Cover Page

The handle http://hdl.handle.net/1887/36475 holds various files of this Leiden University dissertation.

Author: Windhorst, Dina Aafje (Dafna)
Title: Differential susceptibility to parenting : exploring new approaches
Issue Date: 2015-11-24
BEYOND MAIN EFFECTS OF GENE-SETS.
HARSH PARENTING MODERATES THE ASSOCIATION BETWEEN A DOPAMINE GENE-SET AND CHILD EXTERNALIZING BEHAVIOR

Dafna A. Windhorst, Viara R. Mileva-Seitz, Ralph C.A. Rippe, Henning Tiemeier, Vincent W.V. Jaddoe, Frank C. Verhulst, Marinus H. van IJzendoorn, and Marian J. Bakermans-Kranenburg

Manuscript submitted for publication.
ABSTRACT

We extended gene-based and gene-set approaches by examining the interplay of dopaminergic genes and harsh parenting experiences on child externalizing behavior. Genetic variants in 12 autosomal dopaminergic genes were available in an ethnically homogenous part of a population-based cohort. Harsh parenting was assessed with maternal and paternal reports at age three. Externalizing behavior was assessed with the Child Behavior Checklist at age five (71 ± 3.7 months). We examined the effects of paternal, maternal, and pooled harsh parenting (n = 1710, 1881, and 1916 respectively). We conducted gene-set analyses of the association between variation in dopaminergic genes and externalizing behavior, stratified for harsh parenting. The association was statistically significant or approached significance for children without harsh parenting experiences, but was absent in the group with harsh parenting. Similarly, significant associations between single genes and externalizing behavior were only found in the group without harsh parenting. Effect sizes in the groups with and without harsh parenting did not differ significantly. Gene-environment interaction tests were conducted for individual genetic variants, resulting in two significant interaction effects (rs1497023 and rs4922132) after correction for multiple testing. Our findings are suggestive of gene-environment interplay, with associations between dopamine genes and externalizing behavior present in children without harsh parenting, but not in children with harsh parenting experiences. Harsh parenting may overrule the role of genetic factors in externalizing behavior. Gene- and gene-set analyses offer a promising new alternative to analyses focusing on single candidate polymorphisms when examining the interplay between genetic and environmental factors.

Keywords: dopamine; externalizing behavior; gene-environment interaction; parenting; preschoolers; gene-set; harsh parenting; behavior problems dopaminergic system; child development
INTRODUCTION

Young children with externalizing problem behaviors, including oppositional behavior, aggression, and attention problems, are at increased risk for problems later in life. Such problems include poor school performance, peer-relationship issues, aggression, violence, and crime (Campbell, Shaw, & Gilliom, 2000; Campbell, Spieker, Burchinal, Poe, & The NICHD Early Child Care Research Network, 2006; Reef, Diamantopoulou, Van Meurs, Verhulst, & Van der Ende, 2011). Child externalizing behavior is predicted both by child factors, including genetics, and by environmental factors, including parenting behavior (e.g., Haberstick, Schmitz, Young, & Hewitt, 2005; Miner & Clarke-Stewart, 2008; Saudino, Carter, Purper-Ouakil, & Gorwood, 2008). The effects of these factors are not independent of one another. Significant progress has been made in understanding how environmental factors interact with genetic factors to affect child development (Ellis, Boyce, Belsky, Bakermans-Kranenburg, & Van Ijzendoorn, 2011; Moffitt, Caspi, & Rutter, 2005; Rutter, 2006), but the genetic factors are usually considered independently in analyses. In the present study we examine how harsh parenting interacts with genetic variation across multiple child dopamine genes in shaping child externalizing behavior. This is a novel approach to the study of gene-environment interactions (GxE), as it takes into account genetic variation across whole genes and gene-sets (a combination of related genes). This approach complements the more conventional analyses at the level of single nucleotide polymorphisms (SNPs).

The pursuit of genetic underpinnings of complex child problem behaviors, including externalizing behavior and aggression, has yielded limited success. Longitudinal twin studies of childhood aggression show moderate to high estimates of heritability (51-72%) (Hudziak et al., 2003). Common polymorphisms in several candidate genes have been reported in association with childhood aggression and externalizing (Oades et al., 2008; Takahashi, Quadros, De Almeida, & Miczek, 2012; Volavka, Bilder, & Nolan, 2004)), but these effects have not held up after meta-analysis (Vassos, Collier, & Fazel, 2014). The candidate gene approach has been generally criticized for non-replication and potential false discovery, related to publication bias for positive findings, inadequate power to detect true effects, and improper correction for multiple testing (Sullivan, 2007; Tabor, Risch, & Myers, 2002). Furthermore, it has been argued that variation in complex traits such as externalizing behavior is not readily attributable to variation at single genetic loci or on single genes, but rather to variation at multiple genetic loci across the genome (Reif & Lesch, 2003). Thus the lack of robust effects in past candidate gene studies might be partially owing to the fact that aggression is a complex phenotype, and likely has a polygenic nature in humans, as has been demonstrated previously in mice (Brodkin, Goforth, Keene, Fossella, & Silver, 2002). Expanding the search for genetic correlates of aggression-related phenotypes in children and adults via genome-wide association studies (GWAS) has allowed testing for associations between millions of SNPs across the genome and the outcome without an a priori hypothesis. These types of studies treat all SNPs independently, and have provided only suggestive evidence for a genetic main effect on aggressive phenotypes (Dick et al., 2004; Tielbeek et al., 2012; Viding et al., 2013).

One of the factors that can bridge the gap between the substantial heritability estimates and the lack of individual genetic associations is the environment. Genetic effects are now widely accepted
as being open to moderation by environmental influences. Until recently, most studies investigating GxE have focused on specific polymorphisms within candidate genes with functions implicated in the outcome of interest (Manuck & McCaffery, 2014), although candidate GxE studies received similar critiques as studies searching for main effects (Dick et al., 2015; Duncan & Keller, 2011). Given the polygenic nature of complex phenotypes, considering the moderating effects of environment at a multi-gene level becomes important. Candidate genes studied in the context of parenting and externalizing behavior are often related to the dopamine system, a neurotransmitter system involved in motivation and reward processing, and in the regulation of emotion, action, and attention (Keltikangas-Järvinen & Salo, 2009; Robbins & Everitt, 1999). For instance, polymorphisms on the dopamine receptor D2 (DRD2), D4 (DRD4), and dopamine transporter gene (DAT) genes reportedly moderate the association between parenting and child problem behavior (e.g., Bakermans-Kranenburg & Van Ijzendoorn, 2011; Keltikangas-Järvinen & Salo, 2009). Such candidate genes can be considered indicators of underlying genetic pathways.

In contrast to hypothesis-based candidate gene studies, Genome-Wide Environmental Interaction (GWEI) studies investigate interaction effects between individual SNPs located across the genome and environmental factors on an outcome, in a hypothesis-free way similar to GWAS studies (Aschard et al., 2012). Genome-wide analyses require extremely large samples because the high numbers of SNPs tested require conservative corrections for multiple testing. As a consequence, it is difficult to detect small effects of individual SNPs (Hirschhorn & Daly, 2005), particularly on complex traits. Moreover, in very large samples the careful assessment of environmental data is complicated due to practical issues (Khoury & Wacholder, 2009).

These challenges impede GxE research and highlight the need for new methods. One approach used in the search for genetic main effects on behavioral outcomes – the gene-based and gene-set analysis approach – aggregates genetic variance of many SNPs within a gene or within a pre-defined gene-set that includes multiple functionally or biologically related genes (e.g. Fridley & Biernacka, 2011; Lips et al., 2012; Winham & Biernacka, 2013). Testing joint effects of multiple SNPs greatly reduces multiple testing, and consequently improves power. Thus SNPs with individually marginal effect sizes could have a significant combined effect (e.g. Fridley & Biernacka, 2011; Lips et al., 2012; Winham & Biernacka, 2013). Another advantage is that the results are more biologically interpretable. That is more difficult for individual SNPs, as the functional role of many individual genetic variants remains unknown. Although gene-based and gene-set approaches have so far mostly been used to examine genetic main effects, they also provide new opportunities for studying GxE. Unfortunately, most statistical programs do not yet have the option to include estimations for gene-environment interaction effects (Winham & Biernacka, 2013).

In this exploratory study we extend the gene-based and gene-set approaches to include GxE, using a tool for gene-based and gene-set analyses: Joint Association of Genetic Variants (JAG) (http://ctglab.nl/software/jag, Lips, Kooyman, De Leeuw, & Posthuma, 2015). We examine the interplay between harsh parenting and variation across children’s dopamine genes in association with externalizing behavior, in a large ethnically homogenous subsample of a population-based cohort study. We assess aggregated genetic variation across a ‘set’ of multiple dopamine genes. Because the inclusion of an environmental
moderator (presence or absence of harsh parenting) in this gene-set analysis is novel and exploratory, we can only speculate about a directional hypothesis. On the one hand, aggregated genetic variation might have strongest effects in high-risk environments (harsh parenting), which is consistent with the hypothesis that negative environments ‘trigger’ risk alleles (Shanahan & Hofer, 2005). On the other hand, aggregated genetic variance might have strongest effects in low-risk environments (no harsh parenting) in which environmental homogeneity allows for more genetically driven phenotypic differentiation. This is one of the first studies to employ a gene-set analysis in tests of GxE effects on complex behavior. This approach can offer an important alternative or complement to candidate gene and GWEI studies in the search for genetic variation underlying individual differences in behavior.

METHOD

Setting
The current investigation is embedded in the Generation R Study, a prospective cohort study investigating development from fetal life into young adulthood in Rotterdam, the Netherlands (Jaddoe et al., 2012, Tiemeier et al., 2012). Briefly, all pregnant women living in Rotterdam with an expected delivery date between April 2002 and January 2006 were invited to participate. The study has been approved by the Medical Ethical Committee of the Erasmus Medical Center, Rotterdam. Written informed consent was obtained from all adult participants.

Study population
Genetic information and questionnaire data at child age 3 (harsh parenting) and 5 (CBCL) were included in the analyses. We report separate analyses for harsh parenting by the father, the mother, and fathers and mothers combined (referred to as pooled harsh parenting). Genetic data were available for 2,830 Caucasian children, for whom paternal, maternal and pooled harsh parenting data were available for 1,817, 2,005, and 2,042 children respectively. If data were available for multiple siblings, one child per sibling pair was randomly selected and included in the analyses. The final numbers for paternal, maternal and pooled harsh parenting consisted of 1,710, 1,881 and 1,916 children respectively. Sample characteristics are presented in Table 1. The groups with and without harsh parenting differed on a number of characteristics, including gender, parity, birth weight, age of the mother, maternal and paternal educational level, depressive and anxiety symptoms of mother and father, and child externalizing behavior.

Non-response analyses were performed to compare the children with parenting data to the children with missing parenting data. Compared to excluded children, included children had on average a higher birth weight, were more often primiparous, and were younger at CBCL assessment. The mothers of these children and their partners were older at intake, educational level of the mother and partner was higher, parents were more often married or living together, fewer mothers smoked but more mothers drank alcohol during pregnancy. Mothers of included children had higher depressive and anxiety symptom scores compared to mothers of excluded children.
### Table 1 | Sample characteristics

| Group | Paternal Harsh parenting | Maternal harsh parenting | Pooled harsh parenting |
|-------|--------------------------|--------------------------|------------------------|
|       | Total | Without harsh parenting | With harsh parenting | Total | Without harsh parenting | With harsh parenting | Total | Without harsh parenting | With harsh parenting |
| $n$   | 1710  | 1113                     | 597                    | 1881  | 1208                     | 673                    | 1916  | 958                     | 958                    |

| Child characteristics | | | | | | | |
|-----------------------|---|---|---|---|---|---|---|---|
| Gender (% boys)       | 50.40 | 46.00 | 58.60 | ** | 51.10 | 49.50 | 54.10 | 51.00 | 46.90 | 55.20 | ** |
| Parity (% primiparous) | 62.20 | 64.00 | 58.80 | * | 61.60 | 63.40 | 58.30 | * | 61.60 | 64.00 | 58.50 | ** |
| Birth weight          | 3563.11±520.89 | 3542.45±507.73 | 3601.63±542.91 | * | 3553.58±525.39 | 3539.25±519.93 | 3579.30±534.48 | 3553.30±523.15 | 3510.51±508.52 | 3596.09±534.23 | ** |
| Child age at CBCL completion | 71.21±3.65 | 71.16±3.75 | 71.29±3.47 | | 71.21±3.55 | 71.13±3.43 | 71.35±3.74 | | 71.21±3.59 | 71.14±3.60 | 71.29±3.58 |
| CBCL sumscore externalizing behavior | 6.70±5.98 | 5.79±5.31 | 8.40±6.75 | ** | 6.76±6.00 | 5.90±5.45 | 8.31±6.61 | ** | 6.79±6.01 | 5.64±5.33 | 7.95±6.42 | ** |

| Parent characteristics | | | | | | | |
|------------------------|---|---|---|---|---|---|---|---|
| Age mother at intake   | 32.13±3.77 | 32.04±3.84 | 32.30±3.63 | | 32.14±3.84 | 32.28±3.80 | 31.89±3.91 | * | 32.12±3.85 | 32.17±3.94 | 32.07±3.76 |
| Age father at intake   | 34.15±4.72 | 34.18±4.82 | 34.08±4.55 | | 34.14±4.71 | 34.15±4.55 | 34.10±5.00 | | 34.14±4.74 | 34.26±4.75 | 34.01±4.73 |
| Educational level mother (% high) | 77.80 | 78.00 | 77.20 | | 77.00 | 78.20 | 74.00 | * | 76.80 | 78.40 | 75.30 |
| Educational level father (% high) | 75.30 | 75.90 | 74.30 | | 73.70 | 76.40 | 68.90 | ** | 73.60 | 75.40 | 71.80 |
| Marital status mother/father (% married/living together) | 95.10 | 94.70 | 95.80 | | 93.70 | 93.50 | 93.90 | | 93.80 | 93.00 | 94.60 |
| Maternal smoking during pregnancy (% never smoked during pregnancy) | 80.90 | 82.30 | 78.40 | | 80.30 | 80.50 | 80.00 | | 80.30 | 81.10 | 79.40 |
|                  | Paternal Harsh parenting | Maternal harsh parenting | Pooled harsh parenting |
|------------------|--------------------------|--------------------------|------------------------|
|                  | Total                    | Without harsh parenting | With harsh parenting   | Total                    | Without harsh parenting | With harsh parenting   |
|                  | n                        |                          |                        | Total                    | Without harsh parenting | With harsh parenting   |
| Maternal drinking during pregnancy (% never drunk during pregnancy) | 1710 | 1113 | 597 | 1881 | 1208 | 673 | 1916 | 958 | 958 |
| Depressive symptoms mothers BSI | 0.09±0.21 | 0.08±0.20 | 0.11±0.24 | * | 0.09±0.22 | 0.08±0.21 | 0.11±0.23 | ** | 0.09±0.22 | 0.08±0.21 | 0.11±0.23 | ** |
| Depressive symptoms fathers BSI | 0.08±0.20 | 0.05±0.17 | 0.10±0.24 | ** | 0.07±0.20 | 0.07±0.18 | 0.09±0.23 | 0.07±0.20 | 0.05±0.18 | 0.09±0.22 | ** |
| Anxiety symptoms mothers BSI | 0.15±0.25 | 0.14±0.24 | 0.18±0.27 | ** | 0.16±0.26 | 0.14±0.23 | 0.19±0.30 | ** | 0.16±0.26 | 0.13±0.23 | 0.18±0.28 | ** |
| Anxiety symptoms fathers BSI | 0.15±0.24 | 0.13±0.22 | 0.18±0.26 | ** | 0.15±0.24 | 0.14±0.23 | 0.16±0.26 | 0.24±2.50 | 0.12±0.22 | 0.17±0.25 | ** |
| Harsh parenting mother (sumscore 5 items) | 0.73±1.28 | 0.00±0.00 | 2.03±1.40 | ** | 0.73±1.28 | 0.00±0.00 | 1.44±1.50 | ** |
| Harsh parenting father (sumscore 5 items) | 0.70±1.22 | 0.00±0.00 | 2.00±1.28 | ** | 0.70±1.22 | 0.00±0.00 | 1.36±1.41 | ** |

Sample characteristics are presented for paternal harsh parenting, maternal harsh parenting and pooled harsh parenting (total group, group without harsh parenting, and group with harsh parenting). The number of missings varies across variables. Unless otherwise specified, numbers in the table represent mean ± SD. All percentages are valid percentages. Using t-tests and chi-square tests, all variables were compared between the groups with and without harsh parenting. * = p < .05, ** = p < .01
MEASURES

Genetic data

Genotyping of GWAS data

DNA was collected from cord blood samples at birth. All samples were genotyped using Illumina Infinium II HumanHap610 Quad Arrays (Illumina, San Diego, CA, USA) following standard manufacturer’s protocols. Some children (320), for whom DNA was missing, provided blood samples at the age of 6, and these samples were genotyped using the Human 660 Quad Arrays of Illumina. Intensity files were analyzed using the Beadstudio Genotyping Module software v.3.2.32 and genotype calling based on default cluster files. Samples displaying call rates below 95% and mismatch between called and phenotypic gender were excluded. In addition, individuals identified as genetic outliers by the identity-by-state (IBS) clustering analysis (>4 standard deviations away from the HapMap CEU population mean) were considered of other ancestry and were excluded from the analyses. Finally, genotyped SNPs with minor allele frequency (MAF) < 0.01, SNP Call Rate <0.98 and Hardy-Weinberg Equilibrium (HWE) p-value <1x10^{-6} were filtered. SNPs that showed differential missingness between arrays were excluded (for additional information on GWAS data in the Generation R study, see Medina-Gomez et al., 2015). GWAS data were available for 2,830 Caucasian children. The number of genotyped SNPs (build 37) was 489,878.

Gene-set selection and SNP assignment

Based on a wide literature exploring associations between dopamine genes and behavior (e.g. Gardner, Bertranpetit, & Comas, 2008; Gorwood et al., 2012; Nemoda, Szekely, & Sasvari-Szekely, 2011; Talkowski, Bamne, Mansour, & Nimgaonkar, 2007), we selected a gene-set of autosomal dopaminergic genes: DRD1, DRD2, DRD3, DRD4, DRD5, COMT, DBH, DDC, DAT, TH, DARPP-32, VMAT1, VMAT2, ANKK1 (Table 2). All genotyped SNPs that passed quality control were mapped to genes on the basis of NCBI (National Center for Biotechnology Information) human assembly build 37 (Genome Reference Consortium GRCh37, dbSNP release 131, UCSC assembly hg 19). The SNP-to-gene annotation was performed using the software package JAG (Lips et al., 2015). No flanker regions were specified, i.e. the transcription start site and transcription end site were used as boundaries of the genes. The Illumina Array did not yield genotyped SNPs on the DRD4, DRD5 and DARPP-32 genes. The DRD4 48 bp variable number of tandem repeats polymorphism (VNTR), however, was further individually genotyped (see Supporting information S1), and this polymorphism was (manually) added to the GWAS data. The final gene-set consisted of 12 genes including 151 genetic variants. Minor allele frequencies (MAFs) of these genetic variants are presented in Supporting information S2.
Table 2 | Dopamine genes in the gene-set.

| Gene   | Alias      | Entrez Gene ID | Gene Name                                      | Chromosome number | Transcription start site (bp) | Transcription end site (bp) | Strand | Number of SNPs available in GWAS data |
|--------|------------|----------------|-----------------------------------------------|-------------------|------------------------------|----------------------------|--------|--------------------------------------|
| DRD1   |            | 1812           | Dopamine receptor D1                          | 5                 | 174867675                   | 174871163                  | -      | 2                                    |
| DRD2   |            | 1813           | Dopamine receptor D2                          | 11                | 113280317                   | 113346001                  | -      | 15                                   |
| DRD3   |            | 1814           | Dopamine receptor D3                          | 3                 | 113847557                   | 113897899                  | -      | 12                                   |
| DRD4   |            | 1815           | Dopamine receptor D4                          | 11                | 637305                      | 640706                     | +      | 0 (but 1 VNTR manually added)        |
| DRD5   |            | 1816           | Dopamine receptor D5                          | 4                 | 9783258                     | 9785633                    | +      | 0                                    |
| DAT    | SLC6A3, DAT1 | 6531           | Dopamine transporter                          | 5                 | 1392905                     | 1445543                    | -      | 13                                   |
| VMAT1  | SLC18A1    | 6570           | Vesicular monoamine transporter 1 (VMAT1)     | 8                 | 20002366                    | 20040717                   | -      | 20                                   |
| VMAT2  | SLC18A2    | 6571           | Vesicular monoamine transporter 2 (VMAT2)     | 10                | 119000584                   | 119038941                  | +      | 18                                   |
| TH     |            | 7054           | Tyrosine hydroxylase                          | 11                | 2185159                     | 2193035                    | -      | 2                                    |
| DDC    |            | 1644           | Dopa decarboxylase                            | 7                 | 50526134                    | 50633154                   | -      | 32                                   |
| COMT   |            | 1312           | Catechol-O-methyltransferase                  | 22                | 19929263                    | 19957498                   | +      | 14                                   |
| DARPP-32 | PP1R1B   | 84152          | Dopamine- and cAMP-regulated phosphoprotein, 32 kDa | 17 | 37783177 | 37792878 | + | 0 |
| DBH    |            | 1621           | Dopamine beta hydroxylase                     | 9                 | 136501485                   | 136524466                  | +      | 17                                   |
| ANKK1  |            | 255239         | Ankyrin repeat and kinase domain containing 1 | 11                | 113258513                   | 113271140                  | +      | 5                                    |

Note. Gene locations are based on NCBI (National Center for Biotechnology Information) human assembly build 37 (Genome Reference Consortium GRCh37, dbSNP release 131, UCSC assembly hg 19)
Harsh Parenting
Harsh parenting was assessed with a parental self-report questionnaire when the child was 3 years old (36 ± 1.2 months). In a previous study of the same cohort (Jansen et al., 2012), six items of the Parent-Child Conflict Tactics scale (Straus, Hamby, Finkelhor, Moore, & Runyan, 1998) were selected based on factor analysis to constitute a harsh discipline scale. The scale consisted of the following items: ‘shook my child,’ ‘shouted or screamed angrily at my child,’ ‘called my child names,’ ‘threatened to give a slap, but I didn’t do it,’ ‘angrily pinched my child’s arm,’ ‘called my child stupid, lazy, or something like that’. Confirmatory factor analyses indicated good fit for the harsh parenting factor in both mothers and fathers (for additional details, see Jansen et al., 2012). Parents rated their use of discipline types during the past two weeks on a 6-point scale ranging from ‘never’ to ‘five times or more’. The prevalence of the item ‘shouted or screamed angrily at my child’ was very high; therefore, this item was excluded from the sum score. Since JAG does not provide an option to include gene-environment interaction effects, we stratified our sample in two groups, one with and one without harsh parenting experiences, and ran the JAG tool for gene-based and gene-set analyses separately in these groups. This was done for both maternal and paternal harsh parenting. For the pooled maternal and paternal harsh parenting variable, harsh parenting was present if at least one parent reported one or more harsh parenting behaviors. The group sizes of the group without versus with harsh parenting were 1,113 vs. 597 for paternal harsh parenting, 1,208 vs. 673 for maternal harsh parenting, and 958 vs. 958 for pooled paternal and maternal harsh parenting. The association between paternal and maternal harsh parenting dichotomies is provided in the Supporting information S3.

Externalizing behavior
Externalizing behavior of the child was assessed by parental report when the child was five years of age (71 ± 3.7 months). The primary caregiver filled out a Dutch version of the Child Behavior Checklist (CBCL/1.5-5), a widely used questionnaire with 99 items concerning the child’s behavior in the previous two months. The CBCL for ages 1.5-5 was used for all children, because the majority (66 %) of the children were younger than 6 years old at the time of assessment. We considered it important to use only one version of the CBCL to enhance comparability across all children. The 1.5-5 version was chosen because the version for 6- to 18-year-olds contains questions that are less applicable to 5-year-olds, for example questions concerning smoking tobacco. Each item is scored as 0 = not true, 1 = somewhat or sometimes true, and 2 = very true or often true. Seven empirically-based syndrome scales were identified: emotionally reactive, anxious/depressed, somatic complaints, withdrawn, sleep problems, attention problems, and aggressive behavior (Achenbach & Rescorla, 2000). The broadband scale “Externalizing” comprises item scores on the subscales aggressive behavior and attention problems scales. Higher scores indicated more problems. Internal consistency of the CBCL was $\alpha = .90$. CBCL scores on externalizing behavior were missing for between 113 and 137 children (<10%). Missing values on externalizing behavior were imputed by regression imputation using CBCL scores on externalizing behavior collected at 18 and 36 months as predictors.
Covariates

The following covariates were included in all analyses: age of the child when the CBCL was completed, gender of the child and four ancestry-informative principal components. These principal components were obtained by principal component analyses of the GWAS data of the Caucasian children (n = 2,830) and were included to account for population-specific variations in alleles distribution of the SNPs to control for effects due to population stratification.

Analyses

Prior to the main analyses, we ran association tests for all individual SNPs and harsh parenting using PLINK version v1.07 (http://pngu.mgh.harvard.edu/purcell/plink/, Purcell et al., 2007) to check whether allele frequencies of the SNPs differed between the groups with and without harsh parenting experiences, in order to rule out gene-environment correlations.

Gene-set analyses

The gene-set analyses were conducted using JAG (Lips et al., 2015). We ran separate analyses for the total group and for the two groups with and without harsh parenting, for paternal, maternal and pooled harsh parenting. Self-contained tests were conducted to test the null hypothesis that the dopamine gene-set was not associated with externalizing behavior. For every self-contained test, the test statistic was computed using the sum of the -log10 of the p-values for the single SNP associations for all the SNPs in the gene-set. Empirical p-values were obtained with at least 9,000 permutations of the phenotype, keeping linkage disequilibrium and number of SNPs and genes constant. The empirical p-value was calculated as the proportion of test statistics that were equal to or higher than the test statistic of the original data. For the gene-set analyses, an empirical p-value <.05 was considered significant.

In case of a significant empirical p-value in the test for the dopamine gene-set, a competitive test was conducted. Competitive tests show whether the gene-set of interest is more strongly associated with externalizing behavior than randomly generated gene-sets matched for the same number of genes. Competitive tests for polygenic traits like externalizing behavior are essential, because associations with a gene-set that emerge in the self-contained test can be due to the polygenic background. 150 random control gene-sets matching the original dopamine gene-set on number of genes were generated. Self-contained tests similar to those described above were conducted for each of the control gene-sets. Finally, a competitive p-value was calculated as the proportion of self-contained p-values for the random gene-sets that were lower than the empirical self-contained p-value for the dopamine-set. Competitive p-values <0.05 were considered to be significant.

Testing the interaction of harsh parenting and the dopamine gene-set. JAG currently does not allow for a direct test of the interaction between harsh parenting and genetic variance in the dopamine gene set. To test this interaction indirectly, analyses were run on the stratified samples (with and without harsh parenting). Subsequently, the effect sizes for the groups were statistically compared taking sample sizes into account.
Gene-based analyses

On the level of individual dopamine genes, self-contained tests were conducted in a similar way as the gene-set analyses, testing the null hypothesis that each individual dopamine gene is not associated with externalizing behavior. Since the gene-based self-contained tests were conducted for 12 genes, the significance threshold was adjusted for multiple testing. Based on the assumption that genetic distance and the interdependence of genes are correlated with the physical distance between genes (Morton, 1955; Pritchard & Przeworski, 2001), we calculated the number of independent genes that were separated by at least 5Mb. According to this calculation the dopamine gene-set consisted of six independent genes, resulting in a significance threshold of $0.05/6 = 0.0083$.

SNP level analyses

In addition to our main analyses on gene-set and gene levels, we ran individual linear regression analyses for all SNPs using PLINK. The previously mentioned covariates, main effects of SNP genotype (additive effect model) and harsh parenting, and the interaction between the SNP and harsh parenting were included in the regression models. Thresholds for significance were adjusted for multiple testing based on the independent number of SNPs, determined by linkage disequilibrium (LD) based pruning in PLINK (-indep-pairwise 50 5 0.20) of the 151 dopamine SNPs for the total groups of children, for paternal, maternal, and pooled harsh parenting. LD based pruning resulted in 49 independent SNPs, therefore significance thresholds for the analyses at the SNP level were set to $\alpha = 0.05/49 = 0.0010$.

Lastly, we examined whether the gene-set results were specific for harsh parenting or were accounted for by parental educational level, which differed for the maternal harsh parenting groups, see Table 1.

RESULTS

Allele frequencies of the individual SNPs did not differ between groups with and without harsh parenting experiences, indicating that the SNPs were not associated with harsh parenting, see Supporting information S4. This provides evidence against gene-environment correlations of child dopamine SNPs and harsh parenting.

Gene-set analyses

Self-contained tests

The $p$-values for the self-contained gene-set analyses are presented in Table 3 (Manhattan plots and tables presenting single variant associations with externalizing behavior, used for the calculation of the test-statistic, are presented in Supporting information S5 and S6. For plots of the test-statistic distributions of the original data and the permutations see S7). For paternal harsh parenting, variation in the dopamine gene-set and externalizing behavior were significantly associated for the children in the group without harsh parenting experiences ($p = .03$). For maternal harsh parenting and pooled harsh parenting, this association approached significance in the group without harsh parenting ($p = .05$
The association between dopamine gene-set variation and externalizing behavior was far from significant in the group with harsh parenting (*p* = .92-.97).

Finally, the self-contained tests for the total group (groups with and without harsh parenting together) were not significant either (*p* = .17-21). These results seem to indicate that variation in the dopamine gene-set significantly predicts variance in children's externalizing behavior for children who did not experience harsh parenting, but not for children with harsh parenting experiences. However, given the polygenic nature of the trait, a significant self-contained *p*-value may simply reflect this polygenic nature. Therefore competitive tests are run.

**Competitive tests**

Since the self-contained *p*-value for the gene-set analysis in the group without harsh parenting for paternal harsh parenting was significant, a competitive test was conducted. The self-contained *p*-value for the dopamine gene-set (.03) was compared to the self-contained *p*-values for 150 random gene sets. The competitive *p*-value was .03, indicating that genetic variation in the dopamine gene-set was more strongly related to externalizing behavior than genetic variation in random gene-sets.

**Interaction of harsh parenting and the dopamine gene-set**

We compared the effect sizes based on the *p*-values for the self-contained tests of the association between gene-set variance and externalizing behavior in the groups with and without harsh parenting, as an indirect test of the interaction effect. The results of these tests were *Q*<sub>contrast</sub>(1) = 0.94, *p* = .19; *Q*<sub>contrast</sub>(1) = 0.88, *p* = .28; and *Q*<sub>contrast</sub>(1) = 0.70, *p* = .41 for paternal, maternal and pooled harsh parenting respectively – indicating that the effect sizes for the association between externalizing behavior and variance in the dopamine gene-set in the groups with and without harsh parenting were not significantly different, and the interaction was thus not statistically significant.

**Gene-based analyses**

In the gene-based analyses, associations of the individual dopamine genes with children's externalizing behavior were tested. In Table 3, the *p*-values for these self-contained tests are presented; *p*-values below .0083 are considered significant. For maternal harsh parenting, genetic variation in the DRD2 gene was significantly associated with externalizing behavior in the group without harsh parenting (*p* = .0040) but not in the group with harsh parenting or the total group. For paternal harsh parenting, variance in the VMAT1 gene (*p* = .0068) was significantly associated with externalizing behavior, again only in the group without harsh parenting. Finally, for pooled harsh parenting, there were no significant associations between dopamine gene variance and externalizing behavior.

**Interaction of harsh parenting and dopamine genes**

We found two significant associations between externalizing behavior and dopamine genes in the group without harsh parenting experiences. For paternal harsh parenting this association was significant for the VMAT1 gene, for maternal harsh parenting the association was significant for the DRD2 gene. We compared the effect sizes based on the *p*-values for the groups with and without harsh
| Group                     | Paternal harsh parenting | Maternal harsh parenting | Pooled harsh parenting |
|---------------------------|--------------------------|--------------------------|------------------------|
|                           | Total | Without harsh parenting | With harsh parenting | Total | Without harsh parenting | With harsh parenting | Total | Without harsh parenting | With harsh parenting |
|                           | 1710  | 1113                      | 597                    | 1881  | 1208                      | 673                    | 1916  | 958                      | 958                    |
| Gene-set analyses         | .21   | .03 *A                    | .97                    | .19   | .05                       | .95                    | .17   | .08                       | .92                    |
| Gene-based analyses:      |       |                            |                        |       |                            |                        |       |                            |                        |
| COMT                      | 1312  | .10                       | .37                    | .37   | .21                       | .53                    | .23   | .21                       | .67                    | .34                    |
| DBH                       | 1621  | .96                       | .92                    | .72   | .92                       | .99                    | .79   | .92                       | .98                    | .70                    |
| DDC                       | 1644  | .33                       | .11                    | .98   | .25                       | .16                    | .88   | .27                       | .06                    | .85                    |
| DRD1                      | 1812  | .51                       | .10                    | .97   | .33                       | .24                    | .12   | .39                       | .16                    | .57                    |
| DRD2                      | 1813  | .09                       | .33                    | .32   | .14                       | .00 *C                 | .92   | .10                       | .18                    | .42                    |
| DRD3                      | 1814  | .30                       | .24                    | .86   | .18                       | .50                    | .25   | .14                       | .38                    | .38                    |
| DRD4                      | 1815  | .83                       | 1.00                   | .83   | .51                       | .98                    | .23   | .69                       | .59                    | .36                    |
| DRD5                      | 1816  | X                         | X                      | X     | X                         | X                      | X     | X                         | X                      |                        |
| DAT                       | 6531  | .03                       | .11                    | .46   | .03                       | .01                    | .38   | .02                       | .09                    | .30                    |
| VMAT1                     | 6570  | .43                       | .01 *B                 | .35   | .32                       | .15                    | .55   | .42                       | .11                    | .78                    |
| VMAT2                     | 6571  | .92                       | .72                    | .99   | .83                       | .74                    | .98   | .85                       | .42                    | .88                    |
| TH                        | 7054  | .42                       | .01                    | .59   | .35                       | .14                    | .86   | .38                       | .01                    | .72                    |
| DARPP-32                  | 84152 | X                         | X                      | X     | X                         | X                      | X     | X                         | X                      |                        |
| ANKK1                     | 255239| .45                       | .68                    | .40   | .80                       | .52                    | .89   | .70                       | .97                    | .58                    |

Note: For the gene-set, p-values < .05 are considered significant, for the genes, p-values < .0083 are considered significant. A: p = .0250, B: p = .0068, C: p = .0040
GXE of dopamine gene-set and harsh parenting

parenting, to test the interactions indirectly. The results of these tests were $Q_{\text{contrast}} (1) = 0.34, p = .10$ for the VMAT1 gene and $Q_{\text{contrast}} (1) = 0.91, p = .08$ for the DRD2 gene indicating that the interactions between harsh parenting and variance in the dopamine genes were not statistically significant.

SNP level analyses
Results of the interaction tests at SNP level are presented in more detail in the Supporting information, (Manhattan plots S5, table S8). At the level of individual SNPs, significant interaction effects were found between paternal harsh parenting and two SNPs: rs1497023 ($\beta = 0.92, p = .0008$) and rs4922132 ($\beta = 0.92, p = .001$). Both SNPs are located on the VMAT1 gene. For maternal and pooled harsh parenting, no significant GxE interactions were found at the SNP level.

Finally, we tested whether the results found for harsh parenting were accounted for by the lower parental educational level in the harsh parenting group (see Supporting information S9). Results showed that the pattern of results could not be ascribed to differences in educational level.

DISCUSSION

In this longitudinal cohort study, we investigated the interplay of harsh parenting and genetic variation across a set of functionally related dopamine genes, in association with children's externalizing behavior. The pattern of effects suggests that the association between genetic variance in the dopamine gene-set and externalizing behavior depends on harsh parenting experiences: the association between genetic variance in the dopamine gene-set and externalizing behavior was present for children who did not experience harsh parenting and absent for children who did experience harsh parenting. Results were similar for paternal, maternal and pooled harsh parenting. However, the indirect interaction test indicated that the difference between the effects for the two harsh parenting groups did not reach significance. Thus we urge caution in interpreting these results until further replication.

The stratified analyses suggest that unfavorable environmental factors do play a role in explaining the variance in child behavioral outcomes, and that in high-risk conditions environmental factors may overrule genetic factors. In contrast, low-risk environments present less environmental ‘pressure’ on the behavioral phenotype, and therefore greater contributions of underlying genetic factors can be expected. In the present study, this suggests that the variance in children's externalizing behavior is not explained by underlying genetic variation in the dopamine gene-set in children who experience harsh parenting. It is important to note that at this level of analysis, we cannot infer the direction of effects between genotypes and externalizing behavior. In other words, our analyses do not indicate which genetic variants (alleles) confer greater or lesser risk for externalizing behavior, as they aggregate a combined statistical effect for all SNPs in the gene-set. This might be possible at the level of individual SNPs -- but here we should be cautious due to a lack of conclusive functional knowledge for most SNPs. We can however conclude that the genetic variance in a dopamine gene-set in children in a low-risk environment explains a significant proportion of the variance in their externalizing behavior. Allele
frequencies of the individual genetic variants did not differ between the groups with and without harsh parenting experiences, indicating that there were no evocative gene-environment correlations between child dopamine SNPs and harsh parenting.

The analyses for single dopamine genes (Table 3) were consistent with the gene-set findings; significant associations between variance in several single genes were found for the group without harsh parenting. For paternal harsh parenting this association was significant for the VMAT1 gene, for maternal harsh parenting the association was significant for the DRD2 gene. No significant associations between single dopamine genes and externalizing behavior were found for the groups with harsh parenting experiences. At the level of single SNPs, significant interaction effects were found between paternal harsh parenting and two SNPs in VMAT1. In the gene-set analyses, genetic variation across a large number of SNPs over multiple dopamine genes is aggregated. As a result, SNPs with individual small effects can have a significant combined effect at the level of whole genes or gene-sets (Winham & Biernacka, 2013). This is supported by our results, which are suggestive of GxE effects at the gene-set level, while at the SNP level, significant interaction effects were found between paternal harsh parenting and only two single SNPs (after correction for multiple testing). Thus our findings indicate the additional value of including whole gene-sets as alternative or complement to candidate genes and single-SNP analyses that might not reach significance when tested on a one-by-one basis.

The groups with and without harsh parenting differed on a number of variables, including psychopathology symptoms of the parents (anxiety and depression), age of the mother, maternal and paternal education level, child externalizing behavior, gender and birth weight of the child and parity. This could mean that the harsh parenting variable might not have been the only environmental influence. Overall, the group without harsh parenting might be considered a ‘lower-risk group’ while the group with harsh parenting experiences could be seen as a group exposed to relatively higher risk. Nonetheless, our sample is a population sample and therefore not representative of a true high-risk group. Our focus on harsh parenting as an environmental factor stemmed from the assumption that parenting is a more proximal variable that could mediate the effects of more distal environmental influences such as SES on child behavioral outcomes. Additional analyses showed that the results were not determined by differences in SES. However, in future work the gene-set component of the GxE equation might be complemented by an environment-set component to account for the potential multi-factorial influences of the environment. This approach parallels the search for GWAS x EWAS interactions applied to type 2 diabetes mellitus (Patel, Chen, Kodama, Ioannidis, & Butte, 2013) which throws a larger net on the potential pertinent environment and at the same time compensates for multiple testing involved in scrutinizing gene-set x environment-set interactions.

A few issues are important to consider when interpreting our results. First, significant self-contained \(p\)-values for polygenic traits could be caused by this polygenic background. Therefore, competitive tests are needed to see whether the gene-set of interest is more strongly associated with the outcome than randomly generated matched gene-sets. We conducted competitive tests whenever the self-contained test reached significance. This was only the case for paternal harsh parenting, in the group without harsh parenting experiences. The competitive test also proved to be significant for this group, indicating that the association between externalizing behavior and the dopamine gene-set was stronger than it was for 150 random gene-sets. In the group with paternal harsh parenting
experiences, the self-contained test was far from significant. Similarly, for maternal and pooled harsh parenting, none of the self-contained tests reached significance and thus competitive tests were not conducted. Although these results do not allow firm conclusions, it should be noted that the directions of the effects were similar to those of paternal harsh parenting.

A second issue to keep in mind when interpreting our results is multiple testing. We performed multiple tests at the levels of single genes and SNPs in groups based on three related harsh parenting variables. Since the samples in our analyses were largely overlapping (see Supporting information, S3), and the genes and SNPs were not completely independent, we judged it inappropriate to use a highly stringent Bonferroni correction in this first exploratory study of its kind. More conservative approaches might be taken in future replication efforts.

A third issue to consider are sample size differences between the groups with and without harsh parenting. The lack of an association in the groups with harsh parenting experiences might be alternatively explained by smaller sample size.

This study has a number of notable strengths. First, our study sample is a large ethnically homogenous subsample from a longitudinal population-based cohort study. Secondly, we had available data on harsh parenting in both fathers and mothers from the same families. Our study has some limitations as well. Harsh parenting was assessed by self-report, which might be open to bias (e.g. Morsbach & Prinz, 2006). Valid assessment of the environment is shown to be critical for GxE results. Therefore, it is preferable to assess environmental factors by observational measures or interviews in future studies (McGuffin, Alsabban, & Uher, 2011; Uher & McGuffin, 2010). Another limitation is that the specifications of the JAG tool implied restriction to a dichotomous rather than continuous environmental measure. The harsh parenting group thus represents a group with harsh parenting ranging from moderate to extreme. Note, however, that this may partly tackle the limitation of self-report: Parents who did not completely deny but underreported their level of harsh parenting were classified in the group with harsh parenting. Finally, even though our sample was of considerable size for gene-set analyses, larger samples might improve power, particularly when splitting the sample to investigate GxE effects. Replication of our exploratory findings is needed.

In sum, we found suggestive evidence for interplay between harsh parenting and dopamine genetic variance, with an association between dopamine genes and externalizing behavior for children who did not experience harsh parenting. Harsh parenting may overrule the role of genetic factors in externalizing behavior. The approach presented here, involving gene and gene-set analyses, offers a useful and relatively unexplored alternative to studies focusing on single candidate polymorphisms in the search for genetic variation underlying complex behavior.

Additional acknowledgements

The generation and management of genotype data for the Generation R Study were performed at the Genetic Laboratory of the Department of Internal Medicine at Erasmus Medical Center. We thank M. Jhamai, M. Ganesh, P. Arp, M. Verkerk, L. Herrera and M. Peters for their help in creating, managing and performing quality control for the genetic database. The JAG software is developed at the Department of Complex Trait Genetics, (Center for Neurogenomics and Cognitive Research, Neuroscience Campus Amsterdam, VU University Amsterdam). We thank Danielle Posthuma and Anke Hammerschlag for their contributions.
REFERENCES

Achenbach, T. M., & Rescorla, L. A. (2000). *Manual for the ASEBA Preschool forms & profiles*. Burlington, VT: University of Vermont, Research Center for Children, Youth, & Families.

Aschard, H., Lutz, S., Maus, B., Duell, E. J., Fingerlin, T. E., Chatterjee N., ... Van Steen, K. (2012). Challenges and opportunities in genome-wide environmental interaction (GWEI) studies. *Human Genetics, 13*, 1591-1613.

Bakermans-Kranenburg, M. J., & Van IJzendoorn, M. H. (2011). Differential susceptibility to rearing environment depending on dopamine-related genes: New evidence and a meta-analysis. *Development and Psychopathology, 23*, 39–52.

Brodkin, E. S., Goforth S. A., Keene A. H., Fossella J. A., & Silver L. M. (2002). Identification of quantitative trait loci that affect aggressive behavior in mice. *The Journal of Neuroscience, 22*, 1165-1170.

Campbell, S. B., Shaw, D. S., & Gilliom, M. (2000). Early externalizing behavior problems: Toddlers and preschoolers at risk for later maladjustment. *Development and Psychopathology, 12*, 467–488.

Campbell, S. B., Spieker, S., Burchinal, M., Poe, M. D., & The NICHD Early Child Care Research Network. (2006). Trajectories of aggression from toddlerhood to age 9 predict academic and social functioning through age 12. *Journal of Child Psychology and Psychiatry, 47*, 791–800.

Dick, D. M., Agrawal, A., Keller, M. C., Adkins, A., Aliev, F., Monroe, S., ... Sher, K. J. (2015). Candidate gene-environment interaction research: Reflections and recommendations. *Perspectives on Psychological Science, 10*, 37-59.

Dick, D. M., Li, T. K., Edenberg, H. J., Hesselbrock, V., Kramer, J., Kuuperman, S., ... Foroud, T. (2004). A genome-wide screen for genes influencing conduct disorder. *Molecular Psychiatry, 9*, 81-86.

Duncan, L. E., & Keller, M. C. (2011). A critical review of the first 10 years of candidate gene-by-environment interaction research in psychiatry. *American Journal of Psychiatry, 168*, 1041-1049.

Ellis, B. J., Boyce, W. T., Belsky, J., Bakermans-Kranenburg, M. J., & Van IJzendoorn, M. H. (2011). Differential susceptibility to the environment: An evolutionary–neurodevelopmental theory. *Development and Psychopathology, 23*, 7–28.

Fridley, B. L., & Biernacka, J. M. (2011). Gene set analysis of SNP data: Benefits, challenges, and future directions. *European Journal of Human Genetics, 19*, 837-843.

Gardner, M., Bertranpetit, J., & Comas, D. (2008). Worldwide genetic variation in dopamine and serotonin pathway genes: Implications for association studies. *American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 147B*, 1070-1075.

Gorwood, P., Le Strat, Y., Ramoz, N., Dubertret, C., Moalic, J., & Simonneau, M. (2012). Genetics of dopamine receptors and drug addiction. *Human Genetics, 131*, 803-822.

Haberstick, B. C., Schmitz, S., Young, S. E., & Hewitt, J. K. (2005). Contributions of genes and environments to stability and change in externalizing and internalizing problems during elementary and middle school. *Behavior Genetics, 35*, 381-396.

Hirschhorn, J. N., & Daly, M. J. (2005). Genome-wide association studies for common diseases and complex traits. *Nature Reviews Genetics, 6*, 95-108.

Hudziak, J. J., Van Beijsterveldt, C. E. M., Bartels, M., Rietveld, M. J. H., Rettew, D. C., Derks, E. M., & Boomsma, D. I. (2003). Individual differences in aggression: Genetic analyses by age, gender, and informant in 3-, 7-, and 10-year-old Dutch twins. *Behavior Genetics, 33*, 575-589.

Jaddoe, V. W. V., Van Duijn, C. M., Franco, O. H., Van der Heijden, A. J., Van IJzendoorn, M. H., De Jongste, J. C., ... Hofman, A. (2012). The Generation R Study: Design and cohort update 2012. *European Journal of Epidemiology, 27*, 739-756.

Jansen, P. W., Raat, H., Mackenbach, J. P., Hofman, A., Jaddoe, V. W. V., Bakermans-Kranenburg M. J., ... Tiemeier H. (2012). Early determinants of maternal and paternal harsh discipline: The Generation R study. *Family Relations, 61*, 253-270.

Keltikangas-Järvinen, L., & Salo, J. (2009). Dopamine and serotonin systems modify environmental effects on human behavior: A review. *Scandinavian Journal of Psychology, 50*, 574-582.
GXE of dopamine gene-set and harsh parenting

Khoury, M. J. & Wacholder, S. (2009). Invited Commentary: From Genome-Wide Association Studies to Gene-Environment-Wide Interaction Studies-Challenges and Opportunities. *American Journal of Epidemiology, 169,* 227-230.

Lips, E. S., Cornelisse, L. N., Toonen, R. F., Min, J. L., Hultman, C. M., The International Schizophrenia Consortium, … Posthuma, D. (2012). Functional gene group analysis identifies synaptic gene groups as risk factor for schizophrenia. *Molecular Psychiatry,* 17, 996-1006.

Lips, E. S., Kooyma M., De Leeuw C., & Posthuma D. (2015). JAG: A computational tool to evaluate the role of gene-sets in complex traits. *Genes,* 6, 238-251.

Manuck, S. B., & McCaffery, J. M. (2014). Gene-environment interaction. *Annual Review of Psychology,* 65, 41-70.

McGuffin, P., Alsabban, S., & Uher, R. (2011). The truth about genetic variation in the serotonin transporter gene and response to stress and medication. *The British Journal of Psychiatry,* 198, 424-427.

Medina-Gomez, C., Felix, J. F., Estrada, K., Peters, M. J., Herrera, L., Kruithof, C. J., … Rivadeneira, F. (2015). Challenges in conducting genome-wide association studies in highly admixed multi-ethnic populations: The Generation R Study. *European Journal of Epidemiology,* 30, 317-330.

Miner, J. L., & Clarke-Stewart, K. A. (2008). Trajectories of externalizing behavior from age 2 to age 9: Relations with gender, temperament, ethnicity, parenting, and rater. *Developmental Psychology,* 44, 771-786.

Moffitt, T. E., Caspi, A., & Rutter, M. (2005). Strategy for investigating interactions between measured genes and measured environments. *Archives of General Psychiatry,* 62, 473-481.

Morsbach, S. K. & Prinz, R. J. (2006). Understanding and improving the validity of self-report of parenting. *Clinical Child and Family Psychology Review,* 9, 1-21.

Morton, N. E. (1955). Sequential tests for the detection of linkage. *American Journal of Human Genetics,* 7, 277-318.

Nemoda Z., Szekely, A., & Sasvari-Szekely, M. (2011). Psychopathological aspects of dopaminergic gene polymorphisms in adolescence and young adulthood. *Neuroscience & Biobehavioral Reviews,* 35, 1665-1686.

Oades, R. D., Lasky-Su, J., Christiansen, H., Faraone, S. V., Sonuga-Barke, E. J., Banaschewski, T., … Asherson, P. (2008). The influence of serotonin- and other genes on impulsive behavioral aggression and cognitive impulsivity in children with attention-deficit/hyperactivity disorder (ADHD): Findings from a family-based association test (FBAT) analysis. *Behavioral and Brain Functions,* 4, 48.

Patel, C. J., Chen, R., Kodama, K., Ioannidis, J. P. A., & Butte, A. J. (2013). Systematic identification of interaction effects between genome- and environment-wide associations in type 2 diabetes mellitus. *Human Genetics,* 132, 495-508.

Pritchard, J. K. & Przeworski, M. (2001). Linkage disequilibrium in humans: Models and data. *The American Journal of Human Genetics,* 69, 1-14.

Purcell S., Neale B., Todd-Brown K, Thomas L., Ferreira M. A. R., Bender D., … Sham P. C. (2007). PLINK: A toolset for whole-genome association and population-based linkage analyses. *The American Journal of Human Genetics,* 81, 559-575.

Reef, J., Diamantopoulos S., Van Meurs, I., Verhulst, F. C., & Van der Ende, J. (2011). Developmental trajectories of child to adolescent externalizing behavior and adult DSM-IV disorder: Results of a 24-year longitudinal study. *Social Psychiatry and Psychiatric Epidemiology,* 46, 1233-1241.

Reif, A., & Lesch, K. P. (2003). Toward a molecular architecture of personality. *Behavioural Brain Research,* 139, 1-20.

Robbins, T.W., & Everitt, B.J. (1999). Motivation and reward. In M.J. Zigmond, F.E. Bloom, S.C. Landis, J.L. Roberts, & L.R Squire (Eds.), *Fundamental neuroscience* (pp. 1246–1260). San Diego, CA: Academic Press..

Rutter, M. (2006). *Genes and behavior. Nature–nurture interplay explained.* Oxford: Blackwell.

Saudino, K. J., Carter, A. S., Purper-Ouakil, D., & Gorwood, P. (2008). The etiology of behavioral problems and competencies in very young twins. *Journal of Abnormal Psychology,* 117, 48–62.
Shanahan, M. J. and Hofer, S. M. (2005). Social context in gene-environment interactions: Retrospect and prospect. The Journals of Gerontology Series B: Psychological Sciences and Social Sciences, 60, 65-76.

Straus, M. A., Hamby, S. L., Finkelhor, D., Moore, D. W., & Runyan, D. (1998). Identification of child maltreatment with the Parent-Child Conflict Tactics Scales: Development and psychometric data for a national sample of American parents. Child Abuse & Neglect, 22, 249 – 270.

Sullivan, P. F. (2007). Spurious genetic associations. Biological Psychiatry, 61, 1121-1126.

Tabor, H. K., Risch, N. J., & Myers, R. M. (2002). Candidate-gene approaches for studying complex genetic traits: Practical considerations. Nature Reviews Genetics, 3, 391-397.

Takahashi, A., Quadros, I. M., De Almeida, R. M. M., & Miczek, K. A. (2012). Behavioral and pharmacogenetics of aggressive behavior. Current Topics in Behavioral Neurosciences, 12, 73-128.

Talkowski, M. E., Bamne, M., Mansour, H., & Nimgaonkar, V. L. (2007). Dopamine genes and schizophrenia: Case closed or evidence pending? Schizophrenia Bulletin, 33, 1071-1081.

Tielbeek, J. J., Medland S. E., Benyamin B., Byrne E. M., Heath A. C., Madden P. A. F., … Verweij K. J. H. (2012). Unraveling the genetic etiology of adult antisocial behavior: A genome-wide association study. PLoS One, 7, e45086.

Tiemeier, H., Velders, F. P., Szekely, E., Roza, S. J. Dieleman, G., Jaddoe, V. W. V., … Verhulst, F.C. (2012). The Generation R Study: A review of design, findings to date and a study of the 5-HTTLPR by environmental interaction from fetal life onward. Journal of the American Academy of Child and Adolescent Psychiatry, 51, 1119- 1135.e7.

Uher, R., & McGuffin, P. (2010). The moderation by the serotonin transporter gene of environmental adversity in the etiology of depression: 2009 update. Molecular Psychiatry, 15, 18-22.

Vassos, E., Collier D. A., & Fazel S. (2014). Systematic meta-analyses and field synopsis of genetic association studies of violence and aggression. Molecular Psychiatry, 19, 471-477.

Viding, E., Price, T. S., Jaffee, S. R., Trzaskowski, M., Davis, O. S. P., Meaburn, E. L., … Plomin, R. (2013). Genetics of callous-unemotional behavior in children. PLoS One, 8, e65789.

Volavka, J., Bilder, R., & Nolan, K. (2004). Catecholamines and aggression: The role of COMT and MAO polymorphisms. Annals of the New York Academy of Sciences, 1036, 393-398.

Winham, S. J. & Biernacka, J. M. (2013). Gene-environment interactions in genome-wide association studies: Current approaches and new directions. Journal of Child Psychology and Psychiatry, 54, 1120-1134.