CNS-localized myeloid cells capture living invading T cells during neuroinflammation

Beatrice Wasser1*, Dirk Luchtman1*, Julian Löffel1, Kerstin Robohm1, Katharina Birkner1, Albrecht Stroh2, Christina Francisca Vogelaar1,2, Frauke Zipp1**, and Stefan Bittner1**

To study the role of myeloid cells in the central nervous system (CNS) in the pathogenesis of multiple sclerosis (MS) and its animal model, experimental autoimmune encephalomyelitis (EAE), we used intravital microscopy, assessing local cellular interactions in vivo in EAE animals and ex vivo in organotypic hippocampal slice cultures. We discovered that myeloid cells actively engulf invading living Th17 lymphocytes, a process mediated by expression of activation-dependent lectin and its binding partner, N-acetyl-D-glucosamine (GlcNAc). Stable engulfment resulted in the death of the engulfed cells, and, remarkably, enhancement of GlcNAc exposure on T cells in the CNS ameliorated clinical EAE symptoms. These findings demonstrate the ability of myeloid cells to directly react to pathogenic T cell infiltration by engulfing living T cells. Amelioration of EAE via GlcNAc treatment suggests a novel first-defense pathway of myeloid cells as an initial response to CNS invasion and demonstrates that T cell engulfment by myeloid cells can be therapeutically exploited in vivo.

Introduction

Variable roles of resident myeloid cells of the brain have been reported in both the healthy and diseased state (Giles et al., 2018; Schafer and Stevens, 2015; Shemer et al., 2018). Aside from defending against invading pathogens, they also use their phagocytic capacity to clear cellular debris, infectious agents, or neurons undergoing programmed cell death, thereby attempting to maintain tissue homeostasis. In steady state, myeloid cells actively scan their environment by prolonging and retracting their highly motile processes in order to screen the central nervous system (CNS) tissue with a turnover of just a few hours (Nimmerjahn et al., 2005).

In experimental autoimmune encephalomyelitis (EAE), an animal model mimicking many aspects of the neuroinflammatory disease multiple sclerosis (MS), in vivo experiments support a detrimental role of myeloid cells (Heppner et al., 2005). Inhibition of myeloid cell function repressed clinical EAE severity in a bone marrow chimera model and reduced both axonal destruction (Heppner et al., 2005) and secretion of proinflammatory cytokines and chemokines in the spinal cord (Goldmann et al., 2013). However, myeloid cell function was also reported to change properties context dependently in the course of the disease (Giles et al., 2018). In contrast to their well-characterized neurotoxic ability, a potential neuroprotective function of CNS myeloid cells is far less understood (Kerschensteiner et al., 2003). Studies investigating polarized “anti-inflammatory” microglia that are able to secrete anti-inflammatory cytokines such as IL-10 were usually performed under in vitro conditions (Lobo-Silva et al., 2017; Napoli and Neumann, 2010). A beneficial in vivo function of myeloid cells of the CNS was found in cuprizone models, in which effective clearance of myelin debris by myeloid cells was demonstrated (Lampron et al., 2015; Olah et al., 2012; Sierra et al., 2013). Live interaction between myeloid cells of the brain and CNS-infiltrating immune cells in vivo has not been investigated up to now.

Here, we applied intravital two-photon laser scanning microscopy to unravel physiologically meaningful myeloid cell responses in the CNS in vivo. In particular, we were interested in responses to the infiltration of nonresident activated T helper 17 (Th17) cells into the CNS using EAE as a disease model (Aktas et al., 2005; Paterka et al., 2016; Siffrin et al., 2010). CNS myeloid cells and Th17 cells interacted frequently and intensely during the initial phase of the disease, underlining a dynamic cell communication system in inflammatory lesions. Unexpectedly, myeloid cells were able to engulf viable and motile T cells in the CNS, a phenomenon that occurred more frequently after inflammatory myeloid cell activation and was regulated by lectin...
expression on myeloid cells and T cell activity-dependent N-acetyl-D-glucosamine (GlcNAc) exposure. This engulfment led to subsequent cell death in the majority of the engulfed T cells, and EAE was ameliorated when this pathway was enhanced by intrathecal GlcNAc administration.

**Results**

**CX3CR1+ cells engulf invading and pathogenic Th17 cells in EAE**

To analyze how CX3CR1+ myeloid cells (microglia and infiltrating macrophages/dendritic cells) react toward a T cell-driven CNS inflammation, we visualized the dynamic interactions of both cell types using in vivo two-photon laser scanning microscopy in the upper brainstem, an area prone to active inflammation in the inflamed CNS.

CNS-resident myeloid cells are responsible for engulfment processes in organotypic slice cultures

In vivo, peripheral CX3CR1+GFP myeloid cells are able to infiltrate the CNS under EAE conditions (Ajami et al., 2018; Geissmann et al., 2003; Goldmann et al., 2013; Jung et al., 2000) and may thus contribute to the observed interactions between CX3CR1+ cells and T cells. In the CNS of EAE-diseased mice, infiltrating pathogenic T cells were found to intensely interact with CX3CR1+GFP cells, revealing multiple types of interactions.Remarkably, in-depth analyses of the surface-reconstructed images demonstrated the capability of CX3CR1+ cells to fully engulf CNS-infiltrating pathogenic Th17 cells (Fig. 1, A–D; and Video 1). The engulfed T cells either stayed engulfed for up to several hours or were able to escape this engulfment. These observations thus suggest an active, albeit not completely successful attempt of the CNS-resident CX3CR1+ myeloid cells to remove invading T cells in the inflamed CNS.

**Engulfed living T cells either escape or undergo cell death**

To determine the viability of engulfed T cells, we stained for Caspase-3/7 activity before live two-photon imaging, allowing the distinction between viable and apoptotic cells. We found that living T cells (CFP-positive), early apoptotic T cells (low Caspase-3/7 expression and diminished CFP-reporter signal), and apoptotic T cells became engulfed by myeloid cells (Fig. 3 A). Long-term imaging of stably engulfed T cells typically showed a gradual reduction of the CFP fluorescence signal, indicating the degradation of these T cells and their genetically encoded reporter signal (CFP) within myeloid cells (Fig. 3 B and Video 8). By contrast, escaping T cells did not lose their fluorescence signal, and their viability was indicated by an active migration in the tissue after escape (Fig. 3 C and Video 9). Escaped T cells showed comparable T cell motilities before and after engulfment, suggesting an unchanged activation status (Fig. 3 D). Importantly, the percentage of stably engulfed T cells was significantly higher compared with the percentage of T cells that were able to escape from engulfment (Fig. 3 E).

**Engulfment of viable cells is not dependent on PS exposure**

As a reversible expression of “eat-me” signals (such as phosphatidylserine [PS]) on stressed (but otherwise viable) cells might already lead to recognition by phagocytic cells (Brown and Neher, 2014), we analyzed the PS expression of Th17 cells and indeed identified a higher PS expression on activated Th17 cells compared with nonactivated Th17 cells (Fig. 4 A). Consistent with the known phagocytic abilities of CNS myeloid cells (Sierra et al., 2013), we observed Caspase-3/7+ cells to be more frequently engulfed by myeloid cells than viable T cells (Fig. 4, B and F). Nonetheless, a detailed quantification showed that the EP frequency of viable T cells was as high as 13% of all interactions between living non-Caspase-3/7+ T cells and myeloid cells (Fig. 4 C). Treatment with high concentrations of soluble Annexin V, in order to mask PS signals (Lu et al., 2011), significantly reduced both the total EPs and the engulfment rate of apoptotic cells but had no effect on EPs and engulfment rate of living T cells (Fig. 4, D–F). Annexin V treatment also did not impact T cell motility parameters (Fig. 4 G). These data suggest that despite PS expression on living Th17 cells, the engulfment of viable T cells is mediated by a PS-independent mechanism, in contrast to the PS-dependent phagocytosis of apoptotic cells.

**Engulfment of viable cells is dependent on GlcNAc exposure on activated T cells**

To identify the receptor on T cells responsible for the engulfment of living cells, we analyzed other eat-me and “don’t eat-me” signals. Of note, expression of the don’t eat-me signal CD47 was not detected on Th17 cells (data not shown). Furthermore, the recognition of eat-me signals via vitronectin receptors has been discussed to promote the phagocytosis of apoptotic cells (Savill, 1997). We observed no influence on the engulfment of Myeloid cells capture living CNS-invading T cells

Wasser et al.

Myeloid cells capture living CNS-invading T cells
living cells, as inhibition of this pathway by treatment with the tetrapeptide RGDS (Arg-Gly-Asp-Ser) did not reduce the percentages of engulfment (Fig. 5 A). By contrast, GlcNAc, which has been discussed previously to promote the binding of apoptotic thymocytes to macrophages (Duvall et al., 1985), had a significant impact on T cell–myeloid cell interaction (Fig. 5, B–G). To identify GlcNAc expression on T cells, we used fluorescently labeled wheat germ agglutinin (WGA), which has been used to measure GlcNAc levels (Notter and Leary, 1987). Stimulated Th17 cells showed an increased binding of WGA compared with unstimulated cells and the presence of soluble GlcNAc in the staining solution could interfere with the binding of WGA, leading to lower fluorescence (Fig. 5 B). Using different experimental protocols, addition of soluble GlcNAc can be used either to competitively inhibit endogenous GlcNAc–lectin interaction or to enhance GlcNAc exposure on T cells. Importantly, the competitive addition of 20 mM GlcNAc in the 24 h co-culture of organotypic hippocampal slices with Th17 cells significantly decreased the engulfment of T cells by myeloid cells (Fig. 5, C and D). On the other hand, pretreatment of T cells with 20 mM GlcNAc increased GlcNAc exposure on activated T cells (Fig. 5 E). Increased GlcNAc exposure was stable for up to 5 h in co-cultures with organotypic hippocampal slices (Fig. 5 F). This enhanced GlcNAc exposure on T cells increased EPs and showed a strong trend toward more stable engulfments of viable T cells (Fig. 5, G and H). Engulfment of dying cells showed only a minor trend toward higher engulfment after GlcNAc pretreatment (Fig. S1 A).

Of note, myelin oligodendrocyte glycoprotein (MOG)–specific Th17 cells showed similar interactions with myeloid cells to MOG-specific Th2 cells, MOG-specific regulatory T (T reg) cells (Fig. 5, I and J), and polyclonal Th17 cells (Fig. 5 K). Furthermore, the presence of supplemented MOG peptide in co-cultures of hippocampal slice cultures with pathogenic Th17 cells in order to enhance antigen presentation processes did not enhance engulfment-related processes (Fig. 5 L). In addition, we analyzed the engulfment of Th17 cells in Th17 reporter mice, which switch from a red to a red and green double-positive phenotype upon IL-17 expression (Fig. 5 M). With this analysis, we were able to confirm that not only IL-17–producing Th17 cells became engulfed. We conclude that engulfment processes are regulated by the state of T cell activity rather than by T cell subtype or antigen specificity.

Since our observations pointed toward a specific role of T cell activation status as a prerequisite for engulfment by myeloid cells, we addressed the role of the PI3K/Akt pathway as a major regulator of T cell activity. Indeed, inhibition of this pathway by Wortmannin reduced WGA binding on activated T cells (Fig. 6 A) and abrogated Th17 cell effector functions, as could be shown by a significantly lower IL-17 expression in unstimulated and stimulated Th17 cells (Fig. 6 B), and down-regulated the expression of activation markers (CD44 and CD25) in stimulated Th17 cells in vitro (Fig. 6, C and D). Importantly, PI3K inhibition had no influence on migration properties of Th17 cells in organotypic slices (Fig. 6 E) and had no obvious impact on myeloid cell structure or morphology (Fig. 6 F). Treating the hippocampal slice–T cell co-cultures with Wortmannin, we observed a significantly lower engulfment rate compared with control

![Figure 1](image-url)
conditions, indicating an impact of T cell activation status on engulfment probabilities (Fig. 6, F–H). Of note, the activity of T cells not only controls T cell engulfment but might also impact their capacity to escape. While escapes were identified to be low under control conditions (Fig. 3 E), with Wortmannin treatment, there was a nonsignificant trend from stable engulfments to escapes (Fig. 6, I and J).

Amplification of myeloid cell activation enhances engulfment of living T cells

We next aimed to identify the corresponding binding partner on myeloid cells. CD206 has been described previously as a receptor for GlcNAc (Taylor et al., 1992, 2005). Here, we showed that alternative activation of myeloid cells led to a significant up-regulation of lectin expression, which was stronger than the up-regulation of lectin in classically activated myeloid cells (Fig. 7 A). Alternative activation of myeloid cells in the CNS has been described to be important in amelioration of EAE disease (Franco and Fernández-Suárez, 2015; Jiang et al., 2014). Up-regulation of CD206 expression was accompanied by increased engulfment frequencies (Fig. 7 B), indicating that a marked increase in myeloid cell activity supports increased engulfment processes and thus clearance of invading T cells in neuro-inflammation. Interestingly, CD206 expression was reported in published single-cell RNA-sequencing data to be expressed by microglia-like cells in health and EAE disease (Falcão et al., 2018), and in agreement, we detected a minor fraction of CD45loCD11b−Ly6G/G− myeloid cells expressing CD206 in EAE (Fig. 7, C–E). In addition, we observed CD206 expression in CD45hiCD11b+ myeloid cells (Fig. 7, C and E), previously discussed to represent CNS-infiltrating myeloid cells or activated microglia (Ponomarev et al., 2007). Our results emphasize low CD206 expression in EAE conditions in different myeloid cell populations. Of note, these low percentages of CD206 expression (∼2–4%) are comparable to percentages of full acute engulfments (Fig. 4 C). We hypothesize a temporary up-regulation of CD206 during the engulfment process. Indeed, we detected CD206 predominantly on T cell-engulfing CX3CR1+ cells in the spinal cord of EAE diseased mice (Fig. 7 F). We observed CD206 expression in organotypic slices after alternative activation and in EAE diseased mice not only in perivascular myeloid cells but also in the parenchyma (Fig. S1, B and C). Parenchymal CD206 expression in EAE might be based on either CNS-infiltrating myeloid cells or up-regulation of CD206 by microglia.
Antigen recognition (Cahalan and Parker, 2008), were altered over the disease course compared with nonengulfed myeloid cells (Fig. 7, G and H), confirming that especially active myeloid cells are capable of engulfing T cells.

**Engulfment is increased during the initial phase of EAE**

To challenge the in vivo relevance of our findings, we investigated the relationship between T cell engulfment and EAE disease course. In vivo, CNS-invading CD4+ T cells during EAE showed higher GlcNAc expression compared with peripheral CD4+ T cells, underlining the importance of this pathway for activated T cells in the CNS (Fig. 8 A). Importantly, interaction characteristics between CX3CR1+ cells and T cells were not static but depended on clinical disease severity (Fig. 8, B–E; and Video 10). The engulfment of pathogenic T cells was significantly higher in EAE mice exhibiting a clinical score <2 (Fig. 8, B and C). Stable contacts increased in correlation with a higher disease score, while temporary contacts were more abundant at lower disease scores (Fig. 8, B and C). Accordingly, T cell motilities, known to inversely correlate with antigen recognition (Cahalan and Parker, 2008), were altered over the disease course. T cell motility parameters (e.g., “T cell speed” and “T cell displacement rate”) were higher during the early disease course compared with EAE mice exhibiting a clinical score >2 (Fig. 8, D and E). Strikingly, the interactions of CX3CR1+ cells and T cells in organotypic hippocampal slice cultures showed overall characteristics comparable to early CNS inflammation in vivo (Fig. 8, C–E). The engulfment frequencies of T cells (4%) as well as the frequencies of stable (20%) and temporary contacts (70–80%) among the interactions were similar in vivo and in organotypic hippocampal slice cultures over an observation period of 20 min (Fig. 8 C). Likewise, T cells that invade the CNS in early EAE disease showed T cell motility parameters comparable to T cells invading the hippocampal slices (Fig. 8, D and E). It should be noted that in contrast to this apparently low relative number of T cell engulfments, they can be considered as biologically significant. The percentage of the engulfment of viable cells reflects a snapshot of the processes taking place in the CNS and does not provide the total cell count of removed cells. Altogether, our findings point toward T cell engulfment by CX3CR1+ myeloid cells as an early defense mechanism upon initial T cell invasion that is overcome later on in a full-blown inflammatory environment.

**Pharmacological increase of the engulfment pathway in mice ameliorates EAE disease severity**

As increased GlcNAc expression in T cells correlates with engulfment, we aimed next to therapeutically ameliorate EAE by enhancing GlcNAc exposure on T cells in the CNS in vivo. In vitro, GlcNAc pretreatment during T cell stimulation was efficient at enhancing GlcNAc exposure on T cells (Fig. 5 D). Upon intrathecal treatment with 1 µmol GlcNAc in vivo, we indeed found an amelioration of disease severity (Fig. 8 F). This effect was enhanced when GlcNAc treatment was started before disease onset (Fig. 8 G), confirming a high relevance during the early phase of the disease. Analysis of GlcNAc expression on...
T cells isolated from treated mice confirmed a higher expression of GlcNAc on T cells compared with the control group (Fig. 8 H). By contrast, intrathecal GlcNAc treatment had no effect on GlcNAc exposure on peripheral T cells (Fig. 8 H). Thus, GlcNAc treatment specifically affected T cells, which infiltrated into the CNS, excluding peripheral immune effects and demonstrates that T cell engulfment by myeloid cells can be therapeutically exploited in vivo. GlcNAc treatment significantly reduced the T cell count within the CNS, indicating a removal of T cells by engulfing myeloid cells (Fig. 8, I and J).
Engulfment of viable cells is dependent on GlcNAc exposure on activated T cells. (A, C, D, and G–L) Co-cultures of organotypic hippocampal slices from CX3CR1<sup>GFP</sup> pups with pathogenic CFP-labeled T cells were stained with Image-iT LIVE Red Caspase-3 and -7 Detection Kit to identify apoptotic cells and imaged over a time period of 20 min. Only non-apoptotic T cells were considered for further analyses. (A) 24-h co-cultures of organotypic hippocampal slices with Th17 cells were treated with RGDS (1 mM). The pie chart comparing the distribution (%) of interaction modes between myeloid cells and viable B6.2D2.CFP.Th17 cells (n = 5 organotypic slices from four different experiments) and contingency bar reflecting the respective distribution of engulfment, escape, and engulfment attempt are shown. (B) WGA binding to Th17-skewed cells was assessed in the presence or absence of GlcNAc (200 mM) in the staining solution using flow cytometric analysis. Representative histograms (left panel) and mean (± SEM) of the mean fluorescence intensity (MFI, right panel) are shown (n = 5 Th17 cultures from two different experiments). (C) 24-h co-cultures of organotypic hippocampal slices with Th17 cells were treated with GlcNAc (20 mM, competitive GlcNAc). The pie chart shows the distribution (%) of interaction modes between myeloid cells and viable Th17 cells (n = 4 organotypic slices from three different experiments; upper panel), and contingency bar reflects the respective distribution of engulfment and engulfment attempt (lower panel). (D) Mean (± SEM) percentages of engulfment rate among the interactions in GlcNAc (20 mM, competitive GlcNAc) treated co-cultures (n = 4 organotypic slices from three different experiments) normalized to the untreated control group (n = 4 organotypic slices from 3 different experiments). (E–H) Naïve T cells were skewed into Th17 cells in the presence or absence of GlcNAc (20 mM). At day 3 of culture, cells were further stimulated with antiICD3 and antiICD28 with/without GlcNAc (20 mM). (E) Cells were washed to remove GlcNAc in the supernatant, and WGA binding was assessed (n = 7 Th17 cell cultures from seven different experiments). (F–H) Control Th17-skewed cells and GlcNAc-pretreated (GlcNAc enhanced) Th17-skewed cells were co-cultured with organotypic hippocampal slices. (F) WGA-binding to T cells was assessed after 5 h of co-culture (n = 7 Th17 co-cultures from three different experiments). (G) The pie chart shows the distribution (%) of interaction modes between myeloid cells and viable GlcNAc-pretreated Th17 cells (n = 7 organotypic slices from three different experiments; upper panel), and contingency bar reflects the respective distribution of engulfments and engulfment attempt (lower panel). (H) Mean (± SEM) percentages of engulfment rate among the interactions of GlcNAc-pretreated Th17-skewed cells in organotypic hippocampal slices (n = 7 organotypic slices from three different experiments) normalized to the GlcNAc-ununtreated control group (n = 7 organotypic slices from three different experiments). (I–L) Pie charts (upper panel) show the distribution (%) of interaction modes between myeloid cells and viable B6.2D2.CFP.Th2 cells (n = 3 organotypic slices from three different experiments; I), viable B6.2D2.CFP.Treg cells (n = 4 organotypic slices from four different experiments; J), polyclonal B6.CFP.Th17 cells (Th17 BL/6; n = 6 organotypic slices from 2 different experiments; K), or viable B6.2D2.CFPTh17 cells in co-cultures treated with MOG (100 µg/ml; n = 3 organotypic slices from two different experiments; L), and contingency bars reflect the respective distribution of engulfment, escape, and attempt to engulf (lower panel). (M) Representative visualization of 24-h co-cultures of organotypic hippocampal slices from CX3CR1<sup>GFP</sup> pups with pathogenic B6.2D2.RFP.Th17.GFP reporter cells, in which IL-17-producing cells are GFP<sup>+RFP</sup> double positive (orange), while non-IL-17-producing cells are RFP<sup>+</sup> single positive (red; representative of n = 3 co-cultures from three different experiments; left panel). Scale bar = 30 µm. A myeloid cell (green) engulfing at the same time an IL-17-producing cell (marked with an asterisk) and a non-IL-17-producing cell (marked with a number sign) was magnified (middle panel), and red signal was hidden to visualize the green signal of the IL-17-producing T cell (right panel). Scale bar = 15 µm. Statistical analyses were performed using two-sided Student's t tests. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ns, not significant.
Nimmerjahn et al., 2005). Although it is well known that thereby mediating inflammation and neuronal degeneration pathogenic Th17 cells invade the CNS during MS and EAE, hippocampal slices from CX3CR1GFP pups with pathogenic B6.2D2.CFP.Th17 cells were stained with Image-iT LIVE Red Caspase-3 and -7 Detection Kit to context (Giles et al., 2018; Heneka et al., 2014; Jolivel et al., 2015; and tissue homeostatic functions, presumably depending on the continuously scan their environment, mediating both detrimental lesions in EAE and MS, where myeloid cells of the brain con-
first-defense-like process, as is known for their phagocytosis properties toward CNS-invading pathogens. In vivo, CNS-infiltrating myeloid cells may contribute to this T cell capture. However, microglia, which are present from the onset of the disease, are capable of forming the first line of defense against invading lymphocytes. Optimizing this first defense process may therefore represent a valuable therapeutic opportunity in neuroinflammatory diseases. Indeed, we were able to show that intrathecal therapeutic enhancement of GlcNAc exposure on T cells was effective at ameliorating EAE disease symptoms. Similar beneficial effects were reported previously after oral GlcNAc administration in mild EAE (Grigorian et al., 2011), focusing so far only on peripheral effects and without providing mechanistic insights. Here, by applying GlcNAc intrathecally, we excluded peripheral effects and demonstrated that GlcNAc-mediated T cell reduction in the CNS causes disease amelioration. Our findings point toward T cell engulfment by CX3CR1+ myeloid cells as an early defense mechanism upon initial T cell invasion.
Figure 8. Enhanced GlcNAc exposure on T cell ameliorates EAE during the early disease course. (A) Flow cytometric analysis of WGA binding on living CD4+ T cells in the CNS and spleen of EAE diseased mice in actively immunized C57BL/6 mice normalized to splenic T cells (n = 11 mice from two different experiments). (B–E) EAE was induced in CX3CR1.GFPxRag-/- and CX3CR1.GFPxRag+/- mice (myeloid cells green) via the passive transfer of B6.2D2.RFP.Th17 cells (red). Interactions between myeloid cells and T cells as well as T cell motility parameters were imaged in vivo with intravital two-photon imaging and analyzed during different disease states. For hippocampal slice culture experiments, co-cultures of hippocampal slices from CX3CR1.GFP pups with pathogenic B6.2D2.CFP.Th17 cells were stained with Image-iT LIVE Red Caspase-3 and -7 Detection Kit to identify apoptotic cells, and only living T cells were included for further analysis. (B) Dots represent the percentages of engulfments, temporary contacts and stable contacts among the analyzed T cells in a randomly defined area of the CNS (brainstem) of EAE diseased mice in dependency on disease score. P values indicate whether the slope of linear regression is significantly nonzero (n = 9 mice from two different experiments). (C) Means (± SEM) of percentages of engulfments, temporary contacts, and stable contacts among the detected interactions or the detectable T cell count per region of interest were compared between mice exhibiting a score <2 (n = 3 mice from two different experiments), mice exhibiting a score ≥2 (n = 6 mice from two different experiments), and hippocampal slice cultures (n = 12 organotypic hippocampal slice cultures from eight different experiments). Statistical analysis was performed using one-sided Mann–Whitney U test. *, P < 0.05; ns, not significant. (D) Mean (± SEM) of T cell motility parameters in the CNS of EAE diseased mice in dependency on disease score. P values indicate whether the slope of linear regression is significantly nonzero (n = 9 mice, 30 T cells per mouse from two different experiments). (E) Normalization of all T cell tracks to a common starting point in a representative experiment of a hippocampal slice co-culture comparing a mouse exhibiting a clinical score of 1.5 and a mouse exhibiting a clinical score of 4.5. Each line represents one individual T cell. Representative of 9 mice from two different experiments. (F–J) EAE was induced actively in C57BL/6 mice and 5 µl
invasion that is overcome later on in a full-blown inflammatory environment. The reasons for this shift are most likely functional changes due to microenvironment alterations in the course of the disease, since we were able to show that alternatively activated (versus classically activated) myeloid cells possess markedly increased engulfment. Importantly, although T cell removal from the CNS is beneficial during a T cell-mediated neuroinflammatory disease, this T cell removal could be harmful, such as in pathogen-mediated diseases, in which T cells are required to promote defense mechanisms in the CNS. Future studies are needed to analyze these engulfment processes in other disease contexts.

According to in vitro observations, cell-in-cell structures have, to date, been reported mostly for tumor cells (Yang and Li, 2012), and the use of different, mostly vaguely defined terms (e.g., entosis, [suicide] emperipolesis, phagoptosis, cytophagocytosis, or cannibalism) indicates that both the physiological significance and underlying pathways of these so-called cell-in-cell structures are still poorly understood (Brown and Neher, 2014; Overholtzer and Brugge, 2008). In our study, in vivo two-photon imaging as well as long-term follow up of slice cultures revealed full engulfment of viable T cells. Strikingly, some engulfed T cells were able to freely move through the glial cytoplasm, often even followed by escape from glial entrapment. In very few earlier neuropathologic reports based on occasional histological observations, cell-in-cell formation has been mentioned for lymphocytes in neurons (Hughes et al., 1968; Ng and Ling, 1999) and for oligodendrocytes in astrocytes (Shintaku and Yutani, 2004; Wu and Raine, 1992). The presently demonstrated engulfment and residence of T cells within myeloid cells may be reminiscent of entosis, a form of cell cannibalism (Krishna and Overholtzer, 2016). However, entotic cells, in contrast to Th17 cells, are reported to not expose the eat-me signal PS, which may explain why they can be rescued from cell death after engulfment. Furthermore, entosis has so far mostly been assigned to homotypic cell-in-cell interactions (i.e., among the same cell types) and in almost all reports is restricted to tumor cells (both in culture and in vivo). A potential role of entosis in a noncancerous context has not been reported so far.

In our study, inhibition of T cell activity or prevention of myeloid cell binding to GlcNAc on T cells resulted in inhibition of T cell engulfment by myeloid cells. GlcNAc has been discussed previously to promote the binding of apoptotic thymocytes to macrophages and their subsequent phagocytosis (Duvall et al., 1985). Of note, we were able to show that the engulfment of viable T cells was not driven by PS exposure, clearly identifying this process to be distinct from classic PS-mediated phagocytosis. GlcNAc, which we showed here to be responsible for the engulfment of living T cells, is also expressed by bacteria, where it serves as a target molecule for pattern recognition receptors on phagocytes, resulting in the phagocytosis and removal of harmful CNS intruders (Nadesalingam et al., 2005; Zhang et al., 2015). Myeloid cell activation in organotypic slices led to an up-regulation of the lectin CD206, which is a receptor for GlcNAc (Taylor et al., 1992, 2005), and was accompanied by increased engulfment processes, thus indicating that an increase of myeloid cell activity supports increased abilities to capture living T cells. Interestingly, CNS-invading CD4+ T cells during EAE showed higher GlcNAc expression than their peripheral counterparts, thus emphasizing the importance of the GlcNAc-lectin pathway within the CNS.

Enhancement of GlcNAc-dependent engulfment of living cells was sufficient to significantly decrease EAE severity, emphasizing the biological relevance of T cell engulfment by myeloid cells. Although the percentage of engulfments among myeloid cell–T cell interactions was in the single digits, it should be pointed out that these percentages represent only a short temporal snapshot of the engulfment processes, especially compared with the large number of temporal screening contacts taking place in the CNS, thus likely underestimating the total cell count of removed cells.

Altogether, we identified highly dynamic interaction modes between myeloid cells and T cells within the brain, reflecting the attempt of myeloid cells to remove viable potentially pathogenic T cells from the CNS in a GlcNAc-mediated pathway, in order to preserve tissue homeostasis during early inflammation. The unique ability of some T cells to migrate through myeloid cells without subsequent cell death and instead escape from engulfment reveals a newly discovered mechanism highlighting their pathological potency in CNS inflammation and limitation of defense capacity in neuroinflammation. Overall, the findings reported here explore a novel, primarily protective role of myeloid cells in early neuroinflammation and may open new avenues for impacting neurological disorders such as MS.

**Materials and methods**

**Mice**

All mice were kept under specific pathogen-free conditions. B6.CX3CR1+/GFP and B6.CX3CR1+/GFPproGFP/GFP_Rag2−/−/−_mice were used for in vivo imaging as indicated. B6.CX3CR1GFP/GFP mice were used for ex vivo imaging of myeloid cells, as genotyping of pups from heterozygous breeding is not feasible before slice preparation (Jung et al., 2000). We confirmed similar
interactions in heterozygous organotypic slices, gained by crossing B6.CX3CR1\textsuperscript{GFP/GFP} mice with C57BL/6 mice (Fig. S2). Throughout the article, CX3CR1\textsuperscript{GFP} is used for simplicity and better readability. B6.2D2.RFP mice and B6.2D2.CFP mice, in which all CD4\textsuperscript+ T cells are MOG\textsubscript{35-55} specific, were obtained by crossbreeding B6.acRFP and B6.CFP with B6.2D2 mice (Bettelli et al., 2003; Hadjantonakis et al., 2002; Luche et al., 2007). These mice were used for T cell cultures. All animal experiments were approved by local authorities (Landesuntersuchungsamt Rheinlandpfalz) and performed in accordance with German Animal Protection Laws.

**EAE**

For passive-transfer EAE, 10–40 × 10\textsuperscript{6} B6.2D2.RFP cells were injected intravenously into recipient mice (B6.CX3CR1\textsuperscript{+/GFP}, B6.CX3CR1\textsuperscript{+/GFPerGFP/GFP}.Rag\textsuperscript{-/-}, or+/+) and pertussis toxin (400 ng/mouse) was injected on the day of transfer and day 2 following transfer. If Rag\textsuperscript{-/-} mice were used, no pertussis toxin was injected. For active EAE, C57BL/6 mice were immunized subcutaneously with 200 \mu g myelin oligodendrocyte protein MOG\textsubscript{35-55} in 200 \mu l CFA emulsion. Additionally, C57BL/6 mice received 200 ng pertussis toxin (Hook; Hook Laboratories) intraperitoneally at the time of immunization and 24 h later. For intrathecal treatment with GlcNAc (5 \mu l of 200 mM GlcNAc dissolved in PBS) and PBS (5 \mu l), mice were shortly anesthetized using isoflurane. The lumbar region was shaved and disinfected. Application was performed into the spinal canal between L5 and L6 every second day seven times. Mice were scored for clinical symptoms daily, and signs of EAE were translated into clinical score as follows: 0, no detectable signs of EAE; 0.5, tail weakness; 1, complete tail paralysis; 2, partial hindlimb paralysis or ataxia; 2.5, unilateral complete hindlimb paralysis; 3, complete bilateral hindlimb paralysis or severe ataxia; 3.5, complete hindlimb paralysis and partial forelimb paralysis; 4, total paralysis of forelimbs and hindlimbs (mice with a score >3.5 to be killed); and 5, death.

**T cell culture**

Procedures were similar to those reported previously (Siffrin et al., 2009, 2010). In brief, cells from B6.2D2.CFP or B6.2D2.RFP mice were isolated from spleen and lymph nodes (axial and cranial), and magnetic bead–based cell sorts (Miltenyi) were performed. Naive CD4\textsuperscript+CD62L\textsuperscript{hi} cells were stimulated with 2 \mu g/ml \alphaCD3 (1452-CII; BD Biosciences) in the presence of irradiated CD90\textsuperscript+ depleted C57BL/6 splenocytes, and culture medium was enriched with the following cytokines: for Th17 differentiation, 3 ng/ml TGF-\beta, 20 ng/ml IL-23, and 20 ng/ml IL-6 (all \&D Systems); for Th2 differentiation, 10 ng/ml IL-4 (\&D Systems), 10 \mu g/ml \alphaIFN\gamma (XMG12.1; BioXcell), and 10 \mu g/ml \alphaIL-12 (C17.8; BioXcell); for T reg cell differentiation, 3 ng/ml TGF-\beta (\&D Systems), 10 \mu g/ml \alphaIL-12 (C17.8; BioXcell), and 10 \mu g/ml \alphaIFN\gamma (XMG12.1; BioXcell). After 3 d of culture, cells were split, and T cell culture medium was enriched with 50 U/ml IL-2 and 10 ng/ml IL-23 for Th17 or 100 U/ml IL-2 and 10 ng/ml IL-4 (all \&D Systems) for Th2 cells. T reg cells were split and fresh IL-2 (100 U/ml) was added after 2 and 4 d of culture. Th17 cells were either used for co-cultures with hippocampal slices after 5 d of culture or restimulated after 1 week with 2 \mu g/ml anti-CD3 in the presence of irradiated CD90\textsuperscript+–depleted C57BL/6 splenocytes and treated with 0.75 ng/ml TGF-\beta, 20 ng/ml IL-23, and 10 ng/ml IL-6. If restimulated, cells were used 3 d after restimulation for further experiments (co-culture with organotypic hippocampal slices or induction of passive-transfer EAE).

For GlcNAc enhancement experiments, naive CD4\textsuperscript+CD62L\textsuperscript{hi} T cells were treated on day 0 of culture with 20 mM GlcNAc. For Th17 differentiation, 3 ng/ml TGF-\beta, 20 ng/ml IL-23, and 20 ng/ml IL-6 (all \&D Systems) were additionally added. On day 3, T cells were harvested and transferred on an anti-CD3/anti-CD28–precoated 48-well plate, and 20 mM GlcNAc was added to the GlcNAc-pretreated group. WGA binding was assessed on day 5.

**Organotypic hippocampal slice culture**

For organotypic hippocampal slice cultures, postnatal day 3–6 B6.CX3CR1\textsuperscript{GFP/GFP} pups were decapitated, and brains were rapidly removed and transferred to ice-cold preparation medium (MEM) containing 2 mM glutamine, adjusted to pH 7.35. Both hemispheres were separated and the meninges removed, and hippocampi were isolated with entorhinal cortex attached and coronally cut (300 \mu m) using a McILWAIN tissue chopper. Intact hippocampal slices were transferred onto Millicell cell culture inserts (Merck Millipore) positioned in each well of a 6-well plate (Greiner) containing 1.2 ml culture medium (48% MEM, 24% basal medium Eagle, 24% heat-inactivated horse serum, 2 mM glutamine, and 0.6% glucose, pH 7.2) and incubated at 37\degree C, 95%/5% O\textsubscript{2}/CO\textsubscript{2}. The medium was changed 24 h after plating and then every other day. After 7–14 d in vitro, hippocampal slices were used for experiments.

For co-cultures, either 10\textsuperscript{5} Th17 cells or 10\textsuperscript{5} Th2 cells were added in a volume of 10 \mu l directly on top of the hippocampal slices and incubated at 37\degree C, 95%/5% O\textsubscript{2}/CO\textsubscript{2} for 24 h or, in the case of GlcNAc enhancement experiments, 4 h. At the time the T cells were added, slices were also optionally treated with Annexin V (10 \mu g/ml), Wortmannin (2.5–5 \mu M), or MOG\textsubscript{35-55} peptide (100 \mu g/ml; Genscript) as indicated. For LPS (classic activation) and LPS/dexamethasone/IL-4 preactivation (alternative activation), organotypic slices were treated with LPS (1 \mu g/ml) or LPS (1 \mu g/ml), dexamethasone (5 × 10\textsuperscript{-7} M), and IL-4 (10 ng/ml) for 24 h before T cell addition. LPS, dexamethasone, and IL-4 were kept in the medium during the co-culture. To distinguish viable and apoptotic T cells, co-cultures were stained with the Image-IT LIVE Red Caspase-3 and -7 Detection Kit (Thermo Fisher Scientific) 30 to 60 min before imaging. Competitive GlcNAc (20 mM) treatment was performed 24 h and additionally 20 min before T cell addition. GlcNAc was kept in the medium during the co-culture. For GlcNAc enhancement experiments, GlcNAc-pretreated T cells (see T cell culture) were added. RGDS (Arg-Gly-Asp-Ser) treatment (1 mM) was performed 24 h and additionally 20 min before T cell addition. RGDS was kept in the medium during the co-culture.

**Intravital two-photon imaging**

Operation procedures and two-photon laser scanning microscopy were performed as described previously (Herz et al., 2010; Luchtman et al., 2016; Siffrin et al., 2009, 2010). Mice were
anesthetized using 1.5% isoflurane in oxygen/nitrous oxide (1:2) and then continuously respirated with a facemask. Mice were transferred to a custom-built operation and microscopy table and fixed in a hanging position. The brainstem was exposed by the removal of the musculature above the dorsal neck area and removal of the dura mater using a microscope. The head was inclined for access to deeper brainstem regions. A sterile agarose patch (0.5% in 0.9% NaCl solution) was installed on the exposed brain surface. During surgery and microscopy, body temperature was maintained at 37°C. Anesthesia depth was controlled by continuous CO2 measurements of exhaled gas and recorded with a CI-240 Microcapnograph (Columbus Instruments). Imaging was performed in the brainstem. We applied dual near-infrared and infrared excitation of the live brainstem at 850 nm by an automatically tunable Ti/Sa laser (Mai Tai HP; Spectra Physics) and 1110 nm by an optical parametric oscillator (APE) pumped by the Ti/Sa laser. Using a TriMScope I multiphoton system from LaVision Biotec, volumes of 300 µm × 300 µm × 72 µm were acquired over time and exported as TIFFs for 3D analyses.

For the ex vivo slice experiments, slices were transferred onto the stage of a Leica TCS-MP5 multi-photon system with heated, humidified, and gassed (95% O2/5% CO2) chamber within a Ludin enclosure and volumes of 300 µm × 300 µm at variable depth (max 80 µm) acquired with a Leica HCX IRAPO L 25×/0.95 W objective at 1000 nm to visualize the CX3CR1GFP myeloid cells and B6.2d2.RFP.Th17 cells. For imaging CX3CR1GFP myeloid cells and B6.2d2.CFP.Th17 cells that were alive (CFP) or stained with Image-iT LIVE Red Caspase-3 and -7, as well as WGA (Thermo Fisher Scientific), dead cells were identified using either propidium iodide or fixable viable stain-V450 (BD Biosciences). For intracellular staining, cells were fixated with 2% paraformaldehyde after 4 h of stimulation in the presence of Brefeldin A. IL-17 expression was determined using fluorescently labeled αIL-17 (eBioscience). PS exposure was determined via Annexin V staining kit from eBioscience. For flow cytometric analysis of T cells and myeloid cells ex vivo, CNS and spleen tissue was prepared as described previously (Wasser et al., 2016). Cells were stained with fluorescently labeled αCD4 (RM4-5; BD Biosciences), WGA (Thermo Fisher Scientific), αCD11b (M1/70; Thermo Fisher Scientific), αCD45 (30-F11; Thermo Fisher Scientific), αLy6C/G (RB6-8C5; BD Biosciences), αCD206 (C068C2; BioLegend), and IgG2a-BV421 (RTK2758; BioLegend) as indicated. For flow cytometric analysis of myeloid cells in organotypic hippocampal slices (≥7 d in culture), organotypic slices from CX3CR1GFP pups were digested with 10 mg/ml Collagenase/Dispase (Sigma), 10 mg/ml Collagenase (Sigma), and 200 U/ml DNase (Roche). After incubation for 30 min at 37°C, the tissue was removed from the cell culture inserts and put through a mesh (100 µm), and mononuclear cells were separated by 30/70 Percoll; i.e., 70% Percoll (Sigma) in PBS (Sigma) was overlayed with cells resuspended in 30% Percoll in IMDM solution (Gibco) and then centrifuged at 750 g for 20 min. Mononuclear cells were collected at the interface and stained as indicated with fluorescently labeled αCD45 (30-F11; Thermo Fisher Scientific), αCD11b (M1/70, Thermo Fisher Scientific), αCD206 (C068C2; BioLegend), and IgG2a-BV421 (RTK2758; BioLegend). All gating strategies are provided in Fig. S3.

Analysis of two-photon imaging results
TIFF and LIF files were imported into 3D reconstruction software Imaris (Bitplane, version 8.1.2). T cell tracks were created using the tracking tool implemented in Imaris software and manually corrected. Contact types and durations were determined manually with 3D rotation and surface analysis to verify contacts. Temporary contacts were defined as contacts lasting <10 min and stable contacts as contacts lasting >10 min. Engagement-related processes (EPs) were subdivided into engulfment (T cell is fully surrounded by the myeloid cell), escape (T cell moves through the myeloid cell and escapes), and clear attempt to engulf (myeloid cell actively prolongs processes to engulf T cells). On average, 20–100 T cells were analyzed per region of interest (maximum 300 µm × 300 µm × 80 µm) in one organotypic hippocampal slice. For T cell motility analysis, a maximum of 30 cells reflecting the mean velocities per area were included into further analyses. Noise reduction was achieved by the software’s “medium filter.”

Histological staining
Immunohistochemistry was performed with co-cultures of organotypic hippocampal slices from CX3CR1GFP mice with 2D2 T cells and spinal cord sections of EAE diseased CX3CR1.GFP animals. Iba-1/CX3CR1+ myeloid cells were stained with anti-mouse GFP (polyclonal-rabbit; Abcam; 270F3 mouse; Synaptic Systems) or anti-mouse Iba-1 (polyclonal-rabbit; Abcam) and anti-rabbit-AF488 (polyclonal; Thermo Fisher Scientific). CD4 was stained with anti-mouse CD4-AF647 (RM4-5 rat; BD Biosciences). MHC-II was stained with anti-mouse MHC-II (2G9 rat; BD Biosciences), CD206 with anti-mouse CD206 (C068C2 rat; BioLegend) and anti-rat-AF568 (polyclonal, Thermo Fisher Scientific), CD31 with anti-mouse CD31 (polyclonal-rabbit, Thermo Fisher Scientific) and anti-rabbit-AF647 (polyclonal; Thermo Fisher Scientific). The cell nucleus was stained with DAPI (Thermo Fisher Scientific).

Flow cytometry analysis
For flow cytometric analysis of T cells in vitro, Th17 cells were harvested and stained with fluorescently labeled αCD4 (RM4-5; BD Biosciences), WGA (Thermo Fisher Scientific), αCD11b (M1/70; Thermo Fisher Scientific), αCD45 (30-F11; Thermo Fisher Scientific), αLy6C/G (RB6-8C5; BD Biosciences), αCD206 (C068C2; BioLegend), and IgG2a-BV421 (RTK2758; BioLegend) as indicated. For flow cytometric analysis of myeloid cells in organotypic hippocampal slices (≥7 d in culture), organotypic slices from CX3CR1GFP pups were digested with 10 mg/ml Collagenase/Dispase (Sigma), 10 mg/ml Collagenase (Sigma), and 200 U/ml DNase (Roche). After incubation for 30 min at 37°C, the tissue was removed from the cell culture inserts and put through a mesh (100 µm), and mononuclear cells were separated by 30/70 Percoll; i.e., 70% Percoll (Sigma) in PBS (Sigma) was overlayed with cells resuspended in 30% Percoll in IMDM solution (Gibco) and then centrifuged at 750 g for 20 min. Mononuclear cells were collected at the interface and stained as indicated with fluorescently labeled αCD45 (30-F11; Thermo Fisher Scientific), αCD11b (M1/70, Thermo Fisher Scientific), αCD206 (C068C2; BioLegend), and IgG2a-BV421 (RTK2758; BioLegend). All gating strategies are provided in Fig. S3.

Statistical analysis
All data were analyzed using GraphPad Prism 6 (GraphPad Software). Mean group differences were investigated by one-way ANOVA followed by Tukey’s multiple comparison test, Mann-Whitney U test, or independent-sample two-tailed t tests. The significance level was set at 0.05. Pie charts reflect the mean of the relative interaction distributions per analyzed video. If normalization was performed, each parameter was normalized to the mean of the control group from the respective slice preparation.

Online supplemental material
Fig. S1 shows the effect on GlcNAc exposure on the engulfment of apoptotic cells and the distribution of CD206 expression in...
organotypic hippocampal slices and the CNS of EAE diseased mice. Fig. S2 shows the interactions of T cells and myeloid cells in organotypic hippocampal slices. Fig. S3 shows all gating strategies for flow cytometric analysis shown in the main text. Video 1 shows surface-reconstructed intravital two-photon live imaging of an escaping T cell in vivo. Video 2 shows two-photon live imaging of Th17 cell–myeloid cell interactions ex vivo. Video 3 shows two-photon live imaging of a temporary contact ex vivo. Video 4 shows two-photon live imaging of a stable contact ex vivo. Video 5 shows two-photon live imaging of a stable engulfment ex vivo. Video 6 shows two-photon live imaging of an escape following walkthrough ex vivo. Video 7 shows two-photon live imaging of an engulfment attempt. Video 8 shows two-photon live imaging of a long-term engulfment. Video 9 shows two-photon live imaging of a long-term escape. Video 10 shows in vivo dynamics of T cell–myeloid cell interactions during early and chronic disease.

Acknowledgments

The authors thank Christin Liefländer, Heike Ehrengard, Ilse Graevenitz, Christine Oswald, and Jerome Birkenstock for excellent technical assistance and Cheryl Ernest and Rosalind Gilchrist for proofreading and editing the manuscript.

This study was supported by the Deutsche Forschungsgemeinschaft (grant SFB CRC-TR-128 to F. Zipp and S. Bittner, grants SFB 1080 and SFB CRC-1292 to F. Zipp, and grant TP BI1822/1-1 to S. Bittner) and the Hertie Foundation (mylab to S. Bittner, grants SFB 1080 and SFB CRC-1292 to F. Zipp, and grant M11.277814).

Disclosures: The authors declare no competing interests exist.

Submitted: 6 May 2019
Revised: 20 September 2019
Accepted: 17 January 2020

References

Ajami, B., N. Samusik, P. Wieghofer, P.P. Ho, A. Crotti, Z. Bjornson, M. Prinz, W.J. Fantl, G.P. Nelen, and L. Steinman. 2018. Single-cell mass cytometry reveals distinct populations of brain myeloid cells in mouse neuroinflammation and neurodegeneration models. Nat. Neurosci. 21:541-551. https://doi.org/10.1038/s41593-018-0100-x
Akras, O., A. Smorodchenko, S. Brocke, C. Infante-Duarte, U. Schulze Topphoff, J. Vogt, T. Prozorovski, S. Meier, V. Osmanova, E. Pohl, et al. 2005. Neuronal damage in autoimmune neuroinflammation mediated by the death ligand TRAIL. Neuron. 46:421–432. https://doi.org/10.1016/j.neuron.2005.03.018
Betelli, E., M. Pagany, H.L. Weiner, C. Linington, R.A. Sobel, and V.K. Kuchroo. 2003. Myelin oligodendrocyte glycoprotein-specific T cell receptor transgenic mice develop spontaneous autoimmune optic neuritis. J. Exp. Med. 197:1073–1081. https://doi.org/10.1084/jem.20021603
Brown, G.C., and J.J. Neher. 2014. Microglial phagocytosis of live neurons. Nat. Rev. Neurosci. 15:209–216. https://doi.org/10.1038/nrn3710
Cahalan, M.D., and I. Parker. 2008. Choreography of cell motility and interaction dynamics imaged by two-photon microscopy in lymphoid organs. Annu. Rev. Immunol. 26:585–626. https://doi.org/10.1146/annurev.immunol.24.021605.090620
Dendrou, C.A., L. Fugger, and M.A. Friese. 2015. Immunopathology of multiple sclerosis. Nat. Rev. Immunol. 15:545–558. https://doi.org/10.1038/nri3871
Duval, E., A.H. Wylle, and R.G. Morris. 1985. Macrophage recognition of cells undergoing programmed cell death (apoptosis). Immunology. 56:351–358.
Falčo, A.M., D. van Bruggen, S. Marques, M. Meijer, S. Jäkel, E. Agrirre, E.M. Samudyata, E.M. Floridcia, D.P. Vanichkina, C. Fřenč-Constant, et al. 2018. Disease-specific oligodendrocyte lineage cells arise in multiple sclerosis. Nat. Med. 24:1837–1844. https://doi.org/10.1038/s41591-018-0236-y
Franco, R., and D. Fernández-Suárez. 2015. Alternatively activated microglia and macrophages in the central nervous system. Prog. Neurobiol. 131:65–86. https://doi.org/10.1016/j.pneurobio.2015.05.003
Geissmann, F., S. Jung, and D.R. Littman. 2003. Blood monocytes consist of two principal subsets with distinct migratory properties. Immunity. 19:71–82. https://doi.org/10.1016/S1074-7613(03)00174-2
Giles, A.J., J.M. Washnock-Schmid, P.C. Duncker, S. Dahlawi, G. Ponath, D. Pitt, and B.M. Segal. 2018. Myeloid cell plasticity in the evolution of central nervous system autoimmunity. Ann. Neurol. 83:131–141. https://doi.org/10.1002/ana.25128
Goldmann, T., P. Wieghofer, P.F. Müller, Y. Wolf, D. Varol, S. Yona, S.M. Brendecke, K. Kierdorf, O. Stassewski, M. Datta, et al. 2013. A new type of microglia gene targeting shows TAK1 to be pivotal in CNS autoimmunity. Nat. Neurosci. 16:1618–1626. https://doi.org/10.1038/nn.3531
Governan, J. 2009. Autoimmune T cell responses in the central nervous system. Nat. Rev. Immunol. 9:393–407. https://doi.org/10.1038/nri2550
Grigorian, A., L. Araujo, N.N. Naidu, D.J. Place, B. Choudhury, and M. Demetriou. 2011. N-acetylglucosamine inhibits T-helper 1 (Th1)/T-helper 17 (Th17) cell responses and treats experimental autoimmune encephalomyelitis. J. Biol. Chem. 286:40133–40141. https://doi.org/10.1074/jbc.M111.277814
Hadjiantonakis, A.K., S. Macmastery, and A. Nagy. 2002. Embryonic stem cells and mice expressing different GFP variants for multiple non-invasive reporter usage within a single animal. BMC Biotechnol. 2:11. https://doi.org/10.1186/1472-6750-2-11
Heneka, M.T., M.P. Kummer, and E. Latz. 2014. Innate immune activation in neurodegenerative disease. Nat. Rev. Immunol. 14:463–477. https://doi.org/10.1038/nri3705
Heppner, F.L., M. Greter, D. Marino, J. Falsig, G. Ravič, N. Hövelmeyer, A. Waisman, T. Rülicke, A.U. Brandt, T. Leuenberger, H. Radbruch, F. Pitt, and R.A. Niesner. 2010. Expanding two-photon intravital microscopy to the infrared by means of optical parametric oscillator. Biophys. J. 98:715–723. https://doi.org/10.1016/j.bpj.2009.10.035
Hughes, D., C.S. Raine, and E.J. Field. 1968. Invasion of neurones in vitro by non immune lymphocytes. An electron microscopic study. Br. J. Exp. Pathol. 49:356–359.
Jiang, Z., J.X. Jiang, and G.-X. Zhang. 2014. Macrophages: a double-edged sword in experimental autoimmune encephalomyelitis. Immunol. Lett. 160:17–22. https://doi.org/10.1016/j.imlet.2014.03.006
Jolivel, V., F. Bicker, F. Binamé, R. Ploen, S. Ploeg, K. Birkner, G. Raivich, N. Hövelmeyer, A. Rainer, and D.R. Littman. 2000. Analysis of fractalkine receptor CX(3)CR1 gene insertion. MCB. http://doi.org/10.1091/0271-4658.2000.20.11.4106-4114
Kershakenstein, C., M. Stadelmann, G. Dechant, H. Wekerle, and R. Hohlfeld. 2003. Neurotrophic cross-talk between the nervous and immune systems: implications for neurological diseases. Ann. Neurol. 53:292–304. https://doi.org/10.1002/ana.10446
Krishna, S., and M. Overholtzer. 2016. Mechanisms and consequences of entosis. Cell. Mol. Life Sci. 73:2379–2386. https://doi.org/10.1007/s00018-016-2207-0

Wasser et al.

Myeloid cells capture living CNS-invading T cells

https://doi.org/10.1084/jem.20190812

Journal of Experimental Medicine

14 of 15
Lampron, A., A. Larochelle, N. Laflamme, P. Préfontaine, M.-M. Plante, M.G. Sánchez, W.V. Yong, P.K. Stys, M.-É. Tremblay, and S. Rivest. 2015. Inefficient clearance of myelin debris by microglia impairs remyelinating processes. J. Exp. Med. 212:481–495. https://doi.org/10.1084/jem.20141656

Lobo-Silva, D., G.M. Carriche, A.G. Castro, S. Roque, and M. Saraiva. 2017. Interferon-β regulates the production of IL-10 by toll-like receptor-activated microglia. Glia. 65:1439–1451. https://doi.org/10.1002/glia.23702

Lu, Z., M.R. Elliott, Y. Chen, J.T. Walsh, A.L. Kilhanov, K.S. Ravichandran, and J. Kipnis. 2011. Phagocytic activity of neuronal progenitors regulates adult neurogenesis. Nat. Cell Biol. 13:1076–1083. https://doi.org/10.1038/ncb2299

Luche, H., O. Weber, T. Nageswara Rao, C. Blum, and H.J. Fehling. 2007. Faithful activation of an extra-bright red fluorescent protein in “knock-in” Cre-reporter mice ideally suited for lineage tracing studies. Eur. J. Immunol. 37:43–53. https://doi.org/10.1002/eji.200636745

Luchtman, D., R. Gollan, E. Ellwardt, J. Birkenstock, K. Robohm, V. Siffrin, Olah, M., S. Amor, N. Brouwer, J. Vinet, B. Eggen, K. Biber, and H.W. Bodmer. 2009. Gatekeeper role of brain antigen-presenting CD11c+ cells in neuroinflammation. J. Immunol. 182:1247–1258. https://doi.org/10.1016/j.brain.2013.09006

Luchtman, D., R. Gollan, E. Ellwardt, J. Birkenstock, K. Robohm, V. Siffrin, Olah, M., S. Amor, N. Brouwer, J. Vinet, B. Eggen, K. Biber, and H.W. Bodmer. 2009. Gatekeeper role of brain antigen-presenting CD11c+ cells in neuroinflammation. J. Immunol. 182:1247–1258. https://doi.org/10.1016/j.brain.2013.09006

M. Luchtman, D., R. Gollan, E. Ellwardt, J. Birkenstock, K. Robohm, V. Siffrin, Olah, M., S. Amor, N. Brouwer, J. Vinet, B. Eggen, K. Biber, and H.W. Bodmer. 2009. Gatekeeper role of brain antigen-presenting CD11c+ cells in neuroinflammation. J. Immunol. 182:1247–1258. https://doi.org/10.1016/j.brain.2013.09006

Napolitano, I., and H. Neumann. 2010. Protective effects of microglia in multiple sclerosis. Exp. Neurol. 225:24–28. https://doi.org/10.1016/j.expneurol.2009.04.024

Ng, Y.K., and E.A. Ling. 1999. Emperiplois of lymphoid cells in vagal ef-...
Figure S1. Relevance of GlcNAc and CD206 during engulfment. (A) Control Th17-skewed cells and GlcNAc-pretreated Th17-skewed cells were co-cultured with organotypic hippocampal slices. Co-cultures of organotypic hippocampal slices from CX3CR1GFP pups with pathogenic B6.2D2.CFP.Th17 cells were stained with Image-iT LIVE Red Caspase-3 and -7 Detection Kit to identify apoptotic cells and imaged over a time period of 20 min. Only apoptotic T cells were considered for further analyses. Mean (± SEM) percentages of engulfment rate among the interactions of GlcNAc-pretreated Th17-skewed cells in organotypic hippocampal slices (n = 4 organotypic slices from three different experiments) normalized to the GlcNAc-untreated control group (n = 4 organotypic slices from three different experiments). Statistical analysis was performed using two-sided Student’s t tests. (B) Representative visualization of CD31 (magenta), CD206 (red), CX3CR1 (green), and DAPI (blue), in dexamethasone/LPS/IL-4–treated organotypic hippocampal slice cultures (representative for n = 4 reslices from two different experiments). Scale bar = 20 µm. Arrows point to CD206+ myeloid cells. (C) Representative staining of DAPI (blue), CD206 (red), CD4 (magenta), and Iba-1 (green) in EAE diseased animals (n = 3 mice [C57BL/6 or Cx3CR1GFP] from two different experiments). The second panel demonstrates a probable blood vessel. Magnification of a slice view of CD206+ engulfing parenchymal (pc) and perivascular (pv) myeloid cell is shown in the third and fourth panel. Scale bars = 30 µm in first and second panel and 20 µm in the third and fourth panels.

Wasser et al. Journal of Experimental Medicine https://doi.org/10.1084/jem.20190812
Figure S2. Myeloid cells in organotypic hippocampal slices gained from CX3CR1.GFP−/− mice show interactions similar to myeloid cells in organotypic hippocampal slices gained from CX3CR1.GFP+/+ mice. 24-h co-cultures of organotypic hippocampal slices from CX3CR1.GFP−/− pups with pathogenic B6.2D2.CFP.Th17 cells were stained with Image-iT LIVE Red Caspase-3 and -7 Detection Kit to identify apoptotic cells and imaged over a time period of 20 min with two-photon microscopy. The pie chart compares the distribution (%) of interaction modes between myeloid cells and viable 2D2.CFP.Th17 cells (n = 4 organotypic slices from two different experiments), and contingency bars reflect the respective distribution of engulfment, escape, and engulfment attempt. All interactions detectable in organotypic hippocampal slices from CX3CR1.GFP−/− pups (Fig. 4 C) were observed.
Video 1. **Surface-reconstructed intravital two-photon live imaging of an escaping T cell in vivo.** Related to Fig. 1. EAE was induced in CX3CR1<sup>GFP</sup> (green CX3CR1<sup>+</sup> cells) mice via the passive transfer of B6.2D2.RFP.Th17 cells (red). The engulfment and escape of a T cell was followed over time (10 min) in \( n = 12 \) mice from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show a stable contact using Imaris) as well as the 3D-reconstructed version (Imaris). Time is shown in h/min/s.

Figure S3. **Gating strategies. (A–F)** Gating strategies for all flow cytometric analyses used for the presented data are shown. The antibodies used are listed in the Materials and methods (experimental repeats in the respective figures). FSC, forward scatter; PI, propidium iodide; SSC, side scatter.
Video 2. **Two-photon live imaging of Th17 cell–myeloid cell interactions ex vivo.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.CFP.Th17 cells for 24 h and contacts were visualized in n = 4 organotypic slices from three different experiments. Shown here is the original CFP (Th17 cells, false colored here in red) and GFP (myeloid cells) 3D image sequence, smoothened and 3D cropped using Imaris. Time is shown in h/min/s/ms.

Video 3. **Two-photon live imaging of a temporary contact ex vivo.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.RFP.Th17 cells for 24 h and contacts visualized over a time period of 10 min in n = 4 organotypic slices from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show a temporary contact) as well as the 3D-reconstructed version (Imaris; green is myeloid cells, and red is Th17 cells). Time is shown in h/min/s.

Video 4. **Two-photon live imaging of a stable contact ex vivo.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.RFP.Th17 cells for 24 h and contacts visualized over a time period of 10 min in n = 4 organotypic slices from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show a stable contact using Imaris) as well as the 3D-reconstructed version (Imaris). Time is shown in h/min/s.

Video 5. **Two-photon live imaging of a stable engulfment ex vivo.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.RFP.Th17 cells for 24 h and contacts visualized over a time period of 10 min in n = 4 organotypic slices from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show a stable engulfment using Imaris) as well as the 3D-reconstructed version (Imaris). Time is shown in h/min/s.

Video 6. **Two-photon live imaging of an escape following walk-through ex vivo.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.RFP.Th17 cells for 24 h and contacts visualized over a time period of 10 min in n = 4 organotypic slices from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show an escape following walkthrough in Imaris) as well as the 3D-reconstructed version (Imaris). Time is shown in h/min/s.

Video 7. **Two-photon live imaging of an engulfment attempt.** Related to Fig. 2. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.RFP.Th17 cells for 24 h and contacts visualized over a time period of 10 min in n = 4 organotypic slices from three different experiments. Shown here is the original GFP (myeloid cells) and RFP (Th17 cells) 3D image sequence (smoothened and 3D cropped to show engulfment attempts in two independent myeloid cells using Imaris) as well as the 3D-reconstructed version (Imaris). Time is shown in h/min/s.

Video 8. **Two-photon live imaging of a long-term engulfment.** Related to Fig. 3. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.CFP.Th17 cells for 24 h and contacts visualized in n = 4 co-cultures from three different experiments. Shown here is the original CFP (Th17 cells, false colored in red) and GFP (myeloid cells) 3D image sequence, smoothened and 3D cropped with Imaris to show a long-term engulfment. Time is shown in h/min/s.

Video 9. **Two-photon live imaging of a long-term escape.** Related to Fig. 3. Organotypic hippocampal slices from CX3CR1GFP pups were co-cultured with pathogenic B6.2D2.CFP.Th17 cells for 24 h and contacts imaged in n = 4 co-cultures from three different experiments. Shown here is the original CFP (Th17 cells, false colored in red) and GFP (myeloid cells) 3D image sequence, smoothened and 3D cropped with Imaris to show a long-term escape. Time is shown in h/min/s.

Video 10. **In vivo dynamics of T cell–myeloid cell interactions during early and chronic disease.** EAE was induced in CX3CR1GFP (green CX3CR1+ cells) mice via the passive transfer of B6.2D2.RFP.Th17 cells (red). Interactions between T cells and myeloid cells were visualized in mice exhibiting low or high clinical scores. Representative of n = 9 mice from two different experiments. Time is shown in h/min/s/ms.