A novel method to allow noninvasive, longitudinal imaging of the murine immune system in vivo


Introduction

In vivo imaging has revolutionized understanding of the spatiotemporal complexity that suberves the generation of successful effector and regulatory immune responses. Until now, invasive surgery has been required for microscopic access to lymph nodes (LNs), making repeated imaging of the same animal impractical and potentially affecting lymphocyte behavior. To allow longitudinal in vivo imaging, we conceived the novel approach of transplanting LNs into the mouse ear pinna. Transplanted LNs maintain the structural and cellular organization of conventional secondary lymphoid organs. They participate in lymphocyte recirculation and exhibit the capacity to receive and respond to local antigenic challenge. The same LN could be repeatedly imaged through time without the requirement for surgical exposure, and the dynamic behavior of the cells within the transplanted LN could be characterized. Crucially, the use of blood vessels as fiducial markers also allowed precise re-registration of the same regions for longitudinal imaging. Thus, we provide the first demonstration of a method for repeated, noninvasive, in vivo imaging of lymphocyte behavior. (Blood. 2012; 119(11):2545-2551)

Methods

Animals

BALB/c (H-2b), C57BL/6J (CD45.1, H-2b), and Ly5.1/C57BL/6J (CD45.1, H-2b) mice were used as recipients and donors. DO11.10 TCR transgenic (Tg) mice on a BALB/c background were used as a source of Ag-specific T cells for experiments that show the tLNs response to immunization. These TCR Tg T cells recognize ovalbumin323-339 in the context of I-Ad. All animals were specified pathogen free and maintained under standard animal house conditions at Strathclyde University or Glasgow University in accordance with Home Office Regulations.

Adoptive transfer of Ag-specific lymphocytes and in vivo challenge

Adoptive transfers were performed as described previously. In some cases, cells were labeled with the fluorescent dye CFSE (Invitrogen) or Cell...
Tracker Red (CMTPX; Invitrogen) immediately before use.15 Twenty-four hours after the adoptive transfer mice were challenged in the ear pinna with 50 μL of heat-aggregated ovalbumin or PBS as a control. Heat-aggregated ovalbumin was prepared as previously described17 and was administered subcutaneously at a concentration of 100 μg/mouse. Cell Dark red fluorescent (660/680) 0.04-μm carboxylate-modified microspheres (FluoSpheres; Invitrogen) were used in LN transplantation experiments to show lymphatic drainage. PBS-diluted microbeads were injected subcutaneously into the outer edge of the ear pinna with tLNs.

Lymph node transplantation

Lymph nodes were removed from newly weaned 3- to 4-week-old mice, removing any excess fat, and placed into ice cold, sterile PBS (LNs were kept in this manner for no longer than 1 hour before transplantation). Whole LNs were inserted into a skin flap created in the ear pinna of the 8- to 12-week-old recipient. Veterinary-grade glue (Vetbond; 3M), enabling warming of tissue heat mat. The ear of interest was mounted on a purpose-built stand with the LN edge. Videos were acquired for 15-30 minutes at an X-Y pixel increment of 1.5-512 with 1.5-μm increments in Z. Three-dimensional tracking was continuously monitored and maintained by a thermostatically controlled heat mat. The ear of interest was mounted on a purpose-built stand with the use of veterinary-grade glue (Vetbond; 3M), enabling warming of tissue throughout the experiment and maintained at 37°C. At the end of imaging sessions, the veterinary glue was dissolved with 70% ethanol, allowing repeated imaging at a later date. Intravenous injection of nontargeted quantum dots (Qtracker 565 nontargeted quantum dots; Invitrogen) allowed for highlighting of blood vessels. tLNs were routinely 100-150 μm from the LN edge. Videos were acquired for 15-30 minutes at an X-Y pixel resolution 512 × 512 with 1.5-μm increments in Z. Three-dimensional tracking was performed with Velocity 5.5 (Perkin Elmer) after correction for tissue drift, and values representing the mean velocity, displacement, and meandering index were calculated for each object.

Histopathology and immunofluorescence microscopy

Tissue samples in OCT embedding medium (Miles Diagnostic Division) were frozen in liquid nitrogen and stored at −70°C. Tissue sections (8 μm) were prepared on a cryostat (ThermoShandon) at −22°C. Sections were mounted onto SuperFrost slides (BHD) before being air-dried and stored at −20°C until further processing. H&E (Sigma-Aldrich) staining was performed on a Varistain 244 auto-staining processor (ThermoShandon). Tissue samples were then mounted in DPX mounting fluid (VWR International) before visualization by bright field microscopy. For fluorescence microscopy, tissue sections were stained with anti-CCL21–FITC, anti-VCAM-1–biotin, anti-CD4–PerCP, anti-CD45.1–bio, anti-CD45.2–FITC, anti-CD8–APC, anti-CD3–biotin, anti-IgM–Rhodamine Red, anti-BP3–biotin, or anti-gp38–biotin. Streptavidin Alexa Fluor 647 (Invitrogen) or streptavidin-FITC (BD Biosciences) was used in conjunction with biotinylated primary Abs. Fluorescent images of the sections were captured with a Zeiss LSM 510 Meta confocal microscope equipped with a 20×/1.0NA water-immersion objective lens (Zeiss).

Flow cytometry

Flow cytometry was performed as described previously.18 Abs or appropriately labeled isotype controls (anti-CD4–PerCP, anti-CD45.1–bio, anti-CD45.2–FITC, anti-CD8–APC, anti-B220–FITC, FITC-Mouse IgG2a isotype control, and FITC/PerCP/APC-rat IgG2a, κ isotype control; all BD Biosciences) and biotinylated KJ1.26 Ab were added to each sample at a dilution of 1:100 Ab. Biotinylated Abs were detected by incubation with fluorochrome-conjugated streptavidin (BD Biosciences). Samples were analyzed with a FACScanto flow cytometer equipped with a 488-nm Argon laser and a 635-nm red diode laser (BD Biosciences) and analyzed with FlowJo Version 9.4 software (TreeStar).

In vivo multiphoton imaging

Multiphoton imaging was performed with a Zeiss LSM7 MP system equipped with both a 10×0.3 NA air and 20×/1.0NA water-immersion objective lens (Zeiss) and a tunable titanium/ sapphire solid-state 2-photon excitation source (Chameleon Ultra II; Coherent Laser Group). Excised LNs were bound with veterinary-grade glue (Venbond; 3M) onto a plastic coverslip and adhered with grease to the bottom of the imaging chamber and continuously bathed in warmed (36.5°C), oxygenated CO2-independent medium. For in vivo imaging, animals were anesthetized with fentanyl/fluanisone (Hypnorm; Janssen Animal Health) and (Hypnovel; Roche) injectable anesthesia (10 mL/kg of a mix of fentanyl/fluanisone/midazolam)/H11001/0.04%/H9262 m carboxylate-modified microspheres (FluoSpheres; Invitrogen) were used in LN transplantation experiments to show lymphatic drainage. PBS-diluted microbeads were injected subcutaneously into the outer edge of the ear pinna with tLNs.

Statistics

Results are expressed as mean ± SD. Gaussian distribution was confirmed by D’Agostino and Pearson omnibus normality test before significance being determined by 2-way ANOVA. A value of P < .05 was regarded as significant.

Results

tLNs in the ear pinna are anatomically normal

Cervical, axillary, or inguinal C57BL/6J LNs from 3- to 4-week-old donors were engrafted to the ear pinna of 8-week-old syngeneic recipients. Donor LNs were inserted into small subdermal pockets...
on the ventral surface of the ear pinna. Mild, localized inflammation was observed at the transplantation site for ≤ 7 days after the procedure. By 3 weeks after transplantation, the superficial appearance of the ear pinna returned to normal, and the tLN/site of engraftment was not obvious to the naked eye (Figure 1A). The tLN was visible within the lightly dissected pinnae and maintained its original size and shape (Figure 1B). By 3 weeks after transplantation, > 80% of the tLNs had engrafted and were readily identifiable in live mice with the use of soft tissue x-ray (Figure 1C). These tLNs were situated posterior to the auricular elastic cartilage and appeared histologically normal (Figure 1D).

Because the stromal cell network is essential for maintaining LN size and cellularity,\textsuperscript{9} we examined the structural integrity of this compartment after the engraftment procedure. No marked differences were noted comparing control (non-tLN) with tLN for the distribution of BP-3 (Figure 2A), a glycosylated cell surface protein expressed by reticular stromal cells of secondary lymphoid tissue.\textsuperscript{19} We also detected both donor and recipient CD45\textsuperscript{+}CD4\textsuperscript{+}CD3\textsuperscript{+} inducer cells in the tLN (supplemental Figure 1, available on the Blood Web site; see the Supplemental Materials link at the top of the online article), indicating that mechanisms known to be important in maintaining cellular organization and function in normal LNs are present in our tLN model.\textsuperscript{20,21}

Accordingly, conventional anatomic compartmentalization of T and B cells was evident in tLNs (Figure 2B). In addition, the overall proportion of CD4\textsuperscript{+}, CD8\textsuperscript{+} T cells and B220\textsuperscript{+} B cells was similar to control LNs (Figure 2C; supplemental Figure 2). CD11c\textsuperscript{+} staining showed normal distribution of dendritic cells throughout the paracortex (supplemental Figure 3). The conventional structural organization and cellular composition of tLNs indicates their capacity to support normal lymphocyte function, and a number of parameters were next examined to confirm this.

**tLNs support lymphocyte recirculation**

Lymphocyte recirculation is a crucial aspect of efficient immune surveillance.\textsuperscript{22} Naive T and B cells enter LNs through high endothelial venules (HEVs). VCAM-1 is a key molecule that promotes adhesion of lymphocytes to the vascular wall of the HEVs\textsuperscript{23} and was evident in tLNs (Figure 3A). HEVs are also a source of CCL21,\textsuperscript{24,25} which actively recruits T cells to the paracortical area of the LN. CCL21 was present in control and tLN tissue sections, appearing throughout the T-cell area (Figure 3B). The presence of both HEVs and appropriate chemotactic cues implies that tLNs would indeed support lymphocyte recruitment.

To demonstrate this function, LNs were transplanted from C57BL/6J/CD45.1 mice into the ear pinnae of C57BL/6J/CD45.2
recipients. Three weeks after engraftment, tLNs had been repopulated with host CD45.2+ T cells with 82.5% of lymphocytes with tLNs being CD45.2+/H11001 (Figure 3C). The peripheral LNs from the recipient mouse were also pooled and analyzed for the presence of CD45.1+/H11001 lymphocytes originating from the tLN. As might be expected very few CD45.1+ cells could be detected, but a very small population was present (0.065%; Figure 3C). Thus, 3 weeks after transplantation, the tLN lymphocyte population predominantly consisted of CD45.2+ lymphocytes. Normal recirculation by the tLN was also confirmed with the use of the adoptive transfer of fluorescently labeled lymphocytes (supplemental Figure 4).

**Lymphatic drainage to the tLN**

Afferent lymphatics are a critical component of immune surveillance, allowing LNs to selectively localize and trap Ag and APCs trafficked there from peripheral tissues. The presence of lymphatic vasculature supplying the tLN was assessed by examining the expression of gp-38, a cell surface glycoprotein preferentially expressed by stromal cells in T cell–dependent areas of secondary lymphoid organs and lymphatic endothelial cells.26 Lymphatic vessels that express gp-38+/H11001 were evenly distributed throughout the tissue and were associated with areas of CD4+ staining in both control and tLN (supplemental Figure 5A-D), consistent with published reports of this glycoprotein being predominantly expressed in T-cell areas of the LN.26 To directly examine the capacity of tLNs to drain material from its new anatomical location, fluorescent microbeads (0.04 μm) were injected into the tip of the ear pinna. Microbeads were easily detected in control, popliteal LNs after footpad injection (supplemental Figure 5E-F) and 24 hours later were localized within the CD4+ T-cell area of the LN. Similarly, microbeads could be detected in the tLN (supplemental Figure 5G-H) after their injection into the tip of the ear pinna, confirming the ability of the transplant to form functional afferent lymphatics in its new tissue location.

**tLNs support naive T-cell activation**

We used adoptive transfer of CFSE-labeled TCR Tg T cells (DO11.10) to determine whether the tLN could facilitate activation of Ag-specific T-cell responses.15 CFSE dilution and expansion of transferred cells was determined by flow cytometry 72 hours after immunization with Ag. The proportion of CD4+/KJ1.26+ cells in both the control and tLN was 4.82% and 4.78% respectively (Figure 4A-D) and was reflected in the loss of CFSE signal, indicative of cell division (Figure 4E-G). No difference in number of divisions between challenged control or tLNs was observed, indicating equivalent proliferative responses (summarized in Figure 4G). Overall, these data suggest that tLNs have a similar ability to control LNs in supporting Ag-driven lymphocyte activation and proliferation.

**Figure 4. Activation of T cells in the tLN.** Incremental loss of CFSE intensity by KJ1.26+/CD4+ cells 72 hours after heat-aggregated ovalbumin (HAO) challenge in the ear showed proliferation of adoptively transferred DO11.10 T cells in BALB/c mice with established tLNs. Representative flow cytometric plots of PBS-challenged cervical LN (A) and tLN (B). An expanded KJ1.26+/CD4+ population is seen in both cervical (C) and tLN (D) after antigenic challenge. Incremental loss of CFSE intensity of KJ1.26+/CD4+ cells in the cervical LN and tLN (E-F), reflecting Ag-specific proliferation that is summarized graphically (G). Data represent the mean of triplicate samples ± SD (n = 2).
was still significantly lower than in excised LNs ($P < .001$). No significant alteration in meandering index was detected (Figure 5E); however, surgical intervention produced similar displacement rates to those seen in excised LNs (Figure 5F).

Because we did not have to perform surgery to permit imaging the tLN, we next sought to determine the potential of our model to reimage the same tissue at different time points. We chose to use blood vessels as fiduciary markers to guide reimagining of exactly the same region of the tLN. By injecting nontargeted quantum dots intravenously we were able to map blood vessels within the tLN during the initial imaging session. Injection of quantum dots also allowed us to visualize blood flow in vivo in the tLN, further indicating the successful formation of normal vasculature (Figure 6; supplemental Video 4). Importantly, mapping of blood vessels enabled us to re-register the imaging field to the same anatomical location in the tLN (Figure 7) as well as subsequently imaging lymphocyte behavior (supplemental Video 4 for day 1 and supplemental Video 5 for day 3). These findings emphasize the utility of this technique both for noninvasive repeat imaging in longitudinal studies and for imaging at a defined locus.

**Discussion**

Here, we describe a novel technique in which a LN is transplanted into the ear pinna of a mouse, providing a convenient anatomic...
regions are displayed from the same tLN. Images were acquired with a 10x objective lens. Two representative regions are displayed from the same tLN.

Established tLNs showed the key anatomic, cellular, and functional characteristics of control LNs. tLNs exhibited a normal stromal cell network, evidence of blood and lymphatic supply, and the presence of the chemokine CCL21, associated with coordinating localization of CD4+ T cells, CD45+CD4+CD3- LN inducer cells of both host and donor origin were also detected within the tLN. These cells are known to play a critical role in LN development through their interactions with stromal organizer cells, which induces the release of chemokines and growth factors.27 Furthermore, their continued presence in adult LNs is considered to be important in maintaining the organization of the LN. Presumably as a result of all of this, the key cell types required for the initiation of an adaptive immune response, including T cells, B cells, and CD11c+ dendritic cells, were all detected in the usual proportions and locations in the tLN.

We also demonstrated the presence of vascular and lymphatic supplies to the tLN and that these were functional, capable of supporting lymphocyte recirculation and Ag drainage from tissue sites. Lymphocyte recirculation was evidenced by the presence of recipient T cells within tLN 3 weeks after transfer. Whether a tLN develops afferent lymphatic vessels capable of draining Ag from its new anatomic location had never been previously explored and was shown by injecting fluorescent microbeads into the peripheral tissue upstream of the tLN. Previous studies had indicated that neonatal spleens transplanted into the ear pinna were capable of responding to Ag challenge.8 However, unlike LNs, spleens do not possess afferent and efferent lymphatic vessels, and the transplanted spleen was immunized directly with a targeted injection of Ag, thus, circumventing the requirement for tissue-draining lymphatic vessels.8 Growth factors capable of directly inducing the growth of lymphatic vessels have been characterized and are up-regulated in response to environmental stress, including hypoxia, and can regenerate lymphatic vessels after tissue damage.15 Therefore, the tLN system may also be useful in studying angiogenesis.

Flow cytometric analysis showed that immunization with ovalbumin in the ear pinna led to comparable Ag-specific T-cell proliferation in both the tLN and the control LN. This conclusively showed that the tLN is fully functional, undergoing vascularization to receive adoptively transferred T cells, draining Ag from a local injection site and supporting the cellular interactions required for clonal expansion of Ag-specific T cells. To confirm this final point, MPLSM was used to show T-lymphocyte movement within tLNs. Analysis of MPLSM data showed basically similar movement characteristics in tLNs and control LNs. Interestingly, lymphocytes in tLNs had reduced velocity and displacement rate compared with excised LNs, whereas sham surgery near the tLN increased these parameters. It is perhaps unsurprising that local trauma can affect lymphocyte function in draining LNs. Sterile trauma is known to promote inflammatory mediator release,28 with dAMPs being important receptors in this process. In addition, localized tissue trauma increases lymphatic drainage, a process shown to increase intranodal levels of CCL21, potentially affected dynamic cellular migration patterns.29 Our model avoids such effects that may complicate the interpretation of data according to approaches with explantation or exposure of LNs.

Importantly, because surgery was not required to image the tLNs, we also showed, for the first time, longitudinal imaging of the same LNs several days apart. For these studies, the ability to image the same tissue area was particularly important. To achieve this, we used blood vessels as fiduciary markers to re-register the imaging field to the same anatomic location in the tLNs during subsequent imaging sessions, clearly showing the potential of this technique for noninvasive, longitudinal imaging of cellular behavior.

In addition to visualizing spatially or temporally separated events, such as the physical movement of cells entering and leaving LNs, this approach opens up the potential for numerous other studies, including lymphangiogenesis and vascularization; longitudinal imaging of infection, vaccinations, and tumors; and combining this technique with genetically modified animals to rapidly create compound transgenics.

In conclusion, we describe a new approach to imaging LNs in vivo with the use of MPLSM without the need for surgical exposure. This approach has allowed us to perform the first noninvasive, longitudinal imaging of lymphocyte behavior in vivo. We believe this model will provide a powerful tool to investigate a wide range of lymphocyte/lymphocyte and lymphocyte/stromal cell interactions that regulate normal and pathologic immune responses in vivo and the effect of therapeutic intervention on these processes.
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Authorship

Contribution: V.B.G., R.A.B., and K.J.B. performed the experiments; V.B.G. and R.A.B constructed the figures, analyzed the data, and wrote the manuscript; R.A.B performed all of the imaging studies; I.B.M., C.M.R., E.J.J., and G.A. provided technical assistance and helped with experimental design; G.G. and P.M. provided technical and analytical assistance; A.J.W. provided expertise with the confocal imaging; and J.M.B. and P.G. designed the research and contributed to writing the paper.

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References

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