MicroRNA-22 enhances the differentiation of mouse induced pluripotent stem cells into alveolar epithelial type II cells

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Considerable evidence has verified that microRNAs (miRNAs) play important roles in various cellular processes including differentiation. However, the regulatory roles of miRNAs involved in the differentiation of induced pluripotent stem cells (iPSC) into lung epithelial cells are still unknown. In this study, we first evaluated the current protocols to differentiate iPSC into alveolar epithelial type II (AEC II) cells, but the efficiency is low. We next identified that miR-22 can efficiently enhance the differentiation of iPSC into AEC II cells under the stimulation of proper growth factors and growing on appropriate matrix. Moreover, the AEC II cells generated from iPSC with miR-22 overexpression can proliferate and secrete lung surfactant. Here, we discovered a previously unknown interaction between miR-22 and iPSC differentiation but also provide a potential target for the effective derivation of AEC II from iPSCs for cell-based therapy.

Key words: miR-22; iPSC; differentiation; AEC II; cell-based therapy.

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Introduction

The alveolar epithelium plays an important role in remodeling, repair, and homeostasis of the lung under both physiological and pathological conditions and serves as a barrier against pathogens, allergens, and pollutants in the inspired air.1 Acute lung injuries (ALI) as well as chronic lung diseases, such as idiopathic pulmonary fibrosis (IPF) and chronic obstructive lung disease (COPD), constitute a significant health burden worldwide.2,3 Current medications can only reduce symptoms or delay disease progression. Therefore, cell-based therapies become an important new frontier in lung research and disease treatment due to the cells’ ability to orchestrate physiological processes in response to local signaling cues. One possible cell source for cell-based treatment is induced pluripotent stem cells (iPSC). AEC II cells are usually cuboidal in shape and cover only about 7% of the total alveolar surface area.4 AEC II cells are responsible for epithelium repair upon injury and ion transport. In addition, alveolar epithelial type II (AEC II) cells also contribute to lung defense by secreting lipoprotein matrix called surfactant, which is responsible for to reducing the surface tension in the alveoli. Moreover, AEC II cells can self-renew and transdifferentiate into alveolar epithelial type I (AEC I) cells. Thus, AEC II cells become an attractive candidate for cell therapies of lung diseases. MicroRNAs (miRNAs) play critical roles in various cellular processes including differentiation.5 MiR-130a has been found to control bone marrow mesenchymal stem cell differentiation towards the osteoblastic and adipogenic fate.6 MiR-690 also has been validated to regulate iPSCs differentiation into insulin-producing cells by targeting Sox9.7 Moreover, overexpression of miR-375 and miR-122 promotes the differentiation of human induced pluripotent stem cells into hepatocyte-like cells.8 However, it is still unclear through which miRNAs can promote the differentiation of iPSCs into AEC II cells.

In this study, we identified that miR-22 can promote to reducing the surface tension in the alveoli. Moreover, AEC II cells become an attractive candidate for cell therapies of lung diseases. MicroRNAs (miRNAs) play critical roles in various cellular processes including differentiation.5 MiR-130a has been found to control bone marrow mesenchymal stem cell differentiation towards the osteoblastic and adipogenic fate.6 MiR-690 also has been validated to regulate iPSCs differentiation into insulin-producing cells by targeting Sox9.7 Moreover, overexpression of miR-375 and miR-122 promotes the differentiation of human induced pluripotent stem cells into hepatocyte-like cells.8 However, it is still unclear through which miRNAs can promote the differentiation of iPSCs into AEC II cells.

In this study, we identified that miR-22 can promote the differentiation of iPSCs into AEC II cells. These iPSC-derived AEC II cells with miR-22 overexpression are capable to proliferate and secrete pulmonary surfactant. Our findings provide new insight to miR-22 functions and iPSCs differentiation, but also provide a potential target for the derivation of AEC II from iPSCs for cell-based therapy.

Materials and Methods

Culture of mouse iPSCs

The mouse iPSCs (iPS-MEF-Ng-492B-4) were purchased from RIKEN BioResource Center (Japan) and propagated on the mitomycin C (Sigma-Aldrich, Shanghai, China) inactivated Mouse Embryo Fibroblast (MEF) cells (ATCC, Manassas, VA, USA), which can be used as feeder layer to condition the medium for iPSCs. Briefly, MEF cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM) (Sigma-Aldrich, Shanghai, China) supplemented by 10% fetal bovine serum (FBS) (TBD Biotechnology Development, Tianjin, China), 50 U/mL penicillin and 50 mg/mL streptomycin (Sigma-Aldrich, Shanghai, China). After the cell division was halted with Mitomycin C (Sigma-Aldrich, St. Louis, MO), a single MEF cell suspension (5x10^5 cells) was seeded on a 0.1% gelatin (Sigma-Aldrich, St. Louis, MO, USA) coated 25 cm² flask. The next day, MEF medium was removed and 5 mL of mouse iPSC medium consisted of DMEM (high glucose, without sodium pyruvate) (Sigma-Aldrich, Shanghai, China), 15% FBS, 0.1 mM nonessential amino acids (Sigma-Aldrich, Shanghai, China), 0.1 mM 2-mercaptoethanol (Sigma-Aldrich, Shanghai, China), 1000 U/mL mouse LIF (Millipore, Billerica, MA, USA), 50 U/mL penicillin, and 50 mg/mL streptomycin was added.

Immunocytochemistry

Immunostaining was carried out to quantify iPSCs. The purified cells were seeded onto a gelatin-coated 24-well cell tissue culture plate (1 x 10^5 cells/well). On the next day, cells were washed with phosphate-buffered saline (PBS, Solarbio Science & Technology Co., Ltd., Beijing, China), fixed with 4% parafomaldehyde in PBS for 30 min at room temperature, and permeabilized with 0.1% Triton X-100 (CWBIO, Beijing, China) for 15 min. The fixed cells were blocked with 1% bovine serum albumin (Beijing Solarbio Science & Technology Co.) for 1 h and were then incubated with mouse monoclonal anti-Nanog (Cell Signaling Technology, Beverly, MA, USA) or anti-SSEA-1 (Cell Signaling Technology) at 4°C overnight. The cells were washed and incubated with Alexa-Fluor-488-conjugated anti-(mouse IgG) secondary antibodies (Cell Signaling Technology). For the AEC II cell marker, cells were incubated with mouse monoclonal mouse anti-SP-C (Santa Cruz Biotechnology, Santa Cruz, CA, USA) at 4°C overnight. The next day, cells were washed and incubated with Alexa-Fluor-546-conjugated anti-(mouse IgG) secondary antibodies (Cell Signaling Technology). The 4',6-Diamidino-2-phenylindole dihydrochloride (DAPI) (2.5 mg/mL) (Sigma-Aldrich, Shanghai, China) was used as nuclear staining.

RNA isolation and quantitative real-time PCR (qRT-PCR)

Total RNAs were extracted using RNeasy mini kit (Qigen, Beijing, China). Concentration and RNA purity were estimated by a NanoDrop ND1000 spectrophotometer (NanoDrop Technologies, Rockland, DE, USA). First-strand cDNA was synthesized by reverse transcription with first strand cDNA synthesis kit (Roche, Basel Switzerland). The specific gene expression was determined by Taqman qRT-PCR and performed on a QuantStudio 6 Flex system (Thermofisher, Gaithersburg, MD, USA). β-Actin was used as an internal control.

To quantify the mature miRNAs, total RNA was reverse transcribed using the Taqman advanced miRNA cDNA synthesis kit (Applied Biosystems, Bedford, MA, USA) according to manufacturer recommended protocols. For miRNA expression normalization, U6 small nuclear RNA (snRNA) was used as the reference control and reverse transcribed by TaqMan microRNA reverse transcription kit following manufacturer’s protocol (Applied Biosystem, Bedford, MA, USA).

All the specific primers for genes and miRNA expression are
commercially available from Applied Biosystem. The fold change of gene or miRNA transcript levels between different samples was calculated with the equation 2-ΔΔCt.10

miRNA overexpression

The lentiviral miR-22 expression vector was constructed as previous study.11 Lentiviruses were generated by co-transfecting 0.9 µg of lentiviral vector [premiR-22 or empty vectors (VC); System Biosciences, Mountain View, CA, USA] and 2.7 µg of Lenti-X HTX Packaging Mix (Clontech, Mountain View, CA, USA) in 293T cells. Supernatants were collected 48 h after transfection, filtered through a 0.45-µm membrane, tittered, and stored at -80°C until use.

Isolation of mouse primary AEC II

Mouse AEC II were isolated from 8-week male C57BL/6 mice (SLAC Laboratory Animals Company Limited, Shanghai, China) based on the previously reported procedure12,13 with modification. Mice were anesthetized and lungs were perfused with solution (10 mM HEPES, pH 7.4, 0.9% NaCl, 0.1% glucose, 5 mM KCl, 1.3 mM MgSO4, 1.7 mM CaCl2, 0.1 mg/mL streptomycin sulfate, 0.06 mg/mL penicillin G, 3 mM Na2HPO4 and 3 mM NaH2PO4, Sigma-Aldrich, Shanghai, China), followed by instilling 1 mL of above solution with dispase (500 caseinolytic units/mL; Sigma-Aldrich, Shanghai, China) directly through the trachea. The isolated lungs were pooled into a beaker containing the solution with dispase and incubated at 37°C for 45 min. After then, the lungs were chopped into small pieces and further digested with the addition of DNase I (100 μg/mL, Sigma-Aldrich, Shanghai, China) with shaking at 37°C for 45 min. Subsequently, the digested lungs were sequentially filtered through 160-, 37- and 15-µm gauge nylon mesh. The filtrate was centrifuged at 250 g for 10 min. The cell pellet was resuspended in DMEM and incubated in a 100-mm-diameter Petri dish coated with mouse IgG (75 μg/dish) for 1 hour. The cells were spun down at 250 g for 10 min and resuspended in DMEM with 10% FBS. The freshly isolated primary AEC II were used as a positive control for the subsequent proliferation and ELISA assay.

Proliferation assay

5-bromo-2-deoxyuridine (BrdU) cell proliferation assay kit (Millipore, Billerica, MA, USA) was carried out to detect cell proliferation of freshly isolated AEC II and AEC II differentiated from iPSCs by the manufacturer’s protocol.

ELISA for SPC

ELISA was performed on cell culture medium collected before and after stimulation of iPSC-AEC II or freshly isolated primary AEC II to quantify secreted SP-C using SP-C ELISA kit (Lsbio, Seattle, WA USA) following the manufacturer’s instructions. Freshly isolated primary AEC II and AEC II derived from iPSCs were cultured in 96-well plates overnight. Cells were then washed twice by PBS and DMEM with 10% FBS was added. The cells were incubated at 37°C for 30 min. Culture medium was removed as a zero-time control (0 h) for one set of wells. After then, cells were stimulated with lung surfactant secretagogues (100 µM ATP, 0.1 µM PMA and 10 µM terbutaline; ATP+PMA+Terb, Sigma-Aldrich, Shanghai, China) for 2 h at 37°C. Unstimulated cells in parallel cultures were used as controls.

Statistics

Results are shown as means ± SE (standard errors). Unpaired Student's t-test for two groups or one-way analysis of variance (ANOVA) test, followed by Tukey's multiple comparison for multiple groups were used to compare the significance of differences between the mean of different groups. SPSS (SPSS Inc. Chicago, IL, USA) was used to calculate the significance. A p value less than 0.05 is statistically significant.

Figure 1. Identification of mouse iPSCs by immunostaining after removal of MEF. Representative images (20x) for pluripotent markers, Nanog and SSEA-4, expression in the iPSC colonies. DAPI staining was used as a nuclear marker.
Results

Purity of iPSCs

Immunostaining and qRT-PCR were used to determine the characterization of iPSCs. As shown in Figure 1, immunostaining confirmed that iPSCs expressed the pluripotency markers NANOG and SSEA-4. There were around 95±1.34% and 96±0.34% Nanog and SSEA-4 positive cells after removal of MEF (Figure 1). In our studies, only iPSCs with more than 95% purity were used for the subsequent studies.

Differentiation of iPSCs to AEC II cells

Mouse iPSCs were induced to differentiate into functional AEC II cells using the protocol developed previously. Purified iPSCs were seeded on collagen IV coated 6-well plates for 24 h. Next, the iPSC medium was switched to the differentiation medium with or without Activin A for 6 days, and then with or without FGF-2 for another 5 days. As shown in Figure 2A, the levels of SP-B and SP-C mRNA were significantly enhanced after adding activin A and FGF-2 compared with the differentiation medium without activin A and FGF-2. Moreover, immunostaining analysis also confirmed that 37.21±5% of the differentiated iPSCs are positive for SP-C which is one of the AEC II cell markers (Figure 2B). In addition, we also monitored the level of Nanog mRNA in the differentiated iPSCs with Activin A and FGF as well. We found that the Nanog mRNA levels were strongly decreased in comparison with cells cultured in differentiation medium without activin A and FGF-2 (Figure 2C). These data indicated that Activin A and FGF can induce the differentiation of AEC II cells from iPSCs, but the efficiency is low.

MiR-22 efficiently enhanced the differentiation of iPSCs to AEC II in vitro

In our preliminary data, miR-22 was highly expressed in adult lungs compared to fetal lung. Since the differentiation of AEC II occurs at the late stages of fetal lung development and continues until adult, miR-22 overexpression may enhance the differentiation of iPSCs into AECs. So, we subsequently determined whether
miR-22 can enhance the differentiation efficiency of iPSCs into AEC II cells. iPSCs were seeded on collagen IV-coated plates and infected with a lentiviral virus expressing miR-22 or control vector (VC). After 24h, the cells were serum-starved overnight and then cultured in differentiation medium with Activin A for 6 days and then FGF-2 for 5 days (11 days). As shown in Figure 3A, miR-22 significantly enhanced the expression of SP-B and SP-C mRNA compared with VC by qRT-PCR for immunostaining, the SP-C-positive cells were 49.12% in cells with miR-22 overexpression as compared to 35.34% in VC (Figure 3B). The levels of mRNA expression of iPSC marker, Nanog, were also detected. We found the level of Nanog mRNA was remarkably decreased in the cells with miR-22 overexpression compared to VC (Figure 3C).

**iPSC derived AEC II are functional**

To decipher whether the differentiated ACE II cells are functional or not, we next investigated the proliferation of the iPSCs derived ACE II with miR-22 (iPSC-AEC II-miR-22) and VC (iPSC-AEC II-VC) by BrdU assay in vitro. We found that iPSC-AEC II-miR-22 cells strongly enhanced the proliferation compared with iPSC-AEC II-VC, although less compared to mouse primary AEC II cells (Figure 4A).

One of the major functions of AEC II is to secrete lung surfactant. So, we subsequently examined whether iPSC derived AEC II can secrete lung surfactant protein. As shown in Figure 4B, the SP-C secretion of iPSC-AEC II-miR-22 was significantly increased compared with iPSC-AEC II-VC after lung surfactant secretagogues (ATP, PMA and terbutaline) stimulation. Primary AEC II was used as positive control (Figure 4B). These results suggest that iPSC-AEC II-miR-22 behave similarly to the freshly isolated mouse AEC II in vitro.

**Discussion**

Alveolar epithelial injury is a hallmark of a variety of acute and chronic lung diseases. Present medications can only stabilize the disease conditions or delay disease progression. Although lung transplantation is the final treatment option for these advanced-stage lung diseases, donor organ shortage is still a major problem. Therefore, the stem cell therapy provides a new strategy for repairing severe acute and chronic lung injuries.

Numerous studies have demonstrated the therapeutic potential of iPSCs. For instance, it has been reported that a long-lasting, but finite AEC II cell line can be derived from iPSC as a novel cellular model to study alveolar epithelial cell biology in lung health and disease.14 AEC II become an attractive candidate for cell therapies of lung diseases since these cells have recently been identified as alveolar stem cells,15 and secrete a high level of lung surfactant in the distal alveoli. Human iPSCs have also been verified to derive lung progenitor and alveolar epithelial cells and can attenuate hyperoxia-induced lung injury.16 However, it is still a challenge to generate large quantities of pure AEC II. The conditions for directing iPSCs to differentiate into an alveolar epithelial lineage with homogeneity are not fully defined yet.

miRNA causes a fast response in protein expression based on inhibition of mRNA translation and stability. Their ability to regulate different proteins can rapidly impose a dominant phenotypic change in cell destination. It has been reported that miRNAs can affect the differentiation of iPSCs. Inhibition of microRNA-495 has been found to enhance therapeutic angiogenesis of human iPSCs.17 MiR-140 has also been validated to enhance chondrogenesis differentiation of human iPSCs with transforming growth factor beta 3 (TGFβ3).18

In this study, we used a two steps method to differentiate AEC II from mouse iPSCs and found that miR-22 can effectively enhance the differentiation of mouse iPSCs into AEC II. Moreover, these iPSC-AEC II with miR-22 overexpression are capable of proliferating and secreting lung surfactant. However, it is still worthy verifying the downstream targets and signaling pathways of miR-22, and considering the in vivo study in the future. Our studies represent an initial and important step towards the ultimate goal of cell-based therapy of lung injuries.

**References**


