**Interactive impact of severity of childhood maltreatment, depression and age on cortical brain structure: mega-analytic findings from a large multi-site cohort**

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Short title: Childhood maltreatment impacts cortical brain structure

Abstract: 237 words, text: 4498, 2 Figures, 2 Tables

**Abstract**

**Background:** Childhood maltreatment (CM) plays an important role in the development of major depressive disorder (MDD). Aim of the study was to examine whether CM severity and type are associated with MDD-related brain alterations and how they interact with sex and age.

**Methods:** Twelve university partner sites within the ENIGMA-MDD network assessed CM and subtypes of CM using the Childhood Trauma Questionnaire and acquired structural magnetic resonance imaging data from patients with MDD and healthy controls (HC) in a mega-analysis comprising a total of 3,872 participants aged between 13 and 89 years. Cortical thickness and surface area were extracted at each site using *FreeSurfer.*

**Results:** CM severity was associated with reduced cortical thickness in the banks of the superior temporal sulcus and supramarginal gyrus as well as with reduced surface of the middle temporal lobe. Participants reporting both childhood neglect and abuse had lower cortical thickness in the inferior parietal lobe, middle temporal lobe, and precuneus compared to participants not exposed to CM. In males only, regardless of diagnosis, CM severity was associated with higher cortical thickness of the rostral anterior cingulate cortex. Finally, a significant interaction between CM and age in predicting thickness was seen across several prefrontal, temporal and temporo-parietal regions.

**Conclusions:** This study represents the largest effort worldwide to identify cortical brain structure differences related to CM in individuals with MDD. Severity and type of CM may impact cortical thickness and surface area. Importantly, CM may influence age-dependent brain maturation, particularly in regions related to the default mode network, perception and theory of mind.

**Key Words:** major depressive disorder (MDD), cortical thickness, childhood maltreatment, ENIGMA

**Introduction**

According to the Centers for Disease Control and Prevention, childhood maltreatment (CM) is defined as “any act or series of acts of commission or omission by a parent or other caregiver that results in harm, potential for harm, or threat of harm to a child” (Leeb, 2008). CM may be physical, sexual or emotional and may result in inadequate environmental input (e.g., deprivation or neglect) or excessive harmful input (Sheridan and McLaughlin, 2014). About one quarter of all adults have encountered CM in their life (Butchart and Mikton 2014); this statistic may even be higher as a history of childhood adversity is likely under-reported. In fact, a recent meta-analysis has found that over half of children globally had experienced violence in just the past year alone (Hillis *et al.*, 2016). In a large birth cohort in Brazil, it was found that emotional abuse and exposure to domestic violence predicted increased risk for major depression for females, while CM did not predict depression onset in males (Gallo *et al.*, 2017).

Given the prevalence of CM, this is especially alarming as CM is strongly associated with a wide range of adverse consequences, not only causing suffering in the immediate aftermath, but long-term detrimental effects to mental and physical health. Children with a history of CM are more prone to smoking and obesity, as well as of being perpetrators and victims of violence (WHO, 2016, November). Notably, CM is one of the strongest factors in the development of major depressive disorder (MDD) (Bernet and Stein, 1999), the leading cause of disability worldwide according to the World Health Organization, with increasing rates over the past decade (WHO, 2016, November).

As both CM and MDD have a high incidence in the general population, the interplay between these two phenomena is important to investigate, both for prevention and treatment. Depressed patients with CM, for example, respond more poorly to antidepressant treatment than those without CM (Nanni *et al.*, 2012). CM and MDD may be causally linked, as MDD is a disorder characterized by pathological responses to stress (Frodl *et al.*, 2008). Both prospective and retrospective reports of maltreatment were found to be associated with adult psychiatric problems in a recent study, though the strongest associations were found when maltreatment was retrospectively self-reported (Newbury *et al.*, 2018). In that study, it was also shown that young adults who recall being maltreated have a particularly elevated risk for psychopathology.

In experimental studies, chronic social stress induces glucocorticoid-mediated pyramidal dendrite retraction in the hippocampus and changes in dendrite arborization in the prefrontal cortex (PFC) (Kole *et al.*, 2004, Magarinos *et al.*, 1996, Wellman, 2001, Woolley *et al.*, 1990), which may be associated with the behavioral manifestations of stress-related disorders like MDD (Macqueen and Frodl, 2010).

Based on the extant literature, one hypothesis is that CM in humans acts as a chronic stressor contributing to changes of brain structure and function, which in turn may increase vulnerability to psychiatric disorders such as MDD. Supporting this theory, CM was found to be associated with reduced brain volumes in the amygdala, prefrontal cortex and cerebellum (Dannlowski *et al.*, 2012, Edmiston *et al.*, 2011, Frodl *et al.*, 2010, Teicher *et al.*, 2016) - regions also reported to be affected in MDD. In the largest meta-analysis to date performed by the “Enhancing Neuro Imaging Genetics through Meta-Analysis” (ENIGMA) MDD consortium, we found thinner cortices in the bilateral medial orbitofrontal cortex (OFC), fusiform gyrus, insula, rostral anterior cingulate cortex (rostral ACC) as well as posterior cingulate cortex (PCC) and unilaterally in the left temporal gyrus, right inferior temporal gyrus and right caudal ACC in adult patients with MDD compared to controls (Schmaal *et al.*, 2017). These findings are consistent with a recent large-scale meta-analysis of voxel-based morphometry studies (Wise *et al.*, 2016).

A dose-response relationship of CM on medial prefrontal gray matter volume was also detected irrespective of diagnosis with MDD or anxiety (van Harmelen *et al.*, 2010). Previously, in an ENIGMA-MDD mega-analysis focusing on subcortical structures, CM also was found to be associated with lower caudate volume in females. Those alterations were more strongly associated with emotional and physical neglect than with other forms of CM (Frodl *et al.*, 2017a).

Research in animals and humans also suggests important distinctions between types of CM on brain structure. Specifically, researchers theorized that experiences characterized by deprivation (e.g., emotional and physical neglect) compared with experiences characterized by threat (e.g., emotional abuse and physical violence) lead to different effects on neuronal development (McLaughlin *et al.*, 2014a). A community study in 287 adolescents showed that exposure to threat and violence was associated with automatic emotion regulation deficits, but not cognitive control disturbances. In contrast, exposure to poverty was associated with worse cognitive control, but no deficits automatic emotion regulation. On the other hand, both violence and poverty predicted poor inhibition in an emotional context (Lambert *et al.*, 2017). Interestingly, children exposed to severe deprivation in the form of institutional rearing exhibited widespread cortical thinning in the superior and inferior parietal cortex (McLaughlin *et al.*, 2014b), and children exposed to neglect often have deficits in language abilities (Farah *et al.*, 2006). Individuals with a history of deprivation showed smaller gray matter volumes compared with individuals with a history of abuse in the fusiform gyrus and the middle occipital gyrus (Everaerd *et al.*, 2016). Therefore, exploring effects from different types of CM on brain structure was an important goal of the current study.

In this mega-analysis we first aimed to investigate the association between CM severity and cortical brain structure in MDD patients and healthy subjects. We hypothesized that more severe CM would be related to lower cortical thickness and surface area, especially of the OFC, ACC, medial prefrontal cortex and insula - regions affected in adult MDD (Fischl *et al.*, 2002) and involved in emotion regulation (Desikan *et al.*, 2006). We also explored whether reductions of cortical thickness and surface area would be specifically associated with CM severity, depression or both. We hypothesized that MDD patients with a more severe history of CM would show smaller cortical brain measures than healthy controls with a similar history of CM. Prior studies detected effects of CM on hippocampal volume (Frodl *et al.*, 2017a) and dorsomedial prefrontal cortex volume (Frodl *et al.*, 2017b) irrespective of diagnosis, but did not fully consider the severity of CM. Second, we investigated the associations of CM type with brain structures. Third, we investigated interactions of CM with sex and age. We hypothesized these changes to be more prominent in females than males and thus investigated the interactions between sex and CM on brain structure. Furthermore, given the large sample size and wide age range, we aimed to explore interactive effects of CM with age effects on brain structures

**Methods and Materials**

**Samples**

In the current study, twelve international sites participating in the ENIGMA MDD Workgroup with information on CM agreed to participate in the Childhood Adversity Subgroup. Detailed demographics and clinical characteristics for each sample may be found in **eTables 1 and 2.** Most studies used SCID-1, CIDI or another form of standardized interview (**eTable 3**). Exclusion criteria for study enrollment are given in **eTable 3**. In total, we analyzed data from 3,872 participants: 1,284 patients with MDD and 2,588 healthy controls (HC). All participating sites obtained approval from local institutional review boards and ethics committees. In addition, this mega-analysis was approved by the ethics board of the medical faculty of the Otto von Guericke University Magdeburg, Germany. All study participants provided written consent at their local site. In case of adolescent participants, parent/legal guardian provided written consent and the adolescent provided written assent.

**Assessment**

Severity of CM (CM-severity) was measured across all sites with the Childhood Trauma Questionnaire (CTQ) (Bernstein et al., 1994). The short form of the CTQ is a standardized self-report instrument consisting of 28 items containing five major subscales of childhood maltreatment. Each one also features a cut-off to determine the presence of emotional (>=12), physical (>=10) and sexual abuse (>=8) or emotional (>=15) and physical neglect (>=10). Three additional items remained to provide information on responders’ tendencies toward minimization and denial. For our analyses, we assessed CM in two ways. First, based on a score above the cut-off for at least one of the abuse or neglect subscales we divided our participants into 4 groups (CM-type): no CM, neglect (no abuse), abuse (no neglect), abuse+neglect. In a second analysis, we explored the effect of CTQ total sum score as a continuous variable (CM-severity).

Severity of depressive symptoms at time of scanning was measured in some sites with the Hamilton Depression Questionnaire (HDRS-17), in others the Beck Depression Inventory (BDI-II) or Inventory of Depressive Symptomatology-Self Report (IDS-SR) was used. Age of onset and antidepressant medication use at the time of scan were also recorded in 11 and 12 sites respectively.

**Image processing and analysis**

Participants all underwent structural T1-weighted MRI brain scans locally at each site, where scans were analyzed using the fully-automated and validated segmentation software *FreeSurfer* (version 5.0 or higher) (Fischl *et al.*, 2002). Image acquisition parameters and software descriptions for each sample are given in e**Table 4**. Deep brain structure volumes were extracted and visually inspected for segmentation accuracy. Parcellations for cortical thickness and surface area of 68 (34 left and 34 right) regions based on the Desikan-Killiany atlas (Desikan *et al.*, 2006) and left and right hemisphere measures were derived and visually inspected for accuracy following protocol designed to facilitate harmonized image analysis across multiple sites (<http://enigma.ini.usc.edu/protocols/imaging-protocols/>). Association between CM and subcortical measures were previously published (Frodl *et al.*, 2017a).

**Statistical framework of mega-analysis**

Statistical analyses were performed using SPSS statistics (version 24).

We performed ANOVAs, or Kruskal-Wallis tests as appropriate, to compare at scan, age at MDD onset, clinical severity of depression and CTQ scores between groups and cohorts. Chi-squared tests were used to analyze differences between frequencies of males and females.

Then, we built generalized estimating equations (GEE) models with thickness or surface area of each region as the dependent variable. Our models had a linear scale response. All participants were included, irrespective of diagnosis. The independent between-subject variable CM was defined in two ways: as the factor CM-type (0=no CM, 1=neglect, 2=abuse, 3=neglect+abuse) or as CM-severity (continuous: total CTQ). Each of these variables was included in separate models. In all models we included the between-subjects factors diagnosis (factor: 1=patients, 0=healthy controls), sex (factor: 0=males, 1=females) and the within-subject factor hemisphere (left, right). Age (continuous), neuroimaging cohort (factor) and total intracranial volume (continuous) were used as between-subject covariates. *FreeSurfer* version and scanner type were comprised in the factor neuroimaging cohort. As we did not expect CM-severity effects to be lateralized, hemisphere was only included as a main effect. Our prior research showed differential effects of CM-severity in predicting the volume of subcortical structures depending on sex and MDD diagnosis (Frodl *et al.*, 2017b). Therefore, we explored in our models all possible interactions between CM, sex and diagnosis for both surface and thickness. To assess the effect of CM across all brain areas, we first ran analyses on the total thickness and surface across all regions respectively, adding region as a within-subject factor. Then, we repeated the process for each region individually. Finally, we explored the interaction between age and severity of CM while keeping all other terms in the model as main effects.

In all models, Wald chi-squared tests were used to assess the significance of each term. To account for multiple tests (34 regions), a false discovery rate (FDR) correction was computed on the resulting *p*-values. Findings were considered significant if pFDR<0.05. Any significant interactive effects resulting from the models described above were followed up with post-hoc testing.

*Investigation of clinical confounds*

A subset of our MDD cohort (N=965) had more detailed clinical information and allowed us to explore additional potential confounding effects. Therefore, we investigated if thickness or surface of all regions were significantly predicted by clinical severity (continuous: BDI total score, since HAM-D was available only for a minor subset of participants), recurrence (factor: 0=first episode, 1=recurrent episode), current antidepressant use (factor: 0=no, 1=yes), remission (factor: 0=currently remitted, 1=currently depressed), age of depression onset (continuous). We did so by building GEE models that featured these measures as predictors (main effects), together with age, sex, site, hemisphere and total intracranial volume. Detailed information on this subsample is presented in **eTable 14**.

**Results**

**Demographics**

For details on participant’s demographics and clinical features, see **Table 1**. Overall, data significantly differed between centers with respect to sex, age, CTQ scores and clinical features (see **eTable 5**). Frequency of co-occurrence between abuse and neglect is 12.7%. CM severity of influenced by abuse and neglect to a similar extend (beta=0.41 and beta=0.42, respectively).

**Cortical Thickness**

*Main effects of CM-severity*

A summary of all significant findings is reported in **Table 2**. For an overview of the results of the models run on each region, see **eTable 6**.

We detected a significant main effect showing an inverse relation between CM-severity and thickness of the banks of the superior temporal sulcus (Wald chi-squared=14.583, pFDR=0.033, B=-0.001, **Figure 1**). A significant main effect of CM was also present on thickness of the supramarginal gyrus (Wald chi-squared=8.889, pFDR=0.049, B=-0.001, **Figure 1**).

*CM-severity and sex interaction*

When considering all regions, the interaction between CM-severity and sex was significant (Wald chi-squared=5.220, p=0.022). Dividing the data by sex, post-hoc analyses showed a significant negative effect of CM-severity on cortical thickness in females (Wald chi-squared=4.861, p=0.027, B=-0.000649), but not in males (Wald-chi-square=1.287, p=0.257, B=-0.000136).

When running models for each region separately, we found a significant interaction between CM-severity and sex on cortical thickness of the rostral anterior cingulate cortex (Wald chi-squared=13.556, pFDR=0.008). Post-hoc analysis revealed a significant positive effect of CM-severity on cortical thickness of this region in males (Wald chi-squared=14.426, p<0.001, B=0.002, **Figure 1**) but not in females (Wald chi-squared=3.174, p=0.075, B=-0.0006).

*CM-severity and age interaction*

When considering all regions, a significant interaction between age and severity of CM was detected (Wald chi-squared=11.105, p=0.001, B=-0.000035).

Models ran for each region separately indicated that this interaction between age and severity of CM was significant across all participants in the rostral anterior cingulate, isthmus of the cingulate, posterior cingulate, lateral orbitofrontal gyrus, parahippocampal gyrus, inferior frontal gyrus (IFG) pars opercularis, IFG pars triangularis, superior frontal gyrus, banks of the superior temporal sulcus, cuneus, fusiform gyrus, insula, precentral gyrus, precuneus, supramarginal gyrus and transverse temporal gyrus (see **Table 3, eTable 7, Figure 3)**. Similar results were found when using CM type in the analysis (see **Table 3, eTable 11, Figure 3)**.

*Main effects of CM-type*

We found a significant main effect of CM type (**eTable 10**) in the banks of the superior temporal sulcus (Wald chi-squared=19.888, pFDR=0.006), inferior parietal lobe (Wald chi-square=15.273, pFDR=0.023), middle temporal lobe (Wald chi-squared=12.123, pFDR=0.048), precuneus (Wald chi-squared=15.325, pFDR=0.023) and supramarginal gyrus (Wald chi-squared=13.990, pFDR=0.026). In all cases, the neglect+abuse group had lower mean thickness values compared to the no CM group (all p<0.01, Figure 1) and there was no difference between the abuse only as well as neglect only CM types and the no CM group.

*CM-type and age interaction*

The interaction between age and type of CM was significant across all participants for most regions (see **Table 3, eTable 11**). In all cases, the effects of age were more negative in the neglect+abuse group compared to the CM group (all p<0.05).

**Cortical Surface**

A summary of all significant findings is reported in **Table 2**. For an overview of the results of the models run on each region, see **eTable 8**.

*Main effects of CM-severity*

Across all regions, a negative main effect of CM-severity on cortical surface area was observed (Wald chi-squared=4.413, p=0.036, B= -0.414). When running separate models for each region **(eTable 8)**, we detected a significant inverse main effect of CM-severity on surface area of the middle temporal gyrus (Wald chi-squared=12.368, pFDR=0.015, B=-1.504, **Figure 2**).

*CM-type diagnosis and sex interaction*

We found a significant interaction between CM type, diagnosis and sex (**eTable 12**) in the caudal anterior cingulate (Wald chi-squared=17.807, pFDR<0.001). Post-hoc testing revealed that, in depressed males, those having suffered from either abuse or neglect had lower average cortical surface of the caudal anterior cingulate cortex than those who had no history of CM (*p*=0.003 and *p*=0.017 respectively, Figure 2).

**Investigation of clinical confounds**

Our post-hoc investigation in a subset of patients with detailed information showed no significant effects of clinical variables on thickness or surface (all p>0.05). See eTable 15 and eTable 16 for the model effects.

**Discussion**

In the present study, we investigated the association between CM and cortical brain structure in a large sample of MDD patients and healthy subjects. We show that CM has a subtle but widespread effect on cortical thickness and surface, which is likely influenced by sex and age.

Two procedures of describing CM were used. First, the continuous measure of CM severity allowed for a continuous analysis and the second classification in no CM, only neglect, only abuse and both abuse and neglect allowed for an analysis of type of CM. It should be highlighted that participants exposed to both neglect and abuse also had higher total CM values.

Severity of CM was associated with lower mean cortical surface area regardless of region across all MDD patients and healthy controls. In women, higher severity of CM was also associated with thinner thickness across all regions. These findings are consistent with prior research showing widespread effects of severity of CM on the brain irrespective of psychopathological status (Chaney *et al.*, 2014).

Regardless of diagnosis, when individual regions were investigated this effect of CM severity survived correction for multiple testing in temporal and temporo-parietal regions. Specifically, participants with higher CM severity had significantly thinner cortex in the banks of the superior temporal sulcus and the supramarginal gyrus. When considering type of abuse, high severity of CM, represented by concurrent childhood neglect and abuse, was once again associated with reduced thickness in these two areas and additionally in the precuneus, middle temporal lobe and inferior parietal cortex. Moreover, participants with higher severity of CM also showed smaller surface area of the middle temporal gyrus. The magnitude of these negative effects on thickness and surface area pointed towards a reduction around 0.001 mm of thickness and around 1.5-5.1 mm2 of surface with each one-point increase in CTQ score depending on the brain region. This means that an increase of 100 points in the CTQ scale is associated with a 0.1mm (or 4 %) decrease of cortical thickness e.g. in the banks of the superior temporal sulcus. The middle temporal lobe is thought to be essential for our ability to understand actions and semantic associations (Davey *et al.*, 2016). One possibility is that CM may lead to difficulties in semantic retrieval through alterations in regions of temporal cortex and the default mode network. Indeed, other studies have also demonstrated that individuals with higher severity of CM showed reduced cortical surface on the left middle temporal area and lingual gyrus (Kelly *et al.*, 2013). In contrast, in a study of adolescents and young adults exposed to CM, increased cortical volume was observed in the left inferior and middle temporal gyri relative to healthy controls (Lim *et al.*, 2018). In the present study, the other regions we report showing an impact from CM type are located in the temporo-parietal area and around the temporo-parietal junction; both of these regions play a role in theory of mind processing which is important during daily social interactions (Saxe and Kanwisher, 2003). Deficits in these areas might suggest a disadvantage for subjects with a history of CM, in particular those with increased severity and more types of CM. No significant main effects of severity of CM were detected for other regions we hypothesized to be vulnerable. For example, prior studies found a significant main effect of CM in different regions such as fronto-limbic areas, visual cortex, and cerebellum (Kelly *et al.*, 2013, Yang *et al.*, 2017). This might be due to smaller sample sizes and more homogeneity in prior studies: in our analysis, which features a larger sample size of N=3,872, we detected an overall effect of severity of CM on the whole cortex with some prominence in the temporal and temporo-parietal regions.

Another interesting finding was that males, but not females, with a more severe history of CM, regardless of diagnosis, showed distinctly thicker rostral anterior cingulate cortices. These results suggest sex differences in the effects of CM on the structure of this region (Canu *et al.*, 2015, Fallucca *et al.*, 2011). The anterior cingulate cortex is involved in emotional and inhibitory processes (Garavan *et al.*, 2006, Steele *et al.*, 2013). Thus, males seem to be particularly sensitive to CM with regards to thickness in a region relevant for emotion regulation and might show a reactive increase of thickness. Whether this thickness increase of the rostral ACC is adaptive cannot be effectively addressed in the current sample, as longitudinal data and resiliency measures were not available for the bulk of the cohort. In this context, it is interesting that male patients with a history of neglect and abuse had significantly smaller surface areas of the caudal ACC compared to those without CM, pointing towards a negative effect of CM in the caudal ACC in participants who developed MDD. In healthy subjects such an effect of CM was not seen in the caudal ACC and we could tentatively ascribe to resilience (Feder *et al.*, 2009).

**Childhood Maltreatment by Age Interaction**

A novel finding detected in our secondary analysis was that CM severity and age interacted regardless of diagnosis to predict both the thickness and surface area of several regions involved in emotional processing, such as portions of the cingulate, orbitofrontal, insular, dorsolateral prefrontal and medial prefrontal cortices. In these areas, older people including patients and controls with higher CM severity had lower cortical thickness. The orbitofrontal and dorsomedial prefrontal cortices can be seen as separate networks interacting closely with limbic structures, but also showing cortico-cortical interconnections with each other (Ongur and Price, 2000, Phillips *et al.*, 2008). These regions allow the brain to process emotionally salient information and help with the regulation of emotional behavior (Phillips *et al.*, 2008). The insula is closely interconnected with the orbitofrontal cortex and is involved in emotion and executive processing as well as working memory (Levens and Phelps, 2010). The cingulate cortex is also well known to have cognitive and emotional functions: its dorsal parts are involved in emotion evaluation, whereas the ventral parts and the dorsomedial prefrontal cortex are involved in emotion regulation (Etkin *et al.*, 2011). Overall, these results are consistent with previous studies showing that CM-severity impacts regions involved in emotion regulation, including the insula (Teicher *et al.*, 2014).

Our cross-sectional data suggests that cortical thickness might decrease more rapidly with age in individuals with a more severe history of CM. although this still needs to be confirmed by longitudinal analyses. It will be critical for future studies to assess the effects of abuse across multiple timepoints and to put it in relation to the age of participants.

**Diagnosis and CM interactions**

In the present study, no effect of diagnosis was detected and we also did not find a significant interaction between diagnosis and CM. It is possible that we could not replicate the main effect of MDD diagnosis on OFC and ACC thickness because of the smaller sample size of the current study (overall N=3,872) compared to the sample size in our previous ENIGMA MDD meta-analysis that focused on the effects of MDD (overall N=10,105, Schmaal *et al.*, 2017). However, since this is the largest joint mega-analysis concerning cortical thickness and CM to date, this null finding could also suggest that the effects of MDD commonly reported in studies and meta-analyses could be the result of the interaction of several underlying variables. For example, different effects of MDD depending on age and onset were already highlighted in Schmaal *et al.*, 2017. Our findings suggest that CM could be another factor that accounts for part of the structural differences between depressed patients and healthy controls.

**Strengths and Limitations**

A major strength of the study is the large sample size with a relevant control sample allowing inclusion of all 34 left and right cortical brain regions in the analyses. However, a larger sample from different sites also limits the common information collected, since not all sites used the same assessments. Not all patients were drug free and, further, the history of antidepressants use as well as duration, type and dosage of antidepressant treatment was not assessed completely during the lifetime, so we cannot rule out effects of antidepressant exposure influencing our results. Additionally, psychopathology was assessed with different ratings, so that it was not possible to use depression severity as a single covariate in the analysis. Overall, the datasets from the different samples included in the mega-analysis were significantly heterogeneous regarding demographics and clinical features. This is a common limitation of multi-site analyses and we accounted for this effect by adding site as a covariate to all of our models. It is worth noting that we were able to analyze the influence of clinical confounds in a subset of our MDD cohort, where we did not find any significant effect of recurrence, antidepressant medication, remission, severity or age of depression onset in predicting cortical thickness or surface. In particular, currently remitted patients represented only 13.5% of this subset and we consider the impact of their inclusion at some sites negligible. However, measures of socioeconomic status and education have been shown to play a role in brain structure (Ritchie et al. 2018), but were unfortunately unavailable in our sample. Finally, we considered hemisphere as a within-subject effect in our dataset. Our hypothesis was that CM would affect anatomically distinct regions differently rather than be selective for a specific region on a specific hemisphere. Therefore, we believed that including all possible interactions between regions and hemisphere would lead to an unnecessarily complex model. However, it is possible that besides the bilateral effects we report, subtler lateralized effects of CM might exist in specific areas.

Even if our investigation features the broad variation of “real life” clinical populations, future studies are needed to confirm our findings in carefully controlled datasets. Here, we explored the effect of CM in a sample of healthy participants and patients with MDD. Because it is not clear how the severity or type of CM may affect the development of structural brain measures, it will be important to consider the onset and timing of CM in future (Ho *et al.*, 2018). In addition, future longitudinal data are required to establish whether cortical thickness might decrease more rapidly with age in individuals with a more severe history of CM, as our current cross-sectional data may suggest. For this analysis, while it was possible to use extracted cortical measures from specific regions of interest, it was not possible to retrospectively analyze the original MRI datasets to perform a whole-cortex analysis with *FreeSurfer*. A surface-based analysis across the entire cortex may afford more sensitivity in detecting effects of CM and thus could be a future step.

**Conclusions**

The results of our study support the idea that CM-severity appears to affect the structure of temporal and parietal regions in particular. Thus, there are effects in the default mode network and in regions related to the theory of mind as well as to perception. Interestingly, CM may interact with the effect of age on cortical thickness in regions involved in emotion regulation, theory of mind as well as belonging to the default mode network. Thus, future studies should investigate if subjects with a history of CM may be more prone to cortical thinning during ageing or if CM results in changes that mimic ageing.

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Nic van der Wee has consulted for Lilly, Wyeth, Servier, Pfizer and GlaxoSmithKline

Thomas Frodl has received fees for presentations for educational programs with Servier, Janssen and Lundbeck

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Theo G.M. van Erp has consulted for Roche Pharmaceuticals, Ltd., and has a contract with Otsuka Phamaceutical Co., Ltd (OPCJ).

Knut Schnell has consulted for Roche Pharmaceuticals and Servier Pharmaceuticals

**Legends**

**Figure 1:** Effect of CM predicting predicting cortical thickness. Coefficients for the GEE model term CM severity or those for the neglect+abuse group compared to the no-CM group are plotted on an inflated left brain hemisphere (effects were bilateral). Only the neglect+abuse group was different from the no-CM group. CM=childhood maltreatment, GEE=generalized estimating equations.

Figure 2: Effect of CM predicting cortical surface. Coefficients for the GEE model term CM severity or those for the abuse only group compared to the no-CM group are plotted on an inflated left brain hemisphere (effects were bilateral). The neglect group showed a similar result in the same region. CM=childhood maltreatment, GEE=generalized estimating equations.

**Figure 3:** Effect of CM\*Age predicting cortical thickness. Coefficients for the GEE model term CM severity\*Age or those for Age in the abuse + neglect group compared to the no-CM group are plotted on an inflated left brain hemisphere (effects were bilateral). CM=childhood maltreatment, GEE=generalized estimating equations.

**Table 1:** Demographic and clinical Data. CTQ = Childhood trauma questionnaire. ICV= total intracranial volume. BDI#= Beck Depression Inventory. HDRS-17#= Hamilton Depression Rating Scale. Shown are mean values +- standard deviation. †Mann Whitney U test used, #From sites that used these ratings.

**Table 2:** Main findings derived from the GEE models not including the interaction of childhood maltreatment and age. Only significant effects of interest are shown, for all effects see supplemental tables. Coefficients are listed for the model term CM severity or those for the neglect+abuse group, with the exception of †neglect group. CM=childhood maltreatment, FDR=false discovery rate.

**Table 3:** Main findings derived from the GEE models predicting cortical thickness and including the interaction of childhood maltreatment with age. Only significant effects of the interaction are shown, for all effects see supplemental tables. Coefficients are listed for the model term CM severity\*Age or those for the neglect+abuse group\*Age. CM=childhood maltreatment, FDR=false discovery rate.

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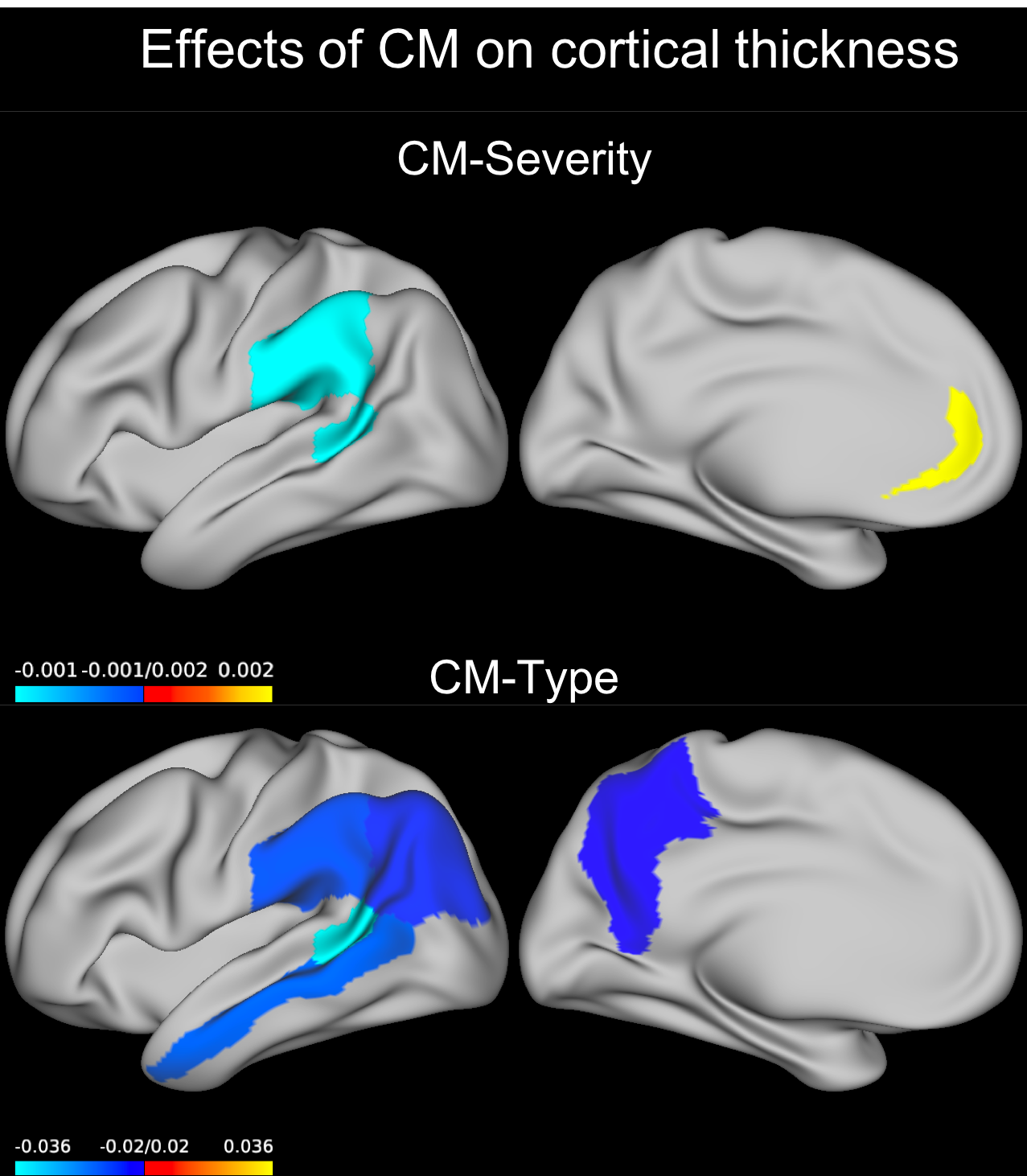
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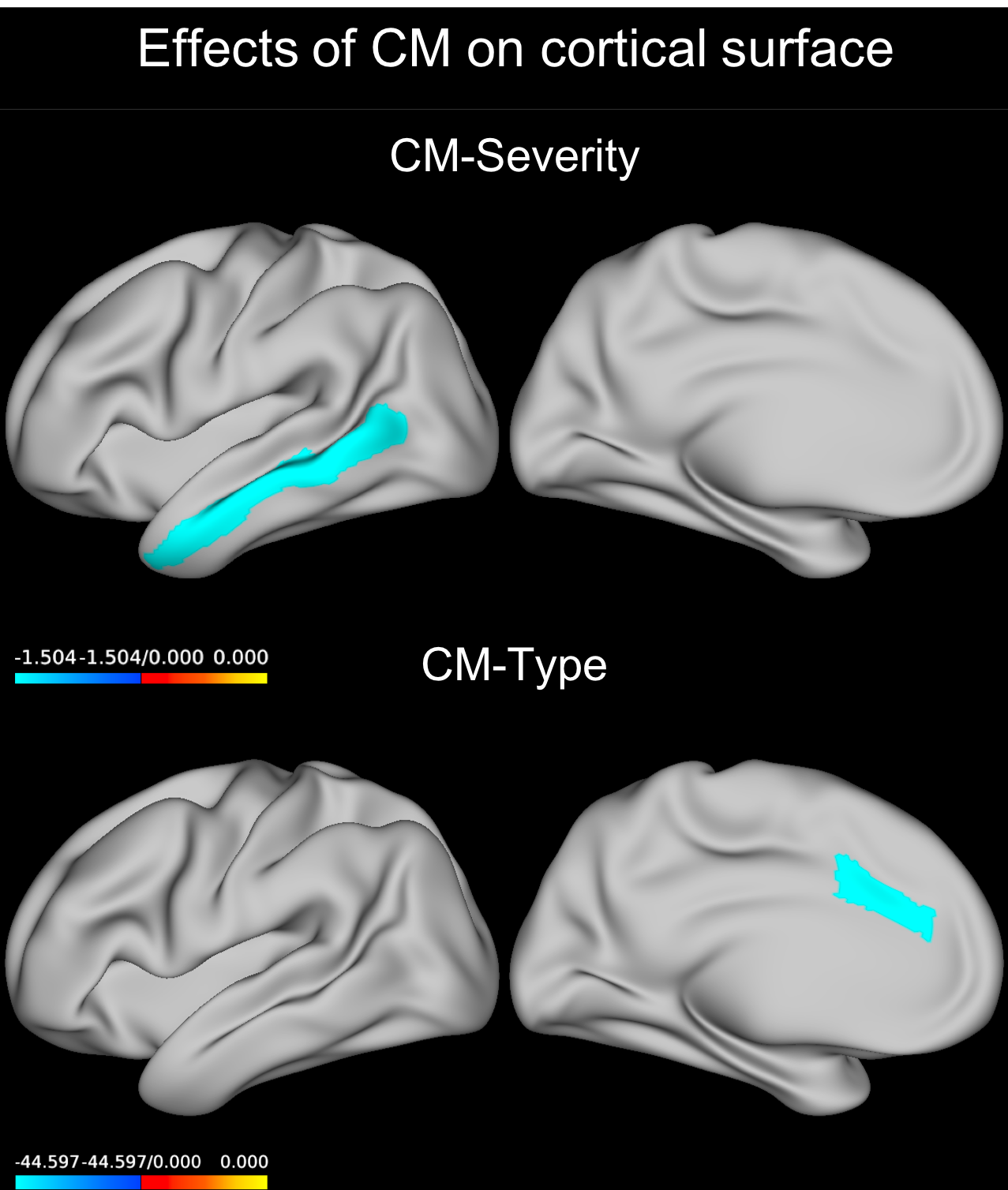
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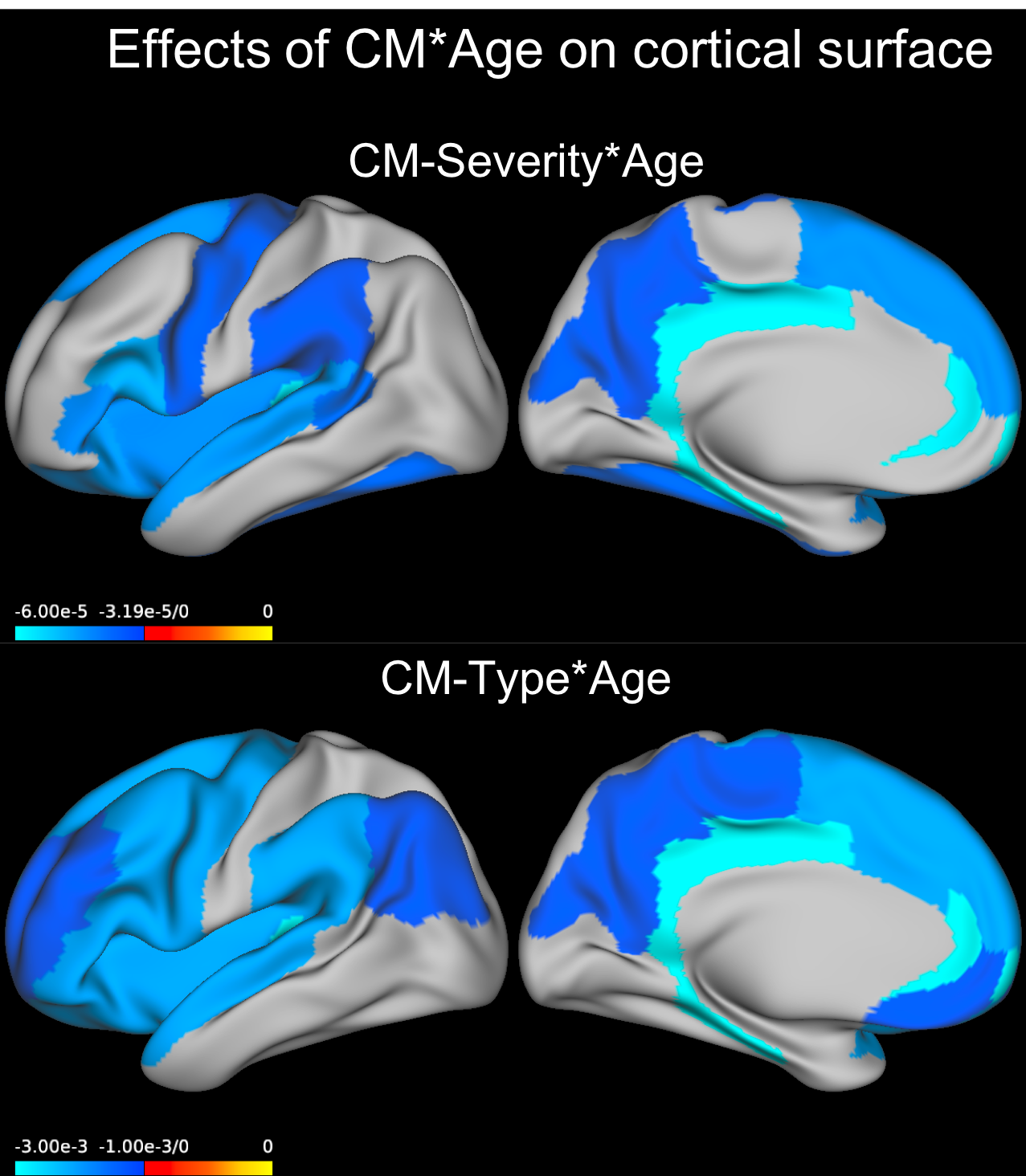
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**Figure 2:** Effect of CM predicting cortical surface. Coefficients for the GEE model term CM severity or those for the abuse only group compared to the no-CM group are plotted on an inflated left brain hemisphere (effects were bilateral). The neglect group showed a similar result in the same region. CM=childhood maltreatment, GEE=generalized estimating equations.



**Figure 3:** Effect of CM\*Age predicting cortical thickness. Coefficients for the GEE model term CM severity\*Age or those for Age in the abuse + neglect group compared to the no-CM group are plotted on an inflated left brain hemisphere (effects were bilateral). CM=childhood maltreatment, GEE=generalized estimating equations.



**Table 1:** Demographic and clinical Data. CTQ = Childhood trauma questionnaire. ICV= total intracranial volume. BDI#= Beck Depression Inventory. HDRS-17#= Hamilton Depression Rating Scale. Shown are mean values +- standard deviation. For CTQ subscales, number of subjects above the cut-off are given in brackets. †Mann Whitney U test used, #From sites that used these ratings.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | All subjects  (n=3872) | Controls  (N=2588) | Patients (N=1284) | Group difference |
| Females | 2116 (54.6%) | 1303 (50.3%) | 813 (63.3%) |  |
| Males | 1756 (45.4%) | 1285 (49.7%) | 471 (36.7%) | χ2=58.2, p<0.001 |
| Age (years) | 42.5 ± 15.5 | 43.3 ± 15.9 | 40.9 ±14.6 | t=4.6, p<0.001 |
| Age of onset (years) | - | - | 29.4 ± 14.0 | - |
| Total CTQ | 36.3 ± 12.7 | 32.6 ± 8.5 | 43.6 ± 16.1 | p<0.001† |
| Sexual abuse | 5.5 ± 2.2 (233) | 5.2 ± 1.2 (75) | 6.2 ± 3.3 (158) | p<0.001† |
| Physical abuse | 6.1 ± 2.5 (297) | 5.6 ± 1.7 (91) | 6.9 ± 3.4 (206) | p<0.001† |
| Emotional abuse | 7.6 ± 4.0 (770) | 6.5 ± 2.5 (246) | 10.0 ± 5.1 (522) | p<0.001† |
| Physical neglect | 7.0 ± 2.6 (243) | 6.6 ± 2.2 (86) | 8.0 ± 3.1 (157) | p<0.001† |
| Emotional neglect | 9.9 ± 4.8 (646) | 8.6 ± 3.8 (202) | 12.5 ± 5.5 (444) | p<0.001† |
| BDI-II# | - | 5.2 ± 4.4 | 18.6 ± 12.1 | p<0.001† |
| HDRS# | - | 2.9 ± 3.1 | 15.6 ± 9.8 | p<0.001† |
| ICV (in mm3) | (1.53±0.19)\*106 | (1.54 ± 0.18)\*106 | (1.527 ± 0.2)\*106 | t=2.7, p=0.007 |

**Table 2:** Main findings derived from the GEE models not including the interaction of childhood maltreatment and age. Wald χ2 and p values of CM severity and type are shown for the regions where they were significant. For all effects see supplemental tables. Effects are coefficients for the model term CM severity or the estimates of the indicated contrast for CM type. CM=childhood maltreatment, FDR=false discovery rate.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | Participants | Wald χ2 | pFDR | Effect |
| Thickness |  |  |  |  |
| *CM severity* |  |  |  |  |
| Overall thickness | Females | 4.861 | 0.027 | -0.001 |
| Rostral anterior cingulate cortex | Males | 14.426 | <0.001 | 0.002 |
| Banks of the superior temporal sulcus | All | 14.583 | 0.004 | -0.001 |
| Supramarginal gyrus | All | 8.889 | 0.049 | -0.001 |
| *CM type* |  |  |  |  |
| Banks of the superior temporal sulcus | Neglect+Abuse > no CM | 19.888 | 0.006 | -0.036 |
| Inferior parietal lobe | Neglect+Abuse > no CM | 15.273 | 0.023 | -0.022 |
| Middle temporal lobe | Neglect+Abuse > no CM | 12.123 | 0.048 | -0.025 |
| Precuneus | Neglect+Abuse > no CM | 15.325 | 0.023 | -0.020 |
| Supramarginal gyrus | Neglect+Abuse > no CM | 13.990 | 0.026 | -0.024 |
| Surface |  |  |  |  |
| *CM severity* |  |  |  |  |
| Overall surface | All | 4.413 | 0.036 | -0.414 |
| Middle temporal lobe | All | 12.368 | 0.015 | -1.504 |
| *CM type* |  |  |  |  |
| Caudal anterior cingulate | Depressed males, Neglect > no CM | 17.807 | 0.003 | -44.597 |
|  | Depressed males, Abuse > no CM | 5.647 | 0.017 | -51.396 |

**Table 3:** Main findings derived from the GEE models predicting cortical thickness and including the interaction of childhood maltreatment with age. Wald χ2 and p values of CM severity\*Age and CM type\*Age are shown for the regions where they were significant. For all effects see supplemental tables. Effect sizes are coefficients for the model term CM severity\*Age or those for Age in the Neglect+Abuse group versus the no-CM group. CM=childhood maltreatment, FDR=false discovery rate.

|  |  |  |  |
| --- | --- | --- | --- |
|  | Wald χ2 | pFDR | Effect size |
| CM severity\*Age |  |  |  |
| Overall thickness | 11.105 | 0.001 | -3.50\*10-5 |
| Banks of superior temporal sulcus | 4.997 | 0.047 | -3.59\*10-5 |
| Cuneus | 7.373 | 0.020 | -3.32\*10-5 |
| Frontal pole | 10.448 | 0.007 | -8.03\*10-5 |
| Fusiform | 6.714 | 0.026 | -3.50\*10-5 |
| Insula | 8.214 | 0.014 | -4.16\*10-5 |
| Isthmus of cingulate | 11.149 | 0.007 | -5.87\*10-5 |
| Lateral orbitofrontal | 8.952 | 0.011 | -4.37\*10-5 |
| Parahippocampal | 6.031 | 0.032 | -6.00\*10-5 |
| IFG pars opercularis | 11.014 | 0.007 | -4.70\*10-5 |
| IFG pars triangularis | 8.583 | 0.011 | -4.14\*10-5 |
| Posterior cingulate | 17.682 | 0.001 | -5.78\*10-5 |
| Precentral | 5.188 | 0.046 | -3.34\*10-5 |
| Precuneus | 6.272 | 0.029 | -3.19\*10-5 |
| Rostral anteriorcingulate | 10.262 | 0.007 | -5.83\*10-5 |
| Superior frontal | 7.301 | 0.020 | -4.22\*10-5 |
| Superior temporal | 8.774 | 0.011 | -4.35\*10-5 |
| Supramarginal | 5.189 | 0.046 | -3.22\*10-5 |
| Transverse temporal | 8.941 | 0.011 | -5.75\*10-5 |
| CM type\*Age |  |  |  |
| Caudal anterior cingulate | 10.155 | 0.030 | -0.002 |
| Caudal middle frontal | 16.297 | 0.002 | -0.002 |
| Cuneus | 15.442 | 0.002 | -0.001 |
| Frontal pole | 16.065 | 0.002 | -0.003 |
| Inferior parietal | 9.848 | 0.034 | -0.001 |
| Insula | 22.037 | <0.001 | -0.002 |
| Isthmus of cingulate | 23.710 | <0.001 | -0.003 |
| Lateral orbitofrontal | 17.174 | 0.002 | -0.002 |
| Medial orbitofrontal | 13.542 | 0.008 | -0.001 |
| Paracentral | 9.383 | 0.039 | -0.001 |
| Parahippocampal | 16.405 | 0.002 | -0.003 |
| IFG pars opercularis | 27.556 | <0.001 | -0.002 |
| IFG pars orbitalis | 8.785 | 0.047 | -0.002 |
| IFG pars triangularis | 20.837 | <0.001 | -0.002 |
| Posterior cingulate | 35.357 | <0.001 | -0.003 |
| Precentral | 17.291 | 0.002 | -0.002 |
| Precuneus | 13.794 | 0.006 | -0.001 |
| Rostral anteriorcingulate | 27.847 | <0.001 | -0.003 |
| Rostral middlefrontal | 9.544 | 0.037 | -0.001 |
| Superior frontal | 28.174 | <0.001 | -0.002 |
| Superior temporal | 22.270 | <0.001 | -0.002 |
| Supramarginal | 17.641 | 0.002 | -0.002 |
| Transverse temporal | 22.386 | <0.001 | -0.003 |