

Polygenic risk scores implicate genetic pathways involved in neurodevelopmental disorders in hearing thresholds and hearing asymmetry in children

Running title:

Genetics of hearing, asymmetry and neurodevelopment

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Abstract

An efficient auditory system contributes to cognitive and psychosocial development. A right ear advantage in hearing thresholds (HT) has been described in adults and atypical patterns of left/right hearing threshold asymmetry (HTA) have been described for psychiatric and neurodevelopmental conditions. Previous genome-wide association studies (GWAS) on HT have mainly been conducted in elderly participants whose hearing is more likely to be affected by environmental effects. We analysed HT and HTA in a children population cohort (ALSPAC, $n = 6,743$, 7.6 years). Better hearing was associated with more advanced cognitive skills and higher socioeconomic status (SES). Mean HTA was negative (-0.28 dB), suggesting a left ear advantage in children but mainly driven by females (-0.48 dB in females v -0.09 dB in males). We performed the first GWAS on HT in children and the very first GWAS on HTA ($n = 5,344$). Single marker trait association analysis did not yield significant hits. Polygenic risk score (PRS) analysis revealed associations of PRS for schizophrenia with HT, which remained significant after controlling for SES and cognitive skills, and of PRS for autism spectrum disorders (ASD) with HTA. Gene-based analysis for HTA reached genome-wide significance for *MCM5*, which is implicated in axon morphogenesis. This analysis also highlighted other genes associated with contralateral axon crossing. Some of these genes have previously been reported for ASD. These results further support the hypothesis that pathways distinguishing the left/right axis of the brain (i.e. commissural crossing) contribute to both different types of asymmetries (i.e. HTA) and neurodevelopmental disorders.

Keywords: hearing threshold, left/right asymmetry, audiometry, air conduction, autism spectrum disorder, schizophrenia, axonogenesis, neuronal migration, axon guidance, laterality, GWAS, ALSPAC

Introduction

The auditory system involves a series of complex distributed cerebral networks and its impairment affects psychosocial, emotional and cognitive development¹. Hearing-impaired children are at increased risk for learning disabilities² and even within the normal range, better hearing has been associated with better reading skills, working memory and nonverbal IQ in a sample of 1,638 UK school children³.

Hearing ability is usually defined as the threshold in decibel (dB) at which a tone is perceived, so that lower values indicate better hearing. An age-related hearing decline is well documented. In a Korean general population sample ($n > 15,000$) the hearing threshold (HT) for medium frequencies declined from 3 dB in adolescents to 38 dB in elderly participants⁴. A sex difference in favour of women was found in adults ($n = 10,145$; 30-69 years old), but not in children and young adults ($n = 3,458$).

Successful hearing requires transformation of changes in air pressure into vibrations in the basilar membrane that is transferred onto sensory hair cells of the inner ear, whose depolarization is initiated by deflection of mechano-sensitive hair bundles⁵. The auditory nerve transmits these signals to the cochlear nucleus in the brainstem. The majority of the input is transmitted to the contralateral superior olivary complex, while a minor part of the input is transmitted ipsilaterally^{6,7}. Greater contralateral medial olivonuclear suppression in the right compared to the left ear⁸ has been suggested as the underlying correlate of a fundamental functional asymmetry between the left and right ear. This hearing threshold asymmetry (HTA) has typically been reported as HT left – HT right so that positive values indicate an advantage of the right and negative values indicate an advantage of the left ear. In a study of more than 50,000 adults⁹, a right ear advantage (HTA between 1 dB and 4 dB) has been reported with more pronounced HTA in males than in females. In a children sample of $n = 1,191$, a right ear advantage has been reported, albeit to a smaller extent than in adults¹⁰. Other authors found a general right ear advantage in males ($n = \sim 400$, HTA between 0.1 dB and 0.5 dB) and a left ear advantage in females for specific frequencies ($n = \sim 400$, HTA between -0.1 and -0.4 dB)¹¹. Smaller studies reported a general left ear advantage in children^{12,13}.

An absence of HTA has been reported in schizophrenia^{14,15} and ADHD¹⁶. Moreover, symmetrical contralateral suppression in the olivary complex in the left and the right ear in schizophrenia¹⁷ is in contrast with the right ear advantage typically found in controls. In children and adolescents, a right ear advantage has been reported in a sample of $n = 22$ with autism spectrum disorder (ASD) while no asymmetry was found in the control group¹⁸. A developmental effect towards stronger HTA has been

reported in controls that was absent in ASD children ($n = 24$)¹⁹. Reduced laterality in processing auditory stimuli was reported in ASD and bipolar disorder (BIP) suggesting that HTA is linked to neurodevelopmental disorders^{20,21}.

Twin and family studies estimated the heritability for HT to range from .26 to .75 with larger environmental effects for the non-dominant ear²²⁻²⁴. Genome-wide association studies (GWAS) focused on age-related hearing loss in subjects ranging from 45 to 75 years of age and identified genes including *GRM7* and *ESRRG*²⁵⁻²⁸. GWAS for normal hearing included subjects from 18 to 92 years of age²⁹⁻³¹ and implicated several genes (i.e. *DCLK1*, *PTPRD*, *GRM8*, *CMIP*, *SIK3*, *PCDH20*, *SLC28A3*), which have partly been associated with neurodevelopmental traits³²⁻³⁷. A case control design based on electronic health records on age-related hearing loss identified SNPs near *ISG20* and *TRIOBP*³⁸, which had previously been associated with prelingual nonsyndromic hearing loss³⁹. In the UK Biobank, 41 and 7 independent loci have been identified for hearing difficulty and hearing aid use, respectively, implicating genes such as *CDH23*, *EYA4*, *KLHDC7B* and again *TRIOBP*⁴⁰. Mutations in *CDH23* have been associated with early-onset hearing loss and Usher syndrome causing early-onset deafness⁴¹.

However, older subjects have had more exposure to environmental factors which might affect hearing, such as extensive noise⁴², medication⁴³, chemicals⁴⁴ and medical conditions⁴⁵. An investigation of HT in 250 monozygotic (MZ) and 307 dizygotic (DZ) twin pairs from 36 to 80 years of age suggests that environmental effects become more significant with age⁴⁶. Despite this age effect, no study has ever investigated genetic factors involved in hearing function in children.

We analysed HT and HTA in children from the Avon Longitudinal Study of Parents and Children (ALSPAC) ($n = 6,743$). Consistent with previous studies we found that better hearing is associated with enhanced cognitive skills and higher SES. We report the first GWAS for HT in children and the very first GWAS for HTA ($n = 5,344$). In addition to single marker trait associations, we conducted gene-based and gene set analyses and tested the effects of polygenic risk scores (PRS) for a range of neurodevelopmental disorders, IQ and educational attainment (EA). Our results suggest that PRS for schizophrenia are associated with HT, while PRS for ASD are associated with HTA. Genes involved in contralateral axon crossing are associated with HTA, linking genetic factors involved in axonogenesis and left/right axis to different types of asymmetries and neurodevelopmental disorders.

Materials and Methods

Cohort

ALSPAC is a longitudinal cohort representing the general population living in the Bristol area. Pregnant women resident in the county of Avon, UK, with expected dates of delivery from 1st April 1991 to 31st December 1992 were invited to take part in the study, resulting in 14,062 live births and 13,988 children who were alive at 1 year of age^{47,48}. From age seven, all children were invited annually for assessments on a wide range of physical, behavioural and neuropsychological traits. Informed written consent was obtained from the parents after receiving a complete description of the study at the time of enrolment into ALSPAC, with the option to withdraw at any time. Ethical approval for the present study was obtained from the ALSPAC Law and Ethics Committee and the Local Research Ethics Committees. The ALSPAC study website contains details of all the data that is available through a fully searchable data dictionary (<http://www.bris.ac.uk/alspac/researchers/data-access/data-dictionary/>).

Phenotypes

Audiometry was performed according to British Society of Audiologists standards. Hearing tests were carried out in a room with minimal external noise. Testing was stopped if the background noise level exceeded 35 dBA. The air conduction threshold, i.e. the lowest intensity in decibels at which a tone is perceived 50% of the time (dBHL, decibel hearing level), was tested using either a GSI 61 clinical audiometer or a Kamplex AD12 audiometer. Lower dBHL values indicate better hearing. For each ear, the air conduction threshold level was tested at 500 Hz, 1 kHz, 2 kHz and 4 kHz. For each frequency, stimuli were first presented on the right and then on the left ear. The average threshold across different frequencies was derived for each ear.

After applying exclusion criteria (supplementary methods), a sample of $n = 6,743$ was available for phenotypic analysis (3,344 females, 3,391 males, 8 missing values for sex, mean age = 7.59 years, SD = 0.32 years).

HT was defined as the average air conduction threshold on the better ear. HTA was defined as the absolute difference in air conduction threshold between the left and right ear. Thus, positive values indicate a right ear advantage, while negative values indicate a left ear advantage. Handedness was assessed in terms of writing hand (5,805 right-handers, 787 left-handers, 151 missing values).

Cognitive skills were assessed using tests for reading ability⁴⁹, communication skills⁵⁰, listening comprehension⁵¹, short term memory⁵², verbal and performance IQ⁵³ and EA measured as capped General Certificate of Secondary Education (GCSE) scores (for

detailed descriptions, see supplementary methods). Maternal highest educational qualification during pregnancy was used as a proxy for SES⁵⁴. Educational qualification was grouped into 'CSE and no education', 'vocational', 'O level', 'A level' and 'Degree'.

Children were assigned to neurodevelopmental and control subgroups as defined for the ALSPAC sample previously³⁷. We specified subgroups for specific language impairment (SLI) ($n = 155$), reading disability (RD) ($n = 141$), ASD ($n = 35$), ADHD ($n = 21$), comorbidity ($n = 49$) and a control sample matched for sex ($n = 2,071$). The strategies for subgroup assignments are reported in the supplementary methods.

Genotype quality control (QC) and imputation

Genotypes were generated on the Illumina HumanHap550-quad array at the Wellcome Trust Sanger Institute, Cambridge, UK and the Laboratory Corporation of America, Burlington, NC, US. Standard QC was performed as described elsewhere⁵⁵. In total, 9,115 subjects and 500,527 SNPs passed QC filtering. Haplotypes were estimated using ShapeIT (v2.r644). QC-filtered autosomal SNPs were imputed using Impute v3 using the HRC 1.1 reference data panel. Poorly imputed SNPs (Info score < 0.8) and SNPs with low minor allele frequency (MAF < 0.05) were excluded from further analysis.

Statistical analysis

Genetic association tests: Genome-wide genotype data were available for 5,344 children with phenotypes (2,691 males, 2,653 females, mean age = 7.58 years, SD = 0.31 years). HT and HTA were inverse rank-transformed to achieve a normal distribution for GWAS. Association testing was performed in Plink v2 under a generalised linear modeling (GLM) framework specifying sex and the first two ancestry-informative principal components as covariates. Overall, 4,875,234 SNPs that were either directly genotyped or imputed and passed QC were tested for association. The genomic inflation factor (λ) was calculated for all SNPs and revealed no evidence of population structure (HT: $\lambda = 1.02$, HTA: $\lambda = 1.00$). For HT, we specifically tested for replication of markers associated with quantitative hearing phenotypes in previous studies ($p < 10^{-5}$ ^{28,31}, $p < 10^{-6}$ ^{29,30}). We also tested for replication of markers showing genome-wide significance in case-control GWAS on age-related hearing loss ($p < 5 \times 10^{-8}$ ^{38,40}). This procedure resulted in 644 unique markers, 502 of which overlapped with the markers tested in this study. The Bonferroni-corrected significance level was thus set to $0.05/502 = .0001$.

Annotation and gene mapping: We applied FUMA v1.3.6⁵⁶ on the GWAS summary statistics. Functional consequences of SNPs were obtained by performing ANNOVAR⁵⁷ using Ensembl genes (build 85). SNPs were mapped to genes based on positional mapping. Intergenic SNPs were annotated to the closest genes upstream and downstream. Input SNPs were mapped to 18,024 protein-coding genes.

Gene-based and gene set analyses: FUMA implements MAGMA v1.07⁵⁸ to summarise SNP associations at the gene level (gene-based analysis) and associate the set of genes to biological pathways (gene set analysis). Gene-based p values were computed using an F-test in a multiple linear principal components regression while accounting for LD between SNPs. Genome-wide significance was defined as $p = 0.05/18,024 = 2.8 \times 10^{-6}$. For gene set analysis, MAGMA converts gene-based p values into z values, which reflect the strength of association. MAGMA performs a competitive gene set analysis, comparing the mean association of genes within a gene set with the mean association of genes not in the gene set while correcting for gene size and density. Gene set p values were computed for 7,343 gene ontology (GO) terms for biological processes obtained from MsigDB v5.2. The Bonferroni-corrected significance level was set to $0.05/7,343 = 6.8 \times 10^{-6}$.

PRS: We carried out PRS analyses to investigate the genetic overlap of markers associated with HT/HTA and relevant traits and conditions using PRSice 2.2.11.b⁵⁹. PRSice uses GWAS summary statistics as training GWAS to build PRS, which are then tested as predictors in the target GWAS. Psychiatric Genomics Consortium summary statistics were downloaded (<https://www.med.unc.edu/pgc/data-index/>) for schizophrenia⁶⁰, ADHD⁶¹, ASD and BIP⁶² as these are the psychiatric and neurodevelopmental conditions often found to be associated with reduced laterality^{15,17,18,21}. Based on associations of HT with cognitive skills, GCSE scores and SES, we downloaded summary statistics for IQ⁶³ from the Complex Trait Genetics lab website (https://ctg.cncr.nl/software/summary_statistics) and EA⁶⁴ from the Social Science Genetic Association Consortium (<https://www.thessgac.org/data>).

SNPs were clumped based on LD ($r^2 \geq 0.1$) within a 250 kb window. PRS were derived as the weighted sum of risk alleles based on odds ratios or beta values from the training GWAS summary statistics. Sex and first two principal components were included as covariates. Results are presented for the optimal training GWAS p value threshold (explaining the highest proportion of phenotypic variance in the target GWAS). For training GWAS p value thresholds and number of SNPs included in the PRS, see Supplementary Table S1. For six training GWAS and two target GWAS (HT and HTA), the Bonferroni-corrected significance level was set to $.05/12 = .004$. For significant

- 1 effects, analyses were repeated for males and females separately and including only
- 2 the first two principal components as covariates.
- 3 Data preparation and visualization was performed using R v.4.0.0. All analysis scripts
- 4 are available through Open Science Framework (<https://osf.io/gewj2/>).

Results

Phenotypes

We first analysed the distribution of HT and HTA in the overall sample ($n = 6,743$) (results for the subset used in the GWAS, $n = 5,344$, are shown in Supplementary Figure S1). HT ranged from -8.75 to 20.00. Mean HT was 6.14 (SD = 3.97) (Figure 1A). Females ($n = 3,344$) showed slightly higher HT (mean = 6.30, SD = 4.06) than males ($n = 3,391$, mean = 5.98, SD = 3.88), $t_{(6709.2)} = -3.31$, $p = .001$ (Figure 1B).

HTA ranged from -18.75 to 20 and revealed a significant left ear advantage for the overall sample (mean = -0.28, SD = 3.75) as determined by one-sample t -test against zero ($t_{(6742)} = -6.17$, $p = 7.37 \times 10^{-10}$) (Figure 1C). Females showed stronger HTA (mean = -0.48, SD = 3.81) than males (mean = -0.09, SD = 3.68), $t_{(6716.9)} = 4.33$, $p = 1.54 \times 10^{-5}$ (Figure 1D). We thus performed one-sample t -tests against zero for females and males separately. While females showed a significant left ear advantage ($t_{(3343)} = -7.30$, $p = 3.50 \times 10^{-13}$), there was no ear advantage in males ($t_{(3390)} = -1.37$, $p = .172$).

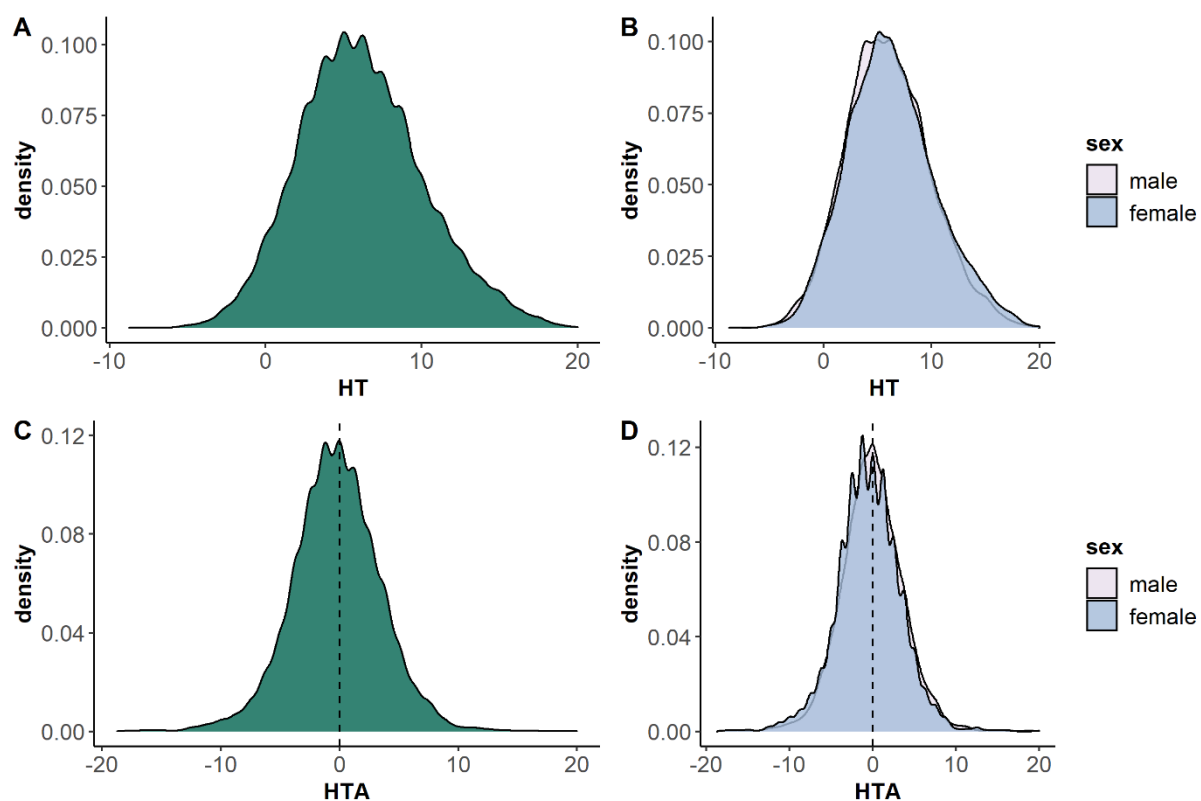


Figure 1: Distribution of HT and HTA. A) Distribution of HT (better ear) in the overall sample ($n = 6743$), and B) as a function of sex. C) Distribution of HTA in the overall sample, and D) as a function of sex. The dotted line represents no asymmetry between the left and right ear.

There was no evidence for an effect of handedness on HTA (right-handers: $n = 5,805$, $M = -0.28$, $SD = 3.75$; left-handers: $n = 787$, $M = -0.24$, $SD = 3.72$, $t_{(1015)} = -0.29$, $p = .776$).

Bivariate Pearson correlations revealed significant positive correlations among the different cognitive measures as previously reported³⁷. There were significant negative correlations after Bonferroni correction (36 comparisons, $p < .0014$) for all cognitive measures but listening comprehension with HT (Figure 2), indicating that lower HT (better hearing) is associated with better cognitive performance (correlation plots are shown in Supplementary Figure S2) and higher GCSE scores. There was no association between HTA and cognitive measures.

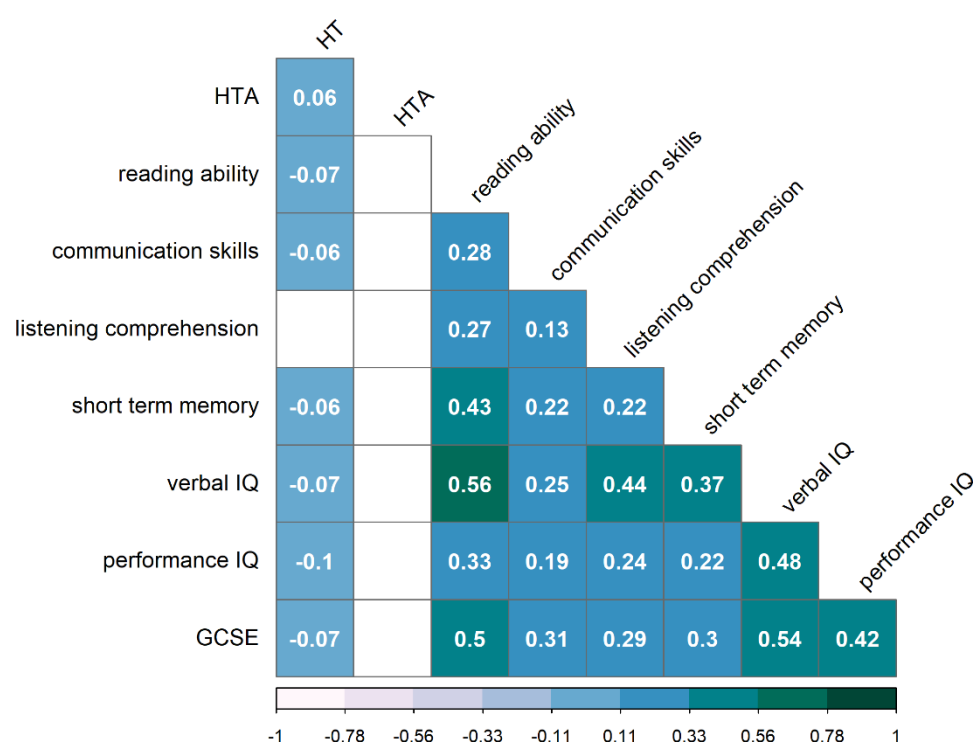


Figure 2: Correlation matrix for HT, HTA and cognitive measures. Correlation coefficients are shown if passing the Bonferroni-corrected significance level ($p < .0014$). Sample sizes range from $n = 4730$ to $n = 6743$ depending on data availability.

One-way between-subjects ANOVAs were conducted to compare the effect of SES on HT and HTA. There was a significant effect of SES on HT ($F_{(4,6137)} = 7.25$, $p = 8.1 \times 10^{-6}$). Post hoc comparisons using the Tukey test indicated significantly lower HT for 'A level' (mean = 6.12, $SD = 3.95$), 'O level' (mean = 5.91, $SD = 3.93$) and 'Degree' (mean = 5.92, $SD = 4.05$) compared to 'CSE' (mean = 6.69, $SD = 4.12$) and significantly lower HT for 'O level' and 'Degree' compared to 'Vocational' (mean = 6.52, $SD = 3.84$), indicating higher SES is associated with better hearing (Supplementary Figure S3). There was no significant effect of SES on HTA ($F_{(4,6137)} = 1.78$, $p = .130$).

Two-sample *t*-tests revealed no difference between children affected by neurodevelopmental disorders and sex-matched controls in HT (Supplementary Table S2) or HTA (Supplementary Table S3). However, there was a consistent pattern across neurodevelopmental subgroups with more negative HTA compared to the control group, indicating more leftward asymmetry.

Genetic association tests

GWAS was performed in $n = 5,344$ children. No individual SNP reached genome-wide significance for HT (Supplementary Table S4). The strongest association was for marker rs11644235 on chromosome 16 ($p = 1.51 \times 10^{-6}$) with each copy of the minor allele (MAF = 0.42) shifting an individual 0.09 standard deviations towards lower HT (i.e. better hearing). In the replication analysis, one marker reached Bonferroni-corrected significance (rs12955474, $\beta = -0.17$, $p = 5.59 \times 10^{-5}$). This marker had been previously reported in a GWAS for the third PC on HT over different frequencies (0.25, 0.5, 1, 2, 3, 4, 6, 8 kHz) ($\beta = -0.10$, $p = 3.57 \times 10^{-7}$)²⁸. Another eight SNPs reached nominal significance with effects in the same direction (Supplementary Table S5), among those five reported on HT at 2 kHz and 4kHz³¹, one reported on PC3²⁹ and two markers reported as genome-wide significant for hearing loss⁴⁰.

No individual SNP reached genome-wide significance for HTA (Supplementary Table S6). The marker rs10434985 on chromosome 7 was the most strongly associated SNP ($p = 7.27 \times 10^{-7}$) with each copy of the minor allele (MAF = 0.21) shifting an individual 0.12 standard deviations towards better hearing on the left ear. Manhattan and QQ plots are shown in Supplementary Figure S4-S6.

Gene-based analysis highlighted three suggestive associations for HT (*AE000662.92*: association $p = 2.15 \times 10^{-5}$; *C6orf201*: association $p = 4.10 \times 10^{-5}$; *FAM217A*: association $p = 6.95 \times 10^{-5}$) (Figure 3).

The *MCM5* gene reached the genome-wide significance level for HTA ($p = 4.30 \times 10^{-7}$) (Figure 3). Other four genes reached the suggestive significance level (*LHX6*, $p = 1.77 \times 10^{-5}$; *NRP1*, $p = 4.50 \times 10^{-5}$; *RUVBL1*, $p = 5.89 \times 10^{-5}$; *EPHA8*, $p = 7.67 \times 10^{-5}$). QQ plots are shown in Supplementary Figures S7-S8.

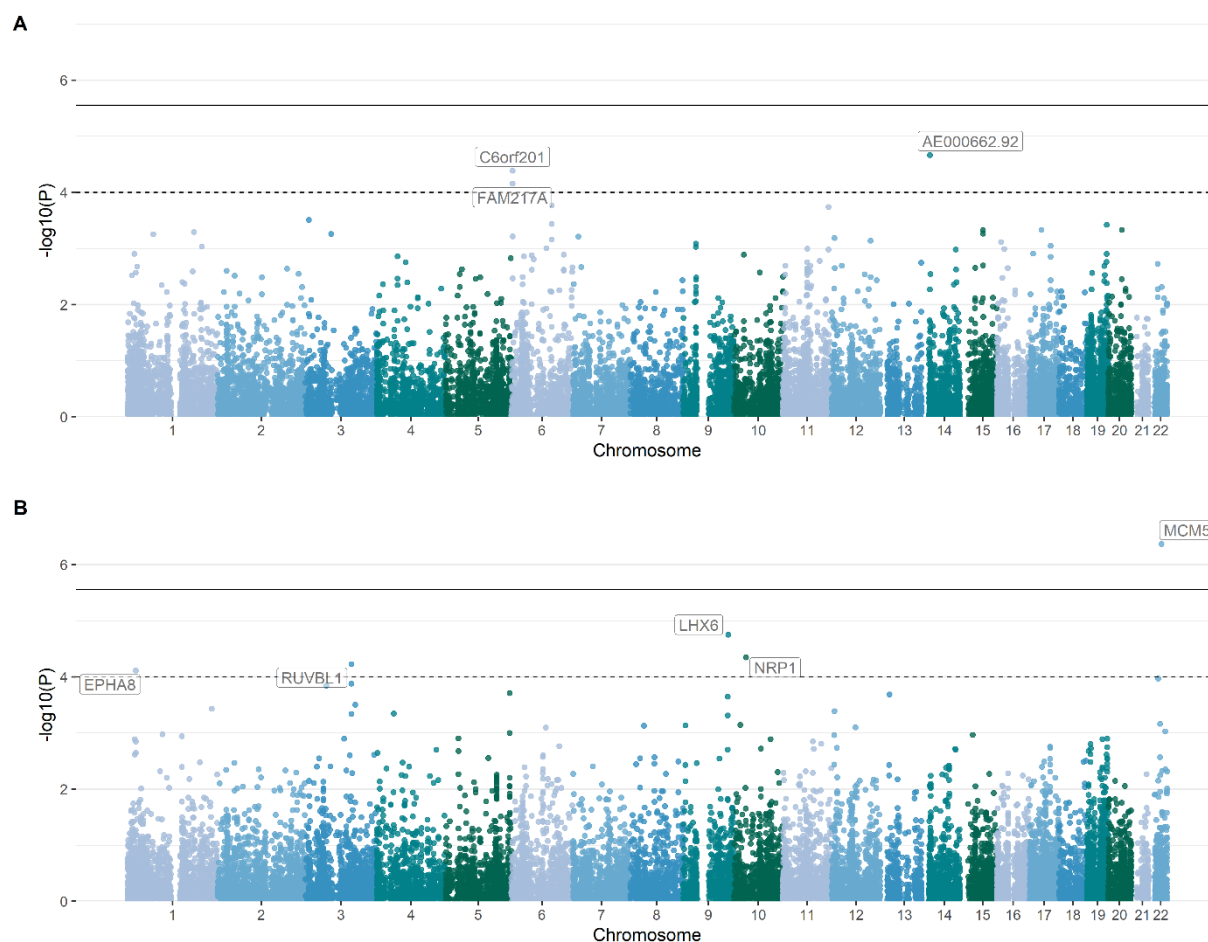


Figure 3: Gene-based analysis results. Manhattan plots are shown for A) HT and B) HTA. Gene-based association p values are plotted against chromosome and position. The solid line represents the genome-wide significance level ($p = 2.8 \times 10^{-6}$), the dotted line represents the suggestive significance level ($p = .0001$).

In gene set enrichment analysis no GO term reached the Bonferroni-corrected level of significance ($p = 6.8 \times 10^{-6}$) for either HT or HTA. “Translational termination (GO: 0006415)” ($p = 1.17 \times 10^{-5}$) and “cerebral cortex tangential migration (GO:0021800)” ($p = 4.17 \times 10^{-5}$) were the strongest associations detected for HT and HTA, respectively.

PRS

PRS were tested for IQ and EA based on the correlations between HT and cognitive measures and GCSE scores (Figure 2). PRS for four neurodevelopmental conditions were tested on the basis of previously reported associations with HTA in the literature^{15,17,18,21}. PRS for ADHD showed an association with HT ($R^2 = 0.003$, $\beta = 314.98$, $SE = 102.78$, $p = .002$, Figure 4), indicating that higher genetic risk for ADHD is associated with higher HT, i.e. worse hearing. This effect was stronger in females than in males (females: $R^2 = 0.004$, $\beta = 571.67$, $SE = 268.11$, $p = .033$; males: $R^2 = 0.003$, $\beta = 275.60$, $SE =$

97.83, $p = .005$). In contrast, schizophrenia PRS showed a negative association with HT, suggesting that higher genetic risk for schizophrenia is associated with lower HT, i.e. better hearing ($R^2 = 0.003$, $\beta = -280.16$, $SE = 85.70$, $p = .001$, Figure 4). This effect was stronger in males than in females (females: $R^2 = 0.003$, $\beta = -100.73$, $SE = 54.99$, $p = .067$; males: $R^2 = 0.004$, $\beta = -408.09$, $SE = 128.37$, $p = .001$). PRS for EA reached borderline significance for HT ($R^2 = 0.003$, $\beta = -1593.45$, $SE = 588.06$, $p = .007$, Figure 4). The negative association suggests that a genetic predisposition towards higher EA is associated with better hearing.



Figure 4: PRS analysis results. $-\log_{10}(p)$ values are reported for PRS analysis on six training GWAS and two target GWAS (HT and HTA). PRS for ADHD and schizophrenia contribute to HT, while PRS for ASD contributes to HTA. BIP = bipolar disorder, SCZ = schizophrenia, EA = educational attainment.

Based on this association and behavioural correlations, we next tested whether the associations between ADHD and schizophrenia PRS with HT were mediated by cognitive skills. We thus reran the PRS analysis on HT with ADHD and schizophrenia as training GWAS and using sex, the first two ancestry-informative principal components and cognitive skills as covariates. This was done separately for reading ability, communication skills, short term memory, verbal IQ, performance IQ and GCSE score. In addition, we tested whether the associations between ADHD and schizophrenia PRS with HT were mediated by SES and EA PRS by using these variables as covariates. This procedure resulted in 16 PRS analyses (two training GWAS, eight covariates). The effect of ADHD PRS on HT was reduced after adjusting for most cognitive skills, SES and EA PRS (β ranging from 2.90 to 309.32, Supplementary Table S7), suggesting that the association is mediated by these variables. In contrast, the effect of schizophrenia PRS on HT remained similar after adjusting for SES, GCSE and EA PRS and was enlarged after adjusting for cognitive skills (β ranging from -302.94 to -422.54).

PRS for ASD showed a negative association with HTA, indicating that higher genetic risk for ASD is associated with more leftward HTA ($R^2 = 0.004$, $\beta = -461.79$, $SE = 158.45$, $p = .0036$, Figure 4). Thus, higher PRS for ASD shift the mean towards the left of the distribution, which suggests stronger asymmetry. This effect was stronger in males than in females (females: $R^2 = 0.003$, $\beta = -404.05$, $SE = 223.93$, $p = .071$; males: $R^2 = 0.003$, $\beta = -982.26$, $SE = 373.22$, $p = .009$). There was no effect of PRS for IQ or BIP on either phenotype.

Discussion

We report a comprehensive study of HT and HTA in children to dissect their relationship with cognitive abilities and neurodevelopmental disorders both at phenotypic and genetic level. We confirm that better hearing is associated with better performance on a range of cognitive abilities (Figure 2). We also report the results of the first GWAS on HT in children. Single marker, gene-based and gene set enrichment analyses did not lead to any statistically significant results. However, PRS for both ADHD and schizophrenia were significantly associated with HT (Figure 4), with genetic risk for ADHD and schizophrenia increasing and decreasing HT, respectively. We also report the very first GWAS for HTA. Although we did not detect any single marker associations, we found that PRS for ASD were statistically associated with HTA (Figure 4), suggesting that higher genetic risk for ASD is associated with a shift towards the left ear, indicating more asymmetry with better left ear performance. Gene-based analysis for HTA highlighted several genes involved in axon guidance and commissural axon crossing in sensory cerebral networks. Some of these genes have previously been implicated in ASD.

At the phenotypic level, previous studies reported lower HT in females⁶⁵. However, this sex difference emerged around the age of 30, while there was no sex difference in children and young adults⁴. Our data ($n = 6,743$) showed slightly higher HT in females (6.30 dB) compared to males (5.98 dB) (Figure 1). Therefore, our data are in agreement with a sex-specific developmental trajectory resulting in better hearing in female adults that is not detectable in children. Consistent with previous studies³, we found that in the normal range of variation, HT is negatively associated with several cognitive skills (Figure 2). We did not detect any association between PRS for IQ and HT suggesting that this association is not mediated by shared biological pathways. A cause-effect relationship would be possible, but cannot be easily explained by our data. The analysis of PRS for ADHD and schizophrenia instead support a role of genes implicated in neurodevelopmental disorders contributing to HT. There was no association between HT and ADHD at the behavioural level, however this analysis was based on a very small sample of children meeting the criteria for ADHD ($n = 21$) in our dataset and therefore the results might not be conclusive (Supplementary Table S2). Hearing deficits in ADHD have been reported in terms of speech perception⁶⁶, but not in air conduction thresholds. Moreover, the effect of ADHD PRS on HT was reduced after adjusting for cognitive skills, PRS and EA PRS (Supplementary Table S7), suggesting that the effect was mediated by cognitive factors. In contrast, we found that higher genetic risk for schizophrenia is associated with better hearing. This effect was still found after adjusting for cognitive skills (Supplementary Table S7). PRS for

schizophrenia have recently been associated with better language skills, but not overall school performance⁶⁷. Thus, the association between better hearing and language development (Figure 2) could be based on shared biological pathways which also increase the risk for schizophrenia. On the phenotypic level, there are no HT differences between individuals affected by schizophrenia and controls⁶⁸. Although no single marker trait associations reached significance in the GWAS for HT (Supplementary Figure S4), the top marker on chromosome 15 (rs1039444) is located in an intron of *RAB8B*, which encodes for a GTPase that is expressed in inner and outer hair cells and is involved in autosomal recessive deafness⁶⁹. Targeted analysis for markers reported in previous GWAS for HT in adults replicated association with only one marker, rs12955474, which is located in an intron of the *CCBE1* gene²⁸. Other markers in this gene have been associated with depression⁷⁰ and left entorhinal cortex volume⁷¹.

Phenotypic analysis for HTA revealed an overall left ear advantage with a lower air conduction threshold of 0.28 dB on average (Figure 1), replicating results from smaller studies in children^{12,13}. This result, which indicates a left ear advantage, was driven by females. Thus, a tendency towards the sex effect on HTA reported in adults seems to be established already in children. It is possible that a developmental shift towards the right ear in both sexes, resulting in reduced asymmetry in female and a rightward asymmetry in males, is driven by environmental factors⁷². This developmental shift towards stronger HTA has not been observed in children with ASD¹⁹. We found that PRS for ASD are associated with HTA. Higher genetic risk for ASD was linked to better hearing on the left compared to the right ear. Although not significant, this is consistent with the behavioural data, where we find a more leftward asymmetry in cases than controls across different subgroups for disorders.

We found more negative HTA in neurodevelopmental conditions including ASD compared to controls on the behavioural level that is congruent with the PRS results. Different types of asymmetry such as structural brain asymmetry⁷³, frontal alpha asymmetry⁷⁴, language processing⁷⁵ and handedness⁷⁶ have been implicated in ASD.

Gene-based analysis on HTA revealed genome-wide significance for the *MCM5* gene (Figure 3). *MCM5* has been linked to ASD by an algorithm modeling genomic data from cortical tissue⁷⁷. In *C. elegans*, homozygous *mcm-5* mutants showed reduced neuron numbers and absent commissures from ventral to dorsal cord neurons⁷⁸. Among the genes reaching the suggestive significance level, *LHX6* is required for neuronal migration⁷⁹ and survival of cortical interneurons⁸⁰. The receptor encoded by *EPHA8* plays a critical role in axonal guidance during neurodevelopment⁸¹. In mice,

Epha8 has been shown contribute to correct formation of crossed commissural axons contributing to the auditory system⁸². *NRP1* encodes for a receptor associated with contralateral axon crossing at the optic chiasm in zebrafish⁸³ and mice⁸⁴. *NRP1* interacts with *TAOK2*, for which downregulation has also been reported to impair axon crossing at the midline⁸⁵ and to decrease the volumes of the *corpus callosum* and anterior commissure⁸⁶. *TAOK2* is located on chromosome 16p11.2, which has been linked to ASD susceptibility⁸⁷. Overall, the top hits of gene-based analysis for HTA have roles in axon morphogenesis and controlling midline crossing. Previous large-scale GWAS for handedness and structural brain asymmetries suggested a role of genes involved in axonogenesis, microtubules and cytoskeleton formation⁸⁸⁻⁹¹. A smaller GWAS meta-analysis (including the ALSPAC sample) on a quantitative measure of relative hand skill implicated genes with a known role in establishing body asymmetries⁵⁵ that also play a role in neurodevelopmental disorders⁹². Of note, *RUVBL1*, one of the genes reaching suggestive significance in the GWAS on HTA, plays a fundamental role in symmetry breaking and cardiac development⁹³. Moreover, *LHX6* inhibits the transcription of *PITX2*⁹⁴, which encodes for a transcription factor that is asymmetrically expressed during development of the heart and other organs and plays a direct role in their asymmetric morphology⁹⁵. Thus, the current results are in line with the idea that shared genetic components contribute to different types of asymmetry and suggest that genes affecting structural asymmetry and contralateral axon crossing in early embryonic development might contribute to laterality and neurodevelopmental traits⁹⁶.

A possible limitation of the current study is that stimulus presentation was not randomised, but the right ear was always tested first. Testing the right ear first has been more common in previous studies than vice versa, which could either result in a learning effect (favouring the left ear) or in a fatigue effect (favouring the right ear)⁹⁷. However, in adults, a right ear advantage is more common even in studies in which the order of stimulus presentation has been randomised⁹⁷, so the effect of stimulus presentation should be minimal. The main limitation of this study is the rather small sample size. It is of note that air conduction thresholds are not routinely collected in large-scale population studies. For example, the UK Biobank includes phenotypic information on hearing ability as self-reported hearing difficulty or use of hearing aids ($n > 300,000$)^{40,98}. Similarly, a GWAS on age-related hearing loss in the Genetic Epidemiology Research on Adult Health and Aging (GERA) cohort ($n > 50,000$) used a case control design, identifying cases based on health records (ICD-9 diagnosis)³⁸. GWAS on quantitative measures of hearing ability were limited to smaller sample sizes below 6,000 subjects for individual samples^{27,29}. Systematic collection of air

1 conduction thresholds in both ears in children would enable larger genetic studies to
2 dissect the links between hearing, asymmetry and neurodevelopment.

3 In summary, our results highlight the importance of HT for cognitive development.
4 We find that PRS for schizophrenia are implicated in HT, while PRS for ASD are
5 implicated in HTA. This is in line with the increasing evidence supporting a role of
6 asymmetries in ASD. Gene-based analysis highlighted genes involved in axon guidance
7 and ASD for HTA suggesting a role of genes involved in contralateral axon crossing
8 at the midline. The results support the hypothesis of shared pathways contributing to
9 different types of asymmetries, neurodevelopment and disorders.

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Conflict of Interest

The authors declare no competing financial interests in relation to the work described.

References

1. Stevenson J. *et al.* Language and reading comprehension in middle childhood predicts emotional and behaviour difficulties in adolescence for those with permanent childhood hearing loss. *J Child Psychol Psychiatry* **59**, 180–190 (2018).
2. Choi S.M.R., Kei J. & Wilson W.J. Learning difficulties and auditory processing deficits in a clinical sample of primary school-aged children. *Int J Audiol*, 1–7 (2020).
3. Moore D.R., Zobay O. & Ferguson M.A. Minimal and Mild Hearing Loss in Children: Association with Auditory Perception, Cognition, and Communication Problems. *Ear Hear*, <https://10.1097/AUD.0000000000000802> (2019).
4. Park Y.H., Shin S.-H., Byun S.W. & Kim J.Y. Age- and Gender-Related Mean Hearing Threshold in a Highly-Screened Population: The Korean National Health and Nutrition Examination Survey 2010–2012. *PLoS ONE* **11**, e0150783 (2016).
5. Schwander M., Kachar B. & Müller U. Review series: The cell biology of hearing. *J Cell Biol* **190**, 9–20 (2010).
6. Felix R.A., Gourévitch B. & Portfors C.V. Subcortical pathways: Towards a better understanding of auditory disorders. *Hearing Research* **362**, 48–60 (2018).
7. Warren E.H. & Liberman M.C. Effects of contralateral sound on auditory-nerve responses. I. Contributions of cochlear efferents. *Hearing Research* **37**, 89–104 (1989).
8. Khalfa S. & Collet L. Functional asymmetry of medial olivocochlear system in humans. Towards a peripheral auditory lateralization. *Neuroreport* **7**, 993–996 (1996).
9. Chung D.Y., Mason K., Gannon R.P. & Willson G.N. The ear effect as a function of age and hearing loss. *J Acoust Soc Am* **73**, 1277–1282 (1983).
10. Eagles E.L. A longitudinal study of ear disease and hearing sensitivity in children. *Audiology* **12**, 438–445 (1973).
11. Kannan P.M. & Lipscomb D.M. Letter: Bilateral hearing asymmetry in a large population. *J Acoust Soc Am* **55**, 1092–1094 (1974).
12. Roche A.F., Siervogel R.M., Himes J.H. & Johnson D.L. Longitudinal study of hearing in children: baseline data concerning auditory thresholds, noise exposure, and biological factors. *J Acoust Soc Am* **64**, 1593–1616 (1978).
13. Rahko T. & Karma P. Pure-tone hearing thresholds in otologically healthy 5-year-old children in Finland. *Arch Otorhinolaryngol* **246**, 137–141 (1989).
14. Gruzelier J.H. & Hammond N.V. Gains, losses and lateral differences in the hearing of schizophrenic patients. *Br J Psychol* **70**, 319–330 (1979).
15. Mathew V.M., Gruzelier J.H. & Liddle P.F. Lateral asymmetries in auditory acuity distinguish hallucinating from nonhallucinating schizophrenic patients. *Psychiatry Research* **46**, 127–138 (1993).
16. Combs J.T. Lack of right ear advantage in patients with attention-deficit/hyperactivity disorder. *Clin Pediatr (Phila)* **41**, 231–234 (2002).
17. Veuillet E., Georgieff N., Philibert B., Dalery J., Marie-Cardine M. & Collet L. Abnormal peripheral auditory asymmetry in schizophrenia. *J Neurol Neurosurg Psychiatry* **70**, 88–94 (2001).
18. Khalfa S. *et al.* Peripheral auditory asymmetry in infantile autism. *Eur J Neurosci* **13**, 628–632 (2001).

19. James A.L. & Barry R.J. Developmental effects in the cerebral lateralization of autistic, retarded, and normal children. *J Autism Dev Disord* **13**, 43–56 (1983).
20. Schmidt G.L., Rey M.M., Oram Cardy J.E. & Roberts T.P.L. Absence of M100 source asymmetry in autism associated with language functioning. *Neuroreport* **20**, 1037–1041 (2009).
21. Reite M., Teale P., Rojas D.C., Reite E., Asherin R. & Hernandez O. MEG auditory evoked fields suggest altered structural/functional asymmetry in primary but not secondary auditory cortex in bipolar disorder. *Bipolar Disord* **11**, 371–381 (2009).
22. Viljanen A. *et al.* Genetic and environmental influences on hearing at different frequencies separately for the better and worse hearing ear in older women. *Int J Audiol* **46**, 772–779 (2007).
23. Gates G.A., Couropmitree N.N. & Myers R.H. Genetic associations in age-related hearing thresholds. *Arch Otolaryngol Head Neck Surg* **125**, 654–659 (1999).
24. Viljanen A., Era P., Kaprio J., Pyykkö I., Koskenvuo M. & Rantanen T. Genetic and environmental influences on hearing in older women. *J Gerontol A Biol Sci Med Sci* **62**, 447–452 (2007).
25. Huyghe J.R. *et al.* Genome-wide SNP-based linkage scan identifies a locus on 8q24 for an age-related hearing impairment trait. *Am J Hum Genet* **83**, 401–407 (2008).
26. Friedman R.A. *et al.* GRM7 variants confer susceptibility to age-related hearing impairment. *Hum Mol Genet* **18**, 785–796 (2009).
27. Nolan L.S. *et al.* Estrogen-related receptor gamma and hearing function: evidence of a role in humans and mice. *Neurobiol Aging* **34**, 2077.e1-9 (2013).
28. Fransen E. *et al.* Genome-wide association analysis demonstrates the highly polygenic character of age-related hearing impairment. *Eur J Hum Genet* **23**, 110–115 (2015).
29. Vuckovic D. *et al.* Genome-wide association analysis on normal hearing function identifies PCDH20 and SLC28A3 as candidates for hearing function and loss. *Hum Mol Genet* **24**, 5655–5664 (2015).
30. Wolber L.E. *et al.* Salt-inducible kinase 3, SIK3, is a new gene associated with hearing. *Hum Mol Genet* **23**, 6407–6418 (2014).
31. Girotto G. *et al.* Hearing function and thresholds: a genome-wide association study in European isolated populations identifies new loci and pathways. *J Med Genet* **48**, 369–374 (2011).
32. Håvik B. *et al.* DCLK1 variants are associated across schizophrenia and attention deficit/hyperactivity disorder. *PLoS ONE* **7**, e35424 (2012).
33. Takaki H., Kikuta R., Shibata H., Ninomiya H., Tashiro N. & Fukumaki Y. Positive associations of polymorphisms in the metabotropic glutamate receptor type 8 gene (GRM8) with schizophrenia. *Am J Med Genet B Neuropsychiatr Genet* **128B**, 6–14 (2004).
34. Zhang L. *et al.* Association analysis of the GRM8 gene with schizophrenia in the Uyghur Chinese population. *Hereditas* **151**, 140–144 (2014).
35. Newbury D.F. *et al.* CMIP and ATP2C2 modulate phonological short-term memory in language impairment. *Am J Hum Genet* **85**, 264–272 (2009).
36. Newbury D.F., Gibson J.L., Conti-Ramsden G., Pickles A., Durkin K. & Toseeb U. Using Polygenic Profiles to Predict Variation in Language and Psychosocial Outcomes in Early and Middle Childhood. *J Speech Lang Hear Res* **62**, 3381–3396 (2019).

37. Scerri T.S. *et al.* DCDC2, KIAA0319 and CMIP are associated with reading-related traits. *Biol Psychiatry* **70**, 237–245 (2011).
38. Hoffmann T.J., Keats B.J., Yoshikawa N., Schaefer C., Risch N. & Lustig L.R. A Large Genome-Wide Association Study of Age-Related Hearing Impairment Using Electronic Health Records. *PLoS Genet* **12**, e1006371 (2016).
39. Diaz-Horta O. *et al.* Whole-exome sequencing efficiently detects rare mutations in autosomal recessive nonsyndromic hearing loss. *PLoS ONE* **7**, e50628 (2012).
40. Wells H.R.R. *et al.* GWAS Identifies 44 Independent Associated Genomic Loci for Self-Reported Adult Hearing Difficulty in UK Biobank. *Am J Hum Genet* **105**, 788–802 (2019).
41. Schultz J.M. *et al.* Allelic hierarchy of CDH23 mutations causing non-syndromic deafness DFNB12 or Usher syndrome USH1D in compound heterozygotes. *J Med Genet* **48**, 767–775 (2011).
42. Stanbury M., Rafferty A.P. & Rosenman K. Prevalence of hearing loss and work-related noise-induced hearing loss in Michigan. *J Occup Environ Med* **50**, 72–79 (2008).
43. Ruhl D.S., Cable B.B. & Martell D.W. Medication associated with hearing loss: 25 years of medical malpractice cases in the United States. *Otolaryngol Head Neck Surg* **151**, 431–437 (2014).
44. Morata T.C. Chemical exposure as a risk factor for hearing loss. *J Occup Environ Med* **45**, 676–682 (2003).
45. Kakarlapudi V., Sawyer R. & Staecker H. The effect of diabetes on sensorineural hearing loss. *Otol Neurotol* **24**, 382–386 (2003).
46. Karlsson K.K., Harris J.R. & Svartengren M. Description and primary results from an audiometric study of male twins. *Ear Hear* **18**, 114–120 (1997).
47. Boyd A. *et al.* Cohort Profile: the 'children of the 90s'--the index offspring of the Avon Longitudinal Study of Parents and Children. *Int J Epidemiol* **42**, 111–127 (2013).
48. Fraser A. *et al.* Cohort Profile: the Avon Longitudinal Study of Parents and Children: ALSPAC mothers cohort. *Int J Epidemiol* **42**, 97–110 (2013).
49. Rust J., Golombok S. & Trickey G. *WORD Wechsler Objective Reading Dimensions Manual* (Psychological Corp: Sidcup, UK, 1993).
50. Bishop D.V.M. Development of the Children's Communication Checklist (CCC): a method for assessing qualitative aspects of communicative impairment in children. *J Child Psychol Psychiatry* **39**, 879–891 (1998).
51. Rust J. *WOLD Wechsler Objective Language Dimensions Manual* (Psychological Corp: London, UK, 1996).
52. Gathercole S.E., Willis C.S., Baddeley A.D. & Emslie H. The Children's Test of Nonword Repetition: a test of phonological working memory. *Memory* **2**, 103–127 (1994).
53. Wechsler D., Golombok S. & Rust J. *Wechsler Intelligence Scale for Children - Third Edition UK Manual* (Psychological Corp: Sidcup, UK, 1991).
54. Rashid V. *et al.* Ethnicity and socioeconomic status are related to dietary patterns at age 5 in the Amsterdam born children and their development (ABCD) cohort. *BMC Public Health* **18**, 115 (2018).
55. Brandler W.M. *et al.* Common variants in left/right asymmetry genes and pathways are associated with relative hand skill. *PLoS Genet* **9**, e1003751 (2013).

56. Watanabe K., Taskesen E., van Bochoven A. & Posthuma D. Functional mapping and annotation of genetic associations with FUMA. *Nat Commun* **8**, 1826 (2017).
57. Wang K., Li M. & Hakonarson H. ANNOVAR: functional annotation of genetic variants from high-throughput sequencing data. *Nucleic Acids Res* **38**, e164 (2010).
58. de Leeuw C.A., Mooij J.M., Heskes T. & Posthuma D. MAGMA: generalized gene-set analysis of GWAS data. *PLoS Comput Biol* **11**, e1004219 (2015).
59. Choi S.W. & O'Reilly P.F. PRSice-2: Polygenic Risk Score software for biobank-scale data. *Gigascience* **8** (2019).
60. Schizophrenia Working Group of the Psychiatric Genomics Consortium. Biological insights from 108 schizophrenia-associated genetic loci. *Nature* **511**, 421–427 (2014).
61. Neale B.M. *et al.* Meta-analysis of genome-wide association studies of attention-deficit/hyperactivity disorder. *J Am Acad Child Adolesc Psychiatry* **49**, 884–897 (2010).
62. Psychiatric GWAS Consortium Bipolar Disorder Working Group. Large-scale genome-wide association analysis of bipolar disorder identifies a new susceptibility locus near ODZ4. *Nat Genet* **43**, 977–983 (2011).
63. Savage J.E. *et al.* Genome-wide association meta-analysis in 269,867 individuals identifies new genetic and functional links to intelligence. *Nat Genet* **50**, 912–919 (2018).
64. Lee J.J. *et al.* Gene discovery and polygenic prediction from a genome-wide association study of educational attainment in 1.1 million individuals. *Nat Genet* **50**, 1112–1121 (2018).
65. McFadden D. A speculation about the parallel ear asymmetries and sex differences in hearing sensitivity and otoacoustic emissions. *Hearing Research* **68**, 143–151 (1993).
66. Fuermaier A.B.M. *et al.* Perception in attention deficit hyperactivity disorder. *Atten Defic Hyperact Disord* **10**, 21–47 (2018).
67. Rajagopal V.M. *et al.* Genome-wide association study of school grades identifies a genetic overlap between language ability, psychopathology and creativity (2020).
68. Prager S. & Jeste D.V. Sensory impairment in late-life schizophrenia. *Schizophr Bull* **19**, 755–772 (1993).
69. Heidrych P. *et al.* Rab8b GTPase, a protein transport regulator, is an interacting partner of otoferlin, defective in a human autosomal recessive deafness form. *Hum Mol Genet* **17**, 3814–3821 (2008).
70. Power R.A. *et al.* Genome-wide association analysis accounting for environmental factors through propensity-score matching: application to stressful life events in major depressive disorder. *American Journal of Medical Genetics Part B: Neuropsychiatric Genetics* **162B**, 521–529 (2013).
71. Zhao B. *et al.* Genome-wide association analysis of 19,629 individuals identifies variants influencing regional brain volumes and refines their genetic co-architecture with cognitive and mental health traits. *Nat Genet* **51**, 1637–1644 (2019).
72. Nageris B.I., Raveh E., Zilberberg M. & Attias J. Asymmetry in noise-induced hearing loss: relevance of acoustic reflex and left or right handedness. *Otol Neurotol* **28**, 434–437 (2007).
73. Postema M.C. *et al.* Altered structural brain asymmetry in autism spectrum disorder in a study of 54 datasets. *Nat Commun* **10**, 4958 (2019).

74. Gabard-Durnam L., Tierney A.L., Vogel-Farley V., Tager-Flusberg H. & Nelson C.A. Alpha asymmetry in infants at risk for autism spectrum disorders. *J Autism Dev Disord* **45**, 473–480 (2015).
75. Herringshaw A.J., Ammons C.J., DeRamus T.P. & Kana R.K. Hemispheric differences in language processing in autism spectrum disorders: A meta-analysis of neuroimaging studies. *Autism Res* **9**, 1046–1057 (2016).
76. Markou P., Ahtam B. & Papadatou-Pastou M. Elevated Levels of Atypical Handedness in Autism: Meta-Analyses. *Neuropsychol Rev* **27**, 258–283 (2017).
77. Liu L. *et al.* DAWN: a framework to identify autism genes and subnetworks using gene expression and genetics. *Mol Autism* **5**, 22 (2014).
78. Wang X., Suh C., Zhu Z. & Fan Q. Minichromosome maintenance protein 5 homologue in *Caenorhabditis elegans* plays essential role for postembryonic development. *Biochemical and Biophysical Research Communications* **359**, 965–971 (2007).
79. Liodis P., Denaxa M., Grigoriou M., Akufo-Addo C., Yanagawa Y. & Pachnis V. Lhx6 activity is required for the normal migration and specification of cortical interneuron subtypes. *J. Neurosci.* **27**, 3078–3089 (2007).
80. Denaxa M. *et al.* Modulation of Apoptosis Controls Inhibitory Interneuron Number in the Cortex. *Cell Rep* **22**, 1710–1721 (2018).
81. Holland S.J., Peles E., Pawson T. & Schlessinger J. Cell-contact-dependent signalling in axon growth and guidance: Eph receptor tyrosine kinases and receptor protein tyrosine phosphatase β . *Current Opinion in Neurobiology* **8**, 117–127 (1998).
82. Park S., Frisén J. & Barbacid M. Aberrant axonal projections in mice lacking EphA8 (Eek) tyrosine protein kinase receptors. *EMBO J* **16**, 3106–3114 (1997).
83. Sakai J.A. & Halloran M.C. Semaphorin 3d guides laterality of retinal ganglion cell projections in zebrafish. *Development* **133**, 1035–1044 (2006).
84. Erskine L. *et al.* VEGF signaling through neuropilin 1 guides commissural axon crossing at the optic chiasm. *Neuron* **70**, 951–965 (2011).
85. Anda F.C. de *et al.* Autism spectrum disorder susceptibility gene TAOK2 affects basal dendrite formation in the neocortex. *Nat Neurosci* **15**, 1022–1031 (2012).
86. Richter M. *et al.* Altered TAOK2 activity causes autism-related neurodevelopmental and cognitive abnormalities through RhoA signaling. *Mol Psychiatry* **24**, 1329–1350 (2019).
87. Weiss L.A. *et al.* Association between microdeletion and microduplication at 16p11.2 and autism. *N Engl J Med* **358**, 667–675 (2008).
88. Cuellar Partida G. *et al.* Genome-wide association study identifies 48 common genetic variants associated with handedness (2019).
89. de Kovel C.G.F. & Francks C. The molecular genetics of hand preference revisited. *Sci Rep* **9**, 5986 (2019).
90. Wiberg A. *et al.* Handedness, language areas and neuropsychiatric diseases: insights from brain imaging and genetics. *Brain* **142**, 2938–2947 (2019).
91. Sha Z. *et al.* The genetic architecture of structural left-right asymmetry of the human brain (2020).
92. Paracchini S., Diaz R. & Stein J.F. Advances in Dyslexia Genetics-New Insights Into the Role of Brain Asymmetries. *Adv Genet* **96**, 53–97 (2016).
93. Hartill V.L. *et al.* DNAAF1 links heart laterality with the AAA+ ATPase RUVBL1 and ciliary intraflagellar transport. *Hum Mol Genet* **27**, 529–545 (2018).

- 1 94.Zhang Z. *et al.* The LIM homeodomain transcription factor LHX6: a transcriptional
2 repressor that interacts with pituitary homeobox 2 (PITX2) to regulate odontogenesis. *J*
3 *Biol Chem* **288**, 2485–2500 (2013).
- 4 95.Norris D.P. Cilia, calcium and the basis of left-right asymmetry. *BMC Biol* **10**, 102 (2012).
- 5 96.Brandler W.M. & Paracchini S. The genetic relationship between handedness and
6 neurodevelopmental disorders. *Trends Mol Med* **20**, 83–90 (2014).
- 7 97.Pirilä T., Jounio-Ervasti K. & Sorri M. Left-right asymmetries in hearing threshold levels
8 in three age groups of a random population. *Audiology* **31**, 150–161 (1992).
- 9 98.Kalra G. *et al.* *Biological insights from multi-omic analysis of 31 genomic risk loci for adult*
10 *hearing difficulty* (2019).
- 11