Technical advance

2-photon imaging of phagocyte-mediated T cell activation in the CNS

Marija Pesic,1 Ingo Bartholomäus,1 Nikolaos I. Kyratsous,1 Vigo Heissmeyer,2 Hartmut Wekerle,1,3 and Naoto Kawakami1,4

1Max Planck Institute of Neurobiology, Department of Neuroimmunology, Martinsried, Germany. 2Helmholtz Zentrum München, Institute of Molecular Immunology, Munich, Germany. 3Hertie Senior Professor and 4Institute of Clinical Neuroimmunology, Ludwig-Maximilians Universität, Munich, Germany.

Autoreactive T cells can infiltrate the CNS to cause disorders such as multiple sclerosis. In order to visualize T cell activation in the CNS, we introduced a truncated fluorescent derivative of nuclear factor of activated T cells (NFAT) as a real-time T cell activation indicator. In experimental autoimmune encephalomyelitis, a rat model of multiple sclerosis, we tracked T cells interacting with structures of the vascular blood-brain barrier (BBB). 2-photon imaging documented the cytoplasmic-nuclear translocation of fluorescent NFAT, indicative of calcium-dependent activation of the T cells in the perivascular space, but not within the vascular lumen. The activation was related to contacts with the local antigen-presenting phagocytes and was noted only in T cells with a high pathogenic potential. T cell activation implied the presentation of an autoantigen, as the weakly pathogenic T cells, which remained silent in the untreated hosts, were activated upon instillation of exogenous autoantigen. Activation did not cogently signal long-lasting arrest, as individual T cells were able to sequentially contact fresh APCs. We propose that the presentation of local autoantigen by BBB-associated APCs provides stimuli that guide autoimmune T cells to the CNS destination, enabling them to attack the target tissue.

Introduction

Brain-specific autoimmune T cells homing to the CNS face a formidable challenge, the blood-brain barrier (BBB), which is a complex composite of a central endothelial tube, concentrically arranged pericytes and phagocytes, and 2 basal laminas (1). This barrier blocks most of the circulating blood components, but its impermeability is not absolute. Most pertinently, the T cells mediating EAE have developed an elaborate set of sequential interactions with different BBB components to access the brain tissue. Most encephalitogenic T cells arrive at the CNS within the leptomeninges, where they attach to the luminal surface of the local small vessels, roll along a short distance, and then crawl before passing through the endothelial wall (2). At this stage, recognition of the autoantigen does not seem to have a major role; however, after leaving the blood vessel, the T cells make serial contacts with perivascular phagocytes and ultimately become activated. Although these phagocytes are distinct from classical DCs, they function as efficient APCs. In particular, as previously shown ex vivo, these phagocytes can present myelin autoantigens acquired from the surrounding CNS tissue (2, 3).

These observations led us to speculate that the presentation of autoantigens by perivascular and interstitial phagocytes provides immigrant T cells with the critical cues that direct them into the CNS parenchyma. However, due to technical limitations, direct evidence connecting interactions of T cells with local APCs and following T cell activation has been lacking to date.

In the present study, we applied a new fluorescent marker of cell activation: a truncated nuclear factor of activated T cells (NFAT) fused to GFP (ΔNFAT-GFP) that contains the polypeptide sequence controlling nuclear translocation. 2-photon imaging resolution was sufficient to determine subcellular localization of ΔNFAT-GFP in vivo, and its translocation kinetics were fast enough to investigate functional T cell interactions with different structures of the target milieu. We used this construct to elucidate the interactions between T cells and APCs within the CNS leptomeninges, the main portal for CNS migrant encephalitogenic T cells. This approach allowed us to demonstrate that perivascular phagocytes, not endothelial cells, activate the incoming T cells. Furthermore, our results emphasized the effect of autoantigen availability on the locomotor behavior and pathological capacity of CNS autoimmune T cells.

Results

ΔNFAT-GFP-expressing T cells. We applied a GFP-labeled, truncated variant of NFAT1 as a “functional” tag to visualize the real-time activation events by which CNS autoimmune T cells cross the leptomeningeal BBB, the essential portal to the CNS parenchyma (3–6). The construct, ΔNFAT-GFP, contained the regulatory domain of NFAT1 that is necessary for phosphorylation, cytoplasmic sequestration, and calcium-induced and calcineurin-mediated dephosphorylation. Dephosphorylation induces a conformational change, which exposes a nuclear localization signal leading to cytoplasmic-nuclear translocation (7, 8). The NFAT construct was truncated to delete the DNA-binding domain of native NFAT (Figure 1A and ref. 9), so as not to interfere with gene regulation by endogenous NFAT. Upon T cell activation, ΔNFAT-GFP was translocated from the cytosol to the nucleus (Figure 1, B and C), similar to native NFAT1 (10). Cytoplasmic-nuclear translocation of ΔNFAT-GFP happened within minutes upon ionomycin stimulation; however, reverse transport from nucleus to cytosol after removing activating stimulus took much longer, around 1 hour (Figure 1, C and D, and Supplemental Video 1; supplemental material available online with this article; doi:10.1172/JCI67233DS1). Most importantly, ΔNFAT-GFP–expressing T cells behaved like their GFP-expressing counterparts; in particular, their encephalitogenic potential remained unimpaired (Figure 1E). We concluded that ΔNFAT-GFP serves as a reliable genetic indicator of T cell activation for 2-photon imaging.
The encephalitogenic potential of autoreactive T cells depends on their antigen specificity. Myelin basic protein–specific (MBP-specific) T cells (referred to herein as TMBP cells) are strongly activated in the CNS and trigger classical acute EAE, thus qualifying as EAEhi T cells. In contrast, myelin oligodendrocyte glycoprotein–specific (MOG-specific) T cells (TMOG cells) display no activation and hardly mediate any clinical defects; these are EAElo T cells (ref. 5 and Figure 1E). This discrepancy suggested that these 2 cell lines represent good candidates for the study of differential T cell activation in vivo.

Activation status of intraluminal T cells. The first contact between encephalitogenic T cells and the BBB occurs on the luminal surface of the endothelial tube. The T cells attach to the intraluminal surface of leptomeningeal vessels, roll along a short distance, and then continue to crawl on the endothelial cells. Both modes of motion differ in the molecules used for the endothelial contacts. While rolling occurs along the blood stream and involves transient interactions between selectins and glycoproteins (11), crawling is often against the direction of blood flow and depends on the binding of integrins VLA-4 and LFA-1 (2). These interactions appear to involve some T cell stimulation (12); however, it is not clear which of the endothelial contacts trigger T cell activation, and what extent of stimulation is provided to T cells within the vascu-
technical advance

Figure 2
Activation status of autoreactive T cells within leptomeningeal blood vessels. (A and B) Intraluminal velocity (A) and average crawling time (B) of TMOG-GFP or TMBP-GFP cells. Results are sum of 5 different experiments per cell line. (C and D) Representative still images of crawling (C; from Supplemental Video 2) and rolling (D; from Supplemental Video 3) TMBP-NFAT-GFP cells. Arrowheads denote ΔNFAT-GFP–spared nuclei. Scale bars: 10 μm. Significance of differences was determined by Mann-Whitney nonparametric t test.

The results of our 2-photon study were unambiguous: regardless of their pathogenic potential (EAEhi versus EAElo), the T cells crawled with similar velocities and for similar durations (Figure 2, A and B), and neither the crawling nor the rolling T cells showed ΔNFAT-GFP in their nuclei (Figure 2, C and D, and Supplemental Videos 2 and 3). We concluded that, if there is any activation of intraluminal T cells, it is not sufficient to translocate NFAT into the nucleus and, implicitly, to initiate gene activation.

T cell/APC contact after extravasation. Having crossed the BBB, autocrine T cells promptly contact local APCs. To characterize these interactions, we initially compared the EAEhi TMBP and EAElo TMOG cells labeled with a traditional, static GFP (referred to herein as TMBP-GFP and TMOG-GFP cells, respectively) to outline their locomotion (2). TMBP-GFP cells moved within the leptomeningeal space with lower velocity and mean square displacement than did TMOG-GFP cells (Figure 3, A and B). We next examined T cell/APC contacts for these 2 T cell lines. These APCs exhibit mostly macrophage and not DC markers, and a substantial proportion expresses MHC class II molecules on their surface, which qualifies them as APCs (2). TMOG-GFP cells largely ignored the local APCs, moving continuously along straight trajectories (white lines in Figure 3C and Supplemental Video 4). In contrast, TMBP-GFP cells halted upon contact with the APCs, drawing trajectories that twisted around an anchoring point (white lines in Figure 3C and Supplemental Video 5). Contacts of TMBP-GFP cells with APCs were significantly longer-lasting than those of TMOG-GFP cells (Figure 3D). Confirming previous reports (2, 5), TMBP-GFP cells in leptomeninges and parenchyma upregulated the activation marker OX-40, but TMOG-GFP cells did not (Figure 3E).

ΔNFAT-GFP translocation in perivascular T cells. To link the contact-dependent changes in T cell locomotion with cell activation directly, we compared EAEhi TMBP and EAElo TMOG cells expressing ΔNFAT-GFP (referred to herein as TMBP-NFAT-GFP and TMOG-NFAT-GFP cells, respectively). In order to enhance visualization of the marker’s subcellular location, we counterstained the cells with SNARF-1 (13), thus labeling both cytosol and nucleus. To avoid dilution of SNARF-1 staining, we labeled CD4+ T cells isolated from spleen 3 days after transfer of ΔNFAT-GFP–expressing T cells, then transferred them into WT recipients (Supplemental Figure 1). Transferred CD4+ T cell–enriched splenocytes penetrated into the CNS meninges more rapidly than did in vitro activated T cells (14). SNARF-1 counterstaining allowed for determination of ΔNFAT-GFP location in up to 90% of all of cells and facilitated the distinction of 3 ΔNFAT-GFP localization patterns (Figure 4A): cytoplasmic, located in the cytosol and not the nucleus (indicative of resting T cells); nuclear, located in the nucleus and not the cytosol (indicative of activated T cells); and localization in both cytosol and nucleus (indicative of transitional T cells).

Importantly, in contrast to their intraluminal counterparts, more than 30% of extravascular TMBP-NFAT-GFP cells exhibited nuclear localization of ΔNFAT-GFP, and another 20% showed the marker in both cytosol and nucleus. Conversely, TMOG-NFAT-GFP cells restricted ΔNFAT-GFP to the cytosol (Figure 4, B and C, and Supplemental Video 6). Nuclear translocation of ΔNFAT-GFP was tightly related to the locomotor behavior of the T cells. Regardless of antigen specificity, T cells with cytoplasmic ΔNFAT-GFP were highly motile, with nearly straight trajectories; in contrast, T cells with nuclear ΔNFAT-GFP showed intercalated segments of coiled tracks (Figure 4D). In addition, T cell velocity directly correlated with ΔNFAT-GFP localization. T cells with cytoplasmic ΔNFAT-GFP moved at an average of 10 μm/min, again regardless of antigen specificity, whereas T cells with nuclear ΔNFAT-GFP traveled at a slow velocity (mean, 4.5 μm/min), and those with both nuclear and cytoplasmic ΔNFAT-GFP moved with an intermediate speed (mean, 6.5 μm/min; Figure 4E). The change in T cell velocity correlated with the change in ΔNFAT-GFP localization (Figure 4F and Supplemental Video 7).

T cell/APC contacts are crucial for T cell activation. Nuclear translocation of ΔNFAT-GFP in the T cells correlated with their contacts to local APCs (Figure 5A and Supplemental Video 8). Furthermore, continuous APC contacts (average duration, 20 minutes) were required to transport ΔNFAT-GFP from the cytosol to the nucleus, whereas brief, transient interactions (average duration, 3 minutes) maintained ΔNFAT-GFP in the cytosol, regardless of T cell antigen specificity (Figure 5B). It is noteworthy that the cytoplasmic-nuclear translocation of ΔNFAT-GFP occurred shortly after contact with an APC (average interval, 4 minutes), whereas the average duration of this contact amounted to 20 minutes (Figure 5C). In contrast to this downstream translocation, the reverse nuclear-cytoplasmic transport of ΔNFAT-GFP required a longer period of time: once detached from the APCs, ΔNFAT-GFP was gradually translocated back to the cytosol, a process requiring approximately 30 minutes (Figure 5, D and E, and Supplemental Video 9). These results correlated to in vitro calculated kinetics of translocation (Figure 1, C and D).
Contacts between T cells and APCs were promiscuous; some APCs could engage several T cells sequentially. The same APC was observed to interact with 2 distinct T cells, in both cases triggering ΔNFAT-GFP translocation (Figure 5A and Supplemental Video 8). In fact, sequential T cell activation by a single APC was not unusual. Conversely, individual T cells visited several APCs over time. Although most enduring APC contacts were established by the T cells showing cytoplasmic-nuclear ΔNFAT-GFP translocation, we also observed extended APC interactions by T cells with pre-existing nuclear ΔNFAT-GFP (average duration, 10 minutes; Figure 5B). These findings suggest that already-activated T cells visit successively local APCs and that these visits either maintain their activation state at a particular level or escalate it continuously.

Finally, we examined sequential versus stable T cell/APC interactions for their capacity to induce T cell activation. Nonactivating and activating contacts were distinguished by ΔNFAT-GFP localization: nonactivating contacts were shorter-lasting (average duration, 3 minutes; Supplemental Figure 2) than the activating ones (average duration, 20 minutes; Figure 5B and Supplemental Figure 2), and during nonactivating contacts, T cells maintained their high velocity (average, 9 μm/min; Supplemental Figure 2). Although we cannot exclude that the cells ultimately receive some stimulus after these sequential APC visits, our results indicate that long-lasting contacts are more important for T cell activation.

Antigen availability and T cell activation in the CNS meninges. The discrepant interactions of the EAE hi and EAE lo T cell lines with CNS APCs may reflect a difference in the availability of presented autoantigen. It is known that the myelin autoantigen is presented within the local tissue, yet the amounts of antigen are suboptimal, below saturating concentrations. In fact, addition of exogenous autoantigen enhances TMBP-GFP and TMOG-GFP cell activation and aggravates clinical EAE (4, 5). To explore the effect of autoantigen availability on TMOG-NFAT-GFP cells, we saturated CNS APCs with the exogenous autoantigen MOG by local microcapillary infusion. The effect of MOG on TMOG-NFAT-GFP cell behavior and ΔNFAT-GFP localization was drastic and immediate. Injection of MOG affected the locomotion of TMOG-NFAT-GFP cells and ΔNFAT-GFP localization in up to 60% of cells analyzed (Figure 6, A–C, and Supplemental Video 10), which was more than in TMBP-NFAT-GFP cells without the exogenous antigen (Figure 4C). Similar to TMBP-NFAT-GFP cells, TMOG-NFAT-GFP cells showed nuclear ΔNFAT-GFP localization displayed reduced velocity (Figure 6D). In contrast, the motility of TMOG-NFAT-GFP cells having cytoplasmic ΔNFAT-GFP did not differ before and after MOG infusion (Figure 6,
ΔNFAT-GFP were contacting the APCs to maintain their activation status, similar to the APC interactions observed with MBP-specific T cells. Because NFAT nuclear translocation leads to gene transcription, we confirmed T cell activation using flow cytometry. As

Figure 4
In vivo T cell activation, shown by nuclear translocation of the ΔNFAT-GFP marker. (A) 3 patterns of ΔNFAT-GFP (green) and SNARF-1 (red) in double-labeled T cells: cytoplasmic (c), nuclear (n), and nuclear/cytoplasmic (nc). Scale bar: 5 μm. (B) Still images of TMOG-NFAT-GFP or TMBP-NFAT-GFP cells in the CNS. Scale bars: 10 μm. (C) Categorization of the ΔNFAT-GFP patterns in TMOG-NFAT-GFP and TMBP-NFAT-GFP cells; relative representation was calculated. na, not analyzable. (D) T cell trajectories from representative movies were superimposed on the coordinates; each line starts at 0. ΔNFAT-GFP location is denoted by line color. (E) Velocity of T cells with different ΔNFAT-GFP locations. (F) Velocity (averaged over 3 time points) of representative TMBP-NFAT-GFP cells changing ΔNFAT-GFP location. Light and dark red shading denotes ΔNFAT-GFP in the nucleus/cytosol and in the nucleus, respectively. Images above show ΔNFAT-GFP localization (green) in a representative SNARF-1–labeled (red) TMBP-NFAT-GFP cell at the indicated times (in minutes; from Supplemental Video 7). Scale bar: 5 μm. Data in C and E are the sum of at least 3 independent experiments and 4 different videos per cell line. ***P < 0.001, 1-way ANOVA followed by Kruskal-Wallis/Dunn multiple-comparison test.

B and D). The contacts that led to cytoplasmic-nuclear ΔNFAT-GFP translocation were long-lasting, as were contacts made by T cells that had already acquired activation (i.e., nuclear localization; Figure 6E). This suggests that the T cells that displayed nuclear ΔNFAT-GFP were contacting the APCs to maintain their activation status, similar to the APC interactions observed with MBP-specific T cells. Because NFAT nuclear translocation leads to gene transcription, we confirmed T cell activation using flow cytometry. As
Figure 5
T cell interaction with local APCs. (A) 2 SNARF-labeled (red) TMBP-NFAT-GFP cells were successively activated (nuclear translocation of ΔNFAT-GFP; green) after contact with the same APC (cyan). Speculated outlines are shown by dotted lines. From Supplemental Video 8. Relative time after start of acquisition is indicated. Closed arrowheads denote T cell/APC interaction; T cells of interest (open arrowheads) are shown in the insets (green and red channels only). Scale bars: 10 μm. (B) Duration of T cell/APC contacts by cells with different ΔNFAT-GFP locations and of contact leading to cytoplasmic-nuclear ΔNFAT-GFP translocation. (C) Time required for cytoplasmic-nuclear ΔNFAT-GFP translocation, starting at the beginning of the T cell/APC contact, compared with the total duration of the same contact. Each color represents the same contact. (D) Time required for nuclear-cytoplasmic ΔNFAT-GFP translocation. Each symbol represents a single cell. (E) SNARF-labeled (red) TMBP-NFAT-GFP (green) cell undergoing nuclear-cytoplasmic ΔNFAT-GFP translocation after detachment from a local APC (cyan). From Supplemental Video 9. Relative time after start of acquisition is indicated. Closed arrowheads denote T cell/APC interaction; T cells of interest (open arrowheads) are shown in the insets (green and red channels only). Scale bars: 10 μm. Results in (B–D) are the sum of 3 independent experiments per cell line. ***P < 0.001, 1-way ANOVA followed by Kruskal-Wallis/Dunn multiple-comparison test.
Our present study describes a genetic marker, ΔNFAT-GFP, that enables mapping — in vivo and in real time — the activation events in autoimmune CD4+ T cells passing through the BBB and the surrounding leptomeningeal space. ΔNFAT-GFP displayed qualities that appear to render it more suitable for 2-photon imaging than previously suggested activation indicators. Among these, TCR complex–related genetic markers, such as a CD3ζ-GFP fusion protein (15–17), or linker for activation of T cells–GFP (LAT-GFP) fusion protein (18), showed T cell activation by clustering at the immunological synapses. However, due to relatively low vertical

shown in Figure 6F, the TMOG cells were activated only after soluble MOG treatment. Taken together, these results indicated that the increased availability of exogenous MOG changed the behavior of TMOG-NFAT-GFP cells from an EAElo to an EAEhi pattern.

Discussion

Immune responses against foreign or self-antigens are the result of complex interactions between immune cells and their surrounding cellular milieu. These interactions centrally involve contacts between neighboring cells, and these contacts critically determine the character of the unfolding immune reaction. Approaches to characterize immune cell communication in different milieus include real-time imaging and isolation of the interacting partner cells for ex vivo analysis, but, owing to the lack of suitable indicators, direct visualization of contacts resulting in cell activation has been futile so far.

Our present study describes a genetic marker, ΔNFAT-GFP, that enables mapping — in vivo and in real time — the activation events in autoimmune CD4+ T cells passing through the BBB and the surrounding leptomeningeal space. ΔNFAT-GFP displayed qualities that appear to render it more suitable for 2-photon imaging than previously suggested activation indicators. Among these, TCR complex–related genetic markers, such as a CD3ζ-GFP fusion protein (15–17), or linker for activation of T cells–GFP (LAT-GFP) fusion protein (18), showed T cell activation by clustering at the immunological synapses. However, due to relatively low vertical
resolution of 2-photon microscopy (typically, 3- to 4-μm spacing), detection of small clusters within 3-dimensional volume is not optimal. In contrast, ΔNFAT-GFP subcellular localization was reliably detectable in more than 90% of the T cells during intravital 2-photon imaging. In transgenic mice, GFP expressed from the immediate early gene locus Nur77 was used as activation indicator (19). This system can distinguish the strength of TCR stimulation according to GFP expression level; however, the GFP protein needs to be transcribed and matured, a process that takes at least few hours, too long for real-time imaging. ΔNFAT-GFP nuclear translocation, on the other hand, occurred within minutes upon stimulation. Finally, calcium indicators had been used to monitor T cell activation in vivo (20); however, small-molecule calcium indicators are rapidly pumped out from T cells and thus fail to label them over extended periods of time. Thus, ΔNFAT-GFP fusion protein qualifies as a reliable indicator to detect rapid T cell activation induced by antigen recognition in vivo.

The members of the NFAT family are expressed in a large number of tissues, most prominently in immune cells (21). In T cells, NFAT acts as a key factor in calcium-controlled gene transcription. After TCR engagement and intracellular calcium increase, calcineurin dephosphorylates NFAT, which then translocates from the cytosol to the nucleus, where—in cooperation with other transcription factors—it binds to DNA to initiate gene expression. However, although ΔNFAT-GFP is a reliable indicator of TCR/calcium-dependent cell activation, it is not absolutely specific. Apart from TCR activation, the pathway may also be triggered via nonimmune receptors; conversely, T cells may be stimulated via alternative pathways, which would not be indicated by ΔNFAT-GFP (22). This must be considered in the case of the intraluminal T cell migration modes for which no ΔNFAT-GFP translocation was observed. The negative finding of NFAT transition may exclude classic TCR-mediated activation, but not necessarily distinct stimulatory pathways with or without TCR participation.

In the present study, we investigated several sets of myelin autoimmune T cells, sharing either genotype or antigen specificity, to correlate their behavior with their discrepant encephalitogenic potentials. Translocation of ΔNFAT-GFP was noted in the brain autoimmune T cells that had crossed the cerebrovascular wall, but was strictly limited to EAEhi T cells. Nuclear ΔNFAT-GFP appeared in the EAEhi TMBP-NFAT-GFP cells during and after contact with the local phagocytes. When isolated ex vivo, these phagocytes present locally produced myelin autoantigens to specific T cells, but the antigenic strength in vitro is weak. The responding T cells are activated in the gut (30), then pass through different milieus in lung and peripheral immune organs, where they are deactivated and reprogrammed (31), before finally passing through the cerebrovascular BBB (2). The use of ΔNFAT-GFP allows for distinguishing activating from other cell-to-cell events. Beyond its use for studying brain autoimmunity, this class of reagents may be applicable to any study of immune milieus, such as during development, infection, and tumor responses.

Methods

Animals. Lewis rats were obtained from and kept and bred in the animal facility at the Max Planck Institutes of Biochemistry/Neurobiology.

Establishment of GFP-labeled antigen-specific T cell lines. The antigen-specific T cell lines were established from the LNs of immunized animals, as described previously (32). Briefly, 6- to 8-week-old male rats were immunized with antigen/CFA. 10 days later, the LN cells were isolated and cultured together with GP+E-86 cells (ATCC no. CRL-9642), which were transfected to produce a GFP gene-coding retrovirus, pMSCVneo-GFP. Alternatively, GP+E-86 cells transfected with pMSCVneo-ΔNFAT-GFP (aa 1–460 of mouse NFAT1) were used. The T cells were positively selected by specific antigen stimulation and negatively using an antibiotic resistance gene included in the retroviral vector. At least 3 rounds of stimulation with the antigen and subsequent expansion in the conditional medium were completed before the cells were used for experiments. For EAE induction, 5 × 10^6 freshly stimulated T cells were transferred into naive animals via the tail vein. Clinical scoring was as follows: 0, no disease; 1, flaccid tail; 2, gait disturbance; 3, complete hind limb paralysis; 4, tetraparesis.

Flow cytometric analysis. T cells were purified at the peak of infiltration from CNS meninges or parenchyma as described previously (2). Briefly, meninges and parenchyma were carefully dissected, and single-cell suspensions were prepared by passing through a 70-μm cell strainer. To purify lymphocytes from the CNS parenchyma, Percoll (GE Healthcare) gradient was used. No further purification was necessary to recover T cells from meninges. The cells were stained with activation marker, CD134 (OX-40; Serotec), followed by rabbit anti-mouse IgG-PerCP (Santa Cruz Biotechnology). Staining was evaluated by FACSAria operated by Cell Quest software (Becton Dickinson). The obtained results were processed by FlowJo 7.6 software (TreeStar).

Confocal microscopy. In vitro cultured resting TMBP-NFAT-GFP cells were labeled with PKH26 Red Fluorescent Cell linker kit for general cell mem-

The Journal of Clinical Investigation  http://www.jci.org  Volume 123  Number 3  March 2013  1199
brane labeling (Sigma-Aldrich). Some cells were treated with 1 μM ionomycin for 3 minutes, after which stimulated and nonstimulated cells were incubated on collagen-coated glass slides for 15 minutes to immobilize them. Cells were fixed in 4% PFA and stained with 5 μg/ml DAPI for 10 minutes. Images were taken with ×40 oil-immersion objective (NA 1.25) using Leica SP2 confocal microscopy.

Fluorescent videomicroscopy. Resting in vitro T<sub>M</sub>NEAT-GFP cells were incubated in a μ-slide I<sup>®</sup> collagen-coated Microscopy Chamber (ibidi) for imaging. T cells were stimulated with 1 μM ionomycin during image acquisition, and imaging continued for 15 minutes after stimulation. Cells were then collected, washed of ionomycin, and plated again in a new Microscopy Chamber, and imaged for another 2–3 hours. Time-lapse recordings were performed using an inverted microscope (Axiovert 200M) equipped with ×40 oil-immersion objective (NA 1.3; Zeiss). Images were acquired using a CoolSnap-HQ camera (Photometrics; Roper Scientific) and processed by MetaMorph (Visitron Systems).

SNARF-1 labeling. ANFAT-GFP–labeled “migratory” T cells were prepared from host spleens 3 days after transfer. The erythrocytes were removed from the suspensions by osmotic lysis, and the macrophages by adhesion on culture dishes. Prior to SNARF-1 labeling, we enriched for CD4<sup>+</sup> T cells using the MagCellect<sup>®</sup> Rat CD4<sup>+</sup> T Cell Isolation Kit (R&D Systems). On average, 20–30% of the T cells were ANFAT-GFP–expressing cells. The T cells were labeled with SNARF-1 (Invitrogen) by incubating for 15 minutes at a concentration of 1.25 μM. SNARF-1 labeling results in a strong signal in the nucleus, whereas cytosol staining is often weaker; therefore, ANFAT-GFP in the cytosol may be detected without SNARF-1 counterstaining.

Intravital imaging. The surgical procedures and technical setup of the 2-photon microscopy were largely as described previously (2, 33). For excitation, the pulsed laser was tuned to 880 or 935 nm, and imaging was performed using a ×20 water-immersion objective (NA 0.95; Olympus) or a ×25 water-immersion objective (NA 0.95; Leica). The emitted photons were collected using nondescanned photomultiplier tubes. Typically, imaging was performed at a 40- to 50-μm stack height (3- to 4-μm z step) in approximately 25-second intervals.

Image analysis. The position of the T cells in the 3-dimensional space was analyzed using Imaris software. Calculation of mean square displacement step (δ<sub>XY</sub>) in the 3-dimensional volume were defined as contacts. All representative videos were processed using ImageJ software. In some videos, the contrast was adjusted by liner rescaling, and the noise was removed using a Median filter and Gaussian blur.

Injection of exogenous antigen. Soluble MOG was injected 3 days after i.v. transfer of T<sub>M</sub>MOG-NEAT-GFP cells. Intrathecal injection into the cisterna magna was performed as described previously (2). Alternatively, during intravital imaging, MOG was instilled locally via a microcapillary (tip diameter, 13 μm; Biomedical Instruments) that was positioned using a microcapillary manipulator (Luis & Neumann). The capillary was pushed through the arachnoida through a hole cut by a high-energy 2-photon laser. For the injection, the microcapillary was filled with a solution of 10 μg/ml MOG protein and 0.2 μg/ml tetramethylrhodamine dextran. We injected a total volume of 3–4 μl of the solution.

Statistics. Statistical evaluation was performed using Prism software (GraphPad). In all experiments, Mann-Whitney nonparametric t test (2-tailed) was used for 1 pair, and 1-way ANOVA followed by Kruskal-Wallis test/Dunn’s multiple-comparison test was used for more than 1 pair. A P value less than 0.05 was considered significant.

Study approval. All experiments were conducted according to the Bavarian state regulations for animal experimentation and were approved by the appropriate authorities.

Acknowledgments

We thank Sabine Kosin and Marko Lazic for technical support. This work was supported by the Max Planck Society, Ludwig-Maximilians University München, Deutsche Forschungsgemeinschaft (SFB571, projects C6 and B10), Hertie Foundation, Novartis Foundation for Therapeutic Research, and German Academic Exchange service (DAAD).

Received for publication October 5, 2012, and accepted in revised form December 6, 2012.

Address correspondence to: Naoto Kawakami or Hartmut Wekerle, Max-Lebsche-Platz 31, 81377 Munich, Germany. Phone: 49.89.7095.8386; Fax: 49.89.7095.8380; E-mail: Naoto.Kawakami@med.uni-muenchen.de (N. Kawakami), hwekerle@neuro.mpg.de (H. Wekerle).


