Lymph-migrating, tissue-derived dendritic cells are minor constituents within steady-state lymph nodes

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Observations that dendritic cells (DCs) constitutively enter afferent lymphatic vessels in many organs and that DCs in some tissues, such as the lung, turnover rapidly in the steady state have led to the concept that a major fraction of lymph node DCs are derived from migratory DCs that enter the lymph node through upstream afferent lymphatic vessels. We used the lysozyme M–Cre reporter mouse strain to assess the relationship of lymph node and nonlymphoid organ DCs. Our findings challenge the idea that a substantial proportion of lymph node DCs derive from the upstream tissue during homeostasis. Instead, our analysis suggests that nonlymphoid organ DCs comprise a major population of DCs within lymph nodes only after introduction of an inflammatory stimulus.

Afferent lymphatic vessels transport interstitial fluid and cells from tissues before terminating in draining lymph nodes. DCs and T cells are the major cellular components of afferent lymph (1–3), but very few DCs enter efferent lymph (3, 4). Inflammatory stimuli increase the rate of afferent lymphatic DC output by approximately an order of magnitude (5). However, inflammatory stimuli are not required for DC and T cell entry into afferent lymphatics, because the recovery of these cells from lymph in the steady state is still substantial (5). Indeed, the projected afferent lymphatic DC output is high enough to fully supply all lymph node DCs, even considering a short DC lifespan (4). Thus, from these early observations, it seemed quite plausible that lymph was the major source of lymph node DCs in general. Reinforcing this possibility were other early studies revealing that DCs purified from the spleen can, upon adoptive transfer, immigrate to lymph nodes through lymphatic vessels but not through the bloodstream (6). Furthermore, revelations that the steady-state turnover of pulmonary DCs is very rapid fit well with the interpretation that this fast turnover was linked to a high rate of DC entry into afferent lymphatics (7). Functionally, DCs that migrate through afferent lymph to lymph nodes in the steady state transport parenchymal antigens from various organs (8–10), and such transport may facilitate the initiation of peripheral immune tolerance (11), possibly working in concert with other mechanisms that give rise to tolerance within lymph nodes (12).

More recently, our understanding of DC homeostasis in lymph nodes has evolved to include not only lymph-migrating DCs but also a role for DCs that enter from blood vessels, the specialized high endothelial venules, to maintain specific DC subsets, notably plasmacytoid DCs (13, 14) and mouse CD8α+ DCs (15, 16). Accordingly, a recent review of DC biology suggested that half of the DCs within lymph nodes enter from the bloodstream, including the full complement of these specialized subsets, but the other half of DCs were still believed to arise from lymph-migrating tissue DCs (for review see reference 17). This conclusion was based in part on the notion that it is possible to readily identify which DCs traveled to lymph nodes through afferent lymph based on combinations of various markers, such as higher expression of MHC II in the steady state (18, 19). However, the relationship between DCs bearing such markers and their role as lymph-migrating DCs was left unclear.

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presumed precursors derived from upstream tissue has not been established.

Our past studies on the role of monocytes as a potential source of lymph-traffic DCs (20–22) fueled our interest in the relationship between nonlymphoid organ DCs and lymph node DCs. Several recent studies have concluded that spleen DCs do not arise from monocytes using adoptive transfer approaches (23–26). In addition, a study in parabiotic mice suggested that monocytes are also not precursors for lymph node DCs, because blood monocytes and lymph node DCs exchanged between parabionts to differing extents (27). Strikingly, one of these studies indicated that, in strong contrast to spleen DCs, monocytes are precursors of peripheral nonlymphoid organ DCs (25), implying for the first time that DCs in lymphoid organs may have a distinct origin from DCs in nonlymphoid organs. However, little lineage analysis has been conducted in lymph nodes, which in contrast to the spleen, are positioned to serve as a nexus wherein lymphoid organ–resident DCs and nonlymphoid organ DCs would converge, presumably in equal proportions (17), to interact with lymphocytes and drive immune responses.

Indications that nonlymphoid organ DCs may arise from monocytes (22, 25, 28), in contrast to other DCs, suggested to us an approach to analyze the proportion of lymph node DCs that might arise from DCs originating in upstream tissues. In this study, we used a popular reporter mouse line that uses the lysozyme promoter, active in many monocytes, to drive Cre recombinase (29) that subsequently removes a stop cassette upstream of enhanced GFP (EGFP) (30). Studies in this model fit well with the evidence that nonlymphoid organ and spleen DCs may be unrelated in origin and, most importantly, reveal that, against expectations and prevailing models in the field, only a minority of lymph node DCs in the steady state arises from DCs residing in upstream nonlymphoid organs. Instead, DCs that migrate through lymphatics from upstream tissues contribute substantially to the pool of lymph node DCs, mainly during inflammatory states.

RESULTS
Illustration of the utility of lysozyme M (LysM)–Cre × Rosa26-stopfl oxEGFP mice to assess DC relationships with monocytes or other DC populations

LysM is expressed by neutrophils and macrophages. Accordingly, LysM-Cre mice (29, 31) have been widely used to generate cell type–selective deletions of floxed genes or expression of floxed reporter markers, especially Rosa26 promoter-driven reporter genes. In these mice, Cre activity removes a stop cassette upstream of the floxed reporter in neutrophils and monocyte-derived macrophages. Neutrophils are not considered candidates for DC precursors, but monocytes may be precursors for at least some DC populations (20–22, 25). If only some DCs are derived from monocytes, and if the lysozyme promoter has low activity in other DC precursors, then the LysM-Cre reporter mice would be a useful tool to trace relationships between monocytes and DCs and possibly between different DC populations in vivo.

When we analyzed blood monocytes in LysM-Cre × Rosa26-stopfl oxEGFP mice in the steady state, the two subsets of circulating CD115lo blood monocytes (32–34) differed in the percentage that was EGFP+. Ly-6Clo monocytes (sometimes called Gr-1hi monocytes) were on average 55% EGFP+, whereas the Ly-6Clo monocyte subset (sometimes called Gr-1lo monocytes) averaged 75% EGFP+ (Fig. 1 A). Since their discovery in mice, these two subsets have not been further subdivided, and accordingly, we found no obvious differences in cell-surface markers between EGFP– and EGFP+ monocyte subpopulations. Furthermore, transfer of EGFP– monocytes into congenic recipient mice revealed that, 1 d later, many of the transferred EGFP– monocytes had become EGFP+ (unpublished data), indicating that EGFP– monocytes do not maintain a distinct population from EGFP+ monocytes. Instead, the failure of EGFP to be uniformly expressed by a given monocyte subset in blood is more likely caused by the stochastic failure of strong lysozyme promoter usage or Cre-mediated excision of the stop cassette in the reporter construct.

In blood, CD115lo cells are granulocytes, mostly high-scatter neutrophils, and 93% of these cells were EGFP+ on average, consistent with the expectation that the lysozyme promoter is most strongly transcribed in these cells (Fig. 1 A). Because of a low level of utilization of the lysozyme promoter by hematopoietic stem cells (31), populations other than neutrophils and monocyte-derived cells, such as lymphocytes, were EGFP+ at a low frequency, but this frequency (6% EGFP+ on average; Fig. 1 A) was substantially less than that of neutrophils or monocytes. As expected, control WT naive mice had no EGFP expression, whereas mice that have EGFP knocked into the Rosa26 locus for constitutive expression (35) showed that >98% of monocytes were EGFP+, thus indicating that the Rosa26 promoter is constitutively active in these cells (Fig. 1 A).

In the bone marrow, monocytes are mainly Ly–6Clo monocytes (36), and these monocytes were EGFP+ at a frequency similar to their blood counterparts, averaging 50% EGFP+ (Fig. 1 B). In contrast, the recently described macrophage DC progenitors (MDPs) (24), identified as CD115lo lineage– cells (37), were EGFP+ to a lower frequency (14% on average; Fig. 1 B).

Although not all monocytes were EGFP+, we reasoned that populations of DC subsets derived from monocytes should express EGFP at a percentage at least equal to the percentage of monocytes that scored positive. If the lysozyme promoter remained or became active during the differentiation of DCs after the monocyte stage, the percentage of DCs positive for EGFP might exceed that of monocytes. However, DCs positive for EGFP at a percentage sufficiently high to meet the criterion for potentially being derived from monocytes would not prove that they originated from monocytes. Nonetheless, as long as the Rosa26 promoter upstream of the EGFP reporter remained active, DC populations expressing a much lower fraction of EGFP than blood monocytes certainly could not be derived primarily from monocytes, and most importantly, EGFP+ DC populations could not be descendants of EGFP+ DCs or monocytes.
Thus, we next analyzed DC subsets in the spleens of LysM-Cre × Rosa26-stopEGFP mice to determine if their propensity to express EGFP may differ from that of monocytes, as might be expected from previous work (23–26). When total CD11c+ cells were considered, EGFP+ frequency more closely matched that of MDPs than monocytes (Fig. 1C). To assess spleen DCs more carefully, we analyzed various DC subsets by further flow cytometric staining and gating. Conventional DCs in the spleen are CD11c+ cells that differentially express CD8α and CD11b, and based on these gates, we separately evaluated EGFP expression in these subsets, labeled as populations 1–3 (Fig. 1C). CD11c+CD11b−B220+ cells were plasmacytoid DCs (Fig. 1C, population 4). Other CD11c− or CD11c− populations that expressed high CD11b were likely splenic monocytes and macrophages (37), and these could be divided in three populations (Fig. 1C, labeled populations 5–7). In our analyses, we used monocytes as a reference population to normalize the data and draw a direct comparison to monocytes. In every mouse analyzed, we calculated the percentage of EGFP expressed in a given...
spleen DC or macrophage population and divided this value by the percentage of Ly-6C<sup>hi</sup> monocytes that were EGFP<sup>+</sup> in the same mouse, because DCs might arise from Ly-6C<sup>hi</sup> monocytes in some tissues (20–22, 25). If the percentage of a given DC subset that was EGFP<sup>+</sup> was similar to Ly-6C<sup>hi</sup> monocytes, then the product of this division would produce a value of 1. Ratios substantially <1 would indicate populations that could not have received a majority of precursor input from monocytes or any other EGFP<sup>+</sup> population. In the spleen, all DC subsets (Fig. 1 C, gated populations 1–4) (37) exhibited a low level of EGFP expression and, fitting with previous conclusions (23–26), did not meet the criterion for a potentially close relationship with monocytes. Failure to meet this criterion was not caused by inactivity at the Rosa26 promoter, because CD11c<sup>hi</sup> and CD11c<sup>int</sup> cells were uniformly misexpressing EGFP<sup>+</sup> in control Rosa26-EGFP knock-in mice (Fig. 1 C). In contrast to DCs, monocytes and macrophages in the spleen (gated populations 5–7) (37) met or exceeded the minimal criterion for possible derivation from monocytes (Fig. 1 C), serving as a positive control for the approach taken.

**Reporter expression by nonlymphoid organ DCs is distinct from spleen DCs in the steady state**

Next, we examined peripheral organ DCs in skin, lung, and small intestine in the steady state. Skin DCs were separated into epidermal DCs (Langerhans cells; Fig. 2 A) and dermal DCs that, as a whole, are CD11c<sup>+</sup> and MHC II<sup>+</sup> (Fig. 2 A, Dermal 1) (38). These tissue DC populations showed a much higher frequency of EGFP<sup>+</sup> cells than that observed for spleen DCs, and overall, the frequency of EGFP<sup>+</sup> skin DCs resembled the frequency seen in Ly-6C<sup>hi</sup> blood monocytes (Fig. 2 A). As noted previously (38), CD11b<sup>hi</sup>CD11c<sup>+</sup> cells in the dermis could be separated into MHC II<sup>+</sup> and MHC II<sup>+</sup> populations (Fig. 2 A, Dermal 2 and 3, respectively). These cells likely include macrophages (38), with considerable heterogeneity in CD11b<sup>hi</sup> MHC II<sup>+</sup> cells. Overall, these populations also showed a frequency of EGFP expression that resembled Ly-6C<sup>hi</sup> monocytes (Fig. 2 A).

Pulmonary DCs are identified as CD11c<sup>+</sup> cells with low autofluorescence, whereas CD11c<sup>+</sup> cells with high autofluorescence are macrophages (22). Two subsets of pulmonary DCs exist, displaying differential expression of CD11b and CD103: CD103<sup>−</sup>CD11b<sup>hi</sup> and CD103<sup>+</sup>CD11b<sup>lo</sup> DCs (22, 39, 40). Both DC subsets, whether recovered from the lung interstitium or bronchoalveolar lavage (BAL), were EGFP<sup>+</sup> to a similar extent as Ly-6C<sup>hi</sup> blood monocytes (Fig. 2 B). Macrophages in the BAL were nearly 100% EGFP<sup>+</sup> (Fig. 2 B), yielding a mean ratio of ~2. Thus, as in the skin, DCs in the lung displayed considerable expression of EGFP, in contrast to spleen DCs and consistent with previous suggestions that blood monocytes may serve as a major precursor source for these cells (22, 25, 28).

In the small intestine, we analyzed DCs and/or macrophages in the lamina propria (LP). Two thirds of MHC II<sup>+</sup> cells expressed high levels of CD11b and intermediate levels of CD11c (Fig. 2 C, gate 3/LP3). Most of these cells were EGFP<sup>+</sup> in LysM-Cre × Rosa26-stop<sup>loxPS</sup>EGFP mice. CD11b<sup>hi</sup> cells with lower levels of MHC II could be separated into two populations based on differential surface expression of CD11b. One of these populations, representing from 4 to 10% of MHC II<sup>+</sup> CD11c<sup>+</sup> cells and expressing low levels of CD11b (Fig. 2 C, gate 1/LP1), was mainly EGFP<sup>+</sup> and stood out as the only potential DC population in any of the nonlymphoid organs examined that expressed EGFP at percentages as low as those observed for spleen DCs. However, this population was also CD8α<sup>+</sup> (Fig. 2 C), resembling a subset of CD11c<sup>−</sup>CD11b<sup>a</sup>CD8α<sup>+</sup> DCs in nearby Peyer’s patches (41) and CD8α<sup>+</sup> lymph node DCs in general. Indeed, this population may not be part of the LP proper but instead may be a contaminant from Peyer’s patches, as we noticed that the population became progressively reduced in frequency (down to 4% of recovered DCs) as our skills in the dissection of the Peyer’s patches away from the LP improved. Another population (Fig. 2 C, gate 2/LP2) expressed intermediate levels of EGFP that did not reach the frequency of EGFP<sup>+</sup> cells observed in monocytes but that maintained a frequency higher than spleen DCs. These cells (Fig. 2 C, gate 2/LP2) are likely the DCs that populate the villi of the intestine and upper LP layers, because in some mice, we recovered DCs from villous tips that were removed while stripping the intestinal epithelium (Fig. 2 C, Villi DC), and the LP DCs recovered from these villi appeared identical in phenotype (not depicted) and EGFP<sup>+</sup> frequency to LP2 (Fig. 2 C).

Considering data from all of these organs together, we conclude that most nonlymphoid organ DCs or their precursors use the lysozyme promoter in the steady state. This feature distinguishes them from spleen DCs and is consistent with, but does not prove, that the origins of nonlymphoid organ DCs may be distinct from spleen DCs, as previously proposed (25).

**Nonlymphoid organ–derived DCs are scarce in lymph nodes**

Because it is thought that lymph nodes receive substantial seeding by DCs that emigrate from upstream nonlymphoid organs through the afferent lymphatics (see Introduction), we expected to find that lymph node DCs, and discrete subpopulations of DCs in particular, would express a higher frequency of EGFP than observed in spleen DCs. However, the frequency of EGFP<sup>+</sup> cells within lymph node DC subpopulations was much like that of spleen DCs, with the exception of skin-draining lymph nodes (Fig. 3, A–C). This could not be attributed to loss of reporter expression through failure to use the Rosa26 promoter, because all lymph node cells and all CD11c<sup>int</sup> and CD11c<sup>hi</sup> cells that would be candidate DCs were EGFP<sup>+</sup> in Rosa26-EGFP knock-in mice (Fig. 3 D).

None of the classical DC populations in the mesenteric or lung-draining lymph nodes mirrored the percentage of EGFP<sup>+</sup> DC populations of the upstream organs (Fig. 3 A, gates 1–3; and Fig. 3 B, gates 1–4; compare with Fig. 2, B and C). As in the spleen, the frequency of EGFP<sup>+</sup> cells was elevated in CD11b<sup>hi</sup>CD11c<sup>−</sup> and CD11b<sup>lo</sup>CD11c<sup>hi</sup> lymph node populations, but these are not typically considered classical DC populations (Fig. 3 A, gates 4 and 5;
organ EGFP + DCs and draining lymph node EGFP + DCs, even in the lung, where it has been shown that DC turnover is especially rapid (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20081430/DC1).

Like the other lymph nodes, most skin-draining lymph node DCs were EGFP + , but certain populations of DCs within skin-draining lymph nodes, especially those that were langerin + , were EGFP + . Expression of langerin, found on all epidermal DCs and a small fraction of dermal DCs (21, 38), is considered a long-lasting marker of DCs that survey the skin and enter skin-draining lymph nodes through lymphatics (42).

Fig. 2. EGFP reporter expression in LysM-Cre × Rosa26-stoploxEGFP mice by DCs and macrophages from skin, lung, and small intestine. (A) Single-cell suspensions were prepared from the epidermis and dermis. Epidermal DCs were CD11c + langerin + , and dermal DCs were gated as CD11c + CD11b + MHC II + (dermal 1), CD11c + CD11b + MHC II + (dermal 2), or CD11c + CD11b + MHC II + (dermal 3). The percentage of EGFP + cells in these populations was normalized to the percentage of EGFP + Ly-6C + monocytes in the same mouse. (B) The percentage of EGFP in the BAL and lung of the two DC subsets (CD103 + CD11b + and CD103 + CD11b + ) and macrophages was normalized to the percentage of EGFP + Ly-6C + monocytes in the same mouse. (C) Gated live cells derived from the small intestine LP (or LP associated with the villous tips during removal of the epithelium) were stained and analyzed to reveal populations expressing CD11c, CD11b, CD8, and MHC II. The LP contained lymphocytes and three populations that resembled DCs and/or macrophages, including CD11c + MHC II + cells that differed with respect to CD11b expression (LP1 and LP2). Population LP2 was also associated with villous tips processed along with the intestinal epithelium (Villi DC). The third population was CD11c + MHC II + CD11b + (LP 3). The percentage of these LP populations that were EGFP + was normalized to the percentage of EGFP + Ly-6C + monocytes in the same mouse. In each graph, all mice studied are compiled, and each data point corresponds to one mouse. Two to three mice per group were handled per experiment. Horizontal bars represent means.
Langerin+ cells in resting LysM-Cre × Rosa26-stoploxEGFP mice represented, on average, 10% of all CD11c+ cells in skin-draining lymph nodes and fell in gates 1 and 4 according to the intensity of CD11c, CD11b, and CD8α staining used to divide the subsets (Fig. 3 C). These langerin+ lymph node DCs and CD11bhi CD11chi CD8α− langerin− DCs (Fig. 3 C, gate 2) expressed a percentage of EGFP that was elevated above the fraction seen in background populations such as lymphocytes, but the EGFP levels were somewhat lower than any of the upstream populations of skin DCs (Fig. 2 A). Collectively, skin-draining lymph nodes contrasted with other lymph nodes examined, as they showed evidence of containing tissue-derived DCs, with up to 20% of the total lymph node DCs (~10% langerin+ cells plus half of the DCs in gate 2 equals ~20%; Fig. 3 C) possibly arriving from skin lymphatics.

DCs highly enriched in MHC II were observed in skin- and lung-draining lymph nodes, present on average in 17 and 9% of total CD11c+ cells, respectively. MHC IIhi DCs were not readily identified in mesenteric lymph nodes or the spleen (Fig. 3 E), although in other studies MHC IIhi cells were found in the spleen (43), indicating that even under specific
pathogen-free conditions the phenotype of DCs in the steady state can vary between different mouse colonies. Because migrating DCs have sometimes been defined by being MHC II+ to a level elevated over other DCs (18, 19), we assessed the frequency of EGFP+ cells within gated MHC II+ lymph node DCs. The frequency of EGFP+ MHC II+ DCs was, like langerin+ DCs, elevated in this gate relative to lymph node DCs overall. However, the percentage of MHC II+ CD11c+ cells that were EGFP+ (Fig. 3 E) fell short of that observed in monocytes and in upstream tissue DCs (compare Fig. 3 E with Fig. 2 A), suggesting that many but not all of the MHC II+ DCs might arise from defined, upstream tissue DC populations. As expected (18), the MHC II+ DC gate was partially redundant with the langerin+ gated DCs, because in our analysis 31 ± 2% of MHC II+ DCs were also langerin+ (Fig. 3 F). There was no difference in the EGFP frequency in either langerin+ or langerin– MHC II+ cells (Fig. 3 F), suggesting that some of the MHC II+ DCs may be migratory dermal DCs that are langerin+ and that overall EGFP+ MHC II+ DCs were equivalent to the sum of langerin+ cells in gates 1 and 4 and DCs in gate 2, as shown in Fig. 3 C.

Given that EGFP+ DCs could not give rise to EGFP– DCs, the failure to observe substantial EGFP expression in the vast majority of lymph node DC populations suggests that upstream organ DCs do not substantially populate lymph nodes in the steady state and certainly cannot account for as many as half of lymph node DCs (17). Clearly, some subpopulations of DCs accumulated in steady-state lymph nodes, most notably langerin+ and MHC II+ DCs within skin-draining lymph nodes, but these subpopulations maintained a relatively low overall frequency.

Contrasting with the steady state, DCs from upstream organs are populous in lymph nodes that drain sites of inflammation and immune sensitization

One caveat to the conclusion drawn from Fig. 3 is the possibility that EGFP– organ DCs preferentially migrate to draining lymph nodes over their EGFP+ counterparts. To consider this possibility and to be certain that migratory DCs could be detected as EGFP+ cells in lymph nodes if they were present, we experimentally induced large-scale DC migration by applying a fluorescent contact sensitizer to the skin, thereby using one of the most established DC migration assays (3, 44). Tetramethylrhodamine-5-(and-6)-isothiocyanate (TRITC) was mixed with dibutylphthalate and acetone and applied epicutaneously to trace DC migration (15, 45). As previously reported (45), the migrating population observed in lymph nodes 1 d after sensitization were mainly langerin+ cells (not depicted), which are thought to arise from dermal langerin+ DCs (45), such as the CD11b+ populations observed in Fig. 2 A. In the draining lymph node, TRITC+ DCs represented >30% of all lymph node DCs (Fig. 4 A) and fell within gates 2 and 4, as shown for skin-draining lymph nodes in Fig. 3 C (compare Fig. 3 C with Fig. 4 A, middle). TRITC+ cells were EGFP+ at the same frequency observed for dermal DCs (compare Fig. 4 A with Fig. 2 A), but TRITC– DCs remained mainly EGFP– (Fig. 4 A). Overall, CD11c+ TRITC+ cells in the lymph node phenotypically resembled and showed a similar EGFP frequency as the Dermal 1 population in Fig. 2 A, and CD11c+ TRITC+ cells most closely matched the Dermal 3 population in Fig. 2 A (Fig. 4 A). These data indicate that migrating DCs retain expression of EGFP during mobilization to downstream lymph nodes and illustrate that the migrating DC population does not favor EGFP– DCs in LysM-Cre × Rosa26-stopΔ6–EGFP mice.

We next examined the effect of tissue inflammation on populations of DCs in lung-draining lymph nodes. As mentioned earlier, CD103 expression marks approximately half of the pulmonary DCs, and its expression persists after the DC migrates to the downstream, lung-draining lymph node (40, 46). We therefore monitored expression of CD103 by CD8α+ DCs in the lung-draining lymph node to determine the frequency of this migrated population therein. CD103+ CD8α+ DCs fell within the gate of CD11c+ CD11b– DCs (Fig. 4 B). These CD103+ DCs in the lymph node reflected a frequency of EGFP expression expected from analysis of upstream tissue CD103+ CD11b+ DCs (Fig. 2 B) but comprised a very small component within the gate (<10% gated, and 4% of lymph node DCs overall) when noninflamed lymph nodes from mice treated with PBS were examined (Fig. 4 B). However, when we treated mice intranasally with LPS to greatly enhance DC migration (5, 40, 47) and examined the lymph node 24 h later, CD103+ CD8α– DCs were the dominant population in the same gate (Fig. 4 B, top left dot plots), and quantification of their absolute (Fig. 4 C) and relative (Fig. 4 D) frequency confirmed that CD103+ DCs were now the majority (60%) of the gated CD11c+ CD11b– DCs (Fig. 4 D) and represented 25% of the lymph node DCs overall. LPS treatment led to an increase in the absolute number of all DC populations (Fig. 4, B and C), but migratory populations such as the CD103+ DCs were disproportionately increased (Fig. 4, C and D; and not depicted). CD103+ DCs were EGFP+ to a similar normalized frequency as their upstream tissue counterparts (Fig. 4 E). LPS increased the proportion of all lymph node DC subsets that expressed EGFP (Fig. 4 F), but because it also increased the frequency of EGFP+ monocytes and tissue DCs, it scarcely altered the normalized data. Taking these observations together, we conclude that inflammatory stimuli, such as LPS or mediators like TNF– or IL–1 induced during contact sensitization (48, 49), are generally necessary to provoke substantial accumulation of upstream tissue–derived DCs within lymph nodes that enter by way of the afferent lymphatic vessels, and that, in contrast and against our expectations, in the steady state the lymph node contains few nonlymphoid organ–derived DCs.

DISCUSSION

Historically, several observations led to the development of the view that lymph node DCs arose substantially from lymph-borne DCs that originated in upstream tissues. First, DCs in cannulated afferent lymph are relatively abundant and are clearly involved in transporting antigen. In contrast, committed
DC precursors are sparse in the circulation of humans (4). Furthermore, when afferent lymphatics are severed, the lymph node becomes nearly void of DCs (50) and antigen presentation is blocked (51). Recently, the idea that specialized DC subsets—plasmacytoid and CD8α+ DCs—enter the lymph node from the bloodstream has been introduced, but this understanding has not modified the belief that many lymph node DCs arise from interstitial nonlymphoid organ DCs that mobilize through lymphatics (16, 17) and may even mature in the steady state as they migrate (19). In this study, we present surprising evidence that most steady-state DCs in lymph nodes that drain a variety of organs by and large do not originate from upstream tissue DCs. Instead, we find that it is only during inflammation that tissue-derived DCs comprise a major fraction of lymph node DCs.

The model system used herein used LysM-Cre × Rosa26-stopfloxEYFP mice that "report" expression of LysM in neutrophils, monocytes, and macrophages by irreversibly inducing EGFP expression. A much lower frequency of expression is observed in T and B cells, stemming from promiscuous activation of the LysM promoter in some hematopoietic stem cells (31). Although macrophages continue to express lysozyme as they mature (31), this gene is not considered to be a marker for DCs and may be turned off developmentally upon induction of DC differentiation. The current study’s comparison between tissue DC populations and those in the lymph node was possible because many tissue DCs, but not most lymph node or spleen DCs, expressed EGFP downstream of lysozyme promoter usage. The discrepancy in the frequency of the EGFP reporter between splenic and nonlymphoid

Figure 4. EGFP reporter expression in LysM-Cre × Rosa26-stopfloxEYFP mice by DCs that migrate into lymph nodes from upstream organs in response to inflammatory stimulation. (A) TRITC was applied to skin epicutaneously in a contact-sensitizing solution, and TRITC+ cells in the draining lymph nodes 18 h later were identified (left dot plot, gated area) and further analyzed. The flow cytometry plot on the right shows an overlay of TRITC+ cells (black) over all live lymph node cells (gray) with respect to the expression of CD11c and CD11b. The percentage of EGFP+ cells within TRITC+ and TRITC− lymph node DCs was normalized to the percentage of EGFP− Ly-6C+ monocytes in the same mouse. Graph shows individual mice per data point. TRITC− DCs refers to a gate on TRITC− CD11c+ DCs that excludes the CD11b+CD11c+ gate that is considered to contain mainly monocytes (reference 37). The TRITC− CD11c+ grouping includes all DCs and the putative monocyte gate, thereby containing all TRITC− CD11c+ and CD11c− cells in the lymph node. (B) PBS or LPS was delivered intranasally (I.N.). 24 h later, live cells from the lung-draining lymph nodes were gated to identify CD11c+ CD11b− cells, and gated CD11c+ CD11b− cells were examined for CD103 or CD8α expression, revealing three subpopulations: CD103+ CD8α− (CD103+) lung-derived DCs, CD8α+ DCs, and double-negative DCs. Shown is the number (C) or percentage (D) of the CD103+ CD8α− (CD103+) DCs, CD8α+ DCs, and double-negative DCs within the CD11c+ CD11b− gated population (from the FACS dot plot, top left) from the lung-draining lymph nodes treated with LPS (black squares) or with PBS (gray circles) to maintain near steady-state conditions. (E) Graph shows the effect of LPS treatment, used to induce DC migration from the lung, on the expression of EGFP by various populations, normalized to Ly6C+ monocytes within the same mouse. BAL and lung were gated on CD11c+CD11b+CD103+ DCs (CD103+), Lung-draining lymph node was gated on CD11c+CD11b+CD103+ DCs (CD103+), Lung-draining lymph node was gated on CD11c+CD11b+CD103+ DCs (CD103+), and plasmacytoid DCs. Horizontal bars represent means. (F) Percentage of blood monocyte subsets and lymph node DC populations that expressed EGFP with or without intranasal LPS administration (from mice in E). In each graph, all mice studied are compiled, and each data point corresponds to one mouse. Two to three mice per group were handled per experiment. Data represent the mean ± SEM.
organ DCs is consistent with the possibility that nonlymphoid organ DCs may derive from monocytes or some other lysozyme-positive precursor, but that most lymph node or spleen DCs do not (25). Both lymph node and spleen DCs more closely expressed the frequency of EGFP observed in bone marrow DC precursors called MDPs (24, 37), which do not inevitably differentiate into monocytes and may shut off EGFP as they differentiate into alternative end-stage cells, including DCs. Although complementing former adoptive transfer studies, (23–25) our approach has the advantage of permitting examination of mice with no previous experimental manipulation, thus truly maintaining the steady state. In addition, we examined whole populations, rather than a small proportion of transferred cells that require extrapolation, to make conclusions about what might have occurred in most members of a given population. Though our data are consistent with the possibility that nonlymphoid organ DCs may arise from monocytes, and indeed we used Ly-6Chi monocytes as a reference population throughout this study, our approach cannot prove an origin from monocytes for any population. However, because EGFP+ cells cannot be precursors for EGFP− cells in these mice, the strength of the model lies in its power to reveal which relationships between populations cannot be true.

Based on the fact that nonlymphoid organ DCs are frequently EGFP+ in LysM-Cre × Rosa26-stop6EGFP mice and lymph nodes DCs are not, we conclude that nonlymphoid organ DCs cannot account for the origin of most steady-state lymph node DCs, with the exception of discrete populations of skin DCs, such as langerin+ DCs, in skin-draining lymph nodes, with the exception of discrete populations of skin DCs, such as langerin+ DCs, in skin-draining lymph nodes. Because lung and airway DCs are well characterized and possess durable markers such as CD103 that independently enable tracking in the steady state, this conclusion is particularly clear and striking when lung-draining lymph nodes are examined (5, 40, 47). In the small intestine, little information is available as to which LP population best migrates into lymph nodes. We found one population of LP DCs with a very low frequency of EGFP+ in LysM-Cre × Rosa26-stop6EGFP mice, an examination of total numbers of CD11bhiCD103+ DCs in steady-state versus inflamed lung-draining lymph nodes (Fig. 4B, bottom graphs) leads to a similar conclusion as that drawn from data that depend on the premises made in analyses of EGFP frequency in LysM-Cre × Rosa26-stop6EGFP mice, an examination of total numbers of CD11bhiCD103+ DCs in steady-state versus inflamed lung-draining lymph nodes (Fig. 4B, bottom graphs) leads to a similar conclusion as that drawn from data that depend on the premises made in analyses of EGFP frequency in LysM-Cre × Rosa26-stop6EGFP mice.

Although we conclude that the presence of nonlymphoid organ DCs is more minimal than expected in lymph nodes, this finding does not necessarily dispel the possibility that there is a relatively high frequency of DCs trafficking through lymph nodes. For example, the lifespan of lymph-migrating DCs, especially in the steady state, may be quite brief within downstream lymph nodes. All of the available evidence suggests that DC turnover in general is very rapid in lymph nodes (15, 27). Skin-resident dermal and epidermal DCs may in particular survive and persist in lymph nodes longer than other DC populations (53, 54). Our finding that these skin DC populations were the most enriched nonlymphoid organ-derived DCs in all lymph node examined corresponds with the idea that lifespan may be the major regulator of the presence of migratory tissue DCs within the lymph node.

Even as ephemeral constituents within lymph nodes, migratory DCs could still play a major role in immunity in the steady state. However, their failure to accumulate in lymph nodes for substantial durations fits best with a role in which they supply antigen, already processed or not, to other DCs within lymph nodes (55, 56). Transfer of antigen from migratory DCs to other DCs may be at least as substantial, or more so, in the steady state as it has been proposed to be during infection (56). The appearance of parietal antigens in CD8α+ DCs (10), a population of so-called lymph node–resident DCs that our data uphold do not derive from upstream tissue DCs, suggests that even large fragments of tissue antigens are not restricted within the lymph node to migratory DCs but are instead readily shared with other DCs. Alternatively, migratory DCs may often possess a phenotype distinct from that expected.
lymph node cells are not considered by most to be classical DCs, and they share this phenotype with Ly-6C\(^+\) monocytes (33, 57), making it tempting to consider all such cells as putative monocytes (37). Cells with this phenotype were abundant in steady-state lymph nodes, were EGFP\(^+\) like upstream tissue counterparts, and may reflect an important DC type with the capacity to directly present antigen or transfer it to other DCs. Particle-bearing, phagocytic monocyte-derived DCs that emigrate from skin to lymph nodes also express low levels of CD11c and very high levels of CD11b (20). Yet, little consideration has been given to the possibility that these cells have an important role in steady-state lymph nodes or, ultimately, in antigen presentation, in part because of the evidence that CD11c\(^+\) cells are required for immune priming, even when CD11c\(^-\) cells are not deleted in CD11c-DTR mice (58).

Because we cannot link lymph node DCs substantially to upstream tissue DCs, more credence may be given to the possibility that lymph node DCs mainly enter lymph nodes during the steady state through high endothelial venules. This notion still requires direct experimental support. Another interesting possibility to be investigated in future work is that the upstream tissue feeds the lymph node with DCs that enter through lymphatics, but that these cells circulate in lymph in an unexpected form, other than as descendants of tissue-resident DCs. The recent evidence that hematopoietic progenitor cells are found in lymph raises the possibility that they might seed the lymph node, although these progenitors were not retained in lymph nodes themselves and were argued to play a more important role in generating tissue DCs during inflammation rather than lymph node DCs in the steady state (59). As our findings challenge long-held views about the role of steady-state trafficking DCs in comprising lymph node populations of DCs, the present work highlights the need to investigate in more detail how lymph nodes are populated by DCs in the steady state, and the relationship between lymph-migrating and nonmigratory DCs in presenting antigen under homeostatic conditions.

MATERIALS AND METHODS

Mice. LysM-Cre mice (29) were crossed to a reporter strain in which the Rosa26 promoter is upstream of a floxed stop cassette and the EGFP gene (30). This cross was provided to us by T. Graf (Centre for Genomic Regulation, Barcelona, Spain) (31), and we subsequently backcrossed the mice 12 generations on the C57BL/6 background before these experiments. Wild-type C57BL/6 and control mice in which EGFP is knocked into the Rosa26 locus for constitutive expression without the need for deletion of a floxed stop cassette (35) were purchased from the Jackson Laboratory. Mice were housed in a specific pathogen-free environment at Mount Sinai School of Medicine and were used in accordance with protocols approved by the Institutional Animal Care and Utilization Committee.

Cell suspension of spleen, lymph nodes, and nonlymphoid organ DCs. Skin-draining lymph node (brachial, axillary, and inguinal), lung-draining lymph node (mediastinal), and mesenteric lymph node were excised, teased with needles and digested in collagenase D (Roche) for 1 h on a shaker. The supernatants that contained a sus- pension of epithelial cells, endothelial lymphocytes, and a significant fraction of villi DCs were collected, washed and filtered. The remaining pieces of intestinal tissue were subjected to further processing. To enrich the LP DC fraction, the mucosal layer of the gut was separated from the rest with the use of a dissection microscope (model SZ51; Olympus) and discarded, whereas the remaining tissue that contained the villi and underlying submucosa (further referred to as the LP) was subjected to tissue digestion with 0.2 mg/ml of type IV collagenase (Sigma–Aldrich) at 37\(^\circ\)C for 1 h. These tissues were homogenized, filtered, and washed. Obtained cell sus- pensions were stained and analyzed by flow cytometry.

Flow cytometry analysis. The following mAbs from BD (except where another vendor is noted) were used for staining: biotinylated mAbs to CD3, B220, Ter119, and I-Ab; PE-conjugated mAbs to CD103 (eBioscience), CD8, B220, I-Ab, or CD115 (eBioscience); PerCP-Cy5.5-conjugated mAbs to Gr-1 (recognizes Ly-6C and Ly-6G) and CD11b; and allophyco- cyanin (APC)-conjugated mAbs to CD11c and pan-NK (eBioscience). Biotinylated mAbs were detected with streptavidin–conjugated APC (Invitrogen). Conjugated isotype-matched control mAbs were obtained from eBiosciences or BD.

To distinguish MDPs from monocytes in bone marrow (37), we stained for two combinations of lineage markers. Lineage stain 1 used a combination of mAbs to CD3, B220, Ter119, and I-Ab; PE-conjugated mAbs to CD103 (eBioscience), CD8, B220, I-Ab, or CD115 (eBioscience); PerCP-Cy5.5-conjugated mAbs to Gr-1 (recognizes Ly-6C and Ly-6G) and CD11b; and allophycocyanin (APC)-conjugated mAbs to CD11c and pan-NK (eBioscience). Biotinylated mAbs were detected with streptavidin–conjugated APC (Invitrogen). Conjugated isotype-matched control mAbs were obtained from eBiosciences or BD.
cells for 20 min at 4°C. Cells were washed twice in 1× Perm/Wash buffer, and the secondary antibody, anti-goat PE (Invitrogen), was diluted in 1× Perm/Wash buffer at a 1:300 concentration and incubated with permeabilized cells for 20 min at 4°C. Finally, cells were washed once in 1× Perm/Wash buffer, followed by a wash in PBS. Cells were resuspended in PBS and analyzed by flow cytometry.

**Fluorescent labeling of DC migration in the skin.** To analyze migration of DCs, TRITC (Sigma-Aldrich) was prepared as a stock solution at a 1:1 acetone/dibutylphthalate mixture (Sigma-Aldrich). The back skin of mice was shaved in three areas, and 25-μl aliquots of the TRITC solution were applied to each area, as described previously (60). 18 h later, brachial, axillary, and inguinal lymph nodes were excised and pooled, stained for flow cytometry, and analyzed.

**Enhanced pulmonary DC migration to the lung-draining lymph nodes.** Intranasal delivery of 5 μg LPS (Sigma-Aldrich) in 30 μl PBS was used to enhance pulmonary DC migration to the lung-draining lymph nodes, as previously described (40). PBS alone, administered similarly by intranasal delivery, was used as a control. 24 h after intranasal delivery, mice were killed and tissues were processed for analysis.

**Online supplemental material.** Fig. S1 analyzes EGFP intensity to determine if EGFP+ DCs might proliferate and dilute EGFP in the process. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20081430/DC1.

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