A Quantitative Proteomic Analysis of Hemogenic Endothelium Reveals Differential Regulation of Hematopoiesis by SOX17

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SUMMARY

The in vitro derivation of hematopoietic stem cells (HSCs) from pluripotent stem cells (PSCs) is complicated by the existence of multiple overlapping embryonic blood cell programs called primitive, erythromyeloid progenitor (EMP), and definitive. As HSCs are only generated during the definitive stage of hematopoiesis, deciphering the regulatory pathways that control the emergence of this program and identifying markers that distinguish it from the other programs are essential. To identify definitive specific pathways and marker sets, we used label-free proteomics to determine the proteome of embryonic-derived and mouse embryonic stem cell-derived VE-CADHERIN+CD45− definitive hematopoietic progenitors. With this approach, we identified Stat1 as a marker that distinguishes the definitive erythroid lineage from the primitive- and EMP-derived lineages. Additionally, we provide evidence that the generation of the Stat1+ definitive lineage is dependent on Sox17. These findings establish an approach for monitoring the emergence of definitive hematopoiesis in the PSC differentiation cultures.

INTRODUCTION

The generation of hematopoietic stem cells (HSCs) from pluripotent stem cells (PSCs) is dependent on our ability to accurately recapitulate the intricate steps of early embryonic hematopoietic development in the differentiation culture. The mammalian embryonic hematopoietic system consists of at least three distinct programs—primitive, erythromyeloid progenitor (EMP)/yolk sac definitive, and intraembryonic definitive—that differ in their potential as well as spatial and temporal patterns of development (McGrath et al., 2011; Medvinsky and Dzierzak, 1996; Palis et al., 1999). Primitive hematopoiesis is the first to develop, initiates in the yolk sac at embryonic day 7.0 (E7.0), and produces a restricted repertoire of hematopoietic cells consisting of primitive erythrocytes that express embryonic forms of globin and macrophages. Shortly after the onset of primitive hematopoiesis, the yolk sac generates a cohort of erythroid, myeloid, and megakaryocytic progenitors that constitute a distinct phase of embryonic hematopoiesis known as the erythromyeloid progenitor program, also referred to as the yolk sac definitive program (McGrath et al., 2011). This program appears to be positioned developmentally between primitive and definitive hematopoiesis. In the mouse, it generates erythroid cells that express adult and embryonic globins but does not give rise to lymphoid lineage cells or hematopoietic stem cells (McGrath et al., 2011; Palis et al., 1999). HSCs are generated during the intraembryonic-definitive stage of hematopoiesis that is specified following the emergence of the two yolk sac programs at different sites within the embryonic arterial vasculature, the most well characterized being the developing aorta, gonad, and mesonephros (AGM) (Medvinsky and Dzierzak, 1996). In addition to HSC and lymphoid potential, murine definitive hematopoiesis is distinguished from the two earlier programs by the fact that the erythroid lineage expresses exclusively adult forms of globin. Although both the EMP and intraembryonic-definitive hematopoietic programs are considered to be “definitive” due to their ability to generate erythroid precursors, to distinguish between them in this study, we use the terms EMP for the yolk-sac-derived progenitors and definitive for those derived from intraembryonic sites.

There is now compelling evidence indicating that both the EMP and definitive programs derive from progenitors known as hemogenic endothelial cells (HECs) that display endothelial markers and are associated with the arterial vasculature (Bertrand et al., 2005; Boisset et al., 2010; Clarke et al., 2013; Kissa and Herbomel, 2010). Live imaging and histological analyses have demonstrated that hematopoietic progenitors develop from the HECs through a process known as the endothelial to hematopoietic transition (EHT) (Boisset et al., 2010; Kissa and Herbomel, 2010). Currently, there are no markers to distinguish EMP-HECs from definitive HECs. As the EMP program develops earlier than the definitive program, the EMP-derived lineages can predominate the PSC differentiation cultures and are often misinterpreted as representing definitive
hematopoiesis. Developing strategies to distinguish definitive- from EMP-derived hematopoiesis in the PSC differentiation cultures is essential for identifying the signaling pathways that regulate the development of definitive HECs and their specification to HSCs.

We have previously shown that SOX17 is expressed in definitive HECs and that it is required for the establishment of the definitive, but not the primitive, hematopoietic program (Clarke et al., 2013). Our analysis of differentiated hematopoietic progeny from HECs generated from SOX17-null mouse embryonic stem cells (mESCs) revealed the presence of primitive erythroid progenitors, myeloid progenitors, and erythroid progenitors that gave rise to large colonies distinct from the primitive erythroid colonies. The presence of this population of erythroid progenitors suggests that the EMP program may also be independent. Our goals in the present study were to define the role of SOX17 in EMP hematopoiesis, understand how SOX17 regulates the generation of definitive hemogenic endothelium, and identify additional markers to distinguish these programs. To achieve these goals, we deciphered the proteome of AGM-derived HECs and of HECs generated from Sox17-wild-type and Sox17-knockout mESCs using label-free quantitative proteomics. The findings from these studies demonstrate the overall utility of analyzing the proteome of small numbers of cells and show that expression of the transcription factor Stat1 and dependency on Sox17 distinguish the definitive erythroid lineage from the primitive and EMP-derived erythroid lineages.

RESULTS

Proteomic Analyses of Wild-Type mESC- and Embryo-Derived HEC-Containing Populations

To identify markers that distinguish the different HEC populations and regulators that control their development, we quantitatively measured the global protein expression profile of mESC- and embryo-derived progenitors using label-free proteomics. In the first set of experiments, we isolated and analyzed the VE-CADHERIN* (VEC) CD45− population from the AGM region of E10.5 and E11.5 mouse embryos. This population contains non-hemogenic endothelial cells (ECs), HECs, and type I pre-HSCs at E10.5 and E11.5 (Rybtsov et al., 2011). The corresponding VEC−CD45− fractions from the same embryos were analyzed as representing the non-EC/HEC populations. For the second set of samples, we focused on mESC-derived populations. We previously described a chemically defined serum-free and feeder-free differentiation protocol for mESCs in which the primitive hematopoietic program is identified as an early arising FLK-1+ population (day 3.25 [D3.25]) and the definitive program as a FLK-1+ population that emerges at D5.25 (Clarke et al., 2013; Irion et al., 2010). When isolated and cultured for an additional 2 days (total day 7 [D7]) these definitive D5.25 FLK-1+ cells give rise to an endothelial VEC+ population that contains HECs that undergo the EHT to generate CD45− hematopoietic cells at D9. The D5.25 FLK-1+ progenitors and derivative D7 VEC+CD45− cells were isolated from two separate mESC lines (mESC-1 and mESC-2) and subjected to proteomic analyses (Figure 1A).

For all analyses, protein digests were measured by nano-liquid chromatography-tandem mass spectrometry (nano-LC-MS/MS) on a Q Exactive mass spectrometer and quantified using label-free methods (Luber et al., 2010). In total, 36,678 peptides corresponding to 4,044 proteins were quantified in both the in vivo and in vitro studies (Table S1). Protein quantification was reproducible across in vitro and in vivo measurements, with a Pearson’s correlation between replicates of 0.89 and 0.90, respectively (Figures 1B and 1C).

To validate that the proteomics analysis could accurately distinguish between the EC/HEC and non-EC/HEC fractions, we analyzed the E11.5 VEC+ and VEC− populations for proteins known to be expressed in HECs, including ALDH1A2 (RALDH2), SCL/TAL1, CBFβ, and ESAM. All were enriched in the VEC+ fraction, confirming the utility of this proteomic approach to accurately identify the HECs within this heterogeneous cell population (Chanda et al., 2013; Van Handel et al., 2012; Yokota et al., 2009) (Figure S1A; Tables S2 and S3). Next, we asked if our proteomics analysis could identify differences between the E10.5 VEC−CD45− and E11.5 VEC−CD45− populations. The E11.5 VEC−CD45− population contains more potent or higher numbers of pre-HSCs than the E10.5 population (Rybtsov et al., 2011; Yokomizo et al., 2012). We identified 85 proteins that were differentially expressed between these populations (Figures S1B and S1C; Table S4). Among the proteins expressed at higher levels in the E11.5 population than in the E10.5 population, we identified STAM2 and SLFN1. STAM2 is a critical adaptor protein downstream of growth factor and inflammatory cytokine signaling pathways (Pandey et al., 2000). The identification of STAM2 in the E11.5 population is of interest, as recent studies indicate that inflammatory signaling plays a crucial role in the emergence of HSCs (Espín-Palazón et al., 2014; Li et al., 2014; Sawamiphak et al., 2014). SLFN1 is a member of the Schlafen family of proteins that are expressed in endothelial and hematopoietic cells and are emerging as important regulators of the immune response (Kuang et al., 2014; Schwarz et al., 1998). By contrast, ESAM (endothelial cell selective adhesion molecule), which is a marker of emerging multipotent stem and progenitor cells, and members of the RAS signaling pathway (RAβ17 and SIPA1) were found at higher levels in the E10.5 population.
Figure 1. Quantitative Proteomic Profiling of Hemogenic Endothelium
(A) Proteomic workflow used to identify differentially regulated protein expression in vitro and in vivo.
(B) Label-free quantification of protein expression in vitro is reproducible. Tryptic digestions of cell lysates were analyzed by nano-LC MS/MS. n = 3 individual measurements; Pearson correlation of 0.89.
(C) Label-free quantification of protein expression in vivo is reproducible. Tryptic digestions of mouse tissue were analyzed as in (B). n = 3 individual measurements; Pearson correlation of 0.90.
(D) Protein expression was analyzed by unsupervised hierarchal cluster analysis (HC). Green denotes high, relative protein expression and blue low, relative protein expression.
(E) Protein expression was analyzed by principle-component analysis.
(F) GO terms linked to proteins enriched in VE-CADHERIN+ populations were analyzed in G-profiler using the cocoa algorithm. See also Figure S1.
Lastly we compared our D7 mESC-derived ECs/HECs to the E10.5 and E11.5 VEC+CD45− ECs/HECs (Figure 1D). Principle-component analysis revealed that the VEC+ populations from the two mESC lines were similar to one another. Component 2 reflects the variance between E10.5 AGM and E11.5 AGM. Based on this analysis, the mESC-derived populations appear to represent a stage of development between them. Component 1 is an indication of the variance between in vivo and in vitro culture conditions. As expected, cells grown in vitro do not perfectly recapitulate in vivo development (Figure 1E). Gene Ontology (GO) analysis showed that proteins linked to anchoring junctions, including calcium ion binding and calcium-dependent phospholipid binding, were more highly expressed in the AGM VEC+CD45− and VEC−CD45− cells than the D7 mESC-derived cells (Figure 1F). Collectively, these findings highlight the power of the label-free proteomics approach for assessing protein expression in a small number of cells, and in doing so, they identify candidate signaling pathways that may play a role in the generation of HSCs from HECs in vitro.

**Determination of the SOX17-Dependent Proteome in mESC-Derived HEC-Containing Populations**

We previously reported that SOX17 is a critical regulator of hemogenic endothelium development in vivo and during the directed differentiation of mESCs in vitro (Clarke et al., 2013). However, the direct targets of SOX17 required for the generation of HECs are unknown. In our previous studies, we demonstrated that the doxycycline-inducible overexpression of SOX17 in FLK-1+ mESC-derived endothelial progenitors increased the formation of HECs while repressing the endothelial to hematopoietic transition. We therefore hypothesized that the quantitative measurement of changes in protein expression following the induced expression of SOX17 would identify both direct and indirect target proteins of SOX17 during the generation of HECs. D5.25 FLK-1+ endothelial progenitors were isolated and treated with doxycycline for 1–3 days to induce the expression of SOX17. Samples were isolated from D6 to D9 and analyzed for total protein expression using label-free proteomics (Figure S2A). In total, SOX17 directly or indirectly regulated the expression of 110 proteins (Figure S2B; Table S5) that we classified into early, mid, or late effectors (Figure S2D). SOX17 has been implicated as both a transcriptional activator and repressor, suggesting that proteins that exhibit a significant increase or decrease immediately following the induction of SOX17 may be direct transcriptional targets of SOX17 in HECs. We not only quantified changes in SOX17 expression (Figures S2B and S2C) but also identified several proteins that have been suggested to be targets of SOX17 in other cell systems, including UCHL1 (extraembryonic endoderm development) (Sinner et al., 2004) and CDKN1C (p57/KIP2, respiratory epithelial cells) (Lange et al., 2009) (Table S5). Consistent with our previous observation that the enforced expression SOX17 promotes the maintenance and expansion of HECs, thus postponing the EHT (Clarke et al., 2013), we identified that SOX17 overexpression decreased the expression of proteins linked to the erythrocyte function during the time course of directed differentiation (Figures S2E and S2F). Taken together, we expect that this proteomic dataset will be a useful resource for researchers seeking to identify regulators of the expansion and maintenance of HECs.

As SOX17 is required for the establishment of definitive, but not primitive, hematopoiesis, we hypothesized that analysis of the SOX17-dependent proteome would reveal previously unidentified insights into the global pathways involved in the regulation and identification of these disparate hematopoietic programs. To investigate this, we next analyzed the proteome of D7 VEC+CD45− cells generated from control and Sox17−/− mESCs (Figure 2A). We found that 344 proteins were differentially expressed between these populations (Figure 2B; Table S6). Among the differences, we noted that the SOX17-null VEC+ cells expressed higher levels of erythrocyte-linked proteins (Figure 2C), specifically those associated with the primitive and EMP hematopoietic programs including the embryonic forms of β-globin, HBB-ε and HBB-βH1 (Table S6). This is consistent with our previous observation that the D7 Sox17−/− VEC+ population is heterogeneous and contains committed erythroid and myeloid progenitors as well as HECs (Clarke et al., 2013). Interestingly, GO analysis identified that Stat transcription factors had decreased expression in the Sox17−/− population compared to the control cells (STAT1, STAT3, and STAT5B) (Figure 2C; Table S6). Notably, STAT1 expression was significantly reduced in the D7 Sox17−/− VEC+ cells, suggesting that it may be a previously unidentified transcriptional target of SOX17. Consistent with the reduction in STAT1, we also observed a significant reduction in its direct transcriptional target ISG15 (Malakhov et al., 2003) (Figures 2D and S3A).

To determine whether SOX17 can directly activate Stat1 transcription, HEK293T cells were transfected with a Sox17 expression plasmid along with a construct containing a 2.5-kb region of the Stat1 promoter that includes four potential SOX17-binding sites upstream of luciferase (JASPER database; data not shown). Transfection of Sox17 did not lead to induction of luciferase activity (Figure S3B). Consistent with this observation, we did not detect binding of SOX17 to the Stat1 promoter by chromatin immunoprecipitation (ChIP) analysis (data not shown). Finally, we were unable to demonstrate an upregulation of Stat1 following the enforced expression of SOX17 in the D5.25 FLK-1+ cells using a doxycycline-inducible mESC line.
Collectively, these findings provide strong evidence that Stat1 is not a direct target of SOX17 in the VEC+ cells and in doing so support an alternative interpretation in which Stat1 is expressed in a definitive hematopoietic progenitor that is regulated by SOX17.

**Expression of Stat1 in Embryo-Derived Erythroid Populations**

The observed differences in globin gene expression between the wild-type and SOX17-null VEC+CD45− populations indicate that they are heterogeneous and additional markers are needed to identify the type of hematopoietic progenitors present within them. As previous studies have shown that CD41 is expressed on emerging primitive and definitive hematopoietic progenitors, including those of the erythroid lineage (Ferkowicz et al., 2003; Mitjavila-Garcia et al., 2002), we next analyzed D7 wild-type and Sox17−/− VEC+CD45− populations for the presence of this marker. Approximately half the wild-type VEC+CD45− population expressed CD41, whereas more than 80% of the SOX17-null cells were positive (Figure S3D). These findings show that the VEC+CD45− populations used for the proteomic analyses contained CD41+ hematopoietic progenitors and raise the interesting possibility that the differences in Stat1 and globin expression between the two reflect differences in their erythroid progenitor content. Specifically, they support the hypothesis that Stat1 is expressed in an erythroid progenitor that is regulated by SOX17. Differential expression of the Stat regulators in different erythroid populations is in line with a recent computational study that provided evidence that STAT3 signaling is required for primitive erythropoiesis, whereas Stat1 is preferentially expressed during the definitive stage (Greenfest-Allen et al., 2013). EMP-derived erythroid progenitors were not examined in these experiments.

To determine if expression of Stat1 distinguishes the erythroid lineages generated from the different embryonic hematopoietic programs, we monitored its expression as well as that of other Stat genes in the primitive, EMP-derived, and definitive erythroid cells (Figure 3A) obtained from wild-type and SOX17-null embryos and mESCs.
differentiated in culture. In the mouse, the primitive erythroid (Ery-P), EMP-derived (Ery-EMP), and definitive erythroid (Ery-D) cells can be distinguished based on distinct β-globin gene expression patterns. Ery-P cells express high levels of the embryonic β-globin HBB-βH1 and HBB-εγ and low levels of the adult forms β-globin Hbb-β1/β2. Ery-EMP cells express low levels of HBB-βH1 and HBB-εγ, and an increased amount of HBB-β1/β2 (McGrath et al., 2011). Ery-D cells express exclusively the adult β-globins HBB-β1/β2. In addition to patterns of globin expression, the development of these erythroid populations can be monitored by the expression of specific transcription factors. EKLF/KLF1 regulates the expression of several erythroid-specific genes, including both embryonic and adult globin, and is present in all three erythroid populations (Basu et al., 2007; Hodge et al., 2006).

\[ \text{EKLF/KLF1} \]

Sox6 and Bcl11A are expressed in the EMP and definitive erythroid cells, where they function to suppress the embryonic globin genes (Sankaran et al., 2009; Yi et al., 2006). Neither is expressed in the primitive erythroid lineage.

E8.5 yolk sacs (YS) and E14.5 fetal livers (FL) were isolated and the cells from each cultured in methylcellulose for the generation of primitive, EMP (E8.5 YS), and definitive (E14.5 FL) erythroid colonies, respectively (Figure 3B). The subsets of colonies were identified based on phenotype, then picked, pooled, and analyzed by real-time qPCR for gene expression profiles (Figure 3A). As expected, primitive erythroid colonies expressed high levels of Hbb-εγ and low levels of Hbb-β1. Additionally, they expressed Stat3, as previously reported (Greenfest-Allen et al., 2013). The primitive erythroid colonies expressed only low levels of the other Stat genes and EKLF. They had no detectable levels of Sox6 or Bcl11A. In contrast, the fetal-liver-derived definitive colonies expressed high levels of adult β-globin (Hbb-β1) but no embryonic (Hbb-εγ, Hbb-βH1) globin. These colonies also expressed Stat1, EKLF, Sox6, and
Bcl11a (Figure 3C). Only low levels of Stat3, Stat5a, and Stat5b were detected in the definitive cells. As a means to categorize the real-time qPCR values, we will consider a deltaCT (ΔCT) level of >50 as “high” for globin genes and >0.02 as high for transcription factors. The EMP-derived erythroid colonies expressed a mixture of low levels of embryonic and adult β-globin, as well as Eklf, Sox6, and Bcl11a. Interestingly, these erythroid colonies showed low expression levels of all STAT proteins. Together, these data show that embryo-derived primitive, EMP, and definitive erythroid cells have distinct gene expression signatures and that the expression of Stat1 distinguishes between EMP and definitive erythroid cells.

Expression of Stat1 in mESC-Derived Wild-Type and Sox17<sup>+/−</sup> Erythroid Populations

We next examined if this gene expression signature can be used to distinguish the ESC-derived erythroid lineages generated from both wild-type and Sox17<sup>+/−</sup> ESCs. The Sox17<sup>+/−</sup> populations were included, as our previous study indicated that development of primitive and EMP programs were SOX17 independent, whereas specification of the definitive lineage required SOX17. In the initial set of experiments, we analyzed the primitive and EMP lineages generated from D3.25 FLK-1<sup>+</sup> mesoderm, a population that we have previously shown gives rise to the equivalent of yolk sac hematopoiesis (Clarke et al., 2013; Iriion et al., 2010). For these studies, the wild-type (control) and Sox17<sup>+/−</sup> FLK-1<sup>+</sup> cells were isolated by fluorescence-activated cell sorting (FACS) and aggregated overnight. The aggregates were cultured for 48 hr (Figure 4A). As an independent method to distinguish the different erythroid progenitors, we analyzed their dependency on NOTCH signaling, as previous studies have shown that the primitive and EMP hematopoietic programs are NOTCH independent whereas specification of the definitive program is dependent on this pathway (Hadaland et al., 2004). To measure NOTCH dependency, we included the NOTCH pathway antagonist gamma secretase inhibitor (γSI) in the aggregate cultures. The resulting populations were assayed in methylcellulose. Both the wild-type and Sox17<sup>+/−</sup> D3.25 FLK-1<sup>+</sup> populations generated typical small primitive erythroid (Ery-P) colonies as well as larger erythroid colonies that resembled those that develop from either EMP or definitive progenitors (Ery-EMP/D) (Figure 4B). Although the erythroid potential was intact in the Sox17<sup>+/−</sup> mesoderm, the cellularity as well as the frequency of Ery-P and Ery-EMP progenitors was significantly lower in the Sox17<sup>+/−</sup> cultures than in the wild-type cultures (Figure 4C). The reduced cellularity in Sox17<sup>+/−</sup> populations is consistent with observations in our previous study (Clarke et al., 2013). The addition of γSI did not significantly affect cell numbers or the generation of Ery-P progenitors from either mESC line (Figure 4D). It did, however, lead to a modest but significant reduction in the number of larger wild-type colonies (Ery-EMP/D) indicating that a subset of them originate from a NOTCH-dependent definitive hematopoietic progenitor or that NOTCH is involved in the proliferation, but not emergence, of EMP hematopoietic progenitors. (Figure 4D, black bars). The number of Sox17<sup>+/−</sup> Ery-EMP/D progenitors was not affected by the addition of γSI (Figure 4D, gray bars).

The wild-type and Sox17<sup>+/−</sup> Ery-P colonies had a gene expression signature similar to that of the E8.5 YS-derived primitive erythroid cells, including high levels of Hbb-εγ and Stat3 (Figure 4E). They also expressed low levels of Eklf and the other Stat genes. Sox6 and Bcl11a were not expressed in these colonies. The expression profile of the wild-type Ery-EMP colonies was also in line with that of the YS-derived colonies and showed significantly lower levels of embryonic globin Hbb-εγ and Stat3 than the Ery-P colonies. As observed with the yolk sac colonies, the mESC-derived Ery-EMP colonies also had low levels of all the other Stat genes tested. These colonies did express Eklf, Sox6, and Bcl11a (Figure 4F, black bars). The Sox17<sup>+/−</sup>-derived colonies had a similar “EMP” profile, including low levels of Hbb-εγ and the Stat genes. These colonies did, however, express lower levels of Hbb-δ1 and Eklf than the wild-type colonies. The reason for these differences is currently not clear.

To generate ESC-derived definitive progenitors, D5.25 FLK-1<sup>+</sup> cells were isolated, aggregated, and cultured overnight (Figure 5A). The aggregates were then plated as a monolayer in the presence of cytokines known to induce the EHT and the generation of CD45<sup>+</sup> hematopoietic progenitors. γSI was included in some of the cultures during the aggregation and Matrigel stages. The wild-type and Sox17<sup>+/−</sup> D9 populations gave rise to erythroid colonies that phenotypically resembled those derived from Ery-EMP or Ery-D progenitors (Figure 5B). Primitive erythroid colonies were not detected in these cultures. The addition of γSI did not impact overall cellularity, but it did lead to a reduction in the proportion of CD45<sup>+</sup> cells and in the number of erythroid colonies generated in the wild-type population, consistent with the interpretation that they represent the definitive lineage (Figures 5C–5E, black bars). In contrast, neither the size of the Sox17<sup>+/−</sup> CD45<sup>+</sup> population nor the number of Sox17<sup>+/−</sup> erythroid colonies was affected by the addition of γSI, indicating that these cells represent the EMP program. Gene expression analysis confirmed these differences as the control erythroid colonies revealed an expression pattern comparable to the FL-derived definitive lineage, including Hbb-β1<sup>hi</sup>, Hbb-εγ<sup>hi</sup>, Stat1<sup>hi</sup>, Eklf<sup>hi</sup>, Sox6<sup>hi</sup>, and Bcl11a (Figure 5F, black bars). The Sox17<sup>+/−</sup> colonies, by contrast, displayed an EMP profile that included significantly lower levels of...
Figure 4. Day 3.25 FLK-1+ Wild-Type and Sox17-alt/mESC-Derived Populations Contain Primitive and EMP Progenitors

(A) Schematic representation of the differentiation protocol for generating different hematopoietic populations from mESCs. (B) Representative image of erythroid colonies from a control D5 population generated from D3.25 FLK-1+ cells after 5 days in culture. (C) Fold change in cell numbers following 48 hr of culture of control and Sox17-alt/mESC D3.25 FLK-1+ cells in the presence and absence of γ-secretase inhibitor (γSI, L685458, Tocris, 10 μM). Cells cultured in the vehicle DMSO represent the control. DMSO p = 0.00958, γSI p = 0.0285. n = 3 individual experiments (SEM). (D) Erythroid potential of the D5 population generated from D3.25 FLK-1+ cells. n = 3 individual experiments (SEM). Control EryP DMSO: Sox17-alt/mESC DMSO p = 0.031, control EryP γSI: Sox17-alt/mESC γSI p = 0.012, control Ery-EMP DMSO: Sox17-alt/mESC Ery-EMP γSI p = 0.024, control Ery-EMP DMSO: Sox17-alt/mESC Ery-EMP γSI p = 0.0039, and control Ery-EMP γSI: Sox17-alt/mESC Ery-EMP γSI p = 0.0032.

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SOX17 Is Not Required for Development of the Primitive and EMP-Derived Erythroid Lineage In Vivo

To evaluate the SOX17 dependency of primitive and EMP erythroid lineages in vivo, we next analyzed colonies generated from the yolk sacs of E8.5 Sox17+/+, Sox17+/−, and Sox17−/− embryos for the above expression signature. As observed with the mESC-derived populations, the SOX17-deficient (Sox17+/− and Sox17−/−) embryos contained a lower frequency of primitive erythroid progenitors than the wild-type littermates, possibly due to defects in the yolk sac vasculature resulting from SOX17 deficiency (Kim et al., 2007). In spite of the reduced numbers, the colonies from the three different sets of embryos had very similar expression profiles that included high levels of Hbb-ε and Stat1 and Stat3 and low levels of the other Stat genes (Figures 6A and 6B). Both the Sox17+/− and Sox17−/− embryos contained progenitors that generated EMP-type colonies. The expression pattern of these colonies was consistent with that of the EMP, with the exception that those derived from the Sox17−/− progenitors expressed lower levels Hbb-ε1 than the colonies from the Sox17+/− and Sox17+/+ progenitors, similar to the pattern seen in the Sox17−/− mESC-derived EMPs (Figure 5F). Collectively, these data show that SOX17 is not required for the emergence of primitive and EMP-derived hematopoietic program in the embryo.

DISCUSSION

Label-free proteomics represents an emerging technology for identifying and quantifying patterns of protein expression. While differences in patterns of protein expression have previously been shown to correlate with differentiated cell types, it has been much more challenging to develop technologies that can detect regulatory proteins such as transcription factors that specify specific cellular fates. Emerging embryonic HSCs and their progeny have been comprehensively examined via global transcriptional profiling, which revealed key insights into hematopoietic specification and leukemogenesis (Boisset et al., 2010; McKinney-Freeman et al., 2012). The direct precursor to the HSC, the HEC, has not been extensively examined, as only an abbreviated transcriptional profile of HECs within the murine embryo was recently elucidated using a Fluidigm-based approach (Swiers et al., 2013). Here, we have utilized label-free proteomics to decipher key regulators of embryonic hematopoiesis both in vivo and during the directed differentiation of ESCs in vitro. This approach to cellular profiling has the distinct advantage of identifying only translated functional proteins present in the cell, thus eliminating the unknown effects of post-transcriptional modification that may affect mRNA translation.

Our first analysis focused on the proteomic landscape of E10.5 and E11.5 VEC+ populations that contain ECs, HECs, and pre-HSCs (Rybtsov et al., 2011). With this approach, we were able to accurately identify and quantify known regulators of the early hematopoietic programs such as SCL/TAL1 and ALDH1A2, as well as a small cohort of proteins that exhibited dramatic expression changes between E10.5 and E11.5, providing a focused and manageable list of candidates to interrogate as putative regulators of the generation of type I pre-HSCs. To date, the generation of PSC-derived HSCs has been difficult, likely due to the undefined nature of PSC differentiation culture conditions and the lack of accurate comparison of the in vitro populations to potential equivalent in vivo embryonic stages. Comparison of D7 mESC-derived HEC populations to the E10.5 and E11.5 HEC populations revealed that our D7 HEC population represents a developmental intermediate. Therefore, interrogation of this dataset will help identify the absence of key regulatory proteins, thus providing a potential explanation for the inability of these cells to faithfully generate HSCs in vitro.

Understanding the regulatory pathways that control the establishment of the different hematopoietic programs and developing methods to identify and isolate progenitors for each are essential to accurately model hematopoiesis in ESC and induced PSC differentiation cultures and ultimately to generate HSCs in vitro. Building on our previous observations that SOX17 is a critical regulator of definitive hematopoiesis (Clarke et al., 2013), in this study, we proceeded to analyze the SOX17-dependent proteome in

Stat1 and Hbb-ε1 expression than observed in the wild-type definitive colonies. Taken together, these findings show that combination of Stat1 and globin gene expression can be used to distinguish the different mESC-derived erythroid programs. Additionally, they confirm our earlier interpretation that the primitive and EMP programs are not dependent on SOX17, whereas generation of the definitive lineage is SOX17 dependent. Finally, they show that the EMP-derived erythroid lineage persists to the definitive stage in the Sox17−/− cultures.

(E and F) Real-time qPCR analysis of the erythroid colonies. n = 3 individual experiments (SEM). Control Ery-EMP:Sox17+/− Ery-EMP Hbb-ε1 p = 0.00012. In addition, but not labeled in the figure, the control EryP and Ery-EMP colonies were significantly different from each other for the following genes; Hbb-ε1 (p = 0.002), Hbb-β3 (p = 0.0023), and Stat3 (p = 0.09). The Sox17−/− EryP and Ery-EMP colonies were significantly different from each other for the following genes; Hbb-ε1 (p = 0.05).
Figure 5. Day 9 mESCs Generate Stat1⁺, NOTCH-Dependent, and SOX17-Dependent Definitive Erythroid Cells

(A) Schematic representation of the differentiation protocol for generating definitive hematopoietic progenitors from mESCs.

(B) Representative image of control erythroid colonies from a D9 population generated from D5.25 FLK-1⁺ cells after 8 days in culture.

(C) Representative flow cytometric profiles of cells in D9 populations stained with CD41.

(D) Fold change in cell numbers of D9 populations generated in the presence and absence of γSI.

(E) Erythroid potential of the D9 populations; n = 3 individual experiments (SEM). Control DMSO: control γSI p = 0.024. Control γSI:Sox17⁻/⁻ γSI p = 0.0028.

(F) Real-time qPCR analysis of the erythroid colonies. n = 3 individual experiments (SEM). Control:Sox17⁻/⁻ Stat1 p = 0.0032, control Stat1:Stat3 p = 0.03, control Stat1:Stat5a p = 0.03, and control Stat1:Stat5b p = 0.03.
HECs and hematopoietic progenitors, and through this approach, we were able to identify potential regulatory pathways of the definitive hematopoietic program and markers to distinguish it from the earlier primitive and EMP programs. Specifically, we identified Stat1 as a marker whose expression distinguishes the definitive erythroid lineage from the primitive and EMP-derived erythroid lineages. The differences observed between the EMP and definitive programs are particularly important, as they are difficult to distinguish by other methods. The observation that the D3.5 FLK-1⁺ progenitors give rise to both the primitive and EMP erythroid lineages that are found in the yolk sac provides further support that the in vitro PSC system accurately recapitulates in vivo embryonic hematopoiesis.

Our STAT protein expression data support the recently published computational prediction of a differential role for STAT signaling in primitive and definitive erythropoiesis (Greenfest-Allen et al., 2013). STAT signaling is central to hematopoietic cell biology, as this pathway is activated in response to numerous cytokines, including erythropoietin (EPO) and stem cell factor (SCF/KL). The observations that EPO activates Stat1, Stat3, Stat5α, and Stat5b in mouse and human cell lines in culture, and that STAT1- and STAT5-deficient fetuses display a wide array of erythroid defects, provide clear evidence that STAT proteins are critical regulators of developmental erythropoiesis and erythroblast maturation (Cui et al., 2004; Halupa et al., 2005; Kirito and Komatsu, 2002; Kirito et al., 1997; Richmond et al., 2005; Socolovsky et al., 1999, 2001). Analysis of the role of STAT3 during erythropoiesis is hampered by the fact that Stat3⁻/⁻ mice are embryonic lethal at E7.5 due to a lack of mesoderm formation. However, small-molecule inhibition of STAT3 activation resulted in decreased numbers of primitive, but not adult definitive, erythroblasts in culture, suggesting a differential role of STAT3 during distinct stages of hematopoiesis (Greenfest-Allen et al., 2013). A mechanism by which STAT1 may preferentially regulate the maturation of definitive erythroid cells is through its transcriptional target ISG15. ISG15 is strongly upregulated during erythroid differentiation in the bone marrow (Margno et al., 2011) and was identified in our proteomics screen as a potential SOX17 target exhibiting a parallel expression pattern to STAT1 (Figure 2D). ISG15 can be covalently linked to lysine residues of target proteins as a method of post-translational modification (Zhang and Zhang, 2011). STAT5 and globin proteins have been shown to be ISGylated in erythroblasts, and ISG15⁻/⁻ erythroblasts are unable to terminally differentiate, thus providing a testable hypothesis by which STAT1 regulates definitive erythroid differentiation. Collectively, the observation
that distinct developmental subsets of erythroid cells express different STAT proteins presents a previously unidentified mode of regulation of embryonic erythropoiesis.

In summary, the data presented in this study demonstrate that quantitative proteomics is an informative and accurate method to identify global patterns of protein expression in small numbers of cells. By applying this approach to early hematopoietic progenitor populations, we have gained insights into potential pathways that regulate embryonic hematopoietic development and uncovered strategies for identifying the definitive hematopoietic program in PSC differentiation cultures. Further comparison of our proteomics analysis to previously published transcriptome datasets will provide a systems biology view of the early stages of embryonic hematopoiesis and allow us to obtain a better understanding of the signaling and transcription factor networks that underlie the specification of HSCs.

EXPERIMENTAL PROCEDURES

Sample Preparation for Mass Spectrometry
50,000 cells underwent FACS. The cells were then washed in 1× PBS and flash frozen. Cell pellets were lysed in 1 M urea and 50 mM ammonium bicarbonate (pH 7.8) and heated to 50°C for 20 min. Cell debris was removed by centrifugation (1,000 × g, 2 min). Following a bicinchoninic acid assay, normalized quantities of protein were reduced with 2 mM DTT, alkylated with 15 mM iodoacetamide, and digested overnight with a 1:50 ratio of trypsin to total protein. The resulting peptides were desalted with 20 min. Cell debris was removed by centrifugation (1,000 × g, 2 min). Following a bicinchoninic acid assay, normalized quantities of protein were reduced with 2 mM DTT, alkylated with 15 mM iodoacetamide, and digested overnight with a 1:50 ratio of trypsin to total protein. The resulting peptides were desalted on Waters Sep-Pak C18 cartridges.

Nano-LC-MS/MS Measurements
Peptides were measured by nano-LC-MS/MS on a Q Exactive (Thermo Scientific). Peptides were separated online by reverse-phase chromatography using a heated 50°C 30-cm C18 columns (75-mm ID packed with Magic C18 AQ 5-μM/100-A beads) in a 120-min gradient (1%-45% acetonitrile with 0.1% formic acid) separated at 250 nl/min. The Q Exactive was operated in the data-dependent mode with the following settings: 70,000 resolution, 300–2,000 m/z full scan, Top 10, and an 1.8-m/z isolation window.

Data Processing and Analysis
Identification and label-free quantification of peptides was done with MaxQuant 1.3.0.5 using a 1% false discovery rate (FDR) against the mouse Swiss-Prot/TrEMBL database downloaded from UniProt on October 11, 2013 (Cox and Mann, 2008). The databases contained forward and reverse mouse sequences as well as common contaminants. We analyzed three replicates per condition. Peptides were searched for variable modification of n-term acetylation, oxidation (M), deamidation (NQ), with a 6-ppm mass error and a match between run window of 2 min. Proteins that were significantly regulated between conditions were identified using a permutation-based t test (S1, FDR 5%) in Perseus 1.4.1.3. GO terms were analyzed in G-profiler (Reimand et al., 2011).

Real-Time qPCR
Total RNA was prepared with the RNAqueous-Micro Kit (Ambion) and treated with RNase-free DNase (Ambion). 500 ng to 1 μg RNA was reverse transcribed into cDNA using random hexamers and Oligo (dT) with Superscript III Reverse Transcriptase (Invitrogen). qPCR was performed on a MasterCycler EP RealPlex (Eppendorf) using Quantifast SYBR Green PCR Kit (QIAGEN). Expression levels were normalized to the housekeeping gene β-actin (Actb). The oligonucleotide sequences are listed in Table S7.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, three figures, and seven tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2015.07.008.

AUTHOR CONTRIBUTIONS
R.L.C., A.M.R., G.K., and R.T.M. designed experiments. R.L.C. and A.M.R. performed experiments and analysis. R.L.C. and A.M.R. wrote the manuscript. G.K. and R.T.M. edited the manuscript.

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