Primary Human Hepatocytes 3D in vitro Culture Model for Studying Hepatic Function

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INTRODUCTION
Primary Human Hepatocyte (PHH) culture provides the closest in vitro model to human liver that can produce a metabolic profile of a given drug very similar to that found in vivo. Hence, PHH culture is the gold standard for studying the in vitro hepatic biology, liver function, and drug induced hepatotoxicity. The conventional way of culturing PHH in 2-dimension (2D) has major pitfalls. The PHH rapidly de-differentiate and lose the specific functions in a week. Therefore, there is a need for more robust in vitro models that reflect in vitro liver biology more accurately and maintain the liver functions for a longer time. 3-dimensional (3D) hepatic in vitro models have gained a lot of attention for their ability to recapitulate the hepatic function and greater longevity.

Recently we have developed an easy-to-assemble user-friendly in vitro Primary Human Hepatocyte (PHH)-3D spheroid model. The 3D-hepatic spheroids are viable for at least 4 weeks in culture and remain phenotypically stable, retaining the hepatocyte-specific functions.

MATERIALS AND METHODS

Spheroid culture
Hepatic spheroids were formed using Gibco™ cryopreserved spheroid-qualified human hepatocytes (Catalog No. HMCP-50) following the user guide [1]. Each well contained PHHs between 750 and 7500 hepatocytes, depending on the experimental conditions. The spheroids formed within 5 days of cell seeding. Starting on day 5, half of the plating medium was changed every 48–72 hours. Following this culture condition the 3D hepatic spheroids can remain viable at least up to 4 weeks.

Metabolic assay
As the liver is the primary site of metabolism for most drugs, primary hepatocytes are the most popular in vitro tool to evaluate hepatic drug metabolism. However, the efficiency of 3D hepatic spheroids for studying drug metabolism is relatively unknown. In order to study the biotransformation of various drugs by Cytochrome P450 (CYP) enzymes in hepatic spheroids, 3000 PHH were seeded in Nunc™ Spherih™ plate and cultured as described above. On Day 9 of the 3D culture 2D hepatocytes were seeded using 80,000 PHH per well as described in the user manual [2]. On day 10, six test articles were evaluated in both 2D and 3D PHH cultures in serum-free Gibco™ William’s E Medium. The compounds were selected such that several CYP enzymes that are important for hepatic drug metabolism could be interrogated. Table 1 lists the identity of each compound and the CYP enzymes primarily responsible for their metabolism, the metabolites analyzed, and the drug concentrations tested. Cell culture samples collected from both 2D and 3D cultures were analyzed for metabolite formation using the Thermo Scientific™ Exactive™ Plus Hybrid Quadrupole-Orbitrap™ Mass Spectrometer.

RESULTS

Figure 1. Work Flow of self assembly and characterization of primary hepatocyte into 3D-spheroid

Figure 2. Optimization of Hepatic Spheroids: Variable Cell Numbers.

Figure 3. Comparison of Spheroid volumes with variable number of cells.

Figure 4. TUNEL Staining of Hepatic Spheroid Sections with Variable Number of Cells

Parafin fixed sections of hepatic spheroids collected at day 10 with variable number of cells were H&E stained and micrographed at 20X optical zoom. These sections also underwent in situ cell death detection (DNA strand breaks) using a fluorometric Click-iT™ TUNEL Assay protocol (Invitrogen C10617). The red fluorescence denotes apoptotic cells, which seemed to be homogenously distributed throughout the spheroids of different sizes.

Figure 5. Elevated Gene Expressions 3D Hepatic Cultures compared to conventional 2D culture.

The metabolites quantified using HRMS were converted to mole amounts based on the standard curves of the respective metabolites. Two different lots of PHH, HuB28X and HuB28X (partially redacted), were used in this assay. Results were normalized to incubation time of individual substrates and number of cells per well in the 2D and 3D cultures. Data is the mean ± SD, n = 3.

CONCLUSIONS

• Gibco™ Primary Human Hepatocytes (HMCP-50) can easily be assembled into a 3D culture in 5 days using either Gibco™ Hepatic Spheroid Kit (A41390) or Nunc™ Spherih™ low attachment U-bottom 96-well Microplates, Gibco™ plating media and plating supplements.

• The Primary Hepatic Spheroids are functionally viable for at 4-weeks, which is a significant progress in primary hepatocyte culture considering the conventional 2D-culture methods.

• The 3D hepatocyte culture requires a significantly lower number of cells than that of the 2D counterpart, which opens new possibilities for high throughput assays using PHH.

• 3D hepatocyte culture is superior with respect to gene expression and metabolic activities compared to conventional 2D hepatic culture system.

FUTURE DIRECTIONS
• Assessment of in vitro CYP induction in 3D Hepatic Spheroids.
• Coculture of PHH with non-parenchymal cells to establish in vitro liver model.
• 3D culture of PHH isolated from diseased livers (such as NAFLD, NASH and hepatic fibrosis).

REFERENCES

Figure 6. Formation of Bile Ducts in Hepatic Spheroids.

Bile duct formation in the primary human hepatic spheroids was analyzed by treating 7-day old spheroids with bile tracer CFDA. HepG2 spheroids were also treated with CFDA as negative control.

Figure 7. Higher Temporal CYP3A4 activity in 3D hepatic cultures.

CYP3A4 activity was measured using P450 ProGo assay (Promega). Cyp3A4 activity of the 3D culture was found to be higher than that of 2D culture, indicating higher metabolic function of 3D culture. Results are mean ± SD, n = 3.

Figure 8. Higher metabolic activities in 3D cultures compared to conventional 2D hepatic culture system.

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