

The User's Manual—Vero SFM01 Pro

Basic Information Introduction

Product Introduction

Vero SFM01 Pro is a Serum-Free basal medium, without protein and any animal-derived components, developed specifically for Vero cell lines, it is suitable for Vero cell adherent static culture and microcarrier suspension large-scale culture, supporting high-density cell growth and viability maintenance, it can be used in the development and production process of influenza vaccine and other related products.



Application Scope

Vero SFM01 Pro can be used in the process of Vero cell thawing, cell passage, adherent static culture, and microcarrier suspension large-scale culture. The basic medium is suitable for scientific research and the production of large-scale biological products based on cell culture, but cannot be directly used in the human body or used as a medication.

Shipping, Storage and Validity Period

| Product | Catalog No. | Storage | Shipping | Validity Period |
|----------------|--------------|----------------------------------|----------------------------------|-----------------|
| Vero SFM01 Pro | LQ22, Liquid | 2°C ~ 8°C, Protect from light | 2°C ~ 8°C, Protect from light | 12 months |
| Vero SFM01 Pro | DP22, Powder | 2°C ~ 8°C, Protect from light | 2°C ~ 8°C, Protect from light | 24 months |

Protocol for Hydration of Powder Medium

1. Fill the mixing container with purified water (25 ~ 30°C) at 90% of the final volume.
2. Slowly add 16.85 g/L of powder medium with gentle stirring. Mix for more than 10 minutes.
3. Adjust the pH to 6.6 ~ 6.8 using 1M HCl or 1M NaOH solution. Mix continuously for 20 minutes until completely dissolved.
4. Slowly add 2000 mg/L NaHCO₃ to the solution with gentle stirring. Mix for 10 minutes.
5. Adjust the solution volume to 100% with purified water. Mix for 10 minutes.
6. Filter immediately the media with a 0.22 μm membrane filter.

Quality Index of Powder and Liquid Media

| Product Index | Vero SFM01 Pro (LQ22), Liquid | Vero SFM01 Pro (DP22), Powder |
|------------------------|-------------------------------|--|
| Appearance | Red, clear liquid | Off- pink powder |
| pH | 7.1 ~ 7.5 | 7.1 ~ 7.3 (pre-filter) |
| Osmolality (mOsmol/kg) | 270 ~ 320 | 270 ~ 320 |
| Solubility | -- | Dissolve well according to the protocol for hydration of powder medium |
| Endotoxin (EU/mL) | < 3 | < 3 |
| Sterility | Negative | -- |
| Bioburden | -- | Aerobic bacteria: < 200 CFU/g Molds and yeasts: < 50 CFU/g |

Reference Cell Culture Protocol

Culture Conditions

| Parameter | | Value |
|---------------------|--|---|
| Culture volume | T25 cell culture flask | 5 ~ 10 mL |
| | T75 cell culture flask | 15 ~ 25 mL |
| | T175 cell culture flask | 40 ~ 55 mL |
| | 1-layer Cell Factory | 130 ~ 200 mL |
| | 2-layer Cell Factory | 260 ~ 400 mL |
| | 10-layer Cell Factory | 1300 ~ 2000 mL |
| Culture environment | Seeding density | 2 ~ 4×10 ⁴ cells/cm ² |
| | Incubation temperature | 37°C |
| | Incubation CO ₂ concentration | 5% |
| | Incubation relative humidity | > 80% RH |

Cell Thawing

1. Pre-warm the medium (Vero SFM01 Pro) in 37°C water bath.
2. Spray the outside of the medium bottle with 75% alcohol and place the bottle into the bio-safety cabinet.

3. Thaw one vial at a time in 37°C water bath. Gently agitate the vial within 1 minute until the ice in the vial melting.
4. Pipet the contents from the vial gently into a centrifuge tube containing 10 mL of pre-warmed medium (Vero SFM01 Pro).
5. Centrifuge 150 g to 300 g for 5 minutes. Discard the supernatant and re-suspend cells in 10 ~ 30 mL fresh pre-warmed medium (Vero SFM01 Pro), then adjust the cell density to $2 \sim 4 \times 10^4$ cells/cm².
6. Sample 0.5 mL of cell suspension and analyze the viable cell density ($\times 10^6$ cells/mL) and viability (%) of the sample using cell counter.
7. If the cell viability > 90%, incubate cells in the specified condition (refer to "culture conditions" table).

Cell Passage

Take the T75 flask as an example:

1. Pre-warm the medium (Vero SFM01 Pro) in 37°C water bath for 20 ~ 30 min.
2. Cells with cell viability $\geq 90\%$, and in the middle of logarithmic growth phase were selected for passage.
3. Aspirate medium from cell monolayer and rinse flask with 7 ~ 10 mL pre-warmed DPBS. Aspirate DPBS.
4. Add 1 mL of Recombinant Trypsin Solution (PDE RecTrypsin-LQ02 (1 \times), PDE-LQ02/ PDE RecTrypsin-LQ03 (10 \times), PDE-LQ03) to flask, infiltrate all culture surfaces, and digest for 3 ~ 5 minutes at room temperature.
5. When the cells fall off from the bottle wall, pat the bottle wall, add the preheated mixture (containing 1 mL of 0.5% trypsin inhibitor and 9 mL of Vero SFM01 Pro medium, a total of 10mL) to pipette and disperse the cells, collect the cells, and centrifuge the cell suspension at room temperature 100 g for 5 ~ 10 minutes.
6. Discard the supernatant and resuspend the cell pellet with Vero SFM01 Pro medium.
7. According to the seed density of $2 \sim 4 \times 10^4$ cells/cm², calculate the amount of total number of seed cells.
8. Seed cells at $2 \sim 4 \times 10^4$ cells/cm² in the T75 flask and add a certain volume of pre-warmed fresh medium.
9. Passage cells with fresh medium according to the above steps every 3 to 5 days.

Adaptation

1. For cells can direct adapt, transfer cells suspension cultures into Vero SFM01 Pro directly, and the seed cell density refer to the cell passage procedure.
2. Cell passage until cell expression steadily.
3. The recommended cell passage seeding density was $2 \sim 4 \times 10^4$ cells/cm², and the cells were subcultured every 4 to 6 days. Generally, three passages of cells could be completely adapted.

Cell Cryopreservation

1. Prepare cells, harvesting in mid-log phase of growth with viability > 90%.
2. Sample and cell counting, calculate the required volume of cell freeze solution to give a final density of 1×10^7 cells/mL.
3. Prepare the cell freeze solution: 90% Vero SFM01 Pro + 10% DMSO, refrigerate at 4°C.
4. Prepare the cell suspension according to the cell digestion procedure.
5. Centrifuge 300 g for 5 minutes, discard the supernatant and re-suspend cells with the cell freeze solution.
6. Immediately dispense aliquots of cells suspension into cryovials according to the specific needs of the project.
7. Achieve cryopreservation in an automated or manual controlled rate freezing apparatus following standard procedures (1°C decrease per minute).
8. Transfer to liquid nitrogen tank for storage.