Selection of Hepatocyte-Like Cells from Mouse Differentiated Embryonic Stem Cells and Application in Therapeutic Liver Repopulation

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Abstract
Background/Aim: Because of the oncogenic risk, it is important to gain the homogeneous and purified cells from differentiated ESCs before transplantation. Here, we aim to select hepatocyte-like cells from differentiated ESCs, and investigate their growth, differentiation and neoplastic formation after intrahepatic transplantation. Methods: Mouse ESCs were primarily induced by Dexamethesone, FGF-4 and HGF sequentially, then placed to a conditioning selection media consisting of 5% cholestatic sera and cultivated for 2 wks. After labeled by CFDA-SE, the selected cells were transplanted into mouse liver in therapeutic liver repopulation models. Results: In the early stage of screening cultivation, most cells were suffered from apoptosis or even death. 1w later, some hepatocyte-like colony-forming units were observed, then the selected cells could grow and tend to be more mature, as assessed by morphological and functional tests. After intrahepatic transplantation, the labeled cells could proliferate and expressed albumin. Moreover, teratoma didn’t form over 3 months. Conclusion: Our conditioning selection media could not only effectively select hepatocyte-like cells from differentiated ESCs, but further promote their growth and differentiation as well. After intrahepatic transplantation in therapeutic liver repopulation models, the selected cells could grow, differentiate and keep partial hepatic function. In particular, the transplantation was safe.
Introduction

The treatment of troublesome or end-stage liver diseases (ESLD), such as hepatitis, liver cirrhosis, primary liver carcinoma, congenital biliary atresia, inherited enzyme deficiency diseases and acute liver failure, is always a global difficulty. Orthotopic liver transplantation (OLT) is an effective treatment for patients with ESLD, but the shortage of organ donors limits the number of liver transplantations [1]. Hepatocyte transplantation (HTx) provides a new way for it, and maybe an alternative substitute therapy for these liver diseases [2]. However, the anticipated clinical efficacy of HTx has not been proved yet, mainly because of the limitations of the source deficiency and the proliferation difficulty of the transplanted hepatocytes [3-4].

The establishment of human embryonic stem cells (ESCs) lines will undoubtedly give much hope to solve the source deficiency of transplantable cells [5]. The distinguishing feature of ESCs is their pluripotent ability to differentiate into a broad spectrum of derivatives of all 3 embryonic germ layers. This ability has drawn attention to ESCs as a novel source of cell populations for new therapeutic strategies such as cell transplantation and tissue engineering. Actually, abundant studies have reported the capacity of ESCs to differentiate into hepatocytes or hepatocyte-like cells by different ways [6-10], but the propensity to develop teratomas about the ESCs-derived hepatocytes and low efficiency are still the dominating problems. So getting the homogeneous and secure cells suitable for transplanting is still important.

In fact, researchers had adopted various methods to select and purify hepatocyte-like cells from differentiated cells [11-14]. For example, Yamada et al. found that indocyanine green (ICG)-staining was a useful marker to identify differentiated hepatocyte-like cells from EBs in vitro. Since ICG is eliminated exclusively by hepatocytes, so ICG-staining cells can be isolated from other kinds of cells [12]. Yin et al. used AFP as a marker. After knocking the GFP gene into the AFP locus of ESCs and differentiating the modified ESCs in vitro, they got a subpopulation of GFP (+) and AFP-expressing hepatic-like cells, thus hepatocyte-like cells were isolated [13]. However, the methods mentioned above are either complex or have a poor isolation efficiency. Therefore, it is essential to establish not only an appropriate culture system to induce ESCs to hepatocytes, but also an useful method for isolating hepatocyte-like cells from differentiated ESCs to eliminate the potential danger of forming aggressive ESCs-derived teratomas in vivo.

Here, we primarily induced mouse ESCs to differentiate into hepatocyte-like cells by adding Dex, FGF-4 and HGF sequentially, in accordance with the research proceeding of liver developmental biology. Then, on foundation of succeeding to select hepatic stem cells from bone marrow stem cells with a conditioning selection media consisting of 5% cholestatic sera [15], we further investigated the effects of this pathological microenvironmental culture system on selecting hepatocyte-like cells from differentiated ESCs. Finally, we transplanted these induced and selected cells into mice liver through portal vein, in therapeutic liver repopulation mice models which were pre-treated with retorsine and 70% partial hepatectomy. We would confirm whether the transplanted cells could be normally incorporated into mice liver parenchymal structure and whether they could further proliferate, keep the hepatic function or form teratomas. Our research may lay a foundation for the clinical application of ESCs as a novel source of HTx.

Materials and Methods

**Culture of ESCs**

E14 mouse ESCs were maintained in feeder-free DMEM (Dulbecco's modified eagle medium) with high glucose (Gibco BRL, Rockville MD), 1000 U/ml recombinant mouse leukemia inhibitory factor (rMLIF) (Chemicon), 10% fetal bovine serum FBS (HyClone), 0.1 mmol/L β-mercaptoethanol (Chemicon), 10ml/L...
Insulin-Transferrin-sodium selenite (Gibco) and 1% nonessential amino acids (Chemicon), 100 u/ml penicillin, 100 μg/ml streptomycin. The density of ESCs was approximately 1×10⁶/ml.

**Initial proliferation and differentiation of ESCs**

When mouse ESCs grew to about 70% of the cultivating bottle, they were dissociated and resuspended in media above without rmLIF. Then we seeded the individual drop of 30ul (0.5×10⁵/ml) onto the surface of 100mm bacterial grade plate and inverted the whole assembly. The handing-drop ESCs were incubated at 37 °C in 5% CO₂ for 2 d. After incubation of 3 d, the embryoid bodies (Ebs) were aspirated onto six-well tissue culture plates for adherent culture. Then several factors were sequentially added to the culture media for hepatocyte maturation as the following: 10⁻⁷ M Dexamethasone (Dex) (Sigma) at d 1, 10ug/L fibroblast growth factor-4 (FGF-4) (Chemicon) at d 3 and 25 ug/L hepatocyte growth factor (HGF) (PeproTech) at d 5.

**Preparation of the conditional selective medium**

Cholestatic serum was prepared according to our previously reported method [15]. Sprague-Dawley rats weighing 200–250 g (Laboratory Animal Research Center of Sun Yat-sen University, Guangzhou, China) underwent ligation and transection of the common bile duct under general anesthesia with ether to induce cholestasis. Ten days after the operation, the rats were sacrificed and whole blood was collected. Serum was isolated from the whole blood and then subjected to liver function testing, inactivated, and sterilized for use in culture. Finally, cholestatic serum was added into the basic differentiating media mentioned above, to achieve 5% concentration. Differentiating media containing 5% cholestatic serum served as the conditional selective media. For the control group, the serum collected from normal rats was added into the basic differentiating media to achieve 5% concentration. In our study, primary differentiated hepatocyte-like cells (9 d after initial induction) were transferred into the conditional selection media consisting of 5% cholestatic sera and cultured for 2 wks. Media was changed every 2 d.

**Morphological changes**

Primary differentiated cells and selected cells were observed under inverted phase contrast microscope. Cells with regular multilateral shape were viewed as hepatocyte-like cells. What’s more, 2 wk after screening cultivation, ultrastructure of the cells was observed with electron microscopy.

**Immunohistochemistry assay**

9 d after initial induction, differentiated ESCs were taken out and washed with PBS twice, then were fixed with 4% paraformaldehyde for 30 min at room temperature. Albumin and CK8/18 level were detected with polyclonal anti-mouse antibody for albumin (1:200) (Accurate Chemical) and polyclonal anti-mouse antibody for CK8/18 (1:200) (Lab Vision). The slides were incubated with respective primary antibodies at room temperature for 2 h in a humidified chamber. Blocking serum and biotinylated secondary antibody were matched with the primary antibody. Immunoperoxidase was stained and counterstained with DAB and Gill-I hematoxylin.

**RT-qPCR**

Total RNA was extracted using Trizol (invitrogen, USA) according to the product specification and the concentration of total RNA was measured with NanoDrop 2000c (Thermo, USA). 2ug total RNA was obtained for cDNA synthesis with RevertAidTM First Strand cDNA Synthesis Kit (Fermentas, USA). Quantification of gene expression was performed using a real-time polymerase chain reaction (PCR) kit (Takara, Japan).

The expression of the following genes was analyzed: ALB, TTR, AAT, G-6-P, CK18, TAT, FXR, Cyp7a1 and GAPDH (as internal reference). All primers used above were listed as follows: ALB sense: 5’- CAG GAT TGC AGA CAG ATA GTC-3’ and antisense: 5’- GCT ACG GCA CAG TGC TTG-3’; G-6-P sense: 5’- CAG GAC TGG TTC ATC CTT-3’ and antisense: 5’- GGT TGC GTA GTC GGG TGG-3’; TTR sense: 5’- CTC ACC ACA GAT GAG AAG-3’ and antisense: 5’- GCC TGA TGC TCTCAA TAC TTT-3’; AAT sense: 5’- AAT GGA AGA CAT TCG AT-3’ and antisense: 5’- AAG ACT GTA GCT GCT GGA GC-3’; CK18 sense: 5’-GTT CCT CTC CTC AAT CTG CTG-3’ and antisense: 5’- CAC ACA GCT CTC ACT CTT TTT-3’ ; Transferrin sense: 5’-GCA GTC GCC AGT TGC TCC TTC TTC TC-3’ and antisense: 5’- TTA AAC AGC AGG TCC TTC TCT GC-3’; FXR sense: 5’-CCA AGT TTG TCT ACC ACC C-3’ and antisense: 5’-CAC ACA GCT CAT CCC CTT T-3’; Cyp7a1 sense: 5’- AGT ATG ACT ATT TCT TTG ATT TGG GGA-3’ and antisense: 5’- CTT CGT TGT CCA AAT GCC TTC TGC GAA-3’.
Immunofluorescence assay

2 wk after screening cultivation, the medium was removed. The selected hepatocyte-like cells were fixed in 4% paraformaldehyde in phosphate-buffered saline (PBS) at room temperature for 20 min. Then they were blocked with PBS containing 0.1% Triton X-100, 10% normal horse serum and 1% bovine serum albumin for 1 h. Cells next were incubated with the primary antibodies against mouse AFP, ALB (1:200) (Accurate Chemical), CK8/18 (1:200) (Lab Vision) and FXR (1:50)(R&D Systems) at 4°C overnight. After 5 washes with PBS, biotinylated secondary antibody (1:100) was added and incubated at 37 °C for 30 min. At last, SABC-Cy3 (1:100) was added and incubated at 37 °C for 30 min. Cells were washed between each step with PBS containing 0.1% bovine serum albumin. As for the control group, the same amounts of cells were collected for immunofluorescence. Cytoplasm red was positive in the fluorescence microscopy.

Glycogen dying assay

2 wk after screening cultivation, the selected hepatocyte-like cells were fixed with 95% ethanol for 10 min and rinsed in dH2O. Afterwards, cells were oxidized in 10 g/L periodic acid for 15 min, rinsed 3 times in dH2O and treated with Schiff’s reagent for 30 min, then rinsed in dH2O for 10 min again. At last, they were stained with Mayer’s hematoxylin for 1 min and rinsed in dH2O. Red granular deposits within cytoplasm with a light microscope were defined as the positive dying.

Indocyanine Green (ICG) uptake assay

Indocyanine green (ICG), a water-soluble dye that does not undergo metabolism or enterohepatic circulation, serves as a measure of hepatic function because of its exclusive elimination via biliary excretion [16]. 2 wk after screening cultivation, selected hepatocyte-like cells were washed with PBS 3 times, 2 ml ICG (Sigma) solution was added to the plates at a final concentration of 50 μM and incubated at 37 °C for 1 h, then rinsed with PBS 3 times. Cytoplasm green was positive. The control group at the same period were not used the conditioning selection media. At last we randomly counted 100 cells and calculated the ICG positive cells, each group were repeatedly done in 6 cases. Positive results to ICG dyeing were displayed with mean + standard deviation.

Assay of urea synthesis function and ammonia elimination

Level of urea nitrogen can be used as an indicator of hepatocytes’ function because of the urea synthesis capability of normal hepatocytes. Experiment was divided into five groups: mouse ESCs, cells induced for 9 days, cells induced for 23 days, cells after 2 wk screening cultivation and the group of normal hepatocytes. About 1×10⁶ cells were taken respectively with 6-well plates, and then discarded the medium; add 2ml culture medium, sucked out 10μl culture medium in the first three days for test. Colorimetric assay of cells’ urea nitrogen were detected according to the urea nitrogen assay kit (Sigma, USA) instructions.

The detoxification capability of hepatocyte-like cells was evaluated by the ammonia elimination assay. Cells were divided into five groups as above and then incubated with serum-free DMEM containing 1 mmol/L ammonium chloride for 4 hours. The decrease in ammonia concentration in the medium during a 24 h period was measured using the automatic biochemistry analyzer (HITACH). We performed all tests including the controls in triplicate.

CYP metabolism assay

For the measurement of CYP enzyme activities, the screened cells and the control cells were cultured in the medium with 50 mM 3-methylcholanthrene for 48 h. Cells were dissociated and incubated with substrate in 200 ml incubation medium at different concentrations for 3 h at 37°C. Then 800 ml cold methanol was added and centrifuged to stop the reaction. The supernatants were collected for measurement of indicated productions by LC–MS/MS (Agilent 1200 HPLC and ABI 4000 massspectrometer). Normal rat hepatocytes were used as a positive control. Total cell protein amount was used to normalize the data. Substrates and metabolic products for standard were commercially purchased of phenacetin and diclofenac (Sigma).

Animal model establishment and intrahepatic transplantation of ESCs-derived hepatocyte-like cells labeled by CFDA-SE

Carboxy fluoresce in diacetate succinimidyl ester (CFDA-SE) (Molecular Probes) was used to label the transplanted cells. Firstly, 2 wk screening cultivation cells were digested and selected with 0.25%
pancreatin and centrifuged at 1000 r/min for 5 min. Then the supernatant was removed, and the cells were re-suspended in PBS containing 10 μmol/L CFDA-SE and incubated at 37 °C for 15 min to make sure the complete absorption of fluorescent tracer. Secondly, the obtained cells were centrifuged at 1000 r/min for 5 min and the supernatant was removed, then the cells were re-suspended in PBS and incubated at 37 °C for another 30 min. Acetyl group was removed from CFDA-SE under the action of esterase in plasma, and CFDA-SE was fluorescent.

24 BALB/c mice, 6-8 wk old, weighing 20-35 g, irrespective of gender, clean grade (Guangzhou Experimental Animal Center) were selected. 12 random mice in the group A were intraperitoneally injected with 50 mg/kg retrorsine (Sigma) once every 2 wk for totally twice. 4 wk after the second injection, about 70% liver was resected. Then, about 1×10^5 hepatocyte-like cells selected by 5% cholestatic serum and labeled by CFDA-SE were transplanted into mouse liver through portal vein. On the other hand, 70% liver of another 12 mice in the group B was resected and the same ESCs-derived hepatocyte-like cells were transplanted into mouse liver. The liver samples were observed under fluorescent microscopy on wk 1 or 2.

All mice were anesthetized with diethyl ether to expose the abdomens. Blood flow of portal vein was blocked for 15 min with forceps clip, and then fluorescence-labeled cells (1×10^5) were transplanted into mouse liver through portal vein.

**Albumin immunoassay after intrahepatic transplantation**

Differentiation of donor hepatocyte-like cells was determined in recipient liver with albumin fluorescence immunoassay (double fluorescence staining). Experimental procedures: 2 wk after transplantation of ESCs-derived hepatocyte-like cells, 3 mice in each group were anesthetized and sacrificed. Liver tissue was rapidly obtained, frozen, and cut into serial sections (5 μm in thickness). The frozen sections were fixed with 4% formaldehyde, defrosted at room temperature for 15-30 min, and hydrated with PBS at room temperature in wet box for 15 min. And then, the samples were added with PBS containing 0.2% Triton X-100 (PBT) and maintained at room temperature for 20 min, added with PBT containing 0.2% BSA and 5% goat serum (or 1% BSA) and maintained at room temperature for 30 min (to block non-specific binding), added with 1 μg antibody 1 (rabbit-anti-mouse albumin monoclonal antibody dissolved in 20 μL PBS) and 5 μL antibody 2 (Zenon A reagent dissolved in antibody 1 solution) and incubated at room temperature for 5 min (albumin antibody was labeled by PE), and added with 5 μL Zenon B reagent and incubated at room temperature for 5 min to block labeling reaction. PBT (3:1) was used to dissolve fluorescent-labeled albumin antibody (immune compound of antibodies 1 and 2). Tissue sections were dripped with fluorescent-labeled albumin antibody and incubated in dark slide at room temperature for 1.0-2.0 h. Subsequently, the samples were washed with PBT twice with each for 10-15 min and with PBS twice with each for 5 min, re-fixed with 4% formaldehyde at room temperature for 15 min, and washed with PBS. Finally, the samples were put under confocal fluorescent microscope to observe fluorescent signals.

**Liver function**

Two weeks after transplantation, blood of group A and B were taken by using routine biochemical check method (Hitachi) to measure serum albumin, total bilirubin, alanine aminotransferase (ALT) and aspartate aminotransferase (AST).

**Reliability evaluation**

Embryonic stem cells-derived hepatic stem cells were poured into liver of remedial liver regeneration mice, and undifferentiated embryonic stem cells were transplanted into subcutaneous tissue in axillary region as the controls to observe neoplastic formation in embryonic stem cells-derived hepatic stem cells over 3 months.

**Statistical analysis**

Statistical analysis was manually calculated by the first author. Measurement data were compared with t test.
Results

Morphological observation

The 3-day embryoid bodies (EBs) (Fig. 1 B-a) were transferred onto six-well tissue culture dishes for further cultivating. To realize the primary differentiation of ESCs into hepatic lineages, several growth factors (including FGF-4 and HGF) and Dex were sequentially added to culture media respectively. 9 d after initial induction, a large number of hepatocyte-like cells were observed, which exhibited a characteristic of large nucleus and abundant cytoplasm outside of it, ranked as streak or Slab-like structure, and had regular triangle or multilateral contours (Fig. 1 B-b). On the other hand, after screening cultivation in a conditioning selection media containing 5% cholestatic sera, most cells were suffered from apoptosis or even death. 4d after screening cultivation in a conditioning selection media containing 5% cholestatic sera, most cells were suffered from serious apoptosis or even death. 1w later, some hepatocyte-like colony-forming units (H-CFU) were observed and the cells showed good uniformity (Fig. 1 B-e). In 2 wk, there was a trend to mature from central to round area. Peripheral cells were more close to maturity liver cells in morphology. After selected culture for 2 wk, few cells could survive in control group (undifferentiated E14 mouse ESCs). Magnifications: a–e:100×; f, g: 40×.

Immunohistochemistry

Firstly, we assessed these hepatocyte-like cells derived from ESCs by immunochemical assay. Both albumin and CK8/18 are the characteristic proteins expressed during hepatocyte
development. 9 d after initial induction, the differentiated ESCs were immunostained with anti-albumin antibody or anti-CK8/18 antibody followed by second antibody. Both albumin (A) and CK8/18 (B) positive staining (brown–yellow particals) were observed (×400). (C): After 9 days of induction, ALB and CK18 protein in HPCs could be detected, but the express level was low, while AFP protein was highly expressed. (D): 23 days after induction, the levels of ALB and CK18 protein were little high comparing with 9d induction HPCs, while AFP was almost the same. (E): 2 wk after screening cultivation, cells were regular polygons with large nucleus and abundant cytoplasm in morphology. And the levels of ALB and CK18 protein were dramatically increased than other two groups, while AFP protein fluorescence was almost invisible. Magnification: 200×. AFP: alpha fetoprotein; ALB: albumin; CK18: Cytokeratin 18.

Immunofluorescence

9 days after induction, cells ALB and CK18 proteins were lowly expressed, while AFP protein was highly expressed (Fig. 2C). 23 days after induction(cells of the control groups),
the levels of ALB and CK18 protein were little high comparing with 9d induction HPCs, while AFP was almost the same (Fig. 2D). 2 wk after screening cultivation, cells were regular polygons with large nucleus and abundant cytoplasm in morphology, and the levels of ALB and CK18 protein were dramatically increased than other two groups, while AFP protein fluorescence was almost invisible (Fig. 2E). This result revealed that after our screening cultivation, the selected cells could express hepatic protein highly.

**Detection of the expression of hepatic special genes by RT-qPCR**

mRNA levels of hepatocytes special genes expression, containing ALB, TTR, AAT, G-6-P, CK18, TAT were detected by real-time PCR. 9 days after induction, the HPCs can express the hepatocytes special genes listed above, but level was very low. Comparing with control group cells, the mRNA levels of hepatocytes special gens of experimental group cells which undergoing 2wk screening cultivation were significantly increased, what's more they were close to the level of normal liver cells (Fig. 3), especially the level of ALB reach 90% of mature hepatocytes. T-tests were carried out respectively between group 3 and 4, all the P values were less than 0.05. ALB: Albumin; TTR: transthyretin; AAT: Alanine Aminotransferase; G-6-P: glucose-6-phosphatase. CK18: Cytokeratin 18.

**ICG uptake, glycogen dying, urea synthesis function tests, ammonia elimination and CYP metabolism assay**

23 days after induction, we examined ICG uptake, which is a liver-specific function to identify differentiated hepatocytes in vitro. These cells took up ICG from the medium and showed little green, while after 2wk screening cultivation cells of the test group showed more dark green, which indicated that the selected cells had better function of ICG up-taking (Fig. 4A). Meanwhile, glycogen dying was detected after 23 days after induction. Little
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Fig. 4. Detection of the hepatic function of the induced cells. ICG uptake test and glycogen dying assay. (A): 23 days after induction, some cells could take up ICG and show little green. 2wk after screening cultivation, most cells took up ICG from the medium and showed dark green. The background colors of the two pictures were deleted. (B): Red granular deposits within cytoplasm indicated the hepatocyte-like cells of test group had the function of glycogen synthesis, the control group expressed little glycogen staining. The experimental group expressed much more glycogen staining. (C): The average ICG uptake and glycogen dying ratio of experimental groups were 76.13% and 70.22% respectively; while the control groups’ were 43.43% and 42.47% respectively ($P < 0.05$ both in ICG and PAS). (D): Urea synthesis function was detected in five groups: Group 1 stands for mouse ESCs; Group 2 represent mouse ESCs induced for 9 days. Group 3 stands for the ESCs induced for 23 days and Group 4 stand for cells after 2 wk screening cultivation, Group 5 for mature mouse hepatocytes. $* P < 0.05$. (E) For ammonia removal rate, cells were divided into 5 groups as urea synthesis. The ammonia removal activity of rat ESCs couldn’t be detected. As the induction going on, it was gradually increased. After 2 wk screening cultivation, the ammonia removal activity was 2.09±0.25µg/10⁶ cells per hour, higher than the control group and reach to 70% of mature mouse hepatocytes. (F, G): CYP metabolic activities of the induced cells and normal rat hepatocyte. Group 1 stand for the control group cells, Group 2 for the cells after 2 wk screening cultivation, Group 3 for the normal rat hepatocyte. The metabolic products of phenacetin (converted to acetaminophen by Cyp1a2; F) and diclofenac (converted to 4'-OH-diclofenac by Cyp2c enzymes; G) were determined by liquid chromatography-tandem mass spectrometry according to standard curve. $* P < 0.05$. The above results indicated that miR-122 could promote the functional maturation of HPCs. Magnification: 100×.

expression of glycogen staining in control group was detected, whereas much more red granular deposits within cytoplasm were observed in the experimental groups, indicating that the cells transfected with miR-122 had better function of glycogen synthesis (Fig. 4B). In
addition, both the ICG-positive and PAS-positive cells number of test group were higher than the control by statistic analysis (Fig. 4C). With the induction going on, the urea synthesis level was gradually increased. After 2 wk screening cultivation, urea synthesis function of cells was significantly increased, and urea synthesis level reached almost 70% of mouse normal liver cells (Fig. 4D). Similarly, the ammonia elimination rates tend to rise with culture time. After 2 wk screening cultivation, the ammonia removal rate was 2.09±0.26 µg/10^6 cells per hour, while the control group was 2.09±0.25 µg/10^6 cells per hour (Fig. 4E). Ammonia removal activity in differentiating ES cells was detected by day 21 and increased with culture time. Finally, the CYP metabolism activities of the control cells, screened cells and normal rat hepatocytes were detected by determining the metabolic products of phenacetin and diclofenac (Fig. 4F,G). The above results indicated that our screening cultivation could promote the functional maturation of HPCs.

**Hepatic FXR gene expression is increased after screening cultivation**

Recent investigations have revealed that bile acids (BA) may be essential in the initiation of the hepatoproliferative response via farnesoid X receptor (FXR) stimulation [17]. To determine whether FXR gene expression is regulated after induction and screening cultivation, hepatic FXR mRNA levels were measured by RT-qPCR. A significant increment of hepatic FXR mRNA expression was observed in screening cultivated cells compared with control group. To evaluate whether increased expression of hepatic FXR mRNA after
induction and screening cultivation is associated with altered FXR target gene expression, Cyp7a1 expression was analyzed. Liver Cyp7a1, a negative target of FXR [18], expression tended to decreased screening cultivated cells (Fig. 5A). Demonstration of FXR protein by immunofluorescence microscopy also confirmed that FXR expression was increased (Fig. 5B) after screening cultivation. These results infer the FXR pathway may be an important factor for liver cells differentiation and development.

Intrahepatic distribution and function of CFDA-SE-labeled cells after transplantation

1 wk after transplantation of CFDA-SE-labeled ESCs-derived hepatocyte-like cells in recipient mice, scattered green fluorescence was distributed in hepatic parenchyma in both groups, and density was generally coincident (Fig. 6A). 2 wk later, scattered green fluorescence was enlarged in hepatic parenchyma in the liver regeneration model + hepatocyte-like cells transplantation group, and the scattered green fluorescence arrayed like hepatic cord structure (Fig. 6B). The area of scattered green fluorescence in the partial liver resection + hepatocyte-like cells transplantation group was not significantly enlarged as compared in the liver regeneration model + hepatocyte-like cells transplantation group, but a hepatic cord-like structure was still observed (Fig. 6C). Normal liver tissue was red under fluorescent microscopy, while CFDA-SE-labeled cells were green under fluorescent microscopy. Besides, 2 wk after transplantation of CFDA-SE-labeled cells, the albumin fluorescent immunohistochemistry (double staining) demonstrated that labeled cells could
express positive signals of albumin in liver tissue of recipient mice in the two groups (Fig. 6D), which showed yellow. However, amount of positive signals in the liver regeneration model + hepatocyte-like cells transplantation group was apparently higher than in the partial liver resection + hepatocyte-like cells transplantation group.

Liver function

Two weeks after transplantation of hepatic stem cells, serum albumin levels were (29.9±1.45) g/L in the group A and (29.2±1.93) g/L in the group B, and there was no significant difference between the two groups (P > 0.05)(Fig. 7 C). However, all of the total bilirubin, ALT and AST levels of group A were lower than group B (P<0.05)(Fig. 7 D,E,F). This results suggested that the liver function of group A was better than group B.

Reliability evaluation

Transplantation of undifferentiated embryonic stem cells in the axillary region could cause formation of teratoma after 6 weeks (Fig. 7 A and B); however, pouring hepatocyte-like cells selected by a pathological microenvironmental culture system consisting of cholestatic sera into therapeutic liver repopulation model mice could not cause the formation of teratoma during 3 months.

Discussion

Although ESCs can differentiate into hepatocyte-like cells either spontaneously or after induction in vitro, the resulting cells still contain multiple heterogeneous lineages which are
not suitable for therapeutic transplantation. To avoid the risk of teratoma formation after cell transplantation, recent research highlights the importance of acquiring functional ESCs-derived hepatocyte with a uniform phenotype very much.

In this study, we first prepared embryoid bodies for the purpose to increase the interaction of ESCs and create 3 levels of mesoderm in vitro. Then we induced the ESCs to differentiate into hepatocytes with Dex, FGF-4 and HGF sequentially. The exogenous factors we selected for promoting hepatic differentiation are in accordance with the liver developmental biology research. Zaret KS et al. found that FGFs was the crucial factor to activate the liver specific gene expression and could elicit embryonic induction of the liver from the mammalian gut endoderm [19]. Hepatocyte growth factor (HGF), as a potent mitogen for hepatocytes, shows mitogenic, motogenic and morphogenic activities for a wide variety of cells that express the HGF receptor c-Met, a transmembrane protein possessing an intracellular tyrosine kinase domain [20, 21]. Dexamethasone (Dex) has the function to up-regulate GATA-4, HNF-3β and Hex during progress from endoderm to ventral foregut, besides this, it also could promote HNF-4α expression [22]. Our results showed that the EBCs could differentiate into hepatocyte-like cells 9 d after initial induction with Dex, FGF-4 and HGF, which exhibited a characteristic of large nucleus and abundant cytoplasm outside of it, ranked as streak or Slab-like structure, and had regular triangle or multilateral contours (Fig. 1). The immunostaining (Fig. 2), RT-PCR results (Fig. 3) and the hepatic-function test (Fig. 4) revealed that the ESCs-derived hepatocyte-like cells not only had the morphological character, but also expressed at least partial hepatocyte function. In short, our initial induction was effective.

At present, technologies used to screen hepatocyte-like cells from differentiated ESCs mainly include: Flow Cytometry cell sorting and magnetic activated cell sorting. However, all the methods mentioned above are either complex or have poor isolation efficiency, mainly because no very specific surface marker of hepatocyte-like cells had been found so far. Our team had reported that a conditioning selection media containing 5% cholestatic serum could select and purify the hepatic stem cells from bone marrow cells in vitro [15]. Here, we adopted the similar culture system containing 5% cholestatic serum to detect if it could select hepatocyte-like cells from differentiated ESCs initially induced with Dex, FGF-4 and HGF. Our results showed that most cells had been suffered from apoptosis or even death in the early stage (Fig. 1 B-c and B-d), but 1w later, some hepatocyte-like colony-forming unit (H-CFU) had been observed, in which most cells exhibited hepatocellular morphological characteristics and showed good uniformity (Fig. 1 B-e), while few cells could survive in control group (undifferentiated E14 mouse ESCs ) (Fig. 1 B-g). Indocyanine Green (ICG) uptake and PAS were positive for most of cells, and the positive ratio of experimental group was significantly higher than that of control group. The results of urea synthesis function tests, ammonia elimination and CYP metabolism assay also demonstrate that our selected cells have better hepatocyte function (Fig. 4). It proved our selection media containing 5% cholestatic serum could effectively select and purify the hepatocyte-like cells from differentiated ESCs in vitro.

Previous experiments had indicated the cholestatic serum contained humoral factors that had a role in recruiting putative liver stem cells and induced them to proliferate and differentiate [23]. Our previous research has demonstrated that the conditional selective medium containing cholestatic serum is optimal to selectively enrich hepatocyte-like cells from mixed differentiated ESCs [24]. Here, we consider the mechanism may be as following: In the context of cholestasis, organism will produce some humoral signal factors reactively through self-regulation to promote liver regeneration. So, there are some beneficial ingredients for liver regeneration in the cholestatic serum, such as phospholipids, cholesterol and some growth factors. On the other hand, cholestatic serum contains some toxic metabolites, such as bilirubin, cholalic acid, endotoxin, ammonia and aromatic amino acids; and these pathological toxic products are only metabolized through liver cells. For example, the hepatocyte-like cells expressing glucuronyl transferase can combine with bile acid and clear away bilirubin and synthetize urea to remove ammonia. When cultured in the cholestatic serum with appropriate concentration, only the functional hepatocyte-like
cells could dispose these toxic metabolites, survive and then selectively proliferate and differentiate in response to the signals characteristic of cholestatic serum, while the non-hepatic cells could not adapt to such a pathological environment and result in apoptosis and death. Therefore, we think the pathological serum containing 5% cholestatic serum has two function: First, it activates hepatocyte-like cells selective proliferation and differentiation signals. Second, it washes out the non-hepatocyte-like cells.

The farnesoid X receptor (FXR, also named RIP14 and HRP1), which belongs to a subcluster of metabolic receptors, could bind to DNA either as a monomer or as a heterodimer with a common partner for NRs or retinoid X receptor (RXR) to regulate the expression of various genes involved in bile acid (BA), lipid and glucose metabolisms [25]. Many ligands can activate the FXR and regulate the expression of various genes [26, 27]. Wang et al. found that activation of FXR by BAs increased the expression of FoxM1b, which is shown to regulate cell cycle progression during liver regeneration, implying that the FXR is a mediator of the effect of BA signaling on liver regeneration [28]. The FXR is a nuclear receptor highly expressed in the liver, intestine, adrenal, kidney [29]. FXR binds to various FXR response elements (FXREs) in different manners [30, 31]. Upon activation by bile acids (BA), its endogenous ligands [32], FXR down-regulates BA synthesis via suppressing the rate-limiting enzyme cholesterol 7a hydroxylase (Cyp7a1) in order to prevent BA overload and toxicity [33]. Previous data have also confirmed the essential roles of BA and FXR in lipid metabolism [34-36]. In addition the modulation of BA flux seems essential for liver regeneration to proceed normally [37]. Overall, bile acid-activated FXR controls bile acid homeostasis. Our result also confirmed the expression of FXR was significantly increased after screening cultivation with BA, and its negative target gene Cyp7a1 was decreased (Fig. 5). Based on the composition of cholestatic serum and our results, we infer the FXR pathway may also be one important factor for liver cells differentiation and development.

Therapeutic liver repopulation (TLR) by transplanted hepatocytes is a promising approach for many inborn errors of metabolism and it has a number of potential advantages over whole organ transplant [38, 39]. Under normal circumstances, hepatocytes of recipient itself proliferates precedently, so transplanted hepatocytes are not able to proliferate to reach the required amount. Otherwise, if retrorsine (RS) is used to pre-inhibit proliferation of recipient hepatocytes, followed by applying exogenous factors which can promote growth and differentiation of hepatocytes or stimulating excretion of exogenous factors through inducing liver injury, transplanted hepatocytes will be promoted to proliferate and nearly completely replace original hepatocytes [40-43]. In this study, we also adopted the similar way to establish therapeutic liver repopulation models for estimating the transplanted hepatocyte-like cells selected with a conditioning selection media consisting of 5% cholestatic sera. To be optimistic, our results demonstrated that these cells could effectively integrate into hepatic plate of host, further grow, differentiate, persist partial hepatic function and would not cause the formation of teratoma during 3 months (Fig. 6, 7). Our study explained the reliability and security of these selected ESCs-derived hepatocyte-like cells.

In summary, we established a simple and efficient method to select hepatocyte-like cells from differentiated ESCs. Our conditioning selection media consisting of 5% cholestatic serum could not only effectively select and purify hepatocyte-like cells from differentiated ESCs initially induced by Dex, FGF-4 and HGF, but also promote the growth and differentiation of hepatocyte-like cells. After intrahepatic transplantation in therapeutic liver repopulation models, the selected cells could further grow, differentiate and persist partial hepatic function without forming teratoma. But, there is still a long way to go if we want to rebuilt a liver organ with ESCs-derived hepatocyte-like cells.

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