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Single-Cell RNA Sequencing of Lymph Node Stromal Cells Reveals Niche-Associated Heterogeneity

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SUMMARY

Stromal cells (SCs) establish the compartmentalization of lymphoid tissues critical to the immune response. However, the full diversity of lymph node (LN) SCs remains undefined. Using dropletbased single-cell RNA sequencing, we identified nine peripheral LN non-endothelial SC clusters. Included are the established subsets, Ccl19^{hi} T-zone reticular cells (TRCs), marginal reticular cells, follicular dendritic cells (FDCs), and perivascular cells. We also identified Ccl19^{lo} TRCs, likely including cholesterol-25-hydroxylase⁺ cells located at the T-zone perimeter, Cxcl9⁺ TRCs in the T-zone and interfollicular region, CD34⁺ SCs in the capsule and medullary vessel adventitia, indolethylamine N-methyltransferase⁺ SCs in the medullary cords, and Nr4a1⁺ SCs in several niches. These data help define how transcriptionally distinct LN SCs support niche-restricted immune functions and provide evidence that many SCs are in an activated state.

Graphical abstract

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AUTHOR CONTRIBUTIONS

Conceptualization, L.B.R. and J.G.C.; Investigation, L.B.R., E.L., C.L.S., and X.W.; Software, L.B.R.; Formal Analysis, L.B.R.; Resources and protocols, M.L.B., B.A.B., A.D.L., S.A.L., and J.G.C.; Writing, L.B.R. and J.G.C.; Supervision, C.J.Y. and J.G.C.; Funding Acquisition, J.G.C.

SUPPLEMENTAL INFORMATION

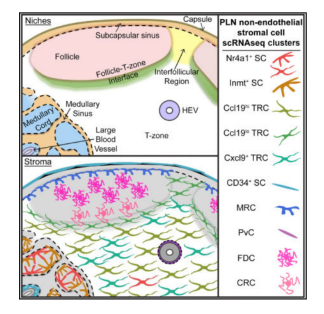
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DECLARATION OF INTERESTS

The authors declare no competing interests.

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In Brief: Lymph node stromal cells support diverse processes, but bulk assessments obscure their niche-specific functions. Rodda et al. identify transcriptional profiles for nine lymph node stromal cell clusters using single-cell RNA sequencing, validate subset markers *in situ*, and suggest niche-restricted functions.



INTRODUCTION

Lymph nodes (LNs) are organized into discrete niches to support the efficient antigen encounter and lymphocyte activation required for an effective adaptive immune response (Mueller and Germain, 2009). The stromal cells (SCs) that pattern these niches in reticular networks contribute to the structural and functional specificity of the niche by influencing lymphocyte migration, survival, antigen encounter, and tolerance (Chang and Turley, 2015; Schulz et al., 2016).

Follicular SCs, including CR2⁺ antigen-trapping follicular dendritic cells (FDCs) and TNFSF11⁺ marginal reticular cells (MRCs), express the chemokine CXCL13 and guide B cells to LN follicles. FDCs are positioned at the follicle center and the MRCs sit at the edge of the follicle adjacent to the subcapsular sinus (SCS) (Cyster et al., 2000; Katakai et al., 2008). During an immune response, light zone (LZ) FDCs and dark zone (DZ) CXCL12-expressing reticular cells (CRCs) emerge and support the respective germinal center (GC) niches (Bannard et al., 2013; Rodda et al., 2015). The transcriptional programs that control the specialized functions of follicular SCs are unknown.

CCL21- and CCL19-expressing SCs in the T-zone, termed T-zone reticular cells (TRCs), attract CCR7⁺ lymphocytes (Cyster, 2005). TRCs also produce trophic factors such as interleukin-7 (IL-7) and maintain the extracellular matrix (ECM)-rich conduit network that allows rapid diffusion of small molecules throughout the LN (Link et al., 2007; Mueller and Germain, 2009). During an immune response, activated B and T cells meet at the follicle-T-

zone interface, but it is unclear whether SCs in this region differ from TRCs (Katakai et al., 2008).

Lymphocytes enter LNs through specialized blood vessels called high endothelial venules (HEVs). Many blood vessels in LNs are associated with ITGA7⁺ perivascular cells (PvCs) (Girard et al., 2012). CD34⁺ adventitial cells (ACs) may further support large vessels in the LN medulla (Díaz-Flores et al., 2014). Lymphocytes exit the LN through the medulla, where lymphatic sinuses weave around medullary cords. The cords are populated by medullary reticular cells (MedRCs), macrophages, and plasma cells. MedRCs highly express CXCL12 (Hargreaves et al., 2001), but little else is known about them.

The Immgen consortium has reported gene expression profiles for two SC and two endothelial cell subsets in LNs: PDPN⁺CD31⁻ fibroblastic reticular cells (FRCs), PDPN⁻CD31⁻ double-negative cells (DNCs), PDPN⁺CD31⁺ lymphatic endothelial cells (LECs), and PDPN⁻ CD31⁺ blood endothelial cells (BECs) (Malhotra et al., 2012). While this work provides important insight into the possible contributions of SCs to LN function, niche-restricted stromal roles remain obscured because TRCs, MRCs, FDCs, and MedRCs all fall into the FRC subset.

To investigate the heterogeneity of LN SCs, we analyzed single-cell RNA sequencing (scRNAseq) of peripheral LN (pLN) non-endothelial SCs. We identified nine clusters of SCs previously partitioned into the FRC and DNC populations. We associated eight clusters with specific niches and validated markers identified in our dataset for MedRCs and FDCs.

RESULTS

Single-Cell RNA Sequencing Reveals Nine pLN Non-endothelial SC Clusters

We performed droplet-based scRNAseq on isolated mouse pLN CD45⁻CD31⁻ cells from uninfected mice and mice at day 15 post-infection with LCMV-Armstrong (Figure 1A). After quality control, we retained 2,870 cells with 2,153 median genes per cell from the uninfected LNs and 12,669 cells with 1,938 median genes per cell from the post-infection LNs. After initial unsupervised clustering with Seurat (Satija et al., 2015) to select the >95.1% of non-endothelial SCs in *Pdgfrb*⁺ and/or *Pdgfra*⁺ clusters for further analysis, we combined the samples to investigate conserved heterogeneity.

We chose to analyze pLN SCs on day 15 after infection with LCMV-Armstrong because the virus induces large GCs with LZ FDCs and DZ CRCs. Although SCs are diminished in the early days after LCMV infection, we decided it was appropriate to combine the samples because by day 15 the virus has been cleared, the percentage of CD45⁻VCAM1⁺PDPN⁺ SCs has almost recovered, the B and T zone organization is normal, and the expression of *Cxcl13, Ccl19*, and *Ccl21* by the post-infection SCs is within 3-fold of the uninfected SCs (Figure S1A; Mueller et al., 2007; Rodda et al., 2015; Scandella et al., 2008). The mean expression of variable genes by cells in the two samples was also highly correlated (r = 0.99, p value < 2.2×10^{-16} , Pearson) (Figure S1B).

Employing diagonal canonical correlation analysis (CCA) to combine the samples, we identified a shared gene correlation structure that explained more than 50% of the variance of 94.1% and 96.8% of cells from the uninfected and post-infection samples, respectively (Butler and Satija, 2017). The 14,243 cells adequately described by the structure were then aligned for clustering analysis. The removed cells came from both samples and 12.2% of the removed cells were enriched for cell cycle gene expression (cell cycle score > 0.1) compared to 0.39% of the remaining cells. Of the removed cycling cells, 98.5% were from the post-infection sample (Figure S1C).

Performing unsupervised clustering on the combined samples revealed nine conserved clusters, which we visualized with tSNE and a hierarchical cluster tree (Figures 1B and S1D). We chose this clustering resolution because higher resolutions did not reveal linearly increasing numbers of clusters. Cells from both samples contributed to each cluster, suggesting that the clustering was not due to sample batch effect (Figure S1E). In addition, the mean expression profile for each cluster was highly correlated between the two samples (r > 0.97, p value $< 2.2 \times 10^{-16}$, Pearson) and the cluster composition of each sample was similar (Figure 1C). We have included the mean gene expression profiles for each cluster for the 16,775 detected genes (Table S1).

We identified differentially expressed genes (DEGs) (log2-fold change > 0.5, FDR < 0.05, proportion of cluster expressing > 0.10) for each cluster calculated as the difference between the average expression by cells in the cluster and the average expression by cells not in the cluster (Tables S2, S3, S4, S5, S6, S7, and S8). While the clusters could represent distinct cell types or cell states, the DEGs suggested assignment of several of the clusters to the previously described niche-associated SC types Ccl21⁺ TRCs, Tnfsf11⁺ MRCs, Itga7⁺ $Acta2^+$ PvCs, and $Cr2^+$ FDCs (Figure 1D). The proportions of these cells corresponded to expectations based on sectioning and flow cytometric studies (Figure 1C; Jarjour et al., 2014; Link et al., 2007). The remaining clusters represented unappreciated heterogeneity and were distinguished as Inmt⁺Cd34⁻Tnfsf11⁻ SCs (Inmt⁺ SCs), Cd34⁺ SCs (CD34⁺ SCs), *Nr4a1*⁺ SCs (Nr4a1⁺ SCs), and three subsets of TRCs: *Ccl21*⁺*Ccl19*th*II7*⁺ TRCs (Ccl19th TRCs), Ccl21+Ccl19oII7+ TRCs (Ccl19lo TRCs), and Ccl21+Ccl19oII7+ Cxcl9+ TRCs (Cxcl9⁺ TRCs) (Figure 1D). Notably, historically used pLN non-endothelial SC markers, such as Pdpn, Pdgfra, and Pdgfrb, were insufficient to distinguish the clusters (Figure 2A). Since each cluster had at least 50 DEGs (Figure 2B), we have highlighted 10 marker candidates per subset using stricter thresholds (log2-fold change > 1, FDR < 0.05, proportion of cluster expressing > 0.20) (Figures 2C and S2). We proceeded to investigate the distinctive features of each pLN non-endothelial SC cluster.

Ccl19^{hi} TRC DEGs Suggest Additional Mechanisms for Lymphocyte Support and Conduit Maintenance

TRCs support the LN T-zone and have been among the most in-depth studied LN SCs (Mueller and Germain, 2009). The Ccl19^{hi} TRC cluster differentially expressed genes previously associated with TRCs as well as unappreciated genes for surface and secreted molecules (Figure S3A, Table S2). While Ccl19^{hi} TRCs expressed *Pdpn* shown to support DC motility and LN expansion (Acton et al., 2014; Astarita et al., 2015), they also distinctly

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expressed *Cx3c11*, the chemokine ligand for CX3CR1⁺ activated CD8⁺ T cells, monocytes, and recently described T-zone macrophages (Figures 2A and S3A; Baratin et al., 2017). CX3CL1 has been detected in the LN T-zone, SCS, and interfollicular regions (IFR), but has not been attributed to TRCs (Johnson and Jackson, 2013; Kanazawa et al., 1999). Ccl19^{hi} TRCs were also enriched for expression of *II13ra2*, an IL-13 decoy receptor that could limit T-zone IL-13 (Fichtner-Feigl et al., 2006), and *II4i1*, an L-amino acid oxidase that can inhibit the CD8⁺ T cell anti-tumor response *in vivo* (Figure S3A; Lasoudris et al., 2011).

TRCs make critical components of the reticular conduit network and themselves ensheath the conduits (Mueller and Germain, 2009). Highlighted previously as expressed by FRCs, Ccl19^{hi} TRCs were enriched for collagen 14 (*Col14a1*) and fibromodulin (*Fmod*), components of the conduit core (Figure S3A; Malhotra et al., 2012). Ccl19^{hi} TRCs expressed desmocolin-2 (*Dsc2*), a cadherin-type protein that links adjacent cells via desmosomes (Figure S3A; Jarjour et al., 2014). A portion of Ccl19^{hi} TRCs expressed the latrophilin adhesion-type GPCR *Adgrl3* and one of its ligands, *Flrt3*, which may support adhesion between Ccl19^{hi} TRCs (Figure S3A, Table S2; O'Sullivan et al., 2012).

Ccl19^{hi} TRC DEGs suggest mechanisms of Ccl19^{hi} TRC development and maintenance. TNF family molecules have strong influences on TRCs (Cyster, 2005) and Ccl19^{hi} TRCs expressed *Nradd* and shared *Relt* and *Cd200* expression with FDCs (Figure S3A). Ccl19^{hi} TRCs distinctly expressed *Fabp7*, a transport protein that may deliver lipids to nuclear hormone receptors and has been shown to be restricted to LN TRCs *in situ* (Figure S3A; Tokuda et al., 2010). Finally, Ccl19^{hi} TRCs were enriched for expression of the transcription factors *SpiC*, *Tead2*, and *Yeats4* among others (Figure S3A). The Ccl19^{hi} TRC DEGs support this subset fulfilling the known TRC roles in chemotaxis to the T-zone, trophic support for T cells and DCs and building the conduit matrix while suggesting additional mechanisms to achieve these functions.

Ch25h-Expresssing Ccl19^{lo} TRCs Populate the Follicle T-Zone Interface and IFRs

Ccl19^{lo} TRCs expressed *Ccl21*, *II7*, and *Cxcl13*, features of both the T cell zone and B cell follicle (Figures 2A and S3A). In addition, relative to Ccl19^{hi} TRCs, they expressed high *Tnfsf13b* (Baff) (Figure 3A), a cytokine critical for B cell survival and non-redundantly produced by non-FDC FRCs (Wang et al., 2012; Cremasco et al., 2014), suggesting that Ccl19^{lo} TRCs might occupy a niche that is engaged by migrating B cells such as the follicle T-zone interface.

Cholesterol-25-hydroxylase (*Ch25h*) was expressed by a portion of Ccl19^{lo} TRCs (log2-fold change = 0.68, FDR = 2.4×10^{-4} , proportion of cluster expressing = 0.21) along with MRCs (log2-fold change = 2.83, FDR = 1.99×10^{-51} , proportion of cluster expressing = 0.51) (Figure 3A, Table S2). Ccl19^{lo} TRCs are not highlighted in the violin plot because the visualization has a higher cutoff for proportion of a cluster expressing a gene than our DEG cutoff. Stromal expression of this enzyme is important for generation of the EBI2 (GPR183) ligand 7a,25-HC to guide activated B cells, T cells, and DCs to the follicle T-zone interface and IFRs (Cyster et al., 2014; Li et al., 2016; Lu et al., 2017). Using RNAscope on pLNs and mesenteric LNs (mLNs), *Ch25h* was detected highly expressed by cells at the follicle-T-zone interface and in IFRs, as has been shown in spleen (Figures 3B and S3B; Lu et al.,

2017). *Ch25h* expression was also found adjacent to the SCS in accordance with MRC expression and in ring-like structures in the T-zone in accordance with HEV expression (Figures 3B and S3B; Lee et al., 2014). *Ch25h*+Ccl19^{lo} TRCs likely correspond to the *Ch25h*-expressing cells positioned at the follicle-T-zone interface and in IFRs, functioning to guide EBI2⁺ cells to these niches. While the low percentage of Ccl19^{lo} TRCs with detectable *Ch25h* expression could reflect low expression of the enzyme, it likely indicates that Ccl19^{lo} TRCs include *Ch25h*⁻ cells that support other T-zone-adjacent niches. Determining whether this is the case will require additional marker assessment.

Cxcl9⁺ TRCs Are Likely Activated TRCs in the T-Zone and IFRs

While Cxcl9⁺ TRCs expressed *Ccl21* and *II*7 and differentially expressed *Ccl19* suggestive of TRCs (Figures 2A and S3A), they were most distinguished by *Cxcl9* and *Cxcl10*, the ligands for CXCR3, and MHCII-related gene expression (Figure 4A, Table S2). To validate pLN stromal *Cxcl9* and *Cxcl10* expression and localize Cxcl9⁺ TRCs in pLNs, we analyzed pLN stroma from REX3 mice, which express *Cxcl9*-driven RFP and *Cxcl10*-driven BFP (Groom et al., 2012). By flow cytometry (FC), portions of each classical subset (FRC, DNC, BEC, LEC) expressed the chemokines and 35.5% (SEM = 0.69%) of non-endothelial SCs (FRC and DNC) expressed both CXCL9 and CXCL10 (Figure 4B). While this is higher than the 10.6% of *Cxcl9*- and *Cxcl10*-expressing cells in our scRNAseq dataset (Table S2), the difference may reflect an underestimate due to the low mRNA capture per cell in scRNAseq.

In REX3 pLN sections, immunofluorescence microscopy (IFM) revealed CXCL9⁺CXCL10⁺PDGFR β^+ CD11c⁻ stroma in the T-zone and IFRs interspersed with CXCL9⁻CXCL10⁻PDGFR β^+ CD11c⁻ stroma (Figure 4C). CXCL9⁻CXCL10⁺PDGFR β^+ CD11c⁻ stroma was also detected in the medullary cords in accord with one-third of nonendothelial SCs expressing CXCL10 alone by FC and our scRNAseq analysis (Figures 4B and 4C). These are likely MedRCs that correspond to the *Cxcl10*-expressing Inmt⁺ SC subcluster of Nr4a1⁺Inmt⁺ SCs discussed in more detail in a following section (Figure 4A). Consistent with these findings, CXCL9 and CXCL10 expression by radio-resistant cells has been reported in REX3 reverse BM chimera popliteal LN T-zone, IFRs, and medulla (Groom et al., 2012).

The Cxcl9⁺ TRC DEGs were dominated by interferon (IFN)-inducible genes such as *Gbp4* and *Gbp5* (Figure 2C). The Cxcl9⁺ TRCs also expressed the type 1 IFN and IFN- γ receptors *Ifnar2, Ifngr1*, and *Ifngr2*, though not exclusively (Table S2). Prior work has shown that during an immune response, IFN- γ -induced CXCL9 and type 1 IFN-induced CXCL10 from myeloid cells and SCs can position CD4⁺ T cells, CD8⁺ memory T cells, and DCs in IFRs and support antigen encounter (Groom et al., 2012; Kastenmüller et al., 2013; Sung et al., 2012). Further study is required to determine whether the Cxcl9⁺ TRCs are an IFN-inducible state of Ccl19^{hi} TRCs or a specialized, stable subset.

Cxcl9⁺ TRCs could also represent the FRCs reported to induce CD4⁺ T cell tolerance through presentation of peptide-MHC (Brown and Turley, 2015) since their DEGs included several MHCII genes (*H2-Aa, H2-Eb1*, and *H2-Ab1*), the invariant chain (*Cd74*), and peptide-loading chaperone (*H2-DMa*), but not co-stimulatory or inhibitory molecules (*Cd80*, *Icosl, Cd274, Tnfsf9, Cd40, Tnfrsf14, Tnfsf18, Tnfsf14, Havcr1, Havcr2, Tnfsf15, Tnfsf8*,

Vsir, *Csf1r*, *Cd86*, *Pdcd11g2*, *Ctla4*, *Btla*, *Tnfrsf4*, *Cd70*, *Cd48*, or *Slamf1*) (Tables S1 and S2). Cxcl9⁺ TRCs may be an activated subset of Ccl19^{hi} TRCs that can tolerize T cells in a resting LN and help position CXCR3⁺ cells in IFRs during an immune response.

MRC Gene Expression Suggests Involvement in Barrier Defense

MRCs are situated at the base of the SCS, where lymph drains antigen from peripheral tissues, and have been described as PDPN

+TNFSF11+MADCAM1+VCAM1+ICAM1+BST1+RELB+CXCL13+ LN stroma (Katakai et al., 2008). One cluster expressed almost all of these markers and was enriched for *Tnfsf11* and *Cxc113* expression, suggesting that the constituent cells are MRCs (Figure 5A, Tables S1 and S3). However, this subset was not enriched for *Madcam1* expression. The basis for this discrepancy is unclear, but MRCs may more efficiently translate *Madcam1* mRNA or maintain the stability of the protein better than the TRCs which had detectable expression in our dataset. Alternatively, *Madcam1* may be expressed by cells closely associated with MRCs, such as LECs (Cordeiro et al., 2016) (Immgen.com). In this regard it is notable that *Madcam1* expression is LTβR signaling dependent while *Tnfsf11* expression is not (van de Pavert and Mebius, 2010).

Little is understood about the role of MRCs in the SCS, a niche critical for antigen capture. The MRC subset had enriched expression of *Enpp2* (autotaxin) (Figure 5B), a secreted enzyme required for the production of lysophosphatidic acid (LPA) that has been linked to T cell high-speed motility (Katakai et al., 2014). This may allow MRCs to influence lymphocyte-antigen interaction in the SCS. In pLN sections, IFM revealed that ENPP2 colocalized with TNFSF11 on the follicle edge of the SCS, consistent with MRC expression. ENPP2 was also detected to a lesser extent in the T-zone and HEVs (Figure 5C) in accord with expression by several other stromal subsets (Figure 5B) and with previous work (Kanda et al., 2008; Katakai et al., 2014).

MRCs may also interact with cells in the SCS to influence the initiation of the immune response and possibly prevent pathogen spread. Sensory nerves containing the neurotransmitter calcitonin gene-related peptide (CGRP) are abundant in lymphoid organs and nerve fibers have been reported in the LN SCS (Iannacone et al., 2010). The MRC subset was enriched for *Ramp1* expression, which directs CALCRL (also expressed) to form the receptor for CGRP (Table S3; McLatchie et al., 1998). The MRCs also expressed embigin (*Emb*), which has roles in nerve terminal sprouting (Table S3; Lain et al., 2009). MRCs may also influence cKit⁺ DCs, BECs, lymphoid tissue inducer cells, and neuronal precursors through expression of the cytokine *Kitl*, which we found differentially expressed (Table S3; Chappaz et al., 2010; Goldstein et al., 2015). Finally, the most differentially expressed gene for the MRC subset is hepcidin antimicrobial peptide 2 (*Hamp2*), which could suggest a direct role in barrier defense (Figure 2C).

Itga7⁺ PvC DEGs Suggest Roles in Endothelial Support and Leukocyte pLN Entry

Non-adventitial PvCs support multiple functions of blood vessels and have been found around some vessels in the pLN cortex and medulla (Chang and Turley, 2015). Non-adventitial PvCs include pericytes and vascular smooth muscle cells, which can be

distinguished *in situ* only by their location surrounding capillaries and large vessels, respectively (Armulik et al., 2011). One particularly transcriptionally distinct stromal cluster in our dataset expresses the hallmarks of non-adventitial PvCs including *Pdgfrb*, *Acta2*, *Itgb1*, *Esam1*, *My19*, *Mcam*, and *Notch3* (Figure S4, Table S4; Armulik et al., 2011; Malhotra et al., 2012). This subset is similar to the *Itga7*⁺ PvCs described as making up more than 50% of isolated LN DNCs (Malhotra et al., 2012). Our analysis did not, however, detect a transcriptionally distinct *Cnn1*⁺*Itga7*⁻*Pdpn*⁻PvC subset as proposed and the canonical PvC gene *Cspg4* was detected in only a few cells, likely reflecting low expression and the transcript capture limitations of scRNAseq (Table S1; Malhotra et al., 2012).

The PvC subset was enriched for expression of thrombospondin (*Thbs1*), *II34*, and endothelin-A receptor (*Ednra*), a gene involved in crosstalk between tissue cells and PvCs in kidneys (Figure S4; Kitazawa et al., 2011). PvCs also distinctly expressed ECM-adhesion factors *Bcam* and *Nexn* and several transcription factors including *Mef2c* and *Fhl2* (Figure S4). The PvCs expressed a constellation of genes that may aid interactions at the vascular portals to the pLN parenchyma.

CD34⁺ SCs Express Features of ACs and Are Positioned in the Capsule and Medulla

CD34⁺ SCs were distinguished by expression of the sialomucin *CD34*, which on CD31⁺ LN BECs can bind L-selectin on passing lymphocytes (Figure 6A; Baumheter et al., 1993). Recent work characterized PDPN⁺BST1⁻ACTA2⁻CD31⁻CD34⁺ SCs surrounding large vessels in the LN medullary cords as ACs based on shared gene expression with ACs in other tissues (Corselli et al., 2012; Sitnik et al., 2016). In accord with CD34⁺ SCs being ACs, the subset expressed *Pdpn* but had no detectable expression of *Bst1*, *Acta2*, or *Cd31* (Table S1). In addition, LN ACs have been described to secrete vasculogenic and angiogenic factors and CD34⁺ SCs expressed *Igf1*, *Igfbp3*, *Igfbp4*, and exclusively *Igfbp6* along with growth factors and ECM factors (Figure 2C, Tables S1 and S5; Sitnik et al., 2016).

IFM on pLNs revealed CD34⁺CD31⁻ cells surrounding blood vessels in the medulla near the efferent lymphatics, in keeping with these being ACs (Figure 6B). However, CD34⁺CD31⁻ cells that were PDGFR β^+ BST1⁻ were most prominently detected in the capsule (Figure 6C). CD34⁺ SCs were the only stromal subset with low desmin (*Des*) expression and DES was not detected on the CD34⁺ cells in either niche (Figures 6A–6C).

CD34⁺ SCs positioning in the pLN capsule is in accord with a study of human LNs showing CD34⁺ cells in the capsule as well as around large vessels (Díaz-Flores et al., 2014). CD34⁺ SCs were also enriched for endosialin (*Cd248*) expression, which is reported in the resting LN capsule and required for LN expansion after immunization (Table S5; Lax et al., 2010). These capsular cells were likely not LECs lining the SCS as LECs do not express *Cd34* (Immgen.org) and expression of the LEC markers *Lyve1*, *Prox1*, and *Ackr4* (Ulvmar et al., 2014) was not detected in CD34⁺ SCs (Table S1). CD34⁺ cells may, however, communicate with LECs as LECs highly express the secreted ligand for CD248, MMRN2 (Immgen.org) (Khan et al., 2017). CD34⁺ SCs could play a role in capsule integrity as the subset was enriched for expression of the ECM component and assembly associated genes *Col5a3*, *Col15a1, Lama2, Vit, Lox11, Mfap5*, and *Fbln1*. This subset was distinctly enriched for the

expression of the transcription factors Peg3 and Ar (Table S5). These data suggest that CD34⁺ SCs include ACs and capsular fibroblasts.

Inmt⁺ SCs Populate Medullary Cords and Likely Include MedRCs

The Inmt⁺ SC cluster expressed intermediate *Cxcl13* and low *Ccl19* and did not distinctly express any known niche- or subset-associated markers (Figure 2A). Instead, these cells differentially expressed indolethylamine N-methyltransferase (*Inmt*), an enzyme that can catalyze the N-methylation of tryptamine and could play a role in cell-cell communication (Mavlyutov et al., 2012), and stanniocalcin-1 (*Stc1*), a LIM-domain protein that may have a role in heterchromatin stability (Figures 2C and 6D, Table S6; He et al., 2013).

To determine whether Inmt⁺ SCs are associated with a specific LN niche, we used RNAscope and detected *Inmt* expression in pLN and mLN medullary cords, identified by DES⁺ ECM patterning and F4/80⁺ macrophages (Figures 6E and S5A). The *Inmt* signal location, density, and pattern is suggestive of MedRCs. An additional distinguishing feature of the Inmt⁺ SCs was low expression of *Bst1* relative to the FDC, MRC, and TRC subsets (Figure 6D). In IFM on pLN sections, BST1 was detected throughout the cortical stroma (FDCs, MRCs, TRCs) but was absent from the MedRCs in accord with previous work (Figure 6F; Link et al., 2007). While *Inmt* was also expressed in other stromal clusters (CD34⁺ SCs and MRCs) and other cell types do populate the medullary cords (B cells, memory T cells, and plasma cells), these cell types are either not found in the medulla with this frequency or do not express *Inmt* (Figure 6D; Gray and Cyster, 2012; Immgen.org). A portion of the Nr4a1⁺ SCs also expressed *Inmt* and are discussed in the next section. Further analysis will be needed to determine whether medullary *Inmt* expression also involves medullary macrophages, which have no available expression data. This differential expression profile will enable investigation of the Inmt⁺ SC function in the medullary cords.

Nr4a1⁺ SCs Likely Consist of Activated Cells from Other Subsets

The Nr4a1⁺ SC cluster could not be readily assigned to a known stromal subset or specific niche. Instead, this cluster was distinguished by expression of early-response genes including *Nr4a1* (*Nur77*), *Fosb, Fos, Junb, Egr1, Nfkbia* (*Ikba*), *Nfkbiz*, and *Zfp36* (Figures 2C and 6G, Table S7; Ullman et al., 1990). To validate that these SCs are present in uninfected pLNs, we isolated pLN stroma from *Nr4a1*-GFP transcriptional reporter mice (Zikherman et al., 2012). Indeed, a portion of FRCs, DNCs, BECs, and LECs expressed *Nr4a1*-GFP greater than the GFP⁻ control, with FRCs and BECs having the highest *Nr4a1*-GFP expression (Figure 6H).

Early-response genes can be activated by TLR, cytokine, mitogen, or growth factor signaling as well as physical stimuli (Ullman et al., 1990). To investigate whether Nr4a1⁺ SCs were activated cells from other subsets, we subclustered the Nr4a1⁺ SCs and compared the subcluster DEGs to those from the other original clusters (Table S7). We distinguished three subclusters with more than 300 DEGs each (log2-fold change > 0.5, FDR < 0.05, proportion of cluster expressing > 0.10) that significantly correlated with Inmt⁺ SCs (r = 0.49, FDR = 8.9×10^{-31} , Pearson; 84.4% of Nr4a1⁺ SCs), Ccl19^{hi} TRCs (r = 0.78, FDR = 1.5×10^{-90} , Pearson; 13.1% of Nr4a1⁺ SCs), and CD34⁺ SCs (r = 0.44, FDR = 7.7×10^{-17} , Pearson;

2.6% of Nr4a1⁺ SCs) (Figures S5B and S5C, Table S7). Cells from both samples contributed to each subcluster (Figure S5D).

To test whether Nr4a1⁺ SCs populate a specific niche or, as our subclustering would suggest, are found in multiple niches, we stained PDGFR β^+ stroma in *Nr4a1*-GFP⁺ pLN sections. PDGFR β^+ *Nr4a1*-GFP⁺ cells were found frequently in the medullary cords and occasionally in IFRs, the follicle-T-zone interface, the T-zone, and the SCS in accord with expression by Inmt⁺ SCs, TRCs, and MRCs (Figure 6I). Medullary CD34⁺ SCs and PDGFR β^- FDCs could not be distinguished with this stain. The medulla is a site of high lymph flow and macrophage-mediated clearance of antigens. Further study is required to determine whether Nr4a1⁺ Inmt⁺ SCs in the medullary cords are activated by lymph-borne material.

FDCs Distinctly Express Pthlh and Tmem119 and Share Sox9 Expression with CRCs

LN FDCs have been difficult to isolate in sufficient numbers for transcriptional profiling because they are rare and FDC markers are lost during tissue digestion. However, with scRNAseq we identified a small cluster highly enriched for well-defined FDC-expressed genes including complement receptor 2 (*Cr2*), Fc receptors (*Fcgr2b* and *Fcer2a*), *Coch* (Py et al., 2013), *Prnp, Madcam1, Vcam1, Cxc113*, and *Mfge8* (Figures 2A and 2C, Table S8; Allen and Cyster, 2008). Matching published estimates for FDCs, this cluster made up 1.7% of the recovered pLN non-endothelial SCs (Figure 1C; Jarjour et al., 2014). The FDCs were the most transcriptionally distinct subset, possibly reflecting their unique dendritic morphology and specialized function in antigen presentation (Figure 2B; Heesters et al., 2014).

The FDC cluster distinctly expressed parathyroid hormone like hormone (*Pthlh* or *Pthrp*), a secreted protein that could be involved in development or maintenance of the follicle or GC (Figure 7A). Initially described as an inducer of hypercalcemia of malignancy, *Pthlh* has since been established to regulate the differentiation of a number of cell types (McCauley and Martin, 2012). RNAscope analysis revealed *Pthlh* expression in the center of pLN primary follicles and in the LZs of mLN GCs in accord with expression by the CR2⁺ FDCs detected in a sequential stain (Figure 7B). FDCs depend on LT and TNF signaling for their maintenance (Allen and Cyster, 2008). Four-day *in vivo* blockade of these cytokines with LTβR-Fc and TNFR1-Fc led to a loss of *Pthlh* signal, consistent with expression by FDCs (Figure 7C). *Pthlh* is thought to largely act through *Pth1r*, though some receptor-independent actions have been proposed (McCauley and Martin, 2012). Of all the SC clusters, only Ccl19^{hi} TRCs expressed *Pth1r* consistent with previous detection in bulk FRCs (Immgen.org) (Figure 7A). It will be of interest to determine whether this factor functions in cross-talk between FDCs and Ccl19^{hi} TRCs.

The FDC cluster was also enriched for *Sox9* expression, a transcription factor that could play a role in FDC differentiation (Figure 7D). SOX9 has roles in the differentiation of multiple cell types including chondrocytes and neural crest cells (Huang et al., 2015). In pLN follicles from irradiated *Cr2*-cre R26-ZsGreen reporter mice (reconstituted with reporter negative hematopoietic cells), SOX9 co-stained ZsGreen⁺ CR2⁺ FDC cell bodies (Figure 7E). SOX9 also stained within GC LZ CR2⁺ FDC networks (Figure 7F). Consistent

with FDC SOX9 expression, the SOX9 detection was lost along with CR2⁺ FDCs after short-term LT and TNF blockade (Figure 7F).

GCs also contain CXCL12⁺CR2^{lo} DZ CRCs, SCs with a frequency and morphology similar to FDCs, but which guide CXCR4⁺ GC B cells to the DZ (Bannard et al., 2013; Rodda et al., 2015). To probe whether SOX9 expression is unique to FDCs or a shared feature of GC stroma, we stained SOX9 in pLNs from immunized *Cxcl12*-GFP reporter mice and detected expression in both CR2^{hi} *Cxcl12*-GFP⁻ LZ FDCs and CR2^{lo} *Cxcl12*-GFP⁺ DZ CRCs (Figure 7G).

While we expect LZ FDCs and DZ CRCs to have distinct transcriptional programs, their similar morphology and shared role in GCs suggests that they may be at least more similar to each other than to other SCs. Thus we attempted to identify distinguishing expression by subclustering the FDCs from the post-infection sample predicted to contain GC stroma (Rodda et al., 2015). However, the subclusters did not pass our significance threshold (more than 50 DEGs relative to the other subclusters and original clusters). Still, among post-infection FDCs, *Cxcl12* expression was negatively correlated with canonical FDC gene (*Fcgr2b* and *Fcer2a*) and *Sox8* expression (Figure S6A). *Sox8* is a transcription factor in the same SoxE family as *Sox9* (Huang et al., 2015). Notably, *Sox9* was less negatively correlated with *Cxcl12* (Figure S6A). More cells or increased depth of mRNA capture are required to determine whether the heterogeneity in the post-infection FDCs reflects the inclusion of DZ CRCs or mRNA sampling variability and whether *Sox8*-directed expression contributes to the distinct phenotypes of LZ FDCs and DZ CRCs.

The FDC cluster also differentially expressed *Tmem119*, a surface protein that could facilitate FDC-lymphocyte interactions in the follicle and GC. *Tmem119* encodes an O-glycosylated surface protein found on microglia and involved in osteoblast development (Figure 7H; Bennett et al., 2016; Kanamoto et al., 2009). Using IFM on pLN primary follicles, TMEM119 selectively labeled CR2⁺ FDC networks (Figure 7I). In addition, short-term blockade of LT and TNF led to a loss of CR2 and TMEM119 staining in accord with FDC expression (Figure 7J). To test for a role of FDC TMEM119 in support of the GC, *Tmem119*^{+/+} and *Tmem119*^{-/-} mice (Bennett et al., 2016) were immunized with NP-CGG in Sigma Adjuvant and analyzed on day 11. While *Tmem119*^{-/-} pLN LZ Fc γ R2b⁺ FDC networks had undetectable TMEM119, there was no change in FDC morphology, GC polarization, or frequencies of GC B cell, NP-specific GC B cell, follicular B cell, plasma cell, and memory-phenotype B cell (Figures 7K and S6B). TMEM119 was therefore not essential for mounting this type of GC response, but could be important in responses to more complex antigens or in shaping specialized features of the GC.

DISCUSSION

Here we used scRNAseq to identify nine pLN non-endothelial SC clusters. We mapped eight clusters to specific anatomical niches and established differential expression profiles for Ccl19^{hi} TRCs (T-zone), Ccl19^{lo} TRCs (*Ch25h*⁺ cells in follicle-T-zone interface and IFR), Cxcl9⁺ TRCs (T-zone and IFR), MRCs (SCS), PvCs (perivascular), CD34⁺ SCs (capsule and medullary adventitia), Inmt⁺ SCs (medullary cords), and FDCs (follicle center). While

the Cxcl9⁺ TRCs and Nr4a1⁺ SCs, found in several niches, are likely distinguished by transient activation programs instead of niche-establishing programs, the niche association of the majority of transcriptional clustering suggests that this resolution of non-endothelial SC heterogeneity will be useful for studying their niche-specific functions.

The differential gene expression profiles of the nine clusters provide a unique opportunity to identify positive markers, develop additional tools, and study functions of rare and poorly studied SC types. Previously published FDC expression profiles have suffered from B cell, myeloid cell, or FRC contamination or required additional manipulation, such as irradiation of the mice or *in vitro* culture (Heesters et al., 2014; Suzuki et al., 2010; Wilke et al., 2010). Our analysis of freshly harvested pLN FDCs revealed *Pthlh*, *Sox9*, and *Tmem119* as markers of FDCs that could be used to develop more specific genetic tools. Staining for SOX9 and TMEM119 on human tonsil and LN GCs supports conserved expression of these genes by human FDCs (Uhlén et al., 2015) (proteinatlas.org). DZ CRC expression of SOX9 in combination with *Cxc112* also suggests an isolation strategy for these rare and poorly understood cells. In addition, MedRCs previously had no distinctive positive markers, but *in situ* validation of top DEGs *Inmt*, *Cxc19*, and *Cxc110* suggests that MedRCs are *Inmt* +*Cxc110*+*Cxc19*-.

How distinct, niche-associated stromal subsets develop and their degree of plasticity in the adult is incompletely understood. Previous work with *Ccl19-* and *Pdpn*-directed lineage reporters have demonstrated the shared lineage of LN SCs across niches (Chai et al., 2013; Onder et al., 2011). While the SC clustering suggests that most pLN non-endothelial SC subsets are transcriptionally similar, the cluster DEGs can guide selection of specific markers and unique transcription factors for additional fate-mapping efforts to address the developmental relationships between niche-associated subsets. Since SCs cultured *in vitro* tend to lose their specific features, studying the cluster-specific transcription factors defined here could help explore non-endothelial SC plasticity and improve *in vitro* models.

While the nine pLN non-endothelial SC clusters are conserved in two biological samples, we expect proportional composition will vary with digestion protocols and more heterogeneity will be found with improved scRNA capture technology. The large proportion of Ccl19^{lo} TRCs suggests that this cluster includes not only the *Ch25h*⁺ cells at the follicle-T-zone interface, but also stroma in another T-zone-adjacent niche such as the T-zone-medulla interface or deep cortex periphery (Sainte Marie, 2010). In addition, our dataset did not resolve versatile SCs, non-FDC *Cr2*-lineage marked cells in the follicle and T-zone (Mionnet et al., 2013), and these may also be included in the Ccl19^{lo} TRC or in another cluster. We anticipate that as more is learned about subset heterogeneity and niche localization, it will be appropriate to revise the draft nomenclature used here. Nevertheless, our transcriptomic analysis of nine niche-associated pLN non-endothelial SC clusters can aid study of their roles in the compartmentalized steps of the immune response as well as facilitate comparisons to SCs at sites of chronic inflammation to identify methods for clinical targeting.

STAR ***** METHODS

Detailed methods are provided in the online version of this paper and include the following:

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rat monoclonal anti-Thy1.2 (clone 30- H12) - Biotin	Biolegend	CAT#105304; RRID:AB_313175
Rat anti-CD45 (clone 30-F11) - MACS microbeads	Miltenyi	CAT#130-052-301
Rat anti-CD45 (clone 30-F11) - PerCpCy5.5	Biolegend	CAT#103132; RRID:AB_893340
Rat anti-CD31 (clone MEC 13.3) - PE	BD Biosciences	CAT#553373; RRID:AB_394819
Syrian hamster anti-gp38 (clone 8.1.1) - APC	Biolegend	CAT#127410; RRID:AB_10613649
Mouse anti-CD45.2 (clone 104) - Pacific Blue	Biolegend	CAT#109820; RRID:AB_492872
Streptavidin - PECy7	Biolegend	CAT#405206
Syrian hamster anti-gp38 (clone 8.1.1) - Alexa488	Biolegend	CAT#127405; RRID:AB_1133992
anti-CD31 PECy7 (clone 390)	Biolegend	CAT#102417; RRID:AB_830756
anti-CD45 BUV395 (clone 30-F11)	BD Biosciences	CAT#565967
Rat anti-CD16/32 (clone 93)	Biolegend	CAT#101320; RRID:AB_1574975
Fixable Viability Dye eFluor780	eBioscience	CAT#65-0865-14
Rat anti-B220 (clone RA3-6B2) - BV785	Biolegend	CAT#103246; RRID:AB_2563256
Rat anti-IgD (clone 11-26c.2a) - BV650	Biolegend	CAT#405721; RRID:AB_2562731
Rat anti-IgD (clone 11-26c.2a) - Alexa647	Biolegend	CAT#405708; RRID:AB_893528
Rat anti-IgD (clone 11-26c.2a) - Pacific Blue	Biolegend	CAT#405712; RRID:AB_1937244
Armenian hamster anti-Fas (clone Jo2) - PE-Cy7	BD Biosciences	CAT#557653; RRID:AB_396768
Rat anti-T- and B Cell Activation Antigen (clone GL7) – Alexa647	Biolegend	CAT#144606; RRID:AB_2562185
Rat anti-T- and B Cell Activation Antigen (clone GL7) – Pacific Blue	Biolegend	CAT#144614; RRID:AB_2563292
NP - PE	Biosearch Technologies	CAT#N-5070-1
Rat anti-CD138 (clone 281-2) - BV421	Biolegend	CAT#142508; RRID:AB_11203544
Rat anti-CD73 (clone TY/11.8) - PerCpCy5.5	Biolegend	CAT#127214; RRID:AB_11219403
Rat anti-CD38 (clone 90) - Alexa647	Biolegend	CAT#102716; RRID:AB_2073334
Rabbit anti-Tmem119	B. Barres	Bennett et al., 2016
Goat anti-ENPP2	R&D Systems	CAT#AF5255; RRID:AB_2277989
Goat anti-TNFSF11	R&D Systems	CAT#AF462; RRID:AB_2206198
Rat anti-CD34 FITC (clone RAM34)	BD PharMingen	CAT#553733; RRID:AB_395017
Rat anti-CD31 (clone MEC13.3) - biotin	Biolegend	CAT#102504; RRID:AB_312911

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Goat anti-IgD	Cedarlane Labs	Code: GAM/IGD(FC)
Goat anti-desmin	R&D Systems	CAT#AF3844; RRID:AB_2092419
Mouse anti-BST-1 (clone BP-3)	BD Biosciences	CAT#552545; RRID:AB_394419
Rabbit anti-PDGFRβ (clone 28E1)	Cell Signaling Technology	CAT#3169T; RRID:AB_2162497
Armenian hamster anti-CD11c (clone N418) - Alexa647	Biolegend	CAT#117312; RRID:AB_389328
Rabbit anti-Sox9	Millipore	CAT#AB5535; RRID:AB_2239761
Rat anti-CD35 (clone 8C12) - biotin	BD Biosciences	CAT#553816; RRID:AB_395068
anti-CD16/32 (clone 2.4G2) - biotin	UCSF Monoclonal Antibody Core	Lot MC080081
Mouse anti-Bcl6 (clone K112-91) - Alexa647	BD PharMingen	CAT#561525; RRID:AB_10898007
Rabbit anti-GFP - Alexa488	Life Technologies	CAT#A21311; RRID:AB_221477
Rat anti-F4/80 (clone C1:A3-1) - biotin	Cedarlane Laboratories	CAT#CL8940B; RRID:AB_10059797
Mouse anti-FITC - Alexa488	Jackson Immunoresearch	CAT#200-542-037; RRID:AB_2339038
Streptavidin - Cy3	Jackson Immunoresearch	CAT#016-160-084; RRID:AB_2337244
Donkey anti-goat IgG - AMCA	Jackson Immunoresearch	CAT#705-156-147; RRID:AB_2340410
Donkey anti-rabbit IgG - Alexa488	Life Technologies	CAT#A21206; RRID:AB_2535792
Donkey anti-rabbit IgG - Alexa647	Jackson Immunoresearch	CAT#711-606-152; RRID:AB_2340625
Streptavidin - Alexa555	Life Technologies	CAT#S-21381; RRID:AB_2307336
Donkey anti-goat IgG - Horseradish Peroxidase	Jackson Immunoresearch	CAT#705-035-147; RRID:AB_2313587
Streptavidin - Alkaline Phosphatase	Jackson Immunoresearch	CAT# 016-050-084; RRID:AB_2337239
Bacterial and Virus Strains		
Lymphocytic choriomeningitis virus (LCMV)-Armstrong	M. Matloubian	Clingan and Matloubian, 2013
Biological Samples		
pLN frozen blocks REX3-Tg, <i>Cxcl9</i> -RFP <i>Cxcl10-BFP</i> (REX3)	A. Luster	Groom et al., 2012
Chemicals, Peptides, and Recombinant Prote	eins	
Collagenase IV	Worthington Biochemical	CAT#LS004188
DNase I, bovine pancreas (single cell stromal cell analysis)	Sigma-Aldrich	CAT#DN25
DNase I grade II, bovine pancreas (REX3 stromal cell analysis)	Roche	CAT#10104159001
Collagenase P	Sigma-Aldrich	CAT#11213857001
Dispase II	Sigma-Aldrich	CAT#D4693
NP(25)-CGG	Biosearch Technologies	CAT#N-5055C-5
Sigma Adjuvant System	Sigma-Aldrich	CAT#\$6322
Sheep red blood cells (SRBCs) (Sheep #0469)	Colorado Serum Company	CAT#38112
mLTβR-huIgG1 (LTβR-Fc)	J. Browning	N/A
TNFR55-huIgG1 (TNFR1-Fc)	J. Browning	N/A

REAGENT or RESOURCE	SOURCE	IDENTIFIER
human IgG1 (hIg)	J. Browning	N/A
Critical Commercial Assays		
Chromium Single Cell 3' Reagent Kit (v2 Chemistry)	10X Genomics	CAT#120267
RNAscope RED 2.5HD manual assay kit	Advanced Cell Diagnostics	CAT#322350
Deposited Data		
Raw and analyzed data	This paper	GEO: GSE112903
Violin plot web tool	This paper	http://scorpio.ucsf.edu/shiny/LNSC/
Experimental Models: Organisms/Strains		
Mouse: C57BL/6	National Cancer Institute, Charles River	CAT#027
Mouse: B6-CD45.1	National Cancer Institute, Charles River	CAT#564
Mouse: <i>B6.Cg-Cxcl12^{tm2Tng}</i> (<i>Cxcl</i> 12-GFP)	T. Nagasawa	Ara et al., 2003; RRID:IMSR_RBRC04200
Mouse: B6.Tg(Cr2-Cre)3Cgn (Cr2-Cre)	K. Rajewsky	Kraus et al., 2004; RRID:IMSR_JAX:006368
Mouse: <i>B6.Cg-Gt(ROSA)</i> 26Sor ^{tm6(CAG-Zsgreen1)Hze/J} (R26-ZsGreen)	The Jackson Laboratory	CAT#007906; RRID:007906
Mouse: 129S5-Tmem119 ^{<tm1lex> /Mmcd</tm1lex>} (<i>Tmem119</i> KO)	B. Barres	Bennett et al., 2016; RRID:MMRRC_041484-UCD
Mouse: <i>Nr4a1</i> -EGFP BAC-transgenic (<i>Nr4a1</i> -GFP)	A. Weiss	Zikherman et al., 2012; RRID:MMRRC_036737-UCD
Mouse: <i>B6.129S6-Ch25h</i> ^{tm1Rus/J} (<i>Ch25h</i> KO)	D. Russell	Bauman et al., 2009; RRID:IMSR_JAX:016263
Mouse: Tg(UBC-GFP)30Scha/J(UBI-GFP)	The Jackson Laboratory	CAT#004353; RRID:IMSR_JAX:004353
Oligonucleotides		
Primers: <i>Inmt</i> -F CCTTCTCTACAGGAGGTGTAGG; <i>Inmt-</i> <i>R</i> GTTCTGCGGGGGTGTAGTCAG	This paper	NA
Primers: <i>Siglec1-F</i> GGTCAGCCAACAGTTCACTC; <i>Siglec1</i> - R GAGACTCCTGTGGGCACC	This paper	NA
Primers: <i>Pdgfrβ-F</i> GCAGAAGAAGCCACGCTATG; <i>Pdgfrβ-</i> <i>R</i> CAGGTGGAGTCGTAAGGCAA	This paper	NA
Primers: <i>Pthlh-F</i> GGAGTGTCCTGGTATTCCTGC; <i>Pthlh-R</i> CCCTTGTCATGCAGTAGCTGA	This paper	NA
Software and Algorithms		
Flowjo (version 9.9.6)	Treestar	https://www.flowjo.com/
Prism (version 5.0a)	GraphPad	https://www.graphpad.com/scientific-software/prism/
10X Cell Ranger package	10X Genomics	https://support.10xgenomics.com
Seurat (version 2.2)	R. Satija	Butler and Satija, 2017; http://satijalab.org/seurat/
R (version 3.4.2) and dependencies	The Comprehensive R Archive Network	https://cran.r-project.org/
RStudio (version 1.0.153)	RStudio, Inc.	https://www.rstudio.com/
DESeq2 (version 1.16.1)	Love et al., 2014	https://bioconductor.org/packages/release/bioc/html/DESeq2.htr

REAGENT or RESOURCE	SOURCE	IDENTIFIER
RNAscope probe: <i>Ch25h</i> (NM_009890.1, targeting bp 115-1240)	Advanced Cell Diagnostics	CAT#424561
RNAscope probe: <i>Inmt</i> (NM_009349.3, targeting bp 3-1017)	Advanced Cell Diagnostics	CAT#486371
RNAscope probe: <i>Pthlh</i> (NM_008970.4, targeting bp 173-1231)	Advanced Cell Diagnostics	CAT#456521

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Jason Cyster (Jason.Cyster@ucsf.edu).

EXPERIMENTAL MODEL DETAILS

Adult C57BL/6 (B6) and B6-CD45.1 mice were purchased from the National Cancer Institute at Charles River. *B6.Cg-Cxcl12^{tm2Tng}* (*Cxcl12*-GFP) gene-targeted mice were provided by T. Nagasawa and were backcrossed to the B6 background more than seven generations. *B6.Tg*(*Cr2-Cre*)*3Cgn* (*Cr2*-Cre) BAC-transgenic mice were fully backcrossed to B6 and provided by K. Rajewsky (Immune Disease Institute, Boston, MA). *B6.Cg-Gt*(*ROSA*) *26Sor*^{tm6(CAG-Zsgreen1)Hze/J (R26ZsGreen) mice have a floxed stop ZsGreen1 reporter genetically targeted to the Gt(ROSA)26Sor locus and were purchased from The Jackson Laboratory. 129S5-Tmem119<tm1Lex > /Mmcd (*Tmem119^{-/-}*) mice were fully backcrossed to B6 and provided by B. Barres (Bennett et al., 2016). *Nr4a1*-EGFP BACtransgenic (*Nr4a1*-GFP) mice (Zikherman et al., 2012) were backcrossed to B6 at least 6 generations from A. Weiss and provided by J. Roose. *B6.129S6-Ch25h^{tm1Rus/J}* (*Ch25h^{-/-}*) mice were fully backcrossed to C57BL/6. *Cxcl9*-RFP *Cxcl10*-BFP transgenic mice (REX3) were made on a B6 background (Groom et al., 2012) and pLNs provided in frozen blocks by A. Luster. *Tg*(*UBC-GFP*)*30Scha/J* (UBI-GFP) transgenic mice were backcrossed to C57BL/6 for more than 8 generations and were from The Jackson Laboratory.}

Bone marrow (BM) chimeras were made as described previously and analyzed after at least 8 weeks (Bannard et al., 2013). UBI-GFP mice were treated intraperitoneally (ip) with 500 µg anti-Thy1.2 (clone 30H12) to ablate radio-resistant T cells before being irradiated and reconstituted with wild-type CD45.1 BM. *Cr2*-Cre R26ZsGreen mice were irradiated and reconstituted with WT CD45.1 BM.

Animals were housed in a specific pathogen–free environment in the Laboratory Animal Research Center at the University of Cal-ifornia, San Francisco, and all experiments conformed to ethical principles and guidelines approved by the University of California, San Francisco, Institutional Animal Care and Use Committee.

METHOD DETAILS

Stromal Cell Preparation and Flow Cytometry—PLN stromal cells (SCs) were prepared for scRNAseq from the inguinal, brachial and axillary LNs of 12 uninfected, adult, female C57BL/6 mice and 9 post-LCMV infected adult, female C57BL/6 mice. PLNs were harvested into DMEM (Fisher Scientific) with 2% FCS (HyClone), 10mM HEPES (Fisher

Scientific) and Pen/Strep (P/S)(Fisher Scientific) on ice and minced with 25G needles (Fisher Scientific). Tissue was transferred to room temperature digestion buffer (DMEM + 2% FCS + HEPES + P/S + 3mg/ml Collagenase IV (Worthington Biochemical) and 40 µg/ml DNase I (Sigma-Aldrich) and incubated at 37°C in a beaker with water and a stir bar gently spinning for 15 min. Tissue was pipetted 50× with a Pasteur pipet and incubated another 15min before being pipetted $100 \times$ and filtered through a 100 µm filter into MACS Buffer (PBS with 2% FCS and 2mM EDTA) on ice. The cell suspension was spun down and resuspended to 1×10^7 cells/ml in MACS buffer with $10 \,\mu$ l/4×10⁷ cells anti-CD45 MACS microbeads (Miltenyi) to rotate for 40min at 4°C. After MACS depletion of CD45⁺ cells, the remaining suspension was stained with anti-CD45 PerCpCy5.5 (30-F11, Biolegend), anti-CD31 PE (MEC 13.3, BD Biosciences), anti-gp38 APC (8.1.1, Biolegend) and DAPI (Invitrogen) for flow cytometry assisted cell sorting for viable CD45⁻ CD31⁻ cells on a FACSAria Fusion into PBS + 0.04% BSA. Cells with the lowest DAPI staining were excluded as SSC-A^{lo} CD45⁻ CD31⁻ PDPN⁻ insufficiently stained CD45⁺ cells. Data was analyzed using FlowJo (Treestar). PLN SCs from individual Nr4a1-GFP mice were processed using the same protocol and additionally stained with anti-CD45 Pacific Blue (104, Biolegend), anti-BST-1 (BP-3, BD Biosciences) conjugated to biotin and streptavidin PECy7 (Biolegend).

For the analysis of REX3 pLN stroma, whole LNs were placed on 70 μ m sterile filters and mechanically disrupted and subjected to digestion in DNase I (100 μ g/ml), Collagenase P (200 μ g/ml), Dispase II (800 μ g/ml), 1% FCS in RPMI. LNs were placed in the pre-warmed enzyme mixture and incubated at 37°C. At 8 minute intervals, any supernatant was removed, added to RPMI/2mM EDTA/1% FCS, and replaced with fresh enzyme media. This was repeated at 8min intervals until no large tissue fragments remained. For flow cytometry (FC) on Rex3 pLN stroma, single-cell suspensions of 2×10⁶ cells underwent staining in PBS/ 0.5% FCS with anti-gp38 Alexa488 (Biolegend), anti-CD31 PECy7 (Biolegend), anti-CD45 BUV395 (BD Biosciences), anti-CD16/32 (Biolegend), and Fixable Viability Dye eFluor780 (eBioscience). Samples were run on BD Fortessa X20 and data was analyzed using FlowJo (TreeStar) and Prism (GraphPad).

Single-cell suspensions of B cells were generated and stained as previously described (Allen et al., 2007). The following antibodies were used for cell staining: anti-B220 BV785 (RA3-6B2, Biolegend), anti-IgD BV650 and Pacific Blue (11-26c.2a, Biolegend), anti-Fas PE-Cy7 (Jo2, BD Biosciences/Fisher), anti-T- and B Cell Activation Antigen Pacific Blue and Alexa647 (GL7, Biolegend), NP-PE (Biosearch Technologies), Fixable Viability Dye ef780 (eBioscience), anti-CD138 BV421 (281-2, Biolegend), anti-CD73 PerCpCy5.5 (TY/ 11.8, Biolegend) and anti-CD38 Alexa647 (90, Biolegend). Samples were acquired and analyzed with a BD LSRII, Flowjo (Treestar) and Prism (GraphPad).

Droplet-based single-cell RNA sequencing—Immediately post-sorting, DAPI[–] CD45[–] CD31[–] pLN SCs were run on the 10X Chromium (10X Genomics)(Zheng et al., 2017) and then through library preparation by the Institute for Human Genetics at UCSF following the recommended protocol for the Chromium Single Cell 3' Reagent Kit (v2 Chemistry). Libraries were run on the HiSeq4000 for Illumina sequencing. Post-processing and quality control were performed by the Genomics Core Facility at the Institute for

Human Genetics at UCSF using the 10X Cell Ranger package (v1.2.0, 10X Genomics). Reads were aligned to mm10 reference assembly (v1.2.0, 10X Genomics). Primary assessment with this software for the uninfected sample reported 2,912 cell-barcodes with 5,542 median unique molecular identifiers (UMIs, transcripts) per cell and 2,148 median genes per cell sequenced to 87.8% sequencing saturation with 116,135 mean reads per cell. Primary assessment with this software for the post-LCMV infection sample reported 12,686 cell-barcodes with 4,477 median unique transcripts per cell and 1,937 median genes per cell sequenced to 59.9% sequencing saturation with 26,050 mean reads per cell.

Infections, Immunizations and Treatments—Mice were infected with LCMV-Armstrong intravenously (iv) at 2.5×10^5 pfu and analyzed on day 15 for scRNAseq and immunocytochemistry.

Mice were immunized with 0.5mg/ml NP(25)-CGG (Biosearch Technologies) in Sigma Adjuvant System (Sigma-Aldrich) in a total of 185 μ l/mouse subcutaneously (sc) at the shoulders, flanks and above the tail and analyzed on day 10.

For LT β R and TNFR1 signaling blockade, *Cxc112*-GFP (for SOX9 assessment) or UBI-GFP reverse chimeric mice (for TMEM119 assessment) were immunized sc at the shoulders, flanks and above the tail with SRBCs (Colorado Serum Company) on day 0 and day 5 and on day 10 treated iv with 100 µL each of 1mg/ml mLT β R-huIgG1 (LT β R-Fc, provided by J. Browning) and 1mg/ml TNFR55-huIgG1 (TNFR1-Fc, provided by J. Browning) or saline. Draining pLNs (axillary, brachial and inguinal) were analyzed 4 days later. C57BL/6 mice were treated on day 0 with LT β R-Fc and TNFR1-Fc and analyzed on day 3 (n = 1) or treated on day 0 and day 4 with LT β R-Fc and TNFR1-Fc (n = 1) or human IgG (hIg)(n = 1) and analyzed day 7. The *n* number listed in the figure legend and here indicates the number of mice examined in each experiment.

Immunofluorescence Microscopy—Tissues expressing GFP or ZsGreen were fixed in 4% PFA for 2 hours at 4°C, washed and sunk in 30% sucrose before freezing in OCT. REX3 tissues were fixed in PLP and processed as previously described (Groom et al., 2012). All other tissues were directly frozen in OCT and slides were fixed in acetone. 7 μ m cryosections and 30 μ m sections were stained as described (Rodda et al., 2015) with primary antibodies: Rabbit anti-Tmem119 (produced and gifted by B. Barres)(Bennett et al., 2016), anti-ENPP2 (polyclonal, R&D Systems), anti-TNFSF11 (polyclonal, R&D Systems), anti-CD34 FITC (RAM34, BD PharMingen), anti-CD31 biotin (MEC13.3, Biolegend), goat anti-IgD (polyclonal GAM/IGD(FC)/7S, Cedarlane Labs), goat anti-desmin (polyclonal, R&D Systems), anti-IgD Alexa647 (11-26c.2a, Biolegend), anti-BST-1 (BP-3, BD Biosciences) conjugated to Alexa647, rabbit anti-PDGFR β (28E1, Cell Signaling), anti-CD11c Alexa647 (Biolegend), Rabbit anti-Sox9 (Millipore), anti-CD35 biotin (8C12, BD PharMingen), anti-CD16/32 biotin (93, Biolegend), Alexa647-conjugated anti-Bcl6 (K112-91, BD PharMingen), anti-GFP Alexa488 (Life Technologies) and anti-F4/80 biotin (Cedarlane Laboratories).

Sections were stained with the following secondary antibodies as previously described (Lu et al., 2017; Rodda et al., 2015): anti-FITC Alexa488, streptavidin Cy3, anti-goat AMCA,

anti-Rabbit Alexa488 (Life Technologies), anti-Rabbit Alexa647, streptavidin Alexa555 (Life Technologies), anti-goat horseradish peroxidase, and streptavidin alkaline phosphatase. All secondary antibodies are from Jackson Immunoresearch unless otherwise noted.

Confocal microscopy on thick sections (30µm) was performed as described previously (Bannard et al., 2013). All other images were captured with a Zeiss AxioObserver Z1 microscope.

RNAscope—Performed as previously described (Lu et al., 2017) on 14µm sections using the RNAscope RED 2.5HD manual assay kit (Advanced Cell Diagnostics)(Wang et al., 2012). The RNAscope probes used targeted: *Ch25h* (NM_009890.1, targeting bp 115-1240), *Inmt* (NM_009349.3, targeting bp 3-1017) and *Pthlh* (NM_008970.4, targeting bp 173-1231).

Quantitative RT-PCR—Total RNA from pLNs was extracted using an RNeasy kit (QIAGEN) and reverse-transcribed. Quantitative PCR was performed as described (Lu et al., 2017) with the following primers: *Inmt*-F CCTTCTCTACAGGAGGTGTAGG; *Inmt*-R GTTCTGCGGGGGTGTAGT CAG; *Siglec1*-F GGTCAGCCAACAGTTCACTC; *Siglec1*-R GAGACTCCTGTGGGGCACC; *Pdgfrb*-F GCAGAAGAAGCCACGCTATG; *Pdgfrb*-R CAGGTGGAGTCGTAAGGCAA; *Pthlh*-F GGAGTGTCCTGGTATTCCTGC; *Pthlh*-R CCCTTGTCATGCAGTAGCTGA; *Hprt*-F AGGTTGCAAGCTTGCTGGT; *Hprt*-R TGAAGTACTCATTATAGTCAAGGGCA. Data were analyzed using the comparative CT (2- Ct) method with *Hprt* as the reference. Data plotted and unpaired t test p value calculated with Prism (GraphPad).

QUANTIFICATION AND STATISTICAL ANALYSIS

Unsupervised clustering of scRNAseq data—We used the Seurat R package (version 2.2)(Butler and Satija, 2017) for graph-based clustering and visualizations, all functions mentioned are from this package or the standard R version 3.4.2 package unless otherwise noted and were used with the default parameters unless otherwise noted. We analyzed only cells (unique barcodes) that passed quality control processing (above) and expressed at least 200 genes and only genes that were expressed in at least 3 cells, leaving us with 2,870 cells (5,573 median unique transcripts detected per cell and 2,153 median genes per cell) and 15,633 genes from the uninfected sample and 12,669 cells (4,482 median unique transcripts detected per cell) and 17,257 genes from the post-infection sample for further analysis. We also removed cells with greater than 5% mitochondrial genes and greater than 30% ribosomal protein genes.

Initially analyzing each sample separately, we applied library-size normalization to each cell with NormalizeData. Normalized expression for gene i in cell j was calculated by taking the natural log of the UMI counts for gene i in cell j divided by the total UMI counts in cell j multiplied by 10,000 and added to 1. To reduce the influence of variability in the number of UMIs, mitochondrial gene expression and ribosomal gene expression between cells on the clustering, we used the ScaleData function to linearly regress out these sources of variation before scaling and centering the data for dimensionality reduction. Principle component analysis was run using RunPCA on the variable genes calculated with FindVariableGenes

(uninfected sample: 872 genes using x = (0.1,6), y = (0.5, 15); post-infection sample: 1193 genes using x = (0.0125,6), y = (0.5, 15) and then extended to the full dataset with ProjectPCA. Based on the PCElbowPlot result we decided to use 24 and 39 principle components (PCs) for the clustering of the uninfected sample cells and post-infection sample cells, respectively.

We ran FindClusters to apply shared nearest neighbor (SNN) graph-based clustering to each sample (0.5 for the uninfected sample and 0.4 for the post-infection sample) and used FindAllMarkers (Wilcoxon rank sum test, min.pct = 0.25, only.pos = True, thresh.use = 0.25) to identify the small clusters of endothelial ($Cd31^+$) and non-SCs ($Pdgfra^- Pdgfrb^-$ and/or CD45⁺) to be removed. These included neutrophils ($Lyz2^+$), mast cells ($Mcpt8^+$) and BECs ($Clnd5^+$). Rare keratinocytes (Krt18⁺ Krt19⁺), LECs (Lyve1⁺ Prox1⁺) and Schwann cells (Mbp⁺ Plp1⁺) were also removed. The non-stromal and endothelial cells may have been collected due to low CD45 expression, low CD31 expression or sorting impurity. The remaining cells in each sample were then normalized and scaled as above.

Canonical Correlation Analysis—To combine both samples and identify a conserved gene correlation structure, we applied diagonal Canonical Correlation Analysis with RunCCA. We used the top 2000 variable genes from each sample (2,817 genes, 59.2% overlap) to calculate the correlation components (CCs) and determined 10 CCs captured the variability in the dataset by inspecting the results of DimHeatmap. Using CalcVarExpRatio to calculate the percentage of variance explained by the CCs for each cell, we retained cells with 50% or more variance explained. Discarded cells were analyzed for sample bias and enrichment for cell cycle signature. The Cell Cycle Score (CC Score) was calculated with CellCycleScoring using 96 canonical cell cycle genes (Tirosh et al., 2016) and cells with a score greater than 0.1 were considered cycling.

The retained cells from both samples were then aligned with AlignSubspace using 10 CC dimensions. Alignment quality was assessed by Pearson correlation of averaged log-normalized variable gene expression from each sample with cortest. We then used FindClusters to apply SNN clustering to the combined cells using the 10 aligned CCs and resolution 0.5. After calculating differentially expressed genes (DEGs), we made sure this choice of resolution meant each cluster had at least 50 DEGs. The clustering was visualized with t-distributed stochastic neighbor embedding (tSNE) dimensionality reduction using RunTSNE (10 aligned CCs) and TSNEPlot. We also used BuildClusterTree to report the hierarchical distance matrix relating an 'average' cell from each cluster.

Differential gene expression—Since single-cell technologies currently capture only a portion of the transcripts in any cell, cells of the same type will not all report expression of exactly the same genes. This creates zero-inflated data making it challenging to determine differential expression of lowly expressed genes. To address this issue, we summed the raw UMI counts for each gene in each cluster over groups of twenty cells from the same sample and then treated these 'pseudobulks' (Lun et al., 2016) as technical replicates for DESeq differential gene analysis between each cluster's pseudobulk samples and all other pseudobulk samples (DESeq2 package, version 1.16.1)(Love et al., 2014). We included the

sample origin of each pseudobulk in the design formula of DESeqDataSetFromMatrix to account for batch effects in the p values. The Benjamini-Hochberg method was applied to adjust the p values for multiple testing (reported as FDR). Normalization was performed by the estimateSizeFactors and estimateDispersions functions as part of the DESeq function. We also calculated the proportion of cells in the cluster of interest expressing each gene to focus on differential expression of only well detected genes (pct_in).

Since there is a 4.5-fold difference in the number of cells between our two samples, we considered that batch effects from the larger sample could unduly influence the differential expression analysis. Therefore, we also used FindConservedMarkers to calculate the DEGs. This method calculates the cluster DEGs for the samples separately and then identifies the DEGs for each cluster conserved between the two sample analyses using the Fisher's combined probability to calculate the p values and the Benjamini-Hochberg method to adjust for multiple testing. We found this method to be more conservative as it reports the minimum log2 fold change and maximum FDR for each gene, but all clusters still had at least 50 DEGs. All gene expression enriched for in a cluster and highlighted in the paper was found by both methods.

Subclustering analysis—The Nr4a1⁺ SCs were further subclustered by reapplying the CCA component analysis and alignment as above to cells in this cluster alone and used 10 aligned CCs for SNN clustering (resolution 0.3). DEGs between the three Nr4a1⁺ SC subclusters (greater than 50 DEGs each) were calculated with DESeq2 on pseudobulks as above with batch included in the design. Pearson correlation of the subclusters with the other subsets was calculated by comparing the differential expression (log2 fold change) of the DEGs for each subcluster with the differential expression (log2 fold change) of the genes for each other original subset.

The post-infection FDCs were analyzed for further subclustering by normalizing and performing unbiased clustering as above without CCA (14 PCs, resolution = 0.3), but the subclusters did not have 50 DEGs when compared to the other subcluster and all other clusters using DESeq2 as above.

Visualization—Log-normalized gene expression data was used for visualizations with violin plots (VlnPlot), tSNE plots (FeaturePlot) and expression comparison plots (GenePlot). Pearson correlations were calculated with cor and cor.test and visualized with plot. Scaled log-normalized gene expression data was used for visualizations with dot plots (DotPlot), heatmaps (DoHeatmap) and bar plots made with Prism (GraphPad). We plotted proportions with barplot. We reported DEG lists with write.table. Additional packages used: readxl (1.0), dplyr (0.7.4), Matrix (1.2-12), ggplot2 (2.2.1), cowplot (0.9.2), pheatmap (1.0.8), SummarizedExperiment (1.6.5), DelayedArray (0.2.7), matrixStats (0.52.2), GenomicRanges (1.28.6), GenomeInfoDb (1.12.3), IRanges (2.10.5), S4Vectors (0.14.7), Biobase (2.36.2) and BiocGenerics (0.22.1).

DATA AND SOFTWARE AVAILABILITY

The two scRNAseq datasets have been deposited in GEO under ID code GSE112903. Violin plot visualization of the datasets is available as a web tool at http://scorpio.ucsf.edu/shiny/LNSC.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

• Single-cell RNA sequencing of lymph node stromal cells reveals nine clusters

- Known subsets TRCs, MRCs, FDCs, and perivascular cells identified
- Five additional stromal cell clusters identified and anatomical locations determined
- Two clusters, Cxcl9⁺ TRCs and Nr4a1⁺ SCs, are defined by activation signatures

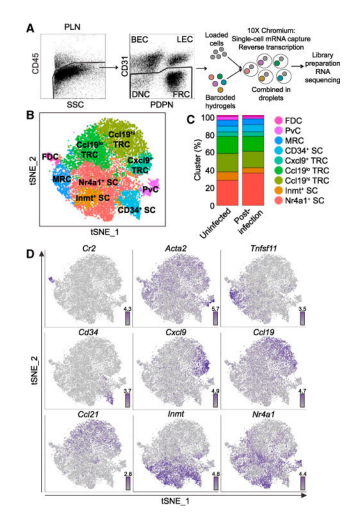


Figure 1. Identification of Nine pLN Non-endothelial SC Clusters by scRNAseq

(A) Droplet-based scRNAseq workflow.

(B) Unsupervised clustering of pLN non-endothelial SCs visualized with tSNE. Each point

is a single cell colored by cluster assignment.

(C) Percent of the nine clusters in two samples: Uninfected and day 15 post-LCMV infection (Post-infection).

(D) Gene expression distinguishing the nine clusters projected onto tSNE plots. Color scaled for each gene with highest log-normalized expression level noted.

See also Figure S1.

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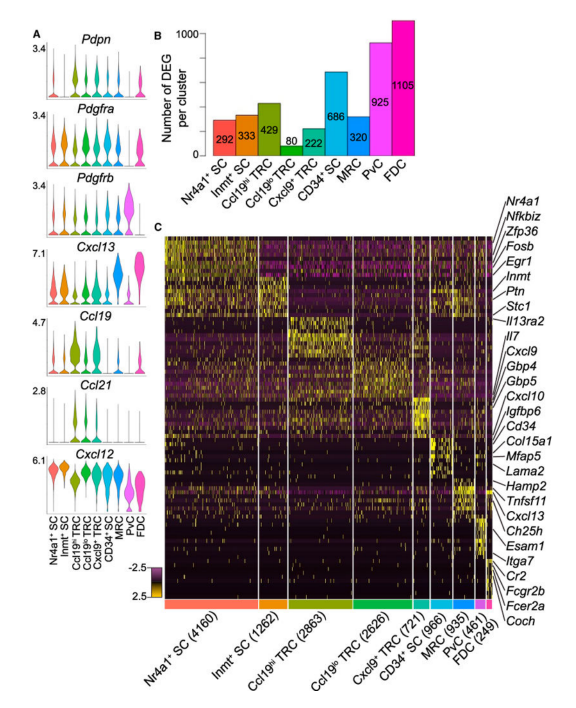
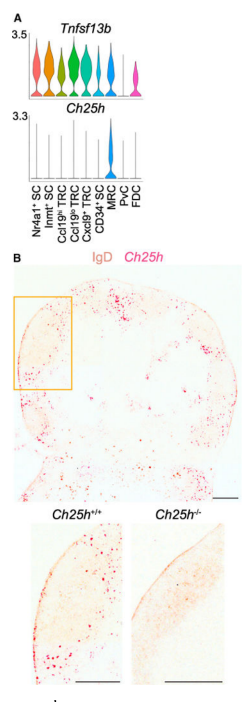


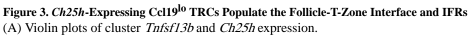
Figure 2. Differential Gene Expression for Nine pLN Non-endothelial SC Clusters (A) Violin plots of canonical SC gene expression by cluster with highest log-normalized expression value labeled.

(B) Number of DEGs per cluster.

(C) Heatmap of each cell's (columns) expression of the top ten DEGs per cluster (rows). Select genes are labeled. Log-normalized expression scaled for each gene. Cluster name and number of cells per cluster displayed below.

See also Figure S2.





(B) RNAscope for *Ch25h* on *Ch25h*^{+/+} (box indicates enlarged area) and *Ch25h*^{-/-} (enlarged only) pLNs counterstained for IgD. Representative of three mice per genotype. Scale bar is 200 μ m.

See also Figure S3.

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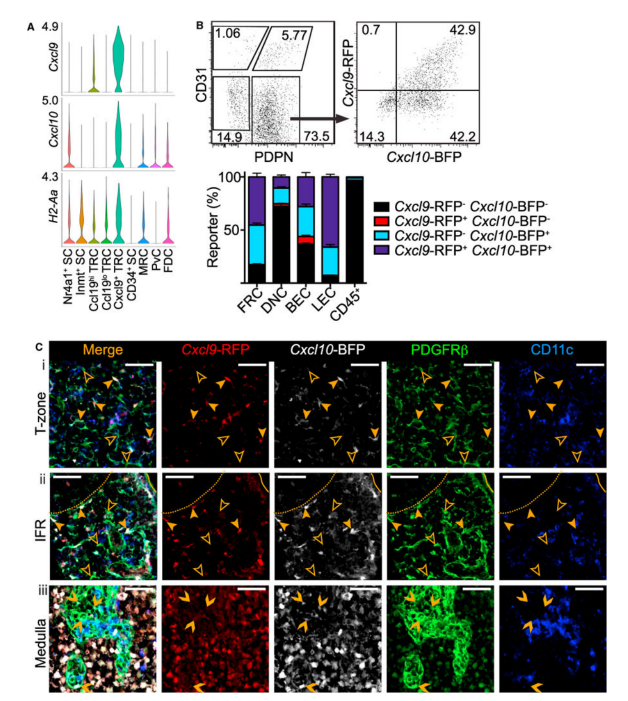


Figure 4. Cxcl9⁺ TRCs Are Located in the T-Zone and IFRs

(A) Violin plots of cluster Cxcl9, Cxcl10, and H2-Aa expression.

(B) Representative gating and percent of REX3 pLN FRCs, DNCs, BECs, and LECs expressing *Cxcl9*-RFP and/or *Cxcl10*-BFP, both or neither by FC. Mean and SEM indicated

(n = 3).

(C) IFM of REX3 pLN (i) T-zone, (ii) IFR, and (iii) medulla stained for PDGFR β^+ CD11c⁻ SCs (representative of 3 mice). Examples of *Cxcl9*-RFP⁺ *Cxcl10*-BFP⁺ SCs (filled

arrowhead), *Cxcl9*-RFP⁻ *Cxcl10*-BFP⁻ SCs (empty arrowhead), and *Cxcl9*-RFP⁻ *Cxcl10*-BFP⁺ SCs (chevron) indicated. Scale bar is 50 μm.

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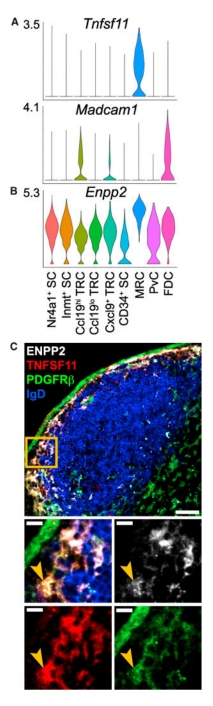


Figure 5. Tnfsf11⁺ MRCs Express *Enpp2* in the SCS

(A and B) Violin plots of cluster *Tnfsf11* (A), *Madcam1* (A), and (B) *Enpp2* expression. (C) IFM of pLN SCS stained for ENPP2 and TNFSF11⁺PDGFR β^+ IgD⁻ MRCs (representative of 3 mice). Scale bar is 50 µm. Box indicates area shown in merged and single channel images beneath. Arrowhead indicates example ENPP2⁺ MRC. Scale bar is 10 µm.

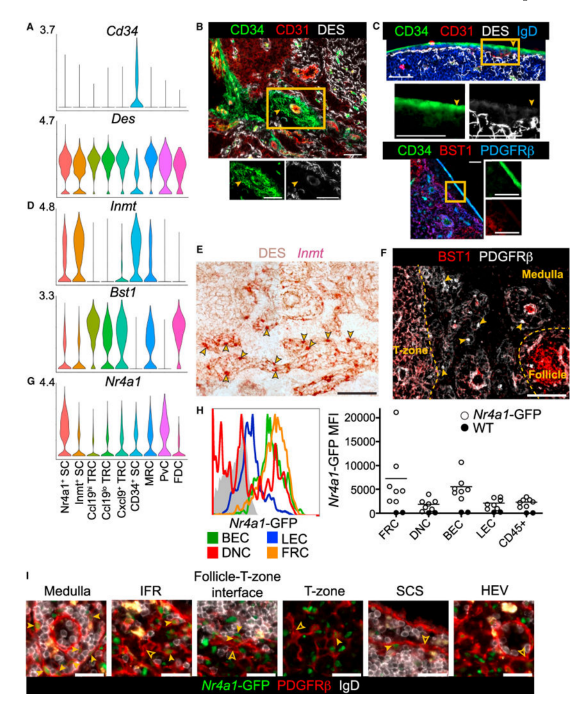


Figure 6. Characterization of CD34⁺ SCs, Inmt⁺ SCs, and Nr4a1⁺ SCs

(A) Violin plots of cluster Cd34 and Des expression.

(B and C) IFM of pLN medulla (B) and capsule (C) for CD34⁺ cells with co-stains indicated. Filled arrowheads indicate examples of CD34⁺DES⁻ staining. Box indicates area shown with individual channels (representative of 3 mice).

(D) Violin plots of cluster Inmt and Bst1 expression.

(E) RNAscope for *Inmt* with DES counterstain on pLN medullary cords (representative of 3 mice). Arrowheads indicate examples of *Inmt*⁺ cells.

(F) IFM of BST1 and PDGFR β on pLN medullary cords (representative of 3 mice). Arrowheads indicate examples of BST1⁻ PDGFR β^+ SCs and yellow dotted lines mark the niche boundaries.

(G) Violin plot of cluster *Nr4a1* expression.

(H) GFP MFI of *Nr4a1*-GFP pLN FRCs, DNCs, BECs, LECs, and CD45⁺ cells detected by FC (n = 7, 3 experiments).

(I) IFM of *Nr4a1*-GFP pLN niches for PDGFR β^+ GFP⁺IgD⁻ SCs (filled arrowheads) and PDGFR β^+ GFP⁻IgD⁻ SCs (empty arrowheads) (representative of 2 mice).

Scale bar is 50 µm in (B), (C), (E), and (F) and 25 µm in (I). See also Figure S5.

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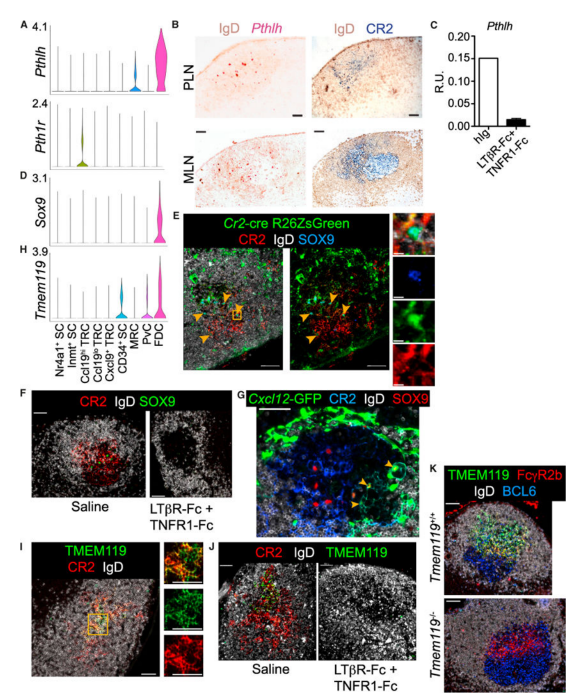


Figure 7. FDCs Express *Pthlh* and *Tmem119* while FDCs and CRCs Express *Sox9*

- (A) Violin plot of cluster *Pthlh* and *Pth1r* expression.
- (B) RNAscope for *Pthlh* with IgD counterstain on pLN primary follicle and mLN GC. Sequential stains for CR2^{hi} FDCs and IgD (representative of 2 mice).
- (C) QPCR of pLN Pthlh expression from mice treated 1 or 2 times with LTBR-Fc and
- TNFR1-Fc (n = 2) or once with human IgG (hIg) (n = 1) (R.U. = relative units).
- (D) Violin plot of cluster *Sox9* expression.

(E) IFM of *Cr2*-cre R26ZsGreen reverse BM chimera pLN follicle (representative of 2 mice). Arrowheads indicate examples of CR2^{hi}ZsGreen⁺SOX9⁺ FDCs. Box indicates area shown with merged and individual channels. Scale bar is 10 μ m.

(F) IFM of SOX9⁺ cells among CR2^{hi} FDC networks in pLN GC from mice immunized with SRBC and on day 10 treated with LT β R-Fc and TNFR1-Fc or saline for analysis on day 14 (representative of 6 pLNs from 1 mouse per treatment).

(G) Thick-section IFM of *Cxc112*-GFP pLN GCs day 10 post-SRBC immunization (representative of 6 pLNs from 1 mouse per treatment). Arrowheads indicate *Cxc112*-GFP +SOX9+CR2^{lo} CRCs.

(H) Violin plot of cluster *Tmem119* expression.

(I) IFM of pLN primary follicle TMEM119⁺CR2^{hi} FDCs (representative of 2 mice). Box indicates area shown with individual channels.

(J) IFM of TMEM119 on CR2^{hi} FDCs in pLN GCs post-immunization and treatment as in (F) (representative of 6 pLN from 1 mouse per treatment).

(K) IFM of TMEM119 on $Fc\gamma R2b^+$ FDCs in BCL6⁺ pLN GCs from *Tmem119^{+/+}* and *Tmem119^{-/-}* mice on day 11 post-NP-CGG immunization (representative of 2 mice per genotype).

Scale bars are 50 µm unless otherwise noted. See also Figure S6.