

INFERENCE OF CELL TYPE COMPOSITION FROM HUMAN BRAIN TRANSCRIPTOMIC DATASETS ILLUMINATES THE EFFECTS OF AGE, MANNER OF DEATH, DISSECTION, AND PSYCHIATRIC DIAGNOSIS

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Abstract

Psychiatric illness is unlikely to arise from pathology occurring uniformly across all cell types in affected brain regions. Despite this, transcriptomic analyses of the human brain have typically been conducted using macro-dissected tissue due to the difficulty of performing single-cell type analyses with donated post-mortem brains. To address this issue statistically, we compiled a database of several thousand transcripts that were specifically-enriched in one of 10 primary cortical cell types, as identified in previous publications. Using this database, we predicted the relative cell type composition for 833 human cortical samples using microarray or RNA-Seq data from the Pritzker Consortium (GSE92538) or publicly-available databases (GSE53987, GSE21935, GSE21138, CommonMind Consortium). These predictions were generated by averaging normalized expression levels across transcripts specific to each cell type using our R-package *BrainInABlender* (validated and publicly-released: <https://github.com/hagenaue/BrainInABlender>). Using this method, we found that the principal components of variation in the datasets were largely explained by the neuron to glia ratio of the samples. This variability was not simply due to dissection – the relative balance of brain cell types was influenced by a variety of demographic, pre- and post-mortem variables. Prolonged hypoxia around the time of death predicted increased astrocytic and endothelial content in the tissue, illustrating vascular upregulation. Aging was associated with decreased neuronal content. Red blood cell content was reduced in individuals who died following systemic blood loss. Subjects with Major Depressive Disorder had decreased astrocytic content, mirroring previous morphometric observations. Subjects with Schizophrenia had reduced red blood cell content, resembling the hypofrontality detected in fMRI experiments. Finally, in datasets containing samples with especially variable cell content, we found that controlling for predicted sample cell content while evaluating differential expression improved the detection of previously-identified psychiatric effects. We conclude that accounting for cell type can greatly improve the interpretability of microarray data.

1 **1. Introduction**

2 The human brain is a remarkable mosaic of diverse cell types stratified into rolling cortical layers,
3 arching white matter highways, and interlocking deep nuclei. In the past decade, we have come to
4 recognize the importance of this cellular diversity in even the most basic neural circuits. At the same time,
5 we have developed the capability to comprehensively measure the thousands of molecules essential for
6 cell function. These insights have provided conflicting priorities within the study of psychiatric illness: do
7 we carefully examine individual molecules within their cellular and anatomical context or do we extract
8 transcript or protein en masse to perform large-scale unbiased transcriptomic or proteomic analyses? In
9 rodent models, researchers have escaped this dilemma by a boon of new technology: single cell laser
10 capture, cell culture, and cell-sorting techniques that can provide sufficient extract for transcriptomic and
11 proteomic analyses. However, single cell analyses of the human brain are far more challenging (1–3) –
12 live tissue is only available in the rarest of circumstances (such as temporal lobe resection) and intact
13 single cells are difficult to dissociate from post-mortem tissue without intensive procedures like laser
14 capture microscopy.

15 Therefore, to date, the vast majority of unbiased transcriptomic analyses of the human brain have
16 been conducted using macro-dissected, cell-type heterogeneous tissue. On Gene Expression Omnibus
17 alone, there are at least 63* publicly-available macro-dissected post-mortem human brain tissue datasets,
18 and many other macro-dissected human brain datasets are available to researchers via privately-funded
19 portals (Stanley Medical Research Institute, Allen Brain Atlas, CommonMind Consortium). These
20 datasets have provided us with novel hypotheses (e.g., (4,5)), but researchers who work with the data
21 often report frustration with the relatively small number of candidate molecules that survive analyses
22 using their painstakingly-collected samples, as well as the overwhelming challenge of interpreting
23 molecular results in isolation from their respective cellular context. At the core of this issue is the inability
24 to differentiate between (1) alterations in gene expression that reflect an overall disturbance in the relative

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25 ratio of the different cell types comprising the tissue sample, and (2) intrinsic dysregulation of one or
26 more cell types, indicating perturbed biological function.

27 In this manuscript, we present results from an easily accessible solution to this problem that
28 allows researchers to statistically estimate the relative number or transcriptional activity of particular cell
29 types in macro-dissected human brain microarray data by tracking the collective rise and fall of
30 previously identified cell type specific transcripts. Similar techniques have been used to successfully
31 predict cell type content in human blood samples (6–9), as well as diseased and aged brain samples (10–
32 12). Our method was specifically designed for application to large, highly-normalized human brain
33 transcriptional profiling datasets, such as those commonly used by neuroscientific research bodies such as
34 the Pritzker Neuropsychiatric Research Consortium and the Allen Brain Institute.

35 We took advantage of a series of newly available data sources depicting the transcriptome of
36 known cell types, and applied them to infer the relative balance of cell types in our tissue samples in a
37 semi-supervised fashion. We draw from seven large studies detailing cell-type specific gene expression
38 in a wide variety of cells in the forebrain and cortex (2,13–18). Our analyses include all major categories
39 of cortical cell types (17), including two overarching categories of neurons that have been implicated in
40 psychiatric illness (19): projection neurons, which are large, pyramidal, and predominantly excitatory, and
41 interneurons, which are small and predominantly inhibitory (20). These are accompanied by the three
42 prevalent forms of glia that make up the majority of cells in the brain: oligodendrocytes, which provide
43 the insulating myelin sheath that enhances electrical transmission in axons (21), astrocytes, which help
44 create the blood-brain barrier and provide structural and metabolic support for neurons, including
45 extracellular chemical and electrical homeostasis, signal propagation, and response to injury (21), and
46 microglia, which serve as the brain's resident macrophages and provide an active immune response (21).
47 We also incorporate structural and vascular cell types: endothelial cells, which line the interior surface of
48 blood vessels, and mural cells (smooth muscle cells and pericytes), which regulate blood flow (22).
49 Progenitor cells were also included in our analysis because they are widely regarded as important for the
50 pathogenesis of mood disorders (23). Within the cortex, these cells mostly take the form of immature

51 oligodendrocytes (17). Finally, the primary cells found in blood, erythrocytes or red blood cells (RBCs),
52 carry essential oxygen throughout the brain. These cells do not contain a cell nucleus and do not generate
53 new RNA, but still contain an existing, highly-specialized transcriptome (24). The relative presence of
54 these cells could arguably represent overall blood flow, the functional marker of regional neural activity
55 traditionally used in human imaging studies.

56 To characterize the balance of these cell types in psychiatric samples, we first compared the
57 predictive value of cell type specific transcripts identified by diverse data sources and then summarized
58 their collective predictions of relative cell type balance into covariates that could be used in larger linear
59 regression models. We find that these “cell type indices” can successfully predict relative cell content in
60 validation datasets, including *in vitro* and post-mortem datasets. We discover that the variability in the
61 relative cell type balance of samples can explain a large percentage of the variation in macro-dissected
62 human brain microarray and RNA-Seq datasets. This variability is driven by pre- and post-mortem
63 subject variables, such as age, aerobic environment, and large scale blood loss, in addition to dissection.
64 Finally, we demonstrate that this method enhances our ability to discover and interpret psychiatric effects
65 in human brain microarray datasets, uncovering known changes in cell type balance in relationship to
66 Major Depressive Disorder and Schizophrenia and potentially increasing our sensitivity to detect genes
67 with previously-identified relationships to Bipolar Disorder and Schizophrenia in datasets that contain
68 samples with highly-variable cell content.

69

70 **2. Methods & Validation**

71

72 **2.1 Compiling a Database of Cell Type Specific Transcripts**

73 To perform this analysis, we compiled a database of several thousand transcripts that were
74 specifically-enriched in one of nine primary brain cell types within seven published single-cell or purified
75 cell type transcriptomic experiments using mammalian brain tissues (2,13–18) (**Suppl. Table 1**). These

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76 primary brain cell types included six types of support cells: astrocytes, endothelial cells, mural cells,
77 microglia, immature and mature oligodendrocytes, as well as two broad categories of neurons
78 (interneurons and projection neurons). We also included a category for neurons that were generically
79 extracted (“neuron_all”). The experimental and statistical methods for determining whether a transcript
80 was enriched in a particular cell type varied by publication (**Figure 1**), and included both RNA-Seq and
81 microarray datasets. We focused on cell-type specific transcripts identified using cortical or forebrain
82 samples because the data available for these brain regions was more plentiful than for the deep nuclei or
83 the cerebellum. In addition, we artificially generated a list of 17 transcripts specific to erythrocytes (red
84 blood cells or RBC) by searching Gene Card for erythrocyte and hemoglobin-related genes
85 (<http://www.genecards.org/>).

86 In all, we curated gene expression signatures for 10 cell types expected to account for most of the
87 cells in the cortex. Our final database included 2499 unique human-derived or orthologous (as predicted
88 by HCOP using 11 available databases: <http://www.genenames.org/cgi-bin/hcop>) transcripts, with a focus
89 on coding varieties. We have made this database publicly accessible within our R package
90 (<https://github.com/hagenaue/BrainInABlender>) and as a downloadable spreadsheet
91 (<https://sites.google.com/a/umich.edu/megan-hastings-hagenauer/home/cell-type-analysis>).

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Citation	Cell Origin	Method	Stringency	Derived Cortical Cell Type Indices	Transcripts/Orthologs
Cahoy et al., <i>J Neuro</i> , 2008.	Forebrain of young transgenic mice	Fluorescent cell sorting using antibodies to deplete non-specific cell types followed by Affymetrix microarray	>20 Fold Enrichment	Astrocyte_All	73
				Neuron_All	80
				Oligodendrocyte_All	50
Zhang et al., <i>J Neuro</i> , 2014	Cortex of young transgenic mice	Fluorescent cell sorting using antibodies to deplete non-specific cell types followed by RNAseq	Top 40 transcripts with >20 Fold Enrichment	Astrocyte_All	40
				Endothelial_All	40
				Microglia_All	40
				Mural_Pericyte	40
				Neuron_All	40
				Oligodendrocyte_Myelinating	40
				Oligodendrocyte_Newly-Formed	39
				Oligodendrocyte_Progenitor Cell	40
Zeisel et al., <i>Science</i> , 2015	Somatosensory cortex and CA1 hippocampus of juvenile mice	Unbiased capture of single cells from whole tissue cell suspension followed by RNAseq	Enriched with 99.9% posterior probability	Astrocyte_All	240
				Endothelial_All	353
				Microglia_All	436
				Mural_All	155
				Neuron_Interneuron	365
				Neuron_Pyramidal_Cortical	294
				Oligodendrocyte_All	453
Darmanis et al., <i>PNAS</i> , 2015	Anterior temporal lobe resected from adult human epileptic patients and cortex from fetuses 16-18 wks postgestation.	Unbiased capture of single cells from whole tissue cell suspension followed by RNAseq	Top 20 enriched transcripts	Astrocyte_All	21
				Endothelial_All	21
				Microglia_All	21
				Neuron_All	21
				Oligodendrocyte_Mature	21
				Oligodendrocyte_Progenitor Cell	21
Doyle et al., <i>Cell</i> , 2008	Cortex, Striatum, Cerebellum, Spinal Cord, Basal Forebrain, and Brain Stem of young transgenic mice	Capture of translated mRNA from specific cell types labeled in transgenic mice using translating ribosome affinity purification (TRAP) followed by microarray.	Top 25 enriched transcripts determined by iterative rank comparisons	Astrocyte_All	25
				Neuron_CorticoSpinal	25
				Neuron_CorticoStriatal	25
				Neuron_CorticoThalamic	25
				Neuron_Interneuron_CORT	25
				Neuron_Neuron_CCK	25
				Neuron_Neuron_PNOC	24
				Oligodendrocyte_All	25
				Oligodendrocyte_Mature	25
Daneman et al., <i>PLOS</i> , 2010	Cortex of young transgenic mice	Fluorescent cell sorting using antibodies to deplete non-specific cell types followed by Affymetrix microarray	>20 Fold enrichment for endothelial, >8 fold enrichment for vasculature	Endothelial_All	49
				Mural_Vascular	50
Sugino et al., <i>Nature Neuro</i> , 2006	Cingulate and Somatosensory Cortices, Basolateral Amygdala, CA1-CA3 Hippocampus, and Dorsal Lateral Geniculate Nucleus of the Thalamus of transgenic mice	Hand-sorting fluorescently-labeled cells followed by amplification and Affymetrix microarray	Enriched with $p < 1.5E-11$	Neuron_GABA	32
				Neuron_Glutamate	67
<i>Gene card</i>	Human	Erythrocyte-related genes	Unknown	RBC_All	17

92

93 **Figure 1. Thousands of transcripts have been identified as specifically-enriched in particular cortical**
 94 **cell types within published single-cell or purified cell type transcriptomic experiments (“reference**
 95 **datasets”). The experimental and statistical methods for determining whether a transcript was enriched**
 96 **in a cell type varied by publication, and included both RNA-Seq and microarray datasets.**

97

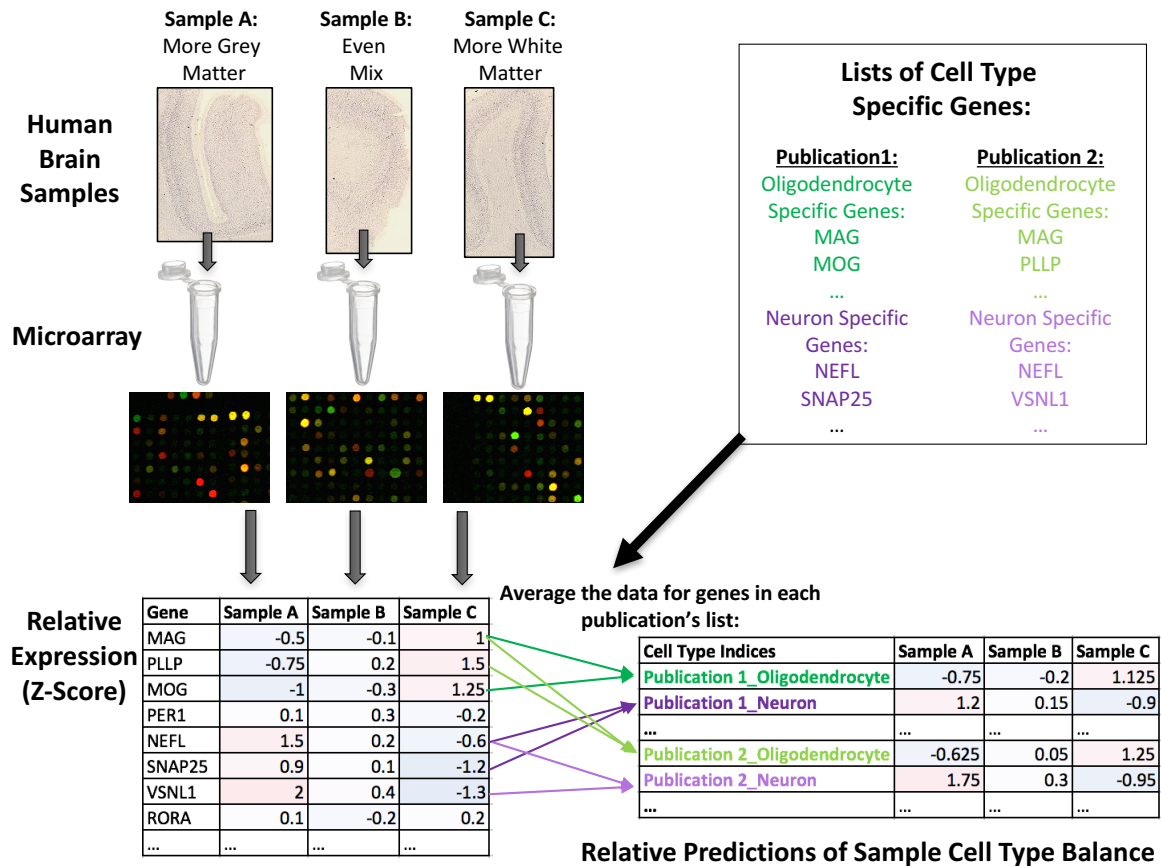
98 **2.2 “BrainInABlender”: Employing the Database of Cell Type Specific Transcripts to Predict**
99 **Relative Cell Type Balance in Heterogenous Brain Samples**

100 Next, we designed a method that uses the collective expression of cell type specific transcripts in
101 brain tissue samples to predict the relative cell type balance of the samples (“BrainInABlender”). We
102 specifically designed our method to be compatible with large, highly-normalized human brain
103 transcriptional profiling datasets such as those used by our neuropsychiatric research consortium
104 (Pritzker). We have made our method publicly-available in the form of a downloadable R package
105 (<https://github.com/hagenaue/BrainInABlender>).

106 In brief, BrainInABlender extracts the data from any particular transcriptional profiling dataset
107 (microarray, RNA-Seq) that represent genes identified in our database as having cell type specific
108 expression in the brain (as curated by official gene symbol). The expression-level data for each of these
109 transcripts (RNA-Seq: gene-level summary, microarray: probe or probeset summary) are then centered
110 and scaled across samples (mean=0, sd=1) to prevent transcripts with more variable signal from exerting
111 disproportionate influence on the results. Then, if necessary, the normalized data from all transcripts
112 representing the same gene are averaged for each sample and re-normalized. Finally, for each sample,
113 these values are averaged across the genes identified as having expression specific to a particular cell type
114 in each publication included in the database of cell transcripts. This creates 38 cell type signatures derived
115 from the cell type specific genes identified by the eight publications (“Cell Type Indices”, **Figure 1**), each
116 of which predicts the relative content for one of the 10 primary cell types in our brain samples (**Figure 2**).

117 Please note that our method was specifically designed to tackle challenges present in the Pritzker
118 Consortium microarray data, but we later discovered that it bears some resemblance to the existing
119 method of Population Specific Expression Analysis (PSEA, (10–12)). A more detailed discussion of the
120 similarities and differences between the techniques can be found in **Section 7.2**.

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121 **Figure 2. Predicting the relative cell type balance in human brain samples using genes previously-**
 122 **identified as having cell type specific expression.** Within macro-dissected brain tissue samples, variable
 123 cell type balance is likely to influence the pattern of gene expression. To estimate this variability, we
 124 extracted the microarray data for probe sets representing genes that had been previously identified as
 125 having cell type specific expression in previous publications (“Lists of Cell Type Specific Genes”, **Figure**
 126 **1**) and then averaged across the transcripts identified as specific to a particular cell type in each
 127 publication to create 38 different “Cell Type Indices” that predicted relative cell content in each of the
 128 brain samples.

129

130 **2.3 Validation of Relative Cell Content Predictions Using Datasets Derived from Purified or**
 131 **Cultured Cells**

132 We validated the method using publicly-available datasets from purified cell types and artificial
 133 cell mixtures (**Supplementary Methods and Results**). We found that the statistical cell type indices easily
 134 predicted the cell type identities of purified samples (datasets GSE52564 and GSE6783; (2,18); **Suppl.**
 135 **Figure 1, Suppl. Figure 2**). This was true regardless of the publication from which the cell type specific
 136 genes were derived: cell type specific gene lists derived from publications using different species (human

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137 vs. mouse), platforms (microarray vs. RNA-Seq), methodologies (florescent cell sorting vs. suspension),
138 or statistical stringency all performed fairly equivalently, with some minor exception. Occasionally, we
139 found that the cell type indices associated with cell type specific gene lists derived from TRAP
140 methodology (15) did not properly predict the cell identity of the samples. In general the cell type indices
141 associated with immature oligodendrocytes were somewhat inconsistent, most likely due to their
142 dependency on developmental stage and experimental conditions.

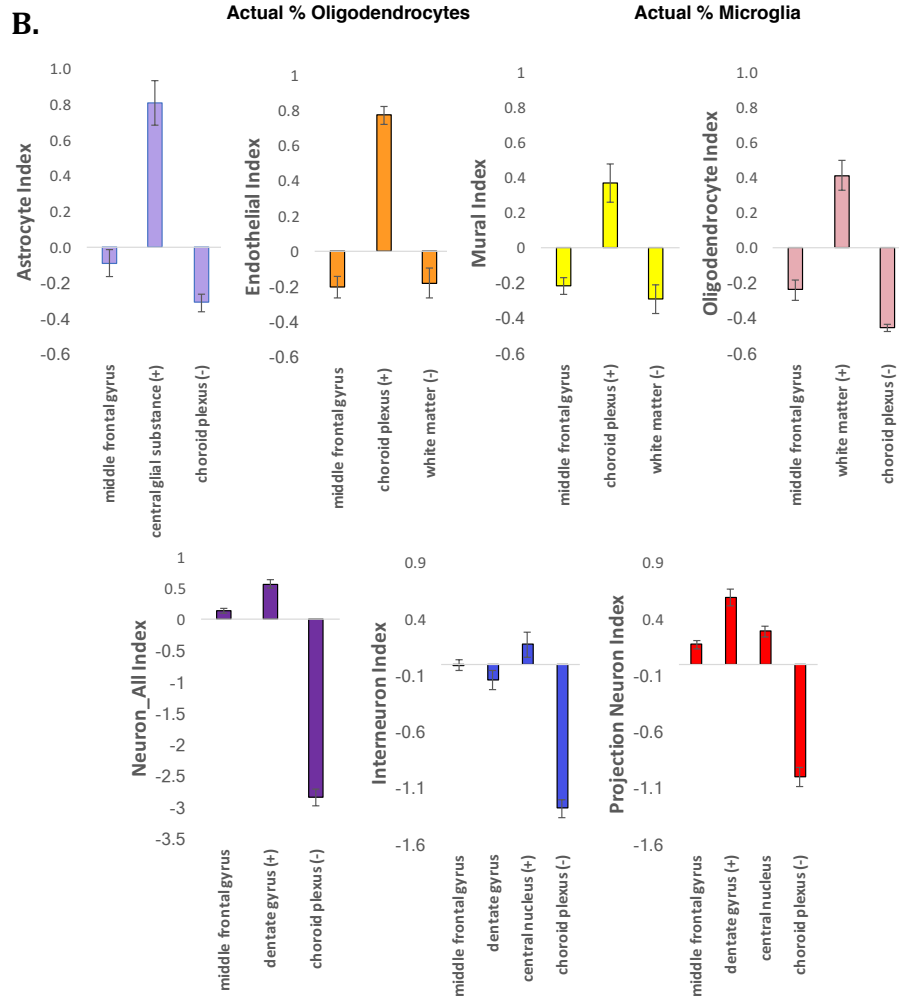
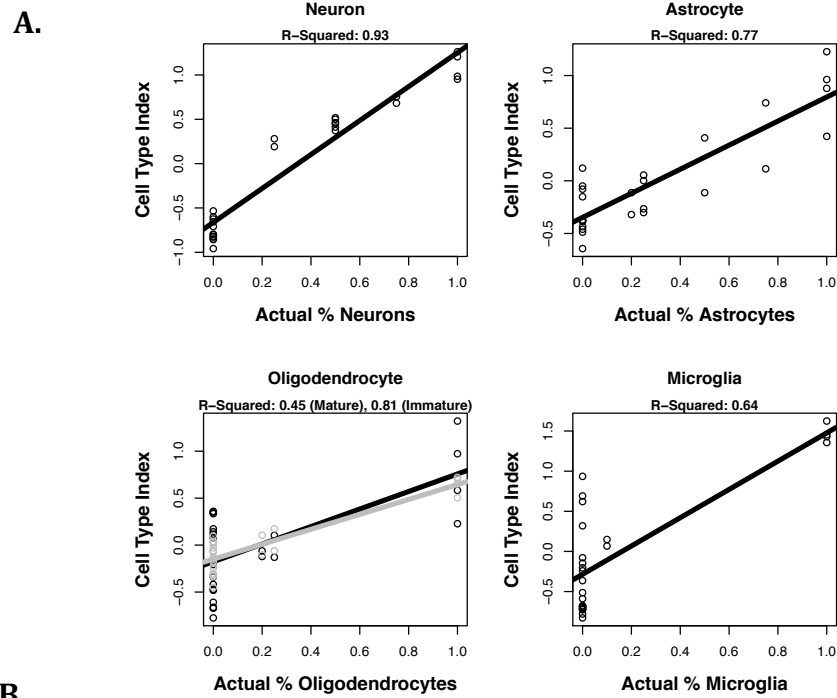
143 Therefore, overall we found substantial support for simply averaging the individual publication-
144 specific cell type indices within each of ten primary categories (astrocytes, endothelial cells, mural cells,
145 microglia, immature and mature oligodendrocytes, red blood cells, interneurons, projection neurons, and
146 indices derived from neurons in general) to produce ten consolidated primary cell-type indices for each
147 sample. To perform this consolidation, we also removed any transcripts that were identified as “cell type
148 specific” to multiple primary cell type categories (**Suppl. Figure 5**). These consolidated indices are
149 included as a output from BrainInABlender.

150 Next, as further validation, we determined whether relative cell type balance could be accurately
151 deciphered from microarray data for samples containing artificially-generated mixtures of cultured cells
152 (GSE19380; (12)). We found that the consolidated cell type indices produced by BrainInABlender
153 strongly correlated with the actual percentage of cells of a particular type included in the artificial
154 mixtures (**Figure 3**, Neuron% vs. Neuron_All Index: R-squared=0.93, $p=1.54e-15$, Astrocyte % vs.
155 Astrocyte Index: R-squared=0.77, $p=5.05e-09$, Microglia% vs. Microglia Index: R-Squared=0.64, $p=$
156 $8.2e-07$), although we found that the cell type index for immature oligodendrocytes better predicted the
157 percentage of cultured oligodendrocytes in the samples than the cell type index for mature
158 oligodendrocytes (Mature: R-squared=0.45, $p=0.000179$, Immature: R-squared=0.81, $p=4.14e-10$). We
159 believe this discrepancy is likely to reflect the specific cell culture conditions used in the original
160 admixture experiment. In a follow-up analysis, artificial mixtures of cells produced *in silico* by averaging
161 randomly-selected data from purified cell types similarly indicated that the cell type indices produced by

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162 BrainInABlender follow a linear relationship with actual cell type balance in mixed samples, even for less
163 prevalent cell varieties (endothelial, **Suppl. Figure 3, Suppl. Figure 4**).

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164 **Figure 3. Validation of Relative Cell Content Predictions.** *A) Using a microarray dataset derived from*
165 *samples that contained artificially-generated mixtures of cultured cells (GSE19380; (12)), we found that*
166 *our relative cell content predictions (“cell type indices”) closely reflected actual known content. B) Our*
167 *cell type indices also easily differentiated human post-mortem samples derived from brain regions that*
168 *are known to contain relatively more (+) or less (-) of the targeted the cell type of interest. Results from*
169 *the middle frontal gyrus are included for comparison, since the rest of the paper primarily focuses on*
170 *prefrontal cortical data. (Bars: average +/-SE).*

171

172 **2.4 Validation of Relative Cell Content Predictions Using a Dataset Derived from Human Post-**

173 **Mortem Tissue**

174 Next, we wanted to see whether the cell content predictions produced by BrainInABlender
175 correctly reflected relative cell type balance in human post-mortem samples. To test this, we applied our
176 method to a large human post-mortem Agilent microarray dataset (841 samples) spanning 160 cortical
177 and subcortical brain regions from the Allen Brain Atlas (**Suppl. Table 2**; (25)). This dataset was derived
178 from high-quality tissue (absence of neuropathology, pH>6.7, post-mortem interval<31 hrs, RIN>5.5)
179 from 6 human subjects (26). The tissue samples were collected using a mixture of block dissection and
180 laser capture microscopy guided by adjacent tissue sections histologically stained to identify traditional
181 anatomical boundaries (27). Prior to data release, the dataset had been subjected to a wide variety of
182 normalization procedures to eliminate technical variation (28) which included log(base2) transformation,
183 centering and scaling for each probe (<http://human.brain-map.org/microarray/search>, December 2015).

184 After applying BrainInABlender to the dataset, we extracted the results for a selection of brain
185 regions that are known to contain relatively more (+) or less (-) of particular cell types (the results for the
186 other brain regions can be found in **Suppl. Table 3**). The results clearly indicated that our cell type
187 analyses could identify well-established differences in cell type balance across brain regions (**Figure 3**).
188 Within the choroid plexus, which is a villous structure located in the ventricles made up of support cells
189 (epithelium) and an extensive capillary network (29), there was an elevation of gene expression specific
190 to vasculature (endothelial cells, mural cells). In the corpus callosum and cingulum bundle, which are
191 large myelinated fiber tracts (29), there was an enrichment of oligodendrocytes- and microglia-specific
192 gene expression. The central glial substance was enriched with gene expression specific to glia and

193 support cells, with a particular emphasis on astrocytes. The dentate gyrus, which contains densely packed
194 glutamatergic granule cells projecting to the mossy fibre pathway (30), was enriched for gene expression
195 specific to projection neurons. The central nucleus of the amygdala, which includes a large number of
196 GABA-ergic neurons (31), had a slight enrichment of gene expression specific to interneurons. These
197 results provide fundamental validation that our methodology can accurately predict relative cell type
198 balance in human post-mortem samples. Moreover, these results suggest that each of the consolidated cell
199 type indices is capable of generally tracking their respective cell types in subcortical structures, despite
200 the fact that our analysis method relies on cell type specific genes originally identified in the forebrain
201 and cortex.

202

203 **2.5 Using Cell Type Specific Transcripts to Predict Relative Cell Content in Transcriptomic Data** 204 **from Macro-Dissected Human Cortical Tissue from Psychiatric Subjects**

205 Next, we examined the collective variation in the levels of cell type specific transcripts in several
206 large psychiatric human brain microarray datasets. The first was a large Pritzker Consortium Affymetrix
207 U133A microarray dataset derived from high-quality human post-mortem dorsolateral prefrontal cortex
208 samples (final sample size of 157 subjects, **Suppl. Table 14**), including tissue from subjects without a
209 psychiatric or neurological diagnosis (“Controls”, n=71), or diagnosed with Major Depressive Disorder
210 (“MDD”, n=40), Bipolar Disorder (“BP”, n=24), or Schizophrenia (“Schiz”, n= 22). The severity and
211 duration of physiological stress at the time of death was estimated by calculating an agonal factor score
212 for each subject (ranging from 0-4, with 4 representing severe physiological stress; (32,33)). Additionally,
213 we measured the pH of cerebellar tissue as an indicator of the extent of oxygen deprivation experienced
214 around the time of death (32,33) and calculated the interval between the estimated time of death and the
215 freezing of the brain tissue (the postmortem interval or PMI) using coroner records. The transcriptional
216 profiling of these samples had originally been processed in batches across multiple laboratories (1-5
217 replicates per sample). Before averaging the replicate samples for each subject, the data was highly
218 normalized to correct for technical variation, including robust multi-array analysis (RMA) (34) and

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219 median-centering (detailed procedure: (35)). Our current analyses began with this subject-level summary
220 gene expression data (GSE92538).

221 We determined the replicability of our results using three smaller publicly-available post-mortem
222 human cortical Affymetrix U133Plus2 microarray datasets (GSE53987 (36), GSE21935 (37), GSE21138
223 (38), **Figure 4**). These datasets were selected because they included both psychiatric and control samples,
224 and provided pH, PMI, age, and gender in the demographic information on the Gene Expression Omnibus
225 website (<https://www.ncbi.nlm.nih.gov/geo/>). To control for technical variation, the sample processing
226 batches were estimated using the microarray chip scan dates extracted from the .CEL files and RNA
227 degradation was estimated using the R package AffyRNADegradation (39). Prior to running
228 BrainInABlender, the probe-level signal data from each dataset was normalized using RMA (34),
229 summarized using a custom .cdf (http://nmg-r.bioinformatics.nl/NuGO_R.html,
230 “hgu133plus2hsentrezgcdf_19.0.0”), and cleaned of any samples that appeared low-quality or
231 misidentified.

232 Finally, we also explored replicability within the recently-released large CommonMind
233 Consortium (CMC) human dorsolateral prefrontal cortex RNA-seq dataset (603 individuals (40)). This
234 dataset was downloaded from the CommonMind Consortium Knowledge Portal
235 (<https://www.synapse.org/CMC>) and analyzed at UCI. The bam files were converted to fastq files and re-
236 mapped to a more recent build of the human genome (GRCh38, (41)). The total reads mapping uniquely
237 to exons (defined by Ensembl) were transformed into logCPM values (42). Prior to data upload, poor
238 quality samples from the original dataset (40) were removed (<50 million reads, RIN<5.5) by the CMC
239 and replaced with higher quality samples. We additionally excluded data from 10 replicates and 89
240 individuals with incomplete demographic data (missing pH), leaving a final sample size of 514 samples.
241 We predicted the relative cell type content of these samples using a newer version of BrainInABlender
242 (v2) which excluded a few of the weaker cell type specific gene sets (15). Later, the expression data were
243 further filtered by expression threshold (CPM>1 in at least 50 individuals), leaving data from
244 approximately 17,000 genes.

245 In general, the code for all analyses in the paper can be found at <https://github.com/hagenaue/> or
 246 https://github.com/aschulmann/CMC_celltype_index.

247

Microarray:

GEO Accession #	Submitter & Date	Published?	Brain Bank	Brain Region	Sample Size (no outliers)	Subjects per group (no outliers)	# of replicates per sample	# of known Batches	AVE pH (+/- SD)	AVE Age (+/- SD)	AVE PMI (+/- SD)	% Female
GSE92538	Hagenaue (2016)	<i>current paper</i>	UC-Irvine	BA9/BA46	337	157: 71 CNTRL, 24 BP, 40 MDD, 22 SCHIZ	1-5 (AVE: 2)	15-36	6.8 (+/-0.3)	52 (+/-15)	24 (+/-9)	27%
GSE53987	Lanz (2014)	Lanz et al. (2015)	PITT	BA46: grey matter	66	66: 18 CNTRL, 17 BP, 17 MDD, 14 SCHIZ	1	0	6.6 (+/-0.3)	46 (+/-10)	20 (+/-6)	45%
GSE21935	Huxley-Jones (2011)	Barnes et al. (2011)	CCHPC	BA22	42	42: 19 CNTRL, 23 SCHIZ	1	2	6.3 (+/-0.3)	70 (+/-19)	8 (+/-5)	45%
GSE21138	Thomas (2010)	Narayan et al. (2008)	MHRI	BA46: grey matter	54	54: 27 CNTRL, 27 SCHIZ	1	5	6.3 (+/-0.2)	45 (+/-17)	40 (+/-13)	17%

RNA-Seq:

Public Data Release	Submitter & Date	Published?	Brain Bank	Brain Region	Sample Size (all)	Subjects per group (all)	# of replicates per sample	# of known Batches	AVE pH (+/- SD)	AVE Age (+/- SD)	AVE PMI (+/- SD)	% Female
Synapse.org	CMC (2016)	Fromer et al. (2016)	MSSM, PENN, PITT	BA9 / BA46 (PITT: grey matter)	621	603: 285 CNTRL, 263 SCZ, 47 BP, 8 AFF	1 (rarely 2)	491	6.5 (+/- 0.3)	65 (+/- 18) (binned 90+)	17 +/- 11	41%

248

249 **Figure 4. We examined the pattern of cell-type specific gene expression in five post-mortem human**
 250 **cortical tissue datasets that included samples from subjects with psychiatric illness. Abbreviations:**
 251 **CTRL: control, BP: Bipolar Disorder, MDD: Major Depressive Disorder, SCHIZ: Schizophrenia, GEO:**
 252 **Gene Expression Omnibus, BA: Brodmann's Area, PMI: Post-mortem interval, SD: Standard Deviation,**
 253 **Brain Banks: UC-Irvine (University of California – Irvine), PITT (University of Pittsburgh), CCHPC**
 254 **(Charing Cross Hospital Prospective Collection), MSSM (Mount Sinai Icahn School of Medicine), MHRI**
 255 **(Mental Health Research Institute Australia), PENN (University of Pennsylvania)**

256

257 **2.6 Does the Reference Dataset Matter? Cell Type Specific Transcripts Identified by Different**
 258 **Publications Produce Similar Predictions of Relative Cell Type Balance**

259 We first confirmed that the predicted cell content for our post-mortem human cortical samples
 260 (“cell type indices”) was similar regardless of the methodology used to generate the cell type specific
 261 gene lists used in the predictions. Within all five of the human cortical transcriptomic datasets, there was
 262 a strong positive correlation between cell type indices representing the same cell type, even when the
 263 predictions were derived using cell type specific gene lists from different species, cell type purification

264 strategies, and platforms. Clustering within broad cell type categories was clear using visual inspection of
265 the correlation matrices (**Suppl. Figure 11, Suppl. Figure 12**), hierarchical clustering, or consensus
266 clustering (**Suppl. Figure 13**, ConsensusClusterPlus: (43), **Suppl. Figure 14, Suppl. Figure 16**). In some
267 datasets, the cell type indices for support cell subcategories were nicely clustered and in others they were
268 difficult to fully differentiate (**Suppl. Figure 11, Suppl. Figure 12**). Clustering was not able to reliably
269 discern neuronal subcategories (interneurons, projection neurons) in any dataset. Similar to our previous
270 validation analyses, oligodendrocyte progenitor cell indices derived from different publications did not
271 strongly correlate with each other, perhaps due to heterogeneity in the progenitor cell types sampled by
272 the original publications.

273 Therefore, for further analyses in the post-mortem human datasets, we consolidated the cell type
274 indices using a procedure similar to our previous validation analyses. To do this, we averaged the 38
275 publication-specific cell type indices within each of ten primary categories: astrocytes, endothelial cells,
276 mural cells, microglia, immature and mature oligodendrocytes, red blood cells, interneurons, projection
277 neurons, and indices derived from neurons in general, with any transcripts that overlapped between
278 categories removed (**Suppl. Figure 17**). This led to ten consolidated primary cell-type indices for each
279 sample.

280

281 **3. Results**

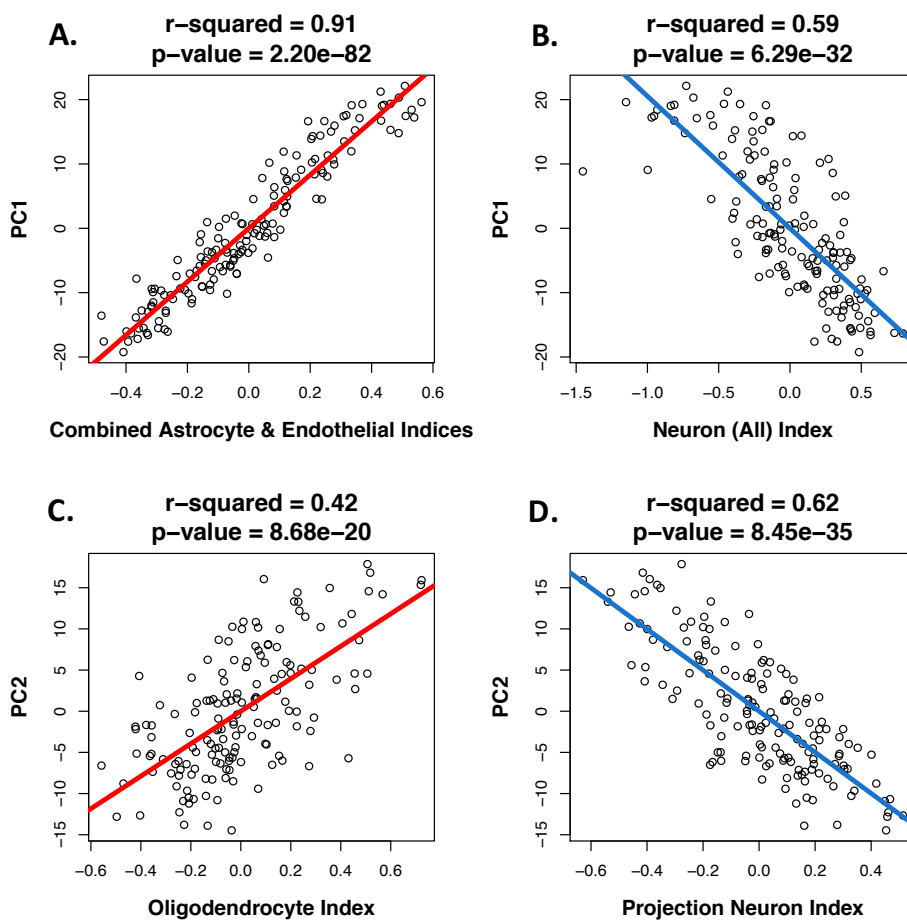
282

283 **3.1 Inferred Cell Type Composition Explains a Large Percentage of the Sample-Sample Variability** 284 **in Microarray Data from Macro-Dissected Human Cortical Tissue**

285 Using principal components analysis we found that the primary gradients of variation in all four
286 of the cortical datasets strongly correlated with our estimates of cell type balance. For example, while
287 analyzing the Pritzker dorsolateral prefrontal cortex microarray dataset, we found that the first principal
288 component, which encompassed 23% of the variation in the dataset, spanned from samples with high

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289 support cell content to samples with high neuronal content. Therefore, a large percentage of the variation
290 in PC1 (91%) was accounted for by an average of the astrocyte and endothelial indices ($p < 2.2e-82$, with a
291 respective r-squared of 0.80 and 0.75 for each index analyzed separately) or by the general neuron index
292 ($p < 6.3e-32$, r-squared=0.59; **Figure 5**). The second notable gradient in the dataset (PC2) encompassed
293 12% of the variation overall, and spanned samples with high projection neuron content to samples with
294 high oligodendrocyte content (with a respective r-squared of 0.62 and 0.42, and respective p-values of
295 $p < 8.5e-35$ and $p < 8.7e-20$).



296

297 **Figure 5. Cell content predictions explain a large percentage of the variability in microarray data**
298 **derived from the human cortex.** As an example, within the Pritzker dataset the first principal component
299 of variation (PC1) encompassed 23% of the variation in the dataset, and was A) positively correlated
300 with predicted “support cell” content in the samples (a combination of the astrocyte and endothelial
301 indices: r-squared: 0.91, $p < 2.2e-82$) and B) negatively correlated with predicted neuronal content (r-
302 squared=0.59, $p < 6.3e-32$). The second principal component of variation (PC2) encompassed 12% of
303 variation in the dataset, and was C) positively correlated with predicted oligodendrocyte content in the

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304 *samples (r-squared: 0.42, $p < 8.7e-20$) and D) negatively correlated with predicted projection neuron*
305 *content (r-squared: 0.62, $p < 8.5e-35$). Examples in other datasets can be found in **Suppl. Figure 20**.*

306
307 When digging deeper, we found that none of the original 38 publication-specific cell type indices
308 were noticeably superior to the consolidated indices when predicting the principal components of
309 variation in the dataset. Human-derived indices did not outperform mouse-derived indices, and indices
310 derived from studies using stricter definitions of cell type specificity (fold enrichment cut-off in **Figure 1**,
311 e.g., (13) vs. (17)) did not outperform less strict indices.

312 Within the other four human cortical tissue datasets, the relationships between the top principal
313 components of variation and the consolidated cell type indices were similarly strong (**Suppl. Figure 20**),
314 despite the fact that these datasets had received less preprocessing to remove the effects of technical
315 variation. Within the GSE21935 dataset (published in (37)) the first principal component of variation
316 accounted for 37% of the variation in the dataset and similarly seemed to represent a gradient running
317 from samples with high support cell content (PC1 vs. endothelial index: r-squared= 0.85, $p < 3.6e-18$, PC1
318 vs. astrocyte index: r-squared= 0.67, $p < 3.6e-11$) to samples with high neuronal content (PC1 vs.
319 neuron_all index: r-squared= 0.85, $p < 3.9e-18$). Within the GSE53987 dataset (submitted to GEO by
320 Lanz, 2014), which had samples derived exclusively from gray-matter-only dissections, the first principal
321 component of variation accounted for 13% of the variation in the dataset and was highly correlated with
322 predicted astrocyte content (PC1 vs. astrocyte index: r-squared=0.80, $p < 4.6e-24$). In GSE21138
323 (published in (39)), which also had samples derived exclusively from gray-matter-only dissections, the
324 first principal component of variation accounted for 23% of the variation in the dataset and was strongly
325 related to technical variation (batch), but the second principal component of variation, which accounted
326 for 14% of the variation in the dataset, again represented a gradient from samples with high support cell
327 content to high neuronal content (PC2 vs. astrocyte: r-squared=0.56, $p < 8.3e-11$, PC2 vs. neuron_all: r-
328 squared=0.54, $p < 2.3e-10$). Finally, within the CMC RNA-Seq dataset, the first principal component of
329 variation accounted for 16% of the variation in the dataset and was highly correlated with projection
330 neuron content (PC1 vs. Neuron_Projection: r-squared=0.54, $p = 5.77e-104$).

331 To confirm that the strong relationship between the top principal components of variation and our
332 cell type composition indices did not originate artificially due to cell type specific genes representing a
333 large percentage of the most highly variable transcripts in the dataset, we repeated the principal
334 components analysis in the Pritzker dataset after excluding all cell type specific transcripts from the
335 dataset and still found these strong correlations (**Suppl. Figure 21**). Indeed, individual cell type indices
336 better accounted for the main principal components of variation in the microarray data than *all other*
337 *major subject variables combined* (pH, Agonal Factor, PMI, Age, Gender, Diagnosis, Suicide; PC1: R-
338 squared=0.4272, PC2: R-squared=0.2176). When examining the dataset as a whole, the six subject
339 variables accounted for an average of only 12% of the variation for any particular probe (R-squared,
340 Adj.R-squared=0.0715), whereas just the astrocyte and projection neuron indices alone were able to
341 account for 17% (R-squared, Adj.R-squared=0.1601) and all 10 cell types accounted for an average of
342 31% (R-squared, Adj.R-squared=0.263), almost one third of the variation present in the data for any
343 particular probe (**Suppl. Figure 22**).

344 These results indicated that accounting for cell type balance is important for the interpretation of
345 post-mortem human brain microarray and RNA-Seq data and might improve the signal-to-noise ratio in
346 analyses aimed at identifying psychiatric risk genes.

347 **3.2 Cell Content Predictions Derived from Microarray Data Match Known Relationships Between** 348 **Clinical/Biological Variables and Brain Tissue Cell Content**

349 We next set out to observe the relationship between the predicted cell content of our samples and
350 a variety of medically-relevant subject variables, including variables that had already been demonstrated
351 to alter cell content in the brain in other paradigms or animal models. To perform this analysis, we first
352 examined the relationship between seven relevant subject variables and each of the ten cell type indices in
353 the Pritzker prefrontal cortex dataset using a linear model that allowed us to simultaneously control for
354 other likely confounding variables in the dataset:

355 **Equation 1:**

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356 Cell Type Index= $\beta_0 + \beta_1 * (\text{Brain pH}) + \beta_2 * (\text{Agonal Factor})$
357 $+ \beta_3 * (\text{PMI}) + \beta_4 * (\text{Age}) + \beta_5 * (\text{Sex}) + \beta_6 * (\text{Diagnosis}) + \beta_7 * (\text{Exsanguination}) + \epsilon$
358

359 We then examined the replicability of these relationships using data from the three smaller
360 publicly-available human post-mortem microarray datasets (GSE53987, GSE21935, GSE21138). For
361 these datasets, we initially lacked detailed information about manner of death (agonal factor and
362 exsanguination), but were able to control for technical variation within the model using statistical
363 estimates of RNA degradation and batch (scan date):

364 **Equation 2:**

365 Cell Type Index= $\beta_0 + \beta_1 * (\text{Brain pH}) + \beta_2 * (\text{PMI}) + \beta_3 * (\text{Age}) + \beta_4 * (\text{Sex}) + \beta_5 * (\text{Diagnosis}) +$
366 $\beta_6 * (\text{RNA Degradation}) + \beta_7 * (\text{Batch, when applicable}) + \epsilon$

367 We evaluated the replicability of these relationships across the four microarray datasets by performing a
368 meta-analysis for each variable and cell type combination. To do this, we applied random effects
369 modeling to the respective betas and accompanying sampling variance derived from each dataset using
370 the *rma.mv()* function within the *metafor* package (44). P-values were then corrected for multiple
371 comparisons following the Benjamini-Hochberg method (*q-value*) using the *mt.rawp2adjp* function
372 within the *multtest* package (45).

373 Finally, we characterized these relationships in the large CMC human post-mortem RNA-Seq
374 dataset. For this dataset, we had some information about manner of death but lacked knowledge of agonal
375 factor or exsanguination. We controlled for technical variation due to dissection site (institution) and
376 RNA degradation (RIN):

377 **Equation 3:**

378 Cell Type Index= $\beta_0 + \beta_1 * (\text{Brain pH}) + \beta_2 * (\text{PMI}) + \beta_3 * (\text{Age}) + \beta_4 * (\text{Sex}) + \beta_5 * (\text{Diagnosis}) +$
379 $\beta_6 * (\text{RNA Degradation}) + \beta_7 * (\text{Institution}) + \beta_8 * (\text{MannerOfDeath}) + \epsilon$
380

381 This analysis uncovered many well-known relationships between brain tissue cell content and clinical
382 or biological variables (**Figure 6, Suppl. Table 4**). First, as a proof of principle, we were able to clearly

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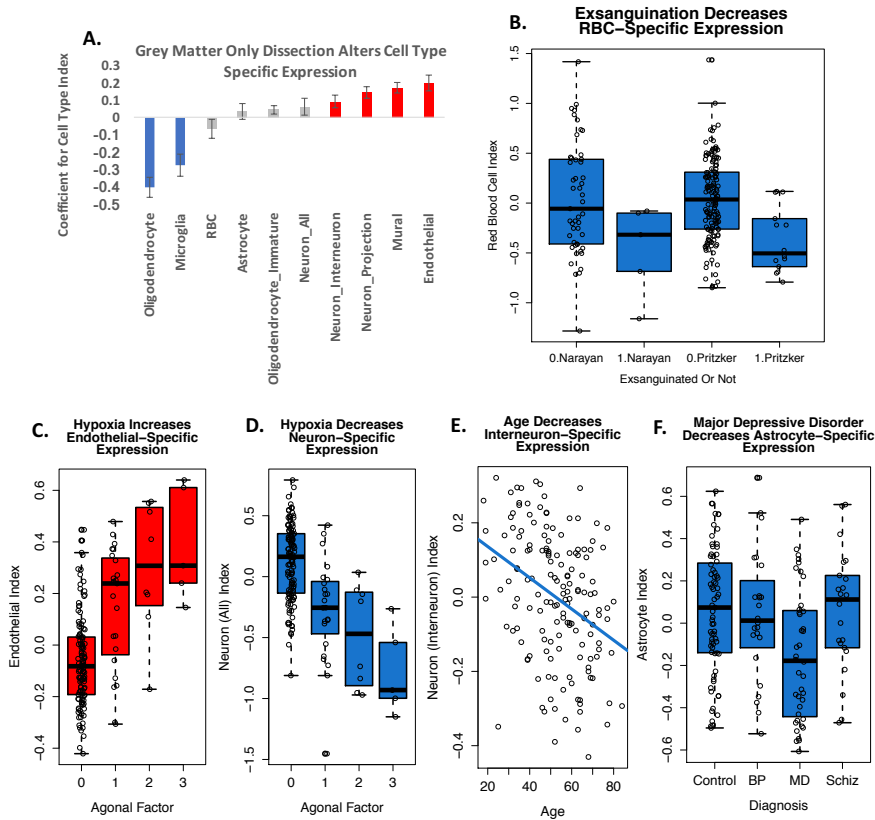
383 observe dissection differences between institutions within the large CMC RNA-Seq dataset, with samples
384 from University of Pittsburgh having a predicted relative cell type balance that closely matched what
385 would be expected due to their gray matter only dissection method (Oligodendrocyte: $\beta = -0.404$, $p = 2.42e-11$,
386 $q = 4.03e-10$; Microglia: $\beta = -0.274$, $p = 3.06e-05$, $q = 2.42e-04$; Neuron_Interneuron: $\beta = 0.0916$,
387 $p = 0.0161$, $q = 0.525$; Neuron_Projection: $\beta = 0.145$, $p = 2.31e-05$, $q = 1.93e-04$; Mural: $\beta = 0.170$, $p = 2.14e-08$,
388 $q = 2.68e-07$; Endothelial: $\beta = 0.200$, $p = 1.12e-05$, $q = 1.12e-04$). Samples from University of Pennsylvania
389 were associated with lower predicted cell content related to vasculature (Endothelial: $\beta = -0.255$, $p = 4.01e-04$,
390 $q = 2.40e-03$; Mural: $\beta = -0.168$, $p = 4.59e-04$, $q = 2.59e-03$; Astrocyte: $\beta = -0.189$, $p = 7.47e-03$, $q = 0.0287$).

391 Predicted cell type content was also closely related to manner of death. For example, within the
392 Pritzker dataset we found that subjects who died in a manner that involved exsanguination had a notably
393 low red blood cell index ($\beta = -0.398$; $p = 0.00056$). Later, we were able to replicate this result within
394 GSE21138 using data from 5 subjects who we discovered were also likely to have died in a manner
395 involving exsanguination ($\beta = -0.516$, $p = 0.052$ **trend*, manner of death reported in suppl. in (38)). The
396 presence of prolonged hypoxia around the time of death, as indicated by either low brain pH or high
397 agonal factor score within the Pritzker dataset, was associated with a large increase in the endothelial cell
398 index (Agonal Factor: $\beta = 0.118$, $p = 2.85e-07$; Brain pH: $\beta = -0.210$, $p = 0.0003$) and astrocyte index (Brain
399 pH: $\beta = -0.437$, $p = 2.26e-07$; Agonal Factor: $\beta = 0.071$, $p = 0.024$), matching previous demonstrations of
400 cerebral angiogenesis, endothelial and astrocyte activation and proliferation in low oxygen environments
401 (46). Small increases were also seen in the mural index in response to low-oxygen (Mural vs. Agonal
402 Factor: $\beta = 0.0493493$, $p = 0.0286$), most likely reflecting angiogenesis. In contrast, prolonged hypoxia
403 was associated with a clear decrease in all of the neuronal indices (Neuron_All vs. Agonal Factor: $\beta = -$
404 0.242 , $p = 3.58e-09$; Neuron_All vs. Brain pH: $\beta = 0.334$, $p = 0.000982$; Neuron_Interneuron vs. Agonal
405 Factor: $\beta = -0.078$, $p = 4.13e-05$; Neuron_Interneuron vs. Brain pH: $\beta = 0.102$, $p = 0.034$; Neuron_Projection
406 vs. Agonal Factor: $\beta = -0.096$, $p = 0.000188$), mirroring the notorious vulnerability of neurons to low
407 oxygen (e.g., (47)). These overall effects of hypoxia on cell type balance replicated in the smaller human
408 microarray post-mortem datasets, despite lack of information about agonal factor (Astrocyte vs. Brain pH

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409 (meta-analysis: $b = -0.459$, $p = 2.59 \times 10^{-11}$, $q = 2.33 \times 10^{-9}$): Narayan et al. 2008: $\beta = -0.856$, $p = 0.00661$, Lanz
410 2014: $\beta = -0.461$, $p = 0.00812$, Neuron_All vs. Brain pH (meta-analysis: $b = 0.245$, $p = 7.72 \times 10^{-4}$, $q = 1.16 \times 10^{-2}$), Neuron_Interneuron vs. Brain pH (meta-analysis: $b = 0.109$, $p = 7.89 \times 10^{-3}$, $q = 5.52 \times 10^{-2}$ **trend*): Narayan
411 et al. 2008: $\beta = 0.381134$, $p = 0.0277$) and partially replicated in the CMC human RNA-Seq dataset
412 (Neuron_Interneuron vs. Brain pH: $\beta = 0.186$, $p = 9.81 \times 10^{-5}$, $q = 6.69 \times 10^{-4}$). In several datasets, we also found
413 that prolonged hypoxia correlated with fewer microglia (Microglia vs. Brain pH: Lanz 2014: $\beta = 0.462$,
414 $p = 0.00603$; CMC: $\beta = 0.286$, $p = 4.66 \times 10^{-4}$, $q = 2.59 \times 10^{-3}$), which may suggest that our microglia cell type
415 index is specifically tracking ramified microglia, although we did not observe a relationship between
416 microglia and death related to infection/parasitic disease (CMC: Microglia vs. CauseOfDeath(infection):
417 $\beta = 0.231$, $p = 0.121$, $q = 0.256$).
418
419

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The most highly replicated effects across datasets:	Pritzker	Lanz 2014	Barnes et al. 2011	Narayan et al. 2008	Meta-Analysis: Nominal P-value	Meta-Analysis: BH Adjusted P-value	CMCC	CMCC: Nominal P-value	CMCC: BH Adjusted P-value	Replication in Microarray Meta-analysis and CMCC
Hypoxia:										
Astrocyte vs. Brain pH	-5.4	-2.7	-1.1	-2.9	2.59E-11	2.33E-09	2.1	3.25E-02	9.55E-02	
Neuron_All vs. Brain pH	3.4	0.1	1.3	1.9	7.72E-04	1.16E-02	0.6	5.60E-01	7.00E-01	*
Neuron_Interneuron vs. Brain pH	2.1	1.0	-0.5	2.3	7.89E-03	5.52E-02	3.9	9.81E-05	6.69E-04	**
Microglia vs. Brain pH	-1.0	2.9	-0.1	1.9	5.41E-01	7.91E-01	3.5	4.66E-04	2.59E-03	*
Oligodendrocyte_Immature vs. Brain pH	-3.0	0.4	-0.5	1.4	1.40E-01	3.71E-01	3.0	2.68E-03	1.22E-02	
Age										
Neuron_All vs. Age	-1.9	-2.1	-1.2	-1.2	1.57E-03	2.02E-02	-4.3	2.27E-05	1.93E-04	**
Neuron_Interneuron vs. Age	-3.4	-0.5	-2.5	-2.5	2.91E-06	6.56E-05	-6.5	2.10E-10	3.15E-09	**
Neuron_Projection vs. Age	-2.8	-3.7	-0.5	-3.1	1.61E-06	4.83E-05	-7.5	2.93E-13	7.33E-12	**
Oligodendrocyte vs. Age	1.8	1.2	0.3	3.1	2.74E-03	2.74E-02	1.6	1.02E-01	2.25E-01	*
Oligodendrocyte_Immature vs. Age	-3.7	-4.7	-0.2	-4.5	5.98E-11	2.69E-09	-11.0	3.32E-25	2.49E-23	**
PMI										
Oligodendrocyte vs. PMI	-3.6	-3.6	-1.6	-0.5	2.23E-05	4.02E-04	-4.1	4.70E-05	3.36E-04	**
Endothelial vs. PMI	-2.0	-0.8	0.4	-0.1	5.51E-02	2.36E-01	-3.9	1.32E-04	8.60E-04	*
Microglia vs. PMI	-1.0	-1.5	-1.2	-0.7	9.72E-02	3.05E-01	-3.5	5.15E-04	2.76E-03	*
Oligodendrocyte_Immature vs. PMI	3.5	1.0	1.6	-0.4	4.81E-03	4.33E-02	-0.4	6.86E-01	8.04E-01	
Neuron_Projection vs. PMI	3.9	1.6	-0.2	-1.2	2.28E-03	2.56E-02	3.1	1.97E-03	9.24E-03	**
Neuron_All vs. PMI	2.5	1.8	-0.4	0.0	1.74E-02	9.81E-02	2.6	1.10E-02	3.88E-02	*
Diagnosis:										
Astrocyte vs. Diagnosis_MDD	-2.6	-1.0			5.88E-03	4.81E-02				
Neuron_All vs. Diagnosis_BP	-0.9	-0.7			2.46E-01	5.39E-01	2.6	8.44E-03	3.17E-02	
RBC vs. Diagnosis_Schiz	-1.2	0.1	-1.0	-0.5	2.04E-01	4.96E-01	-2.5	1.41E-02	4.71E-02	*
Gender:										
Neuron_Interneuron vs. GenderFemale	-0.9	0.0	0.7	0.3	6.65E-01	9.06E-01	-2.5	1.20E-02	4.09E-02	
Neuron_Projection vs. GenderFemale	1.0	-0.3	-0.4	1.6	3.60E-01	6.46E-01	-2.5	1.11E-02	3.88E-02	

421 **Figure 6. Cell content predictions derived from microarray data match known relationships between**
422 **subject variables and brain tissue cell content.** Boxplots represent the median and interquartile range,
423 with whiskers illustrating either the full range of the data or 1.5x the interquartile range. A. Within the
424 CMC dataset, cortical tissue samples that were dissected to only contain gray matter (PITT) show lower
425 predicted oligodendrocyte and microglia content and more neurons and vasculature (bars: $\beta \pm SE$,
426 red/blue: $p < 0.05$). B. Subjects that died in a manner that involved exsanguination ($n=14$) had a notably
427 low red blood cell index in both the Pritzker ($p=0.00056$) and Narayan et al. datasets ($p=0.052$ *trend).
428 C. The presence of prolonged hypoxia around the time of death, as indicated by high agonal factor score,
429 was associated with a large increase in the endothelial cell index ($p=2.85e-07$) matching previous
430 demonstrations of cerebral angiogenesis, activation, and proliferation in low oxygen environments (46).
431 D. High agonal factor was also associated with a clear decrease in neuronal indices ($p=3.58e-09$)
432 mirroring the vulnerability of neurons to low oxygen (47). E. Age was associated with a decrease in the
433 neuronal indices ($p=0.000956$) which fits known decreases in gray matter density in the frontal cortex in
434 aging humans (48). F. Major Depressive Disorder was associated with a moderate decrease in astrocyte
435 index ($p=0.0118$), which fits what has been observed morphometrically (49). G. The most highly-
436 replicated relationships between subject variables and predicted cortical tissue cell content across all five
437 of the post-mortem human datasets. Provided in the table are the T-stats for the effects
438 (red=upregulation, blue=downregulation), derived from a larger linear model controlling for confounds
439 (**Equation 1, Equation 2, Equation 3**), as well as the nominal p-values from the meta-analysis of the
440 results across the four microarray studies, and p-values following multiple-comparisons correction (q-
441 value). Only effects that had a $q < 0.05$ in either our meta-analysis or the large CMC RNA-Seq dataset are
442 included in the table. Asterisks denote effects that had consistent directionality in the meta-analysis and
443 CMC dataset (*) or consistent directionality and $q < 0.05$ in both datasets (**). Please note that lower pH
444 and higher agonal factor are both indicators of greater hypoxia prior to death, but have an inverted
445 relationship and therefore show opposing relationships with the cell type indices (e.g., when pH is low
446 and agonal factor is high, support cell content is increased).

447
448 In the Pritzker dataset, age was associated with a moderate decrease in two of the neuronal indices
449 (Neuron_Interneuron vs. Age: $\beta = -0.00291$, $p = 0.000956$; Neuron_Projection Neuron vs. Age: $\beta =$
450 0.00336 , $p = 0.00505$) and was strongly replicated in the large CMC RNA-Seq dataset (Neuron_All vs.
451 Age: $\beta = -0.00497$, $p = 2.27e-05$, $q = 1.93e-04$; Neuron_Projection Neuron vs. Age: $\beta = -0.00612$, $p = 2.93e-13$,
452 $q = 7.33e-12$; Neuron_Interneuron vs. Age: $\beta = -0.00591$, $p = 2.10e-10$, $q = 3.15e-09$). A similar decrease in
453 predicted neuronal content was seen in all three of the smaller human post-mortem datasets (Neuron_All
454 vs. Age (meta-analysis: $b = -0.00415$, $p = 1.57e-03$, $q = 2.02e-02$): Lanz 2014: $\beta = -0.00722$, $p = 0.0432$,
455 Neuron_Interneuron vs. Age (meta-analysis: $b = -0.00335$, $p = 2.91e-06$, $q = 6.56e-05$): Narayan et al. 2008:
456 $\beta = -0.00494$, $p = 0.0173$, Barnes et al. 2011: $\beta = -0.00506$, $p = 0.0172$, Neuron_Projection vs. Age (meta-
457 analysis: $b = -0.00449$, $p = 1.61e-06$, $q = 4.83e-05$): Lanz 2014: $\beta = -0.0103$, $p = 0.000497$, Narayan et al.
458 2008: $\beta = -0.00763$, $p = 0.00386$). This result fits with known decreases in gray matter density in the frontal
459 cortex in aging humans (48), as well as age-related sub-region specific decreases in frontal neuron

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460 numbers in primates (50) and rats (51). There was also a consistent decrease in immature
461 oligodendrocytes in relationship to age across datasets (Oligodendrocyte_Immature vs. Age (meta-
462 analysis: $b=-0.00514$, $p=5.98e-11$, $q=2.69e-09$): Pritzker: $\beta=-0.00432$, $p=0.000354$, Narayan et al. 2008:
463 $\beta=-0.00721$, $p=5.73e-05$, Lanz 2014: $\beta=-0.00913$, $p=1.85e-05$; CMCC: $\beta=-0.00621$, $p=3.32e-25$,
464 $q=2.49e-23$), which seems intuitive, but actually contradicts animal studies on the topic (52). Since the
465 validation of the Oligodendrocyte_Immature index was relatively weak, this result should perhaps be
466 considered with caution.

467 Other non-canonical relationships between subject variables and predicted cell content can be found
468 in the tables in **Figure 6**. In some datasets, there appears to be an increase in oligodendrocyte index with
469 age (Oligodendrocyte vs. Age (meta-analysis: $b=0.00343$, $p=2.74e-03$, $q=2.74e-02$): Narayan et al. 2008,
470 $\beta=0.00957$, $p=0.00349$) which, at initial face value, seems to contrast with well-replicated observations
471 that frontal white matter decreases with age in human imaging studies (48,53,54). However, it is worth
472 noting that several histological studies in aging primates suggest that brain regions that are experiencing
473 demyelination with age actually show an *increasing* number of oligodendrocytes, which is thought to be
474 driven by the need for repair (52,55).

475 Another prominent unexpected effect was a large decrease in the oligodendrocyte index with longer
476 post-mortem interval (Oligodendrocyte vs. PMI (meta-analysis: $b=-0.00764$, $p=2.23e-05$, $4.02e-04$):
477 Pritzker: $\beta=-0.00749$, $p=0.000474$, Lanz 2014: $\beta=-0.0318$, $p=0.000749$; CMC: $\beta=-0.00759$, $p=4.70e-05$,
478 $q=3.36e-04$). Upon further investigation, we found a publication documenting a 52% decrease in the
479 fractional anisotropy of white matter with 24 hrs post-mortem interval as detected by neuroimaging (56),
480 but to our knowledge the topic is otherwise not well studied. These changes were paralleled by a decrease
481 in endothelial cells (CMC: $\beta=-0.00542$, $p=1.32e-04$, $q=8.60e-04$) and microglia (CMC: $\beta=-0.00710$,
482 $p=5.15e-04$, $q=2.76e-03$) and relative increase in immature oligodendrocytes (Oligodendrocyte_Immature
483 vs. PMI (meta-analysis: $b=0.00353$, $p=4.81e-03$, $q=4.33e-02$): Pritzker: $\beta=0.00635$, $p=0.000683$) and
484 neurons (Neuron_All vs. PMI: Pritzker: $\beta=0.006997$, $p=0.000982$; CMC: $\beta=0.00386$, $p=0.0110$,
485 $q=0.0388$; Neuron_Projection vs. PMI (meta-analysis: $b=0.00456$, $p=2.28e-03$, $q=2.56e-02$): Pritzker:

486 $\beta = 0.00708$, $p = 1.64 \times 10^{-4}$; CMC: $\beta = 0.00331$, $p = 0.00197$, $q = 0.00924$). This result could arise from the
487 zero-sum nature of microarray analysis: due to the use of a standardized dissection size, RNA
488 concentration, and data normalization, if there are large decreases in gene expression for one common
489 variety of cell type in relationship to post-mortem interval (oligodendrocytes), then gene expression
490 related to other cell types may appear increase.

491 Overall, these results indicate that statistical predictions of the cell content of samples effectively
492 capture many known biological changes in cell type balance, and imply that within both chronic (age) and
493 acute conditions (agonal, PMI, pH) there is substantial influences upon the relative representation of
494 different cell types. Thus, when interpreting microarray data, it is as important to consider the chronic and
495 acute demographic factors at the population level as well as cellular functional regulation.

496

497 **3.3 Cell Type Balance Changes in Response to Psychiatric Diagnosis**

498 Of most interest to us were potential changes in cell type balance in relation to psychiatric illness. In
499 previous post-mortem morphometric studies, there was evidence of glial loss in the prefrontal cortex of
500 subjects with Major Depressive Disorder, Bipolar Disorder, and Schizophrenia (reviewed in (57)). This
501 decrease in glia, and particularly astrocytes, was replicated experimentally in animals exposed to chronic
502 stress (58), and when induced pharmacologically, was capable of driving animals into a depressive-like
503 condition (58). Replicating the results of (49), we observed a moderate decrease in astrocyte index in the
504 prefrontal cortex of subjects with Major Depressive Disorder (meta-analysis: $b = -0.132$, $p = 5.88 \times 10^{-3}$,
505 $q = 4.81 \times 10^{-2}$, Pritzker: $\beta = -0.133$, $p = 0.0118$, **Figure 6f**), but did not see similar changes in the brains of
506 subjects with Bipolar Disorder or Schizophrenia. We also observed a decrease in red blood cell index in
507 association with Schizophrenia (CMC: $\beta = -0.104$, $p = 0.0141$, $q = 0.0471$) which is tempting to ascribe to
508 reduced blood flow due to hypofrontality (59). This decrease in red blood cell content could also arise due
509 to psychiatric subjects having an increased probability of dying a violent death, but the effect remained
510 present when we controlled for exsanguination, therefore the effect is likely to be genuinely tied to the
511 illness itself.

512

513 3.4 Discriminating Between Changes in Cell Type Balance and Cell-Type Specific Function

514 Gray matter density has been shown to decrease in the frontal cortex in aging humans (48), and
515 frontal neuron numbers decrease in specific subregions in aging primates (50) and rats (51). However,
516 many scientists would argue that age-related decreases in gray matter are primarily driven by synaptic
517 atrophy instead of decreased cell number (60). This raised the question of whether the decline that we saw
518 in neuronal cell indices with age was being largely driven by the enrichment of genes related to synaptic
519 function in the index. More generally, it raised the question of how well cell type indices could
520 discriminate changes in cell number from changes in cell-type function.

521 We examined this question using two methods. First, we specifically examined the relationship
522 between age and the functional annotation for genes found in the Neuron_All index in more depth. To do
523 this, we evaluated the relationship between age and gene expression in the Pritzker dataset while
524 controlling for likely confounds using the signal data for all probesets in the dataset:

525 *Equation 4:*

$$526 \text{ Gene Expression (Probeset Signal) = } \\ 527 \beta_0 + \beta_1 * (\text{Diagnosis}) + \beta_2 * (\text{Brain pH}) + \beta_3 * (\text{Agonal Factor}) + \beta_4 * (\text{PMI}) + \beta_5 * (\text{Age}) + \beta_6 * (\text{Sex}) + \varepsilon \\ 528$$

529 We used “DAVID: Functional Annotation Tool” ([//david.ncifcrf.gov/summary.jsp](http://david.ncifcrf.gov/summary.jsp), (61,62) to
530 identify the functional clusters that were overrepresented by the genes included in our neuronal cell type
531 indices (using the full HT-U133A chip as background), and then determined the average effect of age
532 (beta) for the genes included in each of the 240 functional clusters (**Suppl. Table 5**). The vast majority of
533 these functional clusters showed a negative relationship with age on average (**Suppl. Figure 13**).
534 However, these functional clusters overrepresented dendritic/axonal related functions, so in a manner that
535 was blind to the results, we identified 29 functional clusters that were clearly related to dendritic/axonal
536 functions and 41 functional clusters that seemed distinctly unrelated to dendritic/axonal functions (**Suppl.**
537 **Table 5**). Using this approach, we found that transcripts from both classifications of functional clusters
538 showed an average decrease in expression with age (dendritic/axonal: $T(28) = -4.5612$, $p = 9.197e-05$, non-

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539 dendritic/axonal: $T(40)=-2.7566$, $p=0.008756$), but the decrease was larger for transcripts associated with
540 dendritic/axonal-related functions ($T(50.082)=2.3385$, $p=0.02339$, **Suppl. Figure 23**). Based on this
541 analysis, we conclude that synaptic atrophy could be partially driving age-related effects on neuronal cell
542 type indices in the human prefrontal cortex dataset but are unlikely to fully explain the relationship.

543 Next, we decided to make the process of differentiating between altered cell type-specific functions
544 and relative cell type balance more efficient. We used our cell type specific gene lists to construct gene
545 sets in a file format (.gmt) compatible with the popular tool Gene Set Enrichment Analysis (63,64) and
546 combined them with two other commonly-used gene set collections from the molecular signatures
547 database (MSigDB: <http://software.broadinstitute.org/gsea/msigdb/index.jsp>, downloaded 09/2017, “C2:
548 Curated Gene Sets” and “C5: GO Gene Sets”, **Suppl. Table 6**). Then we tested the utility of
549 incorporating our new gene sets into GSEA (fGSEA: (65)) using the ranked results (betas) for the
550 relationship between each subject variable (**Equation 4**) and each probeset in the Pritzker dataset. Using
551 this method, we could compare the enrichment of the effects of subject variables within gene sets defined
552 by brain cell type to the enrichment seen within gene sets for other functional categories. In general, we
553 found that gene sets for brain cell types tended to be the top result (most extreme normalized enrichment
554 score, NES) for each of the subject variables that showed a strong relationship with cell type in our
555 previous analyses (Agonal Factor vs. “Neuron_All_Cahoy_JNeuro_2008”: NES=-2.46, $p=0.00098$, $q=$
556 0.012 , Brain pH vs. “Astrocyte_All_Cahoy_JNeuro_2008”: NES=-2.48, $p=0.0011$, $q=0.014$, MDD vs.
557 “Astrocyte_All_Cahoy_JNeuro_2008”: NES= -2.60, $p=0.0010$, $q=0.017$, PMI vs.
558 “GO_OLIGODENDROCYTE_DIFFERENTIATION”: NES=-2.42, $p=0.00078$, $q=0.027$; **Suppl. Table**
559 **7**). Similarly, the relationship between the effects of age and neuron-specific gene expression was ranked
560 #4, following the gene sets “GO_SYNAPTIC_SIGNALING”,
561 “REACTOME_TRANSMISSION_ACROSS_CHEMICAL_SYNAPSES”,
562 “REACTOME_OPIOID_SIGNALLING”, but each of them was assigned a similar p-value ($p=0.001$) and
563 adjusted p-value ($q=0.036$). We conclude that it is important to consider cell type-specific expression
564 during the analysis of macro-dissected brain microarray data above and beyond the consideration of

565 specific functional pathways, and have submitted our .gmt files to the Broad Institute for potential
566 addition to their curated gene sets in MSigDB to promote this form of analysis.

567

568 **3.5 Including Cell Content Predictions in the Analysis of Microarray Data Improves Model Fit And** 569 **Enhances the Detection of Diagnosis-Related Genes in Some Datasets**

570 Over the years, many researchers have been concerned that transcriptomic and genomic analyses
571 of psychiatric disease often produce non-replicable or contradictory results and, perhaps more
572 disturbingly, are typically unable to replicate well-documented effects detected by other methods. We
573 posited that this lack of sensitivity and replicability might be partially due to cell type variability in the
574 samples, especially since such a large percentage of the principal components of variation in our samples
575 were explained by neuron to glia ratio. Within the Pritzker dataset, we were particularly interested in
576 controlling for cell type variability, because there were indications that dissection might have differed
577 between technical batches that were unevenly distributed across diagnosis categories (**Figure 8, Suppl.**
578 **Figure 10**). There was a similarly uneven distribution of dissection methods across diagnosis categories
579 within the large CMC RNA-Seq dataset. In this dataset, the majority of the bipolar samples (75%) were
580 collected by a brain bank that performed gray matter only dissections (PITT), whereas the control and
581 schizophrenia samples were more evenly distributed across all three institutions (40).

582 We hypothesized that controlling for cell type while performing differential expression analyses
583 in these datasets would improve our ability to detect previously-documented psychiatric effects on gene
584 expression, especially psychiatric effects on gene expression that were previously-identified within
585 individual cells, since these effects on gene expression should not be mediated by psychiatric changes in
586 overall cell type balance. To test the hypothesis, we first compiled a list of 130 strong, previously-
587 documented relationships between Schizophrenia or Psychosis and gene expression in particular cell
588 types in the human cortex, as detected by in situ hybridization or immunocytochemistry (reviewed further
589 in (19); GAD1: (66–68); RELN:(66); SST: (69), SLC6A1 (GAT1): (70), PVALB:(67), *suicide*: HTR2A
590 (71)), or by single-cell type laser capture microscopy (**Figure 7, Suppl. Table 8** (1,72,73)).

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Validation Datasets:	# of Genes	Method:	Brain Bank:	# of Subjects	Brain Region	Co-Variates: Controlled?	Co-Variates: Balanced?	Statistical Stringency
Schizophrenia Effects in Particular Cortical Cell Types:								
Reviewed in Lewis & Sweet (2009)	7	ICC/in situ hybridization	Variable, often PITT	Variable	Prefrontal cortex	Variable	Variable	Variable
Arion et al. (2015)	41	LCM-Microarray: Pyramidal Neurons (Layers 3 & 5)	PITT	72	BA9	Direction of effect evaluated, but covariates not included in final model.	Sex, Age, PMI, pH, RIN, tissue storage time, race	Top 40 (FDR<0.1 in both layers, Table 2A), Top 2 in Table 2B, FDR<10E-17 for Layer5)
Pietersen et al. (2014)	47	LCM-Microarray: PVALB Interneurons	HBTRC (MacLean)	16	BA42	Batch. Considered effects of Sex, Age, PMI but not included in final model.	Sex, Age, PMI, pH not significantly different	Top 47 (FDR<0.01, FC>2, Table 3)
Mauney et al. (2015)	35	LCM-Microarray: Oligodendrocyte Precursors	HBTRC (MacLean)	18	BA9	None	Sex, Age, PMI, pH not reported	Top 35 (FDR<0.001, Table S2)
Psychiatric Effects in Macro-dissected Prefrontal Cortex:								
Mistry et al. (2013)	126	Meta-analysis of microarray data: Schizophrenia effects	Stanley Foundation, HBTRC (MacLean), PITT, CCHPC, MSSM, MHRI	306	BA9, BA10, BA46	Model selection procedure included Batch, Age, pH, Study	Sex, PMI	FDR<0.1 (Table S2)
Choi et al. (2011)	367	Meta-analysis of microarray data: Bipolar effects	Stanley Foundation	83	BA46 (grey matter only)	Batch (Scan Date), pH, Psychosis, Medication at TOD	Age, BMI, PMI not reported	FDR<0.05, FC>1.3 (Table S1)

591

592 **Figure 7. Gene lists used to assess whether controlling for cell type while performing differential**
 593 **expression analyses enhances the detection of previously-documented psychiatric effects on cortical**
 594 **gene expression. These lists include genes with documented relationships to psychiatric illness in either**
 595 **1) particular cortical cell types or 2) macro-dissected cortex. The full lists can be found in *Suppl. Table***
 596 **8. Abbreviations: LCM: Laser Capture Microscopy, PVALB: Parvalbumin, BA: Brodmann's Area, PMI:**
 597 **Post-mortem interval, FDR: False detection ratio (or q-value), Brain Banks: PITT (University of**
 598 **Pittsburgh), HBTRC (Harvard Brain Resource Tissue Center), CCHPC (Charing Cross Hospital**
 599 **Prospective Collection), MSSM (Mount Sinai Icahn School of Medicine), MHRI (Mental Health Research**
 600 **Institute Australia).**

601 As a comparison, we also considered lists of transcripts strongly-associated with Schizophrenia
 602 (74) and Bipolar Disorder (75) in meta-analyses of microarray data derived from human frontal cortical
 603 tissue (**Figure 7**). The effects of psychiatric illness on the expression of these transcripts could be
 604 mediated by either psychiatric effects on cell type balance or by effects within individual cells. Therefore,
 605 controlling for cell type balance while performing differential expression analyses could detract from the
 606 detection of some psychiatric effects, but perhaps also enhance the detection of other psychiatric effects
 607 by controlling for large, confounding sources of noise (*e.g.*, dissection variability).

608 Next, we examined our ability to detect these previously-documented psychiatric effects using
 609 regression models of increasing complexity (**Figure 8 B**), including a simple base model containing just

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610 the variable of interest (“Model 1”), a standard model controlling for traditional co-variates (“Model 2”),
611 and a model controlling for traditional co-variates as well as each of the cell type indices (“Model 5”:
612 **Equation 5**). We also used two reduced models that only included the most prevalent cell types
613 (Astrocyte, Microglia, Oligodendrocyte, Neuron_Interneuron, Neuron_Projection; (21)) to avoid issues
614 with multicollinearity. The first of these models included traditional co-variates as well (“Model 4”),
615 whereas the second model excluded them (“Model 3”).

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A. Diagnosis Effects May Be Partially Confounded By Dissection Variability

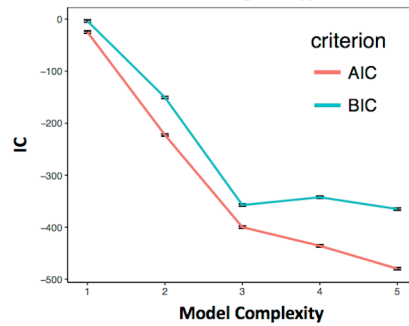
Pritzker:	
Samples	Batch*
Control	All Batches
BP, MDD	1-4, 9-13
Schiz	5-8, 13-15

CMC:	
Samples	Institution*
Controls	All (PITT, MSSM, PENN)
BP	PITT, MSSM
Schiz	All (PITT, MSSM, PENN)

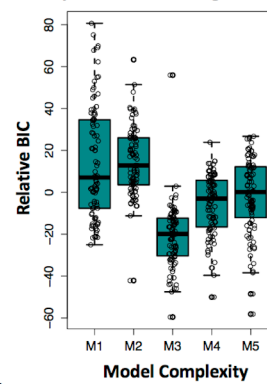
*Batches partially defined by subject cohort *PITT was a grey matter only dissection

B. Model Complexity	
M1	Base Model: Diagnosis Only
M2	Standard Model: Diagnosis + Traditional Co-variables
M3	Diagnosis + Most Prevalent Cell Types
M4	Diagnosis + Traditional Co-variables + Most Prevalent Cell Types
M5	Diagnosis + Traditional Co-variables + All Cell Types

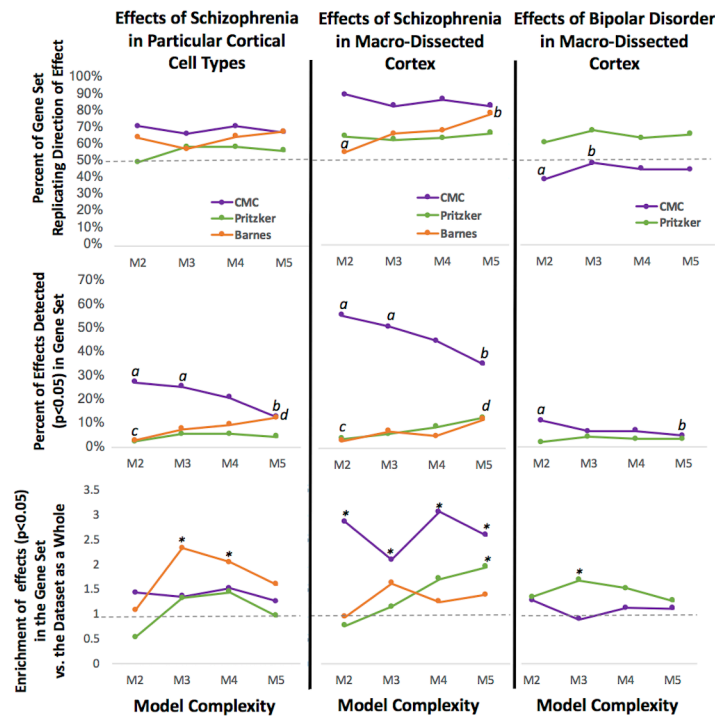
C. All Expressed Genes: Model Fit Improves After Adding Cell Type



D. Genes with Previously-Identified Psychiatric Effects: Model Fit Improves After Adding Cell Type



E. Controlling for Cell Type Variability Enhances Detection of Psychiatric Effects in Some Datasets



617 **Figure 8. Including Cell Content Predictions in the Analysis of Microarray Data Improves Model Fit**
618 **and Enhances the Detection of Previously-Identified Diagnosis-Related Genes in Some Datasets. A.**
619 *Diagnosis effects were likely to be partially confounded by dissection variability within the Pritzker and*
620 *CMC datasets. B: We examined a series of differential expression models of increasing complexity,*
621 *including a base model (M1), a standard model (M2), and three models that included cell type co-*
622 *variates (M3-M5). C. Model fit improved with the addition of cell type (M1/M2 vs. M3-M5) when*
623 *examining all expressed genes in the dataset (example from CMC: points= AVE +/-SE). D. Model fit*
624 *improved with the addition of cell type (M1/M2 vs. M3-M5) when examining genes with previously-*
625 *documented relationships with psychiatric illness in particular cell types (example from Pritzker: BIC*
626 *values for all models for each gene were centered prior to analysis). Boxplots represent the median and*
627 *interquartile range, with whiskers illustrating either the full range of the data or 1.5x the interquartile*
628 *range. E. Evaluating the replication of previously-detected psychiatric effects (Figure 7) in three datasets*
629 *(Pritzker, CMC, and Barnes) using a standard differential expression model (M2) vs. models that include*
630 *cell type co-variates (M3-5). Top graphs: The percentage of genes (y-axis: 0-1) replicating the direction*
631 *of previously-documented psychiatric effects on cortical gene expression sometimes increases with the*
632 *addition of cell type to the model (Barnes (effects of Schiz): M2 vs. M5, CMC (effects of Bipolar*
633 *Disorder): M2 vs. M3). Middle graphs: The detection of previously-identified psychiatric effects on gene*
634 *expression (p<0.05 & replicated direction of effect) increases with the addition of cell type to the model*
635 *in some datasets (Barnes: M2 vs. M5, Pritzker: M2 vs. M5) but decreases in others (CMC: M2 vs. M5,*
636 *M3 vs. M5). Bottom graphs: In some datasets we see an enrichment of psychiatric effects (p<0.05) in*
637 *previously-identified psychiatric gene sets only after controlling for cell type (Barnes: M3, M4, Pritzker:*
638 *M5, M3). For the CMC dataset, we see an enrichment using all models. The full results for all models can*
639 *be found in Suppl. Table 9, Suppl. Table 10, Suppl. Table 11, Suppl. Table 12, and Suppl. Table 13.*

640

641 **Equation 5: A model of gene expression for each dataset, colored to illustrate the subcomponents**
642 **evaluated during our model comparison (#M1-M5). The base model (intercept and variable of interest)**
643 **is presented in green, the typical subject variable covariates included in a standard model are blue, the**
644 **cell type indices for the most prevalent cell types are colored red, and the remaining cell type indices are**
645 **in purple. Model components unique to each dataset are underlined.**

646 **The Pritzker microarray dataset:**

647 Gene Expression (Probeset Signal) =
648 $\beta_0 + \beta_1 * (\text{The variable of interest: Diagnosis})$
649 $+ \beta_2 * (\text{Brain pH}) + \beta_3 * (\text{PMI}) + \beta_4 * (\text{Age}) + \beta_5 * (\text{Sex}) + \beta_6 * (\text{Agonal Factor}) +$
650 $+ \beta_7 * (\text{Astrocyte}) + \beta_8 * (\text{Oligodendrocyte}) + \beta_9 * (\text{Microglia}) + \beta_{10} * (\text{Interneuron}) + \beta_{11} * (\text{ProjectionNeuron})$
651 $+ \beta_{12} * (\text{Endothelial}) + \beta_{13} * (\text{Neuron_All}) + \beta_{14} * (\text{Oligodendrocyte_Immature}) + \beta_{15} * (\text{Mural}) + \beta_{16} * (\text{RBC}) + \epsilon$

652 **The CMC RNA-Seq dataset:**

653 Gene Expression (Probeset Signal) =
654 $\beta_0 + \beta_1 * (\text{The variable of interest: Diagnosis})$
655 $+ \beta_2 * (\text{Brain pH}) + \beta_3 * (\text{PMI}) + \beta_4 * (\text{Age}) + \beta_5 * (\text{Sex}) + \beta_6 * (\text{RIN}) + \beta_7 * (\text{Institution}) + \beta_8 * (\text{CauseOfDeath}) +$
656 $+ \beta_9 * (\text{Astrocyte}) + \beta_{10} * (\text{Oligodendrocyte}) + \beta_{11} * (\text{Microglia}) + \beta_{12} * (\text{Interneuron}) + \beta_{13} * (\text{ProjectionNeuron})$
657 $+ \beta_{14} * (\text{Endothelial}) + \beta_{15} * (\text{Neuron_All}) + \beta_{16} * (\text{Oligodendrocyte_Immature}) + \beta_{17} * (\text{Mural}) + \beta_{18} * (\text{RBC}) + \epsilon$
658

659 **The smaller microarray datasets (GSE53987, GSE21935, GSE21138):**

660 Gene Expression (Probeset Signal) =
661 $\beta_0 + \beta_1 * (\text{The variable of interest: Diagnosis})$
662 $+ \beta_2 * (\text{Brain pH}) + \beta_3 * (\text{PMI}) + \beta_4 * (\text{Age}) + \beta_5 * (\text{Sex}) + \beta_6 * (\text{RNA Degradaion}) +$
663 $+ \beta_7 * (\text{Astrocyte}) + \beta_8 * (\text{Oligodendrocyte}) + \beta_9 * (\text{Microglia}) + \beta_{10} * (\text{Interneuron}) + \beta_{11} * (\text{ProjectionNeuron})$

664 $+\beta_{12}*(\text{Endothelial})+\beta_{13}*(\text{Neuron_All})+\beta_{14}*(\text{Oligodendrocyte_Immature})+\beta_{15}*(\text{Mural})+\beta_{16}*(\text{RBC})+ \epsilon$

665

666 We found that including predictions of cell type balance in our models assessing the effect of
667 diagnosis on gene expression dramatically improved model fit as assessed by Akaike's Information
668 Criterion (AIC) or Bayesian Information Criterion (BIC) (**Figure 8**). These improvements were largest
669 with the addition of the five most prevalent cell types to the model (M3, M4); the addition of less
670 common cell types produced smaller gains (M5). These improvements were clear whether we considered
671 the average model fit for all expressed genes (*e.g.*, **Figure 8B**) or just genes with previously-identified
672 psychiatric effects (*e.g.*, **Figure 8C**).

673 However, models that included cell type were not necessarily superior at replicating previously-
674 observed psychiatric effects on gene expression (**Figure 7**), even when examining psychiatric effects that
675 were likely to be independent of changes in cell type balance. For each model, we quantified the
676 percentage of genes replicating the previously-observed direction of effect in relationship to psychiatric
677 illness, as well as the percentage of genes that replicated the effect using a common threshold for
678 detection ($p<0.05$). Finally, we also looked at the enrichment of psychiatric effects ($p<0.05$) in each of the
679 previously-documented psychiatric gene sets in comparison to the other genes in our datasets. For this
680 analysis, to improve comparisons across datasets, we defined the statistical background for enrichment
681 using genes universally represented in all three datasets (Pritzker, CMC, Barnes).

682 In general, we found that the two datasets that had the most variability in gene expression related
683 to cell type (Pritzker, Barnes: **Results 3.1**) were more likely to replicate previously-documented
684 psychiatric effects on gene expression when the differential expression model included cell type
685 covariates. For example, in the Barnes dataset, adding cell type co-variates to the model increased our
686 ability to detect effects of Schizophrenia that had been previously documented within particular cell types
687 or macro-dissected tissue (**Figure 8E**, Fisher's exact test: M2 vs. M5, $p<0.05$ in both gene sets).
688 Similarly, adding cell type co-variates to the model allowed us to see a significant enrichment of
689 Schizophrenia effects ($p<0.05$) in genes with previously-documented psychiatric effects in particular cell

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690 types (Fisher's exact test $p < 0.05$: M3 & M4). In the Pritzker dataset, we saw that adding cell type co-
691 variates to the model increased our ability to detect previously-documented effects of Schizophrenia in
692 macrodissected tissue (M2 vs. M5: $p < 0.05$). Likewise, adding cell type co-variates to the model allowed
693 us to see a significant enrichment of Schizophrenia and Bipolar effects ($p < 0.05$) in genes with previously-
694 documented psychiatric effects in macro-dissected tissue (Fisher's exact test $p < 0.05$: Schizophrenia: M5,
695 Bipolar: M3). This mirrored the results of another analysis that we had conducted suggesting that
696 controlling for cell type increased the overlap between the top diagnosis results in the Pritzker dataset and
697 previous findings in the literature as a whole (**Suppl. Figure 24, Suppl. Figure 25**).

698 In the large CMC RNA-Seq dataset, the rate of replication of previously-documented effects of
699 Schizophrenia was already quite high using a standard differential expression model containing traditional
700 co-variates (M2). Using a standard model, we could detect 27% of the previously-documented effects in
701 cortical cell types and 55% of the previously-documented effects in macro-dissected tissue (with a
702 replicated direction of effect and $p < 0.05$). However, in contrast to what we had observed in the Pritzker
703 and Barnes datasets, controlling for cell type seemed to actually *diminish* the ability to detect effects of
704 Schizophrenia that had been previously-observed within particular cell types or macrodissected tissue in a
705 manner that scaled with the number of co-variates included in the model (M2 or M3 vs. M5: $p < 0.05$ for
706 both gene sets), despite improvements in model fit parameters and a lack of significant relationship
707 between Schizophrenia and any of the prevalent cell types (**Section 3.3**). Including cell type co-variates in
708 the model did not improve our ability to observe a significant enrichment of Schizophrenia effects in
709 genes with previously-documented psychiatric effects in macro-dissected tissue – this enrichment was
710 present in the results from all differential expression models (Fisher's exact test $p < 0.05$: M2-M5). In
711 contrast, controlling for cell type slightly improved the replication of the direction of previously-
712 documented Bipolar Disorder effects (Fisher's exact test: M2 vs. M3: $p < 0.05$) in a manner that would
713 seem appropriate due to the highly uneven distribution of bipolar samples across institutions and
714 dissection methods, but even after this improvement the rate of replication was still no better than chance
715 (48%), and, counterintuitively, the ability to successfully detect those effects still diminished in a manner

716 that seemed to scale with the number of co-variates included in the model (Fisher's exact test: M2 vs. M5,
717 $p < 0.05$). In a preliminary analysis of the two smaller human microarray datasets that were derived from
718 gray-matter only dissections (GSE53987, GSE21138), the addition of cell type co-variates to differential
719 expression models clearly diminished both the percentage of genes replicating the previously-documented
720 direction of effect of Schizophrenia in particular cell types (Fisher's exact test: Narayan et al.: M2 vs. M4
721 or M5: $p < 0.05$, Lanz et al.: M2 vs. M4 or M5) and the ability to successfully detect previously-
722 documented effects (Fisher's exact test: Narayan et al.: M2 vs. M4 or M5: $p < 0.05$).

723 Therefore, we conclude that the addition of cell type covariates to differential expression models
724 is only recommended when there is a particularly large amount of variability in the dataset associated
725 with cell type balance. For public use we have released the full results for each dataset analyzed using the
726 different models discussed above (**Suppl. Table 9, Suppl. Table 10, Suppl. Table 11, Suppl. Table 12,**
727 **and Suppl. Table 13**).

728

729 **4. Discussion**

730 In this manuscript, we have demonstrated that the statistical cell type index is a relatively simple
731 manner of interrogating cell-type specific expression in transcriptomic datasets from macro-dissected
732 human brain tissue. We find that statistical estimations of cell type balance almost fully account for the
733 top principal components of variation in microarray data derived from macro-dissected brain tissue
734 samples, far surpassing the effects of traditional subject variables (post-mortem interval, hypoxia, age,
735 gender). Indeed, our results suggest that many variables of medical interest are themselves accompanied
736 by strong changes in cell type composition in naturally-observed human brains. We find that within both
737 chronic (age, sex, diagnosis) and acute conditions (agonal, PMI, pH) there are substantial changes in the
738 relative representation of different cell types. Thus, accounting for demography at the cellular population
739 level is as important for the interpretation of microarray data as cell-level functional regulation. This form
740 of data deconvolution was also useful for identifying the subtler effects of psychiatric illness within our

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741 samples, divulging the decrease in astrocytes that is known to occur in Major Depressive Disorder and the
742 decrease in red blood cell content in the frontal cortex in Schizophrenia, resembling known fMRI
743 hypofrontality. This form of data deconvolution may also aid in the detection of psychiatric effects while
744 conducting differential expression analyses in datasets that have highly-variable cell content.

745 These results touch upon the fundamental question as to whether organ-level function responds to
746 challenge by changing the biological states of individual cells (Lamarckian) or the life and death of
747 different cell populations (Darwinian). To reach such a sweeping perspective in human brain tissue using
748 classic cell biology methods would require epic efforts in labeling, cell sorting, and counting. We have
749 demonstrated that you can approximate this vantage point using an elegant, supervised signal
750 decomposition exploiting increasingly available genomic data. However, it should be noted that, similar
751 to other forms of functional annotation, cell type indices are best treated as a hypothesis-generation tool
752 instead of a final conclusion regarding tissue cell content. We have demonstrated the utility of cell type
753 indices for detecting large-scale alterations in cell content in relationship with known subject variables in
754 post-mortem tissue. We have not tested the sensitivity of the technique for detecting smaller effects or the
755 validity under all circumstances or non-cortical tissue types. Likewise, while using this technique it is
756 impossible to distinguish between alterations in cell type balance and cell-type specific transcriptional
757 activity: when a sample shows a higher value of a particular cell type index, it could have a larger number
758 of such cells, or each cell could have produced more of its unique group of transcripts, via a larger cell
759 body, slower mRNA degradation, or an overall change in transcription rate. In this regard, the index that
760 we calculate does not have a specific interpretation; rather it is a holistic property of the cell populations,
761 the “neuron-ness” or “microglia-ness” of the sample. Such an abstract index represents the ecological
762 shifts inferred from the pooled transcriptome. That said, unlike principal component scores or other
763 associated techniques of removing unwanted variation from genomic data, our cell type indices do have
764 real biological meaning - they can be interpreted in a known system of cell type taxonomy. When single-
765 cell genomic data uncovers new cell types (e.g., the Allen Brain Atlas cellular taxonomy initiative (76))
766 or meta-analyses refine the list of genes defined as having cell-type specific expression (e.g., (77)), our

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767 indices will surely evolve with these new classification frameworks, but the power of the approach will
768 remain, in that we can disentangle the intrinsic changes of individual genes from the population-level
769 shifts of major cell types.

770 We found that many variables of medical interest are accompanied by strong changes in cell type
771 composition in naturally-observed human brains. One result from this analysis seems particularly worth
772 discussing in greater depth. It has been acknowledged for a long time that exposure to a hypoxic
773 environment prior to death has a huge impact on gene expression in human post-mortem brains (e.g.,
774 (32,33,78–80)). This impact on gene expression is so large that up until recently the primary principal
775 component of variation (PC1) in our Pritzker data was assumed to represent the degree of hypoxia, and
776 was sometimes even systematically removed before performing diagnosis-related analyses (e.g., (35)).
777 The strong relationship between hypoxia and gene expression in human post-mortem samples was
778 hypothesized to be partially mediated by neuronal necrosis (81) and lactic acidosis (79). However, the
779 magnitude of the effect of hypoxia on gene expression was still puzzling, especially when compared to
780 the much more moderate effects of post-mortem interval, even when the intervals ranged from 8-40+ hrs.
781 Our current analysis provides an explanation for this discrepancy, since it is clear from our results that the
782 brains of our subjects are *actively compensating* for a hypoxic environment prior to death by altering the
783 balance or overall transcriptional activity of support cells and neurons. The differential effects of hypoxia
784 on neurons and glial cells have been studied since the 1960's (82), but to our knowledge this is the first
785 time that anyone has related the large effects of hypoxia in post-mortem transcriptomic data to a
786 corresponding upregulation in the transcriptional activity of vascular cell types (46).

787 This connection is important for understanding why results associating gene expression and
788 psychiatric illness in human post-mortem tissue sometimes do not replicate. If a study contains mostly
789 tissue from individuals who experienced greater hypoxia before death (e.g., hospital care with artificial
790 respiration or drug overdose followed by coma), then the evaluation of the effect of neuropsychiatric
791 illness is likely to inadvertently focus on differential expression in support cell types (astrocytes,
792 endothelial cells), whereas a study that mostly contains tissue from individuals who died a fast death (e.g.,

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793 car accident or myocardial infarction) will emphasize the effects of neuropsychiatric illness in neurons.
794 That said, although both indicators of perimortem hypoxia (agonal factor and acidosis (pH)) showed
795 similar strong relationships with cell type balance, we do recommend some caution when interpreting the
796 relationship between pH and cell type in tissue from subjects with psychiatric disorders, as pH can
797 indicate other biological changes besides hypoxia. For example, there are small consistent decreases in
798 pH associated with Bipolar Disorder even in live subjects (83–85) and metabolic changes associated with
799 pH are theorized to play an important role in Schizophrenia (80). Therefore, some of the relationship
800 between pH and cell type balance may be driven by a third variable (psychiatric illness or psychiatric
801 treatment). It is also possible that a change in the cell content of brain tissue could cause a change in pH
802 (86).

803 We found that including cell type indices as co-variates while running differential expression
804 analyses helped improve our ability to detect previously-documented relationships between psychiatric
805 illness and gene expression in datasets that were particularly affected by variability in cell type balance.
806 This improvement was not seen in datasets that were less affected by variability in cell type balance,
807 despite improvements in model fit and a lack of strong multicollinearity between diagnosis and the cell
808 type indices. This finding was initially surprising to us, but upon further consideration makes sense, as
809 the cell type indices are multi-parameter gene expression variables. Therefore, there is increased risk of
810 overfitting when modeling the data for any particular gene. We conclude that the addition of cell type
811 covariates to differential expression models is only recommended when there is a particularly large
812 amount of variability in the dataset associated with cell type balance, or when there is strong reason to
813 believe that technical variation associated with cell type (such as dissection) may be highly confounding
814 in the result. We strongly recommend that model selection while conducting differential expression
815 analyses should be considered carefully, and evaluated not only in terms of fit parameters but also validity
816 and interpretability.

817 Regarding the importance of model selection for interpretability, it is worth noting that an
818 important difference between our final analysis methods and those used by some previous researchers

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819 (*e.g.*, 10–12) was the lack of cell type interaction terms included in our models (*e.g.*, Diagnosis*Astrocyte
820 Index). Theoretically, the addition of cell type interaction terms should allow the researcher to statistically
821 interrogate cell-type differentiated diagnosis effects because samples that contain more of a particular cell
822 type should exhibit more of that cell type’s respective diagnosis effect. Versions of this form of analysis
823 have been successful in other investigations (*e.g.*, (11,12,87)) but we were not able to validate the method
824 using our database of previously-documented relationships with diagnosis in prefrontal cell types (**Figure**
825 **7**) and a variety of model specifications (*e.g.*, **Suppl. Figure 26**). Upon consideration, we realized that
826 these negative results were difficult to interpret because significant diagnosis*cell type interactions should
827 only become evident if the effect of diagnosis in a particular cell type is different from what is occurring
828 in all cell types on average. For genes with expression that is reasonably specific to a particular cell type
829 (*e.g.*, GAD1, PVALB), the overall average diagnosis effect may already largely reflect the effect within
830 that cell type and the respective interaction term will not be significantly different, even though the
831 disease effect is clearly tracking the balance of that cell population. In the end, we decided that the
832 addition of interaction terms to our models was not demonstrably worth the associated decrease in overall
833 model fit and statistical power.

834 Finally, our work drives home the fact that any comprehensive theory of psychiatric illness needs
835 to account for the dichotomy between the health of individual cells and that of their ecosystem. We found
836 that the functional changes accompanying psychiatric illness in the cortex occurred both at the level of
837 cell population shifts (decreased astrocytic presence and red blood cell count) and at the level of intrinsic
838 gene regulation not explained by population shifts. A similar conclusion regarding the importance of cell
839 type balance in association with psychiatric illness was recently drawn by our collaborators (*e.g.*,(88))
840 using a similar technique to analyze RNA-Seq data from the anterior cingulate cortex. In the future, we
841 plan to use our technique to re-analyze many of the other large microarray datasets existing within the
842 Pritzker Neuropsychiatric Consortium with the hope of gaining better insight into psychiatric disease
843 effects. This application of our technique seems particularly important in light of recent evidence linking
844 disrupted neuroimmunity (89) and neuroglia (*e.g.*, (49,58,90)) to psychiatric illness, as well as growing

845 evidence that growth factors with cell type specific effects play an important role in depressive illness and
846 emotional regulation (for a review see (23,91)).

847 In conclusion, we have found this method to be a valuable addition to traditional functional
848 ontology tools as a manner of improving the interpretation of transcriptomic results. The capability to
849 unravel alterations of cell type composition from modulation of cell state, even just probabilistically, is
850 inherently useful for understanding the higher-level function of the brain as emergent properties of brain
851 activity, such as emotion, cognition, memory, and addiction, usually involve ensembles of many cells.
852 Facilitating the interpretation of gene activity data in macro-dissected tissue in light of both processes
853 provides new opportunities to integrate results with findings from other approaches, such as
854 electrophysiology analysis of brain circuits, brain imaging, optogenetic manipulations, and naturally
855 occurring variation in response to injury and brain diseases.

856 For the benefit of other researchers, we have made our database of brain cell type specific genes
857 (<https://sites.google.com/a/umich.edu/megan-hastings-hagenauer/home/cell-type-analysis>) and R code for
858 conducting cell type analyses publicly available in the form of a downloadable R package
859 (<https://github.com/hagenaue/BrainInABlender>) and we are happy to assist researchers in their usage for
860 pursuing better insight into psychiatric illness and neurological disease.

861

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875

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1199 **7. Supporting Information**

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1202 **Supplementary Material for:**

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1204 **INFERENCE OF CELL TYPE COMPOSITION FROM HUMAN BRAIN TRANSCRIPTOMIC**

1205 **DATASETS ILLUMINATES THE EFFECTS OF AGE, MANNER OF DEATH, DISSECTION,**

1206 **AND PSYCHIATRIC DIAGNOSIS**

1207 *Megan Hastings Hagenauer, Ph.D.¹, Anton Schulmann, M.D.², Jun Z. Li, Ph.D.³, Marquis P. Vawter,
1208 Ph.D.⁴, David M. Walsh, Psy.D.⁴, Robert C. Thompson, Ph.D.¹, Courtney A. Turner, Ph.D.¹, William E.
1209 Bunney, M.D.⁴, Richard M. Myers, Ph.D.⁵, Jack D. Barchas, M.D.⁶, Alan F. Schatzberg, M.D.⁷, Stanley J.
1210 Watson, M.D., Ph.D.¹, Huda Akil, Ph.D.¹

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1214 **7.1 Detailed Methods and Results: Using Cell Type Specific Transcripts to Predict Relative Cell**

1215 **Content in Datasets from Purified Cells and Artificial Cell Mixtures**

1216 To validate our technique, we used the expression of the cell type specific transcripts included in
1217 our database to predict the relative balance of cell types in samples with known cell content (purified cells
1218 and artificial cell mixtures). To do this analysis, we used two RNA-Seq datasets: one derived from from
1219 purified cortical cell types in mice (n=17: two samples per purified cell type and 3 whole brain samples:
1220 <https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE52564>) (18), and one derived from 466 single-
1221 cells dissociated from freshly-resected human cortex
1222 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE67835>) (2). To estimate the limitations and
1223 noise inherent in our technique, we also constructed *in silico* mixtures of 100 cells with known
1224 percentages of each cell type by randomly sampling from each dataset (with replacement).

1225 The RNA-Seq data that we downloaded from GEO (Gene expression Omnibus) was already in
1226 the format of FPKM values (Fragments Per Kilobase of exon model per million mapped fragments) (18)
1227 or counts per gene (2). To stabilize the variance in the data, we used a log transformation (base 2), and
1228 then filtered out the data for any genes that completely lacked variation across samples (sd=0). Within the
1229 mouse dataset (18) this filtering decreased the dataset from 22462 genes to 17148 genes. Within the
1230 human dataset (2), this filtering decreased the dataset from 22085 genes to 21627 genes.

1231 Then, using the methods now found in the BrainInABlender package, we extracted the data for
1232 genes previously identified as having cell type specific expression in our curated database. Within the
1233 mouse dataset, there were data from 2513 genes that aligned with 2914 entries in our database of cell type
1234 specific transcripts (as matched by official gene symbol). Within the human dataset, there were data from
1235 2374 genes that aligned with 2882 entries in our database of cell type specific transcripts (as matched by
1236 gene symbol). We centered and scaled the expression levels for each gene across samples (mean=0, sd=1)
1237 to prevent genes with more variable signal from exerting disproportionate influence, and then, for each
1238 sample, averaged this value across the transcripts identified in each publication as specific to a particular
1239 cell type. This created 38 cell type signatures derived from the cell type specific genes identified by the

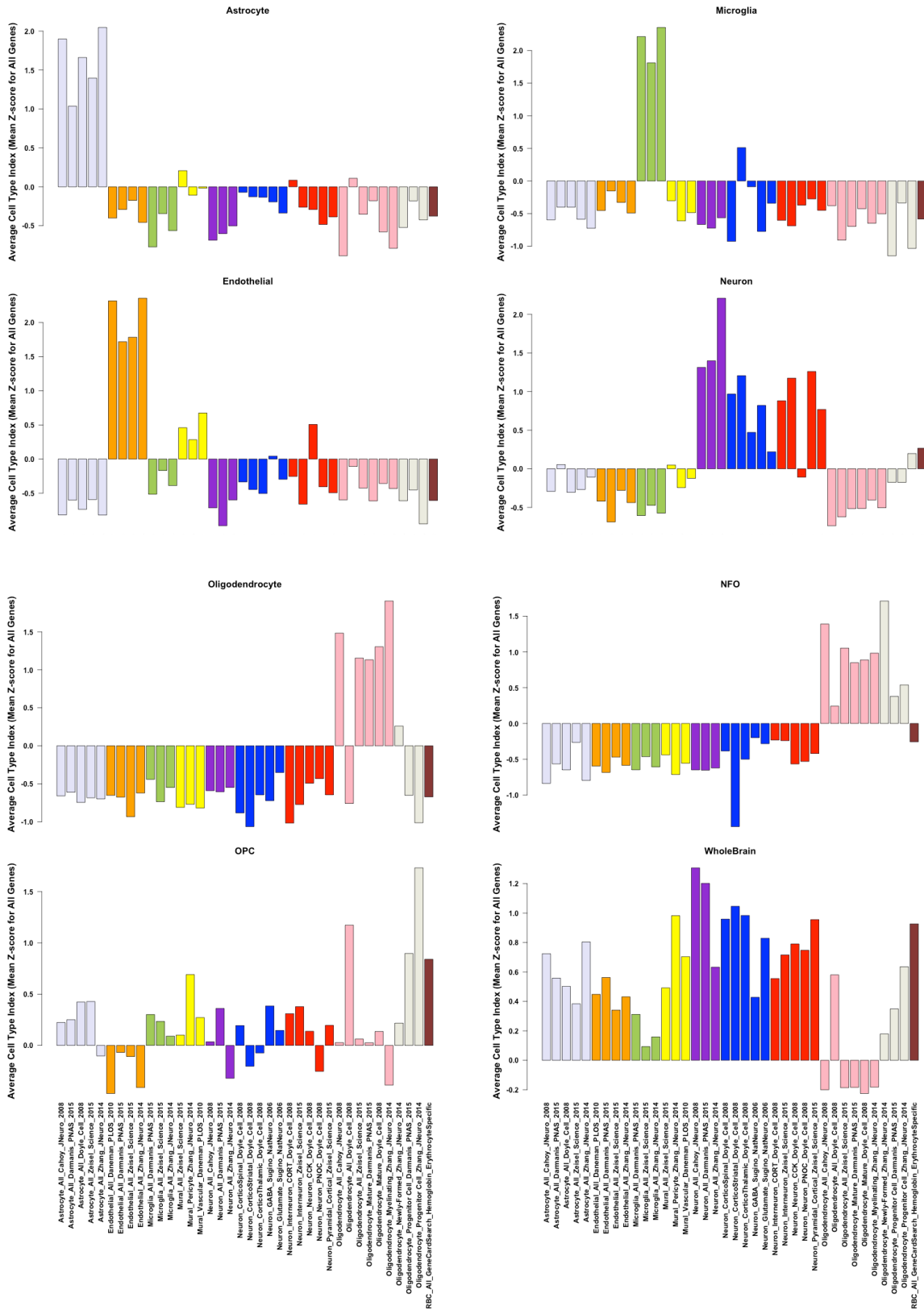
Running Head: PREDICTING CELL TYPE BALANCE

1240 eight publications ("*Cell Type Indices*"), each of which predicted the relative content for one of the 10
1241 primary cell types in our cortical samples. All of the R script documenting these analyses can be found at
1242 https://github.com/hagenaue/CellTypeAnalyses_Darmanis and
1243 https://github.com/hagenaue/CellTypeAnalyses_Zhang.

1244 We found that the statistical cell type indices easily predicted the cell type identities of the
1245 original samples (**Suppl. Figure 1, Suppl. Figure 2**). This was true regardless of the publication from
1246 which the cell type specific genes were derived: cell type specific gene lists derived from publications
1247 using different species (human vs. mouse), platforms (microarray vs. RNA-Seq), methodologies
1248 (florescent cell sorting vs. suspension), or statistical stringency all performed fairly equivalently, with
1249 some minor exception. Occassionally, we found that the cell type indices associated with cell type
1250 specific gene lists derived from TRAP methodology (15) did not properly predict the cell identity of the
1251 samples, and in general the cell type indices associated with immature oligodendrocytes were somewhat
1252 inconsistent. As would be expected, the cell type indices derived from cell type specific genes identified
1253 by the same publication that produced the test datasets (2,18) were (by definition) superb predictors of the
1254 sample cell identity in their own dataset, and were thus excluded from later validation analyses.

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1259 ***Suppl. Figure 1. The cell content predictions derived from cell type specific transcripts originating***
1260 ***from different publications successfully predict sample cell type in mouse purified cell type RNA-Seq***
1261 ***data. The sample cell type in a mouse purified cell type RNA-Seq dataset (18) was predicted equally well***
1262 ***by cell type indices derived from cell type specific transcripts originating from publications using***
1263 ***different species, methodologies, and platforms. The actual sample cell type is indicated in the main***
1264 ***heading above the plot (NFO: “newly-formed oligodendrocyte”), and each bar represents the average***
1265 ***for two samples for each cell type index (identified by primary cell type, subtype, and publication on the***
1266 ***x-axis). The cell type indices that fall within a particular primary category of cell are further identified by***
1267 ***color (lavender: astrocytes, orange: endothelial, green: microglia, yellow: mural, purple: neuron_all,***
1268 ***blue: neuron_projection, red: neuron_interneuron, pink: oligodendrocyte, gray: oligodendrocyte***
1269 ***progenitor cell (OPC), brown: red blood cell (RBC)).***

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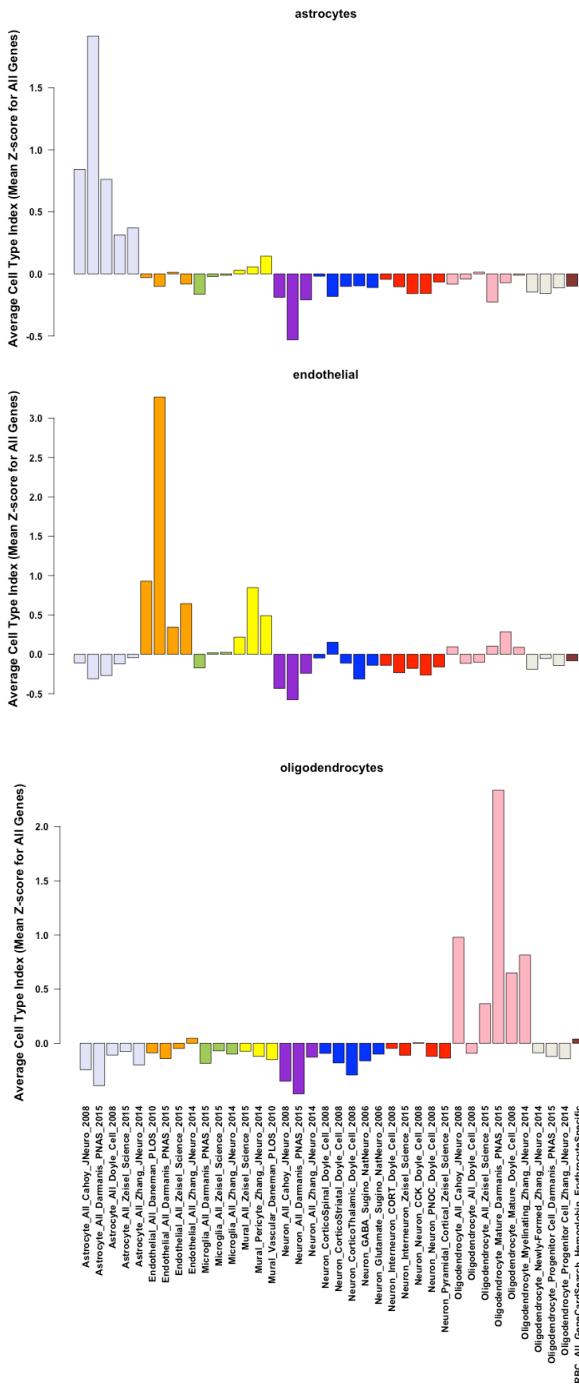
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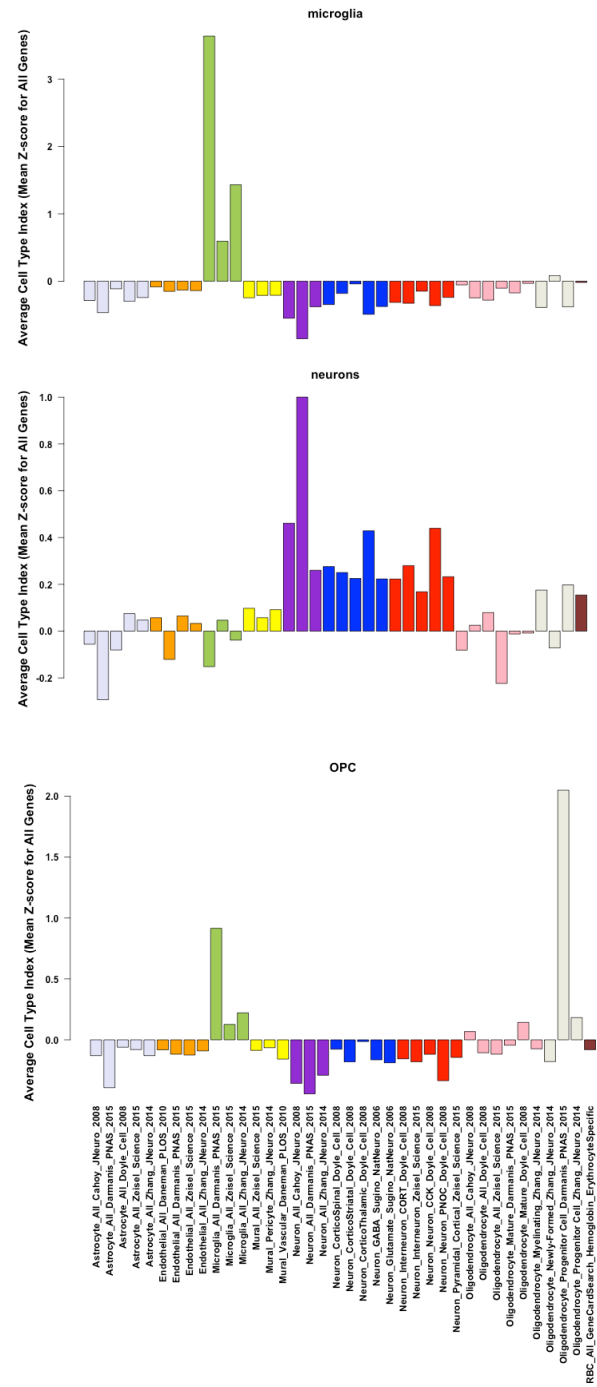
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1279 **Suppl. Figure 2. The cell content predictions derived from cell type specific transcripts originating**
1280 **from different publications successfully predict sample cell type in human single cell RNA-Seq data.**
1281 *The sample cell type in a human single cell RNA-Seq dataset (2) was predicted equally well by cell type*
1282 *indices derived from cell type specific transcripts originating from publications using different species,*
1283 *methodologies, and platforms. The sample cell type (as identified in the publication) is indicated in the*
1284 *main heading above the plot, and each bar represents the average cell type index (identified by primary*
1285 *cell type, subtype, and publication on the x-axis) for all samples of that cell type. The cell type indices that*
1286 *fall within a particular primary category of cell are further identified by color (lavender: astrocytes,*
1287 *orange: endothelial, green: microglia, yellow: mural, purple: neuron_all, blue: neuron_projection, red:*
1288 *neuron_interneuron, pink: oligodendrocyte, gray: oligodendrocyte progenitor cell (OPC), brown: red*
1289 *blood cell (RBC)).*

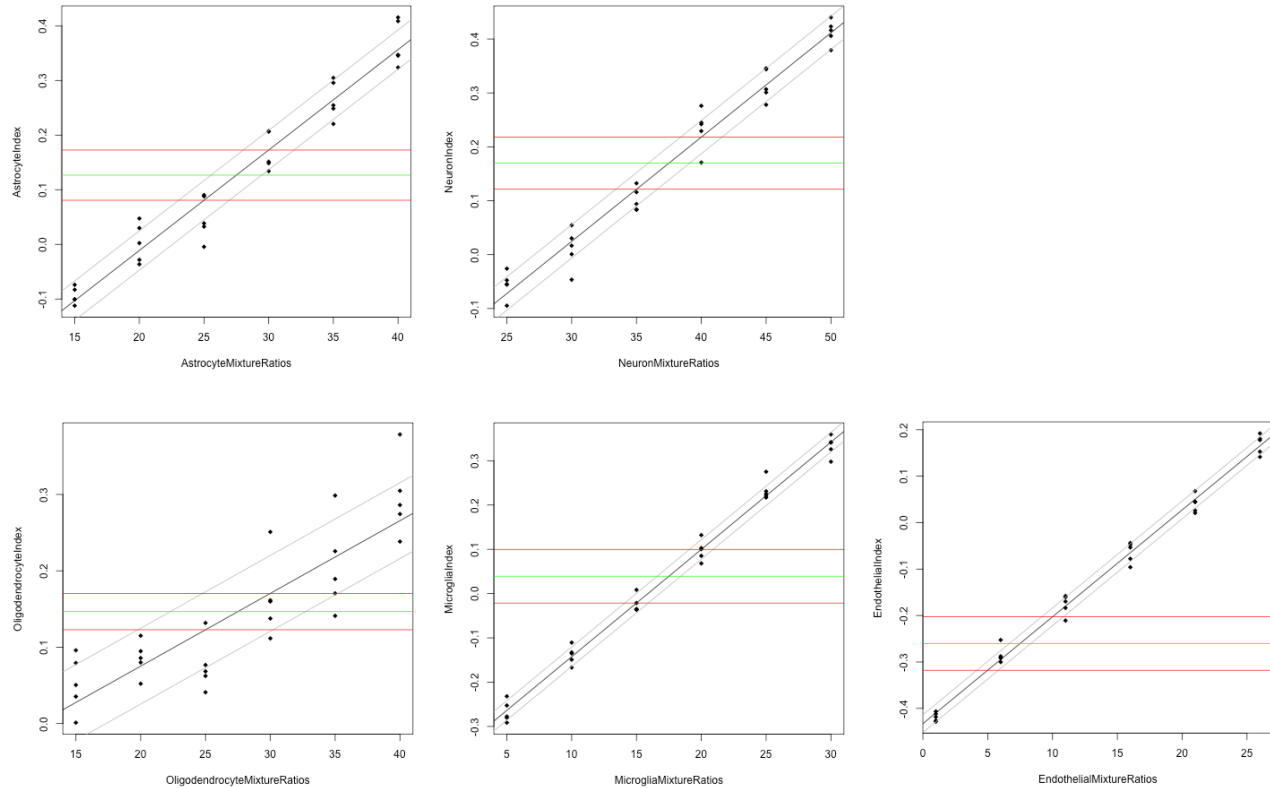
1290

1291 For further analyses, individual cell type indices were averaged within each of ten primary
1292 categories: astrocytes, endothelial cells, mural cells, microglia, immature and mature oligodendrocytes,
1293 red blood cells, interneurons, projection neurons, and indices derived from neurons in general, with any
1294 genes that were identified as being specific to more than one category removed (*e.g.*, a gene identified as
1295 being specifically expressed in both microglia and endothelial cells). This led to ten consolidated primary
1296 cell-type indices for each sample. We then examined the relationship between these consolidated cell type
1297 indices and actual cell content in artificial mixtures of 100 cells generated *in silico* by randomly sampling
1298 from the purified cell datasets (with replacement). We found that the consolidated cell type indices
1299 strongly predicted the percentage of their respective cell type included in our artificial mixtures of 100
1300 cells in a linear manner (**Suppl. Figure 3, Suppl. Figure 4**) across a range of values likely to encompass
1301 the true proportion of these cells in our cortical samples. The amount of noise present in these predictions
1302 varied by data type, with the predictions generated from single-cell data having substantially more noise
1303 than that generated from pooled, purified cells, but even the noisier data was associated with most of the
1304 data (± 1 stdev) falling within $\pm 5\%$ of the prediction. Therefore, we conclude that cell type indices are
1305 a relatively easy manner to estimate relative cell type balance across samples.

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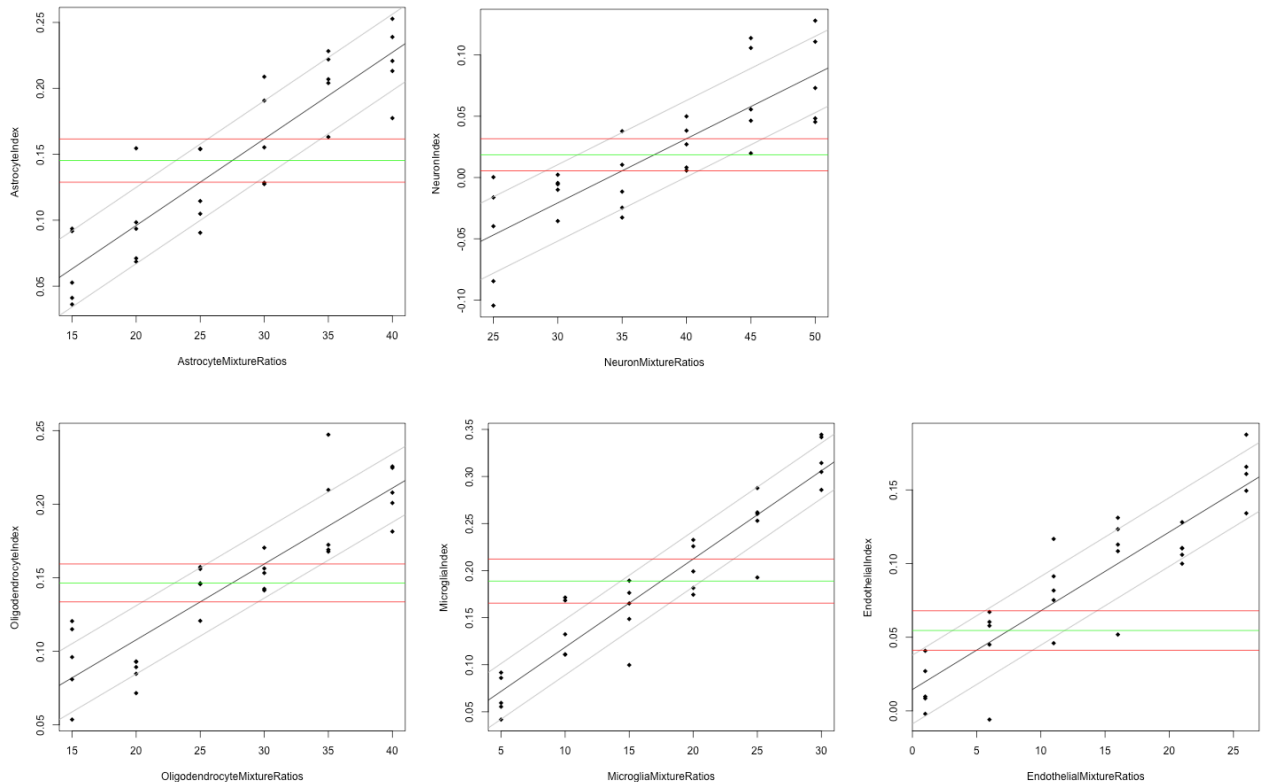
1309 **Suppl. Figure 3. Cell type indices successfully predict the percentage of cells of a particular type in**
1310 **artificial mixtures of 100 cells created using mouse purified cell type RNA-Seq data. Depicted are the**
1311 **cell type indices (y-axis) calculated for mixed cell samples generated in silico using random sampling**
1312 **(with replacement) from a mouse purified cell type RNA-Seq dataset (18). Each sample contains 100 cells**
1313 **total, with a designated percentage of the cell type of interest (x-axis), with the percentages designed to**
1314 **roughly span what might be found in cortical tissue samples. The black best fit line (as defined by a linear**
1315 **model) is accompanied by the standard error of the regression (gray), and the green and red lines are**
1316 **visual guides to help illustrate a 5% increase in the cell type of interest.**

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Running Head: PREDICTING CELL TYPE BALANCE



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1321 **Suppl. Figure 4. Cell type indices successfully predict the percentage of cells of a particular type in**
1322 **artificial mixtures of 100 cells created using human single-cell RNA-Seq data.** Depicted are the cell
1323 type indices (y-axis) calculated for mixed cell samples generated in silico using random sampling (with
1324 replacement) from a human single cell RNA-Seq dataset (2). Each sample contains 100 cells total, with a
1325 designated percentage of the cell type of interest (x-axis), with the percentages designed to roughly span
1326 what might be found in cortical tissue samples. The black best fit line (as defined by a linear model) is
1327 accompanied by the standard error of the regression (gray), and the green and red lines are visual guides
1328 to help illustrate a 5% increase in the cell type of interest. Note the greater amount of variation present in
1329 the predictions for this dataset (based on single-cell data) versus the predictions based on mouse purified
1330 cell data (**Suppl. Figure 3**).

1331

1332 As further validation, we determined whether relative cell type balance could be accurately
1333 deciphered from microarray data for samples containing artificially-generated mixtures of cultured cells
1334 (GSE19380; (12)). The cells used to make these mixtures were cultured from the cerebral cortices of rat
1335 P1 pups. The microarray profiling was then performed using a Affymetrix Rat Genome 230 2.0 Array,
1336 We downloaded the pre-processed gene expression dataset from GEO using the R package *GEOquery*.
1337 According to the methods published on GEO, this data had already undergone probe set summarization
1338 and normalization using robust multi-array averaging (RMA, *affy* package), including background

Running Head: PREDICTING CELL TYPE BALANCE

1339 subtraction, summarization by median polish, log (base 2) transformation, and quantile normalization. We
1340 used the R package *GEOquery* to extract out the description of the cell type mixture associated with each
1341 sample, and then used this data to construct a new matrix that contained the percent of each cell type
1342 (columns: neuron, astrocyte, oligodendrocyte, microglia) found in each sample. We then predicted the
1343 cell content of each sample from the microarray data using BrainInABlender, and plotted these
1344 predictions against the actual percent of each cell type found in the mixtures. We made these plots both
1345 for predictions derived from cell type specific gene lists from particular publications (**Suppl. Figure 6**)
1346 and after averaging these individual cell type indices within each of ten primary categories, with any
1347 genes that were identified as being specific to more than one category removed (*e.g.*, a gene identified as
1348 being specifically expressed in both microglia and endothelial cells, **Suppl. Figure 5**). These results are
1349 included in main text of paper (**Figure 3**). The code for all of these analyses can be found at:
1350 https://github.com/hagenaue/CellTypeAnalyses_KuhnMixtures/tree/master.

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Cell Type Index	Number of probesets in the microarray that represent cell type specific genes according to each publication	Percent that were truly specific to that cell type (not identified as "specific" to another category of cell type in a different publication)
Astrocyte_All_Cahoy_JNeuro_2008	47	87%
Astrocyte_All_Darmanis_PNAS_2015	14	93%
Astrocyte_All_Doyle_Cell_2008	12	100%
Astrocyte_All_Zeisel_Science_2015	181	88%
Astrocyte_All_Zhang_JNeuro_2014	32	78%
Endothelial_All_Daneman_PLOS_2010	34	76%
Endothelial_All_Darmanis_PNAS_2015	14	93%
Endothelial_All_Zeisel_Science_2015	261	90%
Endothelial_All_Zhang_JNeuro_2014	30	83%
Microglia_All_Darmanis_PNAS_2015	17	94%
Microglia_All_Zeisel_Science_2015	305	91%
Microglia_All_Zhang_JNeuro_2014	26	88%
Mural_All_Zeisel_Science_2015	114	93%
Mural_Pericyte_Zhang_JNeuro_2014	32	69%
Mural_Vascular_Daneman_PLOS_2010	36	64%
Neuron_All_Cahoy_JNeuro_2008	60	63%
Neuron_All_Darmanis_PNAS_2015	18	72%
Neuron_All_Zhang_JNeuro_2014	22	68%
Neuron_CorticoSpinal_Doyle_Cell_2008	17	59%
Neuron_CorticoStriatum_Doyle_Cell_2008	16	6%
Neuron_CorticoThalamic_Doyle_Cell_2008	14	64%
Neuron_GABA_Sugino_NatNeuro_2006	23	83%
Neuron_Glutamate_Sugino_NatNeuro_2006	48	81%
Neuron_Interneuron_CORT_Doyle_Cell_2008	13	77%
Neuron_Interneuron_Zeisel_Science_2015	259	90%
Neuron_Neuron_CCK_Doyle_Cell_2008	12	58%
Neuron_Neuron_PNOC_Doyle_Cell_2008	18	67%
Neuron_Pyramidal_Cortical_Zeisel_Science_2015	189	88%
Oligodendrocyte_All_Cahoy_JNeuro_2008	33	94%
Oligodendrocyte_All_Doyle_Cell_2008	19	74%
Oligodendrocyte_All_Zeisel_Science_2015	323	93%
Oligodendrocyte_Mature_Darmanis_PNAS_2015	15	100%
Oligodendrocyte_Mature_Doyle_Cell_2008	18	72%
Oligodendrocyte_Myelinating_Zhang_JNeuro_2014	34	100%
Oligodendrocyte_Newly-Formed_Zhang_JNeuro_2014	31	65%
Oligodendrocyte_Progenitor_Cell_Darmanis_PNAS_2015	15	73%
Oligodendrocyte_Progenitor_Cell_Zhang_JNeuro_2014	32	59%
RBC_All_GeneCardSearch_Hemoglobin_ErythrocyteSpecific	5	100%

1354 ***Suppl. Figure 5. Identifying non-specific “cell-type specific genes”: An example from dataset***
1355 ***GSE19380 of the number of probesets that represented genes identified as cell type specific in each***
1356 ***publication in our database vs. the percentage that were actually found to truly specific to that cell type***
1357 ***(i.e., not identified as "specific" to another category of cell type in a different publication). The data***
1358 ***from genes that were identified as being specific to more than one category of cell type (e.g., a gene***
1359 ***identified as being specifically expressed in both microglia and endothelial cells) was removed before***
1360 ***averaging the individual cell type indices within each of ten primary categories (astrocytes, endothelial***
1361 ***cells, mural cells, microglia, immature and mature oligodendrocytes, red blood cells, interneurons,***
1362 ***projection neurons, and indices derived from neurons in general) to create the ten consolidated primary***
1363 ***cell-type indices used throughout our paper.***

1364

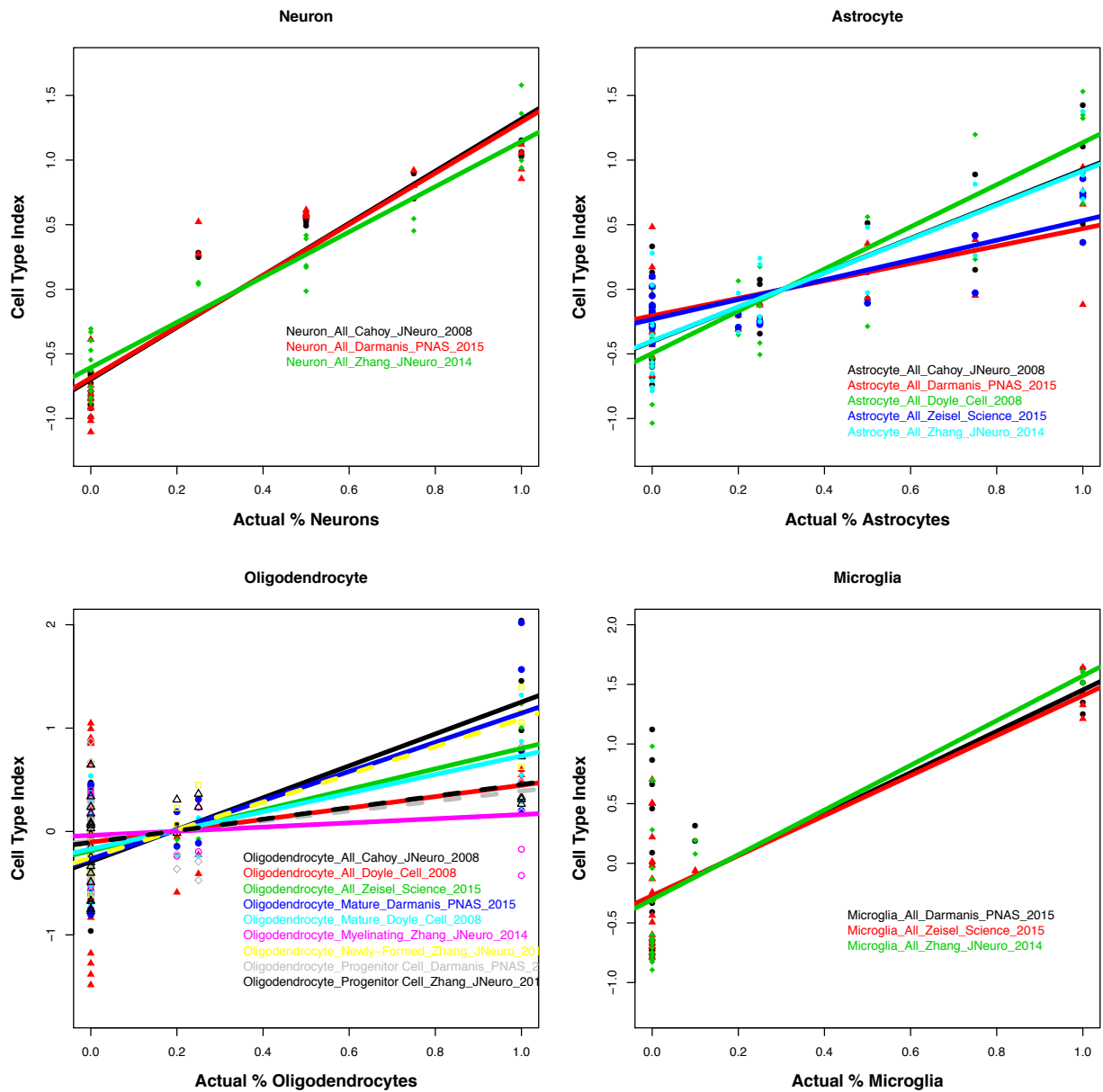
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1371 **Suppl. Figure 6. Validation of Relative Cell Content Predictions.** A) Using a microarray dataset
1372 derived from samples that contained artificially-generated mixtures of cultured cells (GSE19380; (12)),
1373 we found that our relative cell content predictions ("cell type indices") closely reflected actual known
1374 content, except that the percentage of cultured oligodendrocytes included in the mixtures was better
1375 predicted using cell type specific gene lists derived from immature oligodendrocytes instead of mature
1376 oligodendrocytes.

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1382 **7.2 Comparison of Our Method vs. PSEA: Predicting Cell Identity in a Human Single-Cell RNA-**

1383 **Seq Dataset**

1384 Although we generated our method independently to address microarray analysis questions that arose
1385 within the Pritzker Neuropsychiatric Consortium, we later discovered that it was quite similar to the
1386 technique of population-specific expression analysis (PSEA) introduced by (12) with several notable
1387 differences. Similar to our method, PSEA is a carefully-validated analysis method which aims to estimate
1388 cell type-differentiated disease effects from microarray data derived from brain tissue of heterogeneous
1389 composition and approaches this problem by including the averaged, normalized expression of cell type
1390 specific markers within a larger linear model that is used to estimate differential expression in microarray
1391 data (10–12). Analyses using PSEA similarly indicated that individual variability in neuronal, astrocytic,
1392 oligodendrocytic, and microglial cell content was sufficient to account for substantial variability in the
1393 vast majority of probe sets in microarray data from human brain samples, even within non-diseased
1394 samples (12). The differences between our techniques are mostly due to the recent growth of the literature
1395 documenting cell type specific expression in brain cell types. PSEA uses a very small set of markers (4-7)
1396 to represent each cell type, and screens these markers for tight co-expression within the dataset of interest,
1397 since co-expression networks have been previously demonstrated to often represent cell type signatures in
1398 the data (92). This is essential for the analysis of microarray data for brain regions that have not been well
1399 characterized for cell type specific expression (e.g., the substantia nigra), but risks the possibility of
1400 closely tracking variability in a particular cell function instead of cell content (as described in our results
1401 related to aging). Our analysis predominantly focused on the well-studied cortex, thus enabling us to
1402 expand our analysis to include hundreds of cell type specific markers derived from a variety of
1403 experimental techniques.

1404 Our manner of normalizing data also differs: PSEA normalizes the expression values for each gene by
1405 dividing by the average expression of that gene across samples, whereas we use z-score normalization,

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1406 both at the level of the individual transcript and later at the level of the gene level summary data. Due to
1407 the dependence of PSEA on ratios, genes that have average expression values that are close to zero can
1408 end up with normalized values that are extremely high for a handful of samples. For microarray data, this
1409 form of normalization should function well because \log_2 expression values rarely drop below 5.
1410 However, within RNA-Seq, counts of zero are quite common and therefore we suspected that the ratio-
1411 form of normalization used by PSEA might not function optimally for this data type.

1412 Therefore, we decided to run a head-to-head comparison of our method and PSEA using a single-
1413 cell RNA-Seq dataset derived from freshly-resected human cortex (2). To make the comparison as
1414 interpretable as possible, we used the same list of cell type specific genes for both methods: the cell type
1415 specific genes remaining in our database following the removal of all transcripts that were found to be
1416 “specifically expressed” in multiple categories of cell types (e.g., a transcript that is “specific” to both
1417 astrocytes and neurons). In order to avoid circular reasoning, we also did not include any cell type
1418 specific genes that had originally been identified by the publication currently used as the test dataset (2).
1419 Then we extracted the variance-stabilized and filtered data (see **Section 7.1**) for the cell type specific
1420 genes. For PSEA, we downloaded the PSEA package from Bioconductor
1421 (<https://www.bioconductor.org/packages/release/bioc/vignettes/PSEA/inst/doc/PSEA.pdf>) and used the
1422 `marker()` function to calculate the “Reference Signal” for the most common primary categories of cell
1423 types (astrocytes, endothelial cells, microglia, mature oligodendrocytes, and neurons in general). For our
1424 method, we used a procedure similar to that used in the manuscript. We applied a z-score transformation
1425 to the data for each gene (mean=0, sd=1), and then either averaged by the primary cell type category (to
1426 conduct an analysis most similar to PSEA), or averaged the data from the cell type specific genes
1427 identified by each publication, followed by averaging by primary cell type category (to create
1428 consolidated cell type indices similar to those used in most of our manuscript).

1429 To compare the efficacy of each method, we ran a linear model to determine the percentage of
1430 variation in the population “reference signal” (PSEA) or “cell type index” (our method) accounted for by
1431 the cell type identities assigned to each cell in the original publication (2). We found that both the

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1432 population reference signals (PSEA) and cell type indices (our method) for each cell were strongly related
1433 to their previously-assigned cell type identity, but in general the relationship was stronger when using our
1434 method: on average, 34% of the variation in the reference signal for each cell type was accounted for by
1435 cell identity, whereas an average of either 45% or 49% of the variation in our cell type indices was
1436 accounted for by cell identity using either the simplified or consolidated versions of our method,
1437 respectively (**Suppl. Figure 7**). An illustration of this improvement can be found in **Suppl. Figure 8**: note
1438 the presence of extreme outliers in the population reference signal when using the PSEA method. We
1439 conclude that the simple use of a different normalization method is sufficient to make our method a more
1440 effective manner of predicting cell type balance in some datasets. We also find that averaging the
1441 predictions drawn from the cell type specific genes identified by multiple publications into a consolidated
1442 index produces some additional improvement.
1443

1444

Method of deriving a statistical cell type signal:

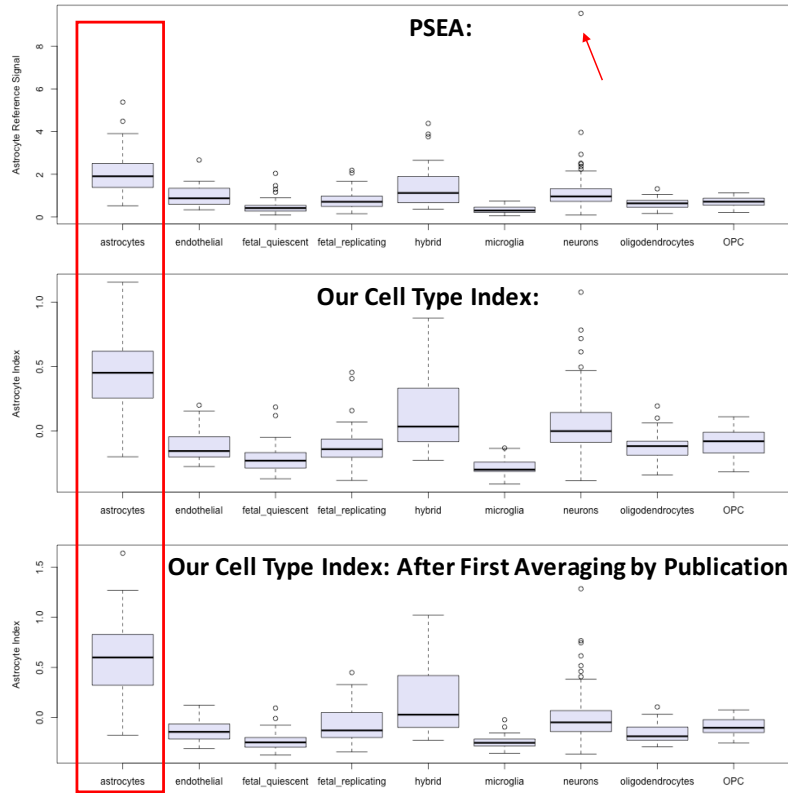
Signal from cell type specific genes for:	PSEA (mean signal ratio average)	Our Cell Type Indices (z-score average)	Our Cell Type Indices: After first averaging by Publication
Astrocytes	34%	52%	57%
Oligodendrocytes	38%	45%	50%
Microglia	36%	42%	51%
Endothelial	30%	28%	33%
Neurons	33%	57%	53%

1445

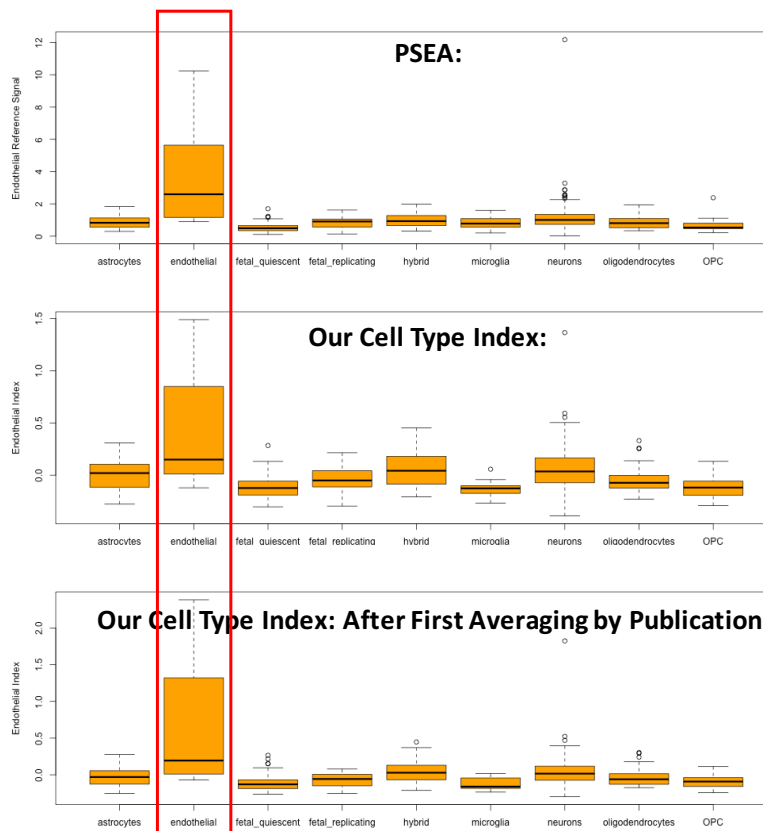
1446 **Suppl. Figure 7. The method for deriving predicted relative cell content determines the strength of the**
 1447 **relationship with sample cell type.** Depicted below is a comparison of the efficacy of three different
 1448 manners of predicting the relative cell content of samples (columns) in a human single-cell RNA-seq
 1449 dataset (2): 1) the “population reference signal” generated by PSEA, 2) a simplified version of our
 1450 method that is meant to be relatively analogous to PSEA (a simple average of the z-score-transformed
 1451 data for all genes specific to a particular cell type in our database), 3) the version of our method used in
 1452 this manuscript, which consolidates the predictions derived from the cell type specific genes identified in
 1453 different publications. For the predicted relative content of each of the major cell types (rows) derived
 1454 using these different methods, the table provides the percentage of variation (*r*-squared) that is accounted
 1455 for by the original cell type identities of the samples provided by the publication (2). Overall, there is a
 1456 strong relationship between the predictions generated by all methods and sample cell type identity, but
 1457 the method used in this manuscript produces predictions that best fit sample cell type.

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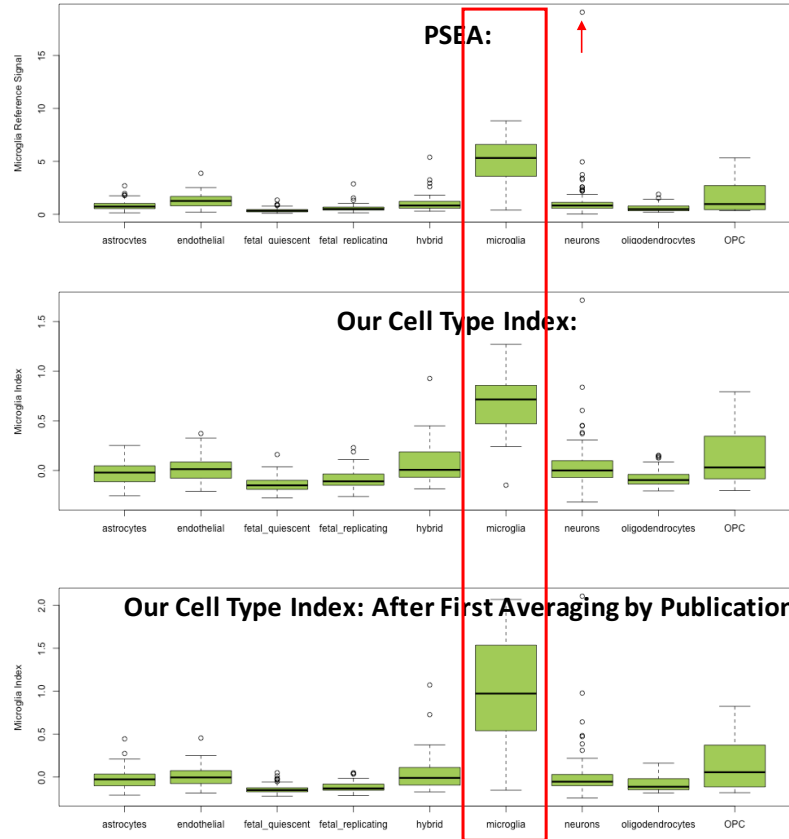


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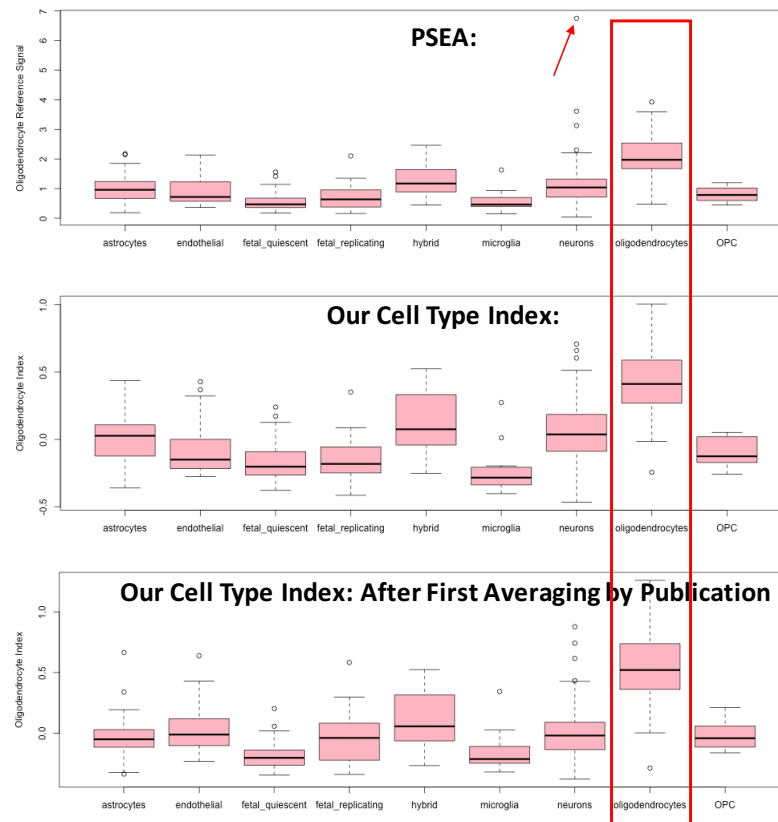


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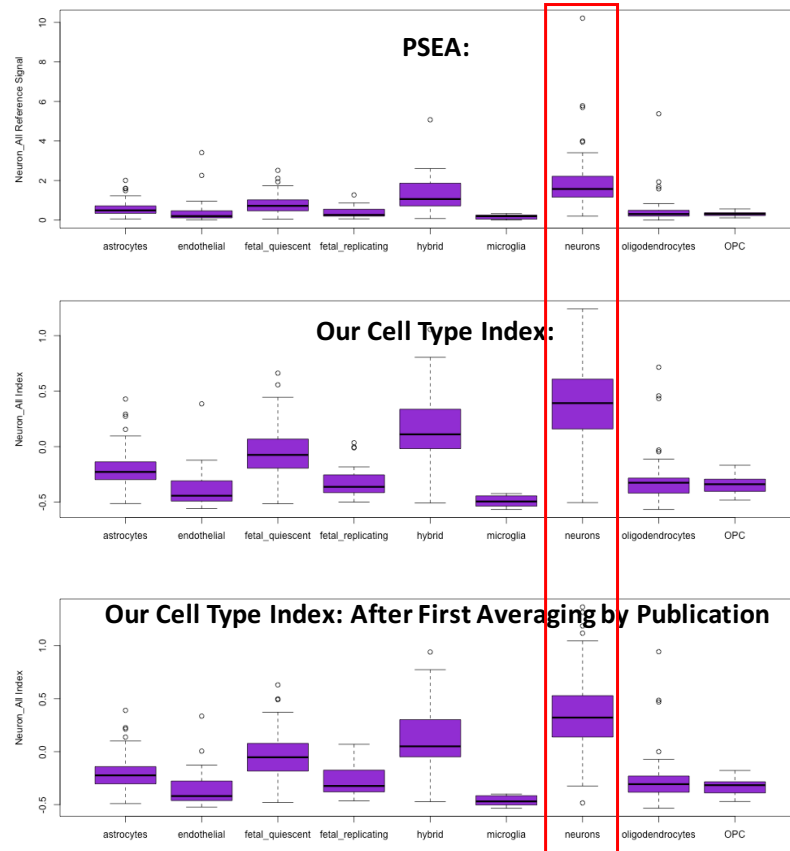


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1464 **Suppl. Figure 8. The method for deriving predicted relative cell content determines the strength of the**
1465 **relationship with sample cell type.** Depicted below is a comparison of the efficacy of three different
1466 manners of predicting the relative cell content of samples (columns) in a human single-cell RNA-seq
1467 dataset (2): 1) the “population reference signal” generated by PSEA, 2) a simplified version of our
1468 method that is meant to be relatively analogous to PSEA (a simple average of the z-score-transformed
1469 data for all genes specific to a particular cell type in our database), 3) the version of our method used in
1470 this manuscript, which consolidates the predictions derived from the cell type specific genes identified in
1471 different publications. Boxplots illustrate the distribution of the relative cell content predictions across
1472 samples identified as different cell types in the original publication (2). Lavender: astrocytes, orange:
1473 endothelial cells, green: microglia, pink: oligodendrocytes, purple: neurons. Note the presence of several
1474 extreme outliers (red) in the predictions produced by PSEA.

1475

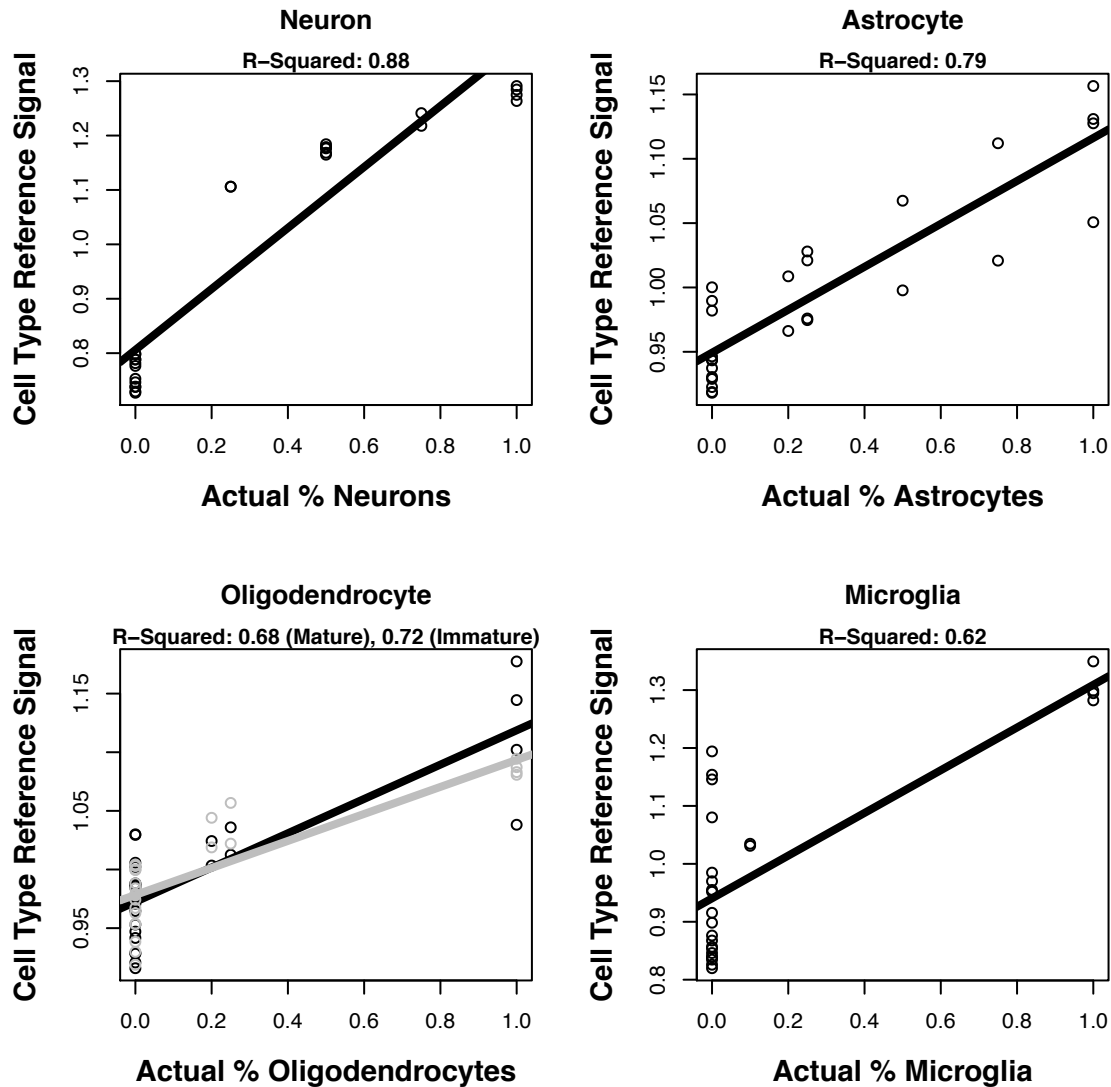
1476 Using similar methodology, we also calculated the population “reference signal” with PSEA for
1477 microarray data from artificially-created mixtures of cultured cells (GSE19380 – see discussion of data
1478 preprocessing in **Section 7.1**). The results strongly tracked the actual cell content of the mixed samples
1479 (**Suppl. Figure 9**) in a manner that was not strikingly better or worse than the predictions made using
1480 BrainInABlender for the same dataset (**Figure 3**). This again drives home the fact that the ratio-based

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1481 normalization methods used in PSEA are particularly incompatible with low count data in RNA-Seq –

1482 results derived from microarray data are fine.

1483



1484

1485 **Suppl. Figure 9. Relative cell content predictions made using PSEA and our cell type specific gene**
1486 **lists.** Using a microarray dataset derived from samples that contained artificially-generated mixtures of
1487 cultured cells (GSE19380; (12)), we found that the relative cell content predictions (“cell type reference
1488 signal”) produced by PSEA closely reflected actual known content, similar to the predictions made by
1489 BrainInAblender (Figure 3).

1490

1491

1492 **7.3 Additional Detailed Preprocessing Methods for the Macro-Dissected Microarray Datasets**

1493

1494 **7.3.1 Pritzker Dorsolateral Prefrontal Cortex Microarray Dataset (GSE92538)**

1495 The original dataset included tissue from 172 high-quality human post-mortem brains donated to
1496 the Brain Donor Program at the University of California, Irvine with the consent of the next of kin.
1497 Frozen coronal slabs were macro-dissected to obtain dorsolateral prefrontal cortex samples. Clinical
1498 information was obtained from medical examiners, coroners' medical records, and a family member.
1499 Patients were diagnosed with either Major Depressive Disorder, Bipolar Disorder, or Schizophrenia by
1500 consensus based on criteria from the Diagnostic and Statistical Manual of Mental Disorders (93). Due to
1501 the extended nature of this study, this sample collection occurred in waves ("cohorts") over a period of
1502 many years. This research was overseen and approved by the University of Michigan Institutional Review
1503 Board (IRB # HUM00043530, Pritzker Neuropsychiatric Disorders Research Consortium (2001-0826))
1504 and the University of California Irvine (UCI) Institutional Review Board (IRB# 1997-74).

1505 As described previously (32,35), total RNA from these samples was then distributed to
1506 laboratories at three different institutions (University of Michigan (UM), University of California-Davis
1507 (UCD), University of California-Irvine (UCI)) to be hybridized to either Affymetrix HT-U133A or HT-
1508 U133Plus-v2 chips (1-5 replicates per sample, n=367). Before conducting the current analysis, the subset
1509 of probes found on both the Affymetrix HT-U133A and HT-U133Plus-v2 chips was extracted,
1510 reannotated for probe-to-transcript correspondance (94), summarized using robust multi-array analysis
1511 (RMA) (34), log (base 2)-transformed, quantile normalized, and gender-checked. Then, 15 batches of
1512 highly-correlated samples were identified that were defined a combination of cohort, chip, and laboratory
1513 **(Suppl. Figure 10).**

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Batch#	Site	Chip	Cohort	Control	BP	MDD	SCHIZ
1	UCD	U133A	Dep Cohort 1 & 2	20	9	11	0
2	UCD	U133A	Dep Cohort 3	11	6	5	0
3	UCD	U133A	Dep Cohort 4	16	4	7	0
4	UCD	U133Plus2	Dep Cohort 5	13	5	10	0
5	UCD	U133A	Schiz Cohort 1	9	0	0	9
6	UCD	U133Plus2	Schiz Cohort 1	8	0	0	8
7	UCD	U133Plus2	Schiz Cohort 2	8	0	0	10
8	UCI	U133A	Schiz Cohort 1	9	0	0	9
9	UM	U133A	Dep Cohort 1	16	10	9	0
10	UM	U133A	Dep Cohort 2	3	2	5	0
11	UM	U133A	Dep Cohort 3 & 4	27	11	11	0
12	UM	U133Plus2	Dep Cohort 5	13	5	10	0
13	UM	U133Plus2	Dep Cohort 6	7	2	9	3
14	UM	U133A	Schiz Cohort 1	9	0	0	9
15	UM	U133Plus2	Schiz Cohort 2	9	0	0	10

1514

1515 **Suppl. Figure 10. The number of microarray chips run in each batch, defined by processing site,**
1516 **Affymetrix chip type, and sample collection cohort. Samples from the four diagnostic categories**
1517 **(Control, Bipolar Disorder, Major Depressive Disorder, Schizophrenia) were unevenly distributed across**
1518 **batches.**

1519

1520 Samples that exhibited markedly low average sample-sample correlation coefficients (<0.85:
1521 outliers) were removed from the dataset, including data from one batch that exhibited overall low sample-
1522 sample correlation coefficients with other batches and was a poor match with their duplicate microarrays
1523 run in a separate laboratory. The batch effects were then subtracted out using median-centering (detailed
1524 procedure: (35)) and the replicate samples were averaged for each subject. Our current analyses began
1525 with this sample-level summary gene expression data (publicly available in the Gene Expression
1526 Omnibus, GEO: GSE92538). We further removed data from any subjects lacking information regarding
1527 critical pre- or post-mortem variables necessary for our analysis, leaving a final sample size of n=157. All
1528 of the R script documenting these analyses can be found at

1529 https://github.com/hagenaue/CellTypeAnalyses_PritzkerAffyDLPFC.

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1535 **7.3.2 Allen Brain Atlas Cross-Regional Microarray Dataset**

1536 The Allen Brain Atlas microarray data was downloaded from <http://human.brain->
1537 [map.org/microarray/search](http://human.brain-map.org/microarray/search) on December 2015. This microarray survey was performed in brain-specific
1538 batches, with multiple batches per subject. To remove technical variation across batches, a variety of
1539 normalization procedures had been performed by the original authors both within and across batches
1540 using internal controls, as well as across subjects (28). The dataset available for download had already
1541 been log-transformed (base 2) and converted to z-scores using the average and standard deviation for each
1542 probe. These normalization procedures were designed to remove technical artifacts while best preserving
1543 cross-regional effects in the data, but the full information about relative levels of expression within an
1544 individual sample were unavailable and the effects of subject-level variables (such as age and pH) were
1545 likely to be de-emphasized due to the inability to fully separate out subject and batch during the
1546 normalization process.

1547 Prior to conducting other analyses, we averaged the expression level of the multiple probes that
1548 corresponded to the same gene, and re-scaled, so that the data associated with each gene symbol
1549 continued to be a z-score (mean=0, sd=1). The 30,000 probes mapped onto 18,787 unique genes (as
1550 determined by gene symbol). We then extracted the z-score data for the list of cell type specific genes
1551 derived from each publication (1608 total). Then, based on our results from analyzing the Pritzker dataset,
1552 we excluded the data for genes that were non-specific (i.e., included in a list of cell type specific genes
1553 from a different category of cells within any of the publications), and then averaged the data from the
1554 cell-type specific genes derived from each publication to predict the relative content of each of the 10
1555 primary cell types in each sample. All of the R script documenting these analyses can be found at
1556 https://github.com/hagenaue/CellTypeAnalyses_AllenBrainAtlas.

1557

1558

1559 **7.3.3 Human Cortical Microarray Dataset GSE53987 (submitted to GEO by Lanz et al. (36))**

1560 The full publicly-available dataset GSE53987 (described in (36)) contained Affymetrix
1561 U133Plus2 microarray data from 205 post-mortem human brain samples from three brain regions: the
1562 DLPFC (Brodmann Area 46, focusing on gray matter only (Lanz T.A., *personal communication*)), the
1563 hippocampus, and the striatum. These samples were collected by the University of Pittsburgh brain bank.
1564 For the purposes of our current analysis, we only downloaded the microarray .CEL files for the
1565 dorsolateral prefrontal cortex samples. We summarized these data with robust multi-array analysis
1566 (RMA) (from the R package *affy* (34)) using a custom up-to-date chip definition file (.cdf) to define
1567 probe-to-transcript correspondence (“hgu133plus2hsentrezgcdf_19.0.0.tar.gz” from [http://nmg-
1569 r.bioinformatics.nl/NuGO_R.html](http://nmg-
1568 r.bioinformatics.nl/NuGO_R.html) (94)). This process included background subtraction, log (base 2)-
1569 transformation, and quantile normalization. Gene Symbol annotation for probeset Entrez gene ids were
1570 provided by the R package *org.Hs.eg.db*. We extracted the sample characteristics from the GEO website
1571 using the R package *GEOquery*. To control for technical variation, the sample processing batches were
1572 estimated using the microarray chip scan dates extracted from the microarray .CEL files (using the
1573 function *protocolData* in the *GEOquery* package), but it appeared that all chips for the DLPFC were on
1574 the same date. RNA degradation was estimated using the R package *AffyRNADegradation* (39). During
1575 quality control, two samples were removed - GSM1304979 had a range of sample-sample correlations
1576 that was unusually low compared (median=0.978) compared to range for the dataset as a whole (median:
1577 0.993) and GSM1304953 appeared to be falsely identified as female (signal for XIST<7). We then
1578 predicted the cell content of each sample from the microarray data using *BrainInAblender*. The code for
1579 all analyses can be found at:

1580 https://github.com/hagenaue/CellTypeAnalyses_LanzHumanDLPFC/tree/master

1581

1582 **7.3.4 Human Cortical Microarray Dataset GSE21138 (submitted to GEO by Narayan et al. (38))**

1583 The publicly-available dataset GSE21138 (described in (38)) contained Affymetrix U133Plus2
1584 microarray data from 59 post-mortem human brain samples from the DLPFC (Brodmann Area 46, gray

1585 matter only (Thomas E.A., *personal communication*) collected by the Mental Health Research Institute
1586 in Victoria, Australia. The procedures for data download and pre-processing were identical to those used
1587 above for GSE53987 with a few minor exceptions. In particular, there were six separate scan dates
1588 associated with the microarray .CEL files, but one of these scan dates was not included as a co-variate in
1589 our analyses because it had an n=1 (“06/14/06”). During quality control, the data for two subjects because
1590 they appeared to be falsely-identified as male (XIST>7, GSM528839 & GSM528840) , and one subject
1591 that appeared to be falsely-identified as female (XIST<7, GSM528880). Data for two more subjects were
1592 removed as outliers due to having an unusually low range of sample-sample correlations (GSM528866,
1593 GSM528873) as compared to the dataset as a whole. The code for all analyses can be found at:
1594 https://github.com/hagenaue/CellTypeAnalyses_NarayanHumanDLPFC.

1595

1596 **7.3.5 Human Cortical Microarray Dataset GSE21935 (submitted to GEO by Barnes et al. (37))**

1597 The publicly-available dataset GSE21935 (described in (37)) contained Affymetrix U133Plus2
1598 microarray data from 42 post-mortem human brain samples from the temporal cortex (Brodmann Area
1599 22) collected at the Charing Cross campus of the Imperial College of London. The procedures for data
1600 download and pre-processing were identical to those used above for GSE53987 with a few minor
1601 exceptions. In particular, there were two separate scan dates associated with the microarray .CEL files,
1602 but they were closely spaced (6/25/04 vs. 6/29/04) and we did not find any strong association between
1603 scan date and any of the top principal components of variation in the data, so we opted to not include scan
1604 date as a co-variate in our statistical models. Quality control did not identify any problematic samples.
1605 The code for all analyses can be found at:

1606 https://github.com/hagenaue/CellTypeAnalyses_BarnesHumanCortex/tree/master.

1607

1608 **7.3.6 CommonMind Consortium Human Cortical RNA-Seq Dataset**

1609 The CommonMind Consortium (CMC) RNA-seq dataset profiled prefrontal cortex samples from 603
1610 individuals (40) collected at three brain banks: Mount Sinai School of Medicine, University of Pittsburgh,
1611 and University of Pennsylvania. This dataset was downloaded as GRCh37-aligned bam files from the
1612 CommonMind Consortium Knowledge Portal (<https://www.synapse.org/CMC>). Tophat-aligned bam files
1613 were converted back to fastq format and mapped to GRCh38 using HISAT2 (41) with default settings.
1614 Reads mapping uniquely to exons were then counted using subread featureCounts with ensembl transcript
1615 models. RNA-seq read counts were analyzed using limma/voom (42); cell type indices were calculated on
1616 logCPM values, and analysis of differential gene expression was performed using limma with observed
1617 precision weights in a weighted least squares linear regression. Prior to upload, poor quality samples from
1618 the original dataset (40) had already been removed (<50 million reads, RIN<5.5) and replaced with higher
1619 quality samples. We further excluded data from 10 replicates and 89 individuals with incomplete
1620 demographic data (missing pH), leaving a final sample size of 514 samples. The dataset was further
1621 filtered using an expression threshold (CPM>1 in at least 50 individuals) which reduced the dataset from
1622 including data from all annotated genes (about 60,000) to data from around 17,000 genes.

1623

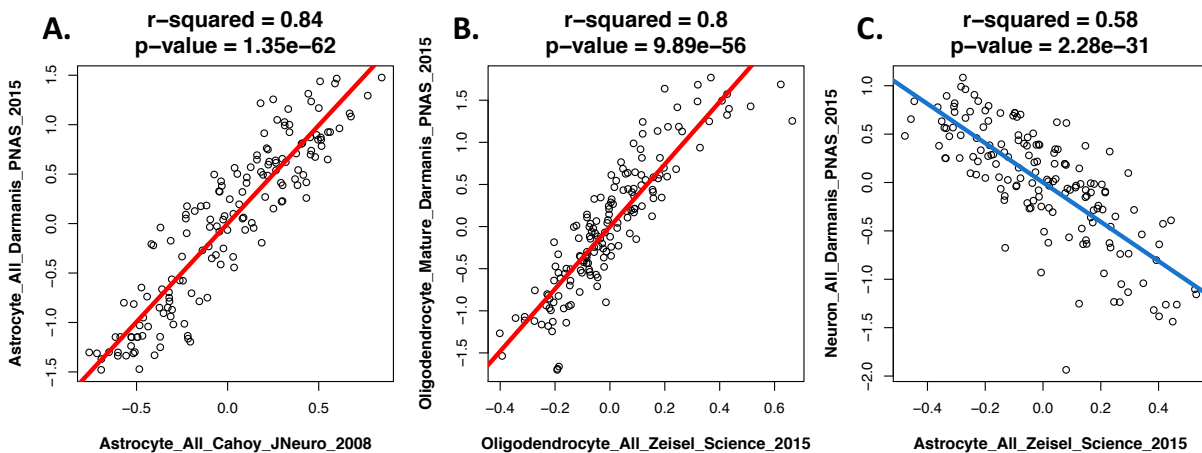
1624

1625 **7.4 Additional figures and results: Does the Reference Dataset Matter? There is a Strong**
1626 **Convergence of Cell Content Predictions Derived from Cell Type Specific Transcripts**
1627 **Identified by Different Publications**

1628 Similar to what we observed during our validation analyses using data from purified cell types,
1629 we found that the predicted cell content for our post-mortem human cortical samples (“cell type indices”)
1630 was similar regardless of the methodology used to generate the cell type specific gene lists used in the
1631 predictions. Within all four of the cortical microarray datasets, there was a strong positive correlation
1632 between cell type indices representing the same cell type, even when the predictions were derived using
1633 cell type specific gene lists from different species, cell type purification strategies, and platforms. In

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1634 general, we found that the pattern of correlations between the 38 cell type indices clearly clustered within
1635 three large umbrella categories: neurons, oligodendrocytes, and support cells (astrocytes, microglia, and
1636 neurovasculature). This clustering was clear using visual inspection of the correlation matrices (**Suppl.**
1637 **Figure 11, Suppl. Figure 12**), hierarchical clustering, or consensus clustering (**Suppl. Figure 13**;
1638 ConsensusClusterPlus: (43)) and persisted even after removing data from genes identified as cell type
1639 specific in multiple publications (e.g., gene expression identified as astrocyte-expression in both
1640 Cahoy_Astrocyte and Zhang_Astrocyte; **Suppl. Figure 14, Suppl. Figure 16**). In some datasets, the cell
1641 type indices for support cell subcategories were also clearly clustered (**Suppl. Figure 11**). In contrast,
1642 clustering was not able to reliably discern neuronal subcategories (interneurons, projection neurons) in
1643 any dataset. Likewise, oligodendrocyte progenitor cell indices derived from different publications did not
1644 strongly correlate with each other, perhaps indicating a lack of significant presence of progenitor cells in
1645 the cortex of the primarily middle-aged subjects.



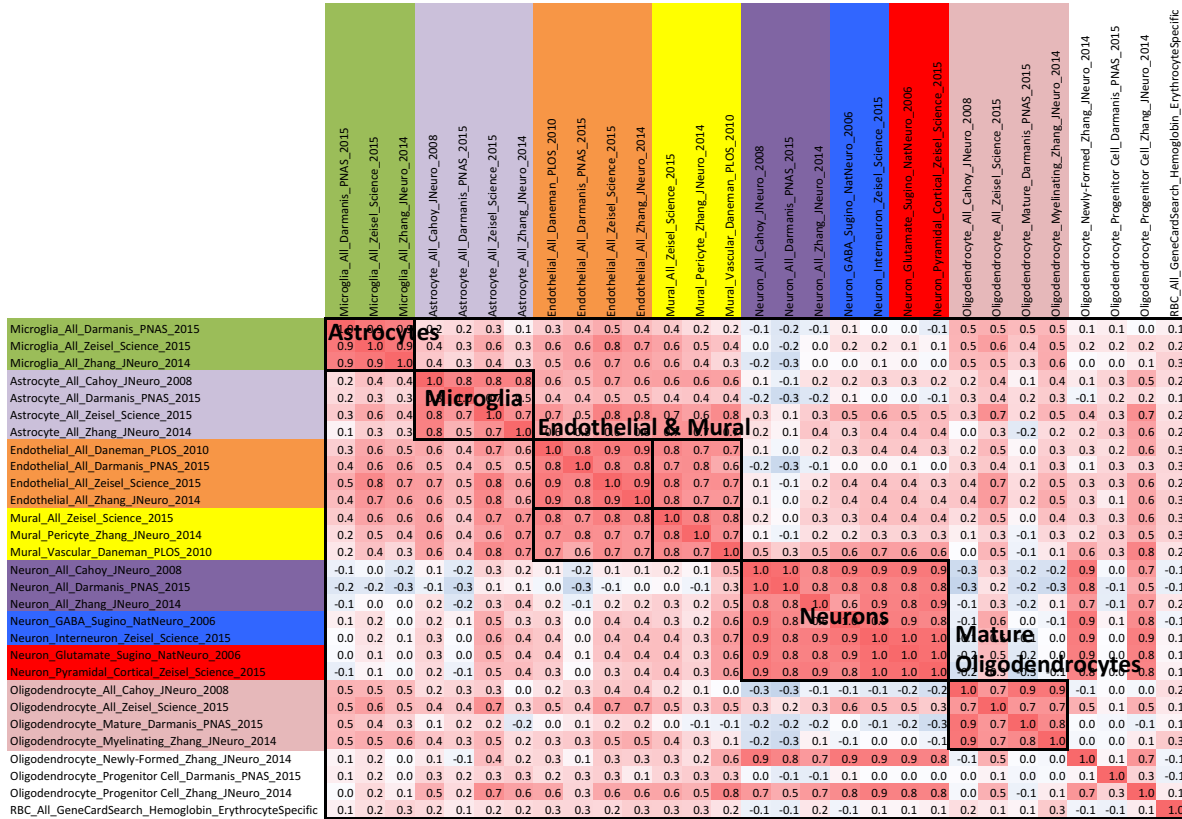
Running Head: PREDICTING CELL TYPE BALANCE

	Astrocyte_AI_Cahoy_Neuro_2008	Astrocyte_AI_Darmanis_PNAS_2015	Astrocyte_AI_Doye_Cell_2008	Astrocyte_AI_Zeisel_Science_2015	Astrocyte_AI_Zhang_Neuro_2014	Microglia_AI_Darmanis_PNAS_2015	Microglia_AI_Zeisel_Science_2015	Microglia_AI_Zhang_Neuro_2014	Endothelial_AI_Darmanis_PNAS_2015	Endothelial_AI_Zeisel_Science_2015	Mural_Pericyte_Zhang_Neuro_2014	Mural_Vascular_Daneman_PLOS_2010	Neuron_AI_Cahoy_Neuro_2008	Neuron_AI_Darmanis_PNAS_2015	Neuron_AI_Doye_Cell_2008	Neuron_AI_Zhang_Neuro_2014	Neuron_Corticospinal_Doye_Cell_2008	Neuron_Corticospinal_Zhang_Neuro_2014	Neuron_GABA_Sugino_NatNeuro_2006	Neuron_Interneuron_CORT_Doye_Cell_2008	Neuron_Interneuron_Zeisel_Science_2015	Neuron_Neuron_CCK_Doye_Cell_2008	Oligodendrocyte_AI_Cahoy_Neuro_2008	Oligodendrocyte_AI_Zeisel_Science_2015	Oligodendrocyte_Mature_Darmanis_PNAS_2015	Oligodendrocyte_Mature_Doye_Cell_2008	Oligodendrocyte_Myelinating_Zhang_Neuro_2014	Oligodendrocyte_Newly-Formed_Zhang_Neuro_2014	Oligodendrocyte_Progenitor_Cell_Darmanis_PNAS_2015	Oligodendrocyte_Progenitor_Cell_Zhang_Neuro_2014	RBC_AI_GeneCardSearch_Hemoglobin_ErythrocyteSpecific
Astrocytes	1.0	0.9	0.9	1.0	0.8	0.8	0.7	0.7	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	
Microglia	0.8	0.7	0.7	0.7	1.0	1.0	1.0	1.0	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	
Endothelial & Mural	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	1.0	1.0	1.0	1.0	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	
Neurons	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	
Mature Oligodendrocytes	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	0.8	

G.

	Astrocyte_AI_Cahoy_Neuro_2008	Astrocyte_AI_Darmanis_PNAS_2015	Astrocyte_AI_Doye_Cell_2008	Astrocyte_AI_Zeisel_Science_2015	Astrocyte_AI_Zhang_Neuro_2014	Microglia_AI_Darmanis_PNAS_2015	Microglia_AI_Zeisel_Science_2015	Microglia_AI_Zhang_Neuro_2014	Endothelial_AI_Darmanis_PNAS_2015	Endothelial_AI_Zeisel_Science_2015	Mural_Pericyte_Zhang_Neuro_2014	Mural_Vascular_Daneman_PLOS_2010	Neuron_AI_Cahoy_Neuro_2008	Neuron_AI_Darmanis_PNAS_2015	Neuron_AI_Doye_Cell_2008	Neuron_AI_Zhang_Neuro_2014	Neuron_Corticospinal_Doye_Cell_2008	Neuron_Corticospinal_Zhang_Neuro_2014	Neuron_GABA_Sugino_NatNeuro_2006	Neuron_Interneuron_CORT_Doye_Cell_2008	Neuron_Interneuron_Zeisel_Science_2015	Neuron_Neuron_CCK_Doye_Cell_2008	Oligodendrocyte_AI_Cahoy_Neuro_2008	Oligodendrocyte_AI_Zeisel_Science_2015	Oligodendrocyte_Mature_Darmanis_PNAS_2015	Oligodendrocyte_Mature_Doye_Cell_2008	Oligodendrocyte_Myelinating_Zhang_Neuro_2014	Oligodendrocyte_Newly-Formed_Zhang_Neuro_2014	Oligodendrocyte_Progenitor_Cell_Darmanis_PNAS_2015	Oligodendrocyte_Progenitor_Cell_Zhang_Neuro_2014	RBC_AI_GeneCardSearch_Hemoglobin_ErythrocyteSpecific							
Astrocytes	1.0	0.925	0.886	0.800	0.363	0.663	0.586	0.624	0.622	0.702	0.677	0.653	0.479	0.616	0.61	0.69	0.45	0.17	0.09	0.444	0.47	0.64	0.56	0.24	0.44	0.11	0.264	0.191	0.127	0.14	0.11	0.246	0.256	0.58	0.796	0.42	0.24	
Microglia	0.886	0.798	0.826	1.0	0.729	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	
Endothelial & Mural	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	1.0	1.0	1.0	1.0	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886
Neurons	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	
Mature Oligodendrocytes	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	0.886	

1646 H.

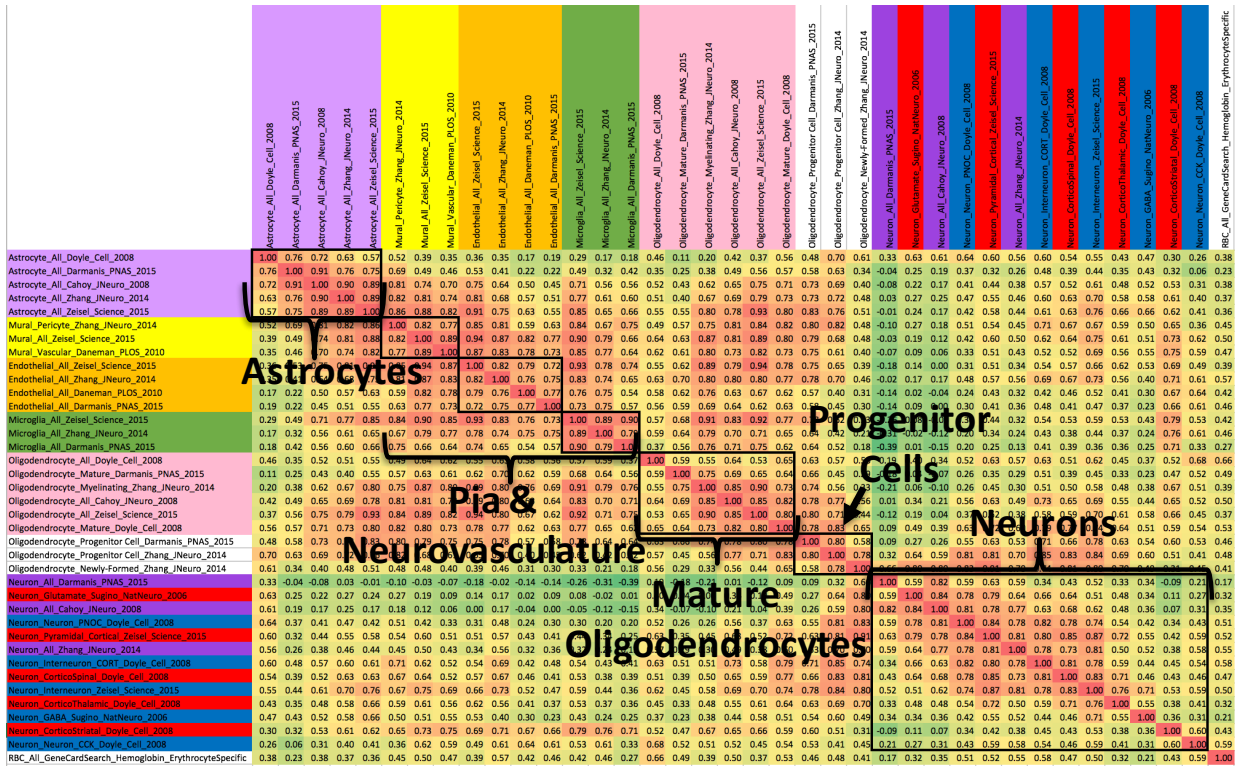


1647

1648 **Suppl. Figure 11. There is a convergence of cell content predictions derived from cell type specific**
 1649 **transcripts identified by different publications. A-B.** Predictions of the relative cell content of our
 1650 human cortical samples (“cell type indices”) for any particular cell type were strongly correlated, even
 1651 when the predictions were based on cell type specific transcripts identified by experiments using very
 1652 different methodology. The examples given above include predictions based on cell type specific
 1653 transcripts originally identified in mouse (x-axis) vs. human (y-axis) tissue. **C.** In contrast, there was a
 1654 strong negative correlation between the predictions for dissimilar cell types, such as neurons and
 1655 astrocytes. **D.** The similarity of different cell type indices in the Pritzker cortical dataset can be visualized
 1656 using a correlation matrix. Within this matrix, correlations can range from a strong negative correlation
 1657 of -1 (blue) to a strong positive correlation of 1 (red), therefore a large block of pink/red correlations is
 1658 indicative of cell type indices that tend to be enriched in the same samples. The axis labels for cell type
 1659 indices representing the same category of cell are color-coded: general neuronal categories are dark
 1660 purple, pyramidal neurons are red, inhibitory interneurons are dark blue, astrocytes are light purple,
 1661 endothelial cells are orange, mural cells are yellow, microglia are green, mature oligodendrocytes are
 1662 pink, and the remaining indices remain white to represent lack of coherent categorization. The number of
 1663 probes included in each index is present in the far left column (also color-coded, with green indicating
 1664 few probes and red indicating many probes). **E-H.** The cell type index correlation matrices for the
 1665 replication cortical datasets: **E.** Narayan et al. (GSE21138), **F.** Lanz et al. (GSE53987), **G.** Barnes et al.
 1666 (GSE21935) **H.** CMC RNA-Seq

1667

Running Head: PREDICTING CELL TYPE BALANCE



1668

1669 **Suppl. Figure 12. The convergence of cell content predictions derived from cell type specific transcripts**
 1670 **within the Allen Brain Atlas dataset. Within the Allen Brain Atlas dataset, the main source of variation**
 1671 **within the data (PC1) was negatively related to all cell types, with an especially strong relationship with**
 1672 **mural cells ($R^2=0.847$), oligodendrocytes ($R^2=0.808$), and endothelial cells ($R^2=0.775$), suggesting that**
 1673 **perhaps the main source of variation in the dataset (PC1) represented general tissue cell density instead**
 1674 **of cell type balance per se. This causes the cell type indices for almost all cell types to be positively**
 1675 **correlated (see below), and therefore this correlation matrix has a slightly different format than Suppl.**
 1676 **Figure 11 and Suppl. Figure 16. The cell types are still coded as in the previous figures, but the**
 1677 **correlation coefficients (R) in the matrix are no longer color coded with blue indicating a negative**
 1678 **correlation and red indicating a positive correlation. Instead, the green to red gradient indicates**
 1679 **increasing percentile from most negative to most positive. The tightest correlations are red, with two**
 1680 **obvious clusters: one cluster representing neurons and another representing glia and support cells. This**
 1681 **differs from the Pritzker dorsolateral prefrontal cortex data, in which oligodendrocytes were found in**
 1682 **their own cluster, perhaps due to a greater variation in the agonal conditions of the subjects (providing**
 1683 **an impetus for the correlated upregulation of astrocytes and neurovasculature) or perhaps due to the**
 1684 **spatial segregation of these cell types within the layered cortex (with an enrichment of vasculature and**
 1685 **astrocytes at the surface of the cortex and an enrichment of white matter under the cortex). Also notable**
 1686 **is the more coherent signature for progenitor cells within the Allen Brain Atlas dataset, perhaps due to**
 1687 **the inclusion of tissue from neurogenic regions.**

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Running Head: PREDICTING CELL TYPE BALANCE

	k=3	k=4	k=6	k=9
Astrocyte_All_Cahoy_JNeuro_2008	1	1	1	1
Astrocyte_All_Darmanis_PNAS_2015	1	2	2	2
Astrocyte_All_Doyle_Cell_2008	1	2	2	2
Astrocyte_All_Zeisel_Science_2015	1	1	1	1
Astrocyte_All_Zhang_JNeuro_2014	1	1	1	1
Endothelial_All_Daneman_PLOS_2010	1	1	1	1
Endothelial_All_Darmanis_PNAS_2015	1	1	1	1
Endothelial_All_Zeisel_Science_2015	1	1	1	1
Endothelial_All_Zhang_JNeuro_2014	1	1	1	1
Microglia_All_Darmanis_PNAS_2015	1	1	3	3
Microglia_All_Zeisel_Science_2015	1	1	1	1
Microglia_All_Zhang_JNeuro_2014	1	1	1	1
Mural_All_Zeisel_Science_2015	1	1	1	1
Mural_Pericyte_Zhang_JNeuro_2014	1	1	1	1
Mural_Vascular_Daneman_PLOS_2010	2	1	1	1
Neuron_All_Cahoy_JNeuro_2008	2	3	4	4
Neuron_All_Darmanis_PNAS_2015	2	3	4	4
Neuron_All_Zhang_JNeuro_2014	2	3	4	5
Neuron_CorticoSpinal_Doyle_Cell_2008	2	3	4	5
Neuron_CorticoStriatum_Doyle_Cell_2008	1	1	3	3
Neuron_CorticoThalamic_Doyle_Cell_2008	2	3	5	6
Neuron_GABA_Sugino_NatNeuro_2006	2	3	4	5
Neuron_Glutamate_Sugino_NatNeuro_2006	2	3	4	5
Neuron_Interneuron_CORT_Doyle_Cell_2008	2	3	4	7
Neuron_Interneuron_Zeisel_Science_2015	2	3	4	5
Neuron_Neuron_CCK_Doyle_Cell_2008	2	3	4	7
Neuron_Neuron_PNOC_Doyle_Cell_2008	2	3	4	7
Neuron_Pyramidal_Cortical_Zeisel_Science_2015	2	3	4	5
Oligodendrocyte_All_Cahoy_JNeuro_2008	3	4	6	8
Oligodendrocyte_All_Doyle_Cell_2008	2	3	5	5
Oligodendrocyte_All_Zeisel_Science_2015	3	4	6	8
Oligodendrocyte_Mature_Darmanis_PNAS_2015	3	4	6	8
Oligodendrocyte_Mature_Doyle_Cell_2008	3	4	6	8
Oligodendrocyte_Myelinating_Zhang_JNeuro_2014	3	4	6	8
Oligodendrocyte_Newly-Formed_Zhang_JNeuro_2014	2	3	4	5
Oligodendrocyte_Progenitor Cell_Darmanis_PNAS_2015	1	1	1	6
Oligodendrocyte_Progenitor Cell_Zhang_JNeuro_2014	2	3	5	6
RBC All GeneCardSearch Hemoglobin ErythrocyteSpecific	2	1	5	9

1692

1693 **Suppl. Figure 13. Consensus clustering indicates the cell type indices clearly cluster into three large**
 1694 **umbrella categories: neurons, oligodendrocytes, and support cells.** The cell type indices were developed
 1695 using cell type specific genes identified by different publications, species, and methodologies, and are
 1696 categorically color-coded in a manner similar to **Suppl. Figure 11**. Each column represents the
 1697 numerical category (cluster) assigned to a cell type index in a k-means clustering algorithm with k
 1698 number of clusters – for example, in the column “k=3”, the algorithm sorted each of the cell type indices
 1699 into 3 clusters based on similarity (defined by Euclidean distance). These 3 clusters are easily identifiable
 1700 as neurons, oligodendrocytes, and support cells. Increasing the number of clusters (k) did not improve the
 1701 ability of the algorithm to detect more specific neuronal subcategories (interneurons, projection neurons)
 1702 or support cell subcategories (astrocytes, endothelial cells, mural cells, microglia), and the immature
 1703 oligodendrocyte indices from different publications showed a notable lack of convergence. The consensus
 1704 clustering was run using 50 bootstraps with a proportion of 0.8 item subsampling and 1.0 feature
 1705 subsampling.

1706

1707

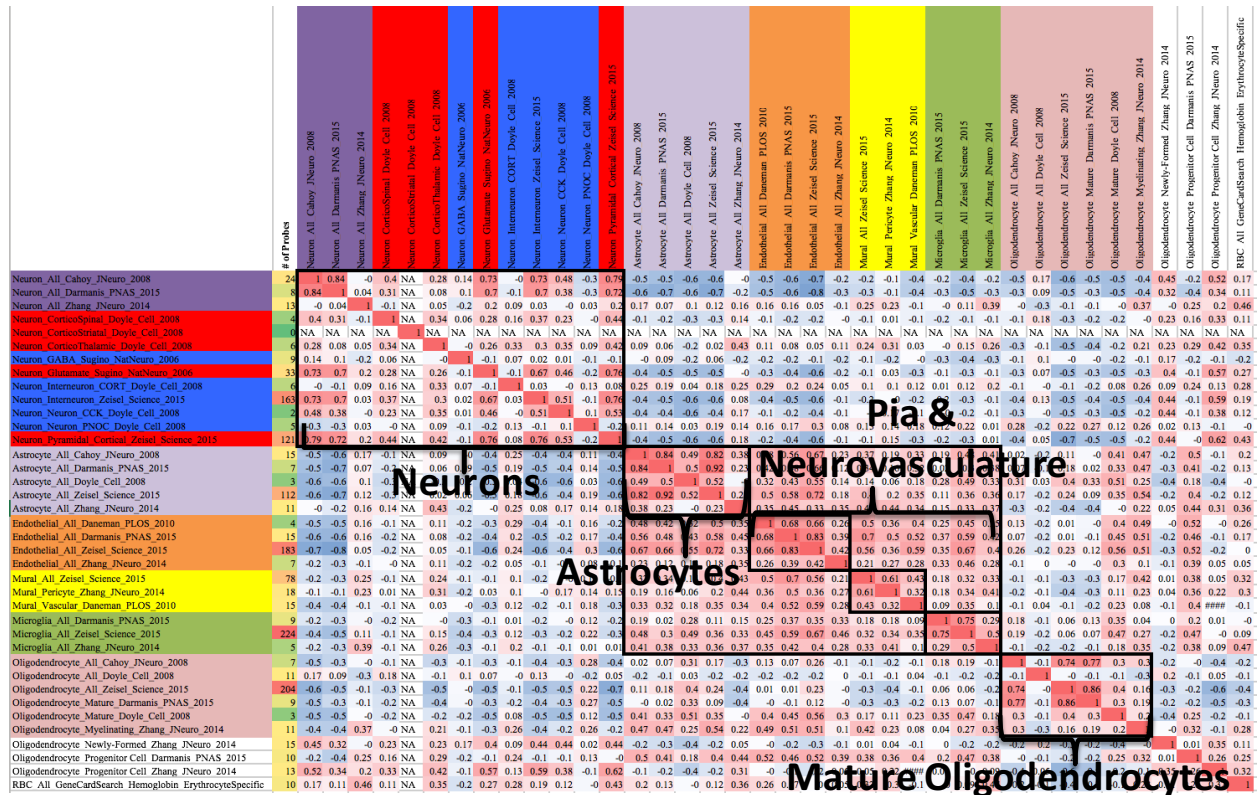
Running Head: PREDICTING CELL TYPE BALANCE

Cell Type Index:	# of Probes (before overlap removal)	# of Probes (after overlap removal)	% remaining:
Astrocyte_All_Cahoy_JNeuro_2008	43	15	0.35
Astrocyte_All_Darmanis_PNAS_2015	13	7	0.54
Astrocyte_All_Doyle_Cell_2008	11	3	0.27
Astrocyte_All_Zeisel_Science_2015	145	112	0.77
Astrocyte_All_Zhang_JNeuro_2014	31	11	0.35
Endothelial_All_Daneman_PLOS_2010	23	4	0.17
Endothelial_All_Darmanis_PNAS_2015	18	15	0.83
Endothelial_All_Zeisel_Science_2015	237	183	0.77
Endothelial_All_Zhang_JNeuro_2014	35	7	0.20
Microglia_All_Darmanis_PNAS_2015	19	9	0.47
Microglia_All_Zeisel_Science_2015	291	224	0.77
Microglia_All_Zhang_JNeuro_2014	24	5	0.21
Mural_All_Zeisel_Science_2015	92	78	0.85
Mural_Pericyte_Zhang_JNeuro_2014	32	18	0.56
Mural_Vascular_Daneman_PLOS_2010	34	15	0.44
Neuron_All_Cahoy_JNeuro_2008	52	24	0.46
Neuron_All_Darmanis_PNAS_2015	15	8	0.53
Neuron_All_Zhang_JNeuro_2014	23	13	0.57
Neuron_CorticoSpinal_Doyle_Cell_2008	18	4	0.22
Neuron_CorticoStriatal_Doyle_Cell_2008	12	0	0.00
Neuron_CorticoThalamic_Doyle_Cell_2008	14	6	0.43
Neuron_GABA_Sugino_NatNeuro_2006	19	9	0.47
Neuron_Glutamate_Sugino_NatNeuro_2006	42	33	0.79
Neuron_Interneuron_CORT_Doyle_Cell_2008	14	6	0.43
Neuron_Interneuron_Zeisel_Science_2015	199	163	0.82
Neuron_Neuron_CCK_Doyle_Cell_2008	10	2	0.20
Neuron_Neuron_PNOC_Doyle_Cell_2008	18	5	0.28
Neuron_Pyramidal_Cortical_Zeisel_Science_2015	154	121	0.79
Oligodendrocyte_All_Cahoy_JNeuro_2008	29	7	0.24
Oligodendrocyte_All_Doyle_Cell_2008	18	11	0.61
Oligodendrocyte_All_Zeisel_Science_2015	255	204	0.80
Oligodendrocyte_Mature_Darmanis_PNAS_2015	15	9	0.60
Oligodendrocyte_Mature_Doyle_Cell_2008	14	3	0.21
Oligodendrocyte_Myelinating_Zhang_JNeuro_2014	28	11	0.39
Oligodendrocyte_Newly-Formed_Zhang_JNeuro_2014	24	15	0.63
Oligodendrocyte_Progenitor_Cell_Darmanis_PNAS_2015	15	10	0.67

1722

1723 **Suppl. Figure 15. Removing probes that represent genes identified as having cell type specific**
 1724 **expression in multiple publications (preparation for the analysis in Suppl. Figure 16).** A summary of
 1725 the full number of probes included in each cell type index before and after removal of the probes that
 1726 overlapped with any other index (color-labeled with pink representing a large number of probes and blue
 1727 representing fewer probes). The percentage of probes retained in the index after full overlap removal is
 1728 also provided, with green indicating a small percentage of probes retained, and red indicating a large
 1729 percentage.

Running Head: PREDICTING CELL TYPE BALANCE



Running Head: PREDICTING CELL TYPE BALANCE

1742

	Freq	Astrocyte	Endothelial	Microglia	Mural	Neuron_ All	Neuron_ Interneuron	Neuron_ Projection	Oligodendrocyte	Oligodendrocyte _Immature	RBC
Astrocyte	243	100%	1%	2%	5%	0%	2%	2%	3%	2%	0%
Endothelial	313	1%	100%	3%	2%	1%	2%	8%	0%	0%	0%
Microglia	334	1%	2%	100%	1%	0%	1%	4%	2%	0%	0%
Mural	158	8%	2%	3%	100%	1%	1%	7%	3%	5%	0%
Neuron_All	90	1%	2%	1%	1%	100%	13%	13%	2%	1%	0%
Neuron_Interneuron	260	1%	2%	1%	1%	5%	100%	3%	1%	0%	0%
Neuron_Projection	240	2%	6%	5%	3%	7%	3%	100%	2%	2%	0%
Oligodendrocyte	359	2%	0%	2%	1%	1%	1%	1%	100%	3%	0%
Oligodendrocyte_Immature	58	7%	2%	0%	9%	2%	2%	9%	16%	100%	0%
RBC	11	0%	9%	0%	0%	0%	0%	0%	0%	0%	100%

1743

1744 *Suppl. Figure 17. For later analyses, individual cell type indices were averaged by primary category,*
 1745 *with any transcripts that overlapped between categories removed. The percent overlap between*
 1746 *transcripts defined as specific to different categories of cell type is illustrated below, color-coded with a*
 1747 *gradient from blue (indicating 0% overlap) to red (indicating 100% overlap). The denominator in the*
 1748 *percentage overlap equation was defined as the cell type category specified by the row. The column on*
 1749 *the far left provides the number of probes included in each cell type category.*

1750

1751

1752

1753 **7.5 Additional figures and results: Cell Type Indices Predict Other Genes Known to Be Cell Type**

1754 **Enriched**

1755 To identify other transcripts important to cell type specific functions in the human cortex, we ran
 1756 a linear model on the signal from each gene probeset in the Pritzker prefrontal cortex microarray dataset
 1757 that included each of the ten consolidated primary cell type indices as well as six co-variables traditionally
 1758 included in the analysis of human brain gene expression data (pH, Agonal Factor, PMI, Age, Gender,
 1759 Diagnosis; **Equation 5**). On average, this model explained 35% of the variation in the data (R^2). Shown in
 1760 **Suppl. Figure 18** are the most significant 10 gene probe sets positively associated with each cell type
 1761 while controlling for the other cell types and co-variables within the model. Additional gene probe sets and
 1762 statistical details can be found in **Suppl. Table 15**.

1763

Running Head: PREDICTING CELL TYPE BALANCE

Astrocyte	Endothelial	Microglia	Mural	Neuron_All	Neuron_Projection	Neuron_Interneuron	Mature Oligodendrocyte	Red Blood Cell (RBC)
NOTCH2	HLA-E	AIF1	TAGLN	VSNL1	PDE2A	TAC3	KLK6	HBD
SDC2	EPAS1	LAPTM5	MYL9	SYT1	USF2	SLC24A3	UGT8	HBB
NTRK2	CLCN7	IRF8	MYH11	SYNGR3	DGKZ	GAD1	MAG	PKLR
CLDN10	CLDN5	FCER1G	CNN1	NEFL	NUAK1	KIT	ELOVL1	PGC
FGFR3	PAK4	PTPRC	MGP	NRXN1	SLC38A7	GAD2	EVI2A	NA
APOE	MYOF	LAIR1	ACTA2	SNAP25	BEGAIN	ERBB4	PLLP	DKK4
EZR	ICAM2	LY86	TP53I11	BCL2L1	KIAA0182	LHX6	MOG	LIPE
SLC1A3	ABCB1	FPR1	COL18A1	MAPK1	KIF21B	SLC6A1	ASPA	SPDEF
CST3	GPR116	C3	TPM2	EEF1A2	PLXNA1	RELN	TF	C19orf57
MLC1	SDPR	ALOX5AP	CRABP1	MEF2C	SLC8A2	ARL4C	MAL	NA

1764

1765 *Suppl. Figure 18. The top 10 transcripts associated with each cell type index include those previously-*
 1766 *identified as cell type enriched in the literature. Transcripts are identified by official gene symbol.*
 1767 *Yellow labels identify transcripts included in the original cell type index, orange transcripts were*
 1768 *previously-identified as cell type enriched in the literature but were not included in the original list of cell*
 1769 *type specific transcripts used to create the index. Additional transcripts and statistical details can be*
 1770 *found in Suppl. Table 15. Please note that not all of the genes listed in the top ten list associated with the*
 1771 *Red Blood Cell index would survive a traditional threshold for false detection threshold ($q < 0.05$).*

1772 Many of the top gene probesets that we found to be related to each of the cell type indices are
 1773 already known to be associated with that cell type in previous publications, validating our methodology.
 1774 Importantly, this is true even when the genes were not included in the original list of cell type specific
 1775 genes used to generate the index. For example, we found that HLA-E (Major Histocompatibility
 1776 Complex, Class I, E) and EPAS1 (endothelial PAS domain protein 1) were both strongly associated with
 1777 our endothelial index, and both are known to be involved in endothelial cell activation (HLA-E, in
 1778 response to immune challenge: (95); EPAS1, in response to lack of oxygen: (96)). NOTCH2 (Notch 2),
 1779 one of the top astrocyte-related genes, promotes astrocytic cell lineage (97), and APOE (Apolipoprotein
 1780 E) is primarily secreted by astrocytes in the central nervous system (98). One of the top interneuron
 1781 genes, LHX6 (LIM Homeobox 6), is specifically enriched in parvalbumin-containing interneurons in the
 1782 human cortex (2). Another top interneuron gene, ERBB4 (Erb-B2 Receptor Tyrosine Kinase 4), controls
 1783 the development of GABA circuitry in the cortex (99). The top neuron-related genes include several
 1784 genes related to synaptic function (SYT1 (Synaptotagmin I), SYNGR3 (Synaptogyrin 3), NRXN1
 1785 (Neurexin 1); <http://www.genecards.org/>). The top projection neuron-related gene, PDE2A

Running Head: PREDICTING CELL TYPE BALANCE

1786 (Phosphodiesterase 2A, CGMP-Stimulated), is preferentially expressed in cortical pyramidal neurons
1787 (100), and KIF21B (Kinesin Family Member 21B) is a kinesin that has been found in the dendrites of
1788 pyramidal neurons (101). We also rediscovered probesets representing genes that were listed as
1789 alternative orthologs to those included in our original cell type specific gene lists (oligodendrocytes:
1790 EVI2A vs.CTD-2370N5.3, microglia: LAIR1 vs. LAIR2, mural cells: COL18A1 vs. COL15A1, ACTA2
1791 vs. ACTG1). Altogether, these results suggest that our cell type indices were associated with the
1792 variability of transcripts in the cortex that represented particular cell types and could re-identify known
1793 cell type specific markers.

1794 As a follow-up analysis, we also outputted a table of the top genes associated with each cell type
1795 within the Allen Brain Atlas dataset (as assessed using the model in **Equation 6**). We found that the
1796 results similarly included a mixture of well-known cell type markers and novel findings (**Suppl. Figure**
1797 **19, Suppl. Table 6**).

1798 ***Equation 6: A model of gene expression for the Allen Brain Atlas dataset, colored to illustrate***
1799 ***subcomponents. The base model (intercept) is presented in green, the cell type indices for the most***
1800 ***prevalent cell types are colored red, and the remaining cell type indices are in purple.***

1801 **Gene Expression (Probeset Signal) =**
1802 $\beta_0 + \beta_1*(\text{Astrocyte}) + \beta_2*(\text{Oligodendrocyte}) + \beta_3*(\text{Microglia}) + \beta_4*(\text{Interneuron}) + \beta_5*(\text{ProjectionNeuron})$
1803 $+ \beta_6*(\text{Endothelial}) + \beta_7*(\text{Neuron_All}) + \beta_8*(\text{Oligodendrocyte_Immature}) + \beta_9*(\text{Mural}) + \beta_{10}*(\text{RBC}) + \epsilon$
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1805

Running Head: PREDICTING CELL TYPE BALANCE

Astrocyte	Endothelial	Microglia	Mural	Neuron_All	Neuron_ Interneuron	Neuron_ Projection	Oligodendrocyte	Oligodendrocyte_ Immature	RBC
GJA1	SDPR	AIF1	SAMD11	SCN2A	GAD1	EGR4	C11orf9	RIMBP2	HBG2
BMPR1B	A_23_P136753	RGS10	PDLIM5	SYT4	GAD2	EXOC6	CPOX	A_32_P176036	HBZ
PON2	LOC390760	LY86	MXRA8	SNAP25	KLHDC5	BAIAP2	FRYL	IL1RAP	TOP2A
PPAP2B	A_32_P234414	TYROBP	ABHD4	NOL4	CRHBP	A_32_P136033	SLC5A11	LARP1	HOXA11-AS
ARHGEF26	CLDN19	GPR34	CBX5	BEX1	UQCRB	SIPA1L1	FAM125B	CHD5	HBG1
FGFR3	PRND	CYBB	APH1B	CRMP1	PPP1CB	A_24_P219094	LAMP2	YWHAG	MELK
TP53BP2	A_32_P181339	ADAM28	A_24_P504050	ADD2	KLHDC1	FAM153A	VWA1	NKD2	PBK
CLDN10	SYNGR1	APBB1P	GBP3	SYN1	CREBL2	A_23_P324706	PKP4	NRG3	A_23_P258666
METTL7A	CLDN2	CD74	STXBPA	A_24_P896765	TADA1	PHYHIP	NCKAP5	STOML1	RHAG
SOX9	CDKN2A	A_32_P223985	DKC1	NRXN1	DPY19L2P3	ITPKA	WIPF1	A_32_P121785	DLGAP5
GPR125	SLC22A8	LST1	PTTG1P	GAP43	SLC32A1	KDM5B	A_24_P540560	ARHGEF4	GAGE2C
SLC7A11	SPTBN2	ALOX5AP	LIMK2	SYT1	PJA1	THRA	FMNL2	NCS1	A_23_P435390
SLC1A3	COL4A2	HLA-DRA	PTPLAD1	SVOP	CTAGE5	FAM153B	RFTN2	CCDC92	A_32_P113110
AGXT2L1	LEFTY2	FCGR1B	RPN1	JPH4	DLX2	VIPR1	HEPACAM	ANO5	PRAMEF10
BBOX1	ITIH5	HLA-DMB	EMILIN1	RAB3C	SERP1	PDZD4	RDX	ODZ2	AHSP
AQP4	ABHD2	CX3CR1	HSPA5	CELF4	RABGEF1	DGKZ	C1orf198	RANGAP1	CENPA
MGST1	FBLN1	P2RY13	A_24_P187407	MEG3	HDGFRP3	SOWAHA	BCAS1	BAI1	A_23_P99653
EFEMP1	KCNJ13	FCGR3A	EIF2AK2	ST8SIA3	ATE1	ATXN7L1	SCD	IDS	TYR
PLTP	KLK11	FYB	COPB2	CUST_422_PI41	TRIQK	STX6	MAP4K4	PTPN14	A_23_P10525
MLC1	A_24_P290114	PTPRC	C22orf42	ACTA1	SLIRP	ANXA11	FGF1	KIF21B	APOH
NTRK2	HPD	BLNK	FTO	KIAA0319	ZZZ3	EFNB2	C10orf90	A_24_P365349	IGJ
A_32_P162494	TMEM86B	LAPTM5	SPARC	L1CAM	SENP6	NPTX1	SLC48A1	NOVA2	HSD17B2
CYBRD1	STEAP4	P2RY12	NUCB2	GABRB3	TATDN3	SYNE1	CLCA4	SLITRK1	CCL20
CNN3	SULT1C2	HPGDS	VAT13	CHGB	A_24_P925241	JPH1	PRRG1	CHST1	APOB
CAMTA1	NOS3	RNASE6	PDIA3	A_32_P77831	CNOT7	MPP7	MAGT1	SHC3	A_24_P334208

1806

1807 *Suppl. Figure 19. The top 25 probes associated with each primary cell type index in the Allen Brain*
 1808 *Atlas dataset. Depicted are the top probes identified in association with each of the primary cell types as*
 1809 *determined by a linear model that included indices for all 10 primary cell types. This model was run*
 1810 *using samples from all 160 brain regions. Similar to the results for the Pritzker dorsolateral prefrontal*
 1811 *cortex data, the genes identified in the Allen Brain Atlas data as having strong relationships with*
 1812 *particular cell types include a mixture of well-known cell type markers and more novel findings.*

1813

1814

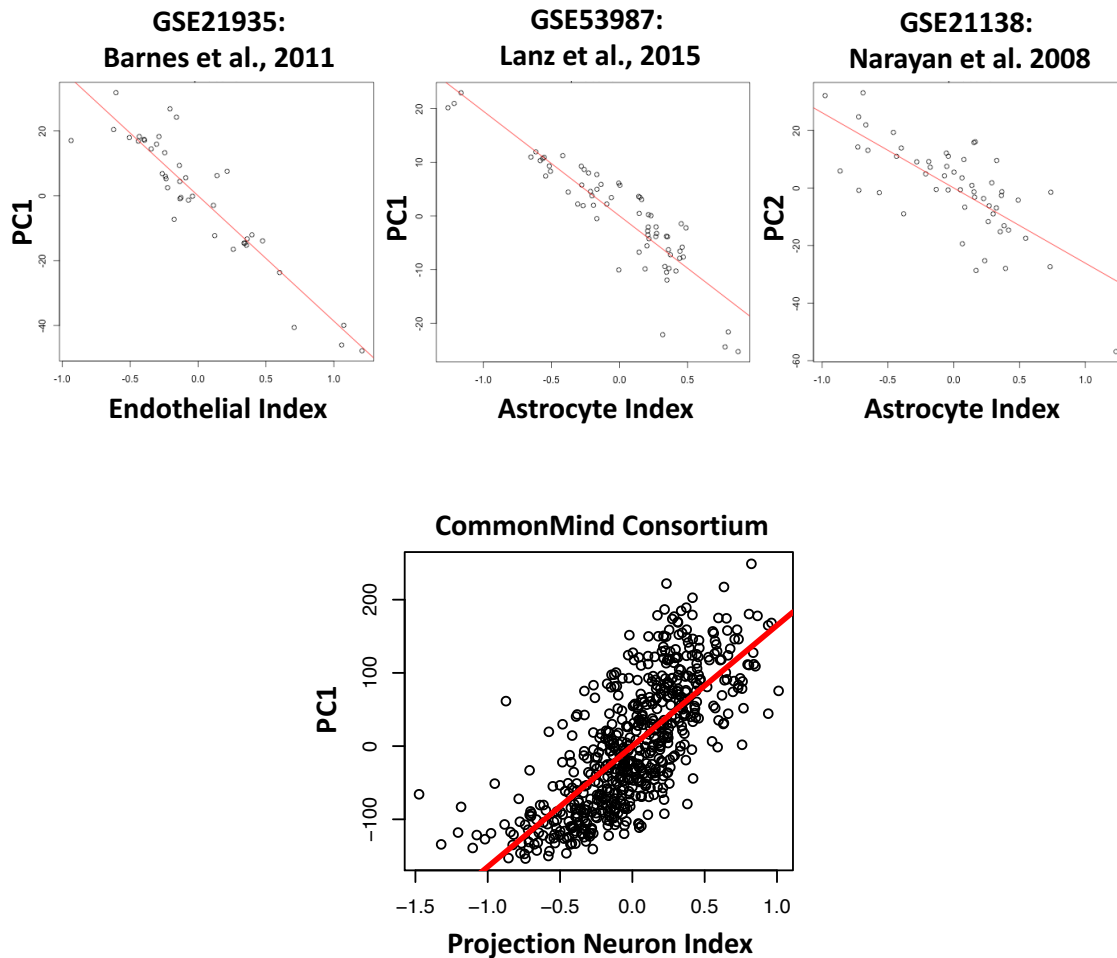
1815 **7.6 Additional figures and results: Inferred Cell Type Composition Explains a Large Percentage of**

1816 **the Sample-Sample Variability in Microarray Data from Macro-Dissected Human Cortical**

1817 **Tissue**

1818

Running Head: PREDICTING CELL TYPE BALANCE



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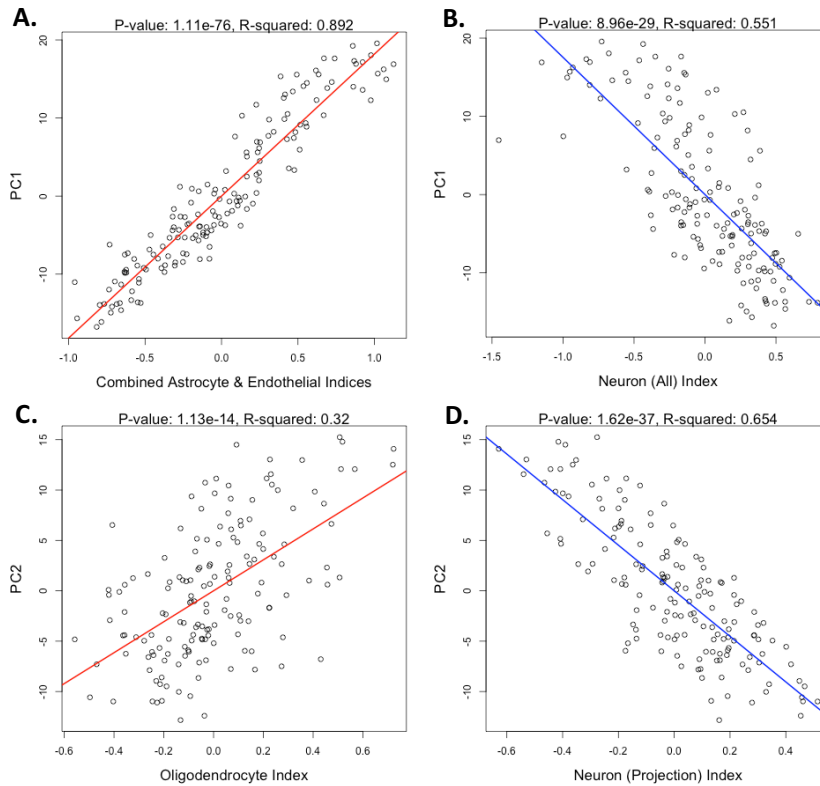
1821

1822 **Suppl. Figure 20. Replication: Cell content predictions explain a large percentage of the variability in**
1823 **microarray and RNA-Seq data derived from the human cortex. The results shown above are from the**
1824 **four other human cortical datasets discussed in the paper. Within the cross-regional Allen Brain Atlas**
1825 **dataset, we also found that the top principal components of variation were overwhelmingly explained by**
1826 **predicted cell type balance, with a model that included all 10 cell type indices accounting for a larger**
1827 **percentage of the variation in the top 4 principal components (PC1: $F(10, 830)=1051, R^2=0.927, p<2.2e-$**
1828 **$16; PC2: F(10, 830)=96.98, R^2=0.539, p<2.2e-16; PC3: F(10, 830)=133.2, R^2= 0.616, p<2.2e-16; PC4:$**
1829 **$F(10, 830)=121.3, R^2= 0.594, p<2.2e-16)$, although the specific relationships sometimes differed from**
1830 **what was seen in the prefrontal cortex.**

1831

1832

Running Head: PREDICTING CELL TYPE BALANCE



1833

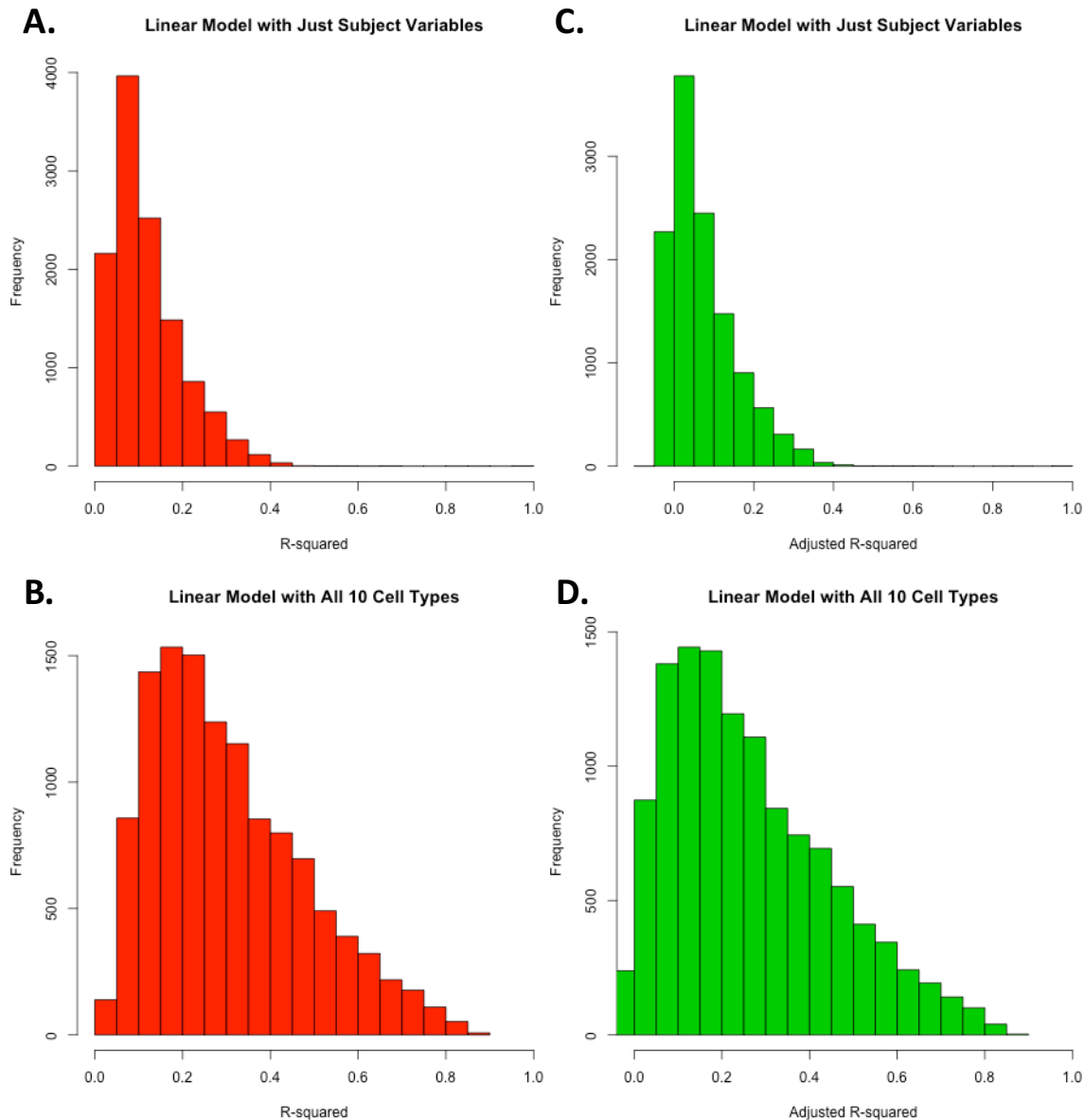
1834 **Suppl. Figure 21. Cell content predictions explain a large percentage of the variability in microarray**
1835 **data from non-cell type specific genes. The results shown here look almost identical to those shown in**
1836 **Figure 5 , except that the principal components analysis in this case was run while excluding all cell type**
1837 **specific genes from the dataset.**

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Running Head: PREDICTING CELL TYPE BALANCE



1841

1842 **Suppl. Figure 22. Predicted cell content accounts for a larger percentage of the variability in the signal**
1843 **from individual probesets than the most commonly examined subject variables.** Shown below are
1844 histograms illustrating the R-squared (A & B) and adjusted R-squared (C & D) for all 11979 probesets in
1845 the Pritzker dorsolateral prefrontal cortex dataset as fit using two linear models: (A & C) A model that
1846 includes diagnosis (MDD, BP, Schiz) and five subject variables commonly used as co-variates in the
1847 analysis of brain microarray data (Brain pH, agonal factor, age, gender, post-mortem interval), (B & D)
1848 A model that includes the consolidated indices for all 10 primary cell types.

1849

1850

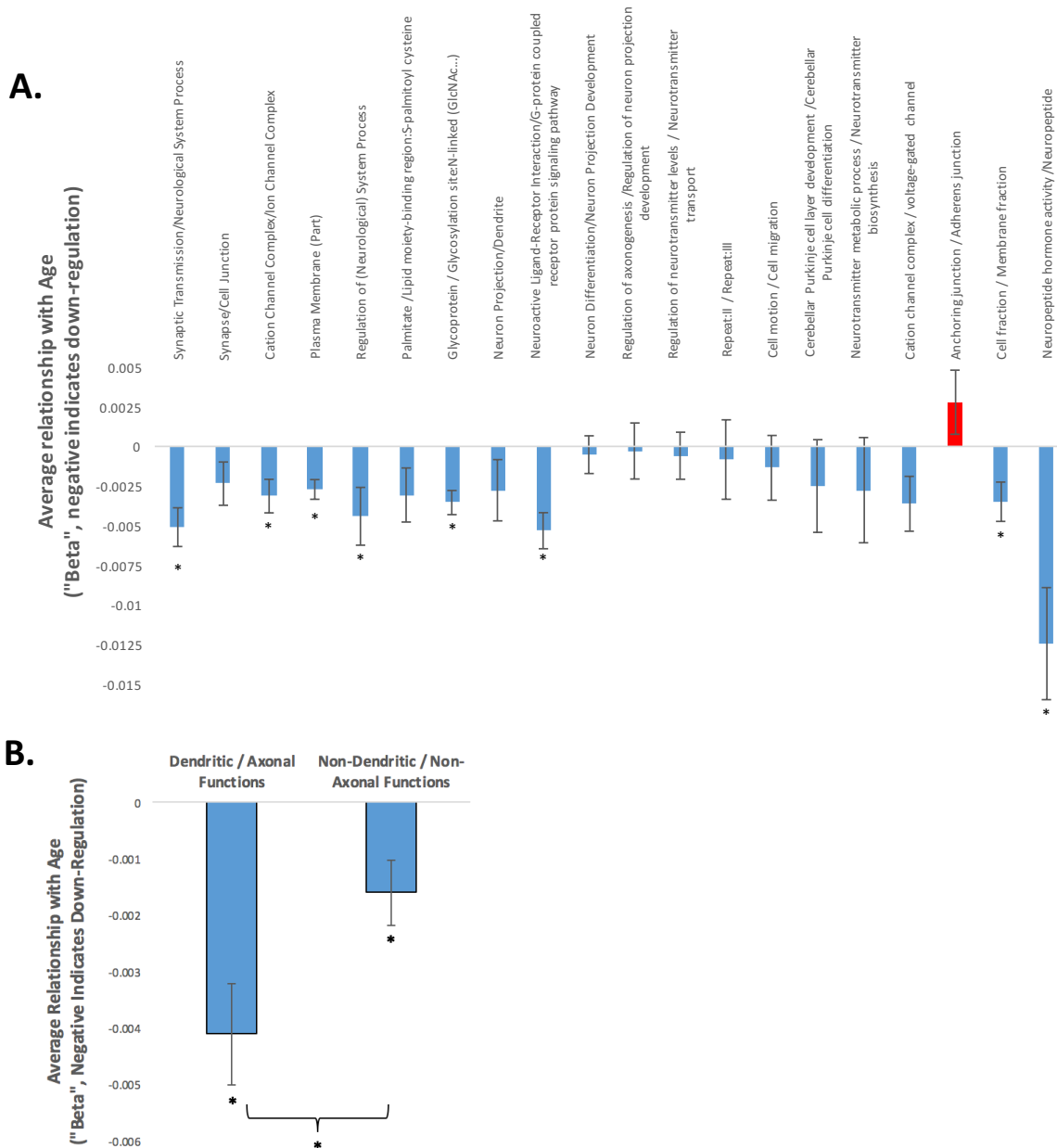
1851

1852

1853 **7.7 Additional figures and results: Discriminating Between Changes in Cell Type Balance and Cell-**

1854 **Type Specific Function**

1855



1856

1857 **Suppl. Figure 23. The predicted decrease in neuronal cell content in relationship to age is unlikely to**
 1858 **be fully explained by synaptic atrophy. Within the list of neuron-specific genes, 240 functional clusters**
 1859 **were identified using DAVID (using the full HT-U133A chip as background). A) The genes in 19 out of**
 1860 **the top 20 functional clusters showed decreased expression with age on average, as determined within a**
 1861 **linear model that controlled for known confounds. Depicted is the average effect of age +/-SE for each**

1862 *cluster (asterisks: $p < 0.05$). Blue represents down-regulation, red is up-regulation. Overall, 76% of all*
1863 *240 functional clusters showed a negative relationship with age on average (Suppl. Table 5). B.) We*
1864 *blindly chose 29 functional clusters that were clearly related to dendritic/axonal functions and 41*
1865 *functional clusters that seemed distinctly unrelated to dendritic/axonal functions. Transcripts from both*
1866 *classifications showed an average decrease in expression with age ($p = 9.197e-05$, $p = 0.008756$,*
1867 *respectively), but the decrease was larger for transcripts associated with dendritic/axonal-related*
1868 *functions ($p = 0.02339$). Depicted is the average effect of age \pm SE for each classification of cluster.*

1869

1870 **7.8 The Top Diagnosis-Related Genes Identified by Models that Include Cell Content Predictions**

1871 **Pinpoint Known Risk Candidates**

1872 Although the inclusion of predicted cell type balance in our model occasionally improved our
1873 ability to detect previously-identified relationships with diagnosis, most relationships still went
1874 undetected in the Pritzker dataset and none of the diagnosis relationships survived standard p-value
1875 corrections for multiple comparisons when included in a full microarray analysis. This could be due to a
1876 variety of factors, including microarray platform and probe sensitivity as well as the possibility that other
1877 cell types in the dataset are showing effects in a competing direction. Therefore, we decided to ask a
1878 complementary question: Of the top diagnosis relationships that we see in our dataset, how many have
1879 been previously observed in the literature? If including predicted cell type balance in our models
1880 improves the signal to noise ratio of our analyses, then we would expect that the top diagnosis-related
1881 genes in our dataset would be more likely to overlap with previous findings. In an attempt to perform this
1882 comparison in an unbiased and efficient manner, we limited our search to PubMed, using as search terms
1883 only the respective human gene symbol and diagnosis (“Schizophrenia”, “Bipolar”, or “Depression”). For
1884 the genes related to MDD in our dataset, we also expanded the search to include two highly-correlated
1885 traits that are more quantifiable and likely to have a genetic basis: “Anxiety” and “Suicide”. Then we
1886 narrowed our results only to studies using human subjects.

1887 Before controlling for cell type, we found that only one of the top 10 genes related to diagnosis
1888 (FOS: (102,103)) or the presence or absence of psychiatric illness (ALDH1A1: (104)) had been
1889 previously noted in the human literature. In contrast, when we used a model that included the five most
1890 prevalent cortical cell types (Model#4), we found that five of the top 10 genes associated with

Running Head: PREDICTING CELL TYPE BALANCE

1891 Schizophrenia had been previously identified in the literature (ARHGEF2: (105), DOC2A: (106), FBX09:
1892 (74), GRM1: (107,108); CEBPA: (89)), and three of the top 10 genes associated with Bipolar Disorder
1893 (ALDH1A1: (104), SNAP25: (109), NRN1:(110); **Suppl. Figure 24, Suppl. Table 9**). This was a
1894 significant enrichment in overlap with the literature when compared to the rate of overlap with the
1895 literature for 100 randomly-selected genes in the dataset subjected to the same protocol (Schizophrenia:
1896 5/10 vs. 7/100, $p=0.0012$; Bipolar: 3/10 vs. 8/100, $p=0.0610$). Likewise, if we replaced diagnosis with a
1897 term representing the general presence or absence of a psychiatric illness, we found that four of the top 10
1898 genes had been previously identified in the literature (ALDH1A1: (104); HBS1L: (4); HIVEP2: (111),
1899 FBX09: (74), **Suppl. Figure 25, Suppl. Table 9**), and 9/10 of the top genes were actually significant with
1900 an $FDR < 0.05$ when using permutation based methods (using the R function `Imp{lmPerm}`,
1901 iterations=9999). The top 10 genes associated with psychiatric illness in models selected using
1902 forward/backward stepwise model selection (criterion=BIC) similarly included five that had been
1903 previously identified in the literature (PRSS16: (112), GRM1: (107,108); ALDH1A1: (104); SNAP25:
1904 (109); HIVEP2: (111), a significant improvement in overlap with the literature than what can be seen in
1905 100 randomly-selected genes in the dataset subjected to the same protocol (Fisher's exact test: 5/10
1906 vs.15/100, $p=0.0168$).

1907 Together, we conclude that including cell content predictions in the analysis of macro-dissected
1908 microarray data can sometimes improve the sensitivity of the assay for detecting altered gene expression
1909 in relationship to psychiatric disease, especially if the dataset is confounded with dissection variation.

1910

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Running Head: PREDICTING CELL TYPE BALANCE

Top Genes Associated with Schizophrenia:

Eq.3: Diagnosis + Confounds:

Probe	Gene Symbol	Beta	Pval	FDR
11330_at	CTRC	-0.13	1.00E-04	4.75E-01
1758_at	DMP1	-0.06	1.37E-04	4.75E-01
10086_at	HHLA1	-0.40	1.70E-04	4.75E-01
23760_at	PITPNB	-0.13	1.96E-04	4.75E-01
55760_at	DHX32	-0.16	2.61E-04	4.75E-01
3397_at	ID1	-0.51	2.73E-04	4.75E-01
1414_at	CRYBB1	-0.12	3.04E-04	4.75E-01
7644_at	ZNF91	-0.29	3.26E-04	4.75E-01
26071_at	FAM127B	0.15	3.84E-04	4.75E-01
4878_at	NPPA	-0.17	3.98E-04	4.75E-01

Eq.6: Diagnosis + 5 Prevalent Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
9181_at	ARHGGEF2	-0.12	3.96E-05	2.66E-01
8448_at	DOC2A	0.18	4.55E-05	2.66E-01
3397_at	ID1	-0.53	6.69E-05	2.66E-01
23760_at	PITPNB	-0.12	8.87E-05	2.66E-01
26268_at	FBXO9	-0.16	2.53E-04	4.48E-01
11330_at	CTRC	-0.10	3.12E-04	4.48E-01
81491_at	GPR63	0.12	4.28E-04	4.48E-01
2911_at	GRM1	0.07	4.71E-04	4.48E-01
55760_at	DHX32	-0.13	4.79E-04	4.48E-01
1050_at	CEBPA	0.15	5.70E-04	4.48E-01

Eq.1: Diagnosis + All Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
3397_at	ID1	-0.54	3.68E-05	2.22E-01
8448_at	DOC2A	0.17	6.26E-05	2.22E-01
9181_at	ARHGGEF2	-0.12	6.78E-05	2.22E-01
23760_at	PITPNB	-0.13	7.41E-05	2.22E-01
5376_at	PMP22	-0.24	1.67E-04	3.64E-01
1414_at	CRYBB1	-0.10	2.34E-04	3.64E-01
4878_at	NPPA	-0.14	2.65E-04	3.64E-01
11330_at	CTRC	-0.10	2.68E-04	3.64E-01
23187_at	PHLDB1	-0.17	4.41E-04	3.64E-01
2263_at	FGFR2	-0.16	4.49E-04	3.64E-01

Top Genes Associated with Bipolar Disorder:

Eq.3: Diagnosis + Confounds:

Probe	Gene Symbol	Beta	Pval	FDR
4725_at	NDUF55	-0.15	7.77E-04	1.00E+00
51042_at	ZNF593	0.16	1.20E-03	1.00E+00
79705_at	LRRK1	0.10	1.64E-03	1.00E+00
10146_at	G3BP1	0.13	1.71E-03	1.00E+00
26664_at	OR7C1	-0.08	1.85E-03	1.00E+00
4677_at	NARS	-0.08	1.89E-03	1.00E+00
2353_at	FOS	-0.63	2.00E-03	1.00E+00
23760_at	PITPNB	-0.10	2.22E-03	1.00E+00
9815_at	GIT2	-0.05	2.44E-03	1.00E+00
7404_at	UTY	0.04	2.87E-03	1.00E+00

Eq.6: Diagnosis + 5 Prevalent Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
216_at	ALDH1A1	-0.37	7.57E-05	9.06E-01
6616_at	SNAP25	-0.20	3.59E-04	1.00E+00
10146_at	G3BP1	0.14	7.61E-04	1.00E+00
4725_at	NDUF55	-0.15	8.07E-04	1.00E+00
51042_at	ZNF593	0.16	1.05E-03	1.00E+00
4677_at	NARS	-0.08	1.07E-03	1.00E+00
8534_at	CHST1	0.21	1.09E-03	1.00E+00
23760_at	PITPNB	-0.10	1.11E-03	1.00E+00
81567_at	TXNDC5	0.14	1.33E-03	1.00E+00
51299_at	NRN1	-0.13	1.42E-03	1.00E+00

Eq.1: Diagnosis + All Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
216_at	ALDH1A1	-0.40	3.05E-05	2.21E-01
6616_at	SNAP25	-0.17	3.69E-05	2.21E-01
8534_at	CHST1	0.22	4.33E-04	9.98E-01
29896_at	TRA2A	-0.15	5.78E-04	9.98E-01
10146_at	G3BP1	0.14	6.58E-04	9.98E-01
90806_at	ANGEL2	-0.09	7.27E-04	9.98E-01
4677_at	NARS	-0.08	1.24E-03	9.98E-01
79705_at	LRRK1	0.10	1.33E-03	9.98E-01
23510_at	KCTD2	0.10	1.34E-03	9.98E-01
81567_at	TXNDC5	0.13	1.41E-03	9.98E-01

Top Genes Associated with MDD:

Eq.3: Diagnosis + Confounds:

Probe	Gene Symbol	Beta	Pval	FDR
23476_at	BRD4	0.12	7.10E-05	4.29E-01
5961_at	PRPH2	0.21	7.16E-05	4.29E-01
9862_at	MED24	0.15	2.08E-04	7.94E-01
10253_at	SPRY2	-0.21	3.20E-04	7.94E-01
10279_at	PRSS16	0.11	3.31E-04	7.94E-01
23493_at	HEY2	-0.15	6.04E-04	9.16E-01
9148_at	NEURL	0.11	6.40E-04	9.16E-01
79570_at	NKAIN1	0.11	1.15E-03	9.16E-01
23163_at	GGA3	0.09	1.59E-03	9.16E-01
139538_at	VENTXP1	-0.03	1.60E-03	9.16E-01

Eq.6: Diagnosis + 5 Prevalent Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
5961_at	PRPH2	0.21	6.96E-05	8.34E-01
23476_at	BRD4	0.11	2.12E-04	9.99E-01
8314_at	BAP1	0.11	2.77E-04	9.99E-01
10279_at	PRSS16	0.10	5.10E-04	9.99E-01
379_at	ARL4D	-0.13	7.57E-04	9.99E-01
9862_at	MED24	0.14	7.86E-04	9.99E-01
79570_at	NKAIN1	0.11	7.97E-04	9.99E-01
9985_at	REC8	0.12	8.57E-04	9.99E-01
2535_at	FZD2	0.08	9.54E-04	9.99E-01
3781_at	KCNN2	-0.14	1.19E-03	9.99E-01

Eq.1: Diagnosis + All Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
23476_at	BRD4	0.12	4.06E-05	4.86E-01
5961_at	PRPH2	0.20	1.20E-04	5.49E-01
2535_at	FZD2	0.08	1.37E-04	5.49E-01
8314_at	BAP1	0.11	4.30E-04	9.99E-01
9985_at	REC8	0.13	5.80E-04	9.99E-01
379_at	ARL4D	-0.13	9.74E-04	9.99E-01
9862_at	MED24	0.13	1.06E-03	9.99E-01
10279_at	PRSS16	0.09	1.29E-03	9.99E-01
10767_at	HBS1L	-0.18	1.33E-03	9.99E-01
79570_at	NKAIN1	0.10	1.40E-03	9.99E-01

1912

1913

1914 **Suppl. Figure 24. When analyzing the full dataset, the top genes associated with diagnosis in models**
 1915 **that include cell content predictions include genes previously identified in the literature. Depicted are**
 1916 **the top 10 genes associated with diagnosis using three different models of increasing complexity, along**
 1917 **with their β 's (magnitude and direction of effect within the model – blue indicates downregulation, pink is**
 1918 **upregulation), nominal p-values, and p-values that have been corrected for false detection rate using the**
 1919 **Benjamini-Hochberg method. Gene symbols that are bolded and highlighted yellow have been previously**
 1920 **detected in the human literature in association with their respective diagnosis in papers identified using**
 1921 **the PubMed search terms “Schizophrenia” (Row 1) and “Bipolar” (Row 2). None of the top genes**
 1922 **associated with major depressive disorder in any of the three models were found to be associated with**
 1923 **“Depression”, “Anxiety”, or “Suicide” on PubMed (Row 3).**

1924

1925

Running Head: PREDICTING CELL TYPE BALANCE

Top Genes Associated with Psychiatric Illness:

Eq.3: Psychiatric + Confounds:

Probe	Gene Symbol	Beta	Pval	FDR
7461_at	CLIP2	0.16	2.18E-04	8.71E-01
26071_at	FAM127B	0.11	2.29E-04	8.71E-01
9862_at	MED24	0.11	3.91E-04	8.71E-01
22864_at	R3HDM2	-0.16	4.43E-04	8.71E-01
55700_at	MAP7D1	0.11	4.61E-04	8.71E-01
216_at	ALDH1A1	-0.30	5.34E-04	8.71E-01
23760_at	PITPNB	-0.08	6.38E-04	8.71E-01
2176_at	FANCC	-0.08	7.08E-04	8.71E-01
64427_at	TTC31	0.08	8.57E-04	8.71E-01
7832_at	BTG2	-0.16	8.87E-04	8.71E-01

Eq.6: Psychiatric + 5 Prevalent Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
379_at	ARL4D	-0.12	1.26E-04	6.37E-01
216_at	ALDH1A1	-0.24	3.02E-04	6.37E-01
7461_at	CLIP2	0.14	4.06E-04	6.37E-01
10767_at	HBS1L	-0.16	4.21E-04	6.37E-01
79778_at	MICALL2	0.09	4.57E-04	6.37E-01
23760_at	PITPNB	-0.08	4.69E-04	6.37E-01
3300_at	DNAJB2	0.12	4.80E-04	6.37E-01
5537_at	PPP6C	-0.07	5.75E-04	6.37E-01
3097_at	HIVEP2	-0.12	5.91E-04	6.37E-01
26268_at	FBXO9	-0.10	7.58E-04	6.37E-01

Eq.1: Psychiatric + All Cell Types & Confounds

Probe	Gene Symbol	Beta	Pval	FDR
379_at	ARL4D	-0.12	9.91E-05	5.12E-01
79778_at	MICALL2	0.08	2.07E-04	5.12E-01
3097_at	HIVEP2	-0.11	2.41E-04	5.12E-01
6616_at	SNAP25	-0.10	3.72E-04	5.12E-01
29896_at	TRA2A	-0.11	3.79E-04	5.12E-01
7461_at	CLIP2	0.14	3.79E-04	5.12E-01
2535_at	FZD2	0.06	4.06E-04	5.12E-01
216_at	ALDH1A1	-0.22	4.33E-04	5.12E-01
6604_at	SMARCD3	0.12	4.46E-04	5.12E-01
8534_at	CHST1	0.16	4.47E-04	5.12E-01

Stepwise Regression:

Top Genes Associated with Psychiatric Illness:

Probe	Gene Symbol	Beta	Pval
9862_at	MED24	0.13	1.83E-05
7461_at	CLIP2	0.17	4.74E-05
10279_at	PRSS16	0.10	8.86E-05
2911_at	GRM1	0.05	1.11E-04
216_at	ALDH1A1	-0.23	1.28E-04
379_at	ARL4D	-0.11	1.37E-04
6616_at	SNAP25	-0.11	1.39E-04
8534_at	CHST1	0.16	1.45E-04
3097_at	HIVEP2	-0.12	1.53E-04
64427_at	TTC31	0.08	1.67E-04

Top Genes Associated with Suicide:

Probe	Gene Symbol	Beta	Pval
8526_at	DGKE	0.035	1.81E-05
64718_at	UNKL	0.106	2.40E-05
65998_at	C11orf95	0.17	6.56E-05
84617_at	TUBB6	0.162	9.41E-05
4752_at	NEK3	-0.08	1.72E-04
9640_at	ZNF592	0.158	2.27E-04
25940_at	FAM98A	-0.11	3.00E-04
80176_at	SPSB1	0.087	3.01E-04
1051_at	CEBPB	0.249	4.04E-04
50515_at	CHST11	0.069	4.18E-04

1926

1927 *Suppl. Figure 25. When analyzing the full dataset, the top genes associated with psychiatric illness in*
 1928 *models that include cell content predictions include genes previously identified in the literature.*
 1929 *Depicted are the top 10 genes associated with psychiatric illness using three different models of*
 1930 *increasing complexity, or associated with psychiatric illness or suicide in models chosen using stepwise*
 1931 *regression. Notably, the results from stepwise regression for the diagnosis term are not included in this*
 1932 *figure because the term was only included in the model for eight genes total (DHX32, ID1, CSRP1,*
 1933 *AKR1B10, TBPL1, HIST1H4F, SETD3, GAL). Included are the β 's (magnitude and direction of effect*
 1934 *within the model – blue indicates downregulation, pink is upregulation), nominal p-values, and p-values*
 1935 *that have been corrected for false detection rate using the Benjamini-Hochberg method. Note that the p-*
 1936 *values associated with stepwise regression are likely to be optimistic due to overfitting. Gene symbols*
 1937 *that are bolded and highlighted yellow have been previously detected in the human literature in*
 1938 *association with their respective diagnosis in papers identified using the PubMed search terms*
 1939 *“Schizophrenia”, “Bipolar”, “Depression”, “Anxiety”, or “Suicide”.*

1940

1941

1942

1943

Running Head: PREDICTING CELL TYPE BALANCE

1944

Eq.8 Interaction Terms: Psych * Prevalent Cell Types:

Psychiatric:

Probe	Gene Symbol	Beta	Pval	FDR
379_at	ARL4D	-0.12	1.54E-04	6.53E-01
3097_at	HIVEP2	-0.13	2.43E-04	6.53E-01
216_at	ALDH1A1	-0.24	2.68E-04	6.53E-01
10767_at	HBS1L	-0.16	2.91E-04	6.53E-01
4677_at	NARS	-0.06	3.83E-04	6.53E-01
7461_at	CLIP2	0.14	4.48E-04	6.53E-01
3300_at	DNAJB2	0.125	4.76E-04	6.53E-01
23760_at	PITPNB	-0.08	5.01E-04	6.53E-01
79778_at	MICALL2	0.086	5.34E-04	6.53E-01
5537_at	PPP6C	-0.07	6.47E-04	6.53E-01

Psychiatric*Astrocyte

Probe	Gene Symbol	Beta	Pval	FDR
28958_at	CCDC56	-0.46	1.75E-05	1.87E-01
23305_at	ACSL6	0.617	3.12E-05	1.87E-01
9929_at	JOSD1	0.438	8.44E-05	3.37E-01
55751_at	TMEM184C	0.312	1.64E-04	4.92E-01
64794_at	DDX31	-0.37	3.55E-04	6.65E-01
58525_at	WIZ	-0.23	3.69E-04	6.65E-01
8492_at	PRSS12	-0.3	3.89E-04	6.65E-01
3709_at	ITPR2	0.22	4.46E-04	6.68E-01
81890_at	QTRT1	-0.48	5.21E-04	6.93E-01
9514_at	GAL3ST1	0.339	6.30E-04	7.55E-01

Psychiatric*Microglia

Probe	Gene Symbol	Beta	Pval	FDR
55308_at	DDX19A	0.324	5.07E-05	3.26E-01
6351_at	CCL4	-0.58	5.44E-05	3.26E-01
23305_at	ACSL6	-0.51	4.48E-04	9.96E-01
79953_at	TMEM90B	-0.39	4.57E-04	9.96E-01
116496_at	FAM129A	0.264	5.55E-04	9.96E-01
26539_at	OR10H1	-0.82	6.99E-04	9.96E-01
9278_at	ZBTB22	-0.32	9.99E-04	9.96E-01
11326_at	VSIG4	0.491	1.05E-03	9.96E-01
1415_at	CRYBB2	-0.34	1.07E-03	9.96E-01
2615_at	LRRC32	-0.45	1.31E-03	9.96E-01

Psychiatric*Interneuron

Probe	Gene Symbol	Beta	Pval	FDR
3638_at	INSIG1	-1.57	3.76E-05	1.91E-01
56937_at	PMEPA1	-0.95	7.23E-05	1.91E-01
50835_at	TAS2R9	-0.4	8.67E-05	1.91E-01
39_at	ACAT2	-1.77	1.06E-04	1.91E-01
3606_at	IL18	-0.31	1.06E-04	1.91E-01
10473_at	HMGN4	0.846	1.24E-04	1.91E-01
50489_at	CD207	0.655	1.34E-04	1.91E-01
79053_at	ALG8	1.179	1.37E-04	1.91E-01
4693_at	NDP	1.462	1.43E-04	1.91E-01
253943_at	YTHDF3	1.522	1.94E-04	2.32E-01

Psychiatric*Projection Neuron

Probe	Gene Symbol	Beta	Pval	FDR
10473_at	HMGN4	-0.56	2.29E-04	9.96E-01
56606_at	SLC2A9	-0.54	3.31E-04	9.96E-01
652_at	BMP4	0.585	3.57E-04	9.96E-01
3586_at	IL10	0.255	4.63E-04	9.96E-01
4649_at	MYO9A	1.102	5.58E-04	9.96E-01
23288_at	IQCE	0.347	5.82E-04	9.96E-01
23385_at	NCSTN	-0.55	6.35E-04	9.96E-01
9582_at	APOBEC3B	0.174	8.10E-04	9.96E-01
50807_at	ASAP1	-0.76	1.01E-03	9.96E-01
80146_at	UXS1	-0.67	1.25E-03	9.96E-01

Psychiatric*Oligodendrocyte

Probe	Gene Symbol	Beta	Pval	FDR
11184_at	MAP4K1	-0.39	2.21E-04	1.00E+00
5936_at	RBM4	0.621	4.58E-04	1.00E+00
10432_at	RBM14	0.513	6.94E-04	1.00E+00
51073_at	MRPL4	-0.39	1.02E-03	1.00E+00
8552_at	INE1	-0.34	1.51E-03	1.00E+00
22934_at	RPIA	0.355	1.56E-03	1.00E+00
10351_at	ABCA8	1.005	1.63E-03	1.00E+00
10428_at	CFDP1	0.487	1.74E-03	1.00E+00
23180_at	RFTN1	0.723	1.86E-03	1.00E+00
58488_at	PCTP	0.289	1.95E-03	1.00E+00

1945

1946

1947 **Suppl. Figure 26. When analyzing the full dataset using a model that includes Psychiatric Illness*Cell**
 1948 **Type interaction terms, the top genes associated with psychiatric illness include genes previously**
 1949 **identified in the literature. Depicted are the top 10 genes associated with psychiatric illness and its**
 1950 **interaction with the five most prevalent cell types in the cortex using the model in Equation 6:**

1951 **Equation 7:**

1952 $Gene\ Expression = \beta_0 + \beta_1*(Astrocyte\ Index) +$
 1953 $+ \beta_2*(Microglia\ Index) + \beta_3*(Neuron_Interneuron\ Index) + \beta_4*(Neuron_Projection\ Neuron$
 1954 $Index) + \beta_5*(Oligodendrocyte\ Index) + \beta_6*(Brain\ pH) + \beta_7*(Agonal\ Factor) + \beta_8*(PMI) +$
 1955 $\beta_9*(Age) + \beta_{10}*(Gender) + \beta_{11}*(Psychiatric\ Illness) + \beta_{12}*(Psychiatric\ Illness)*(Astrocyte$
 1956 $Index) + \beta_{13}*(Psychiatric\ Illness)*(Microglia\ Index) + \beta_{14}*(Psychiatric$
 1957 $Illness)*(Neuron_Interneuron\ Index) + \beta_{15}*(Psychiatric\ Illness)*(Neuron_Projection\ Neuron$
 1958 $Index) + \beta_{17}*(Psychiatric\ Illness)*(Oligodendrocyte\ Index) + \epsilon$

1959
 1960 *Included are the β's (magnitude and direction of effect within the model – blue indicates downregulation,*
 1961 *pink is upregulation), nominal p-values, and p-values that have been corrected for false detection rate*
 1962 *using the Benjamini-Hochberg method. The number of top genes that were found to be previously-*

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1963 *identified in literature does not significantly surpass what was observed in a group of 100 randomly*
1964 *selected genes from our dataset (14/60 vs. 15/100).*
1965

1966 **8. Supplementary Tables**

1967 **Suppl. Table 1. Master Database of Cortical Cell Type Specific Gene Expression.** The attached excel
1968 document contains a single spreadsheet listing the genes defined as having cell type specific expression in
1969 our manuscript, including the species, age of the subjects, and brain region from which the cells were
1970 purified, the platform used to measure transcript, the statistical criteria and comparison cell types used to
1971 define “cell type specific expression”, the gene symbol or orthologous gene symbol in mouse/human
1972 (depending on the species used in the original experiment), and citation. If a gene was identified as
1973 having cell type specific expression in multiple experiments, there is an entry for each experiment – thus
1974 the full 3383 rows included in the spreadsheet do not represent 3383 individual cell type specific genes. A
1975 web-version of this spreadsheet kept interactively up-to-date can be found at
1976 <https://sites.google.com/a/umich.edu/megan-hastings-hagenauer/home/cell-type-analysis>.

1977 **Suppl. Table 2. Microarray data spanning 160 human brain regions downloaded from the Allen Brain**
1978 **Atlas.** Included in this excel file are three worksheets. The first includes all of the sample information,
1979 including the subject identifier and brain region. The second includes all of the probe information.
1980 Finally, the third includes the relative expression for each probe for each sample (z-score), including the
1981 official gene symbol, Entrez gene ID, and gene name. Additional information about the human
1982 microarray dataset can be found on the Allen Brain Atlas website.

1983 **Suppl. Table 3. The average cell type indices for all 160 brain regions included in the Allen Brain Atlas**
1984 **dataset.** This excel file contains two worksheets. The first includes the average cell type index for 10
1985 primary cell types for all 160 brain regions included in the Allen Brain Atlas. More detail about those
1986 brain regions can be found in the first worksheet (Columns_Sample Info) in **Suppl. Table 2**. The second
1987 spreadsheet contains the standard error (SE) for the averages in the first worksheet.

1988 **Suppl. Table 4. Output for the analyses of cell type vs. subject variables for all datasets.** The first
1989 spreadsheet provides the output from the meta-analysis for each cell type vs. subject variable
1990 combination (“b”= the estimated effect, provided in the units for the variable – e.g., the effect of one
1991 year of age, or the effect of one hour of PMI; “SE”= standard error, “p-value”= nominal p-value,
1992 “BH_adj_P-value (q-value)”= the p-value corrected for multiple comparisons). The second spreadsheet
1993 includes the T-statistics for all cell type vs. subject variable combinations for all datasets.

1994 **Suppl. Table 5. Functions associated with genes identified as having neuron-specific expression.** The
1995 first column of the excel spreadsheet is a list of general physiological functions that were identified by
1996 DAVID as associated with our list of neuron-specific genes (relative to the full list of probesets included
1997 in the microarray). We used the functional cluster option in DAVID because it prevents multiple functions
1998 that share a large subset of overlapping genes from dominating the results. We named each cluster by the
1999 top two functions included in it. The second column of the spreadsheet indicates whether an experimenter
2000 blindly categorized the functional cluster as being clearly related or unrelated to synaptic function. The
2001 “Mean Fold Enrichment” column indicates how well on average each of the functions within that cluster
2002 were associated with our list of neuron-specific genes. The next three columns (Top p-value, Top
2003 Bonferroni-corrected p-value, and top BH (Benjamini-Hochberg)-corrected p-value) indicate the
2004 statistical strength of the association between the top function within that cluster and our list of neuron-
2005 specific genes. The number of genes from each functional cluster included in our results is listed in
2006 column G. The next few columns indicate the strength of the relationship between the functional cluster
2007 and age. Columns H-J indicate the mean, standard deviation, and standard error, for the betas for Age
2008 for each gene included in the cluster. The betas indicate the strength and direction of the association with
2009 Age as determined within a larger linear model controlling for known confounds (pH, PMI, gender,
2010 agonal factor). Columns K-M indicate whether, on average, the age-related betas for the genes in that
2011 cluster are statistically different from 0 as determined by a Welch’s t-test (t-stat, df, p-value). The final

2012 *column indicates what percentage of the genes included in the cluster have a negative relationship (β)*
2013 *with age.*

2014 **Suppl. Table 6. A .gmt file created using our database of cell type specific genes for use with Gene Set**
2015 **Enrichment Analysis (GSEA). This file should be in the correct format for usage with either GSEA**
2016 **(<http://software.broadinstitute.org/gsea/index.jsp>) or fGSEA.**

2017 **Suppl. Table 7. Performing Gene Set Enrichment Analysis using a .gmt that includes traditional**
2018 **functional gene sets and cell type specific gene lists indicates that cell type specific gene sets are**
2019 **enriched for effects related to a wide variety of subject variables.** *Gene set enrichment analysis was*
2020 *performed using the results from a differential expression analysis performed on the Pritzker dataset*
2021 *using a model that included diagnosis, pH, agonal factor, age, PMI, and sex. The gene set enrichment*
2022 *results for each variable is included as its own worksheet in the file.*

2023 **Suppl. Table 8. Previously-identified relationships between gene expression and psychiatric illness in**
2024 **the human cortex in either particular cell types or macro-dissected cortex.** *We used this database of*
2025 *previously-identified effects to determine whether controlling for cell type while performing differential*
2026 *expression analyses increased our ability to observe previously-documented effects.*

2027 **Suppl. Table 9. The relationship between diagnosis and all probesets in the Pritzker Dorsolateral**
2028 **Prefrontal Cortex dataset as assessed using models of increasing complexity.** *For all probesets in the*
2029 *dataset, the spreadsheets for Model #2 and Model#4 include the β for all variables in the model (“Beta”:*
2030 *magnitude and direction of the association, with positive associations labeled pink and negative*
2031 *associations labeled blue), the p-value (“Pval_nominal”) and the p-value adjusted for multiple*
2032 *comparisons using the Benjamini-Hochberg method (“BH_Adj”), both labeled with green indicating*
2033 *more significant relationships and red indicating less significant relationships. There are also summary*
2034 *spreadsheets that include just the results for Bipolar Disorder and Schizophrenia for Models#1-5. In*
2035 *these spreadsheets, the formatting is a little different: T-statistics are provided, the β is called “LogFC”,*
2036 *the BH_Adj p-value is called “adj.P.Val”.*

2037 **Suppl. Table 10. The relationship between diagnosis and all genes in the CMC RNA-Seq dataset as**
2038 **assessed using models of increasing complexity.** *There are two summary spreadsheets that include the*
2039 *results for Bipolar Disorder and Schizophrenia for Models#1-5. For all genes in the dataset, each*
2040 *spreadsheet includes the β (“LogFC”: magnitude and direction of the association), the T-statistic, the p-*
2041 *value (“Pval_nominal”) and the p-value adjusted for multiple comparisons using the Benjamini-*
2042 *Hochberg method (“adj.P.Val”) for the effect of diagnosis in each model (#M1-M5).*

2043 **Suppl. Table 11. The relationship between diagnosis and all probesets in the Barnes et al. microarray**
2044 **dataset as assessed using models of increasing complexity.** *There is a summary spreadsheet that*
2045 *includes the results for Schizophrenia for Models#1-5. For all probesets in the dataset, each spreadsheet*
2046 *includes the β (“LogFC”: magnitude and direction of the association), the T-statistic, the p-value*
2047 *(“Pval_nominal”) and the p-value adjusted for multiple comparisons using the Benjamini-Hochberg*
2048 *method (“adj.P.Val”) for the effect of diagnosis in each model (#M1-M5).*

2049 **Suppl. Table 12. The relationship between diagnosis and all probesets in the Lanz et al. microarray**
2050 **dataset as assessed using models of increasing complexity.** *There are two summary spreadsheets that*
2051 *include the results for Bipolar Disorder and Schizophrenia for Models#1-5. For all probesets in the*
2052 *dataset, each spreadsheet includes the β (“LogFC”: magnitude and direction of the association), the T-*
2053 *statistic, the p-value (“Pval_nominal”) and the p-value adjusted for multiple comparisons using the*
2054 *Benjamini-Hochberg method (“adj.P.Val”) for the effect of diagnosis in each model (#M1-M5).*

2055 **Suppl. Table 13. The relationship between diagnosis and all probesets in the Narayan et al. microarray**
2056 **dataset as assessed using models of increasing complexity.** There is a summary spreadsheet that
2057 includes the results for Schizophrenia for Models#1-5. For all probesets in the dataset, each spreadsheet
2058 includes the β (“LogFC”: magnitude and direction of the association), the T-statistic, the p-value
2059 (“Pval_nominal”) and the p-value adjusted for multiple comparisons using the Benjamini-Hochberg
2060 method (“adj.P.Val”) for the effect of diagnosis in each model (#M1-M5).

2061 **Suppl. Table 14. Sample demographics for the Pritzker Consortium Dorsolateral Prefrontal Cortex**
2062 **Affymetrix microarray data.**

2063 **Suppl. Table 15. The relationship between each cell type index and all probes in the Pritzker**
2064 **Dorsolateral Prefrontal Cortex dataset.** The attached excel document (.xlsx) contains multiple
2065 spreadsheets. The first spreadsheet (“Methods”) contains a brief summary of the methods used to
2066 evaluate the relationship between the cell type indices and expression of each probe in the dataset (also
2067 discussed in the body of the manuscript). The second spreadsheet (“GeneByCellType_DF”) contains the
2068 statistical output associated with all cell type index terms in the linear model for all probes in the dataset,
2069 including the β (“Beta”: magnitude and direction of the association, with positive associations labeled
2070 pink and negative associations labeled blue), the p-value from the original model (“Pval”) and the p-
2071 value adjusted for multiple comparisons using the Benjamini-Hochberg method (“AdjP”), both labeled
2072 with green indicating more significant relationships and red indicating less significant relationships. All
2073 other spreadsheets contain the top 100 probes positively associated with each cell type index, including
2074 each of the statistical outputs presented in the full “GeneByCellType_DF” summary spreadsheet, as well
2075 as a column “CellTypeSpecific” which indicates whether the probe was included in one of the original
2076 cell type indices (1=included, 0=not included).

2077 **Suppl. Table 16. The relationship between each cell type index and all probes in the Allen Brain Atlas**
2078 **dataset.** Depicted are the β (magnitude and direction) and p-values for the relationship between the
2079 expression for each probe and each primary cell type across samples from all 160 brain regions as
2080 determined in a large linear model that includes all 10 primary cell types. Please note that the p-values in
2081 this spreadsheet have not been corrected for multiple comparisons. Additional information about the
2082 probes can be found in **Suppl. Table 2.**

2083
2084
2085
2086