Ordering Information

Human Brain Neuroepithelioma Cells SK-N-MC can be ordered from ATCC as a frozen ampoule with >2 x 10^6 per 1mL volume. This is an adherent cell line.

Name: SK-N-MC—Human Brain Neuroepithelioma Cells
ATCC #: HTB-10

Materials List

1. MEM with 2mM L-glutamine and Earle’s salts Medium (Cellgro, Cat# 10-010-CM)
2. Characterized Fetal Bovine Serum (HyClone, Cat# SH30071)
3. Sodium Pyruvate, 100mM (Cellgro, Cat# 25-000-CI)
4. Non-essential Amino Acids, 100X solution (Invitrogen, Cat# 11140-050)
5. Penicillin-Streptomycin Solution (200X) (Cellgro, Cat# 30-001-CI)
6. T25, T75, T225 tissue culture flasks
7. Corning conical centrifuge tubes (15mL and 50mL)
8. Graduated pipets (1, 5, 10, 25, 50mL)
9. Phosphate Buffered Saline (1X PBS) (Cellgro, Cat# 21-040-CM)
10. Accutase Enzyme Cell Detachment Medium (EBiosciences, Cat# 00-4555)
11. Freezing Medium ( Growth medium containing 5% DMSO)
12. DMSO, Hybri-Max (Sigma-Aldrich, Cat# D2650)
13. Cryovials (Nunc, Cat# 368632)
14. Cryo 1°C Freezing Container (Nalgene Cat# 5100-0001)
15. Eppendorf Centrifuge 5810R
16. Revco UltimaII -80°C Freezer
17. Thermolyne Locator 4 Liquid Nitrogen Freezer
18. Hemocytometer
19. Micropipet w/ P20 tips
20. Microscope

Growth Medium for SK-N-MC

MEM with 2mM L-glutamine and Earle’s salts Media
10% Characterized FBS
Sodium Pyruvate (1mM)
Non-essential Amino Acids (1X)
Pen-Strep (1X)

Procedure

A. Receipt of Frozen Cells and Starting Cell Culture
   1) Immediately place frozen cells in liquid nitrogen storage until ready to culture.
   2) When ready to start cell culture, quickly thaw ampoule in a 37°C water bath.
   3) As soon as ice crystals disappear, swab outside surface of the ampoule with 70% ethanol, then dispense contents of ampoule into a T25 flask with 10mL of warm growth media.
4) Allow cells to recover overnight in 37°C, 5% CO₂ humidified incubator.
5) The next morning, the diluted DMSO-containing shipping/cryopreservation medium is aspirated from the cell layer and replaced with fresh medium.

B. Sub-culture
1) Propagate cells until density reaches 70-80% confluence.
2) Aspirate medium.
3) Wash cells with warm 1X PBS.
4) Add 15mL of Accutase and return to incubator for 10-15 minutes, or until cells detach.
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) Perform 1:6 to 1:12 cell split as needed.
8) Record each subculture event as a passage.

C. Maintenance and Generation of Seed Stocks
1) Change media the day after seeding and every 2-3 days thereafter. Use 50mL of growth medium per T225 flask.
2) Following first or second passage after receipt of cells and with sufficient number of cells to continue maintenance and expansion, the major portion of the flasks should be sub-cultured using Accutase as above under “Sub-culture” and a small portion should be set aside as a seed stock. The cell pellet for the seed stock should be resuspended in freezing medium.
3) Cells in freezing medium are dispensed into cryovials (2 million cells per 1 mL aliquot) and frozen at -80°C in a Nalgene Cryo 1°C freezing container overnight.
4) Cryovials are transferred the next day to liquid nitrogen freezer for long-term storage.

D. Harvest
1) Passage cells until the desired number of cells is reached.
2) Remove cells from flasks according to protocol described above under “Sub-culture”.
3) Examine viability using Trypan blue staining (SOP TP-7).