

Primary Cells for Scientific Research

Rat Brain Microvascular Endothelial Cells (Primary cell immortalization)

Cat. No. : CP-R108Y

General Information

Species	Rat/SD rats
Tissue Type	Nervous system
Tissue	Cerebral cortical tissue
Cell Type	Endothelial cells
Morphology	Endothelial-like
Growth Properties	Adherent
Storage Conditions	For long-term cryopreservation, cryovials should be stored in liquid nitrogen at -150°C to -196°C. Storage at -80°C is restricted to short-term interim use only.

Culture Conditions and Handling

Complete Medium	Rat Brain Microvascular Endothelial (Primary cell immortalization) Cell Complete Medium[CM-R108Y]
Population Doublings	Guaranteed to further expand for 250 population doublings
Subcultivation Ratio	First passage at 1:2 ratio, subsequent passages at 1:2 to 1:6 ratio
Dissociation Reagent	0.25% trypsin
Coating Conditions	PLL (0.1 mg/mL) or gelatin coating solution (0.1%)
Medium Renewal	Every 2 to 3 days
Freezing Medium	General Freezing Medium[PB180436]
Incubation Atmosphere	Air, 95%; CO ₂ , 5%
Temperature	37°C
Instructions	1. Check all containers for leakage or breakage. 2. Remove the frozen cells from the dry ice packaging and immediately transfer them to liquid nitrogen (liquid or vapor phase) for long-term cryopreservation. 3. For the first subculture, a 1:2 split ratio is recommended.
Subculturing Procedure	Adherent cell dissociation

Subculturing Procedure

1. Aspirate the culture medium from the T25 flask and wash the cells once with phosphate-buffered saline (PBS).
2. Add 1 mL of 0.25% trypsin solution to the T25 flask. Gently swirl the flask to ensure the solution fully covers the bottom surface, then aspirate 500–700 µL of the trypsin solution. Incubate at 37°C for 1–3 min. Examine the cells under an inverted microscope. Once cells become rounded and retracted, add 5 mL of complete growth medium to neutralize the digestion.
3. Gently pipette to mix and dissociate the cells. Subculture into a new T25 flask at the recommended split ratio, and top up with fresh complete growth medium to a final volume of 5 mL. Incubate the flask in a humidified incubator at 37°C with 5% CO₂.
4. Once the cells have fully adhered, assess cell viability and confluency under a microscope prior to downstream experiments. Subsequently, refresh the medium with fresh complete growth medium per the standard cell maintenance protocol.

Background

Rat brain microvascular endothelial cells (primary cell immortalized) (*Rattus norvegicus*) are derived from primary rat brain microvascular endothelial cells that have been transduced with the SV40T gene via lentiviral transduction, followed by selection using antibiotics and passaging to obtain the immortalized cell line. Rat brain microvascular endothelial cells are isolated from brain tissue. They are monolayer squamous endothelial cells lining the luminal surface of brain microvessels, with well-developed tight junctions. They are the main component of the blood-brain barrier and express tight junction protein ZO-1. Their core functions include: forming the blood-brain barrier to selectively control the passage of substances into and out of brain tissue; regulating cerebral blood flow; and maintaining the homeostasis of the central nervous system microenvironment. Barrier dysfunction of rat brain microvascular endothelial cells is closely associated with stroke and Alzheimer's disease. In vitro-cultured rat brain microvascular endothelial cells serve as a classic cell model for investigating blood-brain barrier function.

Rat brain microvascular endothelial cells (primary cell immortalization) from Procell Laboratory were generated by lentiviral delivery of the SV40T gene into primary rat brain microvascular endothelial cells, followed by antibiotic selection and serial subculture screening. Each vial contains approximately 5×10^5 cells.

Rat brain microvascular endothelial cells (primary cell immortalization) isolated from Procell Laboratory exhibit positive immunofluorescence staining for CD31, with a purity greater than 90%. In addition, the cells test negative for HIV-1, HBV, HCV, mycoplasma, bacteria, yeast, and fungi.

Handling Recommendations for Cryopreserved Cells

Preparations Before Thawing

1. Primary cells have high nutritional demands; therefore, prepare complete culture medium in advance.
2. Preheat the complete culture medium to 37°C for 30 minutes.
3. Prepare 9 mL of complete medium within a sterile centrifuge tube to dilute the cryoprotectant present in the frozen cell suspension.
4. Use sterile gloves to protect the cryovial from contact with the water bath water, and preventing cell contamination.

Thawing Procedure

1. Retrieve the cryovial containing frozen cells from liquid nitrogen storage. Place it in sterile gloves and immediately submerge it in a 37°C water bath.
2. Thaw the cells rapidly (<1 minute) by gently swirling the vial in the 37°C water bath until only a small ice crystal remains.
3. Transfer the vial to a laminar flow hood. Wipe the exterior of the vial with 75% ethanol before opening.
4. Transfer the thawed cells dropwise into a preheated 9 mL centrifuge tube containing complete medium.
5. Centrifuge the cell suspension at approximately 250 × g for 5 minutes. For cells characterized by low density and small volume, such as lymphocytes and suspended cells, the centrifugation speed can be appropriately increased to 400 × g for 8 minutes.
6. After centrifugation, discard the supernatant and retain the cell pellet.
7. Gently resuspend the cells in complete growth medium, then transfer them to an appropriate culture vessel and place it in the recommended culture environment.

Notes

1. The entire recovery process should be completed as quickly as possible.
2. Select the culture vessel size based on the number of cells in the cryovial. Procell single cryovials are recommended for resuspension in 6 cm dishes or T25 flasks.
3. Minimize the time thawed cells are kept at room temperature. DMSO must be immediately diluted or removed by centrifugation.

Precautions After Receipt of Frozen Cells

1. Upon unpacking, inspect the condition of the frozen cells and dry ice, and take photographs immediately. The following after-sales service will be provided based on these photographs, including assessment of the remaining dry ice, verification that the cryovial was fully buried in dry ice, and evaluation of whether the cells thawed and refrozen during transit.
2. Upon receipt, the cells should be transferred to liquid nitrogen immediately or directly resuscitated. If liquid nitrogen is unavailable, the cells can be temporarily stored at -80°C, however, the storage period should be limited to less than one week whenever possible. Prolonged storage at -80°C may gradually reduce post-thaw cell viability, and the extent of this viability loss is unpredictable.
3. Ensure that the operator has sufficient knowledge and experience in cell culture, and the laboratory is equipped with essential instruments, including a biosafety cabinet, CO₂ incubator, inverted microscope, centrifuge, water bath. Carefully review the cell instruction manual to understand key cell characteristics including growth properties (adherent or suspension), morphology, basal medium requirements, serum concentration, cytokine supplementation, subcultivation ratio, medium renewal schedule.
4. After resuscitation, observe the cells under a microscope and record the cell status by taking photographs (1-3 images each at 100× and 200× magnification for 3 consecutive days). These images will serve as supporting documentation for follow-up services. In addition, a small aliquot of cells may be used to assess cell viability by automated cell counting or trypan blue staining.

Note: Cells should not be observed too frequently within the first 24 hours after resuscitation, as this may affect cell growth or adherence. Observation once per day is sufficient.

5. Primary cells have a limited number of passages; it is recommended that they be used as soon as possible.