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Chemically-defined cytokine-free human hematopoietic stem cell expansion

1 2

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- 44
- 45 Keywords: Human hematopoietic stem cell; ex vivo expansion; chemically defined;
- 46 PI3K activator; polyvinyl caprolactam-polyvinyl acetate-polyethylene glycol graft
- 47 copolymer.

48 Abstract:

49 Hematopoietic stem cells (HSCs) are a rare cell type that reconstitute the entire blood and 50 immune systems following transplantation, a curative cell therapy for a variety of hematological diseases^{1,2}. However, the low number of HSCs makes both biological 51 52 analyses and clinical application difficult, and the limited ability to expand human HSCs 53 ex vivo remains a substantial barrier to the wider and safer therapeutic use of HSCT³. 54 While various reagents have been tested in attempts to stimulate human HSC expansion, 55 cytokines have long been thought to be essential for supporting HSCs ex vivo⁴. Here we 56 report the establishment of a novel culture system that supports the long-term ex vivo 57 expansion of human HSCs, achieved through the complete replacement of exogenous 58 cytokines and albumin with chemical agonists and a caprolactam-based polymer. A phosphoinositide 3-kinase activator, in combination with a thrombopoietin-receptor 59 60 agonist and the pyrimidoindole derivative UM171 were sufficient to stimulate expansion 61 of umbilical cord blood HSCs capable of serial engraftment in xenotransplantation assays. 62 Ex vivo HSC expansion was further supported by split-clone transplantation assays and 63 single cell RNA-sequencing analysis. We envision that this chemically-defined expansion 64 culture system will help to advance clinical HSC therapies.

65

66 Main text:

67 Self-renewing multipotent hematopoietic stem cells (HSCs) are a rare bone marrow (BM) cell population that support life-long hematopoiesis⁵⁻⁸ and hematopoietic 68 69 system reconstitution following HSC transplantation (HSCT)¹. HSCs can also be 70 collected from umbilical cord blood (CB), which represents a highly-accessible source 71 for transplantation but often contain too few HSCs for successful engraftment and durable 72 hematopoietic reconstitution. Ex vivo expansion of human HSCs, particularly CB-73 derived HSCs, is therefore a major goal in hematology and one that remains a substantial 74 barrier to the wider and safer therapeutic use of HSCs³.

75 Various recombinant cytokines are commonly added to human HSC cultures in 76 attempts to promote HSC expansion, usually in combination with serum albumin⁴. These 77 cultures generally support short-term maintenance of HSCs but fail to expand functional HSCs. However, two-week ex vivo expansion of human HSCs has been achieved by the 78 addition of small molecules, StemRegenin 1 (SR-1)⁹ and UM171¹⁰. Clinical trials using 79 these approaches to expand CB HSCs prior to transplantation have reported encouraging 80 81 results^{2,11}. Other recent approaches have included use of 3-dimensional zwitterionic hydrogels¹², addition of novel growth factors¹³, or combinations of small molecule 82 83 inhibitors¹⁴. These methods have highlighted the importance of collaboration between 84 chemical biology and stem cell biology to overcome this major barrier in hematology.

85

86 A chemically-defined cytokine-free media

87 Working towards the goal of expanding HSCs ex vivo, we recently established 88 a long-term ex vivo expansion system for functional mouse HSCs by optimizing the 89 concentrations of recombinant stem cell factor (SCF) and thrombopoietin (THPO), and replacing serum albumin for the synthetic polymer polyvinyl alcohol (PVA)^{15,16}. Use of 90 PVA avoided the batch-to-batch variability associated with serum albumin¹⁷ and culture 91 92 contamination with albumin-associated impurities that promote HSC differentiation. 93 While mouse HSCs expanded rapidly in these conditions, human HSC expansion was 94 more limited. When we compared the proliferation of mouse BM Kit+Sca-1+Linage-(KSL) HSPCs with human CB CD34⁺CD38⁻ HSPCs in PVA-based media supplemented 95 with SCF and THPO¹⁵, mouse HSPCs proliferated ~18-fold in 7 days while human 96

97 HSPCs only proliferated ~3-4-fold during the same time (Figure 1a). To examine the
98 difference between mouse and human HSPCs during these cultures, we analyzed the
99 phosphorylation status of major signaling pathways (PI3K, JAK/STAT, MAPK) linked
100 to SCF and THPO signaling^{8,18,19}. Significant decreases in PI3K and AKT were observed
101 in human cells (Figure 1b, Extended Data Figure 1a, b). The PI3K phosphorylation
102 signal was also significantly decreased in human CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺
103 phenotypic HSCs (Extended Data Figure 1c).

104 Based on these results, we hypothesized that we could improve human HSPC 105 expansion by activating PI3K-AKT signaling. We therefore evaluated chemical agonists 106 740Y-P (a PI3K activator) and SC79 (an AKT activator) in human HSPC cultures. While 107 SC79 did not improve expansion efficacy, 740Y-P significantly increased the number of 108 CD34⁺ cells (Figure 1c) and CD34⁺CD45RA⁻ cells in 7-day cultures (Extended Data 109 Figure 1d). Furthermore, addition of 740Y-P significantly increased PI3K 110 phosphorylation in CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cells (Extended Data Figure 111 1e). These results suggested that chemical activation of the PI3K pathway was sufficient to improve human HSPC proliferation. 112

113 Previous studies have shown that SCF stimulates HSC cell cycle entry via the 114 PI3K/AKT/FOXO pathway²⁰⁻²³. We therefore hypothesized that we could replace SCF 115 with 740Y-P in human CD34⁺ HSPCs cultures. No significant differences were observed 116 in the 7-day cell proliferation in THPO and 740Y-P with or without SCF (Figure 1d), 117 suggesting that SCF can be replaced with a PI3K activator. Cell cycle analysis confirmed 118 that the frequency of S/G2/M cells was comparable (Extended Data Figure 1f) while 119 colony forming unit (CFU) assays showed similar increases in multipotent granulocyte-120 erythrocyte-monocyte-megakaryocyte (GEmM) CFUs in the presence or absence of SCF 121 (Extended Data Figure 1g). These results suggested that SCF was replaceable with 122 740Y-P in ex vivo human HSPC cultures.

We previously reported that recombinant proteins could destabilize HSC cultures¹⁵. Recently, we found that chemical THPO receptor agonists (THPO-RAs) could be used to induce human HSC expansion²⁴. We therefore examined whether we could replace recombinant THPO with THPO-RAs in PVA-based media containing 740Y-P. Initial screening for optimal THPO-RAs was performed using a THPO-dependent MPL-

expressing cell line reported to proliferate in THPO-supplemented PVA conditions²⁵. Of 128 129 the three THPO-RAs tested, only butyzamide supported cell proliferation (Extended 130 **Data Figure 1h**). We validated that butyzamide stimulated human CD34⁺ cell 131 proliferation in PVA-based media containing 740Y-P (Extended Data Figure 1i). Surprisingly, when compared to the THPO and 740Y-P cultures, the butyzamide and 132 133 740Y-P cultures displayed significantly improved 7-day proliferation (total, CD34⁺ and CD34⁺CD41⁻CD90⁺CD45RA⁻ cell numbers) and GEmM CFU numbers (Figure 1e, 134 135 **Extended Data Figure 1j, k**). However, there was no additive effect of THPO and 136 butyzamide (Extended Data Figure 1j). In summary, these results confirmed that human 137 CD34⁺ HSPCs could be grown without exogenous cytokines by replacing SCF and THPO 138 with 740Y-P and butyzamide, respectively.

139 We next titrated 740Y-P and butyzamide concentrations for CD34⁺ cell expansion and identified the combination of 1 µM 740Y-P and 0.1 µM butyzamide as 140 141 optimal for cell expansion, in terms of both total and CD34⁺ cell expansion (Figure 1f). 142 We defined this combination of 740Y-P (1 µM) and butyzamide (0.1 µM) as two 143 activators (2a) media. Using this media composition, we next examined long-term 144 stability of CD34⁺ cell cultures. Although total cell numbers increased during 14-day 145 cultures, the number of phenotypic CD34⁺ cells decreased between day 7 and 14, and the cultures became dominated by CD41⁺ cells (Figure 1g, Extended Data Figure 1l). 146 147 Consistent with accumulation of these CD41⁺ megakaryocyte-lineage cells (Extended Data Figure 1m), significant increases in megakaryocyte (MgK) CFUs were observed in 148 the day 14 cultures (Extended Data Figure 1n). Additionally, while 1x10⁴ cells from 7-149 day 2a cultures engrafted robustly in immunodeficient NOD/Shi-scid IL-2Ry^{null} (NOG) 150 151 mice²⁶, chimerism was not detected from 14-day 2a cultures (Extended Data Figure 10, 152 **p**). Together, these results suggested that although 2a cytokine-free media supported 153 human HSCs short-term, it was not sufficient to stabilize longer-term expansion.

154

155 Long-term ex vivo HSC cultures

Based on our 2a culture results, we searched for potential MgK inhibitors to
stabilize long-term ex vivo HSC expansion. We evaluated two reported HSPC expansion
compounds, SR-1⁹ and UM171¹⁰. Addition of UM171 increased the total, CD34⁺ and

159 CD34⁺EPCR⁺ cell numbers after 7-days (Figure 2a, Extended Data Figure 2a). In 14-160 total. CD34⁺. $CD34^{+}EPCR^{+}$ dav UM171 supplemented cultures, and 161 CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells were significantly increased while CD41⁺ cell numbers were reduced (Figure 2b, Extended Data Figure 2b). Furthermore, the 162 163 expansion of CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells was significantly higher with UM171 at 70 nM as compared to than 35 nM (Extended Data Figure 2c). Meanwhile, 164 165 the addition of SR-1 induced apoptosis (Extended Data Figure 2d). This three activator 166 (3a) media cocktail of UM171 (70 nM), 740Y-P (1 μ M) and butyzamide (0.1 μ M) 167 continued to stimulate proliferation over a 30-day culture by ~14-fold (Figure 2c). These 168 results suggested that HSPCs may be stably expanding in the 3a media.

169 To evaluate the in vivo engraftment and differentiation potential of the cultured 170 HSPCs, we performed xenotransplantation assay. We transplanted 1×10^4 CD34⁺ cells 171 before culture (fresh) and after 10-day or 30-day cultures. Significantly higher human 172 CD45⁺ PB chimerism was observed in the 10-day and 30-day culture groups, with 173 chimerism increasing over time (Figure 2d, Extended Data Figure 2e). BM analysis at 174 16- and 24-weeks also identified significantly higher human cell chimerism in the 175 cultured HSPC groups (Figure 2e, Extended Data Figure 2f); human CD45⁺ chimerism 176 from the fresh group was \sim 3%, while 10-day and 30-day culture groups displayed \sim 70% 177 and \sim 85% chimerism, respectively (Figure 2e). The frequency of human CD34⁺ cells in 178 the BM and spleen at 16- and 24-weeks was also significantly higher in the 10- or 30-day 179 culture group (Figure 2f, Extended Data Figure 2f-g, where multilineage output was 180 also observed (Extended Data Table 1a). These results confirmed that our cytokine-free 181 3a media could maintain and expand functional human HSCs for at least one-month ex 182 vivo.

183

184 Caprolactam polymers improve HSC growth

Having established an albumin- and cytokine-free human HSC culture system,
we next aimed to improve the rate of HSPC expansion ex vivo. The PVA-based 3a media
only supported ~10-fold expansion of CD34⁺ cells over 30 days, suggesting that further
improvement was required. We hypothesized that other synthetic polymers might be more
suitable for human HSC expansion. We therefore screened 10 polymers and identified

190 Soluplus[®], a polyvinyl caprolactam-polyvinyl acetate-polyethylene glycol graft copolymer (PCL-PVAc-PEG)^{27,28}, as supportive of significantly higher cell expansion 191 192 (Figure 3a). In PCL-PVAc-PEG-based cultures, the combination of 740Y-P, butyzamide, 193 and UM171 was as effective as in the PVA-based cultures (Extended Data Figure 3a-194 f). However, the toxicity of SR-1 was significantly reduced compared to the PVA condition (Extended Data Figure 3g). PCL-PVAc-PEG-based 3a media also supported 195 196 faster cell proliferation longer-term, with a ~75-fold expansion of total cells and ~55-fold 197 expansion of CD34⁺ cell observed after a 30-day culture (Figure 3b). The addition of a 198 PI3K inhibitor led to cell death, suggesting that cell expansion was dependent on the 199 PI3K/AKT signaling (Extended Data Figure 3h). Furthermore, PVA- and PCL-PVAc-200 PEG-based 3a media also supported ex vivo expansion of adult-PBSC CD34⁺ cells, with 201 a ~8-10-fold expansion of total cells observed after a 10-day culture (Extended Data 202 Figure 3i).

203 To compare in vivo engraftment and differentiation potential of the PCL-PVAc-204 PEG and PVA cultured HSPCs, we performed xenotransplantation assays. We 205 transplanted 1x10⁴ cells per recipient from day-30 3a media cultures containing PVA 206 and/or PCL-PVAc-PEG. Interestingly, similar robust human cell chimerism was 207 observed from all conditions, including in the PB, BM, and spleen (Figure 3c-e, 208 Extended Data Figure 3j-l, Extended Data Table 1b, Supplementary Table 1). 209 Robust human CD45⁺ chimerism was also observed in secondary xenotransplantation 210 recipients (Figure 3f-g, Supplementary Table 2). However, lymphoid bias was observed due to the characteristics of the NOG mice^{26,29}. We re-performed the 211 xenotransplantation assays using human IL-3/GM-CSF-transgenic NOG (NOG IL-212 3/GM-Tg) mice³⁰. In this context, we observed robust engraftment from long-term PCL-213 PVAc-PEG-based 3a cultures, with CD33⁺ myeloid cells at 27% of human CD45⁺ cells 214 215 at 16-weeks post-transplantation (Figure 3h, i).

We next compared 10-day CD34⁺ cell expansion in PCL-PVAc-PEG-based 3a
media against published serum albumin-based media (StemSpan SFEM supplemented
with cytokines and SR1 or UM171). While the total number of cells generated in the 10day cultures was ~50% in 3a media (Extended Data Figure 4a), the frequency and
absolute number of CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells was strikingly increased

221 (Extended Data Figure 4b, c). PCL-PVAc-PEG-based 3a media was also superior to 222 PCL-PVAc-PEG-based cytokine cocktail media (Extended Data Figure 4b, c). 223 Consistent with higher metabolic activity and active cell division, all culture conditions 224 caused an increase in ROS and yH2AX compared to fresh CD34⁺ cells (Extended Data 225 Figure 4d); however, these did not accumulate further in longer-term 30-day 3a cultures 226 (Extended Data Figure 4e). Corresponding with the increased frequency of HSPCs, the 3a cultured cells also demonstrated significantly higher human CD45⁺ PB and BM 227 chimerism following transplantation of 1x10⁴ 10-day cultured cells into recipient NOG 228 229 mice (Extended Data Figure 4f, g). Human CD45⁺ chimerism was detected in the PB 230 and BM of 1/3 secondary transplantation recipients at 24-weeks post-transplantation 231 (Extended Data Figure 4h, i). It is worth noting that our xenotransplantation assay protocol differs to those used in the development of SR-1¹¹, which may account for the 232 233 differences in engraftment. Nonetheless, these results confirmed that 3a media support 234 robust expansion of functionally engraftable human HSCs ex vivo.

235

236 Long-term selective HSC expansion

The robust in vivo engraftment potential of these human HSC cultures suggested that a high frequency of HSCs was maintained within the 3a cultures. To further investigate the composition of these long-term HSPC cultures, we performed flow cytometric analysis of HSC-associated surface markers. This revealed a striking enrichment of phenotypic HSCs after PCL-PVAc-PEG culture, with the CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cell fraction significantly enriched after 7-day cultures (Extended Data Figure 4b-c, Extended Data Figure 5a).

We also sought to characterize the PCL-PVAc-PEG based 3a cultures at the
molecular level. We performed whole exome sequencing on fresh and 10-day cultured
CB CD34⁺ cells and detected seven mutations that could a cause amino acid change and
four mutations that were located on a splice site in cultured cells (Extended Data Table
2). To the best of our knowledge, these mutations are not involved in hematological
malignancies nor clonal hematopoiesis. These results support the safety of our culture
system, and suggest further clinical development is warranted.

We next performed bulk RNA sequencing on CD34^{high}EPCR⁺ 251 and 252 CD34^{high}EPCR⁻ cells from 10-day cultures. EPCR expression is known to mark LT-HSCs 253 in UM171-supplemented media³¹. Consistent with previous studies³², expression of LT-254 HSC markers HLF and AVP were enriched in the CD34^{high}EPCR⁺ fraction (Extended Data Figure 5b). Additional HSC genes, *PRDM16³³* and *FGD5³⁴*, were also upregulated 255 256 in the CD34^{high}EPCR⁺ fraction (Figure 4a) and Gene Set Enrichment Analysis (GSEA) confirmed that HSC gene sets were upregulated in the CD34^{high}EPCR⁺ fraction (Figure 257 258 4b). GSEA also revealed that lysosomal membrane related genes were enriched in 259 CD34^{high}EPCR⁺ cells (**Extended Data Figure 6b**), consistent with a report that lysosomal 260 activity against various external signals has an important role in the self-renewal of human LT-HSCs³⁵. On the other hand, oxidative phosphorylation (OXPHOS) and mitochondrial 261 ribosomes genes were upregulated in CD34^{high}EPCR⁻ cells (Figure 4c, Extended Data 262 263 Figure 6b), consistent with a report that high mitochondrial membrane potential (MMP) 264 HSCs had less intracellular lysosomal contents than quiescent MMP-low LT-HSCs³⁶.

To further resolve the cellular heterogeneity within the HSC cultures, we used 265 single-cell RNA sequencing to compare our 3a conditions with StemSpan SFEM 266 supplemented with cytokines and SR1 or UM171^{2,11}. After integration and analysis of 267 268 these three samples using Seurat, we identified and manually annotated 12 major clusters 269 (Figure 4d, Extended Data Figure 6c). This included a population of cells expressing 270 HSC genes, HLF and AVP, while lacking expression lineage-specific genes (MPO, 271 ITGA2B), which we termed HSPC-HLF (Figure 4d, Extended Data Figure 6c-e). Lower 272 expression of HLF and AVP were also seen in two other populations (HSPC, HSPC-273 cycling), suggesting these to be intermediate stem/progenitor populations (Extended 274 **Data Figure 6c**). Various downstream progenitor cell types, including erythroid, MgK, 275 monocyte (Mon) and granulocyte (Gra) progenitors could also be identified (Figure 4d, 276 Extended Data Figure 6c). Comparing the cellular composition of the 3a media and 277 other two cytokine-based conditions (StemSpan SFEM with SR-1 or UM171), 278 differences were apparent. In particular, high frequencies of the HSPC-HLF cluster and 279 erythroid/MgK progenitors were observed in the 3a media, while the Mon and Gra 280 progenitor clusters were generally depleted (Figure 4e, Extended Data Figure 6f). 281 Similar results for the UM171 cultures were seen when we overlayed published singlecell RNA-seq datasets onto our dataset (Extended Data Figure 6g). These results suggest
that 3a culture conditions were more suitable for selective expansion of LT-HSCs. These
differences in the cellular composition corresponded with the higher engraftment
potential seen in the 3a media (Extended Data Figure 4f, g) and confirmed the highly
selective nature of these cultures. In addition, 10-day cultured cells in 3a media had higher
levels of *HLF* expression as compared to fresh CB CD34⁺ cell samples³² (Extended Data
Figure 6h).

289 Finally, we examined whether 3a media could support the expansion of clonally-290 derived HSC cultures. Single CB-derived CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cells 291 were sorted into 96-well plates and cultured with PCL-PVAc-PEG-based 3a media. After 292 using 7 days, xenotransplantation assays were performed NOD.Cg-Prkdc^{scid}Il2rg^{tm1Sug}Kit^{em1(V831M)Jic}/Jic (W41/W41) recipient mice (Figure 4f), which 293 294 support higher human hematopoietic cell chimerism in xenotransplantation assays 295 (Extended Data Figure 8). Although heterogeneous clonal expansion was observed, 20 296 out of 96 wells (21%) expanded more than 10-fold and 11 out of 96 wells (11%) expanded 297 over 30-fold in 7 days (Figure 4f, Supplementary Table 3). The 10 wells with the 298 highest expansion rate were transplanted into individual recipients. Five out of the ten 299 recipients displayed robust PB, BM, and spleen chimerism with over 5% multilineage 300 human CD45⁺ chimerism in the BM and spleen after 24-weeks (Figure 4h, i, Extended 301 Data Table 1c Supplementary Table 4). Furthermore, we performed split clone 302 experiments by transplanting single HSC-derived cultures into three W41/W41 mice 303 using HSC clones that displayed more than 10-fold expansion by day 7. For three out of 304 six clones, all three recipients showed human cell engraftment (Extended Data Figure 305 **7a**, **b**), confirming that clonal amplification of HSCs is supported by the 3a media. 306 Together, these results confirm that PCL-PVAc-PEG-based 3a media can support both 307 bulk and clonal expansion of human HSCs ex vivo.

308

309 Discussion

Here, we report a recombinant cytokine-free albumin-free long-term expansion
culture system for human HSCs. These conditions selectively expanded functional
HSPCs for at least 30 days ex vivo and also supported clonal HSC expansion, which may

313 contribute to efforts to decipher the heterogeneity of the human HSC compartment in health and disease³⁷. As highlighted by recent clinical trials^{2,11}, the ability to expand CB 314 HSPCs ex vivo has important implications for solving the shortage of donor HSCs for 315 316 allogeneic HSCT. In this regard, 3a media may hold advantages in being recombinant 317 protein-free and chemically defined, which should improve batch-to-batch variability, 318 reduce reagent costs and facilitate rapid clinical translation. However, further comparison 319 of human HSC culture methods are warranted. In conclusion, this culture system provides 320 a powerful platform for both basic scientists and clinicians interested in stem cell biology 321 and HSC therapies.

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- 323

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14

- 415 Figure Legends:
- 416 Figure 1. Chemically-defined cytokine-free media maintains human hematopoietic
- 417 stem/progenitor cells (HSPCs) ex vivo
- 418 (a) Ex vivo proliferation of 1000 mouse bone marrow (BM) c-Kit⁺Sca-1⁺Linage⁻ (KSL)
- 419 HSPCs or 1000 human cord blood-derived (CB) CD34⁺CD38⁻ HSPCs cultured with 10
- 420 ng/ml SCF and 100 ng/ml THPO in polyvinyl alcohol (PVA)-based media. Mean of five

421 independent cultures. **P = 0.0034; ***P = 0.0001; ****P < 0.0001.

- 422 (b) Single cell phosphorylation status of PI3K and AKT in mouse KSL and human CB
- 423 CD34⁺CD38⁻ cultured with PVA-based media containing SCF and THPO. Mean of 30
- 424 cells. AFI: average fluorescence intensity. ****P < 0.0001.
- 425 (c) Fold change in total cell numbers of human-cord-blood-derived CD34⁺CD38⁻ cultured
- 426 with SC79 or 740Y-P in addition to human 10 ng/ml SCF (S) and 100 ng/ml THPO (T)
- 427 in PVA-based media conditions. The starting cell count was 1000. Mean of five
- 428 independent cultures. *P = 0.0145; **P = 0.0051.
- 429 (d) Fold change in total and CD34⁺ cell numbers after a 7-day culture of $2x10^4$ human
- 430 CB CD34⁺ cells in PVA-based media containing 740Y-P and 100 ng/ml THPO (T) with
- 431 or without 10 ng/ml SCF (S). Mean of three independent cultures.
- **432** (e) Fold change in total and CD34⁺ cell numbers after a 7-day culture of $2x10^4$ of human
- 433 CB CD34⁺ cells in PVA-based media containing 740Y-P and THPO (T) or butyzamide
- 434 (Buty). Mean of three independent cultures. $**^{\dagger}P = 0.0020$; $**^{\dagger}P = 0.0054$.
- **435** (f) Fold change in total and CD34⁺ cell number after a 7-day culture of $2x10^4$ human CB
- 436 CD34⁺ cells in PVA-based media containing 0-20 μ M 740Y-P and 0.01-0.5 μ M
- 437 butyzamide. Mean of three independent cultures.
- 438 (g) Cell numbers and phenotypes during the culture of 2.5×10^3 human CB CD34⁺ cells in
- 439 PVA-based media containing 1 μ M 740Y-P and 0.1 μ M butyzamide. Mean <u>+</u> S.D. of
- 440 three independent cultures.
- 441 Statistical significance was calculated using an unpaired two-tailed *t*-test: n.s., not442 significant.

- 443 Figure 2. Long-term ex vivo expansion of human HSPCs in chemically-defined
 444 cytokine-free cultures
- (a) Fold change in total and CD34⁺ cell numbers after a 7-day culture of $2x10^4$ human
- 446 CB CD34⁺ cells in PVA-based media containing 750 nM SR-1 and/or 70 nM UM171 in
- 447 addition to 2a media containing 1 μ M 740Y-P and 0.1 μ M butyzamide. Mean of three
- 448 independent cultures. **P = 0.0095.
- (b) Total, CD34⁺, and CD41⁺ cell numbers after a 14-day culture of $2x10^4$ CB CD34⁺
- 450 cells in PVA-based 2a media with or without 70 nM UM171. Mean of three independent

451 cultures. ***P = 0.004; ****P < 0.0001.

- 452 (c) Fold change in total and CD34⁺ cell numbers during a 30-day culture of $2x10^4$ CB
- 453 CD34⁺ cells in PVA-based media containing 1 μM 740Y-P, 0.1 μM butyzamide, and 70
- 454 nM UM171 (PVA-based 3a media). Mean of three independent cultures.
- 455 (d) Mean human CD45⁺ PB chimerism in recipient NOG mice at 4-, 8- and 12-weeks
- 456 following transplantation of 1×10^4 fresh CB CD34⁺ cells or the cells derived from a 10-
- 457 day or 30-day culture of 1×10^4 CB CD34⁺ cells in PVA-based 3a media. n=5-6 mice per
- 458 group. *P = 0.0401; ***P = 0.0001; ****P < 0.0001.
- (e) Mean 16-week human CD45⁺ cell chimerism in the BM and spleen from mice described in (d). n=3 mice per group. ***P = 0.0003; ****P < 0.0001.
- 461 (f) Mean 16-week human CD34⁺ cell chimerism in the BM and spleen from mice
- 462 described in (d). n=3 mice per group. $*^{\dagger}P = 0.0116$; $*^{\dagger}P = 0.0416$; .**P = 0.0055.
- 463 Statistical significance was calculated using an unpaired two-tailed *t*-test or ANOVA. n.s.,
- 464 not significant.

465 Figure 3. Caprolactam polymer-based 3a media supports efficient human HSC 466 expansion ex vivo

467 (a) Total cell numbers generated after 7-days culture of 2x10⁴ human CB CD34⁺ cells in
468 3a media containing various synthetic polymers (see in *Methods* for details). No polymer
469 (none) was used as a negative control. Mean + S.D. of three independent cultures. n.d.,

- 470 not detected.
- 471 (b) Fold change in total and CD34⁺ cell numbers during a 30-day culture of $2x10^4$ CB
- 472 CD34⁺ in 3a media containing PVA and/or polyvinyl caprolactam-polyvinyl acetate-
- 473 polyethylene glycol graft copolymer (PCL-PVAc-PEG). Mean of three independent
- 474 cultures. *P = 0.0170; ***†P = 0.0190; ***‡P = 0.0012; ***§P = 0.0013; ***|P = 0.0023;
- 475 ***P = 0.0006.
- 476 (c) Mean human CD45⁺ PB chimerism in recipient NOG mice following transplantation

477 of 1×10^4 day-30 cells derived from CB CD34⁺ cell cultured in 3a media containing PVA

478 or PCL-PVAc-PEG (cultures initiated with 1×10^4 cells). n=5 per group. Results from

- 479 replicate experiments shown in Supplementary Table 1. Independent experiments 480 performed with 2-3 human CB samples per experiment. *P = 0.0424.
- (d, e) Mean 16-week human CD45⁺ and CD34⁺ cell chimerism in the BM and spleen
 from mice described in (c). n=5 per group.
- 483 (f, g) Mean human CD45⁺ PB, BM and spleen chimerism in secondary recipient NOG

484 mice following transplantation of 1×10^6 cells derived from primary recipients, as describe

- 485 in (d, e). n=5 per group. Results from replicate experiments shown in Supplementary
- 486 **Table 2**.
- 487 (h) Human PB chimerism and phenotypes in humanized IL-3/GM-CSF-transgenic NOG
- 488 mice at 16-weeks following transplantation of 1×10^4 day-20 cells derived from CB CD34⁺
- 489 cell cultured in 3a media containing PCL-PVAc-PEG (cultures initiated with 1×10^4 cells).
- 490 n=5 per group. ****P < 0.0001.
- 491 (i) Frequency for each human CD45⁺ cell subpopulation in the PB from mice described
- 492 in (h). Mean \pm S.D. of five mice per group.
- 493 Statistical significance was calculated using an unpaired two-tailed *t*-test: n.s., not494 significant.
- 495

496 Figure 4. Long-term selective expansion of functional human HSCs in 3a media

(a) Log₂-fold expression change of indicated HSC-associated genes. Mean ± S.D. of
three independent cultures. EPCR⁺ and EPCR⁻ indicates CD34^{high}EPCR⁺ cells and
CD34^{high}EPCR⁻ cells, respectively. **[†]P = 0.0049; **[‡]P = 0.0043; **[§]P = 0.0022; ***P
= 0.0006.

(b, c) Results from gene set enrichment analysis (GSEA) for genes differentially
expressed between CD34^{high}EPCR⁺ and CD34^{high}EPCR⁻ samples using gene sets for HSC
genes (b) and mitochondrial oxidative phosphorylation-related genes (c). Statistical
significance was calculated using an empirical phenotype-based permutation test
procedure³⁸.

(d) UMAP plot of single-cell RNA sequencing data from 10-day expanded CD34⁺ CB
cells with 12 cell clusters annotated (see in *Methods* for details). Integrated cell map from
cells cultured in PCL-PVAc-PEG based 3a media, StemSpan with SR-1 media, or
StemSpan with UM171 media. Statistical significance was calculated using an empirical
phenotype-based permutation test procedure³⁸.

(e) Cell distribution within PCL-PVAc-PEG based 3a cultures, StemSpan with SR-1
cultures, and StemSpan with UM171 cultures. Black dotted frames indicate the HSPCHLF cell cluster. See Extended Data Figure 6g for a quantification of *HLF* expression.
(f) Schematic of the single HSC expansion assay.

515 (g) Mean number of cells derived from single human CD34⁺CD38⁻CD90⁺CD45RA⁻

516 CD49f⁺ CB cell after 7-days culture (n=96). Results from replicate experiments shown in

517 Supplementary Table 3. Independent experiments performed with 2-3 human CB518 samples per experiment.

- (h, i) Mean human CD45⁺ PB and BM chimerism in recipient NOD.Cg-Prkdc^{scid}
 Il2rg^{tm1Sug} Kit^{em1(V831M)Jic}/Jic (W41/W41) mice following transplantation (n=10), as
 described in (g). Results from replicate experiments shown in Supplementary Table 4.
 Statistical significance was calculated using an unpaired two-tailed *t*-test.
- 523

524 Methods:

Mice. C57BL/6 mice were purchased from Sankyo Lab Service (Tsukuba, Japan) or bred
in-house. Immunodeficient NOD/Shi-scid, IL-2Rγ^{null} (NOG), NOD.Cg-Prkdc^{scid}
Il2rg^{tm1Sug} Kit^{em1(V831M)Jic}/Jic (W41/W41), and human IL-3/GM-CSF-transgenic NOG
(NOG IL-3/GM-Tg) mice³⁰ were purchased from the Central Institute for Experimental
Animals (Kanagawa, Japan).

530 NOG and W41/W41 mice were developed at the Central Institute for 531 Experimental Animals. Kit-mutated NOG-W41 mice were established by genome editing 532 using transcription activator-like effector nucleases (TALENs). Designed TALEN 533 mRNA pairs (Forward; 5'-gtgttccgttctaggcac-3', and Reverse; 5'-atgctctctggtgccatc-3') 534 and 100-bp single-strand oligonucleotide (ssOligo) containing a G to A point mutation in the kinase domain of the c-Kit locus³⁹ were purchased from Thermo Fisher Scientific 535 536 (Waltham, MA, USA). TALEN mRNA (4 ng/μ) and ssOligo (15 ng/μ) were mixed and 537 injected into NOG mouse embryo to generate NOG-W41 mice. All mice were housed in 538 specific-pathogen-free conditions with free access to food and water. All animal 539 experiments were performed in accordance with institutional guidelines and were 540 approved by the Animal Care and Use Committee of the Institute of Medical Science, 541 The University of Tokyo, the Laboratory Animal Resource Center, University of Tsukuba 542 and the Institutional Animal Care and Use Committee of Central Institute for 543 Experimental Animals.

544

545 Isolation of mouse cells. Bone marrow (BM) c-Kit⁺Sca-1⁺Lineage⁻ (KSL) cells were 546 isolated from 8- to 12-week-old mice. Whole BM cells were stained with APC-547 conjugated anti-c-Kit antibody (eBioscience, San Diego, CA, USA) and c-Kit⁺ cells 548 enriched using anti-APC magnetic beads and LS columns (Miltenyi Biotec). The c-Kit-549 enriched cells were then stained with PE-conjugated anti-Sca-1 (eBioscience), and a 550 lineage antibody cocktail (biotinylated CD4, 1:200, CD8, 1:200, CD45R, 1:200, 551 TER119, 1:100, LY-6G/LY-6C 1:200, and CD127; 1:100, all from eBioscience), 552 followed by staining with FITC-CD34 and streptavidin-APC-eFluor 780 (eBioscience, 553 1:200). Cell populations were purified and sorted by FACS AriaII (BD Biosciences, Franklin Lakes, NJ, USA) with BD FACS Diva software using propidium iodide as a
dead cell stain. Antibodies are described in Supplementary Table 5.

556

Human umbilical cord blood cells. Human umbilical cord blood-derived (CB) CD34⁺ 557 558 cells were purchased from StemExpress (Folsom, CA, USA). CD34⁺CD38⁻ cells were 559 purified by staining thawed CD34⁺ cells with PE-Cy7-labeled anti-human CD34 (BD 560 Biosciences, 1:100) and V450-labeled anti-human CD38 (BD Biosciences, 1:100), then 561 sorted as described above. For detailed phenotypic HSC analysis, cells were stained with 562 PerCP-Cy5.5-labeled anti-human CD34 (BioLegend, San Diego, CA, USA, 1:100), 563 BV421-labeled anti-human CD38 (BD Biosciences, 1:100), PE-Cy7-labeled anti-human 564 CD90 (BD Biosciences, 1:100), APC-H7-labeled anti-human CD45RA (BD Biosciences, 565 1:100) and PE-labeled anti-human CD49f (BD Biosciences, 1:100), then sorted as 566 described above. Antibodies are described in Supplementary Table 5. For all 567 experiments, different lots of CBs were used. For the comparison of fresh and expanded 568 cells, a common lot of cord blood was used.

569

570 PVA and cytokine-based cell cultures. Human CB CD34⁺ cells cultures were performed 571 using IMDM (Life Technologies, Carlsbad, CA, USA), 1% insulin-transferrin-selenium-572 ethanolamine (ITSX; Life Technologies), 1% penicillin/streptomycin/glutamine (P/S/G; 573 Life Technologies), 0.1% Good Manufacturing Practice Grade polyvinyl alcohol (PVA; 574 Japan VAM&POVAL CO., LTD, Osaka, Japan), 10 ng/ml recombinant human SCF (PeproTech, Rocky Hill, NJ, USA) and 100 ng/ml recombinant human THPO 575 (PeproTech), at 37°C with 5% CO₂. Cultures were supplemented with 740Y-P (CAS No. 576 577 236188-16-1; synthesized) and SC79 (CAS No. 305834-79-1; Sigma-Aldrich, St. Louis, 578 MO, USA), as indicated. Mouse cell cultures were performed using F12 media (Life 579 Technologies), 1% ITSX, 1% P/S/G, 10 mM HEPES (Life Technologies), 0.1% PVA, 10 580 ng/ml recombinant mouse SCF (PeproTech) and 100 ng/ml recombinant mouse THPO 581 (PeproTech), at 37 °C with 5% CO₂. U-bottomed 96-well tissue culture plates were used 582 in mouse and human comparative experiments. All other cultures were performed using 583 24-well flat-bottomed CellBIND® tissue culture plates (Corning, Corning, NY, USA; 584 Product Number 3337).

586 Signaling analysis. Phosphorylation status of signaling molecules was analyzed by 587 fluorescent immunocytostaining. At indicated timepoints, cells were attached to poly-l-588 lysine-coated slides (Matsunami Glass, Osaka, Japan), then fixed with 4% 589 paraformaldehyde and permeabilized with 0.1% Triton X-100. The cells were stained with phosphorylation-specific anti-PI3K, anti-Stat5, anti-AKT, anti-JAK2, anti-Stat3, 590 591 anti-p38MAPK, and anti-p44/42MAPK antibodies (all from Thermo Fisher Scientific, 592 Waltham, MA, USA). After washing with PBS, cells were stained with Alexa Fluor 488-593 conjugated goat anti-rabbit IgG antibody (CAS No. A11008, Invitrogen) and DAPI. 594 Immunofluorescence images were obtained and analyzed using a Cellomics ArrayScan 595 VTI HCS Reader (Thermo Scientific) as described previously^{18,40}. All experiments were 596 performed using mixture of five cord blood samples.

597

598 PVA-based cytokine-free cultures. Human CB CD34⁺ cell cultures were performed using 599 IMDM, 1% ITSX, 1% P/S/G, 0.1% PVA, 740Y-P and butyzamide (Shionogi, Osaka, 600 Japan) at 37 °C with 5% CO₂. All long-term cultures used 1 µM 740Y-P and 0.1 µM 601 butyzamide, with media changes made every 3 days by manually removing conditioned 602 media by pipetting and replacing pre-warmed and freshly prepared media. Butyzamide is a THPO receptor agonist^{41,42} and has also been used clinically as a lusutrombopag. All 603 604 cell cultures were performed using 24-well flat-bottomed CellBIND® tissue culture 605 plates. Where indicated, cultures were supplemented with 750 nM StemRegenin 1 (SR-606 1; CAS No. 1227633-49-9) and/or 70 nM UM171 (CAS No. 1448724-09-1). As described in the main text, we defined media containing 1 µM 740Y-P, 0.1 µM 607 608 butyzamide and 70 nM UM171 as 3a media.

609

610 *PCL-PVAc-PEG-based cytokine-free cultures.* A step-by-step protocol describing the
611 culture of human CB CD34⁺ cells with PCL-PVAc-PEG-based cytokine-free media can
612 be found at Protocol Exchange⁴³. Human CB CD34⁺ cell cultures were performed using
613 IMDM, 1% ITSX, 1% P/S/G, 1 μM 740Y-P, 0.1 μM butyzamide and 0.1% of polyvinyl
614 caprolactam-polyvinyl acetate-polyethylene glycol graft copolymer (PCL-PVAc-PEG;
615 Soluplus®; BASF, Ludwigshafen am Rhein, Germany) at 37 °C with 5% CO₂. For long-

585

term cultures, media changes made every 3 days by manually removing conditioned 616 617 media by pipetting and replacing pre-warmed and freshly prepared media. For polymer 618 screening, human CB CD34⁺ cells were cultured with IMDM, 1% ITSX, 1% P/S/G, 1 619 μ M 740Y-P, 0.1 μ M butyzamide and 0.1% one of the following chemicals: PVA, 188 620 BIO (Kolliphor® P 188 Bio; BASF), 188 Geismar (Kolliphor® P188; BASF), PCL-621 PVAc-PEG, 407 Geismar (Kolliphor® P407; BASF), 30 Geismar (Kollidon® 30, BASF), 622 17 PF (Kollidon® 17 PF; BASF), 90 F (Kollidon® 90 F; BASF) or 12 PF (Kollidon® 12 623 PF; BASF) at 37 °C with 5% CO₂. In addition, PCL-PVAc-PEG-based cytokine cocktails 624 media consisted of 10 ng/ml recombinant mouse SCF (PeproTech) and 100 ng/ml 625 recombinant mouse THPO (PeproTech) (described in Extended Data Figure 4a).

626

627 UM171 and/or SR-1-based cultures. Human CB CD34⁺ cell cultures for 628 xenotransplantation assays were performed using StemSpan SFEM (Stem Cell 629 Technologies, Vancouver, BC, Canada) supplemented with 100 ng/ml recombinant 630 human SCF (PeproTech), 100 ng/ml FMS-like tyrosine kinase 3 ligand (FLT3, PeproTech), 50 ng/ml recombinant human THPO (PeproTech), 10 µg/ml lipoproteins 631 632 (Stem Cell Technologies) and 35nM UM171 and/or 750nM SR-1 at 37°C with 5% CO₂ (Extended Data Figure 4f-i)¹⁰. In comparison experiments with previously protocols, 633 634 human CB CD34⁺ cell cultures were performed using StemSpan SFEM supplemented 635 with 50 ng/ml recombinant human SCF (PeproTech), 50 ng/ml FLT3-L, 50 ng/ml 636 recombinant human THPO, 50 ng/ml recombinant human IL-6 (PeproTech) and 750nM 637 SR-1 at 37 °C with 5% CO₂ (described in Figure 4e, Extended Data Figure 4a-e, Extended Data Figure 5a, Extended Data Figure 6g)¹¹. 638

639

Analysis of cell cultures. Cultured cells were counted using a hemocytometer or a
CYTORECON cytometer (GE Healthcare, Amersham, UK) before and after culture.
Phenotypic analysis was performed by staining cells with PE-Cy7-labeled anti-human
CD34 (1:100), V450-labeled anti-human CD38 and FITC-labeled anti-human CD41
(BioLegend, 1:100), followed by flow cytometric analysis using a FACS AriaII or FACS
Verse (BD Biosciences) with BD FACS Diva software using propidium iodide as a dead
stain. For detailed phenotypic HSC analysis, fresh or cultured cells were stained with

647 PerCP-Cy5.5-labeled anti-human CD34 (BioLegend, 1:100), BV421-labeled anti-human 648 CD38 (BD Biosciences, 1:100), PE-Cy7-labeled anti-human CD90 (BD Biosciences, 649 1:100), APC-H7-labeled anti-human CD45RA (BD Biosciences, 1:100), PE-labeled anti-650 human CD49f (BD Biosciences, 1:100), FITC-labeled anti-human lineage cocktail (CD2, 651 CD3, CD4, CD7, CD8, CD10, CD11b, CD14, CD19, CD20, CD56, CD235a) (BD 652 Biosciences, 1:200) and FITC-labeled (BioLegend, 1:200) or BV711-labeled (BD 653 Biosciences, 1:100) anti-human CD41, followed by flow cytometric analysis using a 654 FACS AriaIII (BD Biosciences) with BD FACS Diva software using propidium iodide 655 as a dead stain. As a set of markers containing EPCR, cultured cells were stained with 656 APC-labeled anti-human CD34 (BioLegend, 1:100), BV421-labeled anti-human CD90 657 (BioLegend, 1:100), PerCP-Cy5.5-labeled anti-human CD45RA (BioLegend, 1:20), PE-658 labeled anti-human CD49c (ITGA3) (BD Biosciences, 1:200), BV605-labeled anti-659 human CD201 (EPCR) (BD Biosciences, 1:100) and FITC-labeled anti-human lineage 660 cocktail. Antibodies are described in Supplementary Table 5. Results were analyzed 661 with FlowJo software 10.8.1 (Tree Star, Ashland, OR).

662

663 Colony forming unit (CFU) assays. Defined numbers of fresh or cultured cells were
664 sorted by FACS AriaII and subjected to CFU assays using Methocult H4435 (Stem Cell
665 Technologies). Cells were incubated in a humidified atmosphere at 37 °C with 5% CO₂.
666 After two weeks, the number of colonies were counted and types of colonies were
667 validated by cytospin smears stained with Hemacolor (Merck, Darmstadt, Germany).

668

Xenotransplantation assays. Fresh or cultured human CB CD34⁺ cells were transplanted 669 670 by tail artery injection⁴⁴ into sub-lethally (1.5 Gy) irradiated 8-10-week-old immunodeficient NOD/Shi-scid IL-2Rynull (NOG) mice or NOG IL-3/GM-Tg mice. 671 672 Human cell chimerism in the peripheral blood analyzed using V450–labeled anti-mouse 673 CD45.1 (BD Biosciences, 1:200) and APC-Cy7-labeled anti-human CD45 antibodies 674 (BioLegend, 1:100) following red blood cell lysis. Mice were randomly selected and BM 675 and spleen analysis was performed. Human CD34⁺ cell chimerism in the BM and spleen 676 was determined using V450-labeled anti-mouse CD45.1 (1:200), APC-Cy7-labeled anti-677 human CD45 antibodies (1:100), and PE-labeled anti-human CD34 (BioLegend, 1:50). 678 For detailed phenotypic HSC analysis, cells were stained with PerCP-Cy5.5-labeled anti-679 human CD34 (BioLegend, 1:100), BV421-labeled anti-human CD38 (BD Biosciences, 680 1:100), PE-Cy7-labeled anti-human CD90 (BD Biosciences, 1:100), APC-H7-labeled 681 anti-human CD45RA (BD Biosciences, 1:100) and PE-labeled anti-human CD49f (BD 682 Biosciences, 1:100). Human lineage chimerism in the BM and spleen was determined 683 using V450–labeled anti-mouse CD45.1, APC-Cy7–labeled anti-human CD45 antibodies, 684 PE-Cy7-labeled anti-human CD33 (eBioscience, 1:20), PE-labeled anti-human CD3 685 (eBioscience, 1:20), and APC-labeled anti-human CD19 (eBioscience, 1:100). We 686 defined CD33⁺ cells as myeloid cells, CD19⁺ cells as B cells, and CD3⁺ cells as T cells. 687 In xenotransplantation assays using NOG IL-3/GM-Tg mice (described in Figure 3h, i), 688 we used PE-labeled anti-human CD56 (BioLegend, 1:100) and FITC-labeled anti-human 689 CD66b (BioLegend, 1:100) additionally. Flow cytometry analysis was then performed 690 using a FACS AriaII or FACS Verse (BD Biosciences) with propidium iodide as dead 691 stain, and results were analyzed with FlowJo software. For secondary transplantation 692 assays, we collected and pooled bone marrow from all primary recipient mice at 16 weeks 693 and transplanted 1x10⁶ cells into each sub-lethally-irradiated NOG mice as described 694 above, with donor chimerism analyzed as above. In xenotransplantation assays using 695 W41/W41 and NOG mice (described in **Extended Data Figure 8**), 5x10⁴ fresh human 696 CB CD34⁺ cells were transplanted into W41/W41, W41/+ or +/+ mice with or without 697 irradiation, with donor chimerism analyzed as above.

698

699 Clonal HSC expansion assays. Single human CB-derived CD34⁺CD38⁻CD90⁺CD45RA⁻ 700 CD49f⁺ cells were purified by staining thawed PerCP-Cy5.5-labeled anti-human CD34 701 (BioLegend, 1:100), BV421-labeled anti-human CD38 (BD Biosciences, 1:100), PE-702 Cy7-labeled anti-human CD90 (BD Biosciences, 1:100), APC-H7-labeled anti-human 703 CD45RA (BD Biosciences, 1:100) and PE-labeled anti-human CD49f (BD Biosciences, 704 1:100) then sorted single cell into a 96-well flat-bottomed CellBIND® tissue culture plate. 705 After culturing with PCL-PVAc-PEG-based 3a media for 7 days as described above, the 706 top 10 wells with high expansion efficiency were transplanted into sub-lethally (1 Gy) 707 irradiated 8-10-week-old W41/W41 mice by tail artery injection (detailed in Figure 4f). 708 For split clone transplantation assays, after culturing with PCL-PVAc-PEG-based 3a

709 media for 7 days as described above, each of the 6 wells with highest expansion efficiency 710 were transplanted into three sub-lethally (1 Gy) irradiated 8-10-week-old W41/W41 mice

- 711 by tail artery injection (detailed in Extended Data Figure 7a, b).
- 712

713 Thrombopoietin receptor agonist screening. MPL-expressing 32D cells culture were 714 performed using RPMI media containing 1% ITSX, 1% P/S/G, and 0.1% BSA or PVA at 715 37 °C with 5% CO₂. One of the following thrombopoietin receptor agonists was added to 716 each culture: 7 µM eltrombopag (Cayman Chemical Company, Ann Arbor, MI, USA); 3 717 µM avatrombopag (MedChemExpress, Monmouth Junction, NJ, USA); or 0.1 µM 718 butyzamide⁴¹. Human CB CD34⁺ cell cultures were performed using IMDM, 1% ITSX, 719 1% P/S/G, 0.1% PVA, 740Y-P, and one of the following thrombopoietin receptor agonist: 720 7 μ M eltrombopag, 3 μ M avatrombopag, or 0.1 μ M butyzamide at 37 °C with 5% CO₂.

721

722 Preparation and culture of human peripheral blood stem cells. Fresh human peripheral 723 blood stem cells (PBSCs) were obtained from healthy adult donors for allogeneic 724 transplantation in the University of Tsukuba Hospital (approval R02-009). The donors 725 received the treatment of G-CSF before leukapheresis. All donors agreed to experimental 726 use of their PBSCs after informed consent and our study was approved by the ethical 727 committee in the University of Tsukuba. From PBSCs, mononuclear cells (MCs) were 728 separated by Lymphocytes Separation Medium 1077 (PromoCell, CAS No. C-44010). 729 After separation, CD34⁺ cells were enriched using the Human CD34 Microbeads Kit 730 (Miltenyi Biotec Inc., CAS No. 130-046-702) and MACS LS columns (Miltenyi Biotec 731 Inc., CAS No. 130-042-401). Purified CD34⁺ cells were cultured in PVA- and PCL-732 PVAc-PEG based 2a or 3a media. In 3a media cultures, UM729 (1 μM) was used in place 733 of UM171 because UM171 was not commercially available at the time this experiment 734 was performed. All experiments complied with all relevant guidelines and regulations. 735

736 *Exome Sequencing.* We extracted genomic DNA of fresh and 10-day cultured human 737 CB CD34⁺ cells using QIAamp DNA Blood Mini Kit (QIAGEN, CAS No. 51106). After 738 DNA fragmentation, target enrichment by hybrid capture probes was performed using 739 SureSelect Human All Exon V6 (Agilent). Enriched DNA was sequenced on NovaSeq 6000 (Macrogen Inc, Korea). FASTQ files were imported to CLC Genomics Workbench
(ver. 10.1.1) for subsequent analysis. Sequence data was annotated using the reference
genome (GRCh38). After filtering out common variants using the database of Tohoku
Medical Megabank Organization (https://www.megabank.tohoku.ac.jp), we annotated
mutations unique to the culture sample that were located in amino-acid change sites or
splicing sites.

746

747 Bulk RNA sequencing. Human CB CD34⁺ cells were cultured in PCL-PVAc-PEG based 748 3a media at 37°C with 5% CO₂ for 10-days. CD34^{high}EPCR⁺ and CD34^{high}EPCR⁻ cells 749 were then sorted by MoFlo XDP (Beckman Coulter) and processed in TRIZOL-LS 750 (Thermo Fisher Scientific, 10296028). Total RNA was used for rRNA-depletion by 751 NEBNExt rRNA Depletion Kit (New England Biolabs, CAS No. E6310), and next 752 directional library synthesis by NEBNext Ultra Directional RNA Library Prep Kit for 753 Illumina (New England Biolabs, CAS No. E7420). Libraries were sequenced on Illumina 754 NextSeq 5000. We analyzed the data using the edgeR v 3.14^{45} in R (4.1.1). Volcano plots 755 were generated using EnhancedVolcano 756 (https://github.com/kevinblighe/EnhancedVolcano) in R and genes highlighted when the 757 value of \log_2 fold change was >2 and $-\log_{10}P$ was >14. Gene Ontology enrichment 758 analysis was performed by ClusterProfiler⁴⁶. Gene set enrichment analysis (GSEA) 759 software (http://www.gsea-msigdb.org/gsea/index.jsp) was used for comparing our 760 datasets with two previously published datasets.

761

762 Single-cell RNA sequencing. Human CB CD34⁺ cells were cultured for 10-days at 37°C 763 with 5% CO₂ under three different conditions as follows: (1) PCL-PVAc-PEG based 3a 764 media (composition described above); (2) StemSpan SFEM (Stem Cell Technologies, 765 Vancouver, BC, Canada) supplemented with 100 ng/ml recombinant human SCF 766 (PeproTech), 100 ng/ml FMS-like tyrosine kinase 3 ligand (FLT3, PeproTech), 50 ng/ml 767 recombinant human THPO (PeproTech), 10 µg/ml lipoproteins (Stem Cell Technologies) and 35 nM UM171²; and (3) StemSpan SFEM supplemented with 50 ng/ml recombinant 768 human SCF (PeproTech), 50 ng/ml FLT3-L, 50 ng/ml recombinant human THPO, 50 769 ng/ml recombinant human IL-6 (PeproTech) and 750 nM SR-1¹¹. From each CB cell 770

culture, the propidium iodide-negative fraction was sorted by MoFlo (Beckman Coulter)
and single cell Gel Beads-in-Emulsions were generated using the Chromium Controller
(10x Genomics). Libraries were generated using the Single Cell 3' Reagent Kit version
3.1 (10x Genomics) according to manufacturer's instructions.

775 Cells were sequenced on Illumina Hiseq X (Macrogen Inc, Korea). Sequence data 776 was annotated by the reference genome (GRCh38) using the Cellranger v6.1.1 pipeline. 777 Subsequent analysis was performed using Seurat v4.047 in R. Using the Read10X 778 function, we obtained the unique molecular identified (UMI) count matrix of each dataset. 779 This analysis identified 9913 cells for the PCL-PVAc-PEG sample, 5912 cells for the 780 UM171 sample, and 9198 cells for the SR-1 sample. The mean reads per cell was 32205 781 for the PCL-PVAc-PEG sample, 36026 for the UM171 sample, and 30991 for the SR-1. 782 The median number of genes detected per cell was 3292 genes for the PCL-PVAc-PEG 783 sample, 3648 genes for the UM171 sample, and 3300 genes for the SR-1 sample. We 784 filtered out cells that had unique feature counts of over 7500 or less than 200, and cells 785 with >10% mitochondrial counts. The filtered cell count number (cells used for 786 subsequent analysis) was 9572 cells for the PCL-PVAc-PEG sample, 5373 cells for the 787 UM171 sample, and 6991 cells for the SR-1 sample, with 22179 features, 21222 features, 788 and 21645 features detected, respectively.

789 Normalization and scaling were performed with the SCTransform function 790 (method = "glmGamPoi"). At this time, the effect of the mitochondrial gene expression 791 ratio was removed (var.to.regress = "percent.mt"). The SelectIntegrationFeatures 792 (nfeatures = 3000) function was used to select genes for integration of the datasets. After 793 processing the datasets with the PrepSCTIntegration function, FindIntegrationAnchors 794 and IntegrateData functions were used to find anchors for integration and integrate the 795 datasets. To correct for cell-to-cell variation due to the effects of cell cycle, cell cycle 796 scoring and regression was performed with CellCycleScoring function, using a published 797 mouse hematopoietic stem cells dataset⁴⁷. Scaling was then performed with the ScaleData 798 (vars.to.regress = c("S.Score", "G2M.Score")) function. Principal components analysis 799 (PCA) was performed using the RunPCA function (npc = 30). Uniform Manifold 800 Approximation and Projection (UMAP) was performed using the RunUMAP function to 801 reduce the dimension of the dataset of embedded cells into two dimensions.

802 FindNeighbors function was used to determine k-nearest neighbors (KNN) for each cell, 803 and the KNN graph was constructed based on Euclidean distance. Finally, processed cells 804 were clustered based on KNN using the Louvain algorithm (resolution = 0.4) by the 805 FindCluster function. FindMarkers and FindAllMarkers functions were used to identify 806 intercluster differentially expressed genes and select feature genes that characterized 807 specific hematopoietic cell types. This allowed for the follow clusters to be manually 808 annotation: hematopoietic stem/progenitor cells highly expressing HLF (HSPC-HLF), 809 hematopoietic stem/progenitor cells (HSPC), cell-cycle activated hematopoietic 810 stem/progenitor cells (HSPC-Cycling), granulocyte-monocyte progenitor cells (GMP), 811 monocyte progenitor cells (MP), granulocyte progenitor cells (GP), dendritic cell 812 progenitors (DCP), CD34 and GATA2-expressing progenitor cells (CD34⁺GATA2⁺ prog), 813 megakaryocyte and erythroid progenitor cells (MEP), erythroid progenitor cells (EryP), 814 megakaryocyte progenitor cells (MgkP), and mast cell progenitors (MCP).

815 Subsequent analysis was performed using Seurat v4.0⁴⁸ in R. Fresh CB data for the comparison of *HLF* expression in cultured cells (Extended Data Figure 6g) was 816 obtained from GEO (GSE 153370)³². In addition, we compared our datasets with GEO-817 818 deposited scRNAseq data from 7-day cultured CB CB34⁺ cells in StemSpan SFEM + UM171 (GSE 153370). We filtered out cells (with feature counts over 7500 or less than 819 820 200, and those with >10% mitochondrial counts) and used NormalizedData, 821 CellCycleScoring and ScaleData functions to process data. Next, FindTransferAnchors 822 was performed to find anchors between our datasets (as reference data) and the published 823 datasets (as query data). After the RunUMAP function (reduction.model = TRUE) was 824 applied to the reference data, MapQuery function was used to perform Unimodal UMAP 825 Projection (Extended Data Figure 6f).

826

Cell Cycle analysis. Cultured human CB CD34⁺ cells were stained with APC-labeled
anti-human CD34 (BioLegend), PerCP-Cy5.5-labeled anti-human CD45RA (BioLegend),
then washed with phosphate-buffered saline (PBS) twice and pelleted. BD
Cytofix/Cytoperm Fixation/Permeabilization Kit (BD Biosciences, 554714) was then
used to process the samples according to manufacturer's instructions. After fixation and
permeabilization, the cells were stained with FITC-labeled anti-human Ki67 (BioLegend,

1:100) and DAPI (DOJINDO, 1:1000). FITC-labeled IgG2b kappa (BioLegend, 1:100)
was used as an isotype control. Antibodies are described in Supplementary Table 5.
Analysis was performed on a LSR Fortessa Cell Analyzer (BD Bioscience). Data was
analyzed with FlowJo software.

837

838 Apoptosis assay. Suspension of cultured human CB CD34⁺ cells were centrifuged and 839 washed by PBS. Next, annexin binding buffer (10mM HEPES, 140mM NaCl and 2.5mM 840 CaCl₂ diluted in distilled water) was added to the sample. After that, the cells were stained 841 AlexaFluor488 AnnexinV (Invitrogen, 1:40) and propidium by iodide 842 (BioLegend1:1000) and incubated for 15 minutes at room temperature. Antibodies are 843 described in **Supplementary Table 5.** Finally, we resuspended the samples in the annexin 844 binding buffer and analyzed them by LSR Fortessa Cell Analyzer (BD Biosciences).

845

Reactive Oxygen Species (ROS) Assay. Fresh and cultured cells were pre-stained with
APC-labeled anti-human CD34 (BioLegend, 343510). The cells were then processed with
a ROS Assay Kit-Photo-oxidation Resistant DCFH-DA (DOJINDO, R253) according to
the manufacturer's instructions. Cell samples were incubated in the Working Buffer for
30 minutes at 37°C with 5% CO₂ and then washed with HBSS twice. Analysis was
performed on a Attune NxT Flow Cytometer (Invitrogen). Data was analyzed with
FlowJo software.

853

γH2A.X Assay. Fresh and cultured cells were pre-stained with APC-labeled anti-human
CD34 (BioLegend, 343510). The cells were then processed with BD Cytofix/Cytoperm
Fixation/Permeabilization Kit (BD Biosciences, 554714) according to the manufacturer's
instructions and intracellularly stained with FITC-labeled anti-H2A.X Phospho (Ser139)
antibody (BioLegend, 613404, 1:100). Analysis was performed on a Attune NxT Flow
Cytometer (Invitrogen). Data was analyzed with FlowJo software.

861 *Statistical analysis.* Statistical analysis was performed using two-tailed t-testing or
862 ANOVA in Prism 9 software (GraphPad, San Diego, CA, USA).

Data Availability Statement: Exome sequencing data available on BioProject
(PRJNA786760). All RNA-seq data were deposited in the Gene Expression Omnibus
under accessions GSE191338 (bulk) and GSE192519 (single cell). Source data are
provided with this paper.

867

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910 Author contributions: M.S., K.I. and S.Y. conceived, designed and performed 911 experiments, analyzed data and wrote the paper. R.I., T.K., E.M., H.N., K.S., H.J.B. and 912 H.T. designed and performed experiments. M.S. performed cell cultures, colony-forming 913 unit assays and FACS analysis, and xenotransplantation assays. K.I. performed cell 914 cultures, FACS analysis, exome sequencing, RNA sequencing, cell cycle analysis, 915 apoptosis assays, ROS assays, and yH2A.X assays, S.Y performed cell cultures, signaling 916 analysis, xenotransplantation assays, and clonal HSC expansion assays, R.I. performed 917 xenotransplantation assays, T.K. performed exome sequencing and RNA sequencing, 918 E.M., H.N., K.S. H.J.B. and H.T. helped with cell cultures and FACS analysis. S.Y. 919 performed independent replications of the experiments (Supplementary Table 1-4) both 920 in the University of Tokyo, University of Tsukuba, and R.I. performed independent 921 replications of the experiment (Supplementary Table 1-2) in Central Institute for 922 Experimental Animals. A.C.W. and D.G.K. analyzed data and wrote the paper. T.S. 923 provided reagents and discussed the results. K.K., S.T., Y.N., A.I., S.C. and S.O. 924 discussed the results and wrote the paper. H.N. guided and supervised the project. All 925 authors edited and approved the paper. 926

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929	and Celaid Therapeutics. All other authors declare no competing interests.
930	
931	Supplementary Information: This file contains Supplementary Tables 1-5.
932	
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935	
936	Extended data figures and tables: 8 Extended Data Figures and 2 Extended Data Tables
937	available in the online version of the paper.
938	

939 Extended Data Figure Legends:

940 *Extended Data Figure 1: Development of chemically-defined cytokine-free culture*941 *media for human hematopoietic stem/progenitor cells (HSPCs)*

- 942 (a) Single cell phosphorylation status of JAK2, STAT3, STAT5, p38 MAPK, and
- 943 p44/42 MAPK in mouse KSL and human CB CD34⁺CD38⁻ cells cultured with 10 ng/ml
- 944 SCF and 100 ng/ml THPO in PVA-based media. Mean of 30 cells. AFI: average
- 945 fluorescence intensity. ****P < 0.0001.
- 946 (b) Representative image of p-PI3K after 7-days culture cells in mouse and human, as
 947 described in (a). Blue: DAPI, Green: anti-PI3K. Scale bar: 100 μm.
- 948 (c) Single cell phosphorylation status of PI3K in mouse CD34-KSL and human
- 949 CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ CB cells cultured with 10 ng/ml SCF and 100
- 950 ng/ml THPO in PVA-based media for 3 and 7 days. Mean of 31 cells. AFI: average
- 951 fluorescence intensity. ****P < 0.0001.
- 952 (d) CD34⁺CD45RA⁻ cell numbers of human-cord-blood-derived CD34⁺CD38⁻ cultured
- 953 with SC79 or 740Y-P in addition to human 10 ng/ml SCF (S) and 100 ng/ml THPO (T)
- 954 in PVA culture conditions for 7 days. The starting cell count was $2x10^4$. Mean of three

955 independent cultures. **P = 0.0040.

- 956 (e) Single cell phosphorylation status of PI3K in human CD34⁺CD38⁻CD90⁺CD45RA⁻
- 957 CD49f⁺ CB cells cultured in PVA-based media containing 10 ng/ml SCF and 100 ng/ml
- **958** THPO with or without 740Y-P for 7 days. Mean of 31 cells. AFI: average fluorescence **959** intensity. ***P = 0.0002.
- 960 (f) Cell cycle analysis of CD34⁺CD45RA⁻ cells after a 7-day culture of human CB CD34⁺
- 961 cells in PVA-based media containing 740Y-P and 100 ng/ml THPO (T) with or without
- 962 10 ng/ml SCF (S). Mean of three independent cultures. Representative FACS plot was963 shown on the right.
- 964 (g) Fold change in GEmM colony numbers generated from human CB CD34⁺ cells after
- 965 a 7-day culture in PVA-based media supplemented with 740Y-P and THPO (T) with or
- 966 without SCF (S), relative to fresh CD34⁺ cells. Mean of three independent cultures.
- 967 (h) Total cell numbers after 1×10^3 MPL-expressing 32D cells (32D/MPL) were cultured
- 968 for 3-days with various THPO agonists (eltrombopag, avatrombopag or butyzamide) in
- 969 BSA-based or PVA-based media. Mean of two independent cultures. n.d., not detected.

970 (i) Fold change in total and CD34⁺ cell numbers after a 7-day culture of $2x10^4$ CD34⁺

- 971 cells in PVA-based media supplemented with 740Y-P and various THPO agonists
- 972 (eltrombopag, avatrombopag or butyzamide). Mean of three independent cultures. n.d.,973 not detected.
- 974 (j) $CD34^+$ CD41⁻ CD90⁺CD45RA⁻ cell numbers after a 7-day culture of $2x10^4$ of human
- 975 CB CD34⁺ cells in PVA-based media containing 740Y-P and 100 ng/ml THPO (T) and/or
- 976 butyzamide (Buty). Mean of three independent cultures. **P = 0.0020; ***P = 0.0010.
- 977 (k) Fold change in GEmM colony numbers generated from CD34⁺ cells after a 7-day
- 978 culture in PVA-based media supplemented with 740Y-P and THPO (T) or butyzamide
- 979 (Buty), relative to fresh CD34⁺ cells. Mean \pm S.D. of three independent cultures. ***P* = 980 0.0017.
- 981 (I) The frequency of cells during the culture of $2x10^4$ human CB CD34⁺ cells in PVA-982 based media containing 1 μ M 740Y-P and 0.1 μ M butyzamide. Mean <u>+</u> S.D. of three 983 independent cultures.
- 984 (m) Representative image of a day-14 PVA-based culture containing 1 μ M 740Y-P and
- 985 0.1 μM butyzamide (2a media). Representative of at least five experiments. Scale bar:
 986 100 μm.
- 987 (n) Megakaryocytic (MgK) colony numbers obtained from 50 CD34⁺ cells sorted from 988 day 7 and day 14 PVA-based cultures containing 1 μ M 740Y-P and 0.1 μ M butyzamide 989 (2a media). Mean <u>+</u> S.D. of three independent cultures. **P* = 0.0161.
- 990 (o, p) Mean human CD45⁺ peripheral blood (PB) and BM chimerism in recipient 991 NOD/Shi-scid IL-2R γ^{null} (NOG) mice following transplantation of 1x10⁴ cells derived 992 from a 7-day or 14-day culture of 1x10⁴ CB CD34⁺ cells in PVA-based 2a media 993 containing 1 μ M 740Y-P and 0.1 μ M butyzamide. n=5 mice per group. (o) **[†]P = 0.0036; **[‡]P = 0.0037; ***P = 0.0007, (p)***P = 0.0003.
- 995 Statistical significance was calculated using an unpaired two-tailed t-test. n.s., not996 significant.
- 997

998 Extended Data Figure 2: Long-term ex vivo expansion of human HSPCs in chemically999 defined cytokine-free cultures

- 1000 (a) $CD34^+$ EPCR⁺ cell numbers after a 7-day culture of $2x10^4$ human CB CD34⁺ cells
- 1001 in PVA-based media containing 750 nM SR-1 and/or 70 nM UM171 in addition to 2a 1002 media. Mean of three independent cultures. ****P < 0.0001.
- 1003 (b) CD34⁺ EPCR⁺ and CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cell numbers after a 14-
- 1004 day culture of $2x10^4$ CB CD34⁺ cells in PVA-based 2a media with or without 70 nM
- 1005 UM171. Mean of three independent cultures. ****P < 0.0001.
- 1006 (c) The frequency of CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells after a 10-day culture
- 1007 of $2x10^4$ human CB CD34⁺ cells in StemSpan SFEM supplemented with cytokines with
- 1008 35 nM UM171 and PVA-based 2a media with 35 or 70 nM UM171. Mean of three 1009 independent cultures. ****P < 0.0001.
- 1010 (d) Annexin V staining assay of total cells after a 7-day culture of $2x10^4$ human CB
- 1011 CD34⁺ cells in PVA-based media containing 750 nM SR-1 and/or 0.1% BSA in addition
- 1012 to 2a or 10 ng/ml SCF (S) and 100 ng/ml THPO (T). Mean of three independent cultures.
- **1013** *****P* < 0.0001.
- 1014 (e) Mean human CD45⁺ PB chimerism in recipient NOG mice at 24 weeks following
- 1015 transplantation of 1×10^4 fresh CB CD34⁺ cells or the cells derived from a 10-day or 30-
- 1016 day culture of 1×10^4 CB CD34⁺ cells in PVA-based 3a media. n=3 mice per group. Details

1017 described in **Figure 2d**. $*^{\dagger}P = 0.0495$; $*^{\dagger}P = 0.0319$.

- 1018 (f) Mean 24-week human CD45⁺, CD34⁺, CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cell
- 1019 chimerism in the BM from mice described in (e). n=3 mice per group. Details described
- 1020 in Figure 2d. $*^{\dagger}P = 0.0112$; $*^{\ddagger}P = 0.0480$; $*^{\$}P = 0.0187$; **P = 0.0075; ***P = 0.0003;
- 1021 ****P < 0.0001.
- 1022 (g) Mean 24-week human CD45⁺, CD34⁺, CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cell
- 1023 chimerism in the spleen from mice described in (\mathbf{e}) . n=3 mice per group. Details described
- 1024 in Figure 2d. **P = 0.0014.
- 1025 Statistical significance was calculated using one-way ANOVA or an unpaired two-tailed
- 1026 *t*-test. n.s., not significant.
- 1027
- 1028 Extended Data Figure 3: Caprolactam polymer-based 3a media supports efficient
 1029 expansion of human HSCs ex vivo

- 1030 (a) CD34⁺CD45RA⁻ cell numbers of human-cord-blood-derived CD34⁺ cultured with
- 1031 SC79 or 740Y-P in addition to human 10 ng/ml SCF (S) and 100 ng/ml THPO (T) in
- 1032 PCL-PVAc-PEG culture conditions for 7 days. The starting cell count was $2x10^4$. Mean
- 1033 of three independent cultures. *P = 0.0210.
- 1034 (b) $CD34^+CD41^-CD90^+CD45RA^-$ cell numbers after a 7-day culture of $2x10^4$ human CB
- 1035 CD34⁺ cells in PCL-PVAc-PEG-based media containing 740Y-P and 100 ng/ml THPO
- 1036 (T) and/or butyzamide (Buty). Mean of three independent cultures. *P = 0.0256.
- 1037 (c) CD34⁺EPCR⁺ and CD34⁺EPCR⁺CD90⁺CD45RA⁻ cell numbers after a 7-day culture
- 1038 of $2x10^4$ human CB CD34⁺ cells in PCL-PVAc-PEG -based 2a media containing 750 nM
- 1039 SR-1 and/or 70 nM UM171. Mean of three independent cultures. **P = 0.0021; ***P = 1040 = 0.0002.
- 1041 (d) $CD34^+CD41^-CD90^+CD45RA^-$ cell numbers after a 7-day culture of $2x10^4$ of human
- 1042 CB CD34⁺ cells in PCL-PVAc-PEG-based media containing 0-20 µM, 740Y-P, and 0.1
- 1043 μ M butyzamide. Mean of three independent cultures. *[†]P = 0.0214; *[‡]P = 0.0440.
- 1044 (e) The frequency of CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells after a 7-day culture of
- 1045 $2x10^4$ human CB CD34⁺ cells in PCL-PVAc-PEG-based 2a media with 35 nM or 70 nM
- 1046 UM171. Mean of three independent cultures.
- 1047 (f) GEmM colony numbers generated from CD34⁺ cells after a 10-day culture in PVA-
- 1048 and/or PCL-PVAc-PEG-based 3a media. Mean of three independent cultures. ***P = 0.0005.
- 1050 (g) Annexin V staining assay of total cells after a 7-day culture of $2x10^4$ human CB CD34⁺
- 1051 cells in PVA- or PCL-PVAc-PEG-based 2a media containing 750 nM SR-1. Mean of 1052 three independent cultures. **P = 0.0086.
- 1053 (h) The frequency of PI positive cells after a 7-day culture of $2x10^4$ human CB CD34⁺
- 1054 cells in PCL-PVAc-PEG-based 3a media with or without 10 µM LY294002 (Chemscene,
- 1055 CAS No. 154447-36-6), PI3-kinase inhibitor. Mean of three independent cultures. ****P1056 < 0.0001.
- 1057 (i) Total and CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cell numbers after a 10-day culture
- 1058 of $2x10^4$ adult-peripheral blood stem cell (PBSC) CD34⁺ cells in PVA- or PCL-PVAc-
- 1059 PEG-based 3a media including UM729 instead of UM171. Mean of three independent
- 1060 cultures. **P*=0.0153, ***P*=0.0079.

- (j) Mean human CD45⁺ PB chimerism in recipient NOG mice at 24 weeks after
 transplantation of 1x10⁴ day-30 cells derived from CB CD34⁺ cells cultured in 3a media
 containing PVA or PCL-PVAc-PEG. n=3 mice per group. Detailed described in Figure
 3c.
 (k) Mean 24-week human CD45⁺, CD34⁺, CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cell
 chimerism in the BM from mice described in (j). n=3 mice per group. Detailed described
 in Figure 3c.
- 1068 (I) Mean 24-week human CD45⁺, CD34⁺, CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ cell
- 1069 chimerism in the spleen from mice described in (j). n=3 mice per group. Detailed
 1070 described in Figure 3c.
- 1071 Statistical significance was calculated using one-way ANOVA or an unpaired two-tailed1072 *t*-test: n.s., not significant.
- 1073

1074 Extended Data Figure 4: Comparison of human HSC culture protocols

- 1075 (a) Total cell numbers generated from a 10-day culture of $2x10^4$ human CB CD34⁺ cells
- 1076 in PCL-PVAc-PEG or StemSpan SFEM-based cytokine-cocktail media with UM171
- 1077 and/or SR-1 (see in *Methods* for details), or PCL-PVAc-PEG-based 3a media. Mean of

1078 three independent cultures. ***P = 0.0002; ****P < 0.0001.

- 1079 (b) The frequency and (c) absolute number of CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺
- 1080 cells in cultures described in (a). Mean of three independent cultures. (b) ****P < 0.0001.
- 1081 (c) ***P* = 0.0010; ***[†]*P* = 0.0009; ***[‡]*P* = 0.0002; *****P* < 0.0001.
- 1082 (d) The ROS and γ H2AX level of fresh CB CD34⁺ cells (fresh) and CD34⁺ cells in 1083 cultures described in (a). Mean of three independent cultures. **P = 0.0028; ***P =1084 0.0004; ****P < 0.0001.
- 1085 (e) Relative mean fluorescence intensity (MFI) for ROS and yH2AX in fresh CB CD34⁺
- 1086 cells (fresh) and cells from 10 or 30 day PCL-PVAc-PEG-based 3a media cultures. Mean
- 1087 of three independent cultures.
- 1088 (f, g) Mean human CD45⁺ PB and BM chimerism in recipient NOG mice following
- 1089 transplantation of 1×10^4 day-10 cells derived cultures as describe in (a). n=4 mice per 1090 group. ***P = 0.0001; ****P < 0.0001.

- 1091 (h, i) Mean human CD45⁺ PB and BM chimerism in secondary recipient NOG mice
- 1092 following transplantation of 1×10^6 BM cells derived from primary recipient mice, as
- 1093 describe in (**f**, **g**). n=3 mice per group. $*^{\dagger}P = 0.0180$; $*^{\dagger}P = 0.0299$.
- 1094 Statistical significance was calculated using one-way ANOVA: n.s., not significant.
- 1095

1096 Extended Data Figure 5: Gating strategy for HSCs fraction by flow cytometry

- 1097 (a) FACS gating strategy for detecting CD34⁺EPCR⁺CD90⁺CD45RA⁻ITGA3⁺ cells after
- 1098 10-day culture in 3a media containing PCL-PVAc-PEG or using UM171/SR-1.
- 1099

1100 Extended Data Figure 6: Profile of human HSCs expanded in 3a media

- 1101 (a) Volcano plot showing differentially expressed genes (DEGs) detected in bulk RNA-
- 1102 sequencing of CD34^{high}EPCR⁺ (right) and CD34^{high}EPCR⁻ (left) cells after 10-day culture
- 1103 in PCL-PVAc-PEG based 3a media. DEGs are highlighted as red dots ($log_2FC > 2$, -logP
- 1104 Value <14). Gene names are shown in the boxes.
- (b) GO Term cellular component-specific GSEA analysis performed on DEGs, displayedas a dotplot.
- 1107 (c) Expression of key genes within annotated clusters, displayed as a dotplot.
- (d, e) Feature plots showing *HLF* (c) and *AVP* (d) gene expression within the integratedcell map.
- 1110 (f) Ratio of each cell cluster within PCL-PVAc-PEG based 3a cultures, StemSpan with
- 1111 SR-1 cultures, and StemSpan with UM171 cultures, as described in **Figure 4e**.
- 1112 (g) Comparison of scRNAseq data from cells cultured for 10 days in PCL-PVAc-PEG
- 1113 based 3a media with two dataset of cells cultured for 7 days in StemSpan SFEM with
- 1114 UM171 cultures obtained from GEO (GSE 153370).
- 1115 (h) Violin plots displaying *HLF* expression in cells from a 10-day culture using 3a media,
- 1116 cells from UM171/SR-1 cultures, and two fresh CBs from publicly available data (GSE
- 1117 153370).
- 1118

1119 Extended Data Figure 7: Split clone assays

- 1120 (a) Schematic of assay of the single HSC expansion and split clone assay. Single human
- 1121 CD34⁺CD38⁻CD90⁺CD45RA⁻CD49f⁺ CB cells were sorted into 96 wells and expanded

in 3a media containing PCL-PVAc-PEG for 7 days. Individual HSC clones were then
transplanted into three recipient W41/W41 mice.

(b) Human CD45⁺ PB chimerism in recipient W41/W41 mice 24 weeks after
transplantation of day-7 cells derived from single human CD34⁺CD38⁻CD90⁺CD45RA⁻
CD49f⁺ CB cell cultured in 3a media containing PCL-PVAc-PEG (3 mice/well), as

- 1127 described in (**a**).
- 1128

1129 Extended Data Figure 8: NOG-W41/W41 mice display high human hematopoietic cell 1130 chimerism

1131 (a) Mean human CD45⁺ PB chimerism in recipient NOD.Cg-Prkdc^{scid} Il2rg^{tm1Sug}

1132 Kit^{em1(V831M)Jic}/Jic (W41/W41), W41/+, +/+ mice at 4, 8, 12, 16 and 20 weeks following

- 1133 transplantation of $5x10^4$ fresh CB CD34⁺ cells. n=3-4 mice per group. ***P = 0.0004; 1134 ****P < 0.0001.
- 1135 (b) Mean 24-week human CD45⁺ cell chimerism in the BM and spleen from mice 1136 described in (a). n=3-4 mice per group. $**^{\dagger}P = 0.0016$; $**^{\dagger}P = 0.0021$; $***^{\dagger}P = 0.0001$; 1137 $***^{\dagger}P = 0.0006$.
- P = 0.0006.
- 1138 (c) Mean human CD45⁺, CD19⁺, CD33⁺, CD3⁺, CD56⁺ and CD66b⁺ PB chimerism in

recipient non-irradiated or irradiated (0.5 Gy) W41/W41 mice at 4-, 8-, 12- and 16-weeks

1140 following transplantation of $5x10^4$ fresh CB CD34⁺ cells. Mean <u>+</u> S.D of 3-4 mice per

1141 group. $*^{\dagger}P = 0.0257$; $*^{\dagger}P = 0.0335$; $**^{\dagger}P = 0.0030$; $**^{\dagger}P = 0.0060$; ***P = 0.0002.

- 1142 Statistical significance was calculated using one-way ANOVA or an unpaired two-tailed
- 1143 *t*-test: n.s., not significant.
- 1144

1145 Extended Data Table 1: Detailed data of xenotransplantation assays

1146 (a) Mean \pm S.D 16-week frequency of CD33⁺ myeloid cells, CD3⁺ T cells, and CD19⁺ B

- 1147 cells of human CD45⁺ cells in the BM and spleen of mice described in **Figure 2e**.
- 1148 (b) Mean \pm S.D 16-week frequency of CD33⁺ myeloid cells, CD3⁺ T cells, and CD19⁺ B
- 1149 cells of human CD45⁺ cells in the BM and spleen of mice described in **Figure 3d**.
- 1150 (c) Mean \pm S.D 24-week frequency of CD33⁺ myeloid cells, CD3⁺ T cells, and CD19⁺ B
- 1151 cells of human CD45⁺ cells in the PB of mice described in **Figure 4i**.
- 1152

1153 Extended Data Table 2: Whole exome sequencing on uncultured and 10-day cultured

- 1154 *cells.* Whole exome sequencing on fresh CB and 10-day cultured CB CD34+ cells with
- 1155 PCL-PVAc-PEG-based 3a medium.



Figure 1 Sakurai, et al.



Figure 2 Sakurai, et al.



Figure 3 Sakurai, et al.



Figure 4 Sakurai, et al.