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Developmentally Sensitive Interaction Effects of Genes and the Social Environment on Total and Subcortical Brain Volumes

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Abstract

Smaller total brain and subcortical volumes have been linked to psychopathology including attention-deficit/hyperactivity disorder (ADHD). Identifying mechanisms underlying these alterations, therefore, is of great importance. We investigated the role of gene-environment interactions (GxE) in interindividual variability of total gray matter (GM), caudate, and putamen volumes. Brain volumes were derived from structural magnetic resonance imaging scans in participants with (N = 312) and without ADHD (N = 437) from N = 402 families (age M = 17.00, SD = 3.60). GxE effects between DAT1, 5-HTT, and DRD4 and social environments (maternal expressed warmth and criticism; positive and deviant peer affiliation) as well as the possible moderating effect of age were examined using linear mixed modeling. Deviant peer affiliation was associated with lower caudate volume. Participants with low deviant peer affiliations had larger total GM volumes with increasing age. Likewise, developmentally sensitive GxE effects were found on total GM and putamen volume. For total GM, differential age effects were found for DAT1 9-repeat and HTTLPR L/L genotypes, depending on the amount of positive peer affiliation. For putamen volume, DRD4 7-repeat carriers and DAT1 10/10 homozygotes showed opposite age relations depending on positive peer affiliation and maternal criticism, respectively. All results were independent of ADHD severity. The presence of differential age-dependent GxE effects might explain the diverse and sometimes opposing results of environmental and genetic effects on brain volumes observed so far.
Funding support for the IMAGE project was provided by NIH grants R01MH62873 and R01MH081803 to Dr. Faraone and the genotyping of samples was provided through the Genetic Association Information Network (GAIN). The dataset used for the analyses described in this manuscript was obtained from the database of Genotypes and Phenotypes (dbGaP) found at http://www.ncbi.nlm.nih.gov/gap through dbGaP accession number #20726-2. This work was further supported by an NWO Large Investment Grant 175010207010 and NWO Brain & Cognition an Integrative Approach grant (433-09-242) (to Dr. Buitelaar), and grants from Radboud University Nijmegen Medical Center, University Medical Center Groningen and Accare, and VU University Amsterdam. The research leading to these results also received funding from the European Community’s Seventh Framework Programme (FP7/ 2007–2013) under grant agreement number 278948 (TACTICS) and number 602450 (IMAGEEND) as well as from the European Community’s Horizon 2020 Programme under grant agreement n° 643051 (MIND). Dr. Franke is supported by a Vici grant from NWO (grant number 016-130-669), and she and Dr. Buitelaar received funding from the National Institutes of Health (NIH) consortium grant U54 EB020403, supported by a cross-NIH alliance that funds Big Data to Knowledge Centers of Excellence. Dr. Faraone is supported by the K.G. Jebsen Centre for Research on Neuropsychiatric Disorders, University of Bergen, Bergen, Norway, the European Community’s Seventh Framework Programme (FP7/2007-2013) under grant agreement n°602805 and NIMH grants R13MH059126 and R01MH094469.

Competing Interests: In the past year, Dr. Faraone received income, travel expenses, and/or research support from and/or has been on an Advisory Board for Pfizer, Ironshore, Shire, Akili Interactive Labs,CogCube, Alcoba, VAYA Pharma, Neurovation, Impax, NeuroLifeSciences and research support from the National Institutes of Health (NIH). His institution is seeking a patent for the use of sodium-hydrogen exchange inhibitors in the treatment of ADHD. In previous years, he received consulting fees or was on an Advisory Boards or participated in continuing medical education programs sponsored by: Shire, Alcoba, Otsuka, McNeil, Janssen, Novartis, Pfizer and Eli Lilly. Dr. Faraone receives royalties from books published by Guilford Press: Straight Talk about Your Child’s Mental Health and Oxford University Press: Schizophrenia: The Facts. In the past 3 years, Dr. Buitelaar has been a consultant to / member of advisory board of / and/or speaker for Janssen Cilag BV, Eli Lilly, Bristol-Myer Squibb, Shering Plough, UCB, Shire, Novartis, and Servier. He is not an employee of any of these companies, and not a stock shareholder of any of these companies. He has no

Introduction

Smaller total brain and subcortical volumes have been linked to various forms of psychopathology including attention-deficit/hyperactivity disorder (ADHD) [1, 2]. Identifying mechanisms underlying brain volume alterations therefore is of great importance. Both genetic and environmental factors play a crucial role in determining interindividual variability in brain architecture [3]. Twin studies have revealed moderate to high heritability estimates for several brain structures (40–97%) [4, 5] and population-based and case-control studies show effects of specific genetic variants in brain volume variation, e.g. [6, 7, 8]. A number of studies have focused on associations with dopamine- and serotonin-related genes. For example, the short allele of a functional promoter polymorphism in the serotonin transporter gene (SLC6A4/ 5-HTT)—HTTLPR—has been related to smaller anterior cingulate gyrus and amygdala volumes in healthy adults [9], smaller frontal cortex in adults with obsessive-compulsive disorder and healthy controls [9, 10], and to smaller caudate, and both smaller and larger hippocampal volumes in adults with major depression [11, 12] and healthy controls [13]. The 10-repeat variant of a variable number tandem repeat (VNTR) polymorphism in the 3’untranslated region (UTR) of the dopamine transporter gene (SLC6A3/DAT1) has been related to smaller caudate volumes in children with ADHD and controls [14, 15]. Furthermore, variants of a VNTR in exon 3 of the dopamine receptor D4 gene (DRD4) have been associated with thinner frontal and parietal cortex thickness, and less prefrontal gray matter (GM) volume in children with ADHD and healthy controls as well [14, 16].

Besides genetic factors, environmental influences have also been associated with brain volume changes. So far, most studies have focused on severe negative experiences, such as childhood maltreatment and early life stress [17–21]. However, effects of positive influences, such as maternal warmth or support [22, 23] and, in animal studies, enriched environments [24] have also been reported, thereby broadening the view that only severe negative experiences are associated with brain structures. Nonetheless, studies have reported inconsistent results with findings of both smaller and larger volumes for both positive and negative environmental experiences. These inconsistencies could be due to methodological issues, such as differences in sample characteristics. For example, the majority of studies have used relatively small samples, with most including less than 100 participants (see e.g. [21] for a review). Other possible explanations for the inconsistent findings include differences in the timing of environmental exposure on the maturing brain [21], or the moderation of environmental influences or developmental effects by genes.

Genetic and environmental influences are not isolated from one another: genes and environment continuously work together throughout brain development [3, 25]. Gene-environment interactions (GxE) could therefore contribute to the heterogeneous findings when genes and environment are studied in isolation. Evidence for GxE effects on brain volumes is scarce and has come primarily from studies of the interaction of adverse life events with functional variants of the brain-derived neurotrophic factor (BDNF) gene and the serotonin-transporter-linked polymorphic region (HTTLPR). These studies showed that carriers of the Met allele of a single nucleotide polymorphism (SNP)—Val66Met—located in BDNF, or carriers of the HTTLPR short allele, had smaller hippocampal and anterior cingulate cortex (ACC) Gray Matter (GM) volumes when exposed to higher levels of (childhood) adversity [26–32]. Although, besides effects on GM volumes in the caudate and several other brain regions for HTTLPR short allele carriers, one study also found carriers of two HTTLPR long alleles had larger hippocampal and amygdala volumes when exposed to more stressful life events [31]. For BDNF, an interaction with early life stress on amygdala GM volume was found as well [26]. Yet, null findings have also been reported in studies of BDNF x Childhood adversity effects on hippocampal...
and amygdala volumes [32, 33]. Importantly, these studies focused only on genetic vulnerability to the effects of adverse life events. Although this is in agreement with the commonly applied diathesis-stress or dual-risk models [34], recent literature also emphasizes the protective effects of positive environments in combination with specific genetic variants [35–38]. Thus, studies reporting on GxE effects in relation to brain volumes so far have revealed only the negative side of the story.

Because the brain develops throughout the lifespan, one must consider developmental brain maturation when studying the effects of genes, the environment, and their interplay on brain volumes. Longitudinal developmental studies have demonstrated that overall cortical and subcortical GM volumes show curvilinear or ‘inverted-U’-like developmental trajectories with age, with the steepest growth found in early childhood and reductions reported from (pre)adolescence onwards, although there is much heterogeneity in developmental trajectories [39–41]. For example, individuals with ADHD appear to have different developmental trajectories of cortical thickness and subcortical volumes when compared to healthy controls [42–44]. It is likely that both direction and degree of environmental effects on neural structures depend on age, as child and adult studies on early adversity often show different or even opposite effects [19]. Moreover, it is likely that the timing of environmental exposures plays a role in which brain regions are sensitive to environmental effects [21]. Together, these findings suggest that developmental differences should be taken into account when studying GxE effects on brain volumes in childhood and adolescence.

In the present cross-sectional study we set out to investigate possible main and interaction effects of candidate genes and the social environment along with the possible moderating role of age on brain volumes in a large sample of children, adolescents, and young adults with and without an ADHD diagnosis. In addition, we investigated whether these effects depended on ADHD severity. We aimed to advance previous studies on brain volumes by investigating positive and negative environmental influences. Maternal expressed warmth and criticism as well as positive and deviant peer affiliation were chosen, as both parent and peer influences are important social environments during development and have been associated with neural alterations [22,45] and shown to affect child externalizing behavior, such as ADHD [46–48]; although the reverse influence (child behavior shaping social environment) has also been shown [49–51]. Furthermore, while these measures are less extreme in comparison to most environmental measures associated with brain structure differences so far (e.g. maltreatment), recent studies have shown evidence for more subtle measures such as parental warmth as well [22,23]. We focused on candidate variants in the DAT1, 5-HTT, and DRD4 genes. As reviewed above, these genetic variants have been associated with neural structure volumes, and have been associated with ADHD [52,53] and shown to interact with the environment in both children with and without ADHD [54]. We focused on total GM, caudate, and putamen volumes, as these have previously been shown to differ between the participants with ADHD and the (healthy) controls of the present sample [43] (i.e. participants overlapped between studies, with the exception that Greven et al. [43] excluded participants with subthreshold ADHD, while the present study excluded participants without information on EE, peer affiliation or genotyping). Individuals with ADHD had smaller total brain and GM volumes compared to controls (2.5–3.0%), whereas for caudate and putamen volumes the differences had a developmental nature; controls showed a decrease in size over age, while individuals with ADHD did not [43]. Because of the importance of developmental effects on brain maturation, we included a large sample with a broad age-range, which allowed us to explore the modulating role of age on the effects of genes, social environment, and their interaction on total GM, caudate, and putamen volumes.
Materials and Methods

Participants

Participants were selected from a follow-up (2009–2012) of the Dutch part of the International Multicenter ADHD Genetics (IMAGE) study, performed between 2003–2006 (see [55]). At first enrolment in IMAGE, families with at least one child with combined type ADHD and at least one biological sibling (regardless of ADHD diagnosis) were recruited, in addition to control families with at least one (unaffected) child and no formal or suspected ADHD diagnosis in first-degree family members. Inclusion criteria for children into IMAGE were an age between 5–19 years, European Caucasian descent, IQ ≥ 70, and no diagnosis of autism, epilepsy, general learning difficulties, brain disorders, or known genetic disorders (such as Fragile X syndrome or Down syndrome). All families were reinvited for a follow-up measurement with a mean follow-up period of 5.9 years (SD = .74) in Amsterdam or Nijmegen. At this follow-up, a comprehensive assessment protocol was administered, encompassing behavioral questionnaires, a diagnostic interview (assessing ADHD, oppositional defiance disorder [ODD], conduct disorder [CD]), and several neurocognitive measures from all family members, and an extensive MRI scanning protocol in participating children. Participants were asked to withhold use of psychoactive drugs for 48 hours before measurement. To determine ADHD diagnoses at the follow-up measurement, a standardized algorithm was applied to a combination of questionnaires and a semi-structured diagnostic interview. For a detailed description of the assessment protocol, including the diagnostic procedure see [56]. Informed consent was signed by all participants and their parents (for participants under 12 years of age only parents provided consent). The study, including its consent procedure, was approved by the local ethics committees (Centrale Commissie Mensgebonden Onderzoek).

In the current analyses participants were included when information was available on structural MRI, genotype, and maternal expressed emotion (EE) or peer affiliation. The final sample included N = 312 participants with ADHD, N = 80 with subthreshold ADHD (i.e., elevated symptoms of ADHD without meeting the full criteria for an ADHD diagnosis), and N = 357 participants without ADHD, from N = 402 families. Sample size depended in particular on the availability of EE and peer affiliation (N = 360 versus N = 726), as EE could only be assessed when the diagnostic interview was administered. This led to an unequal distribution of participants with or without an ADHD diagnosis in the EE (N = 279 with ADHD, N = 45 with subthreshold ADHD, N = 36 without ADHD) versus peer affiliation selection (N = 293 with ADHD, N = 78 with subthreshold ADHD, N = 355 without ADHD). Participant characteristics for the total sample as well as split for high and low ADHD severity are displayed in Table 1.

Measures

Parental expressed emotion. EE was assessed during the semi-structured diagnostic interview, using codings derived from the Camberwell Family Interview [60]. Only ratings of mothers were used in our study, as the data of fathers were far less complete. Warmth was assessed by the tone of voice, spontaneity, sympathy, and/or empathy toward the child (range 0–3). Criticism was assessed by statements which criticized or found fault with the child based on tone of voice and critical phrases (range 0–4) [61, 62]. Adequate inter-rater reliability has been reported for ratings of warmth and criticism using the Camberwell Family Interview (range .78–.91 and .79–.86, respectively [63]) and during the first measurement wave (the IMAGE study; range .71–1.00 [64]).

Peer affiliation. Peer affiliation was measured with the Friends Inventory [65]. Participants assessed their peers’ behavior on 18 items rated on a 4-point Likert scale (e.g., ‘my friends...
get good grades’, ‘my friends break the rules’; range 1 = ‘none of my friends are like that’ to 4 = ‘all of my friends are like that’). Scores were summed to yield either a positive or deviant peer affiliation score (each 9 items). Both measures have demonstrated good internal consistency reliability (range .78 – .92 [48, 66, 67]), and a mean inter-rater reliability of .71 has been reported between teacher and self-reports [67].

**ADHD severity.** ADHD severity was assessed using the DSM-IV total (raw) scores of a parent rating scale (CPRS-R:L; scale N [68]), and either a teaching rating scale (CTRS-R:L; scale N [69]) applied for children <18 years, or a self-report (CAARS-S:S; scale G [70]), applied...
for children ≥ 18 years. We used the CPRS-R:L and CTRS-R:L or CAARS-S:S as they were assessed in all participants (regardless of diagnostic status). Moreover, using a continuous measure of ADHD severity allowed us to retain as much information as possible, including the variability of scores among unaffected participants.

**Image acquisition and segmentation.** Imaging was conducted at two locations (Amsterdam and Nijmegen) using two similar 1.5 Tesla scanners (Siemens Sonata/Avanto), the same product 8-channel head-coil, and identical scan protocols. The protocol included two high-resolution T1-weighted MP-RAGE anatomical scans (176 sagittal slices, TR = 2730 ms, TE = 2.95 ms, TI = 1000 ms, flip angle = 7 deg, GRAPPA 2, voxel size = 1.0 x 1.0 x 1.0 mm, field of view = 256 mm). MRI scans were manually rated for quality, those that revealed poor quality or motion artefacts (N = 37) were excluded together with scans which yielded relevant incidental findings (N = 18) [71]. Volume estimates were averaged when participants had two good quality scans (N = 741), thereby improving signal-to-noise ratio.

**Brain volumes.** We used volumes that previously have been shown to differ between ADHD and controls included in the present study: of total GM, caudate nucleus, and putamen [43]. The unified procedure of the VBM 8.1 toolbox (http://dbm.neuro.unl-jena.de/vbm/) in SPM (default settings) was used to perform normalization, bias-correction, and segmentation into gray matter, white matter, and cerebrospinal fluid. Total gray and white volumes were calculated by summation of their tissue probability maps. Total brain volume was calculated by summing total gray and white matter volume. For the subcortical volumes, automated FIRST subcortical segmentation was applied to estimate left and right volumes of the caudate and putamen. FIRST, part of FMRIB's Software Library (FSL), performs registration and shape modeling of the above regions in MNI152 standard space [72].

**Genotyping.** For the IMAGE sample (parents and children), DNA was extracted from blood samples or immortalized cell lines at Rutgers University Cell and DNA Repository, New Jersey, USA. Genetic variants in DRD4, DAT1, and 5-HTT were genotyped by the IMAGE consortium [73, 74]. Standard PCR protocols were used for all VNTR markers and amplified products were visualized on 2% agarose under UV light. Additional NeuroIMAGE samples were collected in the form of a saliva sample using Oragene kits (DNA-Genotek; see www.neuroimage.nl). For those, VNTRs were genotyped using standard PCR protocols at the Department of Human Genetics of the Radboudumc, Nijmegen. After the PCR, fragment length analysis was performed on the ABI prism 3730 Genetic Analyser (Applied Biosystems, Nieuwekerk a/d IJssel, The Netherlands) and results were analyzed with GeneMapper® Software, version 4.0 (Applied Biosystems). No deviations from Hardy-Weinberg Equilibrium were found (DAT1 p = .78, 5-HTT p = .13, DRD4 p = .15). For the data analyses, participants were divided into groups based on the presence or absence of the 9-repeat of the DAT1 3'UTR VNTR, the short allele of HTTLPR, or the 7-repeat of the DRD4 exon 3 VNTR, respectively.

**Data Analyses**

**Gene-environment correlations.** The presence of gene-environment correlations (rGE) could bias potential GxE by providing an alternative explanation for the relationship between environmental measures and genes [54, 75]. Therefore, Pearson and Spearman correlation analyses were performed to test for rGE between maternal or adolescent genotype and the environmental predictors.

**Main analyses.** All analyses were performed on the total sample, that is, participants with and without ADHD. Linear mixed model analyses investigated the effects of EE, peer affiliation, genotype, and GxE interactions on each volumetric measure (total GM volume and total, left, and right caudate nucleus and putamen volumes). Models were run with and without the
interaction term separately. Separate models were run for each environmental predictor: warmth, criticism, and positive and deviant peer affiliation, and for each gene (DAT1, 5-HTT, DRD4) as well. Consequently, there were 4 environmental predictors, 3 genes, and 7 outcome measures.

To correct for familial dependency, as a number of participants belonged to the same families, we estimated a random intercept for family in each model. A random intercept accounts for familial dependency by estimating the correlations between cases within families. Age, gender, and collection site were included as covariates. The analyses of the subcortical volumes included total brain volume as an additional covariate. For total GM volume, we added total white matter volume as an extra covariate. Because longitudinal developmental studies have found curvilinear developmental trajectories with age for cortical and subcortical GM volumes, with the steepest growth found in early childhood and reductions reported from (pre)adolescence onwards [39–41], we tested 2- and 3-way interactions with age and age or Age\(^2\) (i.e., Age/ Age\(^2\)xG, Age/Age\(^2\)xE, and Age/Age\(^2\)xGxE) which were dropped from the model when not significant or nominally significant (i.e., did not survive correction for multiple testing). All continuous predictors and covariates were centered around the mean.

**Multiple testing correction.** A multiple comparisons correction was employed which adjusts for correlated tests based on the effective number of independent comparisons (M\(_{eff}\)) [76]. The M\(_{eff}\) was derived from the Eigenvalues of a correlation matrix between the outcome measures adjusted for covariates (age, gender, collection site, and total brain or total white matter volume for subcortical and total GM volumes respectively). In the case of zero correlations between the outcome measures, the M\(_{eff}\) adjusted p-value would be equivalent to a Bonferroni correction. Thus the M\(_{eff}\) procedure is particularly suited for correlated comparisons (such as total, left and right putamen volumes) and corrects for multiple testing balancing between being overly lenient or conservative. The effective number of comparisons was determined to be 3.55 and the adjusted p-value threshold: 0.05/3.55 = .014.

**Sensitivity analyses.** Sensitivity analyses were performed when significant effects that survived the multiple correction threshold were found. First, Regions of significance (RoS), simple slope, and slope difference tests were performed with an application designed for probing 2- and 3-way interactions (http://www.jeremydawson.co.uk/slopes.htm [77]). For interactions with age, slope tests were estimated with non-quadratic age. Second, to investigate the role of ADHD severity, analyses were rerun including main and interaction effects of ADHD severity. Furthermore, separate sensitivity analyses were performed to check whether significant effects were present in participants while sequentially controlling for effects of medication history, estimated IQ, and comorbid ODD or CD diagnosis. All analyses (except for RoS and slope tests) were performed with the Statistical Package for the Social Sciences, version 20.0.

**Results**

Analyses of gene-environment correlations (rGEs) revealed significant correlations between maternal warmth and adolescent DAT1 (r = -.11, p = .045), 5-HTT (r = -.11, p = .040), and DRD4 genotypes (r = -.11, p = .048), and maternal DAT1 genotypes (r = -.15, p = .005). We also found a significant correlation between maternal criticism and adolescent DAT1 genotypes (r = .12, p = .024) (see S1 Table). Considering the size of these associations, there was no reason to believe that these rGEs may have biased significant GxE interactions. In the next sections on main and GxE effects, only results of the mixed model analyses that survived correction for multiple testing are discussed (p < .014). Nominal significant results can be found in the Supporting Information (SI) in S2 and S3 Tables.
Main effects of candidate genes and environments and the effect of age

No significant main effects of maternal warmth or criticism were found (all p-values > .166; see S2 Table). A small main effect of deviant peer affiliation was found on the left caudate volume ($B = -.01, p = .012$), indicating that more deviant peer affiliation was related to smaller caudate volumes. This effect was also present in the total and right caudate volumes, but did not survive correction for multiple testing ($B = -.02, p = .017; B = -.01, p = .031$ respectively). No main gene effects were found that survived the correction for multiple testing (all $p$-values > .015).

Investigation of the effects of linear and non-linear age yielded a significant interaction between deviant peer affiliation and the quadratic effects of age ($age^2$) on total GM volume ($p = .001$). As shown in Fig 1, participants with low deviant peer affiliations (- 2SD) had larger total GM volumes when older ($p = .001$), while participants with high deviant peer affiliations (+ 2SD) showed no association with age ($p = .112$).

GxE interactions and the role of age

For total GM volume we found two 3-way interactions between positive peer affiliation, DAT1, and $age^2$ ($p = .007$), as well as between positive peer affiliation, 5-HTT, and $age^2$ ($p = .012$). Simple slope analyses revealed significant slopes (i.e., different from slope = 0) for carriers of the DAT1 9-repeat or two 5-HTT long alleles when scoring either low or high on positive peer affiliation (- 2SD: $p_{DAT1} = .012, p_{5-HTT} = .034$; + 2SD: $p_{DAT1} = .042, p_{5-HTT} = .017$). These slopes differed significantly from each other as well ($p_{DAT1} = .009, p_{5-HTT} = .014$). As shown in Fig 2, carriers of the DAT1 9-repeat or two 5-HTT long alleles with low positive peer affiliations had larger GM volumes with age, while participants with the same genotype, but high positive peer affiliations had smaller GM volumes with age. Slopes for participants with the DAT1 10/10 genotype or 5-HTT short allele were not significant (- 2SD: $p_{DAT1} = .368, p_{5-HTT} = .153$; + 2SD: $p_{DAT1} = .272, p_{5-HTT} = .346$). Still, slopes of DAT1 9-repeat carriers and 10-repeat homozygotes differed significantly when they scored low (- 2SD, $p = .010$) or high on positive peer affiliation (+ 2SD, $p = .021$). The same was true for slopes of 5-HTT short allele carriers and long allele homozygotes (- 2SD, $p = .012$; + 2SD, $p = .014$).

For putamen volume significant 3-way interactions with age were found as well. Fig 3A and 3B show the interaction between DRD4, positive peer affiliation, and age on the right putamen.
volume ($p = .012$). Significant slopes were found for $DRD4$ 7-repeat carriers scoring either low or high on positive peer affiliation (- $2SD$: $p = .021$; + $2SD$: $p = .003$), and the difference between the two slopes was significant as well ($p = .004$). Thus, 7-repeat carriers showed differential associations between the right putamen volume and age depending on the amount of positive peer affiliation, i.e., a negative association when scoring low on positive peer affiliation, but positive when scoring high on positive peer affiliation. Although slopes of participants without the $DRD4$ 7-repeat allele were not significant (- $2SD$: $p = .492$; + $2SD$: $p = .308$), significant slope differences were found between 7-repeat carriers scoring high on positive peer affiliation and participants without the 7-repeat with either low ($p = .007$) or high positive peer affiliations ($p = .003$).

Finally, an interaction was found between $DAT1$, criticism, and age on total putamen volume ($p = .005$). This effect was present in both left and right putamen volumes (left: $p = .009$; right: $p = .006$). Here, significant slopes were found for the $DAT1$ 10/10 genotype with either low or high maternal criticism (- $2SD$: $p = .026$; + $2SD$: $p = .043$), which differed significantly
from each other as well (p = .016). As can be seen in Fig 3C and 3D, participants with two 10-repeat alleles exposed to high maternal criticism had smaller putamen volumes with age, but participants exposed to low criticism had larger putamen volumes with age. Slopes of DAT1 9-repeat carriers were not significant (-2SD years; p = .095; +2SD years; p = .155). Analyses of slope differences revealed significant differences between DAT1 9-repeat carriers and 10-repeat homozygotes when scoring low (-2SD, p = .023) or high on maternal criticism (-2SD, p = .006). Highly similar results were found when investigating left and right putamen volumes separately.

Sensitivity analyses

Through sensitivity analyses we investigated the possible role of ADHD on the significant main effects and interactions described above. When the analyses were rerun including main
and interaction effects of ADHD severity, no significant interactions with ADHD severity were found (all p-values > .133). Including a main effect of ADHD severity did not change the aforementioned significant main or GxE/Age^2 effects either. The p-value of the interaction between DAT1, positive peer affiliation, and age^2 did drop slightly, thereby becoming nominally significant (p = .018). The same was true for the interaction between 5-HTT, positive peer affiliation, and age^2 (p = .015). Furthermore, rerunning analyses consecutively including IQ, ODD, CD, or medication history in the model yielded highly similar results. Note that for the main effect of deviant peer affiliation, the p-value dropped to nominal significance (all p-values < .037) when including IQ, ODD, or CD. Similarly, p-values dropped slightly when ODD or medication history was included for the interaction between 5-HTT, positive peer affiliation, and age^2 on total GM volume and the 3-way interactions on putamen volume (all p-values < .043).

**Discussion**

We investigated the effects of functional variants in the 5-HTT, DAT1, and DRD4 genes, the social environment, and the interactions between genes and environment on brain volumes in a large sample of children, adolescents, and young adults with and without ADHD. We took a developmental approach, examining the impact of age on main and interaction effects of candidate genes and social environments. As expected, few (i.e., one) main effect was observed, of deviant peer affiliation on left caudate volume. Instead, we observed a complex pattern of the following two-way and three-way interactions: an interaction between deviant peer affiliation and age^2 for total GM volume, and between 5-HTT, DAT1, or DRD4 variants and positive peer affiliation or maternal criticism on total GM and putamen volumes. These findings were independent of ADHD severity. The results extend findings from twin studies [78–80] that genetic effects are developmentally sensitive by showing gene-by-environment interactions appear to be developmentally sensitive as well.

We found different age-effects for total GM and putamen volumes, depending on genotype and/or environmental exposure. In agreement with age-related reductions of total GM volume found in longitudinal studies [39, 40], participants scoring high on positive peer affiliation carrying the DAT1 9-repeat or two HTTLPR long alleles had smaller total GM volumes with age. Moreover, participants with the same genotype, but low positive peer affiliation had larger GM volumes with age. These findings are in line with a longitudinal study reporting regional GM reductions with age in adolescents exposed to high positive maternal behavior, but increased putamen volumes when exposed to maternal aggression [22]. However, we also found positive associations between total GM and age in participants scoring low on deviant peer affiliation, regardless of genotype, while participants with high deviant peer affiliation had no association with age. Similarly, for putamen volume, carriers of the DRD4 7-repeat or DAT1 10/10 genotypes had larger volumes over age when exposed to high positive peer affiliation or low maternal criticism respectively, but opposite patterns, i.e., smaller volumes over age, when exposed to low positive peer affiliation or high criticism.

Although, equivalent to total GM, decreased putamen volumes over age have been reported [40], different age-effects of putamen volume have been found for participants with ADHD in comparison to healthy controls by our group [43]. Thus, one might expect the opposing age patterns found in the present study to be related to ADHD. However, none of the current findings were moderated by ADHD severity, suggesting they contribute to total GM, putamen, and caudate volumes in a more general manner, similar for individuals with and without ADHD. Previous GxE studies on brain volumes have been few and have mostly included (young) adults, leaving GxE effects on child or adolescent brain volumes under-investigated. So far, of
the included candidate genes, only the 5-HTT gene has previously been reported to interact with stressful life events [29–31, 81]. In a previous study using the present sample, stress exposure was associated with less GM volume in the precentral gyrus, middle and superior frontal gyrus, frontal pole, and cingulate gyrus in carriers of the 5-HTT short allele compared to long allele homozygotes [81]. Similar results have been found in adults studies, showing smaller hippocampal volumes for carriers of the HTTLPR short allele when exposed to childhood adversity [29, 30], though Canli et al. [31] reported that only long allele homozygotes had a positive association between stressful life events and hippocampal or amygdala GM volumes. In other regions, such as the ACC or caudate, both short and long allele carriers showed opposite associations with life stress. These opposite effects are in line with the differential effects of DAT1 variants we found on total GM versus putamen volumes. That is, DAT1 9-repeat carriers showed age-dependent associations between positive peer affiliation and GM volumes, while DAT1 10-repeat homozygotes showed differential associations between maternal criticism and putamen volumes. This could suggest that different variants of the same gene are susceptible to different environments, which could further depend on which brain region is focused on. In contrast to what we expected, both genotypes showed the same direction of association with two opposing environments; participants with high positive peer affiliations or high maternal criticism both had smaller GM volumes with age, while participants with low positive peer affiliations or low criticism showed the opposite pattern.

Besides different effects of the same gene, differential effects of positive peer affiliation were found for carriers of specific gene variants as well, i.e., participants scoring low on positive peer affiliation with the DAT1 9-repeat or two HTTLPR long alleles had larger total GM volume, while those with low positive peer affiliation and the DRD4 7-repeat allele had smaller putamen volumes with age, with the opposite pattern found in participants scoring high on positive peer affiliation. This suggests positive peer affiliation can have different effects depending on which brain volume or genotype is focused on. What our findings most consistently show is that (for carriers of specific gene variants) the direction of associations between certain social environments and brain volumes depends on developmental stage. This is illustrated by our finding that, for carriers of the same gene variants, low positive peer affiliation or high maternal criticism was related to smaller total GM or putamen volumes in preadolescents, but to larger volumes in young adults. Such differential effects were found in each of the reported two- and three-way interactions. In agreement, our group has shown that associations between the 5-HTT, DAT1, DRD4 and neurocognitive functioning, such as inhibition and working memory, depended on age as well [82]. These findings highlight the importance of including age when studying genetic and environmental effects on the neural architecture of children, adolescents and young adults, as the direction of associations likely depend on developmental stage.

There are several possible explanations for the finding of developmentally sensitive GxE effects on total GM and putamen volumes. Neurotransmitters have an important role in synaptic and neural plasticity [83], and the age-dependent GxE effects may thus be related to differences in dopamine and serotonin availability associated with DRD4, DAT1 and 5-HTT variants. Specifically, the DRD4 7-repeat and HTTLPR short alleles are associated with decreased transcriptional activity, leading to increased levels of dopamine and serotonin [84, 85]. For DAT1 there are mixed results about whether the 9- or 10-repeat shows increased or decreased expression. However, a recent meta-analysis showed that the 9-repeat allele was associated with increased in vivo striatal dopamine transporter activity in adults independent of the presence of neuropsychiatric disorders [86]. Expression levels of genes can differ across developmental stages [87] and dopamine and serotonin levels in the brain have been shown to differ over age as well [88]. Besides genetic and developmental influences, environmental factors, such as maternal deprivation or environmental enrichment, have also been associated
with differential neurotransmitter levels [89]. Furthermore, other processes during brain development, including neuronal pruning, myelination and hormonal influences are believed to be influenced by genetic and environmental factors as well [24, 90]. Together, these findings reveal many important developmental processes, steered by environmental and genetic factors, which together determine one’s neural architecture.

For caudate volume, one developmentally stable main effect was found. Higher deviant peer affiliation was associated with slightly smaller caudate volumes. This agrees with a previous study that found early life stress was related to smaller caudate volumes in adults [91]. Similarly, smaller volumes of other brain regions, such as the hippocampus, have been associated with adverse psychosocial experiences [18, 19, 21], but findings are not consistent since null-findings, and even larger volumes in relation to adverse environments have been reported [18, 19, 21].

All but one of the reported effects in the current study were found in relation to peer affiliation. While peer influences have been linked to functional brain differences [45, 92] and white matter structure [93], no prior studies have addressed associations with volumetric alterations. Besides peer presence and peer verbal abuse investigated in the aforementioned studies [45, 92, 93], our results indicate that the type of peers seems to be relevant for structural brain differences. The associations between deviant or positive peer affiliation and brain volume alterations could be the result of intrinsic or extrinsic factors not investigated in this study. An example would be that structural brain alterations are first and foremost associated with intrinsic personality traits (such as high sensation seeking or low conscientiousness) that fit well with deviant peer affiliation. Indeed, studies have revealed both positive and negative associations of traits such as extraversion and conscientiousness with (regional) gray and white matter volumes [94–97]. Likewise, circumstances such as neighborhood quality could be an example of extrinsic factors underlying peer affiliation. Future studies are needed to investigate whether the association between peer affiliation and brain volumes reflects an effect of other factors such as personality traits or environmental conditions.

Our findings should be viewed in the light of certain strengths and limitations. Strengths were the use of a large well-characterized sample, inclusion of both positive and negative environments, assessment of both parental- and peer influences, and a developmentally sensitive approach. A limitation has been the cross-sectional MRI study design, only longitudinal MRI studies can clarify the direction of causality. Establishing the direction of effects is particularly difficult when focusing on the social environment as it can also (partly) be driven by child effects. Indeed, both maternal EE and peer affiliations have not only been suggested to influence child behavior, but in turn be influenced by child behavior as well [46–49, 98–101]. Thus, it is most likely that bidirectional effects exist between the social environment and child behavior. Furthermore, not all participants included had an expressed emotion (EE) measurement, as the design of our study was such that EE was only assessed when a full diagnostic interview was administered. This led to loss of power, and unequal numbers and an unequal distribution of ADHD and controls in the EE versus peer affiliation analyses. Nevertheless, sensitivity analyses revealed no significant effects of ADHD severity on the reported findings. In addition, although included measures were chosen a-priori based on previous literature, our findings only shed light on a small part of brain variability based on only a few gene variants. Finally, considering the novel and explorative nature of our study, we did not employ a very stringent correction for multiple testing. If we would have applied the most stringent correction—correcting for the total number of environmental measures (n = 4), genes (n = 3), and outcome measures—the corrected p-threshold would have been p = 0.05/(4*3*3.55) = .0012 instead of p = .014. In this case, all but the interaction between deviant peer affiliation and age would not have survived the correction for multiple testing. Therefore, we emphasize the necessity of independent replication studies.
In conclusion, beside a main effect of deviant peer affiliation on caudate volume, we found multiple developmentally sensitive GxE effects on total GM and putamen volume. Despite previously reported differences in total GM, caudate, and putamen volumes between individuals with ADHD and healthy controls, our results were independent of ADHD severity. Both children, adolescents, and young adults with and without ADHD showed differential sensitivity to environmental influences, depending on genotype and age. This suggests that interactions between genes and the social environment contribute in a general way to the included cortical and subcortical brain volumes and are not specific for ADHD. Nevertheless, variation in these brain volume sizes could be relevant for other clinical or functional outcomes not studied here. The caudate nucleus and putamen, for example, have been linked to cognitive, executive and emotional functioning [102].

While it is clear that our complex findings are in need of replication, our results stress the importance of a developmentally sensitive approach when investigating genetic and social environmental influences on interindividual brain volume variability. The failure to do so could potentially explain the diverse and sometimes opposing results of main environmental and genetic effects on brain volumes reported so far.

Supporting Information

S1 Table. Correlation analyses between environmental measures, maternal or adolescent’s genotype, brain volumes and ADHD severity. Pearson correlation analyses were performed for all variables, except for correlations with peer affiliation and ADHD severity, for which Spearman correlations analyses were performed. a 9-repeat allele present or absent; b short allele present or absent; c 7-repeat allele present or absent short allele present or absent. * Significant at \( p \leq .05 \) ** Significant at \( p \leq .01 \)

S2 Table. Mixed model analyses testing the separate main effects of maternal expressed emotion, peer affiliation and plasticity genes on brain volumes. EE = expressed emotion; PA = peer affiliation. a Reference group: 10/10 absent; b reference group: short allele absent; c reference group: 7-repeat absent. Findings in bold are significant after correction for multiple testing (\( p < .014 \)), findings in bold and italic are nominally significant (i.e. not significant after correction for multiple testing; \( p \leq .05 \)). All analyses were corrected for age, gender, and collection site. Additionally, total brain volume was included in analyses of subcortical volumes, and total white matter in analyses of total gray matter as well.

S3 Table. Mixed model analyses testing interaction effects between plasticity genes and maternal expressed emotion or peer affiliation on brain volumes. PA = peer affiliation. a Reference group: 10/10 absent; b reference group: short allele absent; c reference group: 7-repeat absent. Findings in bold are significant after correction for multiple testing (\( p < .014 \)), findings in bold and italic are nominally significant (i.e., not significant after correction for multiple testing; \( p \leq .05 \)). All analyses were corrected for age, gender, and collection site. Additionally, total brain volume was included in analyses of subcortical volumes, and total white matter in analyses of total gray matter as well. In each model 2- and 3-way interactions with age or age\(^2\) were tested and removed when not significant (\( p > .05 \)) or nominally significant (\( p \geq .014 \)).

Acknowledgments

The authors thank all the families who participated in this study and also the researchers who collected the data.
Author Contributions
Conceived and designed the experiments: JR CH JB AAV. Performed the experiments: JR. Analyzed the data: JR CH. Contributed reagents/materials/analysis tools: JR CH JB AAV SF. Wrote the paper: JR CH. Edited the manuscript and contributed conceptual improvements: CH JB AAV PH BF DH JO SF.

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