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## 31 Abstract

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The human brain undergoes protracted post-natal maturation, guided by dynamic changes in gene expression. Most studies exploring these processes have used bulk tissue analyses, which mask cell type-specific gene expression dynamics. Here, using single nucleus (sn)RNAseq on temporal lobe tissue, including samples of African ancestry, we build a joint paediatric and adult atlas of 75 cell subtypes, which we verify with spatial transcriptomics. We explore the differences between paediatric and adult cell types, revealing the genes and pathways

- 39 that change during brain maturation. Our results highlight excitatory neuron subtypes,
- 40 including the LTK and FREM subtypes, that show elevated expression of genes associated with
- 41 cognition and synaptic plasticity in paediatric tissue. The new resources we present here

42 improve our understanding of the brain during its development and contribute to global43 efforts to build an inclusive brain cell map.

44

## 45 Introduction

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The adult human brain is a complex assembly of diverse cell types, which has been defined 47 48 with unprecedented accuracy using single cell transcriptomics<sup>1-4</sup>. This adult transcriptomic signature is set up over a protracted period of development, which begins in the embryo and 49 50 continues after birth. While the single cell diversity of the embryonic human brain has been explored<sup>5,6</sup>, little is known about how these cell type-specific gene expression profiles change 51 52 during childhood<sup>7</sup>. Most existing studies have used bulk transcriptomic approaches, which 53 revealed a dramatic period of global gene expression change during the late foetal/early 54 infancy transition, that stabilises during childhood (1 to <12-years-old) and adolescence (12 to <20-years-old)<sup>6,8-11</sup>. Bulk transcriptomics, however, cannot reveal the more subtle, cell 55 56 type-specific changes in gene expression that drive brain maturation from childhood, 57 through adolescence to adulthood.

58 Childhood and adolescence are periods of important changes in brain structure, during which 59 neuronal connections are refined and strengthened. While synaptogenesis peaks in the early 60 postnatal period, synaptic pruning activity begins during late childhood, peaks during adolescence, and then gradually decreases<sup>12-14</sup>. These stages therefore represent periods of 61 62 enhanced susceptibility to environmental influence, as well as increased neuropsychiatric risk<sup>15</sup>. Describing the typical cell type-specific gene expression trajectories of the maturing 63 64 brain will allow us to assess the effects of genetic perturbations and early adverse experiences 65 on brain maturation. Furthermore, investigating the driving forces behind cell type-specific maturational processes may help develop targeted therapies for neurological disease<sup>16</sup>. 66

67 To this end, the Paediatric Cell Atlas (PCA)<sup>17</sup>, a branch of the Human Cell Atlas (HCA)<sup>3</sup>, aims to ensure that the benefits of single cell transcriptomics are available to children as well as adults 68 69 from diverse populations<sup>3,17</sup>. Africa has the most genetically diverse<sup>18</sup> and youngest population<sup>19</sup> worldwide and by 2050, 37% of the world's children will grow up in Africa<sup>20</sup>. 70 Consequently, it is essential to include the African paediatric population in the PCA's efforts. 71 72 A reference paediatric brain cell atlas that includes data from African donors will contribute to developing treatments for locally prevalent conditions, such as tuberculosis meningitis 73 (TBM) and HIV<sup>21,22</sup>. In addition, studying the differences in gene expression dynamics between 74 adult and paediatric brains may explain why the manifestation of neurological conditions and 75 responses to therapies differ across the lifespan<sup>17</sup>. 76

To contribute to these endeavours, we present a joint paediatric and adult temporal cortex cell atlas, including samples from eight Southern African donors, annotated using the Allen Brain Map middle temporal gyrus (MTG) cell taxonomy<sup>1</sup>. We validate our annotation using spatial transcriptomics analysis. In addition, we use *de novo* marker gene analysis with 81 machine learning tools to compare our paediatric and adult datasets to the existing MTG cell 82 taxonomy and compare markers that define paediatric versus adult cell states. Using 83 differential gene expression analysis, we highlight 21 cell subtypes that show differential 84 expression of genes involved in neurodevelopment and cognition. Finally, we use our datasets 85 to define the cell type-specific gene expression of putative site-of-disease TBM biomarkers<sup>23</sup>. 86 Overall, we highlight the subtle cell type-specific differences between the paediatric and adult

87 brain and expand the representation of diverse paediatric populations in the HCA.

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- 89 Results
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## 91 A joint paediatric and adult temporal cortex cell atlas

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93 We generated snRNA-seq libraries from five paediatric and three adult donor temporal cortex 94 tissue samples. The majority of our samples were obtained from surgeries to treat epilepsy 95 (Extended Data Table 1). These new libraries were analysed alongside similar published datasets<sup>24</sup>, resulting in a total of 23 snRNA-seq datasets (including technical replicates) from 96 97 12 individuals (six paediatric and six adult) (Fig. 1a). The samples were sequenced to a median 98 depth of 19,853 reads per nucleus, with 176,012 nuclei remaining after removing low quality 99 barcodes (Extended Data Fig. 1, Extended Data Table 2). While our new datasets had a lower 100 average sequencing depth than the co-analysed published datasets, the average number of 101 genes and transcripts detected across datasets was similar (Extended Data Table 2).

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103 Using data integration and clustering we aligned similar cell types across the 23 datasets, 104 yielding 40 clusters (Fig. 1a, Extended Data Fig. 1i-h). Each cluster was assigned to one of the 105 major brain cell types (level 1 annotation) based on marker gene expression (Extended Data 106 Fig. 2a, Extended Data Table 3, Supplementary Figure 1). Additionally, we used label transfer<sup>25</sup> 107 to classify each nucleus according to the Allen Brain Map MTG atlas<sup>1</sup> (level 2 annotation) 108 (Extended Data Table 3). Barcodes with discordant level 1 and level 2 annotations (17.94%) 109 were removed to focus downstream analyses on nuclei with high confidence annotations 110 (Extended Data Table 3). Based on marker gene analysis<sup>1</sup> (Extended Data Fig. 2b), many of 111 these filtered barcodes are likely multiplets or nuclei contaminated with ambient mRNA.

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113 All 75 reference cell types were present in the final filtered dataset of 144,438 nuclei (Fig. 1b; 114 Extended Data Table 3) and expressed the expected marker genes<sup>1</sup> (Fig. 1d). Both neuronal 115 and non-neuronal cell types showed high correlation with the corresponding reference cell types<sup>1</sup> (cosine similarity score > 0.83) and lower correlation to other subtypes within their 116 class (Fig. 1e). This pattern was maintained when we considered either the paediatric or adult 117 118 datasets on their own, with the majority of paediatric cell types showing only slightly lower similarity scores than the adults (Extended Data Table 3), which is likely due to the reference 119 120 dataset only containing adult data. The cell composition of the samples was very similar with

121 no significant differences in cell type proportions between paediatric and adult samples or 122 between biological sexes (Fig. 1c; Extended Data Figure 2c-d; Extended Data Table 3). Similar to the reference atlas<sup>1</sup>, oligodendrocytes were the most common non-neuronal cell type and 123 124 Exc L2-3 LINC00507 FREM3 was the most common neuronal subtype. Neuronal clusters had 125 a greater number of expressed genes and unique molecular identifiers (UMIs) compared to 126 non-neuronal cells (Extended Data Figure 3a), while excitatory neurons had a greater number 127 of genes detected per nucleus than inhibitory neurons (Extended Data Table 3). When 128 comparing the paediatric to adult cell types, there were no significant differences in the 129 number of genes or UMIs between the age categories. Overall, the quality and composition 130 of the paediatric and adult cell atlases were very similar.

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# Spatial mapping of cell types reveals similar tissue cytoarchitecture in adult and paediatric temporal cortex

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Next, we used spatial transcriptomics to explore the positions of our annotated cell types within the temporal cortex. We generated Visium datasets from adult (31-year-old) and paediatric (15-year-old) temporal cortex samples (two sections each; Extended Data Table 1; Extended Data Fig. 4). The four Visium libraries were sequenced to a median depth of 87,178 reads per spot (median of 5,878 UMIs and 2,745 genes per spot) (Extended Data Table 4).

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Using *cell2location*<sup>26</sup>, we calculated cell type abundance estimates for each Visium spot, with our annotated snRNA-seq dataset as a reference. Oligodendrocytes were the most common cell type, while Exc\_L2\_LAMP5\_LTK was the most abundant neuronal cell type (Extended Data Fig. 5a). The annotated cell types mapped to their expected cortical layer locations across all tissue sections (Fig. 2a; Extended Data Fig. 5b), matching the spatial expression of known cortical layer marker genes<sup>1,27,28</sup> (Fig. 2b). These layered expression patterns were verified for a subset of layer-specific marker genes using *in situ* hybridisation (Extended Data Fig. 6).

149 To examine the co-location of cell types within the layered structure of the temporal cortex, 150 non-negative matrix factorization (NMF) was performed resulting in 15 cellular 151 compartments, which were visualised across the Visium samples, revealing their spatial 152 distribution (Fig. 2c-d, Extended Data Fig. 5c). In both the paediatric and adult datasets, there 153 was clear co-location of the expected neuronal cell types within overlapping compartments 154 across the cortical layers. Layer 2 was dominated by Exc L2 LAMP5 LTK (factor 11) and 155 Exc L2-3 LINC00507 FREM (factor 5), layer 3 by Exc L3-4 RORB CARM1P1 (factor 13), layer 4 by the *RORB* excitatory neuron subtypes (factor 12), layer 5 by the *THEMIS* excitatory 156 157 neuron subtypes (factor 10) and layer 6 by the FEZF2 excitatory neuron subtypes (factor 14 158 and factor 1), with the latter extending into the white matter. Inhibitory neurons were 159 primarily associated with factors 6 and 2, which were more widely spread across the layers. 160 Interestingly, these factors were more strongly associated with layers 5/6 in the adult than in

the paediatric samples. The two astrocyte subtypes were confirmed to have distinct distribution profiles, with Astro\_L1-2\_FGFR3\_GFAP (factor\_4) located primarily in layer 1 and the white matter, and Astro\_L1-6\_FGFR3\_SLC14A1 (factor\_9) more widely distributed. The remaining non-neuronal cell types were largely associated with factors located in layer 1 and the white matter.

Overall, our spatial transcriptomic analyses provide support for our annotation approach,
 showing the expected spatial distribution of annotated cell types, and revealing a largely
 similar tissue cytoarchitecture in adult and paediatric temporal cortex tissue.

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## 170 A machine learning approach identifies new temporal cortex cell type markers

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172 To establish a standardized approach for defining cell types, it has been proposed to use the 173 minimum combination of gene markers that can classify a cell type and distinguish it from other cell types<sup>29,30</sup>. Towards achieving this, Aevermann et al. (2021)<sup>29</sup> developed the machine 174 learning tool, NS-Forest V2.0, which they applied to the MTG cell atlas. Ideally, these MTG 175 176 minimal markers would be conserved in similar datasets to facilitate accurate comparisons across different studies<sup>31</sup>. Indeed, we found that the majority of MTG cell atlas minimal 177 markers<sup>29</sup> (~94%) are expressed at significantly higher levels in the expected cell types than 178 179 in other cell types (Extended Data Fig. 7, Extended Data Table 5).

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Application of the NS-Forest V2.0<sup>29</sup> algorithm to our down-sampled snRNA-seq datasets (see 181 182 Methods) revealed 202 paediatric and 196 adult minimal marker genes (Fig. 3; Extended Data 183 Table 5). The median F-beta score per cell type (the measure of the discriminative power of a 184 given combination of marker genes; paediatric=0.55; adult=0.6) and the average binary expression score (a measure of an individual gene's classification power; paediatric=0.9; 185 186 adult=0.89) were comparable across age groups and only slightly lower than that obtained for the MTG cell atlas (0.68 and 0.94 respectively)<sup>29</sup>. 47 paediatric (23.3%) and 45 adult 187 (23.0%) minimal markers overlapped with existing markers<sup>29</sup> (Fig. 3; Extended Data Table 5). 188 189 However, there was a greater overlap in minimal markers between the paediatric and adult datasets, with 68 markers (~34%) present in both lists. MERFISH<sup>32</sup> spatial transcriptomic 190 analysis of a subset of minimal makers that were shared between paediatric and adult 191 datasets confirmed their co-expression with previously described minimal markers<sup>29</sup> in adult 192 193 (31-year-old) and paediatric (15-year-old) temporal cortex samples (Extended Data Fig. 8).

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Our minimal marker analysis revealed improved markers for some cell types when compared to the reference MTG cell atlas. In our datasets, the long non-coding RNA, *LINC01331*, is a minimal marker for Exc\_L2-3\_LINC00507\_FREM3 with a beta score of 1, indicating high specificity. In contrast, one of the existing markers for this cell type, *PALMD*, is more highly expressed in endothelial cells in our datasets (Fig. 3; Extended Data Fig. 9a-b). This 200 discrepancy is likely due to the lower percentage of endothelial cells in the MTG cell atlas compared to our datasets (0.06% vs 0.9%)<sup>1</sup>. Similarly, one of the existing MTG cell atlas 201 202 markers for Exc L5-6 THEMIS CRABP1, OLFML2B, is more highly expressed in other layer 203 5/6 neurons in our dataset, whereas our minimal marker, POSTN, shows greater specificity 204 (Fig. 3; Extended Data Fig. 9c-d). Additionally, UMAP analysis of our annotated datasets using 205 our minimal marker gene list for each age group, in comparison to an equivalent number of 206 random genes, resulted in better grouping of the cell subtypes into clusters, similar to the 207 original UMAP plot (compare Fig. 1a and Fig. 4 a-b). This analysis reveals that our shortlists of 208 ~200 marker genes capture much of the underlying transcriptomic diversity in our datasets.

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210 Gene ontology (GO) analysis of our minimal marker gene lists revealed significant enrichment 211 of GO terms related to development, cell signalling, extracellular matrix and synapse 212 organisation, when considering the paediatric and adult datasets individually or together 213 (Extended Data Table 6). These results suggest that genes involved in neuronal development 214 and signalling are key to neuronal identity as the brain matures and in adult life. To further 215 assess the difference in cell type markers between our paediatric and adult datasets, we 216 expanded our analysis to include all genes with a high NS-Forest binary expression score (> 217 0.7)<sup>29</sup>. For most cell types, the majority of these top markers (>18 genes) were shared between our paediatric and adult datasets (Fig. 4c; Extended Data Tables 7-8). The 218 oligodendrocytes showed the highest number of shared marker genes (53), as well as the 219 220 second highest number of paediatric-specific markers (22). Exc L3-4 RORB CARM1P1 had 221 the highest number of adult-specific marker genes (30), while Exc L2-4 LINC00507 GLP2R 222 had no shared markers. GO analysis of the shared oligodendrocyte marker genes revealed 223 driver terms related to oligodendrocyte structure and function, including "structural constituent of myelin sheath", while the top driver terms for the paediatric-specific markers 224 225 were "oligodendrocyte differentiation" and "myelination" (Extended Data Table 6). Overall, 226 our expanded marker gene analysis suggests that neuronal cell types show greater 227 dissimilarity between their paediatric and adult states than non-neuronal cells. It is likely that 228 more diversity in the non-neuronal marker gene profiles could be revealed with subdivision 229 into further subtypes.

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## Differential gene expression analysis highlights enriched expression of genes associated with neurodevelopment in paediatric samples.

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To identify genes that were upregulated in the paediatric cell populations and thus might be involved in brain maturation, we conducted cell type-specific differential gene expression analysis with *DESeq2<sup>33</sup>*. In total, we detected 165 significantly differentially expressed genes (DEGs) across 21 cell types (123 upregulated in paediatric samples and 42 downregulated), with some DEGs associated with multiple cell types (Fig. 5a; Extended Data Table 9-10). For all DEGs, the change in expression was accompanied by a corresponding change in the

percentage of nuclei expressing the gene (Extended Data Table 10). *BayesSpace*<sup>34</sup> analysis of
a subset of DEGs in our Visium datasets confirmed that the genes were expressed at higher

- levels in the 15-year-old compared to the 31-year-old (Extended Data Fig. 10).
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245 Many of the excitatory neuron subtypes shared DEGs that are known to be developmentally regulated in the mammalian brain (Fig. 5b-e,h). LAMC3, a subunit of the extracellular matrix 246 247 protein laminin, was upregulated in three paediatric subtypes (Exc L3-5 RORB ESR1, Exc L2-3 LINC00507 FREM3, Exc L4-5 RORB FOLH1B) (Fig. 5b-d,h). LAMC3 plays a role in cortical 248 lamination in the mouse<sup>35</sup> and mutations are implicated in human brain heterotopias and 249 gyration defects<sup>36,37</sup>. Similarly, *SOX11*, a transcription factor that plays a role in embryonic and 250 adult neurogenesis in the mouse brain<sup>38</sup> and decreases in expression in the cerebral cortex 251 during development<sup>39,40</sup>, was upregulated in paediatric Exc L3-5 RORB ESR1 and Exc L2-252 253 3 LINC00507 FREM3 (Fig. 5b-c,h). FNBP1L (TOCA-1) was upregulated in Exc L2-254 3\_LINC00507\_FREM3 and Exc\_L2\_LAMP5\_LTK (Fig. 5c,e,h). FNBP1L promotes actin 255 polymerisation, regulating neurite outgrowth, and declines in expression over the course of brain maturation in the rat<sup>41</sup>. Two genes, *STEAP2*, a metalloreductase, and the TNF receptor 256 257 TNFRSF25 (DR3), were higher in adult Exc L3-5 RORB ESR1 and Exc L4-5 RORB FOLH1B 258 subtypes (Fig. 5b,d,h). STEAP2 increases in expression during post-natal hippocampal maturation in mice<sup>42</sup>. *TNFRSF25* is activated post-natally in the mouse brain, where it may 259 play a role in retention of motor control during aging<sup>43</sup>. These findings indicate that previously 260 261 reported expression dynamics for these genes in mammalian models are conserved in the 262 human temporal cortex. Importantly, our analysis reveals these patterns are specific to 263 groups of excitatory neuron subtypes.

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The majority of the DEGs were not shared across the cell types. For example, *FGF13* (*FHF2*) 265 266 and TENM1 were upregulated in paediatric Exc L3-5 RORB ESR1 (Fig. 5b). FGF13 decreases 267 in expression with age in the mouse brain, where it regulates post-natal neurogenesis<sup>44</sup> and axonal formation<sup>45</sup>. *TENM1* is a member of the teneurin transmembrane protein family that 268 269 regulates cytoskeletal organisation and neurite outgrowth, as well as shaping synaptic connections<sup>46-48</sup>. KCNG1, a voltage gated-potassium channel (Kv6.1), was upregulated in 270 paediatric Exc\_L2-3\_LINC00507\_FREM3 neurons (Fig. 5c), while MYO16 (MYR8), an 271 272 unconventional myosin protein, was upregulated in the Exc L2 LAMP5 LTK subtype (Fig. 5e). 273 Both of these genes decrease in expression with age in the mammalian brain<sup>49,50</sup>.

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In line with our minimal marker analyses, fewer genes were differentially expressed in nonneuronal cells (Fig. 5f-g, Extended Data Table 10). In Astro\_L1-6\_FGFR3\_SLC14A1, *PIK3R3* was
upregulated in paediatric samples, while *PFKFB2* was downregulated. *PIK3R3* is involved in
the PI3K-AKT growth signalling pathway, which is implicated in brain growth disorders<sup>51</sup>. *PFKFB2* is a bifunctional kinase/phosphatase that controls glycolysis. In contrast to our
findings, PFKFB2 expression is higher in juvenile rat hippocampal astrocytes than in adults,
where it may support energy demands during learning<sup>52</sup>. In oligodendrocytes, *NOTCH2* and

*RRAS2* were both upregulated in paediatric samples. *Notch2* expression decreases in the rat
 cortex with age<sup>53</sup> and is proposed to regulate glial differentiation<sup>54</sup>. These results provide new
 molecular candidates to expand our understanding of the mechanisms of astrocyte and
 oligodendrocyte maturation.

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To explore the trajectories of DEG expression, we employed *psupertime* pseudotime 287 288 trajectory analysis<sup>55</sup>, focussing on the four excitatory neuron sub-types with the highest 289 number of DEGs. In support of our DESeq2 findings, several of the identified DEGs had non-290 zero psupertime coefficients and therefore represent genes that are relevant to the ordering 291 of the cells in pseudotime<sup>55</sup> (Fig. 5i; Exc\_L3-5\_RORB\_ESR1: 13/47 [28%], Exc\_L2-292 3 LINC00507 FREM3: 16/38 [42%], Exc L4-5 RORB FOLH1B 3/27 [11%] and 293 Exc L2 LAMP5 LTK: 5/18 [28%]; Extended Data Table 11). When considering the pseudotime 294 trajectories for all DEGS in these excitatory neuron subtypes, the direction of the expression 295 matched the DESeq2 results (Supplementary Fig. 2-5). The pseudotime trajectories revealed 296 subtle expression dynamics within the analysed sample groups, showing that the majority of 297 DEGs gradually increase in expression with age from childhood to adolescence, followed by 298 a decrease in expression towards late adulthood.

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300 Genes associated with intelligence quotient (IQ) and educational attainment (EA), as well as 301 those associated with accelerated evolution in humans, have recently been shown to be 302 enriched in adult temporal lobe cortical neurons, especially the Exc L2-3 LINC00507 FREM3 subtype<sup>56</sup>. Since childhood is a key period of cognitive development<sup>57</sup>, we explored whether 303 304 the same genes were found amongst our DEGs. Of the 149 DEGs found in at least one cell 305 type, 20 (13.42%, p=0.02) are known to be significantly associated with  $EA^{58}$ , 6 (4.02%, p=0.7) with IQ<sup>59</sup> and 30 (20.13%, p=3.89E-07) with accelerated evolution in humans<sup>60</sup>. These 306 307 included several genes that are upregulated in paediatric samples, such as MYO16, KCNG1, 308 FGF13 and SOX11 (Extended Data Table 10).

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Overall, we highlight several genes that are upregulated in children/adolescents which have known roles in brain development and have been associated with cognitive ability. Our analysis builds on previous knowledge by implicating specific cell subtypes and provides new candidate genes that likely contribute to cell type-specific maturation processes.

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## 315 Gene pathways involved in cellular respiration and synaptic functioning are enriched in 316 paediatric cell types

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We next used gene set enrichment analysis (GSEA) to conduct a broad analysis of the gene pathways that are differentially regulated across all brain cell types during brain maturation. 2,006 GOBP terms where enriched in the paediatric samples compared to the adults, while 866 were depleted (p<0.01 and q<0.1) (Extended Data Table 12). When focussing on the 25 most frequently enriched terms, the majority (10 terms) were associated with cellular 323 respiration pathways (Fig. 6; Extended Data Table 12). Six were associated with intra-cellular 324 transport, including transport of neurotransmitters, while five were linked to 325 neurotransmitter release and synaptic plasticity. Three terms, including the top enriched 326 term, were associated with protein translation and modification. The majority of depleted 327 terms (10 terms) were associated with synaptic processes (Fig. 6). A further six depleted terms 328 were connected to neuronal morphogenesis, including axon and dendrite morphogenesis. 329 Two of the top depleted terms were associated with axon ensheathment. Interestingly, 330 neither of these terms were significantly enriched in oligodendrocytes or OPCs, while they 331 were associated with neuronal sub types, and microglia.

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Overall, our GSEA analysis points towards putative genetic pathways that may drive maturation in the paediatric brain. Cellular respiration processes needed to support the higher metabolic rates in the brain during childhood<sup>61</sup> may be enriched. Additionally, pathways related to strengthening synapses through neurotransmitter release may be enhanced. On the other hand, as synaptic pruning is underway<sup>62</sup>, pathways that promote synaptic growth may need to be suppressed.

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## 340 Cell type-specific expression of site-of-disease TBM biomarkers

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The PCA aims to create reference atlases that can be used to improve our understanding of cell type-specific responses to disease in children<sup>17</sup>. Here, we used our snRNA-seq datasets to interrogate the cell type-specific expression of putative genetic biomarkers for TBM<sup>23</sup>. These biomarkers are enriched in the ventricular cerebrospinal fluid from children with TBM in comparison to controls with meningitis caused by other brain infections<sup>23</sup>.

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348 66 of the 76 TBM biomarkers were expressed in our dataset, with similar expression across 349 the two age groups, and genes clearly clustering according to their relative expression across 350 the broad cell type categories (Extended Data Fig. 11). The genes with the highest relative 351 expression in our data were expressed by non-neuronal cell types, which is in line with the 352 view that immunological activity of supporting cells and their intercellular signalling 353 interactions are important drivers of the immune response to TBM<sup>63</sup>. Several of these 354 biomarkers (e.g. FADS2, AMOT and ALDH6A1) were enriched in the two astrocyte subtypes, 355 potentially pointing towards a prominent role for astrocytes in the host response to TBM.

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357 Our analyses also clearly revealed subsets of biomarkers that are more highly expressed by neuronal subtypes than non-neuronal cell types. This included biomarkers that were 358 359 associated Exc L2 LAMP5 LTK, Exc L2-4 LINC00507 GLP2R with and Exc L5-6 THEMIS C1QL3 (LYNX1, FAIM2, MAP1A, TUBB4A). This is line with the finding that neuronal 360 361 excitotoxicity is elevated in TBM<sup>23</sup> and suggests that specific excitatory neuron subtypes may 362 be contributing to this signal.

Interestingly, the two most enriched genes in the TBM biomarker dataset, *CXCL9* and *CXCL11*, were either completely absent from our datasets (*CXCL11*) or expressed by very few nuclei (*CXCL9*). The absence of these interferon-inducible chemokines in our datasets from uninfected tissue, supports the proposition that they are indeed biomarkers from the site-ofdisease<sup>64</sup> in both adults and children with TBM, and could also reflect the contribution of peripheral immune cells recruited to the brain during infection.

- 370 371
- 372 Discussion
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The brain is the most complex organ in the human body, which continuously changes as we mature. Here, we begin to unmask the molecular mechanisms guiding these processes in the temporal cortex, using single cell and spatial transcriptomics to compare similar cell types between paediatric and adult datasets.

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379 To facilitate accurate comparisons of cell types across age groups, we used the existing Allen 380 Brain Map MTG cell atlas<sup>1</sup> to annotate our datasets. This demonstrated that the reference 381 atlas, generated from adult snRNA-seq datasets, is indeed generalisable<sup>31</sup>, and can be used 382 to classify cell types from samples of different ages. This generalisability is essential for healthy human reference atlases to serve as a baseline to improve our understanding of 383 human development and disease<sup>3</sup>. Our samples and those in the reference MTG cell atlas 384 385 include neurosurgical tissue from donors with epilepsy, and while the analysed tissue is not from the site of pathology, it is important to view our findings in light of the patient diagnosis. 386 387 Previous research comparing gene expression between the neurosurgical and post-mortem samples used in the reference MTG cell atlas found a strong correlation of expression 388 389 between cell types across conditions<sup>1</sup>. In addition, a comparison of samples from 45 adult 390 donors with epilepsy to the post-mortem samples from the reference MTG cell atlas found a 391 similar number of genes and similar cell abundance per cell subclass across tissue sources, 392 however they did find more variation for these parameters in neurosurgical samples<sup>65</sup>. As 393 more paediatric MTG samples of post-mortem and neurosurgical origin become available, it 394 will be important to conduct similar analyses to determine if these findings hold for the 395 paediatric temporal cortex.

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397 Our machine-learning marker gene analysis shows that while the cell type classifications, 398 which are based on the expression of thousands of genes, can be transferred onto new datasets, the minimal markers that define the cell types do vary across datasets. Only a 399 400 quarter of our NS-Forest minimal markers overlap with the existing MTG cell atlas minimal 401 markers<sup>29</sup>. The differences in the single cell transcriptomics technologies used to generate 402 our dataset and the MTG cell atlas may account for much of this discrepancy. Nonetheless, 403 our analyses suggest that some of our markers may provide better discrimination between 404 cell types than existing markers. These results highlight a challenge that the HCA faces to

revise cell type markers as more datasets are made available to ensure that the cell typeclassification is as widely applicable as possible.

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Similar to analyses of aging in the mouse<sup>66</sup>, our analyses show there is little change in cell type 408 409 composition within the temporal cortex during human brain maturation. However, our 410 differential expression analysis highlights differences in cell states between specific paediatric 411 and adult cell subtypes. Recently, the supragranular excitatory pyramidal neurons in the MTG 412 have been shown to have high transcriptional diversity<sup>1,67</sup>, large arborisations<sup>68</sup> and 413 electrophysiological properties that impact signal integration and encoding<sup>69-72</sup> in ways that 414 may contribute to cognition. Since cognitive ability is a key feature that is established during 415 childhood<sup>68</sup>, our analysis offers an opportunity to explore how cell type-specific gene 416 expression dynamics contribute to cognitive development. Interestingly, two of the 21 417 highlighted cell types were the layer 2/3 excitatory neurons, Exc L2 LAMP5 LTK and Exc L2-3 LINC00507 FREM3, that have recently been associated with human cognition<sup>56</sup>. In line with 418 419 these findings, several of the DEGS associated with these cell types, including *FNBP1L*<sup>73</sup> and 420 *SOX11*<sup>74</sup>, have been implicated in cognitive ability and intelligence. Overall, our data points 421 towards genes that may play roles in cognitive development specifically within these 422 excitatory neurons.

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The relatively low number of genes implicated in our differential expression analysis in 424 comparison to similar studies in mouse<sup>66</sup> suggests that the difference between the paediatric 425 426 and adult brain are subtle. However, the inherent high variability in human gene expression 427 data may mask some of the differential gene expression in our limited sample. Nonetheless, 428 our pseudotime trajectory analyses reveals some of the expression dynamics that may be 429 occurring during childhood, with many genes rising in expression towards adolescence and 430 dropping off in adulthood. As the HCA database for the human temporal cortex expands, it 431 will be important to build on these analyses with more samples. Binning of samples of similar 432 age will provide a higher resolution analysis of cell type-specific gene expression trajectories 433 over the course of brain maturation.

434

Finally, we have provided the first single nucleus gene expression datasets for the brain that includes data from black Southern African donors, thus increasing the diversity of the HCA database. We demonstrate how this resource can be used to deconvolute site-of-disease biomarker analyses for TBM, pinpointing which cell types may be driving altered gene expression profiles in the brain. Importantly, these investigations have the potential to contribute to the development of effective treatments, that are tailored to specific needs of both adult and paediatric patients.

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- 443
- 444 Methods
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#### 446 Human samples

Ethical approval was granted for the collection and use of paediatric and adult human brain 447 tissue by the University of Cape Town Human Research Ethics Committee (UCT HREC REF 448 449 016/2018; sub-studies 146/2022 and 147/2022). The human brain tissue samples used to 450 generate new datasets were obtained by informed consent for studies during temporal lobe 451 surgical resections to treat epilepsy and/or cancer performed at the Red Cross War Memorial 452 Children's Hospital and Constantiaberg Mediclinic in Cape Town, South Africa. The samples 453 used in this study were of temporal cortex origin and represent radiologically 454 and macroscopically normal neocortex within the pathological context (details in Extended 455 Data Table 1). Race was recorded by the clinical teams based on their knowledge of the 456 donors. The category "black South African" includes both black and mixed race ancestries. 457 Upon resection, samples were placed in carbogenated ice-cold artificial cerebral spinal fluid 458 (aCSF) containing in (mM): 110 choline chloride, 26 NaHCO<sub>3</sub>, 10 D-glucose, 11.6 sodium 459 ascorbate, 7 MgCl<sub>2</sub>, 3.1 sodium pyruvate, 2.5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, and 0.5 CaCl<sub>2</sub> (300 mOsm) 460 and immediately transported to the laboratory (~20 minutes). Tissue blocks containing the 461 full span from pia to white matter were prepared and either flash frozen in liquid nitrogen or 462 embedded in optimal cutting temperature compound (OCT) and stored at -80°C. The OCT-463 embedded samples were flash frozen in a 10×10 mm<sup>2</sup> cryomold which was either frozen 464 directly in liquid nitrogen or placed in a container of isopentane (Merck) which was in turn 465 placed in liquid nitrogen at the same level as the isopentane. The publicly available snRNAseq datasets<sup>24</sup>, generated from samples obtained during elective surgeries performed at 466 467 Universitair Ziekenhuis Leuven, Belgium, were downloaded from the Sequence Read Archive 468 database.

469

#### 470 Nuclei isolation for snRNA-seq

Nuclei were isolated according to a protocol adapted from Habib et al. (2017)<sup>75</sup> and the 10X 471 472 Genomics nuclei isolation protocol (CG000124, User Guide Rev E). Frozen brain tissue was 473 homogenised in a dounce-homogeniser containing 2 ml ice-cold lysis solution (Nuclei EZ Lysis 474 Buffer [Sigma-Aldrich, NUC101] or Nuclei PURE Lysis buffer [Sigma-Aldrich, NUC201] with 1 475 mM dithiothreitol [DTT, Promega, P1171, US] and 0.1% Triton X-100 [Sigma-Aldrich, NUC201-476 1KT, US]). Homogenisation was done 20 times with the loose pestle A followed by 20 times 477 with the tight pestle B. An additional 2 ml lysis solution was added, and the sample was 478 incubated for 5 minutes on ice. The sample was centrifuged at 500 x g for 5 minutes at 4°C 479 after which the supernatant was discarded and the nuclei resuspended in 3 ml ice cold nuclei 480 suspension buffer (1x phosphate-buffered saline [PBS, Sigma-Aldrich, P4417-50TAB, US]), 481 0.01% bovine serum albumin [BSA, Sigma-Aldrich, A2153-10G, US], and 0.2 U/µl RNAsin Plus 482 RNase inhibitor [Promega, N2615, US]). Resuspended nuclei were passed through a 40 µm 483 filter and centrifuged at 900 x g for 10 minutes at 4°C. The supernatant was discarded and 484 pelleted nuclei were resuspended in 3 ml blocking buffer (1xPBS [Sigma-Aldrich, P4417-485 50TAB, US], 1% BSA [Sigma-Aldrich, A2153-10G, US], 0.2 U/µl RNAsin Plus RNase inhibitor 486 [Promega, N2615, US]).

#### 487

To remove myelin debris, 30 µl of myelin removal beads [Miltenyi Biotec. 130-096-733, US] 488 489 was added to the solution which was mixed by gently pipetting 5 times. The sample was 490 incubated for 15 minutes at 4°C after which it was mixed with 3 ml blocking buffer and 491 centrifuged at 300 x g for 5 minutes at 4°C. The supernatant was removed and the nuclei were 492 resuspended in 2 ml clean blocking buffer. The sample was transferred to a 2 ml tube and placed on a Dynamag magnet for 15 minutes at 4°C. The supernatant was transferred to a 493 494 new tube and stored on ice. An aliquot of trypan blue stained nuclei was counted using a 495 haemocytometer to determine the nuclei concentration and the volume to use in snRNA-seq 496 library preparation.

497

## 498 **10X Genomics snRNA-seq library preparation**

snRNA-seq library preparation was carried out using the 10x Genomics Chromium Next Gem 499 500 Single Cell 3' Reagent Kit (v3.1) according to manufacturer's protocols (CG000204, User Guide 501 Rev D), targeting 10,000 nuclei per sample. At step 2.2d and 3.5e, the libraries were amplified using 11 cycles and 13 cycles, respectively. Library quality and concentration was assessed 502 using either the TapeStation or Bioanalyser (Agilent) and Qubit (Invitrogen) at the Central 503 Analytical Facility (CAF, University of Stellenbosch). cDNA libraries were sequenced by 504 Novogene (Singapore) on either the Illumina HiSeq or NovaSeq system using the Illumina High 505 Output kits (150 cycles). 506

507

## 508 snRNA-seq read alignment and gene expression quantification

Fastq files were aligned to the human reference transcriptome (GRCh38) and quantified using
the count function from the 10X Genomics Cell Ranger v6.1.1 software (Cell Ranger, RRID
SCR\_017344) (Code availability: script 1). The inclusion of introns was specified in the count
function. An automatic filtering process was performed to remove barcodes corresponding
to background noise which have very low UMI counts.

514

## 515 snRNA-seq quality control

The resulting count matrices were processed using a pipeline adapted from the Harvard Chan 516 Bioinformatics Core (https://hbctraining.github.io/scRNA-seq online/). The filtered gene 517 barcode matrix for each sample was imported into R (V.4.2.0) using the Read10X function 518 from the Seurat (v.2.0)<sup>25</sup>. Nuclei-level filtering was performed to remove poor quality nuclei 519 520 according to their number of UMIs (nUMIs) detected, number of genes detected (nGene), 521 number of genes detected per UMI (log10GenesPerUMI), and the fraction of mitochondrial read counts to total read counts (mitoRatio) (Code availability: script 2). Nuclei that met the 522 following criteria were retained: nUMI > 500, nGene > 250, log10GenesPerUMI > 0.8 and 523 524 mitoRatio < 0.2. Gene-level filtering was performed to remove genes that had zero counts in 525 all nuclei, remove genes expressed in fewer than 10 nuclei, and remove mitochondrial genes 526 from the gene by cell counts matrix. Three doublet removal tools namely

527 DoubletFinder<sup>76</sup>(Code availability: script 3), DoubletDecon<sup>77</sup> (Code availability: script 4), and 528 Scrublet<sup>77</sup> (Code availability: script 5,6) were used to identify doublets for each dataset 529 individually. The sample-specific parameters of each of the tools were adjusted according to 530 the specified guidelines. To achieve a balance between the false positive and false negative 531 rate of the different doublet detection tools, all doublets identified by DoubletFinder as well 532 as the intersection of the doublets identified by DoubletDecon and Scrublet, were removed<sup>77</sup>.

533

## 534 snRNA-seq data normalization, integration and clustering

535 Principal component analysis was performed to evaluate known sources of within-sample 536 variation between nuclei, namely the mitoRatio and cell cycle phase (Code availability: script 7). The UMI counts of the 3000 most variable features were normalised and scaled on a per 537 sample basis by applying Seurat's SCTransform function with mitoRatio regressed out. A 538 539 Uniform Manifold Approximation and Projection (UMAP) analysis was performed on the 540 merged object to assess whether integration was necessary. The datasets were subsequently 541 integrated using Seurat's SelectIntegrationFeatures, PrepSCTIntegration, FindIntegrationAnchors, and IntegrateData functions (Code availability: script 7). To cluster 542 543 the datasets following integration, dimensionality reduction was first performed using UMAP 544 embedding, specifying 40 dimensions (Code availability: script 8). The Seurat FindClusters 545 function was then applied at a resolution of 0.8.

546

## 547 snRNA-seq cluster annotation

548 Two levels of annotation were performed. Clusters were initially annotated as one of the major brain cell types (level 1 annotation) based on the expression of known markers genes 549 550 (Code availability: script 9). Label transfer was then performed using Seurat's TransferData function with Allen Brain Map MTG atlas<sup>1</sup> as a reference dataset (level 2 annotation) (Code 551 552 availability: scripts 10-11). This resulted in each barcode in the query dataset receiving a predicted annotation based on a similarity score to an annotated cell type in the reference. 553 554 Barcodes were then filtered to remove those with discordant level 1 and level 2 annotations (e.g. barcodes with "oligodendrocyte" level 1 annotation and "Exc L4-5 RORB FOLH1" level 555 556 2 annotation) (Code availability: script 12). To validate the annotation, the expression of 557 known marker genes was assessed. Cosine similarity scores were computed to compare the 558 transcriptomic similarity of each of the annotated query cell types to the 75 reference MTG 559 cell types using the SCP package (https://github.com/zhanghao-njmu/SCP) (Code availability: 560 script 13). This was achieved by computing cosine similarity scores for each pair of query and reference cell types using the expression of the top 2000 shared highly variable features 561 between the query and reference datasets. The log normalised expression counts were used 562 for this purpose (RNA assay, data slot). To assess the difference between the paediatric and 563 adult datasets relative to the reference, the above cosine similarity analysis was repeated on 564 the paediatric and adult datasets individually (Code availability: script 13). 565

#### 567 NS-Forest machine learning marker analysis of snRNA-seq datasets

The NS-Forest tool  $(v2.0)^{29,30}$  was used to identify combinations of marker genes uniquely 568 defining each annotated cell type (Code availability: script 14-15) in the paediatric and adult 569 570 datasets separately. The number of nuclei per sample was randomly down-sampled to that 571 of the sample with the fewest nuclei (n=4,865). A random-forest model was used to select a 572 maximum of 15 marker genes per cell type based on them being both highly expressed as 573 well as uniquely expressed within a cell type compared to other cell types (i.e., the top Gini 574 Index ranked features with positive expression values). The number of trees chosen for this 575 model was 30,000, the cluster median expression threshold was set to the default value of 576 zero, the number of genes used to rank permutations of genes by their F-beta-score was 6, 577 and the beta weight of the F score was set to 0.5. The aforementioned parameters were set according to the parameters described in Aevermann et al. (2021)<sup>29</sup>, allowing the outputs to 578 579 be directly compared to their markers and to the Allen Brain Map MTG atlas minimal 580 markers<sup>1</sup>. To assess the relevance of these markers in terms of their capacity to distinguish 581 different cell types in a UMAP analysis, the SCT and integration methods were repeated using either a random set of genes or the NS-Forest markers as anchors<sup>29</sup> (Code availability: script 582 583 16).

584 585

## 586 **DESeq2 age-dependent differential gene expression analysis of snRNA-seq datasets**

587 DESeq2<sup>33</sup> was used to identify genes that were differentially expressed with age (Code 588 availability: script 17). The unnormalized counts were aggregated across all nuclei for each 589 cluster and sample to generate a 'pseudobulk' counts matrix with the counts from technical 590 replicates collapsed to the level of biological replicates. Genes were filtered to only include 591 those expressed in more than 10% of nuclei for a given cell type. Principal component analysis 592 was performed on each cell type separately in order to assess the variation between samples 593 and determine which variables were contributing most to inter-sample variation from a set of possible variables. The collapsed counts served as input into DESeq2's 594 595 DESeqDataSetFromMatrix function in which the design formula ~single cell chemistry + age group was specified to treat the age group (paediatric vs adult) as the variable of interest 596 597 while the effect of single cell chemistry (version2 vs version3 chemistry) was regressed out. 598 A hypothesis test was performed using the Wald test. The null hypothesis for each gene was 599 that there is no difference in gene expression between the sample groups (i.e Log2Fold 600 Change = 0). A Wald test statistic was determined for each gene together with the associated 601 p-value after which the p-values were adjusted for multiple testing using the Benjamini-602 Hochberg method. Positive log2 Fold Changes represent genes which are upregulated in 603 paediatric samples compared to adult samples (p<sub>adi</sub> <0.05).

604

#### 605 Pseudotime trajectory analysis with psupertime

607 To validate the differentially expressed genes identified with DESeq2, a pseudotime trajectory analysis was performed for a subset of excitatory neuron subtypes using the psupertime 608 package<sup>55</sup> (Code availability: script 18). Psupertime is a supervised approach that uses time-609 610 series labels as input to improve the identification of time-varying genes. Each cell type of 611 interest (Exc L3-5 RORB ESR1, Exc L2 LAMP5 LTK, Exc L4-5 RORB FOLH1B, Exc L2-3 LINC00507 FREM3) was individually sub-setted from the Seurat object after which a single cell 612 613 experiment (sce) object was generated using the log normalized counts (RNA assay, data slot). 614 The donor age (4, 5, 7, 9, 15, 20, 24, 26, 31, 41, 50) was included as metadata in the object. 615 The psupertime function was applied to the sce object with the sel genes argument 616 specifying all genes to be used. An automatic filtering step was performed to remove genes 617 expressed in fewer than 10% of cells for each cell type. As an output of the function, the beta 618 coefficients for the association of each gene with pseudotime were extracted and plots were 619 generated showing the expression trajectories of the DESeg2 DEGs with pseudotime. 620 Additionally, the overlap between the DESeq2 DEGs and genes changing as a function of 621 pseudotime (Psupertime-relevant genes) was determined.

622

## 623 Pathway enrichment analysis of snRNA-seq datasets

- 624 GO analysis of NS-Forest marker genes was performed on the gProfiler web server<sup>78</sup> using 625 default settings ( $p_{adj} < 0.05$ ) with "highlight diver terms in GO" selected.
- 626

DEGs identified by DESeq2 (see Extended Data Table 10) that were associated with EA and IQ, as well as those associated with accelerated evolution in humans (HARs), were determined by comparing the list of neuronal DEGS to the EA, IQ and HAR gene lists used by Driessens et al. (2023)<sup>56</sup>, which were subsets of the lists from Lee et al. (2018)<sup>58</sup>, Savage et al. (2018)<sup>59</sup> and Doan et al. (2016)<sup>60</sup> respectively. A hypergeometric test was performed to test the significance of the results relative to chance (Code availability: script 19).

633

634 GSEA on the DESeq2 output for all genes was performed using the Broad Institute's GSEA 635 software (https://www.gsea-msigdb.org/gsea/msigdb) (Code availability: script 20). GSEA 636 aggregates the information from many genes to identify enriched functional pathways, 637 allowing us to interrogate the gene signature changes across all cell types, including those that did not show any significant DEGs<sup>66</sup>. The gene lists for each cell type were queried against 638 639 the C5 GO Biological Processes collection comprising of gene sets derived from the GO 640 Biological Process ontology. The input lists of genes were ranked according to the -log(p-641 value)\*log<sub>2</sub>FoldChange for each gene. The parameters specified to the GSEA function 642 included number of permutations (nperm)=1000, minimum gene set size (set min=15), 643 maximum gene set size (set max=200), excludes genes that have no gene symbols (collapse)= 644 No\_Collapse, value to use for the single identifier that will represent all identifiers for the 645 gene (mode)=Max probe, normalised enrichment score method (norm)= meandiv, weighted 646 scoring scheme (scoring\_scheme) = classic. Positive Normalised Enrichment Scores (NES) 647 represent genes that were upregulated in the paediatric population compared to the adult

648 population (p<0.01 and q<0.1). To visualise the output of universally enriched pathways 649 across multiple cell types, the top 25 most frequently appearing positively and negatively 650 associated terms were plotted. Additionally, for five cell types of interest [which had DEGs 651 meeting the threshold of p<0.05 and abs(log<sub>2</sub>FC)>0.1], the top 5 positively associated terms 652 were plotted.

653

## 654 Analysis of site-of-disease TBM markers

655

The dittoheatmap function from the dittoSeq package<sup>79</sup> was used to generate heatmaps for 656 657 the expression of the TBM biomarkers (upregulated genes listed in Rohlwink et al. 2019, Supplementary Table 5<sup>23</sup>) across cell types in the paediatric and adult datasets individually. 658 Additionally, Seurat's dotplot function<sup>25</sup> was used to visualize the level of expression and 659 proportion of nuclei expressing the markers across cell types (Code availability: script 21). 660 661 Prior to generating the plots, the TBM marker genes were filtered to remove those expressed 662 in 15 nuclei or fewer across all cell types. Gene counts for each marker were aggregated across cell types and scaled. The markers were clustered according to their expression profiles 663 using dittoheatmap's default hierarchical clustering method (Euclidean, complete). The 664 665 clustering order and dendrogram from this output for the peaditaric datasets were used to generate dotplots for both peaditaric and adult datasets (Code availability: script 21). 666

667

## 668 snRNA-seq data plots

Plots were produced with Seurat<sup>25</sup>, ggplot2<sup>80</sup>, ShinyCell<sup>81</sup> and Microsoft Excel.

670

## 671 **10x Genomics Visium library preparation**

672 Frozen OCT embedded temporal cortex tissue samples were scored using a pre-chilled razor 673 blade to fit in the Spatial Gene Expression slide capture areas. 10 µm-thick sections were cut 674 using a cryostat (Leica CM1860/CM1950) and collected onto the Spatial Gene Expression slide 675 capture areas. Two replicate sections of the 15-year-old (10 µm apart) and two replicate 676 sections of 31-year-old (40 µm apart) were collected. The spatial Gene Expression slides with 677 tissue sections were stored in a sealed container at -80°C. Captured sections were 678 Haematoxylin and Eosin (H&E) stained according to the 10x Genomics Demonstrated Protocol 679 Guide (CG000160, Rev B). Brightfield images of the stained sections were captured using an 680 EVOS M5000 microscope (Thermo Fisher Scientific) at 20x magnification without 681 coverslipping. Overlapping images of the sections including the fiducial frame were stitched 682 together using Image Composite Editor-2.0.3 (Microsoft). Visium libraries were prepared from the stained tissue sections following the Visium Spatial Gene Expression Reagents Kit 683 User Guide (CG000239, Rev D). At Step 1.1 the tissue was permeabilised for 12 minutes as 684 determined using the Visium Spatial Gene Expression Tissue Optimisation User Guide 685 (CG000238, Rev D). At Step 3.2, cDNA was amplified using 20 cycles. Library quality and 686 concentration was assessed using TapeStation (Agilent) and Qubit (Invitrogen) at the Central 687

Analytical Facility (CAF, University of Stellenbosch). Libraries were sequenced by Novogene
(Singapore) on the Illumina NovaSeq system using the Illumina High Output kits (150 cycles).

690

## 691 Visium read alignment and gene expression quantification

The H&E images were processed using the 10X Genomics Loupe Browser V4.0 Visium Manual
 Alignment Wizard. 10X Genomics Space Ranger *count* (10X Space Ranger V1.3.0) was used to
 perform alignment of FASTQ files to the human reference transcriptome (GRCh38), tissue
 detection, fiducial detection and barcode/UMI counting.

696

#### 697 cell2location analysis of Visium datasets

The average number of nuclei per Visium spot was determined using Vistoseg<sup>82</sup> (Code 698 availability: script 22). Cell2location (version 0.7a0)<sup>26</sup> was used to spatially map the brain cell 699 types by integrating the Visium data count matrices (Space Ranger output) with the 700 701 annotated snRNAseg datasets (Code availability: script 23). To avoid mapping artifacts, 702 mitochondrial genes were removed from the Visium datasets prior to spatial mapping. 703 Reference signatures of the 75 annotated cell populations were derived using a negative binomial regression model using the default values (Code availability: script 24). 704 705 Unnormalized and untransformed snRNA-seq mRNA counts were used as input in the 706 regression model for estimating the reference signatures (Code availability: script 24. The 707 snRNA-seq mRNA counts were filtered to 14,209 genes and 144,438 cells. The cell2location 708 model for estimating the spatial abundance of cell populations was filtered to 14,197 genes 709 and 14,324 cells that were shared in both the snRNA-seq and Visium data. The following 710 cell2location parameters were used: training iterations = 30,000 cell per location, N<sup>^</sup> = 7 711 (estimated using Vistoseg segmentation results), Normalization (ys) alpha prior = 20 (Code 712 availability: script 25). To visualise the cell abundance in spatial coordinates 5 % quantile of 713 the posterior distribution was used, which represents the value of cell abundance that the 714 model has high confidence in (Code availability: script 26). Cell2location's Non-negative 715 Matrix Factorization (NMF) was used to identify cellular compartments and cell types that co-716 locate from the cell type abundance estimates. NMF was tested using a range of factors (5 to 717 30) for the "n fact" parameter (Code availability: script 26). n fact=15 was chosen as it clearly 718 grouped the oligodendrocyte, astrocyte and excitatory neuron cell sub-types into known 719 tissue zones i.e. the layers of the cortex (Code availability: script 27).

720

## 721 BayesSpace analysis of Visium datasets

The raw gene expression counts from Space Ranger were normalized, log transformed and principal component analysis was performed on the top 2000 highly variable genes. To obtain high-resolution gene expression for selected genes, the principal component values were mapped back to their original log-transformed gene expression space (spot level) using the default BayesSpace<sup>34</sup> regression (Code availability: script 28). To do this the principal components from the original data were used as predictors in training the model for each gene, in which the results were the measured gene expression at the spot level. The trained

model was then used to predict the gene expression at sub spot level using high resolution
 PCs. The high-resolution model was trained using default values except for the following
 parameters: 7 PCs, Number of clusters = 8, nrep = 100,000, burn-in = 10,000. The BayesSpace

- outputs for each sample were quantified for spots with expression level > 0 and displayed as
- 733 boxplots (Code availability: script 29).
- 734

## 735 In situ Hybridisation Chain Reaction (HCR) on frozen human tissue sections

10 µm thick frozen sections were collected on Histobond+ slides (Marienfeld) and stored at -736 737 20°C. The In situ HCR protocol was carried out on tissue sections as detailed in Choi et al. (2016)<sup>83</sup> using reagents, probes and hairpins purchased from Molecular Instruments. Probes 738 739 were ordered for the following genes: RELN (NM 005045.4), FABP7 (CR457057.1), AQP4 740 (NM 001650.5), RORB (NM 006914.4), CLSTN2 (NM 022131.3) and TSHZ2 (NM 173485.6). 741 When necessary to quench lipofuscin autofluorescence, sections were rinsed after HCR in 1x 742 PBS and treated with 200 µl TrueBlack (Biotium) for 30 sec. Slides were rinsed in PBS, stained 743 with Hoescht (Thermofisher) and mounted using SlowFade Gold Antifade Reagent 744 (Invitrogen). Sections were imaged using the LSM 880 Airyscan confocal microscope (Carl 745 Zeiss, ZEN SP 2 software) using the 40X or 60X objective.

746

## 747 MERFISH analysis on frozen temporal cortex tissue sections

748 10 µm thick frozen sections were cut from frozen OCT embedded temporal cortex tissue 749 samples using a cryostat (Leica CM1950). Sections from a peaditaric and adult sample were 750 collected onto the same MERSCOPE coverslip (VIZGEN 2040003), fixed and stored in 70% 751 ethanol following the instructions in the VIZGEN protocol (Fresh & Fixed Frozen Tissue 752 Sectioning & Shipping Procedure Rev A, Doc. number 91600107). The slide was processed on 753 the VIZGEN MERSCOPE system by the MRC Weatherall Institute of Molecular Medicine Single 754 Cell Facility (University of Oxford) within 1 month of storage. Sections were photobleached 755 for 10 hours at 4°C and then washed in 5 ml Sample Prep Wash Buffer (VIZGEN 20300001) in 756 a 5 cm petri dish. Sections were incubated in 5 ml Formamide Wash Buffer (VIZGEN 757 20300002) at 37°C for 30 min and hybridized at 37°C for 36 to 48 hours by using 50  $\mu$ l of 758 VIZGEN-supplied custom Gene Panel Mix according to the manufacturer's instructions. 759 Following hybridization, sections were washed twice in 5 ml Formamide Wash Buffer for 30 760 min at 47°C. Sections were then embedded in acrylamide by polymerizing VIZGEN Embedding 761 Premix (VIZGEN 20300004) according to the manufacturer's instructions. Following 762 embedding, sections were digested in Digestion Pre-Mix (VIZGEN 20300005) and RNase 763 inhibitor (New England Biolabs M0314L) for 3 h at 37°C and then cleared for 16 to 24 hours 764 with a mixture of VIZGEN Clearing Solution (VIZGEN 20300003) and Proteinase K (New England Biolabs P8107S) according to the Manufacturer's instructions. Following clearing, 765 766 sections were washed twice for 5 min in Sample Prep Wash Buffer (PN 20300001) and then stained with VIZGEN DAPI and PolyT Stain (PN 20300021) for 15 min followed by a 10 min 767 768 wash in Formamide Wash Buffer. Formamide Wash Buffer was removed and sections were

769 washed with Sample Prep Wash Buffer during MERSCOPE imaging set up. A mixture of 100 ml of RNAse Inhibitor (New England BioLabs M0314L) and 250 ml of Imaging Buffer Activator 770 771 (PN 203000015) was added to the cartridge activation port to a prethawed and mixed MERSCOPE Imaging cartridge (VIZGEN PN1040004). 15 ml mineral oil (Millipore-Sigma 772 773 m5904-6X500ML) was added on top of the activation port and the MERSCOPE fluidics system 774 was primed according to VIZGEN instructions. The flow chamber was assembled with the 775 section coverslip according to VIZGEN specifications and the imaging session was initiated 776 after collection of a 10X mosaic DAPI image and selection of the 1cm<sup>2</sup> imaging area. MERFISH 777 data was visualised using the VIZGEN MERSCOPE Vizualizer software (version 2.3.3330.0).

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## 1000 Data Availability

All scripts used to analyse the data are indicated in the methods section and are available in the supplementary material. A description of the raw and analysed data files will be made available on the University of Cape Town's <u>ZivaHub</u> data sharing platform on publication. As the data is from living donors, access to the data will be mediated through contact with the corresponding author. A ShinyApp will be made publicly available on publication for exploration of the annotated snRNA-seq data.

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## 1027 Contributions

1028 C.S. and S.F. conducted the snRNA-seq experiments. C.S. conducted the majority of snRNA-1029 seq analyses. R.M. conducted the Visium and HCR experiments and analysis. J.M. conducted 1030 additional HCR experiments. S.Q. provided additional bioinformatics support. M.B.V. liaised 1031 with neurosurgeons and prepared all neurosurgical brain tissue samples. R.M., J.B. and 1032 J.M.N.E conducted the neurosurgeries and provided donor metadata. C.S., T.S.S., M.G. and 1033 D.H. conceptualised, conducted and analysed the MERFISH experiments. U.K.R., M.Z., J.V.R, C.G.D, A.F, and D.H. conceptualised the study and raised funds. D.H., C.S. and R.M wrote the 1034 1035 manuscript. D.H. conducted additional analyses and supervised the project. All authors read 1036 and commented on the manuscript.

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1043 Fig. 1: Annotation of nuclei by label transfer identifies 75 cell types across the 23 datasets. a, Data integration 1044 shows alignment of nuclei across the technical (T) and biological (B) replicates from donors ranging in age from 1045 4 to 50 years. b, UMAP plot annotated to show the 75 cell types from the Allen Brain Map MTG atlas after filtering 1046 to retain nuclei with high confidence annotations. Each cell type is annotated with 1) a major cell class (e.g. Exc 1047 for excitatory neuron), 2) the cortical layer the cell is associated with (e.g. L2 for layer 2), 3) a subclass marker 1048 gene and 4) a cluster-specific marker gene. c, Stacked barplot showing the proportion of nuclei per cell type for 1049 each age category out of the total number of nuclei for each group. The cell types are coloured as in b. See 1050 Extended Data Table 3 for details of statistical tests performed. d, Validation of the high-resolution cell type 1051 annotations shows a high degree of correspondence in the expression of known cell type-specific marker genes 1052 (x axis) with their expected cell type (y axis) (left). The number of nuclei per cell type is shown on the right. e, 1053 Correlation plot showing the cosine similarity scores assessing similarity between the annotated cell types in our 1054 dataset (y axis as in d) and the MTG reference dataset (x axis) based on the log normalized expression counts of 1055 the top 2000 shared highly variable features between query and reference datasets.



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1057 Fig. 2: Visium spatial transcriptomics in the adult and paediatric temporal cortex validates snRNA-seq annotation. 1058 **a**, Estimated cell type abundances (colour intensity) in the 31-year-old and 15-year-old temporal cortex tissue 1059 sections for a selection of cell types including non-neuronal cell types, excitatory neurons (top row) and inhibitory 1060 neurons (bottom row). **b**, Visium gene expression profiles (colour intensity) for a selection of known cortical layer 1061 marker genes in the 31-year-old and 15-year-old temporal cortex tissue sections including *AQP4* (layer 1), *LAMP5* 1062 (layer 2), RORB (layer 4) and *CLSTN2* (layer 5-6). **c,d**, Identification of co-locating cell types using NMF. The dot plot 1063 (**c**) shows the NMF weights of the cell types (rows) across each of the NMF factors (columns), which correspond to 1064 tissue compartments. Block boxes indicate cell types that co-locate within the indicated compartments. Spatial plots 1065 show (**d**) show the NMF weights for selected NMF factor/tissue compartment across the 31-year-old and 15-year-1066 old temporal cortex tissue sections. Panels are displayed in the same order as the dotplot in (**c**), with the dominant 1067 cell types for each factor indicated in brackets. Dashed white lines and numbers indicate estimated cortical layer 1068 boundaries as indicated in the first two panels of **b** and **d**. WM: white matter. See also Extended Data Figs 4-6.





Fig. 3: NS-Forest identifies minimal marker genes distinguishing the cell types in the paediatric and adult temporal cortex snRNA-seq datasets. a,b, Heatmap showing the scaled average normalised expression counts of the NS-Forest minimal marker genes (y-axis) identified for 75 cortical cell types (x-axis) across the six adult (a) and six paediatric (b) datasets. As input into NS-Forest, the nuclei of each sample were randomly down-sampled to the size of the sample with the fewest nuclei. Heatmaps show gene expression values for the down-sampled datasets. The minimal marker genes are annotated (colour codes on the y-axes) according to whether they are unique to a given cell type, whether they are coding/non-coding genes, whether they are unique to the indicated age group, whether they overlap with existing MTG minimal marker gene sets for the same cell type, and according to the cell type they define.







1088 Fig. 4: Validation of NS-Forest minimal markers and assessment of the top NS-forest markers. a,b, Annotated 1089 UMAP plots following data integration using either the minimal marker genes (left) or the equivalent number of 1090 a random set of genes (right) as anchors for the adult (a) and paediatric (b) datasets. The colour scheme for the 1091 cell types is in accordance with the MTG cell taxonomy. c, Overlap of the paediatric and adult NS-Forest markers 1092 with a high binary expression score (> 0.7) per cell type. The bar plot shows the number of shared markers 1093 between paediatric and adult datasets (blue), the number of markers unique to the paediatric datasets (orange), 1094 and the number of markers unique to the adult datasets (grey) for each cell type.

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1102 Fig. 5: Differential expression analysis reveals genes guiding temporal cortex maturation. a, 21 cell types with 1103 DEGS. b-g, Volcano plots showing log<sub>2</sub>FoldChange (x axis) and -log<sub>10</sub>padj values (y axis) for all DESeq2-tested 1104 genes in Exc\_L3-5\_RORB\_ESR1, Exc L2-3\_LINC00507\_FREM3, Exc\_L4-5\_RORB\_FOLH1B, Exc\_L2\_LAMP5\_LTK, 1105 Astro\_L1-6\_FGFR3\_SLC14A1 and Oligo L1-6 OPALIN. Red dots indicate genes that were significantly upregulated 1106 or downregulated in paediatric samples (padj<0.05 & abs(log<sub>2</sub>FoldChange)>10%) and selected genes are 1107 labelled. Red labels indicate DEGs shared between neuronal cell types. Magenta labels indicate DEGS not shared 1108 between cell types that are discussed in the text. Blue dots indicate non-significant genes (padj>0.05 or 1109 abs(log<sub>2</sub>FoldChange)<10%). h, Dot plot showing the scaled average normalised expression across samples for 1110 DEGS shared between Exc\_L3-5\_RORB\_ESR1, Exc L2-3\_LINC00507\_FREM3, Exc\_L4-5\_RORB\_FOLH1B, 1111 Exc L2 LAMP5 LTK, Exc L3-4 RORB CARM1P1 and Exc L3-5 RORB FILIP1L. i, psupertime gene expression trajectories for selected DEGs in the indicated cell types. The x-axis is the calculated psupertime value for each 1112 1113 cell, coloured by sample of origin. The black lines are smoothened curves fit by geom smooth in the R package 1114 ggplot2.

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1120Fig. 6: Pathways that are enriched or depleted across multiple cell types in paediatric samples. GSEA heatmap1121showing the top 25 most frequently enriched (top 25 rows) or depleted (bottom 25 rows) terms appearing across1122all cell types. Abbreviations in bold indicate the following categories as referred to in the text: AE, axon1123ensheathment; CR, cellular respiration; ICT, intracellular transport; NM, neuronal morphogenesis; NR/SP,1124neurotransmitter release/synaptic plasticity; PT/M, protein translation/modification. Only significantly (p < 0.01</td>1125and q < 0.1) terms are shown. NES value represents the normalized enrichment scores. Grey indicates that the</th>1126term was not significantly enriched or depleted in the indicted cell type. See also Extended Data Table 12.





1137 Extended Data Fig. 1: Nuclei quality control (QC) and clustering. a, Number of doublets identified across all 23 1138 datasets by DoubletDecon, DoubletFinder, and Scrublet. Red outline indicates the subset of barcodes called as 1139 doublets that were removed. b, Total number of nuclei per dataset before (yellow) and after (green) QC. c, Mean 1140 number of reads per nucleus (y axis) by dataset before QC split by age group (x axis). p value determined by two-1141 tailed Welch's t-test. d, Number of nuclei (y axis) by sample after QC split by age group (x axis). p value 1142 determined by Brunnermunzel permutation test. e, Violin plots showing the number of unique molecular 1143 identifiers (UMIs) (top) and the number of genes detected (bottom) per nucleus per sample after QC. Black dots 1144 indicate the median value. Error bars show 95% confidence intervals. f,g, Median number of UMIs (2,263 1145 paediatric and 2,011 adult) (f) and the median number of genes (1,372 paediatric and 1,226 adult) (g) detected

bioRxiv preprint doi: https://doi.org/10.1101/2023.09.29.560114; this version posted May 17, 2024. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is per nucleus (y axes) by smapplevalitededdepic Og Yage-group (keawis) op lively as determined by two-tailed
 Brunnermunzel permutation test. h, UMAP plot for the 23 datasets prior to integration. i, UMAP plot showing the resulting clusters determined by the shared nearest neighbour algorithm. Data in all box plots represent mean ± sem for six paediatric and six adult samples. No significant differences were detected between paediatric and adult samples. B, biological replicate; NS, not significant; T, technical replicate. See also Extended Data Table

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1155 Extended Data Fig. 2: Annotation and assessment of cell composition across datasets. a, UMAP plot showing 1156 cluster annotation at the level of major brain cell types (level 1 annotation). b, Examination of known cell type-1157 specific marker genes (x axis) after label transfer classify each nucleus according to the Allen Brain Map MTG 1158 atlas<sup>1</sup> (level 2 annotation) (y axis) (left). Off-target gene expression is evident in several cell types (marked in red), which is likely due to multiplets or nuclei contaminated with ambient mRNA. c-d, Stacked barplots after filtering 1159 1160 to retain nuclei with high confidence annotations showing the proportion of nuclei per cell type (y axis) for each 1161 technical replicate (c) or biological replicate (d) (x axis) out of the total number of nuclei for each group. Samples 1162 with technical replicates showed high degrees of similarity in cell composition between their replicates (c). 1163 Technical replicates from each donor were merged to allow comparisons between the 12 samples (d).

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Extended Data Fig. 3: Assessment of the sequencing metrics for the annotated cell types. a, Violin plots showing the distribution of the number of genes (left) and transcripts (right) detected per nucleus per cell type across all datasets. Black dots indicate the median value. Error bars show 95% confidence intervals. b,c, Boxplots showing the number of genes (b) and the number of UMIs (c) (y axis) detected per cell type per sample (x axis) split by age group (red: adult, grey: paediatric). Data in all box plots represent mean ± sem for six paediatric and six adult samples for each cell type. No significant differences were detected (i.e. padj > 0.05). See Extended Data Table 3 for details of statistical tests performed.

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1190 Extended Data Fig. 5: Spatial mapping of cell types in the human temporal cortex. a, Estimated cell abundance 1191 of 75 cell types across all Visium samples. Shown is a heatmap with the colour indicating the relative cell 1192 abundance of cell types (rows) across the different samples (columns). b, Estimated cell type abundances (colour 1193 intensity) in the technical replicate 31-year-old and 15-year-old temporal cortex tissue sections for a selection 1194 of cell types including non-neuronal cell types, excitatory neurons (top row) and inhibitory neurons (bottom 1195 row). d, Spatial plots show of the NMF weights for selected NMF factor/tissue compartment across the 31-year-1196 old and 15-year-old temporal cortex tissue sections. Panels are displayed in the same order as the dotplot in Fig. 1197 2c, with the dominant cell types for each factor indicated in brackets. T, technical replicate.



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1202 Extended Data Fig. 6: In situ HCR analysis of selected cortical layer marker genes. Expression of a, layer 1 1203 markers AQP4, FABP7 and RELN and b, layer 4-6 markers RORB, CLSTN2 and TSHZ2 in frozen temporal cortex 1204 tissue sections from the same 31-year-old and 15-year-old donor tissue used for Visium. High magnification views 1205 of layer 1 in a indicate AQP4/RELN-positive cells (yellow arrowheads) and FABP7 positive cells (green arrowhead). 1206 In high magnification views of layer 4 in **b** in the 31-year-old tissue section, RORB/CLSTN2-positive (white 1207 arrowhead) and RORB/TSHZ2-positive cells (green arrowhead) are indicated. In high magnification views of layer 1208 4 in **b** in the 15-year-old tissue section RORB/CLSTN2/TSHZ2-positive cells (white arrowheads) are indicated. 1209 Dashed white lines indicate layer boundaries. Solid white line indicates tissue edge. Scale bars are 100  $\mu$ m in low 1210 magnification views (tile scan at 40x) and 20 µm in high magnification views (63x).

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Extended Data Fig. 7: Expression of the reference MTG atlas minimal markers. Heatmap showing the scaled average normalised expression counts of the NS-Forest minimal marker genes identified for the reference MTG cell atlas dataset (y-axis) in each of the 75 query cortical cell types identified in the combined adult and paediatric snRNA-seq datasets (x-axis). The minimal marker genes are annotated (colour codes on the y-axes) according to the cell type they define.

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Extended Data Fig. 8: MERFISH spatial transcriptomics analysis of selected NS-forest markers. a, b Low magnification views of the 31-year-old (a) and 15-year-old (b) MERFISH datasets showing the expression of known layer maker genes in the expected layers as validation of the MERFISH experiment. c-p, High magnification views of 31-year-old (c,e,g,I,k,m,o) and 15-year-old (d,f,h,j,I,n,p) MERFISH datasets showing the overlap of new NS-Forest minimal markers (green) with published NS-Forest minimal markers (magenta) in indicated cells (arrowheads). The cell type that the NS-Forest markers are associated with is indicated in the top left corner. Scale bars: 100 μm.



Extended Data Fig. 9: Evaluation of NS-Forest minimal marker gene expression across cell types in comparison
 to MTG cell taxonomy markers. a-d, Boxplots showing the normalised expression counts for *LINC01331* (a),
 *PALMD* (b), *POSTN* (c) and *OLFML2B* (d) in paediatric (top) and adult (bottom) datasets. The cell types expressing
 the markers at high levels are indicated in bold.

bioRxiv preprint doi: https://doi.org/10.1101/2023.09.29.560114; this version posted May 17, 2024. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a CC-BY-NC-ND 4.0 International license. 31-year-old 2.0 Log-Counts 1.5 FNBP1L 1.00 0.75 0.50 0.25 0.9 0.6 0.3 0.5 0.0 31-T1 31-T2 15-T\* 15-T2 Visium sample KCNG1 2.5 2.0 Log-Counts 1.5 1.00 0.75 0.50 0.25 KCNG1 0.9 0.6 0.3 0.0 0.5 0.0 31-T1 31-T2 15-T1 15-T2 Visium sample SLC35F4 2.5 2.0 Log-Counts 1.5 1.6 1.2 0.8 0.4 0.0 SLC35F4 1.25 1.00 0.75 0.50 0.25 0.5 0.0 31-T1 31-T2 15-T1 15-T2 Visium sample MYO16 2.5 20 Log-Counts 1.5 1.0 MYO16 1.0 0.5 0.5 1.0 0.0 0.0 0.5 0.0 31-T 31-T2 15-T1 15-T2 Visium sample XKR4 2.5 2.0 Log-Counts 1.5 XKR4 1.6 1.2 0.8 0.4 1.0 0.5 0.0 1.0. 0.5 0.0 31-T1 15-T2 15-T Visium sample LAMC3 25. 2.0 2.0 1.5, Conuts 1.0, LAMC3 1.5 1.0 0.5 0.0 1.5 1.0 0.5 0.0 0.5 0.0. 31-T1 15-T1 15-T2

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Extended Data Fig. 10: BayesSpace analysis of differentially expressed genes. High resolution Visium spatial gene expression profiles for selected DEGs using BayesSpace analysis to compare sub-spot level expression intensities between 31-year-old and 15-year-old temporal cortex tissue sections. Barplots show the average gene gene expression (log-counts) across technical replicate (T) samples for the indicated genes for spots with gene expression levels > 0. In all cases, average gene expression is higher in the 15-year-old samples than in the 31year-old samples.

Visium sample



**Extended Data Fig. 11: Cell type-specific expression of putative TBM biomarkers**. **a**. Hierarchical clustering of TBM biomarker genes across the 75 cell types identified in the peaditaric snRNA-seq dataset reveals clusters of genes that are expressed by specific groups of cell types. **b**. Analysis of the same genes across the adult snRNA-seq dataset, using the gene order in (a) reveals very similar patterns of cell type-specific expression across the age-groups. Dashed boxes highlight gene clusters, with associated cell types indicated on the left and right of the right diagram

FGFR3 THEMIS SLC14A1 RORE FEZF2 DIO2 Expression Level Expression Level GFAP CUX1 AQP4 ID3 CUX2 тлс SLC17A7 SLC1A2 SATB2 cluste Inhibitory neurons Microglia PAX6 TYROB LAMPS СЗ VIP CX3CR1 Level Expression Level SST Expression CSF1R PVALB LHX6 **DOCK8** DARB GAD1 ADAM28 clu clust Endothelial cells Oligodendrocytes OPALIN NOSTRIN PLP1 месом MAG CLDN5 evel Level KLK6 HLA-E Expression I cion мовр Expre ABCB1 FLT1 ST18 IGFBP7 MBF HLA-B RNF220 cluste Pericytes Oligodendrocyte precursor cells PDGFRA ATP1A2 COL20A1 DLC1 OLIG2 ITIH5 eve Expression Leve PRRX1 LAMA2 SEMA5A Expre DCN VCAN SLC6A12 TNR SLC19A1 PCDH15 EBF1



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Supplementary Fig. 1: Violin plots showing the expression of known cell type marker genes across the Seurat
 clusters. These data were used for level 1 annotation of each cluster as one of the indicated major brain cell types
 (see also Extended Data Table 3).

18 19 20 21 2 cluster

clust



Supplementary Fig. 2: psupertime gene expression trajectories for all DEGs in Exc\_L3-5\_RORB\_ESR1. The *x*-axis
is the calculated psupertime value for each cell, coloured by sample of origin. The black lines are smoothened
curves fit by geom\_smooth in the R package ggplot2.

ST8SIA2 CELSR1 ADAMTS18 SLC35F4 SPHKAP MEPE GRP CPXM2 LAMC3 KCNG1 RERGL UACA LINC02653 LINC01837 Donor age .2 (years) LINC01965 TIE1 ATP11C SOX11 LINC00922 FNBP1L OTOF . z-scored log2 expression 10-F .2 **GRIN3A FBN3** AC243829.1 HGF AC093898.1 C8orf34 PCAT1 Δ 

HPN

-2

PPARGC1A

AC243829.2

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CFAP58

POPDC3

ZNF229

AF279873.3

С

psuedotime

.2

AL512380.2

ZNF215

HPSE2

Supplementary Fig. 3: psupertime gene expression trajectories for all DEGs in Exc L2-3\_LINC00507\_FREM3. The

x-axis is the calculated psupertime value for each cell, coloured by sample of origin. The black lines are smoothened curves fit by geom\_smooth in the R package ggplot2.



Supplementary Fig. 4: psupertime gene expression trajectories for all DEGs in Exc\_L4-5\_RORB\_FOLH1B. The *x*-axis is the calculated psupertime value for each cell, coloured by sample of origin. The black lines are smoothened curves fit by geom\_smooth in the R package ggplot2.



Supplementary Fig. 5: psupertime gene expression trajectories for all DEGs in Exc\_L2\_LAMP5\_LTK. The *x*-axis is
 the calculated psupertime value for each cell, coloured by sample of origin. The black lines are smoothened
 curves fit by geom\_smooth in the R package ggplot2.

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Extended Data Table 1: Summary of snRNA-seq, Visium and MERFISH sample metadata. Samples are ordered
 by age. The eight "P00" datasets were generated in the Hockman laboratory while the four "Nuc" datasets were
 generated by Thrupp et al. (2020)<sup>24</sup>.

1301

#### Extended Data Table 2: Summary of average quality control metrics for snRNA-seq datasets across nuclei for each sample before and after filtering. Several measures for quality control were evaluated on a per sample basis including the sequencing saturation, the mean number of reads per nucleus, the number of barcodes, the median number of genes detected per nucleus, the median number of UMIs detected per nucleus, and the number of doublets removed.

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1308 Extended Data Table 3: Label transfer annotation of snRNA-seq datasets using the Allen Brain Map MTG atlas 1309 as a reference. Sheet 1, Manual annotation of clusters into major cell types (level 1 annotation). Sheet 2, Number 1310 of nuclei per level 1 annotation category per sample. Sheet 3, Number of nuclei per MTG cell type per sample. 1311 The number of barcodes corresponding to each MTG cell type and sample is shown. Additionally, the total, 1312 minimum, and maximum number of nuclei per cell type and sample was computed. The number of cell types 1313 represented per sample was also determined. Sheet 4, Number of removed nuclei per level 1 annotation 1314 category per sample. Sheet 5, Number of removed nuclei per MTG cell type per sample. Sheet 6, Subtraction 1315 matrix comparing cosine similarity scores (i.e. similarity score for each cell subtype compared to the MTG cell 1316 Atlas as in Fig. 1d) for paediatric dataset to the adult dataset. Values are the paediatric scores minus the adult 1317 scores. Sheet 7, p-values, tests performed for each cell type and padj values (Benjamini-Hochberg method) when 1318 comparing the proportion of nuclei between male and female samples. Sheet 8-10, p-values, tests performed 1319 for each cell type and padj values (Benjamini-Hochberg method) when comparing the proportion of nuclei (sheet 1320 8), number of genes (sheet 9; see Extended Data Fig. 3b) and number of UMIs (sheet 10; see Extended Data Fig. 1321 3c) for each cell type between paediatric and adult samples shown.

1322

Extended Data Table 4: Summary of average quality control metrics for Visium datasets. Several measures for quality control were evaluated on a per sample basis including the sequencing saturation, the percentage of read mapped to the transcriptome, the number of spots under the tissue, the average number of nuclei per spot determined by Vistoseg analysis, the mean reads detected per spot, the median genes detected per spot, the total number of genes detected, the median UMI Counts per Spot and the total number of nuclei.

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1329 Extended Data Table 5: NS-Forest minimal marker analysis. Sheet 1, Statistical tests evaluating the expression of Aevermann et al. (2021)<sup>29</sup> minimal markers (see their Supplementary Tables 1-2) in our datasets. Sheets 2-3, 1330 1331 Metadata for each feature identified by NS-Forest marker in the down-sampled paediatric (sheet 2) and down-1332 sampled adult (sheet 3) datasets describing the cell type, the F-beta score for each marker gene, overlap with 1333 Aevermann et al. (2021) and Hodge et al. (2019), uniqueness to the age group of interest, coding status, and 1334 uniqueness to the associated cell type as shown in Fig. 3. As input to NS-Forest, all datasets (six paediatric and 1335 six adult) were randomly down-sampled such that the total number of nuclei per sample was equal to the sample 1336 with the fewest number of nuclei.

1337

Extended Data Table 6: gProfiler analysis of NS-forest markers. Sheet1-3, Significantly enriched GO terms associated with the paediatric (sheet1), adult (sheet2) and paediatric plus adult minimal marker genes identified by NS-forest. Sheet3-5, Significantly enriched GO terms associated with shared (i.e associated with both adult and paediatric samples) or paediatric-specific minimal marker genes with a binary expression score (> 0.7) for Oligo L1-6 OPALIN. Terms for which "highlighted" is true are driver terms.

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Extended Data Table 7: Summary of metadata for NS-Forest markers with a binary expression score (> 0.7) per cell type across the paediatric and adult datasets. The number of shared markers, the number of markers

1346 unique to paediatric samples, and the number of markers unique to adult samples is shown for each cell type.

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Extended Data Table 8: Overlap NS-Forest markers with a binary expression score (> 0.7) per cell type between the paediatric and adult datasets. NS-Forest markers with a binary expression score (> 0.7) per cell type were extracted for the down-sampled paediatric and down-sampled adult datasets. Each sheet represents 1 of 75 cortical cell types and the NS-Forest features which were shared (intersect) between the paediatric and adult datasets, unique to paediatric datasets, or unique to adult datasets are shown.

down-sampled paediatric datasets, and down-sampled adult datasets.

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1356 Extended Data Table 9: DESeq2 output of all genes tested for differential expression between paediatric and 1357 adult brains per cell type. Sheet 1-75, Differential expression analysis was performed using DESeq2's Wald Test 1358 for each cell type separately. Genes were filtered prior to testing to only include those expressed in > 10% of 1359 nuclei for that cell type across all paediatric and adult datasets. The associated log2FoldChanges, p-adjusted 1360 values (padj, Benjamini-Hochberg method), and description of each feature are shown. Positive log<sub>2</sub>FoldChanges 1361 represent genes upregulated in paediatrics versus adults. See DESeq2 documentation for explanation of NA 1362 values (https://bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html#why-are-1363 some-p-values-set-to-na).

1364

1365 Extended Data Table 10: DESeq2 output of significant DEGs only between paediatric and adult brains in a 1366 subset of cell types. Sheet 1-21, Significant DEGs (padj < 0.05) for cell types shown in Fig. 5a. The associated 1367 log<sub>2</sub>FoldChanges, p-adjusted values (padj), description, percentage of paediatric nuclei expressing the gene, 1368 percentage of adult nuclei expressing the gene, average normalised expression across paediatric nuclei, and 1369 average normalised expression across adult nuclei are shown. The difference in the peaditaric and adult values 1370 for percentage of nuclei and average normalised expression is also shown. Positive log<sub>2</sub>FoldChanges represent 1371 genes upregulated in paediatric versus adults datasets. See DESeq2 documentation for explanation of NA values 1372 (https://bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html#why-are-some-p-1373 values-set-to-na). Sheet 22-24, EA (sheet 6), IQ (sheet7) and HAR (sheet8) associated DEGs and their associated 1374 cell types.

1375

1376 Extended Data Table 11: psupertime coefficients. The calculated psupertime coefficients for each gene for
 1377 indicated excitatory neuron subtypes that showed the highest number of DEGs. Genes with non-zero psupertime
 1378 coefficients represent genes that are relevant to the ordering of the cells in pseudotime.

1379

1380 Extended Data Table 12: GSEA terms associated with each cell type showing enriched or depleted pathways in 1381 paediatric versus adult samples. GSEA was performed using DESeq2's output gene lists for each cell type ranked 1382 according to the log2FoldChange\*-log2(padj) for each gene. All DESeq2-tested genes served as input into GSEA 1383 (genes were expressed in > 10% of nuclei for the cell type of interest). Matrix shows the corresponding positive 1384 (sheet 1) and negative (sheet 2) NES values for each GSEA term (y axis) and cell type (x axis) based on the analysis 1385 using the ranked list of genes for each cell type. Terms were filtered to only include significantly associated terms 1386 (p<0.01, q<0.1). Positive NES values indicate pathways that are enriched in paediatric versus adult samples; 1387 negative NES values indicate pathways that are depleted in paediatric versus adult samples. The total number of 1388 terms per cell type and the total number of cell types associated with a given term are shown.

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