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Spatial and temporal heterogeneity of mouse and human microglia at single-cell resolution

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Microglial heterogeneity by single-cell RNA-seq

ABSTRACT

Microglia play critical roles in neural development and homeostasis. They are also implicated in neurodegenerative and neuroinflammatory diseases of the central nervous system (CNS). However, little is known about the presence of spatially and temporally restricted subclasses of microglia during CNS development and disease. Here, we combined massively parallel single-cell analysis, single-molecule FISH, advanced immunohistochemistry and computational modelling to comprehensively characterize novel microglia subclasses in up to six different regions during development and disease. Single-cell analysis of mouse CNS tissues revealed specific time- and region-dependent microglia subtypes, which were transcriptionally distinct from perivascular macrophages, during homeostasis. Demyelinating and neurodegenerative diseases evoked context-dependent microglia subtypes with distinct molecular hallmarks and diverse cellular kinetics. Diverse microglia clusters were also identified in normal and diseased human brains. Our data provide new insights into the endogenous immune system of the CNS during development, health and disease.

- Key words: microglia, perivascular macrophages, single-cell analysis, immune system,
- 58 human, mouse

INTRODUCTION

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Tissue-resident myeloid cells in the central nervous system (CNS) represent a heterogeneous class of innate immune cells that are essential for the maintenance of tissue homeostasis (1). Parenchymal microglia and the CNS-associated macrophages (CAMs), including leptomeningeal (mM Φ), perivascular (pvM Φ) and choroid plexus macrophages $(cpM\Phi)$, are the organ-specific macrophages of the CNS with pivotal roles in health and disease (2-4). Despite of the similarities that microglia and CAMs share with various other tissue-resident macrophages, the parenchymal and non-parenchymal CNS macrophages have two distinctive properties, namely a restricted prenatal origin and a remarkable longevity (4, 5). It is now generally believed that microglia and CAMs are derived from early yolk sac erythromyeloid precursors in a c-myb- and chemokine receptor (CCR)2-independent fashion (6-8). These specific developmental pathways and anatomical niches make CNSendogenous macrophages distinct from other tissue macrophages, such as those in the aorta, skin, heart, liver, spleen and other organs (9-12). When compared to other hematopoietic cells, microglia and CAMs persist over a very long period of time with low but constant rates of self-renewal (13, 14) coupled to cell apoptosis (15). This longevity necessitates adaptivity of microglia towards environmental challenges (16, 17) and cell perturbations (18). Since microglia act as guardians of the CNS, continuously scavenging for dying cells, pathogens, and molecules through microbialassociated molecular pattern receptor-dependent and -independent mechanisms (1), these highly diverse and specialized functions may be executed by microglia subsets that already pre-exist in situ, or alternatively, by specific development of microglia subsets from a homogeneous pool of cells upon demand. To date, the spatiotemporal heterogeneity of microglia during development, homeostasis and disease has not been studied at the singlecell level. Previous approaches used to analyse microglial diversity have largely relied on

immunophenotyping by flow cytometry complemented with histological analysis of RNA and

proteins in situ (19, 20). More recently, comprehensive transcriptomic (21) and proteomic (22) profiling of bulk populations of large numbers of microglia helped to reveal microglial heterogeneity in the mouse brain. Indeed, different microglia states were identified during development (7, 12, 23-25), homeostasis (26) and disease (27). Although these approaches provided important insights, they have notable limitations. Earlier single-cell analyses of microglia, for instance via flow cytometry, in situ hybridization or immunohistochemistry, were limited to probing a few selected proteins or RNAs. Due to a bias toward candidate genes/proteins, these approaches allow neither analysis of comprehensive expression landscapes nor discovery of previously unrecognized molecules (28). In contrast, transcriptomic analysis of bulk preparations of microglial RNA may conceal the diversity of microglia across different brain regions by relying on ensemble averages (21, 29, 30). During the last few years, the revolution in single-cell genomics has enabled an unbiased genome-wide quantification and multiplex spatial analysis of RNA in single microglia in situ as well as in vitro (31). However, recent single-cell RNA-sequencing (scRNA-seq) studies of microglia either only used pre-sorted myeloid cell populations (32), or whole brain approaches (33) without addressing the question of spatially and temporally restricted subtypes of microglia in several regions of the CNS. Importantly, single-microglia profiling data from humans is not yet available at all, although this knowledge may greatly improve our understanding of the pathogenesis of neuropsychiatric diseases. By combining massively parallel scRNA-seq with single-molecule FISH (smFISH), advanced triple immunohistochemistry, high-resolution microscopy, and computational modelling, we were able to comprehensively characterize microglial diversity in different regions of the mouse and human brain during development and health. We identify molecules that characterize microglial populations involved in neuroinflammatory and neurodegenerative conditions in mice and humans, and highlight context- and time-dependent microglia subsets and their distinct signals. The data provide new potential therapeutic targets and a valuable resource for the study of disease mechanisms in the CNS.

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RESULTS

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122 Distinct cell-specific signatures of individual microglia and pvMΦ in the juvenile 123 mouse brain 124 We first used an unbiased, surface marker-free approach to study the complexity of the CNS 125 of prepubescent juvenile mice and to determine how transcriptionally dissimilar microglia and 126 $pvM\Phi$ are as compared to the other CNS cells. For this purpose, we prepared a CNS cell 127 suspension devoid of meninges and choroid plexus (Fig. 1a). We then performed 128 quantitative scRNA-seg of 3,047 CNS cells as described before (34). Individual RNA 129 molecules were counted using molecular identifiers (UMIs) as performed recently (8, 34), 130 which greatly reduces PCR amplification bias. 131 Dimensionality reduction using t-distributed stochastic neighbor embedding (t-SNE) revealed 132 that both microglia and pvM Φ were transcriptionally related, whereas neurons, 133 oligodendrocytes, astrocytes, endothelial cells and vascular smooth muscle cells (VSMC) 134 had a distinct RNA profile (Fig. 1b). In order to define the transcriptional differences that 135 allow for the distinction of cell types in the CNS, we generated a heat map from 2,996 single 136 sorted cells showing the 49 most variable genes (**Fig. 1c**). Microglia and pvM Φ shared some 137 markers like Aif1, Csf1r and Tyrobp (Extended Data Fig. 1a), but microglia were 138 distinguishable from pvM Φ and other CNS cells on the basis of their expression of Tafbr. 139 Gpr34, Hexb, Selplg, II1a (Fig. 1c). In contrast, pvMΦ expressed higher mRNA levels of 140 Folr2, Lyve1, F13a1, Cbr2, Mrc1, Pf4, Cd163, Ccl24 and Cd209f. Gene ontology (GO) 141 142 involvement of microglia in cell chemotaxis, inflammatory response and regulation of cell 143 adhesion, whereas pvMΦ were involved in inflammatory response, regulation of response to 144 external stimulus, endocytosis and cytokine production (Extended Data Figs. 1b, c). 145 Visualization of scRNA-seg data on t-SNE plots revealed that microglia and pvMΦ 146 populations are distinguishable based on their transcriptomic signature (Fig. 1d). Taken together, these data reveal that microglia and $pvM\Phi$ are transcriptionally distinct myeloid cell populations in the CNS.

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Comprehensive transcriptome analysis microglia during development, homeostasis and disease by single-cell RNA-sequencing

Recent whole transcriptome analysis of microglia development from yolk sac progenitors to adult microglia highlighted the degree to which cells change during this interval, showing dramatic differences in microglial gene expression between early postnatal periods and adulthood (7, 12, 25). However, it is still unknown whether microglia subclasses with distinct transcriptional profiles emerge during development. In order to study microglia heterogeneity on single-cell level during different homeostatic conditions, we collected single microglia cells from multiple anatomical regions of the embryonic (embryonic day E16.5), juvenile (3 weeks) and adult (16 weeks) mouse CNS (Fig. 2a). The areas were selected to match those previously found to exhibit transcriptional differences of microglial bulk RNA on Affymetrix analysis (21). To further compare expression patterns during homeostasis to those under pathological conditions, microglia were also isolated from neurodegenerative (facial nerve axotomy) and demyelinating (cuprizone paradigm) disease models (Fig. 2a). In order to increase the yield, microglia were FACS-sorted from four different CNS regions during embryogenesis and up to six different CNS regions for postnatal time points (Fig. 2a and Suppl. Fig. 1). Following quality control, data from a total of 3,826 single microglia were further analyzed using the RaceID algorithm (Herman JS, 2018) and finally depicted in t-SNE plots (Fig. 2b and Suppl. Fig. 2). Unsupervised clustering gave rise to 13 distinct clusters. resembling ten microglia clusters during development (C1-C10) and one cluster for degeneration (C11) and two clusters for demyelination and remyelination (C12 and C13) (Fig. 2c).

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175 Spatiotemporal specificity of the emergence of microglia subsets during development 176 To investigate microglia diversity during development, we first focused on microglia from non-diseased CNS regions. t-SNE plots visualized two main clouds that clearly segregate 178 embryonic and postnatal microglia (Fig. 3a). Unbiased clustering of the top differentially 179 regulated genes revealed the presence of ten major clusters of microglia (C1-10) with distinct 180 transcriptional profiles (Figs. 3b, c and Suppl. Fig. 3). Among them, the C1-6 clusters predominantly consisted of embryonic microglia, whereas the postnatal microglia constituted 182 C7-10 clusters (Figs. 3a, b). Notably, embryonic clusters (C1-6) were differently distributed 183 across the four embryonic CNS regions tested (Figs. 3d, e). For instance, the C2 cluster was 184 enriched in embryonic forebrain and midbrain, whereas the C6 microglia was predominantly observed in cerebellum and spinal cord (Figs. 3d, e). Likewise, the postnatal clusters 185 186 showed a spatiotemporally variable distribution. For example, , e.g. the C10 cluster was enriched in juvenile cortical and hippocampal microglia (86.0 % and 71.7 % of microglia in the cortex and in the hippocampus, respectively, compared to 25.7 % in the cerebellum, Figs. 3d, e). Furthermore, the minor C7 cluster was more prevalent in cerebellum and 190 corpus callosum during adulthood (for both regions 12.3 % and 8.9 % of microglia in the cerebellum and in the corpus callosum compared to 5.6 % in the cortex). The relative 192 proportion of clusters in the cerebellum didn't change between the juvenile and the adult 193 stages, which is in sharp contrast to what was observed in the cortex and hippocampus, 194 where the C10 microglia decreased at the expense of the C7 and C8 clusters in adulthood (Figs. 3d, e). Overall, adult microglia showed a more homogenous distribution of each 196 cluster across regions than juvenile microglia (Figs. 3d, e). Together, these data suggest that microglia exhibit different subtypes with distinct gene expressional profiles over the 198 course of development with strong variation between different CNS regions that might reflect 199 local maturation differences. 200 Among the top differentially regulated genes during development were the microglial homeostatic genes Tmem119, Selplg and Slc2a5, which were highly induced at postnatal 202 stages (Fig. 4a). In addition, expression of *Malat1*, a long non-coding RNA, increased during

203 development, with the highest expression levesl being observed in adult microglia (Extended 204 Data Figs. 2a, b). In the embryonic clusters, lysosome-related genes Ctsb (encoding 205 cathepsin B), Cstd (encoding cathepsin D), Lamp1 (lysosomal-associated membrane protein 206 1), were strongly induced in C1 and C2 microglia (Fig. 3b and Extended Data Fig. 2c), 207 suggesting enhanced lysosomal activity in these embryonic microglia. In contrast, expression 208 of Apoe, which encodes the myeloid cell activation marker apolipoprotein E (27), was 209 enriched in the C1, C4 and C5 clusters (Fig. 4b). C6 microglia were characterized by high 210 expression levels of *Tmsb4x* (encoding thymosin beta 4), *Eef1a1*, and *Rpl4* (**Fig. 4b and** 211 Extended Data Fig. 2d), We next confirmed the existence of APOE⁺Iba1⁺ microglia and 212 CTSB⁺lba1⁺ microglia in the embryonic forebrain and cerebellum on a protein level by triple 213 immunofluorescence staining (Fig. 4c). These distinct embryonic microglia subpopulations 214 disappeared in the juvenile and adult brains (Figs. 4c, d). On the other hand, postnatal C9 215 and C10 microglia clusters were characterized by high expression of Cst3 (encoding cystatin 216 C, a cysteine protein proteinase inhibition family involved in neurodegenerative diseases of 217 the CNS (36)), and Sparc (encoding secreted protein acidic and rich in cysteine; also known 218 as osteonectin) (Fig. 4e). Immunolabeling for CST3 and SPARC confirmed the presence of 219 CST3*SPARC*Iba1* microglia in the postnatal brains, whereas this population was virtually 220 absent in embryonic forebrains (Figs. 4f,g). Interestingly, expression of CST3 was also 221 detectable in a subpopulation of Aldh111⁺ astrocytes in the adult cerebral cortex (Extended 222 Data Fig. 3). In contrast to the juvenile cerebral cortex, where almost all microglia expressed 223 CST3 and SPARC (Figs. 4f, g), the abundance of this microglia subpopulation slightly 224 diminished in the adult cortex, as CST3⁻SPARC⁺Iba1⁺ microglia emerged (Fig. 4g). In 225 contrast, the proportion of CST3⁺SPARC⁺Iba1⁺ microglia did not change between the juvenile and adult cerebellum (Fig. 4g), although the overall percentage of SPARC-226 227 expressing microglia was lower in the cerebellum than in cortex, and CST3⁺SPARCIba1⁺ 228 microglia made up a significant fraction of juvenile and adult cerebellar microglia (Fig. 4g). 229 Taken together, our data identify novel markers of microglia subsets and demonstrate the

spatiotemporal and phenotypic diversity of microglia subsets during CNS development and homeostasis in the adult brain.

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Identification of microglia clusters unique to demyelination and different from neurodegeneration

To investigate the kinetics of homeostatic microglia clusters and the putative generation of disease-specific microglia populations during CNS pathology, we compared a model of toxic demyelination, the cuprizone model, with a paradigm of neurodegeneration, the unilateral facial nerve axotomy (FNX) lesion (Fig. 2a). The blood-brain barrier remains intact in both models, and a loss of oligodendrocytes in the corpus callosum or a remote neurodegeneration within the facial nucleus, respectively, lead to local microglial activation without recruitment of circulating monocytes (13, 37). The two models allow us to study microglial plasticity following withdrawal of cuprizone or axonal regeneration, respectively (Fig. 2a). On the t-SNE plot, microglia distributed uniformly in a major population, but cells that clustered separately were found 3 days after FNX, and in the 5- and 10-week groups of cuprizone treatment (Fig. 5a-c). The C11-13 microglia predominantly constituted the disease-associated separate clouds on the *t*-SNE map (**Fig. 5c**). Notably, the C11 microglia cluster was specific to neurodegeneration, whereas demyelination induced the diseasespecific clusters (C12 and C13) (Figs. 5d-f). In the FNX model, the 3-day time point revealed a distinct microglia cluster (C11) characterized by stron expression of Ctsc (encoding cathepsin C) (Fig. 5e), whereas microglia from the 14-day time point clustered with the homeostatic microglia population. In contrast, toxic demyelination induced long-lasting transcriptional changes that only slightly recovered at the 10-week time point (Fig. 5f). In sum, our data suggest that homeostatic microglia are able to quickly change their phenotype and gain a discrete context- and time-dependent signature.

When analyzing the data for disease-specific signatures in microglia, a strong upregulation of *Apoe* was noted at all time points after cuprizone treatment (**Fig. 5g**). *Cst7* was more

prevalent in the demyelination-associated microglia (C12) (Fig. 5g), and Cybb was more strongly induced in the remyelination-associated C13 cluster (Fig. 5h and Extended Data Fig. 4g). In addition to Apoe, genes for AxI, Igf, Lyz2, Itgax (encoding CD11c), Gpnmb and Apoc1 were induced during de- and remyelination (Extended Data Fig. 4a), whereas Fam20c, Cst7, Ccl6, Fn1, Ank, Psat1 and Spp1 were enriched to variable degrees in the demyelination-associated C12 microglia (Figs. 5g,h and Extended Data Figs. 4b, c, e). In contrast, the remyelination-associated C13 microglia was characterized by high expression levels of the MHC class II genes *Cd74*, *H2-A2* and *H2-Ab1* (Fig. 5h and Extended Data Fig. 4i). On the other hand, the microglial core marker *Tmem119* was down-regulated following cuprizone treatment (Fig. 5h). Single-molecular fluorescence in situ hybridization (smFISH) validated the disease-associated expression of Fn1, Spp1, Cybb transcripts in Cx3cr1expressing microglia (Extended Data Figs. 4d, f, h). Furthermore, demyelination-associated microglia subtype (SPP1*CD74*Iba1* and TMEM119*CD74*Iba1*) was confirmed on the protein level by triple immunofluorescence staining (Figs. 5i.i). Likewise, remyelinationassociated microglia subtype (SPP1 CD74 ba1 and TMEM119 CD74 ba1) was confirmed by triple immunofluorescence staining (Figs. 5i,i). Overall, our results suggest the emergence of unique microglia subpopulations characterized by distinct signatures under defined disease conditions.

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Microglial diversity in the human brain

In order to extend our studies of microglial heterogeneity from mice to humans, we next analyzed 1,180 cortical microglia isolated from surgically resected human brain tissue without histological evidence of CNS pathology (referred to as "healthy") from five adult individuals with aged 23 to 54 years (**Suppl. Table. 1**). Unbiased hierarchical clustering of individual human microglia revealed four major clusters, hereafter referred to as healthy human clusters (HHu-C) (**Figs. 6a-c and Extended Data Fig. 5**). Detailed analysis of differentially regulated genes across the human microglia clusters revealed similarities with the gene expression profiles of murine homeostatic microglia. For example, *CST3* (enriched

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in mouse clusters C9 and C10) was more highly expressed in HHu-C1 and HHu-C2 than in HHu-C3 and HHu-C4 (Figs. 6b, d). In contrast, the human microglia cluster HHu-C4 showed comparatively high expression of the chemokine genes CCL4 and CCL2, and the zinc finger transcription factors EGR2 and EGR3 (Figs. 6b, d). Interestingly, CCL4 mRNA was rarely expressed in murine microglia even after cuprizone treatment (Extended Data Fig. 4j). Notably, P2RY13 mRNA was highly expressed by human microglia HHu-C1 and HHu-C2 clusters (Fig. 6b, Extended Data Fig. 5c), whereas the gene was not differentially expressed by murine adult microglia at single-cell level. In sum, our analysis identified homeostatic human microglia states with distinct gene expression patterns that partially overlap with adult mouse microglia. Activated microglia have been implicated in disease progression of multiple sclerosis (MS), a debilitating neurological disorder associated with demyelination (38). To examine the presence of disease-specific microglia subpopulations during this pathology in humans, 422 CD45⁺ cells isolated from the brains of five patients with histologically confirmed early active MS (Extended Data Fig. 6) were subjected to scRNA-seg and subsequently analysed together with healthy human microglia (Figs. 6e-i). Unsupervised clustering grouped cells into ten transcriptionally different clusters, which we termed human clusters (Hu-C)1-10 (Figs. 6f-h). Among them, the transcriptome of the Hu-C1 population showed a strong lymphocyte signature (TRAC, TRBC2, CD52, and IL32) (Fig. 6i), and the Hu-C9 and Hu-C10 populations were characterized by a clear monocytic profile (PLAC8, S100A9, CLEC12A, and CCR2) (Fig. 6i); these clusters were therefore excluded from further analysis. The remaining seven myeloid clusters, Hu-C2-8, expressed microglial core genes such as TMEM119, P2RY12, CX3CR1, SLC2A5 and P2RY13 to variable degrees (Fig. 6g). The Hu-C5-7 microglia clusters, which consisted entirely of microglia from healthy brains, showed highest expression levels of the microglial core genes and were therefore considered to represent the homeostatic microglia states (Figs. 6h, j and Extended Data Fig. 7a). Interestingly, the Hu-C4 subset that was shared by microglia from the healthy and diseased human brains revealed reduced expression levels of the core signature genes, but elevated

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levels of CCL2, CCL4, EGR2 and other chemokine/cytokine genes, suggesting a preactivated state of these microglial cells (Figs. 6h, j and Extended Data Fig. 7d). Unbiased clustering further identified two MS-enriched microglia clusters (Hu-C3 and Hu-C8) and one MS-associated microglia cluster (Hu-C2) that were clearly separated from the homeostatic clouds on t-SNE plots (Figs. 6e, f, h, j). The microglia clusters Hu-C2, Hu-C3 and Hu-C8 showed increased expression of APOE and MAFB (Extended Data Fig. 7f), whereas the core microglial genes were down-regulated or absent (Fig. 6i). Immunofluorescence staining of tissue from MS patients confirmed the strong reduction of TMEM119 expression on microglia in demyelinating lesions (Fig. 6k). The MS-associated Hu-C2 microglia was characterized by high expression levels of CTSD, APOC1, GPNMB, ANXA2, FAM20C and LGALS1 genes (Fig. 6j and Extended Data Fig. 7b, f). The Hu-C3 microglia showed increased gene expression of MHC class II-related molecules, such as CD74, HLA-DRA, HLA-DRB1 and HLA-DPB1 (Fig. 6j and Extended Data Fig. 7c). This suggests an immunoregulatory role, reminiscent of the remyelination-associated microglia subtype (C13) in mice (Fig. 5). Finally, the Hu-C8 microglia showed strong expression of SPP1, PADI2 and LPL genes, similar to the demyelination-associated microglia subtype (C12) in mice (Figs. 6g, j and Extended Data Fig. 7e). Of note, pairwise correlation analysis of mouse and human microglia orthologs confirmed that human MS-associated/-enriched microglia clusters (Hu-C2, Hu-C3 and Hu-C8), but not the pre-activated Hu-C4 cluster, are transcriptionally correlated to mouse demyelination-associated (C12) and remyelination-associated (C13) microglia observed after cuprizone treatment (Extended Data Fig. 8) To validate our scRNA-seq results for human microglia from MS patients, we performed immunohistochemical staining of MS brain sections. First, we stained for MRP14, which is known to label infiltrating monocytes but not microglia in early active lesions (39). Human brain sections without CNS pathology were virtually devoid of MRP14⁺lba1⁺ cells, whereas 12 % of all lba1⁺ cells in the MS sections were infiltrating monocytes (healthy: 0.2 ± 0.2 %, MS: 11.6 ± 2.4 %, Fig. 6I). However, this indicates that the vast majority of Iba1⁺ myeloid cells present in these sections were resident human MRP14 ba1 microglia. Next, we

Microglial heterogeneity by single-cell RNA-seq

per	formed	triple	immunofluore	escence	staining	and	identified	CTSD [†] I	MRP14 ⁻ lb	oa1⁺,
SPI	P1 [†] MRP	14⁻lba1⁺	and CD74 [†] M	RP14 ⁻ lba	a1 [†] microg	lia subs	sets as par	t of the H	lu-C2, Hi	u-C8
and	Hu-C3	clusters	s in brain sed	tions fro	<mark>m MS pa</mark>	tients (Figs. <mark>6m</mark> ,	n). In co	ontrast to	the
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DISCUSSION

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Our study provides a high-resolution view of the transcriptional landscape of microglia subtypes across multiple regions of the adult murine CNS. Furthermore, our data reveal a transcriptional continuum between microglia states, with few pre-existing clusters under homeostatic conditions after birth. Initial cell-specific states were rather uniform throughout the CNS during adulthood. In contrast, microglia subtype specification after birth emerged in a region- and disease stage-specific manner with high plasticity (Extended Data Fig. 9). Each brain region appears to be subject to changes in its immunological status, as revealed by regionally distinct states of mature microglia. In fact, classical monogenetic microgliopathies, such as hereditary diffuse leukoencephalopathy with spheroids (HDLS), Nasu-Hakola disease and others, are characterized by variable regional pathologies, suggesting diversity and differential spatial vulnerability of microglia (4, 36). Our data also reveal considerable dynamics of microglia subsets during development. It has been suggested that microglia ontogeny follows a defined stepwise transcriptional program to achieve the full homeostatic signature after birth (7, 25, 40). Indeed, we observed that microglia core genes like *Tmem119*, *Selplg* and *Slc2a5* are abundantly expressed only during adulthood. Since their expression levels increase after birth, it is tempting to speculate that microglia from juveline might not yet have fully matured. Interestingly, the embryonic microglia clusters were characterized by high expression of ApoE (C1, C4 and C5) and Ctsb (encoding cathepsin B) (C1 and C2), suggesting increased microglial activation and phagocytic-lysosomal activity. The developmental upregulation of cathepsin B, a protease activates matrix metalloproteinases (MMPs) and is thereby essential for the proteolysis of extracellular matrix components, might facilitate microglial movement in the growing brain, which depends on MMP8 and MMP9 in vivo (7). The widespread presence of cathepsin B⁺ microglia across different regions of the developing forebrain and cerebellum suggests a general function of these proteins in embryonic microglia. Furthermore, differential enrichment of microglial clusters across CNS regions during development might reflect distinct maturation stages of these regions.

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Previous studies on regional variations in microglial density (41), surface expression of a small panel of immune molecules (20), dependency on interleukin-34 (42), and microarray analysis of microglial bulk RNA (21) suggested diversity of microglia. However, earlier studies based on analysis of bulk cell populations isolated using a small set of surface markers were limited in their resolution of the heterogeneity and complexity of CNS immune cells. scRNA-seq enables unbiased characterization of small cell populations, and was used here to generate a high-resolution picture of microglia heterogeneity in the mouse and human brain. Single-cell analysis can also help to identify novel markers, pathways and regulatory factors that are critical during CNS development, homeostasis and disease. For example, in a recent study combining fate mapping and scRNA-seg, we showed that CAMs. like pvM Φ , are ontogenetically closely related to microglia (8). Despite their ontogenetic resemblance, this study revealed that microglia and pvMΦ are transcriptionally distinct myeloid cell populations in the CNS. Whether microglia and CAMs also originate from distinct progenitor populations in the yolk sac needs to be elucidated in the future. A recent study proposed regional differences in deep brain murine microglia, such as those within the basal ganglia (43). Microglia from the nucleus accumbens, ventral tegmental area and other regions were found to differ in morphology, density and membrane properties. Whether the observed differences in the membrane properties of microglia subsets within basal ganglia are functionally relevant remains unclear. Notably, morphological differences of microglia were accompanied by variations of cell density, with highest numbers of microglia in the midbrain and basal ganglia, as has been described previously (41). These regionspecific features might be due to the specific local microenviroment. In our study, we used scRNA-seq to investigate different CNS regions (excluding the basal ganglia) in the adult mouse brain that are known to exhibit microglia with diverse morphological features; however we did not observe obvious changes at the transcriptional level. The only exception was an enrichment of the C7 and C8 clusters in the cerebellum. Notably, the expression of Sparc, one of the representative genes that can segregate postnatal microglia states, was lower in cerebellar microglia compared to their cortical counterparts. These findings are in line with

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Affymetrix analyses that suggested heterogeneity of murine cerebellar microglia (21). Moreover, recent single-nuclei sequencing of striatal and cerebellar microglia from adult mice revealed epigenetic regulation of microglia clearance activity, with highest clearance activities in cerebellar microglia (Ayata 2018). Previous single-cell analyses identified neurodegeneration-associated microglia subsets in mice (27, 32, 33), but develination- and remyelination-associated microglia subsets were never examined at the single-cell level before. Here, we provide evidence for highly specialized and distinct demyelination- and remyelination-associated microglia subtypes in mice. Notably, we detected transcriptionally similar microglia subclasses in brain tissue from human MS patients, suggesting conserved responses to CNS demyelination. Since microglia down-regulate the expression of core genes during inflammatory conditions, we took advantage of the recently established microglia markers, TMEM119 and P2Y12R, that allow for the distinction of human microglia from infiltrating monocytes (44, 45). Our findings suggest that the highly specialized MS-associated human microglia subpopulations are characterized by enriched expression of SPP1, CTSD and the MHC class II-related molecule *CD74*, providing potential novel targets for MS therapy. Taken together, our study provides the first in vivo comparison of microglia heterogeneity at a single-cell resolution in the mouse and human CNS. Although we detected transcriptionally distinguishable microglia subpopulations, these did not appear as distinct clusters but rather as a transcriptional continuum of the local microglia population. This might represent the transcriptional basis for the ability of microglia to swiftly adapt to environmental changes. Our data further indicate that microglial responses to pathology are not uniform, but are shaped by the underlying pathology. In fact, we found disease-associated microglia subtypes in mice and humans that differed between neurodegenerative conditions (such as FNX) and toxic demyelination (like cuprizone). The appearance of context-dependent microglia subtypes with their own specific transcriptional profiles has potential therapeutic implications. Moreover, by establishing the transcriptional profile of heterogenous microglia populations in

Microglial heterogeneity by single-cell RNA-seq

- healthy and diseased rodents and humans, our study may provide new insights into the
- pathogenesis of CNS diseases.

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AUTHOR CONTRIBUTIONS

TM, RS, OS, CB, LA, CS, SN, PK, GvL, VAC, PCR, AM, US and RG conducted experiments and analyzed the data. MP, CS and JP analyzed the data, contributed to the in vivo studies and provided mice or reagents. TM and MP supervised the project and wrote the manuscript.

MATERIAL AND METHODS

Mice: CD1 mice were used. All animal experiments were approved by local administration and were performed in accordance to the respective national, federal and institutional regulations. Detailed mouse information is provided in Suppl. Table 1.

Analysis of single cell gene expression in diverse CNS cell types: A CNS cell suspension was obtained from a thorough preparation of eight different regions of the juvenile brain (mixed gender) in which meninges and choroid plexus were removed before. Cells were then subjected to single-cell RNA-seq using the C1 AutoPrep instrument (Fluidigm) and STRT/C1 protocol, as previously described (Zeisel et al., 2015; Goldmann et al., 2016). Each single cell was imaged and manually curated, and only single healthy-looking cells without debris were used for the analyses. Data analysis was performed as previously described (doi: DOI: 10.1126/science.aaa1934) using the BackSPIN algorithm.

Single-cell RNA-seq for mouse microglia: Microglia were FACS-sorted from up to six different CNS regions of healthy and diseased brains (see gating strategy shown in supplementary Fig. 1) into a 384-well plate containing a lysis buffer, and were analysed using Smart-seq2 method. Expression profiles were obtained as absolute cDNA molecule counts using the STAR aligner (doi: 10.1093/bioinformatics/bts635) to align raw sequences in conjunction with feature counts as part of the subread package (doi: 10.1093/nar/gkt214) to obtain gene counts. Further analysis and data normalization was performed using the RaceID package (Herman JS 2018). Clusters with more than ten individual cells were retained for further analysis and normalized to "transcripts per million" to compensate for differences in total transcriptome size between cell types. Heatmaps were generated using the online software (https://academic.oup.com/bioinformatics/article/32/18/2847/1743594).

Analysis of microglia from human brains: Human microglia were isolated from histologically healthy brain tissue removed during brain surgery for the treatment of epilepsy in five individuals (these tissues are not part of the epileptic region but are routinely removed to surgically access the epileptic lesion). Histopathological changes were excluded by an experienced neuropathologist, and only histologically healthy specimens were included in this study. Microglia were FACS-sorted into a 384-well plate containing lysis buffer. Single-cell RNA sequencing was conducted using the Cel-Seq2 protocol and processed as previously described (46). Libraries were sequenced on an Illumina HiSeq 3000 System in high output run mode at a depth of ~200,000 reads per cell. Paired end reads were aligned to the transcriptome using bwa with default parameters and all isoforms of the gene counted to a single gene locus (47). Reads that were not uniquely mapped were discarded. The left read contained the barcode information (6 bases corresponded to the cell specific barcode + 6 bases representing the unique molecular identifier (UMI)) and a polyT stretch and was

497 omitted from quantification. The corresponding right read was mapped to the ensemble of all 498 gene loci and used for quantification. Genes were counted based on the number of UMIs per 499 transcript from a given gene locus. The number of UMIs was converted to transcript counts 500 based on a binomial distribution (48). The aggregate of transcript counts with the same cell 501 barcode represented the transcriptome of an individual cell. Data analysis, normalization and 502 visualization was performed using the RaceID2 package (doi:10.1038/nature14966). Clusters 503 with more than 15 individual cells were retained for further analysis and transcript counts 504 were normalized by down sampling to 1500. Detailed human patient information is provided 505 in Suppl. Table.1. 506 Flow cytometry: After transcardial perfusion with PBS, brains were roughly minced and 507 homogenized with a potter in HBSS containing 15 mM HEPES buffer and 0.54 % glucose. 508 Whole-brain homogenate was separated by 70/37/30 % layered Percoll gradient 509 centrifugation at 800 g for 30 min at 4 °C (no brake). The CNS macrophages containing 510 interphase was then collected and washed once with PBS containing 2 % FCS and 10mM 511 EDTA before staining. Cells were stained with primary antibodies directed against CD11b 512 (M1/70, BioLegend), CD45 (30-F11, BD Biosciences), Ly6C (AL-21, BD Biosciences) and 513 Ly6G (1A8, BD Biosciences) for 20 min, and CD206 (C068C2, BioLegend) for 45 min at 4 514 °C. After washing, cells were sorted using a MoFlo Astrios (Beckman Coulter). Viable cells 515 were gated by staining with Fixable Viability Dye (eBioscience). Data were acquired with 516 FACSDiva software (Becton Dickinson). Post-acquisition analysis was performed using 517 FlowJo software, version X.0.7. Immunohistochemistry and cell quantifications: For juvenile and adult mice, after 518 519 transcardial perfusion with PBS, brains were fixed for 4 h in 4 % PFA, dehydrated in 30 % 520 sucrose and embedded in Tissue-Tek® O.C.T. compound (Sakura Finetek Germany GmbH). 521 For embryos, isolated brains were fixed for 4 h in 4 % PFA, dehydrated in 30 % sucrose and 522 embedded in Tissue-Tek® O.C.T. compound. Cryosections were obtained as described 523 previously (26). Sections were then blocked with PBS containing 5 % bovine serum albumin 524 and permeabilized with 0.1% Triton-X 100 in blocking solution. Primary antibodies were 525 added over night at a dilution of 1:500 for Iba-1 (ab178846, Abcam), 1:200 for APOE 526 (AB947, Millipore), 1:200 for CTSB (ab58802, Abcam) 1:200 for CST3 (AF1238, R&D 527 Systems), 1:200 for SPARC (IC942G, R&D Systems), 1:400 for NeuN (MAB377, Millipore), 528 1:1000 for APC (OB80, Millipore), 1:100 for Aldh1l1 (ab87117, Abcam), 1:500 for TMEM119 529 (ab209064, abcam), 1:500 for SPP1 (ab8448, abcam), 1:200 for CD74 (In1/CD74, 530 BioLedend) at 4°C. Secondary antibodies were purchased from Thermo Fisher Scientific added as follows: Alexa Flour® 488 1:500, Alexa Flour® 568 1:500 and Alexa Fluor® 647 531 532 1:500 for 2h at RT. Human tissue blocks were fixed in 4 % PFA overnight and embedded in 533 paraffin. Sections were then blocked with PBS containing 5 % bovine serum albumin and

permeabilized with 0.1% Triton-X 100 in blocking solution. Primary antibodies were treated over night at a dilution of 1:500 for Iba-1 (ab178846, Abcam; ab139590, Abcam; NB100-1028, Novus Biologicals), 1:200 for SPP1 (HPA027541, Sigma), 1:500 for CD74 (ab9514, abcam), 1:500 for CTSD (ab6313, abcam), 1:200 for MRP14 (T-1026, BMA Biomedicals; LS-B12844, LSBio). Secondary antibodies were purchased from Thermo Fisher Scientific added as follows: Alexa Flour® 405 1:500, Alexa Flour® 488 1:500, Alexa Flour® 568 1:500 and Alexa Fluor[®] 647 1:500 for 2 h at RT. Coverslips were mounted with/without ProLong[™] Diamond Antifade Mountant with DAPI (Thermo Fisher Scientific). Images were taken using a conventional fluorescence microscope (Olympus BX-61 with a color camera (Olympus DP71) or BZ-9000 (Keyence, Osaka, Japan) and the confocal pictures were taken with Fluoview FV 1000 (Olympus) using a 20 x 0.95 NA (XLUMPlanFL N, Olympus).

- Facial nerve axotomy and cuprizone model of demyelination and remyelination: Facial nerve was injured as described previously (13, 37). Briefly, mice were anesthetized by subcutaneous injection of a mixture of ketamine (50 mg/kg) and xylazine (7.5 mg/kg), and the right facial nerve was transected at the stylomastoid foramen, resulting in ipsilateral whisker paresis. Cuprizone treatment was used as a model of toxic, demyelination and remyelination (37, 49). For demyelination, mice were fed for 5 weeks with 0.45 % (wt/wt) cuprizone (Sigma, St. Louis, MO) in the ground breeder chow. For remyelination, the cuprizone diet was discontinued after 5 weeks and animals were maintained for further 5 weeks under normal diet to allow spontaneous remyelination. Untreated age-matched mice were used as control.
 - Single molecule fluorescent in situ hybridization (smFISH): Mice were perfused with PBS, followed by 4% paraformaldehyde (PFA). The brain tissues were harvested and immersion-fixed in 4% PFA for 3 h, and subsequently were put into 30% sucrose in 4% PFA at 4°C overnight, and embedded in OCT for sectioning, frozen on dry ice and stored at -80°C until used. 10-µm thick sections mounted on the glass plate were washed 3 times with PBS, and treated with pre-chilled methanol for 10 min at -20 °C. Then the slides were incubated for 10 min at 70°C in Tris-EDTA (pH 8.0), and the sections were washed with SSC 2X and incubated for 4 hr with hybridization buffer containing 250 nM fluorescent label probes (LGC Biosearch Technologies) at 38.5°C. After 4 times washing with 20% formamide wash buffer containing SSC 2X, the slides were mounted with Prolong Gold containing DAPI. Stack Images were taken using a Olympus BX-61 microscope.
- Gene ontology (GO) enrichment analysis: The defined differentially regulated genes were analyzed using the software (available from http://metascape.org/gp/#/main/step1).
- Pairwise correlation analysis: Comparison between human and mouse data was performed by selecting all genes found to differentially expressed by RaceID (adjusted)

Microglial heterogeneity by single-cell RNA-seq

570	p<0.01, log2FC > 1) in any of the identified clusters. For genes from the human dataset
571	mouse orthologs were identified from the NCBI HomoloGene database
572	(https://www.ncbi.nlm.nih.gov/homologene), using the annotationTools R package (Kuhn A
573	2008); the same was done to identify human orthologs for the murine genes. All human
574	genes with a ortholog in the mouse set as well as all murine genes with an ortholog in the
575	human set were kept. Canonical Cluster Analysis as implemented in the Seurat package
576	(Butler A 2018) was then performed on the 768 common genes identified in this manner.
577	Statistical analysis: Statistical significance was determined using one-way ANOVA with
578	post hoc Tukey Multiple Comparison test using GraphPad Prism 5.04 software.
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580	FIGURE LEGENDS
581	
582	Figure 1: Unbiased single-cell RNA-seq of CNS cells reveals specific profiles of murine
583	microglia and pvM Φ .
584	(a) Schematic diagram showing the isolation of single CNS cells from juvenile (3 weeks of
585	age) mice for unbiased sampling and single-cell RNA-seq (scRNA-seq).
586	(b) Cluster analysis using t-SNE of 3,047 individual cells measured by single-cell RNA
587	sequencing and bi-clustering. Each dot represents an individual cell. The populations of
588	microglia and perivascular macrophages (pvM Φ) are marked by a dotted line. Vsmc:
589	vascular smooth muscle cell.
590	(c) Heat map showing clustering of 2,996 single cells, featuring 49 most variable genes.
591	Selected marker genes enriched in each cell-type representing expression levels of
592	selected known and novel markers are shown on the right.
593	(d) t -SNE clustering plots of individual microglia and perivascular macrophages (pvM Φ)
594	showing distinct gene expression pattern between the two cell types in the juvenile CNS
595	(326 cells). Each dot represents a single cell. Microglia are depicted as circles, pvM Φ as
596	<mark>triangles.</mark>
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598	Figure 2: Comprehensive analysis of microglial diversity by single-cell RNA-
599	sequencing.
600	(a) Illustration depicting the workflow for the isolation of microglia from different CNS regions
601	of embryonic (embryonic day E16.5) and juvenile (3 weeks of age) and adult (16 weeks)
602	mice during homeostasis and during pathology, namely facial nerve axotomy (FNX)
603	cuprizone-mediated demyelination for scRNA-seq.
604	(b) t-SNE plot showing all analyzed microglia cells from different conditions tested in this
605	study. Each dot represents a single cell.
606	(c) t-SNE plot depicting 13 clusters for all different conditions. Colors represent each cluster
607	(C).
608	
609	Figure 3: Identification of spatiotemporal subclasses of microglia in the mouse.
610	(a) t-SNE plot of 2,966 individual microglia isolated at different time points of development.
611	Each dot represents a single cell. Colors correspond to the time points investigated.
612	(b) t-SNE plot depicting ten major and three minor microglia clusters at three different
613	developmental stages. Each dot represents a single cell. Colors represent each cluster
614	(C).

- (c) Heat map of top differentially regulated genes that were up- or down-regulated in each
 cluster, including genes such as *Malat1*, *Selplg*, *Tmem119*, *Sparc*, *Cst3*, *Ctsd*, *Lamp1*,
 Ctsb, *Apoe*, *Tmsb4x*, *Eef1a1* and *Rpl4*.
- (d) *t*-SNE plots depicting regional distribution of transcripts from 2966 individual microglia at different developmental time points. Each dot represents a single cell.
- 620 (e) Distribution of microglia clusters among different CNS regions during embryonic and 621 postnatal stages. Colors represent distinct clusters.

- Figure 4: Characteristics of microglial subsets during development.
- (a) t-SNE plots depicting the expression kinetics of the microglial core genes *Tmem119*,
 Selplg and Slc2a5 during ontogeny. Upper right cloud shows E16.5 microglia whereas
 the lower left cloud represents microglia from juvenile and adult mice. Color key indicates
 the expression levels.
- (b) t-SNE plots of embryo-enriched microglia transcripts for Ctsb, Apoe and Tmsb4x.
- 629 (c) Representative immunofluorescence images for apolipoprotein (Apo)E, cathepsin B 630 (CTSB) and ionized calcium-binding adapter (Iba)1 in the embryonic forebrain and 631 juvenile cortex. Dotted frame 1 indicates ApoE⁻CTSB-lba1⁺ embryonic microglia 632 (representing clusters C3, C6). Frame 2 illustrates ApoE⁻CTSB⁺lba1⁺ embryonic 633 microglia (C1, C2) whereas frame 3 shows ApoE⁺CTSB⁺lba1⁺ triple-positive embryonic 634 microglia (C1). Dotted frame 4 depicts ApoE⁺CTSB⁻Iba1⁺ embryonic microglia (C4, C5). 635 Frame 5 illustrates ApoE CTSB bal microglia (white arrowheads) found at the juvenile 636 stage. Yellow arrowheads indicate ApoE⁺Iba1⁻ cells in the juvenile brain. Representative 637 pictures out of four investigated mice are shown. Scale bars: 50 µm (overview), 30 µm 638 (insert).
- (d) Quantification of ApoE and CTSB immunoreactivities in Iba1⁺ microglia from different
 CNS regions during development. Bars represent means ± SEM from four animals (541-853 microglia per region).
- (e) *t*-SNE plots-based distribution of *Cst3* and *Sparc* transcripts in microglia. Color keys indicate the expression levels.
- (f) Representative immunofluorescence images for cystatin C (CST3), secreted protein
 acidic and rich in cysteine (SPARC) and Iba1 in the embryonic forebrain, juvenile and
 adult cortex. Frame 1 indicates CST3⁺SPARC⁺Iba1⁺ microglia (representing clusters C9,
 C10) whereas frame 2 highlights CST3⁻SPARC⁻Iba1⁺ microglia (C7) during adulthood.
 Representative pictures out of four investigated mice are shown. Scale bars: 50 μm
 (overview), 20 μm (insert).

650	(g) Quantification of CST3 and SPARC immunopositivity in microglia from different regions
651	of the CNS at distinct developmental time points. Bars represent means \pm SEM from four
652	animals (569 - 1961 microglia per region).

- Figure 5: Specific disease-associated microglia populations with distinct kinetics during demyelination and neurodegeneration.
- 656 (a) Projection of 1,564 single microglia isolated from different CNS regions during 657 homeostasis or FNX or cuprizone treatment as *t*-SNE plot.
- 658 (b) Heat map of top differentially regulated genes that were up- or down-regulated in each 659 cluster. Highest differentially expressed genes are highlighted.
- 660 (c) t-SNE plot exhibiting 13 clusters for the 1,564 individual microglia isolated from different 661 CNS regions during homeostasis or FNX or cuprizone treatment.
- 662 (d) Left: Kinetics of facial nucleus (FN) microglia subpopulation on a t-SNE map either 663 untreated (FN-normal) or after 3 days post FNX (FNX-d3) or 7 days post FNX (FNX-d14), 664 respectively. Right: Histogram displaying proportion of microglia clusters either untreated 665 (FN-normal) or after FNX-d3 or FNX-d14.
- 666 (e) Clustering of the Ctsc gene expression following FNX. Expression of Ctsc is found to be 667 upregulated in C11 at FNX-d3. The color key indicates the expression levels. Insert: 668 close-up of the C11.
- 669 (f) Persistent transition of corpus callosum (CC) microglia population on a t-SNE map before 670 (CC-normal) and after demyelination (CC-Demyelination) or remyelination (CC-671 Remyelination). Close-ups reveal distribution of clusters after demyelination and 672 remyelination. Right: Histogram showing long-lasting changes in microglia populations 673 following cuprizone treatment.
- 674 (g) Kinetics of Apoe, Cst7 and Cybb expression after cuprizone challenge displayed in t-SNE 675 plots. Color keys represent the respective expression levels.
- 676 (h) t-SNE plots for Tmem119, Spp1, Cd74 after cuprizone treatment. Tmem119 is 677 downregulated following treatment, whereas Spp1 is upregulated in C12, and Cd74 678 mRNA is increased in C13. Color keys indicate the expression levels.
- 679 (i) Left: representative immunofluorescence images for osteopontin (secreted 680 phosphoprotein 1, SPP1), CD74 and Iba1 in the normal and demyelinated corpus 681 callosum. Arrowheads indicate SPP1 CD74 Iba1 (white arrowheads), SPP1 CD74 Iba1 (white arrowheads), SPP1 CD74 Iba1 682 (red arrowheads), and SPP1 CD74 ba1 (blue) parenchymal microglia, respectively. 683 Representative pictures out of three or four investigated mice are shown. Right: 684 Quantification thereof. Bars represent means ± SEM of three to four animals (437 - 825 685 microglia per condition). Each symbol represents one animal. Scale bars: 30 µm.

(j) Left: typical immunofluorescence pictures for transmembrane protein (TMEM) 119, CD74
 and Iba1 in the normal and demyelinated corpus callosum. Arrowheads show
 TMEM119⁺CD74⁻Iba1⁺ (white arrowhead), TMEM119⁻CD74⁻Iba1⁺ (red) and TMEM119⁻
 CD74⁺Iba1⁺ (blue) parenchymal microglia. Representative pictures out of four
 investigated mice are shown. Right: Quantification thereof. Bars represent means ± SEM
 of four animals (808 - 1024 microglia per condition). Each symbol represents one animal.
 Scale bars: 30 μm.

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Figure 6: Presence of distinct subclasses of microglia in healthy human and MS brains.

- (a) *t*-SNE plot of 1,180 individual human microglia isolated from five individual non-pathological brains depicts four major clusters (HHu-C1-4). HHu-C: healthy human microglia cluster. Each dot represents a single cell. Colors correspond to each cluster.
- (b) Heat map of the top differentially regulated genes that were up- or down-regulated in each cluster, including genes such *CST3*, *P2RY13*, *CCL4*, *CCL2*, *EGR2* and *EGR3*.
- 700 (c) Bar graphs representing the relative abundance of microglia cells in the respective clusters from five individual non-pathological brains. Colors represent distinct clusters.
- 702 (d) *t*-SNE plots for *CST3*, *CCL4* and *EGR2 mRNA* expression. *CCL4* and *EGR2* are enriched in the Hu-C4.
- 704 (e) *t*-SNE plot of 422 individual human microglia isolated from five individual non-705 pathological brains and five patients with early active multiple sclerosis (MS). Each dot 706 represents a single cell. Colors correspond to each condition or patient.
- 707 (f) *t*-SNE plot depicting ten major clusters (Hu-C1-10) of microglia from healthy and diseased individuals. Each dot represents a single cell. Colors correspond to each cluster.
- 710 (g) Heat map of all differentially regulated genes that were up- or down-regulated in each cluster.
- 712 (h) Bar graphs representing the relative abundance of microglia in each cluster from healthy 713 and individual MS patients. Colors represent individual patients or conditions.
- 714 (i) *t*-SNE plots representing the core signature genes for lymphocytes, myeloid cells and monocytes in the brains from MS patients. Color keys reflect the expression levels.
- 716 (j) *t*-SNE plots of the microglia cluster representing the top five enriched genes for each cluster. Homeostatic microglial genes are enriched in the microglia clusters Hu-C5-7, whereas Hu-C4 is characterized by the expression of proinflammatory molecules *CCL4* and *CCL2*. Hu-C2, Hu-C3 and Hu-C8 are present in microglia from MS patients.
- (k) Immunofluorescence images for TMEM119 and Iba1 in healthy or MS patient brains.
 Arrowheads indicate TMEM119⁺Iba1⁺ cells (filled) in the healthy brains, and TMEM119⁻
 Iba1⁺ microglia (open) during MS. Scale bar: 50 μm.

(I) Representative immunofluorescence pictures for Iba1*MRP14- (indicating microglia) and lba1⁺MRP14⁺ cells (representing infiltrating early activated monocytes) in the normal and MS brain. Inserts show microglia (first row) and monocytes (second row) in the MS lesion. Right: Quantification thereof. Bars represent means ± SEM. Each symbol 727 represents one patient. Scale bars: 50 µm (overview), 4 µm (insert).

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- (m) Upper panel: representative immunofluorescence images for SPP1, CTSD, Iba1 and MRP14 indicating microglia subsets in normal and MS brains. Representative pictures out of four individuals were chosen. Dotted frames represent SPP1 CTSD MRP14 ba1 microglia (1) and SPP1⁺CTSD⁻MRP14⁻Iba1⁺ microglia (2). Scale bars: 50 µm (overviews), 20 µm (inserts). Lower panel: Quantification of microglia immunoreactivities in healthy or MS brains. Percentages indicate the relation of MRP14 ba-1 microglia subsets in individual brains. 153 - 163 microglia per patient were examined.
- (n) Upper: Immunofluorescence pictures for SPP1, CD74, Iba1 and MRP14 for the characterization of microglia cluster in healthy and MS brains. Colored dotted frames SPP1⁻CD74⁺MRP14⁻Iba1⁺ (white), SPP1⁺CD74⁻MRP14⁻Iba1⁺ (yellow) indicate parenchymal microglia, respectively. Representative pictures out of four individuals are shown. Scale bars: 50 µm (overviews), 20 µm (inserts). Lower left: Quantification of microglia immunopositivities in healthy or MS brains. Percentages indicate the relation of MRP14⁻lba-1⁺ microglia subsets in individual brains. 152 - 200 microglia per patient were investigated. Lower right: Distribution of SPP1 and CD74-reactive Iba1⁺ microglia subsets in the healthy mouse corpus callosum or during cuprizone-induced de-and remyelination as shown in Fig. 5i.

EXTENDED DATA

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- 752 Extended Data Figure 1: Molecular characterization of rodent microglia and 753 perivascular macrophages.
- 754 (a) Heat map showing clustering of 2,996 single cells, featuring ten selected genes enriched 755 both in microglia and perivascular macrophages (pvMΦ) obtained from juvenile (3 weeks 756 of age) mice.
- 757 (b) Gene ontology (GO)-term enrichment analysis for the 274 top genes enriched for 758 microglia.
- 759 (c) GO-term enrichment analysis for the top 317 genes enriched for pvM Φ .

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Extended Data Figure 2: Microglial subpopulations in mice with distinct gene expression during development.

- (a) Distribution of Malat1 gene expression in a t-SNE plot. Color keys represent the respective expression levels. Upper left cloud represents embryonic microglia population whereas lower right cloud combines both juvenile and adult microglia as shown in Fig. 2b.
- 766 (b) Left: smFISH for Malat1 and Cx3cr1 shows the kinetics of Malat1⁺ microglia during 767 development. Scale bar: 10 µm. Representative pictures out of two investigated adult 768 mice are shown. Yellow arrowhead and white indicate Malat1+Cx3cr1+ microglia and 769 Malat1⁺Cx3cr1⁻ non-microglia cells, respectively. Right: Frequency of Malat1⁺ microglia 770 in the forebrain or cortex during development. Bar represents mean ± SEM of 120 771 studied cells from three animals per time points.
- 772 (c) t-SNE plot of Ctsd and Lamp1 gene expression that were enriched in C1 and C2 clusters 773 as shown in Fig. 2c.
- 774 (d) t-SNE plot of Eef1a1 and Rpl4 gene expression that were enriched in C6 cluster as 775 shown in Fig. 2c.

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Extended Data Figure 3: Cst3 is enriched in adult microglia.

- 778 (a-c) Representative sections of the cortex from adult mice using immunofluoresence for 779 cystatin C (CST3, green), NeuN for neurons (red, a), adenomatous polyposis coli (APC) 780 for oligodendrocytes (red, b), respectively. The astrocyte marker Aldh1I1 (red, c) combined with CST3 was used on the hippocampal sections. Scale bars: 50 µm 782 (overviews) and 20 µm (magnifications). Representative pictures out of three investigated 783 mice are shown.
- 784 (d) Quantification of CST3 immunoreactivity in the brain of adult mouse. Bar represents 785 mean ± SEM of three animals (393 microglia, 1817 neurons, 298 oligodendrocytes, 461 786 astrocytes).

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Extended Data Figure 4: Molecular characterization of microglia subpopulations during de- and remyelination.

t-SNE plots showing expression of AxI, Igf1, Lyz2, Itgax, Gpnmb, Apoc1 (a), Fam20c Ccl6, Psat1, Ank (b), Fn1 (c), Spp1 (e), Cybb (q), H2-Aa and H2-Ab1 (i), Ccl4 (j) transcripts after cuprizone challenge. Genes shown in (a) were upregulated in both de-and remyelination, whereas genes depicted in (b, c, e) or (g, i) were increased in demyelination-associated cluster Adt-C5, C6, or in the remyelination- associated Adt-C7, C8, respectively. Color keys represent the respective expression levels. (d) Left and middle panels: single-molecule fluorescent in situ hybridization (smFISH) for Fn1 and Cx3cr1 reveals subpopulations of microglia after 5-week cuprizone treatment in the corpus callosum. "1" indicates Fn1⁺Cx3cr1⁺ microglia (yellow arrowheads: Fn1 mRNA). "2" indicates Fn1 Cx3cr1* microglia. Scale bar: 10 um (overviews) or 3 µm (inserts). Representative pictures out of three investigated mice are shown. Blank arrowheads in the picture of control mice indicate non-specific signals. Right panel: percentage of Fn1⁺ Cx3cr1⁺ microglia in the corpus callosum. Bar represents mean ± SEM of three animals (168 investigated cells). (e) Spp1 mRNA expression after cuprizone challenge. Expression of Spp1 is found to be upregulated in Adt-C5 and C6. The color key indicates the expression levels. (f) Left and middle panels: smFISH for Spp1 and Cx3cr1 reveals subpopulations of microglia after 5-week cuprizone treatment in the corpus callosum. "1" indicates Spp1*Cx3cr1* microglia (yellow arrowheads: Spp1 mRNA). "2" indicates Spp1* Cx3cr1⁺ microglia. Scale bars: 10 µm (overviews) and 3 µm (inserts). Representative pictures out of three investigated mice are shown. Right panel: percentage of Spp1+Cx3cr1+ microglia in the corpus callosum. Bar represents mean \pm SEM of three animals (165 investigated cells). (a) t-SNE plot depicting Spp1 expression after cuprizone challenge. The color key indicates the expression levels (h) Left and middle panels: smFISH for Cybb and Cx3cr1 reveals subpopulations of microglia after 5 weeks cuprizone treatment in the corpus callosum. "1" indicates Cybb⁺Cx3cr1⁺ microglia (yellow arrowheads: Cybb mRNA). "2" indicates Cybb⁻ Cx3cr1⁺ microglia. Scale bars: 10 µm (overviews) and 3 µm (inserts). Representative pictures out of three investigated mice are shown. Right panel: percentage of Cybb⁺Cx3cr1⁺ microglia in the corpus callosum. Bar represents mean ± SEM of three animals (165 investigated cells). The color key represents the expression levels.

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Extended Data Figure 5: Microglial subtypes in healthy human brains.

(a) t-SNE plot of 1180 human microglia showing the distribution of individual microglia from five patients. Each dot represents a single cell. Different colors indicate different patients,(b) Heat map showing the distribution of the healthy human clusters (HHu-C) in each

824 individual patient.

825	(c) t-SNE plot showing the expression of P2RY13. Each dot represents a single cell. Color
826	codes represent expression levels.
827	
828	Extended Data Figure 6: Detailed neuropathological characterization of human MS
829	lesions.
830	Histology of the MS brains (MS-1 patient until MS-5 patient) using hematoxylin and eosin
831	(H&E), luxol fast blue (LFB-PAS), 2',3'-cyclic-nucleotide 3'-phosphodiesterase (CNPase), and
832	myelin basic protein (MBP) for myelin, human leukocyte antigen – DR isotype (HLA-DR) and
833	CD68 for myeloid cells, CD3 for T cells, CD20 for B cells and Bielschowsky (Biel) for axons.
834	Scale bar: 50 µm. Lesions are typical early active MS lesions according to the standard
835	classification system (50).
836	
837	Extended Data Figure 7: Molecular profile of microglia subsets during MS.
838	(a-e) t-SNE plots of genes enriched in cluster Hu-C5-7 (a), Hu-C2 (b), Hu-C3 (c), Hu-C4
839	(d), Hu-C8 (e) are shown. Color codes represent expression levels.
840	(f) t-SNE plots depicting genes upregulated in the clusters Hu-C2, Hu-C3 and Hu-C8. Color
841	codes represent expression levels.
842	(g) t-SNE plots of genes that were upregulated in the disease-associated microglia subsets
843	in the mouse demyelination model, but not in the microglia in the MS patient brains. Color
844	codes represent expression levels.
845	
846	Extended Data Figure 8: Pairwise correlation analysis of scRNA-seq data from mouse
847	and human microglia.
848	(a) Canonical correlation analysis (Seurat alignment procedure) visualizing shared
849	correlation structures (i.e., canonical correlation vectors, CC) between mouse and human
850	data sets. Each dot represents single cell.
851	(b) CC Plot of cells assigned as mouse C7-C13 and human Hu-C1-C10. Mouse
852	demyelination-related microglia clusters (C12 and C13) are transcriptionally close to
853	human MS-associated microglia clusters (Hu-C2, Hu-C3 and Hu-C8). Each dot
854	represents single cell.
855	(c) Violin plots depicting a shared gene correlation structure that is conserved between
856	mouse and human clusters.
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859 860	Extended Data Figure 9: Graphical abstract of experimental findings.
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864	SUPPLEMENTAL FIGURES AND TABLE
865	
866	Supplementary Figure 1: FACS gating strategy for microglia isolation.
867	CNS cells were gated for G1 and G2 (singlets), followed by being gated for living cell (G3,
868	fixable viability dye), CD45 ^{int} CD11b ⁺ (G4), Ly6C ⁻ Ly6G ⁻ (G5), and CD206 ⁻ (G6).
869	
870	Supplementary Figure 2: related to figure 2-5
871	t-SNE plots depicting single microglia from the replicates from different CNS regions of
872	individual embryos, juvenile and adult mice and diseased mice.
873	
874	Supplementary Figure 3: related to figure 3
875	Heat map of all differentially regulated genes that were up- or down-regulated in each cluster
876	microglia during development.
877	
878	Supplementary Figure 4: related to figure 5
879	Heat map of all differentially regulated genes that were up- or down-regulated in each cluster.
880	
881	Supplementary Figure 5: related to figure 6
882	Heat map of top 68 differentially regulated genes that were up- or down-regulated in each
883	cluster of healthy human microglia.
884	
885	Supplementary Figure 6: related to figure 6
886	Heat map of top 148 differentially regulated genes that were up- or down-regulated in each
887	cluster of both healthy and MS-patient microglia.
888	
889	Supplementary Table 1:
890	Sheet 1: Information on mice and cells used in each scRNA-seq analysis including
891	genotype, strain, sex, condition, age, CNS region and cell numbers.
892	Sheet 2: Details on patients and cells used in each scRNAseq analysis including sex,
893	condition, age, CNS region and cell numbers.

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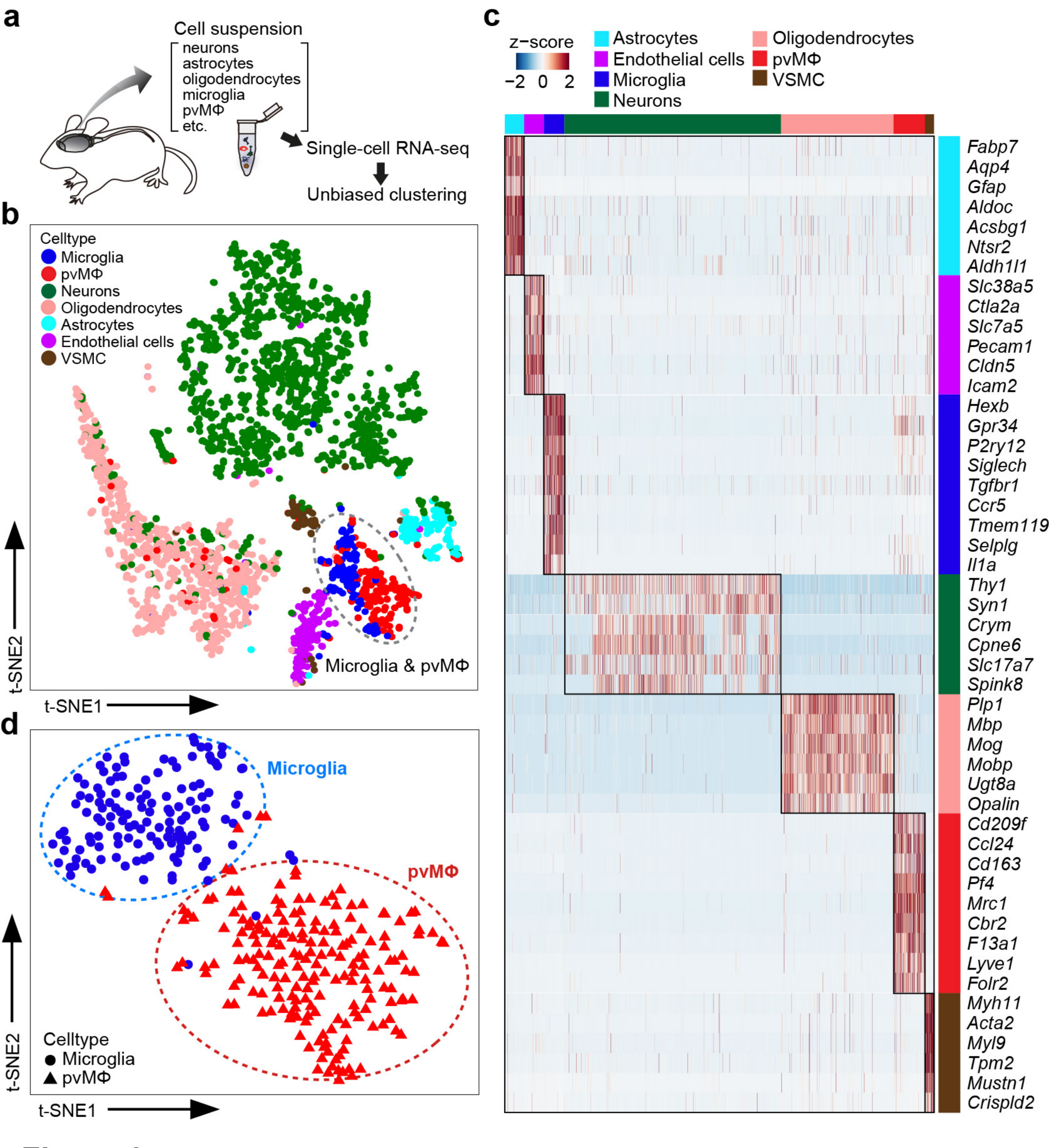


Figure 1

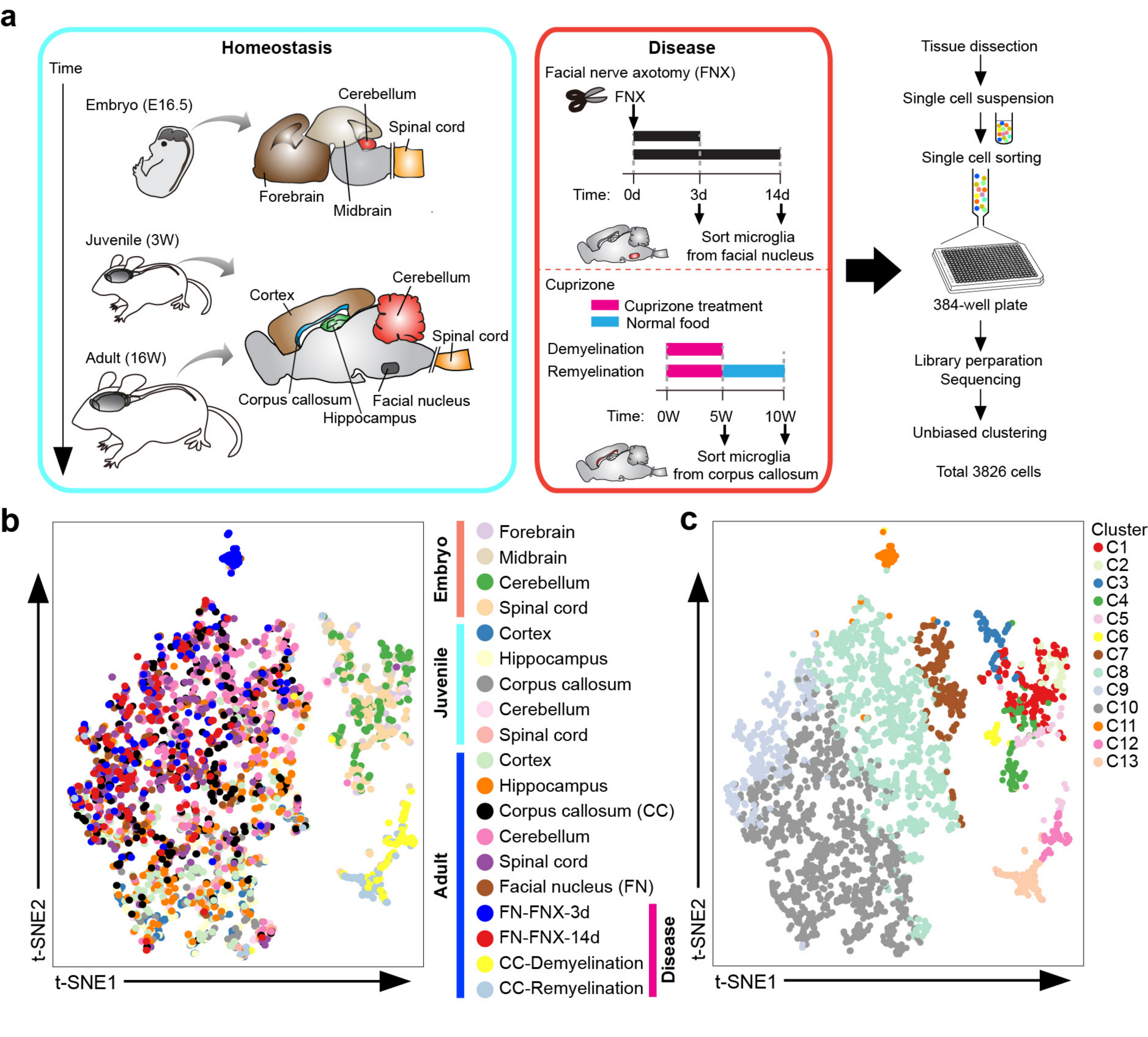


Figure 2

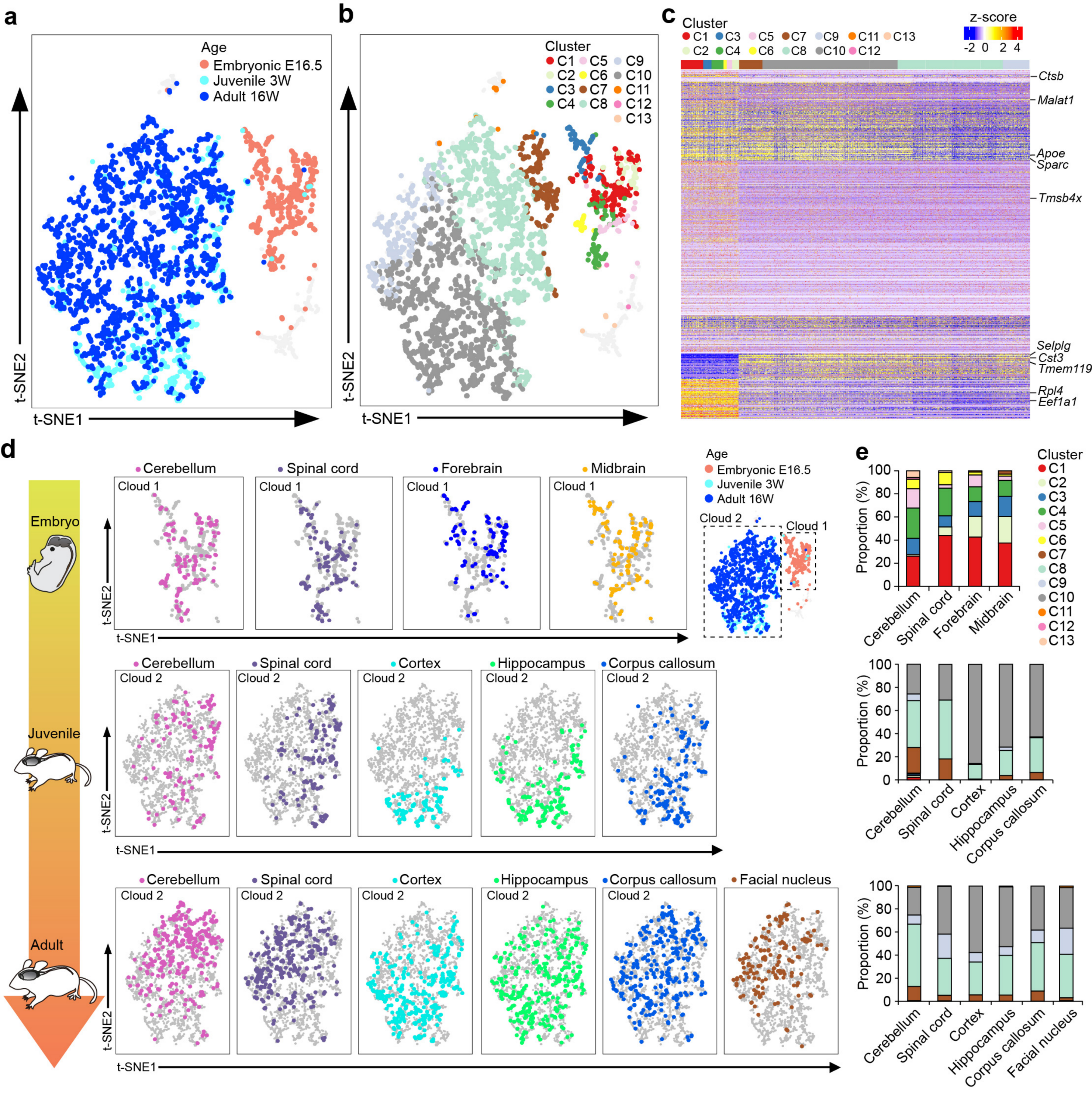


Figure 3

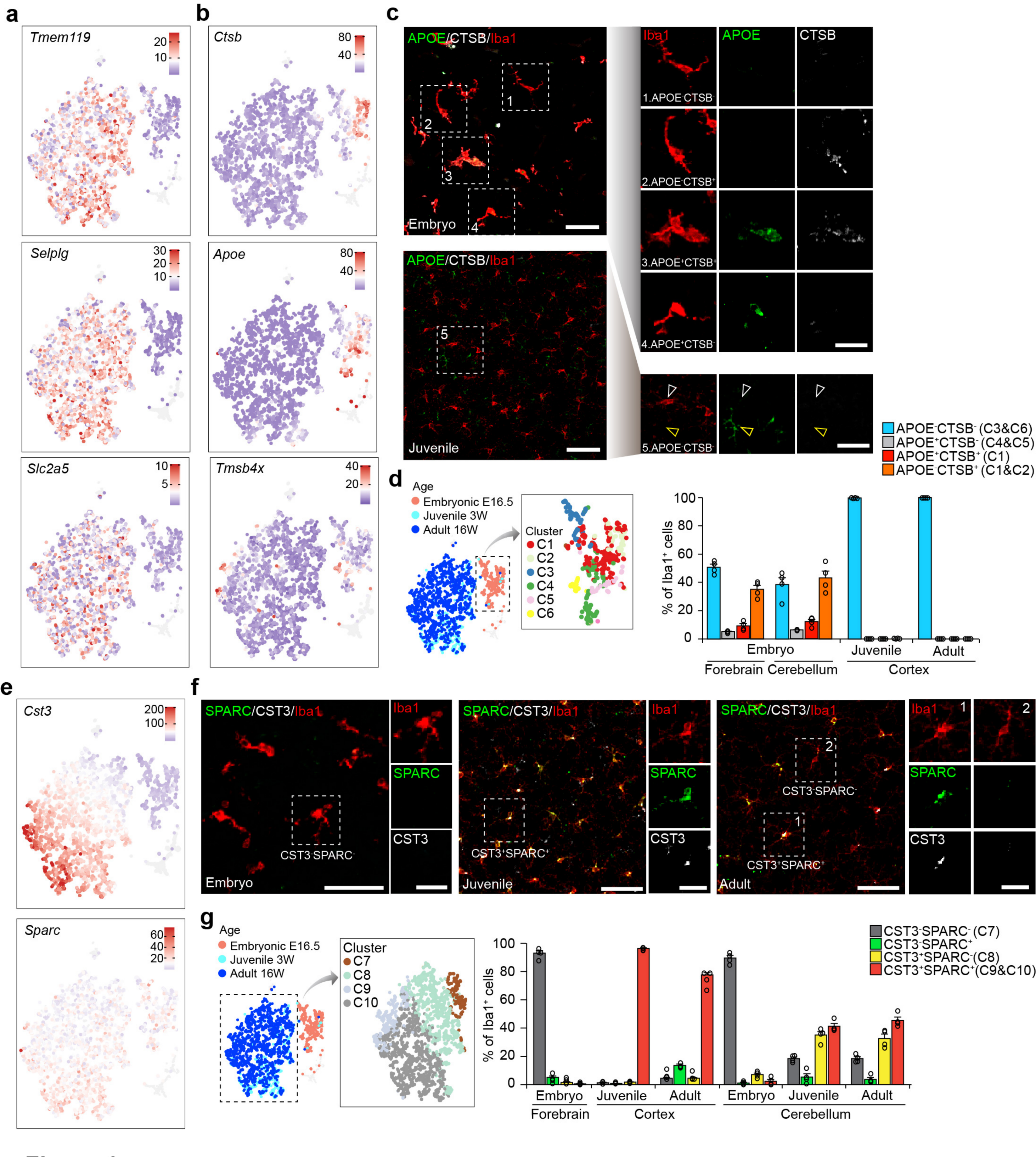


Figure 4

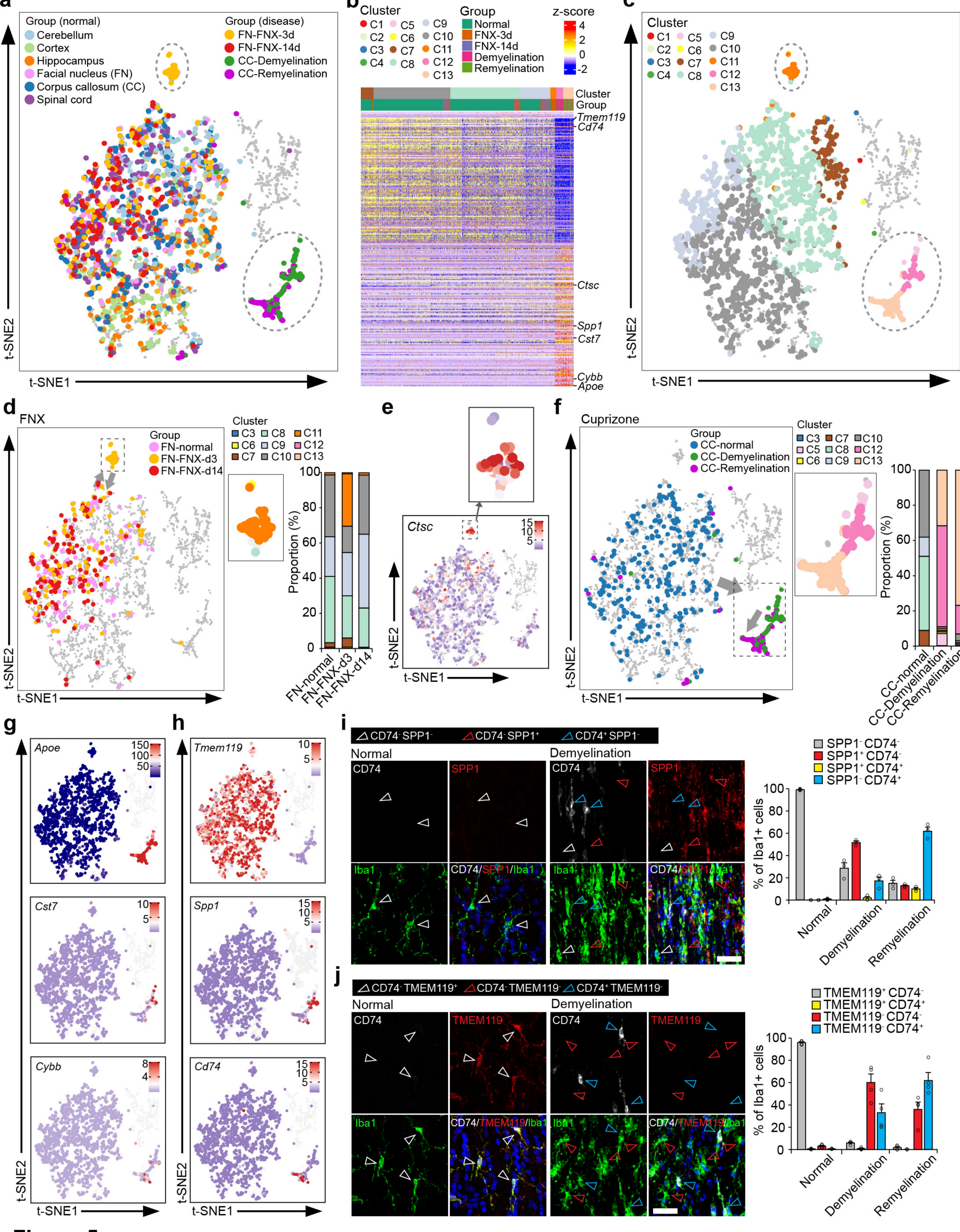


Figure 5

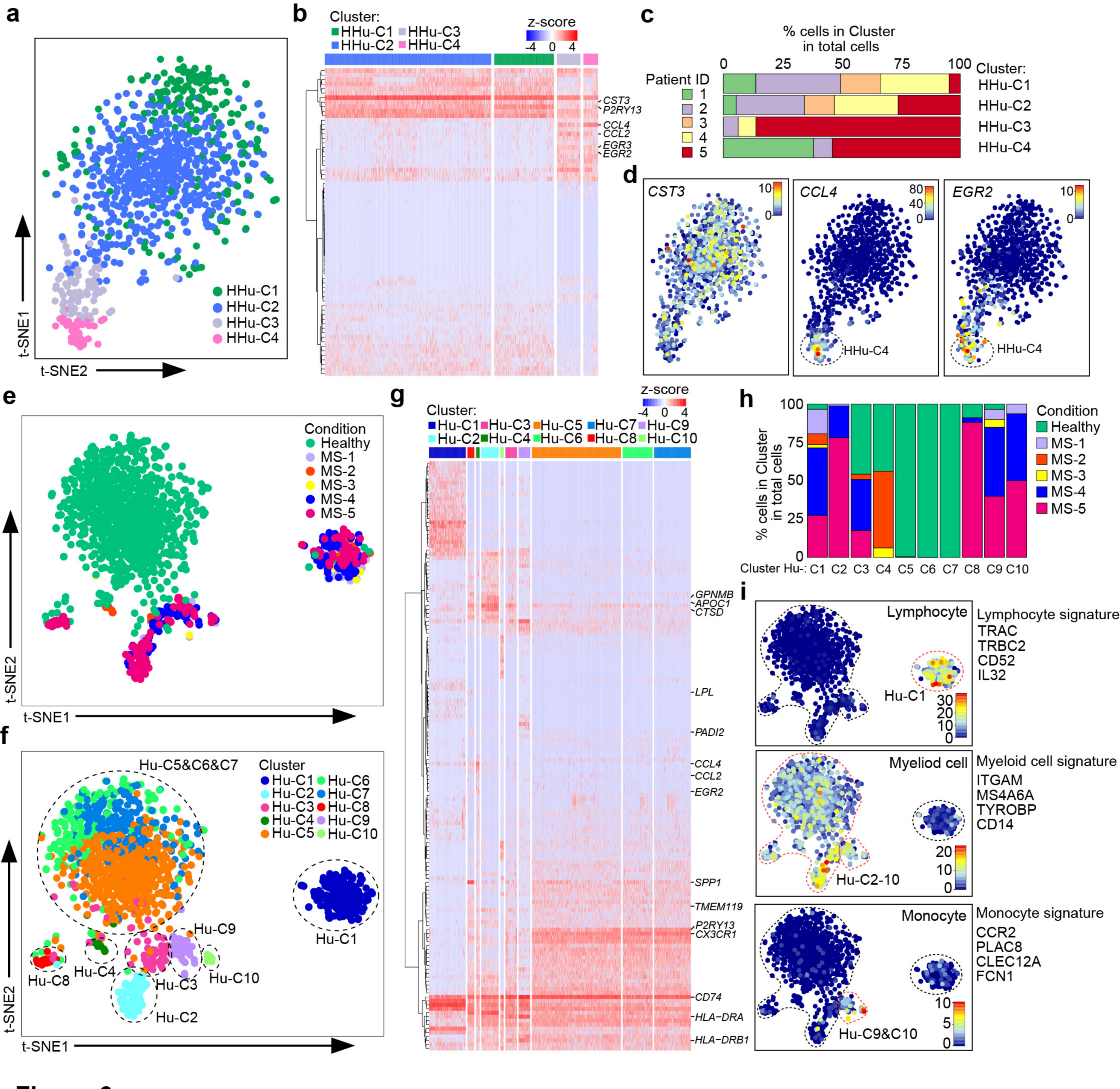


Figure 6

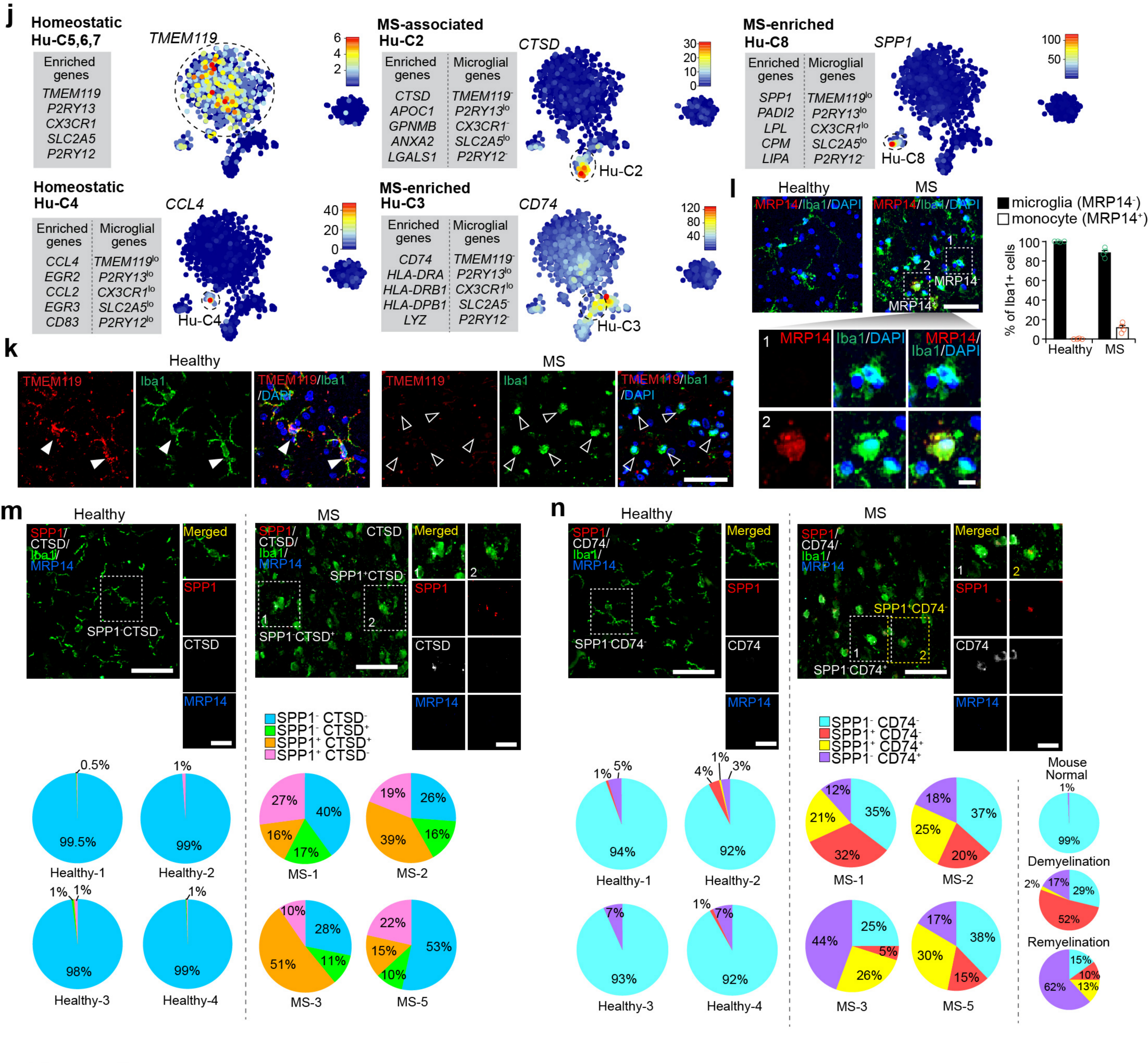


Figure 6