Human yolk sac-like haematopoiesis generates RUNX1- and GFI1/1B-dependent blood and SOX17-positive endothelium

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Summary statement: This study describes the hierarchical requirements for both RUNX1 and GFI1/1B during early human haematopoiesis arising from a yolk sac-like haemogenic endothelial intermediate.
ABSTRACT

The genetic regulatory network controlling early fate choices during human blood cell development are not well understood. We used human pluripotent stem cell reporter lines to track the development of endothelial and haematopoietic populations in an in vitro model of human yolk-sac development. We identified SOX17<sup>−</sup>CD34<sup>+</sup>CD43<sup>−</sup> endothelial cells at day 2 of blast colony development, as a haemangioblast-like branch point from which SOX17<sup>−</sup>CD34<sup>+</sup>CD43<sup>+</sup> blood cells and SOX17<sup>+</sup>CD34<sup>+</sup>CD43<sup>−</sup> endothelium subsequently arose. Most human blood cell development was dependent on RUNX1. Deletion of RUNX1 only permitted a single wave of yolk sac-like primitive erythropoiesis, but no yolk sac myelopoiesis or aorta-gonad-mesonephros (AGM)-like haematopoiesis. Blocking GFI1/1B activity with a small molecule inhibitor abrogated all blood cell development, even in cell lines with an intact RUNX1 gene. Together, our data defines the hierarchical requirements for both RUNX1 and GFI1/1B during early human haematopoiesis arising from a yolk sac-like SOX17-negative haemogenic endothelial intermediate.

INTRODUCTION

Blood cells develop from an endothelial intermediate at multiple stages during embryonic development in vertebrates (Dzierzak and Bigas, 2018; Ivanovs et al., 2017). Studies in the mouse have revealed that this is true for erythro-myeloid progenitors (EMPs) arising from yolk sac endothelium, and for blood cells emerging from aortic endothelium in the aorta-gonad-mesonephros (AGM) region (de Bruijn and Dzierzak, 2017; Frame et al., 2016; Okuda et al., 1996; Swiers et al., 2013; Wang et al., 1996). Similarities between mouse and human haematopoietic development (Dzierzak and Bigas, 2018; Ivanovs et al., 2017) suggest that the same regulatory genes critical for mouse haematopoietic development will play essential roles in blood formation from the human embryo. Since studies of genetically modified human blood cells in the context of a developing human embryo are not possible, haematopoietic differentiation of human pluripotent stem cells has emerged as the most tractable surrogate experimental system.
The methylcellulose-based blast colony forming cell (BL-CFC) assay has been used to interrogate formation of the earliest human blood cells that correspond to the first products of yolk sac haematopoiesis (Kennedy et al., 2007). In this assay, the BL-CFC proliferates and differentiates to form a core structure (D'Souza et al., 2005) that includes haemogenic endothelial cells, that in turn generate blood cells via an endothelial to haematopoietic transition (EHT) (Lancrin et al., 2009). Although haematopoiesis is orchestrated by numerous transcription factors, \textit{RUNX1} is a fundamental regulator of this process (Gao et al., 2018; Thambyrajah et al., 2016b). EMPs from the mouse yolk sac, and HSCs and preHSCs emerging from the mouse AGM all require \textit{Runx1}, whilst the earliest wave of yolk sac erythrocytes still appears in \textit{Runx1}-null mouse embryos (de Bruijn and Dzierzak, 2017; Frame et al., 2016; Okuda et al., 1996; Wang et al., 1996).

Mindful of the ambiguity that surrounds the use of terms ‘primitive’ and ‘definitive’ to describe waves of haematopoiesis, we have endeavoured to use nomenclature based on the embryonic site that blood cells are made (Ivanovs et al., 2017), in this manuscript. The term ‘extra embryonic’ is applied to blood cells similar to those developing in the yolk sac, that do not express \textit{HOXA} genes. This includes ‘primitive’ (erythroid, macrophage and megakaryocytic cells (McGrath et al., 2015a; McGrath et al., 2015b)) and ‘definitive’ waves of yolk sac haematopoiesis (EMPs and yolk sac derived lymphoid cells (McGrath et al., 2015a; Yoshimoto et al., 2011; Yoshimoto et al., 2012)). ‘Intra embryonic’ is applied to blood cells similar to those that develop in the AGM, that express \textit{HOXA} genes in stem cells and progenitors, and include the first repopulating HSCs, their precursors and myeloid and lymphoid progeny (Ivanovs et al., 2017). This is also called ‘definitive’ haematopoiesis in the literature.

In this study, we tracked the emergence of vascular and haematopoietic lineages using a human pluripotent stem cell (hPSC) line in which mCHERRY and GFP report expression of \textit{SOX17} in endothelium and the \textit{RUNX1C} isoform of \textit{RUNX1} in haematopoietic progenitors (Ng et al., 2016). By modelling extra embryonic haematopoiesis in the blast colony assay, we showed that differentiating \textit{SOX17} endothelial-like cells acted like a haemangioblast population, because they gave rise to the majority of blood cells and also to a \textit{SOX17}⁺ endothelium. Blood formation was
**RESULTS.**

To facilitate the dissection of early human haematopoiesis, we utilised a dual reporter hPSC line, \( \text{SOX}^{\text{mCHERRY/w}} \text{RUNX}^{\text{C}}_{\text{GFP/w}} \) (hereafter denoted SOX-RUNX) (Ng et al., 2016), in which GFP targeted to \( \text{RUNX}^{\text{C}} \) marks hematopoietic progenitor cells (Corada et al., 2013; Sroczynska et al., 2009), and mCHERRY targeted to \( \text{SOX}^{17} \) marks vascular endothelium (Burtscher et al., 2012; Challen and Goodell, 2010; Clarke et al., 2013).

**Modelling extra embryonic, yolk sac-like haematopoiesis**

SOX-RUNX cells were differentiated to haematopoietic mesoderm, dissociated and transferred into methylcellulose cultures for blast colony (BL-CFC) assays (Fig. 1A and Materials and Methods). Day 2 (d2) mesoderm cells expressed the mesendodermal marker PDGFR\( \alpha \) (92.5±1.7%, \( n=5 \)) (here and hereafter representing mean ± S.E.M. and the number of independent experiments) (Fig. 1B), from which BL-CFCs derive (Davis et al., 2008), whilst few cells expressed the \( \text{SOX}^{17} \)-mCHERRY reporter (hereafter denoted \( \text{SOX}^{17} \)) (7.2±1.4%, \( n=5 \)) and \( \text{RUNX}^{1C} \)-GFP expressing cells (hereafter denoted \( \text{RUNX}^{1C} \)) were not present (Fig. 1C). There was a 4-fold higher frequency in BL-CFCs generated from d2 compared with d3 differentiated mesoderm cells (Fig. 1D).
Differentiated blast colonies comprised a mix of SOX17-positive and -negative adherent cells, and RUNX1C-positive and -negative haematopoietic cells (Fig. 1E). Tracking haematopoietic and endothelium development revealed that most cells expressed vascular endothelial growth factor receptor 2 (KDR) within the first day of methylcellulose culture, rapidly followed by expression of the haematovascular markers TEK (TIE2), CD31 and CD34 (Fig. 1F). SOX17 (11.4±3.1%, n=5) and CD43 (16.5±3.6%, n=5) expression appeared after two days of methylcellulose differentiation, delineating subsets of endothelial (SOX17+/CD34+/CD43−) and haematopoietic (CD43+) cells. RUNX1C expression (10.6±6.1%, n=5) was observed in one third (33.6±9.8%, n=5) of CD43+ cells (RUNX1C+/CD34+/CD43+) a day later, in a population largely excluding SOX17+ cells (Fig. 1C,F,G). As expected, most RUNX1C+ cells (76±13.3%, n=5) expressed CD43 (Fig. 1G). A small percentage of SOX17+ cells at d3 also expressed CD43 (10.2±3.8%, n=5) implying a modest haemogenic capacity for these cells (Fig. 1G). In summary, these data demonstrated expression of SOX17 in extra embryonic, yolk sac-like endothelium, which appeared at the same time as CD43+ haematopoietic cells, and prior to RUNX1C expression in a subset of the nascent blood.

SOX17-negative endothelium is the major source of the first human blood cells

In order to determine the relationship between the different cell types during differentiation, we flow sorted d2 blast colonies after 3 days in methylcellulose into endothelial (SOX17+/CD34+/CD43− and SOX17−/CD34+/CD43−, hereafter denoted d3 SOX17+/ENDO and d3 SOX17−ENDO) and haematopoietic (CD34+/CD43+ and CD34−/CD43+) populations (Fig. 1H and Fig. S1A). Re-culturing the haematopoietic populations yielded predominantly GYPAn erythroid cells (76.1±4.1%, n=3) from the more mature CD34+/CD43− fraction whilst the CD34+/CD43+ fraction yielded more RUNX1C+ non-erythroid (myeloid and megakaryocytic) cells (65.7±6.8%, n=3) (Fig. 1I and Fig. S1B), suggesting an erythroid bias in the earliest blood cells produced from blast colonies. The d3 SOX17+/ENDO cells generated few CD43+ cells (12.8±8.6%, n=3) (Fig. 1I), while the d3 SOX17−ENDO population generated a very high proportion of CD43+ blood cells (87.8±0.9%, n=3) and infrequent SOX17+ endothelial cells (7.3±1.1%, n=3) (Fig. 1I).
Both d3 SOX17\(^{+}\)ENDO and d3 SOX17\(^{-}\)ENDO cells formed endothelial-like networks in Matrigel\(^{TM}\) within 24 hours (Fig. 1J,K). The SOX17\(^{+}\) cultures retained expression of the mCHERRY reporter, whilst most of the SOX17\(^{-}\) cells remained mCHERRY negative, suggesting that the allocation of cells to a SOX17\(^{+}\) fate was largely complete by d3 in methylcellulose (Fig. 1J,K). Continuing the cultures for a further 24 hours in the presence of stem cell factor generated many haematopoietic cells from SOX17\(^{-}\)ENDO cells but only infrequent foci of blood cells in SOX17\(^{+}\)ENDO cultures (Fig 1L,M), consistent with the results of re-culture experiments (Fig. 1I).

The frequency of haematopoietic progeny from SOX17\(^{-}\)ENDO cells, sorted after 2 days of methylcellulose culture, was determined by limit dilution analysis and compared to the haemogenic frequency of SOX17\(^{-}\)ENDO and SOX17\(^{+}\)ENDO cells sorted from d3 of methylcellulose differentiation (Fig. 1H,N and Supplementary Table 1). The high frequency of haemogenic SOX17\(^{-}\)ENDO cells isolated at d2 or d3 of methylcellulose culture (both 1:11), contrasted with the low frequency (1:2480) observed in d3 SOX17\(^{+}\)ENDO cells (Fig. 1N and Supplementary Table 1).

In order to explore the developmental relationship between SOX17\(^{-}\) and SOX17\(^{+}\) endothelium, we performed live cell imaging over 65 hours of d2 SOX17\(^{-}\)ENDO cells sorted from methylcellulose cultures and re-plated into a Matrigel\(^{TM}\) endothelial network assay (Supplementary Movie 1, Fig. 2 and Fig. S2). It can be seen from time lapse images taken at 10 minute intervals (Supplementary Movie 1 and Fig. 2C) that the endothelia were initially SOX17\(^{-}\), and that individual, SOX17\(^{-}\)cells began to acquire expression of the SOX17 reporter after 6 hours of observation. Importantly, mCHERRY expression was acquired by 24-28 hours, and during this period there was little increase in cell numbers, precluding division of any rare contaminating SOX17\(^{+}\) cells present at the onset of the culture as the reason for the increase in SOX17\(^{+}\) cell numbers (Fig. 2D). After this period the number of blood cells rapidly increased and the number of SOX17\(^{+}\) cells decreased a little and stabilised. Similar kinetics of mCHERRY reporter expression were observed in a second experiment that was not subjected to time lapse imaging (Fig. S2). These data strongly support the premise that d3 SOX17\(^{-}\)ENDO derives from the same d2 SOX17\(^{-}\)ENDO precursor population that also exhibits high haemogenic activity (Fig. 1N).
Since we had observed that a small percentage of SOX17\(^+\) ENDO cells developed into CD43\(^+\) haematopoietic cells (Fig. 1G), we compared them to SOX17\(^-\) ENDO derived CD43\(^+\) cells. We sorted haematopoietic and endothelial cells at d3 of methylcellulose culture that were either SOX17\(^+\) or SOX17\(^-\), confirming that the SOX17\(^+\) cells expressed higher levels of mCHERRY and of SOX17 and providing reassurance that the appearance of SOX17\(^+\)CD43\(^+\) cells was not the consequence of imperfect sorting (Fig. S3A,B). The expression of haematopoietic genes (RUNX1, GFI1, SPI1[PU.1], KLF1 and GATA1) was higher in the d3 SOX17\(^-\) ENDO endothelial cells, probably reflecting their greater haemogenic capacity (Fig. S3B). The CD34\(^+\)CD43\(^+\) derivatives of both SOX17\(^+\) and SOX17\(^-\) endothelium expressed similar levels of haematopoietic transcription factors and globin genes, and displayed similar morphology, frequency and distribution of colony forming cells (Fig. S3B-G). These data suggest that, in human extra embryonic, yolk sac-like cultures, blood cells predominantly derive from SOX17\(^-\) endothelium, with a very small proportion of phenotypically similar cells arising from precursors that are SOX17\(^+\).

**Identification of distinct endothelial and haematopoietic subsets of differentiating blast colonies**

We compared the transcriptional profiles of d2 PDGFR\(\alpha\)^+ mesoderm, d2 and d3 methylcellulose SOX17\(^-\) ENDO cells, d3 methylcellulose SOX17\(^+\) ENDO cells, and d3 methylcellulose CD34\(^+\)CD43\(^+\) and CD34\(^-\)CD43\(^+\) haematopoietic populations (Fig. 3A-C and Supplementary Table 2).

Mesodermal cells, and endothelial/haematopoietic samples clustered into separate cell populations (Fig. 3D). Differential gene expression analysis revealed that up regulated genes in mesoderm were enriched for Gene Ontology terms associated with embryo development (GO:0009790) and gastrulation (GO:0007369) whilst the endothelial/haematopoietic populations were enriched for leukocyte (GO:0050900) or vascular related genes (GO:0001944) (Supplementary Table 3). While very few genes were differentially expressed between d2 and d3 SOX17\(^-\) ENDO cells, several thousand genes were differentially expressed between CD34\(^+\)CD43\(^+\) and CD34\(^-\)CD43\(^+\) haematopoietic cells and d2 SOX17\(^-\) ENDO cells (Fig. S4A,B and Supplementary Table 4).
Patterns of differentially expressed genes between haematopoietic and endothelial cells were also consistent with cells segregating into distinct SOX17-expressing endothelium or CD34^CD43^ haematopoietic fates during blast colony differentiation (Fig. S4C). Specifically, 914/1062 (86.1%) genes up regulated in d3 SOX17^ENDO were down regulated in CD34^CD43^ blood cells, and 711/954 (74.5%) of genes down regulated in d3 SOX17^ENDO were up regulated in CD34^CD43^ blood cells (Fig. S4D and Supplementary Table 4). Similarly, 851/1028 (82.8%) genes up regulated in CD34^CD43^ were down regulated in d3 SOX17^ENDO cells and 1647/1898 (86.8%) of genes down regulated in CD34^CD43^ were up regulated in d3 SOX17^ENDO cells (Fig. S4E and Supplementary Table 4). The data may be summarised to state that the same genes up regulated during the transition from d2 SOX17^ENDO to d3 SOX17^ENDO are down regulated in the transition to CD34^CD43^, and vice-versa. These analyses argue for the presence of a binary ‘switch’ active in the d2 SOX17^ENDO cells that will lead to either a haematopoietic or endothelial fate, and are consistent with a haemangioblast-like function of these cells.

Examining specific genes expressed in the sorted populations, there was selective expression of cell surface proteins (including CDH1, EPCAM, PDGFRA) and transcription factors (including POU5F1, MIXL1, EOMES, T) associated with the primitive streak stage of development (Davis et al., 2008; Hirst et al., 2006) in the mesoderm cells (Fig. 3E,F). Consistent with our previous findings (Ng et al., 2016), HOXB, but not HOXA, genes were up regulated in these extra embryonic, yolk sac-like endothelial populations (Fig. S4F).

Two surface markers identifying blast colony forming cells, KDR (Kennedy et al., 2007) and APLNR (Yu et al., 2012), were expressed in the mesoderm and in their endothelial progeny (Fig. 3E). There was a high concordance in the expression of endothelial cell surface genes (including CD34, PECAM1, CDH5, FLT1, ESAM, EFNB2, CD93) and transcription factors (including SOX7, SOX18, ERG, ETS1, ETS2, HOPX, FLI1) in d2 and d3 SOX17^ and d3 SOX17^ENDO samples (Fig. 3E,F). In addition to up regulated SOX17 expression, we observed reduced expression of cell cycle genes and the proliferation-related transcription factors MYB and MYC in the d3 SOX17^ENDO cells, suggesting that these cells were more quiescent, possibly mediated by higher
levels of NOTCH signalling (Mack and Iruela-Arispe, 2018) (Fig. 3F and Fig. S4F). Expression of a number of genes distinguished the CD43⁺ haematopoietic fractions from their endothelial counterparts, including the surface expressed \textit{SPN} \[CD43\], \textit{ITA2B} \[CD41\], \textit{ITB3} \[CD61\], \textit{CD33} and the transcription factors \textit{GATA1}, \textit{KLF1} and \textit{SPI1} \[\textit{PU.1}\] (Fig. 3E,F). The acquisition of CD43 expression was also associated with a simultaneous down regulation of endothelial cell surface markers (including \textit{APLR}, \textit{CDH5}, \textit{FLT1}, \textit{ESAM}, \textit{EFNB2}, \textit{CD93}) and transcription factors (including \textit{SOX7}, \textit{SOX17}, \textit{SOX18}, \textit{ERG}, \textit{ETS1}, \textit{ETS2}, \textit{HOPX}) (Fig. 3E,F).

Notably, there was variation in expression between \textit{RUNX1} and \textit{GFI1B} in the endothelial populations (Fig. 3F). Higher levels of \textit{RUNX1} and \textit{GFI1B} expression in the d2 and d3 \textit{SOX17⁺ENDO} cells correlated with a high capacity to form haematopoietic cells, whilst low levels of \textit{RUNX1} and \textit{GFI1B} in d3 \textit{SOX17⁺ENDO} marked a largely non-haemogenic endothelium. In order to explore the role of these factors in dictating haemogenic capacity, we characterised differentiation in cell lines in which they were deleted or inhibited.

\textbf{RUNX1 is required for blast colony development}

To examine whether \textit{RUNX1} is a key driver of the EHT in human extra embryonic, yolk sac-like haematopoiesis, we generated \textit{RUNX1-null} hPSCs (denoted \textit{RUNX1-KO}) by excising a portion of the DNA binding domain of \textit{RUNX1} in SOX-RUNX cells (see Materials and Methods and Fig. S5A-C,G). In blast colony assays, \textit{RUNX1-KO} cultures formed vascular cores that expressed the \textit{SOX17} reporter, but did not obviously generate haematopoietic cells (Fig. 4A-E). Flow cytometry analyses confirmed a failure of CD43⁺ blood cells to increase in \textit{RUNX1-KO} cultures (19.9±3.1%, SOX-RUNX; 2.9±0.3%, \textit{RUNX1-KO}; n=4, P<0.01, Student’s t-test) at d2 of methylcellulose culture (Fig. 4F). By d5 (Fig. 4G and Fig. S5E) and d8 (Fig. S5D,E) of methylcellulose culture, more striking reductions in CD43⁺, \textit{GYPA}⁺ and \textit{RUNX1C}⁺ cells were noted in \textit{RUNX1-KO} compared to SOX-RUNX cultures. Gene expression studies confirmed the down regulation of \textit{RUNX1}, \textit{RUNX1C} and \textit{GFI1B}, with similarly reduced expression of erythroid lineage genes (\textit{GATA1}, \textit{KLF1}, \textit{ε-GLOBIN} and \textit{γ-GLOBIN}) and increased expression of endothelial \textit{SOX17} and \textit{CD34} (Fig. 4H and Fig. S5F).
**Primitive erythroid cell generation in RUNX1-KO cells is GFI1-dependent**

The presence of abundant nucleated erythroid cells in Runx1-knock out mouse embryos at E12.5 (Okuda et al., 1996; Wang et al., 1996) argues that the initial wave of yolk sac erythroid differentiation remains intact, although we were initially unable to detect expansion of CD43\(^+\) blood cells in human RUNX1-KO cells after d7 (Fig. 4I,J), or colony-forming cells in the d7 RUNX1-KO differentiation cultures (Fig. 4K).

We then added FGF2 and a low concentration of CHIR from the onset of differentiation (Fig. 5A), because previous studies have shown that WNT agonists synergise with BMP4 to promote differentiation of haematopoietic mesoderm (Gertow et al., 2013), and that early haematopoietic colonies are FGF2-dependent (Choi et al., 2012; Yu et al., 2012). These modifications, combined with culturing EBs at an air-liquid interface, led to the appearance of haemoglobinised cells, more prominent in the RUNX1-KO cultures, after d15 of differentiation (Fig. 5A,B). There were more CD43\(^+\) cells in both SOX-RUNX and RUNX1-KO cultures from d5-d7 under these conditions, with most cells co-expressing GYPA (compare Fig. 5C,D to Fig. 4I).

After d7, an increasing proportion of SOX-RUNX cells expressed CD43, often associated with RUNX1C, and down regulated GYPA. In contrast, no new CD43\(^+\) cells appeared in the RUNX1-KO cultures. These cells down regulated CD43, but retained high levels of GYPA expression, consistent with adoption of an erythroid fate (Fig. 5C,E). These data indicate that RUNX1 is not required for the generation of the first CD43-expressing cells that subsequently differentiate only to erythroid cells. This is consistent with observations in Runx1-null embryos and differentiated Runx1-null mouse ES cells, in which all myeloid cells are absent (Lacaud et al., 2002; Okuda et al., 1996; Wang et al., 1996). We compared the frequency of colony forming cells in differentiating SOX-RUNX and RUNX1-KO cultures. In SOX-RUNX cells, BL-CFC peaked at d2 of differentiation, as noted previously (Fig. 1D), followed by a wave of primarily erythroid colonies at d6-d7 (Fig. 5F). However, in RUNX1-KO cultures, the only clonogenic cells detected were at d2, when a small
number of erythroid colonies were observed arising from vascular cores when cells were cultured at high density in methylcellulose (Fig. 5F,G).

Supporting evidence for the erythroid restriction of \textit{RUNX1-KO} haematopoiesis was provided by RNA seq analysis at d6 of differentiation. At this early time point there were only 17 differentially expressed genes between the parental SOX-RUNX and the \textit{RUNX1-KO} cultures (Supplementary Tables 5 and 7). Notably, 5 of the 10 genes downregulated in the d6 \textit{RUNX1-KO} cell line are expressed in megakaryocytes, myeloid cells or B cells (\textit{HDC, GCSAML, PLEK, RGS18, TSPAN33}) consistent with the hypothesis that differentiation to non-erythroid lineages is compromised. Conversely, 3 of 7 genes with increased expression in \textit{RUNX1-KO} cells are expressed in erythroid cells (\textit{SLC4A1, HBA2, HEMGN}) arguing for a complementary increase in erythroid differentiation, as indeed was observed under these modified differentiation conditions.

We confirmed the critical role of \textit{RUNX1} in specifying intra embryonic, AGM-like haematopoiesis (Chen et al., 2009b; North et al., 2002), demonstrating that CD43\textsuperscript{+} blood cells were not generated in \textit{RUNX1-KO} cultures, and colony formation was not observed, when cultures were differentiated under conditions that facilitated the emergence of AGM-like blood lineages (Ng et al., 2016) (Fig. 5H).

In order to determine the optimum differentiation day at which to transfer embryoid bodies to air-liquid interface cultures for erythroid development, transfers at d2, d3 and d4 were compared (Fig. 6A). Haemoglobinised clusters were most evident in \textit{RUNX1-KO} cells cultured on air-liquid interface from d4, with lesser amounts in SOX-RUNX cultures (Fig. 6A). Flow cytometry confirmed the higher percentage of GYPA expressing cells in the \textit{RUNX1-KO} cultures, and demonstrated that RUNX1C and CD43 expression was confined to the SOX-RUNX cultures (Fig. 6B,C). Analysis of globin expression revealed a higher ratio of \(\zeta/\alpha\) and \(\epsilon/\gamma\) chains in differentiated \textit{RUNX1-KO} cells, indicating the developmentally earlier phenotype of the erythroid cells in these cultures (Fig. 6D). Mirroring the pattern of \textit{RUNX1C} expression, PCR analysis of \textit{SPI1} \([\text{PU.1}]\) revealed higher expression in the SOX-RUNX cultures (Fig. 6E).
RUNX1 and SPI1 are expressed in erythroid progenitor cells, but their levels decline with erythroid maturation. The decline in RUNX1 is the major driver of the reduction in SPI1 expression (Willcockson et al., 2019). Both RUNX1 and SPI1 repress key erythroid transcription factors, exemplified by KLF1 (Kuvardina et al., 2015), and the enforced expression of either gene prevents terminal erythroid differentiation (Willcockson et al., 2019). We hypothesise that in the absence of RUNX1 to drive myeloid and megakaryocytic differentiation, SPI1 levels also eventually fall, and the wave of GYPAC43+ cells seen at d7 (Fig. 5C) defaults entirely to erythroid lineage differentiation. This may explain the greater quantity of primitive erythroid cells generated from RUNX1-KO cells.

The Runx1-target genes Gfi1 and Gfi1b are expressed in haemogenic endothelia in both the E11.5 dorsal aorta and in the E9.5 yolk sac in the mouse (Thambyrajah et al., 2016b). Gfi1/1b form multi-protein complexes that include the co-repressor, Rcor1 (CoREST), the histone demethylase, Kmd1a (Lsd1), and the histone deacetylases, Hdac1/2 (Saleque et al., 2007). It is argued that Runx1 expression in haemogenic endothelium induces Gfi1/1b, which bind to regulatory sequences of endothelial genes, and recruit the CoREST complex that then silences the endothelial program (Lancrin et al., 2012; Thambyrajah et al., 2016b). Inhibition of Lsd1 has been shown to phenocopy the endothelial to haematopoietic transition block observed with deletion of Gfi1/1b (Thambyrajah et al., 2016a). When inhibitors of LSD1 (denoted LSDi) were included in blast colony assays, all blood cell formation was lost, and only cores of endothelial and stromal cells remained, similar to findings with RUNX1-KO cultures (compare Fig. 6F-I to Fig. 4A-E). The endothelium generated under LSDi conditions in the blast colony assay was very similar to endothelium produced by the RUNX1-KO cells and the SOX-RUNX parental cell line. After 10 days differentiation in methylcellulose, there were very few endothelial cells (CD34+CD31*KDR*KDR*KDR*KDR*KDR*) in the SOX-RUNX cultures (<1%) and these were >95% SOX17+ (Fig. S6A,B). In the RUNX1-KO and the LSDi treated SOX-RUNX cultures, approximately 30-40% of the viable cells were SOX17+ endothelial cells (Fig. S6A,B).
However, LSDi-treated cultures, in contrast to *RUNX1*-KO lines, failed to generate or maintain CD43$^+$ or GYP A$^+$ cells (Fig. 6J). PCR analysis indicated that levels of *RUNX1* transcripts were lower in cultures treated with LSDi (compare SOX-RUNX with and without LSDi in Fig. 6K and Fig. S5G), suggesting that *GFI1/1B* may be a regulator, as well as a target, of *RUNX1*. Other haematopoietic transcription factors (*GATA1*, *KLF1*, *SPI1* [*PU.1]*) were also significantly down regulated by LSD1 inhibition (Fig. 6L), consistent with the endothelial to haematopoietic transition block. These differences in transcription factor expression in LSDi-treated cultures were seen from d4 of differentiation, antedating the emergence of CD43$^+$ cells, and therefore excluding major differences in the cellular composition of the cultures as an explanation for this observation. Expression of these genes was similar between SOX-RUNX and *RUNX1*-KO cells, consistent with a primary role of Runx1 in reorganisation, rather than transcriptional regulation, of lineage-specific transcription factor assemblies (Lichtinger et al., 2012; Thambyrajah et al., 2016b) (Fig. 6L). RNA seq analysis in d4 and d6 LSDi treated cultures indicated a down regulation in expression of erythroid lineage genes, associated with Gene Ontology terms oxygen transport (GO:0015671) and haemoglobin complex (GO:0005833) (Fig. 6M, Fig. S5H and Fig. S6C,D, and Supplementary Tables 5 and 6). Genes increased in expression in LSDi most notably included those responsive to the HDAC inhibitor panobinostat (Fig. 6M, Fig. S6C,D and Supplementary Tables 5 and 6). This is consistent with the inhibition of the HDAC containing CoREST complex by LSDi, and with the similar effects on murine haematopoiesis observed between Lsd1 and Hdac inhibition (Thambyrajah et al., 2018).

The human blast colony assay detects predominantly *RUNX1*-dependent haematopoiesis

*RUNX1* is required for the formation of virtually all haematopoietic cells detected in the blast colony assay (Fig. 4), suggesting that this primarily reads out progenitor cells similar to mouse yolk sac EMPs, which are also *Runx1* dependent (Tober et al., 2013). One prediction of this hypothesis is that blast colonies should generate granulocytes, a lineage not observed during the first wave of extra embryonic yolk sac blood formation (Palis et al., 1999). To test this, we differentiated blast colony forming cells in the presence of growth factors that preferentially support erythroid,
macrophage or granulocytic cells (Fig. S7A). May-Grünwald-Giemsa stained cytospin preparations documented that erythroid cells were restricted to cultures supplemented with EPO, macrophages were dependent upon M-CSF, and maturing neutrophils and eosinophils dominated cultures supplemented with G-CSF and GM-CSF (Fig. S7B). These lineage assignments were supported by flow cytometry, showing expression of GYPA on erythroid cells, CD14 expression on macrophages and RUNX1C in granulocytes (Fig. S7C,D). Finally, PCR analysis confirmed globin, KLF1 and GATA1 expression in erythroid cells, CSF1R and SPI1 in macrophages, and EPX, SPI1 and GATA1 in granulocytes (Fig. S7E). Taken together, these data support our hypothesis that the human blast colony assay reads out RUNX1-dependent extra embryonic, yolk sac-like cells with a broad myeloid potential similar to mouse yolk sac EMPs.

**DISCUSSION**

We have modelled extra embryonic human haematopoiesis and dissected the role of the transcription factor RUNX1 in analyses facilitated by the use of a reporter line in which GFP reported cells expressing the haematopoietic specific, C isoform, of RUNX1 and mCHERRY, expressed from the SOX17 locus, marked endothelium. We showed that d2 SOX17- ENDO cells that expressed RUNX1 and GFI1/1B functioned as a haemangioblast-like population. Continued expression of RUNX1 and GFI1/1B was correlated with the emergence of extra embryonic, yolk sac-like blood cells, while the up regulation of SOX17 expression and extinction of RUNX1 and GFI1/1B led to the development of SOX17+ endothelial cells. This complementary expression of RUNX1 and SOX17 in blood and endothelial cells, captured in the RNA seq analysis shown in Figure 3, is reminiscent of the expression pattern of these transcription factors in the human intra embryonic AGM, where endothelial cells transitioning to form intra-aortic haematopoietic clusters expressed lower levels of SOX17 and increasing RUNX1 (Bos et al., 2015). Our data suggests that a similar reciprocal regulation of RUNX1 and SOX17 expression occurs in our model of human extra embryonic yolk sac-like haematopoiesis.
Indeed, the d2 and d3 SOX17 ENDO cells expressed similar key haematopoietic genes to haemogenic endothelia in the mouse and human intra embryonic AGM (Baron et al., 2018; Ng et al., 2016; Solaimani Kartalaei et al., 2015; Swiers et al., 2013), although HOXA expression was absent, as expected, from these extra embryonic, yolk sac-like cells. Our study indicated that the d2 SOX17 ENDO population was the precursor of both CD34+CD43+ haematopoietic cells and a distinct SOX17+ expressing endothelium, although we have not shown that one cell could give rise to both progeny (Fig. 7A). Our data are consistent with prior in vitro mouse ESC differentiation studies that found a transient population of Tie2hi c-Kit+CD41− endothelial cells at d2 of blast colony differentiation that gave rise to CD41+ haematopoietic progeny (Lancrin et al., 2009). Indeed, RNA seq analysis shows that the d2 SOX17 ENDO may be analogous to these mouse cells, in that they also expressed TEK (TIE2) and KIT and were negative for ITGA2B (CD41) (Fig. 3E). Our work extends the mouse findings by demonstrating that this d2 SOX17 ENDO not only gave rise to blood cells, but also gave rise to a largely non haemogenic endothelium, now marked by the acquisition of SOX17 and the loss of RUNX1 expression.

Confirming the requirement for RUNX1 in the haemogenic SOX17- endothelium, we showed that deletion of RUNX1 abrogated all haematopoiesis save for a single wave of extra embryonic erythropoiesis (Fig. 7B). The results of these experiments can be taken to indicate that the blast colony forming assay is not dominated by precursors of the first wave of extra embryonic haematopoiesis, but predominantly reads out progenitors of a second, RUNX1-dependent wave of extra embryonic haematopoiesis, perhaps analogous to a human EMP. This interpretation is supported by our ability to differentiate blast colonies to granulocytic cells, a lineage similarly generated from mouse yolk sac EMPs (Frame et al., 2013). Furthermore, our experiments also confirmed that all human extra embryonic, yolk sac-like macrophages were absolutely RUNX1 dependent, consistent with reports in the mouse (Lacaud et al., 2002; Okuda et al., 1996; Wang et al., 1996).
We observed that loss of RUNX1 lead to reduced levels of GFI1/1B as reported in the mouse (Lancrin et al., 2012; Thambyrajah et al., 2016b), but that expression was not completely lost (Fig. 4H and Fig. S5F). Exploring the role that residual GFI1/1B protein expression might play in the regulation of extra embryonic erythropoiesis, we documented the complete suppression of blood formation by small molecule inhibition of the histone demethylase, LSD1. In the context of haematopoietic differentiation, LSD1 mediates the inhibitory effects of GFI1/1B on the endothelial program and Lsd1 deficiency phenocopies the developmental block in endothelial to haematopoietic transition in Gfi1/1b null mouse embryos (Saleque et al., 2007; Thambyrajah et al., 2016a) (Fig. 7C). Consistent with this action, CD41+ cells in the yolk sacs of double knock out Gfi1/1b mouse embryos maintained endothelial gene expression, preventing their free migration and distribution into embryonic tissues outside the yolk sac. Although haematopoietic colonies formed in vitro from disaggregated yolk sacs, clonogenic cells were not found in the dorsal aorta and intra-arterial haematopoietic cluster formation was lost (Lancrin et al., 2012; Thambyrajah et al., 2016a). In our human pluripotent stem cell model, the complete loss of haematopoiesis in the presence of LSD1 inhibition suggests that the human extra embryonic, yolk sac-like culture modelled in our hPSC in vitro differentiation may be more similar to the mouse AGM, rather than yolk sac haemogenic endothelium. However, an important caveat remains that the effects of the chemical inhibitor may differ from the results that would be obtained from a double knock-out of GFI1/1B genes.

Our data argue for a window of haematopoietic competence during blast colony differentiation in which expression of GFI1/1B reinforced by RUNX1 drives the generation of blood and suppresses an endothelial program (Fig. 7). Such a model fits well with the narrow developmental window during which enforced Runx1 expression in mouse embryonic endothelium is able to drive haematopoiesis (Yzaguirre et al., 2018). The factors that initiate RUNX1 expression during this permissive stage are not known.
In summary, we identified and characterised a population of SOX17- haemogenic endothelial cells that is the dominant source of blood and of SOX17+ endothelium in human extra embryonic, yolk sac-like cultures. We correlated RUNX1 and GFI1/1B expression with increased haemogenic capacity, further identifying that RUNX1 is required for human blast colony development. Finally, our studies also revealed the critical role played by GFI1/1B in the emergence of the first erythroid cells and the absolute dependence of all other blood lineages on RUNX1.

MATERIALS and METHODS

Ethics

Human pluripotent stem cell studies were approved by the Monash University (reference 2002/225MC) and The Royal Children’s Hospital (reference 33001A) Human Research Ethics Committees.

Generation and validation of targeted SOX17mCHERRY/w RUNX1C-GFP/w and SOX17mCHERRY/w RUNX1-/- H9 hPSCs lines

The SOX17mCHERRY/w H9, SOX17mCHERRY/w RUNX1C-GFP/w (SOX-RUNX) H9 hPSC lines have been described previously (Loh et al., 2014; Ng et al., 2016).

To generate the SOX17mCHERRY/w RUNX1-/- (RUNX1-KO) hPSC line, the CRISPR Design Tool (http://tools.genome-engineering.org) was utilised to design 18 nucleotide (nt) single-guide RNAs (sgRNAs) to target two sites within exon 4 of the RUNX1 gene (5’ TGTCGCCGTCTGGTAGGA 3’ (CRISPR SITE 1) and 5’ GGTCGGTCTTCCTAGCTT 3’ (CRISPR SITE 2)) (Fig. S5A)(Ran et al., 2013). The corresponding 18 nt sgDNAs were synthesised, phosphorylated and annealed to complementary sequences to make double stranded (ds) DNA oligos. The oligos were modified to incorporate a 5’CACC and 3’CAAA overhang for BbsI site recognition within the pSpCas9-2A(BB)-GFP (PX458) vector, a gift from Feng Zhang (Addgene plasmid #48138), and 5’G-C base pair (bp) insertion to enable U6 transcription. The pSpCas9-2A-GFP vectors expressing the RUNX1 CRISPR 1 and 2 dsRNAs were electroporated into SOX-RUNX hPSCs. Following 24-72 hours, single GFP-positive cells were flow sorted and clonally expanded on pre-gelatinised 96-well plates.
Clones were screened for the acquisition of a 457bp deletion region of exon 4 of the \textit{RUNX1} locus containing part of the DNA binding domain using primers a and c (Fig. S5A,C). Positive clones were further selected for deletion verification by PCR using primers a and b and Sanger sequenced using primers a and c. Three positive clones (11, 14, 30) contained a complete deletion between CRISPR site 1 and CRISPR site 2.

For all lines, surface markers of undifferentiated hPSCs were expressed and genomic integrity was confirmed by Illumina HumanCytoSNP-12 v2.1 array.

\section*{Culture and differentiation of hPSCs}

H9 hPSCs used in these studies were provided by the WiCell Research Institute. Cell lines were regularly tested to exclude mycoplasma contamination and confirm genomic integrity. Culture and enzymatic passaging of hPSCs lines was performed as previously reported (Ng et al., 2008b).

For the generation of blast colonies and for initial differentiation towards extra embryonic, yolk sac-type haematopoietic cells, hPSCs were differentiated using the spin EB method in APEL medium (Ng et al., 2008a) supplemented for the first 2-3 days with 20ng/ml recombinant human (rh) bone morphogenetic protein 4 (BMP4, R&D Systems), 30ng/ml rh vascular endothelial growth factor (VEGF, PeproTech), 40ng/ml rh stem cell factor (SCF, PeproTech) and 20ng/ml rh ACTIVIN A (R&D Systems).

In later differentiations towards extra embryonic, yolk sac-type haematopoietic cells, hPSCs were differentiated as spin EBs in APEL medium supplemented for the first 4 days with 20ng/ml rh BMP4, 25-30ng/ml rh VEGF, 25-40ng/ml rh SCF, 10-20ng/ml rh ACTIVIN A, 10ng/ml rh fibroblast growth factor-2 (FGF2, PeproTech) and 0.5-1μM CHIR99021 (CHIR, Tocris Biosciences). Where indicated, 1μM GSK-LSD1 (lysine specific demethylase 1) inhibitor (Sigma Aldrich) was added to day2 EBs. For air-liquid interphase cultures 30-60 EBs were transferred in 50μl of growth factor reduced (GFR)-Matrigel™ (BD Pharmingen) onto transwells at days 2, 3 and 4 and supplemented with the medium described above. Differentiation medium was changed after 4 days to APEL medium including 50ng/ml rh SCF, 50ng/ml rh VEGF, 20-25ng/ml rh insulin like growth factor 2 (IGF2, PeproTech), 10ng/ml rh FGF2 and, where indicated, 1μM GSK-LSD1 inhibitor. At 6-7 days of differentiation 20-30 EBs for submerged liquid culture were transferred onto gelatinised or GFR-
Matrigel™ treated (8mg/ml) 6-well plates in APEL medium supplemented with 50ng/ml rh SCF, 50ng/ml rh VEGF, 50ng/ml rh interleukin (IL)-3 (PeproTech), 25ng/ml rh thrombopoietin (TPO, PeproTech), 25ng/ml rh FLT3 receptor ligand (FLT3L, PeproTech), 10ng/ml rh FGF2 and 2-3u/ml rh erythropoietin (EPO, PeproTech). Air-liquid interphase EBs were cultured in APEL medium containing 100ng/ml rh SCF, 100ng/ml rh TPO, 25ng/ml rh VEGF, 25ng/ml rh FLT3L, 25ng/ml rh interleukin (IL)-6 (PeproTech) and 10ng/ml rh IGF2. Medium was changed every 5-7 days for submerged liquid cultures, or every 1-2 days for air-liquid interphase cultures.

For intra embryonic haematopoietic differentiations, cells were differentiated using the spin EB method in APEL medium supplemented for the first 4 days with 20ng/ml rh BMP4, 25ng/ml rh VEGF, 25ng/ml rh SCF and 10ng/ml rh ACTIVIN A, 10ng/ml rh FGF2 and 0.5-1µM CHIR. Between days 2-4, 3µM CHIR99021 and 3-4µM SB431542 (SB, Cayman Chemicals) were added to pattern mesoderm. Medium was changed after 4 days to APEL medium supplemented with 20ng/ml rh BMP4, 50ng/ml rh SCF, 50ng/ml rh VEGF, 10-20ng/ml rh IGF2 and 10ng/ml rh FGF2. After 6-8 days of differentiation, EBs were transferred onto GFR-Matrigel™ treated 6-well plates or air-liquid interphase transwells at 20-60 EBs/well in APEL medium supplemented with 100ng/ml rh SCF, 25-100ng/ml rh FLT3L, 25-50ng/ml rh VEGF, 50-100ng/ml rh TPO, 25ng/ml rh IL-6, 10ng/ml rh FGF2 and 10ng/ml rh BMP4. Medium was changed every 3-5 days thereafter.

For analysis, EBs, liquid, air-liquid interphase and methylcellulose cultures were harvested and dissociated into single cell suspension using TrypLE Select (Invitrogen) or Collagenases Type I and IV (Worthington) and passed through a 21-23-gauge needle and 40µm cell strainer.

**Colonies forming assays and culture of sorted cells**

Blast colony forming cells (BL-CFCs) were identified by culturing 3 x 10³-1 x 10⁴ dissociated cells, or 3 x 10⁴ cells for high density BL-CFC assays, from day 2 or 3 EBs in a formulation designated MC-APEL (1% methylcellulose in APEL medium) or serum-free MethoCult™ (StemCell Technologies) supplemented with 100ng/ml rh SCF, 50ng/ml rh VEGF, 50ng/ml rh IL-3, 50ng/ml rh IL-6, 50ng/ml rh TPO, 20ng/ml rh BMP4, 10ng/ml rh FGF2 and 2u/ml rh EPO. Where indicated, methylcellulose cultures were also supplemented with one of the following inhibitors or cytokines: 1mM GSK-LSD1, 50ng/ml rh macrophage colony-stimulating factor (M-CSF, PeproTech), 50ng/ml
rh granulocyte colony-stimulating factor (G-CSF, R&D Systems) and 50ng/ml rh granulocyte-macrophage colony-stimulating factor (GM-CSF, R&D Systems). Colony formation was scored between 7 to 10 days of differentiation. Following 3 to 7 days of blast colony formation, unsorted or cell sorted populations were cultured on GFR-Matrigel™ coated plates in APEL medium supplemented with 100ng/ml rh SCF, 50ng/ml rh VEGF, 50ng/ml rh IL-3, 50ng/ml rh IL-6, 50ng/ml rh TPO, 20ng/ml rh BMP4, 10ng/ml rh FGF2, 2u/ml rh EPO and where indicated 50ng/ml rh M-CSF, 50ng/ml rh G-CSF and 50ng/ml rh GM-CSF for 3 to 7 days. Later appearing haematopoietic colonies were identified by culturing 1-3 x 10^4 dissociated cells from day 7-25 cultures in a formulation designated MC-APEL or serum-free MethoCult™ supplemented with 100ng/ml rh SCF, 50ng/ml rh VEGF, 50ng/ml rh IL-3, 50ng/ml rh IL-6, 50ng/ml rh TPO, 50ng/ml rh G-CSF, 10ng/ml rh FGF2, 2u/ml rh EPO and 10μg/ml human low-density lipoproteins (LDL, Stem Cell Technologies). Colony formation was scored between 10-15 days of differentiation.

**Limit dilution estimation of frequency of haematopoietic precursor frequency**

To determine the clonal frequency of haematopoietic precursors, day 2+2 and day 2+3 blast colonies were flow sorted on the basis of CD34, CD43 and SOX17 expression and cells deposited by flow cytometer at 1, 3, 10, 30, 100 and 300 cells per well into GFR-matrigel coated 96-well plates. After 5-10 days of culture in APEL medium supplemented with 100ng/ml rh SCF, 50ng/ml rh VEGF, 50ng/ml rh IL-3, 50ng/ml rh IL-6, 50ng/ml rh TPO, 20ng/ml rh BMP4, 10ng/ml rh FGF2 and 2u/ml rh EPO, wells were scored by microscopy for the presence of hematopoietic clusters of greater than 30 cells. The frequency of colony forming cells was estimated using Poisson statistics.

**Endothelial network assay and time lapse imaging**

GFR-Matrigel™ was solidified at 37°C for 30 minutes in wells of a 48-well plate. For the experiments shown in Fig.1, 5 x 10^4, day 2+3 flow sorted SOX17^ENDO or SOX17^-ENDO, cells were seeded onto polymerised GFR-Matrigel™ in APEL medium supplemented with 50ng/ml rh VEGF, 10ng/ml rh FGF2, 5ng/ml rh epidermal growth factor (EGF, PeproTech) and 10^3 Hydrocortisone (StemCell Technologies) and incubated for 24 hours. After 48 hours, APEL medium was supplemented with 100ng/ml rh SCF. For the experiments shown in Fig. 2, Fig. S2
and **Supplementary Movie 1**, 1.2 x 10^5, day 2+1.75 flow sorted SOX17^{ENDO} (SOX17^{CD34^{+}CD43^{-}CD73^{-}}) cells were seeded onto polymerised GFR-Matrigel™ in APEL medium supplemented with 50ng/ml rh VEGF, 100ng/ml rh SCF, 10ng/ml rh FGF2, 5ng/ml rh EGF and 10^{-3} Hydrocortisone. For the experiment in **Fig. S2**, cultures were incubated in a 5%CO_{2} incubator at 37°C and wells were imaged at 24 hours and 48 hours after the endothelial network assay was setup. For the experiment in **Fig. 2**, cultures were incubated in a 5%CO_{2} incubator at 37°C for 6 hours and then placed in an environmentally controlled (37°C, 5% CO_{2} in humidified air) chamber fitted to a Zeiss LSM 780 laser scanning confocal microscope for time lapse imaging.

**Flow cytometry and cell sorting**

Antibodies directed against the following cell surface antigens (fluorochrome, manufacturer, catalogue number, clone (where known), dilution) were used to stain dissociated cells for flow cytometric analysis: CD14 (phycoerythrin-pe-cy7, BioLegend #301814, clone M5E2, 1:50), CD31 (allophycocyanin (apc), BioLegend #303115, clone WM59, 1:50; brilliant violet (bv)-421, BioLegend #303123, clone WM59, 1:30) CD34 (pe-cy7, BioLegend #43515, clone 581, 1:100), CD43 (apc, BioLegend #343206, clone 10G7, 1:50; bv-421, BD Pharmingen #562916, clone 1G10, 1:30), CD45 (bv-421, BioLegend #304032, clone H130, 1:30), CD73 (apc, BioLegend #344006, clone AD2, 1:50), Glycophorin A (GYPA) (apc, BD Pharmingen #551336, clone GA-R2(HIR2), 1:2000), platelet-derived growth factor receptor alpha (PDGFRα) (BD Pharmingen #556001, clone aR1, 1:100), vascular endothelial growth factor receptor 2 (VEGFR2/KDR) (alexa fluor (af)-647, BioLegend #338909, clone HKDR-1, 1:10) and TIE2/TEK (BD Pharmingen #557039, clone 33, 1:100). PDGFRα- and TIE2-unconjugated antibodies were detected with secondary antibodies conjugated with apc (BD Pharmingen #550826, 1:100; BioLegend #405308, clone Poly4053, 1:100) or pe-cy7 (Biolegend #405315, clone Poly4053, 1:100). Flow cytometric analysis was performed using a BD LSR Fortessa™ analyser. Flow sorting used a BD Biosciences Influx™ or BD Biosciences FACSAria™ Fusion cell sorter.

Samples were gated using FSC-A and FSC-H to exclude doublets. In some cases, FSC-W, SSC-A and SSC-H were also used. FSC and Propidium iodide exclusion were used to select live cells. Positive gates for markers of interest were determined by comparing stained samples with those in
which the antibodies were not added. Frequently this could also be corroborated by gating on samples in which the marker under evaluation was not expressed. There are numerous flow cytometry plots in this manuscript. The example chosen to illustrate gating strategy is one of the experiments from which samples were sorted for RNA sequencing analysis. This is shown in Fig. S1A.

**Gene expression analysis**

Total RNA was isolated from hPSCs using the RNA Isolate II Mini or Micro Kits (Bioline) or RNeasy Kit (Qiagen) as specified by the manufacturer. cDNA was reverse transcribed via random hexamer priming and Tetro cDNA synthesis (Bioline) or Superscript III (Invitrogen) kits in accordance with the manufacturers’ instructions. TaqMan™ gene expression probes (Applied Biosystems) and Bioline reagents were used for quantitative real-time PCR using GAPDH as the reference gene to normalise the data.

TaqMan™ assays directed towards the following target sequences were used to detect gene expression: CD34 (Hs00156373_m1), CSFR1 (Hs00234622_m1), EPX (Hs00946094_m1) GAPDH (Hs99999905_m1), GATA1 (Hs00231112_m1), GATA2 (Hs00231119_m1), GFI1 (Hs00382207_m1), GFI1B (Hs01062469_m1), HAEMOGLOBIN-ALPHA (Hs00361191_g1), HAEMOGLOBIN-BETA (Hs00747223_g1), HAEMOGLOBIN-EPSILON (Hs00362215_g1), HAEMOGLOBIN-GAMMA (Hs00361131_g1), HAEMOGLOBIN-ZETA (Hs00923579_m1), KLF1 (Hs00610592_m1), mCHERRY (Custom design by ThermoFisher), RUNX1 (Hs00231079_m1), RUNX1C (Hs01021967_m1), SOX17 (Hs00751752_s1), SPI1 (Hs00231368_m1) and SPN [CD43] (Hs01872322_s1).

**Transcriptional profiling using RNA sequencing**

In the first series of experiments (series 1), differentiated SOX-RUNX (SOX17mCHERRYwRUNX1CGFPw) cell cultures were harvested from EBs at day 2 and from methylcellulose cultures at day 2+2 and day 2+3 of culture, and flow sorted based on their expression of PDGFRα, CD34, CD43 and mCHERRY using a BD Biosciences Influx™ or BD Biosciences FACSAna™ Fusion cell sorter. In a second series of experiments (series 2), cultures
of SOX-RUNX and RUNX1-KO (SOX17<sup>mCHERRY</sup>/wRUNX1<sup>-/-</sup>) cells differentiated for 4 or 6 days in the presence or absence of 1μM GSK-LSD1 inhibitor were harvested without additional sorting. Total RNA was purified from samples (RNA Isolate II Micro Kit, Bioline) and RNA concentration was determined using a Nanodrop 2000 analyser (Thermo Scientific). Total RNA from the differentiated fractions was sequenced at the Australian Genome Research Facility (series 1) or the Murdoch Children’s Research Institute (series 2). In total, 16 samples from series 1 (2-4 independent experiments for each sample) and 24 samples from series 2 (3 independent experiments for each sample) were sequenced. The STAR aligner (v2.4.0h1 or v2.5.2a) in 2 pass mode was used to map single end 100bp reads to the human reference genome (hg38) for series 1, and 75bp paired end reads to the human reference genome (hg39) for series 2 (Dobin et al., 2013). Uniquely mapped reads were summarised using featureCounts (v1.4.6) using Gencode Release 19 comprehensive annotation (Liao et al., 2014). Genes lowly expressed were excluded (less than 10 counts per million in fewer than two samples in series 1, and less than 1 count per million in fewer than three samples in series 2). The data was TMM normalised, voom transformed and differential gene expression was assessed using moderated t-tests from the R Bioconductor limma package (Ritchie et al., 2015; Robinson and Oshlack, 2010). Genes that had a false-discovery rate of less than 5% were called as significantly differentially expressed for the various comparisons of interest. Gene ontology analysis was performed using the ToppGene Suite (Chen et al., 2009a).

Confocal microscopy and image processing

Confocal images (Fig. 1E, J-M; Fig. 2C; Fig. 4A-D; Fig. 5G; Fig. 6F-H; Fig. S2B, C and Supplementary Movie 1) were acquired as single optical sections on a Zeiss LSM 780 laser scanning confocal microscope running Zeiss Black software (Carl Zeiss Pty. Ltd.). Overlay images were created in Adobe Photoshop 2020 release 21.1.0 (Adobe) and figures prepared in Adobe Illustrator 2020 release 24.1 (Adobe). The only image manipulations performed were adjustments of brightness and contrast. The time lapse images, counting of SOX17-mCHERRY fluorescent cells and counting the brightfield cells in the endothelial tube assay for Fig. 2C and Supplementary Movie 1 were performed using Fiji (ImageJ) open source image processing software v2.0.0-rc-69/1.52p (https://imagej.nih.gov/).
Statistical analysis

Experiments were analysed using GraphPad Prism versions 5–8 (GraphPad Software Inc.) and Microsoft Excel for Mac version 13.36 (Microsoft Corporation). Tests for statistical significance are listed with each experiment and included two sided Student’s t-test for paired analyses, multiple t-tests with Sidak-Bonferroni post test, or ANOVA for experiments with multiple comparisons of one or more grouped variables, accompanied by the post tests (Dunnett, Tukey or Holmes-Sidak) indicated as appropriate by the software. No statistical method was used to predetermine sample size.
COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

AUTHORSHIP CONTRIBUTIONS

F.F.B., E.S.N. and A.G.E. designed and performed experiments and interpreted data. A.M., K.V., K.S., R.M., P.M. and L.A. designed and performed experiments. A.R.L., N.M.D. and A.O. performed bioinformatics analysis of the RNA sequencing data. F.F.B., E.G.S. and A.G.E. wrote the manuscript, which all authors edited and approved.

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DATA AVAILABILITY

The RNA sequencing data in this paper have been deposited with Gene Expression Omnibus under accession number GSE124086 (incorporating accession numbers GSE124084 and GSE124085). All figures, with the exception of Figure 7 (i.e., Figs. 1-6 and Figs. S1-S7), are supported by raw data. Source data for figures is available from the corresponding author upon reasonable request.

MATERIALS & CORRESPONDENCE

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Figure 1. SOX17-positive and SOX17-negative endothelium generate haematopoietic cells.

(A) Blast colony differentiation protocol. BL-CFC, blast colony forming cell; MC, methylcellulose; LC, liquid culture. (B) Flow cytometry profile demonstrating that most d2 differentiated cells express PDGFRα (n=5 experiments). Mean±SEM provided in text. The data is shown as a contour plot to aid in the visualisation of the small numbers of events captured. This was also done in...
panels C and G. (C) Flow cytometric analysis of SOX17 and RUNX1C expression in d2 developing blast colonies following one (+d1) to three (+d3) days of methylcellulose (MC) culture (n=5 experiments). Mean±SEM provided in text. (D) Frequency of blast colonies was greater in d2 compared to d3 differentiated cells (mean±SEM, d2, n=4; d3, n=3 experiments). * P=0.018 using Student’s t-test. (E) Brightfield (BF) and merged fluorescent (RUNX1C, SOX17) image of a d6 blast colony with SOX17⁺ endothelial cells (black arrowhead), RUNX1C⁺ (hollow white arrowhead) and RUNX1C⁻ blood cells (white arrowhead). Scale bar, 100µm. (F) Flow cytometry timecourse of differentiating blast colonies from d1 (2+1) to d3 (2+3) in MC culture showing expression of endothelial (KDR, CD31, CD34, TEK, SOX17) and haematopoietic (CD43, RUNX1C) markers (n=3-5 experiments per marker). Mean±SEM provided in text. (G) At d3 of differentiation, RUNX1C identifies CD34⁺CD43⁺ blood cells and SOX17 marks CD34⁺ endothelial cells and some SOX17⁺CD34⁻CD43⁻ haematopoietic cells (n=5 experiments). Mean±SEM provided in text. (H) Sorting strategy to isolate endothelial (SOX17⁺ENDO, SOX17⁻ENDO) and haematopoietic (34⁺43⁺ or 34⁺43⁻) populations from d2 blast colonies at d2 (d2+2) and d3 (d2+3) of methylcellulose (MC) culture (n=8 experiments at d2+2; n=15 experiments at d2+3). (I) Analysis of sorted fractions showing CD34, CD43, GYPA expression after five days of liquid culture (d2+3+5). Data for d2+3+3 and d2+3+7 time points is shown in Fig. S1B (n=3 experiments). (J-M) Images of d2+3 SOX17-sorted endothelial tube networks after 24 (J, K) and 48 hours of culture (L, M), highlighting abundant blood cell formation from SOX17⁻ENDO compared to SOX17⁺ENDO sorted cells (white arrowhead). Scale bar (J, K) 50µm and (L, M) 100µm (n=2 experiments). (n) Limit dilution analyses of d2+2 and d2+3 blast colony sorted fractions, correlating input cell number in each well with the frequency of wells where haematopoietic clones were absent (% negative wells). The frequency of haemogenic and progenitor cells in each population is shown, calculated as the cell input number that gave 37% negative wells using Poisson statistics. Regression lines for d2+2 and d2+3 SOX17⁻ENDO overlie each other. Data represents the sum of three experiments. See Supplementary Table 1.
Figure 2. SOX17-negative endothelium gives rise to SOX17-positive endothelium. (A) Experimental outline. After two days differentiation as spin EBs, cells were harvested and plated in MC in a BL-CFC assay. After 1.75 days in MC, SOX17- ENDO cells were sorted and plated in a Matrigel™ endothelial tube assay. Six hours after seeding cells, time lapse imaging of cells commenced in a humidified chamber at 37°C with 5%CO2 in air. BL-CFC, blast colony forming cell; MC, methylcellulose. (B) Gating strategy outlining the sorting of SOX17-ENDO cells from MC.
at d2+d1.75. The two left-hand panels show the expression of SOX17 and of CD34 at this time point. The right-hand panels show the sequential gating strategy used to sort the SOX17^ENDO (SOX17^CD34^CD43^CD73^-) endothelial cells that were seeded into the endothelial tube assay. (C) Images taken from the time lapse series at illustrative time points from 0 to 3600 min as indicated on each image. Boxed areas and matching fields from intermediate time points are shown at higher magnification. The first faintly red fluorescent SOX17^+ cells are seen from 360 min (6 hr) and are obvious from 720 min (12 hr) onwards. Blood cells appear from 2160 min (36 hr). Scale bar, 100 µm. (D) Graph plotting the number of fluorescent SOX17 cells in the imaged field at hourly intervals, showing a rapid emergence of SOX17^+ cells from 360 min to 1440 min, before a reduction and plateauing of numbers. The field area was 0.9mm^2. (E) Bar graph detailing the number of SOX17^- and SOX17^+ cells and the number of morphologically classified haematopoietic cells at the onset, and at 24 hr and 48 hr after the onset of time lapse imaging. Note that there is little increase in cell number over the initial 24 hr although there is a rapid rise in the number of SOX17^+ cells. See also Supplementary Movie 1 and Fig. S2.
Figure 3. Transcriptional profiling reveals discrete endothelial and haematopoietic populations. (A) Flow sorted fractions (shown in black text) collected for RNA seq analysis. D2 cells sorted from EBs: PDGFRα⁺ mesoderm (MES, grey). D2 MES cells cultured for 2 days in methylcellulose (2+2 MC): SOX17⁻ENDO[CD34⁺CD43⁻] (d2S⁻E, orange). D2 MES cells cultured for 3 days in methylcellulose (2+3MC): SOX17⁺ENDO[CD34⁺CD43⁻] (d3S⁺E, red), SOX17⁻ENDO[CD34⁺CD43⁻] (d3S⁻E, cyan), 34⁺43⁻ immature blood (blue), and 34⁻43⁺ mature blood (green) populations. (B, C) Multidimensional scaling plots of sorted populations showing (B) mesoderm separation from endothelial and haematopoietic populations and (C) excluding...
mesoderm to demonstrate clustering of haematopoietic (34^43^, 34^43^-), SOX17^ENDO (d2S^E, d3S^E) and SOX17^ENDO (d3S^E) fractions. (D) Unsupervised hierarchical clustering of RNA seq data showing that the top 100 differentially expressed genes define mesoderm and haematopoietic/endothelial populations. Genes included in the two clusters are shown. Scale, row Z-score. See also Supplementary Table 3. (E, F) Heatmaps comparing selected genes differentially expressed between sorted populations categorised as (E) cell surface proteins and (F) transcription factors. Scale, log_2 reads per kilobase per million (RPKM). See also Fig. S4 and Supplementary Table 4 for details of differentially expressed genes.
Figure 4. Blast colony development requires RUNX1. (A-D) Blast colony cultures after six days in methylcellulose (d2+6MC), showing (A, C) blast colonies generated from SOX-RUNX cultures and (B, D) SOX17-expressing vascular cores from RUNX1-KO cultures. Scale bar, 100μm (n=9 experiments) (E) Blast colonies were observed only from SOX-RUNX (SR) cultures and endothelial cores only from RUNX1-KO (R1-KO) cultures (mean ± SEM, n=3 experiments). * P<0.02 compared to RUNX1-KO using Student’s t-test. (F,G) Flow cytometry plots of d2 (d2+2MC) and d5 (d2+5MC) SOX-RUNX and RUNX1-KO methylcellulose cultures illustrating the absence of CD43+ blood cells in RUNX1-KO cultures (n=4 experiments at d2+2 MC; n=3 experiments at d2+5
MC). Mean±SEM provided in text. Day 8 (d2+8MC) plots shown in Fig. S5D. (H) Relative gene expression (negative delta (-d) Ct) for selected haematopoietic, genes in d2+5 MC RUNX1-KO cultures compared to SOX-RUNX control cultures (mean±SEM, n=4 experiments). * P<0.03, ** P<0.002, **** P<0.0001 compared to SOX-RUNX, two-way ANOVA, Holm Sidak’s multiple comparisons test. (I) Analysis of mesoderm (PDGFRα, KDR), haematopoietic (CD34, CD43) and reporter (SOX17, RUNX1C) genes at d4, d7 and d12 of differentiation of SOX-RUNX and RUNX1-KO cultures (mean±SEM for n=6 experiments at d4; n=4 experiments at d7, d12). ** P<0.002, **** P<0.0001 compared to SOX-RUNX using two-way ANOVA, Holm Sidak’s multiple comparisons test. (J) Flow cytometry plots illustrating the absence of CD43+ and RUNX1C+ blood cells in RUNX1-KO at d12 compared to the SOX-RUNX control (n=4 experiments). (K) No d7 haematopoietic colonies were observed in RUNX1-KO methylcellulose cultures (mean±SEM, n=3 experiments). * P=0.048 compared to SOX-RUNX control using Students t-test.
Figure 5. *RUNX1*-knockout cultures exclusively generate primitive erythroid lineages. (A) Air-liquid interphase differentiation protocol. (B) *RUNX1*-KO EBs grown on air-liquid interphase (ALI) were more haemoglobinised than SOX-RUNX controls by d20 (d8+12 ALI). Scale bars, left 1000 µm, right 400 µm (n=4 experiments). (C) *RUNX1*-KO cultures maintained GYP43+ erythroid cells but failed to develop *RUNX1C*+CD43+ myeloid cells (n=2-4 experiments per time point) (D) Similar early blood cell differentiation in *RUNX1*-KO and SOX-RUNX cultures, with differences
observed at day 7 (mean±SEM, n=2-4 experiments per time point). ** P<0.01, *** P<0.001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test. (E) SOX-RUNX cultures developed more CD43⁺ myeloid cells than RUNX1-KO cells (mean±SEM, n=2-4 experiments per time point). * P<0.02, ** P<0.01, *** P<0.001, **** P<0.0001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test. (F) Clonogenic frequency peaked at d2 in SOX-RUNX methylcellulose cultures, while RUNX1-KO cultures generated only rare erythroid colonies at d2. Upper and lower panels represent results from two separate series of experiments (mean±SEM, n=2-6 experiments per time point). ** P<0.01, **** P<0.0001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test. (G) Haematopoietic colonies arising from d2 EBs in SOX-RUNX cultures included RUNX1C⁺ myeloid cells, while the rare RUNX1-KO colonies comprised only erythroid cells emerging from a vascular core. The far right panel displays the same RUNX1-KO colony imaged at d8 in methylcellulose (+d8 MC) and after +d16 MC to demonstrate haemoglobinisation (n=6 experiments). (H) Intra embryonic SB CHIR (S/C) differentiated SOX-RUNX cultures consistently generated CD43⁺ blood cells and colonies in methylcellulose, in comparison to RUNX1-KO. Upper and lower panels represent results the same experimental series (mean±SEM, n=2-4 experiments per time point). **** P<0.0001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test.
Figure 6. LSD1 inhibition blocks extra embryonic erythroid differentiation from RUNX1-KO cultures. (A) Day 4 EBs cultured for 14 days on air-liquid interphase (ALI) (d4+14 ALI). RUNX1-KO cultures were more extensively haemoglobinised compared to SOX-RUNX controls (n=3 experiments). Scale bars, left 1000 µm, right 400 µm. (B) RUNX1-KO EBs cultured on air-liquid interphase for 14 to 16 days retained a higher numbers of GYP A⁺ erythroid cells compared to the
SOX-RUNX controls (n=3 experiments). (C) Analysis of the experiments in (B) illustrating the predominance of RUNX1C⁺ and CD43⁺ cells in SOX-RUNX cultures (mean±SEM, n=3 experiments). *** P<0.001 and **** P<0.0001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test. (D, E) Gene expression analysis of experiments shown in (B, C) highlighting (D) higher ratios of ζ/α and ε/γ globin in RUNX1-KO cultures, whilst (E) SOX-RUNX cultures expressed higher levels of the myeloid gene SPI1 (mean±SEM, n=3 experiments). *** P<0.001 and **** P<0.0001 using 2-way ANOVA, Holm Sidak’s multiple comparisons test. (F-H) Images of (F) d2+9 haematopoietic SOX-RUNX colonies and similar vascular core colonies in (G) RUNX1-KO and (H) SOX-RUNX LSD1 inhibitor (LSDi) treated cultures. Scale bars, 50μM (n=3 experiments). (I) Blast colonies (colonies) were confined to SOX-RUNX cultures and vascular cores (cores) in RUNX1-KO and LSDi 1μM (1) and 3μM (3) supplemented SOX-RUNX cultures (mean±SEM, n=3 experiments). * P<0.022, ** P<0.0037 compared to RUNX1-KO cores using Student’s t-test. (J) d3-d7 LSDi treated SOX-RUNX EBs fail to generate CD43⁺ or GYPA⁺ cells (mean±SEM, n=1 experiment for d3; n=3-4 for d4-7). * P<0.05, ** P<0.01, *** P<0.001, **** P<0.0001 compared to SOX-RUNX cells, using 2-way ANOVA, Dunnett’s multiple comparisons test. (K-L) Gene expression analysis of the experiments in (J) showing (K) RUNX1 reduction in LSDi treated cultures and loss in RUNX1-KO cultures and (L) LSDi down regulation of GATA1, KLF1 and SPI1 (mean±SEM, n= 1 experiment for d3, n=2 experiments d7, n=3-4 experiments for d4–d6). * P<0.05, ** P<0.01, *** P<0.001, **** P<0.0001 using 2-way ANOVA, Dunnett’s multiple comparisons test. (M) Heatmaps of selected RNA seq data from d4 and d6 SOX-RUNX (SR), RUNX1-KO (R1KO) cultures showing down regulation of gene expression with LSDi treatment. Scale, log₂ reads per kilobase per million (RPKM). See also Fig. S6D.
Figure 7. Model of endothelial and haematopoietic differentiation in human blast colonies.

(A-C) Proposed differentiation pathway to blood and endothelium in blast colonies, illustrating the roles of RUNX1, GFI1/1B and LSD1 inhibition (LSDi). In the blood lineages, RUNX1^+ cells are shown in green whilst more mature myeloid cells that are RUNX1^- are shown in blue. Erythroid cells are shown transitioning to red. Loss of gene expression or of blood lineages are shown in grey. Reduced levels of GFI1/1B in RUNX1-KO differentiations are shown as increased transparency of text. (A) Regulation of haematopoietic development within SOX-RUNX cultures, highlighting the postulated roles of RUNX1 and GFI1/1B in regulating the pathways to blood and SOX17^- endothelium. Both haematopoietic and SOX17^+ endothelial cells derive from the SOX17^- endothelial precursor, but we have not shown that both populations can arise from one cell. (B) In RUNX1-KO cells, RUNX1^+ cells and mature myeloid cells are absent, and GFI1/1B expression levels are reduced. (C) All blood cells are absent in cultures in which LSDi also blocks the functions of GFI1/1B.
SUPPLEMENTARY TABLES

Table S1. Haematopoietic precursor frequency in blast colony sorted populations.

Click here to Download Table S1

Table S2. RNA-Sequencing analysis of sorted d2 mesoderm and haematopoietic blast colony populations from d2 cultures following d2 (d2+2) or d3 (d2+3) in methylcellulose.

Click here to Download Table S2

Table S3. Gene ontology terms for mesoderm and haematopoietic blast colony populations.

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Table S4. Differentially expressed genes between sorted d2 mesoderm and haematopoietic blast colony populations from d2 cultures after d2 (d2+2) or d3 (d2+3) in methylcellulose.

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Table S5. RNA-Sequencing analysis of control and LSD1 inhibitor treated d4 and d6 SOX-RUNX and RUNX1-KO cultures.

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Table S6. Differentially expressed genes between control and LSD1 inhibitor treated d4 and d6 SOX-RUNX cultures.

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Table S7. Differentially expressed genes between d6 SOX-RUNX and d6 RUNX1-KO cultures.

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Movie 1. Time lapse series of endothelial network assay. Series of images taken at 10 min intervals from 0 to 3920 min (65 hours) as indicated on each image. Images run at 10fps. The first faintly red fluorescent SOX17+ cells are seen from 360 min (6 hours) and are obvious from 720 min (12 hours) onwards. Blood cells appear from 2160 min (36 hours). Scale bar, 100µm. Related to Fig. 2.
Figure S1. Haematopoietic development from SOX-RUNX blast colonies. (A) Example of flow cytometry gating strategy for data shown in Fig. 1H. Top panel of plots describes single and viable cell isolation process, lower panels illustrate unstained controls, SOX17+ and SOX17- cell separation and subsequent surface marker population isolation. (B) Flow cytometric analysis of CD34, CD43, GYP A expression in sorted fractions of d2 blast colonies after 3 days in methylcellulose (d2+3) (see Fig. 1H) following three (d2+3+3) and seven (d2+3+7) days of re-culture. By 7d, d3 SOX17-ENDO fractions remain primarily as endothelium with a few GYP A-CD34- erythroid cells, while the d3 SOX17-ENDO fractions contain RUNX1-CD34+ myeloid and GYP A-CD34+ erythroid cells. The greatest proportion of erythroid cells arises from the first blood cells to emerge, 34-43+ (n=3 experiments). Mean±SEM provided in text for selected markers. Data for d2+3+5 shown in Fig. 1I.
Figure S2. Endothelial network assay showing generation of SOX17-positive from SOX17-negative endothelium. (A) Gating strategy outlining the sorting of SOX17-ENDO cells from methylcellulose at d2+d1.75. The two left-hand panels show the expression of SOX17 and of CD34 at this time point. The right-hand panels show the sequential gating strategy used to sort the SOX17- ENDO (SOX17-CD34⁺CD43⁻CD73⁻) endothelial cells that were seeded into the endothelial network assay. This is an independent experiment to that shown in Fig. 2. (B, C) Images taken from the endothelial network assay at (B) 24 hr and (C) 48 hr time points. SOX17⁺ and SOX17⁻ adherent cells are seen as well as developing clusters of hematopoietic cells. This experiment shows more rapid differentiation than the time lapse experiment in Fig. 2, suggesting that the environmental control achieved in a sealed incubator (this experiment) is superior to differentiation in the controlled climate (37°C, 5% CO₂ in humidified air) chamber used for the time lapse series. The number of SOX17⁺ endothelial cells in each image is indicated. The number of tightly packed hematopoietic cells precluded accurate assessment of the total cell number. Scale bar, 100µm.
Figure S3. (A) Sorting strategy for d2+3 SOX17+ and SOX17- blood (34+43+) and endothelial (ENDO [34+43+]) fractions (n=4 experiments). (B) PCR analysis of d2+3 sorted fractions from (A) correlating SOX17-mCHERRY reporter expression with SOX17, and expression of blood genes (RUNX1, GFI1, SPI1, KLF1, GATA1) in the SOX17+ compared to SOX17- populations (mean±SEM, n=4 experiments). * P=0.048; **** P<0.0001, 2-way ANOVA, Tukey’s multiple comparisons test. (C) SOX17*34*43+ and SOX17*34*43- sorted fractions after d9 of methylcellulose re-culture (d2+3+9) displayed similar haematopoietic colony morphology. Scale bar, 50 µm (n=4 experiments). (D) Clonogenic frequency was similar between the SOX17- and SOX17+ blood (34+43+) fractions (mean±SEM, n=4 experiments). (E) Analysis of sorted fractions after d11 of methylcellulose re-culture (d2+3+11) illustrated similar RUNX1C, CD45 and GYPA expression. Plots are representative of four experiments. (F) Relative gene expression (shown as negative delta (-d) Ct) analysis of haematopoietic (GATA1, SPI1) and globin genes (ZETA, ALPHA, EPSILON, GAMMA, BETA) from d2+3+11 methylcellulose colonies in (D) (mean±SEM, n=4 experiments). (G) Ratio of embryonic globin genes showing a similar ratio of ζ/α and ε/γ indicating a predominance of primitive globins (mean±SEM, n=4 experiments).
Figure S4. SOX17-negative endothelium represents a divergent point between endothelial and blood commitment. (A) Summary of differentially expressed genes between indicated mesoderm (MES, grey), d2 SOX17-ENDO (d2S\(^E\), orange), d3 SOX17-ENDO (d3S\(^E\), red), d3 SOX17-ENDO (d3S\(^E\), cyan) and haematopoietic (34\(^+\)43\(^+\), dark blue; 34\(^+\)43\(^-\), green) sorted fractions. See Fig. 3A for flow-sorting strategy. Genes are listed in Supplementary Table 4. (B) Venn diagram showing up regulated genes comparing d2S\(^E\) to indicated sorted fractions. Genes are listed in Supplementary Table 4. (C) RNA-Seq heatmap of sorted endothelial and haematopoietic populations showing differentially expressed genes compared to the d2S\(^E\) population. In the upper two panels, genes up or down regulated in 34\(^+\)43\(^+\) blood cells tend to be reciprocally expressed in all endothelial populations. In the lower two panels, genes up or down regulated in d3S\(^E\) or were reciprocally expressed in haematopoietic fractions. Thus, the d2S\(^E\) population includes genes in common with both the d3S\(^E\) and 34\(^+\)43\(^+\) populations, Scale, log\(_2\) RPKM. Genes are listed in Supplementary Table 4. (D, E) Venn diagrams illustrating differentially regulated genes in d3S\(^E\) compared to d2S\(^E\) or 34\(^+\)43\(^+\) (D) and 34\(^+\)43\(^+\) compared to d2S\(^E\) or d3S\(^E\) (E). Genes are listed in Supplementary Table 4. (F) RNA-Seq heatmap of sorted fractions showing gene expression of HOX, cell cycle and NOTCH pathway genes. No reads were detected for members of the HOX clusters that are not shown. Scale, log\(_2\) RPKM.
Figure S5. Impaired haematopoiesis in RUNX1-KO cells (A) Schematic representation of the RUNX1C locus showing wild-type and GFP targeted alleles, and sites for CRISPR deletion to create the RUNX1-KO cell line. Exons, e1 to e7, screening primers (a-c), proximal (P) and distal (D) promoters and DNA binding domain (grey) are indicated. (B-C) PCR screening confirmed the homozygous CRISPR deleted region in the RUNX1-KO (KO), compared to SOX-RUNX (SR). PCR product lengths are indicated and screening primer locations are shown in (A). (D) Flow cytometry plots of d2 cells after eight days of methylcellulose (d2+8MC) culture of SOX-RUNX and RUNX1-KO cells illustrate the absence of CD43+ blood cells in RUNX1-KO (n=4 experiments). See also Fig. 4F,G. (E) Summary of flow cytometry analysis, comparing haematopoietic gene expression in SOX-RUNX and RUNX1-KO cells after 5 (d2+5) and 8 (d2+8) days in methylcellulose (d2+5, mean±SEM, n=3; d2+8, mean±SEM, n=4 experiments). * P=0.017, **** P<0.0001 compared to SOX-RUNX, 2-way ANOVA, Holm Sidak’s multiple comparisons test. (F) Relative gene expression, displayed as negative delta Ct (-d Ct), for selected haematopoietic genes in d2+8 methylcellulose RUNX1-KO cultures compared to SOX-RUNX controls, illustrating down regulation of haematopoietic gene expression in RUNX1-KO cultures (mean±SEM, n=4 experiments). * P<0.05, ** P<0.001, *** P<0.0002, **** P<0.0001 compared to SOX-RUNX, 2-way ANOVA, Holm Sidak’s multiple comparisons test. See also Fig. 4H. (G) Integrative Genomics Viewer window showing RNA-Seq reads over the RUNX1 locus in d4 and d6 SOX-RUNX and RUNX1-KO lines with and without treatment with the LSD1 inhibitor (LSDi). Note the absence of transcripts down stream of the deleted exon 4 in the RUNX1-KO lanes but retention of normal levels of transcripts up stream of the deletion. It is clear that LSD1 inhibition reduces RUNX1 transcripts from both the distal and proximal promoters. The CRISPR deleted region in exon 4 (e4) is shown and marked with an arrow in each lane. (n=3 experiments). (H) Integrative Genomics Viewer window showing RNA-Seq reads over the HBE1 and HBZ loci. The levels of embryonic globin transcripts are increased in the RUNX1-KO lanes compared to SOX-RUNX lanes, but transcripts are extinguished following the addition of LSD1 inhibitor.
Figure S6. RNA-Seq of LSD1 inhibitor-treated cultures. (A) Flow cytometry plots of day 10 methylcellulose cultures (d2+10MC) of SOX-RUNX, RUNX1-KO and LSD1 inhibitor treated SOX-RUNX cells, showing that similar endothelium is generated by both the SOX-RUNX and RUNX1-KO cell lines. The proportion of 34^43^-endothelium expressing SOX17 (SX^34^43^-) is highlighted. Plots are representative of three experiments. (B) Summary of flow cytometry analysis of the SOX-RUNX, RUNX1-KO and SOX-RUNX cells supplemented with LSD1 inhibitor shown in (A) (mean±SEM, n=3 experiments). * P<0.05, *** P<0.0002, **** P<0.0001 compared to SOX-RUNX, 2-way ANOVA, Holm Sidak’s multiple comparisons test. (C) Gene ontology (GO) terms related to genes downregulated (LO) in RNA-Seq data from d6 LSD1 inhibitor (LSDi) treated cultures reflected the loss of blood cell differentiation, oxygen transport and hemoglobin complex. GO terms related to genes with higher (HI) expression in the same cultures related to ion transport and, more significantly, genes up regulated by the HDAC inhibitor panobinostat. (D) Heatmaps of RNA-Seq data from d4 and d6 SOX-RUNX (SR) and RUNX1-KO (R1KO) ± LSD1 inhibitor (LSDi) of the most differentially expressed genes down regulated (LO) or up regulated (HI) in LSDi supplemented compared to un-supplemented cultures. Scale, log_2 RPKM. See also Fig. 6M.
Figure S7. Human blast colonies generate RUNX1-dependent yolk sac-like cells. (A) Differentiation protocol for lineage specification. BL-CFC, blast colony forming cell; MC, methylcellulose; LC, liquid culture. (B) Images of May-Grünwald-Giemsa stained cytocentrifuge preparations of d2+7+7 cultures illustrating appearance of erythroid cells in EPO cultures, macrophages in M-CSF supplemented cultures and granulocytes exclusively in G-CSF/GM-CSF treated cultures. Images are representative of three experiments. Scale bar, 100 µm. (C) Flow cytometric profiles of cultures showing expression of GYPA, CD14 and CD43 to distinguish between erythroid, macrophage and granulocytic lineages (D) Quantification of flow cytometric analysis illustrating the predominance of RUNX1C expression in the G-CSF/GM-CSF treated cultures, CD14 in the M-CSF supplemented cultures and GYPA in the EPO treated cultures (mean±SEM n=3 experiments). *** P<0.0002, **** P<0.0001 compared to EPO supplemented cultures, 2-way ANOVA, Holm Sidak’s multiple comparisons test. (E) Relative gene expression (shown as negative delta (-d) Ct) analysis of d2+7+7 cultures demonstrating the expression of EPSILON and GAMMA globin, KLF1 and GATA1 in EPO supplemented cultures, CSF1R and SPI1 in M-CSF supplemented cultures and EPX, SPI1 and GATA1 in G-CSF/GM CSF supplemented cultures, consistent with the development of erythroid, macrophage and granulocytic lineages respectively (mean of n=2 experiments). Data presented in panels B and C are from the same experiment.
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