Video Article Initiating Differentiation in Immortalized Multipotent Otic Progenitor Cells

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Abstract

Use of human induced pluripotent stem cells (iPSC) or embryonic stem cells (ESC) for cell replacement therapies holds great promise. Several limitations including low yields and heterogeneous populations of differentiated cells hinder the progress of stem cell therapies. A fate restricted immortalized multipotent otic progenitor (iMOP) cell line was generated to facilitate efficient differentiation of large numbers of functional hair cells and spiral ganglion neurons (SGN) for inner ear cell replacement therapies. Starting from dissociated cultures of single iMOP cells, protocols that promote cell cycle exit and differentiation by basic fibroblast growth factor (bFGF) withdrawal were described. A significant decrease in proliferating cells after bFGF withdrawal was confirmed using an EdU cell proliferation assay. Concomitant with a decrease in proliferation, successful differentiation resulted in expression of molecular markers and morphological changes. Immunostaining of Cdkn1b (p27^{KIP}) and Cdh1 (E-cadherin) in iMOP-derived otospheres was used as an indicator for differentiation into inner ear sensory epithelia while immunostaining of Cdkn1b and Tubb3 (neuronal β-tubulin) was used to identify iMOP-derived neurons. Use of iMOP cells provides an important tool for understanding cell fate decisions made by inner ear neurosensory progenitors and will help develop protocols for generating large numbers of iPSC or ESC-derived hair cells and SGNs. These methods will accelerate efforts for generating otic cells for replacement therapies.

Video Link

The video component of this article can be found at <http://www.jove.com/video/53692/>

Introduction

The organs of the inner ear, the cochlea, utricle, saccule and three semicircular canals, mediate the ability to hear and balance. Within the cochlea, hair cells convert sounds into electrical signals that are relayed to the spiral ganglion neurons (SGN). The SGNs fire action potentials to propagate neural signals through the auditory circuit. Genetic mutations, ototoxic drugs and exposure to loud sounds contribute to hair cell and SGN death that result in hearing loss¹⁴. Once lost, these cells are not replaced. Use of iPSC and ESC to generate nascent hair cells or SGNs holds great promise for inner ear cell replacement therapies⁵⁻⁸. A flurry of progress has shown that pluripotent stem cells and inner ear derived progenitors can differentiate into hair cells and SGNs at various stages of maturity. Mammalian embryonic stem cells (ESC) and
induced pluripotent stem cells (iPSC) can be used to generate functional hair cells the mammalian inner ear have also been shown to form hair cells and neurons with properties of their *in vivo* cellular counterparts¹ .

Use of iPSC or ESC-derived otic progenitors to replace lost hair cells and SGNs requires efficient differentiation. Improper differentiation or continued proliferation of engrafted stem-derived progenitors in the inner ear can exacerbate inner ear function and pose a tumorigenic risk
such as teratomas formation in the inner ear¹⁷. There is a clear need for devel progenitors. One strategy in developing these methods is to recapitulate cell fate decisions made by neurosensory progenitors during inner ear development. Protocols that prevent proliferation and direct otic progenitors into hair cells or SGNs will help improve safety as well as efficacy of replacement therapies.

During development, the inner ear begins with the thickening of surface ectoderm in a restricted region between rhombomeres 5 and 6 to become the otic placode. As the otic placode invaginates to form an otic cup, a collection of cells in the anterior region of the otic cup gives rise to the neural-sensory-competent domain (NSD), which contains precursors of hair cells and neurons of the inner ear¹⁸. Fate mapping studies from mouse, chicken and zebrafish developing inner ear suggest multiple populations of neurosensory progenitors that give rise to the sensory
hair cells, surrounding supporting cells and otic neurons¹⁹⁻²². The high mobil ear result in the loss of the hair cells, supporting cells and SGNs in the cochlea^{25,26}.

To study otic progenitor cells undergoing cell fate decisions, a fate restricted immortalized multipotent otic progenitor (iMOP) cell line from Sox2 expressing cochlear progenitors was previously established. iMOP cells were originally derived from embryonic E12.5-13.5 cochlea and infected with a c-Myc retrovirus²⁷. IMOP cells can continually proliferate as colony forming cells known as otospheres and have the capacity to differentiate into hair cells, supporting cells and SGNs²⁷. Understanding the capacity of iMOP cells to differentiate into distinct otic lineages allows application of these findings to efficiently generate iPSC or ESC-derived hair cells and SGNs. Efficient differentiation protocols will open new avenues for cell replacement therapies of inner ear diseases that are recalcitrant to conventional treatments. A crucial issue in generating otic cells by *in vitro* cell culture is to have differentiation markers that help determine if cells are undergoing differentiating. Cdkn1b (p27^{KIP}) has been extensively used as an early marker for differentiation in developing inner ear, however, expression of Cdkn1b in iMOP cells and how it correlates to differentiation has not been addressed. In this study, the current culture conditions and how Cdkn1b expression correlates to other markers of iMOP differentiation are described.

Protocol

1. Maintaining Self-renewal in IMOP Cells

- 1. Prepare iMOP culture media: DMEM/F12, 1X B27 supplement, 25 µg/ml carbenecillin and 20 ng/ml bFGF. Make 50 ml of iMOP culture media using sterile reagents. Warm up 49 ml of DMEM/F12 in a 50 ml conical in a 37 °C water bath.
	- 1. Thaw 50X B27 supplement and filter-sterilized 100 mg/ml carbenecillin aliquots for 5 min in a 37 °C water bath. Thaw out 100 µg/ml bFGF aliquot at RT. Add 1 ml 50X B27, 10 µl of 100 µg/ml bFGF and 12.5 µl of 100 mg/ml carbenecillin into DMEM/F12.
- 2. Use 3 ml of media in a 60 mm tissue culture dish for culturing iMOP cells. Add fresh media to cultures every other day by doubling the volume of media. Ensure that concentration of bFGF does not drop below 5 ng/ml.
	- 1. Culture iMOP cells at 37 °C with 5% CO2. Passage cells after 5 7 days in culture as listed in the steps below. Transfer culture to a 15 ml conical using a 10 ml pipette tip.
- 3. Make 1 mM EDTA in HBSS by diluting 0.1 ml of 0.5 M EDTA pH 8.0 into 50 ml of HBSS. Pre-warm 1 mM EDTA made in HBSS in a 37 °C water bath.
- 4. Harvest cells by gravity sedimentation or centrifugation at 200 x g for 5 min. at RT. For gravity sedimentation, place the 15 ml conical containing the cultures in the 37 °C incubator for 5 - 10 min. After that time, observe the otospheres collect at the bottom of the 15 ml conical. NOTE: Excessive centrifugal force can cause cell damage.
- 5. Carefully aspirate spent media using a 2 ml aspirating pipette without disturbing the cell pellet. Add 0.5 ml of pre-warmed 1 mM EDTA HBSS solution to cell pellet. Using a P1000 pipette, gently pipet up and down 2 - 3 times. Place conical at 37 °C and incubate for <5 min to facilitate dissociation into single cells.
- 6. Gently swirl the cell solution to determine if otospheres are dissociated. If no otospheres sediment to the bottom of the conical, the cells are dissociated. The time of incubation may vary.
- NOTE: Prolonged incubation with EDTA will result in excessive cell death.
- 7. Remove cells from 37 °C, add 2 ml of media to neutralize and dilute the EDTA. Collect the cells by centrifugation at 200 x g for 5 min at RT. Aspirate out diluted EDTA using a 2 ml aspirating pipette and add 5 ml of 1X PBS to wash the cells.
- 8. Spin cells down at 200 x g for 5 min at RT and aspirate out 1X PBS using a 2 ml aspirating pipette. Resuspend the cells using a P1000 pipette in 0.5 ml of iMOP culture media by gently pipetting up and down $\tilde{2}$ - 3 times.
- 9. Count cells using a microfluidic particle counter with the appropriate cassettes²⁸. A confluent plate of iMOP cells will contain ~6 X10⁶ cells. Dilute cells 1:100 in media and add 75 ul of cell solution into the cassette for counting. Plate 1 x 10⁶ cells in a 6 cm dish in iMOP culture media (~1:10 dilution). Passage iMOPs every 5 - 7 days. NOTE: Cells that form large otospheres or attach to the bottom of the tissue culture dish will differentiate. Cells will also die if the cultures are over confluent.

2. Freezing and Thawing IMOP Cells

- 1. Prepare synthetic freezing medium by thawing and equilibrating the solution to 4 °C prior to freezing cells.
- 2. Collect cells from a confluent 6 cm dish of iMOP cells. Use a 10 ml pipette and transfer cells (~6 8 x 10⁶ cells) into a 15 ml conical.
- 3. Harvest cells by gravity sedimentation or centrifugation at 200 x g for 5 min at RT. If the otospheres are small, cells will not settle by gravity sedimentation. Optional: Count cells using step 1.9 to determine total cell numbers.
- 4. Aspirate spent media using a 2ml aspirating pipette while leaving the loose cell pellet behind.
- 5. Add 0.25 ml of synthetic freezing media to resuspend cells at a density of ~5 x 10⁵ to 3 x 10⁶ cells/ml. Gently pipette cells with a P1000 pipette. Transfer the cell suspension to cryogenic vials using a P1000 pipette with 1 ml filtered tip.
- 6. Place the vials in an alcohol free freezing container and place the freezing container at -80 °C to reduce the temperature 1 °C per minute until the temperature reaches -80 °C.
- 7. For long-term storage of cells, transfer the vials to the vapor phase of a liquid nitrogen storage tank. To thaw out cells for culture, equilibrate cryogenic vial containing frozen cells at -80 °C.
- 8. Pre-warm iMOP culture media in a 37 °C water bath.
- 9. Thaw the frozen vial quickly by swirling the bottom of the vial in a 37 °C water bath. Add 1 ml of pre-warmed iMOP culture media to thawed cells once the last ice crystal disappears. Transfer cells into a 15 ml conical and add an additional 4 ml of iMOP culture media
- 10. Spin the 15 ml conical for 5 min. at 200 x g and aspirate the spent media. Resuspend the cells in 2 ml of iMOP culture media and plate cells in a 6 cm dish. Incubate cultures at 37 $^{\circ}$ C with 5% CO₂ for expansion.

3. Differentiating IMOP Cells into Sensory Epithelia

1. Prepare 50 ml of iMOP sensory epithelia differentiation media: DMEM/F12, 1X B27 supplement, 25 µg/ml carbenecillin. Make 50 ml of iMOP sensory epithelia differentiation media. Warm up 49 ml of DMEM/F12 in a 50 ml conical in a 37 °C water bath. Thaw 50X B27 supplement and 100 mg/ml carbenecillin aliquots for 5 min. in a 37 °C water bath. Add 1 ml 50X B27 and 12.5 µl of 100 mg/ml carbenecillin into DMEM/ F12.

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- 2. To harvest, dissociate, resuspend and count cells repeat steps 1.4-1.9
- 3. Plate 1 x 10⁶ cells in a 6 cm dish at Day -3 using iMOP culture media. On Day 0, transfer cultures using a 10 ml pipette into a 15 ml conical. Collect the otospheres by gravity sedimentation as stated in step 1.6. Aspirate out spent media using a 2 ml aspirating pipette and leave the otospheres on the bottom of the conical.
- 4. Gently add in 2 ml of sensory epithelia differentiation media. Transfer otospheres to a 6 cm dish using a large bore 10 ml pipette. NOTE: Mechanical shearing from harsh pipetting can dissociate cells from the otospheres.
- 5. Add 2 ml of fresh sensory epithelial differentiation media to cultures every other day. If necessary, cells can be collected by gravity sedimentation and media replaced.
- 6. Collect otospheres at Day 10 by transferring to a 15 ml conical and allowing the otospheres to sediment as stated in step 1.6.Aspirate the spent media using a 2 ml aspirating pipette and leave the otospheres undisturbed.
- 7. Fix otospheres by incubating in 4% formaldehyde in 1X PBS for 15 min at RT. Remove formaldehyde solution, wash otospheres with wash buffer (1X PBS containing 0.1% TritonX-100) before incubating in blocking buffer (1X PBS containing 10% normal goat serum and 0.1% Triton X-100) for 1 hr.
	- 1. Replace buffer and incubate samples in blocking buffer with diluted antibody. Incubate at 4 °C. Wash samples with 1X PBS containing 0.1% Triton X-100 subject the otosphere to immunostaining²⁷. Transfer otospheres into 1X PBS and place otospheres into mounting media on a glass slide using a P1000 pipette. Put a coverslip over the sample and allow mounting media to dry at 4 °C.
- 8. Acquire epifluorescence images using an inverted microscope setup equipped with a 16 bit CCD camera and either a 20X 0.75 air or a 40X 1.3 NA oil immersion objective. Collect fluorescence from different color channels using the listed (excitation and emission) wavelengths: blue $(377 \pm 25 \text{ nm}/447 \pm 30 \text{ nm})$, green $(475 \pm 25 \text{ nm}/540 \pm 25 \text{ nm})$, red $(562 \pm 20 \text{ nm}/$ and $625 \pm 20 \text{ nm}$) and infrared $(628 \pm 20 \text{ nm}/$ and 692 ± 20 nm).

4. Assaying EdU Incorporation

- 1. On Day -3, plate 1 x 10⁶ iMOP cells in a 6 cm dish. Three days later on Day 0, harvest, dissociate, resuspend and count cells by repeating steps 1.4-1.9.
- 2. Plate 2.5 x 10⁵ cells in iMOP culture media and 5 x10⁵ cells in sensory epithelia differentiation media in different wells of a 6 well dish.
- 3. Determine the percentage of cells in S phase by EdU incorporation on day 3 by using an EdU incorporation assay
	- 1. Add EdU stock directly to the iMOP cultures to obtain a final concentration of 1 µM EdU in the culture media.
		- 2. Incubate EdU in iMOP cultures for 2 hr and incubate at 37 °C with 5% $CO₂$. Harvest, dissociate and collect cells repeat steps 1.4-1.9. Add in 4% formaldehyde in 1X PBS for 15 min. at RT to fix cells. Harvest cells by centrifugation at 200 x g for 5 min at RT. Remove formaldehyde solution and properly dispose.
		- 3. Label nuclei of cells with Hoechst and incorporated EdU with green fluorescence dye-azide according to manufacturer's protocol. NOTE: All washes are done with 1X PBS, 3% BSA and 0.1% Tween 20. Wash cells with 1X PBS twice.
		- 4. Mount cells on a slide using antifade reagent and place a 1.5 cover glass over the sample. Acquire fluorescent images of labeled cells using an epifluorescence microscopy.

5. Differentiating IMOP-Derived Neurons

- 1. Make 50 ml of neuronal differentiation media: Neurobasal media, 1X B27 supplement, 2 mM L-Glutamine. Thaw bottle of 50X B27 and 200 mM L-Glutamine in 37 °C water bath for 5 min. Add 1ml 50X B27 and 0.5 ml 200 mM L-Glutamine to 48.5 ml of Neurobasal media.
- 2. Coat coverglass by placing 12 mm round 1.5 glass coverslips in a sterile 10 cm plate and add 70% EtOH to the plate to sterilized and clean the coverslips.
- 3. Gently agitate the coverslips to ensure they are covered in ethanol. Leave the plate for 10 min at RT. Rinse the coverslips 3 times with sterile 1X PBS to wash out the remaining ethanol. Rinse the coverslip once with sterile H₂0 to wash out remaining 1X PBS.
- 4. Aspirate the H20 using a 2 ml aspirating pipette and let the coverslips dry. Expose the coverslips to UV light in the tissue culture hood for 15 min. Store coverslips in a sterile environment if not immediately used.
- 5. Place one 12 mm round coverslip in each well of a 24 well dish. Shake plate gently to ensure that the coverslips lie flat on the bottom of the well.
- 6. Thaw 2 mg/ml poly-D-lysine stock solution at RT and dilute to a 10 µg/ml poly-D-lysine concentration in 1X PBS. Add 0.25 ml of 10 ug/ml poly-D-lysine to the wells. Leave the plate in the 37 °C incubator for 1 hr.
- 7. Wash the wells 3 times with sterile 1X PBS. Thaw 10 mg/ml laminin stock solution at RT and dilute to 10 µg/ml in 1X PBS. Add 0.25 ml of 10 µg/ml laminin working solution into a single well of a 24 multi-well dish. Incubate the plate at 37 °C incubator. Aspirate out the laminin solution using a 2 ml aspirating pipette.
- 8. Wash the coverslip 3 times with 1X PBS by adding in 1 ml of 1X PBS with a P1000 pipette and removing the 1X PBS by aspirating with a 2 ml aspirating pipette. Leave 1X PBS from last wash in the well until cells are ready to be plated. Initiate neuronal differentiation by harvesting, dissociating and counting cells in steps 1.4-1.9. Plate 1 x 10⁶ cells in a 6 cm dish at Day -3.
- 9. On Day 0, harvest, dissociate and count cells by repeating 1.4-1.9. Seed 1 x 10⁵ 1.5 x 10⁵ iMOP cells into 0.5 ml pre-warmed neuronal
- differentiation media per well in a 24 multi-well dish. Aspirate and add pre-warmed neuronal differentiation media to cultures every other day. 10. Fix iMOP-derived neurons in 4% formaldehyde for 15 min. at RT on Day 7. Remove formaldehyde solution, wash iMOP-derived neurons with wash buffer (1X PBS containing 0.1% TritonX-100) before incubating in blocking buffer (1X PBS containing 10% normal goat serum and 0.1% Triton X-100) for 1 hr.
	- 1. Replace buffer and incubate samples in blocking buffer with diluted antibody. Incubate at 4 °C. Wash samples with 1X PBS containing 0.1% Triton X-100 subject cells to immunostaining²⁷. Wash the coverglass with 1X PBS before placing onto mounting media. Allow the mounting media to dry at 4 °C before acquiring epifluorescence images as listed in step 3.8.

Representative Results

bFGF Withdrawal Decreases Proliferation in IMOP Cells

To decrease the proliferative capacity of iMOP cells and initiate differentiation of iMOP cells, bFGF was withdrawn from the cultures. To confirm that growth factor withdrawal decreases proliferation, EdU incorporation was employed as a proliferation assay. The percentage of cells that incorporated EdU from otospheres cultured with iMOP culture media (containing bFGF) and otospheres cultured in sensory epithelia media (without bFGF) was compared. Otosphere cultures were pulsed with the nucleotide analog EdU, harvested and fixed. Incorporated EdU nucleotides were fluorescently labeled with Alexafluor 488 azide using click-chemistry. Cells from otospheres grown in the absence of bFGF showed decreased incorporation of EdU relative to cells cultured with bFGF (**Figure 1A-F**). As the size of the otosphere increased, it was increasingly difficult to visualize all the cells from the otosphere with an epifluorescent microscope. To address this, cells were dissociated and mounted so that they can be unambiguously visualized *en mass*. Even after dissociation and fixation, the cells re-aggregated. To maintain the cells in a dissociated state, detergents that prevented the cells from re-aggregating were tested. Tween 20 was one of the detergents that prevented re-aggregation of the cells. Dissociated cells were washed with 0.1% Tween 20 and mounted onto slides. EdU labeled cells were then visualized by epifluorescence microscopy (**Figure 1G,H**). Representative results from individual experiments (n = 5) showed a significant decrease in the percentage of EdU labeled cells from 48% to 19% after 3 days after bFGF withdrawal (p<0.005). These results confirm a dramatic drop in the percentage of S phase cells and proliferative potential of iMOP cells after bFGF withdrawal.

IMOP-derived Sensory Epithelia Express Cdkn1b and Show Morphological Changes

A timeline of iMOP sensory epithelia differentiation protocol is shown (**Figure 2A**). Otospheres from proliferative cultures were dissociated into single cells and allowed to recover for 3 days in iMOP culture media. This method helps enrich for proliferating cells and selects against postmitotic cells in these starting cultures. After 3 days, newly formed otospheres were seeded in sensory epithelia differentiation media and allowed to undergo unguided differentiation for 10 days. Bright field images of typical cultures containing otospheres at different time points after seeding showed an initial decrease in size before otospheres start increasing in size (**Figure 2B-E**).

Since brightfield microscopy cannot reveal many of the changes that occur during differentiation of iMOP cells, immunostaining with molecular markers was used to highlight morphological and molecular features of the differentiated cells²⁷. To determine if expression of Cdkn1b (p27^{KIP}) can be used as a marker for differentiation, otospheres from proliferative or sensory epithelia differentiated iMOP cultures were compared. Fluorescent markers were visualized and captured by epifluorescence microscopy. Representative images of otospheres from proliferative cultures showed low expression of Cdkn1b outside the nuclei with almost no phalloidin staining (**Figure 3 A-D)**. Otospheres cultured in sensory epithelia differentiation media displayed increased expression of nuclear Cdkn1b concomitant with the appearance of phalloidin labeling in the peripheral edges of cells (**Figure 3 E-H**). In proliferating iMOP otospheres, Cdh1 (E-cadherin) and phalloidin are weakly labeled (**Figure 3 I-L**). In differentiated iMOP otospheres, pronounced phalloidin and Cdh1 labeling highlight the morphological changes in actin filaments and regions of cell-cell adhesion (Figure 3 M-P), similar to previous results²⁷. During sensory epithelia differentiation, the morphological changes in actin filaments parallel the appearance of Cdh1 expression in adhesion sites between cells. In this protocol, the increase in Cdkn1b expression along with the changes in phalloidin and Cdh1 serve as early indicators of sensory epithelia differentiation in iMOP cells.

IMOP-derived Neurons Express Cdkn1b and Tubb3

A schematic of iMOP neuronal differentiation protocol is shown (**Figure 4A**). Similar to the aforementioned protocol, iMOP cells were dissociated and allowed to recovery for 3 days in iMOP culture media. Otospheres were harvested, dissociated into single cells and seeded onto poly-D-lysine and laminin coated coverslips. The cells were allowed to differentiate for 7 days. Representative bright field images at time points displayed progressive morphological changes as they differentiated into iMOP-derived neurons (**Figure 4B-E**). By Day 7, long neurites can be seen extending from cell bodies (**Figure 4E** arrowheads). To determine the changes in expression levels of Cdkn1b during neuronal differentiation, proliferating iMOP cells and iMOP-derived neurons were compared. Otospheres were labeled with Hoechst and Cdkn1b antibodies. iMOP cells from proliferating otospheres had few cells with low nuclear Cdkn1b expression (**Figure 4 F-H**). Furthermore, iMOPderived neurons showed an increase in the number of cells with nuclear Cdkn1b expression 7 days after neuronal differentiation (**Figure 4 I-K**). To determine if the Cdkn1b cells were adopting a neuronal lineage, labeling of iMOP-derived neurons with the neuronal marker Tubb3 was done. Representative images of iMOP-derived neurons showed co-labeling of Cdkn1b and Tubb3 (**Figure 5 A-D**). Magnification and quantification of neurites from these cells showed an average 1.5 neurites associated with each Tubb3 labeled cell (n = 60) (**Figure 5 E-G**). In our neuronal differentiation protocol, immunostaining with Cdkn1b and Tubb3 can be used as an indicator of differentiation into bipolar or pseudo-unipolar iMOP-derived neurons.

Nuclear Cdkn1b Expression Levels Increase after Initiating Differentiation

The morphological changes in otospheres show qualitative changes in iMOP cells as they undergo differentiation. To attain quantitative results from the immunofluorescence images to denote early differentiation events, nuclear fluorescence intensity of Cdkn1b from individual proliferating iMOP cells (n = 239) and iMOP cells undergoing sensory epithelia differentiation (n = 246) were measured. To normalize Cdkn1b fluorescence signals from independent experiments, the ratio of Cdkn1b to Hoechst fluorescence from individual cells was determined. Histograms of the normalized fluorescence intensity of Cdkn1b from proliferating iMOP cells (black) and sensory epithelia differentiating iMOP cells (red) were plotted (**Figure 6A**). An increase in the number of high Cdkn1b expressing cells was observed as a rightward shift of the histogram for sensory epithelia differentiation (red) relative to the histogram for proliferating iMOP cells (black). To determine the increase in the percentage of cells expressing Cdkn1b, a threshold for normalized Cdkn1b fluorescence intensity units (0.65) was set (arrowhead) and the percentage of cells above the threshold was determined. After sensory epithelia differentiation, iMOP cells displayed an increase from 25.3% to 67.1% of cells expressing Cdkn1b (**Figure 6B**). The same quantitative analysis was applied to iMOP-derived neurons. When comparing proliferative iMOP cells (n = 525) and 7 day iMOP-derived neurons (n = 583) a rightward shift in the histogram associated with iMOP-derived neurons was observed relative to the histogram for proliferating iMOP cells (**Figure 6C**). Determining the percentage of cells above the threshold revealed an increase in Cdkn1b expressing cells from 23.5% to 85.5% (**Figure 6D**). Using the described protocols, quantitative analysis of Cdkn1b expression confirmed an increase in the number of cells that upregulate Cdkn1b during sensory epithelia and neuronal differentiation. The increased numbers of Cdkn1b expressing cells can be used to confirm differentiation of iMOP cells in these protocols.

Figure 1. EdU Incorporation as an Assay to Determine Proliferative State of IMOP Cultures. iMOP-derived otospheres cultured in the presence of bFGF. Nuclei of cells were labeled with (**A**) Hoechst and (**B**) EdU Alexafluor 488. (**C**) Merged image of Hoechst and EdU fluorescence. Otospheres cultured 3 days in media lacking bFGF. Cells from cultures were labeled with (**D**) Hoechst and (**E**) EdU Alexafluor 488. (**F**) Merged images of Hoechst and EdU labeling. Merged fluorescence images of Hoechst (magenta) and EdU (green) labeled cells after dissociating otospheres and washing with 1X PBS containing 0.1% Tween 20. Cells were from otospheres cultured (**G**) in the presence of bFGF or (**H**) absence of bFGF. Length of scale bars are 10 µm unless indicated.(**I**) Percent of EdU labeled cells from iMOP cells cultured in the presence or absence of bFGF. The number of individual cells analyzed was denoted within the bar graph and error bars were depicted as standard error of the mean (SEM). Cell counts were from n = 5 independent experiments. The Student's t-test was done to determine statistical significance. [Please click here to view a larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig1large.jpg)

Figure 2. General Schematic of Sensory Epithelia Differentiation. (**A**) Timeline for differentiating iMOP cells into sensory epithelia. The line graph denotes the time of cell dissociation and media changes. (**B**) Typical phase contrast images of otospheres on Day 0, (**C**) 3, (**D**) 7 and (**E**) 10 during unguided sensory epithelia differentiation. Length of scale bar is 100 µm. [Please click here to view a larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig2large.jpg)

Figure 3. Expression of Markers Indicative of Cell Cycle Arrest and Differentiation in IMOP-derived Sensory Epithelia. Otosphere of iMOP cells cultured in iMOP culture media. Nuclei of cells were labeled with (**A**) Hoechst and (**B**) Cdkn1b antibody. Filamentous actin was labeled with (**C**) phalloidin. (**D**) The merged image of a typical otospheres containing proliferating iMOP cells. Otospheres from iMOP cells were cultured in sensory epithelia culture media for 10 days. Cells from otopsheres were marked with (**E**) Hoechst, (**F**) Cdkn1b and (**G**) phalloidin. (**H**) Merged image of otospheres showing increased Cdkn1b and phalloidin labeling. Proliferating otospheres were labeled with (**I**) Hoechst, (**J**) Cdh1, (**K**) phalloidin. (**L**) The merged image shows weak fluorescence from these markers. Otospheres differentiated into sensory epithelia were also labeled with (**M**) Hoechst, (**N**) Cdh1, (**O**) phalloidin. (**P**) The merged image shows typical strong Cdh1 and phalloidin labeling. Length of scale bars are 10 µm unless indicated. [Please click here to view a larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig3large.jpg)

Figure 4. General Schematic of Neuronal Differentiation. (**A**) Timeline for differentiating iMOP cells into neurons. Phase contrast images of iMOP cells undergoing neuronal differentiation at (**B**) Day 1, (**C**) 3, (**D**) 5 and (**E**) 7. Arrowheads point to neurites. Fluorescence images of iMOP cells cultured as otospheres in iMOP culture media labeled with (**F**) Hoechst and (**G**) Cdkn1b. (**H**) Merged image of cells labeled with Hoechst and Cdkn1b. Fluorescence images of iMOP cells after 7 days of culture in neuronal differentiation media. Nuclei of cells were labeled with (**I**) Hoechst and (J) Cdkn1b. (K) Merged image of cells labeled with Hoechst and Cdkn1b. Length of scale bars are 10 µm unless noted. [Please](https://www.jove.com/files/ftp_upload/53692/53692fig4large.jpg) [click here to view a larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig4large.jpg)

Figure 5. Expression of Molecular Markers Indicative of Cell Cycle Exit and Neuronal Differentiation. *iMOP cells 7 days after neuronal* differentiation. Nuclei were labeled with (**A**) Hoechst, (**B**) Tubb3 (neuronal β-tubulin) antibodies to highlight cellular morphology and (**C**) Cdkn1b antibodies. (D) Merged image with labeling of Cdkn1b and Tubb3 in individual cells. Length of scale bar is 20 um. [Please click here to view a](https://www.jove.com/files/ftp_upload/53692/53692fig5large.jpg) [larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig5large.jpg)

Figure 6. Single Cell Quantitative Fluorescence Intensity Analysis of Cdkn1b. (A) Normalized fluorescence intensity of Cdkn1b expression was determined by calculating the ratio of Cdkn1b and Hoechst fluorescence intensity from individual cells. The normalized fluorescence intensity was plotted relative to cell numbers as a histogram. Normalized fluorescence intensity of Cdkn1b from proliferating iMOP cells cultured in the presence of bFGF (black) and iMOP otospheres differentiated into sensory epithelia (SE) (red) are shown. A threshold was set at 0.65 normalized fluorescence intensity units (arrowhead) (**B**) Percentage of iMOP cells expressing Cdkn1b above the threshold value, from proliferating (black) and sensory epithelia differentiated cultures (red). (**C**) Merged fluorescent image of sensory epithelia differentiated iMOP cells labeled with Hoechst and Myo6 antibodies. (**D**) Normalized fluorescence intensity of Cdkn1b expression from proliferating iMOP cells (black) and neuronal differentiated iMOP cells (red) were plotted and a threshold set (arrowhead) as described above. (**E**) Percentage of iMOP cells expressing Cdkn1b from proliferating (black) and neuronal differentiating cultures (red). Individual cells used for analysis were denoted in each bar graph. Results were compiled from different experiments (n = 3) and error bars depicted as SEM. The Student t-test was used to determine the statistical significance. (F) Merged fluorescent image of bipolar or pseudo-unipolar iMOP-derived neurons labeled with Hoechst and Tubb3. Length of scale bars are 20 µm. [Please click here to view a larger version of this figure.](https://www.jove.com/files/ftp_upload/53692/53692fig6large.jpg)

Table 1. Cell Numbers for Maintenance of Differentiation of IMOP Cells in Various Tissue Culture Plate Formats.

Discussion

Monitoring IMOP Cultures

A protocol for maintaining self-renewal and promoting differentiation of a novel iMOP cell line is described and additional plating formats are included (**Table 1**). Several critical steps that help with routine expansion and differentiation of iMOP cells are noted. Similar to pluripotent stem cell cultures, iMOP cells theoretically have an indefinite life-span. To ensure that cell lines are properly maintained, iMOP cultures are routinely monitored for cell viability, self-renewal and differentiation potential. To determine cell viability after dissociation, we employ trypan blue staining. In general, over 70% of cells are viable after dissociation. To monitor self-renewal in iMOP cells, immunostaining with c-Myc and Sox2 antibodies is done. Under proliferative culture conditions, iMOP cells have a doubling time of ~18 hr. Alterations in expression of self-renewal markers or doubling time are potential indicators of altered self-renewal. To monitor the differentiation potential of iMOP cells, cell cycle exit and expression of early stage markers for sensory epithelial or neurons are used to evaluate the differentiation potential of the cells. If iMOP cells do not pass any one of the criterions described above, it would be advisable to terminate the cultures.

Variables that Affect IMOP Cultures

One critical step in culturing iMOP cells is to maintain an active concentration of bFGF in the cultures to promote self-renewal. bFGF is reportedly
highly labile at 37 °C when culturing cells^{29,30}. Growth of cultures ca instability. In the current protocol, fresh media is constantly supplied to the cultures in order to maintain bFGF levels above 5 ng/ml. Stabilization of bFGF levels using controlled release from glycolic and lactic acid polyester microspheres could reduce the frequency of media changes
needed to maintain steady levels of bFGF in media³¹. Addition of the steadily relea maintain bFGF levels in iMOP cultures.

Another critical step in maintaining viability of iMOP cells is to limit exposure to dissociation reagents and mechanical shearing. In the current protocol, dissociation is accomplished by incubation with 1 mM EDTA. Curbing excessive incubation time with EDTA, reducing excessive mechanical shearing caused by vigorous pipetting and limiting multiple centrifugation steps when passaging iMOP cells will help increase cell viability. Attention given to these steps can effectively maintain viability of large numbers of cells during expansion of cultures.

During the routine culture of iMOP cells, several methods for dissociating the cells were tested. Mechanical shearing from pipetting can be used to dissociate otospheres, but does not uniformly or consistently produce single cells. Use of commercially available reagents for cellular dissociation of iMOP cells can effectively generate single cells from otospheres, but these reagents affect the adhesive properties of cells to reform otospheres and to adhere to treated surfaces. In order to use commercial reagents, multiple washes were required to remove the dissociation reagents before employing differentiation protocols.

In the current differentiation protocols, both sensory epithelia and neuronal differentiation conditions resulted in cell death that was observed in the form of single cells or clumps of dead cells. Apoptotic cell death could be due to the lack of appropriate differentiation or survival cues during the prolonged culture *in vitro*. Both differentiation conditions contained B27 supplement to promote survival of cells. Although B27 supplement has been shown to promote survival of primary hippocampal neurons³², it may not be sufficient for promoting survival of iMOP cells. Addition of compounds such as the ROCK inhibitor may help maintain and increase survival of iMOP cells during routine culture and *in vitro* differentiation. ROCK inhibitor has been extensively used in pluripotent stem cell cultures to promote survival of cells in serum free cultures without affecting differentiation³³. Addition of ROCK inhibitor to the culture media can help increase cell survival during long differentiation protocols without affecting the differentiation potential of iMOP cells. Increased cell survival will allow development of protocols to efficiently generate large numbers of mature hair cells or SGNs.

Increased Cdkn1b Expression as an Early Marker of IMOP Differentiation

During early cochlear development, the initial event of cell cycle exit precedes differentiation for hair cells, supporting cells and SGNs³⁴. Terminal mitosis of neuronal progenitors spreads in a wave starting from the base and progressing towards the apex of the cochlea³⁵. In an opposing gradient, terminal mitosis of prosensory progenitors that give rise to hair cells and supporting cells progresses from the apex to base of the cochlea^{34,36}. Although the temporal expression of Cdkn1b correlates well to the post-mitotic state, it is most likely not the sole factor that promotes cell cycle exit in the developing cochlea. In support of this, Cdkn1b mutant animals can still develop post-mitotic hair cells³ . Instead, Cdkn1b can be used as an early indicator to mark the onset of differentiation. In unguided iMOP differentiation cultures, an early marker for differentiation will assist in developing protocols to promote early stages of differentiation before obvious morphological features can be observed.

Similar to otic development, cycle exit initiates differentiation and results in increased Cdkn1b levels in iMOP cells. In proliferating iMOP cells, only low level expression of Cdkn1b can be observed. Other cyclin dependent kinase inhibitors such as Cdkn1a (p21^{CIP}) may be responsible for normal cell cycle progression in iMOP cells. During sensory epithelia and neuronal differentiation of iMOP cells, single cell quantitative analysis revealed a subset of cells with increase expression of Cdkn1b in differentiating iMOP cultures (**Figure 6A, C**). Expression of Cdkn1b in these cells are an indication of iMOP cells undergoing terminal differentiation.

IMOP Cells as a Cellular Platform for Studying Cell Fate Determination

Since iMOP cells can transition from a progenitor state to different otic lineages *in vitro*, they can be utilized to study otic cell fate determination. Currently, the described protocol used unguided differentiation procedures to generate different otic cell types. The iMOP system allows for addition of bioactive molecules, compounds and defined genes that guide iMOP cells towards the mature hair cell, supporting cell and SGN lineages. One use of iMOP cellular platform is to test for candidate transcriptions factors (TF) that promote differentiation. Expression of key TF can be used to drive hair cell or neuronal differentiation. Expression of Cdkn1b along with established markers or morphological features of hair cells and SGNs can be used to determine the contribution of candidate TFs to distinct otic lineages^{38,39} .

Candidate TFs such as Atoh1 are essential for hair cell development and have been repurposed to promote hair cell differentiation in the damaged cochlea^{40,41}. Introduction of Atoh1 into iMOP cells may help promote hair cell differentiation. In the developing cochlea, both Ngn1 and
NeuroD1 are required for proper SGN differentiation *in vivo⁴²⁻⁴⁴.* Expre These experiments are similar to generating induced neuronal cells (iN), where expression of the TFs individually or in combination can convert fibroblasts or pluripotent stem cells into neurons45-48. Introduction of TFs that are normally expressed during development of hair cells or SGNs is a good strategy for guiding iMOP cells towards the hair cell or SGN lineage.

Conditions that promote differentiation of a large number of iMOP-derived hair cells and SGNs will provide a robust cellular platform for disease modelling, small molecule screening and toxicology testing that can be accomplished quickly and cost-effectively before labor-intensive tests in rodents are initiated. The protocol described presents a simple and versatile system that can be utilized for studying otic progenitor differentiation and is amenable to large scale experiments.

Disclosures

The authors have nothing to disclose.

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