T cell progenitor therapy–facilitated thymopoiesis depends upon thymic input and continued thymic microenvironment interaction

Michelle J. Smith,1,2 Dawn K. Reichenbach,1,2 Sarah L. Parker,3 Megan J. Riddle,1 Jason Mitchell,2,3 Kevin C. Osum,2,3 Mahmood Mohtashami,4 Heather E. Stefanski,1 Brian T. Fife,2,3 Avinash Bhandoola,5 Kristin A. Hogquist,2 Georg A. Holländer,6,7 Juan Carlos Zúñiga-Pflücker,4 Jakub Tolar,1 and Bruce R. Blazar1,2

1Division of Blood and Marrow Transplantation, Department of Pediatrics, Masonic Cancer Center, University of Minnesota, Minneapolis, Minnesota, USA. 2Center for Immunology, Department of Medicine, and 3Department of Medicine, University of Minnesota Medical School, Minneapolis, Minnesota, USA. 4Sunnybrook Research Institute and Department of Immunology, University of Toronto, Toronto, Ontario, Canada. 5T-Cell Biology and Development Unit, Laboratory of Genome Integrity, Center for Cancer Research, National Cancer Institute, Bethesda, Maryland, USA. 6Department of Biomedicine, University of Basel, Basel, Switzerland. 7Department of Paediatrics and Weatherall Institute of Molecular Medicine, University of Oxford, Oxford, United Kingdom.

Introduction

Allogeneic hematopoietic stem cell transplantation (HSCT) is the current standard of care for many malignant and nonmalignant hematopoietic diseases (1–4). Prior to infusion of the donor graft, the patient must go through a conditioning regimen involving chemotherapy or chemoradiotherapy. This conditioning inflicts severe damage on the thymic microenvironment and results in delayed peripheral CD4+ and CD8+ T cell reconstitution (5–9), leaving the patient vulnerable to opportunistic infections (10–12). The elements of the thymic microenvironment most sensitive to damage from chemoradiotherapy are the thymic epithelial cells (TECs), and their loss is responsible for prolonged impairment of thymopoiesis following transplant, which depends on successful recruitment of progenitors from the BM graft (13, 14).

TECs are indispensable to T cell development in vivo (15). They support homing of progenitors to the thymus (16, 17), provide critical survival signals (18, 19), direct trafficking of progenitors between the cortex and the medulla (18), and regulate positive and negative selection (20) and egress of single-positive T cells from the thymus (18). Several strategies have been investigated preclinically to preserve TECs during pretransplant conditioning regimens, such as prophylactic administration of keratinocyte growth factor (KGF) (21) and KGF in combination with leuprolide acetate (22) or transient inhibition of p53 prior to irradiation (23). However, these approaches have not yet led to a rapid and full restoration of peripheral T cell recovery early after transplant.
Importantly, TECs also depend on interactions with developing T cells in order to develop, proliferate, and persist (15, 24–26). Conditioning eliminates developing thymocytes, and, in order to generate new T cells, there is a reliance on HSC-derived progenitors to home to and differentiate to T cell progenitors (proTs) within the thymus. Therefore, we utilized in vitro culture systems that use OP9-DL1 cells to generate large numbers of proTs for thymic reconstitution and to speed thymic regeneration. The OP9-DL1 system uses a mouse BM stromal cell line (OP9) expressing Delta-like 1 (Dll1), a Notch1 receptor ligand, to direct and support the differentiation of T cells from hematopoietic (or embryonic) stem/progenitor cells (27, 28), of both mouse and human origin, that range in maturation from CD4+ CD8+ double-negative 1 (DN1) to DN4 cells (27–29). Other approaches, such as TSt-4 cells expressing Notch ligands Dll1 or Dll4 (30) or feeder-free systems based on immobilized Dll4 (31), can be used to achieve similar differentiations. In an allogeneic mouse model of HSCT, proTs differentiated and expanded using these methods when transferred together with HSCT, increased thymopoiesis following transplant (30–32), and conferred protection against tumor challenge (33), without the requirement for MHC matching between the proT donor and recipient. Given that a potential therapy using OP9-DL1–derived proTs is being developed for clinical use, we were interested in the specific effects and the stage of differentiation of proTs prior to adoptive transfer on TEC regeneration in a posttransplant setting. To isolate and characterize these effects, we employed a congenic mouse transplant model in which the in vitro–derived proTs or proTs enriched in DN2 or DN3 cells are adoptively transferred with T cell–depleted (or –deficient) BM into lethally irradiated recipient mice. In this model, we showed that the compartment of the thymus influenced by adoptive transfer of proTs was dependent upon the maturation stage of the input population and that a key factor in the continued benefits derived from this therapy was ongoing recruitment of new progenitors from the BM.

Results

ProTs home to the thymus and immigrate to and persist in the periphery. The initial and rate-limiting step in T-lymphopoietic and TEC recovery after conditioning and transplant requires homing of donor progenitor cells to the thymus (34). Thus, we sought to assess the homing and persistence of proTs in our model as well as compare their migration on both healthy (nonirradiated) and irradiated thymic stromal cells. ProTs were generated from donors expressing a luciferase transgene and transplanted with WT congenically marked BM cells into lethally irradiated recipients. The adoptively transferred cells were predominantly at the DN3 stage of development at the time of transplant (Figure 1A), and photoluminescent signal was monitored and quantified weekly (Figure 1, B and C). We observed a peak in both total body and thymically located signal at week 3 after transplant (Figure 1D).

The degree and nature of thymic recovery depends on the subset of DN T cells that are transplanted. Given that previous work on proT therapy indicated that the DN2 subset of proTs was most effective at increasing thymic size after allotransplant (32), we wanted to examine the differential thymus homing effects of the two dominant DN populations that are produced in OP9-DL1 cell cocultures with our congenic model, DN2 and DN3, as each is attainable in required numbers for adoptive transfer in vivo. Under normal physiological conditions, DN2- and DN3-developing thymocytes are located within the cortical and subcapsular regions of the thymus, while interacting with cortical TECs (cTECs) and undergoing further differentiation into TCR+CD4+CD8+ double-positive cells that then undergo positive selection. As the thymocytes progress through their maturation program, they upregulate CC-chemokine receptor 7 and move to the medulla, wherein medullary TECs (mTECs) direct negative selection. If progenitors survive negative selection and achieve a mature single-positive state, then they upregulate the sphingosine-1-phosphate receptor 1, which directs their egress from the thymus into the periphery (35).

We next tested whether in vitro–derived proTs would interact with the thymic microenvironment in the same way and specifically determined if different developmental stage subsets would have divergent effects on the thymic microenvironment. In order to answer this question, we sorted DN2 and DN3 subpopulations from OP9-DL1 cocultures and transplanted these with congenic T cell–depleted BM cells into lethally irradiated B6 recipients (Figure 2A). Mice that received total body irradiation experienced an 80% to 90% loss of mTECs and cTECs as compared with nontransplanted controls (Figure 2, B and C, and Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.92056DS1). On day 21 after transplant, when proT numbers peak within the thymus, mice given DN3 proTs showed a significant increase in the numbers of mTECs (Figure 2B). In contrast, cTEC numbers were unchanged after injection of either DN2 or DN3 (Figure 2C).
Given the differential effect of these two subsets of DN cells on the thymic microenvironment and the known dependence of TECs on crosstalk with developing thymocytes (36–39), we sought to examine the location of the adoptively transferred DN2 and DN3 subsets within the thymus. On day 7 after proT engraftment, there were similar numbers of DN2 and DN3 proT–derived cells within the thymus, with the majority of them being located in the cortex as expected (Figure 2, D and E). On day 21, there were significantly more thymically located proTs in mice that received the DN2 cells compared with the BM-only group; the mice that received DN3 cells did not display a similar increase (Figure 3A), as measured by flow cytometry and immunofluorescence (Figure 2, D and E). The proTs in the group that received the DN3 subset were located primarily in the medulla, in contrast to proTs in the group that received the DN2 subset that remained primarily within the cortex (Figure 2, D and E).

Total thymic cellularity was increased on day 21 after transplant in the DN2 but not DN3 proT group (Figure 3A). Neither group of proT recipients had increased peripheral T cells in either the spleen or the lymph node (LN) at this time point as compared with BM-only controls. The T cells that were present in the spleen at day 21 after transplant were overwhelmingly derived from the host, which is consistent with homeostatic expansion due to radiation-induced lymphopenia (Figure 3B). Given the dependence of TECs on interactions with developing thymocytes, the difference in localization between the DN2- and DN3-derived proTs, which correlates with the increase in mTECs (CD45–EpCAM+MHCII+UEA1+) in the group that received DN3 proTs, supports a hypothesis that adoptively transferred proTs improve long-term HSC-derived peripheral T cell reconstitution by improving thymopoiesis through interactions with mTECs. Similar to day 21, at day 60 after BM transplant (BMT), this effect was seen in the spleen (Figure 3D) but not in the peripheral LN (data not shown), indicating a defect in homing to the LN,
**Figure 2. Double-negative 3 T cell progenitors stimulate mTEC recovery.** Lethally irradiated CD45.1 recipients were given CD45.2 WT BM alone, with double-negative 2 (DN2) Thy1.1 progenitors (proTs) or with DN3 Thy1.1 proTs. Thymi were harvested at day 21, and their thymic epithelial cells (TECs) were analyzed by flow cytometry. Data are representative of 3 separate experiments, with 3–6 animals per group per time point. Data are shown as mean ± SEM. Data sets were compared using 2-way ANOVA. *P ≤ 0.05, ***P ≤ 0.001, ****P ≤ 0.0001. (A) Representative bulk proTs, DN2-sorted.
proTs, and DN3-sorted proTs. Live cells were selected, and flow cytometry plots show the DN populations of the bulk proTs (left) and DN2 proTs (middle), defined as (CD19, B220, Gr-1, CD11b, CD11c, Ter19, NK1.1) –CD45+CD4–CD8–CD25+CD44hi. DN3 proTs (right) were gated on live cells and then defined as (CD19, B220, Gr-1, CD11b, CD11c, Ter19, NK1.1) CD45+CD4 CD8 CD25+CD44hi. (B) Absolute numbers of mTECs at day 21 after transplant. (C) Absolute numbers of cTECs at day 21 after transplant. (D) DN2 proTs or (E) DN3 proTs (Thy1.1+, blue) in the cortex (BP-1, green) and in the medulla (UEA-1, red, outlined in white) at days 7 and 21 after transplant.

which could be cell intrinsic or due to radiation-induced damage to the LN microenvironment. Consistent with early thymopoiesis, DN2 cells were found at a statistically higher percentage in the cortex following irradiation ($P = 0.0447$, Figure 3E). Although not significant, there was a subsequent increasing trend of DN3 cells migrating into the medulla (Figure 3E). Interestingly, in vitro–derived proTs at the DN3 stage express significantly higher levels of CCR9 than those at the DN2 stage or thymocytes developing in vivo at either the DN2 or DN3 stage (Figure 3F and Supplemental Figure 2C). ProT-derived peripheral T cells expressed similar levels of CD3 as BM-derived and residual host T cells (Supplemental Figure 2, A and B). Taken together, these results indicate that DN3 is the optimum subset to increase peripheral T cells long term after transplant, possibly because of the ability of DN3 proTs to more rapidly boost mTEC numbers, while DN2 proTs provided a greater short-term increase in thymic size, as evidenced by thymocyte number on day 21 (Figure 3, A and C).

**TEC recovery is limited in the absence of developing T cells.** We have shown that improvement in short-term thymopoiesis achieved by adoptively transferred proTs is correlated with an increase in the number of thymically located proTs as compared with BM-only controls (see Figure 3A), whereas increases in elements of the thymic microenvironment, namely mTECs, are correlated with the specific locality of those adoptively transferred proTs within the thymus (Figure 2, B, D, and E). However, in those studies, each group had contributions to thymic recovery from progenitors recruited from the BM graft. While the total number of cells in the thymus peaked at day 21 (Figure 1B), the number of DN progenitors within the thymus peaked at day 14 following adoptive transfer (Figure 4A). The increase in DN cells in the thymus is not only due to the proT-derived cells, but also to increased recruitment of lymphoid progenitors from the engrafted BM (40–42). In fact, progenitors recruited from the BM are known to actively compete with more mature progenitors in the thymus. When comparing mice given either WT or Rag-/-BM with or without proTs, mice that received WT BM with proTs had increased numbers of DN cells overall as compared with those that received WT BM alone, as did mice that received WT BM alone when compared with mice that received Rag-/-BM alone (Figure 4A). When broken down by subset (Figure 4, B–D), this held especially true in DN3 and DN4 subsets (Figure 4, C and D), although a small minority of Rag-/-cells can differentiate past the DN3 stage under some conditions (43). There is less disparity in the DN2 subset of cells present in the thymi of mice transplanted with either WT or Rag-/- BM with or without proTs, the DN2 subset being less proliferative and at a stage of differentiation before the Rag gene is required for successful TCR rearrangements (Figure 4, B and C, and Supplemental Figure 3, A–D). Despite having shown that proT localization in the thymus was correlated with increased TEC recovery (Figure 2) and that progenitors from the BM of a WT graft also increased the number of DN cells within the thymus as compared with mice that received Rag-/-BM with proTs (Figure 4), we wanted to isolate the effects of the proTs from the effects of progenitors recruited from the BM graft on the thymic microenvironment. So, we transplanted Rag-/-BM with or without proTs and assessed TEC recovery and peripheral T cell reconstitution. The group that received the proTs showed a significant increase in the total TEC number on day 21 after transplant as compared with the group that received Rag-/-BM alone (Figure 5A). Additionally, the number of cycling TECs was also significantly increased by day 21 in the group that received the proTs, as measured by BrdU incorporation (Figure 5B) and expression of Ki67 (data not shown). Consistent with our previous experiments, the total increase in TECs was due to an increase in the number of mTECs (Figure 5C), while the proT therapy did not stimulate significant recovery of cTECs (Figure 5D). These results were confirmed by the proportion of immunofluorescent signal due to UEA-1 (Figure 5H), indicating the medulla, in confocal microscopy images of healthy (nonirradiated) control thymi (Figure 5E), thymi of recipients of Rag-/-BM (Figure 5F), and thymi of recipients of Rag-/-BM plus proTs (Figure 5G). However, without the continued recruitment of progenitors from the BM, TECs cease to proliferate (Figure 5B) and their numbers begin to decline after day 21 (Figure 5A).

In the model pairing proTs with Rag-/-BM, we saw a peak in the number of thymocytes at day 21 (Figure 6A) but no significant increase in mature peripheral T cells at this point (Figure 6, B and C). Looking
Figure 3. Double-negative 2 T cell progenitors boost overall thymic size, and double-negative 3 T cell progenitors lead to long-term T cell increases. Lethally irradiated CD45.1 B6 recipients were given CD45.2 WT BM alone, with double-negative 2 (DN2) Thy1.1 progenitors (proTs) or with DN3 Thy1.1 proTs. Thymi and peripheral lymphoid organs were harvested at day 21 and day 60 and were analyzed by flow cytometry. Data are representative of 3 separate experiments, with 4–6 mice per group per time point. Data are shown as mean ± SEM. Data sets were compared using 2-way ANOVA. *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001, ****P ≤ 0.0001. (A) Absolute numbers of thymocytes at day 21 after transplant. (B) Absolute numbers of CD3+ spleen cells at day 21.
After transplant. (C) Absolute numbers of thymocytes at day 60 after transplant. (D) Absolute numbers of CD3+ spleen cells at day 60 after transplant. (E) Percentage of DN3 and DN2 cells present in the cortex, cortical medullary junction (CMJ), and medulla following irradiation and transfer of DN3 and DN2 cells onto thymic slices. (F) Median fluorescent intensity of CCR9 expressed on DN2 and DN3 in vivo thymocytes and in vitro proTs.

Past day 21, proT-derived cells begin to exit the thymus (Figure 6D) and migrate to the periphery by day 35, preferentially residing in the spleen (Figure 6, E and F). The CD4 and CD8 profiles at days 21, 35, and 60 are shown in Supplemental Figure 4. The egress to the periphery correlated with the loss of the beneficial effects of proTs on TECs (Figure 5, A and B). By day 60, we observed no continued benefit to the TEC (Figure 5A), thymocyte (Figure 6G), or peripheral T cell (Figure 6, H and I) compartments. Taken together, these data indicate that TECs damaged by radiation can be stimulated to proliferate by the presence of developing T cells within the thymus but that this effect is temporary in the absence of a continued influx of progenitors from the BM.

Discussion

The presence of thymic progenitors within the thymus is essential to the health of the microenvironment under normal physiological conditions and even more so when recovery from damage is required. In fact, thymic size is regulated by a feedback loop responding to the number of DN precursors and their competition for stromal niches (40). This concept is clearly illustrated by work from the van den Brink group, who first utilized the OP9-DL1 cell coculture system to produce proTs for the purpose of enhancing thymic recovery from radiation damage (32). An allogeneic BMT model was used to show that proTs enhanced thymic reconstitution and produced a broad repertoire of functional mature peripheral T cells, which protected against immunological challenges (32, 33). Furthermore, human in vitro–generated proTs have demonstrated the ability to enhance thymopoiesis after transplant in mice (44). These studies have set the foundation for future clinical trials using human proT protocols.

As these protocols are developed (27, 30, 31), it is important to select the optimal cell product to expand and infuse. To obtain preclinical data that would direct clinical protocols, we used a congenic BMT model to isolate the effects of proT therapy on posttransplant reconstitution from any confounding factors due to allogeneic reactivity, which can impede positive and negative selection (45). We first examined the homing of in vitro–derived proTs to and within the thymus and their persistence in the periphery. By monitoring luciferase signal from proTs, we were able to track their homing to the thymus and egress to the periphery, establishing that numbers of proT-derived cells peaked thymically and in total at day 21, following which there was a gradual decline, reaching a steady-state level in the periphery. Looking at the movement of these cells within the thymus, we found that proTs were not able to migrate normally on irradiated stromal cells as compared with healthy stromal cells, a trend that was also seen in WT thymocytes on the same stromal cells. These data indicate a disruption in the function of the thymic microenvironment, which agrees with the current literature indicating a defect in thymic progenitor homing as a result of disrupted chemokine production following radiation-induced damage (34, 46). This highlights a potential hurdle to the effectiveness of this therapy and also provides a window of time in which strategies to improve thymic homing may be most effective for optimizing proT therapy.

In the current study, we focused on DN3 cells in our homing and migration studies, in contrast to previous studies focusing on proT therapy in the DN2 subset (32). We sought to delineate the differential effects that these two subsets might have on the thymic microenvironment. We found that adoptive transfer of DN2 cells preferentially increased the lymphopoietic compartment of the thymus, while adoptive transfer of DN3 cells stimulated recovery of mTECs, which was associated with a long-term increase in the numbers of T cells found in the spleen. Consistent with this hypothesis was the localization of purified DN2 cells to the cortex in significantly higher numbers than DN3 cells following transfer to irradiated thymus slices and a subsequent increase of DN3 cell egress to the medulla. In this instance the increase was not statistically significant but an upward trend (Figure 3). The reason for these divergent outcomes may lie in the fact that the DN3 cells retained localization within the medulla of the thymus longer than the DN2 cells. Presumably this would facilitate more interactions between the proTs and mTECs. This crosstalk provides important stimulatory signals for mTECs, which turn over every 10 to 14 days (47, 48), through receptor activator of NF-κB, RANKL, CD80/86, LTβR, and CD40 (26, 49–52). The difference in effects on short-term and long-term thymopoiesis and TEC recovery seen between differing input populations of proTs would suggest that using a combination of DN2 and DN3 proTs may be an optimum approach. While mature proT-derived T cells are present in peripheral lymphoid organs soon after
transplant but are likely progressively diluted out, BM-derived T cells in peripheral lymphoid organs are increased more long term. Presumably both the proTs and BM-derived peripheral mature T cells would enhance protection against infection, but the contributions of each cannot be effectively separately tested. Theoretically, the DN2 component of this approach would result in a short-term increase in thymic size, with a resultant early wave of protective peripheral T cells, while the DN3 subset would supplement long-term repair of the thymic microenvironment associated with a long-term boost in peripheral T cell numbers. As such, our work has implications for future clinical protocols.

Similarly, we observed a direct effect on the proliferative state of TECs in relation to the presence or absence of proTs within the thymus by using a model in which the BM graft (Rag-/-) was incapable of producing proTs that could successfully complete differentiation. When proTs were given Rag-/- BM, we observed a significant but transient increase in the proliferation and numbers of TECs, demonstrating the importance of continued strong recruitment of progenitors from the BM and the resultant crosstalk between immature T cells and TECs (24). While proT therapy does enhance recruitment on its
own, it is reasonable to speculate that better recovery of the thymic microenvironment may be achieved by combining proT therapy with other strategies to enhance recruitment, such as pretreatment of donor marrow with thymic homing-associated chemokines (46). This may improve the insufficient support of thymic homing resulting from radiation damage to the microenvironment (53, 54). Therefore, by better understanding critical factors responsible for the effectiveness of proT therapy, we can more successfully adapt it to the clinic and achieve superior patient outcomes.
Methods

Mice. Mice were used at 8 to 12 weeks of age and were all female. C57BL/6 (B6) mice were purchased from Charles River Laboratories. Congenic B6 CD45.1 mice were purchased from the National Cancer Institute. Congenic B6 Thy1.1 mice were purchased from Jackson Laboratories. B6 luciferase–transgenic mice, provided by Christopher Contag (Stanford University, Stanford, California, USA) (55), were bred in our animal colony under the guidance of in-house veterinary staff. Mice were housed in a specific pathogen–free facility.

Isolation of hematopoietic progenitor cells. BM was collected from femurs and tibiae of B6 Thy1.1 mice after crushing them with a mortar and pestle, followed by red blood cell lysis for 5 minutes at room temperature using PharmLyse (BD). Lineage-positive cells were depleted using a mouse lineage cell depletion kit (Miltenyi Biotec) and then enriched for Sca1 using the EasySep Mouse SCA1 Positive Selection Kit (Stem Cell Technologies).
OP9-DL1 cultures. OP9-DL1 cells were provided by the Zühiga-Pfüller lab. OP9-DL1 cultures and OP9-DL1/hematopoietic progenitor cell cocultures were maintained as previously described (56). Briefly, hematopoietic progenitor cells were seeded onto 70%–90% confluent OP9-DL1 10-cm dishes at 1 × 10^6 per dish in the presence of 5 ng/ml Flt3L (R&D systems) and 5 ng/ml IL-7 (Peprotech). After 7 days, nonadherent cells were harvested and split 1:4 (from 1 dish onto 4 dishes containing new OP9-DL1). Phenotype was assessed by flow cytometry. Nonadherent cells were harvested on day 10 and used for transplant or 2-photon imaging.

**HSCT congeneric model with proT adoptive transfer.** B6 recipients were given 11 Gy total body irradiation using an X-RAD 320 (Precision X-Ray) on day −1. On day 0, recipients were injected intravenously with 5 × 10^6 T cell–depleted B6 CD45.1 BM or Rag-/−/BM cells. T cell depletion was accomplished by incubating the marrow with antibodies against CD4 (GK1.5, National Cell Culture Center) and CD8 (2.43, National Cell Culture Center) at 0.02 mg/ml on ice for 30 minutes, followed by the addition of rabbit serum complement at 10% v/v and incubation at 37°C for 45 minutes. The cells were then washed and counted. Some groups also received 8 × 10^6 bulk proTs, DN2-sorted proTs (CD45+CD4 CD8 CD11b CD11c B220 CD19 NK1.1 Gr1 Ter119 CD25 CD44+), or DN3-sorted proTs (CD45+CD4 CD8 CD11b CD11c B220 CD19 NK1.1 Gr1 Ter119 CD25 CD44−) intravenously on day 0.

**Flow cytometry and cell sorting.** For BrdU incorporation experiments, mice received 1 μg BrdU intraperitoneally 1 hour before harvest. For all harvests, thymi, spleens, and LNs were harvested at various time points after transplant, processed into single-cell suspensions, and analyzed by flow cytometry. TECs were isolated as previously described (57). Briefly, thymi were harvested, nicked, and agitated to release hematopoietic cells. They were then transferred to media containing collagenase D and DNase, dissociated using the Program B setting on a Miltenyi GentleMACS tissue dissociator, and incubated for 15 minutes at 37°C. This was repeated once with fresh media and then again with media containing collagenase/dispose and DNase. All samples were stained with a fixable viability dye conjugated to either eFl780 or eFl450 (eBioscience). Fixation and intracellular/intranuclear staining were performed using the eBioscience Foxp3 staining kit. The following antibodies were purchased from BD Biosciences: CD45 (30-F11), CD3 (500A2), CD8 (53-6.7), cKit (2B8), CD62L (MEL-14), CD11b (M1/70), CD11c (HL3), Ter119 (TER119), CD19 (1D3), CD90.1 AF647 (Thy1.1) (Biolegend), BP-1 FITC (eBioscience), and UEA-1 Biotinylated (Vector Laboratories). Single-cell suspensions were interrogated using a BD LSRFortessa flow cytometer. Data were analyzed using FlowJo V10 (Treestar Inc.).

**In vivo bioluminescent imaging.** In vivo bioluminescent imaging was performed as described previously (58). Mice were injected with firefly luciferin substrate (0.1 ml; 30 mg/ml; Promega) intraperitoneally. After 5 minutes, a Xenogen IVIS imaging system was then used to measure luminescent quantification under “auto” exposure. Data were analyzed and quantified with Living Image 3.0 software (Calipers).

**Confocal microscopy.** Thymi were immersed in OCT Compound for Cryostat Sectioning (Tissue-Tek) in plastic molds and frozen in liquid nitrogen. Thymi were cut into 6-micron sections on a cryostat, fixed in room temperature acetone for 5 minutes, and rehydrated with 1 × PBS. Sections were blocked with the Avidin/Biotin blocking kit (Vector Laboratories) according to the kit’s protocol and then blocked with 10% normal donkey serum (Jackson ImmunoResearch). Primary antibodies used were CD90.1 AF647 (Thy1.1) (Biolegend), BP-1 FITC (eBioscience), and UEA-1 Biotinylated (Vector Laboratories). Secondary antibody Streptavidin Cy3 (Jackson ImmunoResearch) was used. Slides were cover-slipped with hard set DAPI (Vector Labs) and examined by confocal fluorescence microscopy (Olympus FluoView FV1000 BX61 Upright Confocal, Olympus Optical). Images were analyzed using the color segmentation plugin in Fiji image processing software (ImageJ, NIH).

**Two-photon microscopy.** Sample preparation and 2-photon microscopy were performed as previously described (59). Cells labeled with either cell tracker violet or cell tracker orange (Thermo Fisher Scientific) were added to the slices and incubated overnight in complete RPMI at 5% CO₂, and 37°C. Samples were excited with a MaiTai TiSaphire DeepSee HP laser (15 W; Spectra-Physics) at 870 nm, and emissions of 440–480 nm (CTV), 500–520 nm (GFP), and 560–630 nm (CTO) were collected. A 4-channel Leica TCS MP microscope with a resonant scanner containing 2 NDD- and 2 HyD- photomultiplier tubes operating at video rate was used for image capture. Images acquired were 50–100 μm below the top of the slice, and 512 × 512 x-y frames were collected at 3.0-μm steps every 30 seconds for 20 minutes. Image stacks were spectrally unmixed with the LAS AF 3.0 dye separation software module.
(Leica Microsystems). Three-dimensional images, time-lapse movies, and cell tracks were generated with Imaris 8.2 ×64 software (Bitplane), and tracks were verified manually. Tracking data were analyzed in Microsoft Excel with a custom macro as previously described (60).

Epifluorescent microscopy. Thymus lobes from irradiated mice were embedded in low-melt agarose and sliced into 800-μm sections using a vibratome as previously described (59, 61). CD45.1 DN2 and Thy 1.1 DN3 thymocytes were sorted and incubated with the slices for 36 hours. The slices were then fixed with 2% PFA, dehydrated with 30% sucrose and then embedded/frozen in OCT. Cryosections of the slices were stained with Alexa Fluor 647 BP-1 (Biolegend), biotinylated UEA-1 (Vector Laboratories), PE CD45.1 (BD Biosciences), and BV480 Thy1.1 (BD Biosciences). Secondary streptavidin BV421 (Biolegend) was used. Images of the sections were tiled using a Leica DM6000B epifluorescent microscope. Images were spectrally unmixed with the LAS AF 3.0 dye separation software module (Leica Microsystems) and analyzed with Imaris 8.2 ×64 software (Bitplane). Surfaces for CD45.1 (DN2), Thy1.1 (DN3), and BP-1 (cortex) were generated, and distances to the cortices of DN2 and DN3 surfaces were calculated by the Imaris distance transformation function. Cells residing in the cortex were determined to be 0–1 μm from the cortex surface, cells in the cortical medullary junction were determined to be 1–10 μm from the cortex, and cells in the medulla were determined to be >10 μm from the cortex. Medulla-residing cells also had to display an UEA-1 MFI higher than background.

Statistics. Data are shown as mean ± SEM. Data sets were compared using 2-way ANOVA. Significance was defined as \( P < 0.05 \).

Study approval. Mice were used with the approval of the University of Minnesota institutional animal care and use committee.

Author contributions
MJS designed and performed experiments, analyzed data, and wrote the manuscript. DKR, SLP, and MJR performed experiments. JM and KCO performed experiments and analyzed data. HES, BTF, AB, KAH, GAH, and JT provided guidance in experimental design and edited the manuscript. MM and JCZP provided guidance in experimental design, provided reagents, and edited the manuscript. BRB designed experiments and edited the manuscript.

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Address correspondence to: Bruce R. Blazar, Pediatrics BMT, MMC 366 Mayo, 8366A (Campus Delivery Code), 420 Delaware Street SE, Minneapolis, Minnesota 55455, USA. Phone: 612.626.2734; E-mail: blaza001@umn.edu.


