Discovery of the first genome-wide significant

risk loci for ADHD

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Abstract

Attention-Deficit/Hyperactivity Disorder (ADHD) is a highly heritable childhood behavioral disorder affecting 5% of school-age children and 2.5% of adults. Common genetic variants contribute substantially to ADHD susceptibility, but no individual variants have been robustly associated with ADHD. We report a genome-wide association meta-analysis of 20,183 ADHD cases and 35,191 controls ascertained from clinical interviews and/or medical records that identifies variants surpassing genome-wide significance in 12 independent loci, revealing new and important information on the underlying biology of ADHD. Associations are enriched in evolutionarily constrained genomic regions and loss-of-function intolerant genes, as well as around brain-expressed regulatory marks. Additional analyses of three replication cohorts; a cohort of diagnosed ADHD, a self-reported ADHD sample and a study of quantitative measures of ADHD symptoms in the population, broadly support these findings while highlighting potential study-specific effects on genetic overlap with educational attainment. The strong concordance with GWAS of quantitative population measures og ADHD symptoms supports the hypothesis that clinical diagnosis of ADHD is an extreme expression of one or more continuous heritable traits.

Background

Attention-Deficit/Hyperactivity Disorder (ADHD) is a neurodevelopmental psychiatric disorder, that affects around 5% of children and adolescents and 2.5% of adults worldwide¹. ADHD is often persistent and markedly impairing with increased risk of harmful outcomes such as injuries², traffic accidents³, increased health care utilization^{4,5}, substance abuse⁶, criminality⁷, unemployment⁸, divorce⁴, suicide⁹, AIDS risk behaviors⁸, and premature mortality¹⁰.

Epidemiologic and clinical studies implicate genetic and environmental risk factors that affect the structure and functional capacity of brain networks involved in behavior and cognition¹, in the etiology of ADHD.

Consensus estimates from over 30 twin studies indicate that the heritability of ADHD is 70-80% throughout the lifespan^{11,12} and that environmental risks are those not shared by siblings¹³. Twin studies also suggest that diagnosed ADHD represents the extreme tail of one or more heritable quantitative traits¹⁴. Additionally, family and twin studies report genetic overlap between ADHD and other conditions including antisocial personality disorder/behaviours¹⁵, cognitive impairment¹⁶, autism spectrum disorder^{17,18}, schizophrenia¹⁹, bipolar disorder²⁰, and major depressive disorder²¹.

Thus far genome-wide association studies (GWASs) to identify common DNA variants that increase the risk of ADHD have not been successful²². Nevertheless, genome-wide SNP heritability estimates range from $0.10-0.28^{23,24}$ supporting the notion that common variants comprise a significant fraction of the risk underlying ADHD²⁵ and that with increasing sample size, and thus increasing statistical power, genome-wide significant loci will emerge.

Previous studies have demonstrated that the common variant risk, also referred to as the single nucleotide polymorphism (SNP) heritability, of ADHD is also associated with depression²⁵, conduct problems²⁶, schizophrenia²⁷, continuous measures of ADHD symptoms^{28,29} and other neurodevelopmental traits²⁹ in the population. Genetic studies of quantitative ADHD symptom

scores in children further support the hypothesis that ADHD is the extreme of a quantitative trait³⁰.

Here we present a genome-wide meta-analysis identifying the first genome-wide significant loci for ADHD using a combined sample of 55,374 individuals from an international collaboration. We also strengthen the case that the clinical diagnosis of ADHD is the extreme expression of one or more heritable quantitative traits, at least as it pertains to common variant genetic risk, by integrating our results with previous GWAS of ADHD-related behavior in the general population.

Genome-wide significantly associated ADHD risk loci

Genotype array data for 20,183 ADHD cases and 35,191 controls were collected from 12 cohorts (Supplementary Table 1). These samples included a population-based cohort of 14,584 cases and 22,492 controls from Denmark collected by the Lundbeck Foundation Initiative for Integrative Psychiatric Research (iPSYCH), and 11 European, North American and Chinese cohorts aggregated by the Psychiatric Genomics Consortium (PGC). ADHD cases in iPSYCH were identified from the national Psychiatric Central Research Register psychiatric and diagnosed by psychiatrists at a psychiatric hospital according to ICD10 (F90.0), and genotyped using Illumina PsychChip. Designs for the PGC cohorts has been described previously^{24,25,31,32,22} (see Supplementary Information for detailed cohort descriptions).

Prior to analysis, stringent quality control procedures were performed on the genotyped markers and individuals in each cohort using a standardized pipeline³³ (Online Methods). Related

individuals were removed, and genetic outliers within each cohort were excluded based on principal component analysis. Non-genotyped markers were imputed using the 1000 Genomes Project Phase 3 reference panel³⁴ (Online Methods).

GWAS was conducted in each cohort using logistic regression with the imputed additive genotype dosages. Principal components were included as covariates to correct for population stratification³⁵ (Supplementary Information), and variants with imputation INFO score < 0.8 or minor allele frequency (MAF) < 0.01 were excluded. The GWAS were then meta-analyzed using an inverse-variance weighted fixed effects model³⁶. The single Chinese cohort had insufficient sample size for well-powered trans-ethnic modelling. (Supplementary Figure 7.B). Association results were considered only for variants with an effective sample size greater than 70% of the full meta-analysis, leaving 8,047,421 variants in the final meta-analysis. A meta-analysis restricted to European-ancestry individuals (19,099 cases, 34,194 controls) was also performed to facilitate secondary analyses.

In total, 304 genetic variants in 12 loci surpassed the threshold for genome-wide significance (P<5×10⁻⁸; Figure 1, Table 1, Supplementary Figure 3.A2 – 3.N2). Results for the European ancestry meta-analysis were substantively similar (Supplementary Figure 2). No marker demonstrated significant heterogeneity between studies (Supplementary Figures 6 and 7.A) and no heterogeneity was observed between the Chinese and European ancestry cohorts (Supplementary Figure 7.B). Conditional analysis within each locus did not identify any independent secondary signals meeting genome-wide significance (Online Methods, Supplementary Table 2).

Homogeneity of effects between cohorts

No genome-wide significant heterogeneity was observed in the ADHD GWAS meta-analysis (Supplementary Information). Genetic correlation analysis (Online Methods) provided further evidence that effects were consistent across cohort study designs. The estimated genetic correlation between the European ancestry PGC samples and the iPSYCH sample from LD score regression³⁷ was not significantly less than one ($r_g = 1.17$, SE = 0.20). The correlation between European ancestry PGC case/control and trio cohorts estimated with bivariate GREML was similarly close to one ($r_g = 1.02$, SE = 0.32).

Polygenic risk scores (PRS)³⁸ were also consistent across target samples. PRS computed in each PGC study using iPSYCH as the training sample were consistently higher in ADHD cases as compared to controls or pseudo-controls (Supplementary Figure 11). Increasing deciles of PRS in the PGC were associated with higher odds ratio (OR) for ADHD (Figure 2). A similar pattern was seen in five-fold cross validation in the iPSYCH cohort, with PRS for each subset computed from the other four iPSYCH subsets and the PGC samples used as training samples (Online Methods; Figure 2). Across iPSYCH subsets, the mean of the maximum variance explained by the estimated PRS (Nagelkerke's R²) was 5.5% (SE = 0.0012). The difference in standardized PRS between cases and controls was stable across iPSYCH subsets (OR = 1.56, 95% confidence interval [CI]: 1.53 – 1.60; Supplementary Figure 9). These results further support the highly polygenic architecture of ADHD and demonstrate that ADHD risk is significantly associated with PRS in a dose-dependent manner.

Polygenic Architecture of ADHD

To assess the proportion of phenotypic variance explained by common variants we applied LD score regression³⁷ and results from the European ancestry meta-analysis (Online Methods). Assuming a population prevalence of 5% for ADHD³⁹, we estimate that the liability-scale SNP heritability $h_{snp}^2 = 0.216$ (SE = 0.014, P = 8.18×10⁻⁵⁴). These estimated polygenic effects account for 88% (SE = 0.0335) of observed genome-wide inflation of the test statistics in the meta-analysis ($\lambda = 1.200$); the remaining inflation, which may reflect confounding factors such as cryptic relatedness and population stratification, is significant but modest (intercept=1.0362, SE = 0.0099, P=2.27 × 10⁻⁴).

To further characterize the patterns of heritability from the genome-wide association data, we partitioned SNP heritability by functional annotations as described in Finucane et al.⁴⁰ using partitioned LD Score regression (Online Methods). The analysis revealed significant enrichment in the heritability from SNPs located in conserved regions ($P = 8.49 \times 10^{-10}$; Supplementary Figure 12), supporting their biological importance. Enrichment of the SNP heritability in cell-type-specific regulatory elements was evaluated using the cell-type-specific group annotations described in Finucane et al⁴⁰. We observed a significant enrichment of the average per SNP heritability for variants located in central nervous system specific regulatory elements (enrichment = 2.44, SE = 0.35, $P = 5.81 \times 10^{-5}$; Supplementary Figures 13 and 14).

Genetic correlation with other traits

Pairwise genetic correlation with ADHD was estimated for 219 phenotypes using LD score regression^{41,42} (Online Methods, Supplementary eTable 5). Fourty-three phenotypes

demonstrated significant genetic overlap with ADHD (P < 2.28 × 10⁻⁴), including major disorder⁴³, anorexia nervosa⁴⁴, educational outcomes⁴⁵⁻⁴⁹, obesity-related depressive phenotypes⁵⁰⁻⁵⁵, smoking⁵⁶⁻⁵⁸, reproductive success⁵⁹, insomnia⁶⁰, and mortality⁶¹ (Figure 3; Supplementary Table 11). In most domains the genetic correlation is supported by GWAS of multiple related phenotypes. For the positive genetic correlation with major depressive disorder $(r_g = 0.42, P = 7.38 \times 10^{-38})$, we also observe a positive correlation with depressive symptoms $(r_g = 0.42, P = 7.38 \times 10^{-38})$ = 0.45, P = 7.00×10^{-19}), neuroticism ($r_g = 0.26$, P= 1.02×10^{-8}) and a negative correlation with subjective well-being ($r_g = -0.28$, $P = 3.73 \times 10^{-9}$). The positive genetic correlations with ever smoked (r_g = 0.48, P= 4.33 \times 10⁻¹⁶) and with number of cigarettes smoked (r_g = 0.45, P = 1.07 \times 10^{-5}) are reinforced by significant positive correlation with lung cancer ($r_g = 0.39$, $P = 6.35 \times 10^{-5}$ ¹⁰). Similarly, genetic correlations related to obesity include significant relationships with body mass index (BMI; $r_g = 0.26$, $P = 1.68 \times 10^{-15}$), waist-to-hip ratio ($r_g = 0.30$, $P = 1.16 \times 10^{-17}$), childhood obesity (r_g = 0.22, P = 3.29 × 10⁻⁶), HDL cholesterol (r_g = -0.22, P = 2.44 × 10⁻⁷), and Type 2 Diabetes ($r_g = 0.18, \, P = 7.80 \times 10^{-5}$). Additionally the negative correlation with years of schooling ($r_g = -0.53$, $P = 6.02 \times 10^{-80}$) is supported by a negative genetic correlation with human intelligence (r_g = -0.41, P = 7.03 \times 10⁻²⁶). Finally the genetic correlation with reproduction include a negative correlation with age of first birth (r_g = -0.612, P = 3.70 × 10⁻⁶¹) and a positive correlation with number of children ever born ($r_g = 0.42$, $P = 8.51 \times 10^{-17}$).

Biological annotation of significant loci

For the 12 genome-wide significant loci, Bayesian credible sets were defined to identify the set of variants at each locus most likely to include a variant with causal effect (Online Methods, Supplementary eTable 1). Biological annotations of the variants in the credible set were then

considered to identify functional or regulatory variants, common chromatin marks, and variants associated with gene expression (eQTLs) or in regions with gene interactions observed in Hi-C data (Online Methods, Supplementary eTable 2). Broadly, the significant loci do not coincide with candidate genes proposed to play a role in ADHD⁶².

Here we highlight genes that are identified in the regions of association (see also Supplementary Table 4). The loci on chromosomes 2, 7, and 10 each have credible sets localized to a single gene with limited additional annotations. In the chromosome 7 locus, *FOXP2* encodes a forkhead/winged-helix transcription factor and is known to play an important role in synapse formation and neural mechanisms mediating the development of speech and learning ⁶³⁻⁶⁵. Comorbidity of ADHD with specific developmental disorders of language and learning is common (7 – 11%)^{66,67}, and poor language skills have been associated with higher inattention/hyperactivity symptoms in primary school⁶⁸. On chromosome 10, the ADHD association is located intronic in *SORCS3*, which encodes a brain-expressed transmembrane receptor that is important for neuronal development and plasticity⁶⁹ and has previously been associated with depression^{43,70} and schizophrenia.

Genome-wide significant loci on chromosomes 12 and 15 have more biological annotations supporting the co-localized genes. The credible set on chromosome 12 spans *DUSP6*, and includes an annotated missense variant in the first exon and an insertion near the transcription start site, though neither is the lead variant in the locus (Supplementary eTable 3). *DUSP6* encodes a dual specificity phosphatase⁷¹, and may play a role in regulating neurotransmitter homeostasis by affecting dopamine levels in the synapses^{72,73}. Regulation of dopamine levels is

likely to be relevant to ADHD since widely used ADHD medications have dopaminergic targets^{74,75} that increase the availability of synaptic dopamine. The chromosome 15 locus is located in *SEMA6D*, and the majority of variants in the credible set are strongly associated with expression of *SEMA6D* in fibroblasts⁷⁶. *SEMA6D* is active in the brain during embryonic development, and may play a role in neuronal wiring⁷⁷. Furthermore, variants in *SEMA6D* have previously been associated with eduational attainment⁷⁸.

Credible set annotations at the remaining loci are more diverse (Supplementary eTable 2). The most strongly associated locus on chromosome 1 (index variant rs112984125) covers a gene-rich 250kb region of strong LD. The index variant is intronic to *ST3GAL3*, and most SNPs in the credible set are strongly associated with expression of *ST3GAL3* in whole blood⁷⁹ (Supplementary eTable 2). Missense mutations in *ST3GAL3* have been shown to cause autosomal recessive intellectual disability⁸⁰. Hi-C and eQTL annotations suggest multiple alternative genes however, including *PTPRF* (Supplementary eTable 3). The locus also includes an intergenic variant, rs11210892, that has previously been associated with schizophrenia³³.

On chromosome 5, the credible set includes links to *LINC00461* and *TMEM161B* (Supplementary eTable 2). The function of *LINC00461* is unclear, but the RNA has highly localized expression in the brain⁸¹ and the genome-wide significant locus overlaps with variants in *LINC00461* associated with educational attainment⁷⁸. Alternatively, a genome-wide significant SNP in this locus (rs304132) is located in *MEF2C-AS1*, of strong interest given previous associations between *MEF2C* and severe intellectual disability,⁸²⁻⁸⁴ cerebral malformation⁸³, depression⁷⁰, schizophrenia³³ and Alzheimer's disease⁸⁵, but the corresponding

variant is not supported by the credible set analysis. Credible set annotations for other significant loci are similarly cryptic.

Analysis of gene sets

Competitive gene based tests were performed for *FOXP2* target genes, highly constrained genes, and for all Gene Ontology terms⁸⁶ from MsigDB 6.0⁸⁷ using MAGMA⁸⁸ (Online Methods). Association results for individual genes are consistent with the genome-wide significant loci for the GWAS (Supplementary Table 5). Three independent sets of *FOXP2* downstream target genes^{89,90} were tested (Online Methods), none of which demonstrated significant association to ADHD (Supplementary Table 7). The lack of association may be caused by unknown functions of *FOXP2* driving ADHD risk, insufficient power to detect relevant downstream genes, or because only a small subset of biological functions regulated by FOXP2 are relevant to ADHD pathogenesis.

Consistent with the partitioning of heritability, a set of 2,932 genes that are highly constrained and show high intolerance to loss of function⁹¹ showed significant association with ADHD (β = 0.062, P = 2.6 × 10⁻⁴). We also find little evidence for effects in previously proposed candidate genes for ADHD⁶²; of the nine proposed genes only *SLC9A9* showed weak association with ADHD (P = 3.4 × 10⁻⁴; Supplementary Table 6). None of the Gene Ontology gene sets were significant after correcting for multiple testing, although the most associated included interesting nominally significant pathways such as "dopamine receptor binding" (P = 0.0010) and "Excitatory Synapse" (P = 0.0088; Supplementary eTable 4).

Replication of GWAS loci

For replication we evaluated the comparison of the GWAS meta-analysis of ADHD with three other independent ADHD-related GWASs: replication of top loci in an Icelandic cohort with ADHD status derived from medical records of ICD codes and medication history by deCODE (5,085 cases, 131,122 controls), a GWAS of self-reported ADHD status among 23andMe research participants (5,857 cases, 70,393 controls) and a meta-analysis of GWAS of childhood rating scales of ADHD symptoms performed by the EAGLE consortium (17,666 children < 13 years of age)³⁰ and QIMR⁹² (2,798 adolescents). Although the phenotyping and cohort ascertainment of the 23andMe and EAGLE/QIMR studies differ from the PGC and iPSYCH ADHD meta-analysis (Supplementary Information), they have clear relevance to understanding how the ADHD GWAS results generalize to closely related phenotypes.

Top loci from the ADHD GWAS showed moderate concordance across the three replication studies. Sign concordance between each of the three replication cohorts and the ADHD GWAS was significantly greater than would be expected by chance (range 72–82% concordant; P < 0.0167 = 0.05/3 replication cohorts; Supplementary Table 12) for nominally associated loci from the ADHD GWAS ($P < 1 \times 10^{-6}$), with the highest concordance observed in EAGLE/QIMR. The deCODE and 23andMe results also permit direct comparisons of the magnitude of effect sizes for the top loci in the ADHD loci (Supplementary Table 13). Regressing effect size estimates from each replication cohort on estimates from the ADHD GWAS adjusted for winner's curse yields significantly positive slopes (deCODE slope = 0.664, $P = 1.2 \times 10^{-4}$; 23andMe slope = 0.417, $P = 1.11 \times 10^{-3}$), although these slopes are less than one, suggesting imperfect replication. Among the genome-wide significant loci, rs9677504 (*SPAG16* locus) in deCODE and

rs112984125 (ST3GAL3/PTPRF locus) and rs212178 (LINC01572 locus) in 23andMe are noteable outlers with weak replication results (Online Methods, Supplementary Figures 15-16).

The genome-wide data available from 23andMe and EAGLE/QIMR showed similar trends for replication. The genetic correlation between EAGLE/QIMR and the ADHD GWAS was extremely strong ($r_g = 0.970$, SE = 0.207, P = 2.66 \times 10⁻⁶) and not significantly different from one (one-sided P = 0.442). Genetic correlation with 23andMe was weaker but still strongly positive ($r_g = 0.653$, SE = 0.114, P = 1.11 × 10⁻⁸), although also significantly less than 1 (onesided P= 1.17×10^{-3}). To explore this lower correlation we evaluated the genetic correlation between 23andMe and traits from LD Hub (http://ldsc.broadinstitute.org/ldhub/)⁴² to potentially identify differences in the profile of genetic correlation compared to the ADHD GWAS (Online Methods). This comparison identified striking differences (Supplementary Table 14), most notably from the 23andMe GWAS showing little to no genetic correlation with college completion ($r_g = 0.056$, compared to $r_g = -0.54$ for the primary ADHD GWAS; approximate P = 1.1×10^{-9} for difference) and other education-related phenotypes. Genetic correlations with obesity-related phenotypes were similarly smaller for the 23andMe cohort. The one domain where 23 and Me exhibited a trend toward stronger genetic correlations was schizophrenia ($r_g =$ 0.27, vs. $r_g = 0.12$ in ADHD, P = 0.053) and bipolar disorder ($r_g = 0.029$, vs. $r_g = 0.095$ in ADHD, P = 0.09), though these trends are not significant with the approximated test of the difference in genetic correlation.

Finally, we meta-analyzed the ADHD GWAS with each replication cohort. For EAGLE/QIMR, we developed a novel model to meta-analyze the GWAS of the continuous measure of ADHD

with the clinical diagnosis in the ADHD GWAS. In brief, we perform a Z-score based metaanalysis using a weighting scheme derived from the SNP heritability and effective sample size for each phenotype that fully accounts for the differences in measurement scale (detailed description in Supplementary Information). This calibration based on the genome-wide estimate of heritability prevents joint meta-analysis of all replication cohorts since genome-wide data is not available for the deCODE study.

Meta-analyses of the ADHD GWAS with each replication identified 10 genome-wide significant loci (P < 5 × 10-8, without multiple testing correction) in meta-analysis with deCODE, 10 significant loci with 23andMe, and 15 significant loci with EAGLE/QIMR (Supplementary eTable 6, Supplementary Figures 17 and 20). Of the 12 significant loci from the primary ADHD GWAS, 4 were significant in all three of these replication meta-analyses: index variants rs11420276 (ST3GAL3/PTPRF), rs5886709 (FOXP2), rs11591402 (SORCS3), and rs1427829 (intergenic). The remaining loci were all significant in at least one of the replication meta-analyses. In addition, ten novel loci reach genome-wide significance in the replication meta-analyses, of which three loci are significant in two of these analyses (Supplementary eTable 6): index variants rs1592757 / rs30266 (Refseq LOC105379109), rs28452470 / rs1443749 (CADPS2), and rs2243638 / rs9574218 (RNF219-AS1). The CADPS2 locus has recently been identified in autism spectrum disorder as a novel locus shared with educational attainment 93.

Meta-analysis with the 23andMe cohort also found genome-wide significant heterogeneity at the lead chromosome one locus from the ADHD GWAS meta-analysis (rs12410155: $I^2 = 97.2$, $P = 2.29 \times 10^{-9}$; Supplementary Figures 18-19). This heterogeneity is consistent with the moderate

sign concordance, effect size replication, and genetic correlation of the 23andMe cohort with the ADHD GWAS. Notably, the lead chromosome 1 locus in the ADHD GWAS overlaps a reported association with educational attainment⁷⁸, suggesting this heterogeneity is consistent with the much weaker genetic correlation between the 23andMe results and published GWAS of education-related outcomes. No genome-wide significant heterogeneity was observed in the replication meta-analyses with deCODE or EAGLE/QIMR (Supplementary Figures 21-22, Supplementary eTable 6).

Discussion

GWAS meta-analysis of ADHD revealed the first genome-wide significant risk loci, and indicates an important role for common variants in the polygenic architecture of ADHD. Several of the loci are located in or near genes that implicate neurodevelopmental processes that are likely to be relevant to ADHD, including *FOXP2*, *SORCS3*, and *DUSP6*. Future work may focus on refining the source of the strong association in each locus, especially the lead locus on chromosome 1 which is complicated by broad LD and substantial heterogeneity between ADHD meta-analysis and analysis of self-reported ADHD status in 23andMe.

The 12 significant loci are compelling, but only capture a tiny fraction of common variant risk for ADHD. The odds ratios for the risk increasing allele at the index SNPs in the 12 significant loci are modest, ranging from 1.077 to 1.198 (Table 1). This is within the range of effect sizes for common genetic variants that has been observed for other highly polygenic psychiatric disorders e.g. schizophrenia³³. A considerably larger proportion of the heritability of ADHD can be explained by all common variants ($h^2_{snp} = 0.22$, SE = 0.01). This is consistent with previous

estimates of h^2_{snp} for ADHD in smaller studies (h^2_{snp} : 0.1 - 0.28) 23,24 , and also comparable to SNP heritability estimates for schizophrenia (h^2_{snp} 0.23 - 0.26) 23,24 . As would be hypothesized for a psychiatric disorder, these effects are enriched in conserved regions and regions containing enhancers and promoters of expression in the central nervous system tissues, consistent with previous observations in schizophrenia and bipolar disorder⁴⁰. On the other hand, we do not observe substantial effects in most previously reported candidate genes for ADHD⁶².

Along with polygenicity, selection and evolutionary pressures may be an important feature of the architecture of ADHD genetics. We observe that ADHD risk variants are strongly enriched in genomic regions conserved in mammals⁹⁴, and constrained genes likely to be intolerant of loss-of-function mutations⁹¹ are associated with ADHD. We also find that common variant risk for ADHD is genetically correlated with having children younger and having more children, in line with epidemiological findings of increased risky sexual behaviour⁹⁵⁻⁹⁷ and increased risk of ADHD for children born to young parents⁹⁸⁻¹⁰⁰. Given the phenotypic^{101,102} and genetic¹⁰³ correlation of ADHD with reduced educational attainment, positive selective pressure on the genetics of ADHD would be consistent with recent work suggesting that variants associated with educational attainment are under negative selection in Iceland¹⁰⁴. Future studies of fecundity and the role of rare and *de novo* variants in ADHD may provide more insight on selective pressures in ADHD-associated loci.

The observed genetic correlations with educational outcomes and other phenotypes suggest a strong genetic component to the epidemiological correlates of ADHD. The significant positive genetic correlation of ADHD with major depressive disorder and depressive symptoms supports

previous findings suggesting a positive genetic overlap between those phenotypes^{24,42}, as well as the broader genetic overlap of psychiatric disorders^{23,24}. Positive genetic correlations between ADHD and health risk behaviors such as smoking and obesity are consistent with the observed increase in those behaviors among individuals with ADHD¹⁰⁵⁻¹⁰⁸ and are indicative of a shared genetic basis for these traits. We also observe a positive genetic correlation of ADHD with insomnia, consistent with reports of sleep disturbances in ADHD¹⁰⁹, but this relationship does not appear to generalize to other sleep-related phenotypes.

These genetic correlations may not generalize to all settings. We observe much weaker genetic correlation of the 23andMe ADHD results with educational attainment, with only partial genetic correlation between 23andMe and the current ADHD GWAS, including significant heterogeneity in the lead chromosome 1 locus. The pattern of replication for the top loci in the deCODE study is stronger but still mixed. These differences may reflect differences in phenotyping (e.g. self-report vs. medical records), exclusion of individuals with comorbid psychiatric disorders (deCODE), study population (e.g. higher average education and socio-economic status among 23andMe research participants possibly under-representing the proportion of individuals with ADHD with poor educational outcomes in the general population), or other study factors that should be a focus of future work.

On the other hand, the replication results from the EAGLE³⁰/QIMR⁹² are much stronger and support the hypothesis that ADHD is the extreme expression of one or more heritable quantitative traits¹¹⁰. We observe strong concordance between the GWAS of ADHD and the previous GWASs of ADHD-related traits in the population, both in terms of genome-wide

genetic correlation and concordance at individual loci. Polygenic risk for ADHD has previously been associated with inattentive and hyperactive/impulsive trait variation below clinical thresholds in the population²⁹. Shared genetic risk with health risk behaviors may similarly be hypothesized to reflect an impaired ability to self-regulate and inhibit impulsive behavior^{111,112}. The observed negative correlation between ADHD and anorexia nervosa may also be related to these behavioral factors.

In summary, we report 12 independent genome-wide significant loci associated with ADHD in GWAS meta-analysis of 55,374 individuals from 12 study cohorts. The GWAS meta-analysis implicates *FOXP2* and other biologically informative genes as well as constrained regions of the genome as important contributors to the etiology of ADHD. The results also highlight strong overlap with the genetics of ADHD-related traits and health risk behaviors in the population, encouraging a dimensional view of ADHD as the extreme end of a continuum of symptoms.

Online Methods

GWAS meta-analysis

Quality control, imputation and primary association analyses were done using the bioinformatics pipeline Ricopili (available at https://github.com/Nealelab/ricopili), developed by the Psychiatric Genomics Consortium (PGC)³³. In order to avoid potential study effects the 11 PGC samples and the 23 genotyping batches within iPSYCH were each processed separately unless otherwise stated (Supplementary Information).

Stringent quality control was applied to each cohort following standard procedures for GWAS, including filters for call rate, Hardy-Weinberg equilibrium, and heterozygosity rates (Supplementary Information). Each cohort was then phased and imputed using the 1000 Genomes Project phase 3 (1KGP3)^{34,113} imputation reference panel using SHAPEIT¹¹⁴ and IMPUTE2¹¹⁵, respectively. For trio cohorts, pseudocontrols were defined from phased haplotypes prior to imputation.

Cryptic relatedness and population structure were evaluated using a set of high quality markers pruned for linkage disequilibrium (LD). Genetic relatedness was estimated using PLINK v1.9^{116,117} to identify first and second-degree relatives ($\hat{\pi} > 0.2$) and one individual was excluded from each related pair. Genetic outliers were identified for exclusion based on principal component analyses using EIGENSOFT^{35,118}. This was done separately for each of the PGC cohorts and on a merged set of genotypes for the iPSYCH cohort (Supplementary Information). Across studies, a total of 20,183 cases and 35,191 controls remained for analysis after QC.

Genome-wide association analyses for the 11 PGC samples and the 23 waves in iPSYCH were performed using logistic regression model with the imputed marker dosages in PLINK v1.9^{116,117}. Principal components were included as covariates to control for population stratification^{35,118}, along with relevant study-specific covariates where applicable (Supplementary Information, Supplementary Table 1). Subsequently the results were meta-analysed using an inverse-variance weighted fixed effects model, implemented in METAL (version 2011-03-25)³⁶. Variants were filtered and included if imputation quality (INFO score) was > 0.8 and MAF > 0.01. Only markers supported by an effective sample size $N_{\rm eff} = 4/(1/N_{\rm cases} + 1/N_{\rm controls})^{119}$ greater than 70% were included. After filtering, the meta-analysis included results for 8,047,421 markers.

Conditional analysis

Twelve independent genome-wide significant loci were identified by LD clumping and merging loci within 400 kb (Supplementary Information). In two of these loci a second index variant persisted after LD clumping. The two putative secondary signals were evaluated by considering analysis conditional on the lead index variant in each locus. In each cohort, logistic regression was performed with the imputed genotype dosage for the lead index variant included as a covariate. All covariates from the primary GWAS (e.g. principal components) were also included. The conditional association results were then combined in an inverse-variance weighted meta-analysis.

Genetic correlations between ADHD samples

Genetic correlation between the European-ancestry PGC and iPSYCH GWAS results was calculated using LD Score regression³⁷. The regression was performed using pre-computed LD scores for HapMap3 SNPs calculated based on 378 European-ancestry individuals from the 1000 Genomes Project (available on https://github.com/bulik/ldsc). Only results for markers with an imputation INFO score > 0.90 were included in the analysis. In addition, a bivariate GREML analysis was conducted using GCTA¹²⁰ in order to estimate the genetic correlation between PGC case/control and trio study designs.

Polygenic Risk Scores for ADHD

The iPSYCH sample were split into five groups, and subsequently five leave-one-out association analyses were conducted, using four out of five groups and the PGC samples as training

datasets³⁸. PRS were estimated for each target sample using variants passing a range of association P-value thresholds in the training samples. PRS were calculated by multiplying the natural log of the odds ratio of each variant by the allele-dosage (imputation probability) and whole-genome polygenic risk scores were obtained by summing values over variants for each individual.

For each of the five groups of target samples, PRS were normalized and the significance of the case-control score difference was tested by standard logistic regression including principal components. For each target group and for each P-value threshold the proportion of variance explained (i.e. Nagelkerke's R^2) was estimated by comparing the regression with PRS to a reduced model with covariates only. The OR for ADHD within each PRS decile group was estimated based on the normalized score across groups (using the P-value threshold with the highest Nagelkerke's R^2 within each target group) (Figure 3). OR was also estimated using logistic regression on the continuous scores for each target group separately and an OR based on all samples using the normalized PRS score across all groups (Supplementary Figure 9). Additionally PRS were evaluated in the PGC samples using the iPSYCH sample as training sample, following the approach described above (Supplementary Information).

SNP heritability and intercept evaluation

LD score regression³⁷ was used to evaluated the relative contribution of polygenic effects and confounding factors, such as cryptic relatedness and population stratification, to deviation from the null in the genome-wide distribution of GWAS χ^2 statistics. Analysis was performed using pre-computed LD scores from European-ancestry samples in the 1000 Genomes Project (available on https://github.com/bulik/ldsc) and summary statistics for the European-ancestry

ADHD GWAS to ensure matching of population LD structure. The influence of confounding factors was tested by comparing the estimated intercept of the LD score regression to one, it's expected value under the null hypothesis of no confounding from e.g. population stratification. The ratio between this deviation and the deviation of the mean χ^2 from one (i.e. it's expected value under the null hypothesis of no association) was used to estimate the proportion of inflation in χ^2 attributable to confounding as opposed to true polygenic effects (ratio = (intercept-1)/(mean χ^2 -1)). SNP heritability was estimated based on the slope of the LD score regression, with heritability on the liability scale calculated assuming a 5% population prevalence of ADHD³⁹.

Partitioning of the heritability

SNP heritability was partitioned by functional category and tissue association using LD score regression⁴⁰. Partitioning was performed for 53 overlapping functional categories, as well as 220 cell-type-specific annotations grouped into 10 cell-type groups, as described in Finucane et al. ⁴⁰. For both sets of annotations we used previously computed LD scores and allele frequencies from European ancestry samples in the 1000 Genomes Project (available on https://data.broadinstitute.org/alkesgroup/LDSCORE/).

Additionally we expanded the cell-type specific heritability analysis by including an annotation based on information about H3K4Me1 imputed gapped peaks excluding the broad MHC-region (chr6:25-35MB), generated by the Roadmap Epigenomics Mapping Consortium^{121,122} (Supplementary Information). The analyses were restricted to the European GWAS meta-analysis results to ensure matching of population LD structure. Results for each functional category were evaluated based on marginal enrichment, defined as the proportion of SNP

heritability explained by SNPs in the annotation divided by the proportion of genome-wide SNPs in the annotation⁴⁰. For each cell-type group and each H3K4Me1 cell-type annotations, the contribution to SNP heritability was tested conditional on the baseline model containing the 53 functional categories.

Genetic correlations of ADHD with other traits

The genetic correlations of ADHD with other phenotypes were evaluated using LD Score regression⁴². For a given pair of traits, LD score regession estimates the expected population correlation between the best possible linear SNP-based predictor for each trait, restricting to common SNPs. Such correlation of genetic risk may reflect a combination of colocalization, pleiotropy, shared biological mechanisms, and causal relationships between traits. Correlations were tested for 211 phenotypes with publically available GWAS summary statistics using LD Hub⁴¹ (Supplementary Information). Additionally, we analysed on our local computer cluster, the genetic correlation of ADHD with eight phenotypes: human intelligence 103, four phenotypes related to education and cognition analyzed in samples from the UK Biobank⁴⁹ (college/university degree, verbal-numerical reasoning, memory and reaction time), insomnia⁶⁰, anorexia nervosa⁴⁴, and major depressive disorder⁴³. The genetic correlation with major depressive disorder was tested using GWAS results from an updated analysis of 130,664 cases with major depressive disorder and 330,470 controls from the Psychiatric Genomics Consortium. As in the previous LD score regression analyses, this estimation was based on summary statistics from the European GWAS meta-analysis, and significant correlations reported are for traits analysed using individuals with European ancestry.

Credible set analysis

We defined a credible set of variants in each locus using the method described by Maller et al. 123 (Supplementary Information), implemented by freely available R script (https://github.com/hailianghuang/FM-summary). Under the assumption that (a) there is one causal variant in each locus, and (b) the causal variant is observed in the genotype data, the credible set can be considered to have a 99% probability of containing the causal variant. For each the 12 genome-wide significant loci, variants within 1MB and in LD with correlation $r^2 > 1$ 0.4 to the index variant were considered for inclusion in the credible set analysis. The credible set analysis was done using the European GWAS meta-analysis to ensure consistent LD structure in the analyzed cohorts.

Biological annotation of variants in credible set

The variants in the credible set for each locus, were annotated based on external reference data in order to evaluate potential functional consequences. In particular, we identify: (a) Gene and regulatory consequences annotated by Variant Effect Predictor (VEP) using Ensembl with genome build GRCh37¹²⁴. We exclude upstream and downstream consequences, and consequences for transcripts that lack a HGNC gene symbol (e.g. vega genes). (b) Variants within 2kb upstream of the transcription start site (TSS) of at least one gene isoform based on Gencode v19¹²⁵. (c) Variants annotated as interacting with a given gene in Hi-C data from samples of developing human cerebral cortex during neurogenesis and migration ¹²⁶. Annotations are considered for both the germinal zone (GZ), primarily consisting of actively dividing neural progenitors, and the cortical and subcortical plate (CP), primarily consisting of post-mitotic neurons. (d) Variants identified as eQTLs based on gene expression in GTEx ¹²⁷ or BIOS⁷⁹.

Expression quantitative trait loci were annotated using FUMA (http://fuma.ctglab.nl/). We restricted to eQTL associations with false discovery fate (FDR) < 1e-3 within each dataset. (e) Chromatin states of each variant based on the 15-state chromHMM analysis of epigenomics data from Roadmap¹²⁸. The 15 states summarize to annotations of active chromatin marks (i.e. Active TSS, Flanking Active TSS, Flanking Transcription, Strong Transcription, Weak Transcription, Genic Enhancer, Enhancer, or Zinc Finger [ZNF] gene), repressed chromatin marks (Heterochromatin, Bivalent TSS, Flanking Bivalent TSS, Bivalent Enhancer, Repressed Polycomb, or Weak Repressed Polycomb), or quiescent. The most common chromatin state across 127 tissue/cell types was annotated using FUMA (http://fuma.ctglab.nl/). We also evaluated the annotated chromatin state from fetal brain.

Gene-set analyses

Gene-based association with ADHD was estimated with MAGMA 1.05⁸⁸ using the summary statistics from the European GWAS meta-analysis (N_{cases} = 19,099, N_{controls} = 34,194; Supplementary Information, Supplementary Information Table 1). Association was tested using the SNP-wise mean model, in which the sum of -log(SNP P-value) for SNPs located within the transcribed region (defined using NCBI 37.3 gene definitions) was used as the test statistic. MAGMA accounts for gene-size, number of SNPs in a gene and LD between markers when estimating gene-based P-values. LD correction was based on estimates from the 1000 genome phase 3 European ancestry samples³⁴.

The generated gene-based P-values were used to analyze sets of genes in order to test for enrichment of association signals in genes belonging to specific biological pathways or processes. In the analysis only genes on autosomes, and genes located outside the broad MHC region (hg19:chr6:25-35M) were included. We used the gene names and locations and the European genotype reference panel provided with MAGMA. For gene sets we used sets with 10-1000 genes from the Gene Ontology sets⁸⁶ currated from MsigDB 6.0⁸⁷.

Targeted *FOXP2* downstream target gene sets were analysed for association with ADHD. Three sets were examined: 1) Putative target genes of *Foxp2* that were enriched in wild type compared to control *Foxp2* knockout mouse brains in ChIP-chip experiments (219 genes), 2) Genes showing differential expression in wild type compared to *Foxp2* knockout mouse brains (243 genes), and 3) *FOXP2* target genes that were enriched in either or both basal ganglia (BG) and inferior frontal cortex (IFC) from human fetal brain samples in ChIP-chip experiments (258 genes). Curated short lists of high-confidence genes were obtained from Vernes et al.⁸⁹ and Spiteri et al⁹⁰.

A set of evolutionarily highly constrained genes were also analysed. The set of highly constrained genes was defined using a posterior probability of being loss-of-function intolerant (pLI) based on the observed and expected counts of protein-truncating variants (PTV) within each gene in a large study of over 60,000 exomes from the Exome Aggregation Consortium (ExAC)⁹¹. Genes with pLI ≥0.9 were selected as the set of highly constrained genes (2932 genes).

Replication of GWAS loci

To replicate the results of the ADHD GWAS meta-analysis we compared the results to analyses from deCODE, 23andMe, and EAGLE/QIMR. We evaluated evidence for replication based on:

(a) sign tests of concordance between the ADHD GWAS meta-analysis and each replication cohort; (b) comparison of bias-corrected effect sizes between the ADHD GWAS and the

deCODE and 23andMe replication cohorts; (c) genetic correlation between the ADHD GWAS and the 23andMe and EAGLE/QIMR replication cohorts; (d) meta-analysis of the ADHD GWAS meta-analysis results with the results from each replication cohort; and (e) tests of heterogeneity between the ADHD GWAS and each replication cohorts.

For the sign test, we first identified the overlapping SNPs present in the ADHD GWAS and each of the three replication analyses (i.e. deCODE, 23andMe, and EAGLE/QIMR). For each replication cohort intersecting SNPs were then clumped for LD ($r^2 > 0.05$ within 1 Mb) for all variants with P < 1 × 10⁻⁴ in the ADHD GWAS (or P < 1 × 10⁻⁵ for the deCODE replication) using 1000 Genomes Phase 3 data on European ancestry populations. After clumping, sign tests were performed to test the proportion of loci with a concordant direction of effect in the replication cohort (π) using a one sample test of the proportion with Yates' continuity correction¹²⁹ against a null hypothesis of π = 0.50 (i.e. the signs are concordant between the two analyses by chance) in R¹³⁰. This test was evaluated separately for concordance in deCODE, 23andMe, and EAGLE/QIMR for loci passing P-value thresholds of P < 5 × 10⁻⁸ (i.e. genomewide significant loci), P < 1 × 10⁻⁷, P < 1 × 10⁻⁶, P < 1 × 10⁻⁵, and P < 1 × 10⁻⁴ in the ADHD GWAS meta-analysis (Supplementary Information).

In addition to testing concordance for the direction of effect, we also evaluate replication for the magnitude of the effect sizes. Specifically, for each of deCODE and 23andMe we regressed the effect size in the replication cohort (i.e. the log odds ratio) on the estimated effect size from the ADHD GWAS after adjustment for winner's curse for loci with P < 1e-6. Winner's curse correction is performed by computing posterior mean estimates of marginal SNP effects β_j after fitting a spike-and-alab distribution

$$\beta_j \sim \begin{cases} 0 & \text{with probability } \pi \\ N(0, \tau^2) & \text{otherwise} \end{cases}$$

by maximum likelihood as described by Okbay et al.⁷⁸ (Supplementary Information). For the regression of effect sizes we oriented all variants in the direction of the risk increasing allele estimated from the ADHD GWAS, constrained the intercept to zero, and weighted the variants proportional to the inverse of their squared standard error from the ADHD GWAS. A regression slope of one indicates "ideal" replication of all loci in the regression, whereas a slope of zero indicates no replication.

Genetic correlation of the ADHD GWAS with the 23andMe and EAGLE/QIMR results was computed using LD score regression³⁷ with pre-computed European ancestry LD scores following the same procedure as described above for other genetic correlation analyses. Genetic correlation could not be computed for deCODE since results were only available for top loci from the ADHD GWAS. To further explore the moderate genetic correlation between the 23andMe results and the ADHD GWAS we also evaluated the genetic correlation between 23andMe and traits from LD Hub (http://ldsc.broadinstitute.org/ldhub/)⁴². To evaluate the magnitude of the observed differences in r_g we consider both the absolute difference (i.e. $|r_{g,ADHD} - r_{g,23andMe}|$) and the test of an approximate Z score for this difference (Supplementary Information):

$$Z = \frac{r_{g,ADHD} - r_{g,23andMe}}{\sqrt{SE_{ADHD}^2 + SE_{23andMe}^2}}$$

We do not expect this to be an ideal formal test for the difference between two genetic correlations, and therefore emphasize caution in interpreting the precise results. Nevertheless, it does offer a useful benchmark for evaluating the magnitude of the difference between the r_g estimates in the context of the uncertainty in those values.

Finally, we meta-analyzed the ADHD GWAS with the results from each replication cohort. For deCODE and 23andMe inverse variance-weighted meta-analyses were performed. For meta-

analysis with the EAGLE/QIMR GWAS of ADHD-related behaviors in childhood population samples we used a modified sample size-based weighting method. Modified sample size-based weights were derived to accounts for the respective heritabilities, genetic correlation, and measurement scale of the GWASs (Supplementary Information). To summarize, given z-scores Z_{lj} and Z_{2j} resulting from GWAS of SNP j in a dichotomous phenotype (e.g. ADHD) with sample size N_l and a continuous phenotype (e.g. ADHD-related traits) with sample size N_2 , respectively, we calculate

$$Z_{j,meta} = \frac{\sqrt{\widetilde{N}_{1j}}Z_{1j} + \sqrt{\widetilde{N}_{2j}}\widetilde{Z}_{2j}}{\sqrt{\widetilde{N}_{1j} + \widetilde{N}_{2j}}}$$

where

$$\begin{split} \widetilde{Z}_{2j} &= sign(r_g) \times \frac{Z_{2j}}{\sqrt{1 + (1 - r_g^2)N_{2j}h_2^2 l_j/M}} \\ \widetilde{N}_{1j} &= N_{1j} \frac{P(1 - P) \phi(\Phi^{-1}[K])^2}{[K(1 - K)]^2} \\ \widetilde{N}_{2j} &= N_{2j} \frac{r_g^2 h_2^2/h_1^2}{1 + (1 - r_g^2) N_{2j} h_2^2 l_j/M} \end{split}$$

The adjusted sample sizes $\widetilde{N_1}$ and $\widetilde{N_2}$ reflect differences in power between the studies due to measurement scale and relative heritability that is not captured by sample size. The calculation of $\widetilde{Z_2}$ reduces the contribution of the continuous phenotype's GWAS to the meta-analysis based on imperfect genetic correlation with the dichotomous phenotype of interest (i.e. ADHD). The adjustments are computed based on the sample prevalence (P) and population prevalence (K) of the dichotomous phenotype, the estimated liability scale SNP heritability of the two phenotypes

 $(h_1^2 \text{ and } h_2^2)$, and the genetic correlation (r_g) between the two phenotypes, as well as the average SNP LD score (l_j) and the number of SNPs (M). Heritability and genetic correlation values to compute these weights are computed using LD score regression. This meta-analysis weighting scheme is consistent with weights alternatively derived based on modelling the joint distribution of marginal GWAS beta across traits¹³¹.

To test heterogeneity with each replication cohort, we considered Cochran's Q test of heterogeneity in the meta-analyses. Specifically, we evaluated the one degree of freedom test for heterogeneity between the ADHD GWAS meta-analysis and the replication cohort.

Availability of results

The PGC's policy is to make genome-wide summary results public. Summary statistics with the results from the ADHD GWAs meta-analysis of iPSYCH and the PGC samples are available on the PGC website (https://www.med.unc.edu/pgc/results-and-downloads). GWA summary statistics with results from the GWAS of ADHD symptom scores analyzed in the EAGLE sample can be accessed at the PGC website (see link above). Summary statistics for the 23andMe dataset can be obtained by qualified researchers under an agreement with 23andMe that protects the privacy of the 23andMe participants.

Availability of genotype data

For access to genotypes from the PGC cohorts and the iPSYCH sample interested researchers should contact the lead PIs (iPSYCH: lead PI Anders D. Børglum; PGC: Benjamin Neale and Stephen Faraone).

URLs

LD-Hub, http://ldsc.broadinstitute.org

LD score regression, https://github.com/bulik/ldsc

Pre-computed European LD scores, https://data.broadinstitute.org/alkesgroup/LDSCORE/

PGC Ricopili GWA pipeline, https://github.com/Nealelab/ricopili

Credible set analysis, https://github.com/hailianghuang/FM-summary

FUMA, http://fuma.ctglab.nl

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature

End notes

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Dr. Neale is a member of Deep Genomics Scientific Advisory Board and has received travel expenses from Illumina. He also serves as a consultant for Avanir and Trigeminal solutions.

Dr. Rohde has received honoraria, has been on the speakers' bureau/advisory board and/or has acted as a consultant for Eli-Lilly, Janssen-Cilag, Novartis, Medice and Shire in the last three years. He receives authorship royalties from Oxford Press and ArtMed. He also received travel award for taking part of 2015 WFADHD meeting from Shire. The ADHD and Juvenile Bipolar Disorder Outpatient Programs chaired by him received unrestricted educational and research support from the following pharmaceutical companies in the last three years: Eli-Lilly, Janssen-Cilag, Novartis, and Shire.

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Figure legends

Figure 1. Manhattan plot of the results from the GWAS meta-analysis of ADHD. The index variants in the 12 genome-wide significant loci are highlighted as a green diamond. Index variants located with a distance less than 400kb are considered as one locus.

Figure 2. Odds Ratio (OR) by PRS within each decile estimated for individuals in the PGC samples (red dots) and in the iPSYCH sample (blue dots). Error bars indicate 95% confidence limits.

Figure 3. Significant genetic correlations between ADHD and other traits reveal overlap of genetic risk factors for ADHD across several groups of traits (grouping indicated by a horizontal line): educational, psychiatric/personality, weight (and possible weight related traits), smoking behaviour/smoking-related cancer, reproductive traits and parental longevity. In total 220 traits were tested. Two significant educational phenotypes are omitted due to substantial overlap with years of schooling. Error bars indicate 95% confidence limits.

Figure 1.

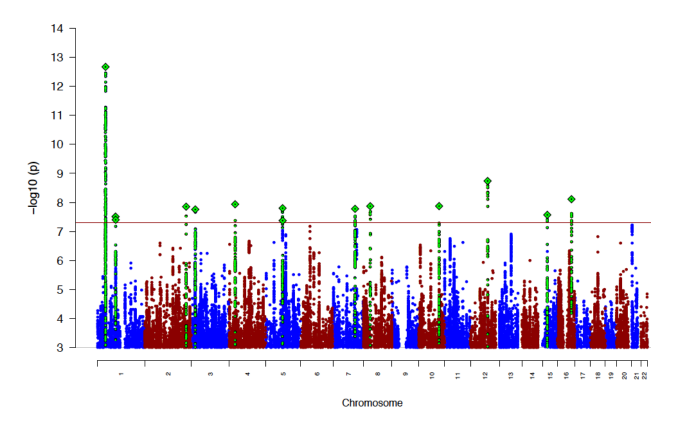


Figure 2.

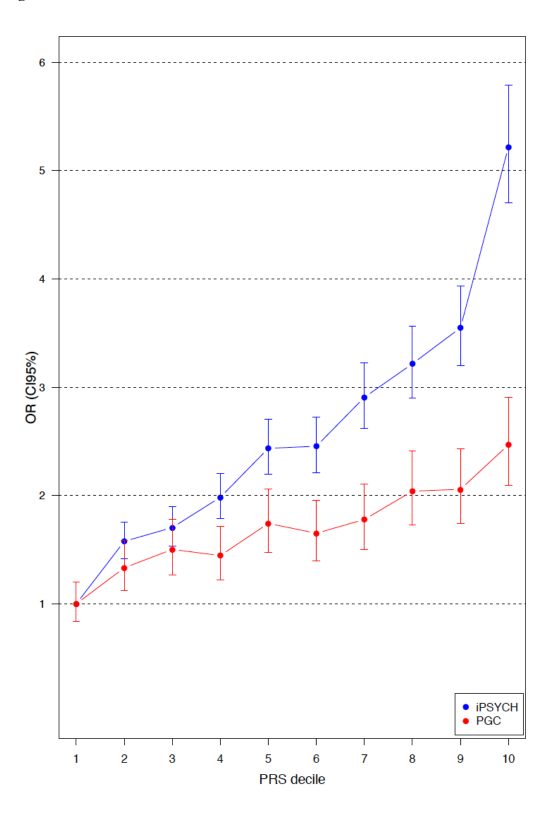


Figure 3.

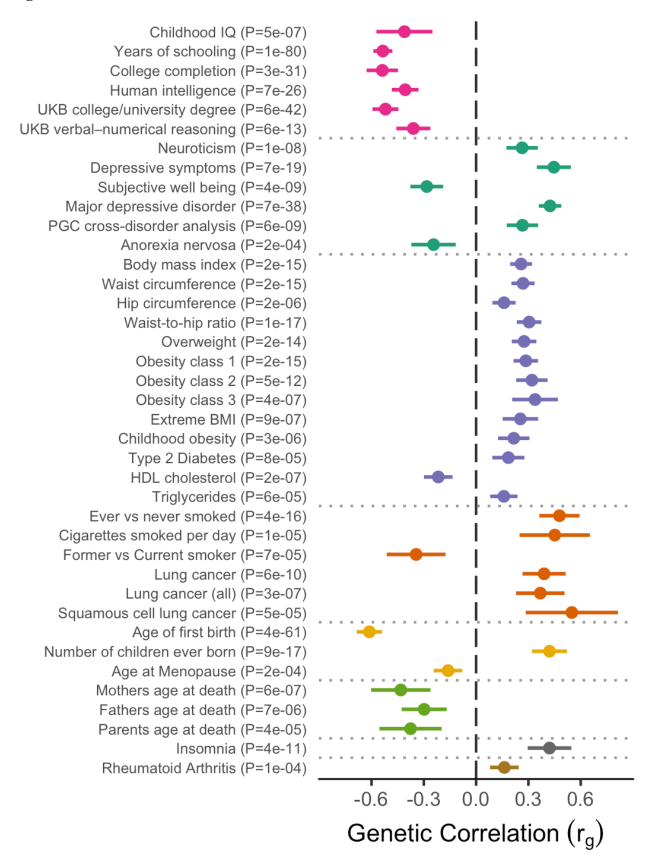


Table 1. Results for the genome-wide significant index variants in the 12 loci associated with ADHD identified in the GWAS metaanalysis. Index variants are LD independent ($r^2 < 0.1$), and are merged into one locus when located with a distance less than 400kb. The location (chromosome [Chr] and base position [BP]), alleles (A1 and A2), allele frequency (A1 Freq), odds ratio (OR) of the effect with respect to A1, and association P-value of the index variant are given, along with genes within 50kb of the credible set for the locus.

Locus	Chr	BP	Index Variant	Genes	A1	A2	A1 Freq	OR	P-value
1	1	44184192	rs11420276	ST3GAL3, KDM4A, KDM4A-AS1, PTPRF, SLC6A9, ARTN, DPH2, ATP6V0B, B4GALT2, CCDC24, IPO13	G	GT	0.696	1.113	2.14 x 10 ⁻¹³
2	1	96602440	rs1222063	Intergenic	A	G	0.328	1.101	3.07×10^{-8}
3	2	215181889	rs9677504	SPAG16	A	G	0.109	1.124	1.39 x 10 ⁻⁸
4	3	20669071	rs4858241	Intergenic	T	G	0.622	1.082	1.74 x 10 ⁻⁸
5	4	31151456	rs28411770	PCDH7, LINC02497	T	C	0.651	1.09	1.15 x 10 ⁻⁸
6	5	87854395	rs4916723	LINC00461, MIR9-2, LINC02060, TMEM161B-AS1	A	С	0.573	0.926	1.58 x 10 ⁻⁸
7	7	114086133	rs5886709	FOXP2, MIR3666	G	GTC	0.463	1.079	1.66 x 10 ⁻⁸
8	8	34352610	rs74760947	LINC01288	A	G	0.957	0.835	1.35 x 10 ⁻⁸
9	10	106747354	rs11591402	SORCS3	A	T	0.224	0.911	1.34 x 10 ⁻⁸
10	12	89760744	rs1427829	DUSP6, POC1B	A	G	0.434	1.083	1.82 x 10 ⁻⁹
11	15	47754018	rs281324	SEMA6D	T	C	0.531	0.928	2.68 x 10 ⁻⁸
12	16	72578131	rs212178	LINC01572	A	G	0.883	0.891	7.68 x 10 ⁻⁹

Extended Data

- eTable 1. Bayesian credible sets of variants for each of the 12 genome-wide significant loci
- **eTable 2.** Summary of the observed annotations for the credible set at each genome-wide significant locus
- eTable 3. Variant-level annotations for the credible set at each genome-wide significant locus
- eTable 4. Results of gene set analyses using sets from Gene Ontology
- eTable 5. Extended results from genetic correlation analyses of ADHD and 219 phenotypes
- **eTable 6.** Genome-wide significant index variants in meta-analyses of iPSYCH, PGC, deCODE, 23andMe and EAGLE