



## **King's Research Portal**

DOI: 10.1016/j.jpsychires.2016.09.010

Document Version Peer reviewed version

Link to publication record in King's Research Portal

Citation for published version (APA):

Cecil, C., Smith, R. G., Walton, E., Mill, J., McCrory, E. J., & Viding, E. (2016). Epigenetic signatures of childhood abuse and neglect: Implications for psychiatric vulnerability. *Journal of psychiatric research*. https://doi.org/10.1016/j.jpsychires.2016.09.010

### Citing this paper

Please note that where the full-text provided on King's Research Portal is the Author Accepted Manuscript or Post-Print version this may differ from the final Published version. If citing, it is advised that you check and use the publisher's definitive version for pagination, volume/issue, and date of publication details. And where the final published version is provided on the Research Portal, if citing you are again advised to check the publisher's website for any subsequent corrections.

#### General rights

Copyright and moral rights for the publications made accessible in the Research Portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognize and abide by the legal requirements associated with these rights.

•Users may download and print one copy of any publication from the Research Portal for the purpose of private study or research. •You may not further distribute the material or use it for any profit-making activity or commercial gain •You may freely distribute the URL identifying the publication in the Research Portal

#### Take down policy

If you believe that this document breaches copyright please contact librarypure@kcl.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.

## Epigenetic signatures of childhood abuse and neglect: Implications for psychiatric vulnerability

### SELF-ARCHIVING VERSION

Charlotte A. M. Cecil<sup>1</sup>\*, Rebecca G. Smith<sup>1,2</sup>\*, Esther Walton<sup>1,3</sup>, Jonathan Mill<sup>1,3</sup>, Eamon J. McCrory<sup>4†</sup> & Essi Viding<sup>4†</sup>

- 1. Department of Psychology, Institute of Psychiatry, Psychology and Neuroscience, King's College London, London, SE5 8AF, UK.
- 2. University of Exeter Medical School, University of Exeter, Exeter, EX1 2LU, UK
- 3. Department of Psychology, Georgia State University, Atlanta, 30302-5010, USA
- 4. Division of Psychology and Language Sciences, University College London, London, WC1H 0AP, UK.

\*Co-first authors <sup>†</sup>Co-senior authors

**Citation:** Cecil, C.A.M., Smith, R.G., Walton, E., Mill, J., McCrory, E.J. & Viding, E. (In Press). Epigenetic signatures of childhood abuse and neglect: Implications for psychiatric vulnerability Assessment. *Journal of Psychiatric Research*.

**Correspondence to**: Charlotte Cecil, Department of Psychology, Institute of Psychiatry, Psychology and Neuroscience, King's College London, De Crespigny Park, London, SE5 8AF, UK. Tel: +44 (0)207 848 0389, E-mail: <u>charlotte.cecil@kcl.ac.uk</u>.

**Acknowledgements:** We would like to thank the young people, teachers and key workers who have taken part in this project. We are grateful to the Urban Academy, Haberdashers' Aske's Hatcham College and Hertswood Academy for their valuable collaboration. We thank Jo Guiney, Yvonne Whelan and Kathryn Hubbard for their assistance with data collection. Research reported in this publication was supported by the Waterloo Foundation (Award Number: R1233/1872; PIs: E.V. and J.M.) and Kids Company charity. C.C. is supported by the Economic and Social Research Council (grant ref: ES/N001273/1). E.W. was supported by the German Research Foundation (Wa 3635/1-1). E.V. is a Royal Society Wolfson Research Merit Award Holder.

### Abstract

Childhood maltreatment is a key risk factor for poor mental and physical health. Recently, variation in epigenetic processes, such as DNA methylation, has emerged as a potential pathway mediating this association; yet, the extent to which different forms of maltreatment may be characterized by unique vs shared epigenetic signatures is currently unknown. In this study, we quantified DNA methylation across the genome in buccal epithelial cell samples from a high-risk sample of inner-city youth (n = 124; age = 16-24; 53% female), 68% of whom reported experiencing at least one form of maltreatment while growing up. Our analyses aimed to identify methylomic variation associated with exposure to five major types of childhood maltreatment. We found that: (i) maltreatment types differ in the extent to which they associate with methylomic variation, with physical exposures showing the strongest associations; (ii) many of the identified loci are annotated to genes previously implicated in stress-related outcomes, including psychiatric and physical disorders (e.g. GABBR1, GRIN2D, CACNA2D4, PSEN2); and (iii) based on gene ontology analyses, maltreatment types not only show *unique* methylation patterns enriched for specific biological processes (e.g. physical abuse and cardiovascular function), but also share a 'common' epigenetic signature enriched for biological processes related to neural development and organismal growth. A stringent set of sensitivity analyses were also run to identify high-confidence associations. Together, findings lend novel insights into epigenetic signatures of childhood abuse and neglect, point to novel potential biomarkers for future investigation and support a molecular link between maltreatment and poor health outcomes. Nevertheless, it will be important in future to replicate findings, as the use of cross-sectional data and high rates of polyvictimization in our study make it difficult to fully disentangle the shared vs unique epigenetic signatures of maltreatment types. Furthermore, studies will be needed to test the role of potential moderators in the identified associations, including age of onset and chronicity of maltreatment exposure.

Keywords: DNA Methylation; epigenome-wide; child abuse; neglect; maltreatment; stress

### Introduction

Childhood maltreatment, encompassing abuse and neglect, is a major public health concern that continues to affect up to one in four children worldwide, with often devastating developmental consequences (WHO, 2014). Children who experience maltreatment are at increased risk for a range of psychiatric problems, including anxiety, depression, post-traumatic stress, and antisocial behaviour (Cicchetti and Toth, 2005). The effects of maltreatment can extend well into adulthood, compromising relationship quality, economic productivity and physical health (Danese *et al.*, 2009).

The theory of latent vulnerability proposes that maltreatment exposure calibrates a range of biological and neurocognitive systems in line with a threatening and unpredictable early environment (McCrory and Viding, 2015). While potentially adaptive in the short term, such changes can increase vulnerability in the long term. Consistent with this view, numerous biological correlates of maltreatment have now been identified, including accelerated cellular ageing, neuroendocrine dysregulation, heightened inflammatory response as well as altered brain structure and function (Danese *et al.*, 2011, McCrory *et al.*, 2012, Shalev *et al.*, 2013). Recent evidence indicates that, as well as affecting common biological pathways, different forms of maltreatment may also exert unique effects. For example, while abuse has been associated with changes in neural circuitry underlying threat processing, neglect has been associated with biological adaptations to low-complexity environments (Sheridan and McLaughlin, 2014).

A key challenge for current research is to understand how, at a molecular level, these environmental exposures are translated into phenotypic variation. Epigenetic processes, such as DNA methylation (DNAm), which control the functional regulation of gene expression are of particular interest in this regard, as mounting evidence suggests they can be modified by environmental factors (Jaenisch and Bird, 2003). For example, animal studies have found that a number of environmental stressors, such as poor maternal care, induce stable alterations in DNAm in the regulatory regions of several HPA axis genes (e.g. the glucocorticoid receptor), which in turn influence responses to future stressors (Turecki and Meaney, 2014). Similarly, a small number of human studies have documented a link between childhood maltreatment and aberrant DNAm in genes important for stress-response, immune function and neurodevelopment (Lutz and Turecki, 2014). DNAm has also been shown to regulate a wide range of neurobiological processes, including neurogenesis, synaptic plasticity, learning and memory (Baker-Andresen *et al.*, 2013, Day *et al.*, 2013) and aberrations in DNAm have been observed in a range of diseased states, including stress-related psychiatric disorders such as post-traumatic stress and major depression (Bergman and Cedar, 2013, Klengel *et al.*, 2014).

To date, most epigenetic studies of maltreatment have focused on variation in the vicinity of a limited set of pre-selected candidate genes (i.e. *GR*, *FKBP5*, *BDNF and 5-HTT*) (Lutz and Turecki, 2014). As such, little is known about the broader effect of maltreatment on DNAm across the genome. This is a substantial limitation in light of the fact that maltreatment impacts multiple aspects of functioning, across psychological, physical, and social domains. Furthermore, existing studies have primarily examined global maltreatment (Labonte *et al.*, 2012, Prados *et al.*, 2015, Suderman *et al.*, 2014, Yang *et al.*, 2013), so that the extent to which different maltreatment types may have common vs distinct epigenetic signatures is unclear. To address these outstanding questions, we explored the relationship

between DNAm and five types of maltreatment in a sample of high-risk youth, using genome-wide DNAm data drawn from buccal epithelial cells.

## **Methods and Materials**

### Participants

The current sample was recruited as part of a larger study examining the effects of developmental adversity on individual functioning (n = 204, age range = 16-24 years). Analyses only included participants for whom DNAm data was available (n = 124). Youth from deprived inner London areas were recruited through multiple channels including innercity colleges, internet websites and a charity providing services and support to self-referred youth. The sample was 53% female and ethnically diverse (49% White, 33% Black, 18% other). The study was carried out in accordance with the latest version of the Declaration of Helsinki. The study design was reviewed and approved by the UCL Research Ethics Committee (*ID No*: 2462/001) and all participants provided informed consent prior to participation, after the nature of the procedures had been fully explained. Further details of the sample and recruitment procedures are available elsewhere (Cecil *et al.*, 2014).

## Measures

<u>Childhood maltreatment</u> – Childhood maltreatment was assessed using the 28-item, selfreport Childhood Trauma Questionnaire (CTQ; Bernstein and Fink, 1998). The CTQ screens for experiences of maltreatment "while growing up" and comprises of 5 continuous subscales: emotional abuse, sexual abuse, physical abuse, emotional neglect and physical neglect. The scales show high internal consistency in our sample ( $\alpha = .70 - .97$ ). For descriptive purposes only, we also classified participants as having experienced maltreatment (i.e. yes/no) if they scored above the 'Low' threshold specified by the CTQ manual for at least one maltreatment type. By including 'I currently feel unsafe at home' as an additional yes/no item we were able to ascertain that none of the participants in the study were currently vulnerable to violence in the domestic environment (e.g. by family or partner). As such, the present study investigates the effects of childhood (i.e. past) maltreatment.

<u>DNA methylation</u> – DNA was extracted from buccal epithelial cells using procedures described in Freeman *et al.* (2003). 500ng of high molecular weight DNA was subjected to sodium bisulfite conversion using the EZ-DNA methylation kit (Zymo Research, Orange, CA, USA) using the manufacturers standard protocol. DNAm was quantified using the Illumina HumanMethylation450 BeadChip (Illumina, USA) with arrays scanned using an Illumina iScan (software version 3.3.28). The Illumina 450K array interrogates >485,000 probes covering 99% of Reference Sequence (RefSeq) genes, with an average of 17 CpG sites per gene region. As the samples were run in a single batch, there was no need for batch correction. To account for potential chip and position effects, we randomized sample chip allocation and placement on the chip. Initial data quality control was conducted using GenomeStudio (version 2011.1) to determine the status of staining, extension, hybridization, target removal, bisulfite conversion, specificity, non-polymorphic and negative controls. Samples that survived this stage were checked for concordance between their reported and assessed sex and then quantile normalised using the *dasen* function within the *wateRmelon* 

package (wateRmelon\_1.0.3; Pidsley *et al.*, 2013) in R. Probes were removed if they were cross-reactive, polymorphic, used for sample identification on the array, had a SNP at the single base extension with a minor allele frequency larger than 5% (i.e. common polymorphisms) or were located on the Y chromosome, leaving a total of 413,239 probes (Chen *et al.*, 2013; Price *et al.*, 2013). DNAm levels are indexed by beta values (ratio of methylated signal divided by the sum of the methylated and unmethylated signal, M/M+U).

### Data Analysis

All analyses were performed within the R statistical environment (version 3.0.1). Methylation data was regressed for sex, age and self-reported ethnicity to account for potential confounding effects (Liang and Cookson, 2014). The analysis proceeded in three steps. First, we ran five independent epigenome-wide association analyses - one for each maltreatment type measured – using linear regression models. Probes were considered significant if they survived a False Discovery Rate (FDR) correction of q < 0.05. Only maltreatment types that were associated with at least one FDR-corrected probe were carried forward to the next step. Second, we identified which probes were most consistently associated with all types of maltreatment, by ranking them in order of average standardized effect size. Third, we examined enriched biological pathways for genes that were associated only with one type of maltreatment vs those that were associated with all maltreatment types, using an optimized gene ontology method that controls for a range of potential confounds, including background probe distribution and gene size (see **OS1** for details). More specifically, genes were considered 'unique' if the probes annotated to them were specifically associated with one maltreatment type and none other (i.e. p<0.005 with one type of maltreatment, and p>0.05 with the other two types of maltreatment). Conversely, genes were considered 'shared' if probes annotated to them were associated across maltreatment types (p < 0.05 consistently across all three forms of maltreatment). To calculate statistical power to detect effects, we used the pwr package with a p-value threshold of 1.00E-06 to reach 80% power (Tsai & Bell, 2015) in the context of a linear regression model. Results showed that, with a sample size of 124 individuals, we could expect to detect an effect size of  $\geq 0.32$ , indicating that the study is appropriately powered for detecting medium-to-large effects.

### Results

Descriptives and correlations between study variables shown in **Table 1**. Based on the CTQ cut-off, the majority of youth in the current study reported having experienced at least one form of maltreatment while growing up (68%; n = 84). The reported experiences were often severe: 75% of physically abused youth were left with bruises or marks as a result of injuries; 50% of sexually abused youth reported being threatened; and 60% of physically neglected youth reported not having enough to eat. Poly-victimization was common, with 74% of maltreated youth reporting two or more forms of maltreatment, consistent with previous studies (Radford *et al.*, 2011).

*Maltreatment types differ in the extent to which they associate with DNAm variation* We first examined associations between DNAm across the genome and individual maltreatment types (measured continuously). No probes were identified as being differentially methylated at a FDR<0.05 for emotional abuse and emotional neglect (OS2); as such, these two types of maltreatment were excluded from further analysis. With regards to other maltreatment types, the number of differentially methylated probes (DMPs) was: 34 for physical abuse, 7 for sexual abuse, and 118 for physical neglect. The 10 top-ranked DMPs for each of these maltreatment types are displayed in **Table 2** (see **OS3** for a complete table of FDR-corrected probes). Associations between DNAm and each maltreatment type are graphically represented in **Figure 1A** (see **OS4** for quantile-quantile plots for each analysis).

<u>Physical abuse</u>: The top-ranked DMP, cg20000641, was significantly hypermethylated with increased exposure to physical abuse (p=3.54E-11, q=1.47E-05; **Table 2A**, **Figure 1B**). This probe is located in the promoter region of *PSEN2*, a gene encoding a presenilin enzyme involved in amyloid precursor protein processing that is robustly implicated in Alzheimer's disease (O'Brien and Wong, 2011). Two top-ranked probes were annotated to *SMC1A* (cg02353937: p = 2.90E-09, q=7.89E-04; cg22311608: p = 8.45E-08, q=0.01), a gene involved in chromosomal maintenance and DNA repair (Kim *et al.*, 2002). Also of interest were DMPs annotated to genes highly expressed in the brain, including *SHC2* (cg19736040: p = 1.38E-06, q=0.05), involved in neurotrophin-activated Trk receptor signaling within cortical neurons and synaptic plasticity in the hippocampus (Epa *et al.*, 2004), and *IMPACT* (cg03013329: p = 2.10E-06, q=0.05), encoding a protein that facilitates neurite outgrowth and modulates kinase activation in neurons (Roffe *et al.*, 2013).

<u>Sexual abuse</u>: The top-ranked DMP associated with sexual abuse (cg17106653, p=4.16E-09, q=1.72E-03; **Table 2B**, **Figure 1B**) was located in the promoter region of the glutamate receptor *GRIN2D*, a gene implicated in CNS plasticity and excitatory synaptic transmission (Traynelis *et al.*, 2010). Other annotated genes of interest include *MGMT* (cg26528551: p = 6.84E-08, q=0.01) implicated in DNA repair mechanisms and *DIP2C* (cg23983710: p = 5.83E-07, q=0.03), a gene primarily expressed in the brain but whose biological function is poorly understood.

<u>Physical Neglect</u>: The top-ranked DMP associated with physical neglect was cg00691266 in *EVPN* (p=1.01E-07, q=0.01; **Table 2C**), encoding a protein involved in epidermal growth. Several probes were annotated to genes related to brain function, including *SYNJ2* (cg27083825: p = 1.02E-07, q=0.01; **Figure 1B**), involved in nervous system development and neuronal vesicle uncoating (Montesinos *et al.*, 2005); and *GABBR1* (cg18116160: p = 4.01E-07, q = 0.02) a GABA class B receptor important for inhibitory synaptic transmission (Kumar *et al.*, 2013). Probes annotated to genes involved in histone regulation were also identified, including *SETDB1* (cg17918089: p = 1.14E-07, q=0.01), *JADE1* (cg09863040: p = 2.10E-07, q=0.01), and *HIST1H1A* (cg08054907: p = 5.12E-07, q=0.02).

<u>Sensitivity analyses:</u> Given that quantile-quantile plots from our epigenome-wide analyses indicated skewing of significant results (**OS4**), we carried out **a set of** sensitivity analyses to test the robustness of the identified associations. **First**, to minimize the influence of unknown

confounders, we ran a surrogate variable analysis (sva package in R; Leek et al., 2012) that enabled us to identify unwanted sources of variation in our DNA methylation data. The analysis identified 11 surrogate variables representing different sources of variation/noise. We then re-ran associations between each form of maltreatment and DNAm additionally controlling for these 11 surrogate variables. While variations in significance levels were observed across a number of probes (both increases and decreases in p-value), all sites remained significantly associated with maltreatment severity (p < 0.05), with the exception of two probes related to physical neglect (see **OS5**). A total of 45 sites (physical abuse: n = 26; sexual abuse: n = 7 and physical neglect: n = 12) also survived genome-wide correction (q < 0.05). Overall, associations with physical and sexual abuse were the least affected by unmeasured confounding, and those with physical neglect were the most affected.

Second, to examine the potential influence of outliers, we applied the winsorize function within the robustHD package to the DNAm data and reran epigenome-wide analyses, controlling for covariates. The winsorizing method uses censoring rather than exclusion, which is preferable with small sample sizes (Sheskin, 2003). Specifically, for each probe, values <5% or >95% of the score distribution were transformed to match the closest value within this percentile, which enables scores to maintain their relative weight without exerting an undue influence on the linear regression model. Although the vast majority of associations remained significant after winsorizing (97%; p<0.05), none of the transformed DMPs survived genome-wide correction (q>0.05; **OS5**). It is important to note, however, that this sensitivity analysis is highly conservative as it transforms, for each probe, scores at the tail ends of the distribution regardless of presence or absence of outliers. As such, this approach reduces the range of methylation scores across all probes, resulting in more limited variability.

Third, we performed bootstrapping as a complement to winsorizing using Mplus (version 6.1.1; Muthen & Muthen, 2011). Bootstrapping is advantageous with small samples as it derives an approximation of the sampling distribution via repeated resampling of the available data to yield bias-corrected 95% confidence intervals (CI). Associations were considered significant if bootstrapped 95% CIs (10,000 times) did not cross zero. The number of DMPs that survived bootstrapping were: n = 29 for physical abuse, n = 1 for sexual abuse, and n = 108 for physical neglect. Hence, sites associated with sexual abuse were the most affected by this sensitivity analysis, potentially reflecting the low number of individuals scoring high on this exposure, resulting in more extreme values.

Based on the above results, we compiled a list of high-confidence associations (see **OS5**), in order to highlight DMPs that were the most robustly implicated across the three sensitivity analyses and as such may be particularly promising candidates for prioritization in future studies. High-confidence associations were defined as having (i) an SVA q-value < 0.05, (ii) a winsorized *p*-value < 0.05, and (iii) significant bootstrapped 95% CIs. The resulting number of high-confidence associations for each exposure were: n = 20 for physical abuse, n = 1 for sexual abuse and n = 10 for physical neglect.

*************	Table 2************************************
*****	Figure 1************************************

*Variability in certain DNAm loci consistently associate with all maltreatment types* The top 20 hyper- and hypomethylated DMPs consistently associated with all three maltreatment types (i.e. physical abuse, sexual abuse and neglect) are shown in **Table 3**, ranked by average standardized effect size. The top-ranked hypermethylated probe, cg08796898 (**Figure 1C**), is located in *HUWE1*, an important regulator of neural proliferation linked to intellectual disability (Vandewalle *et al.*, 2013). Other genes annotated to hypermethylated probes include: (i) *CACNA2D4* (cg27516159), identified as a shared risk locus for multiple psychiatric disorders (Smollen *et al.*, 2013); (ii) *WBSCR17* (cg25579180), a gene widely expressed in the brain and associated with neurodevelopmental delay (Nakamura *et al.*, 2005); (iii) *LRP4* (cg12627354), involved in neuromuscular junction maintenance and acetylcholine signaling; and (iv) *GRB10* (cg26163537), an insulin modulator.

With regard to hypomethylated probes, we identified multiple markers annotated to genes that are involved in response to environmental stimuli, including: (i) two probes in *RPTOR* (cg07870603, cg09596252), a gene involved in cell growth regulation in response to climatic factors, nutrient and insulin levels (Sun *et al.*, 2010); and (ii) *GJD3* (cg22896075), which encodes a connexin thought to facilitate environmental adaptation by regulating physiological processes such as neuronal excitability (Belousov and Fontes, 2013). Another gene, *DNAJB6* (cg27496299; **Figure 1C**), has been found to reduce cellular toxicity and act as a molecular chaperone for neuronal proteins, including huntingtin (Mansson *et al.*, 2014). Finally, *TIAM2* (cg01786585) has been implicated in neurogenesis, particularly in the hippocampus, as well as promoting neural migration in the cerebral cortex (Chiu *et al.*, 1999).

### DNAm variation implicates both maltreatment-specific and shared functional pathways

As a final step, we examined enriched biological pathways for genes that were either (i) annotated to DMPs that associated with only *one* type of maltreatment (p<0.005); or (ii) annotated to DMPs that associated with *all* maltreatment types (p<0.05 across physical abuse, sexual abuse and neglect). Following these criteria, neglect showed 'unique' epigenetic variation in probes annotated to a larger number of genes (n =329) than physical (n =119) and sexual abuse (n = 47). A considerable proportion of genes showed epigenetic variation across all three maltreatment type (i.e. 'shared' genes; n = 2,348; consistent direction of effects).

Gene ontology analyses were performed for the above sets of genes (except sexual abuse, due to the low number of annotated genes), controlling for a range of potential confounds, including background probe distribution and gene size (**Figure 2**; see **OS1** for details). Results indicated that genes associated with DNAm variation specific to *physical abuse* were enriched for biological processes including cardiovascular function (e.g. cardiac muscle hypertrophy, heart rate), fear response, and wound healing (1.04E-12 <*p*<7.65E-03), while those associated with *neglect*-specific variation were enriched for regulation of cholesterol efflux, as well as processes related to cellular function and metabolism (2.91E-11 <*p*<1.04E-02; see **OS6** for a full list of terms). In contrast, genes associated with epigenetic variation *shared* across maltreatment types were significantly enriched for biological processes primarily related to regulation of nervous system development (e.g. neurogenesis, glial proliferation, neurotransmitter biosynthesis, oligodendrocyte development) and

organismal growth (e.g. organ morphogenesis, negative regulation of growth, response to growth factor; see **OS7** for a full list of terms; 4.20E-10<*p*< 1.44E-02).

### Discussion

This study is the first to characterize genome-wide DNAm patterns associated with different types of childhood abuse and neglect. Strengths include the availability of quantitative data on a range of different exposures, the inclusion of a sample of youth featuring high rates of adversity, and the analysis of methylome-wide data. Here, we highlight three key findings: (i) specific types of maltreatment, particularly physical exposures, are associated with DNAm variation at multiple loci; (ii) many of the identified loci are annotated to genes previously implicated in psychiatric and medical disorders; and (iii) gene ontology analyses indicate that, while maltreatment types show distinct patterns of methylomic variation, they also share a common epigenetic 'signature' enriched for biological processes related to neural development and organismal growth. The use of stringent sensitivity analyses further enabled us to identify high-confidence associations, which point to promising candidate loci for prioritization in future studies.

# Findings lend novel insights into the relationship between maltreatment and DNA methylation

Of the maltreatment types investigated, we found that physical exposures – including physical abuse, sexual abuse and physical neglect – all associated with epigenetic variation at multiple loci, with neglect showing the largest number of DMPs after genome-wide correction (despite having comparable prevalence rates to physical abuse). In contrast, no significant loci were identified for emotional abuse and emotional neglect (after multiple correction). This was unexpected given that emotional abuse, in particular, has been recognized as an important independent predictor of poor individual functioning (Rees, 2010). Consequently, more work is required to clarify whether results reflect a true lack of associations or whether other factors may be at play (e.g. challenges with the operationalization of emotional abuse; polyepigenetic effects of smaller magnitude; tissue specificity). It is also interesting to note that we identified an unexpectedly high number of associations between maltreatment types and DNAm sites located on the X-chromosome. Given that sex was controlled for in the analyses, these results suggest that maltreatment exposure may influence these DNAm sites similarly for boys and girls. Although sex differences were not examined in the present study due to sample size limitations, it will be of interest in future to test the potential role of sex in the relationship between maltreatment and DNAm.

Comparability with previous findings on maltreatment and DNAm is limited by the fact that, while other epigenetic studies have examined maltreatment as a global construct, we investigated DNA methylation profiles associated with specific forms of abuse and neglect. As such, a strict test of replication was not possible. Furthermore, past studies have varied widely on factors such as choice of tissue (e.g. saliva, blood or brain), DNAm platform (e.g.

Illumina 450k vs MeDIP), methodology (e.g. filtering of probes, covariates), maltreatment measure (e.g. self-report vs official records), population (e.g. community vs clinical) and sample characteristics (e.g. age, sex), making it difficult to assess our findings in the context of previous work in the area.

Nevertheless, we note here two of the most similar studies to ours, which have also used the Illumina 450k platform in peripheral tissues to investigate methylome-wide associations with childhood maltreatment. The first study (Yang et al. 2013) used DNA from saliva specimens to compare methylomic differences between a sample of maltreated and non-maltreated children, using a case-control design. The second study (Prados et al. 2015) used blood samples to compare DNAm patterns in adults with borderline personality disorder and exposure to high levels of childhood maltreatment vs adults with major depression disorder who had experienced low levels of childhood maltreatment (assessed using the same measure as our study, the CTQ). Comparing findings from our respective genome-wide analyses, we identified one overlapping DNAm site that was associated with maltreatment (after genome-wide correction) across both our study and Yang et al.'s (CILP2cg01487433), and no overlap with the DNAm sites reported in Prados et al's study. In addition, we identified several overlapping genes between studies (29 with Yang's et al.'s study; 1 with Prados et al.'s study) that had at least one genome-wide significant probe annotated to them. However, because these genes varied widely in DNAm probe coverage, it is unclear whether they were identified across studies because they robustly associate with maltreatment or because they may have had a larger number of DNAm probes annotated to them. As a whole, DMPs were found to differ between our study and previous ones. As such, it will be important in future to establish whether discrepancies may reflect the examination of global vs individual forms of maltreatment, methodological differences between studies or the presence of false positives, all issues that should be considered when interpreting the present findings.

We note that the use of sensitivity analyses in the present study (surrogate variable analysis, winsorizing and bootstrapping) enabled us to identify a subset of DMPs that were most robustly associated with maltreatment exposure (i.e. 'high-confidence associations'; n =31, 20% of total DMP set), and consequently may show promise as candidate loci for further investigation. Of interest, DMPs associated with physical neglect were most affected by the surrogate variable analysis, suggesting that the methylomic signature of this maltreatment type may in part reflect other sources of variability (e.g. genetic architecture, unmeasured exposures, confounding sources). In contrast, DMPs associated with sexual abuse were the most affected by the winsorizing and bootstrapping analyses, which may instead reflect the low number of individuals reporting high levels of this exposure. In general, associations between DNAm and physical abuse were the most robust, with 59% of DMPs (n = 20) classified as high-confidence. While these findings may be used in future to inform locusprioritization, it is important to note that the identification of high-confidence associations was based on a stringent set of criteria, so that the extent to which 'lower-confidence' DMPs may be robustly associated with maltreatment exposure will still need to be established via replication in independent samples.

# The identified DNA methylation markers support a molecular link between maltreatment and poor health outcomes

Our analyses identified multiple DMPs annotated to genes previously associated with psychiatric and medical disorders. The top-ranked DMP for physical abuse was located in the promoter region of *PSEN2*, a gene implicated in neurodegeneration and Alzheimer's disease (O'Brien and Wong, 2011). Although we are not aware of any study directly investigating the link between physical abuse and these conditions in advanced age, early life factors, including stress and the quality of the maternal relationship, have been associated with both neuropsychological impairment and dementia (Pechtel and Pizzagalli, 2011, Vaillant *et al.*, 2014). It will therefore be of interest to test whether *PSEN2* methylation mediates the effect of abuse on cognitive function and neurodegenerative risk. Maltreatment-associated DNAm changes were additionally identified in a number of other genes implicated in cognitive deficit and intellectual disability, including *HUWE1*, *WBSCR17* and *SMC1A* (Deardorff *et al.*, 2007, Nakamura *et al.*, 2005, Vandewalle *et al.*, 2013).

Findings also suggest a role of other loci previously implicated in psychopathology. Notably, one of the most hypermethylated markers across maltreatment types was located in CACNA2D4. Genetic variation within this gene has been identified as a shared risk locus for multiple psychiatric conditions that are associated with experience of childhood adversity, including attention deficit-hyperactivity disorder, bipolar disorder, major depressive disorder, and schizophrenia (Smollen et al., 2013). Interestingly, expression levels of this gene have been found to modulate neural hippocampal activity during emotional processing as well as prefrontal activity during executive tasks (Bigos et al., 2010) - functional patterns that are disrupted in maltreated individuals (Hart and Rubia, 2012). Furthermore, genetic variation in SYNJ2, another gene identified in our epigenome-wide analyses, has been linked to corpus callosum abnormalities (Edwards et al., 2014), which are robustly associated with maltreatment (McCrory et al., 2012). Other psychopathology-relevant genes included glutamate and GABA receptors (GRIND2D, GABBR1), which play a key role in excitatory and inhibitory neurotransmission, respectively (Kumar et al., 2013, Yamamoto et al., 2015). Prospective studies will be needed to explicitly test whether the DNAm sites identified in the present study associate with psychopathological outcomes and, if so, whether they may mediate the influence of abuse and neglect on later mental health.

# Gene ontology analyses contribute to a better understanding of biological pathways that may be affected by childhood abuse and neglect.

Gene ontology analyses were performed to explore biological pathways that may be uniquely affected by specific forms of maltreatment vs those 'shared' across maltreatment types. With regards to 'maltreatment-specific' pathways, we found that physical abuse was primarily associated with DNAm variation in the vicinity of genes enriched for cardiovascular processes, including regulation of heart rate and myocardial hypertrophy. This is consistent with evidence from epidemiological studies of elevated risk of heart disease among individuals who were physically abused as children, even after controlling for other childhood stressors, adult lifestyle factors and unhealthy behaviors (Fuller-Thomson *et al.*,

2010, Springer et al., 2007). Other enriched processes of interest related to wound healing, fear response and regulation of stress-activated protein kinase signaling. These pathways are in line with growing evidence from neuroendocrine and imaging studies of an association between physical abuse, HPA axis dysregulation and altered threat processing at neural and behavioral levels (Carpenter et al., 2011, McCrory et al., 2011, Turecki and Meaney, 2014). In contrast, the most enriched biological process specifically linked to neglect was regulation of cholesterol efflux, followed by other processes including DNA damage response, ribosomal function and zymogen activation. Although this finding necessitates replication, it is worth noting that these processes have been shown to respond to diet and nutrient availability (Fenech and Bonassi, 2011, Mizushima et al., 2004), which may be particularly compromised in youth who have experienced neglect (as evidenced in our sample). Together, our data suggest that physical abuse and neglect may affect epigenetic regulation of separate biological pathways, which may in part underlie differential effects observed at a phenotypic level (Sheridan and McLaughlin, 2014). Sexual abuse could not be examined due to the limited number of genes showing 'unique' epigenetic variation, which may reflect lack of power due to lower prevalence rates in our sample.

In addition to showing specific epigenetic variation, maltreatment types were also found to share a common methylomic signature, primarily enriched for processes related to neurodevelopment and organismal growth. This is consistent with a large body of evidence from animal and human studies documenting the impact of maltreatment and early life stress on brain structure, function and development. For example, maltreatment has been consistently associated with reduced volume of the corpus callosum, prefrontal cortex and hippocampus (Hart and Rubia, 2012, McCrory et al., 2012). Broader developmental delay and growth failure is also well established amongst maltreated children (Leslie et al., 2005, Olivan, 2003). Findings are also in line with previous epigenetic studies that have documented an association between global maltreatment exposure and DNAm changes in genes enriched for neural (e.g. Labonte et al, 2012: 'neuron projection', 'dendrite'; Yang et al, 2013: 'neurogenesis', 'axonal guidance') and developmental (e.g. Suderman et al, 2014: 'multicellular organismal development') processes. In future, it will be important to explore whether epigenetic regulation of these 'core' processes in response to maltreatment is functionally relevant at a transcriptomic level, and whether they relate directly to other biological markers of neurodevelopment and growth, such as imaging data and markers of cellular ageing.

### Limitations and future directions

The current findings should be interpreted in light of a number of important limitations. First, although rates of maltreatment in our sample were high, analyses were based on a modestly sized group of inner-city youth, which precluded the possibility of addressing more nuanced research questions (e.g. sex differences). Furthermore, our assessment of maltreatment was based on self-reports that lacked information regarding maltreatment timing and duration, both of which are likely to moderate the association between maltreatment and DNA methylation. In future, it will be important to replicate findings using larger samples, ideally

featuring externally-validated maltreatment histories that will make it possible to investigate how DNAm patterns may vary by age of onset and chronicity of exposure to different forms of maltreatment. Second, due to the high rates of polyvictimization in our sample, it was not feasible to compare methylomic patterns between individuals who experienced single forms of maltreatment in isolation. Instead, we examined methylome-wide associations for each maltreatment type dimensionally, and then explored the presence of unique vs shared signatures at a functional pathway level, as opposed to a DNAm site level. Even though associations were always strongest for the maltreatment type under investigation, it is therefore still possible that the identified sites may have in part reflected the combined effect of exposure to multiple maltreatment types. Furthermore, the identification of unique vs shared biological pathways linked to maltreatment were based on gene ontology analyses, which can be susceptible to bias (Timmons et al., 2015), and consequently will necessitate replication. Third, data on smoking status and medication use were not available in the present study. As these exposures have been shown to alter DNAm patterns (particularly in blood; Gao et al., 2015), it will be important to replicate findings controlling for these potential confounders. It is noteworthy, however, that none of the DNAm sites identified in the present study overlapped with those found to be robustly affected by smoking in a recent systematic review (i.e. DNAm sites associated with smoking in at least three independent reports; Gao et al., 2015). Fourth, findings were based on DNAm from buccal cell samples and can thus only be considered to represent biomarkers of exposure. Although buccal epithelial cells have been shown to converge more strongly with brain methylation patterns compared to other peripheral tissues (e.g. blood; Smith et al., 2015), further investigation will be needed to establish the relevance of our findings to the brain. Because we did not have access to RNA samples in the present study, we were also unable to establish the extent to which the identified DNAm sites associate with gene expression levels. As such, the analysis of transcriptomic data will be important for assessing the functional significance of the observed DNAm changes. Finally, the cross-sectional nature of the study meant that we were unable to establish the causal role of the identified DMPs. Longitudinal assessments featuring repeated DNAm measures will be needed to explore prospective interrelations between maltreatment exposure, DNAm and developmental outcomes.

### **Conclusions**

The current findings shed new light on how maltreatment may alter epigenetic mechanisms that regulate gene expression, providing a possible biological link between early adversity and poor health outcomes. Different forms of maltreatment were found to have distinct as well as shared signatures, pointing to a complex relationship between the nature of early adverse experience and multiple biological processes relevant for healthy normal development.

### References

- Baker-Andresen, D., Ratnu, V. S. & Bredy, T. W. 2013. Dynamic DNA methylation: a prime candidate for genomic metaplasticity and behavioral adaptation. Trends Neurosci 36, 3-13.
- Belousov, A. B. & Fontes, J. D. 2013. Neuronal gap junctions: making and breaking connections during development and injury. Trends Neurosci 36, 227-36.
- Bergman, Y. & Cedar, H. 2013. DNA methylation dynamics in health and disease. Nat Struct Mol Biol 20, 274-81.
- Bernstein, D. P. & Fink, L. 1998. Childhood trauma questionnaire: A retrospective self-report manual. San Antonio.
- Bigos, K. L., Mattay, V. S., Callicott, J. H., Straub, R. E., Vakkalanka, R., Kolachana, B., Hyde, T. M., Lipska, B. K., Kleinman, J. E. & Weinberger, D. R. 2010. Genetic variation in CACNA1C affects brain circuitries related to mental illness. Arch Gen Psychiatry 67, 939-45.
- Carpenter, L., Shattuck, T., Tyrka, A., Geracioti, T. & Price, L. 2011. Effect of childhood physical abuse on cortisol stress response. Psychopharmacology 214, 367-375.
- Cecil, C. A., Viding, E., Barker, E. D., Guiney, J. & McCrory, E. J. 2014. Double disadvantage: The influence of childhood maltreatment and community violence exposure on adolescent mental health. J Child Psychol Psychiatry.
- Chen, Y. A., Lemire, M., Choufani, S., Butcher, D. T., Grafodatskaya, D., Zanke, B. W., Gallinger, S., Hudson, T. J. & Weksberg, R. 2013. Discovery of cross-reactive probes and polymorphic CpGs in the Illumina Infinium HumanMethylation450 microarray. Epigenetics 8, 203-9.
- Chiu, C. Y., Leng, S., Martin, K. A., Kim, E., Gorman, S. & Duhl, D. M. 1999. Cloning and characterization of T-cell lymphoma invasion and metastasis 2 TIAM2, a novel guanine nucleotide exchange factor related to TIAM1. Genomics 61, 66-73.
- Cicchetti, D. & Toth, S. L. 2005. Child Maltreatment. Annual Review of Clinical Psychology 1, 409-438.
- Danese, A., Caspi, A., Williams, B., Ambler, A., Sugden, K., Mika, J., Werts, H., Freeman, J., Pariante, C. M., Moffitt, T. E. & Arseneault, L. 2011. Biological embedding of stress through inflammation processes in childhood. Mol Psychiatry 16, 244-6.
- Danese, A., Moffitt, T. E., Harrington, H., Milne, B. J., Polanczyk, G., Pariante, C. M., Poulton, R. & Caspi, A. 2009. Adverse childhood experiences and adult risk factors for age-related disease: depression, inflammation, and clustering of metabolic risk markers. Arch Pediatr Adolesc Med 163, 1135-43.
- Day, J. J., Childs, D., Guzman-Karlsson, M. C., Kibe, M., Moulden, J., Song, E., Tahir, A. & Sweatt, J. D. 2013. DNA methylation regulates associative reward learning. Nat Neurosci 16, 1445-52.
- Deardorff, M. A., Kaur, M., Yaeger, D., Rampuria, A., Korolev, S., Pie, J., Gil-Rodriguez, C., Arnedo, M., Loeys, B., Kline, A. D., Wilson, M., Lillquist, K., Siu, V., Ramos, F. J., Musio, A., Jackson, L. S., Dorsett, D. & Krantz, I. D. 2007. Mutations in cohesin complex members SMC3 and SMC1A cause a mild variant of cornelia de Lange syndrome with predominant mental retardation. Am J Hum Genet 80, 485-94.

- Edwards, T. J., Sherr, E. H., Barkovich, A. J. & Richards, L. J. 2014. Clinical, genetic and imaging findings identify new causes for corpus callosum development syndromes.
- Epa, W. R., Markovska, K. & Barrett, G. L. 2004. The p75 neurotrophin receptor enhances TrkA signalling by binding to Shc and augmenting its phosphorylation. J Neurochem 89, 344-53.
- Fenech, M. & Bonassi, S. 2011. The effect of age, gender, diet and lifestyle on DNA damage measured using micronucleus frequency in human peripheral blood lymphocytes. Mutagenesis 26, 43-9.
- Freeman, B., Smith, N., Curtis, C., Huckett, L., Mill, J. & Craig, I. W. 2003. DNA from buccal swabs recruited by mail: evaluation of storage effects on long-term stability and suitability for multiplex polymerase chain reaction genotyping. Behav Genet 33, 67-72.
- Fuller-Thomson, E., Brennenstuhl, S. & Frank, J. 2010. The association between childhood physical abuse and heart disease in adulthood: findings from a representative community sample. Child Abuse Negl 34, 689-98.
- Gao, X., Jia, M., Zhang, Y., Breitling, L. P., & Brenner, H. 2015. DNA methylation changes of whole blood cells in response to active smoking exposure in adults: a systematic review of DNA methylation studies. Clinical epigenetics, 7(1), 1.
- Hart, H. & Rubia, K. 2012. Neuroimaging of child abuse: a critical review. Front Hum Neurosci 6, 52.
- Houtepen, L. C., van Bergen, A. H., Vinkers, C. H., & Boks, M. P. 2016. DNA methylation signatures of mood stabilizers and antipsychotics in bipolar disorder. Epigenomics, 8(2), 197-208.
- Jaenisch, R. & Bird, A. 2003. Epigenetic regulation of gene expression: how the genome integrates intrinsic and environmental signals. Nat Genet 33 Suppl, 245-54.
- Johnson, W. E., Li, C. & Rabinovic, A. 2007. Adjusting batch effects in microarray expression data using empirical Bayes methods. Biostatistics 8, 118-27.
- Kim, S. T., Xu, B. & Kastan, M. B. 2002. Involvement of the cohesin protein, Smc1, in Atmdependent and independent responses to DNA damage. Genes Dev 16, 560-70.
- Klengel, T., Pape, J., Binder, E. B. & Mehta, D. 2014. The role of DNA methylation in stress-related psychiatric disorders. Neuropharmacology 80, 115-32.
- Kumar, K., Sharma, S., Kumar, P. & Deshmukh, R. 2013. Therapeutic potential of GABAB receptor ligands in drug addiction, anxiety, depression and other CNS disorders. Pharmacol Biochem Behav 110, 174-84.
- Labonte, B., Suderman, M., Maussion, G., Navaro, L., Yerko, V., Mahar, I., Bureau, A., Mechawar, N., Szyf, M., Meaney, M. J. & Turecki, G. 2012. Genome-wide epigenetic regulation by early-life trauma. Arch Gen Psychiatry 69, 722-31.
- Leek, J.T., Johnson, W.E., Parker, H.S., Jaffe, A.E. and Storey, J.D., 2012. The sva package for removing batch effects and other unwanted variation in high-throughput experiments. Bioinformatics 28, 882-883.
- Leslie, L. K., Gordon, J. N., Meneken, L. E. E., Premji, K., Michelmore, K. L. & Ganger, W. 2005. The Physical, Developmental, and Mental Health Needs of Young Children in Child Welfare by Initial Placement Type. Journal of developmental and behavioral pediatrics : JDBP 26, 177-185.

- Liang, L. & Cookson, W. O. C. 2014. Grasping nettles: cellular heterogeneity and other confounders in epigenome-wide association studies. Human Molecular Genetics 23, R83-R88.
- Lutz, P. E. & Turecki, G. 2014. DNA methylation and childhood maltreatment: from animal models to human studies. Neuroscience 264, 142-56.
- Mansson, C., Kakkar, V., Monsellier, E., Sourigues, Y., Harmark, J., Kampinga, H. H., Melki, R. & Emanuelsson, C. 2014. DNAJB6 is a peptide-binding chaperone which can suppress amyloid fibrillation of polyglutamine peptides at substoichiometric molar ratios. Cell Stress Chaperones 19, 227-39.
- McCrory, E., De Brito, S. A. & Viding, E. 2012. The link between child abuse and psychopathology: a review of neurobiological and genetic research. J R Soc Med 105, 151-6.
- McCrory, E. J., De Brito, S. A., Sebastian, C. L., Mechelli, A., Bird, G., Kelly, P. A. & Viding, E. 2011. Heightened neural reactivity to threat in child victims of family violence. Curr Biol 21, R947-8.
- McCrory, E. J. & Viding, E. 2015. The theory of latent vulnerability: Reconceptualizing the link between childhood maltreatment and psychiatric disorder. Dev Psychopathol 27, 493-505.
- Mizushima, N., Yamamoto, A., Matsui, M., Yoshimori, T. & Ohsumi, Y. 2004. In Vivo Analysis of Autophagy in Response to Nutrient Starvation Using Transgenic Mice Expressing a Fluorescent Autophagosome Marker. Molecular Biology of the Cell 15, 1101-1111.
- Montesinos, M. L., Castellano-Munoz, M., Garcia-Junco-Clemente, P. & Fernandez-Chacon, R. 2005. Recycling and EH domain proteins at the synapse. Brain Res Brain Res Rev 49, 416-28.
- Muthen LK, Muthen BO. MPLUS user's guide, 1998-2010 (6th edn). Los Angeles, CA: Muthen & Muthen; 2011.
- Nakamura, N., Toba, S., Hirai, M., Morishita, S., Mikami, T., Konishi, M., Itoh, N. & Kurosaka, A. 2005. Cloning and expression of a brain-specific putative UDP-GalNAc: polypeptide N-acetylgalactosaminyltransferase gene. Biol Pharm Bull 28, 429-33.
- O'Brien, R. J. & Wong, P. C. 2011. Amyloid Precursor Protein Processing and Alzheimer's Disease. Annual review of neuroscience 34, 185-204.
- Olivan, G. 2003. Catch-up growth assessment in long-term physically neglected and emotionally abused preschool age male children. Child Abuse Negl 27, 103-8.
- Pechtel, P. & Pizzagalli, D. A. 2011. Effects of early life stress on cognitive and affective function: an integrated review of human literature. Psychopharmacology Berl 214, 55-70.
- Pidsley, R., CC, Y. W., Volta, M., Lunnon, K., Mill, J. & Schalkwyk, L. C. 2013. A datadriven approach to preprocessing Illumina 450K methylation array data. BMC Genomics 14, 293.
- Prados, J., Stenz, L., Courtet, P., Prada, P., Nicastro, R., Adouan, W., Guillaume, S., Olie, E., Aubry, J. M., Dayer, A. & Perroud, N. 2015. Borderline personality disorder and childhood maltreatment: a genome-wide methylation analysis. Genes Brain Behav 14, 177-88.

- Price, E. Magda, Allison M. Cotton, Lucia L. Lam, Pau Farré, Eldon Emberly, Carolyn J.
  Brown, Wendy P. Robinson, and Michael S. Kobor. "Additional annotation enhances potential for biologically-relevant analysis of the Illumina Infinium HumanMethylation450 BeadChip array." Epigenetics & chromatin 6, no. 1 (2013): 1.
- Radford, L., Corral, S., Bradley, C., Fisher, H., Bassett, C., Howatt, N. & Collishaw, S. 2011. Child abuse and neglect in the UK today. NSPCC.
- Rees, C. A. 2010. Understanding emotional abuse. Archives of Disease in Childhood 95, 59-67.
- Roffe, M., Hajj, G. N., Azevedo, H. F., Alves, V. S. & Castilho, B. A. 2013. IMPACT is a developmentally regulated protein in neurons that opposes the eukaryotic initiation factor 2alpha kinase GCN2 in the modulation of neurite outgrowth. J Biol Chem 288, 10860-9.
- Shalev, I., Moffitt, T. E., Sugden, K., Williams, B., Houts, R. M., Danese, A., Mill, J., Arseneault, L. & Caspi, A. 2013. Exposure to violence during childhood is associated with telomere erosion from 5 to 10 years of age: a longitudinal study. Mol Psychiatry 18, 576-81.
- Sheridan, M. A. & McLaughlin, K. A. 2014. Dimensions of early experience and neural development: deprivation and threat. Trends Cogn Sci 18, 580-5.
- Sheskin, David J. Handbook of parametric and nonparametric statistical procedures. crc Press, 2003.
- Smith, A. K., Kilaru, V., Klengel, T., Mercer, K. B., Bradley, B., Conneely, K. N., Ressler, K. J. & Binder, E. B. 2015. DNA extracted from saliva for methylation studies of psychiatric traits: evidence tissue specificity and relatedness to brain. Am J Med Genet B Neuropsychiatr Genet 168B, 36-44.
- Smoller, JW, Craddock, N, Kendler, K, Lee, PH, Neale, BM, Nurnberger, JI, Ripke, S, Santangelo, S, & Sullivan, PF 2013. Identification of risk loci with shared effects on five major psychiatric disorders: a genome-wide analysis. Lancet 381, 1371-9.
- Springer, K. W., Sheridan, J., Kuo, D. & Carnes, M. 2007. Long-term physical and mental health consequences of childhood physical abuse: results from a large population-based sample of men and women. Child Abuse Negl 31, 517-30.
- Suderman, M., Borghol, N., Pappas, J. J., Pinto Pereira, S. M., Pembrey, M., Hertzman, C., Power, C. & Szyf, M. 2014. Childhood abuse is associated with methylation of multiple loci in adult DNA. BMC Med Genomics 7, 13.
- Sun, C., Southard, C., Witonsky, D. B., Kittler, R. & Di Rienzo, A. 2010. Allele-specific down-regulation of RPTOR expression induced by retinoids contributes to climate adaptations. PLoS Genet 6, e1001178.
- Timmons, J. A., Szkop, K. J. & Gallagher, I. J. 2015. Multiple sources of bias confound functional enrichment analysis of global -omics data. Genome Biol 16, 186.
- Traynelis, S. F., Wollmuth, L. P., McBain, C. J., Menniti, F. S., Vance, K. M., Ogden, K. K., Hansen, K. B., Yuan, H., Myers, S. J. & Dingledine, R. 2010. Glutamate receptor ion channels: structure, regulation, and function. Pharmacol Rev 62, 405-96.
- Tsai, P.C. and Bell, J.T., 2015. Power and sample size estimation for epigenome-wide association scans to detect differential DNA methylation. International journal of epidemiology, 44(4), pp.1429-1441.

- Turecki, G. & Meaney, M. J. 2014. Effects of the Social Environment and Stress on Glucocorticoid Receptor Gene Methylation: A Systematic Review. Biol Psychiatry.
- Vaillant, G. E., Okereke, O. I., Mukamal, K. & Waldinger, R. J. 2014. Antecedents of intact cognition and dementia at age 90 years: a prospective study. Int J Geriatr Psychiatry 29, 1278-85.
- Vandewalle, J., Langen, M., Zschaetzsch, M., Nijhof, B., Kramer, J. M., Brems, H., Bauters, M., Lauwers, E., Srahna, M., Marynen, P., Verstreken, P., Schenck, A., Hassan, B. A. & Froyen, G. 2013. Ubiquitin Ligase HUWE1 Regulates Axon Branching through the Wnt/β-Catenin Pathway in a <italic>Drosophila</italic> Model for Intellectual Disability. PLoS ONE 8, e81791.
- World Health Organization. 2014. Global status report on violence prevention 2014. Geneva: World Health Organization.
- Yamamoto, H., Hagino, Y., Kasai, S. & Ikeda, K. 2015. Specific Roles of NMDA Receptor Subunits in Mental Disorders. Current Molecular Medicine 15, 193-205.
- Yang, B.Z., Zhang, H., Ge, W., Weder, N., Douglas-Palumberi, H., Perepletchikova, F., Gelernter, J. and Kaufman, J. 2013. Child abuse and epigenetic mechanisms of disease risk. American journal of preventive medicine, 44(2),101-107.

### **Figure legends:**

### Figure 1. Associations between DNA methylation and maltreatment types.

(A) Manhattan plots for physical abuse, sexual abuse and physical neglect; (B) scatterplots of DMPs indicated in (A) for each maltreatment type; and (C) scatterplots of hyper- and hypomethylated DMPs *across* maltreatment types. *N.b.* The dotted line represents FDR correction (i.e. DMPs above the line are significant at q<0.05).

### Figure 2. Unique vs shared enriched biological processes across maltreatment types.

Significantly enriched biological processes for genes uniquely associated with each maltreatment type (physical abuse [green], physical neglect [blue]) vs those shared across maltreatment types (red), based on GO analysis. GO analysis was not run independently for sexual abuse (SA) due to limited gene *n*. Circles represent GO terms that survive FDR correction. The X axis represents  $-\log(10)$  p values. The opacity of the circles indicates level of significance (darker = more significant). The size of the circles indicates the percentage of genes in our results for a given pathway compared to the total number of genes in the same pathway (i.e. larger size = larger %; range = 6.51% - 100%). *Abbreviations*: PA, physical abuse; PN, physical neglect; SA, sexual abuse.

	-			-	-	-	-		-	Ethnicity <sup>b</sup>		icity <sup>b</sup>	
Ма	ltreatment type	M (SD)	% above threshold ( <i>n</i> )	1	2	3	4	Sex	Age	White	Black	Mixed	Asian
1.	Emotional abuse	10.06 (4.99)	50.8 (63)	-				.01	.17	13	.19*	07	.00
2.	Physical abuse	7.82 (4.70)	33.1 (41)	.57***	_			.01	.09	16	.26**	08	06
3.	Sexual abuse	6.14 (3.51)	17.0 (21)	.41***	.25***	_		.05	.13	11	.12	01	04
4.	Emotional neglect	10.47 (4.81)	49.2 (61)	.73***	.55***	.35***	_	03	.18*	20*	.21*	09	.12
5.	Physical neglect	7.55 (3.61)	32.3 (40)	.65***	.60***	.36***	.71***	07	.14	22*	.31***	11	.00

**Table 1.** Descriptives and correlations between study variables.

*N.B.* Intercorrelations between maltreatment types, and their association with socio-demographic characteristics (age, sex and ethnicity). A product-moment correlation is used for associations between two continuous variables, while a point-biserial correlation is used for associations between one continuous and one dichotomous variable (i.e. sex, ethnicity). Bivariate correlations significant at: \*p < .05, \*\*p < .01, \*\*\*p < .001. <sup>a</sup> For descriptive purposes, this column shows the percentage of participants who scored above the 'Low' maltreatment threshold specified by the CTQ manual.

<sup>b</sup> Each ethnic group coded as 1 = yes and 0 = no, based on self-reported ethnicity.

CpG probe	Gene	Chr	Genomic location	Position	B diff. <sup>a</sup>	Std. $B^b$	<b>P-value</b>	FDR (q)		
A. Physical a	buse (N FDR-correcte	d DMPs	s = 34)							
cg20000641	PSEN2	1	TSS1500	227058046	0.14	0.55	2.53E-11	1.47E-05		
cg02353937	SMC1A/RIBC1	Х	TSS1500	53449152	0.09	0.50	2.90E-09	7.89E-04		
cg15440363	GPD1L	3	TSS200	32148023	0.05	0.48	2.06E-08	3.64E-03		
cg04412054	GALNS	16	Body	88897539	-0.12	-0.46	5.69E-08	0.01		
cg22311608	SMC1A/RIBC1	Х	TSS200	53449829	0.09	0.46	8.45E-08	0.01		
cg24365098	C11orf84	11	3'UTR	63594804	-0.12	-0.45	1.43E-07	0.01		
cg03561071	[FAT3]	11		91844355	-0.11	-0.44	4.28E-07	0.03		
cg26454299	PPP3CA	4		102268957	0.10	0.43	5.58E-07	0.03		
cg25047485	[ARHGAP39]	8		145848944	-0.11	-0.43	8.14E-07	0.04		
cg04524770	LMF1	16	Body	946408	-0.11	-0.42	9.96E-07	0.04		
B. Sexual abuse ( <i>N</i> FDR-corrected DMPs = 7)										
cg17106653	GRIN2D	19	TSS1500	48897279	0.22	0.50	4.16E-09	1.72E-03		
cg00974464	PRDM15	21	Body	43254115	-0.12	-0.46	6.02E-08	0.01		
cg26528551	MGMT	10	Body	131445415	-0.09	-0.46	6.84E-08	0.01		
cg10795666	MLNR	13	1stExon	49794635	0.09	0.45	2.23E-07	0.02		
cg02618355	MYOM2	8	Body	2024368	-0.09	-0.44	2.88E-07	0.02		
cg26513050	DIAPH2/RPA4	Х	Body	96138983	-0.18	-0.44	2.91E-07	0.02		
cg23983710	DIP2C	10	Body	370756	-0.16	-0.43	5.83E-07	0.03		
C. Physical n	eglect (N FDR-correct	ted DMI	Ps = 118)							
cg00691266	EVPL	17	Body	74015089	-0.08	-0.46	1.01E-07	0.01		
cg27083825	SYNJ2	6	Body	158453594	-0.07	-0.45	1.02E-07	0.01		
cg17918089	SETDB1	1	Body	150899251	0.08	0.45	1.14E-07	0.01		
cg10094509	MAP2K4P1/CHIC1	Х	TSS1500	72783579	0.09	0.45	1.16E-07	0.01		
cg09863040	JADE1	4	TSS200	129730729	0.02	0.44	2.10E-07	0.01		
cg08796898	HUWE1	Х	5'UTR	53713664	0.14	0.44	3.59E-07	0.02		

**Table 2.** Top 10 DMPs associated with physical abuse, sexual abuse and physical neglect.

cg20668974	ZNF827	4	Body	146857981	0.06	0.44	3.63E-07	0.02
cg18116160	GABBR1	6	Body	29575145	-0.04	-0.44	4.01E-07	0.02
cg09084892	FAT1	4	Body	187557837	-0.12	-0.43	4.35E-07	0.02
cg08054907	HIST1H1A	6	TSS1500	26019358	-0.09	-0.43	5.12E-07	0.02

*N.B.* Gene names in brackets indicate the most proximal genes to the CpG probe based on Genome Studio. Genes with multiple significant probes are highlighted in blue.

<sup>a</sup> Beta differences indicate the overall difference (% methylation change) between the predicted unstandardized minimum and maximum values of the linear model.

<sup>b</sup> Standardized Beta estimates are used as a measure of effect size, where an effect of 0.10 is small effect, an effect of 0.24 is a medium effect, and an effect of 0.37 is a large effect.

CpG probe	Gene	Chr	Genomic	Position	Physical Abuse		Sexu	Sexual Abuse		Neglect		
			location		Std <b>B</b>	<i>p</i> -value	Std <b>B</b>	<i>p</i> -value	Std <b>B</b>	<i>p</i> -value	<b>StdB</b>	
A. Hypermethyla	ited											
cg08796898	HUWE1	Х	5'UTR	53713664	0.38	1.10E-05	0.21	2.19E-02	0.44	3.83E-07	0.34	
cg21239691	GEMIN8	Х	TSS200	14048191	0.42	1.24E-06	0.18	4.38E-02	0.39	7.20E-06	0.33	
cg04704856	FMOD	1	1stExon	203320190	0.32	2.46E-04	0.27	2.61E-03	0.37	2.36E-05	0.32	
cg12986338	[PDE1C]	7		32338950	0.28	1.39E-03	0.31	4.15E-04	0.35	6.27E-05	0.32	
cg23436576	TDH	8	Body	11204132	0.35	7.16E-05	0.23	8.75E-03	0.35	6.17E-05	0.31	
cg25051341	PRDM13	6	Body	100061307	0.27	2.79E-03	0.28	1.40E-03	0.38	1.30E-05	0.31	
cg19637330	[PAX7]	1		19110922	0.26	3.73E-03	0.31	4.51E-04	0.36	3.98E-05	0.31	
cg26163537	GRB10	7	TSS1500	50861592	0.36	3.84E-05	0.19	3.76E-02	0.38	1.37E-05	0.31	
cg25579180	WBSCR17	7	Body	71098623	0.36	4.53E-05	0.32	3.03E-04	0.25	5.24E-03	0.31	
cg12627354	LRP4	11	TSS1500	46940434	0.26	3.29E-03	0.28	1.56E-03	0.38	1.67E-05	0.31	
cg08121755	[KRT80]	12		52545978	0.27	2.15E-03	0.34	1.39E-04	0.29	9.09E-04	0.30	
cg03309770	TVP23A	16	1stExon	10912478	0.38	1.17E-05	0.21	1.66E-02	0.30	6.53E-04	0.30	
cg21453209	[ZFAND1]	8		82635831	0.32	3.47E-04	0.25	5.99E-03	0.33	2.23E-04	0.30	
cg20594607	LOC338799	12	TSS200	122241438	0.33	1.94E-04	0.24	7.13E-03	0.31	3.81E-04	0.29	
cg26840590	PHF1	6	5'UTR	33379330	0.32	2.86E-04	0.22	1.23E-02	0.34	1.39E-04	0.29	
cg04530860	B9D2	19	TSS200	41870213	0.38	1.69E-05	0.24	7.20E-03	0.26	3.45E-03	0.29	
cg17962547	GPR123	10	Body	134918974	0.32	3.57E-04	0.25	5.39E-03	0.31	4.39E-04	0.29	
cg11480019	ADK	10		75936982	0.30	8.66E-04	0.32	3.13E-04	0.25	4.48E-03	0.29	
cg09387749	HOXD3	2	TSS200	177028680	0.31	5.34E-04	0.24	7.14E-03	0.32	3.50E-04	0.29	
cg27516159	CACNA2D4	12	Body	1904847	0.28	1.81E-03	0.22	1.27E-02	0.35	6.03E-05	0.28	

**Table 3.** Top 20 hyper- and hypo-methylated DMPs across maltreatment types (physical abuse, sexual abuse and neglect), ranked by average effect size.

B. Hypomethylat	. Hypomethylated												
cg10494397	CC2D2A	4	Body	15593854	-0.39	7.89E-06	-0.24	6.61E-03	-0.42	1.35E-06	-0.35		
cg27496299	DNAJB6	7	Body	157171202	-0.38	1.56E-05	-0.28	1.84E-03	-0.39	6.90E-06	-0.35		
cg27632471	C20orf96	20	Body	259123	-0.27	2.41E-03	-0.39	8.78E-06	-0.39	9.42E-06	-0.35		
cg07870603	RPTOR	17	Body	78880144	-0.28	1.37E-03	-0.38	1.43E-05	-0.33	2.28E-04	-0.33		
cg09596252	RPTOR	17	Body	78655493	-0.41	3.00E-06	-0.28	1.62E-03	-0.30	7.06E-04	-0.33		
cg01786585	TIAM2	6		155315188	-0.37	2.00E-05	-0.35	8.70E-05	-0.26	3.59E-03	-0.33		
cg23523534	NLRP11	19	5'UTR	56347927	-0.33	1.45E-04	-0.32	3.12E-04	-0.32	3.59E-04	-0.32		
cg16560389	[TCERG1L]	10		133318201	-0.34	9.50E-05	-0.22	1.32E-02	-0.40	3.60E-06	-0.32		
cg05157878	GCNT2	6		10494860	-0.29	1.05E-03	-0.38	1.57E-05	-0.29	1.30E-03	-0.32		
cg21473728	MYOM2	8	Body	2031651	-0.36	4.62E-05	-0.20	2.57E-02	-0.39	7.42E-06	-0.32		
cg21385432	EP400	12	Body	132512858	-0.31	4.33E-04	-0.24	6.55E-03	-0.39	7.04E-06	-0.32		
cg19279265	ACOX3	4	Body	8388670	-0.27	2.83E-03	-0.41	1.73E-06	-0.26	3.66E-03	-0.31		
cg03561071	[FAT3]	11		91844355	-0.44	4.28E-07	-0.22	1.35E-02	-0.28	1.99E-03	-0.31		
cg22896075	GJD3	17	1stExon	38518413	-0.40	5.30E-06	-0.21	2.04E-02	-0.32	3.09E-04	-0.31		
cg13521944	C22orf9	22	Body	45596948	-0.27	2.45E-03	-0.28	1.69E-03	-0.37	2.97E-05	-0.30		
cg11650926	[KIAA0947]	5		5568539	-0.30	8.71E-04	-0.19	3.40E-02	-0.43	6.91E-07	-0.30		
cg21076890	COL4A2	13	Body	110965662	-0.30	8.61E-04	-0.19	3.93E-02	-0.43	5.51E-07	-0.30		
cg03500617	[FA2H]	16		74812570	-0.31	4.27E-04	-0.19	3.11E-02	-0.41	2.86E-06	-0.30		
cg02866700	CARS2	13	Body	111333333	-0.30	7.85E-04	-0.26	3.98E-03	-0.35	5.79E-05	-0.30		
cg13920529	SUGT1P1	9	Body	33402426	-0.34	9.08E-05	-0.21	1.85E-02	-0.35	6.28E-05	-0.30		

*N.B.* Gene names in brackets indicate the most proximal genes to the CpG probe based on Genome Studio. Genes with multiple significant probes are highlighted in blue.





### Figure 2.



### **Supplementary Information**

Epigenetic signatures of childhood abuse and neglect: Implications for psychiatric

### List of supplementary materials:

- **OS1.** Optimized gene ontology (GO) method.
- **OS2.** Top 10 DMPs associated with emotional abuse and emotional neglect.
- **OS3.** Full list of FDR-corrected loci associated with physical abuse, sexual abuse and physical neglect.
- **OS4**. Quantile-quantile plots for each maltreatment type.
- **OS5.** Sensitivity analyses for FDR-corrected DMPs associated with physical abuse, sexual abuse and physical neglect.
- **OS6.** Enriched biological processes for genes associated with epigenetic variation that is unique to physical abuse and physical neglect.
- **OS7.** Enriched biological processes for genes associated with epigenetic variation that is shared across maltreatment types (physical abuse, sexual abuse and physical neglect).

### OS1. Optimized gene ontology (GO) method

A logistic regression approach was used to test if genes in the test list predicted pathway membership while controlling for the number probes annotated to each gene. Pathways were downloaded from the Gene Ontology website and all genes annotated to parent terms were also included. Illumina UCSC gene annotation was used to create a test gene list from probes that shown unique vs shared associations with maltreatment, depending on the analysis performed. All genes with at least one methylation probe annotated and annotated to at least one GO pathway were considered. Pathways were filtered to those with between 10 and 2000 genes in. After applying this method to all pathways, significant pathways (p < 0.05) were taken and grouped where overlapping genes explained the signal. This was achieved by taking the most significant pathway, and retesting all remaining significant pathways while controlling additionally for the best term. If genes in the test list no longer predicted the pathway, the term was said to be explained by the most significant pathway, and hence these pathways were grouped together. This algorithm was repeated, taking the next most significant term, until all pathways had been considered as the most significant or found to be explained by a more significant term. GO terms were interpreted exclusively if they if they contained more than 1 gene and passed FDR-correction, based on the final list of independent (non-redundant) terms.

## OS2. Top 10 DMPs associated with emotional abuse and emotional neglect

CpG probe	Gene	Chr	Genomic location	Position	Std .B	P-value	FDR (q)
A. Emotional	abuse						
cg23436576	TDH	8	Body	11204132	0.38	1.17E-05	1.00
cg10933494	[ZNF311]	6		28983143	-0.38	1.56E-05	1.00
cg00742603	DBNDD1	16	TSS1500	90086301	0.37	1.89E-05	1.00
cg00347856	[FLJ45974]	7		53879253	-0.37	1.95E-05	1.00
cg18116160	GABBR1	6	Body	29575145	-0.37	2.48E-05	1.00
cg11496014	[BUB3]	10		125115762	-0.36	3.43E-05	1.00
cg00436603	CYP2E1	10	TSS200	135340740	0.36	3.87E-05	1.00
cg13778848	LOC283050	10	Body	80746999	-0.35	6.40E-05	1.00
cg04477202	SPAST	2	Body	32289458	0.35	7.76E-05	1.00
cg01067603	DAB1	1	5'UTR	58331989	-0.35	7.78E-05	1.00
B. Emotional	neglect						
cg21964928	[SAP30L]	5		153872958	0.38	1.62E-05	1.00
cg19050351	IL1F5	2	Body	113820090	-0.37	2.48E-05	1.00
cg01630869	CDC42EP2	11		65082120	0.36	3.18E-05	1.00
cg10933494	[ZNF311]	6		28983143	-0.36	3.31E-05	1.00
cg07603484	NUP62	19	TSS1500	50434179	-0.35	5.99E-05	1.00
cg18994566	ZFYVE21	14	Body	104191347	-0.35	6.71E-05	1.00
cg13161621	IQSEC2	Х	Body	53348705	0.35	6.81E-05	1.00
cg13688202	SPTB	14	Body	65230287	-0.35	8.06E-05	1.00
cg09796576	ZNF526	19	Body	42729528	-0.35	8.28E-05	1.00
cg18852899	RMDN2	2		38154797	-0.35	8.33E-05	1.00

N.B. Gene names in brackets indicate the most proximal genes to the CpG probe based on Genome Studio. Standardized Beta estimates are used as a measure of effect size, where an effect of 0.10 is small effect, an effect of 0.24 is a medium effect, and an effect of 0.37 is a large effect.

OS3. Full list of FDR-corrected DMPs associated with physical abuse, sexual abuse and physical neglect

A. Physical abuse (FDR <i>s</i> = 34)           cg20000641         PSEN2         1         TSS1500         227058046         0.55         2.53E-11         1.47E-05           cg02353937         SMCLA/RIBC1         X         TSS1200         52148023         0.048         2.00E-09         7.89E-04           cg15440363         GPD1L         3         TSS200         52148023         0.48         2.00E-08         3.64E-03           cg0412054         GALNS         16         Body         88897539         -0.46         5.69E-08         0.01           cg234365098         C11or/84         11         3'UTR         63594804         -0.45         8.43E-07         0.03           cg24654299         PPP3CA         4          102268957         0.43         5.58E-07         0.03           cg1645633          8          H45848944         -0.43         8.14E-07         0.04           cg1635603         -LMFI         16         Body         451261         -0.42         1.01E-06         0.04           cg1635633          7138640         0.42         1.01E-06         0.04           cg1635029         FBMV12         3         TSS200         14048191	CpG probe	Gene	Chr	Genomic location	Position	Std.B	P-value	FDR (q)
cg20000641         PSEN2         1         TSS1500         227058046         0.55         2.53E-11         1.47E-05           cg03353937         SMCLA/RIBCI         X         TSS1500         53449152         0.50         2.90E-09         7.89E-04           cg15440363         GPD1L         3         TSS200         32148023         0.48         2.00E-08         3.64E-03           cg0412024         GALNS         16         Body         88897539         -0.46         5.69E-08         0.01           cg23511608         SMCLA/RIBCI         X         TSS200         53449829         0.46         8.45E-08         0.01           cg23507485          11          91844355         -0.44         2.8E07         0.03           cg25047485          8          145848944         -0.43         8.14E-07         0.04           cg19736040         SHC2         19         Body         451261         -0.42         9.96E-07         0.04           cg13390589         SAEI         19         TSS200         47634047         0.42         1.37E-06         0.04           cg0330589         SAEI         19         TSS200         47634047         0.42	A. Physical abu	use (FDR $n = 34$ )						
cg02353937         SMCIA/RIBCI         X         TSS1500         53449152         0.50         2.90E-09         7.89E-04           cg15440363         GPD1L         3         TSS200         32148023         0.48         2.06E-08         3.64E-03           cg0412054         GALNS         16         Body         88897539         -0.46         5.69E-08         0.01           cg2311608         SMCLA/RIBCI         X         TSS200         53449829         0.46         8.45E-08         0.01           cg234565098         C/Lor/84         11         3'UTR         63594804         -0.45         1.43E-07         0.03           cg0454270         LMFI         16         Body         946408         -0.42         9.06E-07         0.04           cg04524770         LMFI         16         Body         946408         -0.42         9.06E-07         0.04           cg19736040         SHC2         19         Body         451261         -0.42         1.14E-06         0.04           cg10390589         SAEI         19         TSS200         47634047         0.42         1.37E-06         0.04           cg0315329         IMPACT         18         Body         22006989         0.41 <td>cg20000641</td> <td>PSEN2</td> <td>1</td> <td>TSS1500</td> <td>227058046</td> <td>0.55</td> <td>2.53E-11</td> <td>1.47E-05</td>	cg20000641	PSEN2	1	TSS1500	227058046	0.55	2.53E-11	1.47E-05
cg15440363         GPD1L         3         TSS200         32148023         0.48         2.06E-08         3.64E-03           cg04112054         GALNS         16         Body         88897539         -0.46         8.45E-08         0.01           cg23151608         SMC1A/RIBC1         X         TSS200         53449829         0.46         8.45E-08         0.01           cg03561071          11          91844355         -0.44         4.28E-07         0.03           cg03561071          8          145848944         -0.43         5.58E-07         0.03           cg1645823          8          145848944         -0.43         8.14E-07         0.04           cg16356833          3          87138640         0.42         1.01E-06         0.04           cg1639589         SAE1         19         TSS200         44048191         0.42         1.24E-06         0.04           cg16193029         FBXW12         3         TSS200         44613550         -0.42         1.60E-06         0.04           cg03402073         PMM1         22         Body         22006989         0.41         1.74E-06	cg02353937	SMC1A/RIBC1	Х	TSS1500	53449152	0.50	2.90E-09	7.89E-04
cg04412054         GALNS         16         Body         88897539         -0.46         5.69E-08         0.01           cg23151608         SMCLARIBCT         X         TSS200         53449829         0.46         5.69E-08         0.01           cg2455098         C11or/84         11         3'UTR         63594804         -0.45         1.43E-07         0.01           cg26451299         PPP3CA         4          102268957         0.43         5.58E-07         0.03           cg26452470         LMF1         16         Body         946408         -0.42         1.96E-07         0.04           cg19736040         SHC2         19         Body         451261         -0.42         1.14E-06         0.04           cg19736040         SHC2         19         Body         451261         -0.42         1.37E-06         0.04           cg10390589         SAE1         19         TSS200         47634047         0.42         1.50E-06         0.04           cg16193029         FBXW12         3         TSS200         4481355         -0.41         1.70E-06         0.04           cg0331329         IMPACT         18         Body         22006989         0.41         1.	cg15440363	<i>GPD1L</i>	3	TSS200	32148023	0.48	2.06E-08	3.64E-03
cg22311608       SMCLARIBCI       X       TSS200 $53449829$ $0.46$ $8.45E.08$ $0.01$ cg234365098       CHorf84       11       3'UTR $63594804$ $0.45$ $1.43E.07$ $0.01$ cg26454299       PP3CA       4 $102268957$ $0.43$ $5.58E.07$ $0.03$ cg25047485 $145848944$ $-0.43$ $8.14E.07$ $0.04$ cg164524770       LMF1       16       Body       946408 $-0.42$ $9.96E.07$ $0.04$ cg19736040       SHC2       19       Body $451261$ $-0.42$ $1.14E.06$ $0.04$ cg10390589       SAEI       19       TSS200 $47634047$ $0.42$ $1.37E.06$ $0.04$ cg034530790        5 $150185568$ $-0.42$ $1.50E.06$ $0.04$ cg08479390        5 $1951736$ $-0.41$ $1.70E.06$ $0.04$ cg08479390        19 $13979907$ $0.41$ $1.74E.06$ $0.04$ cg08479390        19       -	cg04412054	GALNS	16	Body	88897539	-0.46	5.69E-08	0.01
cg24365098       CHorf84       11       3'UTR       63594804       -0.45       1.43E-07       0.01         cg03561071        11        91844355       -0.44       4.28E-07       0.03         cg26047485        8        102268957       0.43       5.58E-07       0.04         cg16856833        8        145848944       -0.43       8.14E-07       0.04         cg16856833        3        87138640       0.42       1.01E-06       0.04         cg16390589       SAEI       19       Body       451261       -0.42       1.4E-06       0.04         cg10390589       SAEI       19       TSS200       14048191       0.42       1.24E-06       0.04         cg16193029       FBXW12       3       TSS200       48413550       -0.42       1.60E-06       0.04         cg1639309        19       TSS200       48413550       -0.41       1.70E-06       0.04         cg1639309        19        1951736       -0.41       1.95E-06       0.04         cg03013329       IMPACT       18       Body       183226025       -0.4	cg22311608	SMC1A/RIBC1	Х	TSS200	53449829	0.46	8.45E-08	0.01
cg03561071        11        91844355       -0.44       4.28E-07       0.03         cg26454299       PPP3CA       4        102268957       0.43       5.58E-07       0.03         cg25047485        8        145848944       0.43       8.14E-07       0.04         cg04524770       LMF1       16       Body       946408       -0.42       9.96E-07       0.04         cg16736040       SHC2       19       Body       451261       -0.42       1.14E-06       0.04         cg19736040       SHC2       19       Body       451261       -0.42       1.37E-06       0.04         cg10390589       SAE1       19       TSS200       47634047       0.42       1.50E-06       0.04         cg035090        1        150185568       -0.42       1.60E-06       0.04         cg033013329       IMPACT       18       Body       2206989       0.41       1.74E-06       0.04         cg03402073       PMM1       22       Body       183226025       -0.41       1.95E-06       0.04         cg03402073       PMM1       22       Body       183226025       -0.41	cg24365098	C11orf84	11	3'UTR	63594804	-0.45	1.43E-07	0.01
cg26454299         PPP3CA         4          102268957         0.43         5.58E-07         0.03           cg25047485          8          145848944         -0.43         8.14E-07         0.04           cg16856833          3          87138640         0.42         9.96E-07         0.04           cg16356833          3          87138640         0.42         1.01E-06         0.04           cg10390589         SHC2         19         Body         451261         -0.42         1.14E-06         0.04           cg10390589         SAE1         19         TSS200         47634047         0.42         1.37E-06         0.04           cg16193029         FBXW12         3         TSS200         48413550         -0.42         1.60E-06         0.04           cg03301329         IMPACT         18         Body         22006989         -0.41         1.74E-06         0.04           cg03402073         PMM1         22         Body         183226025         -0.41         1.80E-06         0.04           cg10370025         ABR         17          926264         -0.41         2.12E-06         0	cg03561071		11		91844355	-0.44	4.28E-07	0.03
cg25047485        8        145848944       -0.43       8.14E-07       0.04         cg04524770       LMF1       16       Body       946408       -0.42       9.96E-07       0.04         cg16856833        3        87138640       0.42       1.01E-06       0.04         cg19736040       SHC2       19       Body       451261       -0.42       1.14E-06       0.04         cg10390589       SAE1       19       TSS200       14048191       0.42       1.24E-06       0.04         cg10390589       SAE1       19       TSS200       47634047       0.42       1.50E-06       0.04         cg03250090        1        150185568       -0.42       1.60E-06       0.04         cg03453728        19        1951736       -0.41       1.70E-06       0.04         cg034031329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg03402073       PMM1       22       Body       14974480       -0.41       1.80E-06       0.04         cg10170025       ABR       17        926264       -0.41	cg26454299	<i>РРРЗСА</i>	4		102268957	0.43	5.58E-07	0.03
cg04524770       LMF1       16       Body       946408       -0.42       9.96E-07       0.04         cg16856833        3        87138640       0.42       1.01E-06       0.04         cg19736040       SHC2       19       Body       451261       -0.42       1.14E-06       0.04         cg10390589       SAEI       19       TSS200       47634047       0.42       1.37E-06       0.04         cg10390589       SAEI       19       TSS200       47634047       0.42       1.50E-06       0.04         cg1613029       FBXW12       3       TSS200       48413550       -0.42       1.60E-06       0.04         cg033013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg10130025       ABR       17        926264       -0.41       1.95E-06       0.04         cg07486666        7       Body       151423672       -0.41       2.12E-06       0.04         cg1013020       PRKAG2       7       Body       151423672       <	cg25047485		8		145848944	-0.43	8.14E-07	0.04
cg16856833        3        87138640       0.42       1.01E-06       0.04         cg19736040       SHC2       19       Body       451261       -0.42       1.14E-06       0.04         cg1233691       GEMIN8       X       TSS200       14048191       0.42       1.24E-06       0.04         cg10390589       SAEI       19       TSS200       47634047       0.42       1.37E-06       0.04         cg04554728        1        150185568       -0.42       1.50E-06       0.04         cg0355090        5        1951736       -0.41       1.70E-06       0.04         cg08379300        19        13979907       -0.41       1.80E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg03402073       PMM1       22       Body       183226025       -0.41       2.12E-06       0.04         cg03402073       PMKG2       7       Body       151423672       -0.41       2.12E-06       0.04         cg03402075       ABR       17        926264       -0.41	cg04524770	LMF1	16	Body	946408	-0.42	9.96E-07	0.04
cg19736040       SHC2       19       Body       451261       -0.42       1.14E-06       0.04         cg12339691       GEMIN8       X       TSS200       14048191       0.42       1.24E-06       0.04         cg10390589       SAE1       19       TSS200       47634047       0.42       1.37E-06       0.04         cg04554728        1        150185568       -0.42       1.60E-06       0.04         cg03250090        5        1951736       -0.41       1.70E-06       0.04         cg033013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg034073       PMM1       22       Body       41974480       -0.41       1.89E-06       0.04         cg10370025       ABR       17        926264       -041       2.12E-06       0.04         cg1071020       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg1051055       LOC340094       5        5062128       -0.41       2.47E-06       0.04         cg1051052       DC340094       5        5062128       -0	cg16856833		3		87138640	0.42	1.01E-06	0.04
cg21239691       GEMIN8       X       TSS200       14048191       0.42       1.24E-06       0.04         cg10390589       SAE1       19       TSS200       47634047       0.42       1.37E-06       0.04         cg04554728        1        150185568       -0.42       1.50E-06       0.04         cg16193029       FBXW12       3       TSS200       48413550       -0.42       1.60E-06       0.04         cg03013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg1070025       ABR       17        926264       -0.41       2.12E-06       0.04         cg10710220       PRKAG2       7       Body       151423672       -0.41       2.47E-06       0.04         cg1011025       LOC340094       5        5062128       -0.41       2.48E-06       0.04         cg10210526       TTC40       10        134663864	cg19736040	SHC2	19	Body	451261	-0.42	1.14E-06	0.04
cg10390589       SAE1       19       TSS200       47634047       0.42       1.37E-06       0.04         cg04554728        1        150185568       -0.42       1.50E-06       0.04         cg16193029 <i>FBXW12</i> 3       TSS200       48413550       -0.42       1.60E-06       0.04         cg0330329 <i>IMPACT</i> 18       Body       2206989       0.41       1.74E-06       0.04         cg03402073 <i>PMM1</i> 22       Body       41974480       -0.41       1.80E-06       0.04         cg26151355 <i>KLHL6</i> 3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025 <i>ABR</i> 17        926264       -0.41       2.12E-06       0.04         cg1017020 <i>PRKAG2</i> 7       Body       151423672       -0.41       2.47E-06       0.04         cg16210526 <i>TC40</i> 10        134663864       -0.41       2.48E-06       0.04         cg05959542 <i>RPTOR</i> 17       Body       78655493       -0.41       2.48E-06       0.04         cg16210526 <i>TTC40</i> 10 <t< td=""><td>cg21239691</td><td>GEMIN8</td><td>Х</td><td>TSS200</td><td>14048191</td><td>0.42</td><td>1.24E-06</td><td>0.04</td></t<>	cg21239691	GEMIN8	Х	TSS200	14048191	0.42	1.24E-06	0.04
cg04554728        1        150185568       -0.42       1.50E-06       0.04         cg16193029       FBXW12       3       TSS200       48413550       -0.42       1.60E-06       0.04         cg02350090        5        1951736       -0.41       1.70E-06       0.04         cg03013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg26005482        6        41170679       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.81E-06       0.04         cg0595052       RPTOR       17       Body       78655493       -0.41       2.81E-06       0.04         cg05795849       ITC40       10        134663864       -0.41 </td <td>cg10390589</td> <td>SAE1</td> <td>19</td> <td>TSS200</td> <td>47634047</td> <td>0.42</td> <td>1.37E-06</td> <td>0.04</td>	cg10390589	SAE1	19	TSS200	47634047	0.42	1.37E-06	0.04
cg16193029FBXW123TSS20048413550 $-0.42$ $1.60E-06$ $0.04$ cg0235009051951736 $-0.41$ $1.70E-06$ $0.04$ cg03013329IMPACT18Body22006989 $0.41$ $1.74E-06$ $0.04$ cg0887939019 $13979907$ $-0.41$ $1.80E-06$ $0.04$ cg03402073PMM122Body $41974480$ $-0.41$ $1.95E-06$ $0.04$ cg10370025ABR17 $926264$ $-0.41$ $2.12E-06$ $0.04$ cg12710220PRKAG27Body $151423672$ $-0.41$ $2.33E-06$ $0.04$ cg26054826 $449134$ $-0.41$ $2.48E-06$ $0.04$ cg1610526 <i>ITC40</i> 10 $134663864$ $-0.41$ $2.81E-06$ $0.04$ cg09596252 <i>RPTOR</i> 17Body $78655493$ $-0.41$ $3.00E-06$ $0.04$ cg24399376 <i>LOC100133545</i> 11Body $2005497$ $-0.40$ $3.37E-06$ $0.04$ cg23261530 <i>RPS6KA2</i> 6Body $167017999$ $-0.40$ $3.61E-06$ $0.04$ cg23070311 <i>SMPD1</i> 11TSS1500 $48897279$ $0.50$ $4.16E-09$ $1.72E-03$	cg04554728		1		150185568	-0.42	1.50E-06	0.04
cg02350090        5        1951736       -0.41       1.70E-06       0.04         cg03013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg08879390        19        13979907       -0.41       1.80E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg26151355       KLHL6       3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg12710220       PRKAG2       7       Body       151423672       -0.41       2.47E-06       0.04         cg26005482        6        41170679       -0.41       2.48E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg05595849       ITGA11       15       Body       78655493       -0.40       3.37E-06       0.04         cg23261530       RP56KA2       6       Body       167017999 <t< td=""><td>cg16193029</td><td>FBXW12</td><td>3</td><td>TSS200</td><td>48413550</td><td>-0.42</td><td>1.60E-06</td><td>0.04</td></t<>	cg16193029	FBXW12	3	TSS200	48413550	-0.42	1.60E-06	0.04
cg03013329       IMPACT       18       Body       22006989       0.41       1.74E-06       0.04         cg08879390        19        13979907       -0.41       1.80E-06       0.04         cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg26151355       KLHL6       3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg12710220       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg26005482        6        4419134       -0.41       2.47E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg09596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849 <i>ITGA11</i> 15       Body       48624879       -0.40       3.16E-06       0.04         cg23261530       RP56KA2       6       Body       13382760	cg02350090		5		1951736	-0.41	1.70E-06	0.04
cg08879390        19        13979907       -0.41       1.80E-06       0.04         cg03402073       PMMI       22       Body       41974480       -0.41       1.95E-06       0.04         cg26151355       KLHL6       3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg12710220       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg07486666        7        41170679       -0.41       2.47E-06       0.04         cg26005482        6        449134       -0.41       2.48E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg0556052       RPTOR       17       Body       7855493       -0.41       3.00E-06       0.04         cg05795849       ITPR1       3       Body       4794082       0.40       3.37E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40	cg03013329	IMPACT	18	Body	22006989	0.41	1.74E-06	0.04
cg03402073       PMM1       22       Body       41974480       -0.41       1.95E-06       0.04         cg26151355       KLHL6       3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg10370020       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg07486666        7       Body       151423672       -0.41       2.47E-06       0.04         cg26005482        6        41170679       -0.41       2.48E-06       0.04         cg16210526 <i>LOC340094</i> 5        5062128       -0.41       2.56E-06       0.04         cg09596252 <i>RPTOR</i> 10        134663864       -0.41       2.81E-06       0.04         cg05795849 <i>ITPR1</i> 3       Body       4794082       0.40       3.37E-06       0.04         cg23261530 <i>RPS6KA2</i> 6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311 <i>SMPD1</i> 11       TSS1500       6411	cg08879390		19		13979907	-0.41	1.80E-06	0.04
cg26151355       KLHL6       3       Body       183226025       -0.41       2.01E-06       0.04         cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg1074020       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg07486666        7        41170679       -0.41       2.47E-06       0.04         cg26005482        6        449134       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg059596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITFR1       3       Body       4794082       0.40       3.37E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999	cg03402073	PMM1	22	Body	41974480	-0.41	1.95E-06	0.04
cg10370025       ABR       17        926264       -0.41       2.12E-06       0.04         cg12710220       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg07486666        7        41170679       -0.41       2.47E-06       0.04         cg26005482        6        449134       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg10526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg059596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITGA11       15       Body       68624879       -0.40       3.16E-06       0.04         cg10471356       ATP11A       13       Body       11382760       -0.40       3.59E-06       0.04         cg2051530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396	cg26151355	KLHL6	3	Body	183226025	-0.41	2.01E-06	0.04
cg12710220       PRKAG2       7       Body       151423672       -0.41       2.23E-06       0.04         cg07486666        7        41170679       -0.41       2.47E-06       0.04         cg26005482        6        449134       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg09596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITCR1       15       Body       68624879       -0.40       3.16E-06       0.04         cg10471356       ATP11A       15       Body       2005497       -0.40       3.47E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.59E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04	cg10370025	ABR	17		926264	-0.41	2.12E-06	0.04
cg07486666        7        41170679       -0.41       2.47E-06       0.04         cg26005482        6        449134       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg1920526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg09596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITPR1       15       Body       68624879       -0.40       3.16E-06       0.04         cg10471356       ATP11A       15       Body       2005497       -0.40       3.47E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg12710220	PRKAG2	7	Body	151423672	-0.41	2.23E-06	0.04
cg26005482        6        449134       -0.41       2.48E-06       0.04         cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg09596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITGA11       15       Body       68624879       -0.40       3.16E-06       0.04         cg24399376       LOC100133545       11       Body       2005497       -0.40       3.47E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg07486666		7		41170679	-0.41	2.47E-06	0.04
cg19115205       LOC340094       5        5062128       -0.41       2.56E-06       0.04         cg16210526       TTC40       10        134663864       -0.41       2.81E-06       0.04         cg09596252       RPTOR       17       Body       78655493       -0.41       3.00E-06       0.04         cg05795849       ITGA11       15       Body       68624879       -0.40       3.16E-06       0.04         cg05795849       ITPRI       3       Body       4794082       0.40       3.37E-06       0.04         cg10471356       ATP11A       13       Body       2005497       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg26005482		6		449134	-0.41	2.48E-06	0.04
cg16210526 <i>TTC40</i> 10        134663864       -0.41       2.81E-06       0.04         cg09596252 <i>RPTOR</i> 17       Body       78655493       -0.41       3.00E-06       0.04         cg05960252 <i>ITGA11</i> 15       Body       68624879       -0.40       3.16E-06       0.04         cg05795849 <i>ITPR1</i> 3       Body       4794082       0.40       3.37E-06       0.04         cg10471356 <i>LOC100133545</i> 11       Body       2005497       -0.40       3.47E-06       0.04         cg23261530 <i>RPS6KA2</i> 6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311 <i>SMPD1</i> 11       TSS1500       6411396       0.40       3.64E-06       0.04         g17106653 <i>GRIN2D</i> 19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg19115205	LOC340094	5		5062128	-0.41	2.56E-06	0.04
cg09596252 <i>RPTOR</i> 17       Body       78655493       -0.41       3.00E-06       0.04         cg20560091 <i>ITGA11</i> 15       Body       68624879       -0.40       3.16E-06       0.04         cg05795849 <i>ITPR1</i> 3       Body       4794082       0.40       3.37E-06       0.04         cg24399376 <i>LOC100133545</i> 11       Body       2005497       -0.40       3.47E-06       0.04         cg10471356 <i>ATP11A</i> 13       Body       11382760       -0.40       3.59E-06       0.04         cg23261530 <i>RPS6KA2</i> 6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311 <i>SMPD1</i> 11       TSS1500       6411396       0.40       3.64E-06       0.04         cg17106653 <i>GRIN2D</i> 19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg16210526	TTC40	10		134663864	-0.41	2.81E-06	0.04
cg20560091       ITGA11       15       Body       68624879       -0.40       3.16E-06       0.04         cg05795849       ITPRI       3       Body       4794082       0.40       3.37E-06       0.04         cg24399376       LOC100133545       11       Body       2005497       -0.40       3.47E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg09596252	RPTOR	17	Body	78655493	-0.41	3.00E-06	0.04
cg05795849       ITPR1       3       Body       4794082       0.40       3.37E-06       0.04         cg24399376       LOC100133545       11       Body       2005497       -0.40       3.47E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         g17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg20560091	ITGA11	15	Body	68624879	-0.40	3.16E-06	0.04
cg24399376       LOC100133545       11       Body       2005497       -0.40       3.47E-06       0.04         cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         g17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03 <td>cg05795849</td> <td>ITPR1</td> <td>3</td> <td>Body</td> <td>4794082</td> <td>0.40</td> <td>3.37E-06</td> <td>0.04</td>	cg05795849	ITPR1	3	Body	4794082	0.40	3.37E-06	0.04
cg10471356       ATP11A       13       Body       113382760       -0.40       3.59E-06       0.04         cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         B. Sexual abuse (FDRn = 7)       cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg24399376	LOC100133545	11	Body	2005497	-0.40	3.47E-06	0.04
cg23261530       RPS6KA2       6       Body       167017999       -0.40       3.61E-06       0.04         cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         B. Sexual abuse (FDRn = 7)       cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg10471356	ATP11A	13	Body	113382760	-0.40	3.59E-06	0.04
cg20770311       SMPD1       11       TSS1500       6411396       0.40       3.64E-06       0.04         B. Sexual abuse (FDRn = 7)       cg17106653       GRIN2D       19       TSS1500       48897279       0.50       4.16E-09       1.72E-03	cg23261530	RPS6KA2	6	Body	167017999	-0.40	3.61E-06	0.04
B. Sexual abuse (FDRn = 7)           cg17106653         GRIN2D         19         TSS1500         48897279         0.50         4.16E-09         1.72E-03	cg20770311	SMPD1	11	TSS1500	6411396	0.40	3.64E-06	0.04
cg17106653 <i>GRIN2D</i> 19 TSS1500 48897279 0.50 4.16E-09 1.72E-03	R Sevual abuse	(FDRn - 7)						
1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	cg17106653	$\frac{C(\mathbf{I} D \mathbf{K} n = 1)}{GRIN2D}$	19	TSS1500	48897279	0.50	4 16E-09	1 72E-03
cg00974464 PRDM15 21 Body 43254115 -0.46 6.02E-08 0.01	cg00974464	PRDM15	21	Body	43254115	-0.46	4.10E 09	0.01
cg26528551 MGMT 10 Rody 131445415 -0.46 6.84F-0.8 0.01	cg26528551	MGMT	10	Body	131445415	-0.46	6 84E-08	0.01
cg10795666 MLNR 13 1stExon 49794635 0.45 2.23E-07 0.02	cg10795666	MLNR	13	1stExon	49794635	0.45	2.23E-07	0.01
cg02618355 MYOM2 8 Body 2024368 -0.44 2.88F-07 0.02	cg02618355	MYOM2	8	Body	2024368	-0.44	2.88E-07	0.02
cg26513050 DIAPH2/RPA4 X Body 96138983 -0.44 2.91F-07 0.02	cg26513050	DIAPH2/RPA4	x	Body	96138983	-0.44	2.91E-07	0.02
cg23983710 <i>DIP2C</i> 10 Body 370756 -0.43 5.83E-07 0.03	cg23983710	DIP2C	10	Body	370756	-0.43	5.83E-07	0.02

C. Physical negle	ct (FDR <i>n</i> = 118)						
cg00691266	EVPL	17	Body	74015089	-0.46	1.01E-07	0.01
cg27083825	SYNJ2	6	Body	158453594	-0.45	1.02E-07	0.01
cg17918089	SETDB1	1	Body	150899251	0.45	1.14E-07	0.01
cg10094509	MAP2K4P1/CHIC1	Х	TSS1500	72783579	0.45	1.16E-07	0.01
cg09863040	JADE1	4	TSS200	129730729	0.44	2.10E-07	0.01
cg08796898	HUWE1	Х	5'UTR	53713664	0.44	3.59E-07	0.02
cg20668974	ZNF827	4	Body	146857981	0.44	3.63E-07	0.02
cg18116160	GABBR1	6	Body	29575145	-0.44	4.01E-07	0.02
cg09084892	FAT1	4	Body	187557837	-0.43	4.35E-07	0.02
cg08054907	HIST1H1A	6	TSS1500	26019358	-0.43	5.12E-07	0.02
cg21076890	COL4A2	13	Body	110965662	-0.43	5.18E-07	0.02
cg17742334		1		9489130	0.43	5.31E-07	0.02
cg11650926		5		5568539	-0.43	6.50E-07	0.02
cg13409248	ENTPD3	3	TSS200	40428643	0.43	7.50E-07	0.02
cg09155575	ORMDL3	17	TSS200	38084037	0.43	8.07E-07	0.02
cg07728092	EIF5A2	3	TSS1500	170626980	-0.42	8.55E-07	0.02
cg18458353	CEACAM5	19	TSS1500	42211171	0.42	8.89E-07	0.02
cg21845580	SPRYD3	12	Body	53472882	0.42	9.37E-07	0.02
cg21634218	FLJ26850	19	TSS1500	50553699	0.42	1.01E-06	0.02
cg17693420	ZNF655	7	5'UTR	99157059	0.42	1.06E-06	0.02
cg04480325		2		37422917	-0.42	1.09E-06	0.02
cg06779606	CARD11	7	5'UTR	3041079	-0.42	1.14E-06	0.02
cg10494397	CC2D2A	4	Body	15593854	-0.42	1.28E-06	0.02
cg00311772	PPP1R7	2	Body	242092919	-0.42	1.32E-06	0.02
cg26248645	PELI1	2	5'UTR	64370690	0.42	1.36E-06	0.02
cg18001780		11		128500564	0.42	1.48E-06	0.02
cg18022926	PBLD/HNRNPH3	10	TSS1500	70093071	0.42	1.54E-06	0.02
cg02550027	BLOC1S2	10	TSS1500	102046474	0.41	1.64E-06	0.02
cg17309464	ZNF565	19	5'UTR	36705192	0.41	1.96E-06	0.02
cg04586332		22		45659810	0.41	2.00E-06	0.02
cg07032930	MRPL53	2	3'UTR	74699212	0.41	2.16E-06	0.03
cg02430183	WNK2	9	1stExon	95947766	-0.41	2.17E-06	0.03
cg06850509		2		96314847	-0.41	2.22E-06	0.03
cg24114390	FAM133A	Х	TSS1500	92928730	0.41	2.32E-06	0.03
cg17822007		3		155463780	0.41	2.37E-06	0.03
cg14030997		9		108420639	-0.41	2.49E-06	0.03
cg14679558		13		21652153	-0.41	2.70E-06	0.03
cg12637205	CYLD	16	5'UTR	50776628	0.41	2.71E-06	0.03
cg09090582	CNPY1	7	5'UTR	155324691	0.41	2.77E-06	0.03
cg03500617		16		74812570	-0.41	2.86E-06	0.03
cg10493855	ТТС7В	14	Body	91087586	-0.41	2.89E-06	0.03
cg14945917		4		38145641	-0.41	2.95E-06	0.03
cg16539640	CYFIP1	15	Body	22945028	-0.41	2.98E-06	0.03
cg02564272	MAN1A1	6	Body	119507963	-0.41	3.04E-06	0.03
cg18073471	PRDM8	4	5'UTR	81119198	0.40	3.24E-06	0.03
cg27568885	GCNT3	15	Body	59911449	-0.40	3.45E-06	0.03
cg13468418		4		186048907	0.40	3.54E-06	0.03
cg16560389		10		133318201	-0.40	3.60E-06	0.03

cg16210526		10		134663864	-0.40	3.71E-06	0.03
cg13521097	FERMT2	14	5'UTR	53417433	0.40	3.85E-06	0.03
cg15799267	ALOX15B	17	5'UTR	7942406	0.40	4.08E-06	0.03
cg19270277	CROCC	1	Body	17296995	-0.40	4.10E-06	0.03
cg04375213	DCP1B	12	Body	2113435	0.40	4.12E-06	0.03
cg13109184	LYST	1		236029439	0.40	4.40E-06	0.03
cg22902505	PRDM8	4	5'UTR	81119473	0.40	4.79E-06	0.04
cg08614562	KIF1A	2	Body	241665572	-0.40	5.13E-06	0.04
cg19864758	PCSK2	20	TSS1500	17206720	0.40	5.19E-06	0.04
cg27063720	EP400NL	12	Body	132568987	0.40	5.23E-06	0.04
cg24604043	SLC39A3	19	TSS1500	2740392	0.40	5.34E-06	0.04
cg15136913		10		134893540	-0.40	5.38E-06	0.04
cg19091677	HMGB2	4	TSS1500	174255518	0.40	5.44E-06	0.04
cg10625723		4		6235503	-0.40	5.50E-06	0.04
cg07821496	TDRKH	1	5'UTR	151762942	0.40	5.51E-06	0.04
cg08294772		8		138135392	0.40	5.52E-06	0.04
cg26309691	SCCPDH	1	Body	246887944	0.39	6.14E-06	0.04
cg21074877		14		23481968	-0.39	6.54E-06	0.04
cg19827167	MIR760	1	TSS200	94312206	0.39	6.56E-06	0.04
cg27330720	FBXL20	17	TSS1500	37558981	0.39	6.67E-06	0.04
cg05130304	C3orf23	3	TSS1500	44379503	0.39	6.72E-06	0.04
cg26509269		17		34970306	-0.39	6.89E-06	0.04
cg27496299	DNAJB6	7	Body	157171202	-0.39	6.90E-06	0.04
cg02458885	PRDM8	4	5'UTR	81119249	0.39	6.93E-06	0.04
cg21385432	EP400	12	Body	132512858	-0.39	7.04E-06	0.04
cg06874656	NTM	11	TSS1500	131780353	0.39	7.10E-06	0.04
cg21239691	GEMIN8	Х	TSS200	14048191	0.39	7.20E-06	0.04
cg00895834	PRDM16	1	Body	3147966	-0.39	7.20E-06	0.04
cg21473728	MYOM2	8	Body	2031651	-0.39	7.42E-06	0.04
cg10390589	SAE1	19	TSS200	47634047	0.39	7.64E-06	0.04
cg10246772	IGF1R	15	Body	99250916	-0.39	7.90E-06	0.04
cg02514021	FLJ43663	7	Body	130791171	0.39	8.24E-06	0.04
cg10609984	ZNF25	10	TSS200	38265593	0.39	8.43E-06	0.04
cg25087154	IL1RAP	3	TSS1500	190231338	0.39	8.47E-06	0.04
ch.11.150171F	NUP98	11	Body	3739920	0.39	9.00E-06	0.04
cg23284284	GIGYF2	2	TSS200	233561917	0.39	9.02E-06	0.04
cg01093285	PTPRN2	7	Body	157436506	-0.39	9.38E-06	0.04
cg27632471	C20orf96	20	Body	259123	-0.39	9.42E-06	0.04
cg12411309	SMUG1	12	5'UTR	54582659	0.39	9.51E-06	0.04
cg17341361	TNRC18	7	Body	5414170	-0.39	9.51E-06	0.04
cg07124687		6		28956332	0.39	9.71E-06	0.04
cg12154976	RASGEF1C	5	5'UTR	179585198	-0.39	9.84E-06	0.04
cg12398448	IGF1R	15	Body	99251461	-0.39	9.87E-06	0.04
cg10462751	RIC8B	12	TSS200	107168330	0.38	1.03E-05	0.05
cg04204072	THUMPD1	16	IstExon	20752857	0.38	1.04E-05	0.05
cg10962490	GLIS1	1	5'UTR	54072456	-0.38	1.05E-05	0.05
cg13653274		10	 D 1	132632928	-0.38	1.07E-05	0.05
cg15472403	OSCAR	19	Body	54599455	0.38	1.09E-05	0.05
cg00014152	WDR13	Х	Body	48457128	-0.38	1.10E-05	0.05

cg25247351	HLA-DMB	6	1stExon	32908794	0.38	1.11E-05	0.05
cg03156396	MSLN	16	Body	816177	-0.38	1.13E-05	0.05
cg10650870	TNPO1	5	Body	72144764	0.38	1.15E-05	0.05
cg01487433	CILP2	19	TSS200	19648987	-0.38	1.15E-05	0.05
cg22533689	EPHA2	1	Body	16464479	-0.38	1.16E-05	0.05
cg13735965	ZNF616	19	TSS1500	52643487	0.38	1.18E-05	0.05
cg07307426	DET1	15	5'UTR	89089587	0.38	1.18E-05	0.05
cg25002318	TRPV1	17	Body	3471868	-0.38	1.20E-05	0.05
cg19554389	WNT5A	3	1stExon	55521134	0.38	1.20E-05	0.05
cg16953816	VPS37B	12	3'UTR	123349952	-0.38	1.22E-05	0.05
cg14604000	C5orf13	5	TSS1500	111093875	0.38	1.24E-05	0.05
cg11969467	MRPS14	1	TSS200	174992598	0.38	1.24E-05	0.05
cg03998119	PTPN21	14	Body	88945315	-0.38	1.25E-05	0.05
cg03041046	SPRYD3	12	5'UTR	53473136	0.38	1.27E-05	0.05
cg25051341	PRDM13	6	Body	100061307	0.38	1.30E-05	0.05
cg01407073	NAP1L1	12	TSS200	76478898	0.38	1.30E-05	0.05
cg15874300	CRBN	3	Body	3221199	0.38	1.31E-05	0.05
cg13144843		3		174158740	0.38	1.33E-05	0.05
cg26163537	GRB10	7	TSS1500	50861592	0.38	1.37E-05	0.05
cg20657691	RBM3	Х	TSS1500	48432562	0.38	1.38E-05	0.05
cg06616055	MUPCDH	11	Body	623756	0.38	1.42E-05	0.05

*N.B.* Genes with multiple significant probes are highlighted in blue. Standardized Beta estimates are used as a measure of effect size, where an effect of 0.10 is small effect, an effect of 0.24 is a medium effect, and an effect of 0.37 is a large effect.

### **OS4.** Quantile-quantile plots for each maltreatment type



8

CpG probe	Gene	Chr	Genomic location	Position			Sensitivity Analyses							
					Original	analysis <sup>a</sup>	SVA ar	SVA analysis <sup>b</sup> V		analysis <sup>c</sup>	Bootstrapping <sup>d</sup>			High confidence
					<i>p</i> -value	q-value	<i>p</i> -value	<i>q</i> -value	<i>p</i> -value	<i>q</i> -value	B	95%	o CI	association <sup>e</sup>
A. Physical abu	use $(n = 34)$													
cg20000641	PSEN2	1	TSS1500	227058046	2.53E-11	1.47E-05	4.205E-09	4.91E-04	1.64E-03	0.61	-0.006	-0.010	-0.001	*
cg02353937	SMC1A/RIBC1	Х	TSS1500	53449152	2.90E-09	7.89E-04	1.922E-08	1.35E-03	1.26E-04	0.42	-0.006	-0.012	-0.001	*
cg15440363	GPD1L	3	TSS200	32148023	2.06E-08	3.64E-03	3.819E-09	4.91E-04	0.05	0.76	-0.005	-0.009	-0.001	*
cg04412054	GALNS	16	Body	88897539	5.69E-08	0.01	1.244E-05	0.04	0.01	0.69	-0.005	-0.010	-0.001	*
cg22311608	SMC1A/RIBC1	Х	TSS200	53449829	8.45E-08	0.01	1.486E-05	0.04	1.50E-05	0.22	-0.006	-0.011	-0.001	*
cg24365098	C11orf84	11	3'UTR	63594804	1.43E-07	0.01	8.455E-05	0.07	9.76E-03	0.69	-0.009	-0.014	-0.003	-
cg03561071	[FAT3]	11		91844355	4.28E-07	0.03	1.174E-06	0.01	7.25E-04	0.58	-0.004	-0.006	-0.001	*
cg26454299	PPP3CA	4		102268957	5.58E-07	0.03	1.073E-05	0.04	9.16E-04	0.59	-0.006	-0.010	-0.001	*
cg25047485	[ARHGAP39]	8		145848944	8.14E-07	0.04	1.857E-05	0.04	0.01	0.69	-0.004	-0.006	-0.003	*
cg04524770	LMF1	16	Body	946408	9.96E-07	0.04	1.270E-06	0.01	0.02	0.72	-0.012	-0.020	-0.004	*
cg16856833	[VGLL3]	3		87138640	1.01E-06	0.04	1.605E-05	0.04	1.29E-03	0.60	-0.006	-0.011	-0.001	*
cg19736040	SHC2	19	Body	451261	1.14E-06	0.04	8.149E-06	0.03	3.69E-03	0.64	-0.003	-0.005	-0.001	*
cg21239691	GEMIN8	Х	TSS200	14048191	1.24E-06	0.04	8.405E-04	0.15	1.22E-05	0.22	-0.004	-0.006	-0.001	-
cg10390589	SAE1	19	TSS200	47634047	1.37E-06	0.04	1.177E-06	0.01	4.86E-03	0.65	-0.003	-0.006	0.000	-
cg04554728	[ANP32E ]	1		150185568	1.50E-06	0.04	5.394E-07	0.01	1.08E-03	0.60	-0.008	-0.016	-0.002	*
cg16193029	FBXW12	3	TSS200	48413550	1.60E-06	0.04	8.757E-06	0.04	0.48	0.90	-0.012	-0.022	-0.004	-
cg02350090	[IRX4]	5		1951736	1.70E-06	0.04	5.984E-05	0.06	2.04E-06	0.19	-0.006	-0.010	-0.002	-
cg03013329	IMPACT	18	Body	22006989	1.74E-06	0.04	4.188E-04	0.13	9.24E-03	0.68	-0.003	-0.005	-0.002	-
cg08879390	[NANOS3]	19		13979907	1.80E-06	0.04	6.189E-06	0.03	1.47E-03	0.61	-0.003	-0.005	-0.001	*
cg03402073	PMM1	22	Body	41974480	1.95E-06	0.04	2.855E-05	0.05	7.61E-03	0.67	-0.004	-0.008	-0.001	*
cg26151355	KLHL6	3	Body	183226025	2.01E-06	0.04	1.771E-04	0.10	6.15E-03	0.67	-0.003	-0.006	-0.001	-
cg10370025	ABR	17		926264	2.12E-06	0.04	1.124E-05	0.04	3.74E-03	0.64	-0.002	-0.003	0.000	-
cg12710220	PRKAG2	7	Body	151423672	2.23E-06	0.04	7.008E-06	0.03	7.05E-03	0.67	-0.003	-0.005	-0.002	*
cg07486666	[C7orf10]	7		41170679	2.47E-06	0.04	2.865E-05	0.05	0.02	0.73	0.002	0.000	0.003	-
cg26005482	[IRF4]	6		449134	2.48E-06	0.04	4.799E-05	0.06	2.32E-03	0.63	0.006	0.002	0.011	-

## **OS5:** Sensitivity analyses for FDR-corrected DMPs associated with physical abuse, sexual abuse and physical neglect

cg19115205	LOC340094	5		5062128	2.56E-06	0.04	7.828E-06	0.03	6.99E-05	0.40	0.005	0.001	0.009	*
cg16210526	TTC40	10		134663864	2.81E-06	0.04	8.721E-07	0.01	1.01E-05	0.22	0.002	0.000	0.004	-
cg09596252	RPTOR	17	Body	78655493	3.00E-06	0.04	6.730E-05	0.07	2.63E-04	0.47	0.005	0.002	0.008	-
cg20560091	ITGA11	15	Body	68624879	3.16E-06	0.04	2.024E-05	0.05	2.35E-04	0.47	0.004	0.001	0.007	*
cg05795849	ITPR1	3	Body	4794082	3.37E-06	0.04	3.684E-08	1.84E-03	3.42E-04	0.51	0.005	0.001	0.008	*
cg24399376	LOC100133545	11	Body	2005497	3.47E-06	0.04	5.354E-05	0.06	1.38E-03	0.61	0.004	0.002	0.007	-
cg10471356	ATP11A	13	Body	113382760	3.59E-06	0.04	4.097E-05	0.05	1.53E-03	0.61	0.003	0.000	0.005	-
cg23261530	RPS6KA2	6	Body	167017999	3.61E-06	0.04	3.408E-05	0.05	4.74E-05	0.37	0.004	0.002	0.007	*
cg20770311	SMPD1	11	TSS1500	6411396	3.64E-06	0.04	1.204E-06	0.01	2.48E-03	0.63	0.007	0.002	0.012	*
B. Sexual abus	e (n = 7)													
cg17106653	GRIN2D	19	TSS1500	48897279	4.16E-09	1.72E-03	8.46E-10	1.75E-04	0.03	1.00	0.011	-0.001	0.020	-
cg00974464	PRDM15	21	Body	43254115	6.02E-08	0.01	3.21E-09	3.32E-04	0.15	1.00	-0.006	-0.012	0.000	-
cg26528551	MGMT	10	Body	131445415	6.84E-08	0.01	2.66E-09	3.32E-04	0.04	1.00	-0.005	-0.010	0.000	-
cg10795666	MLNR	13	1stExon	49794635	2.23E-07	0.02	1.21E-07	3.85E-03	1.90E-05	1.00	0.005	0.002	0.009	*
cg02618355	MYOM2	8	Body	2024368	2.88E-07	0.02	3.58E-08	1.85E-03	0.03	1.00	-0.004	-0.011	0.000	-
cg26513050	DIAPH2/RPA4	Х	Body	96138983	2.91E-07	0.02	1.00E-07	3.46E-03	0.30	1.00	-0.009	-0.024	0.001	-
cg23983710	DIP2C	10	Body	370756	5.83E-07	0.03	2.32E-11	9.59E-06	0.05	1.00	-0.008	-0.013	0.000	-
C. Physical neg	glect (n = 118)													
cg00691266	EVPL	17	Body	74015089	1.01E-07	0.01	3.27E-04	0.19	2.75E-05	0.13	-0.005	-0.007	-0.003	-
cg27083825	SYNJ2	6	Body	158453594	1.02E-07	0.01	1.39E-03	0.28	2.50E-04	0.15	-0.005	-0.008	-0.002	-
cg17918089	SETDB1	1	Body	150899251	1.14E-07	0.01	2.80E-09	1.04E-03	3.14E-03	0.19	0.005	0.001	0.009	*
cg10094509	MAP2K4P1/CHIC1	Х	TSS1500	72783579	1.16E-07	0.01	3.82E-05	0.10	6.72E-06	0.12	0.006	0.001	0.011	-
cg09863040	JADE1	4	TSS200	129730729	2.10E-07	0.01	6.01E-07	0.02	7.85E-06	0.12	0.001	0	0.001	-
cg08796898	HUWE1	Х	5'UTR	53713664	3.59E-07	0.02	2.48E-04	0.18	2.23E-05	0.13	0.01	0.005	0.014	-
cg20668974	ZNF827	4	Body	146857981	3.63E-07	0.02	2.13E-06	0.04	4.10E-05	0.13	0.004	0.002	0.006	*
cg18116160	GABBR1	6	Body	29575145	4.01E-07	0.02	8.27E-05	0.13	3.07E-06	0.11	-0.002	-0.004	-0.001	-
cg09084892	FAT1	4	Body	187557837	4.35E-07	0.02	4.65E-05	0.11	0.01	0.23	-0.008	-0.013	-0.002	-
cg08054907	HIST1H1A	6	TSS1500	26019358	5.12E-07	0.02	3.85E-05	0.10	1.13E-04	0.14	-0.006	-0.01	-0.002	-
cg21076890	COL4A2	13	Body	110965662	5.18E-07	0.02	1.09E-03	0.26	4.96E-04	0.16	-0.006	-0.01	-0.002	-
cg17742334	[SLC25A33]	1		9489130	5.31E-07	0.02	1.56E-04	0.15	1.27E-04	0.14	0.003	0.001	0.006	-
cg11650926	[ADAMTS16]	5		5568539	6.50E-07	0.02	1.13E-03	0.26	1.71E-03	0.17	-0.003	-0.006	-0.001	-
													10	

cg13409248	ENTPD3	3	TSS200	40428643	7.50E-07	0.02	9.30E-05	0.13	1.34E-04	0.14	0.002	0.001	0.003	-
cg09155575	ORMDL3	17	TSS200	38084037	8.07E-07	0.02	1.67E-07	0.02	0.09	0.35	0.005	0.001	0.01	-
cg07728092	EIF5A2	3	TSS1500	170626980	8.55E-07	0.02	1.80E-04	0.16	6.75E-05	0.13	-0.002	-0.004	-0.001	-
cg18458353	CEACAM5	19	TSS1500	42211171	8.89E-07	0.02	1.32E-06	0.04	1.57E-06	0.10	0.008	0.005	0.011	*
cg21845580	SPRYD3	12	Body	53472882	9.37E-07	0.02	5.37E-06	0.05	4.57E-06	0.12	0.005	0.003	0.007	*
cg21634218	FLJ26850	19	TSS1500	50553699	1.01E-06	0.02	4.31E-03	0.35	6.95E-04	0.16	0.011	0.005	0.018	-
cg17693420	ZNF655	7	5'UTR	99157059	1.06E-06	0.02	6.95E-04	0.23	3.69E-03	0.19	0.006	0.002	0.009	-
cg04480325	[SULT6B1]	2		37422917	1.09E-06	0.02	6.15E-03	0.38	8.24E-05	0.13	-0.006	-0.012	-0.003	-
cg06779606	CARD11	7	5'UTR	3041079	1.14E-06	0.02	1.59E-04	0.15	4.03E-05	0.13	-0.006	-0.008	-0.003	-
cg10494397	CC2D2A	4	Body	15593854	1.28E-06	0.02	5.73E-03	0.37	1.70E-03	0.17	-0.002	-0.005	-0.001	-
cg00311772	PPP1R7	2	Body	242092919	1.32E-06	0.02	9.47E-06	0.07	7.11E-06	0.12	-0.006	-0.009	-0.003	-
cg26248645	PELII	2	5'UTR	64370690	1.36E-06	0.02	3.21E-06	0.05	5.00E-04	0.16	0.006	0.001	0.012	*
cg18001780	[ETS1]	11		128500564	1.48E-06	0.02	3.98E-05	0.11	2.41E-04	0.15	0.005	0.002	0.007	-
cg18022926	PBLD/HNRNPH3	10	TSS1500	70093071	1.54E-06	0.02	5.38E-06	0.05	0.05	0.31	0.005	0.001	0.011	*
cg02550027	BLOC1S2	10	TSS1500	102046474	1.64E-06	0.02	2.17E-03	0.30	2.38E-05	0.13	0.001	0.001	0.002	-
cg17309464	ZNF565	19	5'UTR	36705192	1.96E-06	0.02	2.67E-07	0.02	7.98E-03	0.21	0.004	0.001	0.008	*
cg04586332	[UPK3A]	22		45659810	2.00E-06	0.02	5.60E-05	0.11	2.58E-04	0.15	0.006	0.001	0.012	-
cg07032930	MRPL53	2	3'UTR	74699212	2.16E-06	0.03	1.38E-04	0.15	2.90E-05	0.13	0.002	0.001	0.003	-
cg02430183	WNK2	9	1stExon	95947766	2.17E-06	0.03	2.84E-04	0.18	0.01	0.22	-0.004	-0.007	0	-
cg06850509		2		96314847	2.22E-06	0.03	7.97E-03	0.40	3.15E-05	0.13	-0.005	-0.008	-0.003	-
cg24114390	FAM133A	Х	TSS1500	92928730	2.32E-06	0.03	2.23E-06	0.04	3.27E-03	0.19	0.003	0.001	0.006	*
cg17822007		3		155463780	2.37E-06	0.03	1.36E-04	0.15	2.11E-05	0.13	0.007	0.003	0.011	-
cg14030997		9		108420639	2.49E-06	0.03	2.42E-02	0.50	4.04E-05	0.13	-0.004	-0.007	-0.002	-
cg14679558		13		21652153	2.70E-06	0.03	1.33E-03	0.27	8.21E-04	0.16	-0.002	-0.004	-0.001	-
cg12637205	CYLD	16	5'UTR	50776628	2.71E-06	0.03	7.45E-04	0.24	4.64E-04	0.16	0.004	0.001	0.007	-
cg09090582	CNPY1	7	5'UTR	155324691	2.77E-06	0.03	2.74E-04	0.18	2.57E-05	0.13	0.009	0.004	0.013	-
cg03500617		16		74812570	2.86E-06	0.03	7.52E-04	0.24	1.85E-04	0.15	-0.005	-0.008	-0.002	-
cg10493855	ТТС7В	14	Body	91087586	2.89E-06	0.03	1.71E-04	0.16	1.64E-04	0.15	-0.008	-0.015	-0.002	-
cg14945917		4		38145641	2.95E-06	0.03	1.72E-04	0.16	6.98E-05	0.13	-0.005	-0.008	-0.002	-
cg16539640	CYFIP1	15	Body	22945028	2.98E-06	0.03	2.54E-03	0.31	5.58E-04	0.16	-0.002	-0.003	-0.001	-
cg02564272	MANIAI	6	Body	119507963	3.04E-06	0.03	1.66E-02	0.46	4.17E-03	0.19	-0.009	-0.018	-0.001	-
cg18073471	PRDM8	4	5'UTR	81119198	3.24E-06	0.03	5.62E-04	0.22	3.23E-03	0.19	0.006	0.002	0.011	-

cg27568885	GCNT3	15	Body	59911449	3.45E-06	0.03	3.08E-08	0.01	3.13E-07	0.10	-0.003	-0.005	-0.002	*
cg13468418		4		186048907	3.54E-06	0.03	1.04E-05	0.07	1.17E-04	0.14	0.003	0.002	0.004	-
cg16560389		10		133318201	3.60E-06	0.03	4.31E-04	0.20	3.16E-05	0.13	-0.003	-0.006	-0.001	-
cg16210526		10		134663864	3.71E-06	0.03	4.93E-05	0.11	3.62E-04	0.16	-0.004	-0.006	-0.001	-
cg13521097	FERMT2	14	5'UTR	53417433	3.85E-06	0.03	8.06E-04	0.24	7.60E-04	0.16	0.002	0.001	0.003	-
cg15799267	ALOX15B	17	5'UTR	7942406	4.08E-06	0.03	2.62E-03	0.32	5.26E-06	0.12	0.007	0.005	0.01	-
cg19270277	CROCC	1	Body	17296995	4.10E-06	0.03	1.64E-04	0.15	2.29E-05	0.13	-0.006	-0.009	-0.003	-
cg04375213	DCP1B	12	Body	2113435	4.12E-06	0.03	4.06E-06	0.05	7.33E-06	0.12	0.001	0.001	0.002	*
cg13109184	LYST	1		236029439	4.40E-06	0.03	4.95E-04	0.21	7.00E-03	0.20	0.005	0.001	0.01	-
cg22902505	PRDM8	4	5'UTR	81119473	4.79E-06	0.04	4.50E-04	0.20	5.07E-05	0.13	0.009	0.005	0.015	-
cg08614562	KIF1A	2	Body	241665572	5.13E-06	0.04	2.42E-03	0.31	9.33E-04	0.16	-0.004	-0.007	-0.001	-
cg19864758	PCSK2	20	TSS1500	17206720	5.19E-06	0.04	2.90E-03	0.32	9.14E-05	0.13	0.003	0.001	0.005	-
cg27063720	EP400NL	12	Body	132568987	5.23E-06	0.04	1.88E-05	0.08	8.97E-05	0.13	0.004	0.002	0.007	-
cg24604043	SLC39A3	19	TSS1500	2740392	5.34E-06	0.04	4.17E-03	0.35	8.92E-03	0.21	0.004	0.001	0.008	-
cg15136913		10		134893540	5.38E-06	0.04	9.07E-05	0.13	8.04E-05	0.13	-0.005	-0.009	-0.002	-
cg19091677	HMGB2	4	TSS1500	174255518	5.44E-06	0.04	2.13E-04	0.17	7.46E-04	0.16	0.004	0.001	0.007	-
cg10625723		4		6235503	5.50E-06	0.04	1.73E-02	0.47	2.08E-03	0.18	-0.002	-0.004	-0.001	-
cg07821496	TDRKH	1	5'UTR	151762942	5.51E-06	0.04	5.99E-04	0.22	3.25E-05	0.13	0.005	0.002	0.007	-
cg08294772		8		138135392	5.52E-06	0.04	9.92E-03	0.42	2.38E-05	0.13	0.007	0.004	0.01	-
cg26309691	SCCPDH	1	Body	246887944	6.14E-06	0.04	5.21E-03	0.36	5.23E-04	0.16	0.003	0.001	0.005	-
cg21074877		14		23481968	6.54E-06	0.04	1.16E-02	0.43	5.31E-04	0.16	-0.002	-0.003	-0.001	-
cg19827167	MIR760	1	TSS200	94312206	6.56E-06	0.04	4.61E-03	0.35	5.63E-03	0.20	0.001	0	0.003	-
cg27330720	FBXL20	17	TSS1500	37558981	6.67E-06	0.04	3.79E-05	0.10	6.64E-03	0.20	0.004	0.001	0.006	-
cg05130304	C3orf23	3	TSS1500	44379503	6.72E-06	0.04	3.64E-04	0.20	9.69E-03	0.21	0.004	0	0.007	-
cg26509269		17		34970306	6.89E-06	0.04	4.28E-04	0.20	1.41E-04	0.14	-0.004	-0.007	-0.002	-
cg27496299	DNAJB6	7	Body	157171202	6.90E-06	0.04	2.07E-03	0.30	1.05E-04	0.13	-0.004	-0.007	-0.002	-
cg02458885	PRDM8	4	5'UTR	81119249	6.93E-06	0.04	2.06E-04	0.17	2.33E-04	0.15	0.009	0.004	0.015	-
cg21385432	<i>EP400</i>	12	Body	132512858	7.04E-06	0.04	4.71E-03	0.36	4.05E-04	0.16	-0.003	-0.006	-0.001	-
cg06874656	NTM	11	TSS1500	131780353	7.10E-06	0.04	1.81E-03	0.29	5.98E-03	0.20	0.004	0.001	0.008	-
cg21239691	GEMIN8	Х	TSS200	14048191	7.20E-06	0.04	7.54E-03	0.39	1.51E-05	0.13	0.006	0.003	0.01	-
cg00895834	PRDM16	1	Body	3147966	7.20E-06	0.04	1.73E-04	0.16	3.73E-05	0.13	-0.005	-0.008	-0.003	-
cg21473728	МҮОМ2	8	Body	2031651	7.42E-06	0.04	1.73E-03	0.29	1.27E-03	0.17	-0.006	-0.009	-0.001	-

cg10390589	SAE1	19	TSS200	47634047	7.64E-06	0.04	4.33E-03	0.35	0.01	0.23	0.002	0	0.004	-
cg10246772	IGF1R	15	Body	99250916	7.90E-06	0.04	3.75E-02	0.54	4.68E-03	0.19	-0.004	-0.008	-0.001	-
cg02514021	FLJ43663	7	Body	130791171	8.24E-06	0.04	6.20E-04	0.22	3.49E-03	0.19	0.006	0.001	0.012	-
cg10609984	ZNF25	10	TSS200	38265593	8.43E-06	0.04	3.31E-05	0.10	1.49E-03	0.17	0.003	0.001	0.006	-
cg25087154	ILIRAP	3	TSS1500	190231338	8.47E-06	0.04	9.19E-04	0.25	2.63E-03	0.18	0.005	0.001	0.008	-
ch.11.150171F	NUP98	11	Body	3739920	9.00E-06	0.04	4.58E-04	0.20	8.88E-04	0.16	0.005	0.002	0.008	-
cg23284284	GIGYF2	2	TSS200	233561917	9.02E-06	0.04	6.68E-03	0.38	0.01	0.23	0.006	0.001	0.01	-
cg01093285	PTPRN2	7	Body	157436506	9.38E-06	0.04	1.68E-03	0.29	4.52E-04	0.16	-0.006	-0.009	-0.002	-
cg27632471	C20orf96	20	Body	259123	9.42E-06	0.04	1.43E-03	0.28	1.67E-03	0.17	-0.004	-0.007	-0.001	-
cg12411309	SMUG1	12	5'UTR	54582659	9.51E-06	0.04	2.22E-03	0.30	3.65E-04	0.16	0.004	0.002	0.006	-
cg17341361	TNRC18	7	Body	5414170	9.51E-06	0.04	1.58E-02	0.46	8.24E-04	0.16	-0.003	-0.004	-0.001	-
cg07124687		6		28956332	9.71E-06	0.04	5.01E-05	0.11	1.57E-04	0.14	0.005	0.002	0.008	-
cg12154976	RASGEF1C	5	5'UTR	179585198	9.84E-06	0.04	9.99E-05	0.13	1.30E-03	0.17	-0.006	-0.01	-0.002	-
cg12398448	IGF1R	15	Body	99251461	9.87E-06	0.04	2.51E-03	0.31	3.70E-05	0.13	-0.007	-0.013	-0.003	-
cg10462751	RIC8B	12	TSS200	107168330	1.03E-05	0.05	6.89E-04	0.23	9.50E-03	0.21	0.005	0.001	0.01	-
cg04204072	THUMPD1	16	1stExon	20752857	1.04E-05	0.05	1.28E-02	0.44	1.61E-04	0.14	0.003	0.001	0.006	-
cg10962490	GLISI	1	5'UTR	54072456	1.05E-05	0.05	4.55E-04	0.20	1.31E-04	0.14	-0.004	-0.006	-0.001	-
cg13653274		10		132632928	1.07E-05	0.05	1.18E-03	0.26	0.04	0.28	-0.006	-0.012	-0.001	-
cg15472403	OSCAR	19	Body	54599455	1.09E-05	0.05	1.22E-03	0.27	8.18E-04	0.16	0.003	0.001	0.005	-
cg00014152	WDR13	Х	Body	48457128	1.10E-05	0.05	2.40E-04	0.18	3.64E-04	0.16	-0.004	-0.007	-0.001	-
cg25247351	HLA-DMB	6	1stExon	32908794	1.11E-05	0.05	1.46E-04	0.15	1.77E-03	0.17	0.003	0.001	0.005	-
cg03156396	MSLN	16	Body	816177	1.13E-05	0.05	1.30E-05	0.07	4.79E-05	0.13	-0.006	-0.01	-0.002	-
cg10650870	TNPO1	5	Body	72144764	1.15E-05	0.05	1.31E-03	0.27	1.76E-04	0.15	0.002	0.001	0.003	-
cg01487433	CILP2	19	TSS200	19648987	1.15E-05	0.05	2.60E-04	0.18	1.78E-05	0.13	-0.002	-0.003	-0.001	-
cg22533689	EPHA2	1	Body	16464479	1.16E-05	0.05	6.08E-03	0.37	0.01	0.22	-0.005	-0.009	-0.001	-
cg13735965	ZNF616	19	TSS1500	52643487	1.18E-05	0.05	5.75E-03	0.37	5.42E-04	0.16	0.005	0.002	0.008	-
cg07307426	DET1	15	5'UTR	89089587	1.18E-05	0.05	2.40E-04	0.18	0.01	0.23	0.004	0	0.008	-
cg25002318	TRPV1	17	Body	3471868	1.20E-05	0.05	1.13E-03	0.26	5.74E-03	0.20	-0.005	-0.009	-0.001	-
cg19554389	WNT5A	3	1stExon	55521134	1.20E-05	0.05	1.92E-03	0.30	0.01	0.23	0.003	0.001	0.005	-
cg16953816	VPS37B	12	3'UTR	123349952	1.22E-05	0.05	7.14E-03	0.39	5.59E-04	0.16	-0.007	-0.01	-0.002	-
cg14604000	C5orf13; NREP	5	TSS1500	111093875	1.24E-05	0.05	7.20E-06	0.06	2.23E-05	0.13	0.003	0.001	0.006	-
cg11969467	MRPS14	1	TSS200	174992598	1.24E-05	0.05	3.45E-04	0.19	0.05	0.31	0.002	0	0.003	-

cg03998119	PTPN21	14	Body	88945315	1.25E-05	0.05	0.06	0.59	0.05	0.31	-0.005	-0.012	0	-
cg03041046	SPRYD3	12	5'UTR	53473136	1.27E-05	0.05	1.10E-05	0.07	9.38E-04	0.16	0.005	0.001	0.009	-
cg25051341	PRDM13	6	Body	100061307	1.30E-05	0.05	0.67	0.86	3.45E-03	0.19	0.002	0	0.005	-
cg01407073	NAPILI	12	TSS200	76478898	1.30E-05	0.05	9.16E-03	0.41	6.63E-04	0.16	0.004	0.001	0.007	-
cg15874300	CRBN	3	Body	3221199	1.31E-05	0.05	1.30E-03	0.27	4.33E-04	0.16	0.003	0.001	0.004	-
cg13144843		3		174158740	1.33E-05	0.05	9.94E-03	0.42	7.50E-04	0.16	0.003	0.001	0.004	-
cg26163537	GRB10	7	TSS1500	50861592	1.37E-05	0.05	5.83E-05	0.11	4.75E-04	0.16	0.004	0.002	0.007	-
cg20657691	RBM3	Х	TSS1500	48432562	1.38E-05	0.05	2.78E-04	0.18	0.01	0.22	0.006	0	0.012	-
cg06616055	MUPCDH	11	Body	623756	1.42E-05	0.05	1.65E-04	0.16	3.76E-05	0.13	-0.007	-0.011	-0.003	-

N.B. Genes with multiple significant probes are highlighted in blue. Standardized Beta estimates are used as a measure of effect size, where an effect of 0.10 is small effect, an effect of 0.24 is a medium effect, and an effect of 0.37 is a large effect. Values in grey represent non-significant estimates.

<sup>a</sup>Original analysis: P-values and genome-wide corrected Q-values from the original epigenome-wide association analyses (i.e. correcting for age, sex and ethnicity) <sup>b</sup>SVA: P-values and genome-wide corrected Q-values after adjusting for the 11 surrogate variables identified via surrogate variable analysis.

**\*Winsorizing:** P-values and genome-wide corrected Q-values after winsorizing <5% and >95% values of score distribution in order to minimize the impact of outliers.

<sup>d</sup>Bootstrapping: Unstandardized betas and 95% Confidence Intervals (CIs) after bootrapping estimates 10,000 times.

**\*High confidence associations** (marked with \*) are defined as having (i) a significant SVA q-value; (ii) a significant winsorized p-value; and (iii) bootstrapped CIs that do not cross zero.

**OS6:** Enriched biological processes for genes associated with epigenetic variation that is unique to each maltreatment type

GO P	athway	<i>N</i> genes in input list	Totan N in pathway	%	<i>p</i> -value	Genes
Physic	cal abuse					
1	regulation of cardiac muscle hypertrophy	3	24	13%	1.04E-12	RGS2;EDN1;ADRB1
2	negative regulation of smooth muscle contraction	2	12	17%	9.01E-12	RGS2;ADRB1
3	superoxide metabolic process	3	31	10%	6.38E-10	EDN1;MT3;PREX1
4	metanephric nephron tubule development	2	15	13%	1.36E-09	CALB1;ACAT1
5	oocyte maturation	2	16	13%	4.15E-09	BRCA2;REC8
6	positive regulation of protein dephosphorylation	2	16	13%	4.90E-09	NELF; CALM1
7	fucose metabolic process	2	16	13%	5.22E-09	B3GALTL;FUT5
8	regulation of keratinocyte proliferation	2	17	12%	1.51E-08	CASK;PRKD1
9	positive regulation of heart rate	2	18	11%	4.04E-08	EDN1;ADRB1
10	male meiosis I	2	19	11%	1.03E-07	BRCA2;REC8
11	regulation of urine volume	2	20	10%	2.19E-07	EDN1;ADRB1
12	glycogen catabolic process	2	20	10%	2.32E-07	ADRB1;CALM1
13	regulation of oxidoreductase activity	4	73	5%	3.26E-07	GLA;EDN1;CALM1;MT3
14	negative regulation of oxidoreductase activity	2	21	10%	4.64E-07	GLA;MT3
15	pyrimidine nucleoside triphosphate biosynthetic process	2	21	10%	4.67E-07	RP2;TBPL1
16	positive regulation of cation channel activity	2	23	9%	1.55E-06	ADRB1;CALM1

17	response to X-ray	2	24	8%	2.49E-06	ERCC8;BRCA2
18	positive regulation of blood vessel endothelial cell migration	2	25	8%	4.29E-06	FGF2;PRKD1
19	fear response	2	27	7%	9.95E-06	ADRB1;GABRA5
20	regulation of wound healing	4	94	4%	1.30E-05	CASK;GP5;FGF2;EDN1
21	negative regulation of stress-activated protein kinase signaling cascade	2	29	7%	2.48E-05	HDAC3;MPP2
22	regulation of glycoprotein biosynthetic process	2	29	7%	2.54E-05	ACAT1;MT3
23	brown fat cell differentiation	2	30	7%	2.83E-05	RGS2;ADRB1
24	positive regulation of osteoblast differentiation	3	62	5%	3.17E-05	BMPR1B;FGF2;PRKD1
25	positive regulation of DNA repair	2	30	7%	3.31E-05	ERCC8;MPP2
26	protein kinase B signaling	2	30	7%	3.43E-05	RPS6KB2;MT3
27	negative regulation of endocytosis	2	30	7%	3.43E-05	PRKD1;PRTN3
28	eye photoreceptor cell development	2	31	6%	4.95E-05	CRB1;BHLHE23
29	activation of adenylate cyclase activity	2	33	6%	7.98E-05	ADRB1;CALM1
30	positive regulation of muscle contraction	2	33	6%	9.09E-05	RGS2;EDN1
31	regulation of cardiac muscle contraction	3	70	4%	9.79E-05	RGS2;ADRB1;CALM1
32	regulation of cytokinesis	2	35	6%	1.49E-04	BRCA2;CALM1
33	regulation of multicellular organism growth	3	81	4%	4.75E-04	HDAC3;ADRB1;GDF5
34	negative regulation of purine nucleotide metabolic process	2	42	5%	6.16E-04	CPEB2;EDN1
35	connective tissue development	5	198	3%	6.26E-04	BMPR1B;FGF2;EDN1;ACAT1;GDF5

36	regulation of phospholipase activity	3	87	3%	7.38E-04	RGS2;FGF2;CALM1
37	regulation of cell development	10	590	2%	8.27E-04	FGF2;EDN1;NELF;ADRB1;LRRC4C;PRKD1;MT3;PREX1;BHLHE23; GDF5
38	regulation of G-protein coupled receptor protein signaling pathway	4	140	3%	8.33E-04	RGS2;EDN1;CALM1;KLK5
39	dorsal/ventral pattern formation	3	94	3%	1.37E-03	DISP1;BMPR1B;EDN1
40	negative regulation of epithelial cell proliferation	3	96	3%	1.62E-03	CASK;BRCA2;GDF5
41	cellular response to abiotic stimulus	5	223	2%	2.26E-03	TNFRSF8;TNFRSF10A;NELF;SNCG;CALM1
42	double-strand break repair via homologous recombination	2	52	4%	2.82E-03	BRCA2;REC8
43	regulation of pH	2	52	4%	2.89E-03	EDN1;ATP5B
44	inositol phosphate metabolic process	2	56	4%	4.02E-03	FGF2;CALM1
45	regulation of cell death	16	1284	1%	4.75E-03	MST4;TNFRSF8;NLRP3;BMPR1B;FGF2;HDAC3;IL20RA;EDN1;NEL;
46	negative regulation of NF-kappaB transcription factor activity	2	62	3%	7.65E-03	ADKB1;PKKD1;GABKA3;KNPS1;M13;BHLHE23;GDF5 ZCCHC11;NLRP3
Physi	ical neglect					
1	regulation of cholesterol efflux	4	17	24%	2.91E-11	ABCA1;LRP1;CETP;APOE
2	negative regulation of multicellular organism growth	3	12	25%	2.12E-09	ANKRD26;ATXN2;BBS2
3	negative regulation of endothelial cell migration	5	31	16%	2.94E-09	CXCL13;ACVRL1;APOH;PTPRM;APOE
4	mitotic G2 DNA damage checkpoint	3	14	21%	4.08E-08	CDC2;CDK1;C10orf90
5	spermatid nucleus differentiation	3	15	20%	1.37E-07	KDM3A;HIST1H2BA;DMRTC2

6	negative regulation of endothelial cell proliferation	4	26	15%	2.25E-07	ACVRL1;APOH;PTPRM;APOE
7	receptor-mediated endocytosis	9	113	8%	1.03E-06	FCHO2;LRP12;LRP1;NEDD4;CETP;COLEC12;DNM2;LDLR;APOE
8	endocytosis	19	372	5%	2.09E-06	CDC42SE1;DNM3;STON1- GTF2A1L;ELMOD2;FCHO2;LRP12;DENND1A;ABCA1;GAPVD1;EEA 1;LRP1;NEDD4;CETP;DLG4;MRC2;COLEC12;DNM2;LDLR;APOE
9	phospholipid transport	5	45	11%	2.72E-06	ATP11B;ABCA1;CETP;LDLR;APOE
10	cerebellar cortex development	5	47	11%	4.86E-06	HSPA5;MDK;ATXN2;GLI1;NEUROD2
11	microtubule bundle formation	4	32	13%	5.31E-06	KIF20A;MAP1B;BBS2;MARK4
12	rRNA transcription	3	20	15%	8.54E-06	GTF3C3;GTF3A;CD3EAP
13	intracellular pH reduction	3	20	15%	8.70E-06	CLIC4;CLCN3;AQP11
14	G-protein coupled glutamate receptor signaling pathway	2	10	20%	1.30E-05	GRM8;GRM5
15	negative regulation of receptor-mediated endocytosis	2	10	20%	1.34E-05	ATXN2;DLG4
16	synaptic vesicle endocytosis	3	21	14%	1.51E-05	DENND1A;EEA1;DLG4
17	polyamine metabolic process	3	21	14%	1.57E-05	ADI1;AZIN1;MTAP
18	response to copper ion	3	22	14%	2.71E-05	ABP1;CDC2;CDK1
19	regulation of endothelial cell chemotaxis	2	11	18%	3.66E-05	CXCL13;HSPB1
20	receptor catabolic process	2	11	18%	3.69E-05	GPRASP1;NEDD4
21	regulation of platelet-derived growth factor receptor signaling pathway	2	11	18%	3.70E-05	LRP1;F7

22	peptidyl-glutamic acid carboxylation	2	11	18%	3.76E-05	F7;F10
23	protein localization to kinetochore	2	11	18%	3.77E-05	CDC2;CDK1
24	negative regulation of muscle cell apoptotic process	3	23	13%	4.24E-05	BAG3;NACA;APOH
25	negative regulation of signal transduction	30	814	4%	5.40E-05	GPRASP1;GPR161;ERRF11;CHRD;CX3CR1;DLK2;DACT2;LATS1;HS PB1;CRHR2;FZD1;FOXH1;HSPA5;MLLT3;DBC1;ANKRD26;GRM5;L GR4;FAM89B;MAPKAPK5;LRP1;GL11;FOXO1;DLK1;DACT1;NEDD4 ;NLRC3;GLIS2;NANOS3;APOE
26	regulation of Cdc42 protein signal transduction	3	24	13%	6.99E-05	FGD1;ABCA1;APOE
27	cellular response to inorganic substance	7	99	7%	7.71E-05	CLIC4;ABP1;ASPH;FOXO1;MT1L;CES1;NEUROD2
28	positive regulation of skeletal muscle tissue development	2	12	17%	8.57E-05	CDON;NACA
29	mitochondrial calcium ion homeostasis	2	12	17%	8.79E-05	CCDC90A;ATP2A1
30	histone H3-K4 trimethylation	2	12	17%	8.80E-05	MLL2;ARID4A
31	methionine biosynthetic process	2	12	17%	8.81E-05	ADI1;MTAP
32	cell migration involved in gastrulation	2	12	17%	8.85E-05	FOXH1;MESP2
33	lipoprotein metabolic process	7	100	7%	8.85E-05	HHAT;ABCA1;LRP1;CETP;MPPE1;LDLR;APOE
34	negative regulation of smoothened signaling pathway	3	25	12%	1.02E-04	GPR161;CHRD;GLIS2
35	transcription from RNA polymerase III promoter	4	42	10%	1.44E-04	GTF3C3;POLR3G;CRCP;GTF3A
36	histone H2A acetylation	2	13	15%	1.77E-04	DMAP1;BRD8

**OS7:** Enriched biological processes for genes associated with epigenetic variation that is shared across maltreatment types (physical abuse, sexual abuse and neglect)

GO	Pathway	N genes in input list	Totan N in pathway	%	<i>p</i> -value	Genes
1	regulation of	133	530	25.09	4.20E-10	DMD;MECP2;NLGN3;SOX3;DAB1;NTRK1;VANGL2;WNT3A;ARHGEF2;LMX1A;LRP8;AGRN;DISC1;SK
	nervous system					I;TNR;NGF;ABL2;CNTN2;REN;HOXD3;OBSL1;GLI2;SOX11;RTN4;NRXN1;SERPINE2;BCL11A;STK25;I
	development					TM2C;SHOX2;PTPRG;EPHB3;ROBO1;ETV5;WNT5A;GHSR;SKIL;ODZ3;SFRP2;NKX6;FGFR3;HTT;SE
						MA6A;ACSL6;NEUROG1;C5orf13;ZSWIM6;OTP;IL6ST;THBS2;SYNGAP1;TFAP2A;PACSIN1;FYN;KIAA
						0319;GLI3;LIMK1;YWHAG;COBL;PTK2B;STK3;PTK2;STMN2;ZHX2;GDF6;TG;LZTS1;NOTCH1;DAB2I
						P;NKX62;KNDC1;CAMK1D;TCF7L2;ITGB1;VAX1;CPEB3;ZNF488;SLIT1;PTEN;PAX2;PAX6;BDNF;OD
						Z4;HCCA2;ASCL2;PVRL1;RELA;CEND1;LHX5;LRP6;FBXW8;FKBP4;SSH1;ULK1;RNF10;STK24;EDNR
						B;PRKCH;FOXA1;BCL11B;AKT1;SIX4;SNW1;SEMA7A;NPTN;LINGO1;ZFHX3;MAPK8IP3;AATK;HOXB
						3;ACCN1;SEZ6;NTN1;STAT3;MED1;TGIF1;ADCYAP1;MBP;BCL2;CACNA1A;PALM;APC2;PLK5P;TCF
						3;NOTCH3;BMP7;FOXA2;CDK5RAP1;PLXNB2;EP300;CELSR1;TSPO;YPEL1

2 regulation of growth

1468

287

19.55

9.48E-10 BCOR;FRMPD4;SHROOM2;RPS6KA3;DMD;MECP2;MID11P1;NLGN3;SH3KBP1;GPM6B;DDX3X;FOX P3;FHL1;AMOT;TMSB15B;DAB1;NTRK1;S100A13;ARF1;GNG4;VANGL2;WNT3A;SGIP1;ARHGEF2;PR KCZ;ARHGEF16;ARHGEF10L;LMX1A;CDC42SE1;LRP8;ESPN;AGRN;CNST;DISC1;MACF1;ASAP3;SL C9A1:TNR:NGF:PEX14:ABL2:CDKN2C:CNTN2:TPR:IGFBP5:LRPPRC:OBSL1:CTNNA2:BIN1:SH3YL1: EML4;RTN4;ALMS1;NRXN1;SH3BP4;SERPINE2;BCL11A;SPTBN1;NCK2;STK25;AAK1;MERTK;ITM2C; SPAST;CGREF1;RALB;SHOX2;PTPRG;TNK2;EPHB3;WWTR1;HYAL2;BCL6;PFN2;ROBO1;ETV5;WNT5 A;GHSR;DPPA2;SKIL;AHSG;ZNF639;GPX1;CTBP1;CAMK2D;MSX1;ODZ3;SHROOM3;SFRP2;FGFR3; APBB2;ODZ2;RUFY1;SLIT3;ACSL6;SCGB3A1;NEUROG1;FGFR4;PLK2;CSF1R;MYO10;TPPP;THBS2;S YNGAP1;TRIM40;DST;FBX05;JARID2;PARK2;SRF;PACSIN1;DDR1;PHF1;PKHD1;FYN;CDKN1A;KIA A0319:FOXK1:MAD1L1:PDGFA:PODXL:CAPZA2:LIMK1:YWHAG:IGFBP3:HOXA13:ARPC1A:GNA12: ELN;TAC1;COBL;PTK2B;ARHGEF10;STK3;MYC;ANK1;PTK2;NSMCE2;NRG1;DLC1;STMN2;CLU;ARC ;TNKS;TRPS1;LZTS1;SOX17;NOTCH1;DAB2IP;GSN;COL5A1;KNDC1;MTG1;ATAD1;CAMK1D;TCF7L2 ;ITGB1;GDF2;CPEB3;SORBS1;PRKCQ;HTRA1;SLIT1;NET1;NRG3;DKK1;PTEN;PAX2;UBE2D1;TNKS2 ;APOA1;RASGRP2;BDNF;PACSIN3;MLL;HCCA2;LRP5;PVRL1;TUB;WT1;BCL9L;CDC42EP2;NCAM1;J AM3;CDCA5;CCDC85B;EPS8;FBXW8;ESPL1;STK38L;LRP1;HRK;FKBP4;FGD6;DYNLL1;SSH1;ANAPC 5:CDKN1B:ULK1:RAPGEF3:NFE2:RB1:TBC1D4:STK24:IFT88:CDC16:PRKCH:KIF26A:FOXA1:INF2:H SP90AA1;AKT1;TRIM9;SIX4;C14orf49;SNW1;THBS1;CYFIP1;SPTBN5;ATP10A;SEMA7A;SMAD6:TPM1: EHD4;NPTN;C15orf23;MAP2K5;NUSAP1;MFGE8;GREM1;LINGO1;FAM38A;CYBA;TSC2;CDH13;FHO D1;CFDP1;SH2B1;ZNF205;ABR;GAS7;COL1A1;MYO1C;TBCD;AATK;SREBF1;MYH10;DLG4;ACCN1;S EZ6;NTN1;TEX14;TADA2A;AXIN2;TRIM37;HNF1B;CDC27;FLCN;XAF1;ADCYAP1;MBP;ACAA2;NEDD 4L;BCL2;CACNA1A;LTBP4;PALM;APC2;C3;INSR;KIAA1543;PLK5P;SMARCA4;MYO1F;BMP7;FOXA2; GTPBP5;L3MBTL;PPP1R16B;NAPB;C21orf66;BRWD1;PLXNB2;EP300;TRIOBP;CELSR1;NF2

3

anatomical

morphogenesis

structure

10

379

1960 19.34

4.54E-08 EFNB1;BCOR;SHROOM2;DLG3;RPS6KA3;DMD;HPRT1;POU3F4;COL4A5;NLGN3;FLNA;SOX3;SLITR K4;FHL1;AMOT;ARX;EDA;L1CAM;DAB1;NTRK1;GNPAT;KIF26B;RYR2;VANGL2;WNT3A;SDCCAG8;G REM2;ARHGEF2;SHC1;ENAH;TBX15;LMX1A;LHX4;KCNO4;LRP8;OBSCN;AGRN;MACF1;SKI;TNR;N GF:SOX13:ABL2:ARTN:CNTN2:NFASC:VAV3:TNN:ZNF281:FHL2:HOXD3:IGFBP5:ZEB2:HOXD8:OBS L1;CTNNA2;ATP6V1B1;GLI2;SOX11;EPAS1;WNT10A;EVX2;RTN4;ANTXR1;NRXN1;OTX1;SLC8A1;ACV R1;HOXD11;SERPINE2;TFCP2L1;EN1;SPTBN1;HOXD9;NCK2;FEZ2;CYP26B1;CREB1;MERTK;CNGA3 :SPAST:SHOX2:FOXP1:TTLL3:SLITRK3:ATG7:THRB:EPHB3:MYLK:WWTR1:HYAL2:BCL6:PROK2:SIA H2:TNNC1:ROBO1:WNT5A:ATR:PAK2:SKIL:IFT122:GPX1:WHSC1:FGFRL1:ABLIM2:PRDM8:IDUA:M SX1:SHROOM3:PITX2:SFRP2:NKX61:UGT8:FGFR3:NPY2R:FAT1:APBB2:FRAS1:HTT:PDGFC:CC2D2 A:ODZ2:IRX4:SEMA6A:SLC12A2:SLIT3:TRIO:FGF18:NEUROG1:PPARGC1B:CSF1R:MYO10:ZSWIM6: HRH2:POU4F3:IRX2:NDST1:CSNK2B:VPS52:T:COL11A2:RPS6KA2:DST:RING1:TFAP2A:GPLD1:FOX P4;SRF;PACSIN1;DTNBP1;DDR1;PKHD1;NOTCH4;HSP90AB1;RPL10A;FYN;FLOT1;ETV7;RADIL;DN AJB6;PDGFA;PODXL;ZP3;GLI3;ZPBP;FAM20C;LIMK1;HIPK2;GBX1;HOXA13;C7orf51;ADCY1;TSPAN 12;GNA12;ELN;LFNG;WNT2;HOXA9;COBL;HOXA6;SOSTDC1;PTK2B;KIAA1688;STK3;SCRIB;EXT1;M YC;ANK1;PTK2;DUSP4;NRG1;LOXL2;DLC1;BMP1;SOX17;CHD7;RDH10;NOTCH1;VAV2;DAB2IP;PTC H1:ROR2:GSN:COL5A1:NTNG2:KNDC1:TCF7L2:ITGB1:VAX1:GDF2:FOXI2:ABLIM1:HMX3:PRKCO:C OL13A1:GFRA1:SLIT1:NRG3:DKK1:PTEN:PAX2:ARHGAP22:UNC5B:PAX6:ALX4:LTBP3:BDNF:ODZ4: MY07A;FLI1;BRSK2;BSX;INPPL1;LRP5;PVRL1;BARX2;HTATIP2;LRP4;WT1;AP2A2;ZBTB16;RELA;NC AM1:JAM3:CEND1:LHX5:APAF1:KDM2B:LRP6:HOXC13:NCOR2:LRP1:PRICKLE1:MYF6:CACNA1C: HOXC11;FRS2;PXN;DYNLL1;SSH1;HOXC9;NAB2;ULK1;UNC119B;IFT81;RAPGEF3;ATP8A2;FARP1;R B1;CDX2;COL4A2;PCDH8;SMAD9;ZIC5;IFT88;MEG3;FOXA1;BCL11B;SLC24A4;JAG2;HSP90AA1;BR F1;RPGRIP1;VSX2;ESRRB;SIX4;LRP10;NR2F2;THBS1;CYFIP1;IGF1R;SPTBN5;RORA;SEMA7A;RGMA; MEGF11;TPM1;ACAN;MFGE8;GREM1;ANKRD11;CACNA1H;CREBBP;IFT140;TSC2;MYH11;ZFPM1; MAPK8IP3;E2F4;CDH13;FOXF1;TUBB3;PSMB10;ABR;TNFSF12;PLCD3;GAS7;COL1A1;MY01C;TBX4 ;MY015A;MYH10;HOXB3;DLG4;NTN1;FOXJ1;STAT3;AXIN2;CCDC40;NPTX1;HNF1B;WNK4;MED1;Z PBP2;TGIF1;NFATC1;ONECUT2;RAX;SALL3;BCL2;LAMA1;CACNA1A;MYO9B;RYR1;INSR;NFIC;B9D2 ;PIP5K1C;TCF3;C19orf20;TRIM28;C19orf51;TNNI3;VASP;NOTCH3;ERCC2;SMARCA4;NRTN;TGM3;G NAS:COL9A3:BMP7:FOXA2:BTBD3:MYLK2:LAMA5:PARD6B:TFAP2C:OTOR:RIPK4:PCNT:COL18A1: PLXNB2:CACNA11:EP300:TBX1:CELSR1:NF2:MAP3K7IP1

#### 4 neurogenesis

274

1283 21.36

5.34E-08 EFNB1;DLG3;RPS6KA3;DMD;MECP2;HPRT1;POU3F4;COL4A5;NLGN3;CDK16;SOX3;SLITRK4;ARX; MCF2;L1CAM;DAB1;NTRK1;PRDM16;VANGL2;WNT3A;ARHGEF2;ENAH;LMX1A;LHX4;LRP8;AGRN; NAV1;DISC1;SK1;TNR;NGF;ABL2;CDKN2C;ARTN;CNTN2;NFASC;TNN;POU3F1;REN;HOXD3;OBSL1; CTNNA2;GLI2;SOX11;WNT10A;RTN4;NRXN1;SERPINE2;BCL11A;EN1;SPTBN1;HOXD9;NCK2;FEZ2;S TK25;DNER;CREB1;CNGA3;ITM2C;SPAST;SHOX2;FOXP1;SLITRK3;ATG7;PTPRG;EPHB3;DGKG;KLF 15;SIAH2;ROBO1;ETV5;WNT5A;PAK2;HDAC11;SKIL;HELT;ABLIM2;PRDM8;ODZ3;PITX2;NKX6-1;UGT8;FGFR3;APBB2;HTT;ODZ2;SEMA6A;SLIT3;TRIO;ACSL6;NEUROG1;C5orf13;MYO10;ZSWIM6; POU4F3;OTP;IL6ST;CSNK2B;SYNGAP1;RPS6KA2;PRDM13;DST;TFAP2A;SRF;PACSIN1;DTNBP1;HSP 90AB1;FYN;FLOT1;MDGA1;KIAA0319;DAGLB;CUX1;GLI3;LIMK1;YWHAG;HIPK2;GBX1;C7orf51;AD CY1:WNT2:OGDH:COBL:PTK2B:ARHGEF10:NRBP2:KIAA1688:STK3:SCRIB:EXT1:ANK1:PTK2:RP1L1 ;NRG1;STMN2;ZHX2;GDF6;LZTS1;NOTCH1;VAV2;DAB2IP;PTCH1;GSN;COL5A1;NTNG2;NKX6-2;KNDC1;KCNMA1;CAMK1D;TCF7L2;ITGB1;PSD;TACC2;VAX1;CHAT;CPEB3;ABLIM1;PRKCO;GFR A1;ZNF488;SLIT1;PTEN;PAX2;UNC5B;PAX6;APOA1;DBX1;OPCML;NTM;BDNF;ODZ4;MYO7A;HCCA 2;BRSK2;KIRREL3;ASCL2;PVRL1;DAGLA;AP2A2;RELA;NCAM1;CEND1;LHX5;LRP6;FBXW8;PHLDA1 ;FKBP4;CACNA1C;FRS2;EIF2B1;SSH1;NAB2;ULK1;RAB35;RNF10;ATP8A2;FARP1;RB1;STK24;COL4A 2;EDNRB;PRKCH;MEG3;FOXA1;BCL11B;JAG2;RTN1;HSP90AA1;RPGRIP1;AKT1;VSX2;SIX4;SNW1;N R2F2:CYFIP1:IGF1R:SPTBN5:RORA:SEMA7A:RGMA:NPTN:LINGO1:CACNA1H:GRIN2A:ZFHX3:MYH 11;MAPK8IP3;NDE1;TUBB3;GAS7;AATK;MYH10;HOXB3;DLG4;SEZ6;NTN1;RTN4RL1;STAT3;NPTX1; MED1;TGIF1;ADCYAP1;MBP;ONECUT2;SALL3;BCL2;LAMA1;CACNA1A;PALM;APC2;PIP5K1C;PLK5 P;TCF3;VASP;NOTCH3;ERCC2;SMARCA4;NRTN;PIGT;COL9A3;BMP7;FOXA2;BTBD3;CDK5RAP1;PA RD6B;TFAP2C;PCNT;NCAM2;PLXNB2;CACNA11;EP300;CELSR1;TSPO;NF2

5 neurological system process

1214

18.95

230

6.41E-08 FRMPD4;SYN1;DLG3;RPS6KA3;SYP;DMD;MECP2;POU3F4;NLGN3;CHM;GPR143;NTRK1;GNG4;KC NAB2;PRKCZ;KCNQ4;ESPN;AGRN;CACNA1E;GPR88;DISC1;HTR6;TNR;NGF;CNTN2;NFASC;ABCA4; GRIK3;GNB1;CTNNA2;ATP6V1B1;EPAS1;NRXN1;SERPINE2;EFEMP1;TRPM8;IL1RN;CREB1;CNGA3; HRH1;ATG7;THRB;EPHB3;PROK2;TNNC1;ETV5;WNT5A;GHSR;MME;GPX1;SORCS2;CAMK2D;CPLX 1;PPP3CA;PDE6B;WDR1;NPY2R;HTT;GRM6;SLC12A2;HMGCR;PLK2;SLC6A3;HRH2;POU4F3;KCNIP 1;PDE6A;GABBR1;GRM4;THBS2;COL11A2;SYNGAP1;ITPR3;RPS6KA2;SLC29A1;TFAP2A;GABRR1;SR F;PACSIN1;DTNBP1;EYS;FYN;STX7;NQO2;GRM1;DPP6;GRM8;CHRM2;DAGLB;POU6F2;PRKAR1B;K CNH2;NPTX2;GBX1;CADPS2;ADCY1;TAS2R3;TAC1;ATP6V0A4;TSNARE1;SCRIB;DLGAP2;PTK2;RP1L 1;NRG1;ARC;CHD7;RDH10;DAB2IP;NKX6-2;ATAD1;KCNMA1;CDH23;CHAT;CPEB3;ABLIM1;HMX3;CAMK2G;GAD2;SLIT1;NET1;PTEN;PAX2;S ORCS1;KCNQ1;SLC6A5;PAX6;ABCC8;BDNF;MY07A;CALCA;PVRL1;DAGLA;MTNR1B;DRD4;SHANK2 ;TUB;AP2A2;CCKBR;RIC3;FOSL1;JAM3;KCNA4;OR52D1;LRP6;PRR4;AAAS;VAMP1;CACNA1C;GRIN2 B;KCNA6;CDKN1B;LIN7A;KCNH3;ALDH2;ATP8A2;FGF14;PCDH8;HTR2A;EDNRB;PRIMA1;KCNK10; SLC24A4;RPGRIP1;CPNE6;VSX2;TRIM9;SSTR1;SIX4;GPR176;APBA2;IGDCC3;RLBP1;GABRG3;CDH8 ;A2BP1;ADCY9;GRIN2A;JPH3;CDH3;NOB1;ABR;COL1A1;MY015A;MYH10;DLG4;ACCN1;SEZ6;KCN

;A2BP1;ADCY9;GRIN2A;JPH3;CDH3;NOB1;ABR;COLIA1;MY015A;MYH10;DLG4;ACCN1;SEZ6;KCN H4;CHRNB1;CAMKK1;CACNG4;NPTX1;DLGAP1;ADCYAP1;BRUNOL4;MBP;LOXHD1;RAX;KCNG2;C ACNA1A;APC2;UNC13A;GNG8;AP3D1;KCNJ14;PDE4A;PIP5K1C;GNG7;KPTN;GRIN2D;C19orf20;TNN I3;CEACAM16;GNAS;DNAJC5;MYLK2;NAPB;TMC2;OTOR;COL18A1;KCNJ6;NCAM2;CACNA11;COMT ;TBX1;TSPO;PANX2;SYN3;RASD2

6	organ morphogenesis	171	791	21.62	1.22E-07	BCOR;SHROOM2;POU3F4;SOX3;FHL1;EDA;KIF26B;RYR2;VANGL2;WNT3A;TBX15;LHX4;KCNQ4;SK I;FHL2;HOXD3;IGFBP5;HOXD8;CTNNA2;ATP6V1B1;GL12;SOX11;WNT10A;RTN4;OTX1;SLC8A1;ACV R1;HOXD11;HOXD9;CYP26B1;CNGA3;SHOX2;THRB;MYLK;HYAL2;TNNC1;WNT5A;IFT122;WHSC1;F GFRL1;IDUA;MSX1;PITX2;SFRP2;NKX6- 1;FGFR3;NPY2R;HTT;PDGFC;IRX4;SEMA6A;SLIT3;FGF18;NEUROG1;PPARGC1B;POU4F3;IRX2;ND ST1;T;COL11A2;RING1;TFAP2A;SRF;ETV7;PDGFA;GL13;FAM20C;HIPK2;HOXA13;TSPAN12;ELN;LF NG;WNT2;HOXA6;SOSTDC1;PTK2;NRG1;DLC1;BMP1;SOX17;CHD7;RDH10;NOTCH1;PTCH1;ROR2; COL5A1;TCF7L2;ABLIM1;HMX3;COL13A1;SLIT1;PTEN;PAX2;PAX6;ALX4;LTBP3;MYO7A;FLI1;INPP L1;LRP5;PVRL1;BARX2;LRP4;WT1;RELA;KDM2B;LRP6;HOXC13;NCOR2;MYF6;HOXC11;FRS2;HOXC 9;NAB2;CDX2;IFT88;FOXA1;BCL11B;SLC24A4;JAG2;RPGRIP1;VSX2;SIX4;LRP10;THBS1;IGF1R;MEG F11;TPM1;ACAN;GREM1;ANKRD11;ZFPM1;MAPK8IP3;E2F4;FOXF1;ABR;COL1A1;TBX4;MY015A;H OXB3;NTN1;FOXJ1;STAT3;AXIN2;CCDC40;HNF1B;WNK4;MED1;ONECUT2;RAX;BCL2;LAMA1;RYR1;
						F11;TPM1;ACAN;GREM1;ANKRD11;ZFPM1;MAPK8IP3;E2F4;FOXF1;ABR;COL1A1;TBX4;MYO15A;H OXB3;NTN1;FOXJ1;STAT3;AXIN2;CCDC40;HNF1B;WNK4;MED1;ONECUT2;RAX;BCL2;LAMA1;RYR1; INSR;NFIC;TNNI3;SMARCA4;TGM3;GNAS;BMP7;FOXA2;MYLK2;LAMA5;OTOR;PCNT;COL18A1;EP3 00;TBX1;CELSR1;NF2;MAP3K7IP1

7 1	regulation of	8	13	61.54	2.85E-07	POU3F4;MSX1;HOXA13;PAX2;WT1;HNF1B;BMP7;TBX1
1	mesenchymal cell					
á	apoptotic process					
8 1	regulation of glial	9	15	60.00	4.00E-07	SKI;SOX11;ETV5;NOTCH1;ASCL2;RNF10;PRKCH;ADCYAP1;TSPO

9	tube development	108	465	23.23	2.32E-06	FOXP3;EDA;KIF26B;RYR2;VANGL2;WNT3A;SDCCAG8;ENAH;SKI;ZEB2;GLI2;SOX11;EPAS1;SLC8A1; ACVR1;HOXD11;CREB1;FOXP1;THRB;ROBO1;WNT5A;DPPA2;IFT122;SHROOM3;PITX2;SFRP2;FGF R3;ADAMTS2;SLC12A2;FGF18;FGFR4;CSF1R;LOX;IRX2;CSNK2B;T;TFAP2A;FOXP4;SRF;DDR1;NOT CH4;BAT3;PDGFA;PODXL;CUX1;GLI3;CFTR;WNT2;COBL;SOSTDC1;STK3;SCRIB;MYC;DLC1;NKX26 ;SOX17;CHD7;RDH10;NOTCH1;DAB21P;PTCH1;GDF2;PAX2;LTBP3;BDNF;LRP5;WT1;INSC;APAF1;K DM2B;LRP6;NCOR2;PRICKLE1;PXN;SMAD9;ZIC5;IFT88;FOXA1;SMAD6;RGMA;CYP1A2;GREM1;TSC 2;CRISPLD2;MAPK8IP3;FOXF1;TBX4;SREBF1;NTN1;FOXJ1;CCDC40;HNF1B;WNK4;MED1;TGIF1;B CL2;LAMA1;CRLF1;VASP;BMP7;FOXA2;LAMA5;TFAP2C;PLXNB2;EP300;TBX1;CELSR1;MAP3K7IP1
10	anterior/posterior pattern specification	54	200	27.00	3.60E-06	VANGL2;WNT3A;SKI;HOXD3;ZEB2;HOXD8;OTX1;HOXD11;EN1;HOXD9;WNT5A;MSX1;SFRP2;HTT;T ;RING1;SRF;GLI3;HIPK2;LFNG;HOXA9;COBL;HOXA6;ARC;SOX17;NOTCH1;ROR2;DKK1;PAX6;ALX 4;MLL;LRP5;WT1;ZBTB16;KDM2B;LRP6;HOXC13;MYF6;HOXC11;FRS2;HOXC9;CDX2;PCDH8;IFT88 ;NR2F2;FOXF1;HOXB3;AXIN2;HNF1B;TCF3;FOXA2;EP300;TBX1;CELSR1
11	muscle structure development	95	426	22.30	6.85E-06	DMD;FOXP3;FHL1;RYR2;WNT3A;OBSCN;SKI;SLC9A1;TAGLN2;FHL2;IGFBP5;OBSL1;CTNNA2;SOX1 1;BIN1;EPAS1;SLC8A1;ACVR1;SPEG;HOXD9;DNER;MAL;SHOX2;FOXP1;CACNA2D2;FLNB;ATG7;MY LK;TNNC1;WNT5A;SKIL;GPX1;FGFRL1;MSX1;PPP3CA;PITX2;HMGCR;SGCD;SYNE1;T;UTRN;SRF;C OL19A1;FOXK1;SGCE;FLNC;ELN;WNT2;NRG1;NKX26;MSC;NOTCH1;TCF7L2;ITGB1;VAX1;CHAT;AL X4;BARX2;WT1;USP2;BCL9L;HINFP;MYF6;RB1;EDNRB;ASB2;AKT1;SIX4;NR2F2;TPM1;GREM1;CAC NA1H;ZFHX3;MYH11;ZFPM1;CDH15;FOXF1;CDH3;MYH10;CHRNB1;MYH3;MAP2K6;MED1;RYR1;T CF3;TNNI3;SMARCA4;MYLK2;LAMA5;SIK1;C21orf66;EP300;TBX1;SMTN;MAPK12
12	neural tube closure	26	76	34.21	7.09E-06	VANGL2;ENAH;SKI;ZEB2;WNT5A;IFT122;SHROOM3;SFRP2;T;TFAP2A;COBL;SCRIB;DLC1;PTCH1;P AX2;APAF1;KDM2B;LRP6;PRICKLE1;ZIC5;RGMA;TSC2;TGIF1;VASP;PLXNB2;CELSR1
13	positive regulation of cell-cell adhesion mediated by integrin	6	10	60.00	8.39E-06	WNT3A;WNT5A;PODXL;FOXA1;FAM38A;FOXA2

14	establishment or maintenance of apical/basal cell polarity	10	22	45.45	1.01E-05	DLG3;VANGL2;ERBB2IP;SCRIB;ANK1;CDX2;FOXF1;DLG4;FOXJ1;LAMA1
15	establishment or maintenance of bipolar cell polarity	10	22	45.45	1.01E-05	DLG3;VANGL2;ERBB2IP;SCRIB;ANK1;CDX2;FOXF1;DLG4;FOXJ1;LAMA1
16	actin filament- based process	91	396	22.98	1.12E-05	IQSEC2;DMD;FLNA;AMOT;TMSB15B;NPHP4;ARHGEF2;PRKCZ;SHC1;ENAH;ESPN;OBSCN;ABL2;OB SL1;ANTXR1;NCK2;FOXP1;IQSEC1;FLNB;BCL6;PFN2;TNNC1;GHSR;ATR;PAK2;SORBS2;ABLIM2;CA MK2D;SHROOM3;FAT1;ARHGAP26;PPARGC1B;LRRC16A;EPB41L2;TNXB;SRF;PACSIN1;DTNBP1;P HACTR1;NEDD9;DNAJB6;PDGFA;ELMO1;CAPZA2;LIMK1;ARPC1A;ELN;COBL;PTK2B;DLC1;GSN;IT GB1;SORBS1;MY07A;HCCA2;BRSK2;INPPL1;CORO1B;CDC42EP2;ARHGEF17;TNNT3;TNN12;EPS8;C ORO1C;FGD6;DYNLL1;SSH1;ACTN1;INF2;CDC42BPB;CFL2;SPTBN5;TPM1;MYH11;FHOD1;ABR;GA S7;MY01C;MYH10;DLG4;FOXJ1;MYH3;BCL2;MY09B;PIP5K1C;TNNT1;KPTN;TNN13;VASP;TRIOBP; NF2
17	negative regulation of ossification	12	32	37.50	2.33E-05	BCOR;AHSG;SMURF1;PTK2B;NOTCH1;LTBP3;CALCA;LRP4;NFE2;LRP10;GREM1;BCL2
18	limb development	45	159	28.30	2.34E-05	SKI;GLI2;SOX11;EVX2;HOXD11;EN1;HOXD9;CYP26B1;SHOX2;WNT5A;IFT122;IDUA;MSX1;PITX2;SF RP2;TFAP2A;GLI3;HOXA13;GNA12;HOXA9;CHD7;RDH10;NOTCH1;PTCH1;DKK1;ALX4;LRP5;LRP4; ZBTB16;LRP6;HOXC13;HOXC11;IFT88;LRP10;NR2F2;GREM1;CREBBP;TBX4;MED1;RAX;SALL3;SMA RCA4;GNAS;BMP7;PCNT
19	appendage development	45	159	28.30	2.34E-05	SKI;GLI2;SOX11;EVX2;HOXD11;EN1;HOXD9;CYP26B1;SHOX2;WNT5A;IFT122;IDUA;MSX1;PITX2;SF RP2;TFAP2A;GLI3;HOXA13;GNA12;HOXA9;CHD7;RDH10;NOTCH1;PTCH1;DKK1;ALX4;LRP5;LRP4; ZBTB16;LRP6;HOXC13;HOXC11;IFT88;LRP10;NR2F2;GREM1;CREBBP;TBX4;MED1;RAX;SALL3;SMA RCA4;GNAS;BMP7;PCNT

20	regulation of neurological system process	74	317	23.34	2.40E-05	<i>SYP;MECP2;NLGN3;GPM6B;NTRK1;GNPAT;ARF1;PRKCZ;LRP8;HTR6;SK1;SLC9A1;TNR;NGF;CNTN2</i> ; <i>NFASC;GRIK3;POU3F1;NRXN1;SERPINE2;MAL;HRH1;CACNA2D2;PXK;PFN2;PPP3CA;UGT8;FGFR</i> <i>3;NPY2R;HTT;PLK2;HRH2;SYNGAP1;ITPR3;SRF;DTNBP1;GRM1;YWHAG;TAC1;ARHGEF10;ARC;TG;</i> <i>NKX62;ATAD1;TCF7L2;CHAT;CPEB3;PTEN;BDNF;ODZ4;CALCA;DRD4;SHANK2;JAM3;GRIN2B;NAB</i> <i>2;RNF10;HTR2A;EDNRB;AKT1;NPTN;GRIN2A;JPH3;DLG4;ADCYAP1;MBP;CACNA1A;APC2;UNC13A;</i> <i>PRX;GRIN2D;ERCC2;NAPB;YPEL1</i>
21	positive regulation of muscle cell differentiation	20	61	32.79	2.61E-05	WNT3A;CTNNA2;CYP26B1;SHOX2;FOXP1;IGFBP3;NRG1;MYF6;MEG3;SETD3;PIAS1;GREM1;ZFHX3; CDH15;CDH3;MAP2K6;BCL2;TCF3;TBX1;MAPK12
22	endothelial cell migration	16	49	32.65	2.98E-05	AMOT;PLEKHG5;ROBO1;GPX1;GPLD1;SRF;PTK2B;PTK2;LOXL2;ITGB1;PTEN;APOA1;CALCA;GRE M1;CDH13;TNFSF12
23	regulation of lipid kinase activity	13	35	37.14	3.22E-05	VAV3;IRS1;FGFR3;PTK2B;PTK2;VAV2;DAB2IP;CCKBR;CD81;RB1;FLT3;RBL1;PIK3IP1
24	regulation of gastrulation	11	27	40.74	3.82E-05	WNT3A;IL1RN;SFRP2;SOX17;DKK1;APOA1;ODZ4;MAP2K5;HNF1B;TGIF1;FOXA2
25	negative regulation of cellular component movement	39	160	24.38	4.42E-05	PTPRU;IGFBP5;IL1RN;HYAL2;SFRP2;MCC;CSNK2B;SRF;PKHD1;IGFBP3;STK3;TRIB1;DLC1;TP53IN P1;NOTCH1;DAB2IP;GDF2;PTEN;GTPBP4;MUC2;IFITM1;LRP1;CDKN1B;STK24;ACTN1;NR2F2;THB S1;TPM1;MAP2K5;GREM1;ABR;STAT3;ALOX15B;HDAC5;FLCN;BCL2;HAS1;ADARB1;NF2
26	base-excision repair AP site formation	5	10	50.00	5.66E-05	MID1;CCNO;SMUG1;MPG;MED1
27	negative regulation of growth	48	215	22.33	6.94E-05	DDX3X;FHL1;GNG4;WNT3A;TNR;CDKN2C;RTN4;ING5;SH3BP4;SERPINE2;BCL11A;CGREF1;MAL;H YAL2;BCL6;WNT5A;MSX1;SFRP2;FGFR3;APBB2;HTT;SLIT3;SCGB3A1;ADRB2;TRIM40;CDKN1A;FOX K1;STK3;PTK2;SOX17;DAB2IP;PTCH1;TCF7L2;GDF2;SLIT1;PTEN;WT1;CCDC85B;CDKN1B;ULK1;G REM1;RA11;AATK;NTN1;ALOX15B;FLCN;BCL2;SMARCA4
28	negative regulation of BMP signaling pathway	14	37	37.84	7.03E-05	SKI;TRIM33;WNT5A;SFRP2;SMURF1;HIPK2;SOSTDC1;NOTCH1;TCF7L2;HTRA1;DKK1;SMAD6;FBN1 ;GREM1
29	regulation of embryonic development	27	93	29.03	7.14E-05	AMOT;NRK;VANGL2;WNT3A;IL1RN;NR2C2;WNT5A;SFRP2;LFNG;WNT2;SOX17;NOTCH1;DKK1;APO A1;ODZ4;LRP6;MEG3;SIX4;MAP2K5;HNF1B;TGIF1;LAMA1;INSR;BMP7;FOXA2;LAMA5;CELSR1

30	multicellular organismal	10	27	37.04	7.40E-05	VPS54;TNNC1;ITPR1;HIPK2;TNNT3;TNNI2;ATP8A2;HSP90AA1;CACNA1A;TNNT1
31	movement musculoskeletal movement	10	27	37.04	7.40E-05	VPS54;TNNC1;ITPR1;HIPK2;TNNT3;TNNI2;ATP8A2;HSP90AA1;CACNA1A;TNNT1
32	hindbrain development	37	130	28.46	8.07E-05	MECP2;DAB1;GNPAT;LMX1A;SDF4;CTNNA2;GLI2;NRXN1;OTX1;SERPINE2;EN1;NR2C2;ATG7;OGD H;DLC1;KNDC1;ITGB1;PAX6;CEND1;LHX5;KDM2B;LRP6;ULK1;SMAD9;SSTR1;RORA;SEZ6L2;MYH1 0;HOXB3;SEZ6;HNF1B;BCL2;CACNA1A;PTPRS;SMARCA4;FOXA2;PCNT
33	positive regulation of skeletal muscle	6	12	50.00	9.39E-05	SHOX2;HMGCR;SOX17;USP2;MEG3;BCL2
34	hair cycle process	22	79	27.85	1.06E-04	EDA;VANGL2;IGFBP5;WNT10A;PDGFA;SOSTDC1;NOTCH1;DKK1;ALX4;BARX2;LRP4;RELA;HOXC1 3;SOX21;AKT1;LRP10;BCL2;ERCC2;TGM3;LAMA5;TFAP2C;CELSR1
35	hair follicle development	22	79	27.85	1.06E-04	EDA;VANGL2;IGFBP5;WNT10A;PDGFA;SOSTDC1;NOTCH1;DKK1;ALX4;BARX2;LRP4;RELA;HOXC1 3;SOX21;AKT1;LRP10;BCL2;ERCC2;TGM3;LAMA5;TFAP2C;CELSR1
36	molting cycle process	22	79	27.85	1.06E-04	EDA;VANGL2;IGFBP5;WNT10A;PDGFA;SOSTDC1;NOTCH1;DKK1;ALX4;BARX2;LRP4;RELA;HOXC1 3;SOX21;AKT1;LRP10;BCL2;ERCC2;TGM3;LAMA5;TFAP2C;CELSR1
37	establishment of nucleus localization	6	11	54.55	1.08E-04	SYNE1;PTK2;SLIT1;SPTBN5;MYH10;NTN1
38	viral entry into host	6	13	46.15	1.36E-04	HYAL2;ZNF639;WWP1;PVRL1;CD81;WWP2
39	cell gland development	67	285	23.51	1.45E-04	SOX3;EDA;WNT3A;HOXD3;IGFBP5;GLI2;WNT10A;IRS1;HK2;SERPINE2;TFCP2L1;HOXD9;CREB1;R OB01;WNT5A;MSX1;PITX2;SLC12A2;CSF1R;SLC6A3;HRH2;OTP;SLC29A1;DDR1;NOTCH4;GLI3;HOX A13;WNT2;HOXA9;SOSTDC1;SCRIB;NRG1;TG;NOTCH1;PTCH1;MGMT;TCF7L2;NRG3;PTEN;PAX6;A POA1;CCND1;BSX;LRP5;SF1;WT1;CCKBR;LRP6;NCOR2;FKBP4;FRS2;FOXA1;AKT1;IGF1R;FOXF1;H OXB3;NTN1;ALOX15B;MED1;ADCYAP1;BCL2;LAMA1;BMP7;LAMA5;TFAP2C;TBX1;TSPO
40	mammary gland morphogenesis	18	52	34.62	1.49E-04	IGFBP5;GLI2;ROB01;WNT5A;SLC12A2;CSF1R;DDR1;GLI3;SOSTDC1;SCRIB;PTCH1;NRG3;BSX;LRP5 ;LRP6;NTN1;MED1;TFAP2C

41	activation of MAPK activity	31	132	23.48	1.62E-04	MAP3K7IP3;IKBKG;IRAK1;SHC1;KIAA1804;ZAK;PROK2;WNT5A;MAPK10;C1QTNF2;DAXX;GRM4;M AP3K5;GRM1;GCK;NOD1;DAB2IP;DRD4;CD81;MAP4K2;FRS2;PXN;THBS1;PTPLAD1;MAP2K5;MAP K8IP3;MAP2K6;MAP4K1;INSR;LPAR2;MAP3K7IP1
42	regulation of sequence-specific DNA binding transcription factor activity	68	344	19.77	1.91E-04	MAP3K7IP3;RLIM;IKBKG;FLNA;IRAK1;FOXP3;EDA;NTRK1;WNT3A;ARHGEF2;PRKCZ;LRP8;NGF;P EX14;TRAF5;HDAC4;PELI1;MAL;WNT5A;MAPK10;NKX6- 1;NEUROG1;TRIM40;SRF;CARD11;HIPK2;WNT2;NOD1;TRIB1;CLU;DAB2IP;PTCH1;TRAF2;CAMK1D ;TCF7L2;TAF3;FOXI2;PRKCQ;PTEN;RELA;FOSL1;LRP6;ERC1;RB1;COMMD6;PRKCH;FOXA1;AKT1; TP53BP1;MAP2K5;GREM1;WWP2;FOXF1;FOXJ1;TRIM37;HDAC5;PIAS4;TCF3;CRTC1;SMARCA4;BM P7;FOXA2;UBE2V1;RIPK4;SIK1;PRMT2;EP300;MAP3K7IP1
43	negative regulation of cell fate commitment	6	13	46.15	2.01E-04	WNT3A;SFRP2;SOSTDC1;SOX17;NKX6-2;DKK1
44	multicellular organismal	20	67	29.85	2.13E-04	MECP2;PPP3CA;NPY2R;TAC1;NET1;PTEN;SLC6A5;BDNF;CALCA;DRD4;GRIN2B;EDNRB;THBS1;HD AC5;ADCYAP1;BCL2;CACNA1A;GNG7;COMT;TSPO
45	neurotransmitter biosynthetic	7	16	43.75	2.16E-04	SLC6A3;DAGLB;CHAT;GAD2;DAGLA;ALDH2;COMT
46	process proximal/distal pattern formation	12	34	35.29	2.17E-04	GL12;HOXD11;EN1;HOXD9;CYP26B1;IRX2;GL13;HOXA9;LRP4;HOXC11;LRP10;GREM1
47	regulation of skeletal muscle tissue development	20	61	32.79	2.96E-04	TSC22D3;MBNL3;HDAC4;ACVR1;CYP26B1;SHOX2;NR1D2;HMGCR;IGFBP3;SOX17;NOTCH1;TCF7L2 ;DKK1;USP2;PRICKLE1;MEG3;ZFHX3;HDAC5;BCL2;TBX1
48	primitive streak	5	10	50.00	3.04E-04	WNT5A;T;SRF;LRP6;FOXA2
49	endocytosis	72	372	19.35	3.40E-04	CLCN5;SYP;NLGN3;SH3KBP1;REPS2;SGIP1;CDC42SE1;LRP8;CD247;PEAR1;CNTN2;VAV3;BIN1;LY7 5;SH3BP4;AAK1;DNER;MERTK;TNK2;RAB7A;ITPR1;AHSG;RUFY1;CYFIP2;MY010;ADRB2;ITPR3;PA CSIN1;HSP90AB1;FYN;ELM01;LIMK1;ARPC1A;ST7;TFR2;SCRIB;PTK2;ARC;VAV2;FNBP1;MY07A;CA LCA;PACSIN3;INPPL1;LRP5;LRP4;TUB;AP2A2;LRP6;COR01C;LRP1;HIP1R;ULK1;HSP90AA1;LRP10; THBS1;EHD4;MFGE8;TSC2;MY01C;DLG4;RAB34;ASGR1;UNC13A;FCH01;PIP5K1C;ABCA7;SH3GL1 ;PIK3R2;ICAM5;SNX5;RAB22A

50	regulation of sister chromatid cohesion	5	11	45.45	3.68E-04	NSMCE2;CDCA5;ESPL1;RB1;AXIN2
51	response to drug	78	395	19.75	3.70E-04	EBP;DAB1;NTRK1;GNPAT;PDE4B;LRP8;HTR6;SLC9A1;NGF;VAV3;BCAR3;REN;ABCG5;HDAC4;IL1R N;CREB1;FOSL2;ATR;PPP3CA;SFRP2;NKX61;HTT;CCN0;SLC6A3;LOX;RPP21;GPLD1;FYN;CDKN1A ;MCM7;PDGFA;KCNH2;ADCY1;CFTR;GNA12;PTK2B;MYC;PTCH1;MGMT;ITGB1;GAD2;NET1;PTEN; KCNQ1;SLC6A5;APOA1;CCND1;BDNF;RELA;FOSL1;CDKN1B;PARP4;HTR2A;THBS1;GABRG3;CYP1 A2;GRIN2A;ABCA3;CYBA;CDH3;SREBF1;RPH3AL;STAT3;HDAC5;MAP2K6;TGIF1;ADCYAP1;BCL2;A PC2;PDE4A;TCF3;TBXA2R;GNAS;NFATC2;COL18A1;COMT;EP300;TSPO
52	myoblast differentiation	12	35	34.29	3.88E-04	EPAS1;HMGCR;T;SRF;NRG1;NOTCH1;TCF7L2;HINFP;RB1;ASB2;GREM1;MAPK12
53	negative regulation of cell motility	36	156	23.08	4.10E-04	PTPRU;IGFBP5;IL1RN;HYAL2;SFRP2;MCC;CSNK2B;SRF;IGFBP3;STK3;TRIB1;DLC1;TP53INP1;NOT CH1;DAB2IP;GDF2;PTEN;GTPBP4;MUC2;IFITM1;LRP1;STK24;NR2F2;THBS1;TPM1;MAP2K5;GREM 1;ABR;STAT3;ALOX15B;HDAC5;FLCN;BCL2;HAS1;ADARB1;NF2
54	activation of transmembrane receptor protein tyrosine kinase	5	11	45.45	4.25E-04	PDGFC;ADRB2;NRG1;NRG3;GREM1
55	response to growth factor	114	569	20.04	4.42E-04	RPS6KA3;NGFRAP1;IRAK1;MCF2;ELK1;L1CAM;NTRK1;ARHGEF2;PRKCZ;ARHGEF16;SHC1;LRP8;P IK3CD;OBSCN;PLEKHG5;SK1;TRIM33;NGF;VAV3;CGN;FMOD;TPR;ARHGEF4;WNT10A;RTN4;IRS1;A CVR1;CREB1;RALB;KALRN;ACAP2;WWTR1;STAG1;HYAL2;WNT5A;ITPR1;SKIL;FGFRL1;FGFR3;TRI O;FGF18;NRG2;FGFR4;NDST1;ITPR3;RPS6KA2;FYN;CDKN1A;TBC1D7;SMURF1;MCM7;PDGFA;PR KAR1B;HIPK2;ADCY1;WNT2;MYC;DUSP4;NRG1;STMN2;NOTCH1;VAV2;DAB2IP;ITGB1;NET1;PTEN; PAX2;UBE2D1;LTBP3;AP2A2;RELA;ARHGEF17;NCOR2;FRS2;CDKN1B;RAB35;MCF2L;COL4A2;SMA D9;ARHGEF7;FLT3;AKT1;THBS1;SMAD6;MEIS2;EHD4;MAP2K5;LINGO1;ADCY9;TSC2;TNRC6A;PAR D6A;E2F4;ABR;SMURF2;COL1A1;MYO1C;STAT3;MED1;TGIF1;ADCYAP1;NEDD4L;LTBP4;INSR;PLK 5P;HAS1;SHC2;PIK3R2;RBL1;GATA5;TBX1;MAPK12;TNRC6B;MAP3K7IP1

56	negative regulation of cell-cell adhesion	10	31	32.26	4.53E-04	TNR;SERPINE2;IL1RN;PODXL;PTK2;GTPBP4;APOA1;MAP2K5;B4GALNT2;NF2
57	extracellular structure organization	71	321	22.12	4.66E-04	DMD;COL4A5;GPM6B;WNT3A;NID1;AGRN;LEPRE1;TNR;OLFML2B;FMOD;PXDN;NRXN1;EFEMP1;I TGB5;ABI3BP;COL29A1;SFRP2;APBB2;COL23A1;SH3PXD2B;ADAMTS2;LOX;SMOC2;COL12A1;COL1 1A2;DST;TNXB;TFAP2A;COL19A1;DDR1;DNAJB6;PDGFA;ELN;CTSB;PTK2;LOXL2;BMP1;COL27A1; COL5A1;ITGB1;COL13A1;TLL2;LTBP3;LRP4;WT1;NCAM1;COL4A2;ACTN1;LRP10;THBS1;FBN1;ACA N;GREM1;MYH11;MMP15;CRISPLD2;FOXF1;COL1A1;MYO1C;TEX14;LAMA1;LTBP4;GLT25D1;PTPR S;HAS1;ERCC2;SMARCA4;COL9A3;BMP7;LAMA5;COL18A1
58	asymmetric protein	7	17	41.18	5.29E-04	SHROOM2;VANGL2;MAL;SHROOM3;ERBB2IP;SCRIB;CELSR1
59	localization positive regulation of lipid biosynthetic process	14	47	29.79	5.39E-04	MID11P1;CREB1;ACSL6;GPLD1;ZP3;SORBS1;SIRT4;HTR2A;AKT1;IGF1R;SREBF1;NR1D1;APC2;ABC G1
60	retina development in camera-type eye	29	113	25.66	5.54E-04	NPHP4;SKI;GNB1;MERTK;CNGA3;PDE6B;LPCAT1;GRM6;PDE6A;TFAP2A;HIPK2;TSPAN12;CHD7;P AX2;LRP5;PVRL1;DRD4;TUB;RRM1;LRP6;MDM1;RPGRIP1;VSX2;MEGF11;MYH10;MED1;TGIF1;LA MA1;SMARCA4
61	cell-type specific apoptotic process	22	83	26.51	5.72E-04	RYR2;NGF;GNB1;HTT;POU4F3;RPS6KA2;TNFRSF21;GLI3;STK3;MYC;GDF6;DAB2IP;APAF1;RB1;JA G2;HSP90AA1;AKT1;AATK;BCL2;PLK5P;PIGT;TBX1
62	positive regulation of B cell	11	38	28.95	5.88E-04	IL13RA1;WNT3A;VAV3;PELI1;MAL;BCL6;CDKN1A;CARD11;BCL2;TCF3;NFATC2
63	oligodendrocyte development	12	37	32.43	7.14E-04	CNTN2;SOX11;HDAC11;FGFR3;GSN;NKX6-2;TCF7L2;ZNF488;PTEN;ODZ4;EIF2B1;ERCC2
64	negative regulation of smoothened signaling pathway	9	25	36.00	7.43E-04	REN;SERPINE2;IFT122;FGFR3;GLI3;PTCH1;RB1;KIF7;SALL3

65	extracellular matrix organization	70	320	21.88	7.60E-04	DMD;COL4A5;GPM6B;WNT3A;NID1;AGRN;LEPRE1;TNR;OLFML2B;FMOD;PXDN;NRXN1;EFEMP1;I TGB5;ABI3BP;COL29A1;SFRP2;APBB2;COL23A1;SH3PXD2B;ADAMTS2;LOX;SMOC2;COL12A1;COL1 1A2;DST;TNXB;TFAP2A;COL19A1;DDR1;DNAJB6;PDGFA;ELN;CTSB;PTK2;LOXL2;BMP1;COL27A1; COL5A1;ITGB1;COL13A1;TLL2;LTBP3;LRP4;WT1;NCAM1;COL4A2;ACTN1;LRP10;THBS1;FBN1;ACA N;GREM1;MYH11;MMP15;CRISPLD2;FOXF1;COL1A1;MYO1C;LAMA1;LTBP4;GLT25D1;PTPRS;HAS1 ;ERCC2;SMARCA4;COL9A3;BMP7;LAMA5;COL18A1
66	ventral spinal cord	6	11	54.55	7.85E-04	GLI2;NKX6-1;GLI3;NKX6-2;PAX6;DBX1
	interneuron					
67	cell fate specification involved in pattern	6	11	54.55	7.85E-04	GLI2;NKX6-1;GLI3;NKX6-2;PAX6;DBX1
68	neural tube patterning	12	34	35.29	9.64E-04	WNT3A;GLI2;EN1;IFT122;GLI3;SOX17;PTCH1;PAX6;KDM2B;LRP6;FOXA1;FOXA2
69	receptor clustering	11	28	39.29	9.93E-04	FLNA;AGRN;NRXN1;ETV5;IFT122;DOK7;SYNGAP1;LRP4;LRP10;DLG4;CACNA1A
70	temperature homeostasis	8	27	29.63	1.16E-03	IL1RN;GPX1;ADRB2;HTR2A;GPX2;FTO;NR1D1;STAT3
71	establishment of mitotic spindle localization	6	16	37.50	1.20E-03	ARHGEF2;HTT;INSC;ESPL1;NUSAP1;NDE1
72	intrinsic apoptotic signaling pathway in response to oxidative stress	6	17	35.29	1.32E-03	GPX1;MAP3K5;STK3;STK24;UACA;BCL2

73	negative regulation of cell proliferation	109	580	18.79	1.43E-03	PIM2;DLG3;PRDX4;FOXP3;NRK;NTRK1;PTPRU;SKI;LEPRE1;CDKN2C;S100A11;REN;RBBP4;IGFBP5 ;SOX11;HDAC4;ING5;PELI1;SH3BP4;SPEG;SERPINE2;NCK2;CGREF1;BCL6;ROBO1;WNT5A;RARRES 1;FGFRL1;CTBP1;MSX1;SFRP2;FGFR3;HTT;SLIT3;MCC;CSNK2B;RPS6KA2;JARID2;TFAP2A;GPLD1; SRF;DDR1;CDKN1A;GLI3;IGFBP3;PTK2B;STK3;MYC;TRIB1;DLC1;TP53INP1;NOTCH1;DAB2IP;PTC H1;WNK2;ROR2;ITGB1;VAX1;GDF2;PTEN;GTPBP4;PAX6;BDNF;MLL;MUC2;INPPL1;ASCL2;SF1;WT 1;ZBTB16;FOSL1;IFITM1;CEND1;KDM2B;LRP6;TESC;CDKN1B;RNF10;ATP8A2;RB1;FLT3;MEG3;PP P2R5C;BCL11B;CDKN3;VSX2;SSTR1;NR2F2;THBS1;SMAD6;GREM1;TSC2;CDH13;FOXJ1;AXIN2;ALO X15B;FLCN;MED1;TGIF1;ADCYAP1;RAX;BCL2;RBM38;BMP7;COL18A1;ADARB1;COMT;TSPO;NF2
74	negative regulation of canonical Wnt signaling pathway	23	91	25.27	1.46E-03	FAM123B;WWTR1;SIAH2;WNT5A;DKK2;SFRP2;MCC;ANKRD6;GLI3;SOSTDC1;STK3;SOX17;NOTCH1 ;DAB2IP;ROR2;TCF7L2;DKK1;LRP4;PRICKLE1;LRP10;GREM1;AXIN2;APC2
75	positive regulation of glycogen biosynthetic process	6	14	42.86	1.74E-03	IRS1;C1QTNF2;GCK;SORBS1;AKT1;INSR
76	activation of	5	13	38.46	1.81E-03	GCK;DAB2IP;MAP4K2;MAP4K1;MAP3K7IP1
77	detection of mechanical stimulus involved in sensory perception	8	23	34.78	1.81E-03	NTRK1;SERPINE2;SLC12A2;FYN;GRIN2B;HTR2A;ACCN1;TMC2
78	cellular protein complex assembly	45	235	19.15	1.82E-03	DMD;MID11P1;WNT3A;PDE4DIP;PEX14;FMOD;NCK2;PDGFC;TPPP;ADRB2;CSNK2B;PFDN6;TAPB P;FBXO5;CADPS2;COBL;PTK2B;PTK2;NRG1;TRAF2;GSN;LZTS2;NRG3;RIC3;PSMD9;TUBA1A;FKBP 4;PXN;HSP90AA1;TMED10;GREM1;CYBA;COG4;MYH11;TUBB3;GAS7;CCDC40;UBE2S;UNC13A;FC HO1;C19orf51;CCDC114;AURKC;TUBB1;PSMG1
79	neural precursor cell proliferation	18	63	28.57	1.82E-03	ARX;WNT3A;DISC1;ARTN;ZEB2;ACSL6;DAGLB;GLI3;TACC2;DAGLA;RRM1;LHX5;LRP6;NCOR2;FRS 2;NDE1;PCNT;PLXNB2
80	nucleotide-binding oligomerization domain containing signaling pathway	8	28	28.57	1.86E-03	MAP3K7IP3;IKBKG;IRAK1;NOD1;RELA;MAP2K6;UBE2V1;MAP3K7IP1
						34

81	endocardium	5	12	41.67	1.91E-03	ACVR1;STK3;NRG1;SOX17;NOTCH1
82	protein polyubiquitination	34	174	19.54	1.91E-03	UBE2A;HUWE1;PSMB2;PELI1;PSMD14;PSME4;RNF168;FBXW11;PSMB9;PARK2;PSMB1;SMURF1;U BE3C;RNF216;TNKS;TRAF2;UBE2D1;TNKS2;PSMD9;CHFR;ANAPC5;CDC16;ASB2;WWP2;MGRN1;PS MB10;CDC27;NEDD4L;BCL2;UBE2S;APC2;FZR1;UBE2V1;PSMF1
83	axis elongation	9	25	36.00	2.07E-03	VANGL2;WNT3A;WNT5A;SFRP2;RDH10;LRP6;MED1;TRIM28;TFAP2C
84	negative regulation of cysteine-type endopeptidase activity involved in apoptotic process	16	66	24.24	2.29E-03	RPS6KA3;DDX3X;HBXIP;MAL;SIAH2;PAK2;GPX1;SFRP2;FNIP1;DNAJB6;INTS1;PAX2;PRDX5;AKT1; THBS1;MAP2K5
85	epithelial cell fate commitment	7	17	41.18	2.35E-03	ARX;ACVR1;FGFR3;HOXA13;NOTCH1;JAG2;NR2F2
86	mitotic cell cycle	113	671	16.84	2.79E-03	PIM2;FLNA;STAG2;PPP1R12B;SDCCAG8;ARHGEF2;PHF13;ENSA;RCC2;CDKN2C;PSMB2;JTB;TPR;C ENPL;RBBP4;EML4;ALMS1;ACVR1;SPTBN1;PSMD14;NCAPH;PSME4;STAG1;MAP4;NUP210;RFC1;C AMK2D;PPP3CA;MLF11P;HTT;PLK2;FBXL7;CSNK2B;DAXX;PSMB9;KIFC1;FBXO5;NUP43;NEDD9;C DKN1A;PSMB1;KIF25;MAD1L1;MCM7;YWHAG;ARHGEF10;NSMCE2;DLC1;TNKS;TAF2;LZTS2;ITGB1 ;TUBGCP2;CAMK2G;PPP2R2D;PTEN;UBE2D1;CCND1;BRSK2;INSC;RRM1;CKAP5;BANF1;CDCA5;H INFP;NCAPD2;EPS8;ESPL1;PSMD9;CHFR;TUBA1A;AAAS;ANKLE2;DYNLL1;ANAPC5;BCAT1;PPP1R1 2A;CDKN1B;RB1;NEK3;CDC16;DYNC1H1;HSP90AA1;CDKN3;C15orf23;NUSAP1;GINS2;E2F4;CDH13 ;NDE1;JMJD5;TUBB3;PSMB10;ZNF830;MYH10;TEX14;TADA2A;CDC27;NFATC1;CTDP1;SEH1L;UBE 2S;APC2;B9D2;FZR1;KIAA0892;PLK5P;AURKC;RBL1;DSN1;PSMF1;PCNT;TRIOBP
87	negative regulation of muscle cell	8	23	34.78	2.95E-03	SFRP2;HMGCR;PTK2B;NRG1;LRP6;IGF1R;ADCYAP1;BMP7

	apoptotic process					
88	phospholipid efflux	4	11	36.36	3.55E-03	APOA1;APC2;ABCA7;ABCG1
89	regulation of energy	4	10	40.00	4.31E-03	SGIP1;PASK;NR1D2;FLCN
	homeostasis					

90	cellular cation	63	383	16.45	4.37E-03	BCAP31;ABCB7;ARF1;RYR2;DISC1;SLC9A1;GNB1;ATP6V1B1;SLC8A1;ATP6V1C2;TRPM8;TTC7A;RAB
	homeostasis					7A;PROK2;TNNC1;XCR1;ITPR1;STIM2;PDE6B;NPY2R;NR3C2;PDE6A;ITPR3;CFTR;TAC1;TFR2;ATP6
						V0A4;GCK;PTK2B;MYC;SLC39A14;KCNMA1;ITGB1;CDH23;GDF2;CBARA1;CALCA;DRD4;SLC39A13;
						CCKBR;RIC3;LRP6;CACNA1C;TMBIM6;HTR2A;GPR12;EDNRB;SLC8A3;SLC24A4;NPTN;JPH3;HMOX
						2;DLG4;ADCYAP1;NEDD4L;BCL2;CACNA1A;RYR1;TBXA2R;TNNI3;LPAR2;SLC4A11;SLC9A8

91	regulation of sodium ion transport	14	51	27.45	4.85E-03	DMD;SERPINE2;MAGI1;CAMK2D;ADRB2;UTRN;CNKSR3;WNK2;DRD4;TESC;FGF14;AKT1;NEDD4L; SIK1
92	negative regulation of	5	12	41.67	4.94E-03	SERPINE2;DAB2IP;PTEN;TSC2;PIK3IP1
	phosphatidylinosito 1 3-kinase signaling					
93	mitotic	5	14	35.71	4.96E-03	PHF13;NCAPH;CDCA5;NCAPD2;NUSAP1
	chromosome condensation					
94	negative regulation	8	28	28.57	5.76E-03	PRDX4;WNT5A;ZP3;SF1;WT1;CDKN1B;BMP7;TPST2
	of reproductive					
95	positive regulation	4	11	36.36	7.12E-03	ZNF639;SNW1;SMARCA4;EP300
	by host of viral					
06	transcription	4	10	22.22	7 155 02	
90	metabolic process	4	12	33.33	7.15E-03	REN;MME;IMED10;UNCISA
97	protein	5	15	33.33	8.74E-03	SKI;TRPM8;SKIL;TRAF2;HSF4
	homotrimerization					
98	transepithelial	4	13	30.77	9.26E-03	SLC12A2;GPLD1;CFTR;RHCG
00	transport	0	27	24.22	0.76E.02	UCD2.MAT2D.NDST1.CCV.EVT1.DDD1D2C.AVT1.CLT25D1.UAS1
99	biosynthetic	9	57	24.32	9.70E-03	UGP2;MA12B;NDS11;GCK;EX11;PPP1K5C;AK11;GL125D1;ftAS1
	process					
100	citrulline metabolic	4	11	36.36	1.44E-02	DDAH1;HTT;DDAH2;ALDH18A1
	process					