Anti-Cancer Drug Screening Based on a Adipose-Derived Stem Cell/ Hepatocyte 3D Printing Technique

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Abstract

Background: During the last several decades, drug screening results based on two-dimensional (2D) cell cultures could hardly be duplicated by animal experiments due to different cell growth environments. With the development of three-dimensional printing (3DP) techniques, in vitro 3D cell cultures show great advantages in many areas. In this study, a 3D drug screening model based on a cell-laden gelatin/alginate/fibrinogen hydrogel was established using a cell 3D printer developed in Tsinghua University.

Methods: 2D and 3D drug screening effects were compared with three anti-liver tumor drugs (i.e. astragalus polysaccharide, 5-Fluorouracil and matrine). Cell survival rates were tested using 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetra-sodium bromide colorimetric and cell counting kit-8 methods. Some other detection methods, such as 4,6-diamidino-2-phenylindole and 5- or 6-(N-Succinimidyl oxycarbonyl)-3',6'-O,-dicetylfluorescein staining, acridine orange/propidium iodide staining, hematoxylin-eosin staining and scanning electron microscope, were applied to assist the analyses.

Results: Cell viability of the 3D adipose-derived stem cell/hepatocyte cocultures was increased. The 3D model enhanced the drug-resistance of hepatocytes.

Conclusion: This in vitro 3D model has promise to be widely used in drug screening.

Keywords: Cell assembly; Drug screening; Adipose-derived stem cells; Hepatocyte; Gelatin/alginate/fibrinogen hydrogel

Introduction

Since the 1990s, the development of new drugs mainly relies on biological detection platforms that are based on molecular and cellular levels [1]. The greatest feature of these platforms is in vitro two-dimensional (2D) cell cultures. Consequently, the drug screening results achieved using 2D cultures have little effect on the vast majority of animals [1]. Especially in the human body many different cell types work in a well-organized way, which is hard to be simulated by 2D cell cultures. It is found that some environmental factors, such as oxygen concentration and temperature, influence the cell growth states, which makes the in vitro simulation particularly challenging.

Recently, a series of new cell culturing techniques, such as optical tweezers [2] and laser-direct writing [3], emerged due to the combination and integration of life science with manufacturing and information technologies. The most obvious disadvantage of these techniques is the difficulty to accurately define and arrange multiple cell types in one construct. 3D printing (3DP), also known as additive manufacturing (AM) or rapid prototyping (RP), is a family of techniques that are able to process nearly all different material systems with computer aided design (CAD) models which are based on object separation-accumulation principles [4]. Especially, biological materials, like cells and cell growth factors, can accurately be put on target positions with quick response speed and high formation precision [5-7]. Several research groups have transformed a variety of RP techniques from industry towards biological laboratory to assemble or print cells under the instructions of CAD models. For example, in Harvard Medical School, a quasi 3D structure including collagen hydrogel, fibroblasts and keratinocytes was printed [8], whereas in University of Cornell a 3D direct-write printing technology was created [9]. In our former studies, a series of cell assembling [10-19] and low temperature deposition manufacturing technologies [20-25] were produced, which are prominent in complex organ manufacturing and high throughput drug screening areas [26-31].

In the present study, we used our cell printing technique to mimic the hepatic lobule structure instead of 2D cell cultures for drug screening. A gelatin/alginate/fibrinogen hydrogel was selected as the extracellular matrix (ECM) to load cells. Hepatocytes were printed separately or together with adipose-derived stem cells (ADSCs) of a Sprague-Dawley (SD) rat following a grid structure. In order to compare the difference between 2D and 3D cell cultures, 1-[4,5-dimethylthiazol-2-yl]-2, 5-diphenyl tetrasodium bromide (MTT) colorimetric and the cell counting kit-8 (CKK-8) methods were used to detect cell viabilities under different anti-cancer drugs, such as astragalus polysaccharide (AP), 5-Fluorouracil (5-FU) and matrine. A series of other biological characterization methods, such as hematoxylin-eosin (HE) staining and acridine orange (AO)/propidium iodide (PI) staining were used for a better understanding of the micro structures and interactions between the hepatocytes and ADSCs.

Materials and Methods

Hepatocytes and adipose-derived stem cells

A new born SD rat (120 g), male, bought from the Laboratory

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Received February 11, 2015; Accepted April 06, 2015; Published April 08, 2015

Citation: Zhao X, Du S, Chai L, Xu Y, Liu L, et al. (2015) Anti-Cancer Drug Screening Based on a Adipose-Derived Stem Cell/Hepatocyte 3D Printing Technique. J Stem Cell Res Ther 5: 273. doi:10.4172/2157-7633.1000273

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Animal Center, Medical Department of Beijing University (approved by the Ministry of Health) was used for the cell separation. The hepatocytes were extracted by infusing the rat liver with a 0.1% type II collagenase (sigma, Aldrich Company, USA) solution \cite{10,11}. Neutralized cell suspension was centrifuged at a speed of 600 r/min for 5 min to get the hepatocytes.

Epididymal adipose tissues from the same rat were excised, washed sequentially in serial dilutions of betadine, and then finely minced in a calcium-free phosphate-buffered saline (PBS) (pH 7.2). The mashed tissues were digested in the 0.1% type II collagenase at a temperature of 37°C for 45 min. Neutralized cells were centrifuged to separate the mature adipocytes from the stromal vascular fraction. The floating adipocytes were removed and pelleted. The stromal cells were filtered through a 100 µm cell strainer before seeding. ADSCs were cultured in Dulbecco's modified Eagle's medium (DMEM) (Gibco, Paisley, UK), containing 10% fetal bovine serum (FBS) (Gibco, Paisley, UK) at 37°C in an atmosphere of 5% CO₂. Cells were grown to subconfluence and passaged using a standard trypsinization method.

Gelatin/alginate/fibrinogen hydrogel

Based on the former research of our laboratory, a mixture of gelatin, sodium alginate and fibrinogen was chosen to be the extracellular matrix (ECM) considering biological compatibility, mechanical behavior and economic factors. The gelatin (Tianjin Green-Island Company, China) was dissolved in calcium-free PBS to form 20% and 10% (w/v) solutions. Sodium alginate (Sigma, Aldrich Company, USA) was dissolved in calcium-free PBS to form 4% (w/v) solution. The gelatin and sodium alginate solutions were pasteurized three times at 72°C for 20 min. Fibrinogen and thrombin (Sigma St. Louis, MO, USA) were dissolved in fetal bovine serum (FBS) (Gibco, Paisley, UK) to form a 5% (w/v) solution. Two concentrations of gelatin and sodium alginate mixtures in a fibrinogen solution were used for cell printing. One was 20% (w/v) gelatin, 4% (w/v) sodium alginate and 5% (w/v) fibrinogen. The other was 10% (w/v) gelatin, 4% (w/v) sodium alginate and 5% (w/v) fibrinogen. Both mixtures consisted of the same ratio between gelatin:alginate:fibrinogen = 2:1:1 (v/v/v).

Controlled cell-laden gelatin/alginate/fibrinogen hydrogel printing and stabilization

The used cell printer is shown in Figure 1a and the printing process in Figure 2. A grid 3D structure of 10 mm × 10 mm × 5 mm was designed as shown in Figure 1b. Hepatocytes were printed separately in the gelatin/alginate/fibrinogen hydrogel at a density of 1 × 10⁶ cells/ml or as a mixture with ADSCs (1 × 10⁶ cells/ml). The printed construct is shown in Figure 1c. After being printed, the construct was stabilized with a 5% (w/v) CaCl₂ solution and a 5% (w/v) thrombin solution separately for 5 min in order to crosslink the alginate and to polymerize the fibrinogen molecules.

Scanning electron microscope observation

After 7 days in vitro culture, some of the samples were fixed in 2.5% glutaraldehyde at 4°C for at least 2 h before they were dehydrated in a series of ethanol (50%, 70%, 80%, 90% and 100%) solutions and a mixture of half alcohol and half tert-butyl alcohol for 10 min. At last the samples were soaked in tert-butyl alcohol, stored under -20°C, freeze-dried and observed using a scanning electron microscope (SEM, Hitachi S-450, Japan).

HE Staining

After 7 days in vitro culture, some of the 3D constructs were firstly fixed in 4% paraformaldehyde overnight, embedded in optimum cutting temperature compound (OCT), frozen and sliced to 14 μm sections. Then the sections were air dried, fixed with cold acetone, washed with PBS for three times, stained with haematine for 5-10 min, and placed in 4% methyl green for 5 min.
in concentrated ammonia for 2 s (for the colored nuclei). Finally, the sections were stained in eosin for 1 min, soaked in 70%, 80%, 90%, 95% (2 times), 100% (3 times) alcohol respectively for 1 min, washed with xylene 3 times, and observed under a microscope. Nucleus appeared in purple and cytoplasm in red color.

4',6-diamidino-2-phenylindole (DAPI)/ 5- or 6-(N-Succinimidylcarbonyl)-3',6'-O,O'-diacetylfluorescein (CFSE) staining

Before cell printing, the concentrations of the hepatocyte and ADSC suspensions were adjusted to 1 × 10^7 cells/ml and 1 × 10^6 cells/ml, respectively. 100 µl DAPI (1 µg/ml) was added to 1 ml of the ADSC suspension and 40 µl CFSE was added to 1 ml of the hepatocyte suspension. The hepatocyte and hepatocyte/ADSC suspensions were placed in a 37°C incubator for 15-20 min. After the cell suspensions were centrifugated, the supernatant liquid was removed, and the precipitated cells were washed with PBS 3 times before they were printed with the prepared gelatin/alginate/fibrinogen hydrogel. After 7 days in vitro culture, the samples were examined using a Zeiss laser scanning confocal microscope (LSCM710). The excitation/emission wavelength for DAPI and CFSE staining were 340 nm/488 nm and 490 nm/530 nm, respectively.

Drug screening

After the 3D constructs were cultured in 24-well petri dishes for one week, three drugs, AP (Bio-medical Assay), 5-FU (Sigma) and matrine (Santa Cruz), were added to the culture medium in three different concentrations according to our former experiences: high concentration (H) AP 4 mg/ml, 5-FU 500 μg/ml, matrine 1.5 mg/ml, middle concentration (M) AP 400 μg/L, 5-FU 50 μg/ml, matrine 0.15 mg/ml, and low concentration (L) AP 40 μg/L, 5-FU 5 μg/ml, matrine 15 μg/ml, for 1 to 4 days. The three concentrations of the three drugs are listed in Table 1. Hepatocytes cultured directly on 2D 96-well petri dishes with the same initial cell density as the 3D constructs and the same drugs were set as a positive control. Untreated 2D plates and 3D cultures were set as negative controls. CCK-8 was used to measure the cell proliferation rate and drug screening effects of the 2D cultures while a MTT colorimeter was used for the 3D cultures.

MTT colorimetric measurement: after the cells were exposed to the drugs for different durations, 100 µl of MTT staining solution was added to the 24-well plates with the 3D samples before incubating them at 37°C for 4 h. The liquid was then removed and 500 µl DMSO was added. The culture plate was rolled 10 min in a shaking table at room temperature. 150 µl of the liquid in each well was transferred to a 96-well. The optical density (OD) of each well was measured at 490 nm wavelength using a Bio-Rad 550 reader (Japan). The ratio of experimental and control groups represents the relative survival rates of the cells.

CCK-8 measurement: 100 µl DMEM and 10 µl CCK-8 solution were added to each well of the 96-well plate with the 2D samples. Subsequently the plate was put to a carbon dioxide incubator for 4 h. The OD of each well was measured at 450 nm using the above mentioned Bio-Rad 550 reader. The ratio of experimental and control groups again represents the relative survival rates of the cells.

| Drug | High concentration (H) | Middle concentration (M) | Low concentration (L) |
|------|------------------------|-------------------------|----------------------|
| AP   | 4 mg/ml                | 400 μg/L                | 40 μg/L              |
| 5-FU | 500 μg/ml              | 50 μg/ml                | 5 μg/ml              |
| Matrine | 1.5 mg/ml            | 0.15 mg/ml              | 15 μg/ml             |

Table 1: Three groups of drug concentrations for screening experiments.
Acridine orange/propidium iodide staining
After being exposed to the drugs for 1-4 days, both the 2D and 3D samples were stained with acridine orange/propidium iodide (AO/PI) fluorescence. 0.01 ml of AO solution (670 µmol/L~0.3 mg/L) and 1 ml PI solution (750 µmol/L~0.5 mg/L) were added to the 24-well plate with the samples before they were put in a 37°C incubator for 10 min. After being washed by PBS for several times the samples were examined using the LSM.C. The emission wavelength of AO is 530/640 nm and PI is 615 nm.

Statistical analysis
Sample values were expressed as means ± standard deviation (SD). Unpaired Student’s t-test was used to compare the differences between the 2D/3D groups or different culture days. SPSS version 13.0 statistic software (SPSS, Chicago, IL, USA) was used to analyse the values. P-values<0.05 were considered significant.

Results
Effect of low and high gelatin concentrations on morphology of cells laden in the gelatin/alginate/fibrinogen hydrogel
Some SEM images of the 3D constructs with 10% (w/v) low and 20% (w/v) high gelatin concentrations are shown in Figure 3. The interconnected go-through pores (channels) were predefined by the CAD model to achieve a larger contact area between the medium and the assembled cell. During the printing stage, gelatin played an important role in the sol-gel transformation of the gelatin/alginate/fibrinogen hydrogel and the stabilization of the printed constructs. With the crosslinking and polymerization of the alginate and fibrinogen molecules, the printed 3D constructs remained stable without collapsing during the long-term in vitro culture. It was found that the gelatin concentration had significant effect on the microstructures of the 3D constructs. In the low concentration group, there were abundant micro-pores in the printed gelatin/alginate/fibrinogen filaments with diameters ranging from 5 µm to 30 µm. These micro-porous were beneficial for nutrient and metabolic substance transportation in the hydrogel, supplying the embedded cells with adequate nutrition and oxygen. In contrast, in the high concentration group the thick gelatin/alginate/fibrinogen hydrogel appeared in sheet like shapes. After 7 days culture, most of the hepatocytes were still wrapped by a dense layer of gelatin/alginate/fibrinogen matrix and were separated from each other (Figures 3d, 3e and 3f). From the comparison of these two groups, we can conclude that in the low concentration group, cell shapes were more distinctive than those in the high concentration group. It is easier for the cells in the low concentration matrix to form connections.

HE staining results of hepatocytes and ADSCs/hepatocytes in the gelatin/alginate/fibrinogen hydrogel
HE staining results of the hepatocyte and ADSC/hepatocyte containing 3D constructs are shown in Figures 4a-4d. After 7 days in vitro culture, many of the cells were in division states with double nuclei. HE staining results containing both hepatocytes and ADSCs are shown in Figures 4e-4h. It is hard to distinguish the hepatocytes and ADSCs from colors. The main differences of the two cell types are shape and size. Most of the hepatocytes have elongated round or elliptical shapes with an average diameter of ~15 µm. Most of the ADSCs are round with an average diameter of 6 µm, which is much smaller than those of the hepatocytes. In the marked area of Figure 4f (by a black oval), cell connections are much clearer than anywhere else where the gelatin/alginate/fibrinogen matrix is prominent. This result indicates that the thick gelatin/alginate/fibrinogen hydrogel encapsulates the cells and prevents them from interacting directly. These results are coincident with the SEM images (Figure 3).

DAPI/CFSE staining results of hepatocytes and ADSCs/hepatocytes in the gelatin/alginate/fibrinogen hydrogel
Figures 5a and 5b are DAPI and CFSE staining results of the 3D construct with only hepatocytes after 7 days culture. DAPI mainly dyes the nucleus and CFSE mainly the cytoplasm. A large number of actively dividing cells were observed from both the staining results, which indicate that the primary hepatocytes in the 3D construct had strong proliferation ability after 7 days in vitro culture. In order to distinguish hepatocytes and ADSCs under a confocal microscope, we applied CFSE to stain the hepatocytes and DAPI to the ADSCs. In Figures 5c and 5d ADSCs/hepatocytes were normal cultured without the addition of drugs. Cell-cell interactions in between the hepatocytes are prevailing with a lot of threadlike structures.
Drug screening results

Table 2 contains the drug screening results (OD) of the AST, and Figure 6 shows the cell survival rates under different culture conditions after exposed to AP, 5-FU and matrine for 1 to 4 days. It was found that the culture conditions have significant effects on the drug screening results. AP shows cytotoxicity for the 2D hepatocyte cultures by reducing the cell survival rate by half, while the cell proliferation ability of the 3D hepatocyte cultures increases approximately 20%. In the 3D coculture model, the hepatocyte number doubled after 4 days exposure to AP. Significant differences were found among the 2D, 3D, 3D coculture groups containing pure hepatocytes or ADSC/hepatocyte mixtures (P<0.05).

Figure 7 show that different concentrations of AP, 5-FU and matrine also have significant influence on the hepatocyte survival rates in the 3D model (pure hepatocytes). For AP, the highest difference appeared on the second day after drug medication. Cell survival rates of the high and the low concentration groups decreased by 20% while the survival rate of the middle concentration group decreased by 13%.
Significant differences were found among the three AP concentration groups containing pure hepatocytes (P<0.05).

Table 3 contains the drug screening results (OD) of the 5-FU. There is no significant difference between the 2D and 3D hepatocyte cultures after medicated by the 5-FU, indicating that the 5-FU has little cytotoxicity to hepatocyte under the concentration of 50 µg/ml. However, the survival rate of the ADSC/hepatocyte cocultures decreases sharply on the second day to 82%, and slowly recovers by the end of the 4th day. Figure 7 the 5-FU concentrations have significant influence on the hepatocyte survival rate of the 3D model. In the low concentration group, the cell survival rate drops continuously from 116% to 59%, while in the middle and high concentration groups the cell survival rate rebounds on the third day. It seems that the higher the used drug concentration, the more obvious is the rebound. These results indicate that the ADSC/hepatocyte cocultures are more likely to develop drug resistance even with the highest 5-FU concentration.

Table 4 contains the drug screening results (OD) of the matrine. During the first two days exposing the proliferation rate of the hepatocytes in the 2D cultures remained nearly the same. After 4 days culture, the cell survival rate drops to ~80%. The survival rate of the 3D hepatocyte cultures fluctuates during the drug action stage, and decreases to ~86% after 4 days exposing. Matrine expresses little cytotoxicity for the 3D ADSC/hepatocyte cocultures on the first day, while the cell proliferation rate increases about 29% after 4 days exposing. Figure 7 also shows the survival rate of the hepatocytes in the 3D model after treated with matrine in different concentrations. The line shapes are similar which reflects that there are no significant differences among the drug concentrations.

Figure 8 is the AO/PI staining results of the 2D model after 7 days in vitro culture and 4 days medication with different anti-cancer drugs in middle concentration (AP 400 µg/L, 5-FU 50 µg/ml, matrine 0.15 mg/ml). Figure 8a is a non-drug-medication control with hepatocytes in division period with two nuclei. Figures 8b-d show the results of the 5-Fu, AP and matrine groups. After treated with 5-FU and matrine, the cell membranes of the hepatocytes were damaged to some extend and the nuclei were stained red. Though the cells were still complete, the number of double nuclei decreased, reflecting that the cell proliferation ability was weakened.

Figure 9 is the AO/PI staining results of the 3D hepatocyte cultures after 7 days in vitro culture and 4 days medication by the three anti-cancer drugs in middle concentration (AP 400 µg/L, 5-FU 50 µg/ml, matrine 0.15 mg/ml). Figure 9a is a negative 3D control without the used drug concentration, the more obvious is the rebound. These results indicate that the ADSC/hepatocyte cocultures are more likely to develop drug resistance even with the highest 5-FU concentration.

Figure 9a, showing some hepatocytes in division period with two nuclei. Figure 8c and 8d are magnifications of Figure 8a. Compared to the control group, cell density declined sharply. In Figures 8c and 8d, the cell membranes are severely damaged and the cells are stained to red. Figures 8e and 8f show the results of the 5-Fu group with different magnifications. Figures 8g and 8h are the matrine group. After treated with 5-FU and matrine, the cell membranes of the hepatocytes were damaged to some extend and the nuclei were stained red. Though the cells were still complete, the number of double nuclei decreased, reflecting that the cell proliferation ability was weakened.

Table 3: Drug screening results (OD) of the 5-Fluorouracil.

| Concentration Culture time (d) | H | M | L | H | M | L | H | M | L |
|-------------------------------|---|---|---|---|---|---|---|---|---|
| 1                             | 0.98 | 1.21 | 1.06 | 0.90 | 1.00 | 1.16 | 0.81 | 1.31 | 0.85 |
| 2                             | 1.97 | 0.95 | 0.94 | 0.64 | 0.97 | 1.07 | 0.73 | 0.82 | 0.99 |
| 3                             | 0.82 | 1.14 | 1.34 | 1.81 | 1.38 | 0.93 | 0.83 | 0.87 | 1.00 |
| 4                             | 1.11 | 0.85 | 0.71 | 1.37 | 0.92 | 0.99 | 0.98 | 1.10 | 1.35 |

Table 4: Drug screening results (OD) of the matrine.

| Concentration Culture time (d) | H | M | L | H | M | L | H | M | L |
|-------------------------------|---|---|---|---|---|---|---|---|---|
| 1                             | 1.07 | 1.07 | 1.00 | 1.02 | 0.93 | 1.11 | 0.78 | 0.67 | 0.84 |
| 2                             | 0.95 | 1.17 | 1.33 | 0.94 | 0.85 | 1.01 | 0.84 | 0.74 | 0.95 |
| 3                             | 0.77 | 0.89 | 0.92 | 1.62 | 1.32 | 1.29 | 1.06 | 1.11 | 1.07 |
| 4                             | 1.35 | 0.80 | 0.86 | 1.03 | 0.86 | 0.87 | 1.33 | 1.29 | 1.41 |
Figure 10 is the AO/PI staining results of the 3D ADSC/hepatocyte coculture after 7 days in vitro culture and 4 days drug medication in middle concentration (AP 400 μg/L, 5-FU 50 μg/ml, matrine 0.15 mg/ml) compared to the only hepatocyte models. Interestingly, cell numbers in the 5-FU (Figures 10e and 10f) and matrine (Figures 10g and 10h) used groups did not change obviously compared to the control group (Figures 10a and 10b). Especially, the number of cells in the AP used group doubled and the biggest cell cluster reaches 200 μm in diameter, indicating that AP could enhance cell proliferation ability of the hepatocyte/ADSC cocultures.

**Discussion**

In vivo, single cells (~10 μm in size) extend to functional subunits (~100 μm – 1 mm) and form hierarchical structures which coordinate tissue or organ functions. Over the last century, cells were isolated from tissues and cultured in 2D systems with neglected cell-cell and cell-matrix interactions. Previously, we have provided hepatocytes with microenvironments similar to the in vivo conditions with a series of 3D cell printing techniques [10,11]. In the present study, hepatocytes cultured in 2D and 3D environments were compared using MTT, CCK-8, HE, SEM, DAPI/CFSE and AO/PI methods. Generally, the 2D cultured cells adhere to the culture dishes with less interconnection. The 3D cultured cells gather together due to the cell migration and adhere to the EMCs, which is spatially more similar to the in vivo environment.

Primary hepatocytes appeared to be the most powerful liver-specific drug screening cell types [32]. Yet cell cocultures have...
and excellent chemical properties [35]. Gelatin turns into colloid by that, gelatin also has moderate crosslink ability, adjustable solubility acid with favorable biodegradability and biocompatibility. Apart from hydrolyzed protein. Its major component of polypeptide contains amino matrix. Gelatin is a kind of incomplete degradation or partially cultures, a gelatin/alginate/fibrin hydrogel was used as a cell laden constructs with the crosslinking of the alginate molecules in the printed cell-laden filaments.

Fibrinogen is an important component to promote the blood coagulation. It is a precursor fiber protein synthesized by the liver and can be fully re-absorbed and replaced by the growing tissues. Fibrinogen is composed of α, β and γ polypeptides, and its peptide chains are connected by disulfide bonds. Under the action of thrombin, α chain and β chain release peptide A and peptide B to generate the fibrin monomer. The monomers are further polymerized by covalent bond and form stable fibrin clot to complete the coagulation process [37]. With the addition of fibrinogen in the gelatin/alginate hydrogel and polymerization process, the 3D constructs were further stabilized. The gelatin/alginate/fibrin hydrogel could act as an ECM for the encapsulated cells and provided the cells with an excellent growth microenvironment. After comparing the different hydrogel concentrations, we found that 10% (v/w) gelatin is beneficial for micro-pore formation enabling the cells to migrate and proliferate inside.

There were two purposes for the series of biological characterizations. One is to observe the cell morphologies and the relationships among cells with the help of HE staining, SEM and DAPI/CFSE fluorescence labeling. The other is to assist the drug screening effects with the help of MTT, CCK-8, and AO/PI staining. DAPI and CFSE double staining were employed to distinguish the hepatocytes and ADSCs. DAPI is a fluorescence stain that can pass through the cell membrane and combine with the double chain DNA to emit purple fluorescence under fluorescence microscope. As no toxicity to cells, it is commonly used in apoptosis detection. CFSE can easily pass through the cell membrane and emit green fluorescence when combined with proteins of living cells. It has no toxicity to cells, and the fluorescence can be divided and emit green fluorescence when combined with proteins of living cells. It has no toxicity to cells, and the fluorescence can be divided and emit green fluorescence when combined with proteins of living cells.

In order to compare the drug screening results of the 2D and 3D cultures, a gelatin/alginate/fibrin hydrogel was used as a cell laden matrix. Gelatin is a kind of incomplete degradation or partially hydrolyzed protein. Its major component of polypeptide contains amino acid with favorable biodegradability and biocompatibility. Apart from that, gelatin also has moderate crosslink ability, adjustable solubility and excellent chemical properties [35]. Gelatin turns into colloid by reversible physical crosslink under 35°C, which makes it useful in our cell printing systems.

Sodium alginate is a natural linear polysaccharide extracted from brown algae, made by β-D-mannuronic acid of 1-4 bond and α-l-guluronic acid with a formula of C₂₁H₂₂O₅Na. It has been used to replace damaged tissues and organs since early 1990s with relatively good biological compatibilities [36]. Alginate solution has a sol-gel transition with Ca²⁺, Mg²⁺ and other bivalent positive ions. With this property, it is easily to stabilize the 3D constructs with the crosslinking of the alginate molecules in the printed cell-laden filaments.

5-Fluorouracil is a common chemotherapy drug for many kinds of cancers. It blocks the synthesis of thymine, an essential raw material of antiviral, anticancer, anti-aging, radiation resistance, stress resistance and oxidation resistance [39]. In the present study, AP enhanced the proliferation capabilities of the 3D cultured pure hepatocytes and ADSC/hepatocyte cocultures, while it weakened the activity of the 2D cultured pure hepatocytes (Figure 6). This phenomenon proved that the 3D construct printed by the cell assembly machine provided better living conditions for the hepatocytes than the 2D cultures.
C, and is recently found to have anti-tumorous effects [41]. It has cytototoxicity to both the 2D and 3D cultured hepatocytes. Nevertheless, cell survival rates in the 3D cocultures increased from 67% to 129% during the 4 days culture (Figure 6).

In the succinate dehydrogenase (SDH) staining method (CCK-8), soluble tetrathionate salt WST-8 is used to reduce the soluble auranus under certain conditions. CCK-8 is a good colorimetric method for living cells, especially in cell proliferation and toxicity measurement [42]. The MTT colorimetric method can be used to detect the cell proliferation rate. The principle of the MTT colorimetric measurement is that mitochondrial succinate dehydrogenase can restore the exogenous MTT to insoluble blue purple crystal formazan and deposit it in cells. However, dead cells do not have this function. DMSO can dissolve the formazan in cells. By using enzyme-linked immune detectors we can measure its absorbance value at λ=490 nm, which is a proxy of the number of living cells. Within a certain amount of cells, the number of MTT crystal is in direct proportion to the cell numbers [43,44].

AO is a fluorescent dye with nucleic acid compatibility, and displays green color under fluorescence microscope. AO has membrane permeability, and has affinity with active DNA. It was difficult for AO to act with apoptotic cells due to the chromatin pyknosis or break. So AO staining was used to distinguish living cells and dead cells under fluorescence microscope. PI is also a nucleus staining material that can dye DNA with red fluorescence. Different from AO, PI cannot penetrate living cell membranes, and is generally used in the detection of cell apoptosis. By integrating AO and PI staining, the living and dead cells were distinguished under fluorescent microscope at the same time. After the exposure to drugs, the membrane permeability of the cells changed and the nucleus of the dying cells were stained red (Figures 8 and 9). Yet in the coculture groups, most of the cell activities remained with hardly any red nucleus (Figure 10).

Similarly, we can clearly see the cell states in the 3D printed hepatocyte/ADSC cocultures using DAPI/CFSE double staining. In the control group (Figure 5), the cytoplasm of the hepatocytes and ADSCs were plump and the nuclei were complete. In the drug used groups different culture conditions had significant influences on the cell survival rates (Figure 6) and morphologies (Figures 8, 9 and 10). Among the three anti-carcinogen drugs, astragalus had apoptosis inhibition ability on the hepatocytes, while 5-FU and matrine were toxic to the hepatocytes. Different drug concentrations also had obvious effects on cell proliferation rates (Figure 7) in the 3D models. Compared to the 2D cultures the 3D models enhanced the hepatocyte proliferation abilities to a certain degree when the anti-tumor drugs were used. Meanwhile, the 3D coculture had the highest cell proliferation rate among all experiment groups.

In addition, with our 3DP techniques, cells can be assembled in a 3D construct using the robotic printer in about 20 min. The following sample handling and data processing can be manipulated repeatedly and quickly with the software control systems. Cells that are cultured in the 3D environments more closely mimic the tissues in the body than those grown in 2D. Consequently, the drug screening platform can be high-throughput, easily scalable and more reliable, which allows researchers to conduct genetic, chemical, or pharmacological tests.

Conclusions

A 3D drug screening model was created through a cell-laden gelatin/alginate/fibrinogen hydrogel printing technique developed by Tsinghua University. And a series of biological characterizations were carried out, including microstructures, cell proliferation abilities, and cell survival status. It was found that pure hepatocyte or ADSC/hepatocyte mixtures were all in good conditions after 7 days culture with many cells in division states. The 3D construct provided cells with a good cell survival environment and was used to compare drug screening abilities with 2D cell culture and 3D coculture systems. Cells easily built connections in the 3D samples that contained a low gelatin concentration. In the ADSC/hepatocyte cocultures, hepatocyte and ADSCs could be distinguished by the staining color and size. After exposed to three anti-tumor drugs some of the cells were damaged with decreased proliferation rate and cell-cell interconnections. The AO/PI staining results were consistent with the MTT results. AP could inhibit the hepatocyte apoptosis to certain extent. 5-FU and matrine both had toxic effects on the hepatocytes. The toxic effect of the 5-FU was relatively stronger within the low concentration group. Compared to the 2D cultures the 3D constructs enhanced the drug-resistance ability of the hepatocytes to a certain degree, while the ADSC/hepatocyte cocultures had the highest cell proliferation rate.

Acknowledgments

The work was supported by grants from the Cross-Strait Tsinghua Cooperation Basic Research (No. 2012TH02-3), Beijing Municipal Natural Science Foundation (No. 3152015), National Natural Science Foundation of China (NSFC) (No. 81271665 & 30970748), International Cooperation and Exchanges NSFC and Japanese Society for the Promotion of Science (JSPS) (No. 81411140040), State Key Laboratory of Materials Processing and Die & Mold Technology, Huazhong University of Science and Technology (No. 2012-P03), and the National High Tech 863 Grant (No. 2009AA043801).

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Citation: Zhao X, Du S, Chai L, Xu Y, Liu L, et al. (2015) Anti-Cancer Drug Screening Based on a Adipose-Derived Stem Cell/Hepatocyte 3D Printing Technique. J Stem Cell Res Ther 5: 273. doi:10.4172/2157-7633.1000273

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