SOP: Propagation of Human Non-pigment Ciliary Epithelial Cells (ScienCell

**Research Laboratories**)

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Modified by: T. Canfield (UW)

# **Ordering Information**

Human Non-pigment Ciliary Epithelial Cells (HNPCEpiC) may be ordered as frozen ampoules. Each vial contains  $>5 \times 10^5$  cells in a 1ml volume.

To order frozen ampoules + media:

Name: HNPCEpiC – Human Non-pigment Ciliary Epithelial Cells

Item #: 6580 (HNPCEpiC - Cryopreserved ampoule)

4101 (EpiCM, Epithelial Cell Medium)

## Notes:

The number of media kits purchased depends on the target number of cells to be generated. A rule of thumb is 10 media kits for every initial cryopreserved ampoule. It is strongly recommended to purchase all of the media that will be required for a complete expansion series, since media supply may be erratic. All culture flasks must be coated with poly-L-lysine (2 ug/cm<sup>2</sup>).

## **Materials List**

- 1. Cell-type specific medium (ScienCell)
- 2. Poly-L-Lysine (#0413, 10mg/ml, ScienCell)
- 3. T225 culture flasks
- 4. Graduated pipets (1, 5, 25, 50mL)
- 5. Pen-Strep solution (if required; ScienCell typically supplies antibiotics)
- 6. Accutase Enzyme Cell Detachment Medium (Cat # 00-4555, EBioscience)
- 7. Hemocytometer
- 8. Micropipet w/ P20 tips
- 9. Microscope

## **Procedure**

## A. Initiation of culture from cryopreserved cells

- 1) Rapidly thaw cells by holding vial and gently rotating in 37°C water bath.
- 2) As soon as ice crystals disappear, dispense contents of vial into a flask at 5,000 cells/cm<sup>2</sup> density.
- 3) Let cells recover for 16 hours in a 37°C, 5% CO<sub>2</sub> humidified incubator.

## **B.** Sub-culture

- 1) Propagate cells until density reaches 90% confluence.
- 2) Aspirate medium.

- 3) Wash cells with warm 1X PBS.
- 4) Add 15mLs of Accutase and return to incubator for 10-15 minutes.
- 5) Immediately remove cells, rinse flask with warm 1X PBS to collect residual cells, and pellet at 500 x g for 5 minutes (4°C).
- 6) Gently re-suspend cell pellet in warm medium.
- 7) Count cells with hemocytometer.
- 8) Add warmed medium to flasks.
- 9) Seed flasks at 5,000 cells/cm<sup>2</sup>. Record each subculture event as a passage.

#### C. Maintenance

- 1) Change media the day after seeding and every OTHER day thereafter.
- Increase media volume as confluency increases (volumes assume the use of T225 flasks):
  - a. 25 % = 1 mL/5 cm 2
  - b.  $25-45\% = 1.5 \text{mL}/5 \text{ cm}^2$
  - c. 45%+ = 2mL/5 cm2.
- 3) Per the above an exemplary schedule might be:
  - a. day 1, plate into T225: use 50 mls of media.
  - b. day 2, change media, use 50 mls of media
  - c. day 4, change media, use 100 mls of media (if confluency is >50%)
  - d. day 6, change media, use 100 mls of media (or harvest if ready)
  - e. day 7 or 8 (harvest when cells reach 6 x 10<sup>6</sup> cells/flask).

## D. Harvest

- 1) Pass cells 3-4 times until the desired cell number is achieved (primary cells will senesce after 4-5 passages).
- 2) Remove cells from flasks according to protocol described above under 'Sub-culture'.
- 3) Examine viability using Trypan blue staining (SOP TP-7).