Cell cycle dynamics and complement expression distinguishes mature haematopoietic subsets arising from hemogenic endothelium

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ABSTRACT

The emergence of haematopoietic stem and progenitor cells (HSPCs) from hemogenic endothelium results in the formation of sizeable HSPC clusters attached to the vascular wall. We evaluate the cell cycle and proliferation of HSPCs involved in cluster formation, as well as the molecular signatures from their initial appearance to the point when cluster cells are capable of adult engraftment (definitive HSCs). We uncover a non-clonal origin of HSPC clusters with differing cell cycle, migration, and cell signaling attributes. In addition, we find that the complement cascade is highly enriched in mature HSPC clusters, possibly delineating a new role for this pathway in engraftment.

Introduction

Hemogenic endothelium (HE) exists during a narrow window (~E9.5 to E11.5) in development.1–7 During this period, these specialized endothelial cells have the ability to generate the first definitive haematopoietic stem and progenitor cells (HSPCs) in the embryo. Hemogenic vascular beds are contained intra-embryonically in the aorta-gonad mesonephros region (AGM),8–10 as well as extra-embryonically in the placenta,11,12 yolk-sac,13–16 and umbilical and vitelline arteries.9,17,18 Hemogenic vascular beds are uniquely characterized by the presence of haematopoietic cell clusters, a distinct group of haematopoietic cells that bud off of the endothelial wall.1,4,10

The dorsal aorta is the most extensively studied hemogenic vascular bed, and is noted to contain several clusters of haematopoietic cells termed intra-aortic haematopoietic clusters (IAHCs).1,2,19,20 These structures have been observed to be as large as ~30 cells and are thought to initially form from a Runx1 dependent endothelial-to-haematopoietic transition (EHT),5,19,21,22 and contain both haematopoietic stem and progenitor cells (hence HSPCs). Cells comprising IAHCs can be identified by surface marker expression of CD31 and CD117, as well as lack of expression of germ line marker SSEA-1.19 Whole embryonic AGM regions have been shown to engraft irradiated adult hosts2,23,24 at E11, but also E10 albeit with very low incidence.24

Further transplantation experiments of isolated IAHC cells have been performed with sorted SSEA-1+/CD31+/CD117+ cells (IAHC populations) from E11.5 AGMs and shown to engraft irradiated adults.19 Thus, the E11 IAHC cell population contains both definitive HSCs and progenitor cells. However, transplantation of the same sorted IAHC cell populations from E10.5 AGMs shows undetectable repopulation activity in adult irradiated mice,19,24,25 but does demonstrate engraftment in neonates.20 The neonatal engraftment at E10.5 suggests that the IAHCs also contain haematopoietic stem and progenitor cells, but that the HSC population is immature or a pre-HSC population.20,23 Overall, these data demonstrate that IAHC populations may only possess adult engraftment potential later in the HE time window, suggesting a maturation process from E10 to E11 that allows for adult engraftment.20,25,26 Thus, during this transient period in development, control of pre-HSC maturation may be spatially and temporally regulated. Understanding the establishment of this multi-step maturation process is critical to efforts of generating HSCs in vitro for clinical applications. To begin to understand IAHC generation and subsequent maturation steps, we evaluated IAHC cell proliferative capacity, cell cycle attributes, clonal origins from HE cells, as well as possible critical signaling pathways. Here we propose that the pre-HSC maturation process includes changes in cell cycle dynamics and activation of the complement system.

Results

Analysis of IAHC cell cycle kinetics reveals a proliferative population following HE emergence

We set out to determine how newly emerged haematopoietic cells form a cluster. To address this, we first defined an IAHC as a group of 3 or more haematopoietic cells attached to the...
endothelium. To identify IAHCs, we used PU.1 as a nuclear marker of IAHCs during E10.5 and E11.5\(^{27}\) (Fig. S1A–C). To determine IAHC size and distribution, we examined the frequency of IAHCs according their size (cell number) at E10.5, the time point that corresponds to the peak of IAHC formation.\(^{19}\) Quantification of IAHCs at E10.5 revealed a wide distribution of IAHC size that ranged from IAHCs with 3 cells to 47 cells (Fig. S1D). Interestingly, at E11.5 the distribution range of IAHC size narrowed from IAHCs with 3 cells to 11 cells (Fig. S1E). Overall, the data suggest that most IAHCs form at E10.5, the time point thought to be peak of IAHC formation.\(^{19}\)

To understand how some IAHCs could contain >20 cells, we next determined the extent cell proliferation contributed to the large number of cells within each IAHC. To determine if IAHCs are proliferative, we injected the nucleoside analog 5-ethynyl-2'-deoxyuridine (EdU) in pregnant females at E10.5 and separately at E11.5, and evaluated the embryos 2 hours later (Fig. 1A). Immunofluorescence for EdU distinguishes cells in the S-phase from cells in other phases of the cell cycle,\(^{28,29}\) and Ki67 labels cells in all active phases of the cell cycle (except G0).\(^{30}\) Our analysis reveals several cells in the IAHCs display markers of proliferation (Fig. 1B and C). Moreover, quantification of IAHCs that express these markers show that ~90% of cells in IAHCs were Ki67\(^{+}\), and ~42% of these cycling IAHC cells were EdU\(^{+}\) (Fig. 1D and E) at E10.5. These percentages were similar upon examination of IAHCs at E11.5 (Fig. 1D and E). Together, these data show that these 2 time points in the hemogenic window contain cells within IAHCs that are proliferative.

Since IAHCs contain proliferating cells, we next determined their average generation time (cell cycle length). We labeled IAHCs with a sequential pulse of nucleoside analogs,
5-ethyl-2'-deoxyuridine (EdU) followed by 5-bromo-2'-deoxyuridine (BrdU). The sequential administration of these compounds allows for the determination of the cell cycle kinetics of proliferating IAHC cells by separating cells into early (EdU–BrdU+), mid (EdU+BrdU+), and late (EdU+BrdU–) S phases3,13,32 (Fig. 1F). To validate the system, we evaluated EdU and BrdU in the neural tube at E10.5 and noted comparable numbers to a previous report31 (Fig. S2A and BrdU in the neural tube at E10.5 and noted comparable numbers to a previous report31 (Fig. S2A–C). To adapt the system for IAHC evaluation, we used PU.1 as a nuclear marker of IAHC cells, which we have previously determined to contain about ~90% Ki67+ cells, as an estimate of the total number of proliferating cells (Fig. 1G and Fig.S1A–C). Analysis of the dual pulse labeling revealed that a proportion of the cells within IAHCs at E10.5 are actively in the S-phase (versus 42% by Ki67/EdU staining, Fig. 1E), while a proportion of cells within IAHCs at E11.5 are positioned in other phases of the cell cycle (Fig. 1H, E10.5 vs. E11.5, p < 0.001). Furthermore, we estimated the total length of the cell cycle (Tc) of IAHC cells at E10.5 to be 5.38 hours and the S-phase length (Ts) at 0.78 hours (Fig. 1F and I). To gain further insight into the cell cycle kinetics of IAHC cells, we extended this analysis to E11.5 and found that the Tc lengthens to 8.16 hours (Fig. 1I, Tc E10.5 vs E11.5, p < 0.003), while Ts remains the same (Fig. 1I, Ts E10.5 0.78 h vs. E11.5 0.74 h). Overall, these data suggest that IAHC cells are capable of dividing 4.5 times within the 24 hour period of time leading to the peak of the hemogenic window (E10.5), such that a single hemogenic endothelial clone cycling at the peak rate could engender up to 22.6 cells or ~23 cells (Table 1). Furthermore, these observations also suggest that cell division alone from a single hemogenic clone could not account for IAHCs with greater than 23 cells in a 24-hour period.

**Clonal analysis of haematopoietic clusters arising from hemogenic endothelium**

We next explored through low frequency labeling/fate tracing of endothelial cells whether IAHCs are derived from one or more hemogenic endothelial clones3,34. To investigate, we initially crossed a constitutive VE-cadherin-Cre to the R26r-mTmG reporter line (VECAD/mTmG) (Fig. S3A). Recombination in this line results in the excision of mTomato (mTom) and expression of mGFP in endothelial cells (CD31+CD117−) and their haematopoietic (CD31+CD117+) derivatives, including IAHCs within the embryo (Fig. S3A). Upon quantification, we found mGFP+ labeling in ~75% of endothelial cells (CD31−CD117−) and ~50% of IAHC cells (CD31+CD117−; Fig. S3A and B). Immunofluorescence also revealed mosaic expression of mGFP and Tomato in the endothelial cells and IAHC cells in the DA (Fig. S3C and D). Interestingly, 39% of IAHCs contained both mGFP+ and mTom+ cells (Fig. S3E). About 53% of the total IAHCs examined (n = 57) were comprised of mGFP+ cells only, while 9% of IAHCs contained cells that were not Cre recombined, and hence mTom+ only (Fig. S3E). Together, these data indicate that distinct HE clones may contribute to the formation of IAHCs. To test this hypothesis, we performed low dose clonal labeling using Cdh5(PAC)-CreERT2 crossed to the R26r-mTmG reporter line (iCdh5/mTmG) (Fig. S3F-H). A range of low dose tamoxifen (50–150µg) was administered to pregnant females at E8.5 before any IAHC formation to label only endothelial clones, and IAHCs were evaluated at E10.5, the peak of IAHC formation,19 for percent recombination in endothelial cells and IAHC cells, Fig. S3F-H. The analysis revealed that a dose of 100 µg allowed for ~10% labeling of the dorsal aorta (ECs and IAHC cells). We then administered the optimal clonal labeling dose of 100µg (0.1mg) at E8.5 and analyzed the clonal labeling of IAHCs at E10.5 (Fig. 2A). Outcomes of the experiment are identified as IAHCs from a single origin (mGFP+only), polyclonal origin (mGFP+/mTom+) or unlabeled (no Cre recombination (mTom+only)) (Fig. 2B). Polyclonal in this context only designating more than one clone. Following induction, we observed mosaic mGFP+ and mTom+ labeling in the endothelium and cells comprising IAHCs of the DA (Fig. 2C, D). For our analysis of clonality, we excluded mTom+ only IAHCs because we could not conclude whether those IAHCs arose from a single clone since no Cre recombination/fate tracing took place. We evaluated 190 IAHCs (n = 5 embryos) and noted 46 IAHCs at E10.5 were either monoclonal (mGFP only) or polyclonal (mGFP+/mTom+). Quantification of single color (mGFP+ only) and mosaic IAHCs (mGFP+ /mTom+) at E10.5 revealed that the majority of IAHCs (41 out of 46) were mosaic (Fig. 2E), while only 5 out of 46 IAHCs were from a mGFP+ only clone. We further extended this analysis to IAHCs at E11.5. We evaluated 59 IAHCs and found 14 IAHCs to be either monoclonal (mGFP+) or polyclonal (mGFP+/mTom+), (Table 2). At this time point we observed a similar trend as at E10.5 where most IAHCs were mosaic (11 out of 14) and minority derived from a single mGFP+ clone (3 out of 14) (Fig. 2F). Together, the data suggest that IAHCs largely derive from more than one clone (polyclonal origin). Next, we wanted to determine whether any difference in clonality existed between small IAHCs (3–9 cells) vs. large IAHCs (≥10 cells). Our analysis revealed that small IAHCs and large IAHCs significantly differed in their clonality, where small IAHCs were more likely to be derived from one clone (Table 3, Fisher’s Exact = 0.0495). To further support these observations, we also used the confetti reporter system by crossing inducible endothelial Cre animals with the confetti reporter line (Fig. 5A). Outcomes of this cross yield random recombination and labeling of endothelial cells and their haematopoietic derivatives that can be identified with 4 distinct fluorescent labels: mCFP, RFP, YFP and GFP (Fig 5B). In terms of inferring clonality, single color IAHCs denote monoclonal origin and multi-color IAHCs denote polyclonal origin (Fig. 5C). Induction with 2 mg of tamoxifen resulted in low percentage of recombination (< 5%) which may be due the inefficiency of the iCdh5-Cre, or perhaps that the relatively short cell cycle length of these IAHC cells

| Table 1. Estimation of cell number from single clonal divisions at E10.5 and E11.5. |
|-----------------|-----------------|-----------------|-----------------|
| Age            | Tc (h)          | Window | Cell divisions | # of cells from single clone |
|----------------|-----------------|--------|----------------|------------------------------|
| E10.5          | 5.38            | 24     | 4.46           | 22.02                        |
| E11.5          | 8.16            | 24     | 2.94           | 7.68                         |
is precludes detectable accumulation of the fluorescent protein after recombination. Quantification of labeled IAHCs (n = 4) at E10.5 showed that IAHCs contained cells in the S phase at E11.5 vs. E10.5, together with the increase in the average cell cycle length (Fig. 1F–I) prompted us to evaluate whether cells comprising IAHCs may be positioned in different cell cycle phases during early (E10.5) and later (E11.5) periods of the hemogenic window. To test this, we used the FUCCI (Fluorescence Ubiquitin Cell Cycle Indicator) reporter mouse lines.35 In this line, CDT1-mKusabiraOrange2 (mKO2) marks cells in G1 and GEMININ-AzamiGreen (mAG) marks cells in S/G2/M (Fig. 3A). We first verified the expression of these markers relative to DAPI incorporation (Fig. S4A) to determine if the fluorescently labeled cells with either mKO2-CDT1 and/or mAG-GEMININ corresponds to appropriate cell cycle stage by DNA content. We then evaluated expression in the developing neural tube and AGM (Fig. S4B–C), and confirmed the pattern of G1 (FUCCI-Red) vs. S/G2/M (FUCCI-Green) in the neural tube. In this region, the somites are labeled with FUCCI-Red denoting that cells are in G1, while the rest of cells in the neural tube are labeled with FUCCI-Green indicating that they are in the S-phase.35 We also evaluated immunofluorescence for the protein GEMININ in the context of the transgenic reporter FUCCI-Green (GEMININ-AzamiGreen (mAG)) within the developing endothelium and IAHC cells (Fig. S4D), and found that the mAG reporter tends to underreport (by 12%) cells that express the protein GEMININ (via immunofluorescence), Fig. S4E. With that in mind, we evaluated cells within IAHCs for their expression of FUCCI-Green and FUCCI-Red, and noted that cells which make up a single IAHC are positioned in different cell cycle phases. Thus, the

IAHCS are positioned in different phases of the cell cycle as the hemogenic window progresses

Our finding that a higher proportion of IAHC cells were mainly in the S phase at E10.5 vs. E11.5, together with the increase in the average cell cycle length (Fig. 1F–I) prompted us to evaluate whether cells comprising IAHCs may be positioned in different cell cycle phases during early (E10.5) and later (E11.5) periods of the hemogenic window. To test this, we used the FUCCI (Fluorescence Ubiquitin Cell Cycle Indicator) reporter mouse lines.35 In this line, CDT1-mKusabiraOrange2 (mKO2) marks cells in G1 and GEMININ-AzamiGreen (mAG) marks cells in S/G2/M (Fig. 3A). We first verified the expression of these markers relative to DAPI incorporation (Fig. S4A) to determine if the fluorescently labeled cells with either mKO2-CDT1 and/or mAG-GEMININ corresponds to appropriate cell cycle stage by DNA content. We then evaluated expression in the developing neural tube and AGM (Fig. S4B–C), and confirmed the pattern of G1 (FUCCI-Red) vs. S/G2/M (FUCCI-Green) in the neural tube. In this region, the somites are labeled with FUCCI-Red denoting that cells are in G1, while the rest of cells in the neural tube are labeled with FUCCI-Green indicating that they are in the S-phase.35 We also evaluated immunofluorescence for the protein GEMININ in the context of the transgenic reporter FUCCI-Green (GEMININ-AzamiGreen (mAG)) within the developing endothelium and IAHC cells (Fig. S4D), and found that the mAG reporter tends to underreport (by 12%) cells that express the protein GEMININ (via immunofluorescence), Fig. S4E. With that in mind, we evaluated cells within IAHCs for their expression of FUCCI-Green and FUCCI-Red, and noted that cells which make up a single IAHC are positioned in different cell cycle phases. Thus, the

### Table 2. Total IAHCs evaluated for clonality at E10.5 and E11.5.

| Age     | Total IAHCs evaluated | Total scored | Total unlabeled |
|---------|-----------------------|--------------|-----------------|
| E10.5 (n = 5) | 190                   | 46 (24%)     | 144 (76%)       |
| E11.5 (n = 3) | 59                    | 14 (24%)     | 45 (76%)        |

### Table 3. Cluster size and clonality at E10.5.

| Cluster type | Clonal | Polyclonal | Fisher’s Exact |
|--------------|--------|------------|----------------|
| Type A IAHCs with 3–9 cells | 5      | 18         | 0.0497 (°)     |
| Type B ≥ 10 cells       | 0      | 21         |                |

Figure 2. Low dose clonal labeling of IAHCs reveal clonal heterogeneity and polyclonal origin. (A) Experimental schema: Cdhs1(PAC)-CreERT2 animals were crossed with R26r-mTmG lines. Pregnant females were induced with a low dose of tamoxifen (0.1mg) at E8.5 before IAHC formation. Embryos evaluated at at E10.5 and E11.5. (B) Potential outcomes following low dose induction: unlabeled (no Cre recombination), mTom—only, monoclonal (mGFP—only) and polyclonal (mGFP—mTom+). (C) Fluorescence of the dorsal aorta (DA) at E10.5 (broken lines) after low dose induction shows mosaic mGFP+ (green) expression in a background of mTom+ (red) cells. PU.1 (gray) labels an IAHC. Scale bar, 20 μm. (D) Higher magnification of the IAHC depicted in (C) which is noted to contain both mGFP+ (green) and mTom+ (red) cells. PU.1 (gray) labels cells in the IAHC and some single haematopoietic cells. Scale bar, 10 μm. (E) Analysis of IAHC clonality at E10.5. Monoclonal (mGFP—only) IAHCs denoted by light blue bars, and polyclonal (mGFP—mTom+) IAHCs denoted by dark blue bars, unlabeled IAHCs denoted by clear bars. A total of 190 IAHCs were analyzed. n = 5 litters. (F) Analysis of IAHC clonality at E11.5. Monoclonal (mGFP—only) IAHCs denoted by light gray bars, and polyclonal (mGFP—mTom+) IAHCs denoted by dark gray bars, unlabeled IAHCs denoted by clear bars. A total of 59 IAHCs were analyzed, n = 3 litters.
cells comprising one IAHC are not necessarily synchronized in their cell cycles (Fig. 3B). To analyze the IAHC cell population as a whole, we used FACS to discern the cell cycle phase of IAHC cells at E10.5 and E11.5. FACS analysis revealed that at E10.5 ~2% of the SSEA-1⁻CD31⁺CD117⁻mKO²⁺ IAHC cells population (IAHCs) resides in G1, while ~28% resides in the S/G2/M (Fig. 3C and D). Interestingly, at ~11% ~35% of the IAHC cell population now resides in G1, and ~11% resides in S/G2/M (Fig. 3C and D). Overall, the data suggest that there is a shift in the cell cycle position of cells within the IAHCs as the hemogenic window progresses during development. This previously unappreciated difference between the cell cycle position of IAHC cells at E10.5 and E11.5 have led us to postulate whether the cell cycle position may influence repopulation potential.

**Cell cycle phase does not fully account for repopulation activity**

Previous studies from Nygren and colleagues which examined the cell cycle kinetics of E14.5 fetal-liver HSCs (FL-HSCs) found that a prolonged cell cycle length is a characteristic of HSCs. In the cell cycle kinetics of E14.5 fetal-liver HSCs (FL-HSCs) found Previous studies from Nygren and colleagues which examined activity Cell cycle phase does not fully account for repopulation potential.

whether the cell cycle position may in of IAHC cells at E10.5 and E11.5 have led us to postulate that these cells were indeed able to reconstitute the different hematopoietic cell lineages (Fig. 4D). Overall, the data suggest that enrichment of E10.5 IAHC cells did not appear to maintain the >1% level of engraftment long-term (Fig. 4B, G1 IAHCs 14.5% vs S/G2/M IAHCs 0.3%). Further evaluation of lineage markers for B and T cells as well as myeloid lineage cells within these animals revealed that these cells were indeed able to reconstitute the different hematopoietic cell lineages (Fig. 4D). Overall, the data suggest that enrichment of E10.5 IAHC cells in G1 does not overcome their inability to engraft adult hosts, as neither does enrichment for cells in S/G2/M. However, our data also support that E11.5 IAHCs contain adult engraftable HSCs (especially considering the low numbers of cells transplanted), and that the E11.5 HSC population may be enriched when selected for the G1 (vs. S/G2/M) cell cycle phase, similar to their fetal and adult HSC counterparts.36,39-41

**Cell cycle-specific transcriptional profiling of E10.5 and E11.5 IAHC cells reveal transcriptional programs related to cell movement and cell migration**

To further define transcriptional differences that underlie the developmental age and maturation of cells comprising the

![Figure 3](image-url)
Figure 4. Direct transplantation of E10.5 and E11.5 FUCCI labeled IAHCs. (A) Transplantation experimental scheme. IAHC cell populations (DAPI−SSEA1−CD31−CD117+) from FUCCI-Red (mKO2-CDT1) and FUCCI-Green (mAG-GEMININ) mice were sorted (cell number = 200–2000 cells per group) and directly transplanted into irradiated adult recipients (E10.5 n = 8–9 per group, E11.5 n = 12 per group). (B) Analysis of engraftment (defined as >1% peripheral blood chimerism) at 8 and 16 weeks depicts E10.5 FUCCI-Red (open circles) and green (open squares) IAHCs with no detectable engraftment (0/8 and 0/9 respectively). E11.5 FUCCI-Red (triangles) and FUCCI-green (hexagons) each exhibit one animal with detectable levels of engraftment (1/12 and 1/12 respectively) at 8 weeks (each labeled with a separate color). Only the recipient of G1 IAHCs retains stable engraftment at 16 weeks (fuchsia triangle). No stable engraftment was seen in E11.5 Fucci-Green at 16 weeks, although there was low level of chimerism detected (0.3%) (fuschia hexagon). Double asterisks indicates the occurrence of mortality in the cohort. (C) Representative FACS plots of E10.5 FUCCI-Red and FUCCI-Green IAHCs show no detectable engraftment by 16 weeks. (D) FACS plots of E11.5 G1 IAHCs (fuschia triangle) demonstrate 15% donor reconstitution and peripheral blood contribution to the T (CD3-FITC), B (B220-PE) and myeloid (CD11b-PeCy7) lineages by 16 weeks and for E11.5 S/G2/M (blue hexagon) 0.3% donor engraftment and lineage contribution to B, T, and myeloid lineages.
IAHCs, we performed RNA sequencing on 4 samples: E10.5 and E11.5 SSEA-1<sup>−</sup>CD31<sup>−</sup>CD117<sup>−</sup>mKO2<sup>+</sup> (G1 IAHCs) and E10.5 and E11.5 SSEA-1<sup>−</sup>CD31<sup>−</sup>CD117<sup>−</sup>mAG<sup>+</sup> (S/G2/M IAHCs) (Fig. 5A). We have identified a total of 774 genes for G1 and 703 genes for S/G2/M that are differentially expressed between E10.5 and E11.5. G1 contained 453 unique genes and S/G2/M contained 381 unique genes (Fig. 5B). To determine the molecular signature of IAHCs that change according to developmental age, only genes with a 2-fold change in expression were analyzed using the Ingenuity Pathway Analysis (IPA,

**Figure 5.** Transcriptional profile of E10.5 and E11.5 FUCCI labeled IAHCs. (A) RNA sequencing experimental scheme. IAHC cell populations (DAPI<sup>−</sup>SSEA1<sup>−</sup>CD31<sup>−</sup>CD117<sup>−</sup>) from FUCCI-Red (mKO2-CDT1) and FUCCI-Green (mAG-GEMININ) mice were sorted and subjected to RNA sequencing. (cell number, n = 300–8000 cells pooled from 2–3 litters per group per age). (B) Venn diagram depicts the number of common and unique genes between G1 (FUCCI-Red) and S/G2/M (FUCCI-Green) IAHCs. We identified 453 unique genes upregulated in G1 E11.5 IAHCs vs. G1 E10.5 IAHCs. Similarly, we identified 381 unique genes upregulated in S/G2/M E11.5 IAHCs vs. S/G2/M E10.5 IAHCs. These 2 groups share 321 common genes. (C) The RNASeq analysis of top biologic functions (z-score > 5, p < 0.05) enriched in E11.5 IAHCs at G1 relative to E10.5 IAHCs at G1 identifies the top 11 increased biologic functions involved a variety of cellular processes including: phagocytosis, organization, migration, immune response, homing, endocytosis, differentiation, chemotaxis, cell viability, cell survival and cell movement. z-score ≥ 5 (black dotted line), p-value (red dots) via Fisher’s exact test. (D) The RNASeq analysis of top biologic functions (z-score > 5, p < 0.05) enriched in E11.5 IAHCs at S/G2/M relative to E10.5 at S/G2/M reveals the top 6 increased biologic functions that are involved in: cell survival, migration, homing, development, cell viability and cell movement. Not surprisingly, these categories are similar to those in E11.5 IAHCs at G1. z-score ≥ 5 (black dotted line), p-value (red dots) via Fisher’s exact test. (E) The analysis of canonical pathways (z-score > 5, p < 0.05) that are active in E11.5 IAHCs at G1 compared with E10.5 IAHCs at G1 identifies the top 10 activated signaling pathways which include: Gamma receptor-mediated phagocytosis, production of NO, leukocyte extravasation, IL-8 signaling, Rac signaling, integrin signaling, dendritic cell maturation, NFAT signaling, NF-Kappa B signaling and Rho GTPase signaling. (F) The analysis of canonical pathways (z-score > 5, p < 0.05) that are active in E11.5 IAHCs at S/G2/M compared with E10.5 IAHCs at S/G2/M identifies the top 10 activated signaling pathways which include: Gamma receptor-mediated phagocytosis, ERK/MAPK signaling, production of NO, PKC delta signaling, leukocyte extravasation, NFAT signaling, p38 MAPK signaling, Tec Kinase signaling, IL-8 signaling and integrin signaling.
Ingenuity Systems, www.ingenuity.com). We used 2 metrics, the p-value and the activation z-score, to identify the most important downstream effects related to age. A positive z-score indicates an increased functional or pathway activity in E11.5 relative to E10.5 for both G1 and S/G2/M samples. The p-value, calculated with Fisher’s exact test, indicates the likelihood that the association between a set of genes in our data set and a biologic function or pathway is significant.

Our analysis revealed that genes overexpressed in E11.5 vs. E10.5 at G1, regulate 11 top functions related to variety of cellular processes (Fig. 5C). Interestingly, the functions that are significantly increased are those that are involved in cell migration, endocytosis, cell movement, phagocytosis, cell organization, immune response, homing differentiation and cell survival. Perhaps not surprising is that the analysis of genes overexpressed in E11.5 vs. E10.5 at S/G2/M, yielded 6 top functions similar to those in G1 with cell migration, cell movement, homing, cell survival, and development being significantly activated (Fig. 5D). Furthermore, examination of top canonical pathways in G1 reveal activation of signaling that regulate phagocytosis, nitric oxide signaling, leukocyte extravasation and IL-8 (Fig. 5E). Our analysis of canonical pathways in S/G2/M show activation of phagocytosis, ERK signaling, nitric oxide signaling and PKC Delta signaling (Fig. 5F). Overall, our data suggest that regardless of the cell-cycle phase, a cell movement and cell migration program is established in IAHC cells by E11.5.

**Molecular differences between E11.5 G1 and S/G2/M IAHC cells point to complement activation for a role in engraftment**

Since our transplantation data suggest that both E11.5 G1 and S/G2/M IAHCs contain cells with different engraftment potential, we wanted to further determine molecular differences related to the cell cycle that may influence engraftment. We identified 106 differentially expressed genes between G1 and S/G2/M IAHCs at E11.5. First, we compared transcripts from E11.5 S/G2/M with E11.5 G1 to identify functions that are upregulated within the G1 phase of the cell cycle. Surprisingly, genes overexpressed in G1 regulate 14 main functions that are involved in different cellular processes. The biologic functions that are most significantly activated include chemotaxis, migration, uptake and cell volume (Fig. 6A). As expected, comparison of E11.5 G1 with E11.5 S/G2/M transcripts to identify functions

![Image of Figure 6](image_url)
upregulated within the S/G2/M phases of the cell cycle yielded functions relating to proliferation, infection and growth (Fig. 6B) with proliferation being the most significantly activated.

Our analysis of upregulated pathways in E11.5 G1 vs. S/G2/M also revealed activation of canonical signaling pathways that regulate the complement system, roles of pattern recognition receptors and LXR/RXR activation (Fig. 6C). Indeed, several complement genes, such as receptors C5AR, C3AR and complement components C1QA, C1QB, and C1QC are actively transcribed in G1 (Fig. 6D). In contrast, examination of transcripts upregulated in S/G2/M reveal signaling pathways regulating the G2/M DNA damage checkpoint, mitotic roles polo-like kinases and melatonin degradation (Fig. 6E). Subsequently, we observe ESPL1, PLK1, CDK1 and TOP2A transcripts associated with the S/G2/M phases of the cell cycle (Fig. 6F). We confirmed the complement component expression via QPCR (Fig. 6G). Overall, when comparing between age groups, we find expression of complement genes in E11.5 G1 IAHC cells suggesting this may be a critical pathway for the maturation of IAHC cells toward definitive HSCs resulting in adult engraftment, chemotactic and migration programs.

Discussion

We set out to determine how IAHCs are formed following their emergence from the endothelium. Our data reveal that the cell cycle length of E10.5 IAHC cells is approximately 5 hours. In addition, our clonal labeling analysis suggests that more than one hemogenic clone, (likely 2) in the dorsal aortic floor contributes to the formation of a single IAHC. Several IAHCs may then be produced from multiple clones, as clonal labeling in the zebrafish suggests the existence of up to 30 HSC clones per aorta. Clonal output is also likely heterogenous, as recent work using limited dilutional analyses suggests, with increased heterogeneous HSPC populations at E10 vs. E11. Thus, IAHC formation is likely driven by the rapid cell proliferation of several hemogenic endothelial clones with differing functional capacities. These findings further support a recent observation that an initial pool of pre-HSCs is established, from which HSCs mature from by E11.5. Correspondingly, the length of the cell cycle in E11.5 IAHC cells increases to about 8 hours. This observation is intriguing as fetal liver (FL) HSCs have been observed to have a mean generation time of 10.6 hours.

The cell cycle of FL and bone marrow (BM) HSCs is tightly associated with their ability to self-renew and differentiate. The progression of HSCs through the cell cycle both in vitro and in vivo is accompanied by notable changes in their engraftment potential. Several lines of evidence suggest that cell cycle position may influence repopulation activity. Specifically, FL and BM HSCs in the G0/G1 appear to engraft adult recipients better than their S/G2/M counterparts. Moreover, a permissive environment is also required for successful engraftment. Arora and colleagues demonstrated that embryonic (AGM) HSCs engraft neonatal recipients better than adult recipients. They also found that adult-like (BM and FL) HSCs more efficiently reconstitute adult recipients than neonates. Here, we investigated whether embryonic HSCs from ontogeny are at a specific cell cycle phase which may affect their engraftment in the adult BM. We note that adjusting for cell cycle phase at E10.5 (by selection and transplantation of the minority G1 population) does not overcome the inability of these cells to engraft adult hosts. However, the lengthening of cell cycle from E10.5 to E11.5 may be commiserate with the E11.5 IAHC cell population ability. Our data show that E11.5 G1 IAHC populations may engraft better than their S/G2/M counterpart. This finding is in agreement with a recent report from Batsivari and colleagues who found that direct transplantation of mature E11.5 DHSs at G0/G1 engrafted better than those in S/G2/M. Furthermore, the seemingly coincident ability of E11.5 IAHC cells to engraft adult irradiated recipients point to the possibility that the lengthening of the cell cycle may influence HSC maturation. The number of IAHC cells detected during the hemogenic window is small with only ~1500 ~ 2000 cells in an E10.5 AGM. In our hands, the average percentage of cells that exhibit either FUCCI-Green or FUCCI-Red fluorescence ranges from 600 to 700 at E10.5, and 800 to 1000 cells at E11.5 per AGM. Thus, a significant limitation of our transplantation study is that only a limited number of cells could be harvested from any single litter for transplantation. We were able to transplant as low as ~200 cells and as much ~2000 cells per adult mouse (~1/8 to 1 IAHC embryo equivalent EE) under these conditions, and thus may be at the detection boundary of the assay.

RNA sequencing of IAHC cells at E10.5 and E11.5 at different phases of the cell cycle revealed a dynamic transcriptional landscape that changes with developmental age and cell cycle phase. Across developmental age by E11.5, an enrichment for genes that regulate “cell movement,” “migration” and “inflammatory response” become evident and further support the findings Mckinney-Freeman and colleagues that IAHC cells may resemble a “macrophage-like signature”, in addition to the recent wealth of information pointing to inflammatory signals being critical for early HSC development. Furthermore, since it has been hypothesized that IAHC cells migrate to the fetal liver after their formation in the AGM, it is likely that the maturation process of E10.5 IAHC cells may involve the activation of cell movement and cell migration signaling pathways via inflammatory signaling that prepares IAHCs for entry into circulation.

Transplantation of E11.5 G1 vs. S/G2/M IAHC cell populations suggested an advantage of G1 IAHC cells to engraft irradiated adult recipients, albeit with low numbers. This observation is in line with earlier findings that HSCs in G1 have an engraftment advantage over their S/G2/M counterparts. Our findings also differed from a recent report that found that pre-HSCs in S/G2/M were capable of enhanced engraftment than pre-HSCs in G0/G1. This discrepancy with recent work may be due to co-culturing which may alter cell cycle states. Transcriptional profiling of E11.5 G1 IAHC cells revealed an enrichment for transcripts that regulated the “immune response,” “chemotaxis” and “cell migration.” While not surprising, profiling of S/G2/M E11.5 IAHC cells demonstrated an activation of signaling pathways that regulated “cell growth” and “proliferation.” Thus, our data suggests that during the G1 phase of the cell cycle, IAHC cells may become sensitive to a multitude of cellular cues which include immune and inflammatory signals that may regulate cell movement, egress...
into circulation, and consequently, migration toward the fetal liver. Furthermore, there is evidence to suggest complement activation plays a role in donor HSPC migration and homing post transplantation. It has been shown that C1q enhances the responsiveness of HSPCs to the chemoattractant CXCL12 (SDF-1). We find that genes of the classical complement cascade which include C1QA, C1QB and C1QC, as well as anaphylatoxin receptors C3AR and C5AR, to be upregulated in E11.5 G1 IAHC cells. Thus, the increased expression of complement system components in E11.5 G1 IAHC cells may provide a method for homing to the adult bone marrow, and possibly explain the engrafment potential of G1 IAHC cells relative to their S/G2/M counterparts. Overall, our data support a model following EHT where at E10.5, IAHCs form rapidly from the proliferation of at least 2 hemogenic clones, and subsequently cells comprising IAHCs undergo prolongation of their cell cycle length and activate transcriptional programs of migration and homing, which may lead to facilitation of transit into the fetal liver (Fig. 7). These findings further lend credence to the establishment of a robust HSC program following EHT, such that the very properties (prolonged cell cycle length, engrafment in G0/G1, lengthening of G1) that define fetal liver and adult HSCs is reflected in the newly emerged embryonic AGM HSCs.

Materials and methods

Mice

Mice were housed under a barrier facility following IACUC procedures and animal protocols under the University of California San Francisco Laboratory Animal Research Committee Guidelines. C57Bl/6 (Wildtype), Gt(Rosa)26Sortm4(Actb-ttdTomato,-EGFP)Lox1/Lox1 (R26r-mTmG),58 Tg(Cdh5-cre)1Sple/J (VEC-Cre),5 and Gt(Rosa)26Sortm1(CAG Brainbow2.1)Cle/J (R26r-confetti)59 mice were obtained from Jackson Laboratories. Cdh5(PAC)-CreERT2 (Tg(Cdh5-cre/ERT2)1Rha)60 mouse line was obtained from Ralph Adams, PhD, CancerUK, and Fucci Green Tg(CAG mAG/GMNN)504Amiy and Fucci Red Tg(CAG-mKO2/CDT1)596Amiy lines from RIKEN.35 Males from this line were crossed to female homozygous Rosa26-mT/mG females. Pregnancies were timed in accordance with the vaginal plug (gestation day 0.5 on the day of the plug). Genomic DNA from adult ear punch, or conceptus yolk sacs was genotyped using MyTaq Extract PCR Kit (Bioline, BIO21127).

Low density clonal labeling

Low density clonal labeling was determined by titrating doses of tamoxifen beginning with 50 μg, 100 μg to 150μg. Tamoxifen was administered at E8.5 before any IAHC formation. After determining the tamoxifen dose, females were injected intraperitoneal with 0.1 mg at E8.5 before cluster formation. Embryos were evaluated at E10.5 (35–39sp) and E11.5 (44–48sp).

Tamoxifen preparation

Tamoxifen was purchased from MP Biologicals LLP. A 10 mg/ml stock was prepared by dissolving 10 mg of tamoxifen in 100 μl of 100% ethanol and mixed with 900 μl sterile sunflower oil (1:9). For low dose clonal labeling experiments, a stock solution of 2.5 mg/ml of tamoxifen was prepared in similar fashion.

Cell cycle length determination

Cell cycle length was determined by sequential administration of EdU and BrdU. First, 0.15 ml of EdU (10 mg/ml) was administered intraperitoneally. After 1.5 hours, 0.20 ml of BrdU (10 mg/ml) was injected. BrdU was detected by treating tissue sections with 2N HCl for 30 minutes at 37°C and incubated with 1:100 anti-BrdU (MOBU-1, Life Technologies #B35128). Samples were washed 3x with 1xPBS + 0.05% Tween (PBST) and a secondary antibody was applied at 1:200 (Invitrogen, #A21203). EdU was detected using a Click-iT EdU Detection System (Life Technologies, #C10337). This technique allows for identification of cells that are in S-phase (S) by EdU incorporation. Incorporation of both EdU+ and BrdU+ designates cells in the S phase (S). Incorporation of EdU+ and not subsequently BrdU+, delineates cells that are leaving the S-phase (L) during the 1.5 hrs between EdU and BrdU pulses. All proliferating IAHC cells are marked by PU.1, (P). The total length of S phase (Ts) is estimated by first calculating the number of cells leaving the S-phase (L) over the total number of cells in the S-phase (S). The initial time (Tl) between EdU and BrdU (1.5 hrs) is then divided by this fraction, thus Ts = Tl/(L/S). The total length of the cell cycle (Tc) is estimated by first determining the number of cells in the S-phase (S) over the total number of proliferating cells (P). The total S-phase length (Ts) is then divided by this fraction, thus Tc = Ts/(S/P).

Flow cytometry

Whole embryos or AGMs were dissociated as described previously (Lizama et al. 2014) and stained for 30 min at 4°C with agitation. Single cell suspensions were sorted in a BD FACS Aria
III. Flow cytometric analyses were performed on a FACS Verse or FACS Aria III with FACSDiva 8.0 software (BD Biosciences) and data analyzed using FlowJo v10.0.7 (Tree Star). Antibodies were purchased from BD Biosciences: CD117-APC (#553356), CD31-PerCp Cy5.5 (#562861), V450-SSEA-1 (#561561), APC-CD45.2 (#558702), FITC-CD45.1 (#553770), PeCy7-CD11b (#552850), FITC-CD3 (#55274), PE-Ter119 (#553673). Isotype controls used were: IgG2b-APC (#55391), IgM-V450 (#560861), IgG1-PerCpCy5.5 (#562861), IgG2b-PeCy7 (#552849), IgG2b-FITC (#553988), IgG2b-PE (#553899). Antibodies were purchased from Biolegend: PE-CD45R/B220 (#400507) and IgG2a-PE (#400507).

**Immunofluorescence and confocal microscopy**

E10.5 – E11.5 embryos were fixed in 4% paraformaldehyde (PFA) solution at room temperature for 20 minutes or in 2% PFA overnight. Samples were frozen in Tissue-Tek OCT Compound (Sakura Finetek, 4583). 20–30 μm cryosections were obtained (Thermo Scientific Micron, HM550). Slides were incubated in blocking buffer (PBS, 0.5% Triton-X100) with 5% pound (Sakura Finetek, 4583). 20 PFA overnight. Samples were frozen in Tissue-Tek OCT Compound (PFA) solution at room temperature for 20 minutes or in 2% E10.5 – Immunofluorescence experiments 1:100 of Rabbit Runx1 (ab35962), Goat Sox17 (R&D Systems #AF1924) Rat CD117 Clone (28B) BD PharMingen (#553352) Rat CD31 Clone (Mec13.3) BD PharMingen (#553370) Rabbit PU.1 Clone (9G7) Cell Signaling (#2258) were used. Secondary antibodies according to host species were used at 1:200 dilutions (Invitrogen: donkey anti-goat 488 #A11055, donkey anti-goat #A21447, donkey anti-rat 594 #A21209, donkey anti-rabbit 647 #A31573, Abcam: donkey anti-rabbit 405 #175649).

**RNA isolation and sequencing**

Samples were pooled from 2–3 litters. Total RNA was isolated and purified from sorted cells using the QIAGEN RNeasy Plus Micro Kit (#74034) following manufacturer’s instructions. RNA integrity was determined using a Bioanalyzer. Samples were ribo-depleted and amplified with Ovation RNA-Seq System V2. Paired-end 2 × 100 bp sequencing was performed using the Illumina HiSeq2000 machine. RNA sequencing analysis was done using Galaxy (https://usegalaxy.org/u/laiumiu/nix/w/rnaseqjoandata) with a False discovery rate (FDR) of 0.085. Transcripts with ≥ 2-fold change expression were further analyzed using the Ingenuity Pathway Analysis to identify key biologic functions and pathways that are expected to be more active in E11.5 than E10.5. To be more stringent, we focused our analysis to functions and pathways with a positive z-score. Data available under GEO accession number: GSE99314.

**Transplantation**

Eight–12 week old CD45.2 C57Bl/6 FUCCI mice were used as donors for cell isolation. E10-E10.5 (30–39sp) and E11-E11.5 (40–48sp) AGMs were sorted for SSEA-1+/CD31+/CD117+ FUCCI-Red or FUCCI-Green as donor cells. 8–12 week old CD45.1 C57Bl/6-Boy/J wild type mice were used as recipients for cell transplantation. Congenic recipient mice (CD45.1) were irradiated with 900 rads, split dose, 3 hours apart using a cesium source. Purified donor cells were injected intravenously with 200,000 spleen helper cells, and haematopoietic reconstitution was monitored over time in the peripheral blood based on CD45.2 expression. Recipients with > 1% donor chimerism were considered reconstituted. Transplanted mice were kept on antibiotic-containing food for 2 weeks. Recipients of helper cells and congenic donor cells were used as negative control, and recipients of 4 × 10⁶ CD45.2 whole bone marrow cells were used as positive control. All mice were maintained at UCSF in accordance with IACUC approved protocols.

**QPCR**

Total RNA was isolated from pooled littermates of wild-type AGM SSEA-1+/CD31+/CD117+ populations (~2000 cells) from E10.5 and E11.5 as described previously. A total of 40 μg of RNA was transformed into CDNA using SuperScript VILO Master Mix (Thermo Fisher Scientific #11755–050) following the manufacturer’s instructions.

**Statistics**

All the data are expressed as mean ± SD. P values were generated either with a Fisher’s exact test or an unpaired t-test with Welch’s correction and considered significant when ≤ 0.05. “n” indicates the numbers of independent experiments performed. Clonality data from the low dose fate tracing experiments were binned according to IAHC size Type A (3–9 cells) and Type B (>10 cells), and fisher’s exact test was performed to evaluate the correlation between IAHC size and clonality.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.
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Author contributions

JPZ and ACZ designed and planned experiments. JPZ performed the experiments, with assistance from COL and KMCR. JPZ and ACZ wrote the manuscript.

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