Identification and characterization of a non-conventional CD45 negative perivascular macrophage population within the mouse brain.

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Article

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**ABSTRACT**

Perivascular macrophages (pvM) are closely associated with cerebral vasculature and play an essential role in drainage of the brain and regulation of the immune response. Here, using reporter mouse models and immunofluorescence on sections and whole brain, flow cytometry and single cell sequencing, we identify a Lyve1+ brain perivascular population lacking classical macrophage markers such as CD45 and Cx3cr1. We named the new non-conventional CD45 negative perivascular macrophages pvM2. These cells have a similar location, morphology and phagocytic function as conventional pvM. The pvM2 are not derived from hematopoietic stem cells, as they are negative in the VavtdT lineage tracing model. They increase in number after photothrombotic induced stroke established by flow cytometry and 3D immunofluorescence analysis. Since CD45 negative cells were typically excluded from macrophage studies, the presence of pvM2 has been previously missed and their role is of importance to assess in the brain disease models.

**INTRODUCTION**

The central nervous system (CNS) has long been considered immune privileged, devoid of immune cells other than microglia and without classical lymphatic vessels. The re-discovery of the dural lymphatic network highlights a route for the drainage of the brain to the periphery \(^1\)-\(^6\). How drainage from the CNS parenchyma occurs towards lymphatics and draining lymph nodes is unknown, but there are indications this happens via the foramen at the base of the skull \(^3\).

Besides microglia, several myeloid populations have now been characterized and shown to be essential for brain homeostasis and in brain diseases \(^7\). These myeloid populations include the so-called non-parenchymal or border macrophages which can be classified in the perivascular, subdural meningeal and choroid plexus macrophages. These macrophages are established during development by embryonic precursors derived from yolk sac precursors and are not replaced by blood monocytes during adulthood, except for the choroid plexus macrophages \(^8\). Specifically, perivascular macrophages (pvM) are located in the perivascular space of the blood vessel, delimited by the vascular basement membrane of blood vessels and glial basement membranes \(^9\). The pvM have been shown to be involved in many processes within the CNS. Early studies demonstrated that pvM are capable of scavenging molecules injected into the cerebral ventricles. They contribute to the blood brain barrier function and mediate the up-take of macromolecules \(^10\)-\(^12\), participate in immune regulation \(^13\). They are involved in a wide variety of brain related disorders such as cerebrovascular and neurocognitive functions in hypertension \(^14\), brain infections, immune activation, Alzheimer’s disease \(^15\),\(^16\) and multiple sclerosis.
Lyve1 is notably expressed on lymphatic endothelial cells (LEC) and macrophages, but not on microglia. Specifically, distinct macrophage populations lining the blood vessels within the periphery have been characterized as Lyve1$^\text{hi}$ and Lyve1$^\text{lo}$. In the mouse brain, myeloid cells are a heterogeneous group of cells localized in specific niches and include parenchymal microglia and non-parenchymal pvM$^\text{7,20}$. Previously, brain pvM were described to be CD45$^+$ and were shown to express canonical macrophage markers such as fractalkine receptor (Cx3cr1), colony-stimulating factor1 receptor (Csf-1R), CD206 and Iba-1, but also the prototypic macrophage markers CD11b and F4/80$^\text{7,8}$. Recent studies highlighting Lyve1$^\text{hi}$ and Lyve1$^\text{lo}$ pvM$^\text{19}$, peripheral nervous system and specific border-associated macrophages$^\text{21,22}$ have shown that the macrophage population within the nervous system is heterogenous. However, all these studies have excluded CD45$^-$ cells. In the current study, using different reporter- and fate-mapping mouse models and employing both (whole-mount) immunofluorescence as well as flow-cytometry and single cell sequencing, we observed a previously uncharacterized CD45$^-$CX3CR1$^-$PU.1$^-$Lyve1$^+$F4/80$^+$ cell population within the mouse brain. We established that these cells were not lymphatic endothelial cells, but non-conventional pvM lining blood vessels, having a similar morphology as conventional pvM. Functionally, they were able to phagocytose macromolecules injected within the ventricle. During photothrombotic-induced stroke, CD45$^-$ pvM increased similar as conventional pvM. Hence, our data characterize a previously overlooked CD45$^-$ pvM cell population, which we name non-conventional CD45$^-$ perivascular macrophages (pvM2).
RESULTS

Identification of a Lyve1\(^+\)CX3CR1\(^-\) cell population in the brain parenchyma

Using whole-mount immunofluorescence of the superior and inferior cortex and cerebellum, we identified Lyve1\(^+\) cells that did not form lymphatic luminized vessels (Fig. 1A-D, whole brain in Video 1 and maximum intensity projection in Supplementary Fig. 1A). Their cellular morphology depended on their location. Within the superior cortex (Fig. 1A) they were small and spread, similar as in the pia mater. Within the inferior cortex, these cells were elongated and more stretched, as they were also within the hippocampus (Fig. 1B-C). In the olfactory bulb, near the cribriform plate, where cerebrospinal fluid (CSF) lymphatic drainage occurs from the sub-arachnoid space into nasal lymphatics, Lyve1\(^+\) cells were also found to be long and stretched (Fig. 1D).

In Cx3cr1\(^{GFP/+}\) and Spi1\(^{GFP/+}\) (gene encoding PU.1) mouse brain sections, we identified conventional Lyve1\(^+\)CX3CR1\(^+\)F4/80\(^+\)Iba1\(^+\) pvM (white arrows in Fig. 1). In addition, we observed a Lyve1\(^+\)CX3CR1\(^-\)F4/80\(^-\)Iba1\(^+\) population expressing F4/80 and Iba1 (Fig. 1E, F) but lacking classical hematopoietic and other macrophage markers such as CD45, Csf1-R and PU.1 (red arrows in Fig. 1E-I), which has not been described before. We confirmed the lack of CX3CR1 and PU.1 on sections using an anti-GFP (Supplementary Fig. 1B,C). Furthermore, these atypical Lyve1\(^+\)CX3CR1\(^-\) cells lacked CD163, CD206, MHCII and CD11b expression (Supplementary Fig. 1D-G). The pvM markers expressed are summarized in Table 2.

Lyve1\(^+\)CX3CR1\(^-\) cells are non-conventional CD45\(^-\) perivascular macrophages (pvM2)

In the light of the recent re-discovery of lymphatic vessels in the meningeal compartment \(^1,6,32\), we investigated whether the Lyve1\(^+\)CX3CR1\(^-\)CD45\(^-\) population in the parenchyma of adult mice was of lymphatic origin. We observed Lyve1\(^+\) cells in very close association with blood vessels (Fig. 2A-D, Video 2) and located outside of a structure called “pia mater cul de sac” (Fig. 2B-D). The pia mater formed invaginations into the brain parenchyma, visualized by Podoplanin \(^9\), surrounding arterioles within the cortex. We observed that the Podoplanin\(^+\) pia was located between the vascular endothelium and the Lyve1\(^+\) cells (Fig. 2B-D, Supplementary Fig. 2A). We also observed that the Lyve1\(^+\) cells were positioned within the Laminin \(\gamma 1\) endothelial cell basement membrane within the perivascular space (Fig. 2E-F). This compartment is defined by the vascular basement membrane on the abluminal side of the vessel wall and by the glia limitans basement membrane on the parenchymal side\(^33\). We observed that Lyve1\(^+\) cells did not express CD31 (Fig. 2A-H), Podoplanin (Fig. 2B-D), VEGFR3 (Fig. 2G-H) or master-regulator for lymphatic endothelial identity Prox1 (Fig. 2I-L), ruling out a lymphatic identity.

We verified non-lymphatic phenotype using Prox1\(^{mOrange2}\) (Supplementary Fig. 2B-C) and
Prox1CreERT2;Rosa26tdT (Supplementary Fig. 2D) reporter mouse models \(^{24,34}\). We analyzed the Prox1CreERT2;Rosa26tdT brain 2 weeks after Tamoxifen injection and did not observe Prox1\(^{+}\) lymphatic endothelial cells, ruling out any LEC identity within the brain parenchyma (Supplementary Fig. 2B).

While the dura mater contains lymphatic vessels, no conventional lymphatics have been described within the parenchyma nor the leptomeninges. Similarly, while we did not observe luminized lymphatic vessels within the pia mater, we did find single Lyve1\(^{+}\)Prox1\(^{+}\) lymphatic endothelial cells in the pia mater (Supplementary Fig. 2B-C) as was also recently described in mammals \(^{35}\).

To further rule out astrocyte, glial or fibroblast identity, we analyzed immunofluorescence staining for Aquaporin-4 (AQP4) and GFAP (Supplementary Fig. 2D), ER-TR7 (Supplementary Fig. 2E), PDGFR\(\beta\) (Supplementary Fig. 2F) and observed these cells to be negative for all. Furthermore, we excluded neural crest cell origin using the Wnt1\(^{+}\);Rosa26tdT reporter mouse model (Supplementary Fig. 2G).

We confirmed the presence of the Lyve1\(^{+}\)CD45\(^{-}\) populations at different stages during life by flow cytometry. Using Cx3cr1\(^{GFP}\) brain parenchyma cell suspension devoid of meninges, conventional pvM were identified as Lyve1\(^{+}\)CD45\(^{int/high}\) while microglia were excluded as they are Lyve1\(^{-}\) (gating strategy in Supplementary Fig. 3A). These Lyve1\(^{+}\)CD45\(^{-}\) cells also lacked CX3CR1, confirming IF section stainings (Fig. 1). After 2 days after birth (P2) we observed a Lyve1\(^{+}\)CD45\(^{-}\)CX3CR1\(^{-}\)F4/80\(^{+}\) population which peaked at P14 and P21 (Fig. 3A-D). Subsequently, we observed lowered numbers in adult and 1-year old brain parenchyma (Fig. 3E-F, absolute Lyve1\(^{+}\)CD45\(^{-}\)CX3CR1\(^{-}\)F4/80\(^{+}\) cell numbers in Fig. 3G). Also in the Spi1\(^{GFP}\) (encoding PU.1) brain parenchyma we observed a CD45\(^{-}\)PU.1\(^{-}\)F4/80\(^{+}\) population, next to the conventional CD45\(^{int/high}\) populations (Supplementary Fig. 3C), albeit less Lyve1\(^{+}\)CD45\(^{-}\)PU.1\(^{-}\)F4/80\(^{+}\) cells were present compared to the Lyve1\(^{+}\)CD45\(^{-}\)CX3CR1\(^{-}\) population observed in the Cx3cr1\(^{GFP}\) brain parenchyma.

We confirmed absence of Ptprc (encoding CD45) expression within the Lyve1\(^{+}\)CD45\(^{-}\) cells by sorting this population and determination of Ptprc mRNA levels by qPCR and observed no Ptprc expression within the Lyve1\(^{+}\)CD45\(^{-}\) population (Fig. 4A, sorting strategy in Supplementary Fig. 4A). To further characterize the transcriptome of the pvM population, we analyzed the single cell RNA sequencing dataset of the non-neuronal cell population in mouse brain cortex (GSE133283), thus including the CD45\(^{-}\) cells (Fig. 4B, C). We identified macrophages expressing the characteristic macrophage markers Mrc1, Cd68 and Fcgr3 (Fig.S4B). This macrophage cluster was segregated into at least two subtypes; Lyve1\(^{+}\) and H2-Aa\(^{+}\) (one of the genes encoding MHCII) (Fig.4C). We observed cells with
lower or undetectable Ptprc or Cx3cr1 expression and fewer cells with lower Spi1 expression (Fig. 4C).

However, these cells were not segregated from other cells by unsupervised clustering analysis, confirming their identity as macrophages.

Since we ruled out a lymphatic identity, established that the cells expressed macrophage markers and clustered together with the conventional pvM in the single cell sequencing, we name these cells non-conventional CD45⁻ perivascular macrophages (pvM2).

PU.1 is considered to be a master regulator for macrophage differentiation. Therefore, we analyzed the $\text{Spi1}^{\text{GFP/GFP}}$ mouse model, in which GFP is knocked into the Spi1 locus (encoding PU.1) rendering this gene inactive. Homozygous mice deficient for Spi1 lack all macrophages and die around birth $^8,^27$. In E18.5 $\text{Spi1}^{\text{GFP/+}}$ embryos, we identified Lyve1⁺PU.1⁺CD45⁺ and Lyve1⁺PU.1⁻CD45⁻ pvM (pvM2) (Fig. 5A) while in E18.5 $\text{Spi1}^{\text{GFP/GFP}}$ (knock-out) embryos we observed a total absence of Lyve1⁺ cells in the brain parenchyma (Fig. 5B, positive staining of Lyve1⁺ cells in the skull in Supplementary Fig. 5). This demonstrates that Lyve1⁺CD45⁻ depend on PU.1 during their development, but lose expression during life (Supplementary Fig. 3C).

We tested a possible hematopoietic stem cell (HSC)–bone-marrow derived hematopoietic origin of this population using the $\text{Vav}^{\text{Cre}}$; $\text{Rosa26}^{\text{tdT}}$ (hereafter named $\text{Vav}^{\text{tdT}}$) reporter mouse model. Vav expression starts around embryonic day 8.5 (E8.5) in the hemogenic endothelium of the dorsal aorta in aorta-gonad-mesonephros (AGM) region, which gives rise to the HSC and monocytes $^36$. By immunofluorescence staining of $\text{Vav}^{\text{tdT}}$ brain sections, we identified a Lyve1⁺CD45⁺Vav-lineage⁺ and a Lyve1⁺CD45⁻Vav-lineage⁻ population (Fig. 5C). Therefore, we concluded that the Lyve1⁺CD45⁻ cells were not derived from HSC or bone-marrow derived monocytes.

**Phagocytosis by pvM2**

Besides their role as source for chemokines and growth factors to regulate an immune response, pvM have a phagocytic function, which could be part of a broader role of these cells for tissue homeostasis $^{19,^37}$. In order to address a role of the pvM2 in fluid drainage or macromolecule clearance, we injected 10kD Dextran-AlexaFluor647 and Acetylated LDL-AlexaFluor594 in the lateral ventricle of $\text{Cx3cr1}^{\text{GFP}}$ mouse brains. To exclude recirculation through the blood stream, we analyzed the brain 10 minutes after injection $^3$. We observed that both Lyve1⁺CX3CR1⁻ and Lyve1⁺CX3CR1⁺ cells phagocytized the injected dyes near the ventricle, in the superior- (Fig. 6A-C) and in the inferior cortex (Fig. 6D-F). We concluded that Lyve1⁺CX3CR1⁻ cells were able to efficiently take up lipoproteins and glycoproteins in the range of at least 3 to 10kDa similar as conventional pvM. The close proximity of the pia mater extending towards the third ventricle (3V) could explain direct drainage from the
injection-site within the lateral and connected 3V towards the pia mater (Fig. 6G). Very few Lyve1 and Prox1mOrange2 positive nuclei were observed within the pia mater (Fig. 6G).

**Involvement in central nervous system diseases**

It was shown that in stroke pvM can influence the blood-brain-barrier function\(^{38,39}\). In patients with intracerebral hemorrhage and focal cerebral ischemia, a CD163\(^+\) cell accumulation around brain blood vessels was observed, which also contained myelin\(^{40}\). Moreover, it was not clear whether these cells are blood-derived monocytes or pvM since they were not characterized in detail (Holfelder K. *et al.*, 2011).

To assess the role of the pvM2 in cerebrovascular pathologies, we investigated a photothrombotic model of ischemic stroke. PT was induced on 8 weeks old male mice. In this model, we observed an increase in Lyve1 staining density in whole-mount stained brains (Fig. 7A, method for quantification of staining density in whole mount brain shown in Supplementary Fig. 6A). At day 14 post stroke induction (P14), we noticed a significant increase of Lyve1 staining density (4.2x, \(p<0.05\)) within the hippocampus, although no significant difference was observed in the superior cortex. At P30, Lyve1 staining density had normalized to control values again (Fig. 7B). We noted an increase in the total Lyve1\(^+\) population, including CD45\(^{\text{int}}\) and CD45\(^{-}\) population by flow cytometry (Fig. 7C). The relative numbers of the CD45\(^{\text{int}}\) and CD45\(^{-}\) populations remained similar compared to the control (not shown), but the absolute number of CD45 \(^{\text{F4/80}}\) Lyve1\(^{\text{F4/80}}\) pvM2 increased in PT (Fig. 7D, E and Supplementary Fig. 6B).
DISCUSSION

We have carefully examined whether lymphatic endothelial cells exist within the central nervous system. However, thorough analysis of different reporter mouse models did not reveal these cells within the parenchyma and few individual lymphatic endothelial cells were observed only within the pial meninges as described before \(^{35}\). Using flow-cytometry, single cell, immunofluorescence and lineage-tracing models, we characterized a novel perivascular cell population and named it non-conventional CD45\(^{-}\) perivascular macrophages (pvM2). This population did not express several classical macrophage markers but still clustered together with other macrophages in single cell sequencing analysis. It closely resembled conventional pvM, in location and function. Similar as pvM, pvM2 located outside the cul-de-sac of the pia-mater and within the Virchow-Robin space, which is likely to be associated with their functioning in drainage \(^{8,39,41}\). Classical brain pvM are derived from the yolk-sac hemogenic endothelium and depend on Csf1R during their maturation \(^{8}\). Based on the absence of labeling in the \(Vav1^{DT}\) model, we conclude that pvM2 are not bone-marrow derived monocytes. As was shown for pvM in the periphery \(^{19}\), pvM2 are also most likely derived from the yolk-sac. Using the \(Spi1\) reporter (encoding the PU.1 protein) and knock-out model, we demonstrate that, similar to myeloid cells, they require PU.1 for differentiation but not for maintenance. They could still have the \(Spi1\) mRNA transcripts, but not he protein or GFP reporter, as we observed more cells positive for \(Spi1\) than for \(Cx3cr1\) or \(Ptprc\) in the single cell sequencing analysis. It was previously noted that some macrophages originated from CX3CR1\(^{+}\) precursors but ceased to express \(Cx3cr1\), such as alveolar macrophages, Langerhans and Kupffer cells. These CX3CR1\(^{-}\) cell coincided with CX3CR1\(^{+}\) cells \(^{42}\), as we also observed in the brain parenchyma. Why pvM2 lacked Csf1R and thus likely do not require Csf1 for maintenance is yet unclear. Further experiments using other lineage tracing models are required to reveal their exact origin in relation to the differentiation of conventional pvM. It was observed before that non-hematopoietic ectodermal cells gave rise to phagocytosing cells, resembling Langerhans cells, in the skin of zebrafish \(^{43}\). However, it was not shown that these cells depend on PU.1 and moreover were located within the skin of the zebrafish. Whether these cells resemble the murine pvM2 is yet unclear.

In adult tissues at steady-state, pvM have important functions related to their perivascular location, such as the regulation of vascular permeability, phagocytosis of blood-transmissible pathogens, antigen presentation or immune regulation \(^{37}\). Here, we observed similar phagocytosis of the pvM2 when compared to conventional pvM. Previously, two perivascular monocyte populations were described, being Lyve\(^{1hi}\) or Lyve\(^{lo}\). MHCII was expressed high on the Lyve\(^{1lo}\) population while the Lyve\(^{1hi}\) population was MHCII low \(^{19}\). However, in past studies both Lyve\(^{1hi}\) or Lyve\(^{lo}\) populations were gated
only from CD45<sup>+</sup> cells. CD45 negative non-conventional pvM, if present, were disregarded. Since the pvM2 did not express CD45, MHCII nor CX3CR1, but were Lyve1<sup>+</sup>, it is not clear how these cells relate to the previously described perivascular monocytes.

Using whole-mount imaging enabled us to establish the pvM contribution specifically within and adjacent to the affected region in ischemic stroke. We established an increase in the total pvM number after stroke, with a concomitant significant increase of the pvM2 population. As the number of the pvM2 increases during stroke, these cells may be involved in the generation of an activated immune status. All previous studies on pvM within the brain analyzed CD45<sup>+</sup> cells. This study now for the first time presents evidence of a CD45<sup>−</sup>CX3CR1<sup>−</sup> population, named pvM2, which is present within the brain and should be considered in the future to better understand the active role of diverse macrophage populations in brain homeostasis and disease.

In conclusion, here we demonstrate the existence and function of non-conventional CD45<sup>−</sup> pvM2, lacking classical macrophage makers such as CD45, CX3CR1, PU.1, CD206, CD163 and CD11b. These cells are consistently observed within the brain and are likely involved in drainage also due to their blood vessel proximity and involved in an activated immune state such as during ischemic stroke. Although these novel pvM2 increase in number during stroke, their specific role during disease remains unclear. Presently, there are no unique markers available to target these cells specifically, but a newly developed marker should allow a targeted approach to study their specific role in brain physiology in the future.
METHODS

Mice

C57BL/6J and Vav1-Cre mice were obtained from Charles River [France]. Prox-CreERT2+/− mice and Wnt1-Cre mice were kindly provided by Dr. Bajénoff (CIML, Marseille, France). Cx3cr1GFP mice were kindly provided by Dr. Lelouard (CIML, Marseille, France). Spi1GFP (PU.1), Csf1R-Cre and Prox1mOrange2 were maintained at the CIML (Marseille, France). Vav1-Cre, Prox-CreERT2+/−, Wnt1-Cre and Cx3cr1-Cre lines were crossed to homozygosity for the tdTomato reporter using Rosa26tdT mice. 5 mg of Tamoxifen was injected IP to a Prox-CreERT2+/−.Rosa26tdT adult 2 weeks before the isolation of the brain. All experiments were reviewed and approved by the local ethics committee of Aix-Marseille University and the Ministère de l’Enseignement Supérieur, de la Recherche et de l’Innovation.

Antibodies Table 1

Cell preparation and flow cytometry

Cx3cr1GFP mouse brains from male mice were collected after perfusion with ice cold PBS-Heparin (after removal of the dura). Brains were cut sagitally in 6-8 pieces, then digested with the Adult Brain Dissociation Kit (Miltenyi, 130-107-677) at 37°C for 30min on the gentleMACS dissociator (Miltenyi). Brain cell suspensions were filtered over 70μm strainers and filters were washed with 10mL HBSS 2%FBS. After centrifugation (5min, 400g, 4°C), debris were removed using a 40% Percoll (Sigma-Aldrich, GE17-0891-02) solution in PBS. Cells were centrifuged (30min, 500g, 4°C) and supernatant was discarded. Red blood cell lysis was performed for 10 min at 4°C using the Miltenyi RBC lysis buffer from the kit. Lysis was stopped with 9mL HBSS 2%FBS and cells were centrifuged (5min, 400g, 4°C). Cells were then blocked (15% normal mouse serum (Jackson Immunoresearch 015-000-120) in FACS buffer (HBSS 2% FBS) for 15 min and subsequently stained for CD45-BUV395 (BD biosciences 564279), Lyve1-eFluor660 (eBioscience 50-0443-82) and F4/80-BV421 (Biolegend 123131) diluted in FACS buffer for 30min on ice. The staining of dead cells with Fixable NIR was performed for another 30 min after the antibody staining in protein free HBSS right after washing the cells. Cells were subsequently washed and resuspended in FACS buffer. Samples were acquired on the LSRFortessa X-20 cytometer (BD Biosciences). Data analyses were done using FlowJo software (version 10, FlowJo, LLC).

Cell sorting
Cells were prepared and stained with CD45, and Lyve1 antibodies and Fixable NIR as described above. Live Lyve1^+ CD45^-, live Lyve1^+ CD45^{int} and live Lyve1^+ CD45^{high} cells were sort-purified separately using the BD FACS ARIA III SORP sorter.

**RNA isolation and gene expression analysis**

After spinning down (7 min, 300g, 4°C), cells were resuspended in 1 mL TRIzol reagent (Sigma T9424). The suspension was transferred to a phase lock gel heavy tube to facilitate the extraction and 200 µL of chloroform was added. The samples were shaken vigorously and incubated for 3 min at room temperature (RT) prior to centrifugation (10 min, 12 000g, 4°C). The clear aqueous phase was transferred to a new tube and 2 µL of GlycoBlue Coprecipitant (Invitrogen AM9515) was added to help with the isolation prior to adding 500 µL of isopropanol. The samples were incubated for 10 min at RT and subsequently centrifuged (30 min, 12 000g, 4°C). The supernatant was discarded and RNA was washed with 800 µL EtOH 70%. The samples were centrifuged (5 min, 12 000g, 4°C) and supernatants were discarded. The RNA was air-dried for 15 min prior to resuspension in 11.5 µL RNase free water. The tubes were subsequently heated at 65°C for 10 min and then put on ice. RNA quantity and quality were assessed by nanodrop. For each sample, 200 ng of RNA was reverse transcribed using the RevertAid Reverse Transcription kit (Thermo Scientific K1691) and adding Oligo (dT)_{12-18} primer (Invitrogen, 12418012). The level of expression of CD45 (Forward CCCCCGGATGAGACAGTTG; Reverse AAAGCCCGAGTGCCTTCCT) gene was assessed by qPCR using Sybr Green (Takara, TAKRR420W) and Hprt was used as reference gene (Forward GGCCAGACTTTGTTGGATT; Reverse CAGATTCAACTTGCCTCAT). Data were analyzed using the delta-delta Ct method and compared to the CD45^- fraction.

Single cell RNA sequencing data of non-neuronal cell populations in mouse brain cortex was adapted from GSE133283 (unpublished). Mrc1-expressing perivascular macrophage population was isolated and reanalysed using Seurat (version 3.2.1) R package. Variable genes were found with parameters of selection.method = vst and nfeatures = 2,000, trimmed for the genes related to cell cycle (GO:0007049) and then used for principal component analysis (RunPCA, npcs=30). Statistically significant principal components were determined by JackStraw method and the first 5 principle components were used for non-linear dimensional reduction (RunTSNE) and clustering analysis (FindNeighbors) with resolution=0.1. FeaturePlot function was used to visualize the specific gene expression on TSNE plot. Blue represents the relative expression level of each gene.

**Whole-mount staining and lightsheet imaging**

Both male and female mice were perfused with PBS-Heparin 5U/ml (Sigma) and subsequently overnight (ON) immersion fixed in paraformaldehyde 4% (PFA-Electron Microscopy Science, ref...
The iDISCO protocol was used for immuno-staining, prior to which brains were dehydrated in increasing methanol (MetOH) concentrations diluted in PBS (20, 40, 60, 80 and twice 100%) for 30 minutes for each concentration at RT. Subsequently specimens were incubated ON in a dichloromethane (DCM, Sigma 270997)-MetOH mixture (2 vol DCM:1 vol MetOH) at RT. After two 10 minutes incubations in absolute MetOH, brains were bleached in 5% H₂O₂ in MetOH ON at 4°C and subsequently rehydrated by a decreasing MetOH series (80, 60, 40, 20% in water), followed by PBS and two washes in PBS-Triton X100 (Tx) 0.2% for 1 hour each. Bleached brains were permeabilized for 2.5 days at 37°C (0.4%Tx, 20%DMSO, 2.3% Glycine in PBS) and subsequently blocked with [PTwH (PBS, Tween20, Heparin), 10% DMSO, 6% serum] for 4.5 days at 37°C. Whole-mount stainings were performed by incubation with primary antibodies for 5 days at 37°C and subsequently with Alexa-dye coupled secondary antibodies diluted in PTwH containing 3% serum for 5 days at 37°C. Following each staining step, samples were extensively washed in PTwH (10min, 15mn, 30min,1h, 2h, and ON at RT). Finally, the samples were again dehydrated by increasing MetOH concentrations diluted in water (20, 40, 60, 80, 2x 100% and ON in absolute MetOH), for 1 hour each step at RT and cleared in a MetOH/BABB mix (1:1) [BABB (benzyl alcohol and benzyl benzoate = 1:2) (Sigma 305197 and Fisher Scientific 10654752)] for 8 hours at RT and finally placed in BABB ON at RT to complete clearing. All incubations were done under mild agitation. After clearing, brains were imaged using a LaVision Ultramicroscope II (LaVision BioTec, Bielefeld, Germany). Stacks were captured with a step size of 5µm at 2.5X magnification using an optic zoom with a NA=0.144.

Vibratome section immunofluorescence staining and confocal imaging

Animals were perfused with PBS/heparin, brains dissected and fixed ON in 4% PFA in PBS at 4°C and subsequently embedded in 1% low melting agarose for generation of 100µm vibratome slices (Leica, VT1000S). Sections were blocked in EBT buffer (EBSS, 0.05% Tx, 1% BSA) containing 10% serum for 2h at RT under agitation. Immunostainings were performed by incubation in primary antibodies for 48h at 4°C in EBT, 3% sera and subsequently with Alexa-fluorochrome coupled secondary antibodies diluted in EBT, 3% sera for 24h at 4°C. Following each staining step, samples were washed several times in PBS-Tx (0.05% Tx in PBS) and in PBS. Sections were finally cleared in
Histodenz (Sigma D2158) medium for 48h at RT and subsequently mounted in Histodenz medium. Confocal images were acquired at RT on a confocal microscope (LSM880, Zeiss, Germany), with a 20x/0.4 Plan-Apochromat objective and using laser lines at 405, 488, 561, and 633nm for the excitation of AlexaFluor405/GFP/AlexaFluor555/AlexaFluor647 respectively. Fluorescence was recorded in individual channels acquired in a sequential mode using a highly sensitive 32-channel GaAsP detector. Channels were respectively detected using these detection bands: A405 (410-470nm), A488 (490-540nm), A555 (565-640nm), A647 (640-690nm). The pinhole was set to 1 airy unit. Z stack were acquired with an optical thickness defined for each image in figure legends, satisfying the Nyquist resolution criterion. Image processing (contrast enhancement, scale bars, etc.) was done with ImageJ (National Institutes of Health) without actions modifying image integrity.

**Infusion of tracers into lateral ventricle**

Both female and male mice were anesthetized (150mg/kg Ketamine and 10mg/kg Xylazine) and fixed in a stereotaxic frame. The skull was thinned with a dental drill at a location of 0.95 mm lateral and 0.22 mm caudal from the bregma. A 30G needle with a silica fiber tip (Phymed) was inserted into the right lateral ventricle 2.35 mm ventral to the skull surface as previously described. Injection of 2.5 µL acetylated LDL-Alexa594 (ThermoFisher) or Dextran-Alexa647 10kDa (ThermoFisher) tracer was done at a speed of 0.5 µL/min using a high precision syringe pump. The needle was left in place for 5 minutes and slowly retracted confirming lack of detectable backflow. To avoid recirculation through the blood stream into the brain, mice were sacrificed 10 min after injection. The mice were perfused with PBS and 4%PFA/PBS, brains isolated and fixed ON in 4% PFA/PBS at 4°C, subsequently washed and then stored at 4°C in PBS until further analysis.

**Photothrombotic (PT) induced stroke**

Stroke using the PT model was induced in male C57BL/6J and Cx3cr1<sup>GFP</sup> mice. Mice were anesthetized by intraperitoneal Ketamine/Xylazine injection as before and the eyes of the mice were covered with Ocry-gel to protect them from light and dehydration. The skin on the skull was incised from the eyes to the neck and retracted to the edges of the skull. After retro-orbital injection of 100 µL Rose Bengal (Sigma, 330000) a Leica, KL 1600LED cold light source was placed in contact with the skull and the illuminated region was precisely adjusted (using a stereotaxic instrument) to 2.5 mm caudal of the Bregma and 2.0 mm to the Lambda. After illuminating a 1 mm diameter area at power 3 for 15 minutes, the skin was put back into place, stitched and the animals returned to their home cages. Food was provided in a plate and Buprenorphine (0.4 mg/ml) was added to the drinking water. Mice were sacrificed between 14- and 30-days post-induction and brains were analyzed by immunofluorescence and flow cytometry.
Statistical analysis

Graphs, average values and standard deviation (SD) shown in all figures were calculated using Prism (version 8.3.0 GraphPad software) software. Unpaired T-Test and One-way Anova test were used to determine significance. The number of individual experiments can be found in the legends of all the figures. Photoshop software (CC2015, Adobe) was used to generate figures.

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Competing Interests
The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Author Contributions
CS, ML, HWJ, MS, LF, AT, performed experiments. SAvdP conceived and supervised the study. CS, ML, HWJ, MS, and SAvdP analyzed data. CS, ML, HWJ, MS, AT, MF, SS, MHS, RS, RHA, SSM, FK and SAvdP contributed to discussion and wrote, illustrated, reviewed, and edited the manuscript. SAvdP acquired funding. All authors approved the submitted version of the manuscript.

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Data Availability Statement
All datasets generated for this study are included in the article/Supplementary Material. Single cell sequencing data is available under GSE133283.
FIGURE LEGENDS

**Figure 1: Identification of a Lyve1^+CX3CR1^- population in the mouse brain.** Confocal microscopy of brain sections stained with αLyve1 showing the different Lyve1^+ cell morphologies in the superior cortex (17µm maximum intensity projection) (A), in the inferior cortex (27µm maximum intensity projection) (B), in the hippocampus (15µm maximum intensity projection) (C) and in the olfactory bulb (61µm maximum intensity projection) (D) of an adult mouse brain. (E-I) Immunofluorescence microscopy on sections of Cx3cr1^GFP and Spi1^GFP (encoding PU.1) in the superior cortex of the mouse brain. (E) Staining of Cx3cr1^GFP sections for Lyve1 (red) and F4/80 (50µm maximum intensity projection), (F) Iba1 (29µm maximum intensity projection), (G) CD45 (46µm maximum intensity projection), (H) Csf1R (34µm maximum intensity projection). (I) Staining of Spi1^GFP (PU.1 – green) sections for Lyve1 (red) and CD45 (37µm maximum intensity projection). White arrows point to conventional Lyve1^+CX3CR1^+ pvM cells, which expressed F4/80^+, Iba1^+, CD45^+ and Csf1R^+. The red arrows indicate Lyve1^+CX3CR1^- cells, which are F4/80^+, Iba1^+, CD45^- and Csf1R^- PU.1^-.

**Figure 2: Perivascular Lyve1^+ cell location.** (A) Maximal projection (200µm maximum intensity projection) of a lightsheet microscope acquisition on a cleared mouse brain labeled for SMA (green), CD31 (red) and Lyve1 (blue). (B) Confocal microscopy of brain sections stained for Podoplanin (green), Lyve1 (red) and CD31 (blue), inserts (C) and (D) zoom in on Lyve1^+ cells lining the blood vessels and placed outside of the « cul de sac of the pia mater » defined by the podoplanin labeling. (E) Confocal microscopy of a superior cortex section stained for CD31 (green), Lamininγ1 (red) and Lyve1 (blue) with insert (F) showing a higher magnification showing the Lyve1^+ cells in the perivascular space. (G) Lyve1^+ cell characterization by immunofluorescence on adult brain sections for Lyve1 (green), VEGFR3 (red) and CD31 (blue) staining. The arrowhead points at a blood vessel and the white arrow shows a Lyve1^+ cell negative for VEGFR3 and CD31. (H) high magnification of the Lyve1^+ cell in (G). (I and J) Lyve1^+ cells (green) did not express Prox1 (red) in the superior cortex (39µm maximum intensity projection) nor in the inferior cortex (50µm maximum intensity projection) (K and L).

**Figure 3: Confirmation of the Lyve1^+CX3CR1^- population by flow cytometry.** Flow cytometric analysis of Cx3cr1^GFP brain parenchyma, pre-gated on living, single and Lyve1^+ cells. Microglia cells were excluded since these are Lyve1^- . Mouse brain were analyzed at (A) P2 (n= 4), (B) P7 (n=4), (C) P14 (n=4), (D) P21 (n=4), (E) Adult (8-12 weeks after birth, n= 4) and (F) 1-year old brains (n=3).
showing subsequent CX3CR1, F4/80 expression for the cells of the CD45$^{\text{int}}$ and CD45$^{-}$ gates. (G) Total number of Lyve1$^{+}$CD45$^{-}$CX3CR1$^{-}$F4/80$^{+}$ cells at the different stages.

**Figure 4: Transcriptional profiling of the pvM subsets.** A) Quantification by RT-QPCR of the relative expression of Ptprc (encoding CD45) within the CD45 cell subpopulations (CD45$^{-}$ for CD45 negative, CD45 I for intermediate and CD45 H for high expressing CD45) isolated from C57BL/6J brain parenchyma (3 sorts with n=5 mice per sort), (B) Single cell RNA sequencing of non-neuronal population, focused on the macrophage subset, revealed two main populations. (D) A segregation occurred between a Lyve1$^{+}$ vs. MHCII (e.g. H2-Aa) macrophage subsets, visualized in the individual gene expressions plotted within the tSNE plots. Ptprc and Cx3cr1 transcripts were present in few cells while Spi1 (encoding PU.1) transcripts were present in more Lyve1 and MHCII cells.

**Figure 5: Origin of Lyve1$^{+}$ cells.** (A) Immunofluorescence on sections of the head of a E18.5Spi1$^{\text{GFP/+}}$ embryo, labeled for Lyve1 (red) and CD45 (60$\mu$m maximum intensity projection) shows the presence of the conventional (white arrows) and the non-conventional pvM population (red arrows) at this stage. (B) No GFP nor Lyve1 fluorescence was observed in the Spi1$^{\text{GFP/GFP}}$ E18.5 superior cortex (33$\mu$m maximum intensity projection). (C) Confocal analysis of Vav$^{\text{tdT}}$ (26$\mu$m maximum intensity projection) brain sections stained for Lyve1 (green) and CD45 (white). White arrows indicate the conventional pvM population, which is tdT positive (red) and the red arrows the CD45$^{-}$CX3CR1$^{-}$ population which is tdT negative.

**Figure 6: pvM2 share phagocytic functioning with conventional pvM.** (A-C) Cx3cr1$^{\text{GFP}}$ mouse brains were injected intra-ventricular with Dextran-Alexa647 (37$\mu$m maximum intensity projection) and (D-F) Acetylated-LDL-Alexa594 (42$\mu$m maximum intensity projection). Ten minutes after injection, mice were sacrificed and brains dissected. Confocal analysis on brain sections shows Lyve1 (white), Dyes (red) and CX3CR1 (green). The pvMs phagocytosed the dyes (red arrows), as well as did conventional pvM (white arrows) (B, C, E and F). (G) Tiled confocal acquisition of a Prox1$^{\text{mOrange2}}$ brain section. Lyve1$^{+}$ cells in green lined blood vessels stained for CD31 in blue. Lyve1$^{+}$ cells in green penetrated the brain to the hippocampal fissure (hif) and all the way towards the third ventricle (3V).

**Figure 7: pvM2 increase in numbers after induced ischemic stroke.** (A) Maximal intensity projections (2660$\mu$m) of lightsheet acquisitions on cleared brains, stained for Lyve1 (red) and CD45 (white), control and after stroke (P14=14 days after induction of ischemic stroke). (B) Quantification
of Lyve1 in the area containing the whole stroke lesion within the superior cortex situated just above
the hippocampus, imaged by lightsheet microscopy. (C) Representative flow cytometry plots from
Cx3cr1GFP brain parenchyma 14 days after induced stroke (P14), pre-gated on living, single and Lyve1+
cells. Microglia cells were excluded since they are Lyve1−. CX3CR1 vs F/80 expression is shown for
the CD45− population. Average percentages for the different subpopulations are shown in the plots
(n=4). (D) Graph representing the total Lyve1+ cell number showing an increase of this population at
P14 after stroke by flow cytometry. (E) Total pvM2 (Lyve1+CD45−CX3CR1−F4/80+) numbers in the
control vs. PT brains at P14.
**SUPPLEMENTARY MATERIALS**

**Table 1**: Ab Resources Table for Imaging (confocal and lightsheet microscopy)

**Table 2**: pvM vs pvM2 markers

**Supplementary Figure 1**: pvM2 characterization. (A) Lightsheet imaging of a cleared adult C57BL/6J mouse brain immunolabeled with αLyve1 using iDisco+ protocol (5550μm maximum intensity projection), related to video 1. (B-C) Immunofluorescence microscopy on brain parenchyma sections of (B) Spi1\textsuperscript{GFP} and (C) Cx3cr1\textsuperscript{GFP} in the superior cortex, using an anti-GFP to enhance the GFP signal (green) with Lyve1 staining labeled in red. (D-G) Brain sections are labeled for Lyve1 (green) and for (D) CD163 (16μm maximum intensity projection), (E) CD206 (48μm maximum intensity projection), (F) MHCII (35μm maximum intensity projection), (G) CD11b (85μm maximum intensity projection) (red). White arrows show the conventional pvM population and red arrows show the non-conventional pvM population.

**Supplementary Figure 2**: Lyve1\textsuperscript{+} cells are neither lymphatic endothelial cells (LEC), nor astrocytes, nor fibroblasts. (A) 1080μm maximum intensity projection of lightsheet microscope acquisition of the superior cortex illustrating the « cul de sac of the pia mater » defined by the podoplanin labeling, structure emanating from the pia mater (CD31 in green, Lyve1 in red and Podoplanin in blue). (C) Confocal analysis of Prox-CreERT2\textsuperscript{+/−}; Rosa\textsuperscript{tdTomato} brain sections stained for Lyve1 (green) confirmed that Lyve1\textsuperscript{+} cells do not express tdTomato (red) (white arrows) (88μm maximum intensity projection). (D) LEC identification (Lyve1\textsuperscript{+} Prox1\textsuperscript{+}) in the pia mater in the superior cortex by Prox1 antibody staining (red) on C57BL/6J mouse brain in combination with Lyve1 (green). Prox1\textsuperscript{+} neuron cellular bodies are observed in the parenchyma (13μm maximum intensity projection). (E) Astrocyte identity was excluded by AQP4 and GFAP (red) staining (24μm maximum intensity projection for the superior cortex and 17μm maximum intensity projection for the inferior cortex), (F) fibroblast identity was excluded by ER-TR7 staining (50μm maximum intensity projection for the superior cortex and 8μm maximum intensity projection for the inferior cortex) and (G) PDGFRβ (red) staining. (H) Confocal analysis of Wnt-Cre; Rosa\textsuperscript{tdTomato} (37μm maximum intensity projection) brain sections stained for Lyve1 (green), containing tdTomato (red). White arrows indicate the Tomato negative CD45\textsuperscript{−}CX3CR1\textsuperscript{−} perivascular cells.
Supplementary Figure 3: Confirmation of the Lyve1⁺CX3CR1⁻ population by flow cytometry

(A) Flow cytometry gating strategy. (B) CD45<sup>High</sup> cells from Cx3cr1<sup>GFP</sup> brain parenchyma, pre-gated on living, single and Lyve1⁺ cells. Microglia cells were excluded as Lyve1⁻. Mouse brain analysis at different ages P2 (n= 4), P7 (n=4), P14 (n=4), P21 (n=4), Adult (8-12 weeks) (n=4), and 1 year old, (n=3) showing subsequent CX3CR1, F4/80 expression for the cells of the CD45<sup>High</sup> gates. Gating of pvM identified as Lyve1⁺CD45<sup>High</sup> isolated from Cx3cr1<sup>GFP</sup> brain parenchyma. (C) Flow cytometry analysis of CD45 cell subpopulations isolated from Spi1<sup>GFP</sup> adult brain parenchyma (8-12 weeks after birth, n= 4), pre-gated on living, single and Lyve1⁺ cells. Microglia cells were excluded as Lyve1⁻. Plots show subsequent PU.1, F4/80 expression for the CD45<sup>Int</sup>, CD45<sup>Neg</sup> and CD45<sup>High</sup> gates.

Supplementary Figure 4: Gene expression profiling

(A) Sort gating strategy for the isolation of the different Lyve1⁺CD45 populations (B). Single RNA sequencing analysis showing different macrophage associated genes within the tSNE plots.

Supplementary Figure 5: Immunofluorescence of the section of a E18.5 head from a Spi1<sup>GFP/⁺</sup> embryo (PU.1-GFP in green), labeled for Lyve1 (red) and CD45 (white). No GFP or Lyve1 fluorescence was observed in the Spi1<sup>GFP/GFP</sup> E18.5 superior cortex (33µm maximum intensity projection) but we observed Lyve1⁺ cells, possibly lymphatic vessels, in the skull.

Supplementary Figure 7: Lightsheet data quantification method using Imaris software. (A) To quantify the Lyve1 staining in cleared brains imaged by lightsheet microscopy, Imaris software was used. Using the “Surface tool” of Imaris, a 3D Rendering function, a first volume was created, named « Surface1 » at different locations (Superior cortex and hippocampus) where we wanted to quantify the staining. Based on Lyve1 labelling, a new channel was created. This new channel was used to create a new volume called « Surface 2 » representing the Lyve1 staining. Imaris calculated the different volumes in µm³. The ratio « Surface2 » / « Surface1 » allows us to normalize the Lyve1 quantification. (B) Gating of conventional pvM identified as Lyve1⁺CD45<sup>Int</sup> and as Lyve1⁺CD45<sup>High</sup> isolated from Cx3cr1<sup>GFP</sup> brain parenchyma at P14 after stroke.

Supplementary Video 1: 3D reconstruction of Lyve1 cells within the mouse brain. A cleared sagittal half of an adult C57BL/6J mouse brain immunolabeled for Lyve1 (red) was acquired on the lightsheet microscope and a 3D reconstruction was made using Imaris. The video zooms in on the inferior cortex and subsequently on the hippocampus to indicate the differences in pvM morphology.
Supplementary Video 2: Z-stack acquisition by confocal microscopy and 3D reconstruction of the pvM morphology adjacent to a blood vessel within the brain parenchyma. A 100μm brain section was stained with αLyve1 (red) and αCD31 (green). 3D reconstruction was done using the Imaris surface tool and the video was created using the animation function of Imaris.
Figure 4

A

Relative expression $Ptprc$

CD45-  CD45 I  CD45 H

B

tSNE_1  tSNE_2

C

Lyve1  H2-Aa  $Ptprc$

Cx3cr1  Spi1
### Table 1
Ab Resources Table for Imaging (confocal and Lightsheet)

| Primary Antibodies | Source             | Identifier |
|--------------------|--------------------|------------|
| Lyve-1             | R&D systems        | AF2125     |
| AQP4               | R&D systems        | AB3594     |
| GFAP               | Biolegend          | 644706     |
| CD45               | eBioscience        | 14-0451-85 |
| MHCIi              | BD Pharmingen      | 562352     |
| F4/80              | Biolegend          | 123122     |
| Iba1               | Abcam              | ab107159   |
| CD163              | Anders             | homemade   |
| CD11b              | eBioscience        | 16-0112-82 |
| CD206              | Thermofisher       | 48-2061-82 |
| CD31               | Thermo             | MA1-40074  |
| Prox-1             | Reliatech          | 102-PA32   |
| VEGFR3             | R&D systems        | AF743      |
| α-SMA              | Thermofisher       | 53-9760-82 |
| PDGFRβ             | Cell Signaling     | 31695      |
| ER-T7R             | Thermofisher       | MA1-40076  |
| Csf-1R             | Santa Cruz         | sc-692     |
| GFP                | AVES               | GFP-1020   |
| Laminin γ1         | Sorokin L (University of Muenster) |           |

| Secondary Antibodies | Source                | Identifier   |
|----------------------|-----------------------|--------------|
| DaChCy3              | Jackson ImmunoResearch| 703-166-155  |
| DaG 488              | Thermofisher          | A-11055      |
| DaG 555              | Thermofisher          | A-21432      |
| DaG 647              | Thermofisher          | A-21447      |
| DaG 790              | Jackson ImmunoResearch| 712-655-153  |
| DaR 488              | Thermofisher          | A-21208      |
| DaR 594              | Thermofisher          | SA5-10028    |
| DaR 647              | Jackson ImmunoResearch| 712-605-153  |
| DaRb 488             | Thermofisher          | A-21206      |
| DaRb 555             | Thermofisher          | A-31572      |
| DaRb 647             | Jackson ImmunoResearch| 711-605-152  |
### Table 2

| Conventional pvM     | pvM2     |
|----------------------|----------|
| Lyve1⁺                | Lyve1⁺   |
| CD45⁺                | CD45⁻    |
| CX3CR1⁺              | CX3CR1⁻  |
| PU1⁺                 | PU1⁻     |
| Csf1R⁺               | Csf1R⁻   |
| F4/80⁺               | F4/80⁻   |
| Iba1⁺                | Iba1⁻    |
| CD163⁺               | CD163⁻   |
| CD11b⁺               | CD11b⁻   |
| CD206⁺               | CD206⁻   |
| MHCII⁺               | MHCII⁻   |
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- pvM2SiretetalSupplFig.pdf
- Video1Lyveinbrainzoomincortexandhc720.mov
- Video2cd31lyve13dvessel720.mov