Myelopoiesis in spleen-producing distinct dendritic-like cells

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Abstract

Dendritic cells (DC) represent a heterogeneous class of antigen presenting cells (APC). Previously we reported a distinct myeloid dendritic-like cell present in spleen, as an in vivo counterpart to cells produced in murine spleen long-term cultures (LTC-DC). These cells, named ‘L-DC’, were found to be functionally and phenotypically distinct from conventional (c)DC, plasmacytoid (p)DC and monocytes. These results suggested that spleen may represent a niche for development of L-DC from endogenous progenitors. Adult murine spleen has now been investigated for the presence of L-DC progenitors. Lineage-negative (Lin) ckitlo and Lin ckithi progenitor subsets were identified as candidate populations, and tested for ability to produce L-DC; in vitro upon co-culture with the spleen stromal line STX3, and in vivo after adoptive therapy into mice. Both subsets colonized STX3 stroma in vitro for L-DC production, indicating that they contained either a common or two distinct progenitors for L-DC. However, only the Lin ckithi subset gave progeny cells after adoptive transfer into lethally irradiated mice. In vivo development was however multilineage and not restricted to L-DC development. Multilineage reconstitution reflects long-term reconstituting haematopoietic stem cells (LT-HSC), suggesting a close relationship between L-DC progenitors and LT-HSC. L-DC were however produced in vivo in much higher number than monocytes/macrophages and cDC, indicating the presence of a specific L-DC progenitor within the Lin ckitlo subset. A model is advanced for development of L-DC directly from haematopoietic progenitors in spleen and dependent on the spleen microenvironment.

Keywords: spleen • haematopoiesis • dendritic cells • myelopoiesis

Introduction

The mouse is a well-recognized model for studying haematopoiesis. At birth, HSC migrate from foetal liver to bone where they remain for the life of the animal. HSC also appear in mouse spleen soon after birth [1] and are maintained there for life. They also exist in extramedullary sites and small numbers mobilize through blood and lymph into tissues like spleen, liver, lung, brain and intestine in the steady-state [2, 3]. The number of haematopoietic stem/progenitor cells (HSPC) in blood, spleen and liver also expands noticeably following acute inflammation or drug treatment [4, 5]. The small number of progenitors in extramedullary tissues in the steady-state should not discount their importance or potential contribution to the immune response. Findings from this laboratory indicate that spleen in the steady-state does contain HSC detectable by their long-term reconstitution ability upon adoptive transfer to irradiated host mice [6]. The importance of bone marrow (BM) in haematopoiesis is clear however since neonatally splenectomized mice can maintain normal BM haematopoiesis [1, 6]. The relative haematopoietic contribution of HSC from spleen and BM of adults is however not known, and it has long been assumed that spleen fills the role of an emergency or backup site in times of stress or disease.

Spleen is also a central organ for development of DC that take up and present antigen to lymphocytes. Multiple subsets have been identified including the CD69- and CD8α+ conventional (c)DC and plasmacytoid (p)DC [7], and the less well-defined regulatory DC [8–10]. Monocyte-derived DC (moDC) are also found in spleen but only under conditions of inflammation [11]. An essential element of DC biology is the definition of progenitors and precursors since this underpins the formation of lineages of cells with distinct function. Initially, cDC and pDC progenitors were defined as a RH3- subset amongst common myeloid and lymphoid progenitors (CMP/CLP) [12]. Current evidence points to a common monocyte/dendritic progenitor (MDP) in BM which gives rise to all monocyte/macrophage and dendritic-type cells [13, 14], and a more committed common dendritic progenitor (CDP) for cDC and pDC in BM [15, 16]. While CDP and MDP are
not present in spleen or blood [17], spleen does harbour cDC precursors, which have a high turnover and are replaced by blood-borne precursors [18]. There is also evidence from para-otic mouse studies to suggest that some splenic DC might arise from endogenous progenitors [19], although others have questioned that result [20].

In this laboratory, a novel dendritic-like antigen presenting cell was discovered in spleen on the basis of similarity with dendritic-like cells developing in splenic long-term cultures (LTC) [21, 22]. ‘LTC-DC’ have a characteristic immature phenotype as CD11c+CD11b+MHC-II+CD8α- cells, distinguishing them from cDC, pDC and monocytes [22, 23]. They are also distinguishable as large, endocytic cells specialized in cross-presentation of antigen to CD8+ T cells [24], a function usually attributed to CD8α+ cDC [25]. They are also distinct in their very weak ability to activate CD4+ T cells, consistent with their MHC-II- phenotype. Since they can be derived from GM-CSF−/− mice [26], they are distinct from moDC [11] or ‘Tip-DC’ [27] that develop in response to inflammatory factors like GM-CSF/TNF-α. The in vivo equivalent ‘L-DC’ subset is readily distinguishable from cDC, pDC and monocytes on the basis of CD11b and CD11c expression, as well as many other markers including CD8α, MHC-II, CD205 and myeloid markers like Ly6G and Mac3 [24]. L-DC show similar antigen cross presenting function as LTC-DC [24] and are functionally distinct from described subsets of regulatory DC which inhibit T cell proliferation [8–10]. The ontogeny and lineage origin of this subset appears to be distinct from other known DC and myeloid subsets in spleen.

It is hypothesized that spleen maintains a lineage of dendritic-like cells, which arise from endogenous haematopoietic progenitors maintained in spleen. Such tissue-specific production of DC has been previously reported for Langerhans cells in skin which are continuously renewed from radio-resistant, skin-derived progenitors [28], only being replaced by blood-borne progenitors under inflammatory conditions [29]. Splenic stromal cells which support haematopoiesis of L-DC in vitro have been shown to have an endothelial origin [30, 31], and L-DC have been shown to arise in co-cultures of BM progenitors or spleen subsets over a splenic stromal cell line [32]. Both neonatal and adult splenocytes contain progenitors that produce L-DC when co-cultured over STX3 spleen stroma [33]. This study identifies and characterizes L-DC progenitors in adult spleen in terms of capacity to produce L-DC in stromal co-cultures and to undergo haematopoiesis for L-DC progenitors in adult spleen in terms of capacity to produce L-DC in inflammatory conditions [29]. Splenic stromal cells which support haematopoiesis of L-DC in vitro have been shown to have an endothelial origin [30, 31], and L-DC have been shown to arise in co-cultures of BM progenitors or spleen subsets over a splenic stromal cell line [32]. Both neonatal and adult splenocytes contain progenitors that produce L-DC when co-cultured over STX3 spleen stroma [33]. This study identifies and characterizes L-DC progenitors in adult spleen in terms of capacity to produce L-DC in inflammatory conditions.

Materials and methods

Animals

C57BL/6J and C57BL/6 S.J.L-Ptpcr<sup>−/−</sup>Pep<sup>−/−</sup>/BoyJ (B6.S.J.L) mice were bred at the John Curtin School of Medical Research (Canberra, Australia) under specific pathogen-free conditions and used at 4–6 weeks of age.

Antibody staining

Antibody staining and flow cytometry were performed to analyse cell surface marker expression as described previously [33]. Non-specific antibody binding via Fc receptors was blocked by incubating cells (×10<sup>6</sup>) with anti-CD16/32 (FcR block) (eBioscience, San Diego, CA, USA). Biotin- or fluorochrome-conjugated antibodies specific for CD11c (N418), CD11b (M1/70), ckit (2B8), IL-7R (A7R34), C045.1 (A20), CD19 (1D3), B220 (RA3-6B2), Thy1.2 (30-H12) and CD34 (RAM34) were purchased from eBioscience. Antibodies specific for CD8 (53-6.7), Sca1 (E13–161.7) and MHC-II (25–9–17) were purchased from Becton Dickinson (San Jose, CA, USA). Isotype control antibodies were purchased from eBioscience. Propidium iodide (PI: 1 μg/ml; Sigma-Aldrich, St. Louis, MO, USA) was added prior to flow cytometry for discrimination of live and dead cells. Flow cytometry was performed immediately on a BD LSRII flow cytometer (Becton Dickinson). Data collected included forward scatter (FSC), side scatter (SSC) and multiple fluorescence channels detecting FITC, CFSE, PE, PI, PE-Cy7, APC and APC-Cy7 (channels FL1-4, FL9-10). BD FACSDiva Software (Becton Dickinson) was used to acquire data. Data analysis involved post-acquisition gating using FlowJo software (Tree Star, Ashland, OR, USA). Cells sorting was performed using a FACSAria cell sorter (Becton Dickinson) as described previously [33].

Enrichment for spleen precursors

Whole splenocytes were enriched for precursors by negative depletion of T cell and B cell populations using antibody-coated magnetic beads as described previously [33], which are specific for CD19 (eBio10D3), Thy1.2 (30-H12) and TER-119, (eBioscience). Recovered cells were washed and then stained with antibody for subsequent isolation of subsets by sorting.

Co-culture assays to assess DC development

Spleen stromal line STX3 is a spleen stromal cell line derived from a long-term culture which ceased production of DC over time with passage [34]. STX3 grows as a confluent monolayer and is passed by scraping and cell transferal. When spleen or BM cells are co-cultured over STX3 (1–5 × 10<sup>5</sup> cells/ml), haematopoiesis is established and myeloid dendritic-like cells are produced. For co-culture maintenance, half medium is exchanged every 3–4 days and non-adherent cells are collected for flow cytometric analysis.

Production of murine chimeras

Chimeras were generated using CD45-allotype distinct donor and host mice as described previously [24]. Haematopoietic cells (CD45.1) were transferred intravenously into lethally irradiated hosts (9.5 Gy) to assess in vivo reconstitution potential of blood cell lineages. Recipients were also given host (CD45.2<sup>+</sup>) BM (10<sup>5</sup> cells) to ensure survival.

Statistical analysis

Data are presented as mean ± S.E., n = 3. With only small sample sizes, a normal distribution cannot be assumed. The Wilcoxon Rank Sum Test was therefore used to assess significance (P ≤ 0.05).
Results

Characterization of L-DC progenitors in adult spleen

Candidate progenitor subsets were identified in adult spleen following staining with markers for HSPC (Fig. 1). A significant fraction of cells (10%) was found to be ckitLoCD11cHi cells, a population reflective of cDC [35]. CD11c− cells expressing ckit were thus excluded from further analysis. The remaining Lin− ckit+ cells could be divided into subsets of Lin− ckitLo and Lin− ckitHi cells, representing ~3% and 1.7% of cells, respectively (Fig. 1A). These were analysed for expression of other known haematopoietic markers [36]. Sca1 was expressed on only ~3% of Lin− ckitHi cells, and the HSPC markers CD34 and Flt3 (not shown) were not expressed on Lin− ckitLo cells. The absence of HSPC in spleen expressing CD34 or Flt3 has been confirmed by others [17, 20]. IL-7R, a marker of lymphoid progenitors in BM [37], was also absent, while a large population of cells were found to be Lin− ckit− CD11b+ (61%), reflecting myeloid precursors.

Differentiation of splenic progenitors in co-cultures

In line with marker analysis shown in Figure 1A, splenocytes depleted of T and B cells (Thy1.2−CD19−) were gated by flow cytometry to exclude CD11c−CD11b− cells and sorted to give Lin− ckit−, Lin− ckitLo and Lin− ckitHi subsets (Fig. 1B). Each of these populations was then co-cultured over STX3 stroma to assess differentiative potential. Lin− ckit− adult spleen precursors failed to produce progeny cells (Fig. 2A and B). In contrast, sorted Lin− ckitHi progenitors generated a high yield of large (FSChi) cells, reflecting two distinct populations of myeloid DC, including CD11b−CD11c−MHCI− L-DC (round gates) and CD11b+CD11c+ MHC-II+ cDC-like cells (square gates). At 15 days after establishment of co-cultures with Lin− ckitHi cells (Fig. 2A), MHC-II+...
cDC-like cells represented a minor population of cells (8.4%), while L-DC accounted for 59% of cells produced. Thereafter, the production of MHC-II⁺ cDC declined 16-fold over 30 days of co-culture (Fig. 2B). The development of MHC-II⁺ DC was thus transient, consistent with development from a preformed precursor, perhaps a ckit⁺CD11c⁻ contaminant. In contrast, co-cultures established with Lin⁻ ckit⁻ precursors produced exclusively L-DC and no MHC-II⁺ cDC-like cells. Although Lin⁻ ckit⁻ splenocytes were restricted in their production of only L-DC, higher yields of L-DC were achieved in Lin⁻ ckit⁻ splenocytes at 15, 24 and 30 days (Fig. 2B). Production of L-DC in co-cultures established with Lin⁻ ckit⁻ and Lin⁻ ckit⁺ precursors was minimal by 15 days and peaked at 24 days. Both Lin⁻ ckit⁻ and Lin⁻ ckit⁺ splenocytes contained self-renewing L-DC progenitors, with a few precursors amongst the Lin⁻ ckit⁺ subset able to develop MHC-II⁺ cDC-like cells transiently.

**Absence of immediate L-DC precursors amongst spleen Lin⁻ ckit⁺ progenitors**

The presence of immediate L-DC precursors amongst the adult spleen Lin⁻ ckit⁺ subsets was further investigated by adoptive transfer of 5 x 10⁶ sorted Lin⁻ ckit⁺ or Lin⁻ ckit⁻ cells into unirradiated hosts, with analysis of donor progeny after 7 and 14 days (Fig. 3). At these times, no donor-type CD11c⁺ dendritic or CD11b⁺ myeloid progeny cells were detected in spleens of recipient mice.
consistent with absence of immediate DC precursors amongst the Lin−ckitlo and Lin−ckithi cell populations. Only control mice given 10^4 (and not 5 × 10^5) unfractionated spleen cells showed donor-type cells in spleen (~0.001%). These were found to be mainly CD11c−CD11b−. The transfer of 5 × 10^5 Lin−ckithi or Lin−ckithi cells represented a 1000-fold enrichment of progenitors over 5 × 10^6 unfractionated spleen cells, reinforcing the conclusion that these subsets contained no immediate precursors of L-DC.

**The spleen Lin−ckithi subset contains LT-HSC and L-DC progenitors**

A unique characteristic of spleen LTC compared with other in vitro DC culture systems is that productivity is sustained for years. One explanation is that self-renewing stem cells maintained in culture differentiate to give L-DC perhaps via formation of an L-DC progenitor [32, 38, 39]. The nature of progenitors amongst the Lin−ckithi and Lin−ckithi subsets was therefore tested by assessing progeny produced following long-term reconstitution of mice with these cell subsets. Adult splenocytes from B6.SJL (CD45.1+) mice were sorted to give Lin−ckithi and Lin−ckithi subsets. These were then cultured with 5 × 10^5 Lin−ckithi or Lin−ckithi cells. The presence of LT-HSC (Fig. 4). Multilineage reconstitution by HSC within the lineage of Lin−ckithi subset was indicated by the detection of donor-derived DC, myeloid cells, T cells and B cells (Fig. S1). No progeny cell reconstitution was achieved with the spleen Lin−ckithi subset (0/3 mice), or with the Lin−ckithi subset (0/4 mice), indicating absence of self-renewing progenitors or LT-HSC within this subset. Furthermore, in mice receiving spleen Lin−ckithi cells, donor-derived cells also reconstituted the corresponding Lin−ckithi stem cell compartment of spleen. Long-term, multilineage reconstitution of hosts by LT-HSC in the spleen Lin−ckithi subset in comparison with BM-derived HSC was also confirmed by analysis of chimeric mice at 54 weeks post-transplantation (Table 1). Three out of four chimeras given donor Lin−ckithi cells showed complete reconstitution with donor myeloid cells, while a fourth chimera showed partial reconstitution. Control chimeras given donor-derived BM alone also gave complete long-term multilineage reconstitution with donor cells in three out of four mice, and also showed new HSC production.

**Capacity of spleen Lin−ckithi progenitors to produce DC in vivo**

The distribution of APC subsets was compared in control mice and in haematopoietic chimeras reconstituted with donor Lin−ckithi spleen cells or donor BM cells. Progeny subsets analysed included CD8α− cDC (CD11c−CD11b−CD8−MHC-II−), CD8α+ cDC (CD11c+CD11b−CD8−MHC-II−), L-DC (FSC−CD11c−CD11b+MHC-II−) and myeloid cells (FSC−CD11c+CD11b+MHC-II−) (Fig. 5A).
Fig. 4 Long-term reconstitution potential of adult spleen Lin<sup>-</sup> ckit<sup>+</sup> cells for myeloid cell subsets. Lin<sup>-</sup> ckit<sup>+</sup>, Lin<sup>-</sup> ckit<sup>lo</sup> and Lin<sup>-</sup> ckit<sup>hi</sup> spleen (SPL) subsets were sorted as described in Figure 1 from spleens of adult B6.SJL mice (CD45.1<sup>+</sup>). Sorted cells (2.5 × 10<sup>5</sup>) were assessed for competitive reconstitution with 10<sup>5</sup> host-type whole bone marrow cells by transfer into lethally irradiated (9.5 Gy) C57BL/6J (CD45.2<sup>+</sup>) recipient mice. Dissociated whole bone marrow (CD45.2<sup>+</sup>) was transferred intravenously alone as a positive control, and age-matched unirradiated C57BL/6J mice were analysed as negative controls. To assess long-term reconstitution, spleens were collected for subset analysis at 15, 16 and 18.5 weeks, enriched for DC by depletion of T and B cells, and stained with fluorochrome-conjugated antibodies to define donor and host cells and their lineage. Prior to flow cytometry, cells were incubated with propidium iodide (PI; 1 μg/ml) for gating live (PI<sup>-</sup>) cells. An example analysis is shown for 16-week chimeras stained to detect CD45.1<sup>+</sup>CD11c<sup>+</sup> progeny cells (top row) and Lin<sup>-</sup> ckit<sup>-</sup> spleen progenitors (bottom row). The frequency of donor cell reconstitution in chimeras after 15, 16 and 18.5 weeks is indicated.

### Table 1

| Animals* (#) | CD8<sup>+</sup>cDC | CD8<sup>-</sup>cDC | L-DC | Myeloid cells | B cells<sup>†</sup> | T cells | HSC<sup>‡</sup> |
|--------------|---------------------|---------------------|------|---------------|-----------------|--------|-------------|
| C57BL/6J control (1) | 0                   | 0                   | 0.2  | 0             | 0               | 0      | 0           |
| Lin<sup>-</sup> ckit<sup>hi</sup> SPL chimeras (1) | 0                   | 0.2                 | 0.2  | 0.1           | 0               | 0      | 0           |
| Lin<sup>-</sup> ckit<sup>hi</sup> SPL chimeras (2) | 78                  | 84                  | 100  | 90            | 17              | 28     | 97          |
| (3) | 99                  | 97                  | 100  | 92            | 98              | 92     | 56          |
| (4) | 96                  | 97                  | 100  | 96            | 73              | 82     | 49          |
| BM chimeras (1) | 8.2                 | 10                  | 23   | 5.7           | 10              | 77     | 13          |
| (2) | 99                  | 99                  | 100  | 99            | 74              | 92     | 97          |
| (3) | 94                  | 94                  | 100  | 98            | 95              | 87     | 100         |
| (4) | 100                 | 100                 | 100  | 100           | 90              | 83     | 81          |

<sup>*</sup>Chimeras were prepared as described in Figures 3 and 4. Analysis of all chimeras was performed at 54 weeks post-reconstitution. <sup>†</sup>B cells and CD8<sup>-</sup> T cells were detected in lymph node. <sup>‡</sup>HSC were identified in spleen as a Lin<sup>-</sup> ckit<sup>Hi</sup>Sca1<sup>+</sup> subset.
shown in terms of percent cells amongst total of T and B cell depleted splenocytes. Distribution of spleen myeloid cell subsets in age-matched, unirradiated mice is shown as a control. Results are shown as the mean ± S.E. of three mice. (A) Representation of donor-derived (CD45.1 

The latter population was also identified in terms of granulocyte and macrophage subsets using specific antibody (data not shown). The population distribution of DC/myeloid subsets in spleens of chimeras was similar to control mice, indicating full haematopoietic reconstitution (Fig. 5B). Each DC or myeloid compartment in spleen was restored to homeostatic levels, with myeloid cells representing the largest population, followed by CD8α+ cDC, CD8α− cDC and L-DC (Fig. 5B). However, only L-DC in spleen showed full reconstitution with donor-derived (CD45.1+) HSC present in spleen Lin− ckit+ cells or in BM (Fig. 5C). The CD8α− cDC, CD8α+ cDC and myeloid cell compartments were only partially replaced by donor-type cells. In chimeras of this type, donor-derived HSC do colonize BM in low number (data not shown). However, there is no clear evidence yet for L-DC development in BM, although a similar but distinct subset of cells is under further investigation.

A comparison of relative numbers of donor- versus host-derived cells confirmed differing levels of chimerism for each cell subset. A distinct trend in relative donor:host levels for different DC subsets was evident across chimeras analysed at 15 to 18.5 weeks, despite variance in overall donor cell reconstitution levels between individual mice. When the fold-increase in donor versus host cell numbers was calculated for each APC subset and standardized to CD8α− cDC (donor:host ratio = 1.0), L-DC consistently exceeded myeloid cells and cDC in terms of relative donor to host cell contribution (Fig. 6). By this analysis, donor-derived progenitors reconstituted CD8α− cDC and CD8α+ cDC subsets to equal levels, but gave ~2-fold more myeloid cells. These results are developmentally consistent with a common progenitor (CDP) for cDC subsets [16], and a common upstream progenitor (MDP) for cDC and macrophages [13]. In contrast, donor-derived progenitors gave rise to significantly higher numbers of L-DC (4–12 fold increase; mean = 8.7) compared with CD8α− cDC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Fig. 6), perhaps indicative of a separate developmental origin for L-DC (Figs 4 and 5), we concluded that the source of HSC was not important. The lodgement of donor-type progenitors into empty niches in spleen would appear to determine the development of L-DC in higher relative numbers than other myeloid/dendritic cell types in spleen.

Discussion

This study addresses myelopoiesis in spleen leading to the development of a novel dendritic-like cell type, namely L-DC, which is unique in terms of its phenotype and immune functional potential [24]. Adult murine spleen is clearly an extramedullary haematopoietic site containing low numbers of multipotent HSC
The hypothesis that extramedullary haematopoiesis mediates the production of tissue-specific APC like L-DC with site-specific functions, is consistent with compartmentalism of the immune response to meet the needs of distinct tissue sites and their respective pathogens. While multiple extramedullary sites clearly contribute to haematopoiesis, the nature of cells produced and the conditions under which this occurs are not yet well defined.

In order to identify an APC subset as distinct it is necessary to show that the lineage origin and progenitor of those cells differs from that of other common dendritic and myeloid subsets. Marker analysis of adult spleen has led to the identification of minor subsets of Lin− ckithi and Lin− ckitlo cells which lack markers like Flt3, CD34 and IL-7R associated with myeloid and lymphoid haematopoietic progenitor subsets in BM (Fig. 1) [17, 20]. There is already evidence that HSC in different tissue sites are different since the marker profile of foetal liver HSC differs from that of BM HSC [40]. Here we have localized the progenitor of L-DC within the Lin− ckithi subset of spleen, which also contains LT-HSC. The presence of LT-HSC in this subset was evident since this subset gave long-term multilineage reconstitution of chimeras out to 54 weeks (Fig. 6). This study therefore identifies a close, if not linked, relationship between LT-HSC and the L-DC progenitor in murine spleen. Further work is under way to obtain the full marker expression profile of splenic HSC, and to determine whether this cell type differs from that of BM HSC with the same differentiative capacity.

Both adult Lin− ckitlo and Lin− ckithi adult spleen subsets were found to contain progenitors which differentiate to give L-DC in co-culture over the STX3 splenic stroma. However, only the Lin− ckithi subset and not the Lin− ckitlo subset contains cells, which reflect self-renewing LT-HSC as demonstrated by their long-term multilineage reconstitution potential in irradiated mice. The differential function of these two subsets in vivo did not mirror their common in vitro differentiative capacity. A first explanation is that they contain a common progenitor, not yet identifiable with available antibodies. The second explanation is that the Lin− ckitlo subset contains L-DC progenitors that derive directly from progenitors within the Lin− ckithi subset of spleen, and that this differentiation occurs when Lin− ckitlo cells are co-cultured over STX3 stroma. The transition of Lin− ckitlo cells into Lin− ckithi cells within co-cultures has proven very difficult to test because of the small size of these subsets in spleen, and the difficulty of recovering enough cells from co-cultures to perform an analysis of phenotypic change.

Indeed, the L-DC progenitor in spleen is phenotypically distinct from other described myeloid and dendritic progenitors present in BM. The CDP and MDP are phenotypically Flt3− cells [13], with no counterpart subset in spleen. The CDP has distinct differentiative capacity for cDC, and the MDP differentiates to give macrophages and DC [17]. Furthermore, neither the splenic Lin− ckitlo nor Lin− ckithi subsets described here resemble the immediate cDC precursor or pre-cDC subset previously identified as ckit+ ckitlo cells, which respond to Flt3L to produce mature cDC and pDC [15, 18]. This pre-cDC subset would have been excluded by our sorting protocol, which specifically gated out Lin+ cells including CD11c+ DC.

L-DC are a distinct CD11c+CD11b+MHC-II+ dendritic-like subset in spleen with strong cross presentation capacity for CD8+ T cell activation [24]. While these cells are phenotypically distinct from monocytes, which are CD11c− and are also unable to cross present antigen [24], it is not yet known whether L-DC share a common lineage relationship with monocytes. Since monocytes, macrophages and cDC/pDC all originate from BM progenitors like MDP and CDP, one expectation is that these cell types might all be reconstituted to similar levels following HSC transfer. This prediction was in fact verified in radiation chimeras shown here, and multiple chimeras demonstrated equal long-term reconstitution of donor-derived splenic monocytes/macrophages, and the CD8α− and CD8α+ cDC populations (Fig. 6). However, these same chimeras showed an average 9-fold increase in donor over host reconstitution of L-DC compared with CD8α− cDC, and a 7-fold...
increase of L-DC over myeloid cells (Fig. 6). Further analyses involving purified progenitors will be necessary in order to establish whether this result indicates independent development of L-DC from macrophage/monocyte lineage cells.

Both in vivo and in vitro evidence now supports the hypothesis that spleen endothelial cells represent a niche for haematopoiesis of L-DC from spleen endogenous self-renewing HSPC. The presence of HSC in spleen has been demonstrated firstly by their ability to reconstitute lethally irradiated hosts [1] (Fig. 4), and secondly since transplanted HSC can home to and engraft spleen [41]. In this laboratory, we have also shown that L-DC progenitors are maintained in spleen LTC, where they appear to continuously self-renew for several years, in a manner strictly dependent on contact with spleen endothelial cells [42]. This study therefore supports a role for spleen as a tissue site supporting haematopoiesis for subsequent production of L-DC. Indeed, a precedent already exists for organ-specific DC haematopoiesis in skin [28]. At this stage, it is not yet known whether L-DC development is an intrinsic property of HSC maintained in spleen, or whether spleen stromal niches can direct DC development from HSC originating from other tissues like BM. Since mice reconstituted with either spleen- or BM-derived HSC also show a strong bias towards donor-derived L-DC reconstitution over cDC and myeloid cells (Fig. 6), the spleen microenvironment rather than the tissue origin of HSC may be the critical factor in directing haematopoiesis of L-DC.

Indeed, an explanation for the bias in development of L-DC over cDC and monocytes can be found in terms of niche space. HSC transplanted intravenously into lethally irradiated mice seed spleen with 37% frequency and BM with 50% frequency [41]. BM contains multiple bone compartments [41] with endothelial and osteoblastic niches, which support HSC development [43]. In contrast, spleen has both lower HSC prevalence [40] and less endothelial niche space than BM, with no osteoblastic niches [43]. Therefore, based on HSC seeding frequency, and relative HSC niche space between BM and spleen, it is likely that intravenously transferred HSC engraft and saturate spleen HSC niches ahead of BM niches, since they are fewer in number but comparable in seeding frequency 1:1.3 ratio [41]. Saturation of niches in spleen by intravenously delivered donor HSC could lead to higher development of donor over host L-DC in spleen, if L-DC development is dependent on splenic endothelial cells.

Evidence is advanced for a spleen-endogenous lineage of dendritic-like cells which develop under steady-state conditions. Both in vivo and in vitro studies support this conclusion that LTC established from neonatal spleen continuously produce L-DC, the adult spleen Lin^- ckit^+ subset contains L-DC progenitors, and adult spleen HSC can reconstitute L-DC to a greater extent than other DC or myeloid subsets. Such a model for spleen-specific DC haematopoiesis will require testing under conditions where cell development is monitored from HSPC engrafted into spleen, but in the absence of BM engraftment.

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Conflict of interest

The authors declare no commercial conflict of interest.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Fig. S1 Shows evidence of long-term multi-potential reconstitution of irradiation chimeras given ckit^hi SPL and bone marrow cells.

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