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1 Title:

2 HSC-independent definitive hematopoietic cells persist into adult life.

3

Michihiro Kobayashi¹, Haichao Wei^{1,2}, Takashi Yamanashi⁴, David J Shih³, Nathalia Azevedo
Portilho¹, Samuel Cornelius¹, Noemi Valiente¹, Chika Nishida¹, Wenjin J Zheng³, Joonsoo Kang⁶,
Jun Seita^{4, 5}, Jia Qian Wu², Momoko Yoshimoto^{1*}

- 7
- 8 ¹Center for Stem Cell and Regenerative Medicine, Brown Institute of Molecular Medicine, and
- ⁹ ²The Vivian L. Smith Department of Neurosurgery, and ³Department of Dept. Biochemistry &
- 10 Molecular Biology, McGovern Medical School, University of Texas Health Science Center at
- 11 Houston, Texas, USA
- ⁴Advanced Data Science Project, RIKEN Information R&D and Strategy Headquarters, Tokyo,
- 13 Japan
- 14 ⁵Center for Integrative Medical Sciences, RIKEN, Kanagawa, Japan
- ⁶Department of Pathology, University of Massachusetts Medical School, Worcester, MA, USA

- 17 *Corresponding author
- 18 Momoko Yoshimoto MD., PhD
- 19 Momoko.Yoshimoto@uth.tmc.edu
- 20
- 21
- 22

23 Summary

24 The stem cell theory that all blood cells are derived from hematopoietic stem cell (HSC) is a 25 central dogma in hematology. However, various types of blood cells are already produced from 26 hemogenic endothelial cells (HECs) before the first HSCs appear at embryonic day (E)11 in the 27 mouse embryo. This early blood cell production from HECs, called HSC-independent 28 hematopoiesis, includes primitive and definitive erythromyeloid progenitors that transiently 29 support fetal blood homeostasis until HSC-derived hematopoiesis is established. Lymphoid 30 potential has traditionally been detected in the extra-embryonic volk sac (YS) and/or embryos 31 before HSC emergence, but the actual presence of lymphoid progenitors at this stage remains 32 unknown. In addition, whether HSCs in the fetal liver are the main source of innate-like B-1a cells 33 has been controversial. Here, using complementary lineage tracing mouse models, we show that 34 HSC-independent multipotent progenitors (MPPs) and HSC-independent adoptive B-lymphoid 35 progenitors persist into adult life. Furthermore, HSCs minimally contribute to the peritoneal B-1a 36 cell pool; most B-1a cells are originated directly from ECs in the YS and embryo and HSC-37 independent for life. Our discovery of extensive HSC-independent MPP and B-lymphoid 38 progenitors in adults attests to the complex blood developmental dynamics through embryo to adult that underpin the immune system and challenges the paradigm of HSC theory in hematology. 39 40

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47 Introduction

48 All blood cells are derived from special endothelial cells (ECs), referred to as hemogenic 49 endothelial cells (HECs) in the extraembryonic volk sac (YS) and para-aortic region of the mouse 50 embryo during a limited time window^{1, 2, 3, 4, 5}. Before the emergence of hematopoietic stem cells 51 (HSCs) from HECs in the aortic regions at E11, multiple waves of blood cell production occur 52 directly from HECs, which contribute to the transient fetal hematopoiesis⁴. During this time, in vitro B-lymphoid potential from HECs in the early YS and embryo has been reported^{6, 7, 8, 9}. 53 54 however, the physiological presence of HSC-independent B-cells has yet to be unequivocally 55 determined. Furthermore, if they exist, how long and to what extent such HSC-independent B-56 cells persist into postnatal life remains unknown. B-lymphocytes are mainly categorized into three 57 subsets; bone marrow (BM) HSC-derived B-2 cells (e.g., splenic follicular (FO) B-cells), marginal 58 zone (MZ) B-cells, and innate-like B-1 cells that reside mainly in the body cavities. CD5⁺ B-1a 59 cells are not replenished by BM HSCs and have generally been considered to be derived from FL HSCs^{10, 11, 12} whereas contrasting results have also been reported^{13, 14}. In addition, it has been 60 61 reported that progenitors at E10.5 AGM region that have biased B-1 cell potential can acquire B-2 potential upon AGM-derived endothelial niche culture¹⁴. Therefore, it is plausible that B-2 cells 62 63 may arise from HECs independently of HSCs.

64 In this study, using HSC- and EC-lineage tracing mouse models, we found that HSCs in 65 the FL slowly produced MPPs and B-lymphoid progenitors after birth and EC-derived HSCindependent MPPs and B-progenitors persisted in adult for more than 6 months. Furthermore, FL 66 67 HSCs minimally contributed to the peritoneal B-1a cells and EC-derived HSC-independent B-1a 68 cells were the major population and maintained for life. Transplantation assays of E11.5 HSC-69 precursors without co-culture demonstrated the presence of transplantable HSC-independent 70 MPPs and B-1a progenitors among this population. Our study resolved the long-lasting 71 controversy of B-1a cells and provide unexpected evidence of HSC-independent MPPs and 72 adaptive B-progenitors in adult life.

73 Results

74 Fetal HSCs do not contribute to the peritoneal B-1a cell pool in a steady state.

Fqd5 is expressed exclusively in LT-HSCs [lin⁻Sca-1⁺c-kit⁺(LSK)CD150⁺CD48⁻ cells] in the FL 75 76 and BM. Fgd5CreERT2:Rosa-TdTomato (iFgd5) mice enable us to label HSCs at a time of Tamoxifen (TAM) injection^{15, 16}. We labeled HSCs in E14.5 FL or postnatal day 2 (P2) BM by TAM 77 78 injection, respectively, and examined Tomato% in various B-cell subsets and BM progenitors over 79 300 days after birth (Fig. 1A, B, Extended Date Fig. 1A, B). There were variations of Tomato 80 labeling efficiencies among animals and timed matings did not always precisely synchronize the 81 actual embryonic age at the time of TAM administration. Therefore, to evaluate the labeling 82 efficiency in a consistent manner, we calculated a Tomato % ratio of each blood cell type to HSCs (Tomato % ratio =Tomato% of a defined cell type /Tomato% of HSC) as previously described¹⁶. 83 84 If a cell population is HSC-derived, Tomato % ratio should become close to 1.0 over time.¹⁶ Surprisingly, Tomato % ratio of B-1a cells stayed very low (<0.2) up to 300 days after birth (Fig. 85 86 1B) even when HSCs were labeled at E14.5 FL stage, indicating that the majority of adult 87 peritoneal B-1a cells were HSC-independent. Furthermore, the Tomato % ratios of MPPs and FO 88 B-cells showed 0.7-0.8 and 0.5, respectively (Fig. 1B, Extended Date Fig. 1A, B). This result raised a question as to whether some of these cells in adults arise independently of HSCs. 89

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91 MPPs and B-progenitors in the FL are HSC-independent and originated at as early as E7.5.

92 It has been reported that FL MPPs, but not LT-HSCs, produced B-1a cells most efficiently upon 93 transplantation^{13, 14}. Since fetal HSCs were not the major drivers of the peritoneal B-1a cells in 94 steady states (Fig. 1B), FL MPPs that have B-1a potential must be derived from precursors at 95 earlier stages than FL HSCs, such as HECs that can produce various hematopoietic cells^{17, 18}. 96 Therefore, we sought the origin of FL MPPs by using an EC-lineage tracing mouse model. *Cdh5* 97 is a specific marker of ECs and *Cdh5CreERT2: Rosa-TdTomato* mice (*iCdh5*) mice are widely 98 used to label ECs at a time of TAM injection¹⁹. First, we tried to label HECs that produce the first 99 de novo HSCs at E11.5. EC-labeling at E11.5 exclusively marked HSCs when we analyzed E15.5 100 FL (Fig. 1C, D). While 18.8±12.7% of LT-HSCs were Tomato⁺, only 2.3±2.8% of MPPs were 101 Tomato⁺, and its Tomato% ratio was only 0.1 (n=3) (Fig. 1D). These results indicate that even the 102 first HSCs produced at E11.5 do not yet differentiate into MPPs in the E15.5 FL, but gradually 103 produce MPPs and FO B-cells after birth (Fig. 1E). Additionally, B-1a cells were not efficiently 104 labeled by E11.5 TAM injection even when analyzed at >300 days (Fig. 1E, Extended Date Fig. 105 1C), in line with the HSC-lineage tracing results that HSCs do not produce B-1a cell efficiently.

106 Next, we sought the origin of FL HSC-independent MPPs at the earlier embryonic stages 107 before HSC emergence. We labeled ECs at E7.5 and examined Tomato⁺ MPPs and other 108 hematopoietic progenitors in E15.5 FL (Fig. 1F). Surprisingly, whereas LT-HSCs were barely 109 labeled, around 10-30% of MPPs were Tomato⁺ (Fig. 1G, Extended Date Fig. 1D). Furthermore, 110 other hematopoietic progenitors including common lymphoid progenitors (CLPs) and CD19⁺ B-111 progenitors showed higher Tomato% than that of LT-HSCs (Fig. 1G, Extended Data Fig. 1D). 112 Because we labeled ECs, not HSCs, in *iCdh5* mice, the Tomato ratio=1.0 indicates that the target 113 cells and HSCs are derived from ECs at the same stage, and >>1.0 or <<1.0 indicates that these two populations are derived from ECs at different time points²⁰. The Tomato % ratios of MPP, 114 115 CLP, and B progenitors to HSCs were much >> 1.0 (Fig. 1G, H), indicating that E15.5 FL MPPs 116 and B-lymphoid progenitors were HSC-independent. Importantly, these HSC-independent MPPs 117 and B-progenitors marked at E7.5 persisted into adult life, more than 300 days after birth (Fig. 1H,

118 l).

When E9.5 ECs were labeled, the Tomato % of MPPs, other progenitors, and HSCs in the E15.5 FL were similar (the ratio was near 1.0) (Extended Data Fig.1E). Considering that HSCs expand rather than differentiate during E10.5 to E15.5^{21, 22}, this result suggests that most progenitors and HSCs were simultaneously produced from HECs at E9.5. Importantly, these E7.5 and 9.5 HSC-independent MPPs persisted more than 300 days after birth because their tomato

ratio kept >1.0, suggesting that HSC-independent MPP-derived hematopoiesis occurs even in the
adult BM (Fig. 1H, I, Extended Data Fig. 1F).

126

127 FL MPPs contain HSC-independent common B-1 and B-2 progenitors

As Tomato⁺E15.5 FL MPPs are HSC-independent (Fig. 1G) and contain B-1a precursors upon transplantation¹⁴, we examined their B-1 and B-2 progenitor potential using modified B-cell colony assays¹⁴. From 500 Tom⁺ MPPs marked at E7.5 in *iCdh5* mice, 29 B-progenitor colonies were detected (Extended Data Fig. 1G). Among them, we found 20 B-1 progenitor colonies and 9 B-1 and B-2 progenitor mixed colonies. These data showed that E15.5 FL HSC-independent MPPs contain B-1 progenitors and common lymphoid progenitors that can differentiate into both B-1 and B-2 cells.

135

136 Most B-1a cells are derived from E7.5-10.5 HECs.

137 Since HSCs after E11.5 showed minimal contribution to the peritoneal B-1a cells (Fig. 1E), we examined at which stage of HECs mark the peritoneal B-1a cells most efficiently. We expected 138 139 that B-1a cells were derived from ECs at early embryonic stages similar to brain macrophage (Extended Data Fig. 1H)²⁰. However, B-1a cells were marked by ECs during E7.5 to E10.5 140 141 (Extended Data Fig. 2, 3). Other B-cell subsets including FO and MZ B cells showed similar 142 labeling patterns (Extended Data Fig. 2, 3). Tomato% ratios of these B-1a and B-2 cells to HSCs 143 were just slightly higher than 1.0 when ECs were labeled at E7.5 or E8.5 (Fig. 1I, J). When E10.5 144 ECs were labeled, almost all B-lymphoid subsets including B-1a cells showed similar Tom % with 145 that of HSCs (the ratio of nearly 1.0) (Fig. 1K). These results suggest that B-1a cells, a part of 146 other B-cell subsets, and HSCs were produced simultaneously from ECs because HSCs after 147 E11.5 contributed to only a part of each B-cell subset (Fig. 1E, Extended Data Fig. 1C). Taken 148 together, the data indicate that most B-1a cells are derived from ECs during E7.5 to 10.5, and a 149 portion of FO and Marginal zone B-cells are also HSC-independent.

150 In order to explain the results from the *iCdh5* mouse (Fig. 1I-K, Extended Data Fig. 1C), 151 we constructed a mathematical model of label tracing experiments by extending a previously established model²³. We constructed three variants of the label tracing model based on competing 152 153 hypotheses for the cell differentiation tree (Extended Data Fig. 4A). The base model M_0 assumes 154 a linear differentiation path from hemogenic EC via HSC, MPP, B-1 progenitor, and finally to B-1 155 cell. The M_1 model hypothesizes that ECs can directly differentiate into MPPs. In addition to M_1 156 model, the M_2 model hypothesizes that ECs can directly differentiate into B-1 progenitors. After 157 model fitting, the M_2 model yielded label tracing predictions that most closely resemble the 158 experimental label tracing data from the *iCdh5* mouse (Fig. 1I-K, Extended Data Fig. 1C, 4, 5). 159 These results indicate that the differentiation tree of the M_2 model can best explain the 160 experimental data, providing additional support that ECs directly differentiate into MMPs and B-1 161 progenitors independently of HSCs during fetal development.

162

Single cell-RNA-sequencing showed heterogeneity and B-lymphoid signatures of pre-HSC and
HSC population

165 Lineage tracing studies demonstrated the multiple waves of HSC-independent 166 hematopoiesis including MPPs, B-progenitors, and B-1a cells. We recently reported that E10.5 HSC-precursor (pre-HSC) population shows B-1 biased repopulating ability¹⁴. Pre-HSCs are 167 168 intermediate precursors between HECs and adult repopulating HSCs detected during E10.5 to 169 11.5 and express VE-cadherin (VC), encoded by Cdh5²⁴. At E11.5, the first adult repopulating HSCs are detected²⁴. To understand the transition from pre-HSCs to HSCs or MPP, and their 170 171 heterogeneity of hematopoietic capability, we performed single-cell (sc) RNA-sequencing of 172 E11.5 AGM and YS VC⁺c-kit⁺EPCR⁺ pre-HSC population, E12.5 FL HSC (CD45⁺LSK⁺EPCR⁺) 173 and E14.5 FL HSCs (CD45⁺LSK⁺CD150⁺CD48⁻) (Extended Data Fig. 6). In parallel, selected 174 sorted subsets were transplanted into sublethally irradiated NSG neonates to validate their 175 hematopoietic capabilities (Extended Data Fig. 6D).

176 We sorted individual cells from above populations and generated single-cell full length 177 transcriptome using SMART-seq. We distributed the genes (read counts >10) in each cell and 178 excluded the cells which expressed less than 2000 genes and genes that were detected in less 179 than 10 cells (Extended Data Fig. 6E). At last, 95 cells and 11,814 genes were used for further 180 analysis. PCA analysis of scRNA-seq showed clear separation of FL HSCs from pre-HSCs (Fig. 181 2A). Notch-related genes were all highly expressed in many AGM cells but were downregulated 182 in FL HSCs (Extended Data Fig. 7) as previously reported transitional requirement of Notch signaling²⁵. Unbiased sc consensus clustering (SC3) of the whole transcriptomes separated these 183 184 cells into four distinct clusters (Fig. 2B). The top 10 differentially expressed genes are depicted 185 in Extended Data Fig. 8. We focused the expressions of HSC- and lymphoid cell-related genes in 186 each cell (Fig. 2C) and examined the trajectory of cell states with ordering of cells from E11.5 187 AGM to E14.5 FL HSCs (Extended Data Fig. 9) using pseudo-time analysis. Trajectory map 188 indicated the progression of E11.5 AGM pre-HSCs to E12.5 and E14.5 FL HSCs (Extended Data 189 Fig. 9). Almost all pre-HSCs and HSCs expressed essential genes for B-cell development, such 190 as *Ikz1* and *Tcf3* in addition to many HSC-related genes (Fig. 2C, Extended Data Fig. 9). Bcl11a, 191 important for fetal and adult B cell development and globulin switching, is also widely expressed 192 among pre-HSCs and HSCs (Fig. 2C, Extended Data Fig. 9). Interestingly, essential BCR 193 signaling related genes, such as CD79b and Btk, were heterogeneously expressed in all cell types, 194 even in FL HSCs, suggesting their biased B-lymphoid potential (Fig. 2C, Extended Data Fig. 9). *Lin28b*, encoding RNA-binding protein and critically important for B-1a cell generation^{11, 26}, is also 195 196 heterogeneously expressed among pre-HSC and HSC populations (Extended Data Fig. 9), which may explain the contrasting results of FL HSCs transplantation assays^{11, 12, 13, 14, 27}. *Pbx1*, the Hox 197 198 cofactor and proto-oncogene, is essential for B-lymphoid lineage commitment and HSC-selfrenewal/maintenance^{28, 29, 30}. While Pbx1 was widely expressed among pre-HSCs, its expression 199 200 was only seen in a small portion of HSCs (Fig. 2C, Extended Data Fig. 9). These data displayed 201 the heterogeneity of highly purified pre-HSC and HSC populations and even genes that have

202 been considered critical for HSC maintenance may not be expressed ubiquitously in FL HSCs. We also performed the velocity analysis^{31, 32} to understand the gene expression status of each 203 cell. Interestingly, the velocity analysis showed two directions of the gene expression status into 204 205 the left and the right shown in Fig. 2D. When we compared the gene expressions between the 206 cells on the left and right sides, 41 genes were differentially expressed (supplementary file). We 207 examined these gene expressions in BM HSPC and B-progenitors from the database at ImmGen 208 (Fig. 2E) and found the difference of these gene expressions reflected the ones found in BM B-209 progenitors or HSPCs. This result suggests that there may be divergence into HSC and B-cell 210 commitment of pre-HSC and HSC populations.

211

212 HSC-independent MPPs and B-1 biased repopulating cells are present in E11.5 Embryo.

213 Upon the heterogenous gene expression signatures of HSPC and B-cell lineages in pre-214 HSC population, we examined the presence of HSC-independent MPPs and B-1a precursors in 215 the pre-HSC population in transplantation settings. We injected 5 -10 cells of CD45⁺Ter119⁻VC⁺c-216 kit⁺EPCR⁺ pre-HSC population isolated from E11.5 aorta-gonad-mesonephros (AGM) region 217 (Extended Data Fig. 6A) into sublethally irradiated NSG neonates. Of total 25 recipient mice, 15 showed donor-derived CD45.2⁺ cells (>0.1%) in the peripheral blood (PB) (Fig. 3A). When we 218 219 analyzed the transplanted mice at 4-6 months post transplantation (Fig. 3B-D), we found there 220 were four types of engraftment patterns; multi-lineage engraftment including BM LSK cells (HSC-221 engraftment, Fig. 3B-D, E-G, mouse #1-5); multi-lineage engraftment without BM LSK cells (MPP 222 engraftment, Fig. 3B-D, H, I, J mouse #6-11); and only B-1 and B-2 cell engraftment (Fig. 3B, C, 223 mouse #12); and only B-1 cell engraftment (Fig. 3B, C, K, mouse #13-15).

Mice #1-5 showed long-term multi-lineage repopulation with significant donor cell % in the PB, peritoneal cavity, spleen, and BM (Fig. 3A-E) and predominant B-2 cell engraftment with B-1a and B-1b cells (Fig. 3B). In the recipient BM, successful donor derived LSK cell repopulation was also confirmed (Fig. 3F). Thus, these mice were categorized as HSC-repopulated mice.

228 Importantly, the secondary recipient BM also showed donor-derived LSK repopulation (Fig. 3G). indicating that LSK-repopulating cells are the functional HSCs that harbor a self-renewal ability. 229 230 In contrast, mouse #6-11 showed long-term reconstitution without donor-LSK cells in the BM (Fig. 231 3A-D, H, I, J), indicating that they were engrafted with HSC-independent progenitors, thus named 232 MPP-engrafted mice. These mice showed predominant B-1 and MZ B cell engraftment with 233 seemingly diminishing B-2 cells in the peritoneal cavity and spleen (Fig. 3B, C) and B-2, T, and 234 myeloid cell engraftment in the PB and BM (Fig. 3D, H, I, J), although the donor percentage in 235 the BM was very low (<0.5%) (Fig. 3D, J). Mouse #13-15 showed B-1a, B-1b, and MZ B-cell 236 engraftment in the peritoneal cavity and spleen but did not show donor-derived cells in the BM 237 (Fig. 3B-D), indicating HSC-independent B-1 cell engraftment. One mouse (#12) showed only B-238 1 and B-2 cell engraftment (Fig. 3B-D, K), suggesting the presence of common B-1 and B-2 239 progenitors.

These results strongly indicate that HSC-independent long-term engraftable MPPs and B-1 precursors are present in E11.5 AGM region, also in line with the scRNA-seq data showing the heterogeneity of these cells. Importantly, all engrafted mice showed B-1a cell repopulation whereas B-2 cells were dominant in HSC-engrafted mice; thus, it seems that B-1a potential is the default within E11.5 pre-HSCs and B-2 cell dominant capacity is a hallmark of functional HSCs.

245

246 Discussion

We demonstrated that definitive hematopoietic cells including MPPs, all B-cell subsets, and HSCs are independently produced from HECs and persist into adult life. While HSC-independent B-1a cells are rarely replaced by HSC-derived cells, HSC-independent MPPs and B-2 cells are gradually replaced by HSC-derived cells. These findings challenge the current paradigm of HSCderived hematopoiesis and finally clarified the longstanding unresolved question regarding the origin and main source of B-1a cells.

253 In adult murine hematopoiesis, it has recently been reported that MPP is a main driver of 254 native hematopoiesis and the discrepancy of HSC- and MPP-derived clones have been 255 observed³³. Our results explain this discrepancy, where the majority of MPPs in young mice are 256 HSC-independent, derived from fetal ECs, and HSC-derived hematopoiesis appears later, This 257 is also in line with the previous report that postnatal HSC-labeling showed gradual increase of HSC-derived-lymphocytes over 32 weeks^{16, 23, 34, 35} and a recent report showing minimal 258 contribution of definitive HSCs to fetal hematopoiesis in a fish model³⁶. In addition, the presence 259 260 of HSC-independent MPPs in the pre-HSC population has also been reported using AGM-EC co-261 culture system¹⁸. Therefore, together with our results, pre-HSC population is essential not only 262 for maturing to HSCs but also containing MPPs that support hematopoiesis in postnatal life.

263 Our lineage tracing results (Fig. 4A), difficult to reconcile with the current classic model 264 (Fig. 4B), instead lead us to propose a multiple hematopoietic wave model (Fig. 4C). In the current 265 classical model (Fig. 4B), YS-derived EMPs maintain hematopoietic homeostasis until perinatal 266 periods³⁷. Once HSCs are produced in the embryo, HSCs expand in the FL, start HSC-derived 267 hematopoiesis, and migrate to the BM just before birth, where HSCs maintain hematopoiesis for 268 life. Our model proposes that multiple waves of HSC-independent hematopoiesis, including EMP, 269 B-1 precursor and MPP production, occur from ECs, and persist into adult life. HSCs are also 270 produced as the final wave of EC-derived blood production. While HSCs expand in the FL, HSC-271 derived hematopoiesis seems to first start after settling the BM, and gradually replaces the HSC-272 independent hematopoietic cells over time. B-1a precursors are produced directly from ECs or 273 via HSC-independent MPPs at early- and mid-gestation but not replaced by HSC-derived cells in 274 steady state.

There are some discrepancies between lineage tracing study and transplantation assays regarding the B-1a cell potential. While E11.5 EC labeling did not mark B-1a cells, E11.5 VC⁺ pre-HSCs showed B-1a repopulating ability. This may be due to the difference of expression timings of Cdh5 transcriptome and VC surface protein. In addition, the first HSCs at E11.5 possess B-1a

- 279 repopulating ability upon transplantation whereas FL HSCs did not. Therefore, HSCs may have
- 280 B-1a cell potential within a limited time window in transplantation or stress settings.
- 281 Taken together, we unveiled unappreciated presence of HSC-independent hematopoietic
- progenitors in adult life and challenge the paradigm of HSC-dogma in hematology.

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292 Some figures were created with BioRender.com.

293

294 Data availability

295 Sequencing data have been deposited in the GEO database under accession number 296 GSE182206.

All original code has been deposited at github and is publicly available as of the date of publication.

298 URLs are listed in the key resources table.

299

300 Author contributions

M.K. conceived, design, and performed experiments and analyzed the results. H.W., T.Y., J.S. and J.Q.W participated in bioinformatics data analysis. D.J.S and W.J.Z calculated the mathematical model, N.A.P., S.C., N.V., C.N. performed experiments, J.K. analyzed the results and edited the manuscript, M.Y. conceived, design, and performed experiments, analyzed the results, wrote and edited the manuscript.

307 Methods

308 Experimental Animals

309 Cdh5(PAC)-CreERT2 mice were obtained from Dr. Ralf Adams. Fqd5CreERT2 mice (Stock No: 310 027789) and Rosa-TdTomato mice (Stock No: 007909) were obtained from Jackson Laboratory. 311 Cdh5(PAC)-CreERT2 mice were crossed with Rosa-TdTomato mice and Cdh5CreE2:Rosa-312 Tomato mice were generated. Similarly, Fgd5CreERT2 mice were crossed with Rosa-TdTomato 313 mice and Fgd5CreERT2:Rosa-Tomato mice were generated. These mice were timed mated with 314 Rosa-Tomato mice and the vaginal plugs were confirmed in the following morning. The noon on 315 the day that the plug was found was counted as embryonic day 0.5. Tamoxifen (15ng/mother 316 body weight) was administrated into timed mated pregnant iCdh5 dams at E7.5, 8.5, 9.5, 10.5, 317 11.5 respectively, or into timed mated pregnant Fgd5 dams at E14.5, or P2 neonates. 10-15 mice 318 for each Tam injection date were examined.

319

For transplantation assays, C57BL/6 mice were timed mated and embryos at E11.5 were harvested from the pregnant dams. AGM region was dissected from the embryo and lin-CD144+ckit+EPCR+ cells were sorted for donor cells. The embryonic age was confirmed by the somite numbers and developmental features of the embryos.

NOD/SCID/II2γc^{null} mice (NSG mice. Jackson Laboratory Stock No: 005557, OD.Cg-*Prkdc^{scid} II2rg^{tm1WjI}*/SzJ mice) were timed mated and day2-5 neonates were used for recipients of transplantation assays. Recipient NSG neonates were sublethally irradiated (150rad) before donor cells were injected into facial vein. For secondary transplantation, 1-2 million BM cells from the primary recipient mice were injected into lethally irradiated adult BoyJ mice.

329 Mice were kept in specific pathogen free condition and all the experimental procedures using the

330 mice were approved by Animal Welfare Committee at UTHealth.

331

332 Lineage tracing experiments

333 Cdh5CreERT2:Tosa-Tomato mice were timed mated. A single dose of Tamoxifen (TAM)(Sigma) 15ng/mother body weight together with Progesterone (7.5ng/mother body weight) solved in corn 334 335 oil was administrated to the pregnant dam by oral gavage at E7.5, 8.5, 9.5, 10.5, and 11.5 336 respectively. TAM usually makes the delivery difficult, therefore, Cesarean section was performed 337 on day 19 pregnant dams to rescue the embryos and these pups were taken care of a surrogate 338 mother prepared in advance. Fgd5CreERT2: Rosa-Tomato mice were used for marking HSCs. 339 therefore, TAM was injected into E14.5 pregnant dam or P2 neonatal mice. We harvested various 340 hematopoietic tissues including peritoneal cells, spleen, thymus, and bone marrow from TAM 341 administrated embryos/mice and examined Tomato⁺ percentages in each hematopoietic subset. 342 We compared relative Tomato+ percentage between the cell population of interest and HSCs. 343 The surface markers used for flow cytometry is listed in the table.

344

345 Mathematical modeling of label tracing data

We extended a previously established mathematical model for label tracing data²³. Consider three successive cell compartments in hematopoiesis: upstream u, reference r, and downstream d. For example, in the differentiation path HEC \rightarrow HSC \rightarrow MPP, compartment urepresents hemogenic EC, compartment r represents HSC, and compartment u represent MPP. The change in cell number N_r over time t is given by:

351

$$\frac{dN_r}{dt} = \beta_r N_r + \alpha_{u,r} N_u - \alpha_{r,d} N_r$$

353

where β_r is the net proliferation rate, $\alpha_{u,r}$ is the differentiation rate from compartment *u* to *r*, and similarly for $\alpha_{r,d}$. Solving this differentiation equation leads to an exponential growth model for N_r (when differentiation rates are set to zero). Biologically, over the lifetime of an organism, a 357 logistic growth model may be more realistic because each cell type would have an upper limit

358 on population size. We therefore modify the label tracing model as follows:

359

$$\frac{dN_r^l}{dt} = \beta_r \left(1 - \frac{\sum_l N_r^l}{\kappa_r}\right) N_r^l + \alpha_{u,r} N_u^l - \alpha_{r,d} N_r^l$$

361

which imposes a carrying capacity κ_r that modulates the net proliferation rate. Integer *l* indexes the cell number N_r in order to distinguish label compartments (Tomato⁻ vs. Tomato⁺). The label compartments share the same parameters (β_r , $\alpha_{u,r}$, $\alpha_{r,d}$), because these parameters depend on the cell type and are independent of label status. Conversely, both label compartments simultaneously experience a common population limit, because the proliferation rate is determined by the total cell number $\sum_l N_r^l$ marginalized over label status, as shown in the above equation.

369

We implemented discrete-time simulations for label tracing models with various differentiation trees (M_0, M_1, M_2) in the R environment (v4.1.1). The κ parameters were initialized to steadystate compartment sizes that were previously determined whenever available (Busch et al., 2015). The simulation models were used to predict cell numbers across time for each cell type and label compartment, from which the label proportions and label ratios were calculated. The model parameters were tuned so that the predicted label ratios resemble the experimentally determined Td-Tomato label ratios.

377

378 **B-progenitor colony forming assay**

Five hundred FL MPPs were plated to methylcellulose containing 10ng/ml IL-7 with 10⁵ OP-9 cells.
Eight days after plating, colony numbers were counted and each colony was picked up and

stained with anti-mouse CD45, AA4.1, CD19, B220, and CD11b to identify B-1 and B-2
progenitors using flow cytometry.

383

384 scRNA-sequencing

E11.5 pre-HSCs, E12.5 and 14.5 FL HSCs were single cell sorted into 96 well plate (1 cell /well). RNA were extracted from each well and converted into cDNA using SAMRT-Seq Single cell kit (Takara). DNA library was made and sequenced at Single Cell Genomic Core at the Baylor College of Medicine. Briefly, following cDNA synthesis, Nextera XT DNA library preparation kit (Illumina) was used to prepare library.120pg of cDNA was simultaneously fragmented and tagged with adapter sequences by transposome. The product was then amplified using 12 cycles of PCR and purified. Final library was sequenced using Illumina Novaseg 600.

392

393 **Bioinformatics analysis of scRNA-sequencing**

394 The quality of all sequenced samples was analyzed using FastQC. Raw reads were aligned to 395 the GRCm38 reference genome using STAR with default parameters. The expression count 396 matrix was generated using htseq-count. We filtered genes whose read counts less than 10 and cells that were less than 2000 genes. Read counts were normalized using DEseq2 with default 397 398 parameters. The normalized matrix was clustered by SC3. We chose K=4 for SC3 as the best 399 represented the heterogeneity in our dataset. Marker genes in each cluster were filtered by the 400 area under the ROC curve (auroc) > 0.85 and the adjusted p-values<0.01. Trajectory and 401 pseudotime analysis were performed by monocle2 package with default parameters.

402

403 PCA analysis was performed using scikit-learn software package (https://scikit-learn.org/stable/).
404 RNA Velocity analysis was performed according to the original article^{31, 32}. Unspliced pre-mRNA
405 counts and mature spliced mRNA counts for each cell were computed from the BAM files
406 generated above., then RNA velocity was computed using scVelo software with default

- 407 parameters. The result was visualized onto the PCA plot based on the expression profile of mature
- 408 spliced mRNA for each cell. Each arrow represents a direction of a cell transition based on the
- 409 RNA Velocity.
- 410
- 411 Statistical analysis
- 412 Non-parametric student-t test was used for statistical analysis.
- 413

414 Figure Legends

Figure 1. B-1a, MPP, and other lymphoid cells arise independently of HSCs and persist into adult.

417 (A) TAM was injected once at E14.5 or P2 into *iFqd5* mice to label HSCs. Tomato⁺ blood cells in 418 the BM, spleen, and peritoneal cavity were examined at different time points after birth. (B) The 419 relative Tomato% ratios of MPP, Follicular (FO) B cells, and B-1a cells to HSCs are shown. TAM 420 was injected to *iFgd5* mice at E14.5 or P2 and mice were examined more than 300 days after 421 birth (n=3-5). (C) TAM was injected into *iCdh5* pregnant mice at E11.5. Tomato⁺ HSPCs were 422 examined at different time points such as E15.5 and after birth. (D) The relative Tomato % ratio 423 of MPPs to LT-HSCs in E15.5 FL when ECs were labeled at E11.5 (n=4). (E) The relative 424 Tomato% ratios of MPPs, splenic FO, and peritoneal B-1a cells to LT-HSCs at different time points 425 (P0-5, days 60-90, days 300<, n=3-6 at each time point). ND: not done. (F) TAM was injected into 426 *iCdh5* pregnant mice at E7.5 and Tomato⁺ HSPCs were examined at different time points such 427 as E15.5 and after birth. (G) The relative Tomato% ratio of each target cell population to LT-HSCs in E15.5 FL, when ECs were labeled at E7.5 (n=6). (H) The relative Tomato% ratio of MPPs to 428 429 LT-HSCs in the FL and post-natal BM at different time points (P0-5, days 60-90, days 300<, n=3-430 5). The relative Tomato% ratios of HSPCs and B cell subsets to LT-HSCs at >300 days after birth 431 when TAM was injected at E7.5 (I), E8.5 (J), and E10.5 (K). N=3-7 for each TAM injection. BM: 432 bone marrow, SPL: spleen, PW: peritoneal wash.

433

Figure 2. scRNA-seq analysis showed HSC and B-lymphoid signatures in of E11.5 preHSCs and E12&14 FL HSCs.

(A) Dimensionality reduction of scRNA-seq data using PCA colored by cell type. E11A, E11.5
AGM pre-HSC, E11Y, E11.5 YS pre-HSC, E12F, E12.5 FL HSC, E14F, E14.5 FL HSCs. (B) SC3
consensus matrix predicted 4 clusters. (C) A heat map depicting the expression of HSC, B, T,
and ILC related genes in E11 AGM&YS pre-HSC populations and E12.5&14.5 FL HSCs. The red,

blue, and yellow intensities indicate high, low, and intermediate expression levels, respectively.
(D) Velocity analysis of scRNA-seq of E11.5 pre-HSC and E12.5 &14.5 HSCs. Small arrows show
the direction of the velocity of single cells. (E) Heat map of gene expressions detected in right and
left directions in the velocity analysis, which were applied to gene expressions in the BM HSPC
and B-progenitors using the database at ImmGen (https://www.immgen.org/).

445

Figure 3. LT-HSCs, MPPs, and B-1 repopulating cells arise independently from CD144⁺ckit⁺EPCR⁺ cells in E11.5 AGM region.

448 Five to fifty pre-HSCs from E11.5 AGM region were injected into sublethally irradiated NSG 449 neonates. (A) CD45.2⁺ donor cell % in the peripheral blood of the recipient mice 4-6 weeks after 450 transplantation. Donor cells % and their composition within the lymphoid subsets in the peritoneal 451 cells (B), spleen (C), and BM (D) are depicted. The donor-derived cell lineages in the recipient 452 PB during the time course of mouse #2 (E, HSC-engrafted), mouse #9 (I, MPP-engrafted), and 453 #12 (K, B-1 and B-2 cell-engrafted). (H) The % of donor-derived lineages in the recipient BM of 454 mouse #6-10 are depicted. Although the donor cell % was low, multi-lineage repopulation was 455 observed in the recipient BM. The representative FACS plots for donor LSK cells in the first (F, 456 mouse #2 and J, mouse #9) and secondary recipient BM (G).

457

Figure 4. The current and proposed models for developmental hematopoiesis in the mouse embryo based on the lineage tracing studies

(A) The summary of the results from HSC- and EC- lineage tracing studies. While HSC-lineage
tracing does not label HSC-independent blood lineages (left), EC-lineage tracing labels blood
cells with different percentages depending on the timing when those blood cells are produced
from ECs (right). EC-labeling at E7.5 marked more HSC-independent blood cell types than HSCs.
(B) The current classical hematopoiesis model during fetus. Hemogenic ECs produce EMP,
possibly lymphoid progenitors (not indicated), and HSCs. These EMPs and HSCs seed the fetal

466 liver where EMPs provide mature definitive erythroid and myeloid cells and HSC-derived hematopoiesis start while HSC self-renew and expand at the same time. (C) Our working model 467 468 proposing EC-derived multiple waves of fetal hematopoiesis. Almost all hematopoietic progenitors 469 including EMP, MPP, and B-1 progenitors are produced from hemogenic ECs during E7.5-10.5 470 independently of HSCs. HSC production is the final wave of EC-derived hematopoiesis. These 471 progenitors and HSCs seed the fetal liver and then bone marrow before birth. While HSCs mainly 472 self-renew and expand in the fetal liver, EC-derived (HSC-independent) blood progenitors 473 maintain hematological homeostasis and provide mature blood cell subsets until HSC-derived 474 progenitors replace them in postnatal life. However, peritoneal B-1a cells are not replaced by 475 HSC-derived progenitors and maintain themselves for life.

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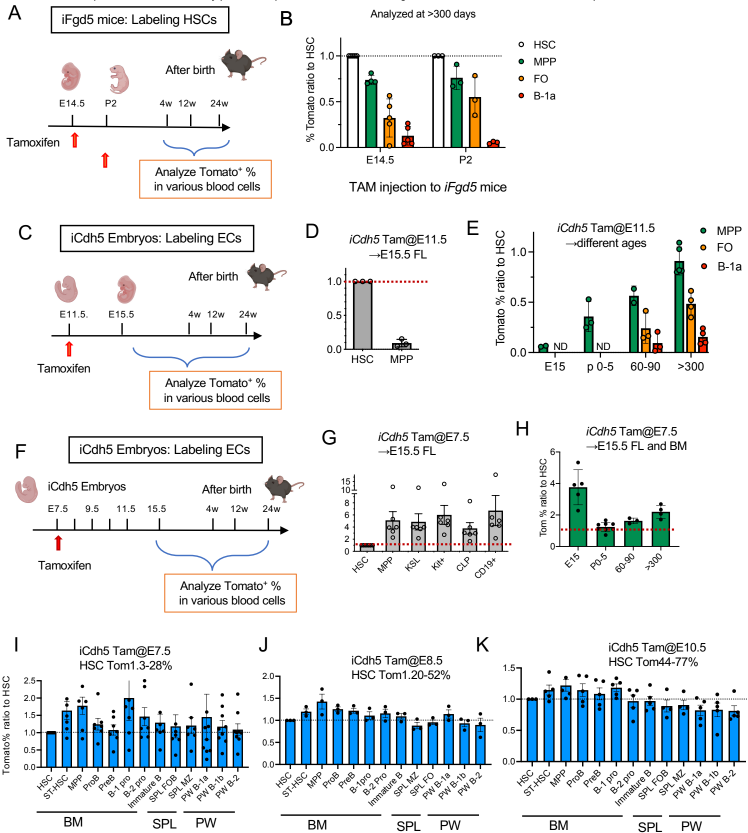
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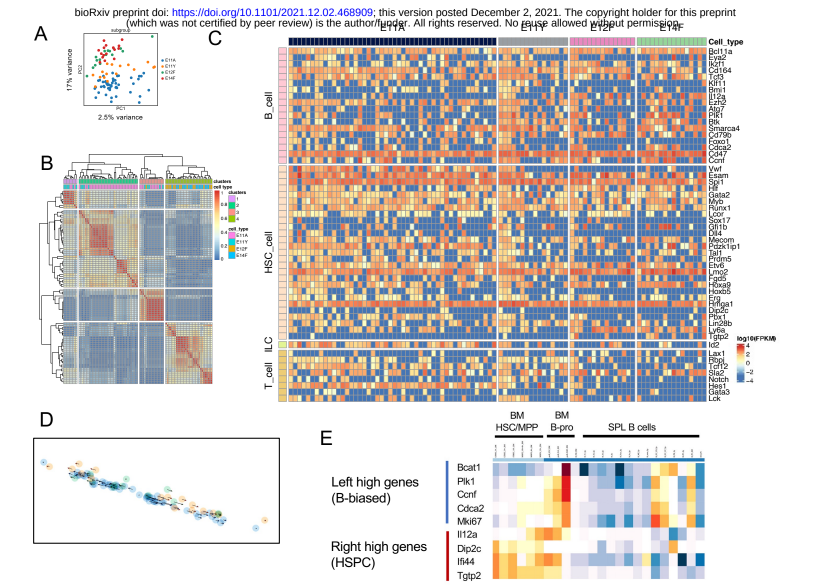
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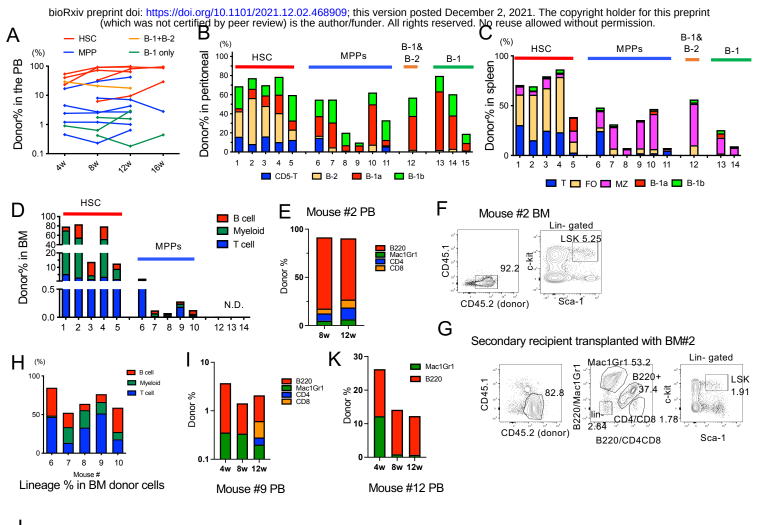
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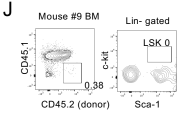


Figure 3

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