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## Evaluating brain cell marker genes based on differential gene expression and coexpression — Source link [2]

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# 1 Evaluating brain cell marker genes based on differential

## 2 gene expression and co-expression

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11

## 12 Abstract

Reliable identification of brain cell types is necessary for studying brain cell 13 biology. Many brain cell marker genes have been proposed, but their reliability 14 has not been fully validated. We evaluated 540 commonly-used marker genes 15 of astrocyte, microglia, neuron, and oligodendrocyte with six transcriptome and 16 proteome datasets from purified human and mouse brain cells (n=125). By 17 setting new criteria of cell-specific fold change, we identified 22 gold standard 18 marker genes (GSM) with stable cell-specific expression. Our results call into 19 question the specificity of many proposed marker genes. We used two single-20 cell transcriptome datasets from human and mouse brains to explore the co-21 expression of marker genes (n=3337). The mouse co-expression modules were 22 perfectly preserved in human transcriptome, but the reverse was not. Also, we 23 proposed new criteria for identifying marker genes based on both differential 24 expression and co-expression data. We identified 16 novel candidate marker 25 genes (NCM) for mouse and 18 for human independently, which have the 26 potential for use in cell sorting or other tagging techniques. We validated the 27 specificity of GSM and NCM by in-silico deconvolution analysis. Our systematic 28 evaluation provides a list of credible marker genes to facilitate correct cell 29 identification, cell labeling, and cell function studies. 30

31

## 32 Introduction

The human brain is a heterogeneous organ with numerous cell types. It has billions of cells including half neurons and half glia<sup>1</sup>. The major classes of glia are astrocyte, microglia and oligodendrocyte. Identifying these cell types is important because it would permit the brain to be understood in greater detail and would be especially useful for studying cellular contributions to the psychiatric disorders. A critical need in neuroscience research, is to develop methods to reliably identify specific brain cell types.

40 A strategy that has been employed to identify specific cell types is the

development of marker genes, which are sets of genes that express specifically 41 in a cell type. Thousands of genes have been proposed as marker genes<sup>2</sup>. One 42 well-known marker gene, RBFOX3 (gene of NeuN), is only expressed in nuclei 43 of most neuronal cell types<sup>3</sup>. Marker genes can be used in several applications. 44 Protein products of marker genes can be used to label different cell types, which 45 may be used in fluorescence activated cell sorting (FACS). Marker genes also 46 can be used to determine cell composition in bulk tissue samples. A 47 computational method known as supervised deconvolution was developed to 48 infer cell proportions in bulk tissue samples based on the expression of marker 49 genes<sup>4-6</sup>. This method has been applied to studying the composition of bulk 50 brain samples<sup>7,8</sup>. High specificity of marker genes is critical for generating 51 reliable results in all of these applications. 52

Differential gene expression (DGE) analysis of transcriptome or proteome 53 data is the most straightforward way to define the specificity of marker genes<sup>9-</sup> 54 <sup>15</sup>. One of the drawbacks of DGE is that the outcomes is study-dependent. The 55 outcomes are affected by many factors such as species, cell or tissue source, 56 and the data generation platform. Human and mouse genomes are 80% 57 orthologous<sup>16</sup>, but differences in gene expression between species are often 58 greater than those between tissues within one species<sup>17</sup>. Within a species, cells 59 isolated from primary culture or acutely from tissue showed different gene 60 expression patterns<sup>18</sup>. Also, the expression estimates of the marker may vary 61 considerably depending on whether mRNA or protein is measured. The 62 statistical variation in transcriptome only explained 40% of the statistical 63 variation in protein level<sup>19</sup>. Besides these biological confounders, the 64 experimental platforms used to quantify gene expression level may also impact 65 marker gene selection. RNA-Seq provides a larger dynamic range for the 66 detection of transcripts and has less background noise, resulting in RNA-Seq 67 being more sensitive in calling cell type-specific genes than microarray 68 platforms<sup>20</sup>. Another weakness of DGE is that relationships among marker 69 genes are not considered in the analysis. Groups of marker genes are often 70 used to describe a cell type, and marker genes work with each other to execute 71 functions in specific cell type. The relationship between marker genes 72 represents their coordinated functions, specificities, and expressions. In DGE 73 analysis, marker genes are defined independently, and the relationship among 74 them is ignored. 75

Co-expression (COE) is a method of identifying interactions among genes by 76 assigning genes with similar expression patterns into a module<sup>21,22</sup>. There was 77 study reported that the co-expression modules in brain enriched cell type 78 marker genes<sup>23</sup>. So it suggested that the co-expression can detected the cell 79 type-specific marker genes, even in the heterogenous samples. The module 80 formed by marker genes indicates their coordinated functions and specificities 81 for a cell type. The correlation of genes with cell type-specific module suggests 82 it's cell specificity. COE has the potential to systematically capture marker 83 genes group that DGE cannot. 84

In this study, we evaluated the specificity of 540 published brain cell marker 85 genes and discovered novel marker genes by DGE and COE analyses. We 86 used six datasets containing transcriptome and proteome data from purified 87 astrocytes, microglia, neurons and oligodendrocytes from both mouse and 88 human brains. We identified 22 brain cell marker genes out of the 540 89 candidates, referred as gold-standard marker genes (GSM), that specifically 90 express in one cell type. We constructed brain cell-related gene co-expression 91 modules for human and mouse, and found large differences among species. 92 We found a statistically significant correlation between cell-specific fold change. 93 a measure developed in this study, and gene membership in the brain cell-94 related coexpression modules. Combining DGE and COE, we identified 16 95 novel candidate marker genes (NCM) in mouse brain and 18 NCM in the human 96 97 brain. Through supervised cell deconvolution analysis, we showed that using GSM and NCM improved the performance of deconvolution. 98

99

## 100 **Results**

To evaluate and discover brain cell marker genes, we performed DGE and COE analysis on transcriptomic or proteomic data (Figure 1). We used six datasets of purified cell populations for DGE analysis (DGEDat) and two single cell datasets for COE analysis (COEDat) (Table 1). The DGEDats included transcriptome and proteome data from human and mouse brain purified cell populations. The COEDats were single-cell RNA sequencing data from both human and mouse brains.

108

## 109 Commonly-used marker genes of four major cell types

We collected 540 marker genes that were commonly used for labeling cells 110 and validating cell isolation (Supplementary Table 1). These marker genes were 111 identified in published literature<sup>9,10,13-15</sup>, company websites<sup>24,25</sup>, and ISH 112 databases, such as the Allen Brain Atlas (ABA) and GENSAT<sup>26-28</sup> for labeling 113 neurons, astrocytes, microglia, oligodendrocytes, and other cell types in the 114 brain. Of 540 candidate marker genes, only eight genes were reported in all 115 data sources while most of the marker genes were source-specific 116 117 (Supplementary Figure 1). Genes annotated as marker genes of more than two cell types by different sources were considered as "conflict marker genes." We 118 found 27 conflict marker genes in the 540 collected genes (Supplementary 119 Table 1). The other genes had no conflict annotations in different data sources 120 and were classified as "consistent marker genes." 121

122

## 123 DGE-based specificity evaluation of commonly-used marker genes

We identified Gold-Standard Marker genes (GSM) that showed cell-type specificity across multiple types of data through DGE analysis. We found that the classical fold-change value, which is typically calculated as the expression in the target cell divided by averaged expression in other cells<sup>14,29</sup>, may produce inaccurate calls of marker genes (Supplementary Figure 2, Supplementary
 Table 2). To avoid this problem, we created a measure of cell-specific fold
 change (csFC). The csFC was defined as equation (1).

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142

$$csFC = \frac{expression in the target cell type}{the highest expression in all other cell types}$$
(1)

To be considered a GSM, the following four criteria had to be met based the 132 datasets we collected: 1) the gene must be detected in the target cell type in all 133 six DGEDats. There were 113 of the 540 candidates that met this criterion. 2) 134  $csFC \ge 2$  in all six DGEDats. 3) Benjamin-Hochberg (BH) corrected p-value of 135 the two-sample Wilcoxon test of expression in the target cell, and expression in 136 other cell types should be lower than 0.05 in more than two of the six DGEDats. 137 4) the gene must be shown to be specific in at least one proteomic dataset. 138 139 Using these criteria, we identified 22 GSM in total. Nineteen of the 22 GSM were from the consistent marker genes group, and three were from the conflict 140 marker genes group (Table 2). 141

## 143 COE analysis of two large single-cell datasets

144 To discover the co-expression of marker genes, we performed weighted gene co-expression network analysis (WGCNA) on human and mouse brain single-145 cell transcriptome data in parallel with DGE. We annotated the co-expression 146 modules using pSI packages<sup>30</sup>, which can identify genes enriched in specific 147 cell populations and test gene overrepresentation by Fisher's exact test. Figure 148 2A shows the p-value of cell type enrichment of each module after correcting 149 for multiple testing by BH. We chose the most significant module in the cell type 150 enrichment analysis as the brain cell co-expression module (BCCM) for each 151 cell type (Table 3, Supplementary Figure 3 and Supplementary Figure 4). We 152 used Gene Ontology analysis to determine the biological functions of each 153 BCCM (Supplementary Table 3). The BCCMs were enriched in biological 154 processes for specific cell types. For example, the oligodendrocyte-related 155 module was enriched in the axon ensheathment pathway. 156

157 Next, we used the module preservation test to compare the BCCMs in 158 human and mouse. The BCCMs of mouse brain were preserved in the human 159 brain co-expression network. However, only the human neuron module was 160 preserved in the mouse brain co-expression network (Figure 2B). Therefore, 161 we analyzed the BCCMs for mouse and human brain separately in subsequent 162 analysis to ensure we discover marker genes tailored specifically for human 163 and mouse.

164

## 165 **DGE-COE relationship of brain cell marker genes**

After the independent analyses of DGE and COE, we explored the relationships between them. We first asked whether marker genes with stronger specificity have a higher probability to enter the BCCMs than those with lower specificity. We tested 107 marker genes covered by six DGEDats and human COEDat. These 107 genes had 72 clustered into the four cell-type specific

BCCMs and 35 into the other non-BCCMs. We found that csFC values of the 171 72 BCCM marker genes were higher than those of the 35 non-BCCM marker 172 genes in all six DGEDats (Figure 3A, p-value of two-sample Wilcoxon test 173 <0.05). In other words, marker genes in the BCCMs were more specific than 174 the marker genes in the non-BCCMs. Significantly higher csFC values of 175 marker genes in BCCMs than in non-BCCMs were also observed in mouse 176 data (Supplementary Figure 5A, p-value of two-sample Wilcoxon test <0.05). 177 This suggests that the highly-specific marker genes are more likely to be placed 178 in a BCCM. 179

Based on the test above, we next hypothesized that the highly-specific 180 marker genes positioned close to the hub of the BCCMs have module 181 membership rankings that are higher than non-GSM in the same BCCM. We 182 divided the 72 marker genes in the human BCCMs into 20 GSM as identified 183 above and 52 non-GSM. To compare the module membership ranking of these 184 two gene groups, we performed a two-sample Wilcoxon test on their module 185 membership (kME). kME is a measurement parameter used to assess the 186 correlation between a gene and the eigengene, the hub of the co-expression 187 module. A gene with high kME means that it has high correlations with other 188 genes and consequently high ranking in the module. The kME values of GSM 189 were significantly higher than those of non-GSM in the human BCCMs (p-value 190 of two-sample Wilcoxon test<0.05, Figure 3B). However, the ranking of GSM in 191 the BCCMs was not significantly higher than non-GSM in the mouse data (p-192 value of two-sample Wilcoxon test = 0.13, Supplementary Figure 5B). 193

These two analyses suggested that a connection did exist between DGE and 194 COE for the marker genes. We further chose csFC representing DGE, and kME 195 representing COE, to study the relationship between them. Significant 196 correlations were observed between csFC values from five of the six DGEDats 197 and kME values from human co-expression network (Spearman rho>0.2, p < 198 0.05; Figure 3C). In the mouse data, kME values of the marker genes were 199 significantly correlated with csFC values in four of the six DGEDats (Spearman 200 rho>0.2, p < 0.05; Supplementary Figure 5C). This indicates that high cell-201 specific fold change and high correlation with other marker genes in the BCCMs 202 are two related properties of marker genes. 203

204

# Novel candidate brain cell marker genes are revealed by integration of COE and DGE

Based on the relationship observed between DGE and COE, we developed 207 new criteria for selecting novel candidate brain cell marker genes (NCM). Since 208 the BCCMs of human and mouse were not completely preserved, NCM was 209 defined in human and mouse separately. The mouse NCM should have 1) csFC 210 equal to or greater than 2 in at least two DGEDats from DGEDat2-DGEDat6 211 212 (BH corrected p-value of two samples of Wilcox test < 0.05), and 2) kME should be greater than 0.6 in COEDat2. We identified 16 mouse NCMs according to 213 the criteria (Table 4, Supplementary Table S4). Because only one DGEDat for 214

the human brain was available for analysis, we set relatively stricter criteria for
human NCM to make more conservative calls. The human NCM should have
1) csFC significantly larger than 4 in the DGEDat1 (BH corrected p-value < 0.05)</li>
and 2) kME should be greater than 0.8 in the COEDat1. We identified 18 human
NCM meeting these criteria (Table 4, Supplementary Table S5).

220

#### 221 GSM and NCM improve the performance of supervised deconvolution

We used supervised deconvolution to examine how the choice of marker 222 genes impacts deconvolution results using mouse data. We hypothesized that 223 including GSM and NCM would improve deconvolution accuracy compared to 224 not having them in the calculations. We downloaded mouse expression data 225 from purified neuron, astrocyte, oligodendrocyte, and microglia, as well as RNA 226 mixtures with known proportions of each cell type<sup>31</sup>. The purified cell expression 227 data was used as a reference profile, and the mixture data was used for 228 deconvolution. We constructed four types of reference gene sets: baseline, 229 GSM plus, NCM plus, and NCM GSM plus. The baseline reference gene set 230 included all the genes except for GSM and NCM. The other references were 231 constructed by adding GSM, NCM, and their combination into the baseline 232 reference. We used the root mean square error (RMSE) between estimated cell 233 proportions and the true proportion to evaluate deconvolution performance. 234 Higher RMSE indicated poorer performance of deconvolution. The optimal 235 number of marker genes for deconvolution was determined (Materials and 236 Methods). We found that the deconvolutions with baseline reference of 400 237 genes had the lowest RMSE, so we used this number of genes to construct the 238 four tested references. 239

We observed that adding either set of GSM or NCM into the reference 240 reduced the RMSE (Figure 4), suggesting that the inclusion of GSM and NCM 241 can improve the performance of deconvolution. The reference including both 242 NCM and GSM performed the best. To prove that the improved performance of 243 the reference with NCM or GSM was not because of a larger number of marker 244 genes used, we completed permutations by constructing three permutated 245 references with randomly selected genes, excluding GSM and NCM. The 246 permutation was repeated 1000 times for each type of permutated reference. 247 Deconvolution using a reference with GSM or NCM outperformed the 248 deconvolution using a permutated reference without GSM or NCM, showing 249 that improved deconvolution performance when GSM and NCM were included 250 was not related to the increased reference size (Figure 4B). 251

252

## 253 **Discussion**

The current study describes the first systematic evaluation of marker gene specificity and their reliability for identifying cell types in human and mouse brains. We not only evaluated the published marker genes but also designed new criteria to discover novel marker genes based on both differential gene expression and co-expression. Applying our proposed novel marker genes to
 deconvolution improved the performance of deconvolution and resulted in more
 accurate cell proportion estimates.

This study identified a set of marker genes to discriminate neurons, 261 astrocytes, microglia, and oligodendrocytes. New brain cell types have recently 262 been identified with the development of single-cell RNA sequencing<sup>32</sup>. The 263 evaluation of marker genes for these new cell types cannot be achieved 264 currently because the multi-omics for these new cell types are not available. 265 We required the cell types in evaluation to be measured at both transcriptome 266 and proteome level, and currently only the four major cell types above satisfied 267 the criteria. Our method will be adaptable to the newly identified brain cell types 268 when multi-omics data are available. 269

270 One of the important outcomes of the current study was validating the specificity of marker genes reported in the literature. Most of the genes 271 (304/540) included in the current study were claimed to be marker genes in a 272 single source, and only eight genes had a consistent claim supported by all the 273 collection sources (Supplementary Figure 1). Some genes that we tested (27 / 274 540) had conflict definitions for different cell types including several well-known 275 marker genes, such as GFAP<sup>33</sup> and ITGAM<sup>34</sup>. Our evaluation refined a list of 276 reliable marker genes and supported using GFAP as a marker of astrocytes 277 and ITGAM as a marker of microglia. 278

We were strict in assessing the specificity of marker genes, which led to 279 removing some genes from commonly used marker gene lists. We compared 280 the classic fold-change and cell type-specific fold-change of consistent marker 281 genes (Supplementary Table 2). Eight marker genes were imprecisely defined 282 in more than three of six DGEDats using the classic fold change. For example, 283 SELENBP1 was a claimed astrocyte marker gene using averaged ranks across 284 comparisons with each of other cell types<sup>13</sup>. However, its expression in 285 microglia is close to, or even higher than expression in astrocytes in DGEDat2-286 DGEDat6. We removed it from the marker gene list because of its similar 287 expression in microglia and astrocyte (Supplementary Figure 2). Most of the 288 candidate marker genes failed to meet our criteria of GSM due to either being 289 expressed at a similar level in more than two cell types (17%) or not being 290 detectable as protein in the target cell type (20%), such as RBFOX3 and 291 TMEM119. These two genes both showed target cell specificity when they 292 could be detected (Supplementary Table 6). We expect that more marker genes 293 including these two genes may be reclassified as GSM when more reliable 294 proteomics data becomes available. 295

We showed a positive correlation between the csFC and kME of marker genes in both human and mouse brain. This is in line with our expectation that good marker genes will have similar expression patterns across cell types and strongly correlate with each other, which forms the core part of the cell module. The most important meaning of the strong correlation is that it suggests COE can be used for discovering marker genes. COE used all cell types, both characterized and uncharacterized, in brain tissue while DGE only used the several measured cell types to identify marker genes. The marker genes identified by COE should be more robust because they showed cell typespecificity across a broader range of cell types. This relationship will help to identify more brain cell marker genes from single-cell sequencing data, a technique that is increasing in popularity.

To explore the potential use of antibodies of NCM for cell labeling, we 308 checked NCM's subcellular localization of expression in the COMPARTMENTS 309 database<sup>35</sup> and the Allen Brain Atlas<sup>36</sup>. Eight human NCM and six mouse NCM 310 are expressed on the plasma membrane, suggesting that antibodies made to 311 these gene products have potential for use in FACS. One human NCM and 312 seven mouse NCM are expressed at the nucleus, suggesting their potential use 313 314 in sorting nuclei. Most of the mouse NCM already had archive ISH data except Elavl4. However, for the human brain, only SNTA1 had ISH data in the database. 315 More experiments are needed to verify the subcellular location of the human 316 NCM. 317

Supervised deconvolution was developed to replace the physical sorting of 318 cell types. Supervised deconvolution infers cell proportion based on the 319 expression of cell marker genes. Consequently, cell-type specificity of marker 320 genes determines the accuracy of estimated proportions<sup>37</sup>. The deconvolution 321 method is relatively well established, but validated marker genes for supervised 322 deconvolution are lacking. NCM we proposed reduced the RMSE of 323 deconvolution from 7.9% to 7.6% and resulted in improved accuracy of cell 324 proportion estimates. The marginal improvement was expected because the 325 baseline reference was composed of 400 genes with > 2-fold csFC. Instead of 326 completing computations with 400 genes, using only the 21 GSM and 13 NCM 327 we identified improved the performance of deconvolution slightly (0.3%) and is 328 less resource intensive. 329

To date, various studies have found similarities and differences between 330 tissue of humans and mice at the transcriptome level<sup>17,38-40</sup>. A study found a 331 high degree of co-expression module preservation between human and mouse 332 brain, and all mouse modules showed preservation with at least one human 333 module whereas there were multiple human-specific modules<sup>41</sup>. The modules 334 enriched in neuronal markers were more preserved between species than 335 modules enriched glial marker genes<sup>41</sup>. This work conducted at the tissue level 336 is consistent with our results showing that mouse shared BCCMs with human, 337 but the BCCMs of the human brain were human-specific, except the neuron-338 related module. Our results also supported a recently published work at the 339 single-cell level by Xu et al. who observed that hundreds of orthologous gene 340 differences between human and rodent were cell type-specific<sup>42</sup>. Our data add 341 to accumulating evidence that human have more cell-specific co-expression 342 modules than mouse. Importantly, this implies that research on brain-related 343 diseases using mouse models may have limited applicability to humans 344 because of the difference between human and mouse brain cells. Furthermore, 345

the definitions of brain cell types should consider species differences.

Our work is limited by the lack of cell-specific gene expression data with a large sample size and replication. This made the criteria for the evaluation less universal and more specific to our data sets. We could only calculate the pvalue for four of six DGEDats due to lack of replication. Another limitation is the data used in the discovery of the relationship between DGE and COE were not from the same samples. This may explain why we did not observe strong correlations in all tested datasets.

Through a comprehensive evaluation of the brain cell marker genes; we developed a new method to identify marker genes, and provide a list of reliable marker genes for brain cells to guide the cell identification. Recently, studies reported methylome<sup>43</sup> and regulome<sup>44</sup> of brain cells, creating the potential to develop marker genes at epigenetics level. It would be meaningful to construct a framework by combining different omics data and methods to fully describe the cell types in the brain.

361

## 362 Materials and Methods

#### 363 DGEdats pre-processing and quality control

We collected six datasets for the DGE-based evaluation. 1) DGEDat1<sup>15</sup>: Cells 364 were isolated from the human temporal lobe cortex by immunopanning. We 365 downloaded the fragments per kilobase of transcript per million mapped reads 366 (FPKM) matrix. Fetal samples and genes with FPKM<0.1 in more than one 367 sample were removed. 2) DGEDat2<sup>14</sup>: Cells were isolated from mouse cerebral 368 cortex by immunopanning and FAC. We downloaded the expression level 369 estimation which was quantified as FPKM. Genes with FPKM<0.1 in more than 370 two samples were removed. 3) DGEDat3<sup>10</sup>: Gene expression of cells isolated 371 from mouse brain cortex were measured by microarray. The microarray data 372 contained 12 cell populations, which made use of the Mouse430v2 Affymetrix 373 platform. We downloaded the raw CEL file. All the CEL files were subjected 374 together to background correction, normalization and summary value 375 calculation using the R package affy<sup>45</sup> ('rma' function). The probes with 'A' or 376 'M' state in more than two samples were removed. 4) DGEDat4<sup>11</sup>: Cells were 377 isolated from E16.5 and P1 mouse brain to culture neuron and glia cells. We 378 downloaded the expression matrix which were quantified as reads per kilobase 379 of transcript per million mapped reads (RPKM). Genes with RPKM<0.1 in more 380 than three samples were removed. 5) DGEDat5 and DGEDat6<sup>11</sup>: both primary 381 cultured cells and acutely isolated cells were collected from four replicates of 9-382 mouse brains. Liquid chromatography-tandem mass week-old whole 383 spectrometry analysis was performed. We downloaded the quantified 384 expression matrix. Genes with one missing value were removed. 385

386

#### 387 COEdats pre-processing and quality control

388 Two large-scale single-cell RNA sequencing datasets from both human and

mouse brain were collected for co-expression analysis. 1) COEDat1. The 389 human single cell transcriptome was from adult human individual's temporal 390 lobes<sup>46</sup>. In total, 332 cells from eight adult human brains (three males and five 391 females) were collected and profiled by Illumina MiSeg and Illumina NextSeg 392 500. Raw sequencing reads were aligned using STAR and per gene counts 393 were calculated using HTSEQ. We downloaded the counts matrix. 2) 394 COEDat2. The mouse single cell transcriptomes of 3005 cells from 395 somatosensory cortex and hippocampal CA1 regions were collected from 396 juvenile (P22 - P32) CD1 mice including 33 males and 34 females<sup>47</sup>. The 397 sequencing platform was Illumina HiSeq 2000. Raw reads were mapped to the 398 mouse genome using Bowtie and the mapped reads were guantified to raw 399 counts. We downloaded the counts matrix. 400

COEDats were pre-processed in Automated Single-cell Analysis Pipeline 401 (ASAP)<sup>48</sup>. Genes with Counts per Million (CPM) lower than 1 in more than ten 402 samples were removed from human brain data, and genes with CPM lower than 403 1 in more than 50 samples were removed from mouse brain data. After quality 404 control, 13941 and 12149 genes were retained for human and mouse brain, 405 406 respectively. The human brain data were normalized by voom function. Mouse data was normalized by scLVM. In total, 57 ERCC spike-ins in mouse data were 407 used for fitting of technical noise. The normalized data were retained. 408

409

#### 410

#### 0 Deconvolution data pre-processing and quality control

Gene expression data of brain samples with known cell proportion from rat was used in cell type-specific deconvolution<sup>31</sup> (GEO accession: GSE19380). This dataset contains four different cell types including neuron, astrocyte oligodendrocyte and microglia, and two replicates of five different mixing proportions (Supplementary Table 7). The platform used was Affymetrix Rat Genome 230 2.0 Array. All the CEL files were subjected together to background correction, normalization and summary value calculation using 'rma' function.

418

#### 419 **Co-expression analysis**

To determine the gene networks of specific cell types, we completed 420 weighted gene co-expression network analysis (WGCNA<sup>22</sup>) on single-cell 421 sequencing data from both human and mouse brain using the signed network 422 type. The parameter settings were as follows: Pearson correlation function, 423 signed Topological Overlap Matrix (TOM) matrix, minimal module size of 20, 424 deepSplit of 4, mergeCutHeight of 0.25 and pamStage of true. The power for 425 human and mouse data was 7 and 6, respectively. The number of modules for 426 human and mouse data was 22 and 10, respectively. The pSI package was 427 used to identify the cell-related modules. The threshold for the enrichment test 428 was BH-corrected p-value<0.05. The GO terms analysis was identified by 429 Gorilla<sup>49</sup>. The expression localizations of genes were provided by 430 COMPARTMENTS<sup>35</sup>. 431

#### 433 Module preservation test

A module preservation test was performed using the modulePreservation<sup>50</sup> 434 function in the WGCNA R package. Zsummary is a measurement to assess the 435 preservation based on the size, density and the connectivity of modules. 436 Zsummary < 2 indicated the module was not preserved, 2 < Zsummary < 10 437 indicated weak to moderate preservation, and Zsummary > 10 indicated high 438 module preservation. We performed the module preservation test twice, once 439 withmouse data as the reference and human data as the test set and once with 440 roles reversed. 441

442

#### 443 Supervised deconvolution

We used function 'Isfit' in CellMix<sup>4</sup> for deconvolution. In each mixture sample, we tested i probes and j cell types. The expression of each probe equals the sum of expression of purified cell types times corresponding cell proportions:

> $A_{11}X_1 + A_{12}X_2 + \dots + A_{1j}X_j = B_1$  $A_{21}X_1 + A_{22}X_2 + \dots + A_{2j}X_j = B_2$

447 448

> 449 450

 $A_{i1}X_1 + A_{i2}X_2 + \dots + A_{ij}X_j = B_i$ 

AX = B

Where A<sub>ij</sub> is an expression signal of probe i in a purified cell j, B<sub>i</sub> is an expression signal of probe i in a mixture of cells, and X<sub>j</sub> is a proportion of cell type j. The formula can be summarized in a matrix equation:

454

where A is the reference matrix of the expression of all probe sets in all cell
types, B is the vector of expression levels of all probe sets in the mixture, and
X is the vector of the proportions of all cell types comprising B. The equation
was solved for X with the R function 'lsfit' (linear least squares algorithm).

The change of reference size was achieved by the following steps: 1) 459 Construct the marker gene pool for four cell types and calculate the csFC. 2) 460 Sort the marker gene pool according to the csFC in descending order. 3) 461 Separate the reference genes into three types: GSM, NCM, and base genes. 462 4) Pick the desired number of marker genes from the base gene pool to 463 construct baseline reference and perform deconvolution. 5) Add the GSM, 464 mouse NCM, or both GSM and NCM into the baseline reference to construct 465 three tested references: gsm plus, ncm plus, gsm ncm plus. 6) perform 466 deconvolution with three types of references separately. 7) Calculate RMSE 467 between the estimated proportion and true proportion using the 'rmse' function 468 in Metrics packages for each type of references. 9) Repeating step 2~step 8 for 469 increasing reference sizes. 470

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# 479 Author contributions

R.D. designed the study, performed the analyses and wrote the paper. Y.C.,
C.J., and J.D. helped with data collection and manuscript writing. C.L. and C.C
created the project, supervised the study, contributed to the interpretation of the
results, and revised the manuscript.

484

## 485 **Competing interests**

486 No competing interests declared.

487

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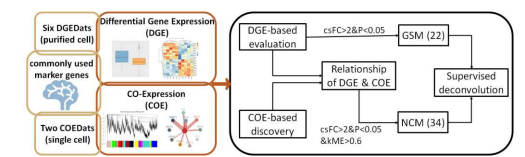
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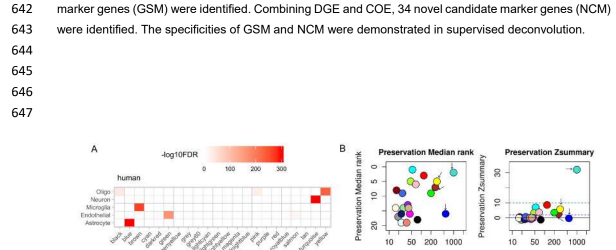
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**Figure 1.** Analysis workflow. Six DGEDats of the purified cell population and two COEDats of single cells were used to evaluate 540 commonly-used brain cell marker genes. Differential gene expression (DGE) was performed on six DGEDats and the cell-specific fold change (csFC) was defined to measure the cell specificity for the marker genes. Co-expression (COE) analyses were performed on two COEDats and cell-specific networks were constructed. The correlation of genes with the module eigengene in the cell network was measured as module membership (kME). Through DGE-based evaluation, 22 gold-standard



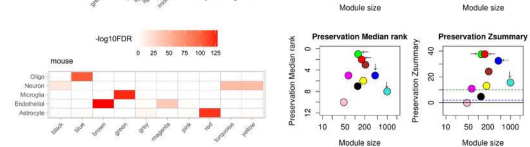


Figure 2. Cell type enrichment and preservation test of co-expression modules for human and mouse brain. (A) Enrichment of brain cell marker genes in human and mouse co-expression modules. The most significantly enriched module was defined as the brain cell co-expression module (BCCM) for each cell type. The human BCCMs are blue (astrocyte), brown (microglia), turquoise (neuron), and yellow (oligodendrocyte). The mouse BCCMs are red (astrocyte), green (microglia), turquoise (neuron), blue (oligodendrocyte). (B) Preservation of BCCMs between human and mouse brain. The top panel is the preservation test of BCCMs of the human brain in mouse data. The bottom panel is the preservation test of BCCMs of the mouse brain in human data. The arrows point to the BCCMs. Zsummary < 2 indicates the module is not preserved, 2 < Zsummary < 10 indicates weak to moderate preservation, and Zsummary > 10 indicates high module preservation.

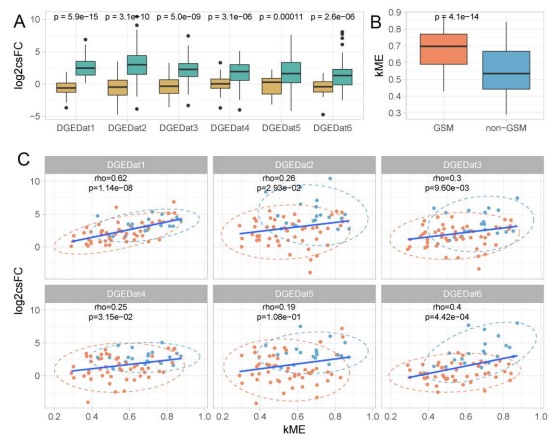


Figure 3. The relationship between DGE and COE of marker genes in human brains. (A) The comparison of csFC of BCCM marker genes and non-BCCM marker genes. The turquoise box denotes the marker genes in BCCMs and the mustard box denotes the marker genes in non-BCCMs (NBCCM = 72, NNON-BCCM = 35). The p-value is from a two-sample Wilcoxon test between csFC of marker genes in BCCMs and non-BCCMs. (B) The comparison of kME of the GSM and non-GSM in the BCCMs. A two-sample Wilcoxon test was used to test the significance of the difference (NGSM=20, Nnon-GSM=52). (C) The Spearman correlation between csFC and kME of marker genes in BCCMs. The blue dot represents GSM and the orange dot represent other marker genes. What are the dashed blue and orange circles?

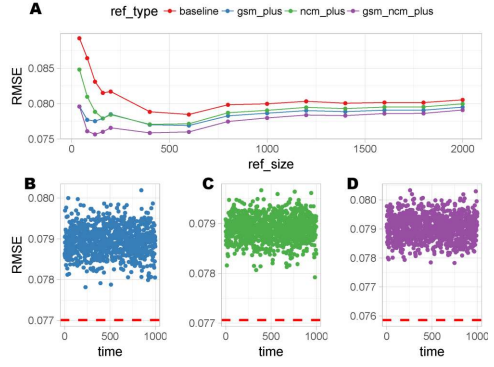


Figure 4. Effect of GSM and NCM in supervised deconvolution. (A) The RMSE between true and estimated cell proportion by supervised deconvolution with different references. The references are defined as follows: baseline = reference without GSM and mouse NCM; gsm plus = baseline + GSM; ncm\_plus = base + mouse NCM; gsm\_ncm\_plus = base + GSM +mouse NCM. With increasing size of the reference, the cell-specific fold change of marker genes included in the reference decreased. The deconvolution performance of permutated references without GSM and NCM where size is equal to the gsm plus (B), ncm plus (C), gsm ncm plus (D). The colors match the five refrences in figure 4A. The red dashed lines indicate the RMSE of deconvolution using gsm\_plus, ncm\_plus, and gsm\_ncm\_plus reference of 400 genes.

#### 720 Table 1 Datasets used

dataset	species	omics	platform	purification	Brain region	#sample/(cells)	study
DGEDat1	human	transcriptome	RNA-seq	isolated*	temporal lobe	45	GSE73721
DGEDat2	mouse	transcriptome	RNA-seq	isolated	cerebral cortex	17	GSE52564
DGEDat3	mouse	transcriptome	array	isolated	forebrain	10	GSE9566
DGEDat4	mouse	transcriptome	RNA-seq	culture*	Whole brain	22	Sharma et al.
DGEDat5	mouse	proteome	MS	culture	Whole brain	27	Sharma et al.
DGEDat6	mouse	proteome	MS	isolated	Whole brain	4	Sharma et al.
COEDat1	human	transcriptome	RNA-seq	isolated	somatosensory cortex and hippocampal CA1	(3005)	GSE60361
COEDat2	mouse	transcriptome	RNA-seq	isolated	temporal lobe	(332)	GSE67835

721 \*seq =RNA-sequencing, array = microarray, MS= mass spectrum, isolated= isolated from tissue, culture = primary culture. The

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CLU         astrocyte         (3.77, 0.6, 3.63e-04)         (3.           ALDOC         astrocyte         (1.62, 0.68, 3.63e-04)         (2.	astrocyte (3.77, 0.6, 3.63e-04)		SLC4A4 astrocyte (3.17, 0.38, 1.81e-03) (4.	SLC1A3 astrocyte (2.79, 0.34, 3.63e-04) (4	ALDH1L1 astrocyte (2.57, 0.3, 1.81e-03) (4.	PPAP2B astrocyte (3.28, 0.56, 3.63e-04) (4.	GJA1 astrocyte (4.5, 0.45, 1.81e-03) (4.	GFAP* astrocyte (3.19, 0.64, 5.76e-04) (2	PTPRC microglia (3.98, 0.24, 2.29e-02) (2.	AIF1 microglia (2.81, 0.46, 2.29e-02) (5.	TLR2 microglia (4.24, 0.27, 2.29e-02) (7.	TLR7 microglia (2.88, 0.25, 2.11e-02) (5.	ITGAM* microglia (3.16, 0.34, 1.91e-02) (6.	RELN neuron (4.74, NA, NA) (8	GAD1 neuron (5.38, NA, NA) (4.	SLC12A5 neuron (3.49, NA, NA) (3.	DCX neuron (1.64, NA, NA) (4.	MBP oligo (3.49, 0.55, 1.46e-03) (9.	SLC44A1 oligo (2.48, 0.26, 1.46e-03) (3.	CNP oligo (2.9, 0.42, 1.46e-03) (6.	PLP1* oligo (5.01, 0.69, 1.63e-03) (10	gene cellType DGEDat1
(2.01, U.37, 9.71e-UZ)	81 0 37 0 71 <sub>0</sub> 00)	(3.65, 0.38, 9.71e-02)	(4.36, 0.61, 9.71e-02)	(4.7, 0.46, 9.71e-02)	(4.11, 0.32, 9.71e-02)	(4.53, 0.46, 9.71e-02)	(4.96, 0.55, 9.71e-02)	(2.7, 0.38, 5.00e-01)	(2.76, 0.62, 9.71e-02)	(5.16, 0.39, 9.71e-02)	(7.09, 0.05, 9.71e-02)	(5.64, 0.64, 9.71e-02)	(6.12, 0.46, 1.25e-01)	(8.4, 0.78, 9.71e-02)	(4.87, 0.29, 9.71e-02)	(3.19, 0.44, 9.71e-02)	(4.76, 0.26, 9.71e-02)	(9.09, 1.16, 9.71e-02)	(3.99, 0.33, 9.71e-02)	(6.52, 0.48, 9.71e-02)	(10.43,0.19, 1.67e-01)	DGEDat2
	(1.02, 0.44, 1.04e-01)	(1.49, 0.34, 1.04e-01)	(2.9, 0.59, 1.04e-01)	(2.65, 0.38, 1.04e-01)	(3.85, 0.26, 1.04e-01)	(2.11, 0.41, 1.04e-01)	(2.44, 0.48, 1.04e-01)	(2.3, 0.54, 2.50e-01)	(7.44, 0.61, 1.04e-01)	(4.69, 0.4, 1.04e-01)	(5.27, 0.2, 1.04e-01)	(5.91, 0.29, 1.04e-01)	(3.48, 0.78, 1.67e-01)	(5.91, 0.32, 1.04e-01)	(5.94, 0.36, 1.04e-01)	(2.42, 0.74, 1.04e-01)	(5.22, 0.32, 1.04e-01)	(2.39, 1.18, 1.04e-01)	(3.51, 0.27, 1.04e-01)	(5.58, 0.34, 1.04e-01)	(5.61, 0.44, 1.67e-01)	DGEDat3
	(1.14, 0.43, 2.42e-03)	(4.66, 0.24, 2.42e-03)	(1.63, 0.19, 2.42e-03)	(3.34, 0.29, 2.42e-03)	(2.26, 0.24, 2.42e-03)	(1.94, 0.29, 2.42e-03)	(3.44, 0.26, 2.42e-03)	(2.87, 0.37, 5.26e-03)	(3.01, 0.18, 2.42e-03)	(3.67, 0.36, 2.42e-03)	(2.1, 0.23, 2.42e-03)	(3.02, 0.14, 2.42e-03)	(3.07, 0.2, 1.05e-02)	(2.83, 0.23, 9.77e-05)	(3.59, 0.42, 9.77e-05)	(2.83, 0.26, 9.77e-05)	(2.81, 0.31, 9.77e-05)	(4.33, 0.51, 8.01e-05)	(1.8, 0.4, 8.01e-05)	(2.63, 0.37, 8.01e-05)	(5.08, 0.47, 4.76e-05)	DGEDat4
	(1.25, 0.49, 3.02e-03)	(3.48, 0.44, 9.89e-03)	(3.06, 0.53, 1.58e-03)	(3.6, 0.49, 9.89e-03)	(2.73, 0.36, 9.89e-03)	(1.77, 0.53, 9.89e-03)	(5.08, 0.67, 1.58e-03)	(3.21, 0.51, 2.39e-03)	(2.87, 0.52, 4.48e-05)	(3.55, 0.68, 7.25e-04)	(3.27, 0.73, 4.48e-05)	(4.13, 0.52, 7.25e-04)	(3.92, 0.54, 9.34e-04)	(4.64, 0.46, 3.77e-06)	(5.21, 0.51, 3.77e-06)	(4.07, 0.31, 3.77e-06)	(3.23, 0.79, 3.77e-06)	(7.54, 0.67, 1.26e-04)	(2.94, 0.52, 1.39e-05)	(3.56, 0.44, 3.77e-06)	(6.05, 0.79, 1.49e-06)	DGEDat5
	(3.47, NA, NA)	(2.45, NA, NA)	(4.34, NA, NA)	(3.43, NA, NA)	(3.76, NA, NA)	(2.98, NA, NA)	(2.96, NA, NA)	(4.62, NA, NA)	(8.05, NA, NA)	(4.89, NA, NA)	(4.87, NA, NA)	(6.32, NA, NA)	(7.09, NA, NA)	(1.46, NA, NA)	(2, NA, NA)	(1.96, NA, NA)	(1.4, NA, NA)	(7.72, NA, NA)	(1.28, NA, NA)	(3.41, NA, NA)	(6.11, NA, NA)	DGEDat6

Wilcoxon tests (log2csFC, SD, p-value); Bold numbers indicate the BH corrected p-value of two-sample Wilcoxon tests is

significant (FDR<0.05); oligo=oligodendrocyte; "\*" denotes this marker gene is a conflict marker gene. The neuron of DGEDat1

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746	and all cell types in DGEDat6 have no replicates so statistical tests were not possible.
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#### 790 Table 3 Brain cell co-expression modules in human and mouse

Species	module	# of genes	cellType	Top three hub genes	Gene ontology (q-value)
human	blue	731	astrocyte	AGXT2L1, GPR98, SLCO1C1	developmental process (3.85
human	brown	377	microglia	C3, ITGAX, LAPTM5	immune system process (1.00E-67)
human	turquois e	111 9	neuron	GABRB2, SNAP25, SYT1	regulation of trans-synaptic signaling (1.73E-19)
human	yellow	370	oligo*	UGT8, ERMN, OPALIN	axon ensheathment (2.39E-1
mouse	red	187	astrocyte	GJA1, AQP4, NTSR2	multicellular organismal process (6.83E-08)
mouse	green	200	microglia	C1QA, C1QB, TYROBP	immune system process (8.79E-59)
mouse	turquois e	639 8	neuron	RAB3A, YWHAB, NDRG4	establishment of localization cell (1.20E-35)
mouse	blue	475	oligo*	UGT8, CLDN11, CNP	axon ensheathment (7.85E-13)

\*oligo=oligodendrocyte; The 'top three hub genes' column displays the top three genes that have the highest kME within BCCM.

The 'gene ontology' column displays the top enriched category for each module.

gene	cellType	species	ISH	location
ABCC9	oligodendrocyte	human	-	Plasma membrane
ACSS1	oligodendrocyte	human	-	Mitochondrial matrix
AHCYL1	oligodendrocyte	human	-	Cytoplasm
CXCR7	oligodendrocyte	human	-	Plasma membrane
DDAH1	oligodendrocyte	human	-	Cytosol
EMX2OS	oligodendrocyte	human	-	-
GNA14	oligodendrocyte	human	-	Plasma membrane
GPR125	oligodendrocyte	human	-	Plasma membrane
IL33	oligodendrocyte	human	-	Nucleoplasm
LRRC16A	oligodendrocyte	human	-	Plasma membrane
MT3	oligodendrocyte	human	-	Nucleus
PAPLN	oligodendrocyte	human	-	Extracellular region
RHOJ	oligodendrocyte	human	-	Plasma membrane
SLC14A1	oligodendrocyte	human	-	Plasma membrane
SNTA1	astrocyte	human	Y	Plasma membrane
TIMP3	astrocyte	human	-	Extracellular region
TPD52L1	astrocyte	human	-	Cytoplasm
WIF1	astrocyte	human	-	Extracellular region
C1qb	microglia	mouse	Y	Extracellular region
Mrc1	microglia	mouse	Y	Plasma membrane
Csf1r	microglia	mouse	Y	Plasma membrane
Ctss	microglia	mouse	Y	Lysosome
Ptpn6	microglia	mouse	Y	Nucleus
Cacna2d1	neuron	mouse	Y	Plasma membrane
Elavl4	neuron	mouse	-	Nucleus
SPin1	neuron	mouse	Y	Nucleus
Gria1	neuron	mouse	Y	Plasma membrane
Nipsnap1	neuron	mouse	Y	Mitochondrion
Slc25a22	neuron	mouse	Y	Plasma membrane
Mapk8	neuron	mouse	Y	Nucleus
Stau2	neuron	mouse	Y	Nucleus
Sirt2	oligodendrocyte	mouse	Y	Nucleus
Bcas1	oligodendrocyte	mouse	Y	Nucleus
Plxnb3	oligodendrocyte	mouse	Y	Plasma membrane

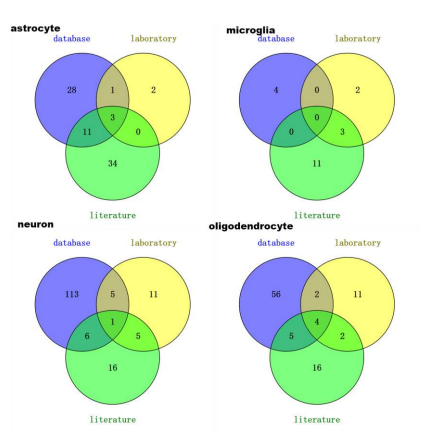
ISH: in situ hybridization image data from Allen Brain Atlas, Y: yes, having ISH image to confirm the locations, -: no ISH image.

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816	Table 4 NCM of human and mouse brain and their cellular locations

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# 826 Supplementary Materials

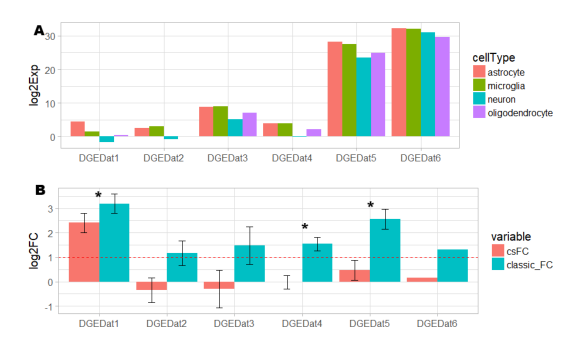
<ul> <li>Supplementary Figure 2. An example to illustrate the difference between cell-specific fold change and classic fold change</li> <li>Supplementary Figure 3. The top 50 hub genes of human brain cell co-expression module</li> <li>Supplementary Figure 4. The top 50 hub genes of mouse brain cell co-expression module</li> <li>Supplementary Figure 5. The relationship between DGE and COE in co-expression</li> <li>analysis of mouse data</li> <li>Supplementary Figure 6. Effect of human GSM in deconvoluting mouse brain tissue</li> <li>Supplementary Table 1. Collected commonly-used brain cell marker gene</li> <li>Supplementary Table 2. The classical fold change and cell type-specific fold change o</li> <li>consistent marker gene</li> </ul>	
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838 Supplementary Table 2. The classical fold change and cell type-specific fold change of	
839 consistent marker gene	F
840 <b>Supplementary Table 3.</b> The GO term of BCCM for human and mouse	
841 Supplementary Table 4. NCM of mouse brain cell	
842 Supplementary Table 5. NCM of human brain cell	
843 Supplementary Table 6. DGE of RBFOX3 and TMEM119	
844 <b>Supplementary Table 7.</b> The true proportion of cell types in the mixture for deconvolution	1
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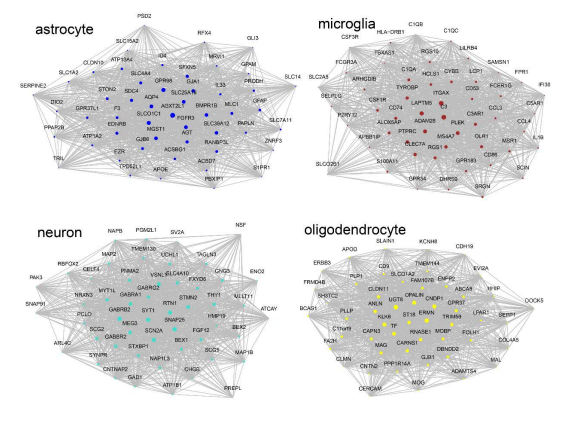
**Supplementary Figure 1** The overlap of marker genes collected from different sources. The commonlyused marker genes we evaluated were collected from three main sources: laboratory catalog, database,

and published literature. The number indicates the number of marker genes belonging to corresponding

- 872 sources.

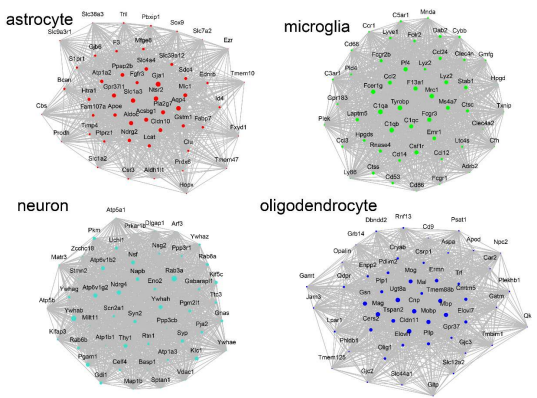


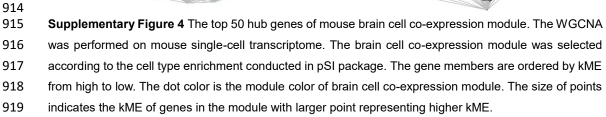
Supplementary Figure 2 An example to illustrate the difference between cell-specific fold change and classical fold change. (A) The expression of SELENBP1. SELENBP1 is an un-validated marker gene of astrocyte. All six DGEDats detected it. Its expression in microglia is very close to even higher that the expression in astrocyte in DGEDat2-DGEDat6. (B) The fold change of SELENBP1. The cell type-specific fold change (csFC) and classical fold change for the SELENBP1 are measured. The red dashed line is the empirical cut-off for the fold change (log2FC=1). The error bar denotes the standard deviation of the fold change. The "\*" indicate the BH-corrected p-value of two-sample Wilcoxon test is lower than 0.05. Since DGEDat6 have no replicates, the standard deviation cannot be calculated. The similar expression in the microglia will be covered up by the classical fold change calculation, while the csFC avoids this situation.

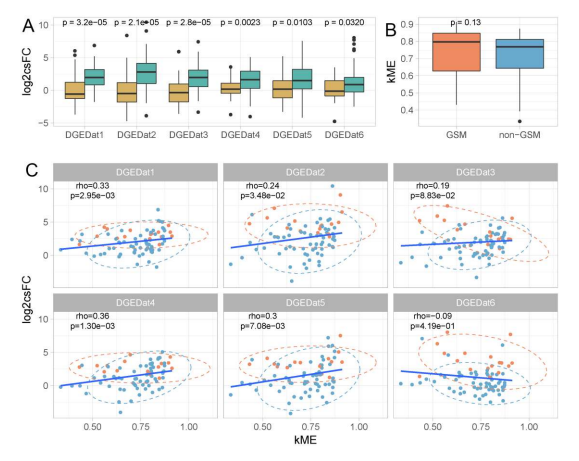


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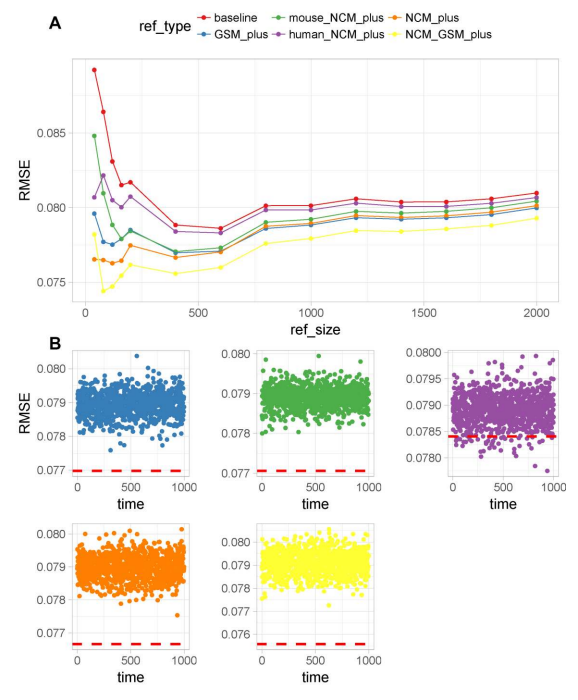
907 Supplementary Figure 3 The top 50 hub genes of human brain cell co-expression module. The WGCNA 908 was performed on human single-cell transcriptome. The brain cell co-expression module was selected 909 according to the cell type enrichment conducted in pSI package. The gene members are ordered by kME 910 from high to low. The dot color is the module color of brain cell co-expression module. The size of points 911 indicates the kME of genes in the module with larger point representing higher kME.







Supplementary Figure 5 The relationship between DGE and COE in co-expression analysis of mouse data. (A) The comparison of csFC of brain cell co-expression module (BCCM) marker genes and non-BCCM marker genes. The turquoise box denotes the marker genes in BCCM and the mustard box denotes the marker genes in non-BCCM (N<sub>BCCM</sub> = 79, N<sub>NON-BCCM</sub> = 28). The p-value is from two-sample Wilcoxon test between csFC of marker genes in BCCMs and non-BCCMs. (B) The comparison of kME of the GSM and non-GSM in the BCCM. two-sample Wilcoxon test was used to test the significance of the difference (N<sub>GSM</sub>=19, N<sub>non-GSM</sub>=88). (C) The Spearman correlation between csFC and kME of marker genes in BCCMs. The blue dot represents GSM and the orange dot represent other marker genes. 



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963 Supplemental Figure 6 Effect of human GSM in deconvoluting mouse brain tissue. (A) The 964 RMSE between true cell proportion and estimated cell proportion by supervised deconvolution 965 with different references. The deconvolution performance of permutated references without 966 GSM and NCM which size is equal to the reference tested above. The colors match the five 967 references in figure 4A. The red dashed lines display the RMSE of deconvolution using tested 968 reference of 400 genes.

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