Hepatocyte culture in a radial-flow bioreactor with plasma polypyrrole coated scaffolds

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ABSTRACT: We have designed and evaluated a radial-flow bioreactor for three-dimensional liver carcinoma cell culture on a new porous coated scaffold. We designed a culture chamber where a radial flow of culture medium was continuously delivered through it. Once this system was established, flow was simulated using flow dynamics software based on numeric methods to solve Navier-Stokes flow equations. Perfusion within cell culture scaffolds was simulated using a flow velocity of 7 mL/min and found that cell culture medium was distributed unhindered in the bioreactor chamber. Afterwards, the bioreactor was built according to the simulated design and was tested with liver carcinoma cells (HepG2) cultured over an L-polylactic acid scaffold whose surface was modified with iodine-doped polypyrrole. The bioreactor was tested under non-flow and in radial flow conditions. Cell density under radial flow conditions was almost double than that under static conditions and both total protein and albumin output was also increased under radial flow conditions.

Introduction

Bioartificial livers (BALs) are based on hepatocytes cultured within a bioreactor and are intended to obtain an efficient temporary therapy for acute hepatic failure (Biederman et al. 2000, Carpentier et al. 2009). BALs utilize cultured hepatocytes within a device providing artificial flow so that plasma and blood from a patient can exchange substances with hepatocytes within the bioreactor, through a semipermeable membrane in order to detox patient blood, in a way that the liver damaged can still secrete important metabolites (Carpentier et al. 2009).

A suitable environment is required to mimic the physiological structure of liver tissue (Carpentier et al. 2009, Chen et al. 2008). Conventional hepatocyte cultures are made on a bidimensional surface and without nutrients flow. These static cultures are subject to local changes in culture medium degradation, pH, accumulation of excretory metabolites, etc., while bioreactors are intended to obtain tridimensional cell cultures, with a better distribution of nutrients, thus enhancing metabolic production and cell growth (Chen et al. 2008, Du et al. 2008). Tissue engineering has also attempted to develop bioreactors in which tridimensional cultures will eventually organize as a fully functional tissue, thus approximating the in vivo conditions (Freshney 2009, Galbusera et al. 2007).

Radial flow bioreactors (RFBs) may solve many of the problems found using conventional artificial liver systems. This is achieved through the use of an extended cylindrical bed matrix made of porous bead microcarriers and a medium flowing continuously from the periphery towards the central axis (Li et al. 2009). This kind of systems supports high density, large scale cell cultures with long-term viability (Kosuge et al. 2007, Miyazawa et al. 2007) based on the
beneficial concentration gradient of gases and nutrients produced by a continuous flow through the matrix, which also prevents shear stress or buildup of waste products.

The design of a bioreactor must consider the hydrodynamic features to meet the flow conditions of the liver tissue (Li et al. 2009). Computational flow dynamics (CFD) is an important tool used for tissue engineering that helps to define the system hydrodynamics in the presence of cellular scaffolds (McKenzie et al. 2008). Bioreactor design is based on the scaffold hydrodynamic insight analysis and some of the controllable variables, such as scaffold morphology (pore size, fiber diameter, etc.) and initial cell distribution. These variables are used to solve Navier-Stockes flow equations to define speed in the bioreactor field lines and to approximate both the scaffold and the bioreactor performance (Li et al. 2009, McKenzie et al. 2008, Morales et al. 2008, Park et al. 2005, Park et al. 2008).

Polyactic acid (PLLA) is a widely used biomaterial for scaffold construction, whose surface can be easily modified to improve its adhesion properties (Ramirez-Fernandez et al. 2012, Sherman 2005, Shi et al. 2004, Shoufeng et al. 2001). Plasma polymerized pyrrole (Ppy) modified PLLA surfaces have shown improved cellular adhesion and increased proliferation and life span of cell cultures (Thomas et al. 2006, Wang et al. 2004, Zhang et al. 2001).

We here present the design, construction and validation of a radial flow bioreactor in which cells from a human hepatocellular carcinoma (HepG2 line) were cultured on iodine-doped polypyrrole (Ppy-I) coated PLLA scaffolds. Such bioreactors may prove useful to build BALs for patients suffering acute hepatic insufficiency.

Materials and Methods

System description

The RFB (Fig. 1) consists of a cylindrical glass recipient (50 mm x 100 mm) bearing a food-grade stainless steel central bracket (T-304, diameter 6.35 mm) with 1mm drills distributed all over the surface so that the cell-bearing scaffold (60 mm x 25 mm x 5mm) can be placed before filling the chamber with medium at 80% of its storage capacity, and placing it in a temperature regulated, 5%CO₂ incubator (Fig. 2). Two peristaltic pumps drive the culture medium through the chamber and across the scaffold's metal bracket (Fig. 2). Gases diffuse into the chamber across the 0.2 µm filters. The hoses (6.35mm diameter) connecting the peristaltic pumps and the sampler are made of medical-grade PVC.

Model design

Data from flow simulation may assist the calculations to avoid zero flow zones within the bioreactor. Hence, a geometrical bioreactor model was designed at SolidWorks (Systems Dessault, inc.). The model was then exported to CosmosFloworks (Systems Dessault, inc.) software to solve Navier-Stokes equations to determine speed fields and work against the scaffold. The physical culture features were: isothermal, Newtonian liquid, 1,000 kg/cm³ density, 0.889mPa viscosity and 7 mL/min continuous flow. For the scaffold a 500 µm pores polyethylene sponge was used, together with Pyrex-type glass and stainless steel (T-304) metallic parts.

Surface-modified scaffold

PLLA sponges were 25-100 mg/cc density and 90% porosity (BIOFELT®, Concordia Fibers; Concordia Manufacturing Corp., Inc., Coventry, RI, USA) and their surfaces were modified with iodine-doped polypyrrole (Ppy-I). Plasma discharge was at 9 x 10⁻² Torr pressure (Morales et al. 2008), measured with a Pirani valve (Edwards Vacuum, Inc., Sonora MX; 50 watts, 13.5 MHz). Pyrrole monomers were used...
(98%, Sigma-Aldrich) and iodine (98%, Sigma-Aldrich). Monomers feeding was alternated in an iodine-pyrrole sequence for 4 min, only pyrrole for 6 min, iodine-pyrrole for 4 min and pyrrole for 6 min. Twenty min was the total reaction time.

Cell culture
HepG2 cell line (ATCC HB8065) was obtained from the American Type Culture Collection and were cultured as a monolayer with type E Williams culture medium (Gibco 12551) supplemented with 10% fetal bovine serum (Gibco 16000), 100 units/mL penicillin and 100 mg/mL streptomycin. Cells were cultured in flasks (Nunc, EE.UU.), 5% humidity, 5% CO\textsubscript{2}, 95% air, and the culture medium was replaced twice weekly. Cells were trypsinized and cultured again every 7 days.

Cell density was measured at the time of seeding and after 15 days culturing in the RFB with 80 mL medium, both at zero flow and at 7 mL/min constant flow.

Protein quantification
Total protein concentration in the outflow was measured from days 9 to 15, using the bicinchoninic acid kit (BCA, ThermoScientific; cat. 23255) using bovine serum albumin as standard; 0.5 mL supernatant samples were obtained daily and measured in a multimodal DTX 880 detector (Beckman Coulter). Triplicate measures were obtained.

Albumin quantification
Elisa test was used to quantify albumin secretion into the supernatant from days 10 to 15 (AssayproEA2201-1); 0.5 mL supernatant samples were obtained daily and measured in a multimodal DTX 880 detector (Beckman Coulter). Triplicate measures were obtained.

Data analysis
Data are presented as means (± SEM) of at least three independent experiments. Multiple comparisons were made using ANOVA followed by the Tukey test, using Origin 8.1 software. A P value of less than <0.05 was considered significant.

Results
Functional simulation of the RFB is shown in Fig. 3 with three different scaffold thicknesses, in both frontal and lateral views. Initially there is a speed gradient inside the central bracket, but speed stabilizes afterwards (lateral views, Fig. 3A, C and E); frontal views show a suction vortex at the RFB exit, and no static zones are observed. Simulations showed that radial flow through the scaffold is independent of scaffold thickness as we can see in the images with velocities of 2 m/s in the inlet, 0.6 m/s across the scaffold and 0.2 m/s in the interchange chamber.
Flow rates cover a wide range of magnitudes in the simulation. However, under equal fluid inlet conditions field velocity decreased when the scaffold thickness increased as 0.6 m/s for 0.5 cm, 0.4 m/s for 1.8 cm and 0.3 m/s for 6 cm; but flow was always maintained perpendicular to the seeded cells. According to this, we can effectively reduce shear stress by choosing appropriate materials, geometry and location, even when the flow rate does not change.

Then we generated characterization curves for scaffold velocity (Fig. 4), that can describe the system response when the medium velocity and the scaffold thickness vary. Those curves gave us an idea of the optimal values that do not generate a high shear stress or a low medium flow.

We then proceeded to the RFB construction (Fig. 5) which was based on the simulation data. The steel RFB parts were processed by a computerized numeric control (CNC) machine.
FIGURE 8. Protein secretion as a function of time. Black circles indicate static cultures, and white circles indicate radial flow cultures. Stars indicate statistically significant differences (P<0.05; Student’s t test).

FIGURE 8. Albumin secretion as a function of time. Black circles indicate static cultures, and white circles indicate radial flow cultures. Stars indicate statistically significant differences (P<0.05; Student’s t test).

Light and scanning electronic microscopy pictures of the cultures with zero flow or with 7 mL/min flow are shown in Figs. 6 and 7, respectively. Cells were attached to the scaffold fibers in both conditions, but more cells were covering the scaffold and were forming cell aggregates under the 7 mL/min flow conditions (Fig. 7).

Fig. 7 also shows the cellular distribution on the Ppy-I scaffold, were cell aggregates are visible around the fibers. This distribution is characteristic of high proliferative rate. The SEM images on Fig. 7 show a continuous cell layer around the fibers, indicative of an adequate adhesion of cells to Ppy-I surface modified fibers.

Cell density under radial flow conditions was almost double than that under static conditions (9.10x10^5 cells/mL and 5.38 x 10^5 cells/mL, respectively). Total protein (Fig. 8) and albumin (Fig. 9) production was also increased under radial flow conditions.

Conclusions

A RFB was designed and built in which the scaffold materials and pore size may be changed while maintaining the flow features and the radial distribution of the culture medium. Three-dimensional cell cultures could be obtained on surface modified PLLA fibers coated with Ppy (Ramirez-Fernandez et al. 2012).

The characterization curves described for the RFB model and the velocity vectors may be used to predict optimal scaffold volume and culture medium velocity for high density cell cultures which result in better total protein and albumin outputs. These approaches may result in better BALs performances.

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