# Brain expression quantitative trait locus and network analysis reveals downstream effects and putative drivers for brain-related diseases

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# 28 Abstract

29	Gaining insight into the downstream consequences of non-coding variants is an essential step
30	towards the identification of therapeutic targets from genome-wide association study (GWAS)
31	findings. Here we have harmonized and integrated 8,727 RNA-seq samples with accompanying
32	genotype data from multiple brain-regions from 14 datasets. This sample size enabled us to
33	perform both cis- and trans-expression quantitative locus (eQTL) mapping. Upon comparing the
34	brain cortex <i>cis</i> -eQTLs (for 12,307 unique genes at FDR<0.05) with a large blood <i>cis</i> -eQTL
35	analysis (n=31,684 samples), we observed that brain eQTLs are more tissue specific than
36	previously assumed.
37	
38	We inferred the brain cell type for 1,515 <i>cis</i> -eQTLs by using cell type proportion information.
39	We conducted Mendelian Randomization on 31 brain-related traits using cis-eQTLs as
40	instruments and found 159 significant findings that also passed colocalization. Furthermore, two
41	multiple sclerosis (MS) findings had cell type specific signals, a neuron-specific cis-eQTL for
42	CYP24A1 and a macrophage specific cis-eQTL for CLECL1.
43	
44	To further interpret GWAS hits, we performed trans-eQTL analysis. We identified 2,589 trans-
45	eQTLs (at FDR<0.05) for 373 unique SNPs, affecting 1,263 unique genes, and 21 replicated
46	significantly using single-nucleus RNA-seq data from excitatory neurons.
47	
48	We also generated a brain-specific gene-coregulation network that we used to predict which
49	genes have brain-specific functions, and to perform a novel network analysis of Alzheimer's
50	disease (AD), amyotrophic lateral sclerosis (ALS), multiple sclerosis (MS) and Parkinson's

disease (PD) GWAS data. This resulted in the identification of distinct sets of genes that show
significantly enriched co-regulation with genes inside the associated GWAS loci, and which
might reflect drivers of these diseases.

54

# 55 Introduction

56 Diseases of the brain manifesting as psychiatric or neurological conditions continue to be a 57 massive global health burden: The World Health Organization estimates that in 2019 globally 58 280 million individuals were affected by depression, 39.5 million by bipolar disorder, and 287.4 59 million by schizophrenia<sup>1</sup>. Likewise, the fraction of 50 million people living with dementia today is expected to rise to 152 million by  $2050^2$ , with similar trajectories for other neurodegenerative 60 61 diseases. While substantial progress has been made in uncovering the genetic basis of psychiatric 62 and neurological diseases through genome-wide association studies (GWAS), much of how the 63 identified genetic variants impact brain function is still unknown.

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To translate from genetic signals to mechanisms, associations with gene expression levels, or 65 66 expression quantitative trait loci (eQTL) have shown great potential. eQTLs can be divided in 67 direct effects of local genetic variants (cis-eOTLs) and indirect effects of distal variants (transeOTLs). Cis-eOTLs and trans-eOTLs can aid interpretation of GWAS loci in several ways. Cis-68 69 eQTLs aid interpretation by identifying direct links between genes and phenotypes through 70 causal inference approaches such as Mendelian randomization (MR) instrumented on QTLs and 71 genetic colocalization analysis, whereas trans-eQTLs expose sets of downstream genes and 72 pathways on which the effects of disease variants converge.

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74 eOTLs are dynamic features and vary with tissue, cell type and additional factors such as 75 response to stimulation. For an optimal interrogation of GWAS loci, it is therefore desirable to perform eQTL analyses in disease-relevant tissues<sup>3</sup>. To help interpret GWAS of 76 77 neurodegenerative and psychiatric diseases, several brain-derived eQTL studies have been published, including meta-analyses by the PsychENCODE<sup>4</sup> and AMP-AD<sup>5</sup> consortia, which 78 79 cover 1,866 and 1,433 individuals, respectively. However, to yield reliable results, statistical 80 approaches such as MR and colocalization require robust effect size estimates from even larger 81 carefully curated eQTL datasets. Large sample sizes are better suited to decompose eQTL effects 82 to specific cell types.

83

84 To maximize the potential of eQTL-based analyses in brain, we here combined and rigorously 85 harmonized brain RNA-seq and genotype data from 15 different cohorts, including 8,727 RNA-86 seq samples from all major brain eQTL studies and publicly available samples from the 87 European Nucleotide Archive (ENA). By leveraging the statistical power across these datasets, 88 we created a gene coregulation network based on 8,544 RNA-seq samples covering different 89 brain regions and performed *cis*- and *trans*-eQTL analysis in up to 2,970 individuals of European 90 descent, with replication in up to 420 individuals of African descent. This sample size enabled us 91 to make inferences on the brain cell types in which eQTLs operate, and to systematically conduct 92 Mendelian Randomization and colocalization analyses to find shared genetic effects between 93 eQTLs and GWAS traits. This prioritized likely causal genes from GWAS loci for 31 brain-94 related traits, including neurodegenerative and psychiatric conditions. Additionally, this 95 identified cell type dependent eQTLs that may be associated with disease risk (Figure 1). 96

# 97 **Results**

## 98 Leveraging public RNA-seq and genotype data to create large, harmonized brain

#### 99 eQTL and gene co-regulation datasets

- 100 We combined 15 eQTL datasets into the 'MetaBrain' resource to maximize statistical power to
- 101 detect eQTLs and to create a brain specific gene coregulation network (Figure 2;

102 Supplementary Table 1, Supplementary Figures 1-5). *MetaBrain* includes 7,604 RNA-seq

- 103 samples and accompanying genotypes from the AMP-AD consortium<sup>6</sup> (AMP-AD MAYO<sup>6</sup>,
- 104 ROSMAP<sup>6</sup> and MSBB<sup>6</sup>), Braineac<sup>7</sup>, the PsychENCODE consortium<sup>8</sup> (Bipseq<sup>4</sup>, BrainGVEX<sup>4</sup>,
- 105 CMC<sup>9</sup>, GVEX and UCLA\_ASD<sup>4</sup>), BrainSeq<sup>10</sup>, NABEC<sup>11</sup>, TargetALS<sup>12</sup>, and GTEx<sup>3</sup>.
- 106 Additionally, we carefully selected 1,759 brain RNA-seq samples from the European Nucleotide
- 107 Archive (ENA)<sup>13</sup>, calling and imputing genotypes based on the RNA-seq alignment
- 108 (Supplementary Note, Supplementary Figure 1). There were 8,727 RNA-seq samples
- 109 remaining after realignment and stringent quality control (Methods and Supplementary Note,
- 110 **Supplementary Figure 2-3**). Using slightly different quality control measures, we created a
- 111 gene network using 8,544 samples (Supplementary Note). We corrected the RNA-seq data for
- technical covariates and defined 7 major tissue groups (amygdala, basal ganglia, cerebellum,
- 113 cortex, hippocampus, hypothalamus and spinal cord): Principal Component Analysis (PCA) on
- the RNA-seq data showed clear clustering by these major tissue groups, resembling brain
- 115 physiology (Figure 2D, Supplementary Figure 4). Genotype data revealed individuals from
- 116 different ethnicities (Figure 2B; Supplementary Figure 5), including 5,138 samples from
- 117 European descent (EUR) and 805 samples from African descent (AFR). We created 6 cis-eQTL
- 118 discovery datasets: Basal ganglia-EUR (n=208), Cerebellum-EUR (n=492), Cortex-EUR
- 119 (n=2,970), Cortex-AFR (n=420), Hippocampus-EUR (n=168) and Spinal cord-EUR (n=108;

120	Supplementary Table 1, Figure 2C). Cis-eQTLs were not calculated for amygdala and
121	hypothalamus tissue groups due to the small sample size (n<100).

- 122
- 123

#### 124 **41%** of the cortex *cis*-eQTL genes are regulated by multiple independent variants

125 Within each discovery dataset, we performed a sample-size weighted *cis*-eQTL meta-analysis on

126 common variants (MAF>1%), within 1 megabase (Mb) of the transcription start site (TSS) of a

127 protein-coding gene. We identified 1,317 (Basal ganglia-EUR), 6,865 (Cerebellum-EUR), 5,440

128 (Cortex-AFR), 11,803 (Cortex-EUR), 990 (Hippocampus-EUR), and 811 (Spinal cord-EUR) cis-

129 eQTL genes (FDR<0.05; Figure 3A; Supplementary Table 2). *Cis*-eQTL effect directions were

130 highly concordant between datasets included in the Cortex-EUR meta-analysis (median

131 Spearman r=0.80; median allelic concordance=89%; **Supplementary Figure 6**), indicating

132 robustness of the identified effects across datasets. We observed that significant *cis*-eQTL

133 findings were sensitive to RNA-seq alignment strategies, and it is difficult to confidently

ascertain *cis*-eQTLs in regions with multiple haplotypes represented on patch chromosomes, like

the *MAPT* locus on 17q21 (Supplementary Note, Supplementary Figures 7-9). We next

136 performed conditional analysis to identify independent associations in each cis-eQTL locus (e.g.,

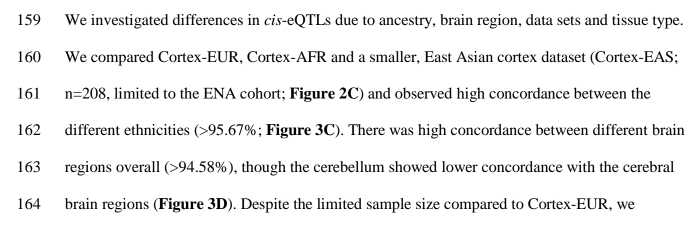
137 secondary, tertiary and quaternary eQTLs). In Cortex-EUR, 4,791 genes had a significant

138 secondary *cis*-eQTL (41% of *cis*-eQTL genes identified in this dataset). 1,658 genes had tertiary

and 598 had quaternary cis-eQTLs. We also identified secondary associations for the other

140 discovery datasets albeit to a lesser extent (Figure 3A; Supplementary Table 2 and 3).

The properties of the Cortex-EUR *cis*-eOTLs conform to studies performed earlier in blood<sup>14</sup> and 142 143 brain<sup>15</sup> (Figure 3B): primary lead *cis*-eQTL SNPs were generally located close (median distance: 144 31 kilobase; kb) to the transcription start site (TSS; Figure 3B) and *cis*-eQTL genes had a lower probability for loss of function intolerance (pLI;  $\chi^2$  p=6.35x10<sup>-147</sup>). Genes with a *cis*-eQTL 145 generally had a higher median expression than those without (Wilcoxon p-value:  $9.96 \times 10^{-12}$ ). 146 147 Contrary to blood, where genes in the highest expression decile are the most likely to have a *cis*-148 eQTL, the third decile of gene expression had the most *cis*-eQTLs in cortex, and higher deciles 149 had increasingly lower proportions of eQTLs (Supplementary Note, Supplementary Figure 150 **10A**). This could suggest that highly expressed genes in the cortex have tighter genetic 151 regulation than highly expressed genes in the blood, although we did not observe differences 152 when comparing variance per gene expression decile between blood and brain (Supplementary 153 Note, Supplementary Figure 10B). Cortex-EUR cis-eQTL genes showed limited functional enrichment for human phenotype ontologies (HPO), GO ontologies and TRANSFAC<sup>16</sup> 154 155 transcription factor motifs (Supplementary Figure 10C and D, Supplementary Table 4). We 156 observed similar patterns for secondary, tertiary and quaternary *cis*-eQTLs (Supplementary 157 Note). 158



165 identified 846 *cis*-eOTLs that were unique to Cerebellum-EUR (**Supplementary Figure 11A**). 166 Of the 846 Cerebellum-EUR unique *cis*-eQTL genes, 184 had low gene expression levels in 167 cortex, which may explain why they did not have a *cis*-eQTL in that tissue (**Supplementary** 168 Figure 11B, C, Supplementary Note). For the remaining 662 genes that were highly expressed 169 in both cortex and cerebellum, we performed functional enrichment of transcription factor 170 binding sites (TFBS; Supplementary Table 5, Supplementary Note) and determined that these 171 genes were enriched for TFBS of 101 distinct transcription factors. Five of these transcription 172 factors had low gene expression in cortex and high expression in cerebellum (EOMES, TFAP2B, 173 TFAP2A, IRX1 and IRX5, Supplementary Figure 11D). These transcription factors might 174 explain the difference in cis-eQTL genes found in cerebellum but not in cortex, while many of 175 these cis-eQTL genes are expressed in both tissues. Next, we compared Cortex-EUR cis-eQTLs 176 with different tissues from the GTEx project (Figure 3E; Supplementary Figure 12, 177 **Supplementary Table 6**). There was high concordance in brain-related tissues (cerebral tissues, 178 >98% and cerebellar tissues, >94%) compared to other tissue types, and the lowest concordance 179 rates were observed in testis (84%) and whole blood (85%). We also compared Cortex-EUR ciseOTLs with eOTLGen<sup>17</sup>, a large blood-based eOTL dataset (n=31.684; majority EUR ancestry) 180 181 and observed a 76% concordance rate (Supplementary Figure 13; Supplementary Table 7) 182 with a moderate correlation of *cis*-eQTL effect sizes ( $R_{b}$ =0.54 including all eQTLs, or  $R_{b}$ =0.62 when pruning genes within 1Mb)<sup>18</sup>, supporting the lower concordance observed in GTEx-blood. 183 184 Since we found that 24% of the shared *cis*-eQTLs between blood and brain showed opposite 185 allelic effects, these results suggest that with larger sample sizes, more tissue specific regulatory 186 variants can be identified. If a causal tissue-specific regulatory variant resides on a haplotype that 187 also contains a variant that is specific for another tissue, it is well conceivable that opposite

allelic effects are going to be observed when contrasting eQTLs for these two tissues<sup>19</sup>. Since the

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189 procedures for eQTL mapping were identical between *MetaBrain* and eQTLGen, our results 190 highlight the relevance of tissue-specific eQTL mapping to accurately assess the directionality of eOTLs, which can elucidate eOTLs with opposite allelic effects<sup>20</sup>. This direct comparison 191 192 illustrates the importance of investigating the appropriate tissue type for the interpretation of 193 GWAS signals. 194 195 8% of Cortex *cis*-eQTLs are mediated by cell type proportion differences 196 Cell type dependent eQTLs can be identified in bulk RNA-seq data by performing cell type deconvolution and determining cell type interaction eOTLs (ieOTLs)<sup>3,21,22</sup>. We predicted five 197 198 major cell types using single cell RNA-seq derived signature profiles<sup>23</sup>. Of these, neurons were 199 the most abundant cell type (median cell proportion: 32.8%), followed by endothelial cells 200 (24.9%), macrophages (17.8%), oligodendrocytes (12.4%) and astrocytes (12.1%; 201 Supplementary Figure 14). We predicted similar proportions for cerebellum as well as other 202 brain regions. We observed that predicted cell proportions are different for spinal cord, showing 203 a relatively low proportion of neuronal cells and high proportions of macrophage and oligodendrocytes compared to other brain tissues, as was previously reported<sup>24</sup> (Supplementary 204 205 Figures 15 and 16). Predicted neuron proportions in both cortex and cerebellum were negatively 206 correlated with the predicted proportions of other cell types, and predicted endothelial cell 207 proportions were negatively correlated with predicted macrophage proportions (Figure 4A). 208 Predicted cell type proportions were positively correlated with immunochemistry (IHC) counts from the ROSMAP cohort<sup>25</sup>, both overall (Spearman r=0.71; Figure 4B) and per individual cell 209

type (Spearman r>0.1; Figure 4B). It is difficult to validate these cell type proportion predictions

due to the small scale of the IHC experiment, but also because IHC and bulk RNA-seq reflect
different aspects of gene or protein expression. Thus, there is a level of uncertainty for the
expected proportion for each cell type<sup>26,27</sup>.

215	With these predicted cell type proportions, we used DeconQTL <sup>22</sup> to identify interaction-eQTLs
216	(ieQTLs) by testing 18,850 cis-eQTLs in Cortex-EUR and 8,347 cis-eQTLs in cerebellum
217	(including primary, secondary, tertiary and quaternary eQTLs). We identified 1,515 significant
218	ieQTLs (8%) in at least one cell type (Benjamini-Hochberg; BH FDR<0.05) for Cortex-EUR
219	(Supplementary Table 8). Of these, 632 (42%) were an ieQTL in neurons, likely because this is
220	the most prevalent cell type. The majority of the ieQTLs (90.2%) were uniquely mapped to one
221	cell type (Figure 4C). Although we observed a lower proportion of ieQTLs in cerebellum (126;
222	1.5%, Supplementary Figure 17, Supplementary Table 8), this is likely a power issue due to
223	the smaller sample size. While we observed the most ieQTLs for neurons in cortex, the majority
224	(n=106; 84%) of ieQTLs in cerebellum were mediated by astrocytes and macrophages.
225	
226	We compared the allelic direction of the identified ieQTLs for each cell type with matching cell
227	types from a single nucleus RNA-seq (snRNA-seq) dataset (ROSMAP cohort, n=39;
228	<b>Supplementary Table 9</b> ) $^{28}$ . When filtering on cell type mediated eQTLs by Decon-QTL
229	(FDR<0.05), we observed a high average concordance in allelic direction for both the eQTL
230	main effect (68%), as well as the direction of the interaction (68%; Supplementary Figure
231	<b>18B</b> ). 106 of the cortex <i>cis</i> -ieQTLs were also significant (BH FDR<0.05) in the snRNA-seq
232	datasets (63 in excitatory neurons and 43 in oligodendrocytes). Of these, 13 excitatory neuron
233	and 21 oligodendrocyte ieQTLs were cell type mediated by the corresponding cell type in bulk

234 with 100% allelic concordance (Decon-OTL; BH FDR<0.05; Supplementary Figure 18D). The 235 ieQTLs replicating in oligodendrocytes included STMN4, NKAIN1, and FAM221A (Figure 4D 236 and E and Supplementary Figure 19A-C), which have previously been identified as oligodendrocyte specific<sup>29</sup>. Additionally, this set of ieOTLs included AMPD3 (rs11042811) and 237 238 CD82 (rs2303865), genes involved in the white matter microstructure<sup>30</sup>, suggesting a role for 239 oligodendrocytes in this pathway. The ieQTLs replicating in excitatory neurons included 240 SLC25A27 (alias UCP4; Figure 4F and Supplementary Figure 19D), a gene principally expressed in neurons<sup>31</sup> that modulates neuronal metabolism<sup>32</sup>. The eQTL SNP for this gene, 241 rs2270450, is in high LD ( $r^2=0.71$ ) with a variant previously associated with schizophrenia<sup>33</sup>. 242 Previous work has suggested a possible role of this gene in Parkinson's disease<sup>34,35</sup>. These results 243 244 suggest that the decomposition of eQTLs to their relevant cell types in *MetaBrain* yields 245 additional valuable information about the underlying biological mechanisms of genes and cell 246 types of interest for genes associated with disease. 247

#### 248 Shared genetic effects between Cortex-EUR *cis*-eQTLs and brain-related traits

As one application of the *MetaBrain* resource, we linked *cis*-eQTLs to variants associated with brain-related traits and diseases. For this, we first evaluated linkage disequilibrium (LD) between the Cortex-EUR *cis*-eQTL SNPs with the strongest association signals and index variants identified in 1,057 GWASs of brain-related traits (**Supplementary Note, Supplementary Table 10**). We observed that 10% of brain-related trait SNPs for 242 eQTL genes were in LD with *cis*eQTL SNPs (r<sup>2</sup>>0.8). This percentage marginally increased to 12% when secondary, tertiary and quaternary eQTL SNPs were included, indicating that the majority of LD overlap is driven by

256	primary eQTL effects: primary eQTLs were 3.3-fold more likely to be in LD with a GWAS SNP
257	(Fisher exact test p-value = $6.2 \times 10^{-16}$ ; Supplementary Note).

259	To more formally test for overlap between GWAS and cis-eQTL signals, we conducted
260	Mendelian randomization (MR) to test for a causal effect between gene expression and 31
261	neurological traits using cis-eQTLs as instruments (Supplementary Table 11). We computed a
262	Wald ratio for each eQTL instrument, from which 1,192 Wald ratios out of 268,030 tested in
263	total passed a suggestive p-value threshold ( $p < 5x10^{-5}$ ; Supplementary Table 12). 120 of the <i>cis</i> -
264	eQTL instruments from these suggestive findings were also cell type ieQTLs. We further
265	prioritized our list of genes with evidence of Wald ratio effects by determining genetic
266	colocalization between GWAS and <i>cis</i> -eQTL signals using coloc <sup>36</sup> . There were 159 significant
267	Wald ratios that passed a strict Bonferroni correction ( $p<1.87x10^{-7}$ ) where the GWAS SNP and
268	eQTL colocalized (PP4>0.7; Figure 5A; Supplementary Figure 20). 69 of these prioritized
269	findings were associated with neurological and neuropsychiatric disease risk (Table 1). Three
270	examples where MR and colocalization pointed to likely causal GWAS genes are reported
271	below, for others, see Supplementary Note, Supplementary Tables 11-16 and
272	Supplementary Figures 21 and 22.
273	

## 274 MR comparison between blood and brain eQTL datasets

MR analysis for multiple sclerosis (MS)<sup>37</sup> identified 102 instruments in 83 genes that passed the
Bonferroni-adjusted p-value threshold (Supplementary Table 12). 20 of these findings passed
colocalization (Table 1; Figure 5B). This included 11 genes for which MR suggested that
increased gene expression and 9 genes where decreased gene expression may confer MS risk.

279	Systematic comparison of the Wald ratio estimates for MS of 5,919 shared cis-eQTL genes
280	between Cortex-EUR and eQTLGen (where the same gene was instrumented but could be with
281	different SNPs) <sup>17</sup> showed opposite directions of effect for 2,291 (38.7%) genes (Supplementary
282	Figure 23, Supplementary Table 14). Agreement improved when the same SNP instrument
283	was compared between studies, but discordance still remained high with 1,891 (26%) out of
284	7,274 MetaBrain Wald ratios showing opposite directionality to eQTLGen (Supplementary
285	Table 15). The notable discordance in the directionality of the blood and brain eQTLs
286	underscore the importance of tissue-specific differences when interpreting transcriptomics data.
287	
288	Of the 135 genes with MR findings in Cortex-EUR for MS, there were 28 genes without a
289	significant eQTLGen instrument, including 3 genes (SLC12A5, CCDC155 and MYNN) for which
290	we found both MR significance and colocalization in MetaBrain (Supplementary Note;
291	Supplementary Table 16. Comparing blood and brain gene expression levels for these genes in
292	GTEx, SLC12A5 had almost no expression in blood, while expression was comparable between
293	tissues for CCDC155 and MYNN (Supplementary Note, Supplementary Figure 24). The
294	discrepancy in MR findings observed between Cortex-EUR and eQTLGen suggest tissue-
295	dependent genetic effects for MS.
296	
297	MR and colocalization analysis links multiple sclerosis GWAS loci to cell type specific
298	eQTLs for CYP24A1 and CLECL1
299	Two MS genes, CYP24A1 and CLECL1, showed cell type specific cis-eQTLs (Figure 5C and
300	<b>D</b> ). Another gene that was previously suggested to be neuron specific <sup>38</sup> , <i>SLC12A5</i> , did not show
301	a significant ieQTL in our data. Our analysis used rs2259735 as the Cortex-EUR eQTL

302	instrument variable and suggested that higher expression of CYP24A1 is associated with
303	increased MS risk (MR Wald ratio= $0.13$ , p= $1.7 \times 10^{-9}$ ). We also observed colocalization of the
304	cis-eQTL and the MS GWAS signal at this region (coloc PP4=0.99), suggesting the same
305	underlying genetic signal. Furthermore, ieQTL analysis showed increasing expression of
306	CYP24A1 with increasing neuronal proportions for the MS risk allele rs2248137 (interaction
307	beta=2.85; interaction FDR= $1 \times 10^{-308}$ ; Figure 5C). Rs2248137 has previously been associated
308	with MS <sup>39</sup> and is in strong LD with SNP rs2259735 ( $r^2=0.9$ ). <i>CYP24A1</i> is a mitochondrial
309	cytochrome P450 hydroxylase that catalyzes the inactivation of 1,25-dihydroxyvitamin $D_3$
310	(calcitriol), the active form of vitamin $D^{40}$ . Loss of function mutations in <i>CYP24A1</i> increase
311	serum calcitriol and cause hereditary vitamin D-mediated PTH-independent hypercalcemia <sup>41,42</sup> .
312	In the brain, vitamin D plays vital functions in regulating calcium-mediated neuronal
313	excitotoxicity, reducing oxidative stress and regulating synaptic activity <sup>43</sup> . Epidemiological
314	studies have proposed vitamin D deficiency as a risk factor for MS <sup>44,45</sup> , which has recently been
315	validated through $MR^{46-48}$ . Our findings are consistent with a previous report of a shared MS
316	GWAS signal and CYP24A1 cis-eQTL signal with frontal cortex but not white matter, using a
317	brain eQTL dataset derived from expression microarrays to confirm the findings in the same
318	direction of effect <sup>49</sup> .

As another MS signal that passed MR and colocalization, decreased expression of *CLECL1* was associated with increased MS risk (MR Wald ratio=-0.16, p=1.58x10<sup>-9</sup>, coloc PP4>0.92). The ieQTL analysis indicated that the rs7306304 allele increased expression of *CLECL1* with increasing macrophage proportion (interaction beta=-3.65; interaction FDR=1x10<sup>-308</sup>, **Figure 5D**), confirming a previous finding of a microglia cell-type specific *cis*-eQTL for *CLECL1* at this 325 MS risk locus<sup>39</sup>. Rs7306304 is in strong LD with the MS lead SNP, rs7977720  $(r^2=0.84)^{39}$ .

326 CLECL1 encodes a C-type lectin-like transmembrane protein highly expressed in dendritic and B

327 cells that has been proposed to modulate immune response<sup>50</sup>. *CLECL1* was previously found to

328 be lowly expressed in cortical bulk RNA-seq data, while having a 20-fold higher expression in a

329 purified microglia dataset<sup>39</sup>, suggesting that decreased *CLECL1* expression increases MS

330 susceptibility through microglia-mediated dysregulation of immune processes in the brain.

331

#### 332 *MetaBrain* allows for the identification of *trans*-eQTLs

333 Trans-eQTL analysis can identify the downstream transcriptional consequences of disease 334 associated variants. However, we have previously observed in blood that trans-eQTL effect-sizes 335 are usually small. Here we studied whether this applies to brain as well. In order to maximize 336 sample size and statistical power, we performed a *trans*-eQTL analysis in 3,111 unique 337 individuals. We reduced the number of tests performed by limiting this analysis to 130,968 338 unique genetic variants: these include variants that have been previously found to be associated 339 with diseases and complex traits through GWAS and variants that were primary, secondary, 340 tertiary or quaternary lead cis-eQTL SNPs from any of the aforementioned discovery datasets. 341

We identified 3,940 *trans*-eQTLs (FDR<0.05), of which 2,589 (66%) were significant after</li>
removing *trans*-eQTLs for which the gene that partially map within 5Mb of the *trans*-eQTL SNP
(Supplementary Note; Figure 6A; Supplementary Table 17). These 2,589 eQTLs reflect 373
unique SNPs, and 1,263 unique genes. 222 (60%) of the *trans*-eQTL SNPs were a *cis*-eQTL
SNP, of which 42 (19%) were a *cis*-eQTL index SNP in Cortex-EUR, and 22 (10%) in tissues

other than cortex. This suggests that *trans*-eQTLs can also be observed for *cis*-eQTLs index
SNPs identified in other tissues (Supplementary Table 17).

349

350 1,060 (83%) of the observed *trans*-eQTL genes were affected by 3 variants at 7p21.3

351 (rs11974335, rs10950398 and rs1990622, LD r<sup>2</sup>>0.95; **Figure 6A and B**; **Supplementary Table** 

352 17). This locus is associated with several brain-related traits, including frontotemporal lobar

353 degeneration<sup>51</sup> and major depressive disorder<sup>52</sup> (**Supplementary Table 17**). The *trans*-eQTL

354 SNP rs1990622 in this locus is the lead GWAS SNP for the TDP-43 subtype of frontotemporal

355 lobar degeneration (FTLD-TDP)<sup>53</sup>, just downstream of *TMEM106B*. Matching previous

356 reports<sup>54,55</sup>, we observed that this locus was associated with predicted neuron proportions

357 (Supplementary Tables 18-20). Moreover, the predicted neuronal proportions were lower in

AD cases than controls (Supplementary Figure 25), which may explain why a large number of

359 trans-eQTLs signals at this region were most pronounced in the AMP-AD datasets and had

360 stronger effect sizes in AD samples (**Supplementary Figure 26 and 27**). We performed

361 functional enrichment on the *trans*-eQTL genes using g:Profiler<sup>56</sup> and observed that upregulated

362 trans-eQTL genes were enriched for neuron related processes such as synaptic signaling

363  $(p=1.3x10^{-28})$  and nervous system development  $(p=2.9x10^{-21})$ . Downregulated genes were

364 enriched for gliogenesis ( $p=1.6x10^{-8}$ ) and oligodendrocyte differentiation ( $p=3.1x10^{-6}$ ;

365 Supplementary Table 21). Surprisingly, 21 of these *trans*-eQTLs were also significant (BH

366 FDR<0.05) in the snRNA-seq data of excitatory neurons with 100% allelic concordance

367 (Supplementary Figure 28; Supplementary Table 22), suggesting that some of these trans-

368 eQTLs might not be driven by differences in neuron proportions. A detailed description of this

369 locus can be found in the **Supplementary Note**.

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570	
371	We observed <i>trans</i> -eQTLs from multiple independent genomic loci for 14 genes, suggesting
372	convergent <i>trans</i> -eQTL effects (Supplementary Table 17). The genes with these convergent
373	trans-eQTL effects were previously associated with immunological phenotypes (HBG2, PIWIL2,
374	and SVEP1), brain-related phenotypes (DAZAP2), immunological and brain-related phenotypes
375	(HMCES, KCNA5, MBTPS1, PRPF19, PTH2R and RFPL2) or other phenotypes (ANKRD2,
376	<i>PEX12, PROM1</i> and <i>ZNF727</i> ).
377	
378	Encouragingly, some of these convergent trans-eQTLs have already been previously identified
379	in blood. For example, two independent variants (rs1427407 on 2p16.1 and rs4895441 on
380	6q23.3) affected hemoglobin subunit gamma-2 ( <i>HBG2</i> ) on 11p15.4 in <i>trans</i> (Figure 6C). These
381	variants have previously been associated with fetal hemoglobin levels <sup>57–59</sup> and various blood cell
382	counts.
383	
384	We also observed converging effects that were not identified in blood. For instance, KCNA5
385	(12p13.32) was affected by variants from three independent loci at 2p23.3 (rs930263), 4p15.32
386	(rs2702575 and rs2604551) and 7p21.3 (rs10950398 and rs11974335) as described in Figure
387	6C; Supplementary Table 17. KCNA5 encodes the potassium voltage-gated channel protein
388	Kv1.5. Potassium voltage-gated channels regulate neuron excitability among other functions, and
389	blockers for these channels have been suggested as a therapeutic target for multiple sclerosis
390	patients <sup>60</sup> . Furthermore, <i>KCNA5</i> has previously been associated with cardiovascular disease <sup>61</sup> ,
391	and has been suggested to modulate macrophage and microglia function <sup>62</sup> . Three <i>cis</i> -eQTLs
392	were associated with rs930263, including ADGRF3, DRC1, and a secondary eQTL on HADHB.

rs930263 was previously associated with sleep dependent LDL levels<sup>63</sup> and several blood
metabolite levels<sup>64-67</sup>. The 4p15.32 locus was previously associated with insomnia and adult
height<sup>68</sup> and the 7p21.3 locus with depression and blood protein levels. These results thus
suggest that several sleep related variants affect potassium voltage-gated regulation of neuron
excitability.

398

This is the first report of *trans*-eQTLs in the brain cortex for many of the variants identified, and our results indicate that many of these signals are brain-specific. We observed the *trans*-eQTL

401 effect-sizes in brain are usually small, similar to what we previously observed in blood,

402 emphasizing the importance of increasing the sample-size of brain eQTL studies.

403

#### 404 Brain co-regulation networks improve GWAS interpretation

We generated brain-region specific co-regulation networks based on the RNA-seq data from 8,544 samples (**Supplementary Note, Supplementary Figures 29-30**). We previously have done this for a heterogenous set of RNA-seq samples spanning across all available tissue types and cell lines  $(n=31,499)^{69,70}$ , which showed that such a co-regulation network can be informative for interpreting GWAS studies<sup>69</sup> and helpful in the identification of new genes that cause rare diseases<sup>70</sup>.

411

We applied a new approach ('*Downstreamer*', in preparation, see Supplementary Note) that
improves upon DEPICT, our previously published post-GWAS pathway analysis method<sup>69</sup>. *Downstreamer* can systematically determine which genes are preferentially co-regulated with
genes that reside within GWAS loci. It does not use a significance threshold for a GWAS, but

416	instead uses all SNP information. In addition, Downstreamer accounts for LD and uses rigorous
417	permutation testing to determine significance levels and control for Type I errors.
418	
419	We applied <i>Downstreamer</i> to schizophrenia (SCZ) <sup>71</sup> , PD <sup>72</sup> , MS <sup>37</sup> , AD <sup>73</sup> and ALS GWAS
420	summary statistics (Supplementary Table 23-27), using three different brain-derived co-
421	regulation networks: one based on all 8,544 brain samples, one limited to 6,527 cortex samples
422	and one limited to 715 cerebellum samples. We observed that there were multiple sets of genes
423	that showed strong co-regulation with genes inside the GWAS loci for these diseases. For MS
424	and AD, these were mostly immune genes, whereas for PD, ALS and SCZ these were genes that
425	are specifically expressed in brain (Supplementary Table 23-27).
426	
427	For ALS, we applied <i>Downstreamer</i> to summary statistics from a recent meta-analysis in
428	individuals from European ancestry (Supplementary Table 28), and a trans-ethnic meta-
429	analysis including European and Asian individuals (EUR+ASN; Supplementary Table 23; van
430	Rheenen et al., manuscript in preparation). To look for contributions of non-neurological cell
431	types and tissues, we first used the previously published heterogenous network <sup>70</sup> that comprises
432	many different tissues and cell types, but did not identify genes that were significantly enriched
433	for co-regulation with genes inside ALS loci. However, when we applied our method to the
434	different brain co-regulation networks, we identified a set of 27 unique co-regulated genes
435	(EUR+ASN summary statistics; Figure 7A; Supplementary Table 23), depending on the type
436	of brain co-regulation network used. HUWE1 was shared between the brain and cortex co-
437	regulation network analysis, while UBR4 was shared between the cortex and cerebellum
438	analysis. UBR4 is a ubiquitin ligase protein expressed throughout the body. A private UBR4

<ul> <li>role in muscle coordination<sup>74</sup>. <i>UBR4</i> interacts with the Ca<sup>2+</sup> binding protein, calmodulin and Ca<sup>2+</sup></li> <li>dysregulation has been linked to proteins encoded by ALS disease genes and motor neuron</li> <li>vulnerability<sup>75</sup>. We observed in the <i>Downstreamer</i> findings that many of these prioritized genes</li> <li>are co-regulated with each other (<b>Figure 7B</b>), and using our recently developed clinical symptom</li> <li>prediction algorithm<sup>70</sup>, there was an enrichment of genes implicated in causing gait disturbances</li> <li>(<b>Figure 7C</b>). These genes are associated with ALS (highlighted in blue), brain-related disorders</li> <li>(including <i>DNAJC5</i>, <i>HTT</i>, <i>HUWE1</i>, <i>TSC1</i> and <i>YEATS2</i>) or muscle-related disorders (including</li> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	439	mutation, segregated with episodic ataxia in a large three-generation Irish family, implicates its
<ul> <li>vulnerability<sup>75</sup>. We observed in the <i>Downstreamer</i> findings that many of these prioritized genes</li> <li>are co-regulated with each other (Figure 7B), and using our recently developed clinical symptom</li> <li>prediction algorithm<sup>70</sup>, there was an enrichment of genes implicated in causing gait disturbances</li> <li>(Figure 7C). These genes are associated with ALS (highlighted in blue), brain-related disorders</li> <li>(including <i>DNAJC5</i>, <i>HTT</i>, <i>HUWE1</i>, <i>TSC1</i> and <i>YEATS2</i>) or muscle-related disorders (including</li> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	440	role in muscle coordination <sup>74</sup> . <i>UBR4</i> interacts with the $Ca^{2+}$ binding protein, calmodulin and $Ca^{2+}$
<ul> <li>are co-regulated with each other (Figure 7B), and using our recently developed clinical symptom</li> <li>prediction algorithm<sup>70</sup>, there was an enrichment of genes implicated in causing gait disturbances</li> <li>(Figure 7C). These genes are associated with ALS (highlighted in blue), brain-related disorders</li> <li>(including <i>DNAJC5</i>, <i>HTT</i>, <i>HUWE1</i>, <i>TSC1</i> and <i>YEATS2</i>) or muscle-related disorders (including</li> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	441	dysregulation has been linked to proteins encoded by ALS disease genes and motor neuron
<ul> <li>prediction algorithm<sup>70</sup>, there was an enrichment of genes implicated in causing gait disturbances</li> <li>(Figure 7C). These genes are associated with ALS (highlighted in blue), brain-related disorders</li> <li>(including <i>DNAJC5</i>, <i>HTT</i>, <i>HUWE1</i>, <i>TSC1</i> and <i>YEATS2</i>) or muscle-related disorders (including</li> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	442	vulnerability <sup>75</sup> . We observed in the <i>Downstreamer</i> findings that many of these prioritized genes
<ul> <li>(Figure 7C). These genes are associated with ALS (highlighted in blue), brain-related disorders</li> <li>(including DNAJC5, HTT, HUWE1, TSC1 and YEATS2) or muscle-related disorders (including</li> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	443	are co-regulated with each other (Figure 7B), and using our recently developed clinical symptom
<ul> <li>446 (including DNAJC5, HTT, HUWE1, TSC1 and YEATS2) or muscle-related disorders (including</li> <li>447 <i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>448 the function of the positional candidate genes within these loci is still unclear. Our</li> <li>449 <i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>450 genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>451 function.</li> </ul>	444	prediction algorithm <sup>70</sup> , there was an enrichment of genes implicated in causing gait disturbances
<ul> <li><i>KMT2B</i>). While various loci have been identified for both familial and sporadic forms of ALS,</li> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	445	(Figure 7C). These genes are associated with ALS (highlighted in blue), brain-related disorders
<ul> <li>the function of the positional candidate genes within these loci is still unclear. Our</li> <li><i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>function.</li> </ul>	446	(including DNAJC5, HTT, HUWE1, TSC1 and YEATS2) or muscle-related disorders (including
<ul> <li>449 <i>Downstreamer</i> analysis identified genes that show strong coregulation with positional candidate</li> <li>450 genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>451 function.</li> </ul>	447	<i>KMT2B</i> ). While various loci have been identified for both familial and sporadic forms of ALS,
<ul> <li>450 genes inside ALS loci, suggesting that these positional candidates must have a shared biological</li> <li>451 function.</li> </ul>	448	the function of the positional candidate genes within these loci is still unclear. Our
451 function.	449	Downstreamer analysis identified genes that show strong coregulation with positional candidate
	450	genes inside ALS loci, suggesting that these positional candidates must have a shared biological
452	451	function.
	452	

453 For MS, the heterogeneous network, including many blood and immune cell type samples, 454 identified 257 unique genes that showed significantly enriched co-regulation with genes inside 455 MS loci (Figure 7D; Supplementary Table 27), and many were immune genes, which is also expected for this disease. However, when we applied the brain co-regulation networks, we 456 457 identified a much smaller set of genes, and these genes showed strong enrichment for genes 458 involved in the neurotrophin signaling pathway (Figure 7E and F). Neurotrophins are 459 polypeptides secreted by immunological cell types. In the brain, neurotrophin concentrations are 460 important to promote the survival and proliferation of neurons as well as synaptic transmission. 461 In MS patients, neurotrophin reactivity is higher in MS plaques, whereby neurotrophins are

462 released by peripheral immune cells directly to the inflammatory lesions, suggesting a protective role of this signaling process<sup>76,77</sup>. Neutrophins are also released by glial cells in the brain, 463 464 including microglia and astrocytes, and their role in stimulating neuronal growth and survival could also contribute to an overall neuroprotective effect<sup>78</sup>. In the heterogeneous network, we 465 466 observed high expression for these genes in immune-related tissues (Supplementary Figure 467 **31A**), supporting the "outside-in hypothesis" that the immune system may be a potential trigger for MS<sup>37,79</sup>. The brain specific network showed high expression in spinal cord and cerebellum 468 469 but lower expression in cortex samples (Supplementary Figure 31B), which could be 470 highlighting the specific biological processes taking place in these CNS regions that lead to 471 disease. For example, the cerebellum is responsible for muscle coordination and ataxia occurs in approximately 80% of MS patients with symptoms<sup>80</sup>. We speculate that both dysregulation of the 472 473 immune system and dysregulation of certain neurological processes is a prerequisite for 474 developing MS.

475

## 476 **Discussion**

We here describe an integrated analysis of the effects of genetic variation on gene expression
levels in brain in over 3,000 unique individuals. This sample size yielded sufficient statistical
power to identify robust *cis*-eQTLs and to our knowledge for the first-time brain *trans*-eQTLs
that emanate from SNPs previously linked to neurodegenerative or psychiatric diseases.

481

482 We compared *cis*-eQTLs in *MetaBrain* to *cis*-eQTLs in eQTLGen from a set of 31,684 blood

483 samples. We observe a large proportion of shared *cis*-eQTLs between brain and blood, most of

484 which have the same allelic direction of effect. Our analysis also permitted us to identify cis-

eQTL effects that are independent of the primary *cis*-eQTLs. Some of these independent effects
reflect SNPs that are also the index variants for several neurological and psychiatric disorders,
making them particularly interesting for subsequent follow-up. Recent observations have
revealed that SNPs with the strongest *cis*-eQTL effects are depleted for GWAS associations<sup>81</sup>.
Thus, secondary, tertiary or quaternary *cis*-eQTL SNPs could potentially be even more
interesting to follow-up than certain primary *cis*-eQTL SNPs to link association signals to
function.

492

493 We studied different regions in the brain, permitting us to identify brain-region specific eQTLs. 494 For this, to exclude spurious differences that may arise from different cell type proportions 495 across brain regions, we first inferred cell type percentages for the major brain cell types. We 496 then applied an eQTL interaction model (i.e., using the cell type percentage x genotype as 497 interaction term), permitting us to identify 1,515 cis-eQTLs that show cell type specificity. Most 498 of these cell type dependent effects were observed for oligodendrocytes and neurons, the two 499 most common cell types in the brain for which statistical power to observe such effects was the 500 strongest. Still, we could identify 461 cell type dependent eQTLs also for macrophages, 501 endothelial cells, or astrocytes. While we found strong concordance with immunohistochemistry 502 results, our findings are largely based on a deconvolution approach, which in future studies will 503 benefit from validation in purified cell types, e.g. using population-based single-cell RNA-seq datasets as they are now becoming available<sup>82,83</sup>. Such single-cell eQTL studies can gain 504 505 substantial statistical power by limiting analyses to the large set of primary, secondary, tertiary 506 and quaternary *cis*-eQTLs our study reveals for bulk brain samples.

508 To our knowledge, this is the best powered Mendelian randomization and colocalization analysis 509 using brain *cis*-eQTLs as instruments for bipolar disease, epilepsy, frontotemporal dementia, 510 multiple sclerosis, cognitive function and years of schooling GWAS outcomes. Interestingly, 511 also for schizophrenia three signals for CILP2, MAU2 and TM6SF2 met our criteria that had not 512 been reported in a recent psychiatric genomics consortium study<sup>84</sup>, further emphasizing the value 513 of our well-harmonized, large eQTL data set in the tissue type of interest (Supplementary 514 **Note**). Our results also identify increased *CYP24A1* expression as associated with multiple 515 sclerosis risk and propose neurons as the most susceptible cell type to CYP24A1 expression 516 changes and likely active vitamin D levels. The potentially novel role of CYP24A1 in brain could 517 play an important role in MS etiology, as may lowered expression of *CLECL1* in microglia. 518 519 The 2,589 identified *trans*-eQTLs allowed us to gain insights into downstream molecular 520 consequences of several disease-associated genetic variants. Our trans-eQTL analysis focused on 521 a single brain region and SNPs with a known interpretation (i.e. trait-associated variants and *cis*-522 eQTL SNPs). We therefore expect that a genome-wide approach will identify many more *trans*-523 eQTLs. 2,218 of the *trans*-eQTLs were located in a 7p21.3 locus and the genes were strongly 524 correlated with neuron proportions, indicating that cell type proportions can heavily impact 525 trans-eQTL identification. However, 21 of these trans-eQTLs replicated in snRNA-seq data, 526 suggesting that some of these *trans*-eQTLs may also exist in single cells. Excluding the 7p21.3 527 locus, we identified 371 *trans*-eQTLs located elsewhere in the genome, which are less likely due 528 to neuron proportion differences. For several neurological and psychiatric conditions, our 529 analyses indicate pathways that may help to elucidate disease causes and putative intervention 530 points for future therapies.

532	We used the brain-specific co-regulation networks to study several brain-related GWAS studies,
533	with the aim to prioritize genes that show significantly enriched co-regulation with genes inside
534	the associated GWAS loci. For ALS this revealed a limited, but significant set of genes which do
535	not map within associated ALS loci, but that link genes within multiple ALS loci. Follow-up
536	research on these prioritized genes might therefore help to better understand the poorly
537	understood causal pathways that cause ALS. While it is tempting to speculate that these
538	prioritized genes might represent genes that could serve as potential targets for pharmaceutical
539	intervention, follow-up research is needed in order to establish whether these genes play a
540	relevant role in ALS.

541

542 Our study had several limitations. For instance, we performed single tissue eQTL analyses that 543 were limited to a single RNA-seq sample per individual, excluding many RNA-seq samples from 544 the analysis. A joint analysis across tissues, including multiple RNA-seq samples per individual using for example random effects models would further improve power<sup>85,86</sup>, which would be 545 546 especially useful for the future identification of *trans*-eQTLs. Additionally, LD overlap analysis, 547 Mendelian randomization and colocalization are sensitive to many factors, including eQTL and 548 GWAS study sample size, effect size, variant density, LD structure and imputation quality. 549 Differences between study designs may consequently influence the results of such analyses. For 550 example, our colocalization and LD overlap analysis did not include the MAPT gene for 551 Alzheimer's disease. The effect sizes of the *cis*-eQTLs for this gene were limited in our study, 552 since our alignment strategy could not account for the different long-range haplotypes in this 553 locus causing the H1/H2 haplotype separating SNP rs8070723 to have a p-value of 0.2

554 (Supplementary Note). We note that this might be an issue for other genes as well. Future 555 studies using graph-based alignment tools or long read sequencing methods would be required to 556 ultimately determine the true effects on such genes. Our approach combined Mendelian 557 randomization and colocalization, as it is possible for the *cis*-eQTL instrument to coincidentally 558 share association with the GWAS trait due to surrounding LD patterns in the genomic region. We opted to perform single SNP MR because other approaches, such as inverse variance 559 560 weighted<sup>87</sup> (IVW) MR, pool the estimates across many SNP instruments, which for many genes 561 were not available. Potentially, methods such as IVW could be applied to our dataset in the 562 future when genome-wide *trans*-eQTL analysis would identify many more independent 563 instruments per gene. However, MR analyses using QTLs could be susceptible to confounding because of horizontal pleiotropy<sup>88</sup>, where a single gene is affected by multiple indirect effects, 564 565 which is likely to be exacerbated by including *trans*-eOTLs. Our colocalization analysis used a 566 more lenient posterior probability (PP4) threshold of >0.7, which we selected because we 567 performed colocalization only in loci having a significant MR signal, limiting potential false 568 positives. However, our colocalization approach assumed the presence of a single association in 569 each locus, which might not be optimal for *cis*-eQTL loci harboring multiple independent 570 variants, such as for the *TREM2* gene (Supplementary Note). Consequently, our approach may 571 have not detected colocalizing signals in some loci. Recently, colocalization methods were published<sup>89</sup> that do not have this assumption, and consequently may improve future 572 573 colocalization results.

574

575 With the numbers of GWAS loci for brain-related traits and diseases steadily climbing, we 576 expect that our resource will prove itself as a highly valuable toolkit for post-GWAS brain

- 577 research and beyond. Among others, we demonstrate how our dataset can be utilized to
- 578 disambiguate GWAS loci, point to causal pathways and prioritize targets for drug discovery. To
- 579 our knowledge, this is the largest non-blood eQTL analysis ever conducted, providing insights
- 580 into the functional consequences of many disease associated variants. We expect that through
- 581 future integration with single-cell eQTL studies that have higher resolution but lower power, our
- results will help to pinpoint transcriptional effects in specific brain cell types for many disease-
- 583 associated genetic variants.

## 585 Methods

#### 586 Dataset collection and description

- 587 We collected human brain bulk RNA-seq datasets from different resources. Briefly, we collected
- 588 previously published samples from the AMP-AD consortium<sup>6</sup> (AMP-AD MAYO<sup>6</sup>, ROSMAP<sup>6</sup>
- and MSBB<sup>6</sup>), Braineac<sup>7</sup>, the PsychENCODE consortium<sup>8</sup> (Bipseq<sup>4</sup>, BrainGVEX<sup>4</sup>, CMC<sup>9</sup>, GVEX,
- and UCLA\_ASD<sup>4</sup>) from Synapse.org using the Python package synapseclient<sup>90</sup>. The NABEC and
- 591 GTEx datasets were retrieved from NCBI dbGaP, and TargetALS data was provided directly by
- the investigators. For an overview of the number of samples per dataset, see **Supplementary**

**593 Table 1**.

594

- 595 Additionally, we collected public brain bulk RNA-seq samples from the European Nucleotide
- 596 Archive (ENA; Supplementary Table 28). To select only the brain samples, we first
- 597 downloaded the SkyMap database<sup>91</sup>, which provides readily mapped read counts and sample
- annotations. We performed rigorous quality control on this dataset, and selected ENA, excluding
- 599 for example brain cell lines, brain cancer samples, and samples with RNA spike ins (See
- 600 Supplementary Note for more details on this method, Supplementary Figure 1), resulting in
- 601 1,759 samples, and 9,363 samples when combined with the previously published datasets
- 602 (Supplementary Table 1).
- 603

#### 604 RNA-seq data

605 RNAseq data was processed using a pipeline built with molgenis-compute<sup>92</sup>. FASTQ files were 606 aligned against the  $GENCODE^{93}$  v32 primary assembly with  $STAR^{94}$  (version 2.6.1c), while

607 excluding patch sequences (see **Supplementary Note**) with parameter settings:

608	outFilterMultimapNmax = 1, twopassMode Basic, and outFilterMismatchNmax = 8 for paired-
609	end sequences, outFilterMismatchNmax = 4 for single-end sequences. Gene quantification was
610	performed by STAR, similar to gene quantification using HTSeq <sup>95</sup> with default settings. The
611	gene counts were then TMM <sup>96</sup> normalized per cohort using $edgeR^{97}$ (version 3.20.9) with $R^{98}$
612	(version 3.5.1).

To measure FASTQ and alignment quality we used FastQC<sup>99</sup> version 0.11.3), STAR metrics, and Picard Tools<sup>100</sup> (version 2.18.26) metrics (MultipleMetrics, and RNAseqMetrics). Samples were filtered out if aligned reads had <10% coding bases (**Supplementary Figure 3A**), <60% reads aligned (**Supplementary Figure 3B**), or <60% unique mapping. 117 of the RNA-seq samples did not pass this filter, mostly from GTEx<sup>97</sup>. The other quality measurements were visually inspected but contained no outliers.

620

621 RNA-sequencing library preparation, and other technical factors can greatly influence the ability 622 to quantify of gene expression. Therefore, for a given sample such factors often influence the 623 total variation. For example, such issues can be caused by problems during RNA-seq library 624 preparation that led to an increased number of available transcripts to quantify, or conversely, a 625 lack of variation in quantified transcripts (compared to other samples in the dataset). We 626 therefore opted to identify RNA-seq outliers that were not explained by poor RNA-seq alignment 627 metrics. For this purpose, we performed PCA on the RNA data prior to normalization: we 628 reasoned that the first two components capture excess or depletion of variation caused by 629 technical problems. We identified 20 samples that were outliers in the PCA plot of the RNA-seq 630 data, where PC1 was more than 4 standard deviations from the mean (Supplementary Figure

3A). Twenty outlier samples were removed and the principal components were recalculated
(Supplementary figure 3B). We detected and removed 45 additional outlier samples. We
confirmed no additional outlier samples in the third iteration and principal component
calculation, (Supplementary Figure 3C) and 8,868 samples were taken through additional QC.

636 We next removed genes with no variation and then log2-transformated, quantile normalized and 637 Z-score transformed the RNA-seq counts per sample. PCA on the normalized expression data 638 showed that datasets strongly cluster together (Supplementary Figure 4A), likely due to dataset 639 specific technical differences (e.g., single-end versus paired-end sequencing). To correct for this, 640 the normalized expression data was correlated against 77 covariates from different QC tools (FastOC<sup>99</sup>, STAR<sup>94</sup>, and Picard Tools<sup>100</sup>), such as percent protein coding, GC content, and 5' 641 642 prime/3' prime bias. The top 20 correlated technical covariates (% coding bases, % mRNA bases, 643 % intronic bases, median 3' prime bias, % usable bases, % intergenic bases, % UTR bases, % 644 reads aligned in pairs, average mapped read length, average input read length, number of 645 uniquely mapped reads, % reads with improper pairs, number of reads improper pairs, total 646 sequences, total reads, % chimeras, number of HQ aligned reads, number of reads aligned, HQ 647 aligned Q20 bases, HQ aligned bases) were regressed out of the expression data using a linear 648 model. After covariate correction, clustering of datasets in PC1 and PC2 were no longer present 649 (Supplementary Figure 4B).

650

Our collection of RNA-seq samples consisted of 36 different tissue labels, many of which were
 represented by only a few samples. Therefore, we next defined major brain regions present in our
 dataset, including samples from amygdala, basal ganglia, cerebellum, cortex, hippocampus and

654 spinal cord. We noted that some samples (especially from ENA) were not annotated with a

655 specific major brain region. To resolve this, we performed PCA over the sample correlation

656 matrix and then performed k-nearest neighbors on the first two PCs (k=7) to classify samples to

the major brain regions. Using this approach, we defined a set of 86 amygdala, 574 basal ganglia,

658 723 cerebellum, 6,601 cortex, 206 hippocampus, 252 hypothalamus and 285 spinal cord samples

659 (Supplementary Table 1, Figure 2A).

660

#### 661 Genotype data and definition of eQTL datasets

The genotype data for the included datasets was generated using different platforms, including genotypes called from whole genome sequencing (WGS; AMP-AD, TargetALS<sup>12</sup>, GTEx<sup>3</sup>), genotyping arrays (NABEC<sup>11</sup>, Braineac<sup>7</sup>), and haplotype reference consortium (HRC)<sup>101</sup> imputed genotypes (PsychENCODE datasets), or were called from RNA-seq directly (ENA dataset; see **Supplementary Note**). In total, 22 different genotyping datasets were available, reflecting 6,658 genotype samples (**Supplementary Table 1**).

668

669 We performed quality control on each dataset separately, using slightly different approaches per 670 platform. For the array-based datasets, we first matched genotypes using GenotypeHarmonizer<sup>102</sup> 671 using 1000 genomes phase 3 v5a (1kgp) as a reference, limited to variants having MAF >1%, 672 <95% missingness and Hardy-Weinberg equilibrium p-value <0.0001. Genotypes were then imputed using HRC v1.1 as a reference on the Michigan imputation server<sup>103</sup>. In all HRC 673 674 imputed datasets, variants with imputation info score <0.3 were removed. For the WGS datasets, 675 we removed indels and poorly genotyped SNPs having VOSR tranche <99.0, genotype quality 676 <20, inbreeding coefficient <-0.3 and >5% missingness, setting genotype calls with allelic depth

677	<10 and allelic balance <0.2 or >0.8 as missing. WGS datasets were not imputed with HRC.
678	Considering the small size of some of the datasets, we decided to focus further analysis on
679	variants with MAF >1% and Hardy-Weinberg p-value >0.0001.

681 In each dataset, we removed genetically similar individuals by removing individuals with pihat >0.125, as calculated with PLINK 2.0<sup>104</sup>. Additionally, we merged genotypes with those from 682 683 1kgp, pruned genotypes with --indep-pairwise 50 5 0.2 in PLINK, and performed PCA on the 684 sample correlation matrix. We performed k-nearest neighbors (k=7) on the first two PCs, using 685 the known ancestry labels in 1kgp, to assign an ancestry to each genotyped sample. The majority 686 of included samples was of EUR descent: 5,138 samples had an EUR assignment, 805 samples 687 had an AFR assignment, and 573 samples were assigned to the other ethnicities (Supplementary 688 Table 1, Figure 2B).

689

690 For the purpose of eQTL analysis, we next assessed links between RNA-seq and genotype 691 samples and noted that some individuals had multiple RNA-seq samples (e.g. from multiple 692 brain regions) or multiple genotype samples (e.g. from different genotyping platforms). In total, 693 we were able to determine 7,644 links between RNA-seq samples and genotype samples 694 (Supplementary Table 1), reflecting 3,525 unique EUR individuals, 624 unique AFR 695 individuals and 510 unique individuals assigned to other ethnicities. We then grouped linked 696 RNA-seq samples based on ethnicity and tissue group to prevent possible biases on eQTL 697 results. For those individuals with multiple linked RNA-seq samples, we selected a sample at 698 random within these groups. Within each tissue and ethnicity group, we then selected unique 699 genotype samples across datasets in such a way to maximize sample size per genotype dataset.

For the eQTL analysis per tissue, we only considered those datasets having more than 30 unique
linked samples available, and for which at least two independent datasets were available. Using
these criteria for sample and dataset selection, we were able to create 7 eQTL discovery datasets:
Basal ganglia-EUR (n=208), Cerebellum-EUR (n=492), Cortex-EUR (n=2,970), Cortex-AFR
(n=420), Hippocampus-EUR (n=168) and Spinal cord-EUR (n=108; Supplementary Table 1,
Figure 2C).

706

#### 707 eQTL analysis

708 Our dataset consists of different tissues and ethnicities, and samples have been collected in 709 different institutes using different protocols. Consequently, combining these datasets to perform 710 eQTL analysis is complicated, due to possible biases each of these factors may introduce. To resolve this issue, we opted to perform an eQTL meta-analysis within each of the defined eQTL 711 712 discovery datasets. To reduce the effect of possible gene expression outliers, we calculated 713 Spearman's rank correlation coefficients for each eQTL in each dataset separately, and then 714 meta-analyzed the resulting coefficients using a sample size weighted Z-score method, as 715 described previously<sup>14</sup>. While we acknowledge that this method may provide less statistical 716 power than the commonly used linear regression, we chose this method to provide conservative 717 effect estimates. To identify *cis*-eQTLs, we tested SNPs located within 1 Mb of the transcription 718 start site, while for the identification of *trans*-eQTLs, we required this distance to be at least 5 719 Mb. For both analyses, we selected variants having a MAF>1%, and a Hardy-Weinberg p-value 720 >0.0001. Using the GENCODE v32 annotation, we were able to quantify 58,243 genes, of which 721 19,373 are protein coding. While non-coding genes have been implicated to be important for 722 brain function<sup>105</sup>, these genes generally have poor genomic and functional annotations, meaning

723	that it is often unknown in which pathway they function, and that there is uncertainty about their
724	genomic sequence. We therefore focused our eQTL analyses on protein coding genes.
725	

726	To correct for multiple testing, we reperformed the cis- and trans-eQTL analyses, while
727	permuting the sample labels 10 times. Using the permuted p-values, we created empirical null
728	distributions and determined a false discovery rate (FDR) as the proportion of unpermuted
729	observations over the permuted observations and considered associations with FDR $< 0.05$ as
730	significant. To provide a more stringent FDR estimate for our <i>cis</i> -eQTL results, we limited FDR
731	estimation to the top associations per gene, as described previously <sup>14</sup> . We note that our FDR
732	estimate is evaluated on a genome-wide level, rather than per gene, and consequently FDR
733	estimates stabilize after a few permutations <sup>106</sup> .

Since *cis*-eQTL loci are known to often harbor multiple independent associations, we performed an iterative conditional analysis, where for each iteration, we regressed the top association per gene from the previous associations, and re-performed the *cis*-eQTL analysis until no additional associations at FDR<0.05 could be identified.</p>

739

Since a genome-wide *trans*-eQTL analysis would result in a large multiple testing burden
considering the billions of potential tests, we limited this analysis to a set of 130,968 variants
with a known interpretation. This set constituted of variants that were either previously
associated with traits, having a GWAS p-value <5x10<sup>-8</sup> in the IEU OpenGWAS database<sup>107</sup> and
EBI GWAS catalog<sup>108</sup> on May 3<sup>rd</sup>, 2020, and additional neurological traits (see Supplementary
Table 17) or were showing an association with FDR<0.05 in any of our discovery *cis*-eQTL

746	analyses (including secondary, tertiary and quandary associations identified in the iterative
747	conditional analysis). Cis-eQTLs in Cortex-EUR were highly concordant when replicated in
748	Cortex-AFR (Figure 3C). Consequently, to maximize the sample size and statistical power, we
749	meta-analyzed Cortex-EUR and Cortex-AFR datasets together. However, for the trans-eQTL
750	analysis we omitted ENA, to prevent bias by genotypes called from RNA-seq samples.
751	Additionally, For the trans-eQTL analysis, we did not correct the gene expression data for 10
752	PCs, since <i>trans</i> -eQTLs can be driven by cell proportion differences <sup>17</sup> , and many of the first 10
753	PCs in the MetaBrain dataset were correlated with estimated cell type proportions
754	(Supplementary Figure 32). To test for <i>trans</i> -eQTLs, we assessed those combinations of SNPs
755	and genes where the SNP-TSS distance was >5 Mb, or where gene and SNP were on different
756	chromosomes. We note that we did not evaluate eQTLs where the SNP-TSS distance was >1 Mb
757	and <5 Mb, which potentially excludes detection of long-range <i>cis</i> -eQTLs or short-range <i>trans</i> -
758	eQTLs. We expect however, that this excludes only a limited number of eQTLs, since we
759	observed that this distance was <31Kb for 50% of <i>cis</i> -eQTLs (Figure 3B), indicating most <i>cis</i> -
760	eQTLs are short-ranged. Additionally, we reasoned that the >5 Mb cutoff would prevent
761	identification of false-positive trans-eQTLs due to long-range LD.
762	

## 763 Estimation of cell type proportions and identification of cell type mediated eQTLs

By leveraging cell type specific gene expression collected through scRNA-seq, a bulk tissue
sample can be modelled as a parts-based representation of the distinct cell types it consists of. In
such a model, the weights of each part (i.e. cell type proportions) can be determined by
deconvolution. In the deconvolution of the *MetaBrain* bulk expression data we used a single-cell
derived signature matrix including the five major cell types in the brain: neurons,

769 oligodendrocytes, macrophages, endothelial cells and astrocytes. This signature matrix was 770 generated in the context of the CellMap project (Zhengyu Ouyang *et al.*; manuscript in 771 preparation). In short, we created pseudo-bulk expression profiles by extracting gene expression 772 values for specific cell types of interest from annotated single cell and single nuclei expression 773 matrices. Using differential expression analysis and applying several rounds of training and 774 testing, we selected 1,166 differentially expressed genes and calculated the average read counts 775 per cell type. We then filtered out genes that had no variation in expression, leaving a total of 776 1,132 genes. We extracted the corresponding TMM normalized gene counts of these signature 777 genes for all European cortex samples in *MetaBrain*. After correcting the counts for cohort 778 effects using OLS, but not for any other technical covariates, we applied log2 transformation on 779 both the signature matrix as well as the bulk gene count matrix. Subsequently we applied nonnegative least squares (NNLS)<sup>109</sup> using SciPy (version 1.4.1)<sup>110</sup> to model the bulk expression as a 780 781 parts-based representation of the single-nucleus derived signature matrix. First introduced by 782 Lawson and Hanson<sup>109</sup>, NNLS method is the basis of numerous deconvolution methods to date. 783 In short, NNLS attempt to find a non-negative weight (coefficient) for each of the cell types that, 784 when summed together, minimizes the least-squares distance to the observed gene counts. 785 Lastly, we transformed the resulting coefficients into cell type proportions by dividing them over 786 the sum of coefficients for each sample. The resulting cell proportions are then used to identify cell type mediated eQTL effects. For this we applied Decon-eQTL<sup>22</sup> (version 1.4; default 787 788 parameters) in order to systematically test for significant interaction between each cell type 789 proportion and genotype, while also controlling for the effect on expression of the other cell 790 types. The resulting p-values are then correct for multiple testing using the Benjamini-Hochberg 791 method on a per-cell-type basis.

#### 793 Cell type specific ROSMAP single-nucleus datasets

794 In order further confirm cell type specific eQTL effects, we used the ROSMAP single-nucleus 795 data, encompassing 80,660 single-nucleus transcriptomes from the prefrontal cortex of 48 796 individuals with varying degrees of Alzheimer's disease pathology<sup>111</sup>. We used Seurat version  $3.2.2^{112}$  to analyze the data. First, we removed the genes that did not pass filtering as described 797 798 previously<sup>111</sup>, leaving us with 16,866 genes and 70,634 cells for further analysis. After this, we 799 normalized the expression matrix on a per individual per cell type basis using sctransform<sup>113</sup> and visualized the normalized expression matrix using UMAP dimensionality reduction<sup>114</sup>. We 800 observed that cell types, as defined by Mathys *et al*<sup>115</sup>., for the majority cluster together 801 802 (Supplementary Figures 33 and 34). We then created expression matrices for each broad cell 803 type (excitatory neurons, oligodendrocytes, inhibitory neurons, astrocytes, oligodendrocyte 804 precursor cells, microglia, pericytes and endothelial cells) by calculating the average expression 805 per gene and per individual basis. We then used these cell-type datasets for eQTL mapping using 806 the same procedure as the bulk data. To correct for multiple testing, we confined the analysis to 807 only test for primary cis- and trans-eQTLs found in MetaBrain cortex, while also permuting the 808 sample labels 100 times. Lastly, we calculated the Spearman correlation between gene expression levels and genotypes and their 95% confidence intervals<sup>116</sup>. 809

810

#### 811 Single SNP Mendelian Randomization analysis

812 Mendelian Randomization (MR) was conducted between the Cortex-EUR eQTLs and 31

813 neurological traits (21 neurological disease outcomes, 2 quantitative traits and 8 brain volume

814 outcomes) (Supplementary Table 11). Cortex-EUR eQTLs at genome-wide significant

815  $(p < 5x10^{-8})$  were selected and then LD clumped to obtain independent SNPs to form our set of 816 instruments. LD clumping was carried out using the ld clump() function in the ieugwasr package<sup>117</sup> on the default settings (10,000 Kb clumping window with  $r^2$  cut-off of 0.001 using 817 818 the 1000 Genomes EUR reference panel). SNP associations for each of the eQTL instruments 819 were then looked up in the outcome GWASs of interest. If the SNP could not be found in the 820 outcome GWAS using a direct lookup of the dbSNP rsid, then a proxy search was performed to extract the next closest SNP available in terms of pairwise LD, providing minimum  $r^2$  threshold 821 822 of 0.8 with the instrument. Outcome GWAS lookup and proxy search was performed using the 823 associations() function in the ieugwasr package. To ensure correct orientation of effect alleles 824 between the eOTL instrument and outcome GWAS associations, the SNP effects were harmonized using the harmonise data() function in TwoSampleMR<sup>87</sup>. Action 2 was selected 825 826 which assumes that the alleles are forward stranded in the GWASs (i.e. no filtering or re-827 orientation of alleles according to frequency was conducted on the palindromic SNPs). Single 828 SNP MR was then performed on the harmonized SNP summary statistics using the 829 mr\_singlesnp() function in TwoSampleMR. Single SNP MR step computes a Wald ratio, which 830 estimates the change in risk for the outcome per unit change in gene expression, explained 831 through the effect allele of the instrumenting SNP. We reported all the MR findings that passed a p-value threshold of  $5 \times 10^{-5}$ , but note that the Bonferroni-corrected p=0.05 threshold for multiple 832 testing correction is  $p=1.865 \times 10^{-7}$ . We did not implement multi-SNP analysis (such as the 833 834 Inverse Variance Weighted method), because there are a small number of instrumenting SNPs 835 available per gene, which could result in unreliable pooled MR estimates for genes.

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### 837 Colocalization

- 838 Following the MR analysis, colocalization analysis was performed on the MR findings that
- passed the suggestive threshold to determine if the eQTL and trait shared the same underlying
- signal. We ran colocalization<sup>36</sup> using both the default parameters ( $p1=p2=10^{-4}$  and  $p12=10^{-5}$ ) and
- parameters based on the number of SNPs in the region  $(p_1=p_2=1/(number of SNPs in the region))$
- and p12=p1/10). We considered the two traits, eQTL and GWAS outcome to colocalize if either
- 843 of the two parameters yielded PP4>0.7. Additionally, colocalization was systematically analyzed
- against one trait to compare to robustness of the Cortex-EUR eQTLs with existing cortex eQTL
- 845 data sets (see **Supplementary Note**).

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# 847 URLs

- 848 **Picard**: http://broadinstitute.github.io/picard/
- 849 **dbGAP**: https://dbgap.ncbi.nlm.nih.gov
- 850 European Nucleotide Archive: http://www.ebi.ac.uk/ena
- 851 ieugwasr package: https://mrcieu.github.io/ieugwasr/
- 852 **TwoSampleMR**: https://mrcieu.github.io/TwoSampleMR/

853

# 854 Accessions

855 **TargetALS**<sup>12</sup> TargetALS data was pushed directly from the NY Genome center to our sftp

server.

- 857 **CMC**<sup>118</sup> CMC data was downloaded from <u>https://www.synapse.org/</u> using synapse client
- 858 (<u>https://python-docs.synapse.org/build/html/index.html</u>). Accession code: syn2759792
- 859 **GTEx**<sup>86</sup> GTEx was downloaded from SRA using fastq-dump of the SRA toolkit
- 860 (http://www.ncbi.nlm.nih.gov/Traces/sra/sra.cgi?cmd=show&f=software&m=software&s=softw
- 861 <u>are</u>). Access has been requested and granted through dbGaP.
- 862 **Braineac**<sup>7</sup> Braineac data has been pushed to our ftp server by Biogen.
- 863 **AMP-AD**<sup>5</sup> AMP-AD data has been downloaded from synapse<sup>13</sup>. Accession code: syn2580853.
- 864 snRNA-seq was collected using Synapse accession code: syn18485175. IHC data:
- 865 <u>https://github.com/ellispatrick/CortexCellDeconv/tree/master/CellTypeDeconvAnalysis/Data</u>
- 866 **ENA**<sup>13</sup> ENA data has been downloaded from the European Nucleotide Archive. The identifiers
- of the 76 included studies and 2021 brain samples are listed in Supplementary Table 29.

868 (	$CMC_{-}$	HBCC:	CMC_	HBCC	data wa	s downl	oaded	from	https	://www.s	yna	pse.org	<u>/</u> usin	g
-------	-----------	-------	------	------	---------	---------	-------	------	-------	----------	-----	---------	---------------	---

- 869 synapse client (<u>https://python-docs.synapse.org/build/html/index.html</u>). Accession code:
- 870 syn10623034
- 871 BrainSeq BrainSeq data was downloaded from <u>https://www.synapse.org/</u> using synapse client
- 872 (https://python-docs.synapse.org/build/html/index.html). Accession code: syn12299750
- 873 UCLA\_ASD UCLA\_ASD data was downloaded from <u>https://www.synapse.org/</u> using synapse
- 874 client (https://python-docs.synapse.org/build/html/index.html). Accession code: syn4587609
- 875 BrainGVEx BrainGVEx data was downloaded from <u>https://www.synapse.org/</u> using synapse
- 876 client (https://python-docs.synapse.org/build/html/index.html). Accession code: syn4590909
- 877 **BipSeq** BipSeq data was downloaded from <u>https://www.synapse.org/</u> using synapse client
- 878 (https://python-docs.synapse.org/build/html/index.html). Accession code: syn5844980
- 879 GTEx GTEx data was downloaded from dbgap. Accession code: phs000424.v7.p2
- 880 NABEC NABEC data was downloaded from dbgap. Accession code: phs001301.v1.p1
- 881 CellMap single-cell and single-nuclei RNA-seq datasets were downloaded from Gene
- 882 Expression Omnibus (GEO), BioProject, the European Genome-phenome Archive (EGA) and
- the Allan Brain Atlas. Accession codes: GSE97930, GSE126836, GSE103723, GSE104276,
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- 885

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## 907 ROSMAP

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- 914 Genotype data: doi:10.1038/mp.2017.20. RNAseq: doi:10.1038/s41593-018-0154-9. snRNA-
- 915 seq: doi:10.7303/syn18485175
- 916

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- 935

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- 976 dbGaP at http://www.ncbi.nlm.nih.gov/gap through dbGaP accession number phs000424.v7.p2
- 977 on 02/27/2020.

978

### 979 **NABEC**

- 980 Data was collected from dbGAP accession phs001301.v1.p1, which was generated by J. R.
- Gibbs, M. van der Brug, D. Hernandez, B. Traynor, M. Nalls, S-L. Lai, S. Arepalli, A. Dillman,
- 982 I. Rafferty, J. Troncoso, R. Johnson, H. R. Zielke, L. Ferrucci, D. Longo, M.R. Cookson, and
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- 1023

1024 1025

# 1026 Author contributions

- 1027 N.K., O.E.G, and H.W. processed the RNA-seq and genotype data. N.K. and H.W. were
- 1028 responsible for data management. N.K. and H.W. were responsible for the *cis*-eQTL analysis.
- 1029 H.W. was responsible for the *trans*-eQTL analysis. M.V., Z.O. and M.I.Z. were responsible for
- 1030 the cell type proportion prediction. N.K. and M.V. were responsible for the cell type interaction
- 1031 analysis. S.D. was responsible for the selection of brain samples from ENA. D.B., Y.H., C.-Y.C.,
- 1032 E.E.M, T.R.G. and E.A.T. were responsible for MR and colocalization analysis and
- 1033 interpretation. P.D., O.B.B. and L.F. were responsible for the Downstreamer analysis. L.F.,
- 1034 E.A.T. and H.R. acquired funding and supervised the study. N.K., E.A.T., M.V., D.B, Y.H., C.-
- 1035 Y.C., O.B.B., H.R., L.F. and H.W. drafted the manuscript. All authors have proof-read the 1036 manuscript.
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# 1307 Figure Legends

1308 Figure 1. Overview of the study. We downloaded publicly available RNA-seq and genotype

1309 data from 15 different datasets consisting 8,727 RNA-seq measurements from 7 main brain

regions in 6,518 individuals. We performed *cis-*, *trans-* and interaction-eQTL analysis, built a

1311 brain-specific gene coregulation network and prioritized genes using Mendelian randomization,

- 1312 colocalization and the co-regulation network.
- 1313

**Figure 2. Overview of the datasets. (A)** The number of samples per included cohort, with each color representing one of the 7 major brain regions. **(B)** The number of genotypes per cohort,

1316 with each color representing a population. (C) The number of individuals per cohort, with each

1317 color representing an eQTL dataset. The number of individuals is different from the intersection

1318 between the number of RNA-seq samples and number of genotypes, because not all samples

1319 with genotypes have RNA-seq samples and vice-versa, and some individuals with genotypes

1320 have multiple RNA-seq measurements. (**D**) PCA dimensionality reduction plot of the normalized

1321 expression data after covariate correction. Each dot represents an RNA-seq sample and is colored

1322 by brain region. The figure shows that the samples cluster mainly on brain region.

1323

1324 Figure 3. Conditional *cis*-eQTLs. (A) The number of conditional *cis*-eQTLs per eQTL dataset. 1325 (B) Comparison of characteristics between primary and non-primary eQTLs, where each row 1326 compares the eQTL genes for that rank with eQTL genes from the previous rank. P-values are calculated using a Wilcoxon test between significant and non-significant genes. (left) The 1327 1328 difference in mean gene expression levels; (middle) the difference in distance between the most 1329 significant SNP-gene combination and the transcription start site (TSS); (right) the difference in probability for loss of function intolerance (pLI) score. For primary, secondary and quaternary 1330 1331 eQTLs, non-significant eQTLs have higher pLI scores. (C) Replication of primary cis-eQTLs 1332 between the cortex eQTLs of different ethnicities and (**D**) the different brain regions for the 1333 European datasets. n indicates sample size of each dataset. Numbers in boxes indicate the 1334 number of eQTLs that are significant in both the discovery and the replication dataset, and the 1335 percentage of those that shows the same direction of effect. (E) Replication of primary *cis*-1336 eOTLs of Cortex-EUR (discovery) in all the GTEx tissues (replication). Each dot is a different 1337 GTEx tissue, the x-axis is the number of eQTLs that is significant in both discovery and 1338 replication, and the y-axis is the percentage that shows the same direction of effect.

1339

Figure 4. Cell type interacting eQTLs. (A) Spearman correlations between the 5 predicted cell count proportions. Lower triangle is within cortex samples, upper triangle is within cerebellum samples. (B) Predicted cell type proportions (x-axis) compared to cell type proportions measured using immunohistochemistry (IHC; y-axis) for 42 ROSMAP samples. Values in the plot are Pearson correlation coefficients. Cell count predictions for most cell types closely approximates actual IHC cell counts, although neurons are underestimated. (C) Number of cell type interacting eQTLs for Cortex-EUR deconvoluted cell types. The majority of interactions are with neurons 1347 and oligodendrocytes. Notably, most interactions are unique for one cell type in 90% of the 1348 cases. (D, E, F) Replication of cell type interacting eQTLs for STMN4 (D), FAM221A (E) and 1349 SLC25A27 (F), consisting of the scatterplot of the interaction eQTL in MetaBrain Cortex-EUR 1350 bulk RNA-seq (left) and a forest plot for the eQTL effect in the ROSMAP snRNA-seq data 1351 (right). Scatterplot: the x-axis shows the estimated cell type proportion, the y-axis shows the 1352 gene expression, each dot represents a sample. Colors indicate SNP genotype, with yellow being 1353 the minor allele. Values under the alleles are Spearman correlation coefficients. Forest plot: 1354 Spearman coefficients with effect direction relative to the minor allele when replicating the 1355 eQTL effect in ROSMAP single nucleus data (n=38). Error bars indicate 95% confidence 1356 interval. Each row denotes a cell type specific dataset: excitatory neurons (EX), oligodendrocytes 1357 (OLI), inhibitory neurons (IN), astrocytes (AST), oligodendrocyte precursor cells (OPC), 1358 microglia (MIC), pericytes (PER) and endothelial cells (END). Cell types highlighted in bold

1359 reflect the equivalent to the cell type used in the interaction eQTL.

1360

1361 Figure 5. Mendelian randomization and colocalization of brain-related traits. (A) Number 1362 of significant Mendelian randomization (MR) signals (blue) and those with both MR and Coloc significant signals for 15 brain-related traits. (B) SNP and effect allele (EA), eQTL beta and 1363 1364 GWAS odds ratio for 20 multiple sclerosis (MS) genes that are both MR and Coloc significant, 1365 and their Wald ratio p-value. Cell type interaction eQTL for CYP24A1 (D) and CLECL1 (E), 1366 showing interactions with predicted neuron, and macrophage proportions respectively. The x-1367 axis shows the estimated cell type proportion, the y-axis shows the gene expression, each dot 1368 represents a sample. Colors indicate SNP genotype, with yellow being the MS risk allele. Values 1369 under the alleles are Spearman correlation coefficients.

1370

1371 Figure 6. Trans-eOTLs in brain. (A) Location of identified trans-eOTLs, with the SNP position (x-axis) and gene position (y-axis) in the genome. Size of the dots indicate the p-value 1372 1373 of the *trans*-eOTL (larger is more significant). 7p21.3, the locus with most (83%) of the *trans*-1374 eQTLs, is highlighted. (B) Three SNPs in the 7p21.3 locus and the number of datasets and 1375 number of up- and down-regulated trans-eQTL genes each SNP has. For rs1990622, a SNP 1376 associated with frontotemporal lobar degeneration, the 35 genes it affects in *trans* and the 1 gene 1377 it affects in *cis* are shown. (C) Two examples of convergent effects, where multiple independent 1378 SNPs affect the same genes in trans. Left: trans-eQTLs of rs1427407 and rs4895441 on HBG2 1379 and right trans-eQTL of rs930263, rs2604551, and rs10950398 on KCNA5.

1380

Figure 7. Gene co-regulation (A) Genes that are co-regulated with genes that are within
amyotrophic lateral sclerosis (ALS) loci. Co-regulation scores between genes are calculated
using all *MetaBrain* samples, *MetaBrain* cerebellum samples, or *MetaBrain* cortex samples.
Except for *URB4*, cortex and cerebellum networks find different co-regulated genes for ALS. (B)
Co-regulation network using all *MetaBrain samples* for all genes prioritized for ALS by *Downstreamer*. (C) Top 5 Human Phenotype Ontology (HPO) enrichments for the

- 1387 *Downstreamer* prioritized ALS genes. (D) Genes that are co-regulated with genes that are within
- 1388 multiple sclerosis loci. Co-regulation scores between genes are calculated using a heterogeneous
- 1389 multi-tissue network, *MetaBrain* cerebellum samples, or *MetaBrain* cortex samples. Most genes
- 1390 are found using a large heterogenous co-regulation network. (E) Co-regulation network of all
- 1391 *MetaBrain* samples for 33 genes prioritized by *Downstreamer* in cortex. Colors indicate the
- 1392 neutrophin signaling pathway enrichment Z-scores. (F) Top 5 KEGG enrichments for the
- 1393 *Downstreamer* prioritized multiple sclerosis genes in cortex.

# 1395 Supplementary Figure Legends

1396

## 1397 Supplementary Figure 1. European Nucleotide Archive brain sample selection. (A)

1398 Principal component (PC) analysis on the expression data of 74,052 samples included in the

1399 SkyMap database shows clustering on tissue type but also many outliers with high PC1 scores.

1400 (B) Coloring on single and paired-end sequencing shows no clear clustering. (C) Coloring single

1401 cell identifies the samples with high PC1 scores as single-cell samples. (**D**) Mean % reads

1402 mapped, number of reads, and max reads per bin of PC1. (E) Re-calculation of PCs on all

samples with PC score <0 in panel A-D, after covariate correction. (F) Brain and Tissue score calculated by correlating expression of known tissue and brain samples to each of the PCs. (G)

- 1405 As panel F, cancer score was calculated by correlating expression of known cancer genes to all
- 1406 PCs.

1407

Supplementary Figure 2. RNA-seq alignment QC. The two main RNA-seq QC metrics used
 for filtering samples. (A) Percentage coding bases colored by dataset and (B) percentage of reads

1410 aligned colored per dataset. Red dotted line is the threshold for filtering (10% for coding bases

1411 and 60% for percentage reads aligned respectively). Triangles are samples filtered out by any of

- 1412 the RNA-seq QC metrics.
- 1413

Supplementary Figure 3. Sample filtering by PCA. Principal component analysis (PCA) plot
 before normalization and covariate removal. For all plots the red line indicates 4 standard

1416 deviations from the mean and red dots are samples to be filtered out. (A) PCA on all samples

1417 after removing alignment QC outliers. (B) PCA on samples after removal of outlier samples

1418 from A. (C) PCA on samples after removal of outlier samples of A and B.

1419

1420 **Supplementary Figure 4. PCA before and after covariate correction.** (A) PC1 and PC2 on

normalized expression data before covariate correction, colored on dataset. (**B**) PC1 and PC2 on

- 1422 normalized expression data after covariate correction.
- 1423

Supplementary Figure 5. Assigning ethnicity through principal component analysis. For
each of the included datasets principal component (PC) scores are calculated on their genotypes.
Samples are clustered with the 1000 genome samples (left). Right panels show dataset genotype
samples without 1000g samples on the right projected on the same PCs. Using k-nearest
neighbors clustering, samples are assigned an ethnicity based on their closeness to the 1000g

1429 samples of a population.

### 1431 Supplementary Figure 6. eQTL Z-score comparison between datasets. The pairwise

- spearman correlation and concordance of direction of the eQTL Z-scores between all cohorts,
- and between each cohort and the meta-analysis Z-score. As two examples, (A) shows the Z-score
- 1434 comparison between Cortex-EUR eQTL datasets EUR-LIBD\_h650 and EUR-UCLA\_ASD, and
- 1435 (B) shows the Z-score comparison between the meta-analysis Z-score and the Cortex-EUR
- 1436 cohort EUR-AMPAD-ROSMAP-V2. (C) shows the correlation for each pairwise combination of
- 1437 cohorts between each other (small dots), and with the meta-analysis Z-scores (large dots). (**D**)
- shows the directional concordance for each pairwise combination of cohorts between each other
- 1439 (small dots), and with the meta-analysis Z-scores (large dots). The dots in (**C**) and (**D**) that
- 1440 correspond to the  $(\mathbf{A})$  and  $(\mathbf{B})$  plots are shown by the grey dottes lines.
- 1441

# 1442 Supplementary Figure 7. Reads mapping on patch chromosome version of MAPT. Number

- 1443 of reads mapped to the MAPT gene located on the primary assembly (ENSG00000186868) and
- the MAPT genes located on the patch chromosomes (ENSG00000276155 and
- 1445 ENSG00000277956). Each dot is an individual, and the color shows if they are homozygous
- 1446 reference (0/0), heterozygous (0/1), or homozygous alternative (1/1) for a SNP (rs34619181)
- 1447 located in the MAPT gene. Left plot compares counts mapped to ENSG00000186868 (ref) to
- those mapped to ENSG00000276155 (patch), middle plot compares ENSG00000186868 (ref)
- and ENSG00000277956 (patch), right plot compares ENSG00000276155 (patch) and
- 1450 ENSG00000277956 (patch).
- 1451
- Supplementary Figure 8. EQTL z-scores in the MAPT locus. Z-scores (y-axis) of the MAPT
   locus (x-axis) for all the datasets used in the Cortex-EUR meta-analysis. Left upper plot shows
- 1454 the meta-analysis Z-score. Blue dots are the SNPs that are in high LD with the top SNP.
- 1455
- 1456 Supplementary Figure 9. Colocalization locus plot for MAPT. Y-axis shows the
- colocalization log10(-p-value). X-axis shows the position of the SNPs (dots). Color is the LDwith rs56240678.
- 1459
- Supplementary Figure 10. (A) Mean of log<sub>2</sub> of the expression (x-axis) and standard deviation of the log<sub>2</sub> of expression for primary, secondary, tertiary, and quaternary eQTL genes. eQTLs that have only one independent SNP effect have higher mean expression but lower standard deviation than genes with multiple independent effects. (B) g:profiler enrichment for all genes with a single independent eQTL effect. (C) g:profiler enrichment for all genes with multiple
- 1465 independent eQTL effects.
- 1466
- Supplementary Figure 11. Properties of cerebellum specific eQTLs. (A) UpSet plot of the
  number of eQTL genes per brain region for European datasets. (B) The distribution of

1469 log2(TMM+1) expression in cortex (x-axis) and cerebellum (y-axis) of the 846 eQTL genes that

- 1470 were only significant in cerebellum. Blue line is the minima of the bimodal distribution and is
- 1471 used as cut-off point in panel C(C) The expression in cortex (x-axis) and cerebellum (y-axis) of

1472 the 846 eQTL genes that were only significant eQTLs in cerebellum. The blue line is the cut-off

- 1473 from panel **B**. (**D**) The expression (dots) and standard deviation (lines) of the transcription
- 1474 factors that are enriched for binding to transcription sites around the 662 genes for cortex (x-
- 1475 axis) and cerebellum (y-axis). The 5 transcription factors that are labelled are lower expressed in
- 1476 cortex and higher expressed in cerebellum.
- 1477

Supplementary Figure 12. Cortex primary eQTL replication in GTEx. The replication
between primary *cis*-eQTLs of Cortex-EUR (discovery) with all the GTEx tissues (replication).
The x-axis is the number of eQTLs that is significant in both discovery and replication, and the

- 1481 y-axis is the percentage that shows the same direction of effect.
- 1482
- 1483 **Supplementary Figure 13.** Comparison of meta-analysis Z-scores for eQTLs detected in the 1484 different *MetaBrain* datasets (x-axis), and eQTLgen (y-axis).
- 1485
- 1486 **Supplementary Figure 14. Distribution of predicted cell proportions.** The distribution of the 1487 predicted cell proportions (x-axis) for cortex and cerebellum samples (y-axis).
- 1488

1489 Supplementary Figure 15. Cell type proportions per brain region are comparable, with the

exception of the spinal cord. Visualization of the cell type proportions with one row per cell
 type and colors indicating brain region. (A) Density plot where the x-axis shows the predicted

1492 cell type proportion, and the y-axis shows the frequency. (**B**) Boxplot of the predicted cell type

- 1493 proportion. Boxes represent the 25<sup>th</sup> and 75<sup>th</sup> percentiles and internal line represents the median.
- 1494 The whiskers represent 1.5 multiplied by the inter-quartile range. Outliers are shown as 1495 individual points.
- 1496

## 1497 Supplementary Figure 16. Cell type fractions per brain tissue shows little differences with

1498 **the exception of the spinal cord.** Visualization of the cell type proportions with one row per

brain region and colors indicating cell types. (A) Density plot where the x-axis shows the

- 1500 predicted cell type proportion, and the y-axis shows the frequency. (**B**) Boxplot of the predicted 1501 cell type proportion. Boxes represent the  $25^{\text{th}}$  and  $75^{\text{th}}$  percentiles and internal line represents the
- median. The whiskers represent 1.5 multiplied by the inter-quartile range. Outliers are shown as
- 1503 individual points.
- 1504

### 1505 Supplementary Figure 17. Cell type mediated eQTLs in cerebellum are mostly mediated by

- astrocytes and macrophages. The number of cell type interacting eQTLs for cerebellum
- 1507 deconvoluted cell types. We did not identify eQTLs that were shared between cell types.
- 1508

1509 Supplementary Figure 18. Replication of cortex cis-eQTLs in snRNA-seq data. Each figure 1510 in this plot represents a comparison between bulk RNA-seq (y-axis) and single-nucleus RNA-seq 1511 (x-axis). Each dot represents one *cis*-eQTL, and the legend shows the Pearson correlation 1512 coefficient. Each column is a comparison between equivalent (and where not possible; similar) 1513 cell types in both datasets. Each row illustrates a different filtering on which eQTLs are shown 1514 and/or a different value on the y-axis. The x-axis always denotes the overall z-score of the eOTL 1515 effect in the single nucleus dataset of that respective column. (A) Meta-analysis eQTL z-score 1516 (y-axis) in Cortex-EUR bulk RNA-seq data, no filtering is applied. (B) Meta-analysis eQTL z-1517 score (y-axis) in Cortex-EUR bulk data, eQTLs are filtered based on the Decon-QTL Benjamini-Hochberg corrected p-value <0.05 in each respective column. (C) same as row **B** but now 1518 1519 showing the log betas of the interaction model on the y-axis. (**D**) Meta-analysis eOTL z-score (y-1520 axis) in bulk data for eQTLs that are significantly replicating in each respective dataset. Dots are 1521 colored if they are significantly cell type mediated (BH FDR<0.05) by the respective cell type in 1522 bulk data. (E) y-axis shows the log betas of the interaction model (y-axis) and filtering eQTLs on 1523 both significantly replicating in each respective dataset, as well as being significantly cell type 1524 mediated in bulk data.

1525

## 1526 Supplementary Figure 19. Bulk interacting eQTLs replicating in single-nucleus ROSMAP.

Replication of cell type interaction eQTLs for *STMN4* (A), *FAM221A* (B), *NKAIN1* (C) and *SCL25A27* (D). First column: Boxplots of the eQTL effect in Cortex-EUR bulk RNA-seq.

- 1529 Second column: Cell type interacting eQTL effect in Cortex-EUR bulk RNA-seq. The x-axis
- 1530 shows the estimated cell type proportion, the y-axis shows the gene expression, each dot 1531 represents a sample, and the colors indicate the SNP genotype, with yellow being the minor
- 1532 allele. Values under the alleles are Spearman correlation coefficients. Third column: Forest plot
- 1533 of the spearman coefficient with effect direction relative to the minor allele when replicating the
- eQTL effect in ROSMAP single nucleus data (n=38). Error bars indicate 95% confidence
- 1535 interval. Each row denotes a cell type specific dataset: excitatory neurons (EX), oligodendrocytes
- 1536 (OLI), inhibitory neurons (IN), astrocytes (AST), oligodendrocyte precursor cells (OPC),
- 1537 microglia (MIC), pericytes (PER) and endothelial cells (END). The bold cell type corresponds to
- the cell type that showed an interaction effect in bulk RNA-seq. Fourth column: Cell type
- interacting eQTL effect in ROSMAP single-nucleus RNA-seq (n=38) of the bold highlighted cell
  type in the third colum.
- 1541
- Supplementary Figure 20. Mendelian Randomization summary. Each plot is for a different
  trait (Intelligence, Intracranial volume, Putamen volume, Years of schooling, Alzheimer's
  disease, Amyotrophic Lateral Sclerosis, Depression (broad), Frontotemporal Dementia,

- 1545 Parkinson's disease, Bipolar disorder, Generalized epilepsy, juvenile myoclonic epilepsy,
- multiple sclerosis and schizophrenia). For each SNP the effect allele (EA) is given, the eQTL 1546
- 1547 beta of the EA on the given gene, the odds ratio (disease traits) or beta (quantitative traits) of the
- 1548 EA on the phenotype, and the Wald ratio p-value of the mendelian randomization analysis.
- 1549

Supplementary Figure 21. Colocalization regional plots for five suggestive MR findings in 1550

- 1551 Cortex-EUR that were replicated in eQTLGen with allelic discordance. Regional plots were
- 1552 made for five MR findings (CASS4 for Alzheimer's disease, TMEM170B for intelligence,
- 1553 GATAD2A for schizophrenia and years of schooling, and ZCWPW1 for years of schooling) in
- 1554 Cortex-EUR (top), eOTLGen (middle) and outcome GWAS (bottom) to show colocalization.
- These five findings all passed suggestive threshold ( $p < 5x10^{-5}$ ) in Cortex-EUR, with eQTL 1555
- effects replicated in eQTLGen (p<0.05), showed colocalization for both Cortex-EUR and 1556
- 1557 eOTLGen but opposite directions of effect.
- 1558

1559 Supplementary Figure 22. Colocalization regional plots for two suggestive MR findings for

1560 multiple sclerosis that showed opposite directions of effect between Cortex-EUR and 1561 eQTLGen. Regional plots were made for two suggestive MR findings for MS (KMT5A,

- 1562 RNF19B), both of which were suggestive signals in Cortex-EUR as well as eOTLGen ( $p < 5x10^{-1}$
- <sup>5</sup>). Opposite directions of effect were observed between Cortex-EUR and eQTLGen but
- 1563
- 1564 colocalization was only found in Cortex-EUR.
- 1565

1566 Supplementary Figure 23. Scatterplots comparing MR effects for multiple sclerosis derived using instruments from the metabrain versus eQTLGen studies. The top panel shows the 1567 1568 WR comparison on the same gene but with the different SNP instruments selected by each study 1569 (matching on the top WR finding if gene instrumented with multiple SNPs in the study) and the 1570 bottom panel the WR comparison between MetaBrain instruments and eQTLGen matching on 1571 both the same gene and SNP instrument. Genes which showed opposite direction of WR effect 1572 between MetaBrain and eQTLGen are colored in red and the genes with the same direction in

- 1573 blue.
- 1574

1575 **Supplementary Figure 24.** Log10 of median expression of brain and blood tissue samples in 1576 GTEx for 28 multiple sclerosis genes for which there are no significant eQTLgen instruments in 1577 brain and blood.

- 1578
- 1579 Supplementary Figure 25. Cell type proportions in Alzheimer's disease patients. Predicted 1580 cell count proportions for the AMP-AD samples that were used in the Cortex-EUR eQTL
- 1581 analysis for individuals with Alzheimer's disease and non-neurological controls. Each dot is the

- 1582 predicted cell proportion for one sample. Numbers under the boxplots indicate the number of
- 1583 samples plotted. Values above the line are p-values from a t-test between groups.
- 1584

Supplementary Figure 26. Forest plots for rs1990622 *trans*-eQTLs. Forest plots for each of the *trans*-eQTL genes associated with rs1990622. Each plot shows the *trans*-eQTL beta and 95% confidence interval for each of the included datasets and the meta-analysis. Effect directions are relative to the A allele of rs1990622. Sizes of dots are relative to sample size of each dataset.

- 1589 *Trans*-eQTL effects are most pronounced in AMP-AD datasets.
- 1590

1591 Supplementary Figure 27. Summary of 7p21.3 locus trans-eQTLs. (A) Forest plots showing

effect sizes for rs1990622 (yellow; beta and 95% confidence interval) for *cis*-eQTL gene

1593 THSD7A, trans-eQTL gene CALB2, and association of rs1990622 with estimated neuron

1594 proportion. Right panel shows average estimated neuron proportions per dataset (blue violin

plots). EQTL and neuron proportion associations are most pronounced in AMP-AD datasets,
while average neuron proportions are comparable. (B) *Trans*-eQTL meta-analysis Z-scores for

rs11974335, rs10950398 and rs1990622 (x-axis), and the correlation of those *trans*-eQTL genes

1598 with predicted neuron proportion (y-axis) are highly correlated. (C) Comparison of *trans*-eQTL

- 1599 Z-scores between Alzheimer's disease patients (x-axis) and neurotypical controls (y-axis) shows
- 1600 that eQTL Z-scores are higher in patients.
- 1601

1602 Supplementary Figure 28. Replication of cortex trans-eOTLs in single-nucleus data. Each 1603 figure in this plot represents a comparison between bulk RNA-seq (y-axis) and single-nucleus RNA-seq (x-axis). Each dot represents one trans-eQTL, and the legend shows the Pearson 1604 1605 correlation coefficient. Each column is a comparison between equivalent (and where not 1606 possible; similar) cell types in both datasets. Each row illustrates a different filtering on which 1607 eQTLs are shown and/or a different value on the y-axis. The x-axis always denotes the overall z-1608 score of the eQTL effect in the single nucleus dataset of that respective column. (A) Meta-1609 analysis eOTL z-score (y-axis) in Cortex-EUR bulk RNA-seq data, no filtering is applied. (B) 1610 Meta-analysis eQTL z-score (y-axis) in Cortex-EUR bulk data, eQTLs are filtered based on the 1611 Decon-QTL Benjamini-Hochberg corrected p-value <0.05 in each respective column. (C) same

1612 as row **B** but now showing the log betas of the interaction model on the y-axis. (**D**) Meta-analysis

- 1613 eQTL z-score (y-axis) in bulk data for eQTLs that are significantly replicating in each respective
- 1614 dataset. Dots are colored if they are significantly cell type mediated (BH FDR<0.05) by the
- respective cell type in bulk data. (E) y-axis shows the log betas of the interaction model (y-axis) and filtering eOTLs on both significantly replicating in each respective dataset, as well as being
- 1617 significantly cell type mediated in bulk data.
- 1618

# 1619 Supplementary figure 29. Comparison of AUC distribution for different eigenvector cut-

1620 offs. The quality of the gene network that we built for *MetaBrain* is measured by an AUC for

- 1621 each gene derived from a leave-one-out procedure. One of the parameters to build the network is
- 1622 the number of eigenvectors to use after PCA over the gene correlation matrix. Here we show for
- 1623 the 6 annotation categories (KEGG, REACTOME, GO Biological Process, GO Molecular
- 1624 Function, GO Cellular Component, and HPO) the AUC mean (dot) and standard deviation (lines)
- 1625 at different eigenvector cut-offs. The red dot and line indicate the eigenvector cut-off that was
- 1626 used for that annotation category.
- 1627
- Supplementary Figure 30. Heatmaps of the Pearson correlation of the AUC values between
   different eigenvector cut-offs. Correlation was calculated between the different eigenvector
- 1630 cutoffs for the 6 annotation categories.
- 1631
- 1632 **Supplementary Figure 31. (A)** UMAP representation of heterogeneous gene network. Immune
- and blood cell types show increased gene expression levels for genes prioritized using
- 1634 *Downstreamer* for multiple sclerosis, while decreased expression is observed in brain related
- 1635 tissues. (B) Within MetaBrain, those same genes show lower expression in cortex, but higher
- 1636 expression in spinal cord and cerebellum.
- 1637

### 1638 Supplementary Figure 32. Spearman correlation heatmap of predicted cell fractions versus

- 1639 principal components calculated using all *MetaBrain* samples. A heatmap showing the first
- 1640 fifty principal components as the columns and the five cell types for which we predicted
- 1641 proportions as rows. Each cell is colored based on the spearman correlation coefficients. Blue
- 1642 denotes a negative correlation, red a positive correlation and white denotes no correlation.
- 1643
- Supplementary Figure 33. SnRNA-seq visualization by cell type. UMAP dimensionality
   reduction plot of 39 snRNA-seq samples from ROSMAP. Each dot represents a single cell
   (n=70,634). The dots are colored by their corresponding cell type: excitatory neurons (EX),
   oligodendrocytes (OLI), inhibitory neurons (IN), astrocytes (AST), oligodendrocyte precursor
- 1648 cells (OPC), microglia (MIC), pericytes (PER) and endothelial cells (END).
- 1649
- Supplementary Figure 34. SnRNA-seq visualization by cell type. UMAP dimensionality
  reduction plot of 39 snRNA-seq samples from ROSMAP. Each dot represents a single cell
  (n=70,634). The dots are colored by their corresponding cell type subcluster: excitatory neurons
  (EX), oligodendrocytes (OLI), inhibitory neurons (IN), astrocytes (AST), oligodendrocyte
  precursor cells (OPC), microglia (MIC), pericytes (PER) and endothelial cells (END).
- 1655
- 1656

# 1657 **Table descriptions**

### 1658 **Table 1. Prioritized genes from the Mendelian Randomization analysis on MetaBrain**

1659 eQTLs versus brain related outcomes. Harmonized eQTL and GWAS SNP effects and single

- 1660 SNP Wald Ratio estimates are reported in the table for all genes with Wald Ratio effects at
- 1661 P<1.865x10<sup>-7</sup>. Columns are **genomic position**, **rsid** and **alleles** for SNP instrument (**EA**: Effect
- allele. NONEA: non-effect allele. proxy SNP: rsid of proxy SNP replacement used for outcome
- 1663 if instrument not present in GWAS), the SNP effects (**beta**, **SE**, **p**) for the MetaBrain eQTLs
- 1664 followed by the SNP effects for the brain related outcomes and then the Wald Ratio effects.

# 1666 Supplementary Table descriptions

### 1667 Supplementary table 1. Number of samples and individuals.

- 1668 Sheet Genotype QC: The number of genotype individuals and samples pre-QC (column C-H)
- and post-QC (column I-N) for the different RNA-seq (column A) and genotype (column B)
- 1670 datasets. Columns are: **PreQC**: Number of initial genotype samples processed for QC. **PostQC**:
- 1671 Number of genotype samples left after QC filtering. **RNA-seq dataset**: Name of the complete
- 1672 dataset. Genotype dataset: Name of the genotype dataset. Some datasets have multiple genotype
- 1673 platforms, or multiple smaller datasets that are part of the larger RNA-seq dataset. **Individuals:**
- 1674 The number of individuals per dataset. **EUR**: Number of genotype samples per dataset of
- 1675 individuals of European population. **AFR**: Number of genotype samples per dataset of
- 1676 individuals of African population. **EAS**: Number of genotype samples per dataset of individuals
- 1677 of East-Asian population. SAS: Number of genotype samples per dataset of individuals of South-
- 1678 Asian population. **AMR:** Number of genotype samples per dataset of individuals of Ad Mixed
- 1679 American population.
- 1680 Sheet **RNA-QC**: The number of RNA-seq samples at different steps of QC and for different
- 1681 brain regions. Cells A2-F18 have the number of samples at different QC steps. Columns are:
- 1682 Dataset: dataset name. Number of RNA-seq samples: Number of RNA-seq samples processed
- 1683 to go through QC. Alignment QC: Number of RNA-seq samples left after filtering on alignment
- 1684 QC (e.g. percent reads aligned). **RNA-seq PCA outliers step 1**: Number of RNA-seq samples
- 1685 left after filtering samples >4SD from mean of PC1. **RNA-seq PCA outliers step 2**: Number of
- samples left after recalculating PCA and again removing samples >4SD fom mean of PC1.
- 1687 Covariate removal: Number of samples left after covariate removal. RNA Tissue grouping:
- 1688 the meta-data across different datasets uses different granularity of tissue annotation. Tissues
- 1689 were grouped accordingly.
- 1690 Sheet Sample Links: RNA-seq samples linked to genotype samples. Left top: numbers of RNA-
- 1691 seq sample linked to a genotype sample per dataset, per population. Top right: number of unique
- 1692 individuals per dataset per population. Middle: number of uniquely linked individuals per
- 1693 dataset, per population and per tissue group. Bottom: numbers of individuals used from each
- 1694 dataset and population for *cis* and *trans*-eQTL analysis.
- 1695

# 1696 Supplementary table 2. Cis-eQTL summary statistics.

- 1697 *Cis*-eQTL summary statistics listing index variant per gene (FDR<0.05). One sheet per eQTL
- 1698 discovery dataset. Genomic positions are GRCh38. eQTL Rank: whether the eQTL is a primary,
- 1699 secondary, tertiary, quaternary, or higher eQTL.
- 1700

# 1701 Supplementary table 3. Number of *cis-* and *trans-eQTLs.* For each dataset the number of *cis-*

- and *trans*-eQTL SNPs, genes, and SNP-gene combinations found at FDR<0.05. Columns are:
- 1703 Basalganglia, Cerebellum, Cortex, Hippocampus, Spinalcord: the five different brain regions
- 1704 for which eQTL calling was done. **EUR:** Number of eQTLs with samples from European
- 1705 population. **AFR**: Number of eQTLs with samples from African population. **EAS**: Number of

- 1706 eQTLs with samples from East-Asian population. EUR+AFR, wo ENA, no PCA: Number of
- 1707 eQTLs with samples from EUR and AFR populations, excluding samples from the ENA cohorts,
- and using gene expression levels that were not corrected for principal components.
- 1709
- 1710 Supplementary table 4. Gene set enrichment summary statistics for primary and higher
- 1711 rank eQTLs. Gene set enrichment summary statistics generated using g:Profiler for genes
- 1712 having a primary eQTL effect (sheet Primary eQTL), and those also having a secondary eQTL
- 1713 (sheet Non-primary eQTL).
- 1714
- 1715 Supplementary table 5. Gene set enrichment summary statistics generated using g:Profiler for1716 genes having an eQTL effect in cerebellum.
- 1717
- 1718 Supplementary table 6. GTEx *cis*-eQTL replication. Replication between *cis*-eQTLs of
- 1719 different MetaBrain regions and all GTEx tissues. Discovery was performed in each MetaBrain
- 1720 dataset while excluding GTEx, and then replicated in each GTEx tissue. **Tested eQTLs:** those
- eQTLs that were also present in the GTEx dataset. **Proportion shared and FDR<0.05**:
- 1722 proportion of tested eQTLs that was also significant in GTEx. Concordant and FDR<0.05:
- 1723 number of tested eQTLs that was also significant and for which the allelic direction was
- 1724 concordant. **Concordance**: proportion of concordant tested and significant eQTLs.
- 1725
- Supplementary table 7. eQTLgen *cis*-eQTL replication. *MetaBrain cis*-eQTLs (FDR<0.05) as</li>
  discovery cohort and eQTLgen eQTLs as replication cohort. Top table: FDR<0.05 in MetaBrain</li>
  discovery only (FDR<1 in eQTLGen). Bottom table: FDR<0.05 in both MetaBrain and</li>
  eQTLgen datasets. Shared: number of shared eQTLs. Concordant: number of shared eQTLs
  that has the same allelic direction of effect. Concordant over total: proportion of concordant
- 1731 eQTLs over the total number of eQTLs discovered. Concordant over shared: proportion of
- 1732 concordant eQTLs over number of shared eQTLs.
- 1733
- 1734 Supplementary table 8. Cell type deconvolution summary statistics. Sheet cortex: All
- 1735 Decon-eQTL results for cortex. Sheet cerebellum: All Decon-eQTL results for cerebellum.
- 1736 Columns for both sheets are: **Gene**: deconvoluted eQTL gene ensebl ID. **Gene symbol**:
- 1737 deconvolution eQTL gene symbol. **SNP**: deconvoluted eQTL SNP. **Alleles**: SNP alleles. **Effect**
- 1738 Allele: the allele to which the betas are directed. Columns ending with p-value: p-value for the
- 1739 cell-type interaction. **Columns ending with beta**: beta for the cell-type proportion term.
- 1740 **Columns ending with beta:GT**: beta for the genotype x cell-type interaction term.
- 1741

### 1742 Supplementary table 9 Replication of the MetaBrain cortex primary *cis*-ieQTLs in

- 1743 **ROSMAP single-nucleus data.** For each of the deconvoluted cell-types, the FDR and betas are
- 1744 listed. For each of the cell types in the single nucleus data, the FDR and eQTL Z-scores are
- 1745 listed. All betas andZ-scores are relative to the Effect Allele.
- 1746

#### 1747 Supplementary table 10. eQTL SNPs in linkage disequilibrium with GWAS SNPs. The 1748 GWAS SNPs that are in high linkage disequilibrium (LD) with the *cis*-eQTL SNPs. Each sheet is 1749 a different *metabrain* eOTL datasets from EUR populations. The sheet Included Traits lists 1750 GWAS traits that were tested. Columns are: eQTL rank: the rank of conditional eQTLs 1751 (1=primary, 2=secondary, etc). GWASID: GWAS ID of the GWAS SNP. Trait: Name of the GWAS trait. Index variant: the GWAS variant. Index Variant P: GWAS p-value. Index 1752 1753 Variant Alleles: Alleles of the GWAS variant. Index Variant Effect: GWAS effect. Linked **EQTL SNP**: the eQTL SNP. LD(rsq): the LD $r^2$ . LinkedEQTLGenes: the eQTL genes that the 1754

- 1755 linked SNP affects. Linked EQTL Gene Symbols: HGNC name of the linked genes. Linked
- 1756 EQTL Alleles: Alleles of the eQTL SNP. Linked EQTL Effect Allele: The allele that is related
- 1757 to the effect direction. Linked EQTL Zscores: Z-scores of the eQTL effect. Linked EQTL P:
- 1758 p-value of the eQTL effect. **GWAS Cluster Size**: Number of GWAS SNPs in LD with Index
- 1759 Variant. SNPs In **GWAS Cluster**: SNPs that are in LD with the Index Variant.
- 1760
- 1761 Supplementary table 11. List of traits used in Mendelian randomization and colocalization1762 analysis.
- 1763

Supplementary table 12 eOTL SNPs which showed evidence of genetic colocalization with 1764 1765 tested brain-related traits. ID, Chromosome, Position, SNP, Effect Allele, Non Effect Allele: 1766 Position of instrumenting SNP with effect allele used during the harmonization procedure. Proxy 1767 used, Proxy SNP: whether proxy lookup had to be performed to find SNP in outcome GWAS 1768 and the rsid of the proxy used. MetaBrain SNP effects: gene name and summary statistics for the instrument-exposure SNP association (MetaBrain eOTL). Outcome SNP effects: outcome 1769 1770 name (neurological trait) and summary statistics for the harmonized instrument-outcome SNP 1771 association. MR effects: single SNP Wald ratio effect between the instrumented eQTL and neurological outcome. Coloc results: colocalization probability of both traits sharing the same 1772 1773 causal variant in the region. Decon-OTL results: eOTL SNP: the SNP that was tested for cell 1774 type mediated effects. In some cases a SNP which is in high LD with the instrument SNP is used 1775 for Decon-QTL. LD R-squared: the LD between SNP and eQTL SNP. Columns listing Decon-1776 OTL results: **beta**: the beta of the interaction term in the Decon-OTL model with respect to the Effect Allele column. FDR: the Benjamini-Hochberg corrected interaction p-value. Mendelian 1777 **Disorders**: overlap of genes with Development Disorder Genotype - Phenotype Database 1778 1779 (DDG2P) and OrphaNet.

### 1781 Supplementary Table 13. Colocalization results for latest AD GWAS loci with *MetaBrain*

1782 **Cortex-EUR primary eQTLs (columns A to P** were adapted from Schwartzentruber *et al.* for

- 1783 comparisons and **columns Q to Y** are *MetaBrain* findings. **Category 1**: previously identified
- and replicated in *MetaBrain* Cortex-EUR, **2**: novel results found by *MetaBrain* Cortex-EUR, **3**:
- 1785 previously identified but not replicated in *MetaBrain* Cortex-EUR.
- 1786

Supplementary table 14. Mendelian Randomization comparison between MetaBrain and eQTLGen on multiple sclerosis outcome. (a) Wald Ratio comparison on the same gene using different SNP instruments. For this analysis, the Wald Ratio effects for the top hit eQTL for each gene within each study were compared. (b) Wald Ratio comparison on the same gene fixing on the same eQTL instrument between studies. For this analysis, the eQTLGen Wald Ratios were re-derived using the second Taylor expansion error term on the same SNP

- 1793 instruments as MetaBrain.
- 1794

Supplementary table 15. Colocalization of MR suggestive hits with high LD but allelic
discordance. This table displays the colocalization results for 31 suggestive MR findings from
Cortex-EUR with eQTL instruments replicated in eQTLGen (p<0.05) but allelic discordance</li>
(opposite directionalities of alleles). Highlighted rows are findings with colocalization in both
Cortex-EUR and eQTLGen.

1800

# 1801 Supplementary table 16. Comparison of MR suggestive hits for MS between metaBrain

1802 and eQTLGen. This table displays 157 suggestive MR signals for multiple sclerosis in Cortex-

1803 EUR and the replication MR and colocalization results of corresponding genes in eQTLGen.

1804

1805 Supplementary table 17. Trans-eQTL summary statistics. Sheet Trans-eQTLs: all trans-1806 eQTLs detected in this study (FDR<0.05). Percentage cross-mapping: percentage of the gene 1807 that can be mapped within 5Mb of the trans-eQTL SNP. Sheet Trans-eQTLs no crossmap: trans-1808 eQTLs that remain significant after cross-mapping eQTLs have been removed. Sheet Trans-1809 eQTLs with cis per trait: in this sheet, trans-eQTLs are annotated with cis-eQTLs for the same 1810 SNP, and subsequently split per trait annotation for the SNP. Consequently, a single trans-eQTL 1811 may be represented by multiple rows. Sheet Convergent *trans*-eOTLs: genes on which multiple 1812 independent loci have a trans-eQTL, split per annotated trait. Sheet TraitsAndNrOfSNPs: list of 1813 traits included in the analysis, and the number of included SNPs per trait.

1814

# 1815 Supplementary table 18. Summary statistics for associations between SNPs and predicted

1816 **cell-type proportions.** Sheet Cortex-EUR: associations (FDR<0.05) while limiting to Cortex-

1817 EUR samples. Sheet Cortex-EUR+AFR-woENA: associations (FDR<0.05) for the analysis

1818 including AFR samples, but excluding ENA samples.

### 1819

### 1820 Supplementary table 19. Differences in predicted neuron proportions between included

datasets. T-test p-values comparing neuron proportions for pairwise comparisons between the
 datasets included in the *trans*-eQTL analysis.

1823

### 1824 Supplementary table 20. Gene-cell count correlations and 7p21.3 *trans*-eQTL Z-scores.

- 1825 Trans-eQTL Z-scores for three SNPs (rs11974335, rs10950398, and rs1990622), and
- 1826 correlations of the *trans*-eQTL genes with predicted neuron proportions.
- 1827

# 1828 Supplementary table 21. Gene set enrichments for 7p21.3 *trans*-eQTL genes. Gene set

1829 enrichments calculated using g:Profiler. Sheet downregulated genes: gene set enrichments for

- 1830 genes that show downregulation due to the 7p21.3 trans-eQTL effect alleles. Sheet upregulated
- 1831 genes: gene set enrichments for genes that show upregulation due to the 7p21.3 trans-eQTL
- 1832 effect alleles.
- 1833

### 1834 Supplementary table 22. Replication of the MetaBrain cortex primary *trans*-ieQTLs in

1835 **ROSMAP single-nucleus data.** For each of the deconvoluted cell-types, the FDR and betas are

- 1836 listed. For each of the cell types in the single nucleus data, the FDR and eQTL Z-scores are
- 1837 listed. All betas and Z-scores are relative to the Effect Allele.
- 1838

## 1839 Supplementary table 23. Downstreamer results for amyotrophic lateral sclerosis in EUR

- 1840 and Asian populations. Sheet overview: lists set of ontologies tested for this phenotype. Sheet
- 1841 GenePrioritization\_MetaBrain: gene prioritization performed in all *MetaBrain* samples. Sheet
- 1842 GenePrioritization\_MetaBrainCortexOnly: gene prioritization performed in *MetaBrain* cortex
- 1843 samples. GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in
- 1844 MetaBrain cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain,
- 1845 GO\_CC\_MetaBrain, GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set
- 1846 enrichments for coregulated genes identified using Downstreamer. Sheets
- 1847 Expression\_MetaBrain, Expression\_HCA, and GtexV8\_relative: expression enrichment using all
- 1848 MetaBrain samples, Human Cell Atlas, and GTEx v8.
- 1849

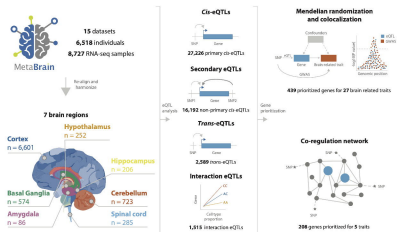
# 1850 Supplementary table 24. Downstreamer results for Parkinson's disease. Sheet overview:

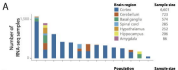
- 1851 lists set of ontologies tested for this phenotype. Sheet GenePrioritization\_MetaBrain: gene
- 1852 prioritization performed in all *MetaBrain* samples. Sheet
- 1853 GenePrioritization\_MetaBrainCortexOnly: gene prioritization performed in *MetaBrain* cortex
- 1854 samples. GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in
- 1855 MetaBrain cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain,

- 1856 GO\_CC\_MetaBrain, GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set
- 1857 enrichments for coregulated genes identified using Downstreamer. Sheets
- 1858 Expression\_MetaBrain, Expression\_HCA, and GtexV8\_relative: expression enrichment using all
- 1859 MetaBrain samples, Human Cell Atlas, and GTEx v8.
- 1860
- 1861 Supplementary table 25. Downstreamer results for schizophrenia. Sheet overview: lists set
- 1862 of ontologies tested for this phenotype. Sheet GenePrioritization\_MetaBrain: gene prioritization
- 1863 performed in all *MetaBrain* samples. Sheet GenePrioritization\_MetaBrainCortexOnly: gene
- 1864 prioritization performed in *MetaBrain* cortex samples.
- 1865 GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in *MetaBrain*
- 1866 cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain, GO\_CC\_MetaBrain,
- 1867 GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set enrichments for
- 1868 coregulated genes identified using Downstreamer. Sheets Expression\_MetaBrain,
- 1869 Expression\_HCA, and GtexV8\_relative: expression enrichment using all MetaBrain samples,
- 1870 Human Cell Atlas, and GTEx v8.
- 1871
- 1872 Supplementary table 26. Downstreamer results for Alzheimer's disease. Sheet overview:
- 1873 lists set of ontologies tested for this phenotype. Sheet GenePrioritization\_MetaBrain: gene
- 1874 prioritization performed in all MetaBrain samples. Sheet
- 1875 GenePrioritization\_MetaBrainCortexOnly: gene prioritization performed in *MetaBrain* cortex
- 1876 samples. GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in
- 1877 *MetaBrain* cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain,
- 1878 GO\_CC\_MetaBrain, GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set
- 1879 enrichments for coregulated genes identified using Downstreamer. Sheets
- 1880 Expression\_MetaBrain, Expression\_HCA, and GtexV8\_relative: expression enrichment using all
- 1881 MetaBrain samples, Human Cell Atlas, and GTEx v8.
- 1882
- 1883 Supplementary table 27. Downstreamer results for multiple sclerosis. Sheet overview: lists
- 1884 set of ontologies tested for this phenotype. Sheet GenePrioritization\_MetaBrain: gene
- 1885 prioritization performed in all MetaBrain samples. Sheet
- 1886 GenePrioritization\_MetaBrainCortexOnly: gene prioritization performed in *MetaBrain* cortex
- 1887 samples. GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in
- 1888 MetaBrain cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain,
- 1889 GO\_CC\_MetaBrain, GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set
- 1890 enrichments for coregulated genes identified using Downstreamer. Sheets
- 1891 Expression\_MetaBrain, Expression\_HCA, and GtexV8\_relative: expression enrichment using all
- 1892 MetaBrain samples, Human Cell Atlas, and GTEx v8.
- 1893

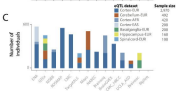
### 1894 Supplementary table 28. Downstreamer results for amyotrophic lateral sclerosis in EUR

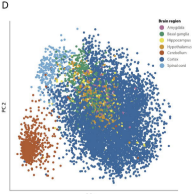
- 1895 **population.** Sheet overview: lists set of ontologies tested for this phenotype. Sheet
- 1896 GenePrioritization\_MetaBrain: gene prioritization performed in all MetaBrain samples. Sheet
- 1897 GenePrioritization\_MetaBrainCortexOnly: gene prioritization performed in *MetaBrain* cortex
- 1898 samples. GenePrioritization\_MetaBrainCerebellumOnly: gene prioritization performed in
- 1899 MetaBrain cerebellum samples. Sheets Reactome\_MetaBrain, GO\_BP\_MetaBrain,
- 1900 GO\_CC\_MetaBrain, GO\_MF\_MetaBrain, KEGG\_MetaBrain, and HPO\_MetaBrain: gene set
- 1901 enrichments for coregulated genes identified using Downstreamer. Sheets
- 1902 Expression\_MetaBrain, Expression\_HCA, and GtexV8\_relative: expression enrichment using all
- 1903 MetaBrain samples, Human Cell Atlas, and GTEx v8.
- 1904
- 1905 Supplementary table 29. ENA accession IDs. List of study accession IDs collected from
- 1906 European Nucleotide Archive. Columns are: **study\_accession:** ID of the study in ENA.
- 1907 **run\_accession:** ID of all the ENA runs included in this study (before quality control)
- 1908

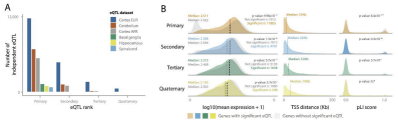




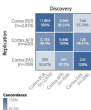




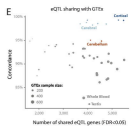


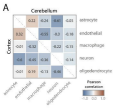


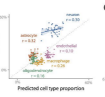


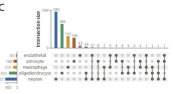


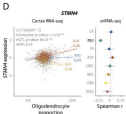
D		Discovery							
	Cortex EUR (n=2,970)	11,803 100%	5,355 94.58%	1,241 99.11%	941 99.79%	747 98.93%			
	Cerebellum EUR (n=492)	6,617 95.75%	6,865 100%	1,119 98.39%	865 98.27%	670 97.01%			
Replication	Basal ganglia EUR (n=208)	3,275 98.35%	2.037 96.96%	1,317 100%	852 99.88%	581 99.31%			
Re	Hippocampus EUR (n=168)	2,920 98.73%		1,106 99.64%	990 100%	568 99.82%			
	Spinal cord EUR (n=108)			863 99.3%	689 99.13%	811 100%			
	Concordance 100% 97.5%	97.12%	n ton son	olifer tost	so con co	10 50 100 M			









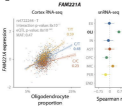


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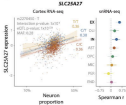
В

cell type proportion

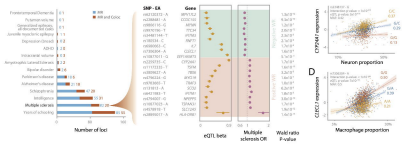
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