**Microglia states and nomenclature: a field at its crossroads**

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**Abstract** **Word limit: 150**

Microglial research has advanced considerably in recent decades yet has been constrained by a rolling series of dichotomies such as “resting *versus* activated” and “M1 *versus* M2”. This dualistic classification of good or bad microglia is inconsistent with the wide repertoire of microglial states and functions in development, plasticity, aging and diseases that were elucidated in recent years. New designations continuously arising in an attempt to describe the different microglial states, notably defined using transcriptomics and proteomics, may easily lead to a misleading, although unintentional, coupling of categories and functions. To address these issues, we assembled a group of multidisciplinary experts to discuss our current understanding of microglial states as a dynamic concept and the importance of addressing microglial function. Here, we provide a conceptual framework and recommendations on the use of microglial nomenclature for researchers, reviewers, and editors, which will serve as the foundations for a future white paper.

**Abbreviations**

AD – Alzheimer’s disease

ARM – activated response microglia

ATM – axon tract-associated microglia

BAM – border-associated macrophage

BBB – Blood-brain barrier

CAM – CNS-associated macrophages

CNS – central nervous system

CSF – cerebrospinal fluid

CSF1R – colony stimulating factor 1 receptor

DAM – disease-associated microglia

HAM – human AD microglia

iPSC – induced pluripotent stem cells

IRM – interferon-responsive microglia

ISF – interstitial fluid

LDAM – lipid-droplet-accumulating microglia in aging mice and humans

MGnD – microglial neurodegenerative phenotype

MIMS – microglia inflamed in multiple sclerosis

MS – multiple sclerosis

PAM – proliferative-region-associated microglia

ROS – reactive oxygen species

scRNASeq – single-cell RNA sequencing

WAM – white matter-associated microglia

**Names, names, names**

*"If the names are unknown knowledge of the things also perishes."1*

*(Carolus Linnaeus)*

And yet, we humans instinctively tend to name things and use that name to define their properties. Biologists are no exception: from the time of 18th century father of taxonomy Carolus Linnaeus, the main purpose of biology has been categorizing the natural world as a way of understanding it. Naming species and grouping them together into taxa served to define evolutionary relationships; even today taxonomy and phylogeny are closely interrelated. But we must never forget that nomenclatures and categories are artificial constructs and biology is seldom black and white, but rather an extended continuum of greys. While giving names is natural and useful, we need to be aware that categorization constrains our thinking by forcing us to fit our observations into established classes. As sociologists say, “categorization spawns expectations”2. This semantic issue has already been acknowledged by immunologists because, in fact, the given names have connotations that often imply a specific function3. In this paper, we extend similar initiatives on macrophages4, dendritic cells3, interneurons5, and astrocytes6 to discuss the widespread problems associated with categorization of microglia using outdated terms such as “resting *versus* activated” (**Box 1**) or “M1 *versus* M2” (**Box 2**).

Dichotomic, rigid categories convey a dualistic idea of good *versus* bad microglia and may actually impede scientific advancement. Widely used terms, such as “neuroinflammation” as a synonym of microglial reactivity (**Box 3**) and naming a panoply of presumed microglial populations and assumed functions arising from single-cell transcriptomics, are misleading and increasingly problematic, especially to those entering the field of glial biology and neuroimmunology. This nomenclature does not address the important question: what are the specific functions of microglia in the contexts of development, health, aging, and disease? It is now clear that microglia exist in diverse, dynamic, and multi-dimensional states depending on the context including local environment (**Figure 1**). We define dimensions as the key variables driving the phenotypic transformations of microglia. These variables are molecularly distinct signaling pathways regulated at multiple levels (e.g., transcriptional, epigenetic, translational, metabolic) that each give rise to distinct microglial functions or properties. In this manner, categorizing microglia based on a historical, one-dimensional nomenclature in the absence of functional data will constrain and stifle future progress and innovation.

To examine and address these issues, we assembled a team of international experts who have made major contributions to microglia research, inclusive of various groups, and balancing gender, geographical distribution, and seniority. Authors from the fields of neuroscience, neurobiology, immunology, neuroimmunology, oncology, and neuropathology, both from academia and industry, discussed their perspectives on the current and future challenges in defining microglial states and nomenclature. A questionnaire (**Supplementary Data**) was created to collect all the authors’ opinions on several nomenclature issues and the importance of directly addressing microglial function. The responses to the questionnaire, an online meeting held in June 2021 and an open session held at the EMBO meeting Microglia 2021 were used as a backbone to develop this paper.

Herein, we summarize our current knowledge about the identity of microglia and discuss best practices for how to define and study microglial state dynamics. We then outline “classical” microglial nomenclatures, highlighting some of the key discoveries that led to the above classifications and their limitations. We intentionally focus on citing studies related to the nomenclature, rather than providing a comprehensive review of the history of microglial research, as it has been done elsewhere7,8. We discuss the overall limitations and conclude with recommendations for the proper usage of microglial nomenclature as research evolves, provide a conceptual framework for discussing microglia, and offer perspectives on the future questions, gaps in knowledge, and challenges to tackle as a field.

**Microglial identity: what we mean about when we talk about microglia**

The origin and identity of microglia was for many years a matter of debate. In the dim and distant past, Ramón y Cajal´s disciple, Pío del Río-Hortega suggested that these cells were of mesodermal origin9. However, over time, an ectodermal origin was also proposed10, sparking controversy until the 1980s. The mesodermal origin took solid hold later with the advance of technical approaches revealing more similarities than differences with the functions and features of macrophages. In 1999, microglia were reported to appear in the brain rudiment as early as embryonic day E8 in mice, and proposed to originate from yolk sac progenitors11. The recent combination of fate mapping studies and transplantation approaches this debate, revealing key aspects of microglial identity and plasticity. In mice, unlike other model organisms such as zebrafish12,13, microglia are now considered to originate from a pool of macrophages produced during primitive hematopoiesis in the yolk sac, which start invading the neuroepithelium at E8.514-17. In humans, microglial precursors invade the brain primordium around 4.5 to 5.5 gestational weeks18.

One key signaling pathway critical for microglial development and maintenance is the CSF1R (colony stimulating factor receptor). Ligands of CSF1R that sustain this pathway include two cytokines with different origins and primary sequences, but similar tridimensional structures and binding to CSF1R: IL-34 and CSF119. IL34 is produced by neurons, while CSF1 is secreted primarily by oligodendrocytes and astrocytes. Accordingly, the two ligands have distinct and non-overlapping functions in the establishment and maintenance of microglia within the grey and white matter20. Microglia have the capacity for self-renewal in certain contexts, allowing them to repopulate the central nervous system (CNS) within one week of depletion, even when more than 99% of microglia are ablated with CSF1R antagonists21,22 or diphtheria toxin22. This process, termed “microglial repopulation” or “microglial self-renewal”23-25 is different from “microglia replacement” which, in contrast, occurs when endogenous microglia are replaced by exogenous cells that can include bone marrow-derived myeloid cells26-29, peripheral blood cells28,30, stem cell- or iPSC-derived peripheral blood cells31, across various experimental or pathological conditions31-33.

Our current definition is that mammalian microglia are yolk sac-derived, long-lived cells within the CNS parenchyma that persist into adulthood, and self-renew without any contribution from bone marrow-derived cells at steady-state.

The identification of microglia is currently based on the expression of specific genes highly enriched in microglia, which represent their transcriptional identity and are commonly employed as “microglial markers” (**Table 1. Microglial markers**). However, the expression of each marker alone is not sufficient to define microglial identity, as levels of expression may change depending on microglial adaptation to local signals. The present consensus is that mammalian microglia can be identified by the expression of transcription factors like Pu.116, cytoplasmic markers such as ionized calcium-binding adapter molecule 1 (IBA1), and surface markers including the purinergic receptor P2YR12, transmembrane protein 119 (TMEM119), and CSF1R34. Based on these markers, genetic tools (such as Cx3cr1CreERT2, P2ry12CreERT2,Tmem119CreERT2 and HexbCreERT2 mouse lines) are available that allow for more specific manipulation or visualization of microglia, although they could also target other populations, including border-associated macrophages (BAMs), also named CNS-associated macrophages (CAMs) and other glial cells35-40. Most recently, a new binary transgenic model relying on co-expression of Sall1 and Cx3cr1 has been introduced that specifically targets microglia in a non-inducible way41.

Nonetheless, many of these markers are downregulated in pathological states, and can be expressed by other brain macrophage populations such as BAMs residing in the perivascular space and leptomeninges42,43, which also derive from the yolk sac44. In addition, caution must be exercised, because many classical microglial markers can also be expressed by cells originating from monocytes or iPSCs, and therefore their presence does not imply *bona fide* microglia. These cells should be more accurately described as monocyte-derived microglia-like or iPSC-derived microglia-like cells (iMGL cells).

As resident macrophages of the brain parenchyma, microglia participate in many critical CNS functions ranging from glio-, vasculo- and neurogenesis to synaptic and myelination, through their process motility, release of soluble factors, and capacity for phagocytosis (**Figure 2**). These functions have been revealed using several constitutive and inducible knock-out models for microglial-specific genes45 and by microglial-depletion paradigms in animal models46, particularly rodents and zebrafish.

The key role of microglia in maintaining CNS health is also supported by the severe phenotype displayed by patients lacking microglia due to loss-of-function CSFR1 mutations. Heterozygous mutations, particularly in the kinase domain of *CSF1R* are associated with ALSP (adult-onset leukoencephalopathy with axonal spheroids and pigmented glia, OMIM:221820) characterized by reduced microglial numbers and white matter atrophy that result in progressive cognitive and motor impairment, dementia, and early death47. Additionally, bi-allelic mutations are reported to cause complete absence of microglia with developmental brain malformation, hydrocephalus, bony lesions, and early death48,49. This phenotype, however, seems in apparent contradiction with the reported absence of gross neurological abnormalities at birth observed in mice with genomic deletion of FIRE, an intra-intronic super enhancer in the *Csfr1*gene enhancer region, whose brains lack microglia50, though more nuanced analyses are needed. Nonetheless, FIRE mice have premature lethality and increased amyloid pathology as early as 5 months of age51. The source of discrepancy between the developmental impact of CSFR1 mutations in humans and mice is not yet fully understood. One possibility is that microglial developmental functions are partly redundant, modified by other environmental factors, or compensated in their absence by other cell types, such as astrocytes52. It will be important to determine how microglia communicate with other glial cells and immune cell populations to support CNS maturation and function in the future.

**(Re)Defining microglial states: DAMs, HAMs, WAMs, and more**

Core markers of cellular identity are useful to identify microglia, but are not necessarily informative about the functional “state” of microglia, which depends on the context (i.e., the physiological conditions in which microglia are found at any given CNS region and time). Microglia have a complex “sensome”53, a series of surface receptors that allow them to detect changes in their environment. Microglial states are thus dynamic, and the outcome of the cell´s epigenome, transcriptome, proteome, and metabolome yields discrete morphological, ultrastructural and/or functional outputs (**Figure 3**). Microglia are anything but static, as they are exceptionally responsive to alterations in their local environment. In the mature healthy CNS, the distribution of microglia is largely uniform and generally regular with little overlap between adjacent territories54. The cell bodies are largely sessile, but their processes are constantly moving and scanning the brain parenchyma55,56. Microglial functions adapt to their location and reciprocal interactions with nearby cells and structures. Their morphology, ultrastructure and molecular profile are similarly dynamic and plastic, resulting in many different cell states. As Conrad H. Waddington, founding father of systems biology, eloquently described: “*Cells are residents of a vast ‘landscape’ of possible states, over which they travel during development and in disease”.57*

Single-cell technologies, multi-omics and integrative analyses of gene and protein expression have helped to not only locate cells on this landscape, but also provide new insight into the molecular mechanisms that shape the landscape and regulate specific cell states in a given context (e.g., development, adult, disease or injury model, etc.). Many diverse and context-dependent microglial states have been observed across species and models. Some examples of these states are the DAM (disease-associated microglia), originally associated with Alzheimer´s disease (AD) pathology models 58; MGnD (microglial neurodegenerative phenotype) documented across several disease models59; ARM (activated response microglia) and IRM (interferon-responsive microglia) in an AD pathology mouse model60; HAM (human AD microglia)61; MIMS (microglia inflamed in multiple sclerosis (MS))62; and LDAM (lipid-droplet-accumulating microglia in aging mice and humans)63, brain tumors (glioma-associated microglia, GAM)64, amyotrophic lateral sclerosis (ALS)-associated signature65 and Parkinson’s disease (PD)-microglial signature66. In the developing and aging brain the WAM (white matter-associated microglia)67; ATM (axon tract-associated microglia)68, and PAM (proliferative-region-associated microglia, related to phagocytosis of developing oligodendrocytes)69, may share some features with the core DAM signature. In the developing human CNS, microglia also express some of the DAM/MGnD/ARM-like profiles70.

While gene expression signatures indicate biological pathways, the functional implications of these states and relationship to one another remain unclear. In fact, the ever-growing list of branding clusters in single-cell RNA sequencing (scRNASeq) experiments and use of acronyms is not consistent across research groups and could hinder future advance of the field without validation and functional experiments to understand their meaning. Moreover, transcriptomic signatures depend on tissue dissection and gating strategies that can lead to isolation artifacts71-74, which, when layered with the technical limitations of single-cell sequencing, can make it difficult to assign state identity across different studies. Another source of complexity comes from evident interspecies differences75-77, which can further hamper comparisons. Advances in computational tools and approaches, which enable the alignment and integration of single-cell datasets, can help solve some of these issues, providing a powerful way to determine microglial state similarities across contexts78,79.

A practical limitation of solely defining functional states by their transcriptional signature is that mRNA expression may not directly predict protein levels80. Protein expression signatures obtained by methods, such as single-cell mass cytometry, have their own technical limitations81 but may better represent true cell states82,83. Importantly, mRNA or protein expression alone do not necessarily predict microglial function, although they can be used to generate functional hypotheses that need to be experimentally tested. There are many methods that allow for the classification of microglia based on their constituent states, including gene expression, protein expression, post-translational modifications, mRNA profiling, morphology and ultrastructure. All these approaches can vary in coverage (e.g., expression of a single cell *versus* whole-transcriptome profiling), which has created overall confusion and mislabeling in the field. Presumably, each microglial state is associated with unique or specialized functions, although the unique roles of any observed state have so far remained elusive. Thus, it is critical that we begin to define microglial states taking into account their specific context within and between species, across sex, space and time (e.g., CNS region and biological age) as well as layers of complexity (e.g., epigenetic, transcriptional, translational, metabolic signatures), which ultimately determine together the cell’s phenome (i.e., motility, morphology, ultrastructure) and function (**Figure 5**).

One major conceptual limitation of the various ‘one-off’ microglial acronyms (e.g., DAM, MGnD, etc.) is that they suggest stable states or phenotypes of microglia associated with a disease context, such as neurodegeneration. Intuitively, this classification system is similar to the concept of neuronal cell types, where neurons cluster into distinct subtypes based on their gene expression or neuroanatomy. However, contrary to microglia, neuronal groupings are considered fixed and terminally differentiated5. We do not know how temporally or spatially dynamic microglial states may be, as microglia are remarkably heterogeneous and plastic. Therefore, these cells are probably not permanently ‘locked’ into any single functional state. From the evidence available so far, microglial states appear dynamic and plastic, possibly transitory, and strongly dependent on the context84. New tools including imaging reporters for microglial states are needed to track transitions within individual cells over time and across the lifespan, following different challenges and perturbations, as well as in response to treatment.

**Microglial heterogeneity: it all depends on the context**

The term “homeostatic” is used to refer to microglia in physiological conditions but there are different interpretations of this nomenclature when describing microglia in health and disease. While homeostatic relates to the ‘physiological’ context assessed in space and time, it does not necessarily correspond to a unique molecular profile because, even without any perturbation, microglia display diverse morphological and functional states, depending on the signals from the CNS microenvironment. This continuous microglial sensing results in multiple transcriptional signatures from development to aging, depending on the specific local signals or challenges to the brain at each developmental stage53. A less responsive microglial state, which in other contexts would be considered more “homeostatic”, might be less effective at responding to damage or pathological cues in aging and disease contexts. For example, in aging and neurodegenerative disease, microglia may have reduced ability to rapidly respond to brain challenges (i.e., removing toxic amyloid, infected, damaged or degenerating neurons), leading to CNS dysfunction and disease progression. Microglia from adult TREM2 knockout mice have been described as ‘locked in a homeostatic state’ as they are less responsive to challenges (such as amyloid) and do not adopt a transcriptional DAM signature in disease contexts85,86. From this example, the term “homeostatic” is not informative if not well-defined and placed in the context of function.

Key modifying factors that lead to microglial heterogeneous states include age, sex, circadian time, local CNS signals and peripheral cues, such as the changes in the microbiota87,88, or other systemic diseases (e.g., asthma)89, in addition to the pathophysiological state of the CNS and overall organism (discussed in more depth in the next section). Age, indeed, has a key influence on the microglial homeostatic state, which goes through several distinct temporal stages (embryonic, perinatal, adult, and aging microglia), each notably characterized by an enrichment of defined regulatory factors and gene expression profiles68,90. After the initial establishment of microglial identity by a network of developmentally programmed and environment-dependent transcription factors75,90, microglia become extremely heterogeneous in their transcriptome during early postnatal development, as determined by scRNASeq68,69,91. In contrast, microglia display a more limited transcriptomic heterogeneity in the adult CNS, where the different microglial scRNASeq clusters fall into a transcriptional continuum instead of representing distinct states68,69,91. Relatively small transcriptional differences may, however, lead to relevant functional differences, as exemplified by the functional variations between hippocampal and cerebellar microglia92,93.

Sex differences due to sex chromosomes and/or gonadal hormones may also impact microglial states in different contexts. A growing body of evidence shows that male and female microglia differ in their transcriptomic, proteomic, and morphological profiles, across brain colonization, maturation and function, in health and disease88,94-96. Of note, the microglial sex-specific transcriptomic signatures appear to be intrinsically determined, being maintained when microglia are transplanted into the brains of mice from the other sex96. Sexually differentiated roles of microglia could critically influence a variety of biological processes, in a time-dependent manner, and thus, emerge as key disease modifiers across various pathological conditions with sexual dimorphism in prevalence, manifestation, and response to treatment97. A well characterized example for sex-specific divergence is the purinergic receptor P2X4R, identified as the male-biased microglial mediator of chronic pain98. Sex differences in sexually dimorphic responses in physiology and pathology likely arise from a combination of Y chromosome-specific genes, sex hormones, neuronal circuit-related factors and epigenetic mechanisms99.

Regardless of the reduced heterogeneity in the mature adult (compared to embryonic) CNS 7,68,90, microglia do differ among CNS areas in terms of their morphology and ultrastructure, transcriptional, proteomic, epigenetic profiles, and functional specialization, suggesting that microglial states are modulated by local cues83,100,101. However, local CNS signals are not sufficient to determine microglial identity because macrophages engrafted in the brain parenchyma can acquire a microglia-like morphology without reaching a transcriptomic signature identical to host microglia, even after prolonged CNS residence26,102,103, supporting the idea that microglia are distinct from peripherally-derived macrophages, even when they colonize a similar niche. In addition, these findings suggest that once their identity is established, microglia assume different functional states in response to local CNS signals. Therefore, both the developmental genetic programs and CNS environment (nature and nurture) collaborate to dynamically determine microglial functional states.

Microglia not only respond to local cues within the brain, but they also receive continuous inputs from the periphery, including signals from the gastrointestinal tract104. In this context, the role of the host microbiota is gaining momentum in controlling microglial maturation and function in the CNS88, with growing evidence that microbiota-derived short-chain fatty acids represent major mediators of the gut-brain axis87,105. Another example of cross-talk between microglia and the periphery is the so called “sickness behavior”, as a result of the central response to peripherally released cytokines produced by peripheral immune cells and tissue resident macrophages detecting specific pathogen-associated molecular patterns (PAMPs)106. This complex and coordinated response, in which the functional role of microglia remains poorly understood, gives rise to adaptive behavioral strategies, including lethargy. Acute systemic inflammation, nevertheless, was extensively shown to impact on microglia107,108 and induce a microglial state associated with robust IL-1 production109.

The concept of the brain as an immune privileged organ has been challenged and definitely revisited in recent years. Indeed, peripherally produced cytokines and immune cells access the CNS and patrol the perivascular space in disease but also in health thus, playing important roles in coordinating central and peripheral immune responses110. It was also suggested that microglia require resident CD4+ T cells in the healthy developing brain for proper maturation and complete fetal-to-adult transition111. Microglia and T cell cross-talk was shown to help maintain homeostasis in the CNS, with dysfunctional regulation occurring in diseases, such as MS112, ALS113, AD114, and encephalitis115. It will be important to continue investigating the influence of the peripheral immune system including B cells, NKs and other cells on microglial states and function in both health and disease.

**Microglial states in the diseased CNS**

Microglia are keen responders and critical players in numerous neurodevelopmental, neurological, and neurodegenerative conditions, as thoroughly reviewed elsewhere. Altered microglial states have been described in the diseased human brain and across various animal models of disease pathology based on morphology and gene expression signature. In addition, these states also differ depending on the timing (i.e., disease stage), genetic background, and local environment. Context-dependent signals vary dramatically during disease progression; they range from apoptotic cells, extracellular debris, toxic proteins (i.e., amyloid, -synuclein), and signals resulting from blood-brain barrier disruption and altered function of neurons and other glial cells. Microglia respond to these challenges by changing their molecular profile, morphology and ultrastructure (**Box 3**), as well as motility and function.

The expression of core microglial markers is also altered over the course of disease, including downregulation of the “homeostatic” microglial signature. A prototypical example is P2RY12, one of the most widely used markers to discriminate microglia from other macrophages, with its reduced expression being one of the salient features of the microglial response to AD pathology and other disease conditions116, as shown in several mouse models of disease (**Figure 4**). The apparent contradiction that core markers do not have a steady expression, as could perhaps be expected, is likely reflecting the functions those proteins have and how they change in the diseased brain. For instance, P2RY12 upregulation in epilepsy may relate to microglial sensing ATP and nucleotides released during seizures117. This seeming paradox strengthens the fact that determining microglial expression profile is far from attributing any function to microglia, as it may only be suggestive of a potential functional identity, which –with unanimous consensus from all the authors– requires experimental validation using appropriate animal models and mutagenesis while using analyses that preserve the environmental influences shaping microglial function.

A microglial state that has received particular focus is the one denoted by the DAM signature, initially identified in a mouse model with mutations within five AD genes (5XFAD)58 and later detected in other AD mouse models and samples from human AD (reviewed in 116) and MS patients62,118. Single cell transcriptomic profiling of human microglial nuclei revealed a tau-associated microglia cluster that had not been identified in mice119, reinforcing the idea that more human studies are needed. The shared DAM signature includes downregulation of CX3CR1 and P2RY12, and upregulation of APOE, AXL, SPP1, and TREM2116, and it has been recently shown that it comprises two ontogenetically different cell lineages, both expressing TREM2: resident microglia and invading monocyte-derived cells (termed disease inflammatory macrophages, DIMs) that accumulate during aging120. Many questions remain open regarding the functional significance of the DAM signature.

Are DAM beneficial, detrimental or both? Several studies, in both mouse and human stem cell-differentiated microglia, demonstrated that the transition to a DAM state is dependent on TREM258,59,85,121. How the TREM2 receptor drives the DAM transcriptional phenotype remains unclear, although the TREM2-ApoE signaling pathway is necessary for the switch from homeostatic to MGnD59. Many questions remain open on TREM2. For instance, is TREM2 a key sensor for amyloid-beta and other AD-related pathology or does its loss of function cause developmental defects in microglia that render them unable to change state? Is TREM2 controlling the microglial state by regulating their energetic and anabolic metabolism?122,123

New bulk and single-cell epigenetic approaches75,124-129 will help answer these questions and ultimately may provide a means to toggle microglial states at will, enabling the field to finally understand the function of distinct microglial states and their impact in different contexts.

Additionally, many genes of the DAM signature were identified across various contexts. For example, a common set of markers including (but not limited to) an upregulation of TREM2, APOE, CD11c, CLEC7A and LPL, and downregulation of TGFβ, CSF1R, P2RY12, and TMEM119 has been recently used to denote a microglial state that associates with myelinating areas in the developing brain, but also with aging and several models of degenerative diseases, such as AD, ALS130, and MS58,67,131. These observations raise the question as to whether the DAM is a signature strictly associated with certain diseases, as the name implies, or perhaps represents a more universal core signature that appears in response to various challenges and may differ between the young/developing *versus* aged/diseased CNS, and across distinct regions. Most likely, the same states that are beneficial in certain contexts may be detrimental in others, strictly depending on the complex interactions between microglia and their surrounding environment. One of the most relevant questions to be addressed is to which extent microglial states identified in the mouse brain are conserved and functionally relevant in the human brain.

**Nomenclature troubles**

Our current understanding of the plasticity of microglial states is at odds with the simplistic scenario established using outdated microglial nomenclature (resting *versus* activated and M1 *versus* M2, **Boxes 1 and 2**). Thus, a systematic, careful naming approach would greatly benefit microglial biology. As a first step to guide the field regarding the use of nomenclature, we generated a questionnaire (**Supplemental Data**) and collected the responses from the co-authors.

Surprisingly, there was more consensus than disagreement that the current nomenclature has severe limitations, and a more useful conceptual framework is needed to properly understand microglial states. There is also agreement that this framework is a first important step to guide the field and should be revisited every five to ten years by an international panel of experts as new discoveries are made. There is also a broad agreement that microglial responses should be framed in a multidimensional space, and should not be simplified as dichotomic good *versus* bad (**Figure 1**). Another point of strong agreement: abandon M1/M2 (and similar) nomenclature once and for all and generally avoid using the vague term ‘neuroinflammation’. Most agree that inflammation is not always detrimental but, instead, represents an adaptive response to damage that can sometimes get out of control (**Box 4**). Quite importantly, a vast majority of authors support the use of “markers” (genes or proteins) to identify cell populations, but not as a readout of cell functions, which need to be addressed directly.

Nonetheless, there were a few points that are still under intense debate. The term “resting” microglia is strongly avoided by some authors, whereas others acknowledge that they still use it even with its limitations, for lack of a better term. “Homeostatic” has more acceptance, although it is recognized that it is based on a very particular gene signature not shared by microglia across all physiological contexts, such as embryonic and postnatal development, and that several homeostatic states likely exist. Thus, the term ‘homeostatic’ should always be accompanied by an accurate description of the context.

The opinion on use of the term “DAM”, on the other hand, is highly polarized. Many authors consider that a core set of transcripts in this signature is common to several pathological conditions and some physiological processes, including the development of white matter, whereas an equal number of authors state there is not enough evidence for “DAM” to be a universal signature of microglial response to damage. Finally, the extent to which microglia are unique or similar to other brain associated or tissue macrophages is evolving with new data and profiling methods: most agree that due to their lineage, microglia are to some extent similar to other macrophages but have unique functions resulting from their longer residence in the CNS environment.

**Recommendations: DOs and DON’Ts**

Based on the collective opinions from the authors, we provide a series of recommendations for researchers, reviewers, and editors. As the field has not yet reached a consensus on several nomenclature topics, including the appropriate use of descriptors for microglial states, it is premature to provide clearer recommendations. Nevertheless, we aim to raise awareness on these issues and stimulate the launch of further initiatives that will guide the field and allow to develop more specific guidelines.

*Classic Nomenclature*

* Consider microglia as highly dynamic and plastic cells that display multivariate morphological/ultrastructural, transcriptional, metabolic and functional states both in the healthy and pathological CNS.
* Describe microglia using as many as possible layers of complexity: ontogeny, morphology/ultrastructure, motility, -omics, and function, always placing them into a species and spatiotemporal context (**Figure 5**).
* Refer to microglia in basal conditions as “homeostatic”, instead of “resting” microglia, considering the limitations discussed above (i.e., that these terms refer to microglia under physiological conditions, not to the function of microglia). Use the term “surveillant/surveilling” to refer to microglia that are engaged in surveillance, but not as a synonym of microglia under normal physiological conditions.
* Refer to microglia in your experimental condition as “reactive to” or “responding to” while describing the particular signals they respond to (i.e., the context), instead of using the widely used broad term “activated”, as microglia are active in both health and disease.
* Disregard simplistic, dichotomic categorizations by providing the observed data and its context.
* Describe profiles of cytokine expression, considering that microglial complexity cannot be reduced to oversimplified and polarized “pro-inflammatory” *versus* “anti-inflammatory” categories. Similarly, do not use M1 *versus* M2 classification.

● When using the term “DAM”, do not use it as a universal term applicable to all diseases, models or challenges. The jury is still out to test whether its full or core signature is common to all or a subset of pathologies, particularly in the human brain.

*Introducing New Terminology*

● Until a consensus is reached about true subtype/s of microglia, with defined ontogeny, physical niches, functions, and transcriptional profiles (whether permanent or transient), use the term “state” rather than “subpopulation.

● Use combinations of gene or protein “markers” to identify putative supopulations but be aware that their expression is plastic and may change over time and under different experimental conditions. Use fate mapping approaches with lineage tracing to track individual microglial cells and assess possible intrinsic differences as well as changes in their state over time84,132.

● In scRNASeq studies, describe the transcriptional signatures (sets or modules of expressed genes) that can be compared with other studies116,133 To describe groups of transcriptionally similar cells in terms of signature, use the term “cluster”.

● Avoid the use of acronyms wherever possible, and only use these once multiple laboratories have defined a stable state with a clearly defined functional role.

● If new terminology needs to be introduced, follow FAIR principles: Findable, Accessible, Interoperable, and Reusable (<https://neuronline.sfn.org/professional-development/data-sharing-principles-to-promote-open-science>). An example of naming cell lines following these principles can be found here134.

*Microglial Markers and Function*

● Use integrative methodological approaches that allow probing of microglia using different levels of analysis (**Figure 5**).

● Follow updated consensus guidelines when using methodologies such as scRNASeq135, RTqPCR136, or digital PCR137.

● Do not use morphology or gene/protein expression as a substitute for directly assessing cell function. Morphology and expression can be used to generate hypotheses about function that need to be specifically tested.

*Grammar Quandary:*

* “Microglia” as a population is a plural noun in English but a singular noun in Latin-derived languages, which occasionally causes confusion. In English texts, microglial cells should always be referred to in the plural form unless referring to an individual cell. For example, “microglia are brain cells” but “this microglia is adjacent to a neuron”.

**Future questions and challenges**

*From words to action:* A key challenge in the field is to match microglial morphological, ultrastructural, transcriptomic, proteomic, metabolomics and emerging lipidomic changes with functional responses (**Figure 3**). In the current single-cell era, an overwhelming wealth of data has been generated, profiling the expression of millions of microglia in different organisms, at different ages, across diverse brain regions. Yet, such ‘omics’ identities are not necessarily linked to functional states, and they often lack spatial resolution. Additionally, many widely used microglial markers are sensome genes, whose expression and activity at the microglial membrane may reflect functional adaptations to a changing environment, and are possibly more indicative of the microglial functional state than the transcription profile.

Transcriptional analysis will benefit from ribosome profiling by RiboSeq138 and from gene-trap insertion profiling by TRAPSeq139. Proteomic approaches combined with *in situ* studies will provide better information in this respect, bridging the gap between expression and function. Further integration of complementary approaches, such as spatial transcriptomics, imaging mass cytometry, and correlative or conjugate electron microscopy in combination with other single-cell approaches, will provide a more comprehensive characterization of microglia. Ultimately, functional studies using specific pharmacological and transgenic approaches in animal models, as well as human-derived cells and organoids are indispensable to understand the multiple roles of microglia within specific spatiotemporal contexts of health and disease.

*How are microglial states coordinated?*

Even as we acquire more data about microglial states, there are still key questions remaining unanswered. To which extent are microglial states plastic and reversible? What is the relationship between microglial state and cellular function? These varied single-cell characterizations ultimately need to be linked to particular functions, to become relevant to development, health, and diseases. How do these states come about? How do signals from the CNS environment get integrated in microglia to produce specific states? New imaging tools and reporters that enable tracking and manipulation of specific microglial states are needed to address these questions.

*How similar are peripherally-derived macrophages and microglia?* A burning question that surely requires further investigation is related to the identity and function of microglia *versus* other brain macrophages. Although recent studies have provided evidence for an intrinsic unique core signature of microglia, their functional resemblances and differences remain undetermined. For instance, could engrafted parenchymal macrophages functionally replace the resident microglia, despite having a different molecular identity, and could they serve as therapeutic vectors?

*The devil is in the details*: Another major caveat is that microglia are incredibly reactive cells and evidence indicates that artifacts are often introduced during sample processing for a variety of methodologies, such as RNA profiling, immunohistochemistry, FACS, *in vivo* imaging, and so on. Hence, we may be missing or confounding important pieces of information because we unintentionally introduce changes in the parameters we are trying to measure. In addition, these artifacts are likely to generate variability across laboratories using different protocols. A future challenge is to increase reproducibility of data across laboratories, by coordinating a shared database of protocols and analysis pipelines curated using STAR methods guidelines. In addition, in the current single-cell multi-omics era, the challenges in big data analysis are exponentially growing140. Statistical methods (including multivariate statistics)141 and artificial intelligence-based data mining approaches (such as machine learning)142 will have to be introduced, to uniformly process and integrate large datasets, as well as extract the biological relevance of the findings.

*Diversity as a source of richness*: Many transcriptional states have been reported during embryonic development, aging, and disease. How many different microglial states can be identified? Within the homeostatic microglia, how many states exist? How do microglia navigate among their many states? Are they related through a transcriptional continuum, or perhaps as a hub-and-spoke set of states, as has been proposed for macrophages4? How dynamic are these states? And how spatially defined are they? Future research will need to address these important questions.

*Male versus female microglia:* Sex differences have been reported to affect the brain colonization, maturation, structure, transcriptomic, proteomic, and functional profiles of microglia, in a time-dependent manner. To what extent these differences may regulate the susceptibility to neurological diseases remains a fascinating question that urgently awaits answers. Investigating the molecular and cellular mechanisms underlying sex-mediated differences in microglial states would advance our understanding of microglial implication in diseases with clear sex-related differences in their prevalence, symptoms, and progression, as well as response to treatments.

*Relevance to humans:* It will be imperative to study developmental and functional differences between human and animal model microglia. To date, most of the studies on microglia were conducted in mice and a direct comparison among brain regions is still missing. Whether microglial states identified in mice also exist in humans is still under debate. Translating and validating these findings across species is critical and will help prevent failure of clinical trials that stem from animal model limitations. In addition, most human microglial studies were performed in Caucasians and only recently data from other groups, such as African American individuals, are becoming available143.

*Towards a unified nomenclature:* The conclusion of this paper is that the community has not yet reached an agreement on what defines microglial identity compared to other cell types; nor consensus on the number, dynamic nature, or definition of microglial states. The community advocates for creating harmonized, curated databases and guidelines for introducing novel terminology; to follow STAR methods; and share data as early as possible. Until such consensus is reached, the community urges all microglial studies to present data with all their layers of complexity and carefully define the context examined to offer clarity instead of confusion, thereby contributing to a more thorough understanding of the many facets of microglial biology. To establish new guidelines for microglial states and nomenclature we call for a community-based approach, whereby the issues and progress are discussed openly in workshops and meetings, with input from diverse researchers across fields and career stages. A useful model to look after are the 10 Human Leukocyte Differentiation Antigen workshops that have taken place since 1982, in charge of renaming CD (cluster of differentiation) antigens (<https://www.sinobiological.com/research/cd-antigens/hlda1>). We lastly advocate for the creation of an international panel/committee of experts in charge of overseeing the guidelines and establishing a specific roadmap to write a white paper in the nearest future.

We would like to conclude with the words of Río-Hortega, who sarcastically identified the problems of microglial nomenclature already 100 years ago: “If we were fond of introducing new nomenclature to describe microglia, as many modern histologists are, who think that enriching nomenclature resolves problems, we would find for microglia names that would indicate their origin, or morphology, or function, in addition to classify all the shapes that acquire when moving and evolving - resulting in the same absurdity that occurs in some branches of Histology and, particularly, Hematology.”144

**Box 1. Resting *versus* activated microglia**

The development of specific silver staining techniques in 1919 allowed Río-Hortega to clearly identify microglia and study their response to experimental manipulations7,145. Early on, Río-Hortega appreciated the striking morphological transformation of microglia following brain damage, but it was in the mid-1970s that the terms “resting” and “activated” microglia first appeared in the literature. These terms were used to morphologically describe cells with affinity for silver staining that were observed in physiological (“resting”) *versus* pathological (“activated”) conditions. This nomenclature consolidated in the 1980s and became widely used during the 1990s146, in parallel with the development and use of histochemical and immunohistochemical techniques, such as lectin staining147, detection of phosphatases and phosphorylases148, and antibodies against the complement receptor CR37. These techniques and nomenclature were pivotal in determining that “resting” microglia were unrelated to astrocytes, as some studies had wrongly concluded149, and that “reactive” microglia shared many characteristics with the blood-borne monocytes10.

As shown by a PubMed search with microglia in all fields, there were only few papers published on the topic before the 1990s, and then a steady increase until the beginning of our century, followed by an exponential growth150. There is a first inflexion point in 2005, with the seminal discovery using non-invasive two-photon *in vivo* imaging that microglia are extremely dynamic in the absence of pathological challenge, continuously surveying the parenchyma with their highly motile processes55,56. The development of non-invasive methods was necessary for our understanding of microglial roles in the healthy brain (reviewed in151). In 2005, microglial extreme dynamism in the intact brain was examined for the first time, through the skull of CX3CR1-GFP mice in which microglia are fluorescently labeled55,56. As a result, microglia are now considered to be the most dynamic cells of the healthy mature brain151. This seminal discovery prompted to rename quiescent or resting microglia as surveying56,152 or surveillant (from the verb to survey)153 microglia, and also led to propose the concept that microglia are never-resting154. Together, these and other *in vivo* two-photon imaging data put into serious doubt the concept of “activated” microglia, which suggests a unique form of response, as in fact microglia are always active, constantly responding (in different ways depending on the context) to the changes in their CNS environment, even under normal physiological conditions. Therefore, microglia do not switch from “resting” to “activated” in response to trauma, injury, infection, disease, and other challenges. Rather, microglia are continuously active and react to the stage of life, CNS region, species, sex, and context of health or disease by adopting different states and performing different functions. Thus, although still widely used, “resting” and “activated microglia” are labels that should be discontinued.

**Box 2. M1 *versus* M2 microglia**

Another terminology emerged in the early 2000s from immunologists classifying macrophages based on findings obtained using *in vitro* models: “M1”, the classical activation, considered pro-inflammatory and neurotoxic, as well as closely related to the concept of “activated” microglia, and “M2”, or alternative activation, considered anti-inflammatory and neuroprotective155. These responses were related to those of T helper lymphocytes (Th1 and Th2) based on their *in vitro* activation by specific immune stimuli that activated differential metabolic programs and changes in cytokine expression156. An associated term is “M0” microglia, which describes their state when cultured in the presence of TGFβ (transforming growth factor beta) and CSF-1 to mimic *in vivo* counterparts157. The terms became widely adopted in microglial research and the 2010s saw a boom of papers phenotyping macrophages and microglia into “M1” and “M2” based on the expression of markers related to these categories, used to indirectly assume a detrimental (“M1”) or beneficial (“M2”) microglial role156. In many cases, editors and reviewers have asked authors to comply with this nomenclature. However, it soon became evident that macrophage responses are more complex than simply “M1” and “M2”158. In the case of microglia, the advent of single cell technologies provided clear evidence that microglia in the living brain do not polarize to either of these categories, often co-expressing M1 and M2 markers159, despite the continued use of M1 and M2 in the literature. We thus recommend to strictly avoid M1 and M2 labels and use more nuanced tools to investigate microglial function (reviewed in160).

**Box 3. Microglial morphological responses across species**

Microglial cells display a profusion of morphologies that have fascinated researchers since the early days of Río-Hortega. Many were tempted to equate morphology with function. Ramified microglia were traditionally associated with the “resting” state, although we now know that ramified microglia actively play many functions during normal physiological conditions. In contrast, “reactive” microglia (rounder cell body, generally with fewer and shorter processes) were called “activated” and equated with an inflammatory response. Only recently, however, a mechanistic link between microglial reduced branching and increased release of the inflammatory cytokine interleukin 1β was reported161. Activation of P2YR12 by tissue damage signals potentiates the tonically active potassium THIK-1 channel, expressed in microglia, leading both to decreased microglial ramifications and activation of the inflammasome machinery processing IL-1β precursors into their mature form161. Another morphology associated with functional changes is “ameboid” microglia, which were thought to be more “phagocytic”, but it is clear now that ramified microglia execute phagocytosis through their terminal or ‘en passant’ branches notably during adult neurogenesis162,163, while in disease conditions such as epilepsy ameboid microglia can display reduced phagocytosis164. Therefore, morphological changes should not be interpreted in functional terms but, rather, taken as a suggestion prompting to investigate further the relationship between microglial structure and function. While the categorization described above is now outdated, the analysis of microglial morphology is considered valuable and still often used across animal model and human *post-mortem* brain studies.

Studies in *post-mortem* brain samples have revealed that human and mouse microglia can adopt similar morphologies. Using the now outdated terms “ramified”, “primed” (larger cell body, ramified processes), “reactive” (ameboid, few ramified processes), and “ameboid” (less than two unramified processes) microglia were described in middle-aged individuals165. In addition, “rod-shaped” microglia (elongated cell body, polarized processes) were found to become more abundant with aging166. Similarly, “dystrophic” microglia, presenting apparently fragmented (but still intact at the ultrastructural level) processes were reported in aging167,168. These different morphological types observed in humans were previously described in rodent models (reviewed in169). Nevertheless, a more sensitive quantitative microglial morphological assessment using a computational pipeline involving cluster analysis revealed differences between mouse and human, with distinct clusters found to be unique to each species170. Subsequently, a high-throughput comparative morphology analysis revealed a generally conserved evolutionary pattern, with some intriguing differences observed between the leech, zebrafish, axolotl, turtle, chicken, gecko, snake, bearded dragon, bat, boar, sheep, whale, hamster, rat, mouse, marmoset, macaque, and human, and across brain regions between mouse and human76. While detailed comparative ultrastructural analyses of microglia between species are currently lacking, the state of “dark microglia” (named based on their increased electron density giving these cells a dark appearance, compared to other microglial states) discovered in 2016, which is defined using electron microscopy by its markers of cellular stress in contexts of aging and disease, was found to be conserved across mouse, rat, and human171,172. New strategies are currently being developed to provide morphological data analyses based on automated pipeline, thus overcoming feature-selection-based biases173. Future studies will show how these varied morphologies correlate with transcriptional and proteomic profiles, and what they imply for the cell’s function. At the molecular level, recent single-cell transcriptome analyses also revealed that human microglia show multiple clusters that indicate a greater heterogeneity than in other mammalian species such as the mouse76,91.

**Box 4. Microglia and the term “neuroinflammation”**

There is a long historical literature stating that inflammation is an important part of recovery from infection, injury, and disease, and it is the lack of resolution of this inflammatory response that is problematic in the context of CNS cell 'reactivity'. Therefore, when the term “neuroinflammation” is encountered in the literature, the reader must be aware that it means different things depending on the context.

While the term “neuroinflammation” is widely used in the field as a synonym of microglial “activation”174, its definition also varies dramatically among authors, according to our survey. Below are representative definitions which are currently used by the authors:

a. Neuroinflammation is inflammation of neural tissue particularly mediated by glial cells.

b. Neuroinflammation is strictly limited to conditions in which leukocytes enter CNS, e.g., in stroke and MS.

c. Neuroinflammation is a mixed cellular response to brain infection or damage involving innate and adaptive responses of resident brain cells and circulating immune cells.

d. The term neuroinflammation is too unclear and imprecise and should be avoided.

Considering that different definitions are used across authors, our main recommendation for the field is to liberate neuroinflammation from microglia and microglia from neuroinflammation, and to use both terms rigorously. The consensus among authors is four-fold. First, protection against tissue damage and extreme departures from homeostasis as well as repair (i.e., ‘inflammation’) encompasses, in the CNS, a highly complex set of local responses, and equally complex interactions with circulating immune cells or with immune cells residing in brain-blood and brain-cerebrospinal fluid interphases. In other words, ‘neuroinflammation’ is not a substitute for ‘microglial reaction’. Second, there are numerous transcriptional states of microglia, astrocytes and oligodendrocytes. The functional outcomes of cells undergoing these transcriptional states remain incompletely understood. Furthermore, it is uncertain which transcriptional states are transient or represent durable cell fate choices. It is also unknown whether changes in states during diseases are ‘inflammatory’ or dedicated to maintaining microglial homeostatic functions. Taking these considerations together, one should exercise extreme caution in simplifying these phenomena as ‘neuroinflammation’, as at least some of these phenomena may represent alternative homeostatic or non-inflammatory reactive states. Third, it is not appropriate to imply that neuroinflammation is invariably deleterious. Rather, it should be recognized that each inflammatory response may exert adaptive or maladaptive effects, contingent on context. To be more specific, research is necessary to explore functions and distinct actions of cytokine-enriched microglia secretomes beyond binary characterizations such as ‘pro-’ and ‘anti-inflammatory’. Fourth, with regards to nomenclature, we recommend the use of modest and precise terms to describe specific phenomena such as: microglial reaction; astrocytic reaction; molecules involved; loss of barrier function at the blood-brain barrier (BBB), etc. All in all, the main message we wish to convey is that inflammation associated with the CNS follows unique rules that need to be fully discerned experimentally and not simply extrapolated from observations in non-nervous tissue.

**TABLES AND FIGURE LEGENDS:**

**Figure 1. Microglial nomenclatures, past and future.** Microglia have been traditionally framed into dichotomic categories but our current integration of epigenetic, transcriptomic, metabolomic and proteomic data favors a multidimensional integration of coexisting states.

**Figure 2. Microglial core properties and functions:** Phagocytosis, surveillance and capacity for releasing soluble factors (inner circle) are core properties through which microglia contribute to key biological functions (outer circle). Created with BioRender.com.

**Figure 3. Microglial identity and states.** The identity of microglia, compared to other CNS-associated macrophages in the perivascular space, choroid plexus and leptomeninges, is established early onfrom yolk sac-derived progenitors. Once they colonize the brain parenchyma and differentiate, they can adopt multiple states depending on the particular spatio-temporal context, as shown in more detail in **Figure 5**. Created with BioRender.com.

**Figure 4. Microglial transcriptomic signatures.** Recent scRNA-Seq studies have identified many microglial transcriptional signatures including but not limited to PAM and ATM in development; DAM, MgnD, ARM, MIMS in disease models of AD, MS, ALS and PD; and WAM, LDAM, HAM in aging, both in mice and human. The key upregulated (red) and downregulated (blue) genes in each signature are indicated. Created with BioRender.com.

**Figure 5. Microglial states defined by their intrinsic and extrinsic determinants, spatiotemporal context, and layers of complexity.** Microglial states depend on intrinsic determinants (such as species, ontogeny, sex, or genetic background) as well as the specific context they inhabit, including age, spatial location, and environmental factors (such as nutrition, microbiota, pathogens, drugs, etc.). All together, these factors impinge on microglia at multiple levels (i.e., epigenomic, transcriptomic, proteomic, metabolomics, ultrastructural and phenomic), which ultimately determine microglial functions. Created with BioRender.com

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | **Marker** | **Specificity** | **Labeled states** | **Staining patterns** | **Main applications** | **Ref.** |
| Antibodies | F4/80 (EMR1) | Macrophages including microglia | Homeostatic conditions and disease-associated.  Expressed in rodents, but presence not yet confirmed in human. | Does not provide a detailed cellular visualization, especially in homeostatic conditions, due to its low basal expression.  Its expression varies significantly between species and is low in human macrophages. | Brightfield or fluorescence analysis of microglial density, distribution, and categorization into morphological states | 175-177 |
| CX3CR1 | Macrophages including microglia | Homeostatic conditions and disease-associated, but downregulated by the DAMs, MGnD, dark microglia, and other pathological states. | CX3CR1-GFP reporter line generally used for visualization, with or without GFP immunostaining. | Brightfield or fluorescence analysis of microglial density, distribution, and categorization into morphological states. | 58,59,178-180 |
| IBA1 | Macrophages including microglia | Homeostatic conditions and disease-associated.  Downregulated in some contexts (e.g., obesity and aging) and by some pathological states (e.g., DAM, dark microglia).  Used to study microglia in early embryonic and postnatal development.  Conserved across several species including human. | Provides exceptional visualization of microglial cell body and processes, including distal extremities.  Diffuses throughout the cytoplasm.  Staining can however be discontinuous in aging. | Brightfield or fluorescence analysis of microglial density, distribution, and morphology.  Ultrastructural studies. | 181,182 58,76,168,183-186 |
| MerTK | Macrophages including microglia | Homeostatic conditions and disease-associated.  Expressed in health and across various contexts of disease, notably in association with the phagocytosis of newborn neurons, amyloid, and myelin. | Partial visualization of microglial cell bodies and diffuse staining of their processes preventing a complete morphological visualization. | Brightfield or fluorescence analysis of microglial density, distribution.  Morphological analysis or categorization into morphological states possible in combination with IBA1. | 187-190 |
| CD11b/c | Macrophages including microglia | Homeostatic conditions and disease-associated.  Used to study microglia in early postnatal development.  Conserved across species including human. | Visualization of microglial cell body and processes.  Low basal expression in adult microglia.  Staining is mainly restricted to the plasma membrane. | Brightfield or fluorescence analysis of microglial density, distribution, and morphology  Ultrastructural studies of subsets downregulating IBA1. | 191 180,192-195 |
| P2RY12 | Largely microglia-specific (not expressed by monocytes), but state-dependent | Homeostatic marker.  Strongly downregulated in disease-associated and reactive states (but upregulated in *status epilepticus*).  Used to study microglia in early postnatal development.  Conserved across several species including human. | Visualization of microglial cell body and processes.  Staining can localize to the plasma membrane or diffuse throughout the cytoplasm and can be more profuse than IBA1 depending on staining conditions. | Brightfield or fluorescence analysis of microglial density, distribution, and morphology.  Ultrastructural studies. | 117,196-198 |
| TMEM119 | Largely microglia-specific, but state-dependent | Homeostatic conditions and disease-associated, but downregulated on reactive microglia in some contexts (e.g., traumatic brain injury and ischemia, MS).  Developmentally regulated.  Conserved across species including human. | Partial visualization of microglial cell bodies and diffuse staining of their processes preventing a complete morphological visualization. | Brightfield or fluorescence analysis of microglial density, distribution.  Morphological analysis or categorization into morphological states possible in combination with IBA1. | 199-203 |
| TREM2 | Macrophages including microglia, state-dependent | Microglial subsets in early postnatal development, aging, and disease conditions (e.g., microglia involved in synaptic pruning or associated with amyloid plaques in AD pathology).  Shown to label monocytes or neurons instead of microglia in human. | Visualization of microglial cell body and processes.  Staining diffuses throughout the cytoplasm. | Brightfield or fluorescence analysis of microglial density, distribution, and categorization into morphological states.  Ultrastructural studies of pathological states downregulating IBA1. | 180,188,201,204,205 |
| Mouse lines | CX3CR1-GFP | Macrophages including microglia | Homeostatic conditions and disease-associated, but downregulated in DAM, MGnD, dark microglia, and other pathological states. | Visualization of microglial cell body and processes.  Fluorescence diffuses throughout the cytoplasm.  Bright enough for two-photon in vivo imaging.  A limitation is that the heterozygous mice used for in vivo imaging are partially deficient in fractalkine signaling, with possible outcomes on the brain and behavior206. The homozygous mice are knockout for CX3CR1 and used to study the outcomes of fractalkine receptor deficiency. | Two-photon in vivo imaging or fluorescence analysis of microglial density, distribution, dynamics, interactions with other parenchymal elements, and categorization into morphological states.  Ultrastructural studies using staining against GFP. | 55,56,178,180,185,207 |
| Iba1-EGFP | Macrophages including microglia | Homeostatic conditions and disease-associated.  Downregulated in some contexts (e.g., obesity and aging) and in some pathological states (e.g., DAM, dark microglia).  Used to study microglia in early embryonic and postnatal development.  Conserved across several species including human. | Visualization of microglial cell body and processes.  Fluorescence diffuses throughout the cytoplasm.  Less bright than fluorescence in CX3CR1-GFP mice, but generally sufficient for two-photon in vivo imaging of cell body and proximal processes.  These mice are not partially deficient in IBA1 in their heterozygous state, which is a main advantage. | Two-photon in vivo imaging or fluorescence analysis of microglial density, distribution, dynamics, interactions with other parenchymal elements, and categorization into morphological states.  Ultrastructural studies using staining against GFP. | 180,184,208 |
| Fms-EGFP or CSF1R-EGFP;  CSF1R-FusionRed | Macrophages including microglia.  CSF1R is expressed by most microglia. | Homeostatic conditions and disease-associated, but considered to be downregulated in DAM and other pathological states. | Fluorescence is less bright than in CX3CR1-GFP mice, and generally sufficient for two-photon in vivo imaging. It also allows for fluorescence-activated cell sorting and fluorescence imaging when combined with immunostaining. These mice are not partially deficient in CSF1R in their heterozygous state, which is a main advantage. | Fluorescence-activated cell sorting and fluorescence analysis of microglial density, distribution, dynamics, interactions with other parenchymal elements, and categorization into morphological states when combined with immunostaining. | 34,162,209 |
| HEXB-TdTomato | Largely overlaps with IBA1 staining but restricted to microglia. Does not label CAMs and other border-associated macrophage populations. | Expression appears stable in homeostatic conditions and disease-associated states. The labeled microglia are also depleted by CSF1R inhibition. | Visualization of microglial cell body and processes.  Fluorescence diffuses throughout the cytoplasm.  Bright enough for two-photon in vivo imaging.  A limitation is that the heterozygous mice used for in vivo imaging are partially deficient in HEXB. However, their microglial gene expression patterns do not appear affected. | Two-photon in vivo imaging or fluorescence analysis of microglial density, distribution, dynamics, interactions with other parenchymal elements, and categorization into morphological states. | 38 |

**Table 1. Main antibody markers and mouse lines used to visualize microglia in rodents and humans from early embryonic development to adulthood and aging.** Other proteins expressed by microglia but whose specificity is not confirmed include APOE, CLEC7A, ITGAX, and LPL.

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