

Healthy Control Human iPSC Line, Female, SCTi003-A

Catalog #200-0511

~1 million viable cells/vial



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Product Description

Healthy Control Human iPSC Line, Female, SCTi003-A, was derived from peripheral blood mononuclear cells (PBMCs) from a 48-year-old donor. Extensive quality control procedures were undertaken in the induced pluripotent stem cell (iPSC) manufacturing process to ensure optimal product performance and reproducibility. SCTi003-A is karyotypically stable, demonstrates trilineage differentiation potential, expresses markers of the undifferentiated state, and was reprogrammed using a non-integrating reprogramming technology. This cell line may be used as a healthy control for a multitude of pluripotent stem cell research applications, including downstream differentiation to lineage-specific cell types and organoids.

SCTi003-A is manufactured with mTeSR™ Plus (Catalog #100-0276) and is fully compatible with STEMdiff™ cell culture media products, allowing for standardized high-quality maintenance and differentiation to various cell types, such as cardiomyocytes, neurons, astrocytes, and microglia.

SCTi003-A was derived from an $\alpha\beta$ T cell and has undergone variable-diversity-joining (VDJ) rearrangement.

Stability and Storage

Cells are frozen in a cryopreservation medium containing dimethyl sulfoxide (DMSO). Product stable at -135°C or colder for 12 months from date of receipt. Thawed samples must be used immediately.

Precautions

Cell Screening: iPSC master cell banks are screened for AAV2, BK virus, Epstein-Barr Virus, Hepatitis A, Hepatitis B, Hepatitis C, Herpes Simplex 1 and 2, Herpes Virus Type 6, 7, and 8, HIV-1, HIV-2, HPV-16, HPV-18, Human Adenovirus, Human Cytomegalovirus, Human Foamy Virus, Human T-Lymphotropic Virus, John Cunningham Virus, LCMV, Parvovirus B19, Sarbecovirus (SARS Virus), Seoul Virus, Corynebacterium Bovis, and Mycoplasma (Human Comprehensive CLEAR Panel) by PCR. Commercial iPSC banks are tested for the absence of Mycoplasma by PCR. As testing cannot completely guarantee that the donor was virus-free, THIS PRODUCT SHOULD BE TREATED AS POTENTIALLY INFECTIOUS and only used following appropriate handling precautions such as those described in biological safety level 2.

Storage of frozen cell products in the vapor phase of a liquid nitrogen storage tank is recommended. Storage in the liquid phase can result in cross-contamination if the vial breaks or is not sealed properly. Storage in the liquid phase also increases the potential for liquid nitrogen to penetrate the vial and cause it to explode when removed from storage. Use of a face shield is required as a safety precaution when transferring cells from one container to another. When handling this product, do not use sharps such as needles and syringes.

STEMCELL cannot guarantee the biological function or any other properties associated with performance of cells in a researcher's individual assay or culture systems. STEMCELL assures the cells will meet the specifications only when assessed immediately after thawing by our test methods.

FOR IN VITRO RESEARCH USE ONLY. NOT APPROVED FOR DIAGNOSTIC, THERAPEUTIC, OR CLINICAL APPLICATIONS.
NOT APPROVED FOR HUMAN OR VETERINARY USE IN VIVO.

Donor Information

Age [†]	48	
Sex [‡]	Female	
Ethnicity and/or Race [†]	Caucasian	
Ancestry [†]	0% African 100% European	0% East Asian 0% South Asian
Diagnosis [†]	Clinically unaffected at donation	
Height [†]	168 cm	
Weight [†]	62.1 kg	
BMI [†]	22.1 kg/m ²	
Blood type [†]	B-	
Tobacco Use [†]	Non-smoker	
HLA Haplotype [†]	HLA Class I: A*24:02:01G, 26:01:01G B*07:02:01G, - C*07:02:01G, - E*01:01:01G, - F*01:01:01G, 01:04 G*01:01:02G, 01:04:01G	HLA Class II: DRB1*15:01:01G, - DRB3*, - DRB4*, - DQA1*01:02:01G, - DQB1*06:02:01G, - DPA1*01:03:01G, - DPB1*02:01:02G, 04:01:01G

[†] Self-declared

[‡] Calculated

Directions for Use

SCTi003-A was manufactured using mTeSR™ Plus, and mTeSR™ Plus is the recommended thawing medium. Thawed cells should be seeded into tissue culture-treated cultureware pre-coated with Corning® Matrigel® hESC-Qualified Matrix. For instructions on preparing complete mTeSR™ Plus and coated cultureware, refer to the Technical Manual: Maintenance of Human Pluripotent Stem Cells in mTeSR™ Plus (Document #1000007757), available at www.stemcell.com, or contact us to request a copy.

NOTE: The following instructions are for seeding cells into coated 6-well plates. If using other cultureware, adjust volumes accordingly.

- Have all tubes, warmed mTeSR™ Plus (15 - 25°C), and coated cultureware ready before starting the protocol to ensure that the thawing procedure is completed as quickly as possible.
NOTE: Do not warm mTeSR™ Plus in a 37°C water bath.
- Wipe the outside of the vial of cells with 70% ethanol or isopropanol.
- In a biosafety cabinet, twist the cap a quarter-turn to relieve internal pressure, then retighten.
- Quickly thaw cells in a 37°C water bath by gently shaking the vial. Remove the vial when a small frozen cell pellet remains. Do not vortex cells.
NOTE: For optimal thawing in a sterile and controlled manner, use the ThawStar® CFT2 Automated Thawing System.
- Wipe the outside of the vial with 70% ethanol or isopropanol.
- Use a 2 mL serological pipette to transfer the contents of the cryovial to a 15 mL conical tube.
NOTE: Using a 2 mL serological pipette instead of a 1 mL pipettor will minimize breakage of cell aggregates.
- Add 5 - 7 mL of warm mTeSR™ Plus dropwise to the 15 mL tube, gently mixing as the medium is added.
- Centrifuge cells at 300 x g for 5 minutes at room temperature.
- Aspirate the medium, leaving the cell pellet intact. Resuspend the cell pellet in 1 mL of mTeSR™ Plus by gently flicking the tube. Avoid pipetting up and down and take care to maintain the cells as aggregates.
- When cells are ready to be plated, aspirate the Matrigel® solution from a coated 6-well plate and add 2 mL of mTeSR™ Plus to each well.
- Aliquot the 1 mL cell suspension into a coated 6-well plate containing 2 mL of mTeSR™ Plus at six different densities: (1) 150 µL, (2) 100 µL, (3) 75 µL, (4) 50 µL, (5) 25 µL, and (6) 15 µL.
NOTE: Gently flick the tube as many times as needed to ensure equal distribution of the cell aggregates between the wells.
NOTE: The remaining 585 µL of cell suspension can be seeded into two wells of a separate 6-well plate to initiate backup cultures.
- Place the plates in a 37°C and 5% CO₂ incubator. Move the plate in several quick, short, back-and-forth and side-to-side motions to distribute the cell aggregates. Do not disturb the plate for 24 hours.

NOTE: Uneven distribution of aggregates may result in increased differentiation of human iPSCs.

13. Perform medium changes as desired using mTeSR™ Plus and visually assess cultures daily to monitor growth and morphology. See Figure 1 for expected growth characteristics during the first seven days after thawing. Medium can be changed daily, every other day, or up to two consecutive days of feeding can be skipped when using mTeSR™ Plus. To skip two consecutive days of feeding, add twice the volume of medium.

NOTE: If only a few undifferentiated colonies are observed after thawing, it may be necessary to select only these colonies for passaging and replat them in the same size well (i.e. without splitting) on a newly coated plate.

14. On Day 6 - 8, select the well with optimal iPSC colony density for passaging (Figure 2). The culture should consist of healthy iPSC colonies that are large, compact, and have centers that are dense compared to their edges.
15. Use a microscope to visually identify regions of differentiation. Mark these using a felt tip or lens marker on the bottom of the plate. Remove regions of differentiation by scraping with a pipette tip.

NOTE: Removing regions of differentiated cells is optional in subsequent passages.

16. Wash each well with 1 mL of D-PBS (Without Ca⁺⁺ and Mg⁺⁺). Aspirate to remove D-PBS from the wells.
17. Passage iPSCs from the optimal well using ReLeSR™. Add 1 mL of ReLeSR™ to each well and aspirate within 1 minute, so that colonies are exposed to a thin film of liquid. For SCTi003-A, incubate the culture at room temperature for 4 to 6 minutes and split at a ratio of 1 in 30 to 1 in 60 every 5 - 7 days.

NOTE: When incubating cells with ReLeSR™, monitor cell detachment under the microscope. To prevent differentiated regions from lifting off the surface of the well, the optimal incubation time for this cell line should be determined in your own laboratory.

NOTE: If the colonies are too dense or too sparse, adjust the split ratio accordingly at the next time of passaging. For complete instructions on passaging human iPSCs cultured in mTeSR™ Plus using ReLeSR™, refer to the Technical Manual: Maintenance of Human Pluripotent Stem Cells in mTeSR™ Plus (see section 5.1, Document #1000007757), available at www.stemcell.com, or contact us to request a copy.

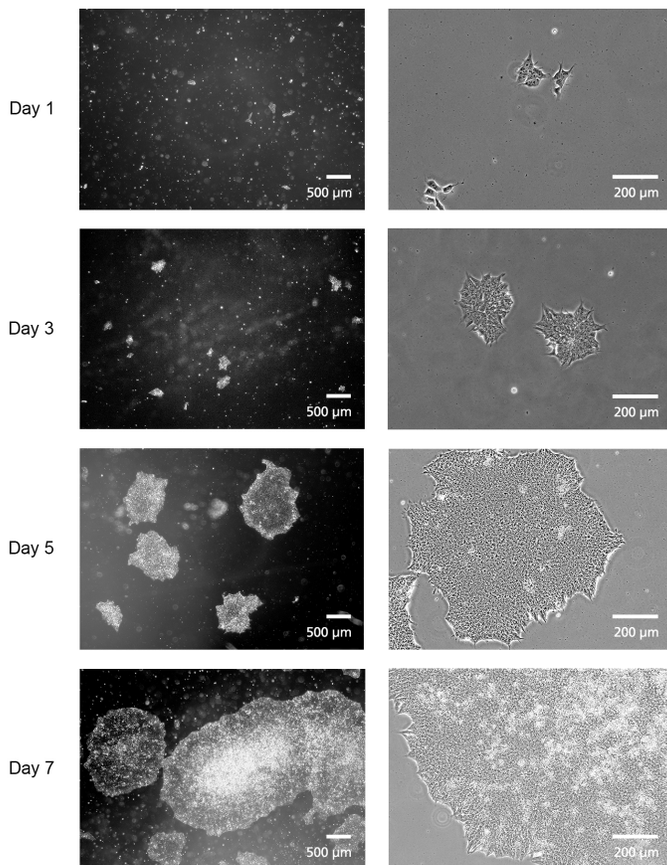
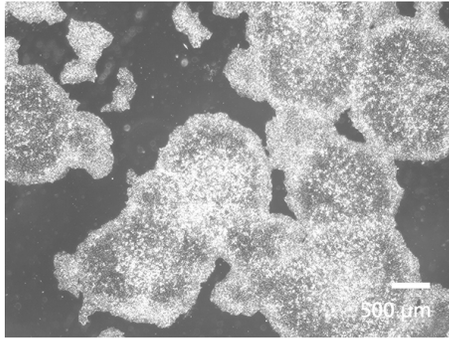


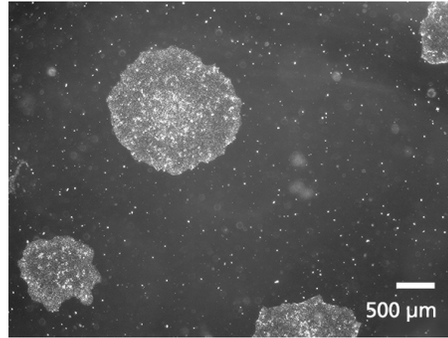
Figure 1. Recovery of SCTi003-A iPSCs on Days 1 - 7 After Thaw.

SCTi003-A iPSCs were recovered in mTeSR™ Plus on Corning® Matrigel® hESC-Qualified Matrix and imaged at a magnification of 20X (left) and 100X (right) for seven days. For this concentration of cellular aggregates at thaw, Day 7 would be the optimal time for passaging.

High Density



Low Density



Optimal Density

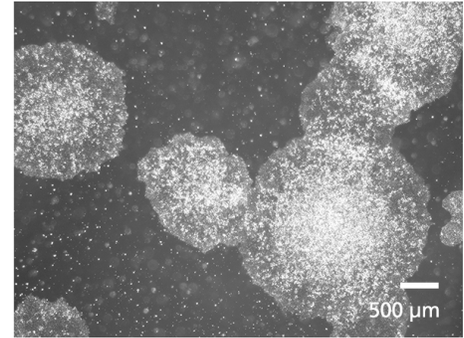


Figure 2. SCTi003-A iPSCs Demonstrate Varying Colony Densities When Seeded at a Range of Starting Concentrations.

SCTi003-A iPSCs were thawed at a range of concentrations and expanded in mTeSR™ Plus for seven days. Final colony densities were imaged at a magnification of 20X. It is recommended that a culture is passaged once it has reached an optimal density consisting of large, multilayered colonies that have begun to merge.

Notes and Tips

The ideal mean cell aggregate size obtained after step 17 of the Directions for Use is approximately 50 - 200 µm. For suggestions on optimizing the ReLeSR™ passaging protocol, refer to the Notes and Tips section of the Product Information Sheet (Document #1000000243) for ReLeSR™.

Accessory Products

PRODUCT NAME	CATALOG #
CloneR™2	100-0691
Corning® Matrigel® hESC-Qualified Matrix	354277
D-PBS (Without Ca++ and Mg++)	37350
Falcon® 6-Well Flat-Bottom Plate, Tissue Culture-Treated	38016
Falcon® Conical Tubes, 15 mL	38009
Falcon® Serological Pipettes, 2 mL	38002
hPSC Genetic Analysis Kit	07550
Human Pluripotent Stem Cell Trilineage Differentiation qPCR Array	07515
ReLeSR™	05872
STEMdiff™ Trilineage Differentiation Kit	05230
ThawStar® CFT2 Automated Thawing System	100-0650
Trypan Blue	07050

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These iPSCs and their modifications (including but not limited to derivatives or differentiated progeny) may not be used for monetization or commercialization purposes, including without limitation, used to, or with the goal to, perform services or supply products or rights, including in the manufacture of cellular therapies or other therapeutics, for monetary gain or the generation of royalties, revenues, sales or other valuable consideration. For clarity, these iPSCs and their modifications (including but not limited to derivatives or differentiated progeny) may not be used for screening compounds, antibodies, proteins or peptides, except for the purposes of target discovery, target validation, or assay development, provided such activities and the results of such activities are not further used for monetization or commercialization purposes. It may be possible to obtain a further license for the prohibited uses referred to in this Limited Use License. Please contact iPSCrequests@stemcell.com for more details.

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