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Association analysis in over 329,000 individuals identifies 116 independent variants influencing neuroticism.

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Neuroticism is a relatively stable personality trait characterised by negative emotionality (e.g., worry, guilt)\textsuperscript{1}; twin study heritability ranges 30 to 50\%\textsuperscript{2}, and SNP-based heritability ranges 6 to 15\%\textsuperscript{3-6}. Increased neuroticism is associated with poorer mental and physical health\textsuperscript{7,8}, translating to high economic burden\textsuperscript{9}. Genome-wide association (GWA) studies of neuroticism have identified up to 11 genetic loci\textsuperscript{3,4}. Here we report 116 significant independent loci from a GWA of neuroticism in 329,821 UK Biobank participants; 15 of these replicated at P<.00045 in an unrelated cohort (N = 122,867). Genetic signals were enriched in neuronal genesis and differentiation pathways, and substantial genetic correlations were found between neuroticism and depressive symptoms ($r_g = .82$, SE=.03), major depressive disorder (MDD; $r_g = .69$, SE=.07) and subjective wellbeing ($r_g = -.68$, SE=.03) alongside other mental health traits. These discoveries significantly advance our understanding of neuroticism and its association with MDD.

Main

Understanding why people differ in neuroticism will provide an important contribution to understanding people’s liability to poor mental health throughout the life course. The strong genetic correlation between neuroticism and mental health, especially anxiety and major depressive disorder\textsuperscript{10,11}, means that exploring the genetic contribution to differences in neuroticism is one way to understand more about these common and burdensome, but aetiologically intractable illnesses. In the largest GWA study of major depressive disorder (MDD; 130,664 cases vs 330,470 controls), 44 independent genetic loci were identified\textsuperscript{12}.

UK Biobank has health, medical and genetic information for over 500,000 individuals aged 39-73 years from the United Kingdom, assessed between 2006 and 2010\textsuperscript{13,14}. We performed a GWA analysis of trait neuroticism in 329,821 unrelated White British adults (152,710 male (46.3%)) with high-quality genotype data (Online Methods). Neuroticism was
measured by the total score of the 12-item Eysenck Personality Questionnaire-Revised Short Form (EPQ-R-S)\(^1\); missing item data (ranging 1.8% to 4.7%) were imputed with reference to age and sex, and individuals with greater than 4 missing items were excluded (Supplementary Note, Supplementary Table 1 and Supplementary Fig. 1). For analysis, the score was residualized for the effects of age, sex, assessment centre, genotype batch, array, and 40 genetic principal components. This score was tested against 18,485,882 bi-allelic single nucleotide polymorphism (SNP) variants, based on the Haplotype Reference Consortium panel\(^2\), with a minor allele frequency $\geq 0.0005$ and an information/imputation quality score of $\geq 0.1$ under an additive model. The distribution of obtained versus expected results under the null hypothesis showed some genomic inflation, with a lambda of 1.15 (quantile-quantile plot shown in Supplementary Fig. 2). Univariate linkage disequilibrium score (LDSC) regression\(^3\) estimates indicated that 96.8% of this inflation was due to the presence of a large polygenic signal with the intercept being close to 1 (1.02, SE = .01). SNP-based heritability of neuroticism was estimated at .108 (SE=.005) using LDSC.

Genome-wide significance (P < 5 x 10\(^{-8}\)) was demonstrated for 10,353 genetic variants with a further 17,668 variants at a suggestive level (P < 1 x 10\(^{-5}\)) (Supplementary Table 2). The Manhattan plot is shown in Figure 1 and gene annotation for the significant SNPs in Supplementary Table 3. SNPs identified in previous neuroticism GWA studies were mostly significant in our sample (Supplementary Note and Supplementary Table 2) and substantial overlap with MDD SNPs (75%) and genes was found (Supplementary Note and Supplementary Table 4). The major histocompatibility complex (MHC) region has been previously linked to schizophrenia, a psychiatric comorbidity trait (including MDD)\(^4,5\) and MDD\(^6,7\). It contained 3 significant independent genetic loci associated with neuroticism, two were in genes \(GABBR1, TNXB\) connected with schizophrenia\(^8,9\). The primary associated SNP, rs2021722, for schizophrenia\(^10,11\), was present in our study and nominally significant (P =
9.42 x 10^{-5}). Supplementary Figure 3 indicates the previous MHC associations in relation to our findings.

116 of the significant SNPs were independent \( (r^2 > 0.1 \text{ and within 500kb of the significant index SNP}) \); these lead SNPs are shown in Supplementary Table 5 with the number of associated SNPs, region size, and genes within the LD interval. 73 lead SNPs were located within genes, 5 were exonic (in \textit{MSRA}, \textit{NOS1}, \textit{PINX1}, \textit{ZCCHC14}, and \textit{C12orf49}) and a further 2 were coding SNPs in \textit{RPP21} (a missense mutation) and \textit{AGBL1} (synonymous), 55 were intronic and 10 were noncoding RNA variants; 42 were intergenic. For the 116 independent SNPs, evidence of expression quantitative trait loci (eQTL) was explored using the GTEx database, 44 were eQTLs (Supplementary Table 5). A Regulome DB score was used to identify SNPs with a likely regulatory function. 33 of the 116 SNPs were included in the Regulome DB database and 8 of these had a score < 3, indicating that they are likely to be involved in gene regulation (Supplementary Table 5).

Replication of the significant association signals in UK Biobank was sought from the results of a GWA meta-analysis of neuroticism that we performed using 23andMe (N = 59,206) \textsuperscript{22} and the Genetics of Personality Consortium (GPC-2; N = 63,661) \textsuperscript{23}. Of the 10,353 genome-significant SNPs in UK Biobank, 10,171 were available in the replication cohorts, and 8,774 of these increased in significance when the replication cohorts were meta-analysed with UK Biobank. This indicated a consistent direction of allelic effect (Supplementary Table 6).

![INSERT FIGURE 1 ABOUT HERE](image-url)

Of the 116 independent associated SNPs, 111 were present in the replication cohort, with 51 nominally significant \( (P < .05; \text{Supplementary Table 5}) \), and 15 at a Bonferroni-corrected level \( (P < .00045; \text{Table 1}) \). One of these, \textit{rs2953805}, was previously associated
with morning chronotype with morning chronotype\textsuperscript{24}, a trait relating to lower neuroticism\textsuperscript{25} and showing allelic effects in the expected direction. The low replication rate (13.5\%) at a strict corrected level reflects the finding that effect sizes are extremely small (up to .02 of a SD increase in neuroticism score per allele) and will thus require similarly large replication samples to confirm their effects. Figure 2a-c shows the regional association plot for chromosomes 8, 11 and 22 in which multiple genes were present in the associated LD region. Of the five chromosome 8 loci only one lead SNP tagged a well-known inversion, previously linked to neuroticism (Supplementary Note and Supplementary Fig. 4), although associations in the broad region had been attributed to the inversion\textsuperscript{4} and so might cautiously be considered as a single locus.

All 69 genes located within the 15 replicated loci were classified in terms of their molecular function, biological process and protein class using the Protein Analysis Through Evolutionary Relationships Classification System which includes 14,710 protein families categorised into 76,032 functionally distinct subfamilies\textsuperscript{26}. Supplementary Figure 4 shows that a large number of genes 1) coded for nucleic acid binding and transcription factors, 2) contributed to metabolic and cellular processes, and 3) had a role in binding and catalytic activity molecular functioning. Transcription factors, in particular, have been implicated in the aetiology of depression\textsuperscript{27,28}, and miRNAs—which have been linked with anxiety\textsuperscript{29} and depression\textsuperscript{30}—might target genes with roles in binding (e.g., \textit{POLR3H}). The PsyGeNET (v2.0) database showed that of the 69 genes, four have been associated with psychiatric disorder (Supplementary Table 7): \textit{DRD2} (bipolar, depression, substance use/dependence, delirium), \textit{EP300} (alcoholic intoxication), \textit{TEF} (depression) and \textit{MSRA} (schizophrenia). Variants in \textit{CACNA1E} have been associated with cross-psychiatric disorder overlap and migraine\textsuperscript{18,31}.

A GTEx database search for the 15 replicated SNPs showed that 9 were associated with significant regulation of 60 genes expressed in a variety of tissues (Supplementary Table
Of the 30 brain expression associations, half of these were in the cerebellum: 4 SNPs regulating 10 genes. Interestingly, MRI studies have shown associations between cerebellar volume and neuroticism, and cerebellar blood flow in response to negative emotional cues. In the BRAINEAC search, all SNPs were identified as eQTLs in at least one brain region at a nominal significance level (P < .05) and 10 were supported at a Bonferroni-corrected level of P < .0003 (Supplementary Table 9). Of potential interest, rs7107356, a novel SNP in an intergenic region of chromosome 11, regulates *MTCH2* in the cerebellar cortex (P = 4.5 x 10^{-6}). *MTCH2* is involved in metabolic pathways and cell function and variants of this gene have been associated with BMI.

Gene-based analysis of the GWA results was performed using MAGMA; 249 genes were significantly associated at a Bonferroni-corrected level (α = 0.05 / 18,080; P < 2.77 × 10^{-6}; Supplementary Table 10). Three of these were genes (*STH, HIST1H3J, HIST1H4L*) containing a single SNP. Of the replicated independent GWA SNPs that were in genes, the following significant genes were corroborated in the gene-based results:

*CACNA1E, XKR6, MSRA, LINGO2, CELF4, ZC3H7B* and *BAIAP2*. SNP rs6981523, previously identified in 23andMe for neuroticism, was an intergenic SNP near *XKR6*; this gene was the second most significant gene in our gene-based analysis (P = 6.55 × 10^{-32}). *L3MBTL2* and *CHADL*, wherein 23andMe’s other significant SNP, rs9611519, resided, showed respective gene-based p-values of 2.40 × 10^{-6} and 1.15 × 10^{-6}.

Pathway analysis in MAGMA highlighted 5 significant gene ontology pathways (family-wise error P < 1.21 × 10^{-6}): neuron spine (cellular), homophilic cell adhesion via plasma membrane adhesion molecules (biological), neuron differentiation (biological), cell adhesion via plasma membrane adhesion molecules (biological), and neurogenesis (biological). See Table 2 for further details. Of note is the neurogenesis pathway, a hypothesis of which exists for depression (and to a lesser extent, anxiety) based on stress.
reducing neurogenesis in the hippocampus and on the action of antidepressants on brain circuitry \(^{37,38}\). Further, variants in \textit{PLXNA2}, potentially involved in adult neurogenesis, have been associated with anxiety and neuroticism \(^{39}\). Cell adhesion molecules have been implicated in neuropsychiatric disorder \(^{40}\), and protocadherins specifically with neuroticism and risk of mood disorder \(^{41}\), which supports the importance of cell adhesion pathways. A further gene-set analysis of genes expressing proteins that can bind to anti-depressant drug molecules was significant (\(P = .005\)) re-affirming the dependency of neuroticism and depression on shared biological pathways. This is consistent, for example, with findings for \textit{CRHRI} (highlighted in our SNP and gene-based analysis), a gene involved in normal hormonal responses to stress (the glucocorticoid pathway being a relevant and well-known target) and associated with anxiety, depression and neuroticism \(^{3,42,43}\). That genes influencing neuroticism reveal pathways involved in currently prescribed and effective antidepressant action suggests that neuroticism could be a potentially useful clinical stratifying factor for effective antidepressant action. There may also be clinical utility in knowing a person’s level of neuroticism after the occurrence of a stressful life event and therefore pre-empting onset of depression via drug therapy in those high in neuroticism. Because our GWA of neuroticism reveals signals associated with the known biological action of existing antidepressants, it may be useful as a means of discovering (or re-purposing) new pharmacological interventions for MDD.

LD score regression \(^{44}\) was used to estimate the genetic correlation between neuroticism and a variety of health traits (Supplementary Tables 11 and 12). The strongest correlation was observed for depressive symptoms (\(r_g = .82, SE = .03\)). Major depressive disorder, subjective wellbeing, and tiredness showed moderate-to-strong correlations (.62-.69). The stronger correlation for depressive symptoms than depressive disorder might be indicative of improved sensitivity of continuous versus dichotomous traits but might also
point to inventory item overlap (greater conceptual similarity) for depressive symptoms and/or noise in MDD diagnosis. Genetic correlations with neuroticism were moderate for self-rated health (.41), moderate-to-low for schizophrenia, ADHD, anorexia nervosa and educational attainment (~.20), and low for bipolar disorder and smoking status (.11). The genetic correlation of one between Eysenck neuroticism and other neuroticism scales (used by 23andMe and the GPC) confirms that GWA meta-analysis based on different measurement instruments is valid. Mendelian randomization was used to determine whether the genetic correlation between neuroticism and non-psychiatric variables (less likely to be influenced by pleiotropy), smoking status and educational attainment, represented a causal relationship from neuroticism. For smoking status, the beta of 0.23 was significant in the inverse variance weighted model (P = .00002) which is preferred in the presence of heterogeneity (P = .001); the MR Egger regression did not show significant directional pleiotropy (intercept = 0.02, P = .10) thus supporting a causal relationship. For educational attainment, the beta of -0.09 was significant (P = 8.35 × 10^-6) in the inverse variance weighted model (heterogeneity P = 5.87 × 10^-7), with no evidence of directional pleiotropy (intercept = 0, P = .23). Although theoretically less plausible, the reverse causal direction should be investigated in UK Biobank once a large number of significant SNPs influencing smoking status and educational attainment have been estimated in non-overlapping samples.

Polygenic profile analyses based on the SNP inclusion threshold with the optimal signal-to-noise ratio (P < .05) indicated that the neuroticism polygenic score explained 2.79% of the variance in neuroticism (β = .19, P = 2.65 × 10^-47) and 0.8% of the variance in depression status (OR = 1.25, P = 1.53 × 10^-8) in Generation Scotland (GS; N = 7,388) 45. Results for polygenic scores in GS based on other SNP significance inclusion thresholds (0.01, 0.05, 0.1, 0.5 and 1) from the UK Biobank GWA can be found in Supplementary Table 13.
The combination, in UK Biobank, of a large ethnically homogenous sample and a well-validated neuroticism scale has afforded the discovery of 15 stringently replicated genetic loci that influence neuroticism levels, four of them novel. Most lead variants were associated with gene regulation, with half of these expressed in the brain; single variant and gene associations overlapped substantially with MDD findings, and genes in antidepressant-targeted pathways were over-represented. There was also support for neuroticism having causal effects on socio-economic markers. These discoveries promise paths to understand the mechanisms whereby some people become depressed, and of broader human differences in happiness, and they are a resource for those seeking novel drug targets for major depression.

After millennia in which scholars and researchers have sought the sources of individual differences in proneness to dysphoria, the present study adds significantly to explaining the (genetic) anatomy of melancholy.

**URLs**

UK Biobank Resource: http://www.ukbiobank.ac.uk

BGENIE: https://jmarchini.org/bgenie/

BRAINEAC: http://www.braineac.org/

Druggable genome: http://dgidb.genome.wustl.edu/

Genotype-Tissue Expression Portal: http://www.gtexportal.org

Gene Ontology: http://geneontology.org

GPC-2 Summary Statistics: http://www.tweelingenregister.org/GPC/

Linkage Disequilibrium Score Regression: https://github.com/bulik/ldsc/wiki

METAL: http://csg.sph.umich.edu/abecasis/metal/index.html

PANTHER: http://pantherdb.org/

PLINK V2: https://www.cog-genomics.org/plink2
Acknowledgements

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Author Disclosure

IJD was a participant in UK Biobank. The other authors declare no conflict of interest.
Author Contributions

M.L. drafted the manuscript with contributions from W.D.H. and I.J.D. G.D., D.C.L.,
R.E.M., M.J.A. and D.M.H. performed quality control of UK Biobank data and/or Generation
performed/assisted with downstream analysis. C.R.G, C.M.L., and A.M.M provided critical
comments on the manuscript draft and analysis. M.L. and I.J.D. co-ordinated the work. All
authors commented on and approved the manuscript.
References for Main text


42. DeYoung, C.G., Cicchetti, D. & Rogosch, F.A. Moderation of the association between childhood maltreatment and Neuroticism by the corticotropin-releasing hormone receptor 1 gene. *Journal of child psychology and psychiatry, and allied disciplines* 52, 898-906 (2011).


**Figure Legends for Main Text**

Figure 1. GWA results for neuroticism in 329,821 UK Biobank individuals.

Figure 2. Regional association plot for suggestive/significant signals in UK Biobank on a) chromosome 8p (site of the inversion polymorphism), b) chromosome 11, and c) chromosome 22. The SNP association p-value is shown on the y-axis and the SNP position (with gene annotation) appears on the x-axis; for each SNP, the strength of LD with the lead SNP is colour coded based on its r². Plots were produced in LocusZoom.
Tables
Table 1. Fifteen independent SNPs associated with neuroticism in UK Biobank most strongly replicated (with consistent allelic effect) in the meta-analysis of 23andMe and the GPC cohorts. Bolded genes were significant in the gene-based tests.

<table>
<thead>
<tr>
<th>Chr</th>
<th>SNP</th>
<th>MAF</th>
<th>Discovery P-value (N=329,821)</th>
<th>Replication P-value (N=122,867)</th>
<th>Nearest Gene</th>
<th>Distance to Gene</th>
<th>Genes within Range Significant in Previous GWA Studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>rs169235</td>
<td>.25</td>
<td>3.97×10^{-9}</td>
<td>2.55E-05</td>
<td><strong>CACNA1E</strong></td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>rs1422192</td>
<td>.17</td>
<td>1.68×10^{-9}</td>
<td>6.54E-07</td>
<td><strong>LINC00461</strong></td>
<td>0</td>
<td><strong>MEF2C</strong></td>
</tr>
<tr>
<td>8</td>
<td>rs2921036</td>
<td>.49</td>
<td>8.04×10^{-26}</td>
<td>3.27E-07</td>
<td><strong>LINC00458</strong></td>
<td>0</td>
<td><strong>CLDN23, ER11</strong>, <strong>MFHAS1</strong>, <strong>SGK223</strong></td>
</tr>
<tr>
<td>8</td>
<td>rs2953805</td>
<td>.47</td>
<td>3.02×10^{-22}</td>
<td>1.26E-08</td>
<td><strong>U3</strong></td>
<td>1292</td>
<td><strong>CLDN23, ER11</strong>, <strong>MFHAS1</strong>, <strong>PPP1R3B</strong></td>
</tr>
<tr>
<td>8</td>
<td>rs6982308</td>
<td>.49</td>
<td>6.46×10^{-21}</td>
<td>2.26E-08</td>
<td><strong>MSRA</strong></td>
<td>0</td>
<td><strong>C8orf74, PINX1, PRSS55, RP1L1, SOX7, XKR6</strong></td>
</tr>
<tr>
<td>8</td>
<td>rs10097870</td>
<td>.47</td>
<td>2.18×10^{-24}</td>
<td>6.51E-07</td>
<td><strong>LINC00208</strong></td>
<td>5665</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>rs1521732</td>
<td>.37</td>
<td>2.91×10^{-9}</td>
<td>4.01E-06</td>
<td><strong>LINGO2</strong></td>
<td>0</td>
<td><strong>AQP2, AGBL2, CIQTNF4, CELF1, DDB2, FAM180B, FNBP4, KBTBD4, MADD, MTCH2, MYBPC3, NDUFS3, NR1H3, NUP160, PSMC3, PTPMT1, RAPSN, SLC39A13</strong></td>
</tr>
<tr>
<td>9</td>
<td>rs72694263</td>
<td>.08</td>
<td>2.12×10^{-8}</td>
<td>0.000237</td>
<td>.</td>
<td>.</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>rs7107356</td>
<td>.50</td>
<td>1.52×10^{-12}</td>
<td>1.34E-05</td>
<td><strong>AGBL2</strong></td>
<td>4973</td>
<td></td>
</tr>
<tr>
<td>SNP</td>
<td>rs Number</td>
<td>P Value (10^-n)</td>
<td>Adjusted P Value</td>
<td>Gene</td>
<td>Genotyped SNP</td>
<td>Located in Inversion Region</td>
<td>Broad region implicated in previous studies</td>
</tr>
<tr>
<td>--------</td>
<td>-----------</td>
<td>-----------------</td>
<td>------------------</td>
<td>------</td>
<td>---------------</td>
<td>-----------------------------</td>
<td>------------------------------------------</td>
</tr>
<tr>
<td>11*</td>
<td>rs7111031</td>
<td>0.36</td>
<td>1.06×10^{-15}</td>
<td>0.000215</td>
<td>.</td>
<td>.</td>
<td>Broad region implicated in previous studies</td>
</tr>
<tr>
<td>15*</td>
<td>rs7175083</td>
<td>0.48</td>
<td>1.16×10^{-9}</td>
<td>0.000297</td>
<td><strong>LINGO1</strong></td>
<td>0</td>
<td>Broad region implicated in previous studies</td>
</tr>
<tr>
<td>17</td>
<td>rs7502590</td>
<td>0.15</td>
<td>2.61×10^{-11}</td>
<td>0.000146</td>
<td><strong>BAIAP2</strong></td>
<td>0</td>
<td>Broad region implicated in previous studies</td>
</tr>
<tr>
<td>18*</td>
<td>rs11082011</td>
<td>0.34</td>
<td>1.25×10^{-16}</td>
<td>2.05E-06</td>
<td><strong>CELF4</strong></td>
<td>0</td>
<td>Broad region implicated in previous studies</td>
</tr>
<tr>
<td>22*</td>
<td>rs11090045</td>
<td>0.30</td>
<td>8.04×10^{-13}</td>
<td>5.40E-07</td>
<td><strong>ZC3H7B</strong></td>
<td>0</td>
<td>Broad region implicated in previous studies</td>
</tr>
</tbody>
</table>

* Genotyped SNP
† Located in Inversion Region
* Broad region implicated in previous studies
** Regulated gene expressed in brain

*Genotyped SNP
†Located in Inversion Region
* Broad region implicated in previous studies
** Regulated gene expressed in brain
### Table 2. Significant gene ontology pathways for neuroticism in UK Biobank

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Number of genes</th>
<th>Beta</th>
<th>SE</th>
<th>P-value</th>
<th>Corrected P</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neuron Spine</td>
<td>147</td>
<td>0.560</td>
<td>0.107</td>
<td>$7.77 \times 10^{-8}$</td>
<td>0.0282</td>
<td>A small membranous protrusion, often ending in a bulbous head and attached to the neuron by a narrow stalk or neck.</td>
</tr>
<tr>
<td>Homophilic Cell Adhesion Via Plasma Membrane Adhesion Molecules</td>
<td>115</td>
<td>0.490</td>
<td>0.0938</td>
<td>$8.81 \times 10^{-8}$</td>
<td>0.0289</td>
<td>The attachment of a plasma membrane adhesion molecule in one cell to an identical molecule in an adjacent cell.</td>
</tr>
<tr>
<td>Neuron Differentiation</td>
<td>1341</td>
<td>0.145</td>
<td>0.0288</td>
<td>$2.36 \times 10^{-7}$</td>
<td>0.0357</td>
<td>The process in which a relatively unspecialized cell acquires specialized features of a neuron.</td>
</tr>
<tr>
<td>Cell Cell Adhesion Via Plasma Membrane Adhesion Molecules</td>
<td>828</td>
<td>0.183</td>
<td>0.0364</td>
<td>$2.72 \times 10^{-7}$</td>
<td>0.0372</td>
<td>The attachment of one cell to another cell via adhesion molecules that are at least partially embedded in the plasma membrane.</td>
</tr>
<tr>
<td>Neurogenesis</td>
<td>195</td>
<td>0.419</td>
<td>0.0859</td>
<td>$5.35 \times 10^{-7}$</td>
<td>0.0439</td>
<td>Generation of cells within the nervous system</td>
</tr>
</tbody>
</table>
Online Methods

Genome-wide association analysis in UK Biobank

An imputed dataset, including >92 million variants, referenced to the UK10K haplotype, 1000 Genomes Phase 3, and Haplotype Reference Consortium (HRC) panels was available in UK Biobank. The current analysis includes only those SNPs available in the HRC reference panel. Quality control filters were applied (see Supplementary Note) which resulted in 18,485,882 imputed SNPs for analysis in 329,821 individuals. The GWA of neuroticism was conducted using BGENIE, a program specifically developed to analyse UK Biobank data in a fast and efficient manner. Further information can be found at the following URL: https://jmarchini.org/bgenie/. A linear SNP association model was tested which accounted for genotype uncertainty. Neuroticism was pre-adjusted for age, sex, genotyping batch, genotyping array, assessment centre, and 40 principal components to speed up analysis.

The number of independent signals from the GWA analysis was determined using LD-clumping in PLINK v1.90b3i (see URLs). The LD structure was based on SNPs with a p-value < 1 × 10^{-3} that were extracted from the imputed genotypes. Index SNPs were identified (P < 5 × 10^{-8}) and clumps were formed for SNPs with P < 1 × 10^{-5} that were in LD (r^2 > 0.1) and within 500kb of the index SNP. SNPs were assigned to no more than one clump.

Meta-analysis of GWA Results

Two meta-analyses were performed. Firstly, to check for replication of the significant (P < 5 × 10^{-8}) GWA signals in UK Biobank, results from a meta-analysis of 23andme (the full GWA summary statistics were made available from 23andMe) and the Genetics of Personality Consortium (GPC-2) (the full GWA summary statistics were publicly available...
see URLs) were used. This meta-analysis was conducted using METAL\textsuperscript{52} and due to the lack of phenotype harmonisation across the cohorts, a sample size weighted meta-analysis was preferred. A second meta-analysis of UK Biobank and the replication cohorts was performed using the same method, but only for the SNPs that were significant in UK Biobank.

Genome-wide Gene-based Analysis

Gene-based analysis of neuroticism was performed using MAGMA\textsuperscript{53}, which provides gene-based statistics derived using the results of the GWA analysis. Genetic variants were assigned to genes based on their position according to the NCBI 37.3 build, with no additional boundary placed around the genes. This resulted in a total of 18,080 genes being analysed. The European panel of the 1000 Genomes data (phase 1, release 3) was used as a reference panel to account for linkage disequilibrium. A genome-wide significance threshold for gene-based associations was calculated using the Bonferroni method ($\alpha=0.05/18,080$; $P < 2.77 \times 10^{-6}$).

Functional annotation and gene expression

For the 116 independent genome-wide significant SNPs identified by LD clumping, evidence of expression quantitative trait loci (eQTL) and functional annotation were explored using publicly available online resources. The Genotype-Tissue Expression Portal (GTEx) (see URLs) was used to identify eQTLs associated with the SNPs. Functional annotation was investigated using the Regulome DB database\textsuperscript{54} (see URLs). Further to GTEx searches, we investigated whether any of the 15 replicated SNPs were brain expression quantitative loci (eQTLs) by entering them into the brain eQTL database BRAINEAC (see URLs), which
contains gene expression data across ten brain regions (cerebellar cortex, frontal cortex,
hippocampus, medulla, occipital cortex, putamen, substantia nigra, temporal cortex, thalamus
and intralobular white matter). The genes located in the region of replicated independent loci
were investigated for protein function using the PANTHER database (Protein ANalysis
THrough Evolutionary Relationships, see URLs) which stores data on the evolution and
function of protein-coding genes from sequenced genomes of diverse species, our focus
here on homo sapiens. Uncharacterized gene function is predicted via phylogenetic branching
information and the resource enables biological pathway annotation.

Pathway Analysis

Biological pathway analysis was performed on the gene-based analysis results. This
gene-set enrichment analysis was conducted utilising gene-annotation files from the Gene
Ontology (GO) Consortium (see URLs) taken from the Molecular Signatures Database
(MSigDB) v5.2. The GO consortium includes gene-sets for three ontologies; molecular
function, cellular components and biological function. This annotation file consisted of 5,917
gene-sets which were corrected for multiple testing correction using the MAGMA default
setting correcting for 10,000 permutations.

To determine whether the genetic targets of antidepressants were enriched for
neuroticism we performed a competitive gene-set analysis using MAGMA. Gene sets
corresponding to the Anatomical Therapeutic Chemical Classification System code N06A
Antidepressants (within the Psychoanaleptics class) were downloaded (see URLs). This
resulted in a set of 110 unique genes corresponding to those that are the targets of the
antidepressants. Enrichment for neuroticism was tested against a set of 5483 ‘druggable’
autosomal genes (see URLs), that is, they code for proteins which can bind to drug-like
molecules. Of the 110 antidepressant genes 86 were found amongst the 5483 druggable
genes.

Linkage Disequilibrium Score Regression

Univariate Linkage disequilibrium Score (LDSC) regression \(^{57}\) was used to test for
residual stratification in our GWAS summary statistics and to derive a heritability estimate.

An LD regression was performed by regressing the GWA test statistics \((\chi^2)\) on to each SNP’s
LD score (the sum of squared correlations between the minor allele frequency count of a SNP
with the minor allele frequency count of every other SNP). This regression allows for the
estimation of heritability from the slope, and a means to detect residual confounders, the
intercept. The percentage inflation in the test statistic due to polygenic signal can be derived
by subtracting the LDSC ratio \(((\text{intercept} - 1)/(\text{mean } \chi^2 - 1))\), which represents inflation due to
population stratification and other confounding, from 1 and multiplying by 100. Bivariate
LDSC regression \(^{58}\) was used to derive genetic correlations between neuroticism and 18
psychiatric and physical health phenotypes (see Supplementary Table 11). For Alzheimer’s
disease, a 500-kb region surrounding \(APOE\) was excluded and the analysis re-run
(Alzheimer’s disease (500kb)). The genetic correlation between neuroticism as measured by
different inventories was also estimated. Further details, including source of GWA summary
statistics can be found in the Supplementary Note. Sample overlap could not be controlled for
in the LDSC analyses because the exact overlap between the UK Biobank data and the health
traits was unknown. In such a case, constraining the intercept to a ‘wrong’ value could lead to
biased estimates. Any sample overlap in the present analyses will only affect the intercept of
the regression and could lead to inflated standard errors, but will not affect the genetic
correlation \(^{12}\).
Two sample Mendelian Randomization (MR) was performed using the TwoSampleMR \textsuperscript{59} package implemented in R. GWA summary statistics from the GWA of smoking status in 74,053 Europeans \textsuperscript{60} was used to create outcome data for the MR between neuroticism and smoking status. 77 independent SNPs associated with neuroticism were available in the smoking status GWA summary data to test for a causal effect of neuroticism on smoking status. There were no significant SNP signals for smoking status to test the reverse causation model. GWA summary statistics from the GWA of educational attainment in 126,559 Caucasians \textsuperscript{61} was used to create outcome data for the MR between neuroticism and educational attainment. 75 independent SNPs associated with neuroticism were available in the educational attainment GWA summary data to test for a causal effect of neuroticism on educational attainment. There were too few significant SNPs available for educational attainment to test for a causal effect of educational attainment on neuroticism. Sensitivity analyses were performed to test for heterogeneity and a further test for horizontal pleiotropy was carried out.

Polygenic Prediction into Generation Scotland

Polygenic profile analyses were performed to predict neuroticism and depression status in Generation Scotland (GS) \textsuperscript{62}. Polygenic profiles were created in PRSice \textsuperscript{63} using the UK Biobank neuroticism SNP-based association results, for 7,388 unrelated individuals in GS. SNPs with a MAF $<0.01$ were removed prior to creating the polygenic profiles. Clumping was used to obtain SNPs in linkage disequilibrium with an $r^2 < 0.25$ within a 250kb window. Individuals were removed from GS if they had contributed to both UK Biobank and GS ($n = 302$). Polygenic profile scores were created based on the significance of the association in UK Biobank with the neuroticism phenotype, at p-value thresholds of 0.01, 0.05, 0.1, 0.5 and
1 (all SNPs). Linear regression models were used to examine the associations between the polygenic profile and neuroticism score in GS, adjusting for age at measurement, sex and the first 10 genetic principal components to adjust for population stratification. Logistic regression models were used to examine depression status, adjusting for the same covariates as in the neuroticism models. The false discovery rate (FDR) method was used to correct for multiple testing across the polygenic profiles for neuroticism at all five thresholds.

Data Availability

The GWA results generated by this analysis are publicly available at http://www.ccace.ed.ac.uk.

References for Online Methods


