addition, we included the gene encoding GR, nuclear recepto-
tor subfamily 3, group C, member 1 (NR3C1), because several studies demonstrated its methylation status as being predictive for childhood abuse (Labonte et al., 2012; McGowan et al., 2009; Perroud et al., 2011; Tyrka et al., 2012). We hypothe-
sized that exposed children (a) report more emotional and be-
havioral problems and (b) display altered epigenetic modifica-
tions in the genes related to HPA axis functioning.

**Methods**

**Procedure**

In the context of a larger research project, a team of Tanzanian and German psychologists conducted structured interviews with a sample of Tanzanian school children (N = 409). Interviewers were taught in interview skills during a 2-week train-
ing session. Furthermore, the Tanzanian interviewers were trained to translate from English to Swahili and back in order to assist the German researchers. All instruments were trans-
lated in written form to Swahili. A valid and accurate trans-
lation into English was ensured through the use of a written, blind backtranslation. In the total sample, 33 interviews were rated by two independent assessors to determine high interrater reliability. Prior to the interviews, we sent a written informed consent form to all parents or caregivers of the children from Class 2 to 7 (age 6–15) explaining the purpose of the study.

Based on these structured interviews, we selected children who had been exposed to high levels of physical and emotional abuse in their homes and those who had been exposed to only low levels of physical and emotional abuse. An a priori power analysis (α = 0.05, power = 0.80, d = 0.80) using G^+^Power software (Faul, Erdfelder, Lang, & Buchner, 2007) indicated a required sample size of n = 26 per group to detect significant group differences. Therefore, we aimed for two groups from the extreme ends of the abuse continuum (no abuse vs. high levels of abuse) of 30 children each. Because many children, particularly younger children, reported a strong fear of drawing blood, due to harmful experiences in the Tanzanian health system, we decided not to include chil-
dren of 8 years or younger. We sent an invitation and in-
formed consent form to 96 parents and caregivers of the se-
lected children clarifying that donating blood and saliva samples would be entirely voluntary and no monetary com-
pensation would be offered. In total, 64% (n = 61 of 96) of the parents and caregivers signed the informed consent. We were unable to recruit enough children who had never been exposed to any type of abuse. This is not too surprising, given that several acts of child abuse are culturally normed and highly prevalent in Tanzania. Almost 75% reported exposure to physical abuse in a nationally representative sample (UNICEF, 2011). Nevertheless, our sampling approach resulted in two extreme groups: one group (n = 35) reporting high levels of child abuse (i.e., six or more different types) and one group (n = 25) reporting low levels of child abuse (i.e., four or fewer different types). In the total sample, 173 (42%) children re-
ported low levels of child abuse with only 8 (2%) reporting no exposure to any form of child abuse. In contrast, 175 (43%) children of the total sample reported high levels of child abuse. Only children with an informed consent signed by their caregivers and who also assented themselves orally were included in the study (only 1 child refused to participate despite parents’ informed consent). A trained nurse from the University of Konstanz with extensive work experience in East Africa collected the blood and saliva samples. The Ethical Review Board of the University of Konstanz approved the study. Other, nonepigenomic parts of the data gathered for the total sample are presented in reports by Hecker et al. (2014); Hermenau, Eggert, Landolt, and Hecker (2015); and Herme-
nau et al. (2014).

**Participants**

The children participating in this study were enrolled at a pri-
mary school in a city of approximately 150,000 inhabitants in southern Tanzania. The high-exposure group consisted of n = 35 children (60% girls) who were on average 11.31 years old (SD = 1.47, range = 9–15). The low-exposure group con-
sisted of n = 25 children (56% girls) who were on average 11.76 years old (SD = 1.20, range = 10–14).

**Measures**

All instruments were applied as a structured interview in Swa-
holi. The first part of the interview consisted of sociodemog-
ographic information, including age, grade, and gender.

**Child abuse.** We assessed exposure to abuse at home using the Maltreatment and Abuse Chronology of Exposure—Pe-
diatric Version (Isele et al., 2015; Teicher & Parigger, 2015). The Maltreatment and Abuse Chronology of Exposure is a structured interview for children consisting of 45 dichoto-
mous (yes/no) questions, measuring witnessed or self-experi-
enced forms of child maltreatment throughout the lifetime. In this study, we only used the 14 items covering possible forms of physical and emotional abuse (see Table 1) by an adult per-
son living in the same household (e.g., parent, relative, or care-
giver) or by a minor living in the same household (e.g.,
housemaid or sibling). In Tanzania many children grow up not only with their parents in one household but also with
other members of their extended families. We also focused on minors in the household because in urban Tanzania many children are raised by an underaged housemaid (12–17) as primary caregiver, while both parents have to work. Using an event checklist, we assessed the presence of different types of abuse but not the frequency. We calculated an abuse score by totaling up all of the question responses. The possible score ranges from 0 to 14. The Cohen \( \kappa \) coefficient measuring the interrater reliability was >0.99 (0.99–1).

**Mental health.** The self-evaluation of internalizing and externalizing problems was assessed with the SDQ (Goodman, Ford, Simmons, Gatward, & Meltzer, 2000; Goodman et al., 1998). We used the self-report version for children in interview form, which consists of 25 statements. The total difficulties score is generated by summing the scores of all items, except the items for prosocial behavior, and ranges from 0 to 40. A score over 16 indicates an enhanced level of internalizing and externalizing problems. In the present sample, the Cronbach \( \alpha \) coefficient was 0.71 and the Cohen \( \kappa \) coefficient was 0.99 (0.94–1).

The University of California at Los Angeles PTSD Reaction Index for Children DSM-IV (Steinberg, Brymer, Decker, & Pynoos, 2004) was used to screen for symptoms of PTSD, again in interview form. For each DSM-IV symptom, the frequency of occurrence within the last month is scored. The maximum score possible is 54. A threshold of 12 has been established as being ideal for identifying children at risk of depression (Kovacs, 2001; Kovacs, Goldstein, & Gastonis, 1993; Traube et al., 2010). In the present sample, the Cronbach \( \alpha \) was 0.81 and the Cohen \( \kappa \) was 0.99 (0.92–1).

**DNA methylation.** Lymphocytes from blood were isolated via a Ficoll gradient and stored in a preservation solution (DNAgard® Tissues & Cells, Biomatrica, San Diego, CA) in order to ensure recovery of high-quality DNA. In addition, saliva samples were collected and stored using the Oragene •DISCOVER (OGR-500) saliva collection kit (DNA Genotek Inc., Ontario, Canada). The tissue samples were subjected to DNA-extraction (DNeasy® Blood & Tissue Kit, Qiagen, Hilden, Germany). Genome-wide analysis of DNA methylation was then conducted at the Barts and the London Genome Centre (Queen Mary University of London). One microgram (1 \( \mu \)g) of genomic DNA was bisulfite converted (EZ DNA Methylation Kit, Zymo) and applied to the Human Methylation 450K array (Illumina). The raw data were preprocessed using both the R package lumi (Du, Kibbe, & Lin, 2008) and beta mixture quantile dilation as suggested elsewhere (Marabita et al., 2013). After preprocessing, DNA methylation was assessed for all of the 41, 26, 14, and 14 cytosine nucleotide–phosphate–guanine nucleotide (CpG) sites associated with the GR gene (NR3C1), the POMC gene, the CRH gene, or the AVP gene, respectively.

**Transcription factor binding site (TFBS).** To reveal potential functional properties associated with the CpG sites included in our study, the respective sequences were submitted for analysis. The presence of transcription factor binding sites was determined using the software TRANSFAC® and JASPAR® database. For each of its 27 items, the children were offered three statements and asked to choose the one which best describes their situation. The maximum score possible is 54. A threshold of 12 has been established as being ideal for identifying children at risk of depression (Kovacs, 2001; Kovacs, Goldstein, & Gastonis, 1993; Traube et al., 2010). In the present sample, the Cronbach \( \alpha \) was 0.81 and the Cohen \( \kappa \) was 0.99 (0.92–1).

Table 1. Occurrence of physical and emotional abuse during the children’s lifetime

<table>
<thead>
<tr>
<th>Physical abuse</th>
<th>Exposure % (n)</th>
<th>High</th>
<th>Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Has any adult intentionally pinched, slapped, punched, or kicked you?</td>
<td>80 (28)</td>
<td>48 (12)</td>
<td></td>
</tr>
<tr>
<td>2. Has any adult spanked you with the palm of his/her hand on buttocks, arms, or legs?</td>
<td>74 (26)</td>
<td>24 (6)</td>
<td></td>
</tr>
<tr>
<td>3. Has any adult spanked you with an object such as a belt, stick, tube, or wooden spoon?</td>
<td>89 (31)</td>
<td>60 (15)</td>
<td></td>
</tr>
<tr>
<td>4. Has any adult hit you so hard that you were injured?</td>
<td>40 (14)</td>
<td>4 (1)</td>
<td></td>
</tr>
<tr>
<td>5. Has any minor intentionally pinched, slapped, punched, or kicked you?</td>
<td>74 (26)</td>
<td>24 (6)</td>
<td></td>
</tr>
<tr>
<td>6. Has any minor spanked you with the palm of his/her hand on buttocks, arms, or legs?</td>
<td>51 (18)</td>
<td>4 (1)</td>
<td></td>
</tr>
<tr>
<td>7. Has any minor spanked you with an object such as a belt, stick, tube, or wooden spoon?</td>
<td>31 (11)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>8. Has any minor hit you so hard that you were injured?</td>
<td>46 (16)</td>
<td>0 (0)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Emotional abuse</th>
<th>Exposure % (n)</th>
<th>High</th>
<th>Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>9. Has any adult called you names or said hurtful things (e.g., fat, ugly, stupid)?</td>
<td>51 (18)</td>
<td>32 (8)</td>
<td></td>
</tr>
<tr>
<td>10. Has any adult yelled or screamed at you?</td>
<td>86 (30)</td>
<td>68 (17)</td>
<td></td>
</tr>
<tr>
<td>11. Has any adult locked you in a dark &amp; narrow place (e.g. basement, closet)?</td>
<td>20 (7)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>12. Has any minor called you names or said hurtful things (e.g. fat, ugly, stupid)?</td>
<td>77 (27)</td>
<td>4 (1)</td>
<td></td>
</tr>
<tr>
<td>13. Has any minor yelled or screamed at you?</td>
<td>60 (21)</td>
<td>8 (2)</td>
<td></td>
</tr>
<tr>
<td>14. Has any minor locked you in a dark and narrow place (e.g. basement, closet)?</td>
<td>3 (1)</td>
<td>0 (0)</td>
<td></td>
</tr>
</tbody>
</table>

**Note:** Adult, Person living in the same household (e.g., parent, relative, or caregiver); minor, person under the age of 18 living in the same household (e.g., housemaid or sibling).
to the Jaspar database (Mathelier et al., 2014) in order to predict known TFBSs. A conservative threshold of 90% sequence identity was applied.

**Data analysis**

For the analyses regarding either mental health or exposure to abuse, parametric Welch \( t \) tests were performed. For DNA methylation, individual 2 (Abuse) × 2 (Gender) analyses of variance (ANOVA) for each CpG site were performed using exposure to abuse and gender as between-group factors. We included gender in these analyses in order to account for potential effects arising from gender on DNA methylation. For three CpGs in blood (cg27107893, cg02079741, cg09916783) and one in saliva (cg23035419), the models did not fulfill the requirement of homogeneity of variances, as indicated by a significant Levene test (Fox & Weisberg, 2011) and are thus not reported. Nonparametric tests could not be performed because these would not control for the potential influence of gender. In addition, we computed individual 2 (tissue) × 2 (gender) ANOVAs for each CpG site using tissue and gender as between-group factors. Because of the heterogeneity of variances, 25 probes were excluded from the analyses (NR3C1: cg06613263, cg08818984, cg08845721, cg10847032, cg18998365, cg19457823, cg23035419, POMC: cg02079741, cg03560973, cg08030082, cg09527270, cg09672383, cg09916783, cg13025668, cg16302441, cg20387815, cg20807790; CRH: cg00603617 cg20327580; and AVP: cg03279206, cg04360210, cg14065127, cg23035419, cg24257309). Nonparametric tests could not be performed because these would not control for the potential influence of gender.

All analyses used a two-tailed \( \alpha = 0.05 \). Our metric for a small effect size was \( d \geq 0.20 \) or \( \eta^2 \geq 0.01 \), for a medium effect \( d \geq 0.50 \) or \( \eta^2 \geq 0.06 \), and for a large effect \( d \geq 0.80 \) or \( \eta^2 \geq 0.13 \). To adjust for multiple testing (for three mental health variables and across the CpG sites for each gene), \( p \) values were computed according to Benjamini–Hochberg (Benjamini & Hochberg, 1995) applying a false discovery rate of 0.05. In an exploratory approach, we also considered the unadjusted \( p \) values. All statistical analyses were performed using IBM SPSS Statistics version 21 for Mac or R for Mac version 3.0.3.

**Results**

**Mental health**

Table 2 displays the descriptive statistics for both groups. In concordance with the sample selection, the high-exposure group reported a substantially higher number of different abuse types than the low-exposure group. The differences between the two groups are especially notable for the items indicating that a minor in the household was the perpetrator of the abuse (see Table 1). All mental health variables (SDQ, UCLA PTSD Reaction Index, and Children’s Depression Inventory) differed significantly between groups with medium to large effects (see Table 2). In total, \( n = 11 \) (31%) children in the high-exposure group showed an enhanced level of internalizing and externalizing problems compared to \( n = 1 \) (4%) in the low-exposure group. Accordingly, \( n = 9 \) (26%) children in the high-exposure group fulfilled the clinical diagnosis for PTSD compared to \( n = 2 \) (8%) in the low-exposure group. In addition, \( n = 10 \) (29%) children in the high-exposure group were at risk of suffering from depression compared to \( n = 1 \) (4%) in the low-exposure group.

**DNA methylation of genes associated with the HPA axis**

We found a group difference between the high-exposure and low-exposure groups in POMC with higher DNA methylation in children with high exposure. This effect was particularly evident in saliva. In the saliva of the high-exposure group, one CpG site was significantly hypermethylated in one-tailed tests at an adjusted significance level of .05, and three additional CpG sites would be significantly hypermethylated an adjusted significance level of .05, and three additional CpG sites were at risk of suffering from depression compared to \( n = 1 \) (4%) in the low-exposure group.
site in POMC displayed a moderate effect size, although unadjusted $p$ values exceeded the significance level of .05.

For the remaining HPA axis genes investigated, we did not find a clear group difference in DNA methylation. In saliva, four CpG sites were differently methylated in GR and one in CRH, displaying moderate effect sizes and unadjusted $p$ values below .05. In the blood of the high-exposure group, one CpG was hypermethylated in CRH at an adjusted significance level of .05, displaying a large effect. If uncorrected $p$ values were considered, one additional differentially methylated CpG could be found in AVP, displaying a medium effect. If only effect sizes were considered, two additional CpG sites associated with GR and one associated with AVP differed between the groups in blood, displaying moderate effect sizes, but no significant $p$ values were obtained.

Because we found the most pronounced effects in POMC, we inspected the 7 and 9 CpGs, which differed between the groups with at least moderate effect size in blood and saliva, respectively, in more detail. A comparison with the Jaspar database (Fox & Weisberg, 2011) revealed that 5 and 6 of these CpGs are either located in or directly flanking a potential TFBS. The potential TFBSs included TFAP2A, ZEB1, THAP1, YY1, BRCA1, E2F, ZNF354C, MZF1, and SPIB. It is interesting that all of these CpGs are located in the 5' promoter, whose methylation status has been shown to modulate transcriptional activity of the POMC gene (Newell-Price, King, & Clark, 2001). Our analyses covered 11 and 12 CpGs in this region in blood and saliva, respectively. In blood, 1 CpG site in this region was excluded from the analyses due heterogeneity of variances. Thus, about one-half of the CpGs in this region differed in their methylation by means of child abuse, and are associated with TFBSs.

**DNA methylation of the POMC gene strengthens children’s self-reports**

Post hoc we hypothesized that we could replicate, on the molecular level, the group allocation that was originally based on children's self-reports. We performed unsupervised hierarchical clustering on methylation of the 26 CpG sites representing the POMC gene using the Euclidean distance metric and the ward clustering method in the hclust package in R. To account for the dispersion differences across the methylation of the CpG sites, data were z-standardized prior to cluster analysis. Both in blood and in saliva, two distinct clusters reflecting the high-exposure and low-exposure groups could be detected (Figure 3). In blood, the analysis allocated $n = 39$ (68%) children into their respective group, and in saliva $n = 35$ (60%). A chi-square test confirmed the significant concordance between the group allocation based on children’s self-report and based on methylation
Figure 2. (Color online) DNA methylation of hypothalamic–pituitary–adrenal axis genes. Mean methylation of all analyzed cytosine nucleotide–phosphate–guanine nucleotide (CpG) sites. CpG sites are ordered according to their genomic location (not drawn to scale). For visual purposes, the data were mean centered. Beneath the scatterplots, the respective CpG sites and their positions in the gene model are displayed. CpG sites, which revealed at least moderate effect sizes comparing the high- and low-exposure groups are highlighted in black and bold font. AVP, Arginine-vasopressin gene; CRH, corticotropin-releasing hormone gene; NR3C1, nuclear receptor subfamily 3, group C, member 1 glucocorticoid receptor gene; POMC, proopiomelanocortin gene. • Adjusted $p < .1$, *adjusted $p < .05$; **adjusted $p < .01$; ***adjusted $p < .001$. Black asterisks/dots depict tissue comparisons, red asterisks/dots (online only) depict comparisons in relation to child abuse in blood, and blue asterisks/dots (online only) depict comparisons in relation to child abuse in saliva.
value in blood ($\chi^2 = 5.95, df = 1, p = .015$) and showed a trend in saliva ($\chi^2 = 3.49, df = 1, p = .062$).

**DNA methylation of HPA axis genes**

We also compared DNA methylation in the four HPA axis genes between the two tissues. Generally, blood tended to show stronger signals of DNA methylation than saliva (Figure 2). The only exception was seen in AVP, in which the pattern was reversed and saliva was characterized by elevated DNA methylation levels compared to blood. This tendency was also revealed in the ANOVAs, as we found 3, 8, 18, and 11 CpG sites in AVP, CRH, POMC, and NR3C1, respectively, which displayed differential methylation between the tissues (online-only supplementary Table S.1).

**Discussion**

Child abuse is known to impair mental health across the entire life span (Carr et al., 2013). However, it has been claimed that the effects of specific forms of child abuse are not as harmful when they take place in societies or cultural groups in which such practices are common, socially accepted, and legal. Lansford et al. (2005), for example, demonstrated that the relation between corporal punishment and mental health problems varied with the perceived normativeness of corporal punishment in the respective country. However, we and others have already demonstrated the detrimental effects of child abuse in such societies (Ani & Grantham-McGregor, 1998; Hecker et al., 2014). In concordance, in the present study children with high exposure to child abuse showed decreased psychological well-being. Furthermore, we demonstrated that this link manifests itself on a molecular level that cannot be manipulated by the subject: child abuse was strongly associated with the methylation of the POMC gene in both blood and saliva. To date, research incorporating child abuse and the methylation of HPA axis genes has focused mainly on the GR gene (de Kloet et al., 2005; Labonte et al., 2014; McGowan et al., 2009; Perroud et al., 2011). Little is known about the physiological and phenotypic consequences of POMC methylation. The POMC gene is characterized by a 5' CpG islands, located at exon 1 and the promoter region, and a 3' CpG island more downstream around the intron 2 and exon 3 boundary (Gardiner-Garden & Frommer, 1994). Research investigating various disease traits or stress exposure has mainly reported differential methylation at the 5' CpG island (Mizoguchi et al., 2007; Muschler et al., 2010; Newell-Price et al., 2001; Stevens et al., 2010), but effects on the 3' CpG island (Kuehnen et al., 2012) have also been reported. In cancer tissue that did not belong to the pituitary
gland that caused Cushing syndrome (hypercortisolism), differential POMC methylation at the 5’ CpG island and increased ACTH levels were reported, suggesting HPA axis dysregulation, a key feature of many mental diseases (Mizoguchi et al., 2007; Newell-Price et al., 2001). Our research supports these previous findings, because the majority of differentially methylated CpGs in our study were located in the 5’ CpG island. Moreover, the respective CpG locates with TFBSs, suggesting transcriptional regulation. These TFBSs include an E2F response element, methylation of which has been shown to suppress POMC promoter activity in vitro (Newell-Price et al., 2001).

In addition to ACTH, the functionally relevant peptides β-endorphin and α-melanocyte stimulating hormone (αMSH) are cleaved from the prohormone POMC. Thus, the possible impairment of other systems than the HPA axis through POMC methylation has to be considered. β-Endorphin has antinociceptive effects that are essential for stress, in particular, the fight–flight situations. It also was reported to have rewarding properties and is considered as a factor in stress-related psychiatric disorders (Merenlender-Wagner, Dikshtein, & Yadid, 2009) and drug abuse (Roth-Deri, Green-Sadan, & Yadid, 2008). POMC methylation was associated with alcohol craving in patients suffering from alcohol dependence (Muschler et al., 2010). Thus, differential POMC methylation by means of child abuse, as found in our study, may heighten the risk of the development of abuse-related mental illness (Carr et al., 2013), including drug abuse (Dube et al., 2003). Abuse seems to affect the methylation of the POMC gene and may lead to increased emotional and behavioral problems in the children, which then increase the likelihood for further abuse. In short, settings of frequent abuse would generate a vicious cycle of further abuse and behavioral problems. Because of the nature of our study, it was not possible to test this idea statistically. Future studies using larger samples and longitudinal designs should test this hypothesis empirically. Nevertheless, our findings are congruent with prior findings that child abuse is related to worse child mental health, even in a society in which specific acts of child abuse are common practice.

DNA methylation profiles appear to be tissue specific (Ollikainen et al., 2010), and several studies indicated a clear separation of samples derived from saliva and blood (Smith et al., 2014; Thompson et al., 2013; Wu et al., 2014). Accordingly, we found significantly different methylation profiles between saliva and blood. Moreover, there was a general trend of hypermethylation in saliva, which has previously been demonstrated. However, despite tissue-specific methylation, we demonstrate

Figure 3. (Color online) Hierarchical clustering dendrogram. Based on the methylation of 26 cytosine nucleotide–phosphate–guanine nucleotide sites present in the proopiomelanocortin gene a hierarchical cluster analysis has been performed. Two distinct clusters were formed in both blood (a) and saliva (b) significantly replicating the two groups that are based on children’s self-reports regarding exposure to child abuse. The parts of the dendrograms highlighted in red (online only) represent the clusters containing mainly children exposed to high levels of child abuse while the turquoise highlighted segments denote the clusters containing mainly children with low exposure. The colored boxes (online only) next to the final branches denote the exposure to childhood abuse based on the self-reports (red, high exposure; turquoise, low exposure).
that childhood abuse is associated with DNA methylation in both saliva and blood. Thus, methylation evoked by adverse experiences seems to be preserved across tissues.

Moreover, parents and caregivers often argue that children tend to overreport the exposure to abuse and the resulting harm. Thus the children’s perception of their experiences is often ignored, because children are not regarded as being mature enough to accurately gauge their situation (Qvortrup, Bardy, Sgritta, & Wintersberger, 1994). Hierarchical clustering based on the methylation of POMC, however, allocated two-thirds of children into their respective group and a subsequent chi-square test confirmed the significant concordance between the group allocations based on children’s self-report and based on methylation value. Therefore, our results strengthen the credibility of children’s self-reports on a molecular level and support the conclusion that children are capable of accurately reporting their exposure to abuse. In the school context of our data assessment, we were unable to include parents’ reports for logistical reasons. Furthermore, we deliberately focused on the credibility of children’s reports, because their view has been often neglected in research thus far. While it is possible that the inclusion of parents’ reports could have further strengthened our findings, previous studies in resource-poor countries cast doubt on the validity of parents’ knowledge about their children’s suffering (Elbert et al., 2009).

The methodologies employed by our study present some limitations. Our data are correlational in nature and thus cannot prove a causal relationship between child abuse and methylation patterns or decreased psychological well-being. Nevertheless, even if certain methylation patterns might increase the likelihood of child abuse, the data still confirm the credibility of children’s subjective reports and with it a wealth of data showing that abused children are more likely to suffer. However, the sample size and our study design using extreme group comparisons limit the generalizability of our findings. In the school context of our data assessment, we were unable to include parents’ reports for logistical reasons. Therefore, we could not gather information regarding the socioeconomic status of our sample. It remains to be tested whether socioeconomic status can impact DNA methylation through other pathways than abuse. Furthermore, it has been suggested that probes containing single nucleotide polymorphisms (SNPs) might result in a biased signal (Price et al., 2013). Based on the 1000 Genomes Project’s database (1000 Genomes Project Consortium, 2012) eight SNPs colocalize with the target sequence of probes associated with POMC. However, the majority of those are very rare in African populations with minor allele frequencies below 0.2% and are thus considered not relevant to our sample. Excluding one CpG, whose respective probe contained a SNP in their target sequence at higher minor allele frequency (i.e., 1.0%), did not markedly change the results (data not shown). Furthermore, this SNP was located more than 10 base pairs away from its target CpG, which does not seem to evoke biased signals (Price et al., 2013). Therefore, we consider our findings to reflect the epigenome of the participants and not as artifacts of their genotype.

In summary, we provide further evidence that in societies or cultural groups in which many specific acts of child abuse are common, legal, and socially accepted, child abuse is nevertheless detrimental for the psychological well-being of affected children. Our evidence for such a link is strengthened by the inclusion of epigenetic information from both blood and saliva. This is the first study reporting the link between child abuse and modifications of DNA methylation of POMC. Epigenetic modifications provide a promising mechanism through which child abuse could act to influence psychological well-being. In addition, on a molecular level, our study strengthens the credibility of children’s self-reports evaluating their exposure to abuse. All in all, our findings underscore the need to inform the population at large about the adverse consequences associated with various forms of child abuse, both those societally accepted and those not. This holds especially true in societies in which such practices are commonly employed and generally regarded as effective.

Supplementary Material
To view the supplementary material for this article, please visit http://dx.doi.org/10.1017/S0954579415001248.


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