

SF1 hMSC Medium

Product Information Sheet

Introduction

SF1 hMSC Medium is a serum free and animal-component free cell culture medium. This medium is designed for the growth and expansion of human mesenchymal stem cells (MSCs) derived from bone marrow, adipose or Wharton's Jelly. Using SF1 hMSC Medium, human MSCs can be expanded quickly for multiple passages while maintaining their ability to differentiate into osteogenic, chondrogenic and adipogenic lineages.

Plate-coating is not needed for the SF1 hMSC Medium, which means that the user can use tissue culture vessels directly and save labor and cost related to coating plate. Information of recommended tissue culture vessels is provided in this sheet.

Cat. No.: MSC-SF-003 includes:

Item	Product	Cat. No	Amount	Storage
1	hMSC SF1 Basal Medium	MSC-SF-004	450 mL	2 to 8°C
2	hMSC SF1 Supplement	MSC-SF-005	50 mL	-20°C

Note: SF1 hMSC Medium is sold as a complete kit; its components are not available separately.

Intended Use

SF1 MSC Medium is intended for human ex vivo tissue and cell culture processing applications.

CAUTION: Not intended for direct administration to humans or animals.

Precaution:

Please read the prepared complete medium procedure before use.

- 1. Thaw the frozen hMSC SF1 Supplement at 4°C refrigerator about 12-20 hrs prior use, do not over 24 hrs.
- 2. Swirl the bottle on the desk gently to mix the hMSC SF1 Supplement solution.
- 3. Warm the hMSC SF1 Basal Medium (Cat. No MSC-SF-004) in a 37°C water bath for 25-30 min.
- 4. Warm the hMSC SF1 Supplement (Cat. No MSC-SF-005) in a 37℃ water bath for 10-15 min. (The warming times please do not longer than 20 min.)
- 5. Swirl warmed hMSC SF1 Supplement and hMSC SF1 Basal Medium on the desk gently.
- 6. Aseptically transferred the entire contains of hMSC SF1 Supplement bottle to hMSC SF1 Basal Medium bottle.
- 7. Swirled the hMSC SF1 Basal Medium bottle gently to ensure complete mixing. **This is the complete medium**.



- 8. Rinse the hMSC SF1 Supplement bottle with complete medium for 3 times, about 25-50 mL per time.
- 9. Swirl the hMSC SF1 complete medium bottle gently to ensure complete mixing.
- 10. Please aliquot the complete medium if you do not once use the medium.
- 11. Please use the aliquot medium within two months.

Cell Culture Notes and Procedure

Tissue culture vessels

- 1. We recommend use Corning CellBind® product line of culture vessels. Corning/COSTAR general type tissue culture product, BD Primaria® product line and Falcon plates (6, 12 well-plate etc.) for MSCs cultured in SF1 hMSC Medium.
- 2. We highly recommend use Corning CellBind® products for initial plating of primary isolated Mesenchymal Stem Cells from bone marrow, adipose or Wharton's Jelly (P0). Corning CellBind® products also are the best choice for the low density seeding related experiment (for example, Colony Forming Unit Assay, seeding density below 5x10²/cm²).

Culture procedure

The following protocols have been used successfully in our laboratories.

- It's highly recommend to <u>warm the SF1 hMSC Medium in 37°C</u> water bath for about 15 minutes and gently mix it before the experiment.
- 2. Remove media from T-25cm² flask. Gently wash flask with 5 mL Dulbecco's PBS (without Ca²⁺ or Mg²⁺) for about 1 minute and removes the solution.
- 3. Add 1 mL pre-warmed Trypsin reagent to flask and dispense evenly over the entire culture surface.
 <u>We recommend products from Life technologies, TrypLE Select cat. no. 12563</u> or 0.05%
 Trypsin-EDTA (0.53mM). But 0.25% Trypsin-EDTA (1mM) is not recommend for the MSCs subculture.
- 4. Incubate the flask at 37°C until cells become rounded (about 2-5 minutes) and are easily dislodged from the surface. Firmly tap the flask as necessary to facilitate cell detachment.
- 5. Once detached, add 6 mL SF1 hMSC medium to flask, mix gently and transfer cell suspension to a 15 mL centrifuge tube. (Note: Don't use Dulbecco's PBS as suspension medium)
- 6. Centrifuge for 5 minutes at 290 x g (~1200 rpm for most table-top centrifuges).
- 7. Remove supernatant and re-suspend cell pellet in 1 mL SF1 hMSC Medium.
- 8. Determine viable and total cell number. Determine cell viability by trypan blue-dye exclusion method, and use a hemocytometer or other counting system (for example, ADAM system or Vi-CELL system) to determine cell number.
- 9. Seed cell to flasks at 0.5-1x10³ viable cells/cm². The seeding density recommended for MSCs derived from adipose is 0.5x10³, for MSCs derived from bone marrow or Wharton's Jelly is 1x10³/cm².
- 10. Incubate cells at 37°C in a humidified chamber containing 5% CO₂.
- 11. Re-fresh medium with pre-warmed SF1 hMSC Medium after 3 days of culture.
- 12. Subculture until cells are 80-90% confluent, which takes about 4-6 days after seeding.



Cryopreservation of cells with SF1 hMSC Medium

- Prepare cryopreservation solution by supplementing SF1 hMSC Medium with 10% Dimethyl Sulfoxide (DMSO).
- 2. Pellet detached cells by centrifugation (290xg, 5minutes), gently resuspend cells in cryopreservation solution to 0.3-1x10⁶ cells/mL, and transfer to cryovials.
- 3. Place cryovials in a freezing container (e.g. Nalgene 5100-0001) and place in a -70 to -80°C freezer overnight.
- 4. Transfer cryovials to liquid nitrogen for long-term storage.

Recovery of Cryopreserved Human MSCs

- 1. Rapidly thaw frozen vial of cells in a 37°C water bath.
- 2. Add 8-9 mL pre-warmed (37°C) SF1 hMSC Medium in one 15 mL conical tube.
- 3. Pipet the entire contents of the cryovial into the conical tube. Tighten the cap and inverse the tube gently to mix the cells and medium.
- 4. Follow the "Culture procedure" (upper section) step 6 to 12 to seed the cells.

Quality Control Testing of the Lot

The performance of SF1 hMSC Medium was verified by cell expansion assay using MSCs derived from adipose. Sterility test and LAL test compliant with USP were performed to verify the absence of bacterial and fungal contamination and in SF1 hMSC Medium.

Manufacture

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