Microglial population expansion occurs in response to neural damage via processes that involve mitosis and immigration of bone marrow-derived cells. However, little is known of the mechanisms that regulate clearance of reactive microglia, when microgliosis diminishes days to weeks later. We have investigated the mechanisms of microglial population control in a well-defined model of reactive microgliosis in the mouse dentate gyrus after perforant pathway axonal lesion. Unbiased stereological methods and flow cytometry demonstrate significant lesion-induced increases in microglial numbers. Reactive microglia often occurred in clusters, some having recently incorporated bromodeoxyuridine, showing that proliferation had occurred. Annexin V labeling and staining for activated caspase-3 and terminal deoxynucleotidyl transferase-mediated dUTP nick end-labeling showed that apoptotic mechanisms participate in dissolution of the microglial response. Using bone marrow chimeric mice, we found that the lesion-induced proliferative capacity of resident microglia superseded that of immigrant microglia, whereas lesion-induced kinetics of apoptosis were comparable. Microglial numbers and responses were severely reduced in bone marrow chimeric mice. These results broaden our understanding of the microglial response to neural damage by demonstrating that simultaneously occurring mitosis and apoptosis regulate expansion and reduction of both resident and immigrant microglial cell populations. (Am J Pathol 2007, 171:617–631; DOI: 10.2353/ajpath.2007.061044)
Acute activation of microglia as a result of neural injury or pathology quickly leads to reactive microgliosis, a cardinal feature being expansion in the number of microglia in the affected region. Increase in cell number originates in part from recruitment of myeloid cells, proliferation, or migration from juxtaposed regions. The state of reactive microgliosis dissolves days to weeks later, according to an inherently tightly regulated schedule, which has been suggested to involve microglial apoptosis. The cellular population control of immigrating and resident microglia should be comparable if immigrating bone marrow (BM)-derived cells are to take part fully in regular microglial tasks.

To address these fundamental questions, we investigated whether resident and immigrating microglia are governed by similar mechanisms of population control, i.e., cellular multiplication by mitosis and reduction of the cellular population by apoptosis. Reactive microgliosis was induced in the dentate gyrus and hippocampus in unmanipulated and green fluorescent protein (GFP)-BM-chimeric mice by transection of the perforant pathway (PP) projection in the entorhinal cortex. Our results show that microglial expansion is balanced by simultaneously occurring mitosis and apoptosis. The axonal lesion-induced mitotic activity of resident microglia supersedes that of BM-derived immigrant microglia, whereas the kinetics of lesion-induced apoptotic responses are comparable.

Materials and Methods

Animals

C57BL/6J mice (Taconic, Skensved, Denmark; or Harlan, Allerød, Denmark) were used for studies of nonchimeric mice. For BM-chimeric studies, C57BL/6 congenic B6.SJL-PtprcaPepcb/BoyJ mice (The Jackson Laboratory, Bar Harbor, ME), which express the CD45.1 allotype and are hence abbreviated B6(CD45.1), were used as transplant recipients and C57BL/6-Tg(UBC-GFP)30Scha/J mice (The Jackson Laboratory) as BM donors. This combination of mice was chosen so that immigrating cells could be identified in chimeric mice by two criteria: positive expression of GFP and CD45.2, which is normally expressed in C57BL/6 mice. Expression of the GFP transgene, however, proved by itself to be a sufficiently strong and reliable signal. A group of nonirradiated B6(CD45.1) mice (The Jackson Laboratory) was included to control for potential differences from C57BL/6 mice. Mice were housed and bred and animal experiments performed at the Laboratory of Biomedicine at the University of Southern Denmark and at the Department of Medical Microbiology and Immunology at Aarhus University. All animal experiments were performed according to Danish law and protocols approved by the Danish Ethical Animal Care Committee.

PP Lesion

Microglial activation in the hippocampus and dentate gyrus was induced by the anterograde axonal and terminal degeneration of presynaptic elements resulting from transection of the entorhino-hippocampal PP projection, performed as previously described. Mice were anesthetized with a mixture of ketamine and xylazine (for flow cytometry) or fentanyl citrate, fluanisone, and diazepam (for histology) and fixed in a stereotaxic frame (Stoelting, Wood Dale, IL). The PP transection was made with a wire knife (David Kopf Instruments, Tujunga, CA) angled 10 degrees lateral and rotated 15 degrees rostral and the nosebar set at −3 mm. The closed wire knife was inserted 2.1 mm lateral to the lambda and 0.3 mm caudal to the lambdoid suture through a drilled trepanation. A 3.1-mm-long cut was made in the entorhinal cortex starting 3.4 mm ventral to the meninges. Mice were supplied with eye ointment and postoperative injections of saline and buprenorphine. Animals were placed in a warm environment for postoperative recovery.

BM Transplantation

Recipient mice were irradiated with 9.5 Gy in a single dose from a 137Cs source (Rise National Laboratory, Roskilde, Denmark) and were reconstituted with BM cells obtained from GFP transgenic donor mice. Donor BM cells were harvested by flushing the medullary canal of the femoral and tibial diaphysis with RPMI 1640 medium (Gibco, Paisley, UK). After centrifugation, the cell suspension was filtered and transplanted by intravenous injection. The transplanted mice were supplied with oxytetracycline (2 g/L Terramycin veterinary 20%; Plizer, Amoise, France) in the drinking water for 3 days after transplantation. Chimerism was assessed by expression of GFP in blood cells at the time of sacrifice. Positive GFP expression was determined using autofluorescence levels in blood cells from non-Tg mice as negative controls. Flow cytometric analysis showed that 91.6 ± 1.1% (mean ± SEM) of blood cells were GFP*, indicating that successful reconstitution had occurred.

Real-Time Polymerase Chain Reaction (PCR)-Based Testing for UbC-GFP Zygosity

Genomic DNA was extracted from external ear tissue, obtained from labeling of animals for identity, by alkaline lysis in 0.1 mol/L KOH for 1 hour at 95°C followed by KH2PO4 neutralization. Extracts were diluted 20-fold, and 2 μl were applied for 25-μl real-time PCR. The reaction mixture contained 1× RealQ master (Ampliqon; Bie and Berntsen A/S, Rosdovre, Denmark) and for transgene PCRs 300 nmol/L of each of the two primers (5'-ATTGT-CCGCTAATTTCTGGCGGT/5'GCTCGACCAGGATGG-GCACCC-3') (TAG, Copenhagen, Denmark) and SYBR Green diluted 20,000-fold. Lymphotixin-α (LT-α) reference gene PCRs contained 400 nmol/L of the two primers (5'-GTCCAGCCTTCTCCTCCTAAT/GTCTTTTG-AAATTCCGGATACAC-3') and 50 nmol/L TaqMan probe.
(5'-CCTCCATGCGCTCTCCCTGAGC-3'). For PCR and detection in real time, an iCycler (Bio-Rad, Herlev, Denmark) was used. The thermocycling protocol was 95°C for 15 minutes to activate the RealQ enzyme followed by 40 cycles of 95°C for 15 seconds, 62°C for 30 seconds, and 72°C for 45 seconds.

Real-Time PCR-Based mRNA Analysis

RNA was extracted using RNaseasy Protect mini kits (Qiagen, Hilden, Germany) or TRizol (Invtrogen, Taastrup, Denmark), as previously described for PP-lesioned hippocampus samples.22,23 Complementary DNA was synthesized from 0.4 µg of total RNA by reverse transcription, using previously established conditions.23 For real-time PCR, a Bio-Rad iCycler was used. Each reaction contained a total volume of 25 µl, 5 of which were diluted cDNA. The other components were RealQ master mix (1.5 mmol/L magnesium at 1°C; Ampliqon; Bie and Berntsen A/S), 1 to 2 mmol/L of additional MgCl2, SYBR Green, fluorescein, and 300 nmol/L of each PCR primer. Newly designed primer sets were validated to specifically amplify the respective target only. These were M-CSF (5'-CGGTCCTTCTCTGCATG/ACGTCGTCGGACACCTTGT-3'; annealing temperature, 60°C; magnesium concentration, 3.5 mmol/L), M-CSFR (5'-GCATT-ACACCTGGACCTAAG/AGACCTGGAATGTGACCTGTAT-3'; annealing temperature, 55°C; magnesium concentration, 3.0 mmol/L), GM-CSF, which was targeted by two different primer sets (5'-CATGTAAGGCACTTCAAAGAG/AAGACTTCACTACCTCAAACTCAA-3'; annealing temperature, 56°C; magnesium concentration, 3.5 mmol/L) and 5'-GGGCAATTTCACCAAACTCAGTGCG-3' (annealing temperature, 60°C; magnesium concentration, 2.5 mmol/L). Amplification of CD11b and HPRT1 was done using previously published primer and probe sequences, and FAM- or HEX-labeled TaqMan probes instead of SYBR Green, respectively.23 Each cDNA was subjected to triplicate real-time PCR analysis. Calibrator samples were included on all plates. Data were normalized using HPRT1 reference gene, averaged, and calibrated against the calibrator ratio. Data are presented as relative values.

Labeling of Proliferating Cells in Vivo

For in vivo detection of microglial mitosis, proliferating cells were labeled with 5-bromo-2'-deoxyuridine (BrdU), which is incorporated into DNA during mitotic S phase. Each mouse was injected with a 90-mg/kg dose of BrdU dissolved in phosphate-buffered saline (PBS; 10 mg/ml) intraperitoneally three times at 8-hour intervals for the last 24 hours before perfusion. Mice used for BrdU histology were injected with 50 mg/kg 1 hour before sacrifice.

Flow Cytometry

Mice were sacrificed under pentobarbital anesthesia by exsanguination and intracardiac perfusion with 20 ml of PBS. The brains were removed, and the hippocampus and dentate gyrus were dissected out en bloc from the contralateral and lesioned hemispheres. These samples did not include tissue surrounding the wire knife lesion because this was in the entorhinal cortex, and any adherent choroid plexus was removed.24 Hippocampal tissue was homogenized through a 70-µm cell strainer (BD Falcon, Franklin Lakes, NJ) in RPMI 1640 medium (Gibco) containing 10% fetal bovine serum (FBS; Gibco). After centrifugation, cells were incubated with anti-FCyRIIa receptor antibody (BD Biosciences, Erembodegem, Belgium) and 50 µg/ml of Syrian hamster Ig (Jackson Immunoresearch, West Grove, PA) in RPMI 1640 medium with 10% FBS at room temperature for 45 to 60 minutes to block nonspecific staining.11,22

For annexin V analysis, surface antigen labeling was performed by incubating cells for 30 minutes at room temperature with phycoerythin (PE)-conjugated anti-CD45 antibody and fluorescein isothiocyanate-conjugated anti-CD11b (BD Biosciences) in RPMI 1640 medium with 10% FBS. For studies in chimeric mice, anti-CD11b was necessarily omitted because the FL1 channel was occupied by GFP fluorescence. Essentially all CD45dim cells coexpress CD11b, and relative levels of CD45 can distinguish microglia (CD45dim) from leukocytes (CD45high).25-27 Cells were then washed once in RPMI 1640 medium with 10% FBS and twice in annexin V-binding buffer (10 mmol/L HEPES, 140 mmol/L NaCl, and 2.5 mmol/L CaCl2, pH 7.4) before incubation for 15 minutes at room temperature in a mixture of APC-conjugated annexin V (BD Biosciences), which labels apoptotic cells, and the viability marker 7-aminomycin D (7-AAD), which allows exclusion of dead cells (BD Biosciences).

For BrdU analysis, cells from chimeric and nonchimeric mice were washed in staining buffer (Hanks’ balanced salt solution containing 2% FBS and 0.1% sodium azide) and for surface antigen labeling incubated with PE-conjugated anti-CD45 and PerCP-Cy5.5-conjugated anti-CD11b antibodies (BD Biosciences). Cells were then washed in staining buffer and fixed, permeabilized, and stained with APC-conjugated anti-BrdU antibody for intracellular BrdU detection according to instructions provided with the APC BrdU flow kit (BD Biosciences). All stainings were analyzed on a BD FACSCalibur flow cytometer and BD CellQuest Pro software (BD Biosciences, San Jose, CA). Isotype, antigen omission (for BrdU), and autofluorescence (for GFP, 7-AAD, and annexin V) controls served to determine fluorescence levels for positive staining. Specificity of annexin V binding was additionally tested by blocking with recombinant annexin V.

For quantification of flow cytometry data, cells were gated on side scatter (SSC) versus CD11b and SSC versus either intermediate expression of CD45 to identify parenchymal microglia (CD45dim/CD11b+), or high levels of CD45 to identify macrophages (CD45high/CD11b+). Note that identification of apoptotic microglia in chimeric
mice was based solely on SSC versus CD45dim expression for reasons outlined above. All microglia/macrophages included in annexin V analyses were additionally gated to show only viable cells, by excluding late apoptotic and necrotic cells positive for 7-AAD from the analysis. Note that certain flow cytometry profiles included in figures are shown for descriptive purposes only, and may have alternate gating strategies; this information is specified in the figure legends.

Numbers of microglia or macrophages in one hippocampus were calculated as the number of gated CD45dimCD11b+ microglia or gated CD45highCD11b+ macrophages multiplied by the reciprocal fraction of the sample volume used and multiplied by two. This last step was taken because each hippocampal sample was split in two, ie, half was used for annexin V analysis and the other half for BrDU incorporation. For C57BL/6 mice, the data are presented as the average of these duplicates. For chimeric mice, the data are presented from our analysis of BrDU-stained cells because both CD45 and CD11b antibodies could only be included there. Note that number data generated from groups of B6(CD45.1) and C57BL/6 mice used for comparison with chimeric mice were also estimated using this approach. Numbers of microglial and macrophage subpopulations were calculated by applying the proportion of the subset to the total number estimate. Preparation and staining of hippocampal homogenate inevitably leads to cells loss, and the cell numbers and proportions presented are based on cells remaining in suspension.

Histology

For stereological quantification of total microglial numbers in the dentate gyrus, vibratome sections were processed for immunohistochemical visualization of CD11b+ microglia. PP-lesioned mice were sacrificed under pentobarbital anesthesia by exsanguination and intracardiac perfusion with 5 ml of Sørensen’s phosphate buffer (25 mmol/L K2HPO4 and 125 mmol/L Na2HPO4, pH 7.4) followed by 20 ml of Sørensen’s phosphate buffer containing 4% paraformaldehyde. The brains were additionally fixed 1 hour in Sørensen’s phosphate buffer containing 4% paraformaldehyde on ice and 21 hours in 4% paraformaldehyde, and 300 g of sucrose diluted in a mixture of 300 ml of NaPO4 buffer for long-term storage. Sections selected for CD11b labeling were rinsed in Tris-buffered saline (TBS; 50 mmol/L Trisma base and 110 mmol/L NaCl, pH 7.4), endogenous peroxidase activity was blocked in TBS (TBS containing 1% Triton X-100 and 10% FBS) for 1 hour at room temperature. Incubation with the anti-CD11b antibody (Serotec, Hamar, Norway) in staining buffer was done overnight at 4°C. Subsequent incubations with secondary goat anti-rat biotinylated antibody (Amersham Biosciences, Little Chalfont, UK) and horse-radish peroxidase conjugated streptavidin in staining buffer were performed at room temperature for 1 hour. Labeling was visualized with 3,3’-diaminobenzidine, and sections were mounted on gelatinized microscope slides, dried overnight, stained with toluidine blue, dehydrated in graded ethanol, cleared in xylene, and coverslipped in Depex mounting medium.

Visualization of apoptotic CD11b+ microglia expressing activated caspase-3 was done in 16-μm-thick cryostat sections mounted on microscope slides. Tissue sections were rinsed in TBS and in TBS containing 1% Triton X-100 (TBS-T) before incubation in staining buffer for 1 hour at room temperature. The tissue was subsequently incubated with rat anti-mouse CD11b antibody (Serotec) and rabbit anti-human activated caspase-3 antibody (Cell Signaling Technology, Boston, MA) diluted in staining buffer overnight at 4°C. Afterward the tissue was rinsed in TBS-T and incubated with Alexa Fluor 568-conjugated goat anti-rat IgG antibody (Invitrogen) and Alexa Fluor 488-conjugated goat anti-rabbit IgG antibody (Invitrogen). The tissue was finally rinsed in TBS and stained with 4’,6-diamidino-2-phenylindole (DAPI). After a final rinse in TBS, sections were coverslipped in ProLong Gold anti-fade medium (Invitrogen). Visualization of activated caspase-3-positive cells in the free-floating vibratome sections was done as described for CD11b using 3,3’-diaminobenzidine as chromogen.

Fragmented DNA in apoptotic microglia was visualized applying terminal deoxynucleotidyl transferase-mediated dUTP nick end-labeling (TUNEL) together with CD11b and DAPI staining. Free-floating 70-μm-thick vibratome sections were rinsed in TBS and TBS-T before preincubation in staining buffer at room temperature and incubation with CD11b antibody (Serotec) at 4°C overnight. Sections were then rinsed in TBS-T and incubated for 1 hour at room temperature with an Alexa Fluor 568-conjugated goat anti-rat IgG antibody (Invitrogen) and Alexa Fluor 488-conjugated goat anti-rabbit IgG antibody (Invitrogen). The tissue was finally rinsed in TBS and stained with 4’,6-diamidino-2-phenylindole (DAPI). After a final rinse in TBS, sections were coverslipped in ProLong Gold anti-fade medium (Invitrogen). Visualization of activated caspase-3-positive cells in the free-floating vibratome sections was done as described for CD11b using 3,3’-diaminobenzidine as chromogen.

For confocal analysis of microglial clusters, free-floating 70-μm-thick vibratome sections were rinsed in TBS and TBS-T before preincubation in staining buffer at room temperature. The tissue was then incubated with rat anti-mouse CD11b antibody (Serotec) overnight at 4°C. Sections were rinsed in TBS-T and incubated for 1 hour at room temperature with an Alexa Fluor 568-conjugated goat anti-rat IgG antibody. After rinsing in TBS-T fluorescein isothiocyanate-conjugated dUTP nick end labeling was performed according to kit instructions (Roche Diagnostics, Hvidovre, Denmark). Finally, sections were rinsed in PBS, mounted on gelatinized microscope slides, dried, stained with DAPI, and coverslipped in ProLong Gold anti-fade medium.

For visualization of mitotic microglia, 16-μm-thick cryostat sections were rinsed in TBS and in TBS-T before
immersion in 1 mol/L HCl for 10 minutes on ice, 2 mol/L HCl for 10 minutes at room temperature, and 2 mol/L HCl for 20 minutes at 37°C. Tissue was rinsed in TBS-T and incubated with staining buffer for 1 hour at room temperature followed by incubation with biotinylated tomato lectin (Sigma, Copenhagen, Denmark) and rat anti-BrdU antibody (Abcam, Cambridge, UK) diluted in staining buffer at 4°C overnight. The tissue was then rinsed in TBS-T and incubated with Alexa Fluor 568-conjugated goat anti-rat IgG antibody and Alexa Fluor 488-conjugated streptavidin. The tissue was finally rinsed in TBS and stained with DAPI, and sections were coverslipped in Prolong Gold anti-fade medium.

In situ visualization of GFP 

BM-derived microglia was done in 16-μm-thick cryostat sections. Sections were rinsed in TBS before preincubation in TBS containing 10% FBS for 1 hour. The tissue was incubated overnight with rat anti-mouse CD11b antibody (Serotec) in TBS containing 10% FBS and 0.5% Triton X-100. The sections were subsequently rinsed in TBS and incubated with an Alexa Fluor 568-conjugated rabbit anti-rat IgG antibody for 1 hour, rinsed again in TBS, and coverslipped in Prolong Gold anti-fade medium with DAPI. Fluorescence stainings were analyzed using an Olympus BX51 fluorescence microscope and DP70 digital color camera.

**Stereology**

Histological quantification of microglia was based on the protocol for counting neurons in the hippocampus, by applying the optical fractionator adapted for unbiased counting of microglia in the dentate gyrus as previously described. All counting was performed using the computer-assisted stereological toolbox (CAST)-Grid software (Visiopharm, Hørsholm, Denmark) connected to an Olympus BX50 microscope equipped with motorized stage and focus units. Each brain was sectioned horizontally in four parallel series of sections. One was used for stereological counting giving a section sampling fraction ssf of 1/4. Cells were sampled in counting frames of 644 to 988 μm² [a(frame)] moved in x and y steps of 100 μm × 100 μm [a(step)]. The area sampling fraction asf was calculated as a(frame)/a(step). The thickness sampling fraction tsf was calculated as the height of the optical dissector probe h (8 or 10 μm) divided by the average height of the sections t (tsf = h/t). Total cell number N was estimated using the equation: N = ΣQ⁻¹ · 1/tsf · 1/ssf, ΣQ⁻¹ being the number of cells counted. In the case of microglial clusters, each cluster was counted as one cell and identified by the most clearly defined nucleus. Coefficients of error and variation were calculated as previously described.

**Statistics**

Data are presented as mean ± SEM. Groups of contralateral and deafferented hippocampi were analyzed with a paired t-test. For comparison of the proportions of GFP⁺ immigrant microglia between the contralateral and lesioned hippocampi, the level of statistical significance was set at a one-tailed P value of <0.05. The level of statistical significance for other analyses was set at a two-tailed P value of <0.05. For comparison of three or more groups, a one-way analysis of variance was applied and individual groups compared using Bonferroni’s post test. P values are indicated as follows: *P < 0.05, **P < 0.01, and ***P < 0.001.

**Results**

**Reactive Changes in the CD11b⁺ Microglial Population in the Dentate Gyrus and Hippocampus**

The degeneration of the PP axons and their presynaptic terminals induced reactive microgliosis in the deafferented areas of the dentate gyrus and hippocampus (Figure 1), as previously described. Reactive microgliosis was observed in the PP termination areas in the outer part of the dentate molecular layer, the CA3 stratum moleculare, and the stratum lacunosum moleculare of the ipsi- and contralateral CA1 (Figure 1), the latter attributable to transection of the ipsilateral and crossed temparoammonic tracts. Microglial activation was therefore also observed in white matter in the alveus and fimbria fornix attributable to degeneration of transected commissural axons (Figure 1). Three days later, microglia had acquired a characteristic dense hyperramified morphology and formed distinct clusters of reactive cells throughout the deafferented areas (Figure 2, A and B). At 5 and 7 days after lesion, the microglial reaction was almost confluent and separation of individual microglial cells unclear (Figure 2B). Using confocal laser-scanning microscopy on clusters identified by immunofluorescence staining for CD11b and propidium iodide as a nuclear marker, it was determined that these clusters represented aggregates of microglial cells, i.e., nuclei separated by a rim of CD11b-stained plasma membrane (Figure 2C). Analysis of ~50 microglial clusters revealed that most clusters represented two reactive microglial cells, but clusters consisting of three to four cells were also observed (Figure 2C, displayed 3 days after lesion).

**Axonal Lesion-Induced Increase in Microglial Numbers**

A cardinal feature of reactive microgliosis after PP lesion is microglial mitosis giving rise to an increase in the number of microglia in the deafferented area. We have previously used stereological methods to estimate that there are 12,300 resting, ramified CD11b⁺ microglia in the molecular and granular cell layer of the dentate gyrus of normal mice. Similar numbers have been generated by others. Here, the same method was applied to estimate the total number of reactive microglia in the molecular and granular cell layer of the deafferented dentate gyrus. Numbers of microglia were significantly increased 3, 5, and 7 days after PP transection (22,700,
Comparison of the groups with 3-, 5-, or 7-day survival after lesion showed no statistically significant differences between groups (Table 1), indicating that there was no clear peak in numerical expansion in the dentate gyrus at these times. Reactive microglia that occurred in multicellular clusters (Figure 2, B and C) were necessarily counted as a single cell, because individual cells within the clusters could not easily be resolved by conventional light microscopy. This probably underestimated the total number of microglia in the deafferented dentate gyrus, as well as the fold increase compared with unmanipulated dentate gyrus, where only individual resting microglia were observed. To determine the proportion of clustered microglia, microglial counts were classified in three categories: 1) reactive microglia with one nucleus; 2) microglial clusters; and 3) resting, ramified microglia (Table 2). Depending on the survival time after lesion, microglial clusters constituted 72 to 76% of the counted microglia. The actual number of microglia could then potentially be twofold higher or even more than the estimates listed in Table 1.

Although microglia is the most prominent myeloid cell population in the PP-deafferented dentate gyrus, CD11b has been shown to additionally stain several smaller populations of myeloid cells after PP lesion.10,11,22 By flow cytometry, microglia and macrophages can be identified and quantified based on their expression of CD45 and CD11b, as illustrated in the flow cytometry dot plots in Figure 3A. Resting and reactive microglia express CD11b (CD11b⁺) and intermediate levels of CD45 (CD45dim) and are hence referred to as CD45dimCD11b⁺, whereas macrophages express considerably higher levels of CD45 (CD45highCD11b⁺). The dot plots that resulted from analysis of isolated whole hippocampus including the dentate gyrus indicate that the number of microglia increased 3 and 7 days after lesion and that macrophages contributed to the response, especially at 3 days after lesion (Figure 3A). This is consistent with previous flow cytometry studies showing that whereas microglial proportions increase between 2 and 5 days after lesion, macrophage proportions decline.22,24 This lesion-induced increase in microglial and macrophage numbers in the hippocampus was quantified. Because a known fraction of hippocampal cells were analyzed by flow cytometry, this allowed for estimation of the total number of microglia and macrophages from unlesioned and contralateral and deafferented hippocampi at different times after lesion (Figure 3). Number estimates generated from flow cytometric analyses include deafferented as well as unaffected hippocampal subregions and reflect inevitable cell loss during the preparation and staining of cell suspensions. Numbers of microglia increased 3, 7, 14, and 28 days after PP transection (41,000, 38,900, 30,100, and 31,100, respectively), compared with unlesioned mice (20,300) (Figure 3B). As with our stereological estimates, no significant difference was detected in deafferented hippocampus between 3 and 7 days. By 14 days, however, microglial numbers had begun to decline. For macrophages, the lesion-induced increase in numbers was only observed 3, 7, and 14 days after lesion (Figure 3C). Three days after lesion only 750 macrophages were present in deafferented hippocampus compared with 41,000 microglia (<2%; Figure 3, B and C).
fewer macrophages were already observed by 7 days, which then comprised only \( \sim 0.5\% \) of the CD11b\(^+\) population.

Table 1. Estimates of Total CD11b\(^+\) Microglial Cell Numbers

<table>
<thead>
<tr>
<th>Animal</th>
<th>Day 3 ( N ) (CE)</th>
<th>Day 5 ( N ) (CE)</th>
<th>Day 7 ( N ) (CE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>24,300 (0.106)</td>
<td>26,000 (0.100)</td>
<td>17,300 (0.094)</td>
</tr>
<tr>
<td>2</td>
<td>23,100 (0.088)</td>
<td>25,100 (0.086)</td>
<td>25,900 (0.120)</td>
</tr>
<tr>
<td>3</td>
<td>22,200 (0.078)</td>
<td>21,900 (0.100)</td>
<td>25,500 (0.111)</td>
</tr>
<tr>
<td>4</td>
<td>20,200 (0.097)</td>
<td>16,300 (0.102)</td>
<td>26,900 (0.093)</td>
</tr>
<tr>
<td>5</td>
<td>23,600 (0.090)</td>
<td>22,900 (0.083)</td>
<td>24,700 (0.112)</td>
</tr>
<tr>
<td>Mean</td>
<td>22,700 (0.092)</td>
<td>22,400 (0.095)</td>
<td>24,100 (0.107)</td>
</tr>
<tr>
<td>SD</td>
<td>1600</td>
<td>3800</td>
<td>3900</td>
</tr>
<tr>
<td>CV</td>
<td>0.070</td>
<td>0.170</td>
<td>0.161</td>
</tr>
</tbody>
</table>

Microglial Proliferation after Axonal Lesion

The peak microglial proliferative response was observed 3 days after PP transection with 27\% (11,000) BrdU\(^+\) microglia in the deafferented hippocampus (Figure 4, A–C). Seven days after PP transection a lesion-induced...
microglial proliferative response was still present compared with the contralateral hippocampus when evaluated by the number of BrdU$^+$ microglia (Figure 4C). There was no statistically significant lesion-induced proliferative response in the macrophage population at any of the investigated times of observation evaluated by the percentage of BrdU$^+$ macrophages (Figure 4A, upper quadrants; and data not shown). This suggests that microglia proliferate readily in response to PP lesion, whereas macrophages rarely do. Because microglial clusters were prominent 3 days after lesion (Table 2), coinciding with microglial peak proliferation (Figure 4, A–C), it was investigated whether these clusters represented nests of proliferating microglia. PP-lesioned mice were pulse-labeled with BrdU 1 hour before perfusion, and the proliferation status of clustered tomato lectin$^+$

Figure 3. Microglia and macrophages in normal and deafferented hippocampus. A: Flow cytometry profiles, showing all viable (7-AAD-negative) cells in the forward/side scatter gated population, illustrate microglia (CD45dimCD11b$^+$, bottom right quadrants) and macrophages (CD45highCD11b$^+$, top right quadrants) in unmanipulated mice or 3 and 7 days after PP transection. Quantification of flow cytometry profiles reveals increases in both total numbers of microglia (B) and macrophages (C) in lesioned (L) hippocampi from C57BL/6 mice compared with contralateral (C) or unmanipulated (U) mice. Lesioned: 3 days ($n = 5$), 7 days ($n = 7$), 14 days ($n = 7$) and 28 days ($n = 6$). Unlesioned, $n = 6$. *$P < 0.05$, **$P < 0.01$, and ***$P < 0.001$.

Figure 4. Microglial proliferation after PP lesion in C57BL/6 mice. A: Flow cytometry profiles, gated to show all CD45$^+$CD11b$^+$ cells, illustrate significant proliferation by BrdU$^+$ mitotic microglia (bottom quadrants) and little incorporation of BrdU$^+$ by macrophages (top quadrants) 3 days after PP lesion. Quantification of the proportion (B) and number (C) of BrdU$^+$ microglia are displayed in bar graphs, after gating on CD45dimCD11b$^+$ cells only. Microglial mitosis was visualized in histological sections using a combination of tomato lectin to identify microglia (green) and BrdU$^+$ staining (red). D: As shown in the merged photograph, BrdU has been incorporated into several nuclei in both a juxtavascular cluster (white arrow) and an intraparenchymal cluster (blue arrow). **$P < 0.01$, and ***$P < 0.001$. Scale bar = 10 μm.
microglia was assessed by conventional fluorescence microscopy. BrdU incorporation was observed in some nuclei within the microglial clusters (Figure 4D).

Microglial Apoptosis after Axonal Lesion

The apoptotic response was investigated using annexin V as marker for apoptosis (Figure 5, A–C). Annexin V binds to phosphatidylserine exposed in the outer leaflet of the plasma membrane of apoptotic cells, and hence only works in intact cells because phosphatidylserine is normally present in the inner leaflet of the plasma membrane. Dead cells were excluded from the flow cytometric analysis by applying the viability marker 7-AAD. A lesion-induced apoptotic response was observed 3, 7, and 14 days after PP transection and was reflected in an increase in the total number of annexin V+ microglia (Figure 5C). At 28 days, numbers of annexin V+ microglia were significantly elevated versus unlesioned hippocampi, but not compared with the contralateral hippocampus. However, because of the changing number of microglia a lesion-induced increase in the proportion of apoptotic annexin V+ microglia was only observed 3 days after lesion (Figure 5B). In the macrophage population a lesion-induced apoptotic response was only observed in the number of annexin V+ macrophages 14 days after lesion compared with the contralateral hippocampus but not versus unlesioned hippocampi (Figure 5A, upper quadrants at 3 days; and data not shown).

Based on observations of activated caspase-3+ process-bearing cells in deafferented hippocampal regions in PP-lesioned mice 3, 5, and 7 days after lesion (n = 2 to 3 per group), apoptotic microglia were visualized in the molecular layer of the dentate gyrus 3, 5, and 7 days after PP lesion. Activated caspase-3 was present throughout the cytoplasm of CD11b+ microglia (Figure 5D). A void in the staining was apparent where the condensed DAPI-stained nuclear remains were situated. Plasma membrane blebbing and cell rounding was apparent (Figure 5D), both phenomena occurring as the cytoskeletal proteins are broken down by activated caspases. Microglial DNA fragmentation as an indication of apoptosis in the molecular layer was also visualized using TUNEL and co-labeling with CD11b in the deafferented molecular layer 5 days after lesion (Figure 5E). The DNA fragmentation was clearly visualized by the dUTP labeling of the condensed nucleus, which overlapped with the DAPI DNA staining. These microglia also displayed the plasma membrane blebbing and cell rounding characteristic of apoptotic cells (Figure 5E). Taken together, the data therefore suggest that occurring in parallel with the lesion-induced reactive proliferation of microglia is microglial cell death by apoptosis.

Increases in Macrophage Colony-Stimulating Factor (M-CSF) and M-CSFR mRNA Levels Coincide with Elevations in CD11b mRNA

M-CSF and its receptor, M-CSFR, have been shown to be essential for microglial proliferation in other neural lesion models. Granulocyte-macrophage colony stimulat-
ing factor (GM-CSF) signaling via the GM-CSF receptor (GM-CSFR) has also been implicated in microglial proliferation. Changes in levels of M/GM-CSF and M/GM-CSFR mRNA as well as CD11b mRNA were therefore investigated at different intervals from 12 hours to 28 days after lesion (Figure 6). Levels of M-CSF and M-CSFR mRNA levels were significantly elevated at 5 days (Figure 6, A and B), which coincided with the time of maximal CD11b mRNA expression (Figure 6C). GM-CSF mRNA was only detected at very low levels in unmanipulated and deafferented hippocampi, with real-time PCR cycle thresholds >38. No significant changes in mRNA levels of GM-CSF/GM-CSFR were observed (data not shown). Preliminary screenings did not reveal any changes in mRNA levels at 3 or 6 hours either (data not shown).

Lesion-Induced Immigration of BM Cells

Based on previous observations that BM-derived cells contribute to reactive microgliosis in PP-lesioned mice, it was investigated whether proliferative and apoptotic mechanisms of population control applied equally to resident and immigrant microglial cells using GFP BM chimeric mice (Figure 7A). A lesion-induced increase in the proportion of immigrating GFP+ microglia was observed 7 days after PP lesion (Figure 7B). At this time, ~5% of CD45dimCD11b+ cells expressed GFP. Significant lesion-induced immigration of GFP+CD45dimCD11b+ microglia was not observed at 3 or 5 days. As expected, 70 to 75% of CD45highCD11b+ macrophages detected after PP lesion were GFP+ (data not shown). To verify the integration of the BM-derived cells into the neural parenchyma, BM chimeras were analyzed histologically (n = 4), 7 days after lesion. Immigrating GFP+ BM-derived microglia specifically infiltrated the deafferented areas of the dentate gyrus, CA3 and CA1 (Figure 7C, shown for the dentate gyrus), as previously shown in other types of BM chimeric mice.

Impaired Microglial Responses in BM Chimeric Mice

Having observed a significant increase in the number of microglia in the PP-deafferented hippocampus 3 days after lesion, we anticipated a similar increase in BM chimeric mice. Surprisingly, the lesion-induced increase in microglial numbers was absent in BM chimeric mice 3 days after lesion (Figure 8A). Microglial numbers were significantly elevated compared with the contralateral hippocampus 5 and 7 days after lesion. Microglial responses to PP lesion in chimeric and C57BL/6 mice 3 days after PP lesion were therefore compared. Included in this analysis was a group of B6(CD45.1) mice because these congenic mice were used as BM transplant recipients. As expected, a significant increase in the number of microglia was observed in the PP-deafferented hippocampus compared with contralateral hippocampus in C57BL/6 and B6(CD45.1) mice (1.6- and 1.9-fold, respectively; Figure 8A). In doing this comparison, it also became evident that the estimated total number of microglia in the contralateral hippocampi in the BM chimeric mice was significantly smaller compared with C57BL/6 and B6(CD45.1) mice (Figure 8A).

Microglial mitotic activity was then assessed because mice used in flow cytometric analyses had been injected with BrdU three times throughout the past 24 hours before perfusion. In accordance with the observed differences in microglial numbers, the number of proliferating BrdU+ microglia in the PP-deafferented hippocampus in C57BL/6 mice was 11,300 and 14,800 in B6(CD45.1) mice. Both were significantly higher than the 600 BrdU+ microglia observed in the BM chimeras (Figure 8B). In all groups of mice, numbers and proportions of proliferating microglia in the deafferented hippocampi were significantly higher compared with contralateral hippocampi (Figure 8B). Whereas low proportions of microglia were proliferating in the contralateral hippocampus of C57BL/6, B6(CD45.1) and chimeric mice (4, 2, and 2%, respectively), the proportion of BrdU+ microglia was 6% in deafferented hippocampi of chimeric mice compared with the significantly higher 34% in C57BL/6 mice and 33% in B6(CD45.1) mice. In chimeric mice, numbers and proportions of proliferating BrdU+ microglia remained...
significantly elevated at 5 and 7 days after lesion (Figure 8B and data not shown). Observed increases in mitotic microglial numbers were lesion-induced because numbers were significantly smaller in contralateral hippocampi (Figure 8B).

Microglial apoptotic responses were also investigated in the chimeric mice, using annexin V as marker for apoptosis (Figure 8C). As before, dead cells were excluded from the analysis by applying the viability marker 7-AAD. No statistically significant differences were observed in numbers of apoptotic microglia at 3 or 5 days after lesion, although a lesion-induced increase in the proportion of apoptotic microglia was observed after 3 days. By 7 days, both the number and proportion of annexin V\(^+\) microglia were significantly increased (Figure 8C and data not shown). Numbers of apoptotic microglia were significantly lower in chimeric mice than in C57BL/6 or B6(CD45.1) mice (Figure 8C), although the relative proportions of annexin V\(^+\) microglia in deafferented hippocampi were not significantly different between groups (5.2% versus 5.6% and 6.8%, respectively). Microglial mitotic and apoptotic responses were comparable in C57BL/6 and congenic B6(CD45.1) mice 3 days after lesion, although the total number of microglia was slightly higher in B6(CD45.1) mice.

**Mitotic and Apoptotic Responses in Microglial Populations in BM Chimeric Mice**

The contribution of immigrant GFP\(^+\) and resident GFP\(^-\) microglia to the proliferative response after lesion was subsequently analyzed (Figure 9A). Mitotic expansion of the microglial population predominantly took place among the resident GFP\(^-\) microglia (Figure 9B). The estimated number of proliferating immigrant GFP\(^+\) microglia was small but increased significantly during the later phase of the microglial reaction 7 days after lesion (Figure 9C). Overall, the observation of steadily increasing microglial numbers (Figure 8A), and the almost constant numbers of BrdU\(^+\) and GFP\(^+\) BrdU\(^+\) microglia observed in the same mice (Figure 9, B and C), was reflected in a decreasing proportion of all microglia and resident GFP\(^-\) microglia undergoing proliferation during the period of observation. Whereas 6% of resident GFP\(^-\) microglia proliferated 3 days after lesion, this proportion was significantly reduced to 3% 7 days after lesion. By contrast, a significantly larger proportion of GFP\(^+\)BrdU\(^+\) immigrant microglia were induced to proliferate at 7 days after lesion (0.20%) than 3 days after lesion (0.04%).

**Figure 7.** Immigrant BM-derived GFP\(^+\) microglia infiltrate the PP deafferented hippocampus. Microglia were defined as CD45\(^{dim}\)CD11b\(^+\) in flow cytometry and the percentage of immigrant GFP\(^+\) microglia was analyzed in contralateral and lesioned hippocampi. A: Arrows indicate the percentages of GFP\(^+\) microglia in the contralateral and lesioned samples. B: Percentages of GFP\(^+\) CD45\(^{dim}\)CD11b\(^+\) microglia in contralateral and lesioned hippocampi. C: Micrographs showing lesion-specific infiltration of immigrant GFP\(^+\) microglia visualized by the expression of green GFP fluorescence and labeled by CD11b immunohistochemistry in red. DG, dentate gyrus; g, granular cell layer; h, hilus; ml, molecular layer. Scale bars: 200 μm (C, left), 20 μm (C, right). *P < 0.05.
Finally, when analyzing apoptotic resident GFP$^{-}$ and immigrating GFP$^{+}$ microglia independently there was no statistically significant lesion-induced apoptosis 3 or 5 days after lesion in either population by numbers (Figure 9, D–F) or proportions (data not shown). Seven days after PP deafferentation, however, a statistically significant lesion-induced apoptotic response was observed in resident and immigrant microglia. The present work also demonstrates the microglial population and the lesion-induced mitotic activity of resident microglia to be severely reduced in BM chimeric mice.

Discussion

This study aimed to investigate the cellular population control of resident and immigrant microglia by combining the use of BM chimeric mice and a classic model for axonal lesion-induced reactive microgliosis in the dentate gyrus and hippocampus. The present work demonstrates that the lesion-induced increase in microglial numbers from 3 to 7 days after PP transection is balanced by simultaneously occurring mitosis and apoptosis but also that microglial numbers remain elevated for weeks after the injury. In the BM chimeric mice, the resident microglial population displayed a lesion-induced mitotic response, whereas a comparable lesion-induced apoptotic response was observed in resident and immigrant microglia. The present work also demonstrates the microglial population and the lesion-induced mitotic activity of resident microglia to be severely reduced in BM chimeric mice.

BM chimeric mice have been used extensively for studies of microglial turnover and replacement$^{42,43}$ and in studies focusing on the possibility of using BM-derived cells for gene therapy.$^{10,44}$ However, the model has the disadvantage of whole-body irradiation of the host to eradicate host BM cells before transplantation, which could potentially induce microglial activation or cell death. The reduction of the microglial population in the BM chimeras to 30% of that observed in the normal mice after PP lesion was unexpected, as was the observation that the mitotic activity of microglia in BM chimeras was severely impaired. Still, a significant but delayed lesion-induced increase in mitosis of resident microglia accounted for almost all of the mitosis observed.

Measuring apoptosis in cells obtained from tissue using flow cytometry is difficult because the relatively extended period of time cells are processed ex vivo and the processing itself could potentially increase the number of apoptotic cells. This, however, was partly compensated for because the contralateral and deafferented hippocampi from the same animal were processed concurrently. In this way, any difference between the deafferented hippocampus and the contralateral hippocampus would be attributable to reactive microgliosis, although the actual percentage of apoptotic microglia measured could be artificially high. Annexin V binding is a sensitive marker for apoptosis but has also been demonstrated to bind to viable B cells at intermediate levels and can therefore not be regarded as a marker only for apoptosis. Dead or dying cells, however, express higher levels of annexin V binding.$^{45}$ Apoptotic annexin V$^{+}$ cells are therefore only to be gated as cells expressing the highest levels of annexin V for the purpose of excluding any potentially false-positive cells.

Stereology is a quantitative method available for obtaining an unbiased estimate of the number of cells in three-dimensional structures in histological sections, and it is unaffected by deafferentation-induced shrinkage and
deformation of the dentate gyrus. Therefore, it was the method of choice for answering the question of global changes in microglial cell numbers. The limited resolution of the light microscope in the immunohistochemically stained thick tissue sections, however, proved to be a limitation to the level of cellular detail that could be obtained. To make up for the loss of cellular resolution, it was taken into consideration that reactive microgliosis during the stereological and confocal analysis often were observed to occur in multicellular microglial clusters. As earlier mentioned, the actual number of microglia would therefore be at least twofold and possibly threefold higher, as also calculated in studies based on density estimations. This would also be in accordance with the observation of an approximately fourfold up-regulation of CD11b mRNA levels in PP-deafferented isolated hippocampi 5 days after lesion.

Unexpectedly, flow cytometry might be superior to answer questions about changes in total microglial cell numbers because the dissociation procedure makes all microglia accessible to analysis. A disadvantage was that nonaffected areas were also included in the analysis, and that there is inevitably cell loss during the preparation procedure. Still, a 74% increase in microglial cell numbers was detected in deafferented compared with contralateral hippocampus 3 days after lesion in C57BL/6 mice. Flow cytometry also proved its strength in the functional analyses of mitotic and apoptotic responses in microglial subpopulations, because of the high rate at which multiple parameters could be acquired on a single cell basis, which enabled investigation of small subpopulations. Analysis of rare events such as BrdU and annexin V cells in the immigrating GFP population of microglia could introduce uncertainties, but again, comparing the deafferented hippocampus with the contralateral offers a good internal control. Parenchymal microglia express lower levels of CD45 (CD45dim) compared with leukocytes (CD45high). Perivascular cells are known to readily turn over with blood cells and it cannot be excluded that some perivascular cells might fall into the CD45dim gate in flow cytometric analysis.

Because the phenomenon of lesion-induced enhanced immigration of BM-derived microglia became generally accepted an increasing number of studies have been published on the subject. The phenomenon of BM-derived cells homing to areas of brain damage and reactive microgliosis opens the perspective of using a naturally occurring event to target cells engineered for a specific purpose into a pathological area of the CNS, an approach demonstrated to be plausible. For a therapeutic technique like this to be safe and applicable for possible use in humans, the population control of such immigrating BM-derived cells would have to be efficient to avoid induction of malignancies or widespread necrosis in these areas of infiltrating cells. Microglial mitosis is an essential part of microglial turnover and reactive microgliosis opens the perspective of using a naturally occurring event to target cells engineered for a specific purpose into a pathological area of the CNS, an approach demonstrated to be plausible.

For a therapeutic technique like this to be safe and applicable for possible use in humans, the population control of such immigrating BM-derived cells would have to be efficient to avoid induction of malignancies or widespread necrosis in these areas of infiltrating cells. Microglial mitosis is an essential part of microglial turnover and reactive microgliosis opens the perspective of using a naturally occurring event to target cells engineered for a specific purpose into a pathological area of the CNS, an approach demonstrated to be plausible. For a therapeutic technique like this to be safe and applicable for possible use in humans, the population control of such immigrating BM-derived cells would have to be efficient to avoid induction of malignancies or widespread necrosis in these areas of infiltrating cells. Microglial mitosis is an essential part of microglial turnover and reactive microgliosis opens the perspective of using a naturally occurring event to target cells engineered for a specific purpose into a pathological area of the CNS, an approach demonstrated to be plausible.
The present study has demonstrated that apoptosis does play a role in control of the microglial population in brain regions displaying a dense anterograde axonal and terminal degeneration in both resident and immigrating microglia as investigated by identification of phosphatidylserine externalization by annexin V binding. Furthermore, it was shown that microglia indeed can undergo apoptosis in a classic sense with apoptotic morphology, condensed fragmented DNA, and cytoplasmic expression of activated caspase-3. This does point toward immigrating microglia being governed by a similar mechanism of cellular clearance by apoptosis as resident microglia.

At present, the key molecules and signaling pathways involved in microglial mitosis and apoptosis in the differentiated dentate gyrus are unknown, but likely candidates are macrophage colony-stimulating factor (M-CSF) and granulocyte macrophage colony-stimulating factor (GM-CSF). After peripheral axotomy, microglia in the facial motor nucleus increase their expression of M-CSF receptors and increase their binding of M-CSF and GM-CSF before or at the onset of microglial proliferation. Although the presently observed up-regulation of M-CSF and GM-CSF receptor mRNA levels 5 days after lesion occurs relatively late compared with the peak microglial proliferation 3 days after lesion, it should be taken into consideration that changes in mRNA levels were determined based on analysis of whole hippocampi, not allowing direct assessment of up-regulation of these molecules or the GM-CSF and GM-CSF receptor in the areas of microglial mitosis. M-CSF also plays an essential role in the maintenance of the microglial cell population in the normal adult CNS, which is in line with studies of cultured microglia showing that deprivation of M-CSF results in microglial apoptosis. Overall, this study broadens our understanding of microglial response to neural lesion by showing that simultaneously occurring mitosis and apoptosis regulate expansion and reduction of resident as well as immigrant microglial populations after acute neural damage.

Acknowledgments

We thank Christina Fenger, Mads D. Pedersen, and Lene Sylvest for assistance with experiments; Lene Jørgensen and Susanne Petersen for technical assistance; and Werner Vach for statistical counseling.

References

29. Schwendig JD, Schwenk S, Imrich H, Dorries R, Butcher GW, Termeur V: Isolation and direct characterization of resident microglial
cells from the normal and inflamed central nervous system. Proc Natl Acad Sci USA 1991, 88:7438–7442


46. White CA, McCombe PA, Pender MP: Microglia are more susceptible than macrophages to apoptosis in the central nervous system in experimental autoimmune encephalomyelitis through a mechanism not involving Fas (CD95). Int Immunol 1998, 10:935–941
