Induced bronchus-associated lymphoid tissue serves as a general priming site for T cells and is maintained by dendritic cells

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Mucosal vaccination via the respiratory tract can elicit protective immunity in animal infection models, but the underlying mechanisms are still poorly understood. We show that a single intranasal application of the replication-deficient modified vaccinia virus Ankara, which is widely used as a recombinant vaccination vector, results in prominent induction of bronchus-associated lymphoid tissue (BALT). Although initial peribronchiolar infiltrations, characterized by the presence of dendritic cells (DCs) and few lymphocytes, can be found 4 d after virus application, organized lymphoid structures with segregated B and T cell zones are first observed at day 8. After intratracheal application, in vitro-differentiated, antigen-loaded DCs rapidly migrate into preformed BALT and efficiently activate antigen-specific T cells, as revealed by two-photon microscopy. Furthermore, the lung-specific depletion of DCs in mice that express the diphtheria toxin receptor under the control of the CD11c promoter interferes with BALT maintenance. Collectively, these data identify BALT as tertiary lymphoid structures supporting the efficient priming of T cell responses directed against unrelated airborne antigens while crucially requiring DCs for its sustained presence.

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via the respiratory route in a nonhuman primate model (Corbett et al., 2008). In mice, MVA delivered via the intranasal (i.n.) route has been shown to induce long-lasting and protective antibody and T cell immune responses (Gherardi and Esteban, 2005; Kastenmuller et al., 2009). However, little is known about the immunological events after respiratory MVA infection.

The present report demonstrates that a single i.n. application of the replication-deficient MVA is sufficient to induce the long-lasting presence of BALT and that the lung-specific depletion of DCs interferes with BALT maintenance. Ex vivo imaging of antigen-specific T cell–DC interactions within BALT by two-photon microscopy indicates that, independent of the specific antigenic challenge inducing its formation, BALT can function as a general priming site for T cell responses directed against antigens that reach the lower respiratory tract.

RESULTS AND DISCUSSION
A single i.n. application of replication-deficient MVA suffices to induce highly organized BALT
We have recently shown that i.n. infection with the mouse γ-herpesvirus MHV-68 induces BALT in mice and that BALT actually serves as a reservoir of latent MHV-68 (Kocks et al., 2009). To address the question of whether latent or continuous reinfection of host cells with virus is required for the induction and/or maintenance of BALT, we tested the replication-deficient poxvirus MVA for its ability to induce BALT. Although the peribronchiolar space in lungs of untreated mice contained only few CD11c+ cells and hardly any lymphocytes (Fig. 1 A and not depicted), a massive perivascular and peribronchiolar infiltration of CD11c+ cells could be observed 4 d after i.n. application of 10⁷ infectious units (IU) MVA (Fig. 1 B). At day 8 after infection, B and T cells had been recruited to CD11c+ cell-rich areas (Fig. 1, C and D). Although in some areas B and T cells were still scattered diffusely around vessels and bronchioles (Fig. 1 C) at this time point, we also frequently observed areas in which B cells started to segregate toward the center of BALT structures (Fig. 1 D). By day 12, large areas of BALT had been formed. Frequently, CD11c+ cells were found surrounding the structures (Fig. 1 E), whereas B and T cells exhibited various degrees of clustering (Fig. 1 F). At day 28 after infection, the MVA-induced BALT displayed a pronounced segregation into T and B cell–rich areas (Fig. 1 G). In this differentiated BALT, DCs were found to reside primarily in the T cell areas (Fig. 1 H). Quantitative analysis revealed that BALT structures were most abundantly present between days 8 and 12 after infection but could still be detected at relatively high frequencies even 28 d after virus application (Fig. 1 I). Collectively, these data demonstrate that a single i.n. application of the replication-deficient MVA is sufficient to induce the long-lasting presence of highly organized BALT, and that the accumulation of CD11c+ cells around vessels and bronchioles represents an early event in its formation.

Figure 1. Induction of BALT after mucosal infection with MVA. (A–H) C57/BL6 mice were infected i.n. with 10⁷ IU MVA, and at the time points indicated (days after infection), lung sections were analyzed by fluorescence microscopy applying mAb and DAPI as indicated. Representative sections from three to six mice analyzed for each time point derived from two to three different experiments are shown. Bars, 100 μm.

Virus-encoded gene expression and recruitment of inflammatory cells after i.n. application of MVA
To monitor productive MVA infection and viral gene expression in the lung, we used an MVA strain encoding an enhanced GFP (EGFP; Lehmann et al., 2009). Within 5–7 h after i.n. infection, high numbers of GFPbright cells could be detected in the bronchoalveolar lavage (BAL) of MVA–GFP–infected but not MVA–WT–infected animals (Fig. 2, A and B).
Although up to 50% of all live cells in the BAL expressed GFP at 7 h after infection, only a very small proportion of cells were still GFP+ at 24 h after infection, and no GFP signals were detectable at later time points (Fig. 2 B). Because MVA cannot produce infectious virions in mammals (Ramírez et al., 2000), these data indicate that MVA infectivity is rapidly inactivated in vivo. Further analysis revealed that 65% of all BAL cells isolated 5–7 h after infection represented alveolar macrophages (AMs; CD11c+MHCII+; Fig. 2 C). More than 92% of all GFP+ cells showed this phenotype, with only 4% of all GFP+ cells being conventional DCs (CD11c+MHCII-; Fig. 2 D). Importantly, the analysis of single-cell suspensions prepared by enzymatic digestion of whole-lung tissue yielded largely identical results (unpublished data).

The rapid decrease of the GFP signal observed shortly after infection with MVA-GFP prevented its further use in directly tracking the fate of MVA-encoded antigenic material. To overcome this limitation, Cy5-labeled MVA (MVA-Cy5) was subsequently used for i.n. infection. As early as 20 min after MVA-Cy5 inoculation, numerous large CD11c+ cells could be detected carrying MVA-Cy5 and having the typical location of AMs within the lumen of alveoli and airways (Fig. 2 E). In accordance with our data derived from i.n. MVA-GFP infection, we could also detect few Cy5+CD11b+CD11c+ DCs (Fig. 2 F) in the lungs of MVA-Cy5–infected animals. However, the CD11c+ DC network surrounding the conducting airway system did not harbor detectable MVA-Cy5 fluorescence (unpublished data). Analyzing BAL 3 d after infection, a significant increase in the number of DCs (CD11c+MHCII+; Fig. 2 G) as well as inflammatory CD11c-CD11b+ cells (Fig. 2 G), such as recruited monocytes/macrophages and granulocytes, could be observed. At this time point, >20% of the total BAL cells still contained MVA-Cy5 (Fig. 2 H). These MVA-Cy5+ cells were now exclusively CD11c+ and/or CD11b+ cells (Fig. 2 I, right), with most of them showing a phenotype of DCs.

Figure 2. Initial host cell tropism of MVA in the lung. Mice were i.n. infected with 10^7 IU MVA, and BAL cells were analyzed by flow cytometry. (A) GFP expression of DAPI+ BAL cells 7 h after infection with MVA-GFP or MVA-WT. (B) GFP expression of BAL 0–72 h after i.n. infection with MVA-GFP (mean ± SD; n = 2–3 mice/time point). (C and D) Expression of CD11c and MHCII on all (C) or GFP+ (D) DAPI+ BAL cells 7 h after infection with MVA-GFP. (E) Immunohistology of lung sections 6 h after i.n. application of Cy5-labeled MVA using antibodies and dyes as indicated. (F) As in E, we obtained sections from lungs 30 min after MVA-Cy5 application. Bars, 25 μm. (G–I) FACS analysis of BAL cells 3 d after MVA-Cy5 application using antibodies and virus as indicated. (G and H) Gate on all DAPI+ cells. (I) Gate on all DAPI+ MVA-Cy5+ cells. The FACS plots and immunohistology shown are representative of three independent experiments, each with two to three mice analyzed per time point. *, P < 0.05; **, P < 0.01.
Collectively, these data indicate that predominantly AMs and some DCs are efficiently infected with MVA within hours after i.n. application. Most of the infected cells rapidly disappear from the lung within a few hours, presumably undergoing apoptosis and/or necrosis, similar to processes found after in vitro infection with MVA (Liu et al., 2008). However, (CD11c+ MHCII+) or inflammatory cells (CD11b+CD11c−; Fig. 2 I, left).

Figure 3. i.n. vaccination with recombinant MVA induces a distinct population of antigen-specific cytotoxic T cells in the lung. (A) Mice i.n. received 10⁷ IU MVA-Cy5, and 3 d later the phenotype of Cy5+ cells in the draining LN was determined by flow cytometry. (B) Expression of CD86 on CD11c+ MHCII+ brLN DCs before and 2 d after i.n. instillation of MVA (shaded area, isotype control). (C-G) 1 d after the i.v. transfer of 200 CD45.1+ CD8+ OT-I T cells into CD45.2+ recipients, mice were i.n. or i.m. infected with 10⁷ IU MVA-OVA. At the indicated time points (days after infection) the absolute number of OT-I cells was determined in the brLNs (C), BAL (D), lung (E), and spleen (F). (G) The percentage of OT-I cells expressing CD69 in the lung and spleen. (H) Expression of CD3 and CD69 in BALT at 12 d after infection. Bar, 50 μm. Data in A and B are representative of four mice analyzed in two independent experiments. Data shown in C–G are pooled from two independent experiments with two to three mice per time point (means + SD). *, P < 0.05; ***, P < 0.001.
Recombinant MVA induces a distinct population of antigen-specific cytotoxic T cells in the lung after i.n. application

AAfter i.n. instillation of MVA-Cy5, we could readily detect Cy5+ cells in the lung-draining bronchial LNs (brLNs). These cells were primarily DCs expressing high levels of CD11c and MHCII (Fig. 3 A). The majority of these DCs were found to express either CD103 or CD11b (Fig. 3 A). These observations, together with the finding that brLN DCs of i.n. MVA-infected mice express higher levels of CD86 than those of noninfected mice (Fig. 3 B), indicate that the i.n. vaccination with MVA can elicit a productive immune response in the lung as well as in the lung-draining brLNs. We therefore hypothesized that the quality of an adaptive immune response directed against the model antigen OVA in the lung might differ between mucosal (i.n.) and systemic (i.m.) application of MVA encoding chicken OVA (MVA-OVA). To closely mirror the physiological situation of a small starting population of naive T cells being specific for a given antigen, we adoptively transferred by i.v. injection 200 OT-I T cells (CD45.1+) that recognize the SIINFEKL peptide derived from OVA in an MHC class I-restricted manner into CD45.2 recipients 1 d before infection with MVA-OVA via the i.n. or i.m. route.

After i.n. MVA-OVA application, OT-I T cells could be detected at day 4 after infection in the brLNs and spleen (Fig. 3, C and F). These cells were also found in the lung and BAL (Fig. 3, D and E) starting at day 6. In contrast, after i.m. application, only very few OT-I T cells were present in brLNs or BAL at any time point investigated (Fig. 3, C and D), whereas these cells appeared with similar kinetics and numbers in the spleen but less frequently in the lung when compared with i.n.-treated mice (Fig. 3, E and F). Of interest, after i.n. but not i.m. application, persistent expression of the early activation marker CD69 were observed on a large proportion of OT-I as well as endogenous CD8+ T cells in the lung and BAL (Fig. 3 G and not depicted). In contrast, hardly any expression of CD69 could be observed on OT-I T cells residing in the spleen independent of the route of MVA application (Fig. 3 G). Immunohistology of lung sections revealed that CD69+ T cells were present in BALT (Fig. 3 H).

These observations indicate that the application of MVA via the respiratory route can induce a local immune response that is characterized by the presence of activated antigen-specific as well as nonspecific CD8+ T cells in the airways (BAL) and BALT that continuously express high levels of CD69. The transient up-regulation of CD69 has recently been suggested to inhibit lymphocyte egress from LNs by retaining sphingosine 1-phosphate receptor 1, a receptor known to be required for lymphocyte exit from LNs into effenter lymphatics, within intracellular compartments (Shiow et al., 2006).

Essential role for DCs in BALT maintenance

The infiltration of CD11c+ DCs observed within the lungs of MVA-infected mice probably represents an early step during BALT induction (Fig. 1 B). To address the role of DCs in the maintenance of MVA-induced BALT in further detail, we i.n. applied diphtheria toxin (DT) to CD11c-DOG mice (Hochweller et al., 2008), which express the DT receptor (DTR) under the control of the CD11c promoter. The i.n. application of DT at days 8 and 10 after MVA infection resulted in an efficient depletion of CD11c+ cells from the lung tissue (Fig. 4 A and not depicted). A quantitative
Figure 5. Visualization of antigen-specific T cell–DC interactions during T cell priming within BALT. TAMRA-labeled CD8+ OT-I T cells and CMAC-labeled polyclonal CD8+ WT T cells were injected i.v. into BALT-bearing recipients. 24 h later, SIINFEKL-loaded EGFP+ bone marrow–derived DCs (Ag-DCs) were i.t. transferred into the same animals, and lungs were explanted an additional 24 or 48 h later. (A) Cryosections of paraformaldehyde (PFA)-fixed lungs isolated 24 h after i.t. transfer of Ag-DCs into MVA-treated WT mice. (left) EGFP+ Ag-DCs and TAMRA+ OT-I T cells localize into BALT. (right) Higher magnification image of a BALT structure harboring Ag-DCs, OT-I T cells, and polyclonal CD8+ WT T cells. (B) Visualization of BALT within lungs of MVA-treated WT (left) or CCR7−/− (right) mice by ex vivo two-photon microscopy. Maximum intensity projections of three-dimensional imaging volumes (left: eight Z-slices, 7.5-μm spacing; right: seven Z-slices, 6-μm spacing). Collagen fibers surrounding the basal surface of the bronchial epithelium (dashed line) are visualized by SHG. Asterisks represent the bronchial lumen, and arrowheads indicate a blood vessel. (C) Analysis of T cell–DC interactions within BALT of MVA-treated WT mice by two-photon microscopy (excitation wavelength = 780 nm). 24 h after i.t. transfer of EGFP+ Ag-DCs, TAMRA+ OT-I T cells exhibit a highly confined migration behavior in the vicinity of Ag-DCs. In contrast, CMAC+ polyclonal WT control T cells display a much higher motility (Video 1). (D) The experiment was performed as in C but imaged at 865 nm (Video 2). (E) Motility parameter analysis for OT-I and polyclonal CD8+ WT T cells migrating within BALT 24 h after transfer of Ag-DCs. (left) Dots represent individual cell tracks, and bars indicate median average track speed. (middle) Mean displacement plots. (right) Motility coefficient (means + SD). **, P < 0.01. (F) 24 h after i.t. transfer of Ag-DCs into CCR7−/− recipients, EGFP+ DCs were found to enter BALT by directional interstitial migration (arrowhead; Video 3). In contrast, DCs remain largely stationary during Ag presentation within BALT (dashed yellow box; Video 3). Data in A–F are representative of two to four mice per group in two independent experiments.
immunohistological analysis showed that the area of CD11c⁺ pixels per lung section was reduced by >90% 2 d after the second DT treatment (Fig. 4 B), whereas flow cytometry revealed that DCs in mesenteric LNs were not affected by the local application of DT (Fig. 4 C). Of interest, these studies also revealed that BALT was strongly reduced in size in the DT-treated group (Fig. 4 D), whereas the total number of BALT structures per lung was not reduced by DT treatment (not depicted). Collectively, these data indicate that DCs play an essential role in the maintenance of MVA-induced BALT. Because AMs are efficiently depleted by this approach as well (unpublished data), we currently cannot rule out that these cells also contribute to BALT maintenance. However, this hypothesis seems unlikely, as AMs are readily depleted within 1 d after MVA infection (Fig. 2), whereas the DT treatment was not started before day 8 after MVA application.

**BALT serves as a general priming site for T cells**

The data described so far indicate that the virus-induced formation of BALT participates in generating antivirus immune responses. However, although MVA cannot replicate in mammals and is rapidly cleared from the lung tissue, MVA-induced BALT is maintained for several weeks (Fig. 1 I). We therefore tested the hypothesis that induced BALT might function comparably to a regular lymphoid organ, being able to support the de novo priming of naive T cells bearing TCR specificities that are not related to the BALT-inducing pathogen. To this end, BALT was induced in WT animals by i.n. application of MVA-WT. 18 d later, these mice, as well as untreated CCR7⁺/− animals, which constitutively develop BALT, received by i.v. injection 2–2.5 × 10⁷ TAMRA-labeled CD8⁺ OT-I T cells together with 1.5–2 × 10⁶ CMAC-labeled polyclonal CD8⁺ T cells isolated from C57BL/6 donors. 24 h later, the animals additionally received by intratracheal (i.t.) application LPS-matured EGFP⁺ Ag-DCs as well. Ag-DCs localized to BALT remained largely stationary but exhibited extraordinary probing activity, displaying far-range sweeping dendrite movements during antigen presentation (Fig. 5, C and D; and Videos 1 and 2). At 48 h after i.t. transfer of Ag-DCs, many OT-I T cells were clearly enlarged, exhibiting slow swarming movements in the vicinity of EGFP⁺ Ag-DCs (unpublished data), probably indicating the start of their proliferative activity (Mempel et al., 2004). In contrast, CMAC-labeled polyclonal CD8⁺ WT T cells did not engage in stable interactions with Ag-DCs at either time point (Fig. 5 C, Video 1, and not depicted). The quantitative analysis of T cell motility parameters revealed much lower cellular velocities and displacements, and consequently highly reduced motility coefficient values, for OT-I cells when compared with the polyclonal CD8⁺ T cell population (Fig. 5 E and not depicted), further corroborating the antigen-specific nature of the observed OT-I T cell–DC interaction behavior in BALT.

Although we sometimes observed EGFP⁺ Ag-DCs to enter BALT by means of a rapid directional interstitial migration during ex vivo imaging (Fig. 5 F and Video 3), it seems likely that DCs can reach BALT via afferent lymphatics as well. Ag-DCs localized to BALT remained largely stationary but exhibited extraordinary probing activity, displaying far-range sweeping dendrite movements during antigen presentation (Fig. 5, D and F; and Videos 2 and 3). We next quantitatively analyzed lung cryosections for the maximum nucleus diameters of TAMRA⁺ OT-I T cells as well as TAMRA⁺ CD3⁺ endogenous T cells present in BALT 48 h after i.t. transfer of Ag-DCs (Fig. 5 G). OT-I T cells displayed significantly larger nuclei compared with resting endogenous T cells, consistent with the idea that the proliferation of antigen-specific T cells activated within BALT had already started at this time point. Finally, we analyzed the proliferation of OT-I T cells by in vivo CFSE dilution at day 4 after the i.t.
transfer of Ag-DCs (Fig. 5 H). The recipient animals were additionally treated with FTY720 to block lymphocyte egress from lymphoid organs. The observation that OT-I T cells isolated from the lungs had proliferated as efficiently as those residing in the bronchus further supports the idea that BALT can indeed function as a local priming site for naive T cells.

Within this study, we demonstrate that a single i.n. application of the replication-deficient vaccine virus MVA, which efficiently infects AMs and DCs, is sufficient to induce long-lasting and highly organized BALT in normal WT mice. DCs in particular seem to represent key organizing cells in BALT biology, as they are among the first cells to be recruited into sites of developing BALT (Fig. 1 B) and depletion of these cells leads to a pronounced regression in BALT size (Fig. 4). Furthermore, antigen-loaded DCs are actively migrating into preexisting BALT where they can present antigen to naive T cells.

Thus, we conclude that induced BALT represents an organized lymphoid structure for antigen presentation by DCs in the lung that is capable of supporting long-lasting T cell–DC interactions as well as the efficient priming of naive T cells. Importantly, the antigenic specificity of the observed priming events (the OVA I peptide SIINFEKL) was completely different from the antigenic nature of the stimulus that had initially induced the formation of BALT (MVA), and occurred long after the complete clearance of the inducing pathogen. Therefore, given its sustained presence, induced BALT can obviously function as a general priming site for T cell responses in the lung.

MATERIALS AND METHODS

Mice. Mice were bred at a local animal facility or purchased from Charles River. CD11c-DOG mice expressing DTR under the control of the CD11c promoter were described previously (Hochweller et al., 2008). B6.129P2(Crj)-Ccr7tm1Rfor/J mice (CCR7−/−) were backcrossed on the C57BL/6 background for >15 generations (Förster et al., 1999). In some experiments, DCs were differentiated from the bone marrow of MVA.

Organ preparation. BAL fluid was obtained by flushing the lung two times with 1 ml of cold PBS. For lung histology, mice were perfused with 5 ml of ice-cold PBS. The trachea was cut and a 1:1 mixture of Tissue-Tek OCT (Sakura) and PBS was instilled. The lung was removed and snap frozen in OCT. 8-μm sections were performed with a cryostat (CM3050; Leica), as previously described (Hintzen et al., 2006).

Flow cytometry. Single-cell suspensions were prepared by mechanical disruption. Lungs were cut into small pieces and digested with 0.5 mg/ml collagenase D and 0.025 mg/ml DNase I in RPMI 1640 for 30 min at 37°C. Cells were blocked with 5% rat serum, stained, and analyzed with a cytometer (LSRII; BD). Data were analyzed using WinList 6.0 (Venty Software House). The following antibodies were used: CD103-FITC (M290), CD11c-PE (HL3), MHCIId-bio (AF6-120.1), detected with streptavidin–Alexa Fluor 750, CD11b-PeCy7 (M1/70), CD11b-PE (M1/70), CD86-allophycocyanin (APC; GL-1), anti-CD11c-APC (isotype control), CD69PerCp-Cy5.5 (H1.2F3), CD69 (H1.2F3), CD127–Alexa Fluor 488 (A7R34), IFN-γ–PE (XM1G1), CD3-PE (145–2C11), B220–Cy5 (TIB 146), F4/80–PE (BM8), and CD11c–bio (HL3) detected with streptavidin–Alexa Fluor 488.

Epifluorescence microscopy. Sections were fixed in cold acetone, rehydrated in Tris-buffered saline with 0.05% Tween 20, blocked with 2.5% rat and 2.5% mouse serum, and stained at room temperature for 30 min. Pictures were taken with a fluorescence microscope (Axiovert) using AxioVision 4.6 software (both from Carl Zeiss, Inc.).

BALT area analysis. Overview pictures of lungs were recorded with the mosaic function of the AxioVision software and whole central sections of the lungs were analyzed. The total number of BALT structures was measured as described previously (Kocks et al., 2007). ImageJ (available at http://rsweb.nih.gov/ij/) was used to determine the percentage of CD11c+ pixel area on pictures taken with identical exposure and display settings.

i.t. DC transfer. EGFP+ bone marrow–derived DCs were generated from the bone marrow of β-actin–EGFP transgenic mice (Okabe et al., 1997) by in vitro culture (Ohl et al., 2004). After overnight LPS maturation and loading with OVA I peptide (SIINFEKL), 2–6 × 105 DCs were resuspended in 60 μl PBS and transferred i.t.

Two-photon microscopy. Mice were anesthetized with ketamine/xylazine and bled to death by cutting the venacava before harvesting the lungs. Horizontal lung slices (thickness = 1–2 mm) were cut with a cryotome blade (Leica), immobilized in an imaging chamber (Fig. S1) using tissue adhesive (Abbott), and superfused with oxygenated (95% O2/5% CO2) RPMI 1640 medium (Invitrogen) containing 1% penicillin/streptomycin, 25 mM Hepes, and 5 g/liter glucose. Imaging was performed on a TriM Scope (LaVision Biotec) equipped with an upright microscope (BX51; Olympus) fitted with a 20× 0.95 NA water immersion objective and a pulsed infrared laser (MaiTai Ti:Sa; Spectra-Physics), tuned to 865 nm for optimal excitation of EGFP and TAMRA and to 780 nm for additional visualization of CMAC. To generate time-lapse series, Z-stacks of up to 10 images were acquired every 20 s.

Data analysis of two-photon imaging was performed as described (Worbs et al., 2007), using Imaris 6.2.1 (Bitplane). Only tracks with durations >60 s were included in the analysis. Average track speed values were calculated using Imaris, whereas mean displacement plots and motility coefficients were calculated using Visual Basic for Applications (Excel; Microsoft).

Statistical analysis. Statistical analysis was performed with Prism 4 (GraphPad Software, Inc.). All significant values were determined using the unpaired two-tailed t test and all error bars represent SDs. Statistical differences for the mean values are as follows: *, P < 0.05; **, P < 0.01; and ***, P < 0.001.
Cutting edge: mucosal application of a lyophilized viral vector vaccine vectors.


**Figure S1.** Custom-built incubation chamber. Prewarmed, oxygenated medium first enters a prechamber by gravity flow (tubing on the left side) to minimize pulsation artifacts. A constant steady medium flow within the central part of the imaging chamber, containing the immobilized lung slices, is directed from the bottom left to the top right. The temperature within this section is monitored by a thermosensor (blue cable) positioned directly adjacent to the imaged tissue and maintained at 37°C. After flowing into the third subchamber, the superfusion medium is pumped out by means of a conventional rotary pump (tubing on the right side).

**Video 1.** T cell migration and interaction behavior within MVA-induced BALT 24 h after i.t. transfer of Ag-DCs. 24 h after i.t. transfer of SIINFEKL-loaded EGFP+ DCs (green) into MVA-treated WT recipients, many TAMRA+ OT-I T cells (red) exhibit a confined migration behavior within BALT, forming long-lasting contacts with Ag-DCs (middle). In contrast, CMAC+ polyclonal WT control T cells (cyan) display a higher motility (right). [left] Three-color overlay. Maximum intensity projection of a 132 × 132 × 54 μm three-dimensional imaging volume (six Z-slices) with a 300x time lapse.

**Video 2.** DC probing and T cell–DC interaction dynamics within MVA-induced BALT 24 h after i.t. transfer of Ag-DCs in the presence of unlabeled competitor OT-I cells. The experimental setup was as described for Video 1, but using an excitation wavelength of 865 nm. This allows the imaging of SHG signals (blue) as well as providing much better visualization of fine dendrite processes of EGFP+ Ag-DCs (green) intensively probing their environment (left). In addition to TAMRA-labeled OT-I T cells (red), mice received an equal number of unlabeled OT-I T cells. This approach largely facilitates the long-term tracking of individual TAMRA-labeled cells (right). Maximum intensity projection of a 200 × 200 × 36 μm three-dimensional imaging volume (four Z-slices) with a 300x time lapse.
Video 3. Directional DC migration toward BALT and active DC dendrite movement during T cell priming 24 h after i.t. transfer of Ag-DCs. 24 h after i.t. transfer of SIINFEKL-loaded EGFP+ DCs (green) into CCR7−/− recipients, Ag-DCs were observed to enter BALT by directional interstitial migration (left panel, top left corner). In contrast, during antigen presentation to TAMRA+ OT-I T cells (red) within BALT, Ag-DCs remained largely stationary (left panel, right half; and right panel). SHG signal (blue). (left) Maximum intensity projection of a 282 × 282 × 60 μm three-dimensional imaging volume (eight Z-slices) with a 300x time lapse. (right) Detail with twofold magnification.