

PRIME-XV T Cell Expansion XSFM

Catalog #	Product	Size
91141	PRIME-XV T Cell Expansion XSFM	1 L liquid

Intended Use

For research or further manufacturing use only.

Product Description

PRIME-XV T Cell Expansion XSFM is a xeno-free, serum-free medium optimized for the activation and expansion of human T lymphocytes. This medium contains gentamicin and requires additional cytokines according to desired applications.

Quality Assurance

All quality control test results are reported on a lot specific Certificate of Analysis (COA) which is available at www.irvinesci.com or upon request.

Shipping

This product is shipped at 2-8°C. Upon receipt, store immediately at 2-8°C and protect from light.

Storage Instructions and Stability

Handle using aseptic techniques to avoid contamination. Unopened liquid medium is stable for 18 months from date of manufacture, as indicated on the label, when stored at 2-8°C and protected from light. PRIME-XV T Cell Expansion XSFM should be used within one month after opening when stored at 2-8°C and protected by light. PRIME-XV T Cell Expansion XSFM can be aliquoted and stored at or below -10°C in a manual defrost freezer for up to 3 months. Repeated freeze-thaw cycles should be avoided. When ready to use, thaw this medium overnight at 2-8°C in the dark. This product should be used within one month after thaw when stored at 2-8°C and protected by light. This product has not been validated for use beyond the unopened expiry shelf life.

Precautions

The safety and efficacy of this product in diagnostic or other clinical uses has not been established. Please refer to the Safety Data Sheet (SDS) for information regarding hazards and safe handling practices. PRIME-XV T Cell Expansion XSFM should be handled as biohazardous and potentially infectious. Safe laboratory procedures should be observed and appropriate personal protective equipment should be worn when handling this medium. The acute and chronic effects of over-exposure to this product are unknown.

Directions for Use

The following protocol has been utilized for the activation and expansion of negatively selected CD3⁺ T cells obtained from human peripheral blood and for human PBMCs with PRIME-XV T Cell Expansion XSFM. Further optimization may be required depending on the desired application.

Coating Procedure for T Cell Activation

This following is a guideline for T cell activation using anti-CD3 and anti-CD28 antibodies. Alternative approaches to T cell activation (e.g. beads, mitogens, activator antigens, or other T cell receptor antibodies) may be used. Types and amounts are dependent on the experimental design of each individual investigator.

1. Coat culture vessel with anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibodies at 1 µg/mL in PBS (PN# 9236) for 2 hours at 37°C or overnight at 2-8°C with Parafilm® to prevent evaporation.
2. Aspirate coating solution and discard.
3. Wash culture vessel twice with PBS (PN# 9236) before the addition of cells.

Recovery of Cryopreserved Cells

The following is a guideline based on the cryopreservation of negatively selected CD3⁺ T cells obtained from human peripheral blood and for human PBMCs with PRIME-XV T Cell Expansion XSFM. Specific procedures provided by the respective cell supplier may also be utilized according to the tissue source and user preference and need.

1. Pre-warm sufficient amount of PRIME-XV T Cell Expansion XSFM in a 37°C water bath.

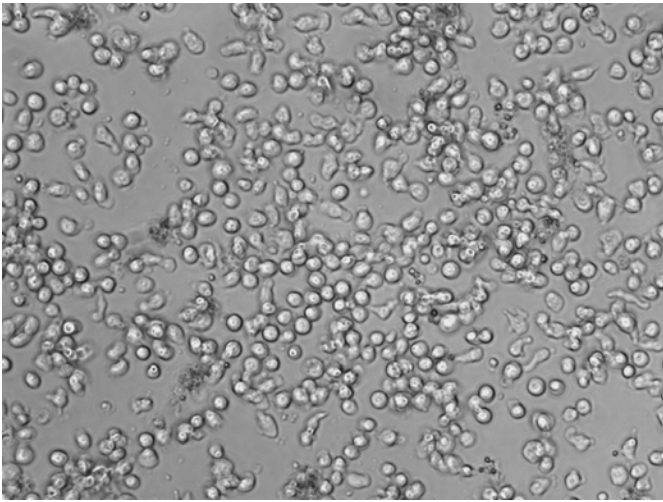
NOTE: To avoid temperature cycling, determine the total volume needed before warming the media.

2. Carefully remove the frozen vial of T cells from liquid nitrogen and thaw by gently stirring the vial in a 37°C water bath for 1 minute.
3. Carefully transfer the entire content of the vial into a 15 mL conical tube containing 10 mL of pre-warmed PRIME-XV T Cell Expansion XSFM.
4. Spin cells down at 400 x g for 10 minutes.
5. Carefully aspirate supernatant leaving a minimum volume of media covering the cell pellet.
6. Supplement PRIME-XV T Cell Expansion XSFM with 33 U/mL (1,800 IU/mL) of IL-2 (PN# 95118).
7. Resuspend the cell pellet with the supplemented PRIME-XV T Cell Expansion XSFM and transfer cells onto the pre-coated culture vessel (see Coating Procedure) at a seeding density of 0.5 x 10⁶ cells/mL (or according to suggested density recommendations from supplier).
8. Incubate the cells in a 37°C, 5% CO₂ humidified incubator.
9. Replace with fresh pre-warmed PRIME-XV T Cell Expansion XSFM supplemented with 33 U/mL (1,800 IU/mL) of IL-2 every 2-3 days of culture. Volume amount of feed should be 70% of the original culture volume.

Harvesting of T cells

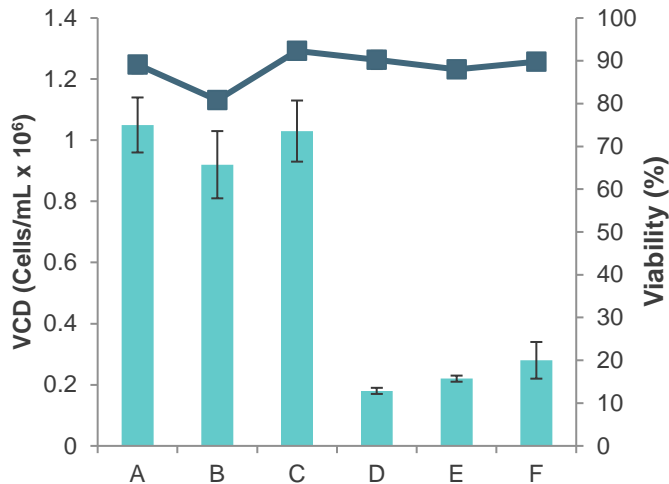
1. Use a cell scraper to gently remove T cells off the culture vessel.
2. Wash with pre-warmed PRIME-XV T Cell Expansion XFSM.
3. Cells are ready for next desired application or analysis.

Sample Data



PRIME-XV T Cell Expansion XFSM maintains characteristic morphology of activated T cell blasts.

Naive CD3⁺ T cells were cultured in IL-2 supplemented PRIME-XV T cell Expansion XFSM on anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibody coated plates at a seeding density of 0.5×10^6 cells/mL. Images were taken on day 2 of culture at 40X magnification.

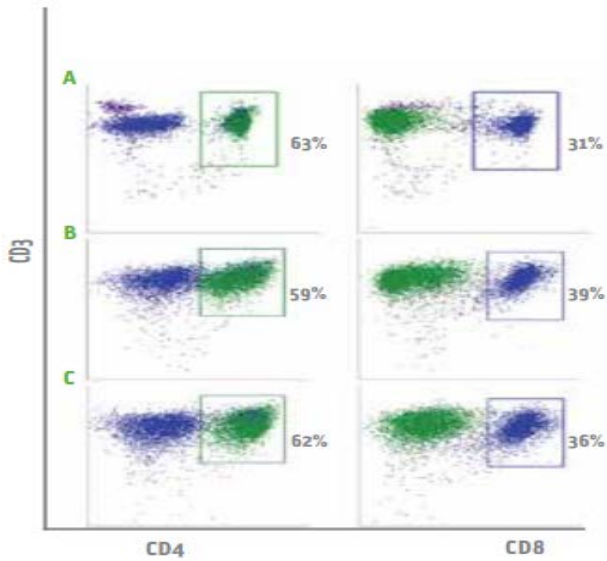


PRIME-XV T Cell Expansion XFSM supports growth of negatively selected purified T cells.

CD3⁺ T cells were cultured in IL-2 supplemented PRIME-XV T Cell Expansion XFSM (A), serum-containing medium (F), and in four leading supplier's serum-free medium (B through E) on anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibody coated plates at a seeding density of 0.5×10^6 cells/mL. Cells were harvested, counted and stained with trypan blue for viability on the fourth day.

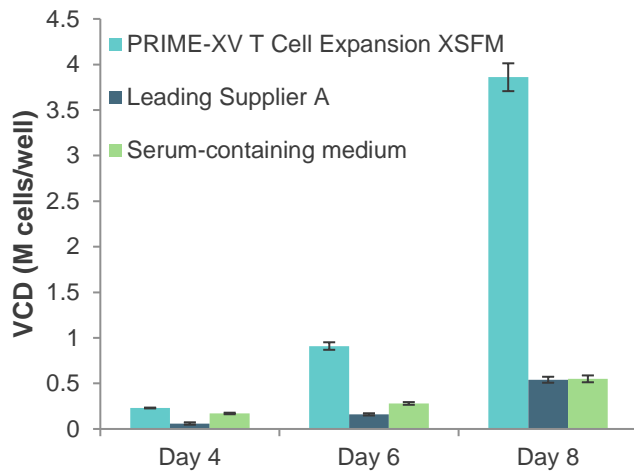
PRIME-XV T Cell Expansion XSFM supports expansion of both CD4⁺ and CD8⁺ T cells.

Naïve CD3⁺ T-cells were cultured in IL-2 supplemented PRIME-XV T Cell Expansion XSFM and a leading supplier's serum-free medium on anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibody coated plates. Phenotypic analyses were performed for the naïve CD3⁺ T-cell after the initial thaw of the cells (A), after four days of culture in PRIME-XV T Cell Expansion XSFM (B) and in the leading supplier's medium (C). The cells were stained with APC-conjugated mouse anti-human CD3 (Clone UTHC1), PE-conjugated mouse anti-human CD8 (Clone SK1), and PerCP-Cy5.5 conjugated mouse anti-human CD4 (Clone OKT4). CD4⁺CD3⁺ cells are highlighted in the green box and CD8⁺CD3⁺ are shown in the blue box.



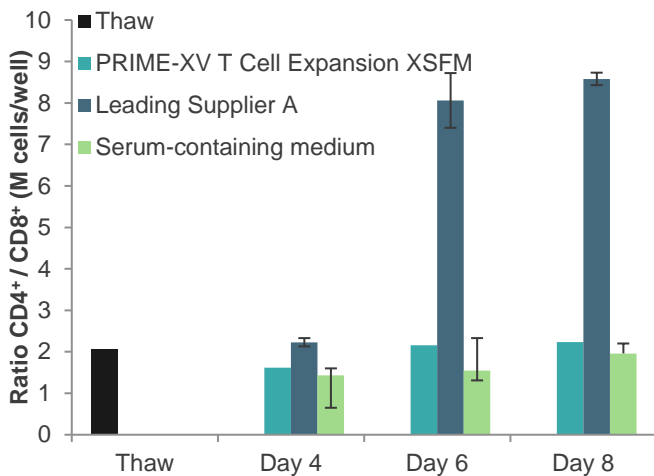
PRIME-XV T Cell Expansion XSFM supports growth of PBMCs.

PBMCs were cultured in IL-2 supplemented PRIME-XV T cell Expansion XSFM, serum-containing medium, and in a leading supplier's serum-free medium on anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibody coated plates at a seeding density of 0.1 x 10⁶ cells per well. Cells were harvested, counted, and stained with trypan blue for viability on the fourth, sixth, and eighth day.



PRIME-XV T cell Expansion XSFM maintains CD4⁺/CD8⁺ ratio in PBMCs.

PBMCs were cultured in IL-2 supplemented PRIME-XV T cell Expansion XSFM, serum-containing medium, and in a leading supplier's serum-free medium on anti-human CD3 (clone UCHT1) and anti-human CD28 (clone CD28.2) antibody coated plates at a seeding density of 0.1 x 10⁶ cells per well. Cells were harvested, counted, and stained with anti-CD4⁺ and anti-CD8⁺ antibodies on the fourth, sixth, and eighth day. Initial CD4⁺/CD8⁺ ratio was measured post-thaw of PBMC.



Related Products

Catalog #	Product	Size
95118	Recombinant Human IL-2 ACF	10 µg
95113	Recombinant Human IL-3 ACF	10 µg
95114	Recombinant Human IL-4 ACF	20 µg
91139	PRIME-XV FreezIS Solution	10 mL and 100 mL liquid
9160	RMPI Medium 1640 w/o L-Glutamine	100 mL, 500 mL liquid
9161	RMPI Medium 1640 with L-Glutamine	100 mL, 500 mL liquid
9236	PBS 1X-Dulbecco's Phosphate Buffered Saline Solution	500 mL, 1 L liquid

Technical Support

CONTACT US

For more information or assistance contact Customer Service at:

- Email: fisitmrequest@fujifilm.com
- Direct line: +1 800 577 6097

WEBSITE RESOURCES

Visit the website at www.irvinesci.com for technical resources and information including:

- Safety Data Sheets (SDS)
- Certificate of Analysis (CoA) (when available)
- FAQs
- Product literature
- Complete list of offices and contact information by country

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