A Cell Culture Platform to Maintain Long-term Phenotype of Primary Human Hepatocytes and Endothelial Cells

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SUMMARY

We show that in contrast to functionally declining cocultures containing primary human hepatocytes and primary human liver sinusoidal endothelial cells, tricultures containing primary human hepatocytes, 3T3-J2 fibroblasts, and liver sinusoidal endothelial cells display stable hepatic and endothelial phenotypes for 3 weeks in vitro.

BACKGROUND AND AIMS: Modeling interactions between primary human hepatocytes (PHHs) and primary human liver sinusoidal endothelial cells (LSECs) in vitro can help elucidate human-specific mechanisms underlying liver physiology/disease and drug responses; however, existing hepatocyte/endothelial coculture models are suboptimal because of their use of rodent cells, cancerous cell lines, and/or nonliver endothelial cells. Hence, we sought to develop a platform that could maintain the long-term phenotype of PHHs and primary human LSECs.

METHODS: Primary human LSECs or human umbilical vein endothelial cells as the nonliver control were cocultivated with micropatterned PHH colonies (to control homotypic interactions) followed by an assessment of PHH morphology and functions (albumin and urea secretion, and cytochrome P-450 2A6 and 3A4 enzyme activities) over 3 weeks. Endothelial phenotype was assessed via gene expression patterns and scanning electron microscopy to visualize fenestrations. Hepatic responses in PHH/endothelial cocultures were benchmarked against responses in previously developed PHH/3T3-J2 fibroblast cocultures. Finally, PHH/fibroblast/endothelial cell tricultures were created and characterized as described previously.

RESULTS: LSECs, but not human umbilical vein endothelial cells, induced PHH albumin secretion for ~11 days; however, neither endothelial cell type could maintain PHH morphology and functions to the same magnitude/longevity as the fibroblasts. In contrast, both PHHs and endothelial cells displayed stable phenotype for 3 weeks in PHH/fibroblast/endothelial cell tricultures; furthermore, layered tricultures in which PHHs and endothelial cells were separated by a protein gel to mimic the space of Disse displayed similar functional levels as the coplanar tricultures.

CONCLUSIONS: PHH/fibroblast/endothelial tricultures constitute a robust platform to elucidate reciprocal interactions between PHHs and endothelial cells in physiology, disease, and after drug exposure. (Cell Mol Gastroenterol Hepatol 2018;5:187–207; https://doi.org/10.1016/j.jcmgh.2017.11.007)

Keywords: Micropatterned Cocultures; Tricultures; LSECs; HUVECs; 3T3-J2 Fibroblasts.
The parenchymal hepatocytes in the liver interact with liver sinusoidal endothelial cells (LSECs), which comprise ~70% of the total nonparenchymal cell (NPC) types in the liver.\(^1\) Separated from the hepatocytes by the space of Disse, a thin layer of extracellular matrix (ECM) proteins, LSECs contain numerous fenestrations arranged in sieve plates and play important roles in liver biology. For instance, LSECs clear waste molecules (including viruses) entering portal venous blood\(^1\); are important for induction of CD8\(^+\) T-cell tolerance\(^3\); synthesize the critical coagulation cofactor, factor VIII\(^1\); contribute to liver regeneration via increased production of cytokines (eg, hepatocyte growth factor)\(^1\); release exosomes that modulate antiviral activity in hepatitis C virus–infected hepatocytes in coculture\(^5\); can become insulin resistant and defenestrated in nonalcoholic fatty liver disease\(^6\); and experience toxicity to certain drugs.\(^7\) Therefore, developing novel liver models containing LSECs and hepatocytes can aid in drug discovery efforts for liver diseases and enable a better understanding of how reciprocal LSEC-hepatocyte interactions modulate cellular phenotypes in liver physiology.

Significant species-specific differences in drug metabolism and other liver pathways\(^8\)–\(^10\) necessitate the supplementation of animal data with human-relevant in vitro assays for drug development.\(^11\) Given their physiological relevance, isolated primary human hepatocytes (PHHs) are widely considered to be ideal for building human liver models. However, when cultured in the presence of ECM proteins (eg, collagen) alone, PHHs rapidly (hours to days) decline in critical phenotypic functions, such as cytochrome P-450 (CYP450) enzyme activities,\(^12\) insulin responsiveness,\(^13\) and expression of the master liver transcription factor, hepatocyte nuclear factor 4a.\(^14\) Similarly, when cultured alone, LSECs lose their characteristic fenestrations and undergo apoptosis within a few days.\(^15\) In contrast to hepatocyte monocultures, coculture with both liver- and nonliver-derived NPC types can enhance hepatocyte functions in culture.\(^16\) Endothelial cells have been previously explored toward transiently enhancing hepatocyte functions in cocultures relative to declining hepatocyte monocultures. However, many such hepatocyte-endothelial coculture studies use rodent cells\(^17\)–\(^21\) that do not completely suffice for modeling human liver biology. Furthermore, the use of abnormal cancerous cell lines\(^22\)–\(^24\) and/or nonliver endothelial cells\(^17\)–\(^21\),\(^25\) may provide a first approximation of hepatocyte-endothelial interactions but needs to be complemented with the use of primary cells from human liver tissue to determine similarities and differences in observed cell responses. Indeed, the Yarmush group has created in vitro cocultures of PHHs and primary human LSECs, which showed high level of low-density lipoprotein uptake in PHHs in the presence of LSECs\(^26\) and increased (~1.3-fold) hepatic CYP1A activity in serum-free coculture with endothelial cells under high (95%) oxygen levels.\(^27\) However, it is not clear from these short-term (≤24 hours) data sets whether LSECs can induce high levels of phenotypic functions in PHHs over long-term (weeks) culture as compared with PHH monocultures. Additionally, the differential effects of LSECs on PHH functions relative to nonliver vascular endothelial cells remain to be elucidated.

To address the limitations with the previously mentioned hepatocyte-endothelial coculture studies, here we sought to first elucidate the effects of primary human LSECs on the long-term functions of PHHs with comparisons to nonliver endothelial cells (human umbilical vein endothelial cells [HUVECs]) and PHH monocultures. We benchmarked the effects of endothelial cells on PHHs to the effects of 3T3-J2 murine embryonic fibroblasts, a cell type that expresses hepatocyte-supporting molecules present in the liver\(^28\) and is known to induce high levels of functions in PHHs for 4–6 weeks in vitro.\(^12\) To maintain consistent homotypic PHH contacts/interactions across the various coculture configurations toward isolating the effects of the heterotypic interactions between PHHs and NPCs, we used a previously developed semiconductor-driven micro-patterning technique to create so-called micropatterned cocultures in which PHHs are first clustered onto circular collagen-coated domains of precise/reproducible diameters and subsequently surrounded by an NPC layer containing 1 or more cell types.\(^29\) Lastly, we determined PHH/fibroblast/endothelial triculture configurations, including those in which the LSECs were separated from PHHs via an ECM protein gel as in the space of Disse in vivo, which promoted stable PHH and LSEC phenotype over several weeks toward developing a model system that has utility in drug development and to better understand PHH and LSEC interactions in liver physiology and disease.

# Materials and Methods

## Endothelial Culture

Primary human LSECs\(^3\) and TMNK immortalized liver endothelial cells\(^19\) were obtained from Dr. Hugo Rosen of the University of Colorado-Denver School of Medicine. A second donor of LSECs was purchased from ScienCell (Carlsbad, CA) and used to verify the trends observed with the first LSEC donor. Pooled-donor HUVECs were purchased from Lonza (Williamsport, PA). Primary endothelial cells were cultured at 37°C, 10% CO\(_2\) in EGM-2 BulletKit medium (Lonza) on tissue culture polystyrene coated with 2\(\mu\)g/cm\(^2\) fibronectin (Corning Life Sciences, Manassas, VA). TMNK cells were cultured at 37°C, 10% CO\(_2\) in a high-glucose Dulbecco’s modified eagle medium base (Corning) supplemented with 10% (vol/vol) bovine serum and 1% (vol/vol) penicillin-

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**Abbreviations used in this paper:** CD31, cluster of differentiation 31; CD54, cluster of differentiation 54; cDNA, complementary DNA; CYP450, cytochrome P-450; ECM, extracellular matrix; F8, factor VIII; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; HUVECs, human umbilical vein endothelial cells; LSECs, liver sinusoidal endothelial cells; NPCs, nonparenchymal cells; PHHs, primary human hepatocytes; SEM, scanning electron microscope; vWF, von Willebrand factor.

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streptomycin. Primary endothelial cell types were passaged no more than 6 times; before coculture with PHHs as described later, the endothelial cells were treated with 0.05% (m/vol) trypsin for 5 minutes to release cells into suspension, centrifuged, and resuspended in fresh culture medium.

**Micropatterned Coculture and Triculture Fabrication**

Cryopreserved PHHs were commercially obtained from Triangle Research Labs (Durham, NC); donors included HUM4011 (26-year-old white male who died of cardiac arrest) and HUM4055A (54-year-old white female who died of stroke). PHHs were thawed, counted, and viability was assessed as previously described. PHHs were subsequently micropatterned on collagen-coated domains as previously described. Briefly, adsorbed rat tail collagen I (Corning) was lithographically patterned in each well of a 24-well plate to create 500-μm diameter circular domains spaced 1200 μm apart, center-to-center. PHHs selectively attached to the collagen domains leaving ~30,000 attached PHHs on ~90 collagen-coated islands within each well of a 24-well plate. The next day after adherent PHHs had spread to fill in the collagen domains, the cultures were incubated with fibronectin (2 μg/cm²) to coat the remaining surface area with ECM protein to enable attachment of NPC types, which included either 3T3-J2 murine embryonic fibroblasts or endothelial cells at ~90,000 per well; the endothelial cells included were either HUVECs or primary human LSECs.

To create coplanar micropatterned tricultures, ~6000 endothelial cells (HUVECs or LSECs) were mixed into a suspension of ~84,000 3T3-J2 fibroblasts, and then this mixture was seeded onto micropatterned PHH colonies (~30,000 total PHHs) in each well of a 24-well plate. The 1:5 endothelial/PHH ratio corresponds to the approximate ratio in the liver. To create layered micropatterned tricultures, micropatterned cocultures containing PHH colonies surrounded by the fibroblasts were first overlaid with a thin gel (250 μg/mL) of Matrigel (Corning) and then ~6000 endothelial cells were seeded on top of the Matrigel the following day. Culture medium containing 40 ng/mL recombinant vascular endothelial growth factor (Thermo Fisher Scientific, Waltham, MA) in a high-glucose Dulbecco’s modified eagle medium base (Corning) was replaced on cocultures and tricultures every 2 days (300 μL/well). Other culture medium components were described previously.

**Gene Expression Analysis**

Total RNA was isolated, purified, and reverse transcribed into complementary DNA (cDNA) as previously described. Briefly, RNA was extracted from the cultures with the RNeasy kit (Qiagen, Germantown, MD) and homogenized via centrifugation through homogenizing columns (Omega Bio-Tek, Norcross, GA). Genomic DNA was removed with a 1-hour treatment with DNase I (New England Biolabs, Ipswich, MA), and cDNA was synthesized with the High-Capacity cDNA Reverse Transcription Kit (Applied
Biosystems, Foster City, CA). Then, 250 ng of cDNA was added to each quantitative polymerase chain reaction along with the Taqman master mix (Thermo Fisher Scientific) and predesigned Taqman human-specific primer/probe sets per manufacturer’s protocols; primer/probe sets were selected for endothelial genes including cluster of differentiation 31 (CD31), cluster of differentiation 54 (CD54), factor VIII (F8), and von Willebrand factor (vWF). The primer/probe sets were selected to be human-specific without cross-reactivity to 3T3-J2 mouse DNA; however, Taqman primer/probe sequences are proprietary to the manufacturer. Gene expression was normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH).

Cell Morphologic and Functional Assessments

Cell morphology was monitored using an EVOS FL microscope (Thermo Fisher Scientific) with standard 4×, 10×, and 20× phase contrast objectives. PHH projected area from phase contrast pictures was quantified using ImageJ software. Concentrations of
human albumin and urea in collected cell culture supernatants were assayed using previously published protocols. Briefly, albumin secretions were assessed using a competitive enzyme-linked immunosorbent assay, whereas urea production was measured via a colorimetric reaction with diacetyl monoxime, acid, and heat (kit from Stanbio Labs, Boerne, TX). CYP450 enzyme activities were measured by first incubating cultures in substrates for 1 hour at 37°C and then detecting either the luminescence or fluorescence of metabolites using previously described protocols. CYP2A6 activity was measured by the modification of coumarin to fluorescent 7-hydroxycoumarin (Sigma-Aldrich, St. Louis, MO), and CYP3A4 activity was measured by cleavage of luciferin-IPA into luminescent luciferin (Promega, Madison, WI). Live cultures containing PHHs were incubated with 2 μg/mL 5 (and 6)-carboxy-2',7'-dichlorofluorescein diacetate for 10 minutes in serum-free and phenol-free culture medium at 37°C, washed 3 times in serum-free and phenol-free culture medium, and imaged using the green fluorescent protein (470 nm excitation, 510 nm emission) light cube on the microscope to visualize functional bile canaliculi.

**Scanning Electron Microscopy**

Cultures were fixed in glutaraldehyde (Sigma-Aldrich) diluted to 2.5% (vol/vol) in phosphate-buffered saline (1X PBS, Corning) solution for 20 minutes at room temperature, and fixed cell cultures were washed 3 times with 1X PBS. Then, fixed samples were dehydrated through a series of washes using molecular-grade ethanol (Fisher Scientific, Pittsburgh, PA) to dry the samples thoroughly without damaging the cellular structures; washes were performed for 10 minutes each at 35%, 50%, 75%, 85%, 95%, 100%, and 100% (vol/vol) ethanol. Following the final
ethanol wash, cultures were dried chemically with hexamethyldisilizane (Alfa Aesar) for 10 minutes, excess hexamethyldisilizane was removed, and samples were air dried for 10 minutes at room temperature. Prepared samples were secured to a specimen mount using carbon adhesive tabs. A scanning electron microscope (SEM) coating unit E5100 Series II (Polaron Instruments, London, England) was used to sputter coat samples with a 4.125-nm-thick gold/palladium layer, and samples were then imaged using a Hitachi S-3000 variable pressure SEM (Tokyo, Japan).

Data Analysis

Each experiment was carried out in triplicate wells for each condition. Studies were repeated in 2 PHH and 2 LSEC donors to confirm trends. Data processing and visualization were performed using Microsoft Excel and GraphPad Prism (La Jolla, CA). Gene expression data were calculated as fold changes with respect to pure LSECs using the ΔΔCt method with GAPDH as the housekeeping gene. Statistical significance was determined with a 1- or 2-way analysis of variance followed by a Bonferroni pair-wise post hoc test ($P < .05$).

Results

Comparison of Primary Human Hepatocytes/Endothelial and Primary Human Hepatocytes/Fibroblast Cocultures

Primary human LSECs and primary HUVECs displayed prototypical endothelial morphology for up to 6 passages in vitro (Figure 1) and could be subsequently used for cocultivation with PHHs. Micropatterned cocultures of PHHs and endothelial cells (either LSECs or HUVECs) were compared with cocultures of PHHs and 3T3-J2 fibroblasts.
in such cocultures, 30,000 PHHs were surrounded by 90,000 NPCs. Both PHH/LSEC and PHH/HUVEC cocultures displayed a loss of prototypical hepatic morphology (i.e., polygonal shape, multinucleation, and visible bile canaliculi) over 2 weeks (Figure 2B). In contrast, PHH/fibroblast cocultures maintained hepatic morphology over at least 2 weeks. Micropatterned PHH monocultures spread out (dedifferentiated) as expected from a previous study.12

At the functional level, PHH/fibroblast cocultures outperformed PHH/endothelial cell cocultures. Albumin (Figure 3A) and urea secretions (Figure 3B), CYP3A4 activity (Figure 3C), and CYP2A6 activity (Figure 3D) in PHH/endothelial cell cocultures had steady-state values that were ~5%–20% of the values in PHH/fibroblast cocultures. The PHH/fibroblast cocultures maintained stable levels of all measured functions over 3 weeks, whereas most functions in PHH/endothelial cell cocultures declined over time. Modulating the PHH:endothelial ratio to 5:1 (physiologic32) or 1:4 in cocultures did not rescue the PHH phenotype (data not shown). Furthermore, cocultures created using a second donor of LSECs followed similar trends in albumin secretions, urea secretions, CYP3A4 activity, and morphology (Figure 4). Nonetheless, PHH/LSEC cocultures had statistically higher albumin secretions than PHH/HUVEC cocultures (~3.7-fold on Day 7 and ~10-fold on Day 11).

Figure 5. PHH/endothelial cell cocultures created using the immortalized human liver endothelial cell line (TMNK) relative to PHH/fibroblast control cocultures. Cocultures were created as depicted in Figure 2A (all culture models shown contained micropatterned PHHs) followed by an assessment of PHH morphology in PHH/TMNK cocultures after 1 week (A) and 2 weeks (B). PHH morphology in PHH/fibroblast cocultures is shown in Figure 2B. Scale bar = 400 μm. Albumin (C) and urea (D) secretions were also measured from the PHH/TMNK and PHH/fibroblast cocultures over time. Error bars represent standard deviations (n = 3 wells).
and PHH monocultures (undetectable levels) until 11 days in culture, whereas urea secretion and CYP450 enzyme activities were statistically similar (Figure 3). Finally, immortalized human liver endothelial cells (TMNK line), previously used for coculture with PHHs, were also not able to stabilize PHH morphology or functions (Figure 5).

The effects of endothelial cells on PHH functions in cocultures were observed irrespective of the culture medium formulation because subjecting the PHH/endothelial cocultures to endothelial culture medium (as that used for expanding pure endothelial cells) or hepatocyte culture medium or a 1:1 mixture of the 2 types of media did not enable long-term rescue of the PHH phenotype (data not shown); thus, supplementing the hepatocyte medium with vascular endothelial growth factor, a previously developed hepatocyte-endothelial culture medium, was used for the remainder of the studies. Therefore, LSECs, but not HUVECs, can induce transient and statistically significant albumin secretion in PHHs for ~11 days; however, neither endothelial cell type can induce functions to the same magnitude and stability as the 3T3-J2 fibroblasts irrespective of PHH/endothelial ratio and medium formulation.

**Hepatic Phenotype in Coplanar Primary Human Hepatocytes/Fibroblast/Endothelial Tricultures**

As opposed to PHH/endothelial cocultures, PHH/fibroblast/endothelial tricultures were seeded as shown in Figure 6A. Briefly, after PHHs attached and spread on micropatterned collagen-coated domains, a mixture of endothelial cells (LSECs or HUVECs) and 3T3-J2 fibroblasts was seeded and filled the remaining area around the PHH colonies. A PHH/endothelial cell ratio of 5:1 was selected to match that observed in vivo. Over the course of 3 weeks, PHH/fibroblast/LSEC and PHH/fibroblast/HUVEC tricultures displayed a hepatic morphology comparable with that observed in PHH/fibroblast cocultures with respect to morphology in the PHH monocultures. All scale bars = 400 μm.

**Figure 6.** Morphology of PHH/fibroblast/endothelial cell tricultures (containing either LSECs or HUVECs) relative to PHH/3T3-J2 fibroblast control cocultures and PHH monocultures. (A) Schematic depicting the creation of the tricultures; cocultures were created as depicted in Figure 2A. (B) Morphology of tricultures models over the course of 2 weeks in comparison with the PHH/fibroblast cocultures and pure PHH monocultures. Note the prototypical PHH morphology (ie, polygonal shape, multinucleation, and presence of visible bile canaliculi) in the tricultures/cocultures and spread-out (dedifferentiated) morphology in the PHH monocultures. All scale bars = 400 μm.
multinucleation, polygonal shape, and the presence of visible bile canaliculi between hepatocytes (Figure 6B). At the functional level, when treated with the bile canaliculi stain, 5 (and 6)-carboxy-2′,7′-dichlorofluorescein diacetate (substrate for transporters, multidrug resistance-associated proteins 2/335), adjacent PHHs in the tricultures displayed functional bile canaliculi as in the PHH/fibroblast cocultures, whereas PHH monocultures displayed little to no functional bile canaliculi (Figure 7). Furthermore, PHH/fibroblast/endothelial cell tricultures secreted albumin (Figure 8A) and urea (Figure 8B) at steady-state levels of \( \sim 1.4 \mu g/h/10^6 \) cells and \( \sim 12 \mu g/h/10^6 \) cells, respectively, for 3 weeks; these secretion levels were statistically similar to the levels measured in PHH/fibroblast cocultures. Similarly, CYP3A4 (Figure 8C) and CYP2A6 (Figure 8D) enzyme activities were statistically similar across the tricultures and PHH/fibroblast cocultures. However, PHH monocultures displayed a severe decline in functions (Figure 8). Furthermore, the previously mentioned morphologic and functional trends in tricultures relative to cocultures were also observed with a second primary LSEC donor (Figure 9). However, in contrast to primary human LSECs and primary HUVECs, the use of the immortalized TMNK line in tricultures with fibroblasts did not lead to stable PHH morphology or functions (Figure 10), likely because of overgrowth of the TMNK cells.

**Endothelial Phenotype in Coplanar Primary Human Hepatocytes/Fibroblast/Endothelial Tricultures**

The presence of endothelial cells in tricultures over time was confirmed via gene expression analysis. Specifically, CD31 (Figure 11A) and CD54 (Figure 11B) gene expression levels increased in PHH/fibroblast/endothelial cell tricultures, whereas F8 (Figure 11C) and vWF (Figure 11D) expression levels were relatively stable in tricultures over the course of 3 weeks. In PHH/endothelial cell cocultures, all the previously mentioned gene expression markers were detected over 3 weeks; however, similarities and differences with the tricultures were observed. CD31 gene expression in PHH/endothelial cell cocultures was generally lower than tricultures over 3 weeks, CD54 gene expression in cocultures was higher than tricultures, F8 gene expression in PHH/LSEC cocultures was lower than PHH/fibroblast/LSEC tricultures but similar between PHH/HUVEC cocultures and PHH/fibroblast/HUVEC tricultures, and vWF expression in cocultures was higher than tricultures. We also verified using SEM the presence of fenestrations on LSECs in pure cultures (Figure 12A), when cocultured with 3T3-J2 fibroblasts but without PHHs (Figure 12B), in PHH/LSEC cocultures (Figure 12C), and in PHH/fibroblast/LSEC tricultures (Figure 12D). LSECs displayed noticeable fenestrations regardless of the culture conditions.
The previously mentioned endothelial characterization results show that both LSECs and HUVECs display prototypical markers over several weeks in both cocultures and triculture configurations; however, the PHH phenotype is enhanced only in the presence of the fibroblasts in cocultures and tricultures.

**Characterization of Layered Primary Human Hepatocytes/Fibroblast/Endothelial Tricultures**

To mimic the space of Disse, we created a layered version of the triculture model by overlaying a PHH/fibroblast coculture with a thin gel (250 μg/mL) of Matrigel and subsequently seeding endothelial cells (LSECs or HUVECs) on top of the gel (Figure 13A). As in the PHH/fibroblast/endothelial cell coplanar triculture model, PHHs in the layered configuration maintained prototypical hepatocyte morphology with polygonal shape, multinucleation, and visible bile canaliculi between adjacent PHHs (Figure 13B). However, micropatterned PHHs overlaid with Matrigel but without any NPC types deteriorated in morphology.

At the functional level, PHH/fibroblast/endothelial cell layered tricultures secreted albumin (Figure 14A) and urea (Figure 14B) at steady-state levels of ~1.4 μg/h/10^6 cells and ~12 μg/h/10^6 cells, respectively, for 3 weeks; these secretion levels were statistically similar to the levels measured in control PHH/fibroblast cocultures with a Matrigel overlay. Similarly, CYP3A4 (Figure 14C) and CYP2A6 (Figure 14D) enzyme activities were ~20 RLU/h/10^6 cells and ~32 μM 7-hydroxycoumarin/h/10^6 cells for both PHH/fibroblast/endothelial cell layered tricultures and PHH/fibroblast cocultures with a Matrigel overlay. Overall, the layered tricultures were highly similar in hepatic functional capacity compared to control cocultures.

**Figure 8.** Hepatocellular functions in PHH/fibroblast/endothelial cell tricultures (containing either LSECs or HUVECs) relative to PHH/3T3-J2 fibroblast control cocultures and PHH monocultures. Cocultures and tricultures were created as depicted in Figures 2A and 6A (all culture models shown contained micropatterned PHHs), respectively, followed by an assessment of hepatic functions over time, including albumin secretion (A), urea secretion (B), CYP3A4 enzyme activity (C), and CYP2A6 enzyme activity (as measured by the production of 7-HC) (D). Error bars represent standard deviations (n = 3 wells).
morphology and functional levels over several weeks as the coplanar tricultures, suggesting that separating the endothelial cells from PHHs did not lead to higher induction of hepatic phenotype (compare Figures 6 and 8 with Figures 13 and 14).

**Comparison of Primary Human Hepatocytes Projected Surface Area in Various Culture Formats**

To confirm qualitative observations of PHH spreading out or lack thereof in various culture formats, we quantified the projected surface area of PHHs cultured in all models developed here (Figure 15). PHHs in monocultures or in PHH/endothelial co-cultures tended to spread out and occupy more projected surface area. However, PHHs in cultures containing 3T3-J2 fibroblasts (PHH/fibroblast cocultures, PHH/fibroblast/endothelial coplanar tricultures, or PHH/fibroblast/endothelial layered tricultures) maintained their polygonal shape and occupied less projected surface area.

**Discussion**

In this study, we developed the first-of-its-kind cell culture platform that induces high and stable levels of phenotypic functions in both PHHs and primary human LSECs over the course of several weeks. We initially created cocultures of PHHs and LSECs, while using HUVECs as the non-liver-endothelial cell control; micropatterning was used to control PHH homotypic interactions toward isolating the effects of PHH/NPC interactions. Fibronectin was used here for enabling endothelial cell attachment because this ECM protein has been shown previously to facilitate the
attachment of LSECs.\textsuperscript{15} However, fibronectin is not amenable to the plasma ablation micropatterning technique used here (data not shown). Therefore, we used adsorbed collagen I for creating micropatterned PHH colonies as in previous studies\textsuperscript{29,36}; the use of collagen I for PHH attachment is also widespread in the field of PHH culture.\textsuperscript{11} Hepatic albumin secretion was statistically higher in PHH/LSEC cocultures for \( \sim 11 \) days than in PHH/HUVEC cocultures and PHH monocultures. However, the effects of both endothelial cell types on PHH phenotype were transient and functions declined over time irrespective of PHH-to-endothelial ratios and medium formulations tested.

Our observations with PHH/endothelial cocultures are not entirely consistent with previously published data in cocultures of rat hepatocytes and endothelial cells, which showed relatively stable functions for several weeks. For instance, the Noh group has shown that primary rat hepatocytes displayed relatively stable urea secretion for \( \sim 30 \) days when cocultivated with immortalized bovine aortic endothelial cells.\textsuperscript{17,21} In contrast, our use of an immortalized human liver endothelial cell line (TMNK) did not lead to induction of high and stable PHH functions; we suspect that species-specific differences may be important in the discrepancies observed across the 2 studies, although we cannot entirely rule out the differences between the use of endothelial cells from different organ systems. In a more directly comparable study using primary rat hepatocyte and primary rat LSEC cocultures, Bale et al\textsuperscript{18} showed relatively stable albumin and urea secretions, and CYP1A activity over 4 weeks. Some key differences with our study include the use of high concentration (~1 mg/mL) collagen I gels by Bale et al\textsuperscript{18} to sandwich hepatocytes and LSECs in various layers and species-specific differences in hepatocyte-LSEC interactions. Here, we did not use high concentration collagen I gels because they are (1) difficult to miniaturize into 24-well and smaller plate formats (ie, tend to be inconsistent in gelation height throughout the well and tend to peel off over a few days because of cell contraction), (2) can bind drugs when being used for screening assays, and (3) may represent a more fibrotic state of the liver in which the diverse collagens in the liver get replaced with high levels of collagen I.\textsuperscript{37,38} Thus, because of such limitations, collagen I gel–based liver models are not routinely used in pharmaceutical practice.\textsuperscript{39,40} Ultimately, our approach/model is human-specific and is more amenable to high-throughput applications in the drug development pipeline.

In contrast to declining functions in PHH/endothelial cocultures, all the PHH functions measured (albumin and urea secretions, and CYP2A6 and CYP3A4 enzyme activities) were significantly higher and stable over 3 weeks in cocultures of PHHs and 3T3-J2 murine embryonic fibroblasts. This mouse fibroblast cell line can induce functions in PHHs at levels closer to those observed in freshly isolated PHHs.
from the same donors.\textsuperscript{41} More broadly, coculture with both liver- and nonliver-derived NPC types, including those of mouse 3T3 fibroblast origin,\textsuperscript{28,42–48} has been long known to induce functions in primary hepatocytes from multiple species, including humans, which suggests that the molecular mediators underlying the coculture effect are relatively well-conserved across species.\textsuperscript{18,49} Although the complete mechanism underlying the effects of 3T3-J2 fibroblasts on PHHs remains to be elucidated, 3T3-J2 fibroblasts express various molecules found in the liver, such as decorin,\textsuperscript{28} vascular endothelial growth factor-D,\textsuperscript{28} and T-cadherin,\textsuperscript{50} which have been implicated in the ability of these fibroblasts to induce functions in hepatocytes from multiple species.\textsuperscript{40} Furthermore, 3T3-J2 fibroblasts have a lack of detectable liver functions, display contact inhibition that avoids overgrowth, and are propagated easily before inclusion in coculture.\textsuperscript{12,16,28,51} More importantly, the use of 3T3-J2 fibroblasts in coculture with PHHs does not prevent

**Figure 11.** Endothelial gene expression patterns in different culture models. PHH/endothelial cell cocultures and PHH/fibroblast/endothelial cell tricultures were created as depicted in Figures 2A and 6A, respectively, followed by evaluation of endothelial (HUVECs or LSECs) gene expression patterns over time using reverse-transcription quantitative polymerase chain reaction, including $\text{CD31}$ (A), $\text{CD54}$ (B), $\text{F8}$ (C), and $\text{vWF}$ (D). Data plotted are fold changes with respect to pure LSECs calculated with the $\Delta\Delta C_T$ method using $\text{GAPDH}$ as the housekeeping gene. Error bars represent standard deviations ($n = 3$ wells). $^* P < .05$, $^** P < .01$, and $^*** P < .001$ between the coculture or triculture condition relative to pure LSECs.
the effective use of the stabilized PHHs for many applications in drug development, such as drug clearance prediction,\textsuperscript{36,52} drug metabolite identification,\textsuperscript{53} drug-transporter interactions,\textsuperscript{54} drug hepatotoxicity,\textsuperscript{16} hepatitis B/C viral infections,\textsuperscript{25,56} malaria infection,\textsuperscript{37,28} and steatosis and insulin resistance caused by hyperglycemia as in diabetes.\textsuperscript{13,31}

We used the previously mentioned coculture effect here to create a triculture model in which 3T3-J2 fibroblasts were used to stabilize PHHs to functional levels closer to physiological outcomes than possible with endothelial cells (LSECs or HUVECs), and endothelial cells were introduced within the fibroblast monolayer at a physiologic ratio (1 endothelial cell:5 PHHs) to allow PHH/endothelial interactions as \textit{in vivo}. Tricultures were created in both coplanar (ie, all 3 cell types could interact via paracrine and contact signaling) and layered (separation of PHHs and LSECs via a gelled Matrigel layer to mimic the space of Disse) configurations. Matrigel is widely used to overlay PHHs with an ECM gel because it has diverse components, many of which are present in the liver (eg, different collagens types instead of collagen I alone and laminins).\textsuperscript{11,59–61} The projected surface area, functions (albumin and urea secretions, and CYP3A4/2A6 enzyme activities), and active bile canaliculi of PHHs in both triculture configurations were remarkably similar to those observed in PHH/fibroblast cocultures, suggesting that the ability of the fibroblasts to induce and stabilize functions in PHHs is not compromised by inclusion of endothelial cells; such an approach enables a well-differentiated PHH phenotype independently (via the 3T3-J2 fibroblast support) of the PHHs’ ability to interact with endothelial cells. Both the coplanar and layered tricultures were statistically similar with respect to PHH functions, suggesting that the \textit{in vivo}–like separation between endothelial cells and PHHs does not induce greater levels of functions than when the cell types can interact via both paracrine and contact signaling. We selected the coplanar triculture configuration for all other studies because the lack of protein gels is preferred for drug development applications as described previously. Nonetheless, the layered configuration can be highly useful when subjecting the tricultures to microfluidic perfusion in liver-on-a-chip platforms because it exposes the endothelial cells to shear stress while protecting the PHHs from shear stress as \textit{in vivo}.

Although we are the first group to our knowledge to create a human-relevant triculture platform using PHHs, fibroblasts, and primary human endothelial cells (LSECs and HUVECs), our findings are consistent with triculture data obtained in rat liver platforms. For instance, March et al\textsuperscript{15} combined primary rat hepatocytes and primary rat LSECs with 3T3 fibroblasts on a mechanically actuated dynamic substrate; this triculture model better preserved the phenotype of hepatocytes and fenestrations of LSECs relative to cultures without fibroblasts. In another study, Liu et al\textsuperscript{62} found that their triculture system containing primary rat hepatocytes with 3T3 fibroblasts and HUVECs functionally outperformed the hepatocyte/endothelial coculture control. These findings suggest that 3T3 fibroblasts can stabilize hepatocyte and endothelial functions across rodent and human species, which bodes well for use...
of hepatocyte/fibroblast/endothelial cell tricultures for elucidating specific-specific mechanisms underlying physiological and pathophysiological phenomena. However, for drug development applications, well-documented differences in drug metabolism and toxicity pathways between rodents and humans necessitate the use of human-relevant liver models; our platform now provides the avenue for predicting the effects of drugs that act on PHHs and/or primary human LSECs.

The sourcing of endothelial cells is a major consideration for building in vitro models of the human liver. Ideally, freshly isolated LSECs from human liver tissue would be used in all applications because they are the closest representation of human liver physiology; however, the routine use of this gold standard cell type is not practical for drug screening applications that necessitates creation of on-demand cultures from the same donors via the use of cryopreserved cells toward mitigating donor-to-donor variability when testing a large number of compounds longitudinally. Because of such limitations with freshly isolated primary human LSECs, most other groups rely on either immortalized human endothelial cells or endothelial cells from other species and/or organ systems when developing culture platforms for drug development. Several studies have cocultivated hepatocytes with the TMNK immortalized human liver endothelial cells because of their ease of propagation; however, here we show that in contrast to primary endothelial cells, TMNK cells cause a severe decline in PHH morphology and functions with or without fibroblasts, likely because of overgrowth. We were able to passage primary human LSECs from multiple donors at least 6 times and use them subsequently in coculture and

![Figure 13. Morphology of layered PHH/fibroblast/endothelial cell tricultures (containing either LSECs or HUVECs) relative to PHH/3T3-J2 fibroblast control cocultures and PHH monocultures. (A) Schematic depicting the creation of the layered tricultures; PHH/fibroblast control cocultures and PHH monocultures were created as depicted in Figure 2A and subsequently overlaid with Matrigel to account for the effects of the protein gel on PHH functions. (B) Morphology of layered tricultures models over the course of 2 weeks in comparison with the Matrigel-coated PHH/fibroblast cocultures and Matrigel-coated pure PHH monocultures. Note the prototypical PHH morphology (ie, polygonal shape, multinucleation, and presence of visible bile canaliculi) in the tricultures/cocultures and spread-out (dedifferentiated) morphology in the PHH monocultures. All scale bars = 400 μm.](image-url)
triculture studies. Nonetheless, because it is not trivial to commercially source primary human LSECs from more than a few donors, we also evaluated the effects of HUVECs, which are readily available from many donors and have a precedence for use in cocultures with primary hepatocytes and stem cell–derived hepatocyte-like cells.22,67–70 Our results here indicate that PHH/fibroblast/endothelial cell tricultures containing HUVECs display similar levels and longevity of PHH functions over 3 weeks, and thus such tricultures can serve as a first approximation when modeling PHH/endothelial interactions, whereas primary human LSECs can be used in select studies to elucidate similarities and differences with reciprocal interactions between PHHs and nonliver-endothelial cells.

To determine endothelial phenotype in tricultures, we first evaluated gene expression of CD31, CD54, F8, and vWF, because these markers have been shown to be consistently expressed in human liver slices and isolated LSECs.71–74 CD31 is a member of the immunoglobulin superfamily known to be expressed in many types of endothelial cells, but specifically in the cytoplasm of LSECs.67,75 CD54 is a member of the immunoglobulin superfamily expressed on the surface of endothelial cells and implicated in various signaling pathways, including some immune pathways.76,77 vWF is a multimeric glycoprotein that mediates platelet adhesion and thrombus formation during vascular injury,78 whereas F8 is a coagulation factor that is carried by vWF in circulating blood that leads to normal arrest of bleeding and thrombus formation.78,79 All of the previously mentioned gene expression markers were detected in both PHH/fibroblast/LSEC tricultures and PHH/fibroblast cocultures.

Figure 14. Hepatic functions in layered PHH/fibroblast/endothelial cell tricultures (containing either LSECs or HUVECs) relative to PHH/3T3-J2 fibroblast control cocultures and PHH monocultures. Matrigel-coated tricultures and cocultures were created as described in Figure 13, followed by an assessment of hepatic functions over time, including albumin secretion (A), urea secretion (B), CYP3A4 enzyme activity (C), and CYP2A6 enzyme activity (as measured by the production of 7-HC) (D). Error bars represent standard deviations (n = 3 wells). *P < .05, **P < .01, and ***P < .001 between the PHH/fibroblast/LSEC tricultures and PHH/fibroblast cocultures.
the exchange of soluble and particulate material between the blood and the space of Disse.80,81 The pattern of fenestration has been extensively studied in rat and mouse LSECs,15,81–84 but has been shown to be similar in human LSECs.85 The distribution of fenestrations is highly dynamic, both with respect to location in the liver and grouping with adjacent fenestrations.81,86 Furthermore, the number and diameter of fenestrations have been shown to correlate strongly with liver conditions (eg, fatty liver, hepatitis, and hepatectomy) and exposure to xenobiotic and environmental agents (eg, ethanol, nicotine).81 It has also been suggested that the diameter of fenestrations affects the uptake and transport of lipoproteins in the sinusoid, which may greatly impact the pathogenesis of atherosclerosis.81 In this study, we detected the presence of fenestrations on LSECs via SEM in pure cultures, PHH/LSECs cell cocultures, and PHH/fibroblast/LSEC tricultures; however, we were not able to elucidate quantitative differences across the culture configurations using SEM analysis. Nonetheless, coupled with gene expression analysis, the presence of fenestrations suggests that both cocultures and tricultures can maintain the survival of endothelial cells irrespective of functional levels in PHHs. However, in contrast to PHH/endothelial cocultures that display declining PHH functions, tricultures are better suited to evaluate reciprocal interactions between stable endothelial cells and PHHs in physiologic and pathophysiologic conditions. For instance, our tricultures could be used to model in vivo–like paracrine signaling between hepatitis C virus–infected PHHs and LSECs, because infected LSECs are known to release exosomes and inhibit viral replication in infected hepatocytes, albeit from a cancerous origin.85 Furthermore, some drugs are known to cause toxicity to LSECs,7,87 which can lead to downstream effects in PHHs because of the release of apoptotic factors from the LSECs; our tricultures can serve to elucidate such crosstalk following drug exposure.

Our goal in this study was to determine how primary human endothelial cells (LSECs and HUVECs) affect long-term PHH functions relative to 3T3-J2 murine embryonic fibroblasts and then construct a triculture platform that can enable stable phenotypes of both PHHs and endothelial cells for several weeks. However, other NPC types in the liver, such as hepatic stellate cells and Kupffer cells/macrophages, also interact with PHHs and LSECs in vivo. The use of the 3T3-J2 fibroblasts to stabilize PHH functions allows us to introduce specific liver NPC types within and around the fibroblast monolayer to study interactions with PHHs, whereas the use of PHH micropatterning allows us to control for PHH homotypic contacts that are critical for establishment of hepatocyte polarity, such as the formation of cell junctions (eg, cadherins) and bile canaliculi. Indeed, Nguyen et al88 showed that primary human Kupffer cells can be cultured atop pre-established micropatterned cocultures containing PHHs and fibroblasts to study the effects of Kupffer cell activation on hepatic CYP450s. Similarly, Davidson et al89 showed that primary human activated (fibrogenic) hepatic stellate cells can be cultured within the fibroblast monolayer surrounding the PHH micropatterned colonies to model an early nonalcoholic steatohepatitis-like phenotype in the PHHs, which could be alleviated with clinically relevant drugs. Thus, PHH/fibroblast/liver NPC triculture configurations offer robust in vitro tools to elucidate reciprocal interactions between PHHs and liver NPCs in physiological/disease contexts and for drug screening.

In conclusion, we showed here that neither primary human LSECs nor nonliver HUVECs can stabilize the PHH phenotype over several weeks, which necessitated the use...
of 3T3-J2 fibroblasts in a PHH/fibroblast/endothelial cell triculture configuration that subsequently led to high levels of prototypical hepatic functions and endothelial phenotype for at least 3 weeks in vitro. Separating the endothelial cells from the PHHs via a thin protein gel (Matrigel) to mimic the space of Disse as in vivo did not lead to better PHH functions than the coplanar tricultures, although both configurations can have utility for addressing specific liver-specific hypotheses as detailed previously. Ultimately, coupling PHH/fibroblast/liver NPC tricultures with cultures created from other tissues on a microfluidic chip (ie, body-on-a-chip) will allow a systems-level exploration of disease progression and the effects of drugs on multiple interacting organ systems.

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