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Mathematical modeling reveals differential effects of erythropoietin on proliferation and lineage commitment of human hematopoietic progenitors in early erythroid culture

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ABSTRACT

rythropoietin is essential for the production of mature erythroid cells, promoting both proliferation and survival. Whether erythropoietin and other cytokines can influence lineage commitment of hematopoietic stem and progenitor cells is of significant interest. To study lineage restriction of the common myeloid progenitor to the megakaryocyte/erythroid progenitor of peripheral blood CD34+ cells, we have shown that the cell surface protein CD36 identifies the earliest lineage restricted megakaryocyte/erythroid progenitor. Using this marker and carboxyfluorescein succinimidyl ester to track cell divisions *in vitro*, we have developed a mathematical model that accurately predicts population dynamics of erythroid culture. Parameters derived from the modeling of cultures without added erythropoietin indicate that the rate of lineage restriction is not affected by erythropoietin. By contrast, megakaryocyte/erythroid progenitor proliferation is sensitive to erythropoietin from the time that CD36 first appears at the cell surface. These results shed new light on the role of erythropoietin in erythropoiesis and provide a powerful tool for further study of hematopoietic progenitor lineage restriction and erythropoiesis.

Introduction

The question of whether extrinsic signals, in particular cytokines, have a deterministic role in lineage commitment of hematopoietic stem and progenitor cells has been much debated. It is difficult to distinguish a definite role in commitment from the well-characterized roles of cytokines in promoting proliferation and survival. Recent work has provided strong evidence that lineage commitment of granulocyte/monocyte progenitors (GMP) to granulocytes or macrophages can be specified by the appropriate colony stimulating factor (CSF)¹⁻³ but it is less clear whether cytokines can control the lineage restriction of less mature stem and progenitor cells. ^{4,5} Common myeloid progenitors (CMPs) are immature hematopoietic progenitors that become restricted to either the erythroid lineage as megakaryocyte/erythroid progenitors (MEP), giving rise to red blood cells and megakaryocytes/ platelets, or the myeloid lineage as GMP, giving rise to neutrophils, eosinophils, mast cells and macrophages. Mutations in regulators of lineage commitment from CMP are often found in leukemia.⁶⁻⁹ highlighting the importance of understanding this key regulatory event.

The erythroid master regulator erythropoietin (epo) is fundamental to the control of both homeostatic and stress erythropoiesis. ¹⁰ Erythroid cell proliferation is dependent on epo during the early S phase of the second colony forming unit-erythroid (CFU-E) division, coincident with the appearance of the erythroid marker glycophorin A (GPA). ^{11,12} Epo is also required for erythroid precursor cell survival

Table 1. Parameters of the mathematical model. Symbols and physiological meaning of the parameters used in the full (37 parameter) mathematical model, and (column 3) their values in the simplified (20 parameter) model.

Parameter	Description	Simplification
α_i	Proliferation rate: CMP generation $i \ge 0$	
δ_{i}	Proliferation rate: MEP generation $i \ge 0$	
$\mu_{\scriptscriptstyle \parallel},\;\mu_{\scriptscriptstyle \parallel}$	Maturation rates (initial, final): CMP to MEP generation 0	
$\mu_{ m l}$	Maturation rate: CMP to MEP generation <i>i</i> >1	$\mu_{i\geq l}=\mu_{\mathrm{F}}$
μi	Logistic function parameters (bias, decay rate): CMP proliferation generation 0	Fixed by dataset A
$Q_{\delta} R_{\delta}$	Logistic function parameters (bias, decay rate): MEP proliferation generation 0	Fixed by dataset A
Q_{μ} , R_{μ}	Logistic function parameters (bias, decay rate): CMP to MEP maturation generation 0	Fixed by dataset A
$ au_{ extsf{C}}, au_{ extsf{M}}$	Proliferation initiation delays: CMP, MEP generation 0	Fixed by dataset A
Δ_i	'b' compartment delay: CMP generation i ≥1	$\Delta_i = \Delta$
Γ_i	'b' compartment delay: MEP generation i ≥1	$\Gamma_{i\! eq} = \Gamma_3$

through signal transducer and activator of transcription (Stat-5) mediated activation of the apoptotic inhibitor Bcell lymphoma-extra large (Bcl-XL). 13 However, the role of epo in controlling proliferation, maturation and survival of the earliest erythroid progenitors is not completely understood. In vivo, epo is essential from the CFU-E stage onward as mutants defective for epo or epo receptor do not produce mature red cells.14 Such mutants are able to produce both early and late erythroid progenitors, burst forming unit-erythroid (BFU-E) and CFU-E, respectively, suggesting that epo signaling is not absolutely required for erythroid commitment. However, a recent study has suggested that high levels of epo can bias commitment toward the erythroid lineage over the myeloid lineage both in vivo and in vitro. 15 To clarify these apparently contradictory findings on the role of epo in lineage commitment and erythropoiesis, we have determined the effect of epo on the kinetics of proliferation and lineage restriction during the maturation from CMP to MEP.

Population dynamics of hematopoietic culture is a complex output generated by the interplay of proliferation, maturation and cell death. Proliferation and cell death can be followed directly using the carboxyfluorescein succinimidyl ester (CFSE) division-tracking dye and Annexin V/propidium iodide staining, respectively, but maturation associated with lineage restriction can only be analyzed indirectly as a measure of the number of mature cells that arise in culture which cannot be accounted for by proliferation or death. To disentangle the individual contributions of these different cell behaviors, we have developed a mathematical model of the early stages of erythroid culture during which cells become committed to the megakaryocyte/erythroid lineage.

Mathematical models have been developed as tools to assist in the analysis of population dynamics during hematopoiesis, ¹⁶⁻¹⁹ and to determine the transcription factor regulatory interactions that control hematopoietic differentiation pathways. ²⁰ To understand how proliferation integrates with lineage commitment and impacts on the overall population dynamics, a number of mathematical approaches for cell division have also been proposed, taking advantage of CFSE division-tracking. In particular, the Smith-Martin model that takes into account progression of cells through the cell cycle, has been successfully used to predict and model population dynamics of *in vitro* erythropoiesis. ²¹⁻²⁵

We have combined immunophenotyping, proliferation analysis and cytokine dependence to refine the analysis of early erythroid culture of CD34⁺ cells isolated from human peripheral blood (PB). Isolated PB CD34+ cells are CMP, and the appearance of detectable surface levels of the erythroid marker CD36 is the earliest identifier of progenitors restricted to the megakaryocyte/erythroid lineage.26 Interestingly, CMP give rise to MEP independently of cell division in erythroid culture and most of the starting CMP become lineage-restricted prior to the onset of proliferation. We have developed an adaptation of the Smith-Martin mathematical model that describes the population dynamics of erythroid culture during lineage restriction from CMP to MEP. The model suggests that cells become responsive to epo as soon as they are committed to the megakaryocyte/erythroid lineage, but epo does not control lineage restriction.

Methods

Antibodies and reagents

Antibodies used were CD34-BV421, CD36-APC, CD90-PE, CD123-PE/Cy5, CD38-APC/Cy7, CD135-PE and CD45RA-BV421 (Biolegend) and CD36-PE, CD45-APC/Vio770, CD235 (GPA)-APC, CD61-APC/Vio770 (Miltenyi Biotec). AnnexinV-BV421 (Biolegend) was used for analysis of dying cells. Monastrol (Merck Chemicals Ltd, UK) was used at 100 μM to block cells in M phase of the cell cycle.

Erythroid culture

Peripheral blood mononuclear cells (PBMCs) were isolated from leukocyte cones by density purification over Histopaque (Sigma) from healthy donors with informed consent. Isolated cells were cultured in erythroid medium (EM); Stem Span (Stem Cell Technologies) + 10 ng/mL stem cell factor (SCF) (for Day 0 to Day 4) and 50 ng/mL (for Day 5 to Day 11), 40 ng/mL insulin-like growth factor1 (IGF-1), 1 ng/mL interleukin-3 (IL-3), 1 µM dexamethasone and with or without 2 U/mL epo (details in *Online Supplementary Methods*). Research was reviewed and approved by Southmead and Bristol Research Ethics Committee Centre (08/H0102/26 and 12/SW/0199, respectively).

Cell cycle analysis

K562 cells were grown in Iscove modified Dulbecco medium + 20% fetal calf serum, fixed in ice cold 70% ethanol and rehydrated

Table 2. Parameter values of the mathematical model.

	α_{0}	α_{1}	Ω_2	α^3	α_{4}	α_{5}	μ_{I}	μ_{f}	
Dataset A Dataset B Dataset C Dataset D	0.018 0.023 0.014 0.012	0.054 0.066 0.045 0.042	0.108 0.120 0.098 0.086	0.181 0.183 0.198 0.174	0.213 0.200 0.264 0.255	0.195 0.185 0.299 0.269	0.016 0.017 0.025 0.020	0.004 0.002 0.003 0.005	
	$\delta_{ extsf{o}}$	$\delta_{\mathtt{1}}$	$\delta_{\scriptscriptstyle 2}$	$\delta_{\scriptscriptstyle 3}$	$\delta_{\scriptscriptstyle 4}$	$\delta_{\scriptscriptstyle{5}}$			
Dataset A Dataset B Dataset C Dataset D	0.128 0.143 0.118 0.088	0.123 0.150 0.121 0.114	0.119 0.124 0.158 0.119	0.127 0.125 0.181 0.146	0.129 0.121 0.279 0.206	0.110 0.125 0.269 0.199 Γ_3			
Dataset A	τ _c 30.394	τ _м 27.705	9.800	14.549	9.817	8.017			
Dataset B Dataset C Dataset D	31.119 29.708 28.841	27.800 26.279 25.765	9.602 10.742 10.623	13.892 14.599 12.453	10.688 10.540 9.521	8.264 10.316 10.893			
	$\delta_{\scriptscriptstyle{6}}$	δ_7	$\delta_{ extsf{8}}$	$\delta_{ extsf{g}}$	$\delta_{ exttt{10}}$	$\delta_{\mathtt{11}}$	$\delta_{\scriptscriptstyle 12}$	$\delta_{\scriptscriptstyle 13}$	
Dataset E Dataset F	0.314 0.234	0.35 0.259	0.386 0.284	0.422 0.309	0.458 0.334	0.494 0.359	0.53 0.384	0.566 0.409	

Rows 1,5 and 9: parameters fitted using dataset A, used in simulations shown in Figure 5A-D, and Online Supplementary Figures S4 and S5 (red lines). Rows 2,6 and 10: re-identified parameters used for validation (dataset B), simulation shown in Online Supplementary Figure S5 (black lines). Rows 3,7 and 11: fitted parameters on the control data sets (dataset C), simulations shown in Figure 6C, E and F, and Online Supplementary Figure S6A-C. Rows 4,8 and 12: fitted parameters on the minus epo data sets (dataset D), simulations shown in Figure 6C, E and F, and Online Supplementary Figure S6A-C. Rows 13 and 14: parameters 6-13 are the extrapolated proliferation rates used to predict the MEP cell counts from 90 h to 186 h (Figure 6E, datasets E and F.plus and minus epo, respectively), calculated by assuming a linear relationship between MEP proliferation rates above. Proliferation rates are divisions per hour and maturation rates are transitions per hour.

in phosphate buffered saline (PBS). Fixed cells were stained in PBS (Sigma) + 0.1% Triton X-100 + 10 $\mu g/mL$ propidium iodide + 100 $\mu g/mL$ RNase and analyzed on the MACSQuant VYB flow cytometer.

Flow cytometry and fluorescence assisted cell sorting

Flow cytometry and fluorescence assisted cell sorting (FACS) was used to analyze cells stained with anti-human-specific antibodies on the MACSQuant and post-acquisition analysis was carried out with FlowJo v.7.6.5 for proliferation analysis and FlowJo v.X0.7 for all other analysis (details in *Online Supplementary Methods*).

CFSE tracking

CD34 $^{\circ}$ cells were labeled on the day of isolation with CFSE (Biolegend) at 2 μ M in PBS for 15 min at 37 $^{\circ}$ C. Excess CFSE was quenched by incubating in Stem span + 10 $^{\circ}$ 6 fetal calf serum for 5 min at 37 $^{\circ}$ C. Cells were washed in PBS and transferred into EM at a density of 2x10 $^{\circ}$ 6 cells/mL in multi-well dishes so that each time point was taken from a separate well. For extended cultures beyond Day 4, cells were fed by 1:2 partial medium change into EM with 100 ng/mL SCF.

Results

CD36 expression marks megakaryocyte/erythroid restricted progenitors

At the time of isolation, CD34 $^{+}$ cells from human PBMNC, are 85% CD34 $^{+}$ CD38 $^{+}$ IL3R α -CD45RA-CD90 $^{-}$ CD45lo (Figure 1A), previously defined as CMP 27,28 and 15% of CD34 $^{+}$ cells are CD34 $^{+}$ CD45lo IL $^{-}$ 3Rlo CD38 $^{+}$ CD90 $^{-}$ CD45RA $^{+}$, GMP as previously described, 29,30 which do not persist in erythroid culture (*Online Supplementary Figure S1*). CD34 $^{+}$ cells cultured in serum-free medium supplemented with appro-

priate cytokines including SCF, IL-3 and epo mature into erythroid cells. The appearance of erythroid precursors and erythroblasts is easily detected by the appearance of the erythroid marker GPA (Ter119) at Day 7 to Day 10 of culture, depending on conditions. To identify the less mature erythroid progenitors, MEP, at the point of lineage restriction from CMP, prior to GPA expression, we examined the change during culture of the previously defined markers interleukin-3 receptor (IL-3R/CD123) and Fmslike tyrosine kinase 3 (Flt-3/CD135), which are present at low levels on CMP and GMP and absent on MEP isolated from human bone marrow and cord blood. 27,31,32 Surface levels of both markers increase over the first two days in culture and then decrease to Day 4 on the bulk population, so they do not provide useful markers for early lineage restriction in erythroid culture (Figure 1B). The erythroid marker CD36 was originally identified as a cell surface marker of erythroblasts33 and later shown to be detectable prior to the appearance of the erythroid differentiation marker GPA.³⁴ At isolation, PB CD34+ cells do not express CD36, but give rise to CD34⁺CD36⁺ cells within the first few days of culture (Figure 1C). To determine whether surface expression of CD36 marks the lineage restriction event, CD34+ cells were cultured for 2-4 days and then separated by FACS into CD34+CD36- and CĎ34⁺CD36⁺ populations. CD34⁺CD36⁻ cells give rise to both erythroid (burst forming unit-erythroid, BFU-E) and myeloid [colony forming unit-granulocyte/monocyte (CFU-GM)] colonies in methylcellulose, indicating these cells are the CMP population. CD34⁺CD36⁺ cells give rise only to BFU-E (Figure 2A). BFU-E colonies contain small numbers of megakaryocytes (Figure 2B) consistent with the assignment of CD34+CD36+ cells as bipotent MEPs. This is further supported by the finding that CD34+CD36+ cells are CD61¹⁶ (Figure 2C, population I,) reflecting megakaryocyte potential.36

Maturation of CMP to MEP is independent of cell division

Previous work had suggested that PB CD34⁺ cells divided asymmetrically dependent on unequal distribution of the determinants Notch and Numb.⁵⁷ One possible explanation for the observed population dynamics of erythroid culture, in particular the retention of CMP concurrent with MEP production (*Online Supplementary Figure S2*), is that CMP might divide asymmetrically to produce 1 MEP and 1 CMP. To test this, we first examined whether maturation of CMP to MEP was coincident with cell division by following proliferation of PB CD34⁺ cells by CFSE tracking. Cells were CFSE-labeled at the time of isolation and the number of cells in each generation was determined separately for CMP and MEP by gating on CD36. After 24 h in culture, all

cells in both CMP and MEP populations remain in generation 0 [G(0)] and do not progress to G(1) until the second day of culture, indicating a lag phase for both CMP and MEP from the time of isolation to the initiation of proliferation between Days 1 and 2 (Figure 3A, top panels). Maturation of CMP to MEP begins immediately, as is clear from the emergence of CD36* MEP on Day 1, before the appearance of G(1) cells (Figure 3B). This indicates that maturation does not require cell division, at least for G(0) CMP, and rules out the possibility that MEP arise from CMP by asymmetric division. This was confirmed by demonstrating that inhibiting cell cycle progression with Monastrol does not block maturation of CD34* cells to MEP (Online Supplementary Figure S3).

By Day 2, 75% MEP and 24% CMP have undergone a

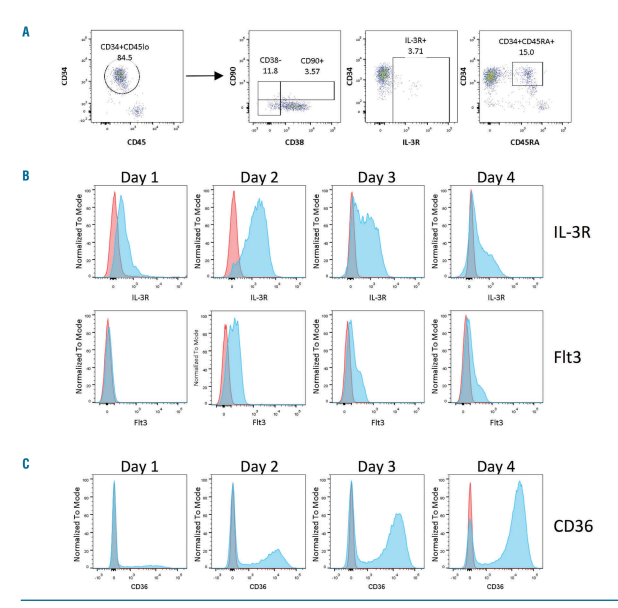


Figure 1. Immunophenotyping of peripheral blood (PB) CD34* cells in erythroid culture. (A) Surface expression of CD34, CD45, CD38, IL-3R (CD123), CD45RA and CD90 on PB CD34* cells on the day of isolation. All populations were gated first on Propidium iodicie incorporation and scatter properties. Cells were gated for CD34* and CD45RA were analyzed on the total live population. Gates were set with isotype controls. (B) Surface expression of IL-3R and Flt3 on CD34* cells cultured in EM for four days. Stained cells in blue and isotype controls in orange. Expression of both markers increases in the total population during the first three days of culture and then decreases. (C) Surface expression of CD36 over four days of culture demonstrating that CD36* and CD36* cells comprise distinct populations. Stained cells in blue and isotype controls in orange.

first division and are in G(1) (Figure 3A). Overall, CMP proliferation is heterogeneous with an even distribution of cells over 5 generations from G(0-4). By contrast, all of the MEP cells divide within the first three days as indicated by the absence of cells in G(0) and, by Day 4, 84% MEP are found in 3 generations G(3-5) with a distinct peak (35%) in G(4), suggesting a largely synchronously proliferating population (Figure 3C).

To determine whether cell death has a significant effect on population dynamics, dying and dead cells were quantified over a 4-day time course by Annexin V staining and Propidium Iodide incorporation, respectively. The total number of dead cells was insignificant compared to live cells (Figure 3D). Annexin V staining was undetectable.

Mathematical model of population dynamics of erythroid culture

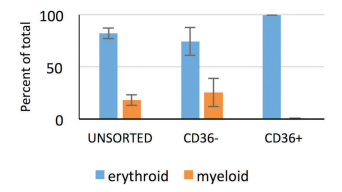
Understanding the role of extrinsic factors on CMP to MEP maturation and progenitor proliferation requires a means of measuring the relative parameters independently, which is not possible using population dynamics alone. Instead, we have developed a mathematical model that simulates the population dynamics, in terms of CMP and MEP numbers, dependent on a set of parameters (Table 1) derived from our experimental data, which include total cell number, CMP:MEP ratio and cell division by CFSE tracking with sampling every eight hours during the first four days of culture

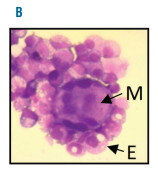
Our model is an adaptation of the Smith-Martin model. It separates the cell cycle into two distinct compartments

'a' and 'b', representing the G_1 ('a') and the combined S, G_2 and M phases ('b') of the cell cycle. We incorporated maturation of CMP to MEP, as shown schematically in Figure 4A. Cells originate in the 'a' compartment, with their transition governed by a stochastic process. This gives rise to rates α_i and δ_i for generation i CMP and MEP cells, respectively, transitioning to the 'b' compartment, and rate μ_i for generation i CMP maturing to generation i MEP. Cells remain in the 'b' compartment for a fixed time (Δ_i and Γ_i for CMP and MEP, respectively) before they divide and then re-enter the 'a' compartment as a next generation cell.

In order to simplify the model whilst maintaining the key dynamics that relate to lineage restriction, we made the following assumptions based on data presented in Figure 3: 1) death rate plays an insignificant role over the time course; 2) cell maturation initiates prior to the onset of CMP and MEP proliferation and, thus, does not require transition through the cell cycle; 3) GMP are excluded from the analysis as this population does not contribute significantly to the population dynamics in the first four days of culture (Online Supplementary Figure S1). The CMP and MEP populations in each generation evolve according to a set of coupled delay differential equations (Figure 4B). To account for the observation that maturation initiates prior to the onset of proliferation (Figure 3B), we allowed the first generation proliferation and maturation rates to be time dependent. Constant rates for proliferation and maturation in G(0) do not provide a good qualitative or quantitative match to data, while piecewise constant rates (a single step function for each rate) substantially improve

A Colony forming capacity of sorted cells





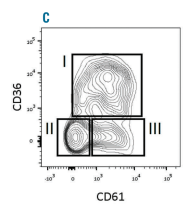


Figure 2. Surface expression of CD36 marks the transition from CMP to MEP in erythroid culture. (A) CD34⁺ cells cultured in EM for two or four days were sorted by FACS into CD34*CD36- and CD34*CD36* populations. Sorted cells were cultured in methylcellulose for two weeks and colonies enumerated. Results are from $\ensuremath{\mathsf{3}}$ independent experiments, 2 tests per experiment. Error bars are standard deviation. (B) Cytospin from an individual erythroid colony stained with May Grünwald/Giemsa showing a megakaryocyte (M) with multiple nuclei and surrounding erythroblasts (E). (C) The megakaryocyte marker CD61 is absent on cells at isolation (data not shown) and heterogeneous at Day 2 with low expression on CD36+ MEP (population I, 46% CD34+CD36+CD6110/-) and largely absent on CD36⁻ CMP (population II, 37% CD34⁺CD36⁻ CD61-). There is a small population of CD61- cells (population III, 17% CD34+CD36+CD61-) which may be megakaryocyte progenitors directly descended from CMP as previously described for mouse multipotent progenitors.

the fit. To maintain physiological plausibility, we model G(0) rates as smoothed step functions, in the form of generalized logistic functions (Figure 4C).

We then used optimization routines to find the set of parameters that minimizes the root mean square error (RMSE) to the experimental data (CFSE tracking) (see Online Supplementary Methods). The system was fitted to experimental data (Online Supplementary Table S1, dataset A) initially in the most general case, where all 37 parameters are generation dependent (Online Supplementary Table S2). This provided insight as to the system behavior and allowed us to make simplifications with no significant

increase in RMSE (see *Online Supplementary Methods*), and reduce the system to 20 parameters: 12 proliferation rates $(\alpha_{(0.5)}, \delta_{(0.5)})$, 2 maturation rates (μ_l, μ_F) , 4 compartmental delays $(\Delta, \Gamma_{(1.5)})$, and 2 proliferation initiation delays (τ_C, τ_M) .

We observed very good quantitative agreement between model and experiment for all generations (Figure 5A and B and *Online Supplementary Figure S4A and B*). To quantify the experimental error, standard deviation was calculated and normalized over total cell counts (shaded regions in Figure 5B). The standard deviation found within the experimental data was 222.44 (3.07%) for CMP, and 279.46 (2.75%) for MEP. The RMSE error between exper-

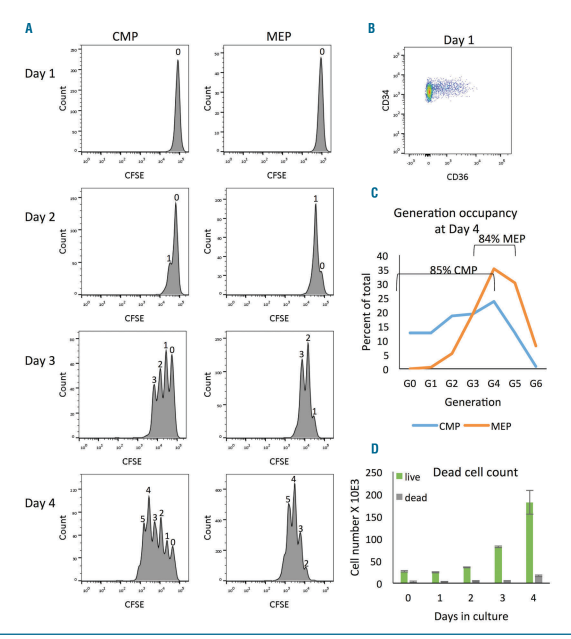


Figure 3. CFSE tracking highlights key features of early erythroid culture. CD34° cells were labeled with CFSE on the day of isolation and cultured in EM for four days. (A) Cells were gated on CD34°CD36° (CMP) or CD34°CD36° (MEP) and generation occupancy was determined by flow cytometry. Numbers above peaks indicate generation. After one day in culture, cells have not divided and all cells are in G(0), (top panels). Proliferation begins for CMP and MEP between Day 1 and Day 2 as demonstrated by the appearance of a peak at half the mean fluorescence intensity (MFI) of the original peak (G(0) MFI=63000 for G(0) and MFI=34000 for G(1)). This generation analysis was done manually in FlowJo vX0.7. (B) CD36 staining on Day 1 of the same cells as shown in the left panels demonstrates the appearance of CD36° cells prior to the onset of proliferation. (C) Generational occupancy at Day 4 shows that MEP proliferation is more synchronized than CMP with 84% MEP within 3 generations, G(3-5), and 85% CMP evenly distributed in 5 generations, G(0-4). (D) Cell death is insignificant during the first four days of culture as measured by propidium iodide staining to identify dead cells (gray) as compared to live cells (green). Error bars are standard deviation.

imental data and model predictions for optimal parameter values (Figure 5B) was 101.02 (1.38%) for CMP and 138.29 (1.85%) for MEP. Table 2 shows the rates (divisions per hour) and delays (hours) obtained from the optimization algorithm.

The model was validated using a second, independent experimental data set (dataset B) in two different ways (Online Supplementary Methods and Online Supplementary Figure S5), and indicated high confidence in the model and parameter values. Further in silico experiments were conducted on an extensive set of simulated data, confirming robustness of the parameter identification methods used (Online Supplementary Methods and Online Supplementary Table S3).

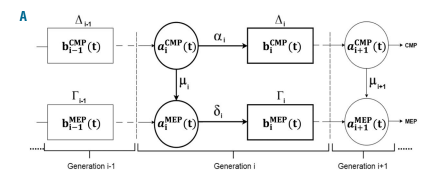
The identified parameters for dataset A (Table 2) show that the average MEP proliferation rate (δ) is 0.12 divisions per hour, and that this rate does not vary greatly from the onset of proliferation to Day 4 (Figure 5C). The rate of proliferation of CMP cells is initially 10% of the proliferation rate of MEP, and increases with each generation such that CMP transit from G(5) to G(6) is approximately twice the average MEP rate (Figure 5C). The maturation rate for G(0) ($\mu_0(t)$) decreases over time, from 0.016 to 0.004 transitions per hour (Figure 5D). Importantly, the maturation rate, even at the beginning of the culture when it is at its highest, is an order of magnitude smaller than the MEP proliferation rate. These parameters show important features of the relative contributions of proliferation and maturation to overall population dynamics at different stages in the culture: while maturation plays a crucial role at the beginning, proliferation is the main effector of later culture stages.

The role of epo in maturation from CMP to MEP

To investigate a possible role for epo in the earliest stages of erythropoiesis in vitro, CD34+ cells were isolated and cultured under standard conditions or in the absence of epo. After one day in culture, prior to the onset of proliferation, 20% CMP have matured to MEP under both conditions (Figure 6A), suggesting that CMP to MEP transition during the first 24 h is not responsive to epo in G(0). However, over nine days in culture, there is a dramatic difference in the number of MEP in control and minus epo cultures and by Day 9 there is over 20 times more MEP

produced in cultures with epo (Figure 6B).

The lower MEP production in the absence of epo could be due to defects in maturation, proliferation, or both. To resolve the individual effects of epo on maturation and proliferation, we used the model to fit parameters from generational CMP and MEP counts of 4-day cultures with and without epo (datasets C and D). The results are shown in Figure 6C and Online Supplementary Figure S6A and B. Parameter fitting shows that the proliferation rate of MEP (δ) is 21% lower in the absence of epo, whereas there is less than 10% difference in proliferation rates of CMP (α) (Table 2). The traces from CFSE tracking also support this result in that a distinct delay in generation progression for MEP is detectable in cultures without epo (Figure 6D). Also, CFSE tracking shows no significant effect of epo on CMP generation occupancy (Figure 6D); this result, taken together with insignificant change in proliferation rate of CMP (Table 2), indicates that CMP proliferation is not epo-sensitive. Model validation, using fitted parameters for δ to reproduce MEP amplification in 2 additional experimental datasets (Online Supplementary Table



$$\begin{split} \mathbf{B} & \quad \dot{C}_0(t) = -\mu_0(t)C_0(t) - \alpha_0(t)C_0(t) \\ \dot{C}_i(t) &= 2\alpha_{i-1}C_{i-1}(t-\Delta_i) - \mu_iC_i(t) - \alpha_iC_i(t) \\ \dot{M}_0(t) &= \mu_0(t)C_0(t) - \delta_0(t)M_0(t) \\ \dot{M}_i(t) &= 2\delta_{i-1}M_{i-1}(t-\Gamma_i) + \mu_iC_i(t) - \delta_iM_i(t) \end{split}$$

$$\begin{split} \mathbf{C} & \alpha_0(t) = \frac{\alpha_0}{(1 + Q_\alpha \mathrm{e}^{-R_\alpha(t - \tau_\mathrm{C})})^2} \\ & \delta_0(t) = \frac{\delta_0}{(1 + Q_\delta \mathrm{e}^{-R_\delta(t - \tau_\mathrm{M})})^2} \\ & \mu_0(t) = \mu_\mathrm{I} + \frac{\mu_\mathrm{F} - \mu_\mathrm{I}}{\left(1 + Q_\mu \mathrm{e}^{-R_\mu(t - \tau_\mathrm{C})}\right)^2} \end{split}$$

Figure 4. Mathematical model of the system. (A) Schematic diagram of erythroid cell culture. The cell cycle is modelled as a 2-stage process, composed of compartments 'a' approximating the $\mbox{\bf G}_{\mbox{\tiny 1}}$ phase of the cell cycle, and 'b' approximating the $S/G_2/M$ phases. CMP and MEP both originate in the 'a' compartment. Their transition is governed by a stochastic process. with rates α_i and δ_i for generation i CMP and MEP cells. respectively, to transition to the 'b' compartment, and a rate μ_i of maturing (transitioning) from CMP into samegeneration MEP cells. Cells remain in the 'b' compartment for a fixed time (Δ_i and Γ_i for CMP and MEP, respectively) before they divide and then re-enter the 'a' compartment as a next generation cell. (B) The size of the CMP and MEP populations in each generation at each time point are given by 4 delay differential equations (DDEs) where $C_i(t)$ and $M_i(t)$ are the generational populations of CMP and MEP, respectively. (C) G(0) proliferation and maturation rates are time-dependent, taking the form of generalized logistic functions, where P_{a} is the maximum proliferation rate for G(0) CMP, P_{a} is the maximum proliferation rate for G(0) MEP, and μ_1 and $\mu_{\rm F}$ are the initial and final maturation rates for G(0). The parameters $Q_{\rm a}$, $R_{\rm a}$, $Q_{\rm b}$, $R_{\rm b}$, $Q_{\rm \mu}$, $R_{\rm \mu}$, $\tau_{\rm C}$ and $\tau_{\rm M}$ control the dynamics of the generalized logistic functions and were optimized for. Equations were solved using MATLAB®

S1, datasets E and F), further supports this result (*Online Supplementary Figure S6C*).

We have used the parameters calculated from 4-day cultures with and without epo to simulate MEP population expansion to Day 8 (Table 2 and Figure 6E, datasets E and F). For control cultures, there is good agreement between the predicted cell numbers and experimental data. For cultures without epo, there is good agreement for the first 5-6 days (up to time 130 h in Figure 6E), and then the experimental data show a significantly slower proliferation than predicted. This suggests that there is a change in epo sensitivity between Days 5 and 6, after which MEP proliferate very slowly without epo.

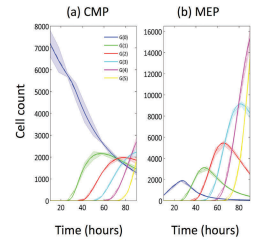
The maturation rate for G(0) ($\mu_0(t)$ (Table 2) is not significantly different with or without epo (datasets C and D). Although the final maturation rate is slightly higher in the absence of epo, this cannot account for the decrease in MEP formation, suggesting that epo effects on population dynamics and MEP production are not mediated by

altered maturation rates. To determine whether epo confers an erythroid bias, we used CFSE tracking to quantify CMP maturation and showed that epo does not affect the total number of CMP that mature to MEP (Figure 6F). These results provide strong evidence that neither the rate nor the bias of the CMP to MEP transition is influenced by epo.

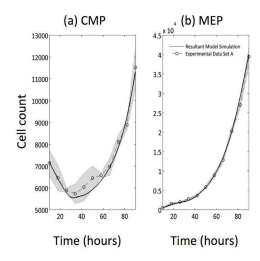
Discussion

Understanding the fundamental mechanisms controlling lineage commitment, proliferation and survival during erythropoiesis is becoming an achievable goal thanks to the advent of novel high throughput technologies to explore the transcriptome, proteome and epigenome. It is order to maximize the specificity of the information afforded by these techniques, it is essential to start with well-defined populations. A recent report pro-

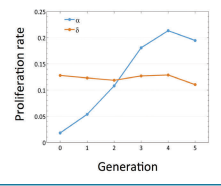
Model results (generational cell counts)



Model results (total cell counts)



CMP vs MEP proliferation rates



G(0) CMP maturation rate

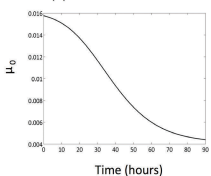


Figure 5. Fitted model simulations provide proliferation and maturation parameters and recapitulate population dynamics. (A) Fitted model simulations (solid lines) of CMP (left panel) and MEP (right panel) cell numbers in each generation. Experimental data (dataset A) are dashed lines, \pm standard error (shaded regions). (B) Model simulation of total CMP (left panel) and MEP (right panel) cell numbers using model derived parameters (solid lines) together with experimental data (dashed lines) \pm standard error (shaded regions). (C) Proliferation rates dataset A for CMP (α) in blue, and MEP (δ) in orange, as divisions per hour over the first 5 generations. (D) Changes for dataset A in maturation rate (μ 0) of G(0) CMP in the first 90 h of culture expressed as CMP to MEP transitions per hour.

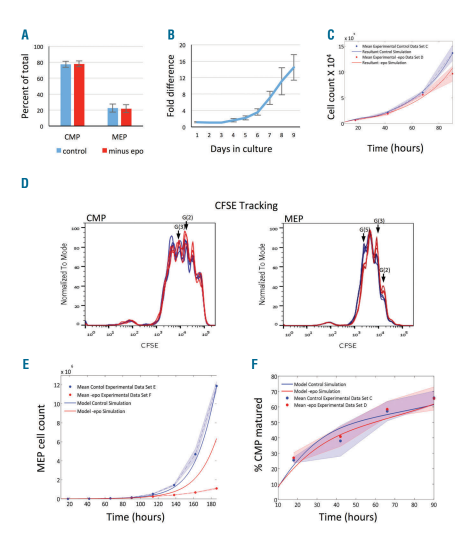


Figure 6. MEP proliferation rate is epo-sensitive but CMP maturation is independent of epo. (A) Cell numbers for CMP and MEP after one day in control (blue) and minus epo (red) culture ± standard deviation. Data are from 4 experiments, 3 samples per experiment. (B) Relative difference in MEP numbers in control and minus epo cultures (fold difference = N. control/N. minus epo MEP ± standard deviation) from 3 experiments. (C) Model fitting simulations for control (blue) and minus epo (red) cultures compared to experimental data from datasets C and D (dots). Shaded regions represent standard error in the experimental data. (D) CFSE in CMP and MEP after four days in control culture (blue) or minus epo (red). Traces from 3 samples are overlaid. Generations 2, 3 and 5 are labeled. (E) Extrapolation of MEP cell counts from datasets E and F to Day 8 (186 h) using best-fit parameters in Table 2 for both control and minus epo populations. Additional proliferation rates (δ_{6-13}) predicted by assuming a linear relationship between the optimized proliferation rates, found from the Day 0-4 fitting for G(0) to G(5) MEP, allowing us to extrapolate forward for G(6) onwards. (F) Maturation over four days expressed as percentage of CMP matured from experimental data (circles) and model predictions (lines). Experimental values were calculated as the number of CMP taking into account cell division. CMP matured = N. CMP in G(0) + sum over 6 generations N. CMP G(i)/2i. Model predicted maturation was tracked from all generations of CMP throughout the simulations.

vided an excellent protocol to identify populations at distinct stages of erythroid differentiation, ⁴⁶ but early erythroid culture has proved more challenging. To achieve this, it is necessary to measure the transition event under experimental conditions where proliferation and survival can be controlled and/or measured simultaneously. We identified CD36 as a cell surface marker that distinguishes CMP and MEP as distinct populations.

Identification of CD36 as an MEP marker provides a powerful tool to investigate the influence of extrinsic factors on lineage restriction of CMP to MEP. Epo is essential for steady-state and stress erythropoiesis and is used therapeutically to increase red cell production in clinical anemia. It is of considerable interest, therefore, to understand the precise role of epo in hematopoiesis. There is a significant body of work demonstrating that an essential role of epo is to promote proliferation of erythroid progenitors and survival of differentiating erythroblasts. 47,48 This raises the question of whether epo acts on less mature progenitors. Early work from Lodish et al.14 showed that epo and the epo receptor are not required to generate early erythroid-specified progenitors in vivo, suggesting that epo does not have an instructive role in early hematopoiesis. On the other hand, a more recent study¹⁵ showed that early progenitors are responsive to epo, and that high levels of epo suppress GMP formation through transcriptional reprogramming, both in vivo and in vitro. Interestingly, this study showed that although there is a relative increase in erythroid progenitors compared to non-erythroid progenitors, the absolute number of erythroid progenitors is not affected by high levels of epo in vivo. Our results using mathematical modeling correspond well with these studies, demonstrating that the time-dependent maturation rate of CMP is not affected by epo. Importantly, equal numbers of CMP convert to MEP with and without epo, further supporting the idea that epo does not bias CMP commitment toward the erythroid lineage. Population dynamics of cord blood CD34+ cells in erythroid culture show similar dynamics to PB CD34+ cells (Online Supplementary Figure S7) suggesting that these results are likely to be applicable to CMP from other sources. This provides validation for the proposed model in demonstrating consistency with in vivo data.

It is important to note that, while our results clearly indicate that the MEP population is homogeneous, the CMP population behaves heterogeneously in terms of proliferation. It is, therefore, possible that there is an earlier restricted progenitor within the CMP population that is indistinguishable from other CMP using known markers. Developing a transcriptomic fingerprint of CD34*CD36*

cells would allow us to investigate this possibility using single cell analysis applied to the CMP population.

Although it is well established that epo is required for proliferation of late erythroid progenitors, a possible influence of epo on earlier progenitors has not been described. To test this, we determined the progenitor proliferation rates using the model. This showed that, from the initial stages of megakaryocyte/erythroid lineage commitment, cells are epo responsive and proliferation is slower in the absence of epo. By disentangling the independent contributions of proliferation and maturation to population dynamics, the model allowed a high-resolution quantification of proliferation rate that revealed the sensitivity of MEP to epo. This means of quantification will be highly beneficial to the efforts to amplify erythroid cells in culture for the production of red cells for clinical transfusion; a 4-5 day culture significantly decreases the cost and simplifies screening of changes in culture conditions as compared to end point analysis from 2-3 week cultures.

Model simulations using a proliferation rate derived from the first four days of culture and extrapolating to Day 8 show that the derived proliferation rate faithfully reproduces the experimental data. This confirms the power of the model to describe erythroid culture. Interestingly, simulating MEP amplification to Day 8 in the absence of epo using the proliferation rate derived from the first four days of culture shows a marked change after Day 5. This suggests that there are two MEP stages with different epo sensitivity: an early stage (Days 1-4) with a small increase in proliferation rate in response to epo and a later stage (Days 4-8), which is increasingly dependent on epo for proliferation. The increased resolution of cell behavior parameters given by the mathematical model provides a valuable tool to further investigate both the mechanism of lineage restriction and proliferation behavior of hematopoietic progenitors in culture.

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