

# Cultrex® HA-R-Spondin1-Fc 293T Cells

Cat# 3710-001-01

Stably transfected 293T cells that express murine Rspo1 with an N-terminal HA epitope tag and fused to a C-terminal murine IgG2a Fc fragment.

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#### I. Introduction

Roof plate-specific Spondin-1 (R-Spondin 1 or RSPO1), also known as CRISTIN3, is a 27 kDa secreted activator protein that belongs to the R-Spondin family. R-Spondins positively regulate Wnt/beta-catenin signaling, most likely by acting as a ligand for LGR4-6 receptors and an inhibitor for ZNRF3. R-Spondin-1 induces proliferation of intestinal crypt epithelial cells, increases intestinal epithelial healing, and supports intestinal epithelial stem cell renewal [1-5]. The 293T cell line is stably transfected to express murine Rspo1 with an N-terminal HA epitope tag and fused to a C-terminal murine IgG2a Fc fragment.

#### II. Precautions and Limitations

- Successful and consistent results are dependent upon the quality and degree of characterization of the cells under investigation. Highly passaged cells may undergo both genotypic and phenotypic changes that render them an inadequate in vitro model for specific investigations. We recommend for all studies that highly qualified low passage number cells are used to ensure reliable and reproducible results.
- 2. For Research Use Only. Not for use in diagnostic procedures.
- 3. This cell line is not known to harbor any agent known to cause disease in healthy adult humans. Handle as a potentially biohazardous material under at least a Biosafety Level 1 containment. This cell line has NOT been screened for Hepatitis B, human immunodeficiency viruses or other adventitious agents. Trevigen recommends that appropriate safety procedures be used when handling all cell lines, especially those derived from human or other primate material. Trevigen assumes no liability for damage resulting from handling or contact with these products.

## III. Materials Supplied

<u>Component</u>	<b>Quantity</b>	<u>Storage</u>	Catalog #
HA-R-Spondin1-Fc	1 Vial (10 <sup>6</sup> Cells)	Liquid Nitrogen***	3710-001-01
293T Cells			

<sup>\*\*\*</sup>Shipped on Dry Ice, immediately thaw for use, or for long term storage place in vapor phase of liquid nitrogen.

## IV. Materials/Equipment Required But Not Supplied

#### Equipment

- 1. 1 20 μl, 20 200 μl, and 200 1000 μl pipettors
- Class 2 Biosafety Hood
- 3. 37°C CO<sub>2</sub> incubator
- 4. 37°C Water Bath
- 5. 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 that allows for slow freezing of cells (e.g. Fisher Scientific cat#15-350-50)

#### Reagents

- Cell Culture Medium: DMEM High Glucose (Life Technologies cat# 11965-092 or equivalent)
- 2. Cell Harvesting Reagent (Trypsin, Dispase, or equivalent)
- 3. Fetal Bovine Serum
- 100X Penicillin-Streptomycin supplement for Media (Life Technologies cat# 15140-122 or equivalent)
- 5. 100 mg/ml Zeocin (Life Technologies cat# 250-01)
- 6. GlutaMAX (Life Technologies cat# 35050061)
- 7. CD293 Medium (Life Technologies cat# 11913-019)
- 8. PBS (Mg<sup>2+</sup>, Ca<sup>2+</sup> free) or HBSS, tissue culture grade
- 9. Trypan blue or equivalent viability stain
- 10. DMSO, tissue culture grade
- 11. 70% Ethanol
- 12. Sterile ddH<sub>2</sub>0
- 13. Protein A Agarose Purification Kit (KPL cat# 553-50-00 or equivalent)

#### **Disposables**

- 1. Cell culture flasks, 25 cm<sup>2</sup>, 75 cm<sup>2</sup>, or 185 cm<sup>2</sup>
- 15 ml tubes
- 3. 0.22 µm Filter Unit
- 4. 1 200 μl and 200 1000 μl pipette tips
- 5. 2, 5 and 10 ml serological pipettes
- 6. gloves

## V. Reagent Preparation

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

#### 1. Basal Growth Medium

For 500 ml of Medium:

DMEM Medium:	440 ml
Fetal Bovine Serum	50 ml
100X Penicillin-Streptomycin	5 ml
GlutaMAX	5 ml

Filter sterilize medium and store at 4°C for one month. Ensure medium is at room temperature or 37°C prior to use.

#### 2. Selection Growth Medium (containing 300 µg/ml Zeocin)

For 50 ml of Medium:

Basal Growth Medium 50 ml 100 mg/ml Zeocin 150 µl

Prepare immediately before use. Scale as needed.

#### 3. Freeze Medium

For 10 ml:

Basal Growth Medium: 5.0 ml FBS: 3.0 ml DMSO: 2.0 ml

#### 4. Expression Medium

For 50 ml of Medium:

CD293 Medium 1,000 ml GlutaMAX 10 ml

#### VI. Protocol

These procedures should be performed in a biological hood utilizing aseptic technique to prevent contamination. Vessels should be sprayed down with 70% ETOH before placing in Tissue Culture Hood.

### A. Thawing HA-R-Spondin1-Fc 293T Cells:

- Pre-warm Basal Growth Medium (Section V.1) to 37°C by placing in H<sub>2</sub>O bath or in Tissue Culture Incubator.
- Immediately before use, remove the vial of cryopreserved cells from liquid nitrogen freezer and thaw quickly (3-4 minutes) in a 37°C H<sub>2</sub>O bath. Ensure cells are completely thawed before proceeding and do not leave cells at 37°C past thawing.
- 3. Aseptically, transfer the thawed cells to an empty 15 ml conical tube. Wash ampoule with 1 ml of warm Basal Growth Medium and add to thawed cells. Add 1 ml of warm Basal Growth Medium to 15 ml conical tube containing cells, gently swirling to mix between drops. Total Volume should be 3 ml.
- 4. Centrifuge cells at 200 x g for 3 minutes.
- Remove supernatant gently to avoid disturbing cell pellet and resuspend cell pellet in 6 ml of fresh Basal Growth Medium.
- 6. Transfer cell suspension to a sterile T25 (25 cm²) Tissue Culture Flask.
- 7. Place Tissue Culture Flask/Dish in 5% CO<sub>2</sub> Tissue Culture Incubator at 37°C.
- 8. Change medium in flasks using freshly prepared and pre-warmed Selection Growth Medium (Section V.2) the next day.

#### B. Passaging HA-R-Spondin1-Fc 293T Cells:

Note: HA-R-Spondin1-Fc 293T Cells may be sequentially passaged under selection into larger culture vessels to expand cell numbers. We recommend that cultures be maintained at densities between 40 – 90% confluent for optimal growth and survival.

- Selection Growth Medium should be changed every 2-3 days. Cells should be passaged when 80-90% confluent for optimal growth rate/efficiency. We recommend splitting cells at a density of 1:4 to 1:6. Cells have a doubling time of approximately 24 hrs.
- Prepare Selection Growth Medium on day of use. Warm Basal Growth Medium to 37°C by placing in 37°C H<sub>2</sub>O bath or in Tissue Culture Incubator. In a sterile container, add the required volume (10-12 ml/flask) of Basal Growth Medium and Zeocin (final concentration of 300 μg/ml).
- Remove medium from T25 flask containing the HA-R-Spondin1-Fc 293T Cells.
- Gently wash flask with 5 ml of sterile 1X PBS (Ca<sup>2+</sup> and Mg<sup>2+</sup> free) and remove PBS.
- 5. Add 1 ml of warmed Trypsin to the flask and place at 37°C for 3-5 minutes until cells are no longer attached to plate. Add 3 ml of Basal Growth Medium to flask to inactivate Trypsin and transfer to 15 ml conical tube.
- 6. Centrifuge cells at 200 x g for 3 minutes.
- Remove supernatant gently to avoid disturbing cell pellet and resuspend cell pellet in 2 ml of fresh Selection Growth Medium.
- 8. Add 1 ml of cell suspension to 11 ml of Selection Growth Medium and transfer to one T75 flask.

#### C. Freezing Cells

- In general, one confluent T75 flask will provide cells for 10-12 vials of 1 x 10<sup>6</sup> cells per vial.
- 2. Prepare 2X Freeze Medium, (see Section V.3) according to volume required. Typically, 0.5 ml of 2X Freeze Medium is mixed with 0.5 ml of 2x10<sup>6</sup> cells.
- 3. To prepare cells remove medium from T-75 flask containing HA-R-Spondin1-Fc 293T Cells.
- 4. Gently wash flask with 5-10 ml of sterile 1X PBS (Ca<sup>2+</sup> and Mg<sup>2+</sup> free) and remove PBS.
- Add 2 ml of pre-warmed Trypsin to each flask, and place at 37°C for 2-3 minutes until cells are no longer attached to the plate. Add 4 ml of Basal Growth Medium to flask to inactivate Trypsin and transfer to 15 ml conical tube.
- 6. Centrifuge cells at 200 x g for 3 minutes.
- Remove supernatant gently to avoid disturbing cell pellet and resuspend cell pellet in 2 ml of Basal Growth Medium.
- Count cells on hemocytometer (per standard protocol) and dilute cells to 2 x 10<sup>6</sup> cells/ml in Basal Growth Medium.
- Add equal volume of 2X Freeze Medium to the cells, mix gently and aliquot 1 ml of cells into labeled cryovials.
- For initial storage, first place cryovials on ice for 15-30 minutes. Transfer cells to specialized cell freezing container and place in -80°C freezer overnight.
- 11. For long term storage, transfer cells (next day) to liquid nitrogen freezer to ensure long-term viability.

#### D. HA-R-Spondin1-Fc Expression and Purification

 After at least five days in Selection Growth Medium, HA-R-Spondin1-Fc cells may be expanded in Basal Growth Medium until they reach 80% confluence

- in the desired culture vessel. A 3-layer flask (Falcon 353143) has a surface area of 525 cm $^2$ , accommodates 150-200 ml of medium, and can produce up to 300  $\mu$ g of HA-R-Spondin-1-Fc.
- Once the cells reach 80% confluence, change the medium to CD293
  medium containing GlutaMAX, and continue to culture for 7-10 days. The
  cells will detach from the cell culture vessel and grow in suspension after
  several days.
- Collect the supernatant, and centrifuge at 3,000 rpm for 15 minutes at 4 °C to remove cells and debri.
- 4. Filter the supernatant through 0.22 µm filter at 4 °C.
- 5. Purify HA-R-Spondin1-Fc using the Fc tag via Protein A Agarose Purification.

#### VII. References

- 1. Ootani, A., et al., Sustained in vitro intestinal epithelial culture within a Wnt-dependent stem cell niche. Nat Med, 2009. **15**(6): p. 701-706.
- 2. Barker, N., et al., *Lgr5+ve Stem Cells Drive Self-Renewal in the Stomach and Build Long-Lived Gastric Units In Vitro.* Cell Stem Cell, 2010. **6**(1): p. 25-36.
- Sato, T., et al., Single Lgr5 stem cells build crypt–villus structures in vitro without a mesenchymal niche. Nature, 2009. 459(7244): p. 262-265.
- 4. Sato, T. and H. Clevers, *Growing Self-Organizing Mini-Guts from a Single Intestinal Stem Cell: Mechanism and Applications.* Science, 2013. **340**(6137): p. 1190-1194.
- 5. Jung, P., et al., *Isolation and in vitro expansion of human colonic stem cells*. Nat Med, 2011. **17**(10): p. 1225-7.

VIII. Troubleshooting

PROBLEM	CAUSE	ACTION
		Ensure medium is added slowly to re- equilibrate the cells from freeze medium
Poor viability from initial freeze	Improper thawing of cells	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 warmed to 37°C
	Fetal Bovine Serum not optimal for cell growth	Try alternative lot/source of FBS
	Media not optimal for cell growth	Ensure medium is of the proper formulation
	Frequency of medium change	Ensure medium is changed every 2-3 days
Poor proliferation		Ensure pH of medium fresh medium has not changed
	CO <sub>2</sub> incubator not humidified	Add sterile water to CO <sub>2</sub> incubator per manufactures instructions
	No gas exchange is allowed by flask	Ensure cap is loosened to allow air gas or use vented flask
	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
Contamination of Cells	Hood is working improperly	Check to make sure blower is on and functioning
		Ensure hood is currently certified
		Wipe down hood with 70% ethanol
	Contaminated CO <sub>2</sub> Incubator	Ensure CO <sub>2</sub> incubator is free of microbial growth
Loss of HA-R-Spondin-	Not growing cells in Zeocin	Ensure Zeocin was added to basal medium just before addition to cells
1-Fc expression	Frequency of Medium Change	Ensure medium is being changed every 2-3 days.

## IX. Related Products Available From Trevigen

### Related Products:

Catalog#	Description	Size
3432-005-01	Cultrex® Basement Membrane Extract, PathClear®	5 ml
3433-005-02	Cultrex® Reduced Growth Factor BME, PathClear®	5 ml
3532-005-02	Cultrex® Basement Membrane Extract, Type 2, PathClear®	5 ml
3533-005-02	Cultrex® Reduced Growth Factor BME, Type 2, PathClear®	5 ml
3632-005-02	Cultrex® Basement Membrane Extract, Type 3, PathClear®	5 ml
3445-005-01	Cultrex <sup>®</sup> 3-D Culture Matrix <sup>™</sup> BME, PathClear <sup>®</sup>	5 ml
3446-005-01	Cultrex <sup>®</sup> 3-D Culture Matrix <sup>™</sup> Laminin I	5 ml
3447-020-01	Cultrex <sup>®</sup> 3-D Culture Matrix <sup>™</sup> Collagen I	100 mg
3434-005-02	Cultrex® Stem Cell Qualified RGF BME, PathClear®	5 ml
3415-001-03	Cultrex® Stem Cell Qualified Human BME, PathClear®	1 mg
3400-010-03	Cultrex® Stem Cell Qualified Laminin I, PathClear®	1 mg
3420-001-03	Cultrex® Stem Cell Qualified Human Fibronectin, PathClear®	1 mg
3420-001-03	Cultrex® Stem Cell Qualified Human Vitronectin, PathClear®	200 µg
3400-010-01	Cultrex <sup>®</sup> Mouse Laminin I	1 mg
3400-010-02	Cultrex® Mouse Laminin I, PathClear®	1 mg
3410-010-01	Cultrex <sup>®</sup> Mouse Collagen IV	1 mg
3440-100-01	Cultrex <sup>®</sup> Rat Collagen I	100 mg
3442-050-01	Cultrex <sup>®</sup> Bovine Collagen I	50 mg
3420-001-01	Cultrex® Human Fibronectin, PathClear®	1 mg
3416-001-01	Cultrex® Bovine Fibronectin, NZHD*	1 mg
3421-001-01	Cultrex <sup>®</sup> Human Vitronectin, PathClear <sup>®</sup>	50 μg
3417-001-01	Cultrex® Bovine Vitronectin, NZHD*	50 µg

**Related Assays and Kits:** 

Catalog#	Description	Size
3500-096-K	Cultrex <sup>®</sup> 3D Spheroid Cell Invasion Assay	96 samples
3510-096-K	Cultrex® 3D Spheroid Fluorometric Proliferation/Viability Assay	96 samples
3511-096-K	Cultrex <sup>®</sup> 3D Spheroid Colorimetric Proliferation/Viability Assay	96 samples
3470-096-K	Cultrex <sup>®</sup> In Vitro Angiogenesis Assay, Tube Formation Kit	96 samples
3471-096-K	Cultrex <sup>®</sup> In Vitro Angiogenesis Assay, Endothelial Cell Invasion Kit	96 samples
3450-048-SK	Cultrex <sup>®</sup> Directed In Vivo Angiogenesis Assay (DIVAA <sup>™</sup> ) Starter Kit	48 samples
3450-048-K	Cultrex <sup>®</sup> DIVAA <sup>™</sup> Kit	48 samples
3450-048-IK	Cultrex <sup>®</sup> DIVAA <sup>™</sup> Inhibition Kit	48 samples
3465-024-K	Cultrex <sup>®</sup> 24 well Migration Cell Assay	24 inserts
3455-024-K	Cultrex® 24 well BME Cell Invasion Assay	24 inserts
3456-024-K	Cultrex <sup>®</sup> 24 well Laminin I Cell Invasion Assay	24 inserts
3457-024-K	Cultrex® 24 well Collagen I Cell Invasion Assay	24 inserts
3458-024-K	Cultrex® 24 well Collagen IV Cell Invasion Assay	24 inserts
3465-096-K	Cultrex <sup>®</sup> 96 well Migration Cell Assay	96 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® 96 well Collagen I Cell Invasion Assay	96 samples
3458-096-K	Cultrex® 96 well Collagen IV Cell Invasion Assay	96 samples
3448-020-K	Cultrex® 3-D Culture Cell Harvesting Kit	20 samples

<sup>\*</sup>New Zealand Herd Derived

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## 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

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