Extramedullary hematopoiesis: mesenchymal stromal cells from spleen provide an in vitro niche for myelopoiesis

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Abstract
Murine spleen has been shown to harbour stromal cells that support hematopoiesis with production of myeloid antigen–presenting cells. Similar stromal lines have now been isolated from long-term cultures (LTC) of human spleen. When human progenitor populations from spleen, bone marrow and cord blood were employed as a source of progenitors for co-culture above splenic stromal lines, myelopoiesis was supported. Human splenocytes gave production of predominantly myeloid dendritic-like cells, with minor subsets resembling conventional dendritic cells (cDC) cells, and myeloid or monocyte-derived DC. Human bone marrow progenitors gave rise to myelopoiesis from hematopoietic progenitors, while human cord blood supported limited myelopoiesis from existing myeloid precursors. Transcriptome analysis compared two stromal lines differing in myelopoietic support capacity. Gene profiling revealed both stromal lines to reflect perivascular reticular cells with osteogenic characteristics. However, the 5C6 stroma which failed to support hematopoiesis uniquely expressed several inhibitors of the WNT pathway. Combined data now show that splenic stroma of both human and murine origin provides a mesenchymal stromal cell microenvironment which is WNT pathway–dependent, and which supports in vitro myelopoiesis with production of specific subsets of myeloid and dendritic-like cells.

Keywords Spleen · Myelopoiesis · Stroma · Niche

Background
Multiple stromal cell types in bone marrow and lymphoid tissues constitute the cellular niche essential for blood cell development (Dorshkind 1990). Within the bone marrow, stromal cells which contribute to hematopoietic niches have been classified generally as perivascular reticular cells. These lie in close proximity to hematopoietic stem/progenitor cells (HSPC), and adjacent to sinusoids (Crane et al. 2017). Extramedullary hematopoiesis in spleen has now been attributed to HSPC localised to the red pulp region (Inra et al. 2015). Stromal cells in spleen red pulp which support extramedullary hematopoiesis have been identified as mesenchymal PDGFRβ+ cells which secrete CXCL12 and KITL, such that they resemble the perivascular/perisinusoidal reticular cells described originally in the bone marrow hematopoietic niche (Inra et al. 2015).

To date, very few studies report stromal lines which support in vitro hematopoiesis. Murine fetal liver stromal lines were shown to maintain HSPC in vitro (Wineman et al. 1996), and murine splenic stromal cell lines maintain HSPC and also support in vitro myelopoiesis (Despars and O’Neill 2006a; Despars and O’Neill 2006b; Periasamy et al. 2009; Periasamy and O’Neill 2013; Periasamy et al. 2013; Petvises and O’Neill 2014b). In this lab, evidence that stromal cells in spleen support hematopoiesis evolved from...
long-term cultures (LTC) of murine spleen which continu-
ously produced myeloid cells reflecting several dendritic cell
(DC) types (Quah et al. 2004). Further studies showed that
these cells arose from HSPC maintained within the stromal
microenvironment (Wilson et al. 2000). Most studies on the
role of splenic stroma in hematopoiesis have involved mice.
When murine bone marrow HSPC were co-cultured above
selected splenic stromal lines, continuous myelopoiesis
ensued, with production of two main populations of antigen-
presenting cells (APC) distinguishable by different abilities
to activate T lymphocytes (Periasamy et al. 2013, 2009). A
novel population of dendritic-like ‘L-DC’ was continuously
produced and shown to be the majority cell type. This was
distinguishable from a transient, minority population resem-
bling conventional DC (cDC)–like cells (Periasamy et al.
2009; Periasamy and O’Neill 2013; Periasamy et al. 2013).
This novel APC was found to be distinct from the commonly
known subsets of cDC and plasmacytoid (p)DC through lack
of MHC-II expression, high capacity for endocytosis and
ability to activate CD8+ but not CD4+ T cells (Periasamy
et al. 2009; Periasamy and O’Neill 2013; Periasamy et al.
2013). In contrast, cDC-like cells were produced in low
numbers for only 2–3 wk of co-culture, to express MHC-II
and to activate both CD4+ and CD8+ T cells (Periasamy
and O’Neill 2013; Periasamy et al. 2013). An in vivo equivalent
subset of L-DC was subsequently characterised in vivo in
both murine and in the human spleen (Tan et al. 2011; Petvises
et al. 2015; Hey et al. 2016).

Spleen from humans and mice are similar in terms of the
representation and function of dendritic and myeloid sub-
sets (Mittag et al. 2011; Petvises et al. 2015). We have now
investigated the human spleen further for the presence of
stromal cells similar to those of murine spleen. The human
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separation according to the manufacturer’s protocol. Cell separation was achieved by placing the column in a SuperMACS® II Separator (Miltenyi Biotec). The column was then washed so that flow-through cells could be collected. Flow cytometry of Lin− cells was used to confirm the efficiency of depletion of ≥ 95%.

**Human long-term cultures (LTC)** To establish LTC from the human spleen, a suspension of splenocytes was depleted of only red blood cells, plated in 25cm² flasks and left undisturbed for 10 d at 37 °C, 5% CO₂ in air and 95% humidity. The medium was then changed by removal and replacement of supernatant every 3 to 4 d. Cells were maintained in this flask for at least 7 wk. This procedure was originally developed in mice (O’Neill et al. 2004).

**Establishment of co-cultures** Stromal cells were plated for co-culture following dissociation of confluent stromal cultures using 0.25% trypsin/2 mM EDTA and plating at 10⁶ cells/ml. Stromal cell lines were grown as a monolayer to 80–90% confluency. Lin− splenocytes, bone marrow or cord blood cells were then added at 1–5 × 10⁴ cells/ml as an overlay. Co-cultures were held at 37 °C, 5% CO₂ in air and 97% humidity. Medium changes were performed every 3–4 d by removal of half volume and replacement with sDMEM. At 7-d intervals, non-adherent cells were collected through removal and replacement of supernatant. Cell yield was determined and cell subsets identified through analysis of surface marker expression by antibody staining and flow cytometry.

**Antibody staining** The procedure used to stain cells with multiple fluorochrome-conjugated antibodies has been described in detail previously (Periasamy and O’Neill 2013; Periasamy et al. 2013; Petvises and O’Neill 2014b). ‘Fc block’ specific for FcγII/IIIIR (eBioscience) was absorbed to cells ahead of antibody to block non-specific Fc receptor binding. Antibodies used to stain human cells were obtained from Biolegend and were specific for hCD3(OKT3), hCD14(HCD14), hCD19(HIB18), hCD56(HCD56) and hCD45RB (MEM-55), binding mature T cells, B cells, natural killer cells, monocytes/macrophages and granulocytes.

**Microscopy** Stroma alone, LTC, or co-cultures, were photographed using a DM IRE2 inverted research microscope (Leica: North Ryde, NSW, Australia) equipped with DFC digital camera (Leica) to obtain phase-contrast photomicrographs. Images were processed using Leica IM software v4.0.

**Transcriptome analysis** Total RNA was isolated from stromal cell lines using the RNeasy mini kit and following the manufacturer’s protocol (Qiagen). Double-stranded cDNA was synthesised in a two-step process. The first strand of cDNA was synthesised using T7-(dT)₂₄ primers and SuperScript II reverse transcriptase (Invitrogen Life Technologies: Mount Waverley, VIC, Australia), followed by second-strand cDNA synthesis. Double-stranded DNA was purified using phenol–chloroform using phase-lock gels (Brinkmann Instruments: Westbury, NY, USA). Subsequent procedures were performed by the Biomolecular Resources Facility (JCSMR, ANU). In vitro transcription and biotin labelling were performed using the BioArray High Yield RNA Transcript Labelling Kit (Affymetrix). cRNA was cleaned using RNeasy Spin columns (Qiagen), fragmented and then labelled with biotin. Fragmented and labelled cRNA was hybridized to Human Gene 1.0ST GeneChips® following the manufacturer’s procedure (Affymetrix: Santa Clara, CA, USA). These were washed and stained on the fluidics station (Affymetrix) ahead of scanning and image analysis using a Gene Array Scanner (Affymetrix). Scanned images of GeneChips were processed using Microarray Suite 5.0 software (MAS5.0; Affymetrix).

**Analysis of microarray data** Gene expression data was analysed using Partek to give average signal values and p values by Stephen Ohms (Biomolecular Resource Facility: ANU, Canberra, Australia). Microsoft Excel files containing all information on probeset numbers, genes, signal values and p-values were prepared for principal components analysis. Data mining on the basis of average signal values was used to determine gene expression associated with known
functions or lineages. Heatmap analysis and agglomerative hierarchical clustering using the Lance-Williams dissimilarity formula involved the use of R project (http://www.r-project.org/). Panther pathway analysis was used to classify the protein class of expressed genes (https://pantherdb.org/).

Statistical analysis When replicates could be prepared, data are presented as mean ± S. E. for sample size n. The Wilcoxon rank-sum test was used to assess significance (p ≤ 0.05).

Results

Establishment of human spleen long-term cultures Human spleen LTC were established and maintained for at least 7 wk. At biweekly medium change, a 5 ml culture yielded between 10^4 and 10^5 non-adherent cells. Both non-adherent cells and fibroblast-like stromal cells were clearly visible after 7 wk (Fig. 1A). Non-adherent cells were collected weekly and stained with antibodies for flow cytometric analysis to determine cell production. At 4 wk, three distinct subsets were readily distinguishable through the expression of the hCD11c DC marker and variable HLA-DR expression (Fig. 1B). The majority of cells in the hCD11c^{hi}HLA-DR^{+} subset expressed hCD86 and hCD11b as well as hCD80, so reflecting myeloid DC. The hCD11c^{lo}HLA-DR^{+} subset contained cells expressing hCD86, hCD80 and hCD11b, so reflecting cDC-like cells. The hCD11c^{lo}HLA-DR^{-} subset contained cells expressing hCD86, although not hCD80 or hCD11b. These resemble immature DC, and potentially L-DC (Fig. 1B). No plasmacytoid DC (hCD123^{+}) were observed (data not shown). A population of hCD11c^{-}HLA-DR^{-} cells represents myeloid precursors and any residual lymphoid cells maintained in culture. This result was reflective of many LTC analysed. By comparison with DC subsets produced in murine co-cultures (Periasamy and O’Neill 2013; Periasamy et al. 2013; Periasamy et al. 2009), cells produced in human co-cultures express lower levels of hCD11c. Lower level staining for hCD11b was also detected, but the CD11b marker is known to be a better marker of the murine than the human DC lineage (Robinson et al. 1999; Ziegler-Heitbrock et al. 2010).

In order to isolate pure stromal cell lines, non-adherent cells were washed out of LTC, and stromal cells isolated and passaged several times to further deplete any remaining hematopoietic cells. HD248 stroma was then cloned by single-cell sorting and deposition, and clones expanded in culture. Multiple (95) splenic stromal clones were derived from the parent HD248 line and assessed for in vitro growth. These lines, like the hu7B2 line shown in Fig. 2A, retain a fibroblastic-like appearance. Stroma could be maintained as a continuous cell line and passaged regularly either by scraping or enzyme dissociation of cells when replicate cultures were needed for experimentation.

Spleen and bone marrow contain progenitors for DC production Several stromal lines were selected for their superior growth capacity. To test their ability to support hematopoiesis, stromal lines were plated, and Lin^{−} human splenocytes overlaid above the stroma. Three independent co-cultures were established above the hu7B2 stroma and non-adherent cells collected weekly for flow cytometric analysis. After just 7 d, three subsets of cells could be delineated on the basis of hCD11c and HLA-DR expression (Fig. 2B). A hCD11c^{lo}HLA-DR^{+} subset contained cells with high antibody staining, gates were set on bivariate plots using FMOCs. Numbers on gates reflect % cells. Staining for HLA-DR and hCD11c delineated 3 subsets (shown by rectangular gates) which were further analysed for marker expression. Cell production shown by flow cytometry is reflective of LTC established from 3 individual human spleens.
expression of hCD86 and hCD11b, with some cells expressing hCD14, so reflecting myeloid or monocyte-derived DC (Katoh et al. 2005). A hCD11c<sup>−/lo</sup>HLA-DR<sup>+</sup> subset contained cells expressing hCD86 but not CD14, and some with hCD11b, so reflecting cDC-like cells. The hCD11c<sup>−/lo</sup>HLA-DR<sup>+</sup> subset showed some expression of hCD86, such that cells reflect human immature DC and maybe developing L-DC (Fig. 2B). The profile of cells produced after only 7 d of co-culture using a highly enriched population of hematopoietic progenitors from spleen was reflective of cells produced in 4-wk LTC shown in Fig. 1 established with whole splenocytes. Over 21 d, hCD11c<sup>−/lo</sup>HLA-DR<sup>−</sup> cells became the majority population produced in co-cultures, consistent with evidence for L-DC as the majority population produced in murine co-cultures over time (Periasamy and O’Neill 2013; Periasamy et al. 2013). Subsets of hCD11c<sup>hi</sup>HLA-DR<sup>+</sup> and hCD11c<sup>−/lo</sup>HLA-DR<sup>−</sup> cDC-like cells were produced in far lower numbers with time and disappeared after 21 d (Fig. 2C). These co-cultures provide evidence that splenic stroma can support myelopoiesis in spleen HSPC. Myeloid cells expressing the hCD14 marker of monocytes or monocyte-derived DC were produced early, but then dropped significantly after 7 d, suggesting transient production of these cells from pre-existing precursors (Fig. 2D). Production of a predominant immature L-DC-like population, with transient production of cDC-like cells and monocyte-derived DC, mirrors the outcome of murine spleen co-cultures reported previously (Periasamy and O’Neill 2013; Periasamy et al. 2013; Petvises and O’Neill 2014a). Again, subset identification was made more difficult by the low expression of both CD11c and CD11b on human DC subsets.
The human bone marrow was then assessed as a source of progenitors for hematopoiesis in human stromal co-cultures. Productive co-cultures were established by the overlay of Lin− human bone marrow containing a highly enriched population of hematopoietic progenitors above the hu7B2 but not the hu5C6 stromal line (Fig. 3). Non-adherent cells were collected and analysed flow cytometrically. The analysis involved gating to detect cells expressing hCD11c and HLA-DR initially. For the hu7B2 stromal co-cultures, analysis as early as 7 d revealed a single population of cells heterogeneous in their expression of HLA-DR, and with very few cells expressing the DC marker CD11c. A majority of cells showed expression of CD11b, but no cells expressed the DC markers hCD80 and hCD86, the myeloid marker hCD14 or the plasmacytoid DC marker hCD123 (Fig. 3). The human stromal line hu7B2 rapidly supports early myelopoiesis within 7 d of co-culture with human bone marrow progenitors. The hu5C6 stromal line was considered a non-supporter of early myelopoiesis, and the hu7B2 line was selected as a good supporter for further gene profiling.

Human stromal lines were also shown to support hematopoiesis in murine bone marrow (data not shown). After only 14 d, each of 6 distinct stromal lines showed outgrowth of a main cell type with a common phenotype as CD11bhiCD11clo cells which did not stain for MHC-II or B220 but were uniformly F4/80+ cells. These are reflective of the previously described dendritic-like L-DC population of antigen-presenting cells derived in murine spleen stromal co-cultures (Periasamy and O’Neill 2013; Periasamy et al. 2013; Petvises and O’Neill. 2014a, b).

Cord blood as a poor source of hematopoietic progenitors

In comparison with spleen, cord blood was a poor source of progenitors. Co-cultures were established by the overlay of Lin− cord blood cells above hu7B2 stroma, and cells collected and analysed flow cytometrically at 7-d intervals. As for spleen co-cultures shown in Fig. 2, three main subsets of dendritic-like cells were identifiable (Fig. 4A). The hCD11c−HLA-DR− subset which showed no expression of hCD86 with some expression of hCD11b expression reflected immature DC or possibly the L-DC subset described in spleen. Production of these cells reached a peak of ~17% of cells after 14 d, although numbers declined to 3.9% by 21 d (Fig. 4A). The hCD11c+HLA-DR+ subset reflected cDC-like cells with some staining for hCD86 and hCD11b, representing only 4% of cells after 21 d. A subset of hCD11c+HLA-DR+ cells expressing hCD86 and hCD11b and reflecting myeloid DC was only detectable at 21 d (Fig. 4A). Overall, the human cord blood did not appear to contain progenitors to sustain myelopoiesis in co-cultures past 14–21 d.

The human cord blood from four individuals was assessed further for the content of progenitor types (Fig. 4B). Flow cytometric analysis involved initial gating to remove cells expressing the hCD45RB hematopoietic marker as well as lineage markers. The remaining cells were then assessed for hCD34 and hCD38 expression, essential markers for isolation of human HSC (Baum et al. 1992; Kondo et al. 1997). hCD34 versus hCD38 delineates CMP and upstream progenitors (Majeti et al. 2007; Seita and Weissman 2010). The hCD34+hCD38− subset can be divided to give hCD90+ HSC and hCD90− multipotential progenitors (MPP) (Chao

7 day Lin− huBM co-cultures

![Figure 3. The human bone marrow provides progenitors for myelopoiesis in co-cultures. Co-cultures of Lin− human bone marrow cells (huBM) were established over the cloned hu7B2 and hu5C6 stromal lines. Total non-adherent cells produced after 7 d were collected for quantitation and phenotypic identification through flow cytometry. PI−FSC+ live cells were gated initially. Gates were set on bivariate plots using FMOCs, and numbers on gates reflect % cells. A single subset of live, large (high FSC) cells (rectangular gate) was further assessed for marker expression.](#)
et al. 2008; Seita and Weissman 2010; Doulatov et al. 2012) (Fig. 4B). Replicate cord blood samples were found to be a poor source of HSC, with a higher representation of MPP and common myeloid progenitors (CMP) (Fig. 4B).

The average number of HSC in the human cord blood was 0.0021% of nucleated cells, while average numbers of MPP and CMP were 0.085% and 0.279%, respectively (Fig. 4B).

A previous study in mice identified self-renewing HSC as the source of L-DC produced continuously in bone marrow co-cultures over spleen, with a role for MPP in the short-term production of these cells (Petvises and O’Neill 2014b). The absence of HSC in the human cord blood and the low level of MPP are consistent with the inability to sustain myelopoiesis in stromal co-cultures. While CMP are present in higher numbers, their contribution to myelopoiesis in vitro is not supported well by splenic stroma.

**Analysis of human splenic stroma through gene expression analysis** Transcriptome analysis using Affymetrix Human Gene 1.0ST GeneChips® was used to investigate the lineage origin of human stromal lines which support hematopoiesis. The analysis compared the hu7B2 and the hu5C6 lines which were found to differ in capacity to support myelopoiesis in the human bone marrow (BM) (Fig. 3). Distinct differences existed between these two lines in terms of gene expression indicated by principal components analysis (PCA). For example, variability in gene expression for the first principal component was 44% (Fig. 5A). One hypothesis is that these two lines reflect distinct lineages or cell types within a lineage. The alternate hypothesis is that differences are due to natural variation developing after in vitro culture. Hierarchical clustering revealed variability reflected in 4 main clusters which are also evident in heat maps depicting the expression of the 500 most variable genes (Fig. 5B). Despite this variability, data mining revealed a common lineage origin for each of the two lines (Fig. 5C). This involved retrieval of signal values for sets of genes selected from the literature related to lineage and function of endothelial cells, osteogenic cells, mesenchymal stem cells and perivascular reticular cells. The most commonly expressed genes were related to osteogenesis like *SPP1, COL1A2, MMP2, FN1, CDH11, CD90 (THY1)* and *BMP2*, as well as genes expressed by perivascular reticular cells including *MMP3, CXCL12* and *THY1*, with no expression of genes specific to endothelial cells or to mesenchymal stem cells except *COL1A1*, which is also a marker of osteogenic cells (Fig. 5C). These data show that both these two human spleen stromal cells are reflective of perivascular or perisinusoidal reticular cells.
with osteogenic differentiation potential. Similar lineage findings were made with murine spleen stromal lines which support myelopoiesis (O’Neill et al. 2019).

Genes differentially expressed by the two lines were then identified through the selection of the 50 most variable genes. Panther Pathway analysis to identify Protein Class and then selection of genes which encode cell surface molecules. Only genes expressed by hu5C6 were identified under these criteria and these are shown in Fig. 5D. hu5C6 did not support hematopoiesis in overlaid human bone marrow progenitors, although it did support myelopoiesis in overlaid murine bone marrow progenitors (data not shown). It also reflects a perivascular reticular cell type which is an important cellular component of HSC niches in bone marrow and spleen. Three of the four genes identified are known regulators of the WNT/β-catenin signalling pathway which is crucial for HSC self-renewal (Luis et al. 2009), and is a powerful regulator of hematopoiesis in murine spleen co-cultures (Lim et al. 2018). These genes encode proteins which inhibit or modulate the WNT/β-catenin signalling pathway and could maintain HSC in a state of quiescence, so limiting their differentiation to myeloid cells. FBLN5 encodes an inhibitor of the WNT/β-catenin pathway and is a suppressor of cell invasion (Chen et al. 2015), while SFRP1 is highly expressed by stromal cells which maintain HSC and supports homeostasis through extrinsic inhibition of β-catenin. (Renstrom et al. 2009). Finally, FGF10 encodes a key regulator of signalling for cell differentiation from stem cells and is a modulator or inhibitor of WNT signalling (Watson and Francavilla 2018). SPON1 encodes Spondin-1, an extracellular matrix protein with a role in neuronal regeneration, but is not known to regulate hematopoiesis (Woo et

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**Figure 5.** Gene profiling to characterise splenic stromal lines. Differential gene expression analysis was performed on stromal clones hu5C6 and hu7B2 which differ in capacity to support myelopoiesis in the human bone marrow. Transcriptome analysis was performed in duplicate using Affymetrix Human Gene 1.0ST GeneChips®. (A) Principal components analysis (PCA) showed variability in gene expression for each line. (B) Hierarchical clustering identified several main clusters amongst the top 500 most variable genes. (C) Data mining was used to plot signal values for genes specific for endothelial cells (ENDO), osteogenic cells (OSTEO), mesenchymal stem cells (MSC) and perivascular reticular cells (PVRC). (D) Panther pathway analysis was used to classify the protein class of the 50 most variable genes and to identify cell surface and secreted proteins which are differentially expressed.
Indeed, hu5C6 appears to reflect a hematopoietic niche although it differs from hu7B2 in ways unrelated to cell lineage, and more in terms of the ability to regulate or specifically inhibit HSPC proliferation and differentiation.

Discussion

Human splenic stromal cell lines have been isolated from long-term spleen cultures and characterised in terms of their hematopoietic support capacity for myeloid and dendritic lineage cells in line with previous studies in mice (Despars et al. 2004; Periasamy et al. 2018; O’Neill et al. 2019). Indeed, this represents a first report of human splenic stromal lines reflecting perivascular or perisinusoidal reticular cells which provide hematopoietic support for HSPC from the human bone marrow (Corselli et al. 2013) and spleen (Inra et al. 2015; Oda et al. 2018). Human splenic LTC can be readily established and maintain the production of myeloid cells for an extended period as demonstrated previously in the mouse model (Periasamy et al. 2009). Stromal cells can be isolated, cloned and expanded to achieve multiple independent lines. Selected lines were shown to provide hematopoietic support capacity for overlays of human spleen progenitors, and then for progenitors from the human and murine bone marrow. To date, we have not attempted to derive stromal lines from the human bone marrow or cord tissue and to assess the capacity to support hematopoiesis. However, previously, we showed that stromal cell line derivation out of spleen long-term cultures was uniquely specific to spleen, and not to murine thymus, lymph node or blood (Ni and O’Neill 1998). Some long-term cultures of murine bone marrow did support the outgrowth of stroma (Ni and O’Neill 1998), but these did not support myelopoiesis, which appears to be a unique property of spleen tissue (Ni and O’Neill 2001).

The human spleen provides HSPC which can seed stroma and differentiate to give long-term production of a novel dendritic-like cell (L-DC), and short-term production of cDC-like cells and some myeloid DC (Fig. 2C). This result reflects the production of DC in human LTC (Fig. 1). It also reflects previous studies in mice which demonstrate the presence of HSPC in murine spleen which can self-renew in stromal co-cultures to continuously produce the novel subset called L-DC (Petvises and O’Neill 2014b). Human Lin− BM is a rich source of HSPC and seeds human splenic stromal lines like hu7B2 with the production of dendritic-like cells within 7 d (Fig. 3). In contrast to the human bone marrow and spleen, the human cord blood was found to be a very poor source of progenitors for seeding hu7B2 co-cultures (Fig. 4). Upon analysis of progenitor subsets, cord blood was shown to contain very few HSC and far higher numbers of MPP and CMP. The low representation of HSC amongst all progenitors in cord blood would appear to be the reason for the inability to sustain myelopoiesis past 14 to 21 d (Fig. 4A). The possibility however exists that cord tissue contains stromal cells which may support distinct cord blood hematopoietic progenitors or precursors, but this remains to be analysed.

In this study, it has been possible to compare human tissues as a source of hematopoietic progenitors/precursors. Human Lin− spleen is an enriched source of HSPC. When co-cultured above the hu7B2 stromal line, a majority population of immature DC as hCD11c−/HLA-DR−/hCD11b−/lo cells was produced with only minor representation of HLA-DR+cDC-like cells and myeloid or monocyte-derived DC. The immature DC subset appears to represent a counterpart to murine L-DC (O’Neill et al. 2004; Tan et al. 2011; Periasamy and O’Neill 2013). Variability in hCD11b expression has also been observed in human bone marrow co-cultures (Robinson et al. 1999), suggesting that hCD11b is a less reliable marker for identification of human DC in vitro compared with mice. Indeed, CD11b is expressed on a range of cell types including granulocytes, monocytes, DC, NK cells and some T and B cells (Stewart et al. 1995). hCD86 expression was also low and variable and so appears to be a more highly predictive marker for murine DC than human DC. While spleen co-cultures transiently support the production of a minority population of hCD11c+hCD11b+HLA-DR+hCD14+ cells resembling monocyte-derived DC (Kampgen et al. 1994; Romani et al. 1994; Salvato and Lanzavecchia 1994; Caux et al. 1997, 1996), it is not yet known which precursors in human spleen give rise to these cells, or their relationship to the L-DC and cDC-like cells also produced in co-cultures.

Transcriptome analysis of two human splenic stromal lines addressed their lineage origin and their different hematopoietic support capacity. These cell lines do not appear to differ in lineage origin as stromal cells. Both the 7B2 and 5C6 lines express genes specific for perivascular reticular cells like MMP3, THY1 and CXCL12 (Fig. 5C). As with similar murine cell lines, human splenic stromal lines described here express genes reflective of osteogenesis. In the case of mice, stromal lines were also shown to have the potential to undergo osteogenic differentiation (O’Neill et al. 2019). This result is consistent with perivascular reticular cells as osteogenic progenitors, a phenotype which may relate to their hematopoietic support capacity.

Differential gene expression analysis identified several cell surface proteins expressed by hu5C6 but not by hu7B2 which are regulators of the WNT signalling pathway. This is significant in light of evidence that WNT signalling is essential for HSC self-renewal and maintenance within the niche and Wnt3a−/− mice are severely compromised in hematopoiesis (Luis et al. 2009). The combined roles of proteins encoded by FBLN5, FGF10 and SFRP1 is to maintain homeostasis amongst HSC through negative regulation of
the WNT/β-catenin signalling pathway, so inhibiting differentiation. Previous studies in this lab also identified a role for WNT signalling in in vitro hematopoiesis in co-cultures ofmurine splenic stroma (Lim et al. 2018). That study used specific pathway inhibitors to show in vitro hematopoiesis dependent on WNT and Notch signalling, as well as the CXCL12/CXCR4 and SCF/c-KIT receptor-ligand interactions (Lim et al. 2018). Subsequent addition of inhibitory antibodies and small molecule inhibitors to spleen stromal co-cultures confirmed CXCL12, CSF1, VCAM1 and SPP1 as potential hematopoietic regulators (Lim et al. 2018). These are known pathways and factors which regulate hematopoietic niches in bone marrow (Ogawa 2002; Sugiyama et al. 2006; Ding et al. 2012; Corselli et al. 2013). Gene profiling of murine splenic stromal cell lines which support in vitro myelopoiesis (Periasamy et al. 2018; O’Neill et al. 2019) also identified perivascular reticular lineage cells with the capacity to undergo osteogenic differentiation in vitro (O’Neill et al. 2019). Some murine stromal lines were shown to support ectopic myelopoiesis in vivo following engraftment under the kidney capsule of mice (O’Neill et al. 2019).

Combined studies in mice and humans now reinforce the role of mesenchymal perivascular/perisinusoidal reticular cells in extramedullary hematopoiesis, and implicate these cells as essential niches for myelopoiesis in spleen.

Conclusions

Extramedullary hematopoiesis has been described for spleen although little knowledge exists for the role of spleen stromal cells as a niche supporting hematopoietic stem and progenitor cell differentiation. Here we describe a mesenchymal stromal cell type in the human spleen which supports hematopoiesis with the production of specific subsets of myeloid and dendritic cells unique to spleen. The human splenic stroma resembles murine stroma and represents a perivascular niche which supports myelopoiesis in spleen.

Abbreviations

APC: Antigen-presenting cells; BM: Bone marrow; cDC: Conventional dendritic cells; CMP: Common myeloid progenitors; DMEM: Dulbecco’s minimal essential medium; FMOC: Fluorescence minus one controls; HSC: Hematopoietic stem cells; HSPC: Hematopoietic stem progenitor cells; LTC: Long-term cultures; L-DC: Dendritic-like cells produced in LTC; MPP: Multipotential progenitors; OVA: Ovalbumin; pDC: Plasmacytoid dendritic cells; sDMEM: Supplemented DMEM

Funding

Open Access funding enabled and organized by CAUL and its Member Institutions This work was supported by project grants to H. C. O. from the Australian Research Council (#DP13010307) and the National Health and Medical Research Foundation of Australia (#585443). S. P. was supported by a postgraduate scholarship from the Royal Thai Government. V. T. and Y-Y. H. were supported by an Australian National University Postgraduate Scholarship. J. T. was supported by a C. J. Martin Fellowship from the National Health and Medical Research Foundation of Australia. The funders had no role in the study design, data collection and analysis, decision to publish or preparation of the manuscript.

Declarations

Competing interests

The authors declare no competing interests.

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