Regulation of myelination by microglia
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Interactions between microglia, the resident macrophages of the central nervous system (CNS), and myelin, the glial sheath on nerve fibers essential for rapid neural impulse transmission, are commonly studied in the context of neurotrauma and disease. However, interactions between microglia and myelin under normal physiological conditions have been largely overlooked. This review summarizes recent research indicating that the unique properties of microglia evident in disease states also enable microglia to regulate myelination during development and throughout life. This includes phagocytosis of cells and myelin membrane as well as the release of trophic factors, cytokines, and chemokines. The ability of microglia to sense neuronal activity and molecular features of the microenvironment enables them to optimize myelination by influencing early oligodendrocyte differentiation, and removal of aberrantly targeted myelin. Understanding how microglia participate in myelination under normal conditions provides a new perspective that will increase understanding of developmental abnormalities.

INTRODUCTION
Microglia are the resident immune cells of the brain and spinal cord. They survey all regions of the brain including the fiber tracts, called white matter, that connect neurons into circuits. The white appearance derives from a multilayered, highly compacted cell membrane on nerve fibers (axons), called myelin, which is wrapped around axons by non-neuronal cells called oligodendrocytes. Severe dysfunction results when myelin is damaged because the myelin sheath is essential for normal conduction of nerve impulses.

Microglia respond to neural injury and neurodegeneration by undergoing pronounced morphological and cell biological transformations to clear myelin debris and promote repair (1), but in autoimmune disorders, such as multiple sclerosis, microglia can contribute to disease progression by releasing inflammatory cytokines (2) and attacking the myelin sheath (3). The transition from a ramified “resting state” to an amoeboid morphology with high phagocytic activity is classically associated with microglia activation during disease (4), but as early as 1978, Imamoto and Leblond (5) observed that microglia in developing white matter take on an amoeboid cell morphology (Fig. 1). These early findings sparked a question—what functions might these specialized microglia serve in the development of white matter?

Microglia are mechanistically poised to sense and respond to the changing conditions in the developing brain through a plethora of signaling pathways and phagocytic activity, yet we are only beginning to understand the multifaceted functions that microglia have during early myelination events. Efficient and highly organized myelin deposition is necessary for effective brain function, and the temporal and mechanistic features of early microglia uniquely position the population to provide essential regulation of myelination during white matter development. These cells contribute to myelination through the phagocytosis of excess oligodendrocytes and myelin membrane that forms abnormally. In addition, microglia support oligodendrocyte survival and differentiation and regulate apoptosis of oligodendrocytes by secreting several trophic factors and other intercellular signaling molecules, including chemokines and cytokines. Collectively, this review summarizes the rapidly expanding body of evidence that begins to answer the nearly 50-year-old question as to what role microglia may be playing in the formation and organization of myelin during early white matter development.

EMERGENCE OF GLIAL POPULATIONS IN THE DEVELOPING WHITE MATTER
Infiltration, development, and diversification of microglia
Cell lineage tracing reveals that microglia arise from a primitive erythromyeloid progenitor (EMP) population (Fig. 2) present in the yolk sac before embryonic day 8 (E8) in mice and reach the brain by E9.5 (6). Microgliogenesis and brain infiltration are dependent on a number of factors including expression of matrix metalloproteinases and the lineage-specific transcription factors Irf8 and Sfpi1 (7). In mice, these brain-resident macrophages proliferate steadily within the prenatal brain before sharply increasing their population size by approximately 10-fold during the first 2 weeks of postnatal development (8). It is widely accepted that the vast majority of adult microglia originate from EMP, but there is evidence of a perinatal wave of infiltrating peripheral monocytes that may contribute to the rapid expansion of the postnatal microglia population (9). Early primative macrophage colonization is also observed in human development with cells expressing Iba1 (a calcium binding protein commonly used as a microglia marker) penetrating the cerebral parenchyma.
wall at 4.5 weeks of gestation, followed by notable periods of prenatal proliferation (10).

Although microglia share a common origin, there is substantial heterogeneity within the population. Microglia undergo coordinated transcriptional changes throughout neurodevelopment, and microglia found in the postnatal corpus callosum have distinct gene expression profiles that differ from adult microglia (11), as well as from microglia in other regions of the postnatal brain (12). Recent work by Hagemeyer et al. (13) identifies a population of microglia in the early postnatal corpus callosum and white matter tracts of cerebellum that display markers of microglial activation such as Mac3, and an up-regulation of genes related to activated microglia including Igf1 and ligax (also known as Cd11c). The studies show that these cells arise from central nervous system (CNS) endogenous precursors and have no contribution from circulating blood monocytes. The cells retain an amoeboid morphology transiently from postnatal day 1 (P1) through P9 to P10 in mice. These amoeboid microglia then transition toward a ramified state later in development (14). In contrast, microglia outside white matter tracts quickly take on a ramified morphology after birth and do not show signs of activation (13).

Microglia arise from the yolk sac EMP population. After EMP-derived cells are incorporated into the CNS around E9.5 and take on microglia-specific profile, microglia of the early white matter begin to exhibit an amoeboid morphology with retracted processes. This is similar to the morphology of “activated” microglia that respond to injury and disease. However, in normally developing white matter tracts, amoeboid microglia appear shortly after birth (observed as early as P1 in mice) (14) and remain prevalent in the postnatal brain until transitioning toward a ramified state later in development (approximately P14 in mice) (43). Under inflammatory conditions, monocytes from the bloodstream can also infiltrate the brain.

Separately, Wlodarczyk and colleagues (15) detect a subpopulation of microglia localized specifically to the corpus callosum and cerebellum white matter tracts of healthy mouse brains from P2 to P7, which expresses the integrin complement receptor CD11c. CD11c is a microglia marker rarely observed in the healthy adult CNS, but it is up-regulated in a TREM2-dependent manner in disease-associated microglia populations that arise during neuroinflammatory and neurodegenerative disease models such as the experimental autoimmune encephalomyelitis (16), aging, Alzheimer’s, and amyotrophic lateral sclerosis (17). Such disease-associated microglia have been shown to have a protective function by clearing degenerating neurons, myelin debris, and pathological tau accumulations (1,18,19). The discovery of this subpopulation of activated microglia within white matter regions during early postnatal development, coinciding with a period of extensive oligodendrogenesis and the initiation of myelination (20), has sparked in-depth investigation of the function of microglia in the development of white matter myelin.

**Oligodendrogenesis and myelination**

CNS myelination is an essential developmental process characterized by the production and organization of myelin membrane into multilayered ensheathments along axons (Fig. 3). The dense lipid membrane layers act as an electrical insulator for neurons, promoting rapid saltatory conduction along axons by reducing transmembrane current leakage and electrical capacitance (21) and constraining sodium channels to the gaps between myelin segments known as nodes of Ranvier (22). In addition to increasing the conduction velocity of neuronal signal transmission, myelin also provides metabolic support to neurons through lactate transport (23, 24). Furthermore, oligodendrogenesis and proper myelination are essential to the complex cognitive functions of a healthy brain. Mice with irregular myelination induced by oligodendrocyte-specific knockdown of cyclin-dependent kinase 5 (Cdk5) have significantly impaired long-term memory consolidation, fear-conditioned learning, and motor skill learning (25). Even in adults, when inhibition of de novo oligodendrogenesis is disrupted with no impairment of pre-existing myelin, mice struggle to learn complex motor tasks (26) and to recall spatial and fear memories (27).

Oligodendrocytes differentiate from a population of cells known as oligodendrocyte progenitor cells (OPCs) that initially populate the brain in three distinct waves beginning during embryogenesis (28). The first and second waves of OPCs (occurring at approximately E11.5 and E15 in mice) divide asymmetrically from neural progenitor cells (NPCs) in the telencephalon ventricular zones and...
migrate outward to disperse evenly throughout the developing cortex; the final wave of OPCs arises from within the cortex near the time of birth (28). As oligodendrocytes mature, each cell can extend 20 to 60 myelinating processes that contact individual axons and wrap multiple compacted layers of membrane around them (29). Although myelination in the CNS begins in late fetal development and is most active in the early postnatal period, it is a slow process that is not completed in some CNS regions until early adulthood. In adulthood, myelin remains plastic (30), and myelin in the cerebral cortex begins to decrease in later life (31).

The myelinating processes of a mature oligodendrocyte are guided by physical (32) and biochemical signals toward axons, selectively myelinating in a neuronal subtype–specific manner (33). Not all nerve axons are myelinated, and the extent of myelination varies greatly, even along an individual axon, within a neuronal subtype, or region of the brain (34). Targeting of axons for myelination is directed, in part, by electrical activity in axons. The activity-dependent release of the neurotransmitter glutamate promotes the formation of an axoglial signaling complex and stimulates local synthesis of myelin basic protein (MBP) in the adjacent oligodendrocyte cell process (35) to initiate myelination preferentially on electrically active axons (36). Once an axon has been targeted, the oligodendrocyte process wraps around the axon, leading with an inner tongue of myelin membrane that extends underneath previously deposited layers of membrane. Simultaneously, the membrane must expand laterally to define the length of the sheath, while membrane compaction and interlamellar proteins help stabilize external layers of myelin (37). The highly specific patterning of myelin deposition is believed to be integral to long-distance neuronal communication and the synchronization of spike-time arrivals across varied synaptic inputs (38, 39). The complex structural organization of myelin is not easily coordinated and is prone to errors. Recent studies have documented a number of ultrastructural myelin abnormalities, such as outfoldings, bulging, fragmentation, and splitting, that arise during early myelination but are rapidly resolved as development progresses. This research indicates that microglia are critical in removing abnormal myelination (40).

Myelination is a complex biological event, which is highly dependent on intercellular interactions. Oligodendrocytes must recognize the appropriate cells to myelinate and the correct cellular region to myelinate (only axons, rather than dendrites or neuronal cell bodies). Myelination requires specialized synthesis of enormous amounts of membrane and complex cellular motility and dynamics to wrap membrane around axons and exclude the cytoplasm from each concentric membrane layer. Microglia are particularly well suited to guide the process of oligodendrogenesis, modulate myelin organization, and correct errors in myelin formation. In the following sections, we will explore the functional importance of these mechanisms of microglia activity in the development of healthy white matter.

REGULATION OF OLIGODENDROCYTE-LINEAGE CELL POPULATIONS

Oligodendrocyte apoptosis and clearance by microglia

Phagocytosis by microglia is an important mechanism for the removal of cellular debris following injury, and recent studies are finding that microglia perform a similar function during development. Although we commonly think of development as a time of cell proliferation, a marked degree of cell death is observed in white matter tracts of the developing CNS. In the rat optic nerve, there is a steep increase in oligodendrogenesis after birth; however, this is closely followed by a spike in apoptosis of oligodendrocytes and late-stage OPCs (41). Similarly, as much as 20% of the immature oligodendrocyte population in the cerebral cortex shows signs of fragmentation and degeneration in young rats (42). Single-cell deep sequencing has identified a transient early postnatal subset of amoeboid microglia in the mouse corpus callosum and cerebellar white matter with a gene expression profile that is notably similar to that of microglia found in adult neurodegenerative disease states (43). These microglia are observed in the white matter from P4 to P14 in mice, and they demonstrate high phagocytic activity as compared to microglia from other brain regions (43). Microglia from early postnatal white matter contain fragments of the apoptotic marker cleaved caspase 3 (CC3) colocalized with MBP, a marker of mature oligodendrocytes (43). These results indicate that an early postnatal population of amoeboid microglia contribute to white matter development by actively phagocytosing dying oligodendrocyte-lineage cells. Li et al. (43) also observe amoeboid microglia in early white matter associated with nonapoptotic oligodendrocytes.

Phagocytosis of living oligodendrocyte-lineage cells

Microglia can also regulate the oligodendrocyte-lineage cell population through phagocytosis of living cells. Amoeboid microglia in the corpus callosum during early postnatal development have recently been shown to engulf NG2+ OPCs over a narrow window between P4 and P11, peaking at P7, in mice (Fig. 4) (44). Intriguingly, only 10 to 15% of the NG2+ OPC cells phagocytosed by these microglia express apoptotic markers, while the remaining 85 to 90% of phagocytosed OPCs are not apoptotic (44). Microglia are therefore not simply responding to apoptotic signals and removing the debris but rather actively modulating cell population dynamics in the developing white matter.

While the purpose of the overproduction of oligodendrocyte cells in developing white matter has yet to be determined conclusively, each of the three distinct waves of migrating OPCs identified

Fig. 4. Microglia phagocytose live OPCs during postnatal white matter development. Time-lapse confocal imaging captures a CX3CR1+ amoeboid microglia (green) phagocytosing a live NG2+ OPC (red) in the corpus callosum of a P7 mouse brain. Reprinted from Nemes-Baran et al. (44) under Creative Commons licensing.
white matter tracts of mice have also been observed with myelin breakdown by oligodendrocytes do contribute to the removal of some myelin sheaths, in 2020, Hughes and Appel (49) reported that microglia assist in the removal of aberrantly deposited myelin, leading the authors to hypothesize that these retraction events may be evidence of corrective action following aberrant myelin targeting. While endogenous retraction and breakdown of myelin by oligodendrocytes do contribute to the removal of some myelin sheaths, in 2020, Hughes and Appel (50) reported that microglia assist in the removal of aberrantly deposited myelin. Furthermore, these studies found that myelin sheaths that were phagocytosed were longer than sheaths that were removed through endogenous retraction, leading the authors to hypothesize that phagocytic pruning functions extend the window for myelin refinement by allowing removal of segments that have already begun to bind to the axon (50). Microglia that populate developing white matter tracts of mice have also been observed with myelin phagocytosis of live OPCs, while others may differentiate into mature oligodendrocytes, only to find that they lack necessary resources, undergo apoptosis, and eventually become targets of phagocytosis.

The need for such cellular elimination becomes evident when investigating the potential effects of an overcrowded oligodendrocyte-lineage population on overall myelin health. When microglial phagocytosis of live OPCs is inhibited experimentally in mice, the OPC population is increased, and there is a decrease in myelin thickness in studies on the corpus callosum. Inhibition of microglial phagocytosis was accomplished by knocking out the fractalkine receptor CX3CR1, an important receptor for microglial recognition of OPCs (44). This suggests an optimal number of OPCs for optimal myelin ensheathment of axons. Another study finds that pharmacological stimulation of excessive oligodendrogenesis increases the incidence of ectopic myelination, in which myelin sheaths were incorrectly targeted to wrap neuronal cell bodies (46). It is hypothesized that overcrowding of oligodendrocyte-lineage cells may lead to an imbalance in the ratio of oligodendrocytes to axons, resulting in aberrant myelin targeting and perhaps a reduction in the health of oligodendrocyte-lineage populations as competition over trophic resources increases. These findings support the conclusion that microglia help ensure correct myelin targeting by carefully maintaining a balanced oligodendrocyte-lineage cell population.

PHAGOCYTOSIS OF MYELIN MEMBRANE

In addition to engulfing whole cells, microglia have been observed to specifically phagocytose myelin sheaths. Several groups have documented the elimination of nascent myelin deposits during development. Live imaging of myelination initiation in the zebrafish spinal cord indicates that most of the myelin sheaths are rapidly deposited over a narrow period during early development; however, several of the initial sheaths are later retracted in a slower process (47). Similar studies report that early myelin retraction events occurred more frequently on smaller caliber axons (48) and axons with inhibited vesicular release (49). These early studies hypothesize that these retraction events may be evidence of corrective action following aberrant myelin targeting. While endogenous retraction and breakdown of myelin by oligodendrocytes do contribute to the removal of some myelin sheaths, in 2020, Hughes and Appel (50) reported that microglia assist in the removal of aberrantly deposited myelin. Furthermore, these studies found that myelin sheaths that were phagocytosed were longer than sheaths that were removed through endogenous retraction, leading the authors to hypothesize that phagocytic pruning functions extend the window for myelin refinement by allowing removal of segments that have already begun to bind to the axon (50). Microglia that populate developing white matter tracts of mice have also been observed with myelin MBP inclusions (40). Abnormalities in the ultrastructure of myelin membrane have been found in the optic nerve of healthy P10 mice, but these aberrations are resolved by adulthood, indicating that occasional myelin structural errors are likely a normal part of development (40). Both studies used methods of microglial inhibition and ablation to investigate the impact on myelination and found a marked increase in the prevalence of abnormal myelin structures and ectopic myelination events as compared to wild-type controls (40, 50). Together, these recent findings demonstrate a role for phagocytic microglia in refining and remodeling myelin structures during development.

MICROGLIA INTERACT WITH OLIGODENDROCYTES THROUGH RELEASE OF BIOCHEMICAL SIGNALING MOLECULES

Microglia do not act on the early CNS exclusively through their phagocytic function; they also participate in complicated intercellular signaling networks through the production and secretion of biochemical factors. Some molecules released by microglia can be inflammatory or cytotoxic to oligodendrocytes, while other factors are essential to oligodendrocyte survival, differentiation, and successful myelination.

Microglial trophic factors support oligodendrocyte-lineage cell survival, differentiation, and myelination

We have discussed the importance of microglia clearance of some oligodendrocytes to maintain population balance, given the myelination requirements and resources available in a specific region; however, microglia also act to fortify the remaining oligodendrocyte population. Oligodendrocyte-lineage cells die rapidly when cultured in isolation, but apoptosis is markedly reduced by the addition of growth factors, suggesting the need for intercellular support for sustained survival of oligodendrocyte-lineage cells (41). A plethora of factors have been shown to support healthy oligodendrocyte development and maturation, many of which cannot be produced by oligodendroglia but rather must be provided by other cells in the surrounding environment. Microglia-released soluble factors collected in conditioned media have been shown to support OPC survival and oligodendrocyte differentiation (51), increase the expression of the myelin markers MBP and proteolipid protein (PLP), and promote in vitro myelination (52). While astrocyte-conditioned media also support oligodendrocyte survival, microglia-released factors are particularly beneficial for oligodendrocyte differentiation and myelination (51).

One microglial factor that has received attention for its role in supporting oligodendrocyte health is insulin-like growth factor 1 (IGF-1). IGF-1 is found at high levels in microglia-conditioned media (51) and significantly reduces oligodendrocyte apoptosis in culture (41). IGF-1 has been localized to amoeboid microglia within the developing rat corpus callosum (53), and CD11c+ microglia from the corpus callosum produce IGF-1 at a level sevenfold greater than CD11c+ microglia found elsewhere in the postnatal brain (15). Mice lacking IGF-1 have reduced OPC and oligodendrocyte populations as well as reduced myelination as compared to controls (54). Notably, the same population of microglia that display highly activated phagocytic activity in developing white matter also show elevated expression of IGF-1 (43). Specific knockout of IGF-1 from CD11c+ microglia results in irregular myelination, characterized by decreased Mbp, Plp, Mag, and Mog gene expression, and thinner myelin sheaths in the
corpus callosum (15). Therefore, early white matter microglia that act to remove unhealthy and extraneous oligodendrocytes simultaneously contribute to the health of surviving oligodendrocyte-lineage population by providing trophic support. It is not yet clear from the available literature whether these dichotomous actions are taken on by separate subsets of the microglia or whether individual cells actively switch their activities based on localized needs.

Other factors produced by microglia have been found to support the health and maturation of OPCs and oligodendrocytes in the developing CNS. Microglia-derived IGF-2 has been shown to prevent tumor necrosis factor–α (TNF–α)-induced apoptosis of oligodendrocytes in vitro (55). In addition, IGF-2 was up-regulated following IGF-1 knockout, perhaps as compensation in the absence of the usual important trophic factor. Notably, however, this study did not investigate the source of IGF-2; therefore, any inhibitory benefits for oligodendrogenesis in this experiment cannot be conclusively attributed to microglia (54). Furthermore, inhibition of microglial activation by minocycline treatment impairs maturation of oligodendrocytes in vivo, by reducing levels of the microglial cytokines interleukin-β (IL-1β) and IL-6 (56). A pathway dependent on the interaction between microglia-derived transglutaminase-2 (TG2) and the extracellular matrix protein laminin has been found to support OPC proliferation in the corpus callosum (57). In addition, neuropilin-1 (NP1), a transmembrane molecule expressed by activated microglia, has been recently shown to promote OPC proliferation in response to platelet-derived growth factor AA (PDGF AA) through its activation of the PDGFRα receptor (58). Collectively, these findings illustrate the complexity of microglial signals promoting the development of healthy oligodendrocyte populations and proper myelin formation.

Microglial signals involved in induction of apoptosis of mature oligodendrocytes

Disease-state microglia are well known cytotoxic actors in the CNS, capable of inducing cell death through a myriad of inflammatory cytokines, proteinases, and reactive oxygen intermediates, and microglia may use similar cytotoxic factors in regulation of oligodendrocyte populations in development. Microglia in the early postnatal white matter are transcriptionally similar to microglia found in disease states (43). These microglia have elevated proinflammatory cytokine and chemokine gene transcripts, including elevated Mif, Ccl3, Ccl4, CCL6, and Ccl9, as compared to other early microglia populations (43), suggesting that some of the mechanisms of microglial cytotoxicity may be used by microglia under physiologically normal developmental conditions. Although microglia can contribute to pathology, they also have known protective functions in neurodegenerative disease states (17, 19). Testing these hypotheses is an important direction for future research.

MULTIDIMENSIONAL SURVEILLANCE OF THE NEUROENVIRONMENT

The complex microglia-mediated intercellular interactions taking place during development are not entirely preprogrammed processes; rather, microglia acutely sense the need for specific mechanistic actions through a multitude of receptors and signaling molecules. These signals can be produced specifically by an individual cell to target microglia action, or these signals can be broader indicators of cellular population dynamics in the tissue and network functionality in developing neural circuits. This versatility allows microglia to sense and respond to a diverse array of needs and to efficiently provide support or regulation where required.

Sensing the health of cellular networks

The rapid motility of macrophage cells allows microglia to scour large portions of the CNS for signals indicative of the health and density of regional cell populations. Because microglia are responsible for maintaining balanced oligodendrogenesis while simultaneously regulating against overproliferation and crowding, they must have a mechanism to sense oligodendrocyte population dynamics. Chemoattractants are soluble signals that do not target specific cells for removal but serve to indicate need for microglia regulation. Following injury or insult, oligodendrocytes are known to release chemoattractant molecules, including CXCL10, CCL2, CCL3, and CCL5, which recruit microglia to the site of disruption and initiate cell removal (59). Inversely, oligodendrocytes experiencing cellular stress can also produce β-crystallin, which attracts microglia that, in turn, act to promote improved health of the oligodendrocytes (60). While it is evident that microglia are capable of sensing oligodendrocyte health in response to disease, it is unknown whether similar recruitment molecules are released in response to nonpathological cellular stressors that can occur during development, such as overcrowding or mild resource scarcity due to competition. Furthermore, once microglia are alerted to a regional need for action, it would be important to determine how they target specific cells for elimination while selecting others for survival.

Signals driving cell-specific microglia action

Phagocytosis of apoptotic and degrading cells is a common function of microglia throughout life. Although signals uniquely targeting dying oligodendrocytes for phagocytosis have not been identified, phosphatidylserine translocation to the cell surface is a common apoptotic event across many cell types and functions as a phagocytosis signal for microglia (61). In the brain, externalized phosphatidylserine interacts with soluble TAM receptor ligands such as GAS6 and protein S to initiate microglial phagocytosis through the TAM receptor tyrosine kinases Mertk and Axl (62). This is one of the same signals that mark synapses for pruning during development (63), and abnormal myelin structures appearing during postnatal myelination also express phosphatidylserine on their surface to attract phagocytic microglia (40).

The accumulation of evidence supporting microglia phagocytosis of live oligodendrocyte-lineage cells and myelin has led investigators to question what signals might be directing microglia to phagocyte a specific target as compared to a neighboring cell or myelin sheath segment. The fractalkine ligand CX3CL1 is expressed by developing OPCs, and fractalkine receptor CX3CR1-deficient microglia are less efficient at OPC phagocytosis despite having an equivalent number of microglia and OPC-microglia contact events when compared with wild-type mice (44). This finding suggests that the CX3CL1-CX3CR1 signaling pathway is involved in targeting live OPCs for phagocytosis; however, the mechanisms driving OPC expression of CX3CL1 have yet to be elucidated.

Sensing neuronal activity changes and regulation through myelin modification

Microglia are capable of sensing neuronal activity both at the synapse (64) and along axons, allowing these cells to gather information
about the neural circuit functionality at a broad and local level, and to control the level of neuronal activity through a negative feedback mechanism (65). A number of neurotransmitters and purinergic molecules act either directly or indirectly to initiate microglial responses to changes in neuronal activity (66). Neuron-derived CX3CL1 is an essential regulatory signal that interacts with the microglial receptor CX3CR1 to induce microglia-mediated synapse remodeling in response to changes in neuronal activity (67). Signaling through interaction between local axonal release of potassium and the microglia receptor THIK-1 also promotes microglia-neuron interaction. In experiments by Ronzano et al. (68), neuronal activity is reported to stabilize microglia activity and hold microglia at the node of Ranvier, while inhibition of neuronal activity or microglia-nodal interactions increases dynamic activity of microglia. As myelination alters the conduction velocity, synchronization, and overall functionality of neural networks, microglia modulation of myelination may be an indirect mechanism through which microglia can sense and respond to changes in neuronal activity.

Myelin targeting and thickness is influenced by neuronal activity such that active axons are preferentially myelinated over inactive axons (36) and increased neuronal activity stimulates thicker myelin (69). Recent findings open the possibility that selective myelination of active neurons may be refined after initial myelination deposition through a microglia-mediated mechanism. Neuronal activity may bias microglia selection of myelin segments for phagocytosis, as phagocytosis in zebrafish studies is elevated when neuronal activity is reduced in neighboring axons, while myelin elimination is reduced when regional neuronal activation is increased (50). This recent study examined the effects of shifts in neuronal activity across an entire neuroanatomical structure (optic tectum) however, microglia are also capable of sensing changes in activity at the level of individual axons and nodes. It remains unclear how such highly localized shifts in activity

### Table 1. Effects of microglia ablation or inhibition on oligodendrogenesis and myelination

| Method(s) for microglia ablation/inhibition | Organism and ROI | Effects on oligodendrogenesis | Effects on myelination | Reference |
|------------------------------------------|-----------------|-------------------------------|------------------------|-----------|
| Csf1r<sup>−/−</sup> transgenic | P20 mouse CC and SVZ | High degree of OPC apoptosis; mature OL population reduced | No noted effects on myelination | (73) |
| Intrapertoneal administration of minocycline to prevent microglia activation | P5 rat SVZ | Increased OPC population but reduced premyelinating and mature OL numbers; no difference in OL-lineage proliferation labeling | Number of MBP<sup>+</sup> cells decreased | (56) |
| CSF1R inhibitor BLZ945 | P20 and P40–P42 mouse WM | Reduced number of OLs in P20 CC | P20: Reduced expression and labeling of myelin proteins in CC and cerebellum; decrease in % of myelinated axons; no difference in myelination thickness; P40–P42: no difference in PLP or MBP labeling | (13) |
| Transgenic deletion of lglf-1 gene in CD11c<sup>+</sup> microglia | P21 mouse CC | No noted effects on oligodendrogenesis | Decreased expression on Plp, Mag, and Mog; reduced myelin staining; thinner myelin sheaths | (15) |
| Csf1r<sup>−/−</sup> transgenic | 3-week rat CNS | Decrease of oligodendrocyte gene expression | Reduced myelination in the CC; decrease of myelin-associated gene expression | (72) |
| Pediatric-onset leukoencephalopathy patients (homozygous Csf1r mutations) | Human brain tissue (10 months to 25 years) | No noted effects on oligodendrogenesis | Periventricular white matter abnormalities; cerebellar and CC hypoplasia/atrophy | (71) |
| PLX5622 and PLX3397 (CSF1R inhibitors) | 8- to 10-week mouse CNS | Reduced OPC viability in vitro and ex vivo; regional reduction of OPCs in vivo | No difference in mature OL populations or PLP labeling | (74) |
| (i) Blockade of Irf8 translation (ii) CSF1R inhibitor GW2580 (iii) Microglia expression of nitroreductase + actuator | 4 DPF zebrafish optic tectum and SC | No difference in number or distribution of OLs | Oligodendrocytes produced more sheaths and sheaths were shorter in length; increased incidence of ectopic myelination | (50) |
| Microglia ablation via Csf1r double mutant | 10 DPF zebrafish SC | Reduced number of OLs | Reduced total myelinated area, increased number of myelin structure pathologies, and no difference in sheath length | (40) |

OL, oligodendrocyte; SC, spinal cord; SVZ, subventricular zone; WM, white matter.
might alter microglia-myelin interactions. Such investigation is pertinent to our understanding of development because neuronal activity rarely undergoes marked changes under physiological conditions; rather, small local shifts often accumulate over time to drive the complex higher-order processes such as learning and memory.

**WHITE MATTER WITHOUT MICROGLIA**

To further investigate the influence of microglia on developing white matter, many groups have used methods of pharmacological inhibition and transgenic knockdown of microglia in animal models. Colony-stimulating factor 1 receptor (CSF1R) is essential to microglia development and survival (70), and therefore, manipulation of this receptor is a popular method of reducing the microglia population. Mutation of the CSF1R gene has been observed in a clinical setting with similar detrimental effects on human microglia populations; white matter abnormalities were observed along with a history of seizure and developmental delays (71). Experimental methods exploring the impact of microglia on oligodendrocyte populations and myelin formation are summarized in Table 1.

Collectively, the results of these experiments indicate that microglia are significant regulators of oligodendrogenesis and myelination, but the results differ depending on the experimental approach. Multiple studies show changes in oligodendrogenesis following microglia ablation with abnormalities in myelination of specific regions of interest (13, 40, 56, 72). Other studies report myelination deficits in the absence of an effect on oligodendrocyte cell numbers (15, 50, 71). In contrast, some studies find no myelination deficits but do report that oligodendrogenesis is impaired (73, 74). The resilience of the microglia-oligodendrocyte relationship across a multiplicity of models can be expected given the importance of myelination to essential brain functions. For example, despite a decrease in oligodendrocyte density and myelin protein in microglia-deficient P20 corpus callosum, myelin protein levels are restored at a later time in development (13). Furthermore, microglia populations can recover from ablation rapidly (70), making the timing of experimental manipulation and tissue collection potential sources of variability. Many of these studies rely on global ablation of microglia; thus, indirect effects of ablating microglia beyond the white matter cannot be ruled out. Results differ, depending on the experimental approach, but as a whole, these experiments indicate that microglia are dynamic contributors to multiple processes involved in the formation and organization of healthy white matter.

**SUMMARY AND CONCLUSION**

Microglia have been portrayed historically as passive surveyors of the healthy CNS, simply cleaning up cellular debris, unless activated by a pathogen or disease; however, microglia in the healthy early postnatal brain sense the surrounding microenvironment proactively, detecting neuronal activity levels and oligodendrocyte population dynamics to determine the need for modulation or stabilization of oligodendrogenesis and myelination. Recent experiments indicate that when microglia detect that oligodendrocyte population dynamics are unsustainable, or when aberrant myelin structures or changes in axons are sensed, microglia selectively target oligodendrocyte-lineage cells and myelin structures for removal. Alternatively, if microglia sense a need for increased or homeostatic maintenance of myelination of active axons, microglia play a supportive role by fortifying oligodendrocyte-lineage populations and promoting oligodendrogenesis and myelin formation (Fig. 5). In summary, studies examining the impact of microglia deficiency on oligodendrogenesis highlight the importance of microglia in proper myelin formation for normal brain function.

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**Fig. 5. Microglia actively regulate the oligodendrogenesis and myelination of the developing white matter.** Microglia sense and act on the early white matter by (1) clearing cellular debris from apoptotic oligodendrocytes through phosphatidylserine signaling, (2) phagocytosing live OPCs to regulate oligodendrocyte population dynamics, (3) removing abnormal myelin ultrastructure, (4) eliminating aberrantly targeted myelin, (5) promoting oligodendrogenesis, (6) fortifying mature oligodendrocytes and promoting myelination, and (7) sensing changes in neuronal activity to determine need for modification of myelin organization.
Currently, there is a rapidly expanding body of work investigating the microglia-oligodendrocyte relationship in development, but there is much that remains to be understood. Among these questions, it will be important to determine whether specialized types of microglia exist in white matter to influence myelination, or instead plasticity of microglia enables the same cells to carry out different functions as required. Notably, the extent to which white matter microglia are distinct from microglia in gray matter or differ from those that carry out immunological responses requires further investigation. The possible involvement of microglia in maintenance, remodeling, plasticity, and de novo myelination in the healthy adult brain, as distinct from during development, is a neglected but promising area of investigation. Cell-cell and intracellular signaling pathways in microglia and oligodendrocytes need to be identified and interpreted in the context of interactions among other brain cells, including neurons and astrocytes, to influence oligodendroglia and myelination. The mechanisms by which microglia are targeted to act on appropriate axons, oligodendrocytes, and subcellular domains to modify myelin must be further identified.

The growing awareness that microglia are critical in myelination in the healthy brain illuminates how disease and other nervous system insults that activate microglia during critical periods could impair myelination with long-term consequences. There is a large body of research indicating that viral infection during pregnancy, for example, can increase the probability of offspring developing cognitive impairments or mental illnesses, including schizophrenia, in later life (75). Other work confirms white matter abnormalities in such conditions (76). Similarly, substance abuse, including alcohol consumption during fetal development (77), and hypoxia/ischemia at birth result in white matter deficits (45), which could, in part, reflect damage to microglia in regulating myelination when the cells become activated in an inflammatory state. There are currently no treatments for such white matter injuries from such insults in the perinatal period, but as knowledge of microglia-oligodendrocytes in the healthy brain increases, new understanding and new treatments may be forthcoming.

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