Influenza A Virus (H1N1) Infection Induces Microglial Activation and Temporal Dysbalance in Glutamatergic Synaptic Transmission

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ABSTRACT Influenza A virus (IAV) causes respiratory tract disease and is responsible for seasonal and reoccurring epidemics affecting all age groups. Next to typical disease symptoms, such as fever and fatigue, IAV infection has been associated with behavioral alterations presumably contributing to the development of major depression. Previous experiments using IAV/H1N1 infection models have shown impaired hippocampal neuronal morphology and cognitive abilities, but the underlying pathways have not been fully described. In this study, we demonstrate that infection with a low-dose non-neurotrophic H1N1 strain of IAV causes ample peripheral immune response followed by a temporary blood-brain barrier disturbance. Although histological examination did not reveal obvious pathological processes in the brains of IAV-infected mice, detailed multidimensional flow cytometric characterization of immune cells uncovered subtle alterations in the activation status of microglial cells. More specifically, we detected an altered expression pattern of major histocompatibility complex classes I and II, CD80, and F4/80 accompanied by elevated mRNA levels of CD36, CD68, C1QA, and C3, suggesting evolved synaptic pruning. To closer evaluate how these profound changes affect synaptic balance, we established a highly sensitive multiplex flow cytometry-based approach called flow synaptometry. The introduction of this novel technique enabled us to simultaneously quantify the abundance of pre- and postsynapses from distinct brain regions. Our data reveal a significant reduction of VGLUT1 in excitatory presynaptic terminals in the cortex and hippocampus, identifying a subtle dysbalance in glutamatergic synapse transmission upon H1N1 infection in mice. In conclusion, our results highlight the consequences of systemic IAV-triggered inflammation on the central nervous system and the induction and progression of neuronal alterations.

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**IMPORTANCE** Influenza virus (IAV) causes mainly respiratory tract disease with fever and fatigue but is also associated with behavioral alterations in humans. Here, we demonstrate that infection with a low-dose non-neurotrophic H1N1 strain of IAV causes peripheral immune response followed by a temporary blood-brain barrier disturbance. Characterization of immune cells uncovered subtle alterations in the activation status of microglia cells that might reshape neuronal synapses. We established a highly sensitive multiplex flow cytometry-based approach called flow synaptometry to more closely study the synapses. Thus, we detected a specific dysbalance in glutamatergic synapse transmission upon H1N1 infection in mice. In conclusion, our results highlight the consequences of systemic IAV-triggered inflammation on the central nervous system and the induction and progression of neuronal alterations.

**KEYWORDS** influenza A virus, microglial activation, glutamatergic synapse transmission, synaptosomes, flow synaptometry, influenza, interorgan communication, microglia, neuronal synapses

Influenza A virus (IAV) causes infection of the respiratory tract and repeatedly appears throughout the lifetime of almost every individual. Subtypes of IAV, such as H1N1, often contribute to seasonal epidemics or may even cause pandemics, representing a major cause for morbidity and mortality worldwide (1). Approximately 8% of the U.S. population becomes infected with IAV each season (2), eventually resulting in over 30,000 IAV-associated deaths per year (3). Based on these assumptions, it has been extrapolated that influenza infections sum up to 4 to 50 million annual cases and 15 to 70,000 deaths in the EU (4, 5). The resolution of IAV infection requires the activation of innate and adaptive immunity and is initiated in lung epithelial cells, which are primary targets of IAV. Here, IAV is detected by cytosolic pattern recognition receptors such as endosomal toll-like receptors or retinoic acid inducible gene I, resulting in the initiation of NF-κB-dependent release of a plethora of cytokines and chemokines, including interleukin (IL-1β), IL-6, tumor necrosis factor (TNF), and type I interferons (IFN) that induce an antiviral state via IFNAR1 and IFNAR2 receptor activation. The induction of interferon-regulated pathways is characterized by the expression of various GTPases such as MX proteins that interfere with virus growth in infected cells (6). Potent activation of innate immunity results in the subsequent establishment of adaptive cellular immunity mediated by virus-specific CD8+ cytotoxic T cells and CD4+ T helper cells and humoral immunity via antibody-producing B cells (7). As a consequence of the mounted immune response, IAV infection is associated with high serum levels of proinflammatory cytokines, resulting in distinct behavioral alterations in patients, a phenomenon also referred to as sickness behavior (8). Although the profound mechanism for the induction of sickness behavior is not fully understood, previous reports have highlighted that this effect is caused by a transport of peripheral cytokines into the brain parenchyma via the blood-brain barrier (BBB) (9, 10). Thus, it has been speculated that infections such as IAV pneumonia act as one of the initial triggers contributing to the development of neurological disorders and therefore have been studied in experimental models over past years (11–14).

As brain-resident macrophages, microglial cells represent the first line of defense against pathogens. Under homeostatic conditions, microglial cells possess a highly ramified morphology, constantly scanning their environment for pathogens as well as supporting neuronal function and synaptic plasticity (15–18). Microglia become activated upon sensing of pathogens and inflammatory signals, releasing cytokines, chemokines, and other inflammatory mediators such as nitric oxide and complement factors (19). Therefore, microglial activation has often been linked to neuropathologies like Alzheimer’s disease (AD) or schizophrenia and, moreover, to aggravated phagocytosis of synapses (20, 21). Although previous reports indicate that peripheral inflammation induced by viral infection or polyinosinic:polycytidylic acid (poly(I:C)) and lipopolysaccharides (LPS) administration leads to microglial activation and behavioral
changes (22–24), the underlying pathways of neurological alterations have not been fully understood. Here, we demonstrate that respiratory IAV/PR8/H1N1 infection results in a distinct peripheral inflammatory pattern that is associated with transient dysbalance in BBB homeostasis. Even though initial histological examination of brain sections did not reveal evidence for pathological changes, detailed flow cytometric characterization highlighted distinct region-specific activation of microglia in the early phase of infection. We detected that expression of markers associated with synaptic pruning peak at 14 days post-infection (dpi) alongside diminished mRNA levels of the presynaptic glutamate transporter VGLUT1. Notably, the establishment of a novel flow cytometry-based approach (“flow synaptometry”) enabled us to assess the composition of excitatory and inhibitory synapses, demonstrating a significant reduction of VGLUT1 in excitatory presynapses at 21 dpi, which is further accompanied by changes in neurotrophin gene expression. Our findings provide detailed insights into temporal effects of peripheral inflammation on brain homeostasis and microglial-neuronal interaction that specifically shape synapse physiology over the course of IAV infection, possibly contributing to persisting neurological alterations.

RESULTS

Infection with influenza A virus (H1N1) alters brain homeostasis of mice and temporarily affects blood-brain barrier function. To explore the effects of virus-induced peripheral inflammation on microglial activation and neuronal alterations, we infected mice with a low dose of the mouse-adapted influenza A virus (IAV)/PR8/H1N1 and monitored their body weight and cytokine levels in serum and lung throughout the infection (Fig. 1a and b). After 4 to 5 days post-infection (dpi) with 0.32 of the median tissue culture infection dose (TCID₅₀) of IAV, the relative body weight of infected mice started to decline, and this trend became significant after 7 dpi (naive, 100.8 ± 0.7 g; IAV-infected, 89.8 ± 1.9 g, \( P < 0.05 \)) while reaching its peak at 8 dpi (naive, 102.2 ± 0.6 g; IAV-infected, 86.7 ± 2.5 g, \( P < 0.001 \)) (Fig. 1b). Beyond this point, mice started to recover and reached their initial weight by 14 dpi (naive, 107.9 ± 0.7 g; IAV-infected, 102.9 ± 1.9 g, \( P < 0.81 \)). These observations corresponded well to the levels of cytokines and chemokines we detected in these mice. In the bronchoalveolar lavage (BAL) fluid we collected from infected lungs, cytokines increased strongly after 4 dpi and remained elevated until 9 dpi (see Fig. S1 in the supplemental material).

Although not reaching equally high concentrations, we also found serum levels of IFN-γ (156.0 ± 34.4 pg/ml, 6 dpi), IL-6 (37.7 ± 11.5 pg/ml, 6 dpi), and CCL2 (21.0 ± 7.3 pg/ml, 9 dpi) to be higher in IAV-infected mice, whereas granulocyte-macrophage colony-stimulating factor (GM-CSF) and TNF remained unaltered (Fig. 1c to j). These observations match those from previous reports of our group, where the viral burden in the lungs of infected animals shows a strong decrease between 7 and 9 dpi and a complete clearance by 14 dpi (25).

In recent years, several studies have shown that systemic infection with pathogens or the application of pathogen-associated substances alone, such as endotoxin or poly(lC), can induce sickness behavior and lead to disturbances in brain homeostasis (8, 12, 26, 27). Given the high abundance of peripheral cytokines, we wondered whether this effect could also be seen in IAV-infected mice. Based on initial results, the time points of 7 dpi (first day of significantly different body weight), 10 dpi (last day of significantly different body weight), 14 dpi (restoration of initial body weight), and 21 dpi (to evaluate potential long-term effects of IAV infection) were selected for further analyses. Consequently, to evaluate the expression of cytokines and chemokines in the brains of naive and infected mice, brains were collected at the above-mentioned time points and dissected into cortex (CTX) and hippocampal formation (HFF). Reverse transcription quantitative PCR (RT-qPCR) analysis revealed that upon acute IAV infection, the expression of Il1b (IL-1β), Il6 (IL-6), Tnf (TNF), and Ccl2 (CCL2) remained unchanged (Fig. 2a to d). Although previous studies have demonstrated IAV/PR8/34 to be non-neurotropic (12, 13), we evaluated the viral load in the cortex, hippocampus, and olfactory bulb at 7 and 10 dpi to exclude direct viral effects in the central nervous system. Differential transcription quantitative PCR (RT-qPCR) analysis revealed that upon acute IAV infection, the expression of Il1b (IL-1β), Il6 (IL-6), Tnf (TNF), and Ccl2 (CCL2) remained unchanged (Fig. 2a to d). Although previous studies have demonstrated IAV/PR8/34 to be non-neurotropic (12, 13), we evaluated the viral load in the cortex, hippocampus, and olfactory bulb at 7 and 10 dpi to exclude direct viral effects in the central nervous system.
system (CNS) of infected mice. In line with published data, we did not detect viral copies in these brain regions (Fig. 2e); however, despite the absence of IAV in the CNS, mRNA levels of Ifnb1 (IFN-β) were elevated in the cortex at 10 dpi ($P < 0.055$) and significantly increased by 14 dpi. A similar trend was also observed for Ifng (IFN-γ) (Fig. 2f and g). As the induction of interferons often results in a plethora of immune responses, ranging from the establishment of an antiviral state in neighboring cells to modulation of the adaptive immunity, we sought to investigate for an altered regulation of interferon-stimulated genes (ISGs). Indeed, we found the expression of the inducible GTPase Irgm1 (IRGM1) to be significantly increased in the cortex 10 dpi, whereas expression of other ISGs, such as Igtp (IGTP), Mx2 (MX2), or Rsad2 (RSAD2), remained unaffected (Fig. 2h to k). In addition to antiviral state induction, type I and type II interferons have been identified recently as key modulators of immune cell entry to the CNS during physiological but also pathological conditions by affecting leukocyte trafficking via the blood-brain barrier (BBB) or the blood-cerebrospinal fluid barrier (BCSFB) located in the choroid plexus (28). Interestingly, our data revealed that the gene expression of barrier-associated tight junction proteins Cldn5 (Claudin-5) and Tjp1 (ZO-1) (29) was significantly altered in the cortex and hippocampus upon IAV infection (Fig. 3a to c). However, the expression level of the chemokines CXCL9 and CXCL10, known attractants of leukocytes to the CNS (30, 31), showed no significant differences (Fig. 3d and e). In summary, peripheral infection with the influenza A virus led...
to increased gene expression of interferons in the brain combined with a reduction of tight junction proteins, suggesting an impaired functionality of the BBB and BCSFB.

**IAV infection results in activation of brain-resident microglial cells.** Since the increased expression of cytokines can be an indicator of an induced immune response, we sought to determine whether IAV infection resulted in the emergence of microglial cell activation and neuroinflammation. Under homeostatic conditions, the majority of the brain’s immune cell population is represented by microglial cells that perform a variety of tasks to support neuronal functions and are also known producers of type I interferons (32, 33). During steady state, microglia are characterized by their highly ramified morphology with numerous thin processes continuously surveilling the brain parenchyma. Upon encountering pathogen- or damage-associated molecular patterns,
these cells become activated, retract their extensions, and become highly mobile while migrating toward the sites of inflammation (15).

Thus, sagittal paraffin sections of brains from naive and IAV-infected mice were stained for ionized calcium binding adapter molecule 1 (IBA1) and complement receptor 3 (CD11b). Compared to naive mice, IBA1-positive cells in cortices and hippocampi of IAV-infected animals did not differ in numbers or their ramified morphology (Fig. 4a and c). Similarly, histological examination of CD11b-positive cells also revealed no prominent differences from controls at 14 or 21 dpi (Fig. 4b and d). Furthermore, staining for neuronal markers MAP2 and NeuN did not indicate neuronal changes as results of IAV infection-induced neuroinflammation (Fig. S2). However, IBA1 and CD11b expression is not exclusive to brain-resident microglial cells, as these markers can also be expressed by infiltrating monocytes or border-associated macrophages. To confirm whether the cells observed during histopathological examination indeed represent microglia, we employed immunofluorescence microscopy of cryosections with staining for the microglia-specific marker transmembrane protein 119 (TMEM119) (34). Even though TMEM119 has been reported to be a marker with robust expression, its signal did not allow a detailed evaluation of microglial morphology, as it was present on their cellular processes but not the soma (Fig. 4e). However, due to the substantial overlap of signal between IBA1 and TMEM119, we concluded that the majority of IBA1-positive cells in our samples constitute microglia (Fig. 4f). Hence, these cryosections from naive
FIG 4 Histopathological and immunohistochemical examination of brain tissue does not reveal alterations upon infection with IAV. (a and b) Histopathological overview of representative sagittal paraffin sections from brains of (Continued on next page)
and IAV-infected mice were further used to compare microglial morphology in more detail. Coherent with the previous histological observations, microglia in the cortex and hippocampus retained their ramified morphology with thin extensions after 10 and 14 dpi. In summary, histology and fluorescence microscopy revealed no obvious evidence of pathological changes in the brain following respiratory IAV infection.

Microglia were previously shown to possess a diverse phenotypic spectrum with multidimensional activation profiles that highly depend on the microenvironmental stimuli (35, 36). Accordingly, we isolated these cells from brains of naive and IAV-infected mice to characterize their phenotype at higher resolution via multiparametric flow cytometry. After the initial removal of debris, we subjected all cells to unsupervised t-distributed stochastic neighbor embedding (t-SNE), resulting in three different clusters of cells (Fig. 5a) that were highly distinct in their surface expression levels of CD45, CD11b, and CX3CR1 and showed minor differences in the expression of MHC class I and II, CD80, CD86, and F4/80. Using a subsequent manual gating approach, cells within the three different clusters were identified as brain-resident microglial cells (CD45low CD11b+) and peripheral immune cell subsets consisting of CD45hi CD11b- and CD45hi CD11b+ cells (Fig. 5b and c). Unexpectedly, we detected a minor but significant increase in the frequency of CD45hi CD11b- but not CD45hi CD11b+ cells at 7 dpi in the brains of IAV-infected mice (Fig. 5d to f). As the infection progressed (10 dpi), not only CD45hi CD11b+ but also more CD45hi CD11b- cells were found in the hippocampus and cortex (CTX; P < 0.06) (Fig. 5g to i). Upon resolution of the peripheral infection (14 dpi), the number of CD45hi CD11b- cells returned to levels of naive controls, whereas the CD45hi CD11b+ cell population remained elevated in the hippocampus of infected mice (Fig. 5j to l). However, microglia constituted by far the largest population of immune cells at all time points and, thus, confirmed our previous immunofluorescence results.

After identification of microglia, we subsequently analyzed changes in their surface marker expression upon IAV infection. We discovered an upregulation of several immunological molecules in both cortex and hippocampus 10 dpi (Fig. 5m, Fig. S3): microglia of infected mice expressed higher levels of MHC class I and II, CD80, and F4/80, whereas expression of the fractalkine receptor CX3CR1 was reduced. At 14 dpi, microglial activation was still evident in both brain regions examined, with significantly increased expression of MHC I and F4/80 and decreased expression of CX3CR1, respectively. However, the alterations were not as pronounced as before, suggesting a return to baseline levels after resolution of peripheral IAV infection. Taken together, flow cytometric analysis supported our previous findings of an altered BBB by revealing an elevated number of recruited peripheral immune cells in the brain parenchyma. Furthermore, activation of brain-resident microglia was increased upon IAV infection and remained high throughout the course of the infection.

**Altered synaptic pruning upon IAV infection-induced microglial activation.**

Microglial shape neuronal connections via pruning of excessive synapses (37, 38), a process by which several studies have highlighted a hallmark of different neurological disorders (39–41). Of note, it has been previously demonstrated that influenza infection caused cognitive dysfunction and led to an altered neuronal architecture in mice (12). Following the aforementioned activation of microglia, we tested whether IAV infection further reactivates microglia-mediated synaptic pruning by analyzing the expression of phagocytosis-associated receptors in the cortex and hippocampus (Fig. 6). While the

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**FIG 4 Legend (Continued)**

naive mice stained against IBA1 or CD11b. (c and d) Panels show the cortex (CTX, left) and hippocampal formation region (HPF, right) from naive and infected mice (14 and 21 dpi) upon staining for IBA1 or CD11b. (e) Immunohistochemical preparation of cryosections shows representative images of microglial cells stained with antibodies for IBA1 and TMEM119, and white arrows indicate strongly overlapping signals in both channels. (f) Comparison of microglial morphology in cryosections from naive (top) and IAV-infected (middle and bottom) mice during the late phase of IAV infection (10 to 14 dpi). Sections were stained with antibodies against IBA1, and confocal microscopy images were generated for cortex (CTX, left) and hippocampal formation (HPF, right). Insets in bottom right corners show selected microglial cells with higher detail. Scale bars, 50 μm.
FIG 5 Characterization of immune cell subsets and microglial activation in the brains of IAV-infected animals. Immune cells were isolated from perfused brains of naive and infected animals as described in the text and subjected to flow cytometric analysis. (a) Unsupervised clustering of immune cell subsets was performed by t-distributed stochastic neighbor embedding (t-SNE), and cells within clusters were subsequently identified by manual gating. Further, differential expression of surface markers CD45, CD11b, major histocompatibility complex (MHC) classes I and II, CD80, CD86, F4/80, and CX3CR1 are shown for the generated clusters. (b and c) Representative strategy for manual gating. First, cells were selected based on the forward-scatter/side-scatterplot (FSC/SSC) before exclusion of dead cells as well as doublets (not shown). Immune cell populations then were separated by their expression of the surface markers CD45 and CD11b into brain-resident CD45<sup>lo</sup> CD11b<sup>hi</sup> microglial cells and recruited CD45<sup>hi</sup> CD11b<sup>lo</sup> or CD45<sup>hi</sup> CD11b<sup>hi</sup> cells. (d to f) Bar charts show the frequencies of identified immune cell populations in the brains of naive (white bars) and IAV-infected animals (black bars) at 7 dpi. (g to l) Bar charts show the frequencies of identified immune cell populations in specific brain regions of naive (white bars) and IAV-infected animals (black bars) at 10 and 14 dpi. (m) Heat map plot of relative microglial surface expression of MHC I, MHC II, CD80, CD86, F4/80, and CX3CR1 in naive and infected mice at 10 and 14 dpi. Median fluorescence intensities of expressed markers were normalized to their overall mean, and data were plotted using R with “lattice” package. Data are shown as means ± SEM, and groups were compared via Student’s t test with Welch’s correction. Significant differences are indicated: *, P < 0.05; ***, P < 0.001.
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Gene expression of the scavenger receptor \textit{Cd36} and lysosome-associated protein \textit{Cd68} was unaffected at 10 dpi, their expression was significantly increased upon sustained proinflammatory triggers (14 dpi) (Fig. 6a and b). Since aberrant synaptic pruning further requires the upregulation of complement components to tag synapses (42), we consequently examined the expression of \textit{C1qa} and \textit{C3}, which likewise increased 14 dpi in cortex and hippocampus, providing the prerequisites for synapse elimination (Fig. 6e and f). In contrast, mRNA levels of \textit{Trem2}, an innate immune receptor implicated in cell activation and phagocytosis (43), and the inducible nitric oxide synthase (\textit{Nos2}) remained unaffected upon IAV infection (Fig. 6c and d). In conclusion, the data presented suggest the dysregulation of synaptic pruning and altered synaptic function following IAV infection.

**Temporally dysregulated glutamatergic synaptic transmission and neurotrophin gene expression following resolution of peripheral IAV infection.** Previously, we have demonstrated that infection-induced neuroinflammation affects synaptic protein composition with detrimental outcome for glutamatergic neurotransmission (44). To determine whether infection with IAV affects synaptic protein composition, we analyzed the gene expression of the presynaptic glutamate transporter \textit{Slc17a7} (VGLUT1) over the course of infection (Fig. 7a). Compared to naive samples, gene expression did not differ significantly at 10 and 14 dpi; however, mRNA levels pointed toward a reduction at 14 dpi \((P < 0.29)\). When assessing the expression levels at 21 dpi, we detected not only a return to baseline expression but also significant overcompensation. Thus, we concluded that the systemic inflammation driven by peripheral IAV infection causes a
FIG 7 New tool to analyze synaptic proteins via flow cytometry. (a) Gene expression of VGLUT1 (Slc17a7) was analyzed in whole-brain homogenate of naive (white bars) and IAV-infected mice at 10, 14, and 21 dpi (black bars). Relative gene expression was examined by RT-qPCR as described in the text, and expression of the target gene was normalized to the expression level of Hprt. Subsequently, relative expression was normalized to the mean of naive animals. Groups were compared via Student’s t test with Welch’s correction. (b) Synaptosomes were isolated from brain regions of perfused animals as described in the text, and the synaptosomal fraction (31) was subjected to electron microscopy. Red arrows indicate intact synapses formed by a pre- and postsynaptic compartment. Scale bar in the bottom left corner indicates 500 nm. (c) Graphical illustration of synaptosome selected in panel b. Presynaptic (Continued on next page)
functional disturbance in excitatory neurons in mice, an assumption well in line with previous reports showing altered neuronal morphology and cognitive impairment following IAV infection (12, 13). To study synaptic changes of excitatory neurons more in depth, we directly analyzed glutamatergic synapse composition. We therefore developed a refined approach by taking advantage of synaptosomes, i.e., sealed presynaptic nerve terminals often containing opposite postsynaptic elements, providing a well-established model to investigate synapses stripped from their surrounding tissue. Protocols to isolate synaptosomes have existed for several decades and describe the purification of synaptic material from brain tissue using discontinuous density gradient centrifugation. Upon adaption of these protocols to our samples, we first examined the content of our isolates via electron microscopy (Fig. 7b and c) and detected single fragments of membranes and intact presynapses adjacent to thickened postsynapses. The imaged synaptosomes show diameters of 350 to 700 nm at the presynaptic side and contain zero to one mitochondrion and many small, clear synaptic vesicles. In addition, postsynaptic densities and the synaptic cleft are well noticeable, allowing the conclusion that our technique ensures the isolation and purification of synaptosomes from brain tissue. Second, we isolated proteins from our synaptosome samples and determined the protein levels of VGLUT1 by Western blotting before and after IAV infection (Fig. 7d and e). Here, levels of VGLUT1 were partially diminished at 14 dpi ($P < 0.09$) and significantly reduced 21 dpi, confirming previous findings by RT-qPCR.

So far, synaptosomes have mostly been analyzed in batches (45), and only a few studies are known to employ quantitative approaches (46, 47). To investigate synapse composition at the single synapse level, we established flow synaptometry, a novel, flow cytometry-based approach allowing a high-throughput analysis. Reportedly, synaptosomes are rather small objects with diameters from 0.5 to 2 μm (48, 49). To allow the size discrimination of detected events in a flow cytometer, we first established a size gate ranging from 0.3 to 1 μm by utilizing fluorescent silica beads with defined diameters (Fig. 7f). Typically, conventional flow cytometers display a poor forward-scatter light (FSC) resolution for events of such small scale. This cannot be compensated by increasing the detector sensitivity alone, as this also favors the amplification of other buffer-residing objects small enough to pass through the 0.22-μm pores of conventional sterile filters. Thus, we employed the lipophilic styryl dye FM4-64 that becomes highly fluorescent when bound to lipid bilayers, enabling us to distinguish cellular components from inorganic residues in the buffer and noise (Fig. 7g). This step was coupled to the replacement of the standard FSC-triggered detection by fluorescence-triggered detection in the BL3 channel of our flow cytometer (not shown), now favoring only FM4-64-stained events while neglecting everything below the threshold. Besides increasing the flow cytometer’s sensitivity for small events, this procedure also reduces the chance of falsely detecting aggregates of multiple synaptosomes as one event (45). Finally, we compared the frequencies of events positive for VGLUT1 (Fig. 7h) and i and consistently detected a significant reduction for this protein at 21 dpi.

**FIG 7 Legend (Continued)**

nerve endings still contain synaptic vesicles and mitochondria, whereas postsynaptic domains are characterized by their region of high postsynaptic density (PSD). (d and e) Protein content of VGLUT1 from synaptosomes of whole-brain homogenate was assessed via Western blotting. Bar charts show the relative optical density of the protein bands from naive (white bars) and infected (black bars) animals at 14 and 21 dpi upon normalization to β-actin expression. Values were further normalized to the mean of naive animals. Groups were compared via one-way ANOVA with Holm-Sidak post hoc correction. (f to h) Isolated synaptosomes were subjected to flow cytometry, and the representative gating strategy is shown. (f) First, a gate with a size range from 300 to 1,000 nm was established in the FSC channel using silica beads. (g) Second, separation of biological particles from residues in the buffer was facilitated using the styryl dye FM4-64 that integrates into the lipid membranes of biological organelles. FSC-triggered detection then was replaced by fluorescence-triggered detection with FM4-64 in the BL3 channel with a fluorescence threshold set above the noise at 0.3 × 103 (not shown). (h) Lastly, events detected in the size range of 300 to 1,000 nm were further gated for their expression of VGLUT1. (i) Bar chart shows the frequency of VGLUT1+ events in the brains of naive (white bars) and IAV-infected animals (black bars) at 14 and 21 dpi. In all cases, data are shown as means ± SEM and significant differences are indicated. *, $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$; **** $P < 0.0001$. 
whereas no differences appeared 14 dpi. In light of these substantial changes, we aimed to narrow down the synaptosome analysis on cortex and hippocampus of IAV-infected mice (Fig. 8).

**Flow Synaptometry**

(a) Synaptophysin (Syp) events were selected and subsequently gated for Homer1+ or Gephyrin+, respectively. (c and d) Bar charts show the frequencies of Syp+ Homer1+ and Syp+ Gephyrin+ subpopulations from naive (white bars) and IAV-infected animals (black bars) at 14 and 21 dpi. (e and f) Intact synaptosomes from excitatory synapses consisting of a pre- and postsynaptic terminal (Syp+ Homer1+) were further examined for their expression of VGLUT1 and GluR1.

**Brain RT-qPCR**

(g to j) RNA was isolated from brains of perfused animals as described in the text. Relative gene expression levels of BDNF (Bdnf), NGF (Ngf), TrkB (Ntrk2), and p75NTR (Ngfr) were examined by RT-qPCR at different time points after IAV infection (10 to 21 dpi). Expression of target genes was normalized to the expression level of Hprt. Subsequently, relative expression was normalized to the means of naive animals. For all graphs, data are shown as means ± SEM, and groups were compared via Student’s t-test with Welch’s correction. Significant differences are indicated. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.

Influenza Alters Glutamatergic Synapses

**FIG 8** Flow cytometric synaptosome analysis and gene expression of neurotrophins and neurotrophin receptors in naive and IAV-infected mice. (a to f) Synaptosomes were isolated from cortex (CTX) and hippocampal formation (HPF) of naive and IAV-infected mice and subjected to further flow cytometric analysis as shown in Fig. 7. (a and b) After size gating and removal of unspecific events via fluorescence-triggered detection (not shown), Synaptophysin+ (Syp+) events were selected and subsequently gated for Homer1+ or Gephyrin+, respectively. (c and d) Bar charts show the frequencies of Syp+ Homer1+ and Syp+ Gephyrin+ subpopulations from naive (white bars) and IAV-infected animals (black bars) at 14 and 21 dpi. (e and f) Intact synaptosomes from excitatory synapses consisting of a pre- and postsynaptic terminal (Syp+ Homer1+) were further examined for their expression of VGLUT1 and GluR1. (g to j) RNA was isolated from brains of perfused animals as described in the text. Relative gene expression levels of BDNF (Bdnf), NGF (Ngf), TrkB (Ntrk2), and p75NTR (Ngfr) were examined by RT-qPCR at different time points after IAV infection (10 to 21 dpi). Expression of target genes was normalized to the expression level of Hprt. Subsequently, relative expression was normalized to the means of naive animals. For all graphs, data are shown as means ± SEM, and groups were compared via Student’s t-test with Welch’s correction. Significant differences are indicated. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.
synaptosomes. Here, results from IAV-infected animals were consistent with previous findings and did not differ from naïve controls at 14 dpi but displayed a substantial loss of Syp⁺ Homer1⁺ VGLUT1⁺ synaptosomes in the cortex and hippocampus at 21 dpi (Fig. 8e). In contrast, frequencies of Syp⁺ Homer1⁺ GluR1⁺ synaptosomes from either cortex or hippocampus did not appear significantly altered in the course of an IAV infection, although minor fluctuations were detectable.

Finally, we addressed the question of whether the evident synaptic changes might, at least in part, be the result of impaired neurotrophin levels. These proteins act as important mediators of neuronal survival and differentiation in the CNS and therefore play a crucial role during the development of the brain and synaptic plasticity (50, 51). Hence, cortices and hippocampi of naïve and infected mice were compared by RT-qPCR at different time points after IAV infection. The levels of brain-derived neurotropic factor (Bdnf) and nerve growth factor (Ngf), both important mediators of neuronal differentiation and survival, did not vary until the late phase of IAV infection but changed significantly at 21 dpi (Fig. 8g to j). While expression of Bdnf showed a substantial reduction in cortex and hippocampus, Ngf was increased. Furthermore, the expression of the BDNF-specific neurotrophin receptor tropomyosin receptor kinase B (TrkB/Ntrk2) was elevated in both brain regions 21 dpi. The pan-neurotrophin receptor p75NTR (Ngfr), known to be differentially expressed during various neurodegenerative diseases (52, 53), remained unaltered throughout the course of IAV infection.

In summary, our data show that mRNA and protein levels of synapse components were altered upon respiratory IAV infection in mice. Furthermore, we established a novel approach to quantify the composition of synapses in brains by applying flow cytometry onto antibody-labeled synaptosomes. Thus, we highlighted that upon IAV infection, the frequency of inhibitory synapses is increased while glutamatergic neurotransmission is impaired. Consistent with this, gene expression of neurotrophins and their receptors was also altered in the brains of infected animals 21 dpi as a possible consequence of a disturbed neurotransmission.

DISCUSSION

Infections with influenza A virus affect all age groups and accumulate in annual epidemics. Usually, the infection is associated with fever, cough, and a sore throat but also induces symptoms of sickness behavior, such as weakness or decreased interest in surroundings (54). Moreover, cases of behavioral alterations in the form of narcolepsy (55) and the development of major depression (56) have been connected with IAV infection, while other studies showed impaired hippocampal neuron morphology and cognition in the presence of neuroinflammation in mice (12, 13). Overall, various findings suggest a link between peripheral infection and the development of neuropsychiatric disorders; however, the underlying mechanisms remained largely elusive. To address this question, we infected mice with a low dose of IAV that aimed to mimic the disease progression in humans and monitored an elevated concentration of cytokines in lungs and sera during the first days of infection. In particular, we detected increased levels of IL-6 and IFN-γ between 6 and 9 dpi, which coincided with a prominent loss of body weight. These data are in line with our previous observations (25) and are the result of a cellular and humoral immune response facilitated by cytotoxic CD8⁺ T cells and IFN-γ-producing NK and CD4⁺ T helper cells that leads to diminished viral burden in the lungs (7). Recently, the induction of peripheral inflammation through administration of LPS, poly(I:C), or IAV has been shown to cause elevated serum levels of IL-6 and TNF, which led to increased expression of inflammatory cytokines in the brain (12, 22, 57, 58). Consequently, we assessed mRNA levels of various inflammatory cytokines but were unable to detect prominent changes for Ifnb1, Il6, Tnf, or Ccl2. Although our data and other previous studies could show that IAV PR8/A/34 (H1N1) lacks the ability to infect the brain (12, 13), we found a moderate but significant increase of type I interferons (Ifnb1) and the IFN-γ-inducible GTPase Irgm1 in the cortex of infected mice at 10 dpi. This observation may be a consequence of activated...
microglia in response to high serum titers of inflammatory cytokines (32, 33) infiltrating the CNS via the BBB (9, 10, 59). However, the magnitude of Ifnb1 expression levels remained considerably low and did not lead to a more restricted permeability of the BBB as observed in previous studies (60, 61). In contrast, we found a diminished expression of genes associated with tight junction proteins of the BBB (29) at 7 dpi, shortly after serum levels of IFN-γ peaked in IAV-infected mice. The dysregulated state of BBB and BCSFB was further indicated by the marginally increased frequencies of peripheral immune cells we detected in cortex and hippocampus between 7 and 14 dpi. These results are in line with several reports demonstrating the ability of type II interferons to downregulate the expression of tight junction proteins (62, 63). Although IFN-γ is further able to modify the expression of chemokines CXCL9 and CXCL10 by the BBB and BCSFB (64, 65), gene expression of Cxcl9 or Cxcl10 in the brain homogenate of IAV-infected mice revealed no difference in our studies, in contrast with a previous report on poly(I:C)-induced release of CXCL10 by brain endothelial cells. In the latter study, the authors associated the induction of sickness behavior in mice with BBB- and BCSFB-derived CXCL10, further leading to aggravated hippocampal synaptic plasticity via CXCR3 signaling in neurons (27). Of note, both poly(I:C) and IAV infection result in RIG-I signaling cascade induction. However, only IAV is able to avoid immune responses by directly interfering with cellular signaling pathways or host gene expression that ultimately disturb induction of interferons and antiviral proteins (66–68).

Thus, poly(I:C)-induced inflammation can only partially translate into effects observed in our IAV infection model; therefore, further investigation on the transcriptome and the contribution of brain endothelia and epithelial cells to chemokine release upon IAV infection is required.

Despite the altered cytokine expression in the brain pointing toward the occurrence of neuroinflammation upon peripheral infection with IAV, histological examination did not show inflammatory foci in cortices and hippocampi of infected mice. In this respect, microglial number and their ramified morphologies remained unaltered throughout the course of IAV infection, which partially collides with data of Jurgens et al. (12) showing increased IBA1 staining and reduced ramification of microglia already at 7 dpi. However, those findings may just be the result of a different genetic background (BALB/c versus C57BL/6Jr) and a higher IAV inoculation dose, as both loss of body weight and locomotive activity of mice appeared at least 2 days earlier than in our studies, using only 0.32 TCID50 of IAV to induce the infection. Presumably, the immune system responds proportionally to the infection dose, and in our experiments, animals had only a mild loss of body weight and did not succumb to the infection. However, whether the indirect activation of microglia by peripheral challenge is dose-dependent is an interesting point that awaits further elucidation. Regardless, microglial activation can appear on a multidimensional scale (35, 36), and, in contrast to morphologic analyses, detailed investigation via flow cytometry allowed us to detect a moderate but significant region-specific activation indicated by increased expression of MHC I and F4/80 at 10 dpi and again at 14 dpi. These observations point toward a delayed onset of neuroinflammation following a peripheral immune response. Moreover, our data suggest that IAV infection induces a rather subtle and short-term inflammation in the brain, which might remain unnoticed using common histopathological approaches. Indeed, when mice infected with IAV PR8/A/34(H1N1) were assessed for possible long-term effects at 30 dpi, no signs of neuroinflammation were apparent at all (13). In this context, a growing body of evidence has emphasized the connection between inflammation-induced activation of microglia and the recurring pruning of synapses in neurological disorders (39–41). In addition to the prerequisite of activated microglia, synaptic pruning depends on components of the complement system, which tag synapses for a possible elimination (42, 69). In fact, we found increased expression of phagocytosis-related genes as well as upregulated mRNA levels of C1QA and C3 in IAV-infected animals. Interestingly, these results were apparent only at 14 dpi, indicating once again temporary repercussions in the nervous system upon
peripheral inflammation. In this regard, our group has previously shown the effects of neuroinflammation on synapse integrity (44, 70). Given the reduced expression of genes associated with synapse neurotransmission, we decided to test for adverse effects of IAV infection on synapse integrity. Establishing a novel flow cytometry-based approach to quantify the composition of synaptosomes derived from cortices and hippocampus allowed us to detect compelling effects on presynaptic glutamatergic signaling in cortices and hippocampi at 21 dpi, while the fraction of inhibitory synapses was increased.

To date, investigation of isolated synaptosomes represents a convenient tool to study the function and physiology of synapses freed from their surrounding environment (48). However, only a few studies have utilized flow cytometric approaches to assess synaptosomes and, in most cases, only make use of one synaptic component (47, 71, 72). Other reports focused on sorting of synaptosomes derived from fluorescent neurons (46) or applying mass cytometry and mass-coupled antibodies (73), but reporter mouse models or mass cytometers are not always featured equipment of a neuroscientific laboratory. Thus, our profound technique represents an easy-to-use alternative to previous approaches, as we demonstrate that simple multiplexing is feasible using a conventional isolation procedure and standard flow cytometer. To our knowledge, our study is the first to extend previous attempts of flow synaptometry by simultaneously quantifying the relative abundance of pre- and postsynaptic components in synaptosomes from different types of neurons from different brain regions. Moreover, choosing flow synaptometry over conventional Western blotting enables us to focus specifically on intact synapses, resulting in a high sensitivity while requiring only small amounts of sample material. In addition, high-throughput acquisition allows the characterization of a large number of samples within the same session and, thus, facilitates a high robustness of data compared to Western blot analysis.

So far, little is known about the direct consequences of peripheral infections or inflammation on glutamatergic signaling, but models using the viral mimic poly(I:C) discovered changed extracellular glutamate levels and synaptic transmission (74). Furthermore, certain analogies can be found with respect to pathophysiological inflammation-induced depression. Here, low levels of the neurotransmitter serotonin are assumed to play a major role and rely mainly on tryptophan availability, which in turn is limited by the activity of indoleamine 2,3-dioxygenase (IDO) as a part of the kynurenine pathway (75, 76). IDO is inducible via IFN-γ, IL-1β, and IL-6 (77–79) or upon IAV infection (80). Degradation of tryptophan occurs through IDO in peripheral organs such as liver, intestine, and spleen (81) but also in brain-resident microglia, astrocytes, or recruited monocytes (54, 82). Altered levels of serotonin have been associated with a modulation of glutamatergic and GABAergic neurotransmission by altering the release of glutamate and GABA at the presynaptic side while further suppressing long-term potentiation (LTP) via inhibition of N-methyl-D-aspartate receptor (NMDA) glutamate receptor activation (83). Although we did not detect an upregulation of IDO mRNA levels in the brains of IAV-infected mice (not shown), studies on models of LPS-induced depressive-like behavior pointed toward a considerable role of the kynurenine pathway metabolites rather than altered brain levels of serotonin (84). Blood-derived metabolites of the kynurenine pathway can enter the brain via the large neutral amino acid transporter and cause excitotoxic effects (85). For example, levels of quinolinic acid are elevated upon peripheral inflammation (54), and it functions as an NMDA agonist and is found responsible for LPS-induced depression (86). Thus, the observed change in glutamatergic neurotransmission in our study might be a consequence of dysregulated serotonin levels or the underlying tryptophan metabolism. Next to adverse effects on glutamatergic neurotransmission, IAV infection led to altered expression of neurotrophins and their receptors. Previously, reduced expression levels of BDNF and GDNF have been discovered in brains of IAV-infected mice (12, 87), and models of poly(I:C) and LPS administration demonstrated reduced expression of BDNF, TrkB, and NGF in hippocampus and cortex of mice (22, 88). On the one hand, reduced
gene expression of BDNF has been associated with the development of depressive symptoms (89) and has been shown to modulate glutamatergic synaptic transmission by affecting presynaptic Ca2+ levels and glutamate release as well as phosphorylation of postsynaptic NMDA receptors (90). On the other hand, studies have shown that glutamate receptor activity has a reciprocal effect on neurotrophin production, indicating a strongly intertwined relationship (22). Notably, inflammation-mediated dysbalance of glutamatergic signaling has been described as a central mechanism in several neurologic disorders, such as schizophrenia (91), autism spectrum disorders (92), obsessive-compulsive disorder (93), or mood disorders, and further as a comorbidity in atherosclerosis or rheumatoid arthritis (94). Overall, this study shows that peripheral IAV infection is followed by temporal effects in the brain but without direct manifestation in neurodegenerative processes, as shown by our histopathological examination of neuronal markers. Nevertheless, our findings contribute to understanding how peripheral inflammation affects CNS integrity. Of note, reinfection of humans with influenza is likely to occur in intervals of 10 to 20 years (1) and may culminate in neurological implications (95). In this respect, a growing body of evidence supports the connection between peripheral infections and epigenetic changes in microglia that, in turn, render these cells more proinflammatory (96). As a consequence, synergistic effects of multiple infections acquired throughout life could lead to a previously unappreciated contribution to the development of chronic neuroinflammation, contributing to the initiation and progression of neurodegenerative diseases such as in AD or Parkinson disease (PD) (97, 98). Correspondingly, previous reports identified IAV infection as a risk factor for PD or the development of PD-like symptoms (99–101).

Altogether, in this study we are able to demonstrate that respiratory infection with IAV PR8/A/34(H1N1) results in the activation of microglia, dysbalanced glutamatergic neurotransmission, and neurotrophin gene expression. Furthermore, we highlight the influence of peripheral inflammation on the immune cell homeostasis of the brain influencing neuronal integrity. Lastly, we provide a novel and highly sensitive approach to characterize the composition of synaptosomes via flow cytometry. Establishing our new method allowed us to gain further insight into the mechanisms involved in the development of neuronal alterations described before but more importantly allows an application to various other research domains in a future perspective. We propose flow synaptometry as an additional tool, besides immunofluorescence microscopy and histological examination, to quantitatively assess the synapse integrity with high sensitivity under homeostatic conditions and to unravel modest synaptic changes during time-limited neuroinflammatory processes or the onset phases of neuroneurodegenerative conditions.

**MATERIALS AND METHODS**

**Animals.** The experiments were performed with female C57BL/6J mice (8 weeks old; purchased from Janvier, Cedex, France). Animals were group-housed under specific-pathogen-free conditions in individual ventilated cages with a 12-h day/night cycle with free access to food and water. All animal care was in accordance with institutional guidelines, experiments were performed in accordance with the German national guidelines for the use of experimental animals, and the protocol was approved by the Landesverwaltungsamt Saxony-Anhalt.

**IAV infection.** The virus stock of mouse-adapted IAV PR8/A/34(H1N1) was derived from Madin-Darby canine kidney cells as described previously (102). First, mice were anesthetized by intraperitoneal injection of ketamine (1%)–xylazine (10 mg/ml), and ointment was applied to keep eyes hydrated. Subsequently, mice were intranasally (i.n.) inoculated with 0.32 TCID50 of IAV diluted in phosphate-buffered saline (PBS), whereas mock-infected animals received PBS only (25).

**Isolation of BAL fluid, blood serum, tissue, and cells.** To collect bronchoalveolar lavage (BAL) fluid, the trachea was punctured and a vein catheter was carefully inserted. BAL was performed using 1 ml ice-cold PBS. BAL fluid was spun down at 420 × g for 10 min. Aliquots of BAL supernatants were stored at −80°C until further use. Tissue and blood serum collection and cell isolation were performed as described previously (103). In brief, mice were deeply anesthetized with isoflurane (CP Pharma, Burgdorf, Germany), and blood was taken from the inferior vena cava and incubated at 37°C. Upon centrifugation for 10 min at 1,500 × g, serum was collected and aliquots were stored at −80°C. Next, mice were perfused intracardially with 60 ml sterile PBS before brain extraction. For subsequent analysis by flow cytometry, brains were dissected into specific regions (cortex [CTX] and hippocampal formation [HPF]) according to the Allen mouse brain atlas (104), collected in separate tubes, and homogenized in a buffer containing Hanks’ balanced salt...
solution (HBSS; Gibco, New York, NY, USA), 13 mM HEPES (pH 7.3; Thermo Fisher Scientific, Waltham, MA, USA), and 0.68% glucose before sieving through a 70-μm cell strainer. The homogenate was fractioned on a discontinuous 30 to 70% Percoll gradient (GE Healthcare, Chicago, IL, USA). Immune cells were collected from the 30/70% Percoll interphase, washed in PBS, and immediately processed for subsequent flow cytometric analysis. For RNA isolation or synaptosome preparation, perfused brains were dissected as described above and either stored in RNAlater solution (Thermo Fisher Scientific) at −20°C or snap-frozen in liquid nitrogen and stored at −80°C until further use.

For immunohistochemistry (IHC) or immunofluorescence (IF), mice were perfused with 40 ml sterile PBS followed by 20 ml of 4% paraformaldehyde (PFA) solution. Brains were extracted and postfixed in 4% PFA for 4 h (IF) or 24 h (IHC) before being transferred to a 30% sucrose solution. After 2 days, IF samples were slowly frozen to −280°C in Tissue-Tek O.C.T. compound (Sakura Finetek Europe B.V., Alphen aan den Rijn, Netherlands) using liquid nitrogen and 2-methylbutane, whereas IHC samples were moved to PBS plus 0.1% NaN₃ and stored at 4°C until further use.

Cytokine immunoassay. Serum and BAL fluid cytokine levels were assessed using the LEGENDplex mouse inflammation panel (BioLegend, San Diego, CA, USA) according to the manufacturer’s instructions. Flow cytometry was performed with an LSR Fortessa instrument (BD, Franklin Lakes, NJ, USA), and data were analyzed using the LEGENDplex data analysis software (v8.0; BioLegend).

RNA isolation. Tissue samples were collected and stored in RNAlater solution as described above and prepared for RNA isolation as follows. Isolated brain regions were homogenized in lysis buffer using BashingBeads lysis tubes (Zymo Research Europe, Freiburg, Germany) and isolated using an AllPrep DNA/RNA minikit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Concentration and purity of isolated RNA were determined using a NanoDrop 2000 spectrophotometer (Thermo Fisher) and stored at −80°C until further use.

RT-qPCR. Gene expression levels of cytokines, inflammatory mediators, tight junction proteins, neurotrophins and neurotrophin receptors, synapse-associated proteins, and complement system were assessed in duplicates using either 30 ng isolated RNA or 600 ng for absolute quantification of viral load. Amplification was carried out with a TaqMan RNA-to-CT 1-step kit (Applied Biosystems, Foster City, CA, USA) or Power SYBR green RNA-to-CT 1-step kit (ThermoFisher Scientific) and LightCycler 96 (Roche, Basel, Switzerland) as previously described (105). Thermal cycling parameters were set as reverse transcription (48°C, 30 min), inactivation (95°C, 10 min) followed by 55 cycles of denaturation (95°C, 15 s), and annealing/extension (60°C, 1 min). In the case of the SYBR green RT-PCR, amplification was followed by a melting curve analysis. Utilized TaqMan gene expression assays (Applied Biosystems) are listed in Table 1. Self-designed primers were synthetized by Tib MolBiol (Berlin, Germany) and used at 100 nM final concentration (listed in Table 2). For relative quantification, expression of Hprt was chosen as a reference, and relative target gene mRNA levels were determined by the target gene/reference gene ratio and subsequently normalized to mean values of the control group. For absolute quantification of viral load, a standard curve was established by using a reference plasmid standard with known numbers of IAV nucleoprotein (NP) copies per sample (1.5 × 10¹ to 1.5 × 10⁹) (25).

### Table 1 TaqMan assays used for RT-qPCR analyses

| Protein | Gene | Assay ID |
|---------|------|---------|
| BDNF    | Bdnf | Mm04230607_s1 |
| C1qa    | C1qa | Mm00432142_m1 |
| C3      | C3   | Mm01232779_m1 |
| CCL2    | Ccl2 | Mm00441242_m1 |
| CD36    | Cd36 | Mm00432403_m1 |
| CD68    | Cd68 | Mm03047343_m1 |
| Claudin-5 | Cldn5 | Mm00727012_s1 |
| HPRT    | Hprt | Mm01545399_m1 |
| IDO     | Ido1 | Mm00492586_s1 |
| IFN-β   | Ifnb1 | Mm00439552_s1 |
| IFN-γ   | Ifng | Mm00801778_m1 |
| IGTP    | Igtp | Mm00497611_m1 |
| IL-1β   | Il1b | Mm00434228_m1 |
| IL-6    | Il6  | Mm00446190_m1 |
| iNOS    | Nos2 | Mm00404085_m1 |
| NGF     | Ngf  | Mm00443039_m1 |
| IRGM1   | Irgm1 | Mm00492596_m1 |
| p75NTR  | Ngfr | Mm00130963_m1 |
| TNF     | Tnf  | Mm00443258_m1 |
| TREM-2  | Trem2 | Mm04209422_m1 |
| TrkB    | Ntrk2 | Mm0435422_m1 |
| VGLUT1  | Slc17a7 | Mm00812886_m1 |
| ZO-1    | Tjp1 | Mm00493699_m1 |
**TABLE 2** Self-designed primer sequences used for RT-qPCR analyses

| Protein       | Gene     | Species | Sequence (5' to 3') |
|---------------|----------|---------|---------------------|
| Claudin-1     | Cldn1    | Mus musculus | ACTCTTGCTGAATCTGAACAGT |
|               |          |         | GGACAAGAGATTGGCATGAG |
| CCL5         | Cxcl9    | Mus musculus | GAGTTCAAGACCCTTGAAGT |
|               |          |         | AAAGTGGTGCTCTTGAGG  |
| CCL10        | Cxcl10   | Mus musculus | AACTGATCCATATCATGAGAC |
|               |          |         | GTGGCAAGTTGCTCAACAC |
| HPRT         | Hprt     | Mus musculus | GTCAAATTTCTCGTACCTGAGT |
|               |          |         | AATTACCTTTATGCTCCCGTGACTG |
| MX2          | Mx2      | Mus musculus | TCACAGAATGCGAGTGAGG   |
|               |          |         | CATTCCCTCTCTGACACATT |
| RSAD2        | Rsad2    | Mus musculus | GTCCATTTGCTGCTGAAAT  |
|               |          |         | GCCACGCTTCGAAACATCT |
| Nucleoprotein | NP       | Influenza A virus | GAGGGTGAATGCGACGAAAAAC |
|               |          |         | CAGGCAGCAGGAGGACTT  |

**Histopathology and immunohistochemistry.** For histopathology, brains collected from naive and infected mice were first dehydrated in zinc fixative (BD) followed by incubation in ethanol at concentrations increasing from 70% to 100%. Brains were then embedded in paraffin, and sagittal sections of 3 μm were mounted on object slides before deparaffinization in xylool and alcohol at concentrations decreasing from 100% to 70%. For hematoxylin and eosin staining, slides were first stained in Mayer's hemalum solution (Sigma-Aldrich, St. Louis, MO, USA) and rinsed with tap water and 0.1% HCl before applying eosin Y solution (Sigma-Aldrich) and rinsing in distilled water. Staining with antibodies against CD11b (dilution, 1:100; Synaptic Systems) and IBA1 (dilution, 1:200; Synaptic Systems, Göttingen, Germany), IBA1 (dilution, 1:200; Synaptic Systems, Göttingen, Germany) was performed at room temperature for 1 h after antigen retrieval using 10 mM citrate plus Tween 20 (pH 6.0) and blocking of endogenous peroxidase activity. Subsequently, slides were incubated with matching biotinylated secondary antibodies and developed using a VECTASTAIN ABC-HRP kit (PK-4000; Vector Laboratories, Burlingame, CA, USA). Upon incubation with DAB substrate, samples were counterstained with hematoxylin, rinsed with tap water, and then dehydrated in alcohol at increasing concentrations, propanol, and xylol before mounting. Finally, sections were imaged and analyzed using a Leica V1200 virtual-slide microscope (Olympus Life Science, Waltham, MA, USA) equipped with an Olympus V5 camera and a 40 × objective and Olympus V5-ASW imaging software (version 2.9.2, build 17565; Olympus Life Science).

To obtain samples for immunohistochemistry, sagittal sections of 20 μm from frozen brain tissue were transferred to SuperFrost plus (Thermo Scientific) slides. Antigen retrieval (10 mM citrate buffer, pH 6.0, 0.1% Tween 20) was performed at 96°C for 30 min. Blocking and permeabilization was performed in PBS plus 0.3% (vol/vol) Triton X-100 with 5% normal goat serum and unconjugated Fab(ab')2 goat anti-mouse IgG (H+L) antibody (1:500; Thermo Scientific) at 4°C for 2 h. Subsequently, samples were incubated with the following primary antibodies at 4°C overnight: anti-IBA1 (dilution, 1:200; HS-234 017; Synaptic Systems), and NeuN (dilution, 1:2,000; 188 011; Synaptic Systems), and NeuN (dilution, 1:2,000; 266 008; Synaptic Systems) was performed at room temperature for 1 h after antigen retrieval using 10 mM citrate plus Tween 20 (pH 6.0) and blocking of endogenous peroxidase activity. Subsequently, slides were incubated with matching biotinylated secondary antibodies and developed using a VECTASTAIN ABC-HRP kit (PK-4000; Vector Laboratories, Burlingame, CA, USA). Upon incubation with DAB substrate, samples were counterstained with hematoxylin, rinsed with tap water, and then dehydrated in alcohol at increasing concentrations, propanol, and xylol before mounting. Finally, sections were imaged and analyzed using an Olympus V9010 virtual-slide microscope (Olympus Life Science, Waltham, MA, USA) equipped with an Olympus V5 camera and a 40 × objective and Olympus V5-ASW imaging software (version 2.9.2, build 17565; Olympus Life Science).

**Flow cytometric analysis.** For flow cytometric analysis of cell phenotypes, freshly isolated cells were first incubated with Zombie NIR fixable dye (BioLegend, CA, USA) for live/dead discrimination and with anti-FcγRI/II receptor antibody (clone 93) to prevent unspecific binding of antibodies. Cells were further stained with the following fluorochrome-conjugated antibodies against cell surface markers in FACS buffer (PBS containing 2% fetal bovine serum and 0.1% sodium azide): eFluor 450-CD45 (clone 30-F11) and FITC-MHC I (clone 28-14-8A), APC-CD11b (clone M1/70) (purchased from ebioscience, San Diego, CA, USA), Brilliant Violet 421-CD86 (clone GL-1), Brilliant Violet 510-F4/80 (clone BM8), Brilliant Violet 605-F4/80 (clone BM8), Brilliant Violet 605-CD11b (clone M1/70), PerCP/Cy5.5-CD80 (clone 16-10A1), PE/Dazzle594-MHC II (I-A/I-E) (clone M5/114.15.2), and PE/Cy7-CX3CR1 (clone SA011F11) (purchased from BioLegend). Cells were acquired using a SP6800 spectral cell analyzer (Sony Biotechnology, San Jose, CA, USA), and obtained data were analyzed using FlowJo software (version 10.5.3; FlowJo LLC, OR, USA). Fluorescence minus one (FMO) controls were used to determine the level of autofluorescence.

**Preparation of synaptosomes from frozen brain samples.** Synaptosomes were obtained from cortex and hippocampal formation according to protocols published elsewhere (107) but with slight modifications. After the first homogenization steps, the crude membrane pellet P2 was fractioned on a discontinuous sucrose gradient with layers of 5 mM Tris-HCl pH 8.1, containing 0.32 M, 1.0 M, or 1.2 M sucrose at 80,000 × g at 4°C for 2 h. Subsequently, synaptic material was collected from the 1.0/1.2 M interphase and washed with SET buffer (320 mM sucrose, 1 mM EDTA, 5 mM Tris, pH 7.4) at 100,000 × g at 4°C for 1 h. Finally, the pelleted
Flow synaptometry. Aliquots of frozen synaptosomes were thawed in a water bath at 37°C and centrifuged for 10 min at 14,000 × g to remove sucrose-containing buffer. Supernatant was removed gently, and pellets were resuspended in fixation buffer (FoxP3 transcription factor staining buffer set, 00-5523-00; eBioscience) and incubated on ice for 45 min. Subsequently, samples were centrifuged again for 10 min at 14,000 × g, resuspended in permeabilization buffer (FoxP3 transcription factor staining buffer set) reconstituted with 10 % normal goat serum (NGS; ThermoFisher), and stained with primary antibodies against Gephyrin (ab136343; Abcam), GluR1 (ABN241; Sigma-Aldrich), Homer1 (MAB6889; R&D Systems, Minneapolis, MN, USA), synaptophysin (101 004; Synaptic Systems), and VGLUT1 (135 303; Synaptic Systems). Following incubation, samples were washed and resuspended in permeabilization buffer plus 10% NGS and stained with matching secondary antibodies: goat anti-mouse Alexa Fluor 405 (A31553; ThermoFisher), goat anti-rabbit Alexa Fluor 488 (ab150581; Abcam), goat anti-chicken Alexa Fluor 546 (A11040; ThermoFisher), and goat anti-guinea pig Alexa Fluor 647 (A21450; ThermoFisher). Finally, samples were washed once more and resuspended in SET buffer before adding the styrly dye FM4-64 (T13320; ThermoFisher) to a final concentration of 0.2 μg/ml (45). Samples were acquired using the Attune NxT flow cytometer (ThermoFisher) equipped with 405-, 488-, 561-, and 633-nm lasers. Voltages for forward-scatter light (FSC), side-scatter light (SSC), and fluorescence detection channels were set as the following: FSC, 400 V; SSC, 500 V; FL1, 400 V; FL2, 380 V; FL3, 440 V. For optimal acquisition of synaptosomes, the FSC-triggered detection was replaced by a fluorescence-triggered detection with FM4-64 in the BL3 channel (threshold set to 0.3 × 10⁴ to select only FM4-64-positive events). Furthermore, the event rate was kept below 300 events/s by utilizing the slowest flow rate in combination with an adequate dilution of the sample prior to measurement to reduce coincident particle detection. A size range from 300 to 1000 nm was applied to detect events in the FSC channel using red fluorescent silica beads with a diameter of 300 nm (DNG-L020; Creative Diagnostics, Shirley, NY, USA) and 1000 nm (DNG-L026; Creative Diagnostics) (45). Obtained data were analyzed using FlowJo software (version 10.5.3; FlowJo LLC, Ashland, OR, USA). FMO controls were used to determine the level of autofluorescence.

Western blot analysis of synaptic proteins. For Western blot analysis, aliquots of isolated synaptosomes were thawed in a water bath at 37°C and centrifuged for 10 min at 14,000 × g to remove sucrose-containing buffer. Pellets were lysed for 30 min at 37°C in radioimmunoprecipitation assay lysis buffer containing protease inhibitors, 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 % IGEPAL CA-630, 0.25 % Na-deoxycholate, and 1 mM NaF. Subsequently, samples were centrifuged for 30 min at 100,000 × g at 4°C. The supernatant was removed and used for further separation on a 12.5 % SDS-PAGE with loading buffer (50 mM Tris-HCl, pH 6.8, 100 mM dithiothreitol, 2 % SDS, 1.5 mM bromophenol blue, 1 M glycerol). Next, proteins were transferred to nitrocellulose membranes and incubated with antibodies against VGLUT1 (dilution, 1:1000; Synaptic Systems) at 4°C overnight. For the loading control, membranes were incubated with a 1:1,000 dilution of anti-β-actin antibody (4970; Cell Signaling Technology, Cambridge, UK). Following incubation, membranes were incubated with matching horseradish peroxidase-conjugated secondary antibodies at room temperature for 2 h before revealing bound antibodies using enhanced chemiluminescence assay. Densitometric analysis of blots was performed using ImageJ with Fiji distribution (106).

Electron microscopy of synaptosomes. Electron microscopic analysis of isolated synaptosomes was performed according to Breuel et al. (108) but with minor modifications. Synaptosomes were fixed in 0.1 M cacodylate buffer (pH 7.4) with 2.5% paraformaldehyde and 2.5 % glutaraldehyde in the refrigerator overnight. The suspension then was postfixed in 0.1 M cacodylate buffer containing 1% osmium tetroxide for 30 min and rinsed with distilled water subsequently. This was followed by a stepwise dehydration in a graded series of ethanol (50% to 100%) for 5 min each. Finally, the suspension was embedded in Durcupan ACM (Honeywell Fluka, Morristown, NJ, USA) by dropping it carefully into the tubes. The resin polymerized at 70°C for 3 days. For sectioning, the tubes were cut off, and the complete block of Durcupan with the pellet of synaptosomes at the tip was directly placed in an ultramicrotome (Ultracut E; Reichert-Jung, Wetzlar, Germany). Ultrathin sections of 50 to 70 nm were collected on Formvar-coated slot grids of copper and examined with a LEO 912 transmission electron microscope (Carl Zeiss, Oberkochen, Germany) and imaged with a MegaScan 2K charge-coupled device camera (Gatan Inc., CA, USA) using DigitalMicrograph software (version 2.5).

Statistical analysis. The body weight was compared by multiple t tests with Holm-Sidak post hoc correction. Data from flow cytometry and RT-qPCR were compared by Student’s t test with Welch’s correction and Western blot analysis by one-way analysis of variance (ANOVA) with Holm-Sidak post hoc correction using GraphPad Prism 7 (GraphPad Software, CA, USA) and R (version 4.0.3) (109) with the “lattice” package (110). Data shown are representative of three independent experiments. In all cases, results are presented as arithmetic means and were considered significant, with a P value of <0.05.

Ethics approval. The study was performed in accordance with the German national guidelines for the use of experimental animals, and the protocol was approved by the Landesverwaltungsamt Sachsen-Anhalt. Food and water were available ad libitum. All efforts were made to minimize the suffering of mice used in this investigation.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, EPS file, 2.5 MB.
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