Lipid Phosphate Phosphatase 3 Enables Efficient Thymic Egress

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Lipid phosphate phosphatase 3 enables efficient thymic egress

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Abbreviations used: CD4SP, CD4 single-positive; DN, double negative; DP, double positive; LPP3, lipid phosphate phosphatase 3; MFI, mean fluorescence intensity; pIpC, polyinosinic:polycytidylic acid; S1P, sphingosine-1-phosphate; S1P, receptor 1; SPP1, sphingosine-1-phosphate; S1PR1, S1P receptor 1; SPDE3, S1P-degrading enzyme.

The signaling lipid sphingosine-1-phosphate (S1P) plays critical roles in mammalian biology. Extracellular S1P binds five G protein–coupled receptors, S1P receptors 1–5 (Hla, 2004), and intracellular S1P binds an expanding set of proteins (Hait et al., 2009; Alvarez et al., 2010; Puneet et al., 2010). The concentration of S1P is high in circulatory fluids, and plasma S1P stabilizes junctions between vascular endothelial cells (Lee et al., 1999; Kono et al., 2008; Camerer et al., 2009). The concentration of extracellular S1P is low in lymphoid tissues compared with blood and lymph, and this difference directs lymphocyte egress from lymphoid organs into circulation (Schwab and Cyster, 2007). Although extracellular S1P in lymphoid organs, and likely most tissues, is low in homeostasis, S1P may increase upon inflammation. Elevated S1P has been shown to promote angiogenesis and to enhance proinflammatory responses of innate and adaptive immune cells (Rivera et al., 2008; Liu et al., 2010). Drugs targeting S1P signaling are in clinical trials as immune suppressants (Carroll, 2010). In fact, FTY720 has just been approved by the United States Food and Drug Administration for treatment of multiple sclerosis (Brinkmann et al., 2010). By blocking exit from lymphoid organs, these drugs prevent activated T cells from trafficking to organs under autoimmune attack. These drugs may also have direct antiinflammatory effects.

Despite its importance, little is known about how the distribution of S1P is controlled in vivo. This question is particularly interesting because it is difficult to understand how a ubiquitously made lipid functions as a signal that requires precise spatial and temporal control. We have found that lipid phosphate phosphatase 3 (LPP3) enables efficient export of mature T cells from the thymus into circulation, and several lines of evidence suggest that LPP3 promotes exit by destroying thymic S1P. Although five additional S1P-degrading enzymes are expressed in the thymus, they cannot compensate for the loss of LPP3. Moreover, conditional deletion of LPP3 in either epithelial cells or endothelial cells is sufficient to inhibit egress. These results suggest that S1P generation and destruction are tightly regulated and that LPP3 is essential to establish the balance.
sphingolipid metabolism (Hannun and Obeid, 2008); metabolic S1P must be destroyed before it accumulates in the interstitial space. Second, plasma S1P must be excluded from the tissue. S1P concentrations in plasma are in the micromolar range, and although tissues are constantly bathed with transudated plasma to bring nutrients and remove waste, S1P in the interstitial fluid of lymphoid organs has been estimated to be sub-nanomolar (Schwab and Cyster, 2007). S1P’s abundance suggests the possibility that levels could be changed rapidly in response to physiological cues and the requirement that its concentration be tightly regulated.

Many factors are undoubtedly required to maintain low tissue S1P but one key is likely how S1P is degraded. Strikingly, not one but six enzymes are known to destroy S1P in vitro. These enzymes fall into two classes. The first consists of S1P lyase, which cleaves S1P to 2-hexadecenal and phosphoethanolamine, and S1P phosphatases 1 and 2, which dephosphorylate S1P to sphingosine (Saba and Hla, 2004). These three enzymes are thought to be highly specific for S1P and reside predominantly in the endoplasmic reticulum. S1P lyase is required to maintain low lymphoid organ S1P (Schwab et al., 2005). Although the source of S1P destroyed by S1P lyase is not known, it is likely to be S1P made in the course of sphingolipid metabolism; S1P lyase’s intracellular location positions it well for this function. The second class of S1P-degrading enzymes consists of three phosphatases: lipid phosphate phosphatases 1, 2, and 3 (Roberts et al., 1998; Brindley et al., 2000; Pyne et al., 2004). These enzymes can dephosphorylate a range of substrates, including S1P, ceramide-1-phosphate, lysophosphatidic acid, and phosphatidic acid. The lipid phosphate phosphatases have been shown in many cell types to localize to the plasma membrane with their active site facing the tissue. S1P enters the tissues from the blood or secreted by tissue-resident cells. The role of the lipid phosphate phosphatases in controlling tissue S1P is largely unknown.

In this paper, we ask what is the role of lipid phosphate phosphatase 3 (LPP3; encoded by Ppap2b) in regulating extracellular S1P within lymphoid organs. We found that LPP3 is essential to enable efficient export of mature T cells from the thymus to the blood, and several lines of evidence suggest that LPP3 promotes egress by maintaining low thymic S1P. Although all six S1P-degrading enzymes are expressed in the thymus, the loss of LPP3 alone in either epithelial cells or endothelial cells is sufficient to delay exit.

RESULTS
LPP3 is required to keep thymic S1P low
Because LPP3 is essential in embryonic development (Escalante-Alcalde et al., 2003), we used Ppap2b−/− mice to study the function of LPP3 in lymphoid organs (Escalante-Alcalde et al., 2007). We crossed in Cre recombinase driven by the interferon-inducible Mx1 promoter (Mx1-Cre; Kühl et al., 1995). Pups were treated with polyinosinic-polycytidylic acid (pI-pC) 3–5 d after birth to activate the Mx1 promoter and Cre expression. Adults were analyzed at least 6 wk later. This induction of Mx1-Cre mediates efficient deletion in a wide range of tissues and lineages, including hematopoietic cells and a subset of endothelial cells (Pappu et al., 2007). For simplicity, we refer to pI-pC–treated Ppap2b−/− Mx1Cre− and Ppap2b+/− Mx1Cre+ mice as LPP3 deficient. Littermate controls were pI-pC treated at the same time as LPP3-deficient mice and retained at least one intact copy of LPP3. To assess whether LPP3 regulates thymic S1P we examined three parameters that are dependent on S1P signaling: the rate of exit of mature T cells from the thymus, the level of surface CD69 on mature thymocytes, and the level of surface S1P receptor 1 (S1PR1) on mature thymocytes.

LPP3 deficiency impairs thymic egress
We first asked whether egress of mature T cells from the thymus was impaired in LPP3-deficient mice. T cells must express S1PR1 to exit lymphoid organs into circulatory fluids (Allende et al., 2004; Matloubian et al., 2004; Weinreich and Hogquist, 2008). Furthermore, disruption of the S1P gradient, either by removing circulatory S1P (Pappu et al., 2007) or by raising lymphoid tissue S1P (Schwab et al., 2005), blocks egress. We expected that if thymic S1P were elevated in the absence of LPP3, exit would be slowed, resulting in an accumulation of mature T cells in the thymus.

We observed a 2.5-fold increase in the percentage of mature CD4 single-positive (CD4SP) T cells (CD4+CD8− CD62L+CD69−) among total CD4SP (CD4+CD8−) thymocytes in LPP3-deficient mice compared with littermate controls (Fig. 1 A and B). There is also an increase in the percentage of CD4SP T cells among total thymocytes. When we calculated absolute numbers, we found a 4.5-fold increase of mature CD4SP T cells in the thymus of LPP3-deficient mice without any change in the number of immature CD4SP precursors (Fig. 1 C). The absolute numbers are more variable than the percentages, as the total size of the thymus varies with many factors including age and stress. Nonetheless, in each case the number of mature CD4SP was larger in the LPP3−/− mice than its littermate control. We saw no change in the number of double-negative (DN) T cells and a 30% reduction in the number of double-positive (DP) T cells in the thymus of LPP3−/− mice compared with littermate controls (Fig. S1 A). As one might predict from a slowing of thymic egress, we saw a moderate reduction in the number of T cells in the periphery (Fig. S1 B). Mature CD8 single-positive T cells similarly accumulate in the thymus of LPP3−/− mice (Fig. S1 C).

The accumulation of mature T cells suggests an egress block but could also result from enhanced proliferation of mature T cells. Developing T cells mature from the DN to the DP to the SP stage. In WT mice, thymocytes proliferate extensively as they transition from DN to DP and relatively infrequently at the SP stage (Egerton et al., 1990). To assess the possibility that mature SP T cells divide more frequently in LPP3-deficient mice, we treated mice continuously with the thymidine analogue BrdU in the drinking water to label proliferating cells. After 11 d of BrdU treatment, virtually all
Figure 1. LPP3 is required for efficient thymic egress. (A) CD4SP thymocytes (CD4+CD8lo/hi Mx1Cre− or Ppap2b−/− Mx1Cre+; treated with 0.8 mg/ml BrdU in the drinking water for 11 d) and littermate controls (B) Percentage of mature CD4SP thymocytes (CD4+CD8lo/hi Mx1Cre−) among total CD4SP thymocytes. (C) Total number of mature CD4SP thymocytes and immature CD4SP thymocytes (CD4+CD8− CD62LhiCD69−) in LPP3-deficient mice and littermate controls. (D) GFP levels of mature CD4SP thymocytes from a representative Rag2p-GFP+ LPP3-deficient (Δ) mouse, mature CD4SP thymocytes from its Rag2p-GFP+ littermate control (F), and naive (CD69hi/CD69lo) CD4 T cells from the blood of the LPP3-deficient Rag2p-GFP+ mouse, measured by flow cytometry. Data are representative of four pairs of mice analyzed in two experiments. (F) Sulfo-NHS biotin was injected into the thymus of LPP3-deficient mice and littermate controls. 16 h later, biotin-labeled mature CD4SP T cells in the periphery were enumerated. The graph shows the ratio of the number of labeled cells in the periphery to the number of labeled cells remaining in the thymus. Each point represents an individual mouse and bars show the mean. The graph compiles data from seven LPP3-deficient mice and four littermate controls analyzed in two experiments. Mature T cells in the thymus of LPP3-deficient and littermate controls were labeled with similar efficiency (mean 57 and 62%, respectively).

mature CD4SP T cells in control were BrdU+ (Fig. 1 D). In contrast, in LPP3-deficient mice a substantial number of mature CD4SP T cells remained unlabeled (Fig. 1 D). The increased fraction of BrdU− mature SP T cells is inconsistent with enhanced proliferation of mature SP thymocytes in LPP3-deficient mice. Instead, it is consistent with delayed egress; the BrdU− cells likely developed before the labeling period began and remained in the thymus for an extended period. Mature CD8SP T cells had a similar pattern (Fig. S1 D).

Although an accumulation of mature T cells in the absence of proliferation is consistent with a failure to exit, it is also possible that this could reflect enhanced reentry of peripheral T cells into the thymus. To distinguish between these possibilities, we took advantage of Rag2p-GFP reporter mice (Yu et al., 1999; Boursalian et al., 2004). In these mice, GFP transcription is driven by the Rag2 promoter. After RAG is turned off at the DP stage, GFP protein levels decay with a half-life of ~56 h (McCaughtry et al., 2007). GFP acts as a clock, with its abundance reflecting the amount of time that has passed since positive selection. Analysis of LPP3-deficient Rag2p-GFP+ mice and Rag2p-GFP+ littermate controls revealed that mature CD4SP LPP3-deficient thymocytes have lower GFP than mature CD4SP control thymocytes but higher GFP than naive CD4 T cells circulating in the periphery (Fig. 1 E). Mature CD8SP T cells behave similarly (Fig. S1 E). This pattern is consistent with prolonged thymic residence and delayed egress but not with reentry of peripheral T cells.

Finally, we directly tested the efficiency of thymic egress by injecting sulfo-NHS biotin into the thymus of LPP3-deficient mice and littermate controls and tracking the appearance of labeled cells in the periphery. To minimize inflammation, we used ultrasound to guide the injection needle (Blair-Handon et al., 2010). After 16 h, we found that the ratio of labeled CD4+ T cells in the periphery (pooled blood, spleen, mesenteric, inguinal, axillary, and brachial LNs) to labeled mature CD4SP T cells, as well as naive blood CD4+ T cells, was measured by flow cytometry. Data are representative of three pairs of mice analyzed in three experiments. (E) GFP levels of mature CD4SP thymocytes from a representative Rag2p-GFP+ LPP3-deficient (Δ) mouse, mature CD4SP thymocytes from its Rag2p-GFP+ littermate control (F), and naive (CD69hi/CD69lo) CD4 T cells from the blood of the LPP3-deficient Rag2p-GFP+ mouse, measured by flow cytometry. Data are representative of four pairs of mice analyzed in two experiments. Mature T cells in the thymus of LPP3-deficient and littermate controls were labeled with similar efficiency (mean 57 and 62%, respectively).

LPP3 deficiency causes down-modulation of CD69 on mature T cells

Our second indication that LPP3 regulates thymic S1P was its requirement for efficient thymic egress. LPP3-deficient mice have lower CD69 than mature CD4SP thymocytes but higher GFP than naive CD4 T cells circulating in the periphery (Fig. 1 E). Mature CD8SP T cells behave similarly (Fig. S1 E). This pattern is consistent with prolonged thymic residence and delayed egress but not with reentry of peripheral T cells.

Finally, we directly tested the efficiency of thymic egress by injecting sulfo-NHS biotin into the thymus of LPP3-deficient mice and littermate controls and tracking the appearance of labeled cells in the periphery. To minimize inflammation, we used ultrasound to guide the injection needle (Blair-Handon et al., 2010). After 16 h, we found that the ratio of labeled CD4+ T cells in the periphery (pooled blood, spleen, mesenteric, inguinal, axillary, and brachial LNs) to labeled mature CD4SP T cells, as well as naive blood CD4+ T cells, was measured by flow cytometry. Data are representative of three pairs of mice analyzed in three experiments. (E) GFP levels of mature CD4SP thymocytes from a representative Rag2p-GFP+ LPP3-deficient (Δ) mouse, mature CD4SP thymocytes from its Rag2p-GFP+ littermate control (F), and naive (CD69hi/CD69lo) CD4 T cells from the blood of the LPP3-deficient Rag2p-GFP+ mouse, measured by flow cytometry. Data are representative of four pairs of mice analyzed in two experiments. Mature T cells in the thymus of LPP3-deficient and littermate controls were labeled with similar efficiency (mean 57 and 62%, respectively).
reproduced in mice treated with the S1PR1-selective agonist SEW2871 (Alfonso et al., 2006). CD69 and S1PR1 interact, and S1PR1 mediates CD69 internalization in a pertussis toxin-sensitive manner (Shiow et al., 2006). CD69 is down-modulated on mature but not immature SP T cells. S1pr1 mRNA is >10-fold more abundant in mature than in immature CD4SP thymocytes (Matloubian et al., 2004), and surface S1PR1 protein is readily detectable on mature but not immature SP T cells (Zachariah and Cyster, 2010). We expected that if S1P were elevated in the thymus of LPP3-deficient mice, surface CD69 on mature SP T cells would decrease.

We observed down-modulation of CD69 on the surface of mature SP T cells in LPP3-deficient animals compared with littermate controls (Fig. 1 A). To pool experiments, we calculated the ratio of CD69 mean fluorescence intensity (MFI) of littermate control mature CD4SP thymocytes/CD69 MFI of LPP3-deficient mature CD4SP thymocytes. This ratio was $\sim 3$ (Fig. 2 A). The down-modulation of CD69 on mature T cells in LPP3-deficient mice is not simply secondary to a longer residence time, as it was also seen on the recently mature GFPhi cells in LPP3-deficient Rag2p-GFP mice (Fig. 2 B). The decreased CD69 on mature T cells is also not the result of a failure to up-regulate fully CD69 during positive selection, as we saw no difference in the level of CD69 on immature CD4SP T cells from LPP3-deficient mice and littermate controls (Figs. 1 A and 2 C).

**LPP3 deficiency causes internalization of S1PR1**

Our third test of whether LPP3 is required to maintain low extracellular thymic S1P was the level of surface S1PR1 on mature SP T cells. Mature SP T cells normally express abundant S1PR1, enabling them to exit the thymus efficiently (Matloubian et al., 2004). S1PR1 is internalized upon binding S1P (Liu et al., 1999; Schwab et al., 2005), and we expected that if LPP3-deficient mice had elevated thymic S1P, surface S1PR1 on thymic T cells would be reduced.

We found that the level of S1PR1 is two- to threefold lower on the surface of mature SP thymocytes of LPP3-deficient mice than littermate controls (Fig. 2 D). S1pr1 transcript is comparable between LPP3-deficient and littermate control mice, suggesting that the decrease is posttranscriptional (Fig. S2 A). CCR7 surface protein also does not differ between LPP3-deficient and littermate control mice, suggesting that the decrease does not reflect a global down-modulation of G protein–coupled receptors (Fig. S2 B).

Collectively, the egress block, CD69 down-modulation, and reduction in surface S1PR1 strongly suggest that the S1P-degrading enzyme LPP3 is required to maintain low thymic S1P. Using mass spectrometry, we were unable to detect elevated S1P in whole thymic tissue of LPP3-deficient mice (Fig. S2C), which is not surprising given the incomplete down-modulation of S1PR1. The concentration of S1P in the thymus of control animals is $\sim 0.5$ pmol/mg wet weight, which translates to 500 nM assuming that the density of the thymus is the same as water (an underestimate). This is consistent with previous measurements of S1P in the whole embryo (Mizugishi et al., 2005). In contrast to the total level of S1P in the thymus, 10 nM of extracellular S1P is sufficient to decrease surface S1PR1 on mature thymocytes to below our limit of detection (Schwab et al., 2005). The S1P measured by mass spectrometry of whole thymic tissue is largely inaccessible to S1P receptors, consistent with S1P’s role as an intracellular metabolic intermediate. Against the variability of our measurements of S1P in control thymus, we are unable to detect an increase in extracellular S1P that causes partial receptor internalization.

**LPP3 is required in radiation-resistant cells**

To determine which cell type requires LPP3 to maintain low thymic S1P, we began by generating BM chimeras.

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**Figure 2.** LPP3 deficiency causes down-modulation of CD69 and S1PR1. (A) The ratio of CD69 MFI of mature CD4SP thymocytes (F)/CD69 MFI of mature CD4SP thymocytes from LPP3-deficient mice (Δ). Each point represents one pair of mice, and the bar represents the mean. The graph compiles 13 experiments with a total of 21 pairs of mice. (B) Dot plots (left) show expression of GFP versus forward scatter (FSC) by mature CD4SP T cells from a representative Rag2p-GFP LPP3-deficient mouse (Δ) and Rag2p-GFP littermate control (F), and they indicate the GFP+ gate. The GFP MFI within the GFP+ gate differs by $<100$ bu-tween littermate controls and LPP3-deficient mice. The graph compiles data from four pairs of mice analyzed in two experiments. (C) The ratio of CD69 MFI of immature CD4SP thymocytes from littermate controls (F)/CD69 MFI of immature CD4SP thymocytes from LPP3-deficient mice (Δ). Each point represents one pair of mice, and each bar represents the mean. The graph compiles six experiments with a total of eight pairs of mice. (D) Surface S1PR1 levels on mature CD4SP thymocytes. The thin line shows staining with a control antibody. Data are representative of three pairs of mice analyzed in three experiments.
LPP3 is expressed by multiple radiation-resistant cell types

To determine which radiation-resistant cell type must express LPP3 to maintain low thymic S1P, we began by asking which cells express $Ppap2b$ mRNA using quantitative PCR on sorted cells from WT mice. Consistent with the importance of LPP3 on radiation-resistant cells, we found that $Ppap2b$ mRNA is enriched 100-fold in CD45+ thymic stroma compared with T cells. However, there are no dramatic differences in expression between EpCAM+ epithelial cells, CD31+ endothelial cells, and other CD45+ stromal cells (Fig. S3 A). We also asked whether any of these cell types differ in expression of the other five S1P-degrading enzymes, suggesting that they might be uniquely dependent on LPP3. We found that LPP1 ($Ppap2a$) and LPP2 ($Ppap2c$) have a similar pattern to LPP3 (Fig. 3 A) and that the S1P-specific enzymes S1P lyase (Spgl1), S1P phosphatase 1 (Spp1; Sgpp1), and SPP2 (Sgpp2) are also expressed by all subsets of radiation-resistant cells (Fig. 3 B).

Our next attempt to narrow down the cell type that must express LPP3 was to ask whether Mx1-Cre deletes preferentially in any radiation-resistant cell types in the thymus. We crossed mice expressing Mx1-Cre with an enhanced (E) YFP reporter strain, R26R-EYFP, which has EYFP in the ubiquitously expressed Rosa26 locus preceded by a floxed transcriptional stop (Srinivas et al., 2001), treated the pups with pL-Cre at 3–5 d after birth, and analyzed the mice as adults. Analysis of disaggregated thymocytes by flow cytometry revealed deletion in approximately two thirds of epithelial cells, one third of endothelial cells, and 10% of other stroma, as well as >90% of hematopoietic cells (Fig. 4 C). Because the strong YFP expression by lymphocytes obscured analysis of YFP expression by radiation-resistant cells using confocal microscopy, we also reconstituted lethally irradiated adults with WT BM. Analysis of thymic sections from the chimeras confirmed that Mx1-Cre deletes in multiple radiation-resistant cell types, including epithelial cells and endothelial cells (Fig. 4 D).

Staining of frozen thymic sections with an antibody to LPP3 revealed robust protein expression by medullary epithelial cells, which was largely absent in LPP3-deficient mice (Fig. 4 E). LPP3 seemed to ring the endothelial wall of blood vessels, consistent with results showing that LPP3 localizes to the basolateral side of cultured MDCK cells (Jia et al., 2003). This staining was less frequent in LPP3-deficient mice (Fig. 4 E). LPP3 protein was also expressed by a range of other stromal cells, likely including pericytes (Fig. S4).

LPP3 is expressed in both endothelial cells and epithelial cells

We then began a candidate approach to determine which cell type must express LPP3. Because we observed high expression of LPP3 by endothelial cells and partial deletion in endothelial cells by Mx1-Cre, and because endothelial cells sit at the border of the high S1P environment of the blood and the low S1P environment of the thymus, we considered the possibility that LPP3 was required in endothelial cells to maintain low thymic S1P. To delete LPP3 in vascular endothelial cells, we crossed $Ppap2b^{fl/fl}$ mice with mice carrying Cre recombinase driven by the Tie2 promoter (Kisanuki et al., 2001).
We had no live-born Ppap2b\(^{-/-}\)Tie2-Cre\(^{+}\) pups (unpublished data), which was not surprising as Ppap2b\(^{-/-}\) embryos suffer severe vascular defects (Escalante-Alcalde et al., 2003). To circumvent embryonic lethality, we used the inducible CreERT2 recombinase under the endothelial VE-Cadherin (Cdh5) promoter (Wang et al., 2010). We treated Ppap2b\(^{-/-}\) mice with pI-pC 3–5 d after birth and analyzed at least 6 wk later. YFP expression by disaggregated thymic cell populations was measured by flow cytometry. CD4\(^{+}\) cells are primarily T cells, as they were isolated by mechanical disruption. Each point represents an individual mouse, and bars show the mean. The graph compiles two mice analyzed in two experiments.

We saw a twofold accumulation of mature T cells in Ppap2b\(^{-/-}\)-Cdh5(PAC)-CreERT2\(^{+}\) mice compared with littermate controls, with tamoxifen treatment. We observed at most a slight down-modulation of S1PR1 on mature T cells (Fig. S5 B). We noted a decrease of surface CD69 on mature T cells, which is, again, significant but less dramatic than in mice in which deletion was mediated by Mx1-Cre (Fig. 5, C and D; CD69 is higher in Ppap2b\(^{-/-}\)-Cdh5(PAC)-CreERT2\(^{+}\) than in Ppap2b\(^{-/-}\) or \(\beta\) Mx1Cre\(^{+}\) mice with \(P < 0.01\)). Finally, we observed at most a slight down-modulation of S1PR1 on mature T cells (Fig. 5 E). We suspect that down-modulation is difficult to detect as a result of the relatively mild phenotype and limited sensitivity of our antibody.

To determine the pattern of deletion, we crossed mice expressing Cdh5(PAC)-CreERT2 with the YFP reporter strain. Confocal microscopy of thymic sections revealed specific deletion in endothelial cells (Fig. 5 F and Fig. S5 A), which was confirmed by analysis of disaggregated thymic cells by flow cytometry. We saw variable excision, ranging from 20% to 70%, whereas deletion in other cell types was undetectable (Fig. 5 G). It is possible that the intermediate effect of LPP3 deletion within endothelial cells is a result of incomplete excision. It is also possible that the loss of LPP3 in endothelial cells causes a very small increase in thymic S1P but that...
the elevated S1P is precisely where T cells most need a sharp gradient to egress as they cross the blood vessel wall into circulation.

Because we also observed high expression of LPP3 by thymic epithelial cells and efficient deletion in thymic epithelial cells by Mx1-Cre, we considered the possibility that LPP3 was required in these cells. To delete LPP3 in epithelial cells, we used the Cre recombinase under the Keratin 14 promoter (Dassule et al., 2000). We bred Ppap2b<sup>f/k</sup>-Krt14-Cre<sup>e</sup> mice. Littermate controls were Ppap2b<sup>f/k</sup>-Krt14-Cre<sup>e</sup>, Ppap2b<sup>f/k</sup>, or Ppap2b<sup>f/k</sup>-.

Mature T cells in the thymus of Ppap2b<sup>f/k</sup>-Krt14-Cre<sup>e</sup> mice have a 3.5-fold accumulation (Fig. 6, A and B), accompanied by a small reduction of T cells in the periphery (Fig. S6 B). Mature T cells in the thymus of Ppap2b<sup>f/k</sup>-Krt14-Cre<sup>e</sup> mice also display down-modulation of CD69 (Fig. 6, C and D) and a twofold decrease in S1PR1 compared with littermate controls (Fig. 6 E).

To determine the pattern of deletion, we crossed mice expressing Keratin 14-Cre with the YFP reporter strain. In both the LPP3 and YFP reporter crosses, Cre recombinase was carried only by the father to avoid potential widespread deletion as a result of expression in the oocyte (Hafner et al., 2004).

Confocal microscopy of thymic sections revealed deletion primarily in medullary epithelial cells (Fig. 6 F, Fig. S6 A). This pattern was also seen in analysis of disaggregated thymic cells by flow cytometry. We observed excision within ~75% of EpCAM<sup>+</sup> cells (Fig. 6 G).

DISCUSSION

We found that LPP3 is essential to enable efficient thymic egress, and several lines of evidence suggest that LPP3 promotes exit by maintaining low thymic S1P. In the absence of biochemical evidence of increased S1P, uncertainty remains in the interpretation of our results; measurements of total thymic S1P by mass spectrometry are uninformative because the background of intracellular S1P (500 nM), combined with measurement variability, is too high to detect an increase in extracellular S1P predicted to cause partial S1PR1 down-modulation (<10 nM or 2% of total). Nonetheless, because S1P is the only LPP3 substrate that has been implicated in egress, CD69 down-modulation, or S1PR1 internalization,
the simplest interpretation of our data is that LPP3 is functioning to limit thymic S1P.

This is the first time to our knowledge that a phosphatase has been implicated in lymphoid organ S1P regulation in vivo. The Drosophila melanogaster homologues of LPP3, wunner and wunner2, are required to guide germ cell migration in the developing embryo. Germ cells and the surrounding tissue compete for hydrolysis of a lipid phosphate, although the identity of this lipid is unknown (Renault and Lehmann, 2006). Ppap2b<sup>−/−</sup> mice die at or before embryonic day 10.5 with failure to form a chorioallantoic placenta and yolk sac vasculature (Escalante-Alcalde et al., 2003). It is not known whether this phenotype is the result of an imbalance in phospholipids and, if so, which phospholipid, or whether it is the result of a phosphatase-independent function of LPP3 (Huntsoe et al., 2003). Ppap2b<sup>−/−</sup>Nestin-Cre<sup>+</sup> mice have increased S1P in the cerebellum (López-Juárez et al., 2011). LPP3 is needed on both endothelial and epithelial cells, revealing a novel role of epithelial cells in enabling thymic egress.

We further found that although LPP1 and LPP2 have a similar expression pattern to LPP3, and S1P lyase and SPP1 are widely expressed in the thymus, they cannot compensate for the loss of LPP3. Moreover, deletion of LPP3 in either epithelial cells or endothelial cells is sufficient to cause an increase in thymic S1P and an egress block. These observations suggest that S1P generation and destruction are finely balanced. Little is known about LPP3 regulation, although LPP3 expression by endothelial cell lines is increased upon exposure to stimuli including VEGF and TNF (Huntsoe et al., 2003; Wary and Huntsoe, 2005; Senda et al., 2009), and LPP3 expression by epithelial lines is increased upon exposure to stimuli including EGF (Kai et al., 1997; Huntsoe et al., 2003). We propose that S1P levels could be rapidly increased in response to physiological signals, not only by increasing phosphorylation of sphingosine but also simply by inhibiting degradation. This may be one reason that a ubiquitous lipid evolved to be a signaling molecule.

One question raised by these findings is what is the normal spatial distribution of S1P within the thymus and how is it altered in the absence of LPP3? On average, extracellular S1P is lower in the thymus than in circulation (Schwab and Cyster, 2007), but whether there are local variations remains unknown. For example, whether a gradient of S1P guides T cells toward egress structures or whether T cells encounter S1P only as they probe the blood vessels is an open question. We hypothesize that in the absence of LPP3 in epithelial cells, S1P may increase only proximal to blood vessels. Although the loss of LPP3 in endothelial cells has a

Figure 6. LPP3 expression by epithelial cells is required to maintain low thymic S1P. Mice in which LPP3 was deleted by Keratin 14-Cre (TEC-Δ, Ppap2b<sup>−/−</sup>Krt14Cre<sup>+</sup>) or littermate controls (TEC-F; Ppap2b<sup>−/−</sup>Krt14Cre<sup>−</sup>, Ppap2b<sup>−/−</sup> or Ppap2b<sup>−/−</sup>) were analyzed at 6 wk or older. (A) Percentage of mature CD45<sup>SP</sup> thymocytes among total CD45<sup>SP</sup> thymocytes. (B) Total number of mature CD45<sup>SP</sup> thymocytes and immature CD45<sup>SP</sup> thymocytes. In A and B, each point represents an individual mouse and bars show the mean. Graphs compile 11 experiments with a total of 13 mice per group. (C) Expression of CD68 and CD62L on CD45<sup>SP</sup> thymocytes. (D) The ratio of CD69 MFI of mature CD45<sup>SP</sup> thymocytes from littermate controls/CD69 MFI of mature CD45<sup>SP</sup> thymocytes from TEC-Δ mice. Each point represents one pair of mice and the bar represents the mean. The graph compiles 13 pairs of mice analyzed in 11 experiments. (E) Surface S1PR1 expression on mature CD45<sup>SP</sup> thymocytes. Thin line shows staining with a control antibody. Data are representative of three pairs of mice analyzed in three experiments. (F) Frozen thymic sections from R26R-EYFP-Krt14Cre<sup>+</sup> mice were stained for EpCAM to mark epithelial cells and CD8 to distinguish the cortex from the medulla and visualized by confocal microscopy. Bar, 100 μm. Image is representative of two mice in two experiments. (G) YFP expression by disaggregated thymic cell populations from R26R-EYFP-Krt14Cre<sup>+</sup> mice was measured by flow cytometry. CD45<sup>−</sup> cells are primarily T cells, as they were isolated by mechanical disruption. Each point represents an individual mouse and bars show the mean. The graph compiles two mice analyzed in two experiments.
minimal effect on average T cell S1PR1, it may desensitize T cells to S1P just where they most need a sharp gradient to cross into circulation. LPP3 is also expressed by other stromal cells, likely including pericytes. Whether S1P is regulated differently in the thymic parenchyma, at the border with blood, and in the perivascular space warrants further investigation.

A second, related, question raised by these findings is what is the source of S1P that LPP3 destroys? LPP3 may be responsible for destroying excess S1P made intracellularly during membrane sphingolipid metabolism and secreted at some rate. LPP3 may also prevent plasma S1P from entering tissues. Vascular endothelial cells sit at the boundary of the high S1P environment of the blood and the low S1P environment of the tissue, and endothelial LPP3 may prevent S1P from crossing the blood vessel wall. Hematopoietic cells, particularly erythrocytes, are the major source of plasma S1P, but endothelial cells themselves may also contribute (Pappu et al., 2007; Hla et al., 2008; Venkataraman et al., 2008). Blood vessel endothelial cells efficiently secrete S1P in culture (Venkataraman et al., 2008), and lymphatic endothelial cells are the main source of lymph S1P (Pham et al., 2010). Intriguingly, LPP3 has been shown to localize to the basolateral side of cultured MDCK cells (Jia et al., 2003), and our LPP3 antibody stain seems to ring blood vessels. The subcellular organization of S1P metabolic enzymes may enable vascular endothelial cells to play a dual role in S1P generation and destruction. Thymic vessels are unique in that they have a double-walled structure with vascular endothelial cells surrounded by epithelial cells (Ushiki, 1986; Kato, 1997; Ushiki and Takeda, 1997; Mori et al., 2007), so a requirement for LPP3 in epithelial cells is also consistent with a role in destroying plasma S1P.

Although our original hypothesis was that LPP3 is destroying metabolic or plasma S1P, it is also possible that LPP3 is responsible for preventing the spread of S1P made and secreted for a highly localized signaling. Like vascular endothelial cells, epithelial cells express high levels of the S1P generating enzyme sphingosine kinase 1 (unpublished data). A role for S1P production by thymic epithelial cells has not been shown, although many are plausible. Epithelial cell S1P secretion may facilitate T cell egress, leading them across the first basement membrane of the double-walled vessels. This would be analogous to the role of pericytes, which sit in the perivascular space and whose S1P secretion promotes exit (Zachariah and Cyster, 2010). Epithelial cell S1P may affect T cell motility or shape T cell maturation; T cell S1PR1 has been shown to inhibit development of regulatory T cells (Liu et al., 2009), but the source of S1P within the thymus has not been identified. Finally, it is tempting to speculate that S1P may play a more general role in epithelial cell junctions, analogous to its role in endothelial barrier function. In this regard, it is interesting to note that LPP3 was first cloned in an attempt to identify genes required for intestinal epithelial cell differentiation (Barillà et al., 1996). Currently there are no techniques to measure changes in extracellular S1P in nonlymphoid tissues, but LPP3 may play a role in regulating not only thymic S1P but also S1P in heavily epithelial tissues, such as the lung, small intestine, and skin and may be an attractive target to manipulate S1P selectively in these organs. The observation that S1P levels are very tightly regulated offers the hope that highly specific drugs can be developed to manipulate S1P signaling, lymphocyte trafficking, and inflammation.

MATERIALS AND METHODS

**Mice.** *Ppap2b* (Escalante-Alcalde et al., 2007), R26R-EYFP (Srinivas et al., 2001), Rag2cre-GFP (Yu et al., 1999), Mx1-cre (Kühn et al., 1995), Keratin 14-cre (Dausé et al., 2000), Foxn1-cre (Gordon et al., 2007), Tie2-cre (Kitamura et al., 2001), and Cdh5(PAC)-creERT2 (Wang et al., 2010) mice have been previously described. CD45.2+ B6 and CD45.1+ congenic B6 mice were obtained from the National Cancer Institute.

To induce deletion by Mx1-cre, 3–5-d-old mice received a single intraperitoneal injection with 50–100 µl pl-g-p-c (Midland Scientific) at a concentration of 10 mg/ml in sterile PBS. To induce deletion by VE-Cadherin creERT2, 3–4-wk-old mice received three intraperitoneal injections on three consecutive days with 50 µl tamoxifen (Sigma Aldrich) at a concentration of 20 mg/ml in maize oil.

Mice were housed in specific pathogen-free conditions at the Skirball Institute animal facility. All animal experiments were performed in accordance with protocols approved by the New York University Institutional Animal Care and Use Committee.

**BM chimeras, BrdU labeling, intrathymic injection, and deoxypyridoxine treatment.** For BM chimeras, recipients were lethally irradiated by two 6.5 Gray doses of γ irradiation from a cesium source separated by 3 h and received 2–10 × 10^6 BM cells by intravenous injection. Chimeras were analyzed at least 7 wk after transplantation. For BrdU labeling, 0.8 mg/ml of BrdU (Sigma-Aldrich) was added to the drinking water and changed daily.

For intrathymic injection, mice were anesthetized with 4% isoflurane in medical air and maintained under anesthesia using a nose cone with 1.5% isoflurane in medical air. Hair was removed from the thorax using depilatory cream, and animals were placed on a heat pad for the injection. Thymus visualization was performed in real time using a 30-MHz 707B ultrasound probe (VisualSonics). The injection needle (Hamilton syringe with 30-gauge needle) was mounted on a 3D micromanipulator and the tip of the needle placed in the imaging plane. 20 µl (10 µl in each lobe) sulfo-NHS biotin (Thermo Fisher Scientific) at a concentration of 5 mg/ml in PBS was injected. For 4-deoxypyridoxine-HeC1 (DOP) treatment, 30 mg/liter DOP (Sigma-Aldrich) and 10 g/liter sucrose (to overcome taste aversion) were added to the drinking water. Control mice received drinking water with 10 g/liter sucrose.

**Cell preparation.** Lymphocytes were isolated by mechanical disruption and filtration through a 70-µm cell strainer and enumerated with a cell counter (Z1 Particle Counter; Beckman Coulter) set to detect nuclei between 3.5 and 7 µm. Thymic stromal cells were isolated as previously described (Gray et al., 2002). In brief, the thymus was diced into small pieces and rotated for 30 min at 4°C in RPMI 1640 supplemented with 10 mM Hepes (RPMI/H). The supernatant containing lymphocytes was removed, the fragments were briefly agitated in ice-cold RPMI/H, the supernatant containing lymphocytes was removed, and the remaining fragments were cultured with a Pasteur pipette and the supernatant was discarded. In the middle and at the end of the second digestion, the fragments were disaggregated with a 22-gauge needle. In the middle and at the end of the third digestion, the fragments were disaggregated with a 26-gauge needle. The supernatants from the final two digestions and remaining fragments were pooled and incubated for 10 min on ice in PBS 1% FBS 5 mM EDTA to yield a mixture enriched in stromal cells. Cells were stained and analyzed using a FACScalibur (BD) or LSRII (BD) or sorted using a MoFlo (Beckman Coulter) or FACSAria (BD). Data were analyzed using FlowJo (v. 8.8.1; Tree Star).
Quantitative RT-PCR. Total RNA was extracted from sorted cell populations using TRIZOL (Invitrogen) according to the manufacturer’s instructions. Before RT, RNA was treated with DNase I (Invitrogen). The RNA was converted to cDNA with the Superscript III First Strand Synthesis System (Invitrogen) or the USB First-Strand cDNA Synthesis kit (Affymetrix) according to the manufacturer’s instructions. To control for DNA contamination, primers were designed to span a large intron when possible, and a reaction without reverse transcription was performed in parallel for each sample/primer pair. To control for nonspecific amplification, the size of the reaction products was analyzed by agarose gel electrophoresis. All primer pairs were tested for linear amplification over two orders of magnitude. Primers used were the following: Hprt sense, 5'-AGGTGTCAGGCTTGCAGTGT-3'; Hprt antisense, 5'-TCGAGTCTATTTGACGAGCA-3'; Sip1 sense, 5'-AGCCCACGATGTCCCTG-3'; Sip1 antisense, 5'-GCGAGATCCATTGTCTGAACGGT-3'; Ppap2b sense, 5'-GGTGGGCCCTGCTCAGATC-3'; Ppap2b antisense, 5'-TCTGTTCTGAATGCCTACCGG-3'; Ppp2r2a sense, 5'-CATACTGGGGTCGTGCT-3'; Ppp2r2a antisense, 5'-TCCTAGCCGTCTTTTAAAT-3'; Ppp2r5 sense, 5'-CCGCTCTGCTGGGATTGC-3'; Sip1 sense, 5'-GGGCGATTATTTGAGTTC-3'; Sip1 antisense, 5'-TTCTATCCGTAGTCTCTAGGC-3'; Sgpp1 sense, 5'-GGGCGCATATTTGTAAGTTTGG-3'; Sgpp1 antisense, 5'-GAGTCCGGAGATCAGGAG-3'; Spgl1 sense, 5'-TTGTGCCACAAATATGAGCC-3'; and Spgl1 antisense, 5'-CTGTGTTGCTGATCTACGTCC-3'.

Antibodies. S1PR1-specific antibody and control were previously described (Lo et al., 2005). Some stainings were done with a second batch of antibodies used were the following: Foxn1 specific antibody, 5'-GAGTCTGAGGCTGATGACAC-3'; EPCR sense, 5'-TTCTATCCGTAGTCTCTAGGC-3'; S1pr1 sense, 5'-GCGGACCGTGAGGACGCCG-3'; Ppap2b sense, 5'-GGTGGGCCCTGCTCAGATC-3'; Ppap2b antisense, 5'-TCTGTTCTGAATGCCTACCGG-3'; Ppp2r2a sense, 5'-CATACTGGGGTCGTGCT-3'; Ppp2r2a antisense, 5'-TCCTAGCCGTCTTTTAAAT-3'; Ppp2r5 sense, 5'-CCGCTCTGCTGGGATTGC-3'; Sip1 sense, 5'-GGGCGATTATTTGAGTTC-3'; Sip1 antisense, 5'-TTCTATCCGTAGTCTCTAGGC-3'; Sgpp1 sense, 5'-GGGCGCATATTTGTAAGTTTGG-3'; Sgpp1 antisense, 5'-GAGTCCGGAGATCAGGAG-3'; Spgl1 sense, 5'-TTGTGCCACAAATATGAGCC-3'; and Spgl1 antisense, 5'-CTGTGTTGCTGATCTACGTCC-3'.

Flow cytometry and confocal imaging. Flow cytometry was done using standard conditions. All staining was performed on ice, except for CCR7 staining which was at 37°C. Staining for S1PR1 was done with 0.05% sodium azide in all buffers. Confocal imaging was done using standard conditions. In brief, the thymus was excised, fixed overnight at 4°C in periodate-lysine-paraformaldehyde medium (PLP; 0.05 M phosphate buffer, pH 7.4, 0.1 M L-Lysine, 10 mg/ml paraformaldehyde, and 2 mg/ml NaIO4) and progressively dehydrated at 4°C in sucrose (10, 20, and 30% in PBS). Tissues were snap frozen in OCT compound (Tissue-Tek; Sakura). 10-µm-thick tissue sections were rehydrated (rehydration and staining buffer was PBS with 4% mouse serum, 4% rat serum, 1% Triton X-100, and 0.1 M Tris), stained, and imaged using a confocal microscope (LSM510; Carl Zeiss) with a 40× oil immersion objective. Images were processed using ImageJ software (version 1.41; National Institutes of Health).

Mass spectrometry. Before taking thymus, mice were perfused with 10 ml PBS. Samples were snap frozen in liquid nitrogen and stored at −80°C. Lipids were purified and mass spectrometry was performed as previously described (López-Juárez et al., 2011).

Online supplemental material. Fig. S1 shows the effect of LPP3 deletion on the numbers of DN and DP T cells in the thymus, that LPP3 deletion is associated with reduced numbers of CD4 T cells in LNs and spleen, and that LPP3 deletion inhibits egress of mature CD8 T cells from the thymus. Fig. S2 shows that LPP3 deletion does not detectably alter Sip1 transcript in mature thymocytes or surface CCR7 expression on mature thymocytes, and that an increase in total thymic SIP is not detectable by mass spectometry. Fig. S3 shows that chimeras generated with LPP3-deficient BM do not have elevated thymic SIP and that inhibition of SIP lyase further increases thymic SIP in LPP3-deficient mice. Fig. S4 shows expression of LPP3 and the pericyte marker PDGFR-β by confocal microscopy. Fig. S5 shows specificity of deletion by Cdhs5 (PAC)-CreERT2 and a reduction in the number of T cells in the periphery of Ppap2b+ Cdhs5 (PAC)-CreERT2 mice. Fig. S6 shows specificity of deletion by Krt14-Cre, a reduction in the number of T cells in the periphery of Ppap2b+ Krt14-Cre+ mice, and an accumulation of mature T cells in the thymus of Ppap2b−/− Foxn1−/− mice. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20102551/DC1.

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