

frequently detached from the culture dishes with gentle pipetting (data not shown). This observation suggested that multilayered hiPSC adhesion on substrate was weakened, resulting in cell detachment from the dish. Because the cells reattached onto the culture dish, however, the expression of cell surface integrin, which mediates adhesion against vitronectin, might not have decreased in multilayered cells. One possible explanation of this spontaneous detachment is the energetic stability of cell aggregates. Supposing that the cell–cell adhesion is stronger than the cell–vitronectin adhesion, cells floating in spherical aggregates would be more energetically stable than cells adhering as dome aggregates onto a small area when the number of composing cells becomes large.

Our present method presents 4 advantages compared with previously reported methods.<sup>9,10,22,30</sup> (1) Because low-pressure plasma treatment using perforated masks enables stable and equal patterning on a wide area, it is suitable for creating a large number of homogeneously patterned spots reproducibly. (2) Single-step coating of a mixture of proteins easily enables the production of small patterns. In our previous research,<sup>10</sup> the extracellular matrix was coated before the perforated mask was removed. Using this method, we sometimes found air bubbles in the perforated holes, which prevented coating in those positions. Moreover, similar methods of producing cell patterning required additional steps, including multistep protein coating such as BSA coating followed by extracellular matrix coating.<sup>22</sup> Comparatively, our present method is quite simple. (3) Because the cost-effectiveness and availability of BSA are higher than those of  $\gamma$ -globulin, BSA is a good candidate for practical use in cell patterning.<sup>9</sup> (4) There are many types of microfabrication tools that can be used to make equally sized spheroids, although most of them are expensive and difficult to use in cell culture laboratories.<sup>30</sup> Our method is an easy and cost-effective way to fabricate hiPSC discs and spheroids.

## V. CONCLUSION

A combination of patterned plasma oxidation with perforated PMMA masks, single-step coating of a mixture of vitronectin and BSA, and serum-free and feeder-free culture conditions enables the simple production of equally sized, small hiPSC discs on a broad PDMS area. We also succeeded in fabricating hiPSC spheroids, which formed spontaneously from the discs. Because our method is simple and cost-effective, it can be used in practical experiments to mimic early human embryonic development. We believe that our technique for patterning hiPSCs will be useful for the development of new bioengineering devices in the search for effective cell differentiation methods and in the testing of drug safety for early human embryonic development, contributing to future medical applications.

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# Prediction of interindividual differences in hepatic functions and drug sensitivity by using human iPSC-derived hepatocytes

Kazuo Takayama<sup>a,b,c</sup>, Yuta Morisaki<sup>a</sup>, Shuichi Kuno<sup>a</sup>, Yasuhito Nagamoto<sup>a,c</sup>, Kazuo Harada<sup>d</sup>, Norihisa Furukawa<sup>a</sup>, Manami Ohtaka<sup>e</sup>, Ken Nishimura<sup>f</sup>, Kazuo Imagawa<sup>a,c,g</sup>, Fuminori Sakurai<sup>a,h</sup>, Masashi Tachibana<sup>a</sup>, Ryo Sumazaki<sup>g</sup>, Emiko Noguchi<sup>i</sup>, Mahito Nakanishi<sup>e</sup>, Kazumasa Hirata<sup>d</sup>, Kenji Kawabata<sup>j,k</sup>, and Hiroyuki Mizuguchi<sup>a,b,c,l,1</sup>

<sup>a</sup>Laboratory of Biochemistry and Molecular Biology; <sup>b</sup>iPS Cell-based Research Project on Hepatic Toxicity and Metabolism, <sup>d</sup>Laboratory of Applied Environmental Biology, <sup>e</sup>Laboratory of Regulatory Sciences for Oligonucleotide Therapeutics, Clinical Drug Development Project, and <sup>k</sup>Laboratory of Biomedical Innovation, Graduate School of Pharmaceutical Sciences, Osaka University, Osaka 565-0871, Japan; <sup>c</sup>Laboratory of Hepatocyte Regulation, and <sup>l</sup>Laboratory of Stem Cell Regulation, National Institute of Biomedical Innovation, Osaka 567-0085, Japan; <sup>f</sup>Research Center for Stem Cell Engineering, National Institute of Advanced Industrial Science and Technology, Ibaraki 305-8562, Japan; <sup>g</sup>Laboratory of Gene Regulation, <sup>h</sup>Department of Child Health, and <sup>i</sup>Department of Medical Genetics, Faculty of Medicine, University of Tsukuba, Ibaraki 305-8575, Japan; and <sup>j</sup>The Center for Advanced Medical Engineering and Informatics, Osaka University, Osaka 565-0871, Japan

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**Interindividual differences in hepatic metabolism, which are mainly due to genetic polymorphism in its gene, have a large influence on individual drug efficacy and adverse reaction. Hepatocyte-like cells (HLCs) differentiated from human induced pluripotent stem (iPS) cells have the potential to predict interindividual differences in drug metabolism capacity and drug response. However, it remains uncertain whether human iPSC-derived HLCs can reproduce the interindividual difference in hepatic metabolism and drug response. We found that cytochrome P450 (CYP) metabolism capacity and drug responsiveness of the primary human hepatocytes (PHH)-iPS-HLCs were highly correlated with those of PHHs, suggesting that the PHH-iPS-HLCs retained donor-specific CYP metabolism capacity and drug responsiveness. We also demonstrated that the interindividual differences, which are due to the diversity of individual SNPs in the CYP gene, could also be reproduced in PHH-iPS-HLCs. We succeeded in establishing, to our knowledge, the first PHH-iPS-HLC panel that reflects the interindividual differences of hepatic drug-metabolizing capacity and drug responsiveness.**

human iPSC cells | hepatocyte | CYP2D6 | personalized drug therapy | SNP

**D**rug-induced liver injury (DILI) is a leading cause of the withdrawal of drugs from the market. Human induced pluripotent stem cell (iPSC)-derived hepatocyte-like cells (HLCs) are expected to be useful for the prediction of DILI in the early phase of drug development. Many groups, including our own, have reported that the human iPSC-HLCs have the ability to metabolize drugs, and thus these cells could be used to detect the cytotoxicity of drugs that are known to cause DILI (1, 2). However, to accurately predict DILI, it will be necessary to establish a panel of human iPSC-HLCs that better represents the genetic variation of the human population because there are large interindividual differences in the drug metabolism capacity and drug responsiveness of hepatocytes (3). However, it remains unclear whether the drug metabolism capacity and drug responsiveness of human iPSC-HLCs could reflect those of donor parental primary human hepatocytes (PHHs). To address this issue, we generated the HLCs differentiated from human iPSCs which had been established from PHHs (PHH-iPS-HLCs). Then, we compared the drug metabolism capacity and drug responsiveness of PHH-iPS-HLCs with those of their parental PHHs, which are genetically identical to the PHH-iPS-HLCs.

Interindividual differences of cytochrome P450 (CYP) metabolism capacity are closely related to genetic polymorphisms, especially single nucleotide polymorphisms (SNPs), in CYP genes (4). Among the various CYPs expressed in the liver, CYP2D6 is responsible for the metabolism of approximately

a quarter of commercially used drugs and has the largest phenotypic variability, largely due to SNPs (5). It is known that certain alleles result in the poor metabolizer phenotype due to a decrease of CYP2D6 metabolism. Therefore, the appropriate dosage for drugs that are metabolized by CYP2D6, such as tamoxifen, varies widely among individuals (6). Indeed, in the 1980s, polymorphism in CYP2D6 appears to have contributed to the withdrawal of CYP2D6-metabolized drugs such as perhexiline from the market in many countries (7). If we could establish a panel of HLCs that better represents the diversity of genetic polymorphisms in the human population, it might be possible to determine the appropriate dosage of a drug for a particular individual. However, it is not known whether the drug metabolism capacity and drug responsiveness of HLCs reflect the genetic diversity, including SNPs, in CYP genes. Therefore, in this study we generated HLCs from several PHHs that have various SNPs on CYP2D6 and then compared the CYP2D6 metabolism capacity and responses to CYP2D6-metabolized drugs between the PHH-iPS-HLCs and parental PHHs.

## Significance

**We found that individual cytochrome P450 (CYP) metabolism capacity and drug sensitivity could be predicted by examining them in the primary human hepatocytes–human induced pluripotent stem cells–hepatocyte-like cells (PHH-iPS-HLCs). We also confirmed that interindividual differences of CYP metabolism capacity and drug responsiveness that are due to the diversity of individual single nucleotide polymorphisms in the CYP gene could also be reproduced in the PHH-iPS-HLCs. These findings suggest that interindividual differences in drug metabolism capacity and drug response could be predicted by using HLCs differentiated from human iPSC cells. We believe that iPSC-HLCs would be a powerful technology not only for accurate and efficient drug development, but also for personalized drug therapy.**

Author contributions: K.T. and H.M. designed research; K.T., Y.M., and S.K. performed research; K.T., Y.M., Kazuo Harada, M.O., K.N., K.I., M.N., and Kazumasa Hirata contributed new reagents/analytic tools; K.T., Y.N., N.F., F.S., M.T., R.S., E.N., K.K., and H.M. analyzed data; and K.T. and H.M. wrote the paper.

The authors declare no conflict of interest.

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Data deposition: The DNA microarray data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, [www.ncbi.nlm.nih.gov/geo](http://www.ncbi.nlm.nih.gov/geo) (accession no. GSE61287).

<sup>1</sup>To whom correspondence should be addressed. Email: [mizuguch@phs.osaka-u.ac.jp](mailto:mizuguch@phs.osaka-u.ac.jp).

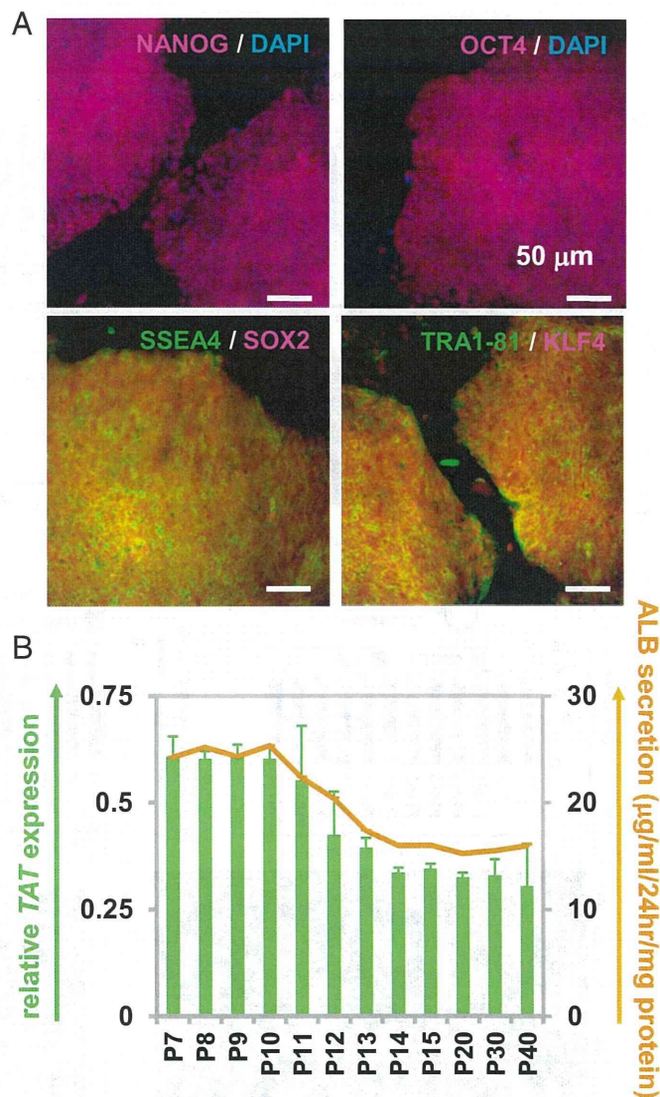
This article contains supporting information online at [www.pnas.org/lookup/suppl/doi:10.1073/pnas.1413481111/-DCSupplemental](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1413481111/-DCSupplemental).

To this end, PHHs were reprogrammed into human iPSCs and then differentiated into the HLCs. To examine whether the HLCs could reproduce the characteristics of donor PHHs, we first compared the CYP metabolism capacity and response to a hepatotoxic drug between PHHs and genetically identical PHH-iPS-HLCs (12 donors were used in this study). Next, analyses of hepatic functions, including comparisons of the gene expression of liver-specific genes and CYPs, were performed to examine whether the hepatic characteristics of PHHs were reproduced in the HLCs. To the best of our knowledge, this is the first study to compare the functions between iPSC-derived cells from various donors and their parental cells with identical genetic backgrounds. Finally, we examined whether the PHH-iPS-HLCs exhibited a capacity for drug metabolism and drug responsiveness that reflect the genetic diversity such as SNPs on CYP genes.

## Results

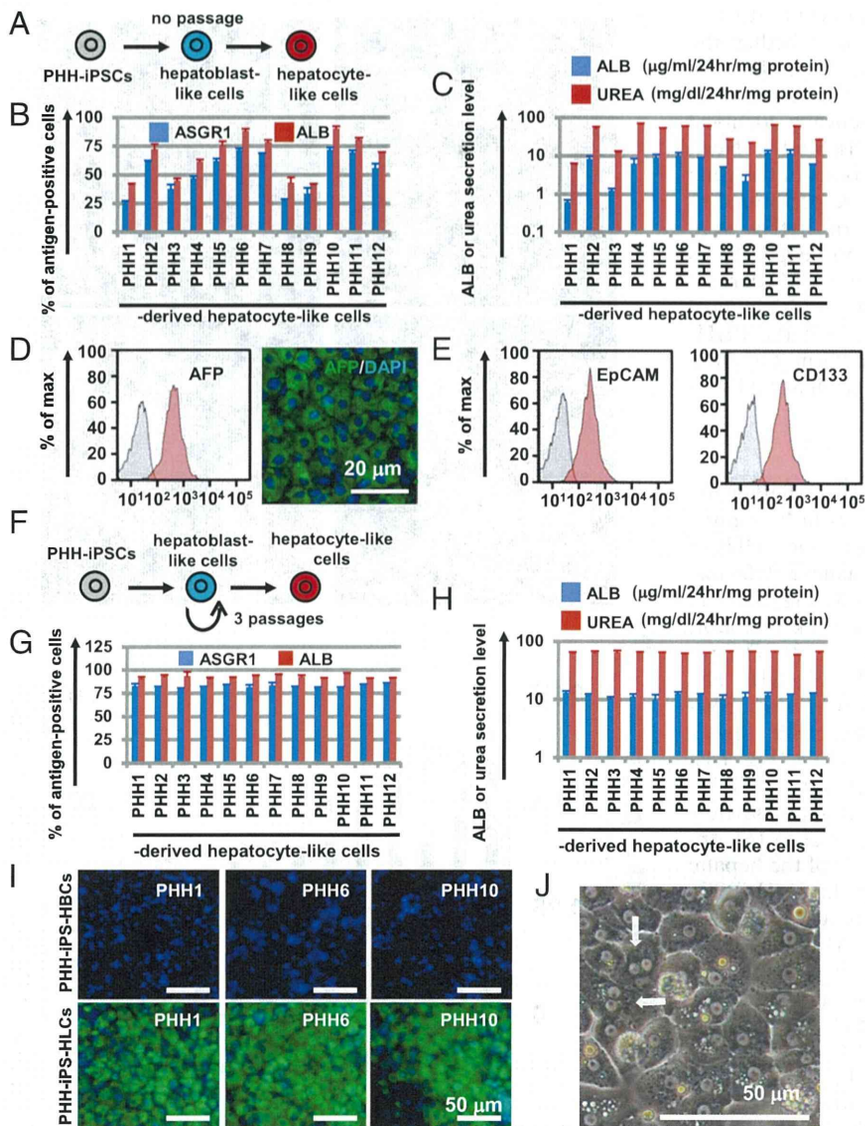
**Reprogramming of PHHs to Human iPSCs.** To examine whether the HLCs could reproduce interindividual differences in liver functions, we first tried to generate human iPSCs from the PHHs of 12 donors. PHHs were transduced with a Yamanaka 4 factor-expressing SeV (SeVdp-iPS) vector (*SI Appendix, Fig. S1A*) in the presence of SB431542, PD0325901, and a rock inhibitor, which could promote the somatic reprogramming (8). The reprogramming procedure is shown in *SI Appendix, Fig. S1B*. The human iPSCs generated from PHHs (PHH-iPSCs) were positive for alkaline phosphatase (*SI Appendix, Fig. S1B, Right*), NANOG, OCT4, SSEA4, SOX2, Tra1-81, and KLF4 (Fig. 1A). The gene expression levels of the pluripotent markers (*OCT3/4*, *SOX2*, and *NANOG*) in the PHH-iPSCs were approximately equal to those in human embryonic stem cells (ESCs) (*SI Appendix, Fig. S1C, Left*). The gene expression levels of the hepatic markers [*albumin (ALB)*, *CYP3A4*, and  *$\alpha$ AT*] in the PHH-iPSCs were significantly lower than those in the parental PHHs (*SI Appendix, Fig. S1C, Right*). We also confirmed that the PHH-iPSCs have the ability to differentiate into the three embryonic germ layers in vitro by embryoid body formation and in vivo by teratoma formation (*SI Appendix, Fig. S2 A and B*, respectively). To verify that the PHH-iPSCs originated from PHHs, short tandem repeat analysis was performed in the PHH-iPSCs and parental PHHs (*SI Appendix, Fig. S2C*). The results showed that the PHH-iPSCs were indeed originated from PHHs. Taken together, these results indicated that the generation of human iPSCs from PHHs was successfully performed. It is known that a transient epigenetic memory of the original cells is retained in early-passage iPSCs, but not in late-passage iPSCs (9). To examine whether the hepatic differentiation capacity of PHH-iPSCs depends on their passage number, PHH-iPSCs having various passage numbers were differentiated into the hepatic lineage (Fig. 1B). The *tyrosine aminotransferase (TAT)* expression levels and albumin (ALB) secretion levels in early passage PHH-iPS-HLCs (fewer than 10 passages) were higher than those of late passage PHH-iPS-HLCs (more than 14 passages). These results suggest that the hepatic differentiation tendency is maintained in early passage PHH-iPSCs, but not in late passage PHH-iPSCs. In addition, the hepatic functions of late passage PHH-iPS-HLCs were similar to those in the HLCs derived from late passage non-PHH-derived iPS cells (such as dermal cells, blood cells, and Human Umbilical Vein Endothelial Cells (HUVEC)-derived iPS cells) (*SI Appendix, Fig. S3*). Therefore, PHH-iPSCs, which were passaged more than 20 times, were used in our study to avoid any potential effect of transient epigenetic memory retained in parental PHHs on hepatic functions.

**HLCs Were Differentiated from PHH-iPSCs Independent of Their Differentiation Tendency.** To compare the hepatic characteristics among the PHH-iPS-HLCs that were generated from PHHs of



**Fig. 1.** Establishment and characterization of human iPSCs generated from PHHs. (A) The PHH-iPSCs were subjected to immunostaining with anti-NANOG (red), OCT4 (red), SSEA4 (green), SOX2 (red), TRA1-81 (green), and KLF4 (red) antibodies. Nuclei were counterstained with DAPI (blue) (*Upper*). (B) The *TAT* expression and ALB secretion levels in the PHH-iPS-HLCs (P7–P40) were examined. On the y axis, the gene expression level of *TAT* in PHHs was taken as 1.0.

the 12 donors, all of the PHH-iPSCs were differentiated into the HLCs as described in Fig. 2A. However, the differences in hepatic function among PHH-iPS-HLCs could not be properly compared because there were large inter-PHH-iPSC line differences in the hepatic differentiation efficiency based on ALB or asialoglycoprotein receptor 1 (ASGR1) expression analysis (Fig. 2B). In addition, there were also large inter-PHH-iPS-HLC line differences in ALB or urea secretion capacities (Fig. 2C). These results suggest that it is impossible to compare the hepatic characteristics among PHH-iPS-HLCs without compensating for the differences in the hepatic differentiation efficiency. Recently, we developed a method to maintain and proliferate the hepatoblast-like cells (HBCs) generated from human ESCs/iPSCs by using human laminin 111 (LN111) (10). To examine whether the hepatic differentiation efficiency could be made uniform by generating the HLCs following purification and proliferation of the HBCs, the PHH-iPS-HBCs were cultured on LN111 as

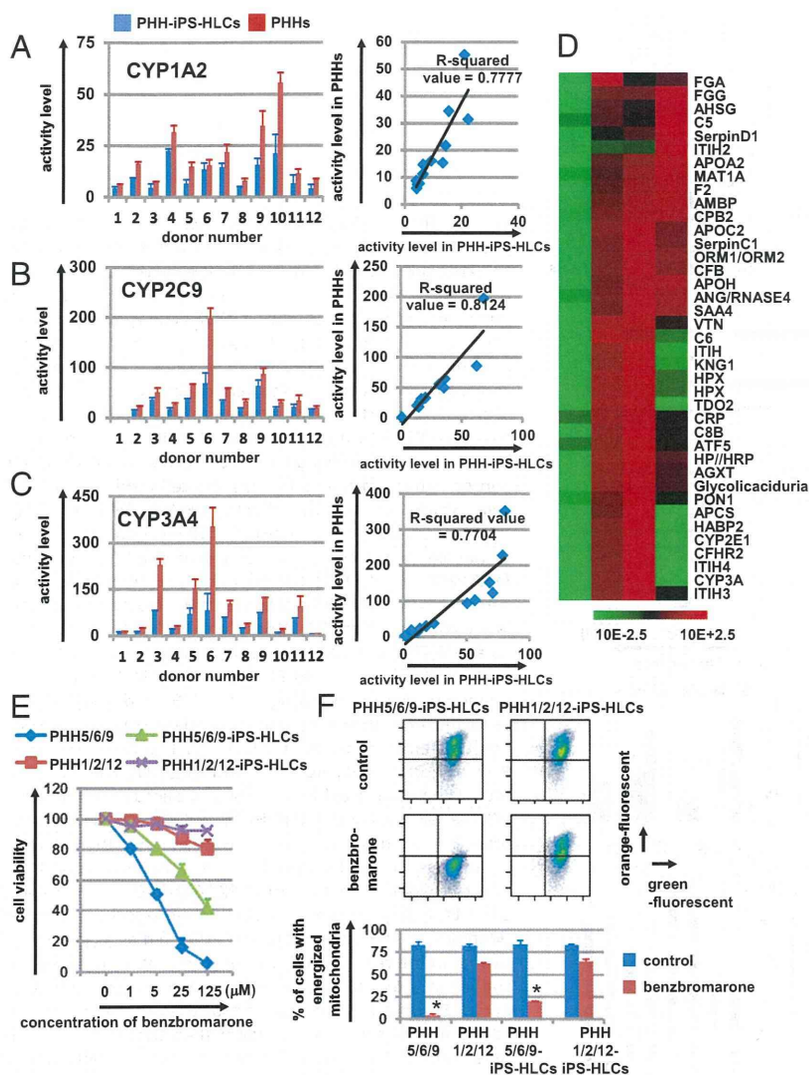


**Fig. 2.** Highly efficient hepatocyte differentiation from PHH-iPSCs independent of their differentiation tendency. (A) PHH-iPSCs were differentiated into the HLCs via the HBCs. (B) On day 25 of differentiation, the efficiency of hepatocyte differentiation was measured by estimating the percentage of ASGR1- or ALB-positive cells using FACS analysis. (C) The amount of ALB or urea secretion was examined in PHH-iPS-HLCs. (D) The percentage of AFP-positive cells in PHH-iPS-HBCs was examined by using FACS analysis (Left). The PHH-iPS-HBCs were subjected to immunostaining with anti-AFP (green) antibodies. Nuclei were counterstained with DAPI (blue) (Right). (E) The percentage of EpCAM- and CD133-positive cells in PHH-iPS-HBCs was examined by using FACS analysis (Left). (F) PHH-iPSCs were differentiated into the hepatic lineage, and then PHH-iPS-HBCs were purified and maintained for three passages on human LN111. Thereafter, expanded PHH-iPS-HBCs were differentiated into the HLCs. (G) The efficiency of hepatic differentiation from PHH-iPS-HBCs was measured by estimating the percentage of ASGR1- or ALB-positive cells using FACS analysis. (H) The amount of ALB or urea secretion in PHH-iPS-HLCs was examined. Data represent the mean  $\pm$  SD from three independent differentiations. (I) The PHH1-, 6-, or 10-iPS-HBCs and -HLCs were subjected to immunostaining with anti- $\alpha$ AT (green) antibodies. Nuclei were counterstained with DAPI (blue). (J) A phase-contrast micrograph of PHH-iPS-HLCs.

previously described (10), and then differentiated into the HLCs. Almost all of the cells were positive for the hepatoblast marker [ $\alpha$ -fetoprotein (AFP)] (Fig. 2D). In addition, the PHH-iPS-HBCs were positive for two other hepatoblast markers, EpCAM and CD133 (Fig. 2E). To examine the hepatic differentiation efficiency of the PHH-iPS-HBCs maintained on LN111-coated dishes for three passages (Fig. 2F), the HBCs were differentiated into the HLCs, and then the percentage of ALB- and ASGR1-positive cells was measured by FACS analysis (Fig. 2G). All 12 PHH-iPS-HBCs could efficiently differentiate into the HLCs, yielding more than 75% or 85% ASGR1- or ALB-positive cells, respectively. In addition, there was little difference between the PHH-iPSC lines in ALB or urea secretion capacities (Fig. 2H). Although there were large differences in the hepatic differentiation capacity among the PHH1/6/10 (Fig. 2B), PHH1/6/10-iPS-HBCs could efficiently differentiate into the HLCs that homogeneously expressed  $\alpha$ AT (Fig. 2I). After the hepatic differentiation of the PHH-iPS-HBCs, the morphology of the HLCs was similar to that of the PHHs: polygonal with distinct round binuclei (Fig. 2J). These results indicated that the hepatic differentiation efficiency of the 12 PHH-iPSC lines could be rendered uniform by inducing hepatic maturation after the establishment of self-renewing HBCs. Therefore, we expected

that differences in the hepatic characteristics among the HLCs generated from the 12 individual donor PHH-iPS-HBCs could be properly compared. In addition, the hepatic differentiation efficiency could be rendered uniform not only in the PHH-iPSC lines but also in non-PHH-iPSC lines and human ESCs by performing hepatic maturation after the establishment of self-renewing HBCs (SI Appendix, Fig. S4). In Figs. 3 and 4, the HLCs were differentiated after the HBC proliferation step to normalize the hepatic differentiation efficiency.

**PHH-iPS-HLCs Retained Donor-Specific Drug Metabolism Capacity and Drug Responsiveness.** To examine whether the hepatic functions of individual PHH-iPS-HLCs reflect those of individual PHHs, the CYP metabolism capacity and drug responsiveness of PHH-iPS-HLCs were compared with those of PHHs. PHHs are often used as a positive control to assess the hepatic functions of the HLCs, although in all of the previous reports, the donor of PHHs has been different from that of human iPSCs. Because it is generally considered that CYP activity differs widely among individuals, the hepatic functions of the HLCs should be compared with those of genetically identical PHHs to accurately evaluate the hepatic functions of the HLCs. The CYP1A2, -2C9, and -3A4 activity levels in the PHH-iPS-HLCs were  $\sim$ 60% of

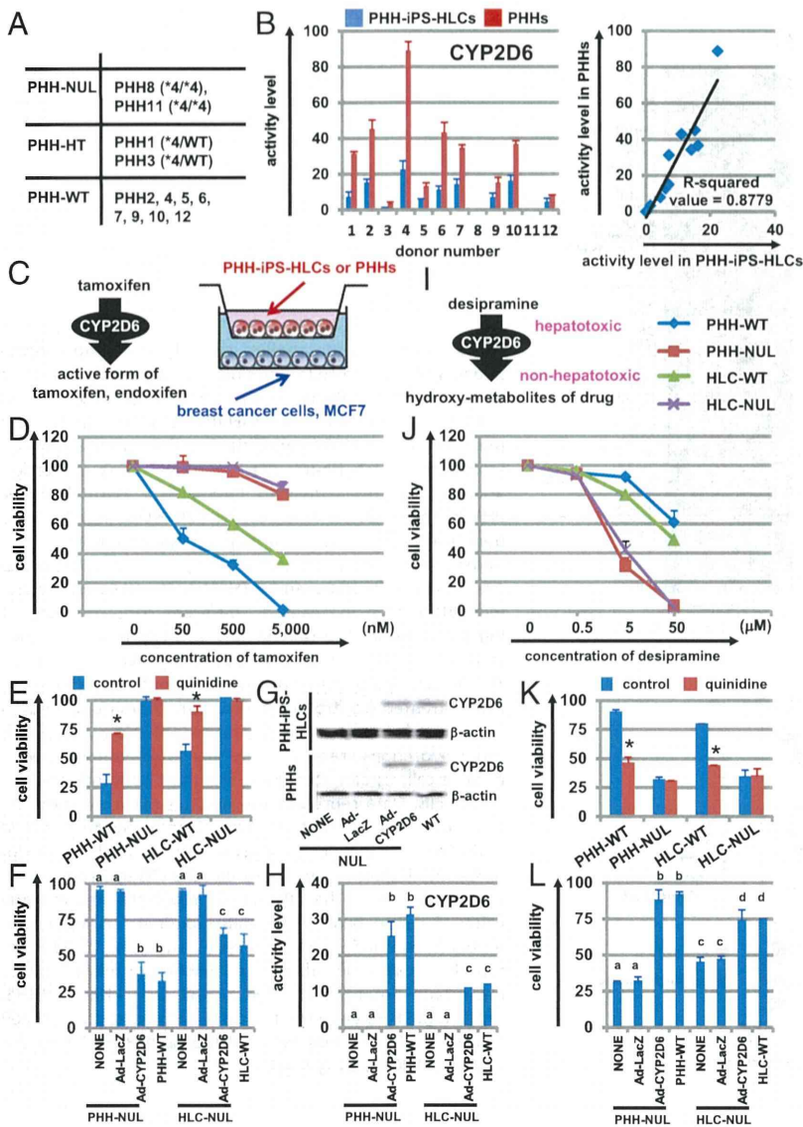


**Fig. 3.** The drug metabolism capacity and drug responsiveness of PHH-iPS-HLCs were highly correlated with those of their parental PHHs. (A–C) CYP1A2 (A), -2C9 (B), and -3A4 (C) activity levels in PHH-iPS-HLCs and PHHs were measured by LC-MS/MS analysis. The R-squared values are indicated in each figure. (D) The global gene expression analysis was performed in PHH9-iPS-HLCs, PHH9-iPS-HLCs, PHH9s, and HepG2 (PHH-iPS-HLCs, PHH-iPS-HLCs, and PHHs are genetically identical). Heat-map analyses of liver-specific genes are shown. (E) The cell viability of PHH5/6/9, PHH1/2/12, PHH5/6/9-iPS-HLCs, and PHH1/2/12-iPS-HLCs was examined after 24 h exposure to different concentrations of benzobromarone. The cell viability was expressed as a percentage of that in the cells treated only with solvent. (F) The percentage of cells with energized mitochondria in the DMSO-treated (control, *Upper*) or benzobromarone-treated (*Lower*) cells based on FACS analysis. Double-positive cells (green+/orange+) represent energized cells, whereas single-positive cells (green+/orange–) represent apoptotic and necrotic cells. Data represent the mean  $\pm$  SD from three independent experiments (*Lower Graph*). Student *t* test indicated that the percentages in the “control” were significantly higher than those in the “benzobromarone” group ( $P < 0.01$ ). The “PHH5/6/9” represents the average value of cell viability (E) or mitochondrial membrane potential (F) in PHH5, PHH6, and PHH9. The “PHH1/2/12” represents the average value of cell viability or mitochondrial membrane potential in PHH1, PHH2, and PHH12. PHH5, PHH6, and PHH9 were the top three with respect to CYP2C9 activity levels, whereas PHH1, PHH2, and PHH12 had the lowest CYP2C9 activity levels.

those in the PHHs (Fig. 3 A–C and *SI Appendix*, Fig. S5). Interestingly, the CYP1A2, -2C9, and -3A4 activity levels in the PHH-iPS-HLCs were highly correlated with those in the PHHs (the R-squared values were more than 0.77) (Fig. 3 A, B, and C, respectively). These results suggest that it would be possible to predict the individual CYP activity levels through analysis of the CYP activity levels of the PHH-iPS-HLCs. Because the average and variance of CYP3A4 activity levels in PHH-iPS-HLCs, non-PHH-iPS-HLCs, and human ES-HLCs were similar to each other (*SI Appendix*, Fig. S6), the drug metabolism capacity of PHH-iPS-HLCs might be similar to that of nonliver tissue-derived iPS-HLCs and human ES-HLCs. Therefore, it might be possible to predict the diversity of drug metabolism capacity among donors by using nonliver tissue-derived iPS-HLCs and human ES-HLCs as well as PHH-iPS-HLCs. On the other hand, the CYP induction capacities of PHH-iPS-HLCs were weakly correlated with those of PHHs (*SI Appendix*, Fig. S7 A–C). To further investigate the characteristics of the HLCs, DNA microarray analyses were performed in genetically identical undifferentiated iPSCs, PHH-iPS-HLCs, and PHHs. The gene expression patterns of liver-specific genes, CYPs, and transporters in the PHH-iPS-HLCs were similar to those in PHHs (Fig. 3D and *SI Appendix*, Fig. S7 D and E, respectively). Next, the hepatotoxic drug responsiveness of PHH-iPS-HLCs was compared with that of PHHs. Benzobromarone, which is known to cause

hepatotoxicity by CYP2C9 metabolism (11), was treated to PHH5/6/9 and PHH5/6/9-iPS-HLCs, which have high CYP2C9 activity, or PHH1/2/12 and PHH1/2/12-iPS-HLCs which have low CYP2C9 activity (Fig. 3E). The susceptibility of the PHH5/6/9 and PHH5/6/9-iPS-HLCs to benzobromarone was higher than that of PHH1/2/12 and PHH1/2/12-iPS-HLCs, respectively. These results were attributed to the higher CYP2C9 activity levels in PHH5/6/9 and PHH5/6/9-iPS-HLCs compared with those in PHH1/2/12 and PHH1/2/12-iPS-HLCs. Because it is also known that benzobromarone causes mitochondrial toxicity (12), an assay of mitochondrial membrane potential was performed in benzobromarone-treated PHHs and PHH-iPS-HLCs (Fig. 3F). The mitochondrial toxicity observed in PHH5/6/9 and PHH5/6/9-iPS-HLCs was more severe than that in PHH1/2/12 and PHH1/2/12-iPS-HLCs, respectively. Taken together, these results suggest that the hepatic functions of the individual PHH-iPS-HLCs were highly correlated with those of individual PHHs.

**Interindividual Differences in CYP2D6-Mediated Metabolism and Drug Toxicity, Which Are Caused by SNPs in CYP2D6, Are Reproduced in the PHH-iPS-HLCs.** Because certain SNPs are known to have a large impact on CYP activity, the genetic variability of CYP plays an important role in interindividual differences in drug response. CYP2D6 shows the large phenotypic variability due to genetic polymorphism (13). We next examined whether the PHHs used



**Fig. 4.** The interindividual differences in CYP2D6 metabolism capacity and drug responsiveness induced by SNPs in CYP2D6 are reproduced in the PHH-iPS-HLCs. (A) SNPs (CYP2D6\*3, \*4, \*5, \*6, \*7, \*8, \*16, and \*21) in the CYP2D6 gene were analyzed. (B) The CYP2D6 activity levels in PHH-iPS-HLCs and PHHs were measured by LC-MS/MS analysis. (C) The pharmacological activity of tamoxifen-dependent conversion to its metabolite, endoxifen, by the CYP2D6. The coculture system of breast cancer cells (MCF-7 cells) and the PHH-iPS-HLCs are illustrated. (D) The cell viability of MCF-7 cells was assessed after 72-h exposure to different concentrations of tamoxifen. (E) The cell viability of MCF-7 cells, which were cocultured with PHH-WT, PHH-NUL, HLC-WT, and HLC-NUL, was assessed after 72-h exposure to 500 nM of tamoxifen in the presence or absence of 3 nM quinidine (a CYP2D6 inhibitor). (F) The cell viability of MCF-7 cells cocultured with Ad-CYP2D6-transduced PHH-NUL and HLC-NUL was examined after 72-h exposure to 500 nM of tamoxifen. (G and H) The CYP2D6 expression (G) and activity (H) levels in Ad-CYP2D6-transduced PHH-NUL and HLC-NUL were examined by Western blotting and LC-MS/MS analysis. (I) The detoxification of desipramine-dependent conversion to its conjugated form by the CYP2D6. (J) The cell viability of PHH-WT, PHH-NUL, HLC-WT, and HLC-NUL was assessed after 24-h exposure to different concentrations of desipramine. (K) The cell viability of the PHH-WT and HLC-WT was assessed after 24-h exposure to 5 μM of desipramine in the presence or absence of 5 μM of quinidine (a CYP2D6 inhibitor). (L) The cell viability of the Ad-CYP2D6-transduced PHH-NUL and HLC-NUL was examined after 24-h exposure to 5 μM of desipramine. The cell viability was expressed as a percentage of that in the cells treated with only solvent. Data represent the mean ± SD from three independent experiments. In E and K, Student t test indicated that the cell viability in the “control” was significantly higher than that in the “quinidine” group ( $P < 0.01$ ). In F, H, and L, statistical significance was evaluated by ANOVA followed by Bonferroni post hoc tests to compare all groups. Groups that do not share the same letter are significantly different from each other ( $P < 0.05$ ).

in this study have the CYP2D6 poor metabolizer genotypes (CYP2D6 \*3, \*4, \*5, \*6, \*7, \*8, \*16, and \*21) (5). PHH8 and -11 have CYP2D6\*4 (null allele), whereas the others have a wild type (WT) or hetero allele (*SI Appendix*, Table S3 and Fig. 4A). Consistent with this finding, the PHH8/11-iPS-HLCs also have CYP2D6\*4, whereas the others have a wild type or hetero allele. As expected, the CYP2D6 activity levels in the PHH8/11 (PHH-NUL) and PHH8/11-iPS-HLC (HLC-NUL) were significantly lower than those in the PHH-WT and HLC-WT, respectively (Fig. 4B). The pharmacological activity of tamoxifen, which is the most widely used agent for patients with breast cancer, is dependent on its conversion to its metabolite, endoxifen, by the CYP2D6 (Fig. 4C). To examine whether the pharmacological activity of tamoxifen could be predicted by using PHHs and HLCs that have either the null type CYP2D6\*4 allele or wild-type CYP2D6 allele, the breast cancer cell line MCF7 was cocultured with PHHs or HLCs, and then the cells were treated with tamoxifen (Fig. 4D). The cell viability of MCF7 cells cocultured with PHHs-NUL or HLCs-NUL was significantly higher than that of MCF7 cells cocultured with PHHs-WT or HLCs-WT. The decrease in cell viability of MCF7 cells cocultured with PHHs-WT or HLCs-WT was rescued by treatment with a CYP2D6 inhibitor, quinidine (Fig. 4E). We also

confirmed that the cell viability of MCF7 cells cocultured with PHHs-NUL or HLCs-NUL was decreased by CYP2D6 overexpression in the PHHs-NUL or HLCs-NUL (Fig. 4F). Note that the expression (Fig. 4G) and activity (Fig. 4H) levels of CYP2D6 in CYP2D6-expressing adenovirus vector (Ad-CYP2D6)-transduced PHHs-NUL or HLCs-NUL were comparable to those of PHHs-WT or HLCs-WT. These results indicated that the PHHs-WT and HLCs-WT could more efficiently metabolize tamoxifen than the PHHs-NUL and HLCs-NUL, respectively, and thereby induced higher toxicity in MCF7 cells. Similar results were obtained with the other breast cancer cell line, T-47D (*SI Appendix*, Fig. S8 A–D). Next, we examined whether the CYP2D6-mediated drug-induced hepatotoxicity could be predicted by using PHHs and HLCs having either a null type CYP2D6\*4 allele or wild-type CYP2D6 allele. PHHs and HLCs were treated with desipramine, which is known to cause hepatotoxicity (Fig. 4I) (14). The cell viability of PHHs-NUL and HLCs-NUL was significantly lower than that of PHHs-WT and HLCs-WT (Fig. 4J). The cell viability of the PHHs-WT or HLCs-WT was decreased by treatment with a CYP2D6 inhibitor, quinidine (Fig. 4K). We also confirmed that the decrease in the cell viability of the PHHs-NUL or HLCs-NUL was rescued by CYP2D6 overexpression in the PHHs-NUL or HLCs-NUL (Fig. 4L). Similar

results were obtained with the other hepatotoxic drug, perhexiline (*SI Appendix, Fig. S8 E–H*). These results indicated that the PHHs-WT and HLCs-WT could more efficiently metabolize imipramine and thereby reduce toxicity compared with the PHHs-NUL and HLCs-NUL. Taken together, our findings showed that the interindividual differences in CYP metabolism capacity and drug responsiveness, which are prescribed by an SNP in genes encoding CYPs, were also reproduced in the PHH-iPS-HLCs.

## Discussion

The purpose of this study was to examine whether the individual HLCs could reproduce the hepatic function of individual PHHs. A Yamanaka 4 factor-expressing SeV vector was used in this study to generate integration-free human iPSCs from PHHs. It is known that SeV vectors can express exogenous genes without chromosomal insertion, because these vectors replicate their genomes exclusively in the cytoplasm (15). To examine the different cellular phenotypes associated with SNPs in human iPSC derivatives, the use of integration-free human iPSCs is essential.

We found that the CYP activity levels of the PHH-iPS-HLCs reflected those of parent PHHs, as shown in Fig. 3 A–C. There were few interindividual differences in the ratio of CYP expression levels in the PHH-iPS-HLCs to those in PHHs (*SI Appendix, Fig. S5*). Together, these results suggest that it is possible to predict the individual CYP activity levels through analysis of the CYP activity levels of the PHH-iPS-HLCs. In the future, it will be necessary to confirm these results in skin or blood cell-derived iPSCs as well as PHH-iPSCs, although donor-matched PHHs and blood cells (or skin cells) are difficult to obtain. In addition, the comparison of hepatic functions between genetically identical PHHs and PHH-iPS-HLCs (Fig. 3 A–C) would enable us to accurately ascertain whether the HLCs exhibit sufficient hepatic function to be a suitable substitute for PHHs in the early phase of pharmaceutical development. Because the drug responsiveness of the individual HLCs reflected that of individual PHHs (Fig. 3 E and F), it might be possible to perform personalized drug therapy following drug screening using a patient's HLCs. However, the R-squared values of the individual CYP activities differed from each other (Fig. 3 A–C), suggesting that the activity levels of some CYPs are largely

influenced not only by genetic information but also by environmental factors, such as dietary or smoking habits.

The interindividual differences of CYP2D6 metabolism capacity and drug responsiveness that were prescribed by SNP in genes encoding CYP2D6 were reproduced in the PHH-iPS-HLCs (Fig. 4). It was impossible to perform drug screening in the human hepatocytes derived from a donor with rare SNPs because these hepatocytes could not be obtained. However, because human iPSCs can be generated from such donors with rare SNPs, the CYP metabolism capacity and drug responsiveness of these donors might be possible to predict. Further, it would also be possible to identify the novel SNP responsible for an unexpected hepatotoxicity by using the HLCs in which whole genome sequences are known. We thus believe that the HLCs will be a powerful tool not only for accurate and efficient drug development but also for personalized drug therapy.

## Experimental Procedures

**DNA Microarray.** Total RNA was prepared from the PHH9-iPSCs, PHH9-iPS-HLCs, PHH9, and human hepatocellular carcinoma cell lines by using an RNeasy Mini kit. A pool of three independent samples was used in this study. cRNA amplifying, labeling, hybridizing, and analyzing were performed at Miltenyi Biotec. The Gene Expression Omnibus (GEO) accession no. for the microarray analysis is GSE61287.

**Flow Cytometry.** Single-cell suspensions of human iPSC-derived cells were fixed with 2% (vol/vol) paraformaldehyde (PFA) for 20 min, and then incubated with the primary antibody (described in *SI Appendix, Table S1*), followed by the secondary antibody (described in *SI Appendix, Table S2*). In case of the intracellular staining, the Permeabilization Buffer (eBioscience) was used to create holes in the membrane thereby allowing the antibodies to enter the cell effectively. Flow cytometry analysis was performed using a FACS LSR Fortessa flow cytometer (BD Biosciences).

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# Plasma Elevation of Vascular Endothelial Growth Factor Leads to the Reduction of Mouse Hematopoietic and Mesenchymal Stem/Progenitor Cells in the Bone Marrow

Katsuhisa Tashiro,<sup>1,\*</sup> Aki Nonaka,<sup>1,2,\*</sup> Nobue Hirata,<sup>1</sup> Tomoko Yamaguchi,<sup>1</sup>  
Hiroyuki Mizuguchi,<sup>3-6</sup> and Kenji Kawabata<sup>1,2</sup>

Vascular endothelial growth factor (VEGF) is reported to exhibit potent hematopoietic stem/progenitor cell (HSPC) mobilization activity. However, the detailed mechanisms of HSPC mobilization by VEGF have not been examined. In this study, we investigated the effect of VEGF on bone marrow (BM) cell and the BM environment by intravenous injection of VEGF-expressing adenovirus vector (Ad-VEGF) into mice. A colony assay using peripheral blood cells revealed that plasma elevation of VEGF leads to the mobilization of HSPCs into the circulation. Granulocyte colony-stimulating factor (G-CSF) is known to mobilize HSPCs by decreasing CXC chemokine ligand 12 (CXCL12) levels in the BM. However, we found almost no changes in the CXCL12 levels in the BM after Ad-VEGF injection, suggesting that VEGF can alter the BM microenvironment by different mechanisms from G-CSF. Furthermore, flow cytometric analysis and colony forming unit-fibroblast assay showed a reduction in the number of mesenchymal progenitor cells (MPCs), which have been reported to serve as niche cells to support HSPCs, in the BM of Ad-VEGF-injected mice. Adhesion of donor cells to the recipient BM after transplantation was also impaired in mice injected with Ad-VEGF, suggesting a decrease in the niche cell number. We also observed a dose-dependent chemoattractive effect of VEGF on primary BM stromal cells *in vitro*. These data suggest that VEGF alters the distribution of MPCs in the BM and can also mobilize MPCs to peripheral tissues. Taken together, our results imply that VEGF-elicited egress of HSPCs would be mediated, in part, by changing the number of MPCs in the BM.

## Introduction

**H**EMATOPOIETIC STEM CELLS (HSCs) sustain blood production throughout life. In a steady state, HSCs exist within the bone marrow (BM) and remain largely quiescent and self-renew at a low rate to avoid their exhaustion. By contrast, HSCs can actively proliferate, differentiate into progenitor cells, or egress from the BM into the circulation in some situations, such as tissue damage-induced cell death and increased plasma levels of hematopoietic cytokines, including the granulocyte colony-stimulating factor (G-CSF). These dynamic behaviors of HSCs are controlled by a local specific microenvironment called niches [1–8]. The non-hematopoietic cells, such as endosteal osteoblasts and perivascular mesenchymal progenitor cells (MPCs), are reported

to function as niche cells by supplying several HSC maintenance factors, including the CXC chemokine ligand 12 (CXCL12). Indeed, previous studies have shown that decreased levels of CXCL12 in the BM caused hematopoietic stem/progenitor cell (HSPC) mobilization, indicating the pivotal role of CXCL12 signaling in HSPC egress [9,10].

Not only is the vascular endothelial growth factor (VEGF) a well-known factor in angiogenesis, but it also plays an important role in the growth and differentiation of hematopoietic cells. Homozygous or heterozygous deletion of VEGF in mice leads to early embryonic lethality because of impaired vascular angiogenesis and hematopoiesis [11,12]. By conditional deletion of VEGF in hematopoietic cells, but not in stromal cells, Ferrara and colleagues clearly showed that VEGF is required for survival and repopulation

<sup>1</sup>Laboratory of Stem Cell Regulation, National Institute of Biomedical Innovation, Osaka, Japan.

Laboratory of <sup>2</sup>Biomedical Innovation and <sup>3</sup>Biochemistry and Molecular Biology, Graduate School of Pharmaceutical Sciences, Osaka University, Osaka, Japan.

<sup>4</sup>iPS Cell-Based Research Project on Hepatic Toxicity and Metabolism, Graduate School of Pharmaceutical Sciences, Osaka University, Osaka, Japan.

<sup>5</sup>Laboratory of Hepatocyte Regulation, National Institute of Biomedical Innovation, Osaka, Japan.

<sup>6</sup>The Center for Advanced Medical Engineering and Informatics, Osaka University, Osaka, Japan.

\*These two authors contributed equally to this work.