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Citation for published version:
https://doi.org/10.5966/sctm.2015-0324

Digital Object Identifier (DOI):
10.5966/sctm.2015-0324

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Stem cells translational medicine

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Enforced Expression of HOXB4 in Human Embryonic Stem Cells Enhances the Production of Hematopoietic Progenitors but Has No Effect on the Maturation of Red Blood Cells

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Key Words. CD34+ • Cell surface markers • Colony formation • Differentiation • Electroporation • Embryonic stem cells • Erythroid differentiation • Transcription factor

ABSTRACT

We have developed a robust, Good Manufacturing Practice-compatible differentiation protocol capable of producing scalable quantities of red blood cells (RBCs) from human pluripotent stem cells (hPSCs). However, translation of this protocol to the clinic has been complicated because the RBCs produced are not fully mature; thus, they express embryonic and fetal, rather than adult globins, and they do not enucleate efficiently. Based on previous studies, we predicted that activation of exogenous HOXB4 would increase the production hematopoietic progenitor cells (HPCs) from hPSCs and hypothesized that it might also promote the production of more mature, definitive RBCs. Using a tamoxifen-inducible HOXB4-ERα expression system, we first demonstrated that activation of HOXB4 does increase the production of HPCs from hPSCs as determined by colony-forming unit activity and the presence of CD43+CD34+ progenitors. Activation of HOXB4 caused a modest, but significant, increase in the proportion of immature CD235a+/CD71+ erythroid cells. However, this did not result in a significant increase in more mature CD235a+/CD71+ erythroid cells. RBCs produced in the presence of enhanced HOXB4 activity expressed embryonic (ε) and fetal (γ) but not adult (α) globins, and the proportion of enucleated cells was comparable to that of the control cultures. We conclude that programming with the transcription factor HOXB4 increases the production of hematopoietic progenitors and immature erythroid cells but does not resolve the inherent challenges associated with the production of mature adult-like enucleated RBCs.

SIGNIFICANCE

As worldwide blood donations decrease and transfusible transmitted infections increase, intense interest has ensued in deriving red blood cells (RBCs) in vitro from alternative sources such as pluripotent stem cells. A translatable protocol was developed to generate RBCs; however, these RBCs have an immature phenotype. It was hypothesized that the transcription factor HOXB4 could enhance their production and maturation. Although HOXB4 increased the production of erythroid progenitors, it did not promote their maturation. Despite the remaining challenges, a robust system has been established to test other candidates and add to the knowledge base in this field.

INTRODUCTION

Cell-based therapies such as bone marrow transplantation and blood transfusion are used to treat diseases of the hematopoietic system; however, these procedures are completely reliant on a limited supply of donor tissue. Thus, one goal has been to produce therapeutic hematopoietic cells from a bankable and limitless source of human embryonic stem cells (hESCs) or induced pluripotent stem cells (iPSCs) [1–3]. Many studies have been reported on the production of hematopoietic progenitors cells (HPCs) and mature hematopoietic cells from ESCs and iPSCs; however, it has been widely acknowledged that significant challenges exist in this field. The differentiation protocols that have been used to date are relatively inefficient, and it has proved particularly difficult to produce the most potent hematopoietic stem
cells (HSCs) capable of long-term reconstitution (LTR-HSCs) and fully mature red blood cells (RBCs) that express adult β-globin and enucleate [4–6]. Most of the strategies used to produce erythroid cells have required the use of stromal cell feeder layers and/or ill-defined serum, which would present significant safety issues in clinical translation [7–11]. Feeder-free hematopoietic differentiation strategies have been developed, but many of these required lengthy embryoid body stages that have proved difficult to control and replicate effectively [8, 12, 13]. The production of fully mature adult-like erythroid cells that undergo appropriate globin switching has been particularly challenging, with most studies detecting embryonic (ε, δ) and fetal (γ) but not adult (β) globin [8, 11, 14]. Although some studies have reported the production of some enucleated cells, none of these have demonstrated a convincing and robust degree of enucleation in pluripotent stem cell-derived RBCs [9, 12].

The first wave of erythropoiesis in vivo occurs in the yolk sac and is responsible for the production of primitive nucleated erythrocytes. A second wave initially arises in the aorta-gonad-mesonephros region, where the first definitive HSCs arise that are capable of long-term multilineage reconstitution and production of adult nucleated erythrocytes [15]. We hypothesized that the limited production of nucleated erythrocytes that can be achieved from pluripotent stem cells in vitro might reflect the “primitive” state of the progenitors produced in this system. This is supported by the finding that the production of LTR-HSCs in vitro from pluripotent stem cells capable of multilineage long-term reconstitution has proved particularly difficult to achieve. We reasoned that HOXB4 might be a transcription factor that could program hPSC-derived hematopoietic cells to a more “definitive” phenotype, because it has been shown to confer long-term reconstitution ability on primitive yolk sac cells and mouse ESCs [16–18]. Furthermore, our studies of the murine ESC system have demonstrated that HOXB4 could promote a paracrine effect; thus, its actions could mimic the action of stromal cell coculture [19–21]. The effects of exogenous HOXB4 on human ESCs has been less clear, with contradictory outcomes likely related to the range of the differentiation protocols used and the different expression strategies used, leading to variations in the level and timing of expression [19]. In the present study, we used a controllable HOXB4 expression system in which we could activate HOXB4 at different time points in a well-defined differentiation protocol.

We have developed a stepwise, serum- and feeder-free clinical grade protocol for both the growth and the maintenance of hESCs and hiPSCs and their subsequent expansion and differentiation into erythroid cells [22]. We tracked the emergence of HPCs using colony-forming unit culture (CFU-C) and flow cytometry and then tested the effects of tamoxifen-inducible expression of HOXB4 in this defined and robust hematopoietic differentiation protocol. In keeping with the results from murine studies, we found that enforced expression of HOXB4 resulted in a specific increase in the production of multilineage HPCs. We also noted a modest increase in the production of immature erythroid cells, but, overall, no significant effect was found for their subsequent maturation. The erythroid cells that were produced in the presence of HOXB4 failed to undergo efficient enucleation and expressed embryonic (ε) and fetal (γ), but not adult (β), globin. Thus, enforced expression of HOXB4 did not overcome the major difficulty in the production of mature enucleated cells required for transfusion.

**Materials and Methods**

**Maintenance and Differentiation of hESCs**

Human pluripotent stem cell lines, including H1 [23] and Runx1C-GFP [24], were routinely maintained in StemPro hESC serum-free medium on CELLstart matrix (Thermo Fisher Scientific Life Sciences, Waltham, MA, http://www.thermofisher.com) with the addition of basic fibroblast growth factor (bFGF) 20 ng/ml (PHG0021; Thermo Fisher Scientific Life Sciences), and passaged (1:4) when 70%–80% confluent, using StemPro EZPassage tools (Thermo Fisher Scientific Life Sciences). Differentiation was performed in a stepwise, serum- and feeder-free protocol, as described [22]. In brief, one confluent well of human ESCs was cut and transferred to two wells (six-well plates; cell repellant surface; catalog no. 657970; Greiner Bio-One, Kremsmünster, Austria, http://www.greinerbioone.com) and then cultured in 3 ml per well of Stemline II Hematopoietic Stem Cell Expansion Medium (catalog no. S0192; Sigma-Aldrich, St. Louis, MO, http://www.sigmaaldrich.com) with the addition of bone morphogenetic protein 4 (BMP4) 10 ng/ml (catalog no. 314-BP-010; R&D Systems, Minneapolis, MN, http://www.rndsystems.com), vascular endothelial growth factor (VEGF) 10 ng/ml (catalog no. 293-VE010; R&D Systems), Wnt3a 10 ng/ml (catalog no. 5036-WN010; R&D Systems), activin A 5 ng/ml (catalog no. 338-AC010; R&D Systems), and pig placenta extract (PEG) 2 μl (catalog no. A014418; Merck Millipore, Billerica, MA, http://www.merckmillipore.com). EBs formed spontaneously, and after 2 days, the following cytokines were added in 500 μl of Stemline II per well: BMP4 20 ng/ml, VEGF 30 ng/ml, Wnt3a 10 ng/ml, activin A 5 ng/ml, fibroblast growth factor-α (FGF-α) 10 ng/ml (catalog no. PHG0014; Thermo Fisher Scientific Life Sciences), stem cell factor (SCF) 20 ng/ml (catalog no. PHC 2111; Thermo Fisher Scientific Life Sciences), and GSK inhibitor VII 2 μM. EBs were disaggregated on day 3 with 0.5 ml per well StemPro Accutase Cell Dissociation Reagent for 3 minutes at 37°C, before gently pipetting 10 times and adding prewarmed medium. After centrifugation, the cells were replated at 2 × 10^5 cells per well (six-well plates; Corning, Corning, NY, http://www.corning.com) in 3 ml of Stemline II, with the addition of BMP4 20 ng/ml, VEGF 30 ng/ml, FGF-α 10 ng/ml, SCF 30 ng/ml, insulin-like growth factor 2 (IGF2) 10 ng/ml (catalog no. 292-G2; R&D Systems), thrombopoietin (TPO) 10 ng/ml (catalog no. 288-TPN-25; R&D Systems), heparin 5 μg/ml (catalog no. H3149; Sigma-Aldrich), and isobutylmethylxanthine (IBMX) 100 μM (catalog no. IS879; Sigma-Aldrich). On day 7, the floating cells and medium were carefully removed and centrifuged before replating all the cells back into the same dishes (containing any adherent cells) in 3 ml of fresh Stemline II medium containing BMP4 20 ng/ml, VEGF 30 ng/ml, FGF-α 10 ng/ml, SCF 30 ng/ml, IGF2 10 ng/ml, TPO 10 ng/ml, heparin 2.5 μg/ml, and IBMX 100 uM. On day 9, the same day-7 cytokines were topped up in 500 μl of Stemline II per well. On day 10, all the cells were aspirated from the well, centrifuged, and placed in 500 μM Accutase solution 37°C for 3 minutes, and 3 × 10^5 cells were replated per well in IBIT medium (240 ml of Iscove Basal Medium; catalog no. FG-0465; Biochrome, Merck Millipore), 1% bovine serum albumin (BSA; catalog no. G10008-01; Thermo Fisher Scientific Life Sciences), 10 μg/ml insulin (catalog no. N9278; Sigma-Aldrich), 0.2 mg/ml transferrin (catalog no. T0665; Sigma-Aldrich), and 500 μl β-mercaptoethanol (catalog 31550-010; Thermo Fisher Scientific Life Sciences), with the addition of hydrocortisone 10 μM, SCF 50 ng/ml, FMS-like tyrosine kinase 2.5, and LIF 5 ng/ml (catalog no. PHL086; R&D Systems).
receptor ligand (Flt3L) 16.7 ng/ml (catalog no. 300-19; Peprotech, Rocky Hills, NJ, http://www.peprotech.com), BMP4 6.7 ng/ml, interleukin (IL)-3 6.7 ng/ml (catalog 213-13; PeproTech), IL-11 6.7 ng/ml (catalog no. 200-11; PeproTech), erythropoietin (EPO) 3 U/ml (catalog no. 287-TC-500; R&D Systems), and IBMX 100 μM. Every 2 days, the cytokines were topped up in a volume of 500 μl per well. On day 17, the cells were centrifuged and counted, and 1 × 10^6 cells were plated per well in IBIT with the addition of hydrocortisone 10 μM, SCF 20 ng/ml, IGF1 20 ng/ml (catalog no. 100-11; PeproTech), IL-3 6.7 ng/ml, IL-11 6.7 ng/ml, and EPO 3 U/ml. The cytokines were replenished at the same concentrations every 2 days in a volume of 500 μl per well.

Flow Cytometry

The cells were harvested, and 2 × 10^5 cells were placed in aliquots in phosphate-buffered saline (PBS) containing 1% BSA (PBS/BSA) and centrifuged (200g) for 5 minutes. Antibodies were added directly to cell pellets that were then resuspended to a final volume of 100 μl, incubated for 30 minutes on ice, washed in 5 ml PBS/BSA, and resuspended in 300 μl. The samples were analyzed using an LSRFortessa (BD Biosciences, Franklin Lakes, NJ, http://www.bdbiosciences.com) using FACSDiva acquisition (BD Biosciences) and FlowJo analysis software (FlowJo, Ashland, OR, http://www.flowjo.com), or sorted on a FACSARIA (BD Biosciences). Anti-human antibodies included CD71-FITC (catalog no. 11-0719; eBioscience, San Diego, CA, http://www.ebioscience.com), CD43-APC (catalog no. 17-0439-42; eBioscience), glycophorinA (M) (CD235a), efluor450 (catalog no. 48-9884; eBioscience), CD34-PE (catalog no. 12-0349; eBioscience), CD41a-FITC (BD Biosciences), CD45-V450 (BD Biosciences), and CD144-PE (Beckman Coulter, Brea, CA, http://www.beckmancoulter.com). Mouse IgG1 APC isotype control (catalog no. 17-4714-81; eBioscience) and mouse IgG1 PE isotype control (catalog no. 12-4714-81; eBioscience) were used (supplemental online Fig. 2).

CFU-C Assay

Single cells (5 × 10^3 or 10^4) were plated into 1.5 ml of Methocult medium containing SCF, granulocyte colony-stimulating factor (CSF), granulocyte macrophage CSF, IL-3, IL-6, and erythropoietin (StemCell Technologies, Inc., Vancouver, BC, Canada, http://www.stemcell.com) in 35-mm-dish low-attachment plates (Greiner Bio-One), incubated at 37°C in a humid chamber, and scored for hematopoietic colony formation after 12–15 days.

Magnetically Activated Cell Sorting

Magnetically activated cell sorting (MACS) was performed according to the manufacturer’s instructions (Miltenyi Biotec, Bergisch Gladbach, Germany, http://www.miltenyibiotech.com). In brief, the cells were resuspended in PBS containing 2 mM EDTA, incubated in FC blocking agent for 10 minutes at room

Figure 1. Monitoring hematopoietic progenitor production throughout the differentiation protocol. Representative flow cytometry analysis of cells from days 0 to 24 of the differentiation protocol using antibodies against CD34 and CD43 (A) and the quantification of the percentage positive (B) was performed from at least three independent experiments. Absolute cell numbers of the differentiation represented in (A) are shown for days 3 and 10 (C). Representative colonies generated from day 7 (D–IX) and day 10 (D–VIII) cells are shown, CFU-C activity having been assessed by methylcellulose assays on days 7 and 10 (E). These data represent three independent experiments, and bars indicate the SEM. Flow cytometry analysis to assess the expression of Runx1C-GFP in the CD34/CD43 double-positive cell population; Runx1C-GFP cells are shown in green and Runx1C-GFP cells in red (F). Original magnification ×60. Abbreviations: CFU, colony-forming unit; G, granulocyte; GFP, green fluorescent protein; GM, granulocyte macrophage; M, macrophage; Q, quartile.
temperature, combined with microbead-labeled antibody (Miltenyi Biotec; CD43: 130-091-333 or CD34: 130-046-702) for 20 minutes at 4°C, then washed and resuspended in PBS/EDTA. Labeled cells were then applied in 250-μl aliquots to the MACS column, left to stand in the magnet for 5 minutes, and then washed three times with 0.5 ml PBS/EDTA. The cells were released from the column according to the manufacturer's instruction, and the purity of sorted cells was assessed by flow cytometry.

**hESC Transfection**

H1 hESCs were fed with fresh StemPro medium containing 20 ng/ml bFGF (R&D Systems) and 10 μM ROCK inhibitor (catalog no. B-8300; Calbiochem, EMD Millipore, Billerica, MA, http://www.emdmillipore.com) at least 1 hour before electroporation, as described previously [25]. Single cell suspensions were generated using Accutase, then the cells were washed and resuspended in StemPro medium (10^7 cells per 0.5 ml) with 30 μg of linearized pCAG-HOXB4ER<sup>12</sup>-IRES-PURO vector [20]. The cells were electroporated (Gene Pulser Xcell: 320 V, 250 μF; Bio-Rad, Hercules, CA, http://www.bio-rad.com), plated on CELLstart matrix (Thermo Fisher Scientific Life Sciences) in StemPro medium containing 20 ng/ml bFGF (R&D Systems) and 10 μM ROCK inhibitor (catalog no. Y-27632; Calbiochem). Resistant colonies were selected in 0.6 μg/ml puromycin for 10 days, then single colonies were picked, expanded, and screened for expression of HOXB4 by Western blot (supplemental online Fig. 1).

**High-Performance Liquid Chromatography**

High-performance liquid chromatography (HPLC) globin chain separation was performed using a protocol modified from Lapillonne...
Erythroid cells were washed three times in PBS, lysed in 50 μl of 0.1% trifluoroacetic acid (TFA) in water. The cells were then centrifuged at 13,000 g at 4°C for 10 minutes, and the supernatant was collected for HPLC analysis. Globin chain separation was performed by injecting 10 μl of the supernatant onto a 1.0 × 250-mm C4 column (Phenomenex, Macclesfield, U.K., http://www.phenomenex.com) with a 42%–56% linear gradient between mixtures of 0.1% TFA in water and 0.1% TFA in acetonitrile at a flow rate of 0.05 ml per minute for 50 minutes on a HPLC Ultimate 3000 system (Dionex, Thermo Fisher Scientific Life Sciences). The column temperature was fixed at 50°C during analysis, and the UV detector was set at 220 nm. Lysates from adult peripheral blood and fetal liver were used as positive controls. The elution times of the peaks generated were compared with those of the control peaks for identification. The area under the curve was used to calculate the proportion of each globin peak from each sample.

**RESULTS**

**Monitoring Hematopoietic Progenitor Production**

We monitored the production of HPCs throughout the differentiation protocol by flow cytometry and CFU-C assays (Fig. 1). The proportion of cells expressing CD34 and CD43 peaked between day 6 and 10 (Fig. 1A, B), and the appearance of cells carrying these HPC markers coincided with the presence of CFU-C colonies (Fig. 1E). Most CFU-C generated from the day 7 cells were small, primitive-like colonies (Fig. 1Di–3Div) with larger, more robust colonies produced by cells in the day 10 cultures (Fig. 1Dv–1Dviii). The absolute numbers of the CD34+ and CD43+ populations within the cultures at days 3 and 10 were calculated. The most abundant population at day 10 was the CD34+/CD43+ double-positive population (Fig. 1C). To further characterize the HPCs generated in our differentiation protocol, we used the RUNX1CGFP/w hESC line carrying a green fluorescent protein (GFP) reporter under the control of the endogenous Runx1C promoter [24], which is considered to be a marker of definitive hematopoietic stem cells [26]. Most of the double-positive, CD34+CD43+ cells at day 10 also expressed Runx1C-GFP, confirming their definitive progenitor-like phenotype (Fig. 1F). To determine which cell population contained the hematopoietic activity, we used MACS to differentiate ESCs at day 7 according to CD34 expression (Fig. 2A) and at day 10 according to CD43 expression (Fig. 2C) and then analyzed the enriched populations for hematopoietic colony activity (Fig. 2B, 2D).
The maximum number of CFU-Cs was found within the CD34hi population at day 7 and CD43+ population at day 10. Further analyses of the fluorescence activated cell-sorted day 10 cells demonstrated that CFU-C activity was contained within the CD43+CD34+ and the CD43+CD34− (Fig. 2E, 2F).

### Enforced Expression of HOXB4

HOXB4 has been widely reported to enhance the production of hematopoietic cells from murine ESCs; however, its effect on the differentiation of hESCs is less clear. The variable findings from hESC studies likely result from the variety of differentiation strategies and the different systems used to express HOXB4, resulting in differing expression levels [19]. Based on our previous studies, we hypothesized that activation of exogenous HOXB4 would increase the production of HPC during the differentiation process [20] and might also act to promote a more “definitive-like progenitor” capable of differentiating into mature adult-like RBCs that could express adult globins and enucleate efficiently. To assess the effects of HOXB4 in our defined differentiation protocol, we transfected H1 hESCs with a tamoxifen-inducible HOXB4-ER T2 transgene under the control of the CAG promoter, as previously described [20]. We have previously confirmed and reported that the HOXB4-ERT2 fusion protein can localize to the nucleus with the addition of tamoxifen and that it is functional in this context by demonstrating the induced expression of the HOXB4 target gene, Frzb, with the addition of tamoxifen [20]. The expression of the HOXB4-ER T2 protein and the hematopoietic differentiation potential of hESCs carrying this transgene were confirmed by Western blotting and CFU-C assays (supplemental online Fig. 1).

We first tested the effects of HOXB4 activation on the production of HPCs in differentiating hESCs carrying this transgene
in the presence and absence of tamoxifen for 10 days by CFU-C assays and flow cytometry (Fig. 3). We observed a significant increase in the number of CFU-Mix and CFU-granulocyte macrophage colonies in the presence of tamoxifen in cells expressing the HOXB4-ERT2 transgene but not in control H1 ESCs (Fig. 3B). No significant difference was found in the number of CFU-macrophage (CFU-M) or burst-forming units-erythroid (BFU-E) in the presence and absence of tamoxifen in either cell line, indicating that enforced expression of HOXB4 preferentially enhanced the production, proliferation, or survival of multipotent HPCs. Consistent with the CFU-C data, the proportion of CD34+/CD43+ double-positive cells was increased when HOXB4 was activated by tamoxifen (Fig. 3C, 3D). The overall proportion of cells expressing CD43 did not change when HOXB4 was activated; thus, the increase in double-positive CD34+/CD43+ cells resulted from an increase in the number of cells expressing CD34 (Fig. 3C).

To assess the effects of enforced HOXB4 expression on RBC maturation, we generated HPCs in the absence of tamoxifen and then added tamoxifen to the later stages of differentiation (days 10–24; Fig. 4A). We observed no effect of tamoxifen on the overall proportion of erythroid cells that expressed CD235a at day 24, but we noted differences in the proportion of those CD235a+ cells that coexpressed CD71 (Fig. 4B–4D). The CD71 marker is lost as erythroid cells mature, and we noted that the proportion of the less mature CD235a+/CD71+ cells was significantly increased with HOXB4 activation (Fig. 4C). However, no significant difference was found in the proportion of the more mature cells (Fig. 4D). Hemoglobinized cells were observed in cultures in both the presence and the absence of tamoxifen, and the globin profile of these cell populations was analyzed by HPLC at day 24 (Fig. 5A, 5B). b-globin was detected in control fetal and adult blood samples, but hESC-derived RBCs generated predominantly embryonic (α) and fetal (γα and γγ)
production of CFU-C clonogenic system to assess the effects of specific transcription factors for differentiating hiPSCs using the OP9 coculture system has shown that iPSC-derived RBCs can undergo terminal differentiation when transplanted in vivo, the nucleated state would preclude the clinical use of these cells [29]. In contrast, RBCs differentiated in vitro from adult, fetal liver, or umbilical cord blood cells enucleated efficiently, demonstrating that it is possible to create a microenvironment in vitro that will support the enucleation process [30, 31]. Adult stem cell-derived enucleated RBCs have been used in a proof-of-principal clinical transfusion study [31]. Although coculture with fetal liver stromal cells can improve the enucleation efficiency of PSC-derived cells, the proportion of enucleated cells remains prohibitively low, and this type of strategy could not be translated into the clinic. A recent study that analyzed the molecular profile of enucleating cells derived from hESCs identified a microRNA (microRNA-30a) as a potential regulator of the enucleation process, demonstrating that it is also possible to modulate the process by altering the intrinsic genetic environment [32].

From our previous findings that enhanced expression of HOXB4 could exert a paracrine effect in differentiating ESCs [20] and that combining coculture with HOXB4 expression did not have an additive effect [21], we sought to test the hypothesis that enhanced expression of this transcription factor could create a microenvironment that would allow for the production of hematopoietic progenitors capable of differentiating into mature erythroid cells. We used a defined, feeder-free culture system designed for the production and expansion of erythroid cells [22].

Production of Hematopoietic Progenitors

We first monitored the production and phenotype of hematopoietic progenitors produced in this defined stepwise protocol. More than 60% of cells expressed HPC markers between days 8 and 10, and the time course of CD43^+CD34^- cell production correlated with the production of CFU-C clonogenic progenitors. At this stage, CD43^-CD34^- cells predominantly expressed the Runx1C-GFP reporter, supporting their progenitor-like phenotype. The phenotype and potential of PSC-derived hematopoietic progenitors is consistent with that described for differentiating hiPSCs using the OP9 coculture system [33–35]; thus, our robust stepwise protocol is an appropriate system to assess the effects of specific transcription factors on the differentiation process.

Enhanced Expression of HOXB4

We hypothesized that the deficiency in erythroid maturation could result from the primitive nature of hematopoietic progenitors. At this stage, CD43^-CD34^- cells predominantly expressed the Runx1C-GFP reporter, supporting their progenitor-like phenotype. The phenotype and potential of PSC-derived hematopoietic progenitors is consistent with that described for differentiating hiPSCs using the OP9 coculture system [33–35]; thus, our robust stepwise protocol is an appropriate system to assess the effects of specific transcription factors on the differentiation process.

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progenitors that are generated in vitro. We considered that HOXB4 might be a transcription factor that could program hPSC-derived hematopoietic cells to a more definitive phenotype, because it had been shown to confer long-term reconstitution ability on primitive yolk sac cells and mouse ESCs [16, 17]. The results described in previous studies using human PSCs were contradictory, likely because of differences in the expression strategies and differentiation protocols that were used [19]. To resolve the discrepancies in the published reports, we used a defined differentiation protocol without feeder cells and an inducible expression system in which we could control the timing of the expression of HOXB4. We have demonstrated that enforced expression of HOXB4 increases the production of multilineage hematopoietic progenitors but has no significant effect on the production and maturation of RBCs. The erythroid progenitors that were produced in the presence of HOXB4 failed to undergo efficient enucleation and expressed embryonic (α) and fetal (γ) but not adult (β) globin. In addition, we observed a detrimental effect of HOXB4 activity on cell expansion, negating any small increases in cell populations seen by inducing HOXB4.

**CONCLUSION**

The differentiation system we have defined in the present study generates hematopoietic progenitors and is robust enough to test the effects of the numerous transcription factors shown to program hematopoietic development [36–38]. We have demonstrated that HOXB4 increased the proportion of hematopoietic progenitor production. However, enhanced activity of this transcription factor did not promote red blood cell maturation.

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**ACKNOWLEDGMENTS**

We thank the following funding agencies: Wellcome Trust, Scottish Funding Council, the Australian Stem Cell Centre, Stem Cells Australia, the National Health and Medical Research Council (NHMRC) of Australia, and the Victorian Government’s Operational Infrastructure Support program. A.G.E. and E.G.S. are senior research fellows of the NHMRC. The work was performed as part of the Novosang Consortium (available at http://www.novosang.co.uk).

**AUTHOR CONTRIBUTIONS**


**DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST**

The authors indicated no potential conflicts of interest.


