Colony-Stimulating Factor 1 Receptor (CSF1R) Regulates Microglia Density and Distribution, but Not Microglia Differentiation In Vivo

Highlights

- *csf1ra* and *csf1rb* together regulate microglia density in the adult zebrafish brain
- *csf1r* haploinsufficient microglia are normally differentiated and show normal signature
- *CSF1R* haploinsufficiency causes reduced microglia density and widespread depletion
- Microglia loss may be an early pathogenic event contributing to leukodystrophy

Authors

Nynke Oosterhof, Laura E. Kuil, Herma C. van der Linde, ..., Elly M. Hol, Mark H.G. Verheijen, Tjakko J. van Ham

Correspondence
t.vanham@erasmusmc.nl

In Brief

Oosterhof et al. show that colony-stimulating factor 1 receptor (CSF1R) primarily regulates microglia density and not their normal differentiation. In addition, they find widespread depletion of microglia in CSF1R-haploinsufficient zebrafish and leukodystrophy patients, also in the absence of pathology, indicating that microglia depletion may contribute to loss of white matter.
Colony-Stimulating Factor 1 Receptor (CSF1R) Regulates Microglia Density and Distribution, but Not Microglia Differentiation In Vivo

Nynke Oosterhof,1 Laura E. Kuil,1 Herma C. van der Linde,1 Saskia M. Burm,2 Woutje Berdowski,1 Wilfred F.J. van Ijcken,3 John C. van Swieten,4,5 Elly M. Ho,2,6 Mark H.G. Verheijen,7 and Tjakko J. van Ham1,8,*

1Department of Clinical Genetics, Erasmus MC, University Medical Center Rotterdam, Wytemaweg 80, 3015 CN Rotterdam, the Netherlands
2Department of Translational Neuroscience, Brain Center Rudolf Magnus, University Medical Center Utrecht, Utrecht University, Utrecht, the Netherlands
3Department for Biomics, Erasmus MC, University Medical Center Rotterdam, Wytemaweg 80, 3015 CN Rotterdam, the Netherlands
4Department of Neurology, Erasmus MC, University Medical Center Rotterdam, Rotterdam, the Netherlands
5Department of Clinical Genetics, VU Medical Center, Amsterdam, the Netherlands
6Department of Neuroimmunology, Netherlands Institute for Neuroscience, an Institute of the Royal Netherlands Academy of Arts and Sciences, Amsterdam, the Netherlands
7Department of Molecular and Cellular Neurobiology, CNCR, Amsterdam Neuroscience, VU University, Amsterdam, the Netherlands
8Lead Contact
*Correspondence: t.vanham@erasusmc.nl
https://doi.org/10.1016/j.celrep.2018.06.113

SUMMARY

Microglia are brain-resident macrophages with trophic and phagocytic functions. Dominant loss-of-function mutations in a key microglia regulator, colony-stimulating factor 1 receptor (CSF1R), cause adult-onset leukoencephalopathy with axonal spheroids and pigmented glia (ALSP), a progressive white matter disorder. Because it remains unclear precisely how CSF1R mutations affect microglia, we generated an allelic series of csf1r mutants in zebrafish to identify csf1r-dependent microglia changes. We found that csf1r mutations led to aberrant microglia density and distribution and regional loss of microglia. The remaining microglia still had a microglia-specific gene expression signature, indicating that they had differentiated normally. Strikingly, we also observed lower microglia numbers and widespread microglia depletion in postmortem brain tissue of ALSP patients. Both in zebrafish and in human disease, local microglia loss also presented in regions without obvious pathology. Together, this implies that CSF1R mainly regulates microglia density and that early loss of microglia may contribute to ALSP pathogenesis.

INTRODUCTION

Microglia are specialized brain macrophages whose functions in the brain include phagocytosis and provision of trophic support (Paolicelli et al., 2011; Safajian et al., 2016; Stevens et al., 2007; Tremblay et al., 2010; van Ham et al., 2012). Mutations in several genes that are highly expressed in microglia cause progressive white matter brain diseases (Meuwissen et al., 2016; Paloneva et al., 2002; Prinz and Priller, 2014; Rademakers et al., 2011). For example, dominant loss-of-function mutations in colony-stimulating factor 1 receptor (CSF1R) cause adult-onset leukoencephalopathy with axonal spheroids and pigmented glia (ALSP), also known as hereditary diffuse leukoencephalopathy with axonal spheroids (HDLS) (Konno et al., 2017; Wider et al., 2009). Even though low expression of Csf1r has been reported in some neurons in the hippocampus, the expression of CSF1R is almost exclusive to microglia, suggesting that ALSP pathogenesis involves microglia dysfunction (Luo et al., 2013). But where one study showed reduced microglia numbers in cortical layers 3 and 4 in postmortem end-stage ALSP brain sections, another showed increased microglia numbers during earlier ALSP disease stages (Oyanagi et al., 2017; Tada et al., 2016). The mechanism whereby heterozygous CSF1R mutations affect microglia and, consequently, brain homeostasis is still unknown. Insight into ALSP pathogenesis will therefore contribute to our understanding of microglia function in the vertebrate brain and of microglia involvement in other brain diseases.

Even though CSF1R signal transduction has been studied extensively in macrophages, it is not entirely clear how defective CSF1R signaling affects microglia in vivo. Activation by one of the two CSF1R ligands (CSF-1 or macrophage colony-stimulating factor [M-CSF]) or interleukin 34 (IL-34) leads to auto-phosphorylation of the tyrosine kinase receptor. In vitro, downstream activation of signal transduction pathways regulates the production, survival, differentiation, and function of macrophages (Dai et al., 2002; Erblich et al., 2011; Ginhoux et al., 2010; Wang et al., 2012). Genetic evidence for the consequences of CSF1R activation in vivo indicates that CSF1R primarily plays a homeostatic role in regulating the viability and proliferation of microglia (Cecchini et al., 1994; Jenkins et al., 2013). Indeed, genetic deficiency of CSF1R signaling reduces protection against bacterial infection, mainly by limiting macrophage supply (Cheers et al., 1989; Pagán et al., 2015; Teitelbaum et al., 1999; Wang et al., 2012). In contrast, by showing that
Csfr1<sup>−/−</sup> macrophage precursors have the same lineage potential as those in the wild-type, differentiating efficiently into macrophages but failing to form colonies, a recent study concluded that Csfr1 deficiency has little effect on myeloid differentiation in vivo (Endele et al., 2017).

Loss of Csfr1 in mice leads to an almost complete absence of microglia and also to severe developmental abnormalities and a shorter lifespan (Dai et al., 2002; Erblich et al., 2011; Ginhoux et al., 2010). Csfr1<sup>−/−</sup> brains show widened cerebral ventricles, which is also observed in ALSP patients. Mice lacking microglia also show cerebrovascular defects and reduced numbers of oligodendrocyte lineage cells (Erblich et al., 2011; Hagemeyer et al., 2017; Nandi et al., 2012). In addition, postnatal pharmacological inhibition of CSF1R in mice reduces the number of oligodendrocytes and oligodendrocyte precursor cells (OPCs) in a region-dependent manner (Hagemeyer et al., 2017). The latter effect could predispose to myelination defects later in life.

To understand the effect of CSF1R haploinsufficiency on microglia, we used the zebrafish as a model organism. Zebrafish are an upcoming genetic model organism to study brain diseases, including leukoencephalopathies (Zhang et al., 2016). They are highly suitable for in vivo imaging because they develop externally and are transparent at early stages (Haud et al., 2011; Oosterhof et al., 2015; Zhang et al., 2016). Previously, we identified the zebrafish microglia transcriptome, which shares high similarity with mouse and human microglia transcriptomes (Gosselin et al., 2017; Oosterhof et al., 2017). Zebrafish express two homologs of human CSF1R: csf1ra and csf1rb. We found that zebrafish csf1ra mutants show reduced microglia numbers only during development, partially mimicking mouse mutants. This suggests that the cellular functions of CSF1R are highly conserved between species but that zebrafish csf1rb and csf1ra are likely partially redundant. In the present study we therefore created an allelic series of zebrafish csfr loss-of-function mutants in which we observed local loss of microglia, a general reduction in microglia numbers, and an aberrant distribution of microglia. Because we found that dysregulation of microglia density was a primary consequence of csfr haploinsufficiency, we next investigated whether CSF1R haploinsufficiency also affects microglia density in postmortem brain tissue of ALSP patients. This revealed widespread depletion of microglia and a general reduction in microglia density. In humans and zebrafish alike, changes in microglia density and distribution in the absence of obvious myelin pathology implied that loss of microglia may be an early event in ALSP pathogenesis.

**RESULTS**

**Zebrafish Csf1ra and Csf1rb Together Are Functionally Homologous to Mammalian CSF1R**

To study how CSF1R mutations affect microglia and the brain, we exploited the fact that zebrafish have two homologs for human CSF1R: Csf1ra and Csf1rb. Both of these are highly expressed in adult zebrafish microglia (Figure 1A; Oosterhof et al., 2017). Unlike Csf1r knockout mice, which are almost completely devoid of microglia, zebrafish with homozygous loss-of-function mutations only in csf1ra (from here on called csf1ra<sup>−/−</sup>), show reduced microglia numbers only during early development (Herbomel et al., 2001). This suggests that csf1rb and csf1ra share a role in microglia development.

To test this, we introduced a premature stop codon in exon 3 of the csf1rb gene by transcription activator-like effector nuclease (TALEN)-mediated genome editing and assessed microglia numbers by neutral red staining (Figures 1B and S1A), which can be used to label microglia in zebrafish larvae in vivo. Although the microglia numbers in homozygous csf1rb mutants were a little lower than in the wild-type, mutants deficient in both csf1ra and csf1rb (from here on called csf1r<sup>PM</sup>), were almost completely devoid of microglia (Figures 1C and 1D). The absence of microglia in csf1r<sup>PM</sup> mutants was confirmed in larval and adult zebrafish by immunostaining for L-plastin (Figures 1E, 2A, and 2B). Although microglia were almost completely absent in csf1r<sup>PM</sup> adult animals, other macrophage populations were still present in adult organs, including the skin and the intestine (Figure S1B). Adult csf1r<sup>PM</sup> animals were viable and, in-cross mating of csf1r<sup>PM</sup> adult animals, produced viable homozygous mutant offspring (data not shown). However, after around 3 months of age, mutant animals occasionally showed seizure-like behavior, and their survival rate was lower than that of wild-type animals (data not shown). Some csf1r<sup>PM</sup> brains displayed signs of cerebral hemorrhaging that were consistent with the hemorrhages previously reported in Csfr1<sup>−/−</sup> mice (Erblich et al., 2011). These data show that zebrafish Csfr1a and Csfr1b both regulate the development of the microglia population, and both are thus functionally homologous to mammalian CSF1R.

**Csfr1 Regulates Microglia Density and Distribution Independent of Brain Pathology**

Previous studies indicate that the density of tissue macrophages, including microglia, is affected by reduced CSF1R signaling (Naito et al., 1991; Sasaki et al., 2000; Umeda et al., 1996; Wegiel et al., 1998). To validate this in zebrafish, we used neutral red labeling and immunohistochemistry to assess microglia numbers in a series of csf1r mutant zebrafish larvae consisting of csf1ra<sup>−/−</sup>, csf1ra<sup>−/−</sup>, csf1rb<sup>−/−</sup>, csf1ra<sup>−/−</sup>;bs<sup>−/−</sup>, and csf1rb<sup>−/−</sup> animals. At the larval stage, a gradual reduction in the number of csf1r alleles resulted in a corresponding decrease in microglia numbers (Figures 1C–1E). The greater reduction in microglia numbers in csf1ra<sup>−/−</sup> mutants than in csf1rb<sup>−/−</sup> mutants suggests that csf1ra is more important during early development. In adult zebrafish, however, microglia numbers in csf1rb<sup>−/−</sup> mutants were strongly reduced, whereas, in csf1ra<sup>−/−</sup> mutants, they were more comparable with those in the wild-type (Figure 2C), suggesting differential requirements of csf1ra and csf1rb in microglia at different developmental stages. Surprisingly, in 5-month-old adult csf1ra<sup>−/−</sup>;bs<sup>−/−</sup> mutants, we observed that, although microglia were absent in the dorsolateral side of the optic tectum, they appeared to accumulate in the underlying deep brain regions (Figures 2A and 2B).

To investigate whether any pathological hallmarks of ALSP are also present in csf1r mutant zebrafish, we assessed tissue and white matter integrity in adult csf1ra<sup>−/−</sup>, csf1ra<sup>−/−</sup>;bs<sup>−/−</sup>, and csf1r<sup>PM</sup> mutants. H&E labeling did not reveal signs of brain pathology (data not shown), nor did immunolabeling for Claudin K (Cldnk)—which labels myelin tracts throughout the zebrafish brain—reveal major loss of myelin, even in csf1r<sup>PM</sup> mutants.
Figure 1. Microglia Numbers during Development Are csfr Dosage-Dependent
(A) Counts per million (CPM) expression values of csf1ra and csf1rb from our previous RNA sequencing study in adult zebrafish microglia (Oosterhof et al., 2017).
(B) Schematic representation of the csf1rb mutation introduced with TALEN-mediated genome editing.
(C and D) 5 days post fertilization (dpf), WT, csf1ra−/−, csf1rb−/−, csf1ra−/−;b+/−, and csf1rDM larvae were treated with neutral red for 2.5 hr. Images were acquired with a stereomicroscope, and microglia numbers were determined by counting the number of neutral red dots. n is at least 15 zebrafish/genotype for (C) and at least 7 for (D).
(E) 4 dpf, WT, csf1ra−/−, csf1ra−/−;b+/− and csf1rDM were labeled with an antibody against L-plastin (Spangenberg et al., 2016), and L-plastin-positive cells were quantified in the optic tecti. n is at least 6 zebrafish/genotype.

LBD, ligand-binding domain; TMD, transmembrane domain; TKD, tyrosine kinase domain. Error bars represent SD. **p < 0.01, ***p < 0.001 (one-way ANOVA, Bonferroni multiple testing correction). Scale bar, 40 μm.
To determine whether csf1r mutants display more subtle myelin abnormalities, such as degeneration, hypomyelination, or hypermyelination, we analyzed their white matter by electron microscopy (EM). We observed highly myelinated regions in the midbrain containing multilayered myelin sheets that resembled those in mammals but no apparent abnormalities in the multilayered myelin sheets in csf1r mutants (Figures S2B and S2C). Immunolabeling for Sox10 also indicated normal numbers of oligodendrocyte lineage cells in csf1r mutants (Figure S2D). Together, this indicates that csf1r deficiency in zebrafish does not result in overt myelin degeneration at this adult stage.

To establish whether loss of csf1r causes more subtle pathological changes, we performed RNA sequencing on brains of adult csf1r mutant zebrafish that were ~8 months old (Figure 3A). Multidimensional scaling of gene expression data showed clustering of the samples based on the csf1r mutation status (wild-type [WT], csf1ra−/−, csf1ra−/−:b+/-, and csf1rDM), indicating csf1r-dependent changes in gene expression (Figure 3B). Differential gene expression analysis between WT and csf1rDM mutant brains revealed 154 differentially expressed genes, 85 of which (e.g., spi1b, irf8, csf1ra, and csf1rb) we had previously identified as part of the zebrafish microglial transcriptome (Figures 3C–3E; Table S2; Oosterhof et al., 2017). Hierarchical clustering of the samples on the basis of 154 differentially expressed genes revealed that csf1ra−/−:b+/- mutants clustered with csf1rDM mutants, whereas csf1ra−/− mutants clustered with the WT (Figure 3D). This suggests that loss of csf1r leads mainly to reduced expression of microglia-specific genes, which indicates that loss of csf1r in zebrafish predominantly affects microglia. The downregulated genes that were not specifically expressed in microglia included the cysteine-glutamine exchanger slc7a11 and growth hormone 1 as well as many poorly annotated genes (Table S2). This indicates that csf1r deficiency and, thus, loss of microglia causes very few molecular changes and no obvious myelin-related pathology in ~8-month-old adult zebrafish.

Csfr-Deficient Microglia Increase the Expression of Genes Involved in Chemotaxis and Migration

To assess in more detail how csf1r deficiency affects microglia independently of brain pathology, we performed RNA sequencing on microglia that were sorted by fluorescence-activated cell sorting (FACS) from WT, csf1ra−/−, and csf1ra−/−:b+/- mutant brains that were dissected from ~9-month-old zebrafish (Figure 4A). Multidimensional scaling revealed clustering of the samples on the basis of csf1r mutation status, indicating csf1r-dependent changes in microglial gene expression (Figure 3B). Based on our microglia density measurements and the importance of csf1rb for adult microglia, we reasoned that csf1ra−/−; b+/- mutant microglia could mimic the CSF1R haploinsufficiency that occurs in ALSP patient microglia and compared the microglia gene expression of these mutants with that of controls. We identified 1,466 genes that were differentially expressed between csf1ra−/−;b+/- and WT microglia (Figure 4C; Table S3). Interestingly, the normalized expression values of 750 of the 1,466 differentially expressed genes in csf1ra−/− mutant microglia lay in between those of csf1ra−/−;b+/- and WT microglia (Figures 4C and 4D; Table S3). Because more than half of the genes differentially expressed between WT and csf1ra−/−;b+/- show csf1r-dependent changes in expression, this indicates that these genes are regulated by Csfr signaling, and their altered expression could be a primary consequence of csf1r deficiency. Gene ontology analysis of genes that showed csf1r-dependent changes in expression revealed that downregulated genes were associated with brain and nervous system development and with regulation of neuronal differentiation (Figure 4E). Upregulated genes were mainly associated with immune response, immune system process, and leukocyte chemotaxis (Figure 4F). The differentially expressed genes in the gene ontology classes associated with the upregulated genes were mainly chemokines and chemokine receptors (e.g., cxcl12a, ccl25b, ccl19a.1, and cxcr4b) (Figure 4G). In fact, the expression of most chemokines and chemokine receptors in zebrafish microglia was higher in csf1ra−/−:b+/- mutants than in the WT (Figure 4G), which may explain the aberrant microglia distribution in csf1ra−/−;b+/- mutants.

To test whether the expression changes observed in csf1ra−/−;b+/- microglia and the brain indicated a general microglia differentiation defect, we investigated whether adult csf1r mutants showed a loss of microglia-specific gene expression or a gain in gene expression associated with immature microglia or macrophages (Figures 4 and 5). Only 8 of the 300 most microglia-specific genes in zebrafish (many of which are also included in the mouse and human homeostatic microglia signature; e.g., slco2b1, pdgfbα, and scn4bβ) were significantly downregulated in csf1ra−/−;b+/- microglia, suggesting that there is no loss of a homeostatic microglia signature (Figures 5A, 5D, and 5E; Butovsky et al., 2014; Gosselin et al., 2017; Oosterhof et al., 2017; Zhang et al., 2014). Next we analyzed the expression of 378 zebrafish orthologs for genes that are strongly downregulated during microglia differentiation in the mouse brain to assess whether csf1r mutant microglia fail to downregulate genes specific to immature microglia (Matcovitch-Natan et al., 2016). Expression of only 10 of these 378 genes was increased in csf1ra−/−;b+/- mutant microglia compared with the expression in WT microglia (Figures 5B and 5F). We also found no evidence for increased expression...
of genes that discern microglia and macrophages (Figure 5C; Bennett et al., 2016). Additionally, csf1ra−/−; b+/− microglia were still highly ramified and showed no signs of activation (Figure 5G). This suggests that the csf1r-dependent changes in microglial gene expression are largely independent of differentiation status. Together, these data imply that the changes in the expression of genes involved in chemotaxis and cell migration in csf1r mutants are a specific consequence of csf1r deficiency and not of a global differentiation defect.

The Damage-Induced Proliferative Response of csf1r Mutant Microglia Is Delayed

Microglia respond quickly to damage by migration and proliferation, and CSF1R has been linked to this proliferative response of microglia (Gómez-Nicola et al., 2013). Therefore, to assess whether proliferation defects could be linked to aberrant microglia localization and, possibly, to microglia migration, we used our previously established neuronal ablation model. In this model, metronidazole (MTZ) treatment in zebrafish with brain-specific transgenic expression of nitroreductase (NTR) results in neuronal cell death (van Ham et al., 2012, 2014). We have shown previously that increasing the local demand for microglia by inducing neuronal cell death causes a strong local proliferative response by microglia (Oosterhof et al., 2017). To investigate whether microglia proliferation depends on csf1r dosage, we used proliferating cell nuclear antigen (Pcna) as a cell proliferation marker to assess microglia proliferation upon induction of neuronal ablation. One day after treatment, control NTR transgenic larvae showed that the microglia numbers had increased from 25 to 32 locally, with a corresponding increase in the fraction of Pcna+ microglia (Figure 6A). In contrast, csf1r mutant microglia, upon MTZ treatment, showed a much larger increase in microglia numbers, respectively, from 5 to 24 and from 2 to 12, but had not yet increased significantly in the fraction of Pcna+ microglia (Figure 6A). Therefore, the increased microglia numbers in treated csf1r mutants cannot be explained by the fraction of
dividing microglia relative to the control. Based on the much stronger increase in L-plastin+ cells in either of the mutants, however, one would expect a much higher fraction of Pcna+ cells in MTZ-treated cells versus controls if the increase is due to proliferation alone. Intriguingly, 2 days after treatment, the fractions of Pcna+ microglia were increased to similar levels in control and in csf1r mutants (Figure 6B). This showed that, although csf1ra−/− and csf1ra−/−;b+/- mutant microglia were able to mount a proliferative response, the proliferative response was delayed. Nevertheless, because numbers still increased, it seems that potential initial proliferation deficiencies were compensated through microglia recruitment. Therefore, the aberrant distribution of microglia upon csf1r deficiency (Figures 2A and 2B) may be a compensatory mechanism intended to meet the brain’s local demand for microglia.

Severe Microglia Depletion in Gray and White Matter of Postmortem ALSP Patient Brains

It has been reported that degenerated white matter in the brains of ALSP patients contains many CD68+ myeloid cells...
Figure 5. Differential Gene Expression of csf1r-Deficient Microglia Shows Normal Microglia Differentiation

(A) Volcano plot showing expression changes of the 300 most highly expressed microglia-specific genes in csf1ra<sup>−/−</sup>;b<sup>+/−</sup> mutant microglia (Oosterhof et al., 2017).

(B) Volcano plot showing the expression changes in csf1ra<sup>−/−</sup>;b<sup>+/−</sup> mutant microglia of normally downregulated genes during differentiation (Matcovitch-Natan et al., 2016) and of genes normally expressed in other macrophages in the CNS (Bennett et al., 2016).

(C) Volcano plot showing the expression changes of non-microglia myeloid genes.

(D) Expression values of zebrafish microglia-specific genes.

(E) Expression values of downregulated microglia-specific genes.

(F) Expression values of upregulated microglia-specific genes.

(G) Representative images of microglia (5-month-old fish) in the ventral part of the optic tectum labeled with an antibody against L-plastin. Scale bar, 20 μm.
Figure 6. The Response to Neuronal Cell Death of csf1r Mutant Microglia Depends More on Recruitment Than on Proliferation

(A and B) We used our previously described conditional neuronal ablation model (van Ham et al., 2012, 2014), in which treatment with metronidazole (MTZ) leads to selective ablation of neurons with transgenic expression of NTR (the nsfB gene encoding NTR). WT, csf1r<sup>-/-</sup>, and csf1r<sup>-/-</sup>;b<sup>+/</sup>/ larvae were [legend continued on next page]
and reduced numbers of IBA1-positive microglia (Konno et al., 2014; Oyanagi et al., 2017; Tada et al., 2016). We wanted to investigate whether altered microglia density and distribution, as we identified in zebrafish, would recur in non-degenerated brain tissue of ALSP patients. By immunohistochemistry, we therefore analyzed microglia morphology, distribution, and density in gray matter, normal-appearing white matter (NAWM; occipital lobe), and degenerated white matter (middle frontal gyrus and cingulate gyrus) of two ALSP patients and age-matched controls (Figures 7 and S4A). As in previous studies, we observed numerous HLA-DR+ cells and large, rounded CD68+ cells in degenerated white matter, whereas, apart from a few IBA1/CD68-double-positive cells, IBA1+ cells were almost completely absent (Figures 7A, S3A–S3C, and S4B). Because we also observed a few CD68+ cells with a low level of IBA1 expression (Figure S3D).

Most IBA1+ microglia still present in the degenerated white matter appeared in clusters of ~10–100 cells (Figure 7B). Interestingly, with the exception of sparse microglia clusters, microglia in NAWM and gray matter were also severely depleted (Figures 7B, 7C, S3B, and S3C). Many of these IBA1+ microglia clusters were located at the border between the gray and the white matter (Figure 7B). Although the few IBA1+ cells present in the white matter looked like foam cells, microglia in the gray matter and NAWM either looked activated or had a normal ramified morphology (Figure 7B). In all brain areas examined, we also observed areas of gray matter in which IBA1+ microglia had a normal distribution and a ramified morphology. However, the density of these microglia was ~50% lower in ALSP patient brain sections than in controls (Figure 7D). This is reminiscent of the findings in our zebrafish experiments, where we also observed regional differences in microglia density in unaffected brain tissue (Figures 2 and 7B and 7D). The loss of IBA1+ microglia and the aberrant microglia distribution and altered morphology in microglia clusters—not only in the gray matter but also in NAWM—indicate that microglial changes could precede white matter degeneration.

**Discussion**

Although mutations in genes that are particularly important for microglia can cause severe brain disorders, it is still unclear whether pathogenesis involves a gain or loss of specific microglia activities. Here we used zebrafish to investigate the effect of a gradual reduction in functional csf1r alleles on microglia numbers and microglia differentiation status and their response to tissue damage. We found that Csfr1 haploinsufficiency was correlated with a lower density of microglia in the dorsal part of the optic tectum. Additionally, we observed altered microglia distribution and local microglia depletion in the ventral and dorsolateral part of the optic tectum, respectively. Loss of three csf1r alleles did not severely impede the proliferative capacity of microglia to dying neurons, nor did it affect the homeostatic microglia signature. Instead, in response to increased phagocytic demand, csf1r mutant microglia initially increased their numbers locally through recruitment rather than proliferation. Accordingly, the expression of migration and chemotaxis genes in csf1r mutants was also increased. We also showed that, in the absence of extensive white matter degeneration in the occipital lobe of the cortex, CSF1R haploinsufficiency results in widespread depletion and aberrant distribution of IBA1+ microglia in humans. These findings support the presence of a disease mechanism in which CSF1R haploinsufficiency reduces microglia density, causes microglial relocation, and results in depletion of functional microglia.

Although the focus of this study is brain microglia, CSF1R haploinsufficiency could potentially affect other macrophages, such as those in the gut, or even neurons, because a low level of Csfr1 expression was reported in a few scattered neurons in the mouse hippocampus (Luo et al., 2013). Because the composition of the gut microbiome or perturbed barrier function of the gut has major effects on microglia and on the CNS (Emry et al., 2015; Sampson et al., 2016) it is possible that, in ALS patients, defects in other macrophage populations, such as those in the intestine, play a role in pathogenesis. Nevertheless, gastrointestinal symptoms related to perturbed gut barrier function were not reported in ALSP patients (Konno et al., 2017). Additionally, our zebrafish data do not imply an increased inflammatory response or response to microbial infection. Given that a potentially protective role of Csfr1 has been described in neurons, it is possible that these protective effects are reduced due to CSF1R deficiency and could be a contributing factor in disease. Because neuronal loss is not obvious in ALSP patients, however, and severe phenotypes of Csfr1−/− mice are largely rescued by hematopoietic cells, we speculate that this is unlikely to play a major part (Bennett et al., 2018).

The CSF1R coding sequence and function are well conserved across species. CSF1R-deficient zebrafish, rodents, and, most likely, humans lack microglia, are osteopetrotic, and occasionally show cerebral hemorrhages (data not shown; Dai et al., 2002; Erblich et al., 2011; Ginhoux et al., 2010; Monies et al., 2017). Our data indicate that csf1r haploinsufficiency leads to a local loss of microglia, possibly through maldistribution of microglia. This is similar to the aberrant distribution of microglia and widespread loss of microglia we observed in the NAWM and gray matter of postmortem ALSP patient brains. Interestingly, based on neuropathological analysis of different ALSP stages, in the early stages, microglia numbers were predicted to be higher than in controls, and microglia appear to be activated in specific brain regions (Oyanagi et al., 2017). This is reminiscent of the increased microglia density we observed in deep brain regions of csf1r−/−; b+/- haploinsufficient zebrafish (Oyanagi et al., 2017). Similarly, microglia numbers are also higher in some brain regions in heterozygous Csfr1 mutant mice than in

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treated with MTZ at 5 dpf for 16 hr and fixed for immunohistochemistry (whole-mount) at 6 dpf (A) and 7 dpf (B). Immunostaining was performed for dividing (Pcna+) microglia (L-plastin+), and the entire forebrain (dotted lines) was imaged and quantified. Scale bars, 40 μm. Group sizes were at least n = 10 zebrafish larvae (A) and at least n = 4 (B). Error bars indicate SD. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 (one-way ANOVA, Bonferroni multiple testing correction).
control animals, but it is unclear whether microglia density is reduced in other areas or at later stages in the mouse (Chitu et al., 2015). Between them, these observations indicate that CSF1R haploinsufficiency causes aberrant microglia distribution, where some regions become devoid of microglia.

Although microglia are efficient phagocytes that clear dead cells, dysfunctional synapses, and myelin (Ling, 1979; Safaiyan et al., 2016; Sierra et al., 2010), the accumulation of myelin debris in microglia can compromise their phagocytic capacity and can also lead to microglial senescence (Boven et al., 2006; Neumann et al., 2009; Safaiyan et al., 2016). Based on the size and morphology of large numbers of CD68+ cells in degenerated white matter, the accumulation of debris could have preceded their presence. We cannot exclude that CD68+ cells include infiltrated macrophages because other macrophage and microglia markers, including IBA1 and P2RY12, appear to be lost or very low in these cells (Tada et al., 2016). Regardless, accumulation of myelin debris, as occurs during normal aging, may contribute to the progressive loss of functional IBA1+ microglia over the course of the disease. In fact, it was shown in a tuberculous infection model that, because of reduced csf1r signaling, the loss of macrophages was driven by a failure to meet phagocytic demand (Pagan et al., 2015). Consistent with this idea, the morphology of microglia among clusters in ALSP patients ranged from ramified to completely round and foamy in appearance, most likely because of the accumulation of phagocytized myelin debris in microglia (Boven et al., 2006). Simultaneously, it is possible that one functional CSF1R copy is not sufficient to sustain both normal microglia survival and proliferation because microglia turn over in humans in adulthood (Askew et al., 2017; Reu et al., 2017). Together, this indicates that CSF1R-dependent loss of microglia in ALSP patients may be progressive.

The absence of overt neuropathology or myelin pathology in csf1r mutant zebrafish may be related to their relatively young age, the fact that the CNS of the zebrafish is smaller and less complex than that of humans, or the time needed for the pathology to develop in humans. The pathological hallmarks of ALSP are observed primarily in the neocortex, which is unique to mammals and has expanded immensely during evolution, particularly in primates and humans (Florio and Huttner, 2014; Hofman, 2014). Because the neocortex is rich in white matter, it may be more susceptible to pathology than the zebrafish brain, in which there is relatively little white matter (Merrifield et al., 2017). Consistent with this, mutations that result in a relatively mild pathology in mice can lead to severe leukodystrophy in humans (Choquet et al., 2017). Additionally, it takes about 30–40 years before ALSP becomes symptomatic, whereas mice and zebrafish live only a few years (Konno et al., 2017). Even though the csf1r mutant zebrafish brain is relatively unaffected, the direct effects of csf1r mutations on microglia as described here were very similar to those in humans.

Although it is still unknown how long-term depletion of microglia in adulthood would affect brain homeostasis and how it might cause pathology, white matter degeneration is a hallmark of several other brain disorders classified as microgliopathies. For example, mutations in the microglia genes TREM2 and TYROBP cause Nasu-Hakola disease (NHD), which is also characterized by white matter pathology. Even though the precise pathogenic mechanisms remain elusive, these disorders support the idea that microglia are critical to the maintenance of myelin in adulthood. In fact, it was recently shown in the adult brain that lower microglia numbers lead to a reduction in the numbers of oligodendrocytes or OPCs in many brain regions (Hagemeyer et al., 2017). We anticipate that a progressive depletion of microglia occurs in ALSP that could lead to a lower number of myelinating cells in adulthood and could contribute to ALSP pathogenesis.

Like tissue macrophages, microglia influence the development and repair of organs by secreting trophic factors, including insulin-like growth factor 1 (IGF1), and by mediating signaling between cells (Eom and Parichy, 2017; Wlodarczyk et al., 2017; Wynn et al., 2013). Local depletion of microglia could lead to a failure to provide such trophic factors, which could contribute to ALSP pathogenesis; for example, by affecting the capacity to form new oligodendrocytes.

In conclusion, the greatest effect of CSF1R haploinsufficiency seems to be a general reduction in microglia density in addition to large areas completely devoid of microglia. The partial or complete lack of microglia occurs in normal-appearing gray matter and NAWM, which suggests that loss of microglia may eventually result in ALSP pathology. Our gene expression data in an allelic series of csf1r-deficient microglia and brains therefore provide an opportunity to further delineate not only the function of csf1r in microglia but also the consequences for the brain. Elucidating these is crucial for a more comprehensive understanding of the physiological functions of microglia and microglia-dependent disease mechanisms. Several studies have shown that pharmacological inhibition of CSF1R causes microglia depletion and that, in mouse models, it ameliorates neurodegenerative disease-like symptoms by depleting microglia or diminishing their proliferation and activation (Elmore et al., 2014; Olmos-Alonso et al., 2016; Spangenberg et al., 2016). Because microglia depletion may underlie and contribute to

Figure 7. ALSP Patient Brains Show Widespread Microglia Depletion.

Shown are representative images of IBA1 and CD68 staining of microglia in white and gray matter of postmortem brain tissue of two ALSP patients and two age-matched control donors.

(A) IBA1 and CD68 double-labeling in the white matter of the cingulate gyrus of ALSP patient and control tissue.
(B) Clusters of IBA1+ microglia are apparent at the borders between the gray matter and the white matter.
(C) Severe depletion of IBA1+ microglia in the gray and white matter of the occipital cortex.
(D) The ramified morphology of microglia in gray matter areas of ALSP patients was similar to that in controls. Quantification of microglia numbers in gray matter areas of the cingulate cortex, frontal cortex, and occipital cortex showed a homogeneous distribution of ramified microglia (5 gray matter areas, 1.5 mm² in size, per brain region per patient). Error bars indicate SD.

Scale bars, 50 µm (A), 500 µm (B, low magnification), 30 µm (B, high magnification), 100 µm (C), 50 µm (D). WM, white matter; GM, gray matter.
the development and progression of ALSP, this raises the question of whether long-term inhibition of CSF1R in neurodegenerative diseases like Alzheimer’s disease is a viable treatment option (Olmos-Alonso et al., 2016; Spangenberg et al., 2016). This warrants further studies to determine how the brain is affected by loss of microglia interactions and microglia-derived factors and to devise ways of promoting the supply of functional microglia.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental Information includes four figures and three tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.06.113.

**ACKNOWLEDGMENTS**

This work was sponsored by an Erasmus University Rotterdam fellowship, a ZonMW VENI grant (016.136.150), a Marie Curie Career Integration grant (232368), and an Alzheimer Nederland fellowship (WE.15-2012-01) (to T.J.v.H.) and an MKMD ZonMW grant (to E.M.H.). We thank Dr. B. Giepmans (322368), and an Alzheimer Nederland fellowship (WE.15-2012-01) (to E.M.H.). We thank Dr. B. Giepmans and A. Wolters (UMC Groningen) for advice regarding electron microscopy, T. van Gestel and Dr. G. Schaaf (Erasmus MC) for help with flow cytometry, the Netherlands Brain Bank for human brain tissue (coordinator Dr. I. Huitinga, Amsterdam, the Netherlands), and J. Wortel (VUMC) for her contribution to electron microscopy preparations.

**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no competing interests.

Received: March 21, 2018
Revised: May 23, 2018
Accepted: June 27, 2018
Published: July 31, 2018

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in VPS11 Causes an Autosomal Recessive Leukoencephalopathy Linked to
## STAR METHODS

### KEY RESOURCES TABLE

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(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Tjakko van Ham (t.vanham@erasmusmc.nl).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animal models
All zebrafish were maintained under standard conditions. Adult animals were kept in groups on a 14 hr light and 10 hr dark cycle. They were fed brine shrimp twice a day. Embryos were kept at 28°C in 10 mM HEPES-buffered E3 embryo medium. In larvae analyzed between 3-5 dpf sex-determination has not yet occurred. For microglia RNA-sequencing experiments groups of 3 animals were used, containing at least 1 male and 1 female. For whole brain RNA-sequencing groups of 3 animals, 1 male and 2 females, were used. The following existing transgenic or mutant zebrafish lines were used: Wild-type AB, Tg(mpeg1:EGFP) fish expressing GFP under the control of the mpeg1 promoter (Ellett et al., 2011), Tg(shhb:KalTA4,UAS-E1b:mCherry), UAS-E1b:NTR-mCherry,
microglia response to neuronal death was quantified in the larval forebrain. Immunostaining on adult brain slices was performed largely as described previously (Oosterhof et al., 2017). Fish were euthanized in ice water, after which the skull (containing the brain) was fixed in 4% PFA at 4°C for at least two days. They were then rehydrated to PBS (0.1% Triton X-100) and incubated in acetone at 30°C for 15 min. Afterward, larvae were washed with PBSTw (2x 10 min) and dH2O (2x 5 min) and incubated in acetone at 30°C for 20 min. This was followed by several washing steps in PBSTw (6x 5 min); dH2O (2x 5 min); and PBSTw (2x 5 min). The larvae were then incubated in blocking buffer (10% goat serum, 1% Triton X-100, 1% BSA in PBSTw) at 4°C for 3h. Primary-antibody labeling (1h) in 1% goat serum, 0.8% Triton X-100, 1% BSA in PBSTw was done at 4°C for three days. Larvae were washed in PBS-TS (10% goat serum, 1% Triton X-100 in PBS) (3x 1h); PBST (1% Triton X-100 in PBS) (2x 10 min); and PBS-TS (2x 1 min). The larvae were then incubated with the secondary antibody and Hoechst at 4°C for 2.5 days. Before imaging, the larvae were washed with PBS-TS (3x 1 h); and PBSTw (2x 1 h). Primary antibodies: PCNA (1:250, Dako); and L-plastin (1:500; gift from Yi Feng, University of Edinburgh). Secondary antibodies: DyLight alexa 488 and DyLight alexa 647 (1:500, ThermoFisher Scientific, Waltham, US); and alexa 594 (1:250, Invitrogen, Waltham, US). Imaging was done with a Zeiss LSM700 confocal microscope using a 20x water dipping lens (NA = 0.75).

**METHOD DETAILS**

**Vital dye labeling**

Neutral red labeling was performed as described previously (Herbomel et al., 2001). For all experiments in larvae, embryos were kept in E3 medium containing 0.003% 1-phenyl 2-thiourea (PTU) (Sigma-Aldrich) from 22 hpf onward. To assess microglia numbers, 5 or 6 dpf larvae were treated with 2.5 μg/ml neutral red in E3+PTU medium at 28°C for two hr. After two hr, larvae were incubated in E3+PTU for at least 30 min before imaging. For imaging, larvae were anesthetized with 0.016% MS-222 (Sigma-Aldrich) and embedded in 1.8% low melting point (LMP) agarose (VWR BDH Prolabo) with the dorsal side facing upward. Neutral red images were acquired by using a Leica M165FC stereo microscope.

**Conditional neuronal cell death**

For neuronal ablation, neuro-NTR transgenic zebrafish were used as described previously (Davison et al., 2007; van Ham et al., 2014). Larvae were treated with 2 mM metronidazole (MTZ) dissolved in DMSO at 28°C for 16 hr. MTZ was washed away with E3+PTU medium. Larvae were euthanized and fixed at different times after the start of treatment, at 1 day post treatment (20 h) and 2 days post treatment (48 h).

**Immunofluorescence staining**

Whole mount larvae This was usually done as described previously (Inoue and Wittbrodt, 2011). Larvae were fixed in 4% PFA at 4°C overnight, dehydrated to 100% MeOH, and kept at −20°C for at least two days. They were then rehydrated to PBSTw (PBS + 0.1% Tween). Antigens were retrieved by incubating the larvae in 150 mM Tris-HCl pH = 9.0 for 5 min, followed by incubation at 70°C for 15 min. Afterward, larvae were washed with PBSTw (2x 10 min) and dH2O (2x 5 min) and incubated in acetone at −20°C for 20 min. This was followed by several washing steps in PBSTw (6x 5 min); dH2O (2x 5 min); and PBSTw (2x 5 min). The larvae were then incubated in blocking buffer (10% goat serum, 1% Triton X-100, 1% BSA in PBSTw) at 4°C for 3h. Primary-antibody labeling (1h) in 1% goat serum, 0.8% Triton X-100, 1% BSA in PBSTw was done at 4°C for three days. Larvae were washed in PBS-TS (10% goat serum, 1% Triton X-100 in PBS) (3x 1h); PBST (1% Triton X-100 in PBS) (2x 10 min); and PBS-TS (2x 1 min). The larvae were then incubated with the secondary antibody and Hoechst at 4°C for 2.5 days. Before imaging, the larvae were washed with PBS-TS (3x 1 h); and PBSTw (2x 1 h). Primary antibodies: PCNA (1:250, Dako); and L-plastin (1:500; gift from Yi Feng, University of Edinburgh). Secondary antibodies: DyLight alexa 488 (1:500); and DyLight alexa 647 (1:500). Hoechst was used for nuclear staining. Images were taken with a Leica SP5 confocal microscope using a 20x water dipping lens (NA = 1.0). Total microglia numbers were quantified for the entire brain. The microglia response to neuronal death was quantified in the larval forebrain.

**Adult brain sections**

Immunostaining on adult brain slices was performed largely as described previously (Oosterhof et al., 2017). Fish were euthanized in ice water, after which the skull (containing the brain) was fixed in 4% PFA at 4°C overnight. Subsequently, the brains were carefully removed and dehydrated with a 25%, 50%, 75%, 100% MeOH series and kept at −20°C for at least 12 hr. After rehydration, brains were embedded in 4% w/v low melting point agarose in PBS and cut into 80 μm horizontal sections using a Microm HM 650V vibratome (ThermoFisher Scientific). Immunostainings on free-floating sections were performed as described. The sections were incubated in blocking buffer (10% goat serum, 0.5% Triton X-100 in PBS) at room temperature for 75 min, followed by incubation with the primary antibody in blocking buffer at 4°C overnight. The sections were thoroughly washed with PBST (0.5% Triton X-100) before incubation with the secondary antibody and Hoechst at 4°C overnight. The brain slices were then washed with PBST (5 x 30-60 min) and mounted on microscope slides using vectashield mounting medium H1000 (Vector Laboratories). Primary antibodies: L-plastin (1:1000; gift from Yi Feng, University of Edinburgh); ClaudinK (1:1000; gift from Thomas and Catherina Becker, Edinburgh); and Sox10 (1:500, Genetex). Secondary antibodies used were DyLight alexa 488 and DyLight alexa 647 (1:500, ThermoFisher Scientific, Waltham, US); and alexa 594 (1:250, Invitrogen, Waltham, US). Imaging was done with a Zeiss LSM700 confocal microscope using a 20x lens (NA = 0.75).
**Immunohistochemistry zebrafish paraffin sections**

4.5 month old zebrafish were euthanized in ice water and fixed in 4% PFA at 4°C over the weekend. Thereafter fish were washed in PBS before transferring them to 20% EDTA at room temperature for weeks for the bone to decalcify. Fish were embedded in paraffin and 10μm sections were cut and dried overnight at 37°C. Sections were deparaffinized and rehydrated to distilled water. Antigen retrieval was performed by heating in 0.1M sodium citrate pH = 6 for 13 min. After cooling slices were rinsed with PBS before blocking endogenous peroxidase activity by 30 min incubation in 0.6% H₂O₂, 1.5% sodium azide in PBS at room temperature. Slides were rinsed in PBS+ (0.5% protifar, 0.15% glycine) before L-plastin antibody incubation (1:1000) overnight at 4°C in PBS+. Slides were rinsed with PBS+ before incubation with conjugated secondary antibody (anti-rabbit-HRP 1:100) for 60 min at room temperature. After washing in PBS+ and PBS slides were incubated in DAB-substrate (DAKO liquid DAB substrate-chromogen system). After washing with distilled water a counterstaining with hematoxylin was performed for 5 min. After washing in distilled water slices were dehydrated and mounted with Entellan (Merck).

**Paraffin sections human brain**

Tissue sections were characterized for the presence of microglia by staining for IBA1, HLA-DR and CD68 as previously described (Hovens et al., 2014). Paraffin sections seven μm in thickness were collected on Superfrost Plus glass slides (VWR international, Leuven, Belgium) and dried at 37°C overnight. Tissue sections were deparaffinized and rehydrated to distilled water. Endogenous peroxidase was quenched in 0.3% H₂O₂ (Merck, Darmstadt, Germany) in phosphate-buffered saline (PBS), and antigen was retrieved by heating the slides in citrate buffer (10 mM; pH 6.0). The sections were then incubated with normal horse serum blocking buffer (TBS supplemented with 2% horse serum, 1% bovine serum albumin (Sigma-Aldrich, St. Louis, USA); 0.1% Triton X-100 and 0.05% Tween (Merk)) for 30 min at RT. Thereafter, sections were incubated with rabbit anti-human IBA1 antibodies (1:1000 TBS-BSA 1%; WAKO Chemicals, Richmond, USA) overnight at 4°C. After rinsing, the sections were incubated with biotinylated goat anti-rabbit antibodies (1:400 in TBS-BSA 1%; Vector Laboratories, Burlingame, USA) for 1 h at RT; this was followed by incubation with the avidin-biotin complex (1:800 in TBS; Vector Laboratories) for 45’ at RT. The sections were then developed with 3,3’-diaminobenzidine (DAB; 0.05 mg/ml, Sigma-Aldrich). After washing in distilled water, slides were dehydrated and embedded in Entellan (Merck). In between all steps, sections were extensively rinsed in TBS. Whole slides were digitalized using a NanoZoomer digital slide scanner (Hamamatsu) and the slides were analyzed using Hamamatsu software NDPview2. The numbers of IBA1+ microglia were quantified in 5 areas of 1.5 mm² per brain region per patient.

For the CD68/IBA1 immunofluorescent double-staining, tissue sections were deparaffinized and rehydrated to distilled water. Antigen retrieval was performed by heating the slides in citrate buffer (10 mM; pH 6.0). Sections were then rinsed in blocking buffer (PBS, 0.5% protifar (Nutricia), 0.15% glycine (Sigma-Aldrich, St. Louis, USA), 0.4% Triton X-100) for 2 times 2 min at RT. Thereafter, sections were incubated with blocking buffer containing rabbit anti-human IBA1 antibodies (1:500, WAKO Chemicals, Richmond, USA) and mouse anti-human CD68 (1:50, clone KP-1) antibodies overnight at 4°C. After rinsing with blocking buffer, sections were incubated with secondary anti-rabbit Alexa 488 (1:200, ThermoFisher Scientific, Waltham, US) and anti-mouse Cy3 antibodies (1:200) for 1h at RT. Autofluorescence was blocked by incubating slides in Sudan Black solution for 5 min at RT. After washing the slides were embedded in Mowiol containing DAPI.

**Luxol fast blue staining**

Paraffin sections seven μm in thickness were collected on glass slides (Superfrost Plus, VWR international, Leuven, Belgium) and dried at 37°C. Sections were deparaffinized and hydrated to 95% ethanol and incubated in Luxol fast blue solution (0.1% w/v luxol fast blue, Sigma-Aldrich, in 95% ethanol) overnight at 55°C. The sections were then quickly rinsed in 95% ethanol and incubated in lithium carbonate solution (0.05% w/v, Sigma-Aldrich, in distilled H₂O) for 30 s. Next, the samples were rinsed in 70% ethanol and subsequently in ddH₂O. The sections were dehydrated to 100% ethanol and mounted with Entellan (Merck). Whole slides were digitalized using a NanoZoomer digital slide scanner (Hamamatsu), and the slides were analyzed using Hamamatsu software NDPview2.

**RNA sequencing**

**Whole brain**

For whole brain transcriptomic analysis brains were dissected as described previously (Oosterhof et al., 2017) and snap frozen in liquid nitrogen. Zebrafish between 6-12 mpf were used. Zebrafish were euthanized in ice water and heads were severed behind the gills, and the lower jaw, gills, and eyes were removed. The brains were taken out of the skull and cut using scalpels. Brains were homogenized followed by total RNA isolation using Trizol. Total RNA quality for triplicates of WT brain, csf1ra−/− brain, csf1ra−/−;b+/− brain and csf1rΔm brain was assessed on an Agilent Technologies 2100 Bioanalyzer using a RNA nano assay. All samples had RIN value greater than 9.10. Triplicate RNA-seq libraries were prepared according to the illumina TruSeq stranded mRNA protocol (https://www.illumina.com). Briefly, 200 ng of total RNA was purified using poly-T oligo-attached magnetic beads to end up with poly-A containing mRNA. The poly-A tailed mRNA was fragmented and cDNA was synthesized using SuperScript II and random primers in the presence of Actinomycin D. cDNA fragments were end repaired, purified with AMPure XP beads, A-tailed using Klenow exo-enzyme in the presence of dATP. Paired end adapters with dual index (illumina) were ligated to the A-tailed cDNA fragments and
purified using AMPure XP beads. The resulting adaptor-modified cDNA fragments were enriched by PCR using Phusion polymerase as followed: 30 s at 98°C, 15 cycles of (10 s at 98°C, 30 s at 60°C, 30 s at 72°C), 5 min at 72°C. PCR products were purified using AMPure XP beads and eluted in 30 μl of resuspension buffer. One microliter was loaded on an Agilent Technologies 2100 Bioanalyzer using a DNA 1000 assay to determine the library concentration and for quality check.

**Bridge amplification, sequencing by synthesis and data analysis**

Cluster generation was performed using the Illumina TruSeq SR Rapid Cluster kit v2 (cBot) Reagents Preparation Guide (https://www.illumina.com/). Briefly, 12 RNA-seq libraries were pooled together to get a stock of 10 nM. One microliter of the 10 nM stock was denatured with NaOH, diluted to 6 pM and hybridized onto the flowcell. The hybridized products were sequenced, amplified, linearized, and end-blocked according to the Illumina Single Read Multiplex Sequencing user guide. After hybridization of the sequencing primer, sequencing-by-synthesis was performed using the HiSeq 2500 with a single read 50-cycle protocol followed by dual index sequencing. Reads were aligned against the GRCz10 genome using HISAT2 (version 2.0.4) (Kim et al., 2015). Counts were generated for each gene from the Ensembl (version 85) transcriptome analysis of GRCz10, using htseq-count (version 0.6.0) (Anders et al., 2015).

**Microglia**

Tg(mpeg1:EGFP) zebrafish were euthanized in ice water. The heads were severed behind the gills, and the lower jaw, gills, and eyes were removed. The brains (4-5 per sample) were taken out of the skull and cut using scalpels. This was followed by dissociation in 0.25% trypsin and 0.1% EDTA in PBS for 2 hr at 4°C, while resuspending regularly. Upon complete dissociation of the brain, trypsin was inactivated by adding 1/6 volume of a stop solution (6 mM CaCl2 in PBS). The cells were collected in a 22% Percoll solution after being run through a 70 μm cell strainer, and ice-cold PBS was placed on top of the cell suspension and centrifuged at 1000 x g at 4°C for 45 min. The remaining cell pellet was resuspended in suspension solution (high-glucose DMEM without phenol red, 0.8 mM CaCl2). The cell suspension was transferred to FACs tubes with 35 μm cell strainer caps and FAC-sorted using a FACsaria III cell sorter (BD biosciences, New Jersey, USA). Dapi was added to exclude dead cells.

**RNaseq library synthesis and bioinformatics analysis**

Microglia were lysed and processed for RNA-seq largely as described (Oosterhof et al., 2017). FAC-sorted microglia were collected and lysed in RNase-free water containing 0.2% v/v Triton X-100 and RNase inhibitor. PolyA+ RNA was reverse transcribed using an oligo(dT) primer. Template switching by reverse transcriptase was achieved by using an LNA containing TSO oligo. The reverse-transcribed cDNA was pre-amplified with primers for 18 cycles followed by clean-up. Tagmentation was performed on 500 pg of the pre-amplified cDNA with Tn5 followed by gap repair. The fragmented library was extended with Illumina adaptor sequences by PCR for 14 cycles and then purified. The resulting sequencing library was measured on Bioanalyzer and equimolar samples were loaded onto a flowcell and sequenced according to the Illumina TruSeq v3 protocol on the HiSeq2500 with a single-read 50 bp and dual 9 bp indices. The sequencing reads were mapped against the GRCz10 zebrafish genome using the HiSat2 aligner (Pertea et al., 2016). To quantify the aligned and filtered data, the Bioconductor package Genomic Ranges was used (Lawrence et al., 2013). Differential gene expression analysis was performed using the Bioconductor package edgeR (Robinson et al., 2010). To assess the differentiation status of csf1r mutant microglia, the differential gene expression list was compared with previously published microglia gene expression profiles obtained in zebrafish and mice (Bennett et al., 2016; Matcovitch-Natan et al., 2016; Oosterhof et al., 2017). We used the Biocoutool Biomat to find high-confidence zebrafish orthologs for the genes from studies performed in mice (Durinck et al., 2009). For gene ontology analysis the Biocoutool package goseq was used (Young et al., 2010).

**Electron microscopy**

Electron microscopy was performed largely as described previously (van Ham et al., 2014). 4 to 5-month-old zebrafish were euthanized in ice water. The brains were carefully removed and incubated in Zamboni’s fixative (4% PFA, 2% glutaraldehyde, 0.2% picric acid, 0.1M cacodylate, pH = 7.4) at 4°C overnight. Brains were washed in cacodylate (3x5 min) and postfixed in a solution containing 1% osmium tetroxide (OsO4) and 1.5% potassium ferrocyanide (K4Fe(CN)6) on ice for 2 hr. The brains were then washed in milliQ water (3x5 min) and dehydrated in series of 30%, 50%, 70% ethanol (10 min each), followed by 3 x 20 min incubation in absolute ethanol. Brains were rinsed in acetone, followed by incubation in a 1:1 EPON:acetone solution overnight. Next, the brains were incubated in pure EPON for 1 hr. New EPON was added followed by incubation in EPON for 2 hr. Samples were put at 200 mbar vacuum for 30 min at 37°C. Polymerization took place at 58°C for 3 days. Ultra-thin sections were subsequently cut, collected on formvar-coated single-slot grids, and stained with a 1% aqueous uranyl acetate solution for 20 min and subsequently for 1 min with lead citrate. Photographs were obtained using a JEOL 1010 electron microscope. For each myelinated axon present the axonal diameter (defined by the inner limit of the myelin sheath) and total fiber diameter (defined by the outer limit of the myelin sheath) were measured using FIJI and the g-ratio was calculated by dividing the axonal diameter by the total fiber diameter (Camargo et al., 2017). Each group consisted of two to three zebrafish, and contained at least 240 myelinated fibers per genotype.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

GraphPad Prism 6 software (La Jolla, CA) was used for statistical tests including one-way ANOVA with Bonferroni multiple testing correction and data are presented as mean ± SD as indicated. For Figures 1 and 5, quantification was done blinded as larval progeny.
were genotyped after quantification, and sample sizes depend on the Mendelian distribution of genotypes. For analysis of adult brains sample sizes (n = 3–4) are similar to those in previous publications (Oosterhof et al., 2017). A p value smaller than 0.05 was considered as significant.

DATA AND SOFTWARE AVAILABILITY

The accession number for the gene level RNA-sequencing data from freshly isolated microglia and brain from control and csf1r mutant zebrafish reported in this paper is GEO: GSE116217.