



Rat Mesenchymal Stem Cell Replenisher Kit

Cat # 5000-001-R

Qualified medium and serum to support undifferentiated growth of Rat Mesenchymal Stem Cells

Rat Mesenchymal Stem Cell Replenisher Kit

Cat# 5000-001-R

Qualified medium and serum to support undifferentiated growth of Rat Mesenchymal Stem Cells

Table of Contents

		Page
I.	Introduction	1
II.	Safety Information	1
III.	Description of Equipment	1
IV.	Materials Required But Not Supplied	1
٧.	Reagent Preparation	2
VI.	Protocol	2
VII.	References	5
VIII.	Troubleshooting	5
IX.	Related Products Available From Trevigen	6

©2009, Trevigen Inc. All rights reserved. Trevigen, Cultrex, CultreCoat and PathClear are registered trademarks of Trevigen, Inc. Corning is a registered trademark of Corning, Inc.

I. Introduction

Mesenchymal Stem Cells (MSC), also known as marrow stromal cells, are a self-renewing population of multipotent cells present in bone marrow and many other adult tissues. (1,2) MSC can be isolated from bone marrow by adherence to plastic, (1,4) and can differentiate into multiple lineage-specific cells that form bone, fat, cartilage, muscle, neuronal cells and tendon. (1,4) Due to their multilineage potential, they can be useful tools for a wide range of therapeutic and basic research, including transplantation studies and studies examining the repair of cardiac tissue, bone, cartilage, and tendons often using 3-D matrices. (1,4)

Trevigen's Qualified RMSC Medium (cat# 5000-500-03) supplemented with Qualified RMSC FBS (cat# 5000-050-02) has been evaluated to support undifferentiated growth of Trevigen's RMSC (cat#5000-001-01) or induction of adipogenic (cat# 5010-024-K) or osteogenic (cat# 5011-024-K) phenotype when supplemented with reagents contained in the Trevigen's differentiation kits.

II. Precautions and Limitations

- 1. For Research Use Only. Not for use in diagnostic procedures.
- The physical, chemical, and toxicological properties of these products may not yet have been fully investigated; therefore, Trevigen recommends the use of gloves, lab coats, and eye protection while using these chemical reagents. Trevigen assumes no liability for damage resulting from handling or contact with these products.

III. Materials Supplied

Component	Quantity	Storage	Catalog#
Qualified RMSC Medium	500 ml	4°C	5000-500-03
Qualified RMSC FBS	50 ml	-20°C ¹	5000-050-02

¹ After initial thaw, FBS should be aliquoted to avoid repeated freeze/thaws. FBS should be thawed at 4°C overnight prior to first use.

IV. Materials/Equipment Required But Not Supplied

Equipment

- 1. 1 20 μl, 20 200 μl, and 200 1000 μl pipettors
- 2. Laminar flow hood or clean room
- 3. 37°C CO₂ incubator
- 4. 37°C Water Bath
- Hemocytometer or other means to count cells
- 6. Inverted standard or phase microscope
- 7. Pipette aid
- 8. Liquid Nitrogen Storage
- 9. Low speed swinging bucket centrifuge and tubes for cell harvesting
- 10. Cell freezing container

Reagents

- 1. Rat Mesenchymal Stem Cells (Trevigen cat# 5000-001-01 or equivalent)
- 2. Cell Harvesting Reagent, trypsin, dispase, etc.
- 3. Antibiotic Supplement for Media (optional)
- 4. Sterile PBS (Mg²⁺, Ca²⁺ free) or HBSS
- 5. Trypan blue or equivalent viability stain
- 6. DMSO
- 7. 70% Ethanol

Disposables

- 1. Cell culture flask, 25 cm², 75 cm², or 185 cm²
- 2. 15 ml conical tubes
- 3. 0.22 µm Filter Unit (optional)
- 4. 1 200 μl and 200 1000 μl pipette tips
- 5. 1, 5 and 10 ml serological pipettes
- 6. gloves
- 7. Cryovials

V. Reagent Preparation

These procedures should be performed in a biological hood utilizing aseptic technique to prevent contamination.

1. Mesenchymal Complete Growth Medium

For 250 ml of Medium:

Qualified RMSC Medium (cat# 5000-500-03): 225 ml Qualified RMSC FBS (cat# 5000-050-02): 25 ml

Optional: media can be filter sterilized before use

Store media at 4°C for one month

Ensure media are at room temperature or 37°C prior to use

2. 2X Mesenchymal Freeze Medium

For 10 ml of Medium:

Qualified RMSC Medium: 4 ml Qualified RMSC FBS: 4 ml DMSO: 2 ml

Mix 1:1 with Growth Medium before use

VI: Protocol

These procedures should be performed in a biological hood utilizing aseptic technique to prevent contamination.

A. Thawing Mesenchymal Stem Cells:

1. Prepare Mesenchymal Complete Growth Medium (Section V.1).

- 2. Prewarm Complete Growth Medium to 37 °C by placing in 37°C H₂O bath or in Tissue Culture Incubator.
- 3. Immediately before use, remove vial of cryopreserved rat mesenchymal stem cells (RMSC) from liquid nitrogen freezer.
- 4. Thaw frozen RMSC quickly in a 37°C H₂O bath.
 - a. Ensure cells are completely thawed before proceeding.
 - b. Do not leave cells at 37°C past thawing.
- 5. Spray down bottle of Complete Growth Medium and ampoule (containing thawed cells) with 70% EtOH before placing in Tissue Culture Hood.
- Aseptically, transfer the thawed cells to a 15 ml conical tube with a 5 ml pipette.
- 7. Wash ampoule with 1 ml of warm medium using a 5 ml pipette.
- 8. Add drop-wise to 15 ml conical tube containing cells, gently swirling to mix between drops.
- 9. Add 1 ml of warm medium to 15 ml conical tube containing cells, gently swirling to mix between drops. Total Volume should be about 3 ml.
- 10. Centrifuge 15 ml conical tube at 200 x g for 3 minutes.
- 11. Remove supernatant gently to avoid disturbing cell pellet.
- 12. Resuspend cell pellet in 1 ml of fresh Mesenchymal Complete Growth medium.
- 13. Count cells on hemocytometer (per standard protocol).
- 14. Plate cells at a density of 5.4 x 10³ cells/cm². For a T-75 tissue flask add 4.05 x 10⁵ cells in 12-15 ml complete growth medium.
 - a. Recommend Corning[®] Tissue Culture Treated Plastic.
 - b. One vial is sufficient to seed two T-75 or one T-185 flasks.
- 15. Place Tissue Culture Flask/Dish in 5% CO₂ Tissue Culture Incubator at 37°C.
- 16. Change medium in flasks next day.

B. Growing Mesenchymal Stem Cells:

- 1. Medium Change (every 3-4 days is recommended).
 - a. Warm Complete Growth Medium to 37 °C by placing in 37°C H₂O bath or in Tissue Culture Incubator.
 - b. Spray down bottle of Complete Growth Medium with 70% EtOH before placing in Tissue Culture Hood.
 - c. Remove medium from T-75 flask containing Mesenchymal Stem Cells.
 - d. Add 12-15 ml of fresh Complete Growth Medium.
 - e. Discard used medium appropriately.

2. Passaging Rat Mesenchymal Stem Cells

Note: When cells became 70-80% confluent, they are ready to be split. If allowed to over-grow, these cells will lay down a matrix and start to differentiate; as a result, the cells will peel off of the plastic which markedly reduces the ability to passage them.

- a. Warm Complete Growth Medium and Trypsin solution to 37 °C by placing in 37 °C H₂O bath or Tissue Culture Incubator.
- b. Spray down bottles containing Complete Growth Medium, or trypsin with 70% EtOH before placing in Tissue Culture Hood.
- c. Remove medium from T-75 flask containing Mesenchymal Stem Cells.

- d. Gently wash flask with 5-10 ml of sterile 1X PBS (Ca²⁺ and Mg²⁺ free)
- e. Remove PBS.
- f. Add 3 ml of Trypsin solution to each flask; place at 37°C for 3-5 minutes (until cells are no longer attached to plate, should take less than 5 minutes) in Tissue Culture Incubator.
- g. Add 5 ml of Complete Growth Medium to flask.
- h. Transfer cells to 15 ml conical tube.
- Centrifuge 15 ml conical tube at 200 x g for 3 minutes.
- j. Remove supernatant gently to avoid disturbing cell pellet.
- k. Resuspend cell pellet in 2 ml of fresh medium.
- I. Count cells on hemocytometer (per standard protocol).

3. Plating

 Plate cells at a density of 5.4 x 10³ cells/cm². For a T-75 tissue flask add 4.05 x 10⁵ cells in 12-15 ml Complete Growth Medium.

C. Freezing Cells

- a. Warm Complete Growth Medium and Trypsin solution to 37 °C by placing in 37 °C H₂O bath or in Tissue Culture Incubator.
- b. Make 2X Freeze Medium, (see Section V.2) adjust volume according to volume needed (*Will be mixed 1:1 with Growth Medium*).
- Spray down bottles containing Growth Medium, Trypsin, and the 2X Freeze Medium tube with 70% EtOH before placing in Tissue Culture Hood.
- Remove medium from T-75 flask containing Mesenchymal Stem Cells from incubator.
- e. Gently wash flask with 5-10 ml of sterile 1X PBS (Ca²⁺ and Mg²⁺ free)
- f. Remove PBS.
- g. Add 3 ml of Trypsin to each flask place at 37 °C for 2-3 minutes (until cells are no longer attached to the plate, which should take less than 5 minutes) in Tissue Culture Incubator.
- h. Add 5 ml of Complete Growth Medium to flask.
- i. Transfer cells to 15 ml conical tube.
- j. Centrifuge 15 ml conical tube at 200 X g for 3 minutes.
- k. Remove supernatant gently to avoid disturbing cell pellet.
 - i. Resuspend cell pellet in 2 ml of Complete Growth Medium.
 - ii. Count cells on hemocytometer (per standard protocol).
 - iii. Dilute cells to a desired concentration for freezing.

Notes: We recommend a concentration of no less than 1 x 10^6 cells/ml. This concentration is enough to seed one T-75 flask (remember, the cells will be diluted 1:1 with freeze medium, final concentration 0.5×10^6 cells/ml).

One T-75 flask is sufficient for 1-3 vials of 5×10^5 cells per vial.

- I. Add equal volume of 2X Freeze Medium to the cells, mix gently.
- m. Aliquot 1 ml of cells into labeled cryovials.
- n. Place on ice for 15-30 minutes.
- o. Transfer to cell freezing container and place in -80°C freezer overnight.

E4/14/09v1

p. Transfer to liquid nitrogen freezer for long term storage.

Note: Vapor phase is recommended to maintain viability.

VII. References

- Phinney DG, Prockop DJ. 2007. Concise Review: Mesenchymal Stem/Multipotent Stromal Cells: The State of Transdifferentiation and Models of Tissue Repair-Current Views. Stem Cells 25: 2896-2902
- 2. Kolf CM, Cho E, Tuan RS. 2007. Biology of Adult Mesenchymal Stem Cells: Regulation of Niche, Self-Renewal and Differentiation. *Arthritis Research and Therapy* 9:204-214
- Javazon EH, Colter DC, Schwarz EJ, Prockop DJ. 2001. Rat Marrow Stromal Cells are More Sensative to Plating Density and Expand More Rapidly from Single-Cell-Derived Colonies than Human Marrow Stromal Cells. Stem Cells 19:219-225
- Li Yi, McIntosh K, Chen J, Zhang C, Gao, Q, Borneman J, Ragniski K, Mitchell J, Shen L, Zhang J, Lu D, Chopp M. 2006. Allogeneic bone marrow stromal cells promote glial-axonal remodeling without immunologic sensitization after stroke in rats. *Experimental Neurology* 198:313-25

VIII. Troubleshooting

PROBLEM	CAUSE	ACTION
Poor Viability from initial	initial	Ensure medium is added slowly to reequillibrate the RMSC from freeze medium
freeze		Ensure cells were removed from freeze medium immediately after vial has been thawed
		Ensure Vial of cells was thawed at 37°C; Fresh medium was prewarmed to 37°C
Poor	Frequency of Medium Change	Ensure medium is changed every 3-4 days
proliferation		Ensure pH of medium fresh medium has not changed
	CO ₂ Incubator not humidified	Add sterile water to CO ₂ incubator per manufactures instructions
	No gas exchange is allowed by flask	Ensure cap is loosened to allow air gas or use vented flask
	Tissue Culture Labware not ideal for RMSC	Used Corning or Nunc Treated Labware
Cells were	Cells were allowed to become over	Extend time in trypsin
clumpy after passaging, limited recovery	confluent and lay down matrix	Tirtrate cells to remove as many cells as possible from matrix
of single cells		Remove visual matrix aggregated from tube before spinning (will reduce cell recovery)
		Pass cell suspension through cell strainer (will reduce cell recovery)

5

DDODLEM	CALICE	ACTION
PROBLEM	CAUSE	ACTION
Poor Cell Recovery from flask (for cell growth)	Cell seeding density too high	Passage cells at lower confluency
Contamination of Cells	Contaminated Medium	To prevent contamination, filter medium through a 0.22 µm filter before use
		Never use contaminated medium once cloudy or after microorganisms are visible under the microscope
	Improper aseptic technique	Spray down hands, reagents and hood with 70% ethanol before opening any flasks
	Hood is working improperly	Ensure hood is currently certified; blower is on and functioning
		Wipe down hood with 70% ethanol
	Contaminated CO ₂ Incubator	Ensure CO ₂ incubator is free of microbial growth

IX. Related Products Available From Trevigen

Contact Trevigen for details of our unique product line for studying DNA damage and repair. All of Trevigen's kits include highly qualified enzymes, substrates, buffers, full instructions for use, and a synopsis specific for your kit.

Differentiation:

Catalog #	Description	Size
5000-001-01	Cultrex [®] Rat Mesenchymal Stem Cells	1 vial
5000-001-K	Cultrex® Rat Mesenchymal Stem Cell Starter Kit	1 kit
5010-024-K	Cultrex® Adipogenic Differentiation Kit for RMSC	24 samples
5011-024-K	Cultrex® Osteogenic Differentiation Kit for RMSC	24 samples

3D Culture Kits:

Catalog #	Description	Size
3445-096-K	Cultrex® 3D Culture BME Cell Proliferation Assay Kit	96 tests
3446-096-K	Cultrex® 3D Culture Laminin I Cell Proliferation Assay	96 tests
3447-096-K	Cultrex [®] 3D Culture 96 Well Collagen I Cell Prolif Assay	96 tests
3448-020-K	Cultrex [®] 3D Culture Cell Harvesting Kit	96 tests

Invasion/Migration Kits:

Catalog#	Description	Size
3455-024-K	Cultrex® 24 Well BME Cell Invasion Assay	24 inserts
3460-024-K	CultreCoat® 24 Well BME-Coated Cell Invasion Assay	24 inserts
3465-096-K	Cultrex [®] 96 Well Cell Migration Assay	96 samples
3465-024-K	Cultrex [®] 24 Well Cell Migration Assay	12 samples
3455-096-K	Cultrex® 96 well BME Cell Invasion Assay	96 samples
3456-096-K	Cultrex [®] 96 well Laminin I Cell Invasion Assay	96 samples
3457-096-K	Cultrex [®] Collagen I Cell Invasion Assay	96 samples
3458-096-K	Cultrex® Collagen IV Cell Invasion Assay	96 samples
3471-096-K	In vitro Angiogenesis Assay Endothelial Cell Invasion	96 samples

6

Accessories:

Catalog#	Description	Size
4870-500	10X PBS (Ca ²⁺ , Mg ²⁺ free)	500 ml
5000-050-02	Cultrex® Qualified RMSC FBS	50 ml
5000-500-03	Cultrex® Qualified RMSC Medium	500 ml
3400-010-01	Cultrex [®] Mouse Laminin I	1 mg
3440-100-01	Cultrex [®] Rat Collagen I	100 mg
3442-050-01	Cultrex [®] Bovine Collagen I	50 mg
3430-005-02	Cultrex® BME with phenol red, PathClear®	5 ml
3431-005-02	Cultrex® BME with phenol red, reduced growth factor PathClear®	5 ml
3432-005-02	Cultrex [®] BME no phenol red, PathClear [®]	5 ml
3433-005-02	Cultrex [®] BME no phenol red, reduced growth factor PathClear [®]	5 ml
3430-005-01	Cultrex® BME with Phenol Red	5 ml
3432-005-01	Cultrex [®] BME; no Phenol Red	5 ml
3431-005-01	Cultrex® BME with Phenol Red; Reduced Growth Factors	5 ml
3433-005-01	Cultrex® BME no Phenol Red; Reduced Growth Factors	5 ml
3416-001-01	Cultrex [®] Bovine Fibronectin	1 mg
3420-001-01	Cultrex [®] Human Fibronectin, PathClear [®]	1 mg
3417-001-01	Cultrex® Bovine Vitronectin	50 μg
3421-001-01	Cultrex [®] Human Vitronectin, PathClear [®]	50 μg
3438-100-01	Cultrex [®] Poly-L-Lysine	100 ml
3439-001-01	Cultrex® Poly-D-Lysine	100 ml

The product accompanying this document is intended for research use only and is not intended for diagnostic purposes or for use in humans.

Trevigen, Inc.

8405 Helgerman Ct. Gaithersburg, MD 20877 Tel: 1-800-873-8443 • 301-216-2800 Fax: 301-560-4973 e-mail: info@trevigen.com

www.trevigen.com



Please Recycle

7



211 bis Avenue Kennedy - BP 1140 03103 Montluçon - France 33 (0) 4 70 03 88 55 Fax 33 (0) 4 70 03 82 60 e-mail interchim@interchim.com Agence Paris - Normandie 33 (0) 1 41 32 34 40 Fax 33 (0) 1 47 91 23 90 e-mail interchim.paris@interchim.com