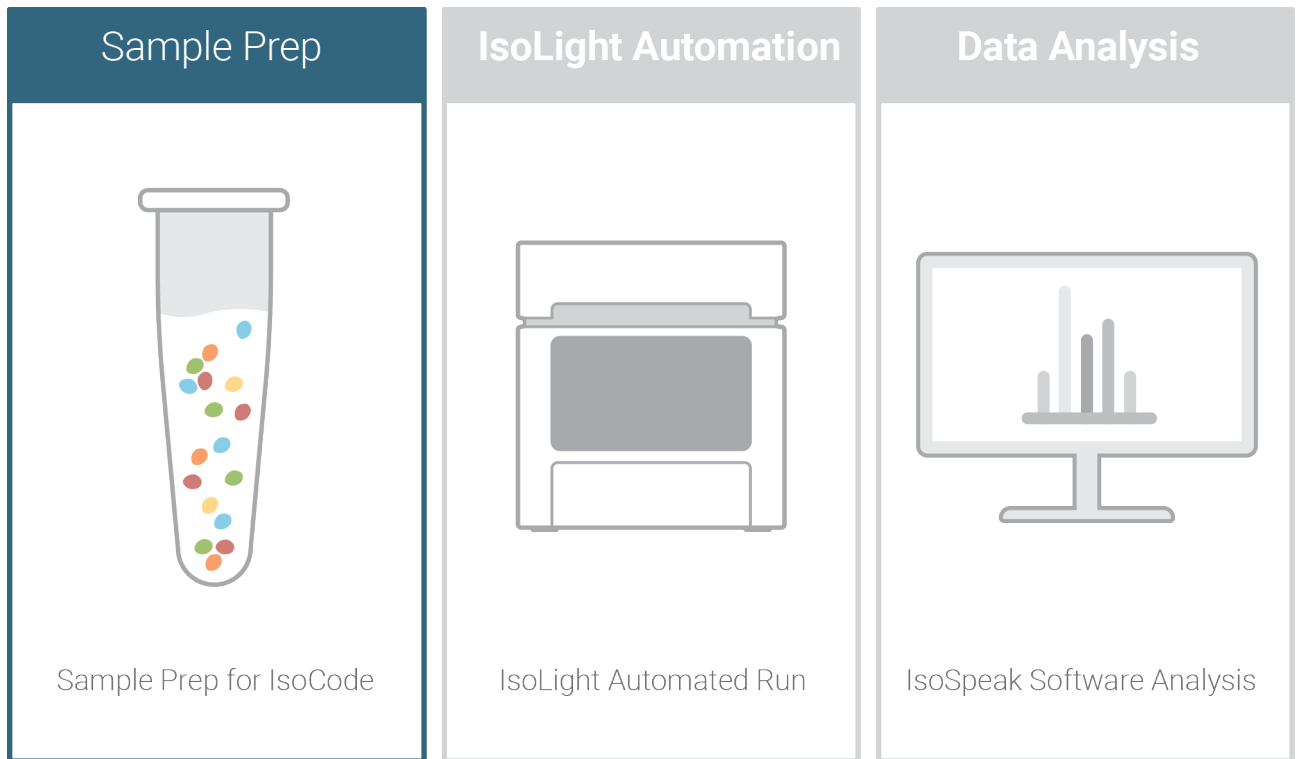


IsoCode Single-Cell Adaptive Immune: Human PBMC Protocol

Ensure you achieve the maximum benefit from our systems and generate useful data ASAP that relates to in vivo immune activity



Key: ● TIP, ● CRITICAL, ● OPTIONAL

PRO-1 REV 1.0 (Formerly 704-00021-01 REV 1.10)

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Contents

A. Overview	3
Overview of Protocol	3
Safety Warnings	4
Required Reagents, Consumables and Equipment	4-7
B. Before Getting Started	8
Important Precautions	8
Reagents to Be Prepared Before Starting	8-9
C. Protocol	10
Chapter 1: Getting Started	10
Chapter 2: Coat Culture Plates with anti-CD3	10-11
Chapter 3: Recovery of Cryopreserved Cells	11-12
Chapter 4: Pre-Sample Enrichment	12-13
Chapter 5: CD8 Sample Enrichment	13-15
Chapter 6: CD4 Sample Enrichment	16-18
Chapter 7: Cell Stimulation	18-19
Chapter 8: Chip Thaw	19
Chapter 9: Cell Staining	20-24
Chapter 10: Chip Loading	24
D. Appendix	25
D1 Protocol: Cell Quantification & Viability	26
D2 Protocol: Dead Cell Removal Using Ficoll	27
Troubleshooting and References	27-28

A. Overview

Overview of Protocol

Day 1: Cryopreserved cells are thawed and cultured overnight in the presence of IL-2.

Day 2: **Enrichment** and **Stimulation** of CD4⁺ and/or CD8⁺ T cells for 24 hours.

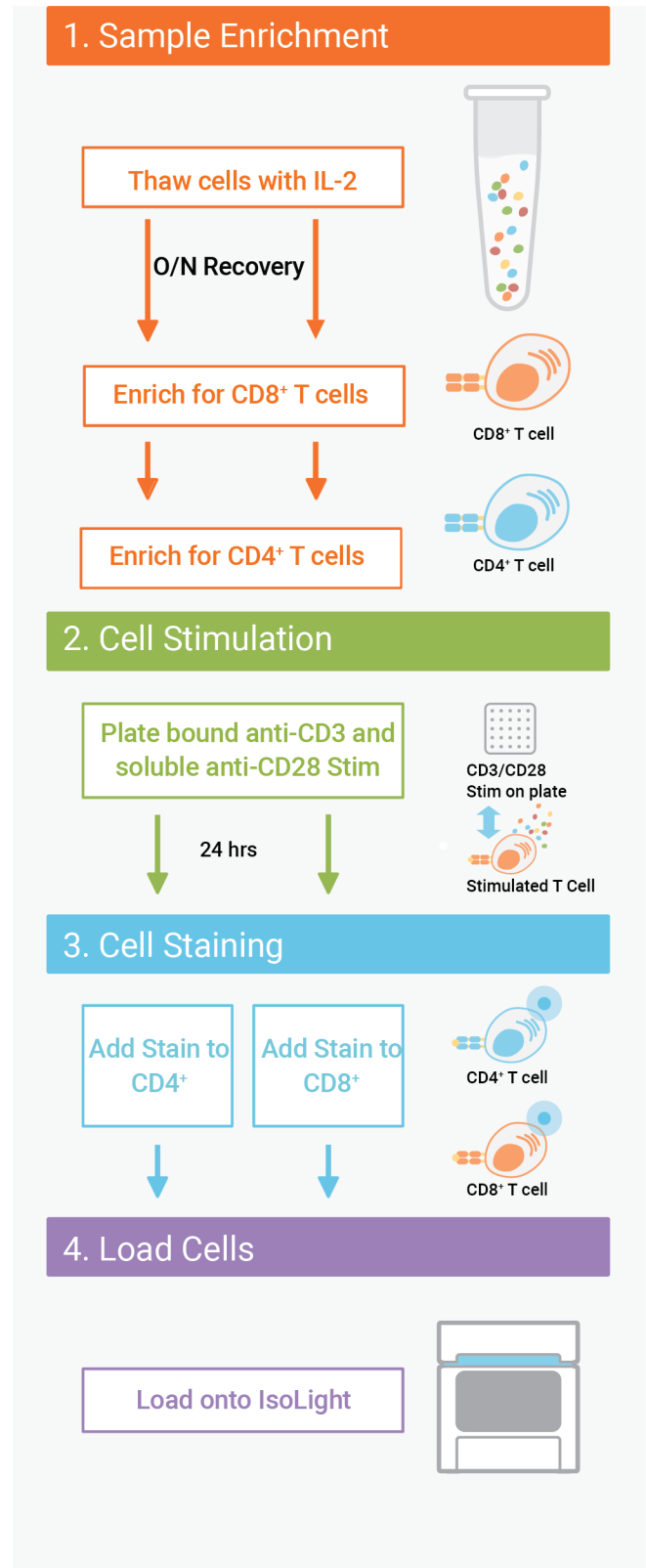
Day 3: **Staining** and **Loading** of T cells onto IsoCode chip

NOTE:

This protocol outlines the standard method for thawing and culturing of human T cells only and may not be valid for other species or cell types.

NOTE:

Using stains and protocols other than recommended stain and protocols might result in failed runs. Stains and staining procedures not approved by IsoPlexis will require validation prior to use. Please consider IsoPlexis' IsoPACE™ program to assist in custom marker and protocol validation.



Safety Warnings

- Read MSDS documents of all materials prior to use.
- Laboratory workers should wear standard PPE including disposable gloves, protective eyewear, and laboratory coats.

Required Reagents, Consumables and Equipment

Table 1: Required Reagents and Consumables Provided by IsoPlexis

Item	Catalog Number	Quantity	Comment
IsoCode Kit	Please see website (https://isoplexis.com/) for available kits or talk to IsoPlexis' Customer Service team for details	One chip per sample/cell type/condition	Subcomponents stored at 4°C and -20°C

IsoCode Kit Components

IsoCode Reagent Box (4°C)

- 15 mL Tube A
- 15 mL Tube B
- 1.5 mL Tubes A/B: Cocktail A in Micro-Tube (Green Cap) and Cocktail B in Micro-Tube (Red Cap)
- 50 mL Tubes containing Reagents 1, 2, 3, 4, 5, 6, 7, 8
- 1 Bag of Disposable Reagent Sippers
- Allophycocyanin anti-human CD45RO stain (**APC-CD45RO**) [ordered separately]
- Alexa Fluor 647 anti-human CD4 stain (**AF647-CD4**) [ordered separately]
- Alexa Fluor 647 anti-human CD8 stain (**AF647-CD8**) [ordered separately]

IsoCode Chip Set (-20°C)

- Boxes of IsoCode Chips (2 per box) to make up 4, 6, or 8 chips
- Membrane Stain [ordered separately]
- Membrane Stain Diluent (DMSO) [part of membrane stain kit]

Table 2: Required Consumables Not Supplied by IsoPlexis

Consumable	Type	Source	Catalog Number
T25 Flask	N/A	Corning	353108
T75 Flask	N/A	Corning	430641U
96 Well Plate Flat-Bottom	N/A	Corning	353072
MACS LS Column	N/A	Miltenyi	130-042-401
Centrifuge Tubes	Polypropylene, 15 mL	VWR	CA62406-200
Lo-Bind Microcentrifuge Tubes, Sterile	1.5 mL	USA Scientific	4043-1081
Pipette Tips (Filtered)	10 µL Graduated Filter Tips 100 µL Graduated Filter Tips 1000 µL XL Graduated Filter Tips	USA Scientific	1181-3710 1183-1740 1182-1730
Serological Pipette	2 mL Pipette 5 mL Pipette 10 mL Pipette	USA Scientific	1072-0510 1075-0110 1071-0810
Syringe with BD Luer-Lok Tip	10 mL	VWR	75846-756
0.2 µm Syringe Filter with Acrylic Housing	N/A	VWR	28145-501
Fisherbrand™ Disposable PES Filter Units (0.20 µm)	500 mL	Fisher Scientific	FB12566504

Table 3: Required* Reagents Not Supplied by IsoPlexis

Consumable	Stock Concentration	Source	Catalog Number
RPMI	1x	Fisher	MT10040CV
Penicillin-Streptomycin-Neomycin Solution Stabilized	100x	Sigma	P4083-100mL
Glutamax	100x	Thermo	35050061
FBS	1x	Sigma	F2442-6X500mL
Recombinant IL-2	200 µg/mL	Biolegend	589104
Bovine Serum Albumin (BSA), lyophilized powder	N/A	Sigma-Aldrich	A9647-10G
Phosphate Buffered Saline (1XPBS) without Calcium or Magnesium	1x	Invitrogen	20012-027
RoboSep buffer	1x	StemCell Tech	20104
Ficoll Paque Plus	N/A	GE Healthcare	17-1440-03
Miltenyi CD8 Microbeads, Human, 2mL	N/A	Miltenyi	130-045-201
Miltenyi CD4 Microbeads, Human, 2mL	N/A	Miltenyi	130-045-101
Trypan Blue	0.4%	Gibco	15250-061
CD3 Monoclonal Antibody (OKT3), Functional Grade	1 mg/mL	ThermoFisher/Invitrogen	16-0037-85
CD28 Monoclonal Antibody (CD28.2), Functional Grade	1 mg/mL	ThermoFisher/Invitrogen	16-0289-81
Reagent Alcohol 70%	N/A	VWR	BDH1164-4LP

*Reagents have been validated by IsoPlexis and no alternatives may be used.

Table 4: Cell Staining Reagents

Test Material	Catalog Number	Color
Membrane stain	STAIN-1001-1	Violet
Allophycocyanin anti-human CD45RO (APC-CD45RO)	STAIN-1004-1*	Red
Alexa Fluor 647 anti-human CD4 (AF647-CD4)	STAIN-1002-1	Red
Alexa Fluor 647 anti-human CD8 (AF647-CD8)	STAIN-1003-1	Red

*STAIN-1004-1 kit contains both APC-CD45RO stain and membrane stain.

Table 5: Required Equipment

Equipment	Source	Catalog Number
IsoLight Instrument	IsoPlexis	ISOLIGHT-1000-1
Culture Hemocytometer	(Fisher) Hauser Levy	02-671-55A
Hemocytometer Cover Glass	(Fisher) Hauser Levy	02-671-53
MidiMACS Separator	Miltenyi	130-042-302
MACS MultiStand	Miltenyi	130-042-303

Table 6: General Equipment

Equipment	Requirements
Pipette	P10, P100, P200, P1000
Pipettor	Ability to pipette between 1 and 10 mL
Incubator	37°C, 5% CO ₂
Tabletop Centrifuge	Temperature controlled*; swinging bucket rotor; ability to centrifuge 15 mL conical tubes
Microcentrifuge	Temperature controlled*; fixed rotor; ability to centrifuge 1.5 mL microcentrifuge tubes
Mini centrifuge	Ability to spin micro sample sizes
Water Bath	Ability to heat to 37°C
Microscope	Inverted light microscope with 10x and 20x objectives

* Temperature controlled centrifuges are required so that centrifuging steps can be conducted at room temperature without risk of overheating. Temperature on centrifuges should be set to 21°C.

B. Before Getting Started

1. Important Precautions

Read MSDS documents of all materials prior to use.

Working with Biohazardous Reagents

Please refer to your institute's guidelines and obtain proper training to handle potentially biohazardous samples. It is also strongly recommended that any lab personnel handling human samples should be vaccinated against HBV if the individual does not have sufficient HBV antibody titer.

Additional precautions need to be taken when working with samples that potentially contain an EID agent:

1. Laboratory workers should wear standard PPE including disposable gloves, protective eyewear, and laboratory coats.
2. Any procedure or process that cannot be conducted in the designated EID BSC should be performed while wearing gloves, gown, goggles and a fit tested N-95 mask.
3. Work surfaces should be decontaminated on completion of work with appropriate disinfectants. This includes any surface that potentially comes in contact with the specimen (centrifuge, microscope, etc.).
4. All liquid waste produced in the processes must be treated to a final concentration of 10% bleach prior to disposal.

2. Reagents to Be Prepared Before Starting

Table 7: Complete RPMI Recipe

- **CRITICAL:** Complete RPMI media has been validated for use by IsoPlexis. Using alternative media may result in failed runs. Please contact FAS for additional information.

Ingredient	Stock Concentration	Final Concentration	Amount for 500 mL	Vendor/Catalog
Penicillin-Streptomycin- Neomycin Solution Stabilized	100x	1x	5 mL	Sigma P4083-100mL
Glutamax	100x	1x	5 mL	Thermo/35050061
FBS	1x	10%	50 mL	Sigma/F2442-6X500 mL
RPMI	1x	1x	440 mL	Fisher/MT10040CV

Note | Sterile-filter through 0.20 μ m filter before use. Store complete RPMI Media at 4°C and warm up to 37°C in water bath prior to use.

Table 8: 1% BSA Recipe

Ingredient	Stock Concentration	Final Concentration	Amount for 100 mL	Vendor/Catalog
Bovine Serum Albumin (BSA), lyophilized powder	N/A	1%	1 g	Sigma-Aldrich/A9647-10G
Phosphate Buffered Saline (1XPBS) without Calcium or Magnesium	1X	1X	99 mL initially*	Invitrogen/20012-027

*Rotate solution until BSA powder is dissolved and then bring final volume up to 100 mL with 1X PBS.

Table 9: Working Stock of Recombinant IL-2 (1 µg/mL) Recipe

Ingredient	Stock Concentration	Final Concentration	Amount for 10 mL	Vendor/Catalog
1% BSA in 1X PBS (sterile filtered)	1%	1%	9.95 mL	Table
Recombinant IL-2	200 µg/mL	1 µg/mL	50 µL	BioLegend/589104

Note | Sterile-filter through a 0.2 µm PES filter before use.

- **CRITICAL:** Prepare 200 µL IL-2 aliquots and freeze at -20°C for no longer than 1 month. Aliquots are single use only and are to be thawed immediately prior to their usage. If there is any remaining volume in an aliquot, do not refreeze but discard.

C. Protocol

Chapter 1: Getting Started

Kit Contents

IsoCode Reagent Box (4°C)

15 mL Tube A

15 mL Tube B

1.5 mL Tube A/B: Cocktail A in Micro-Tube (Green Cap) and Cocktail B in Micro-Tube (Red Cap)

50 mL Tubes Containing Reagents 1, 2, 3, 4, 5, 6, 7, 8

1 Bag of Disposable Reagent Sippers

Allophycocyanin anti-human CD45RO stain (**APC-CD45RO**) [ordered separately]

Alexa Fluor 647 anti-human CD4 stain (**AF647-CD4**) [ordered separately]

Alexa Fluor 647 anti-human CD8 stain (**AF647-CD8**) [ordered separately]

IsoCode Chip Set (-20°C)

Boxes of IsoCode Chips (2 Per Box) to Make Up 4, 6, or 8 Chips

Membrane stain [ordered separately]

Membrane stain diluent (DMSO) [part of Membrane stain kit]

Chapter 2: Coat Culture Plates with anti-CD3

Materials Required

Sterile 1X PBS (room temperature)
Anti-CD3 (4°C) 10 µg/mL
15 mL Centrifuge Tube (<i>anti-CD3 Dilution</i>)
96 Well Plate Flat Bottom (<i>PBMC, Date, Time</i>)

All the following steps should take place in a sterile tissue culture hood.

Methods

- 1. Vortex the anti-CD3 tube at a slow speed for 10 seconds. **TIP: Ensure contents are well suspended.**
- 2. Spin anti-CD3 in mini centrifuge for 10 seconds. **TIP: Ensure that contents are all in the bottom of the vial.**
- 3. Preparation of anti-CD3 dilution in 1X PBS to a final concentration of 10 µg/mL in a 15 mL centrifuge tube:
 - a. Dilute 15 µL of 1 mg/mL anti-CD3 into 1.5 mL of 1X PBS.
 - b. Use 1 mL pipette to mix.
- 4. Pipette 100 µL of anti-CD3 dilution to each well of a 96 well flat bottom plate.
- **CRITICAL: Be careful not to create air bubbles.**

5. Incubate covered plate at 4°C overnight.

Chapter 3: Recovery of Cryopreserved Cells

Materials Required

Complete RPMI (37°C)
Recombinant IL-2 at 1 µg/mL (-20°C)
Cryopreserved PBMC
15 mL Centrifuge Tube
Plate and/or Flask
For > 10 M cells, T75 Flask
For 6 - 9.9 M cells, T25 Flask
For < 6 M, 6 Well Plate

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. Pipette 5 mL of complete RPMI into a 15 mL centrifuge tube, labeled *Thawed PBMC*.
- 2. Using proper PPE, remove cells from liquid nitrogen storage and thaw cells. **TIP: Be careful of contamination.**
3. Quickly move vials into a water bath (37°C) to thaw. While thawing, swirl the vial in the water until a single ice crystal remains in the vial. Be sure to prevent (to the best of your ability) any of the water from the water bath from getting under the cap and into the sample.
4. When the sample is nearly thawed, remove the vial and immediately spray vial with 70% alcohol before bringing into the hood. It is important to allow the alcohol to evaporate before opening the vial.
- 5. Slowly pipette thawed cells into 5 mL of complete RPMI in 15 mL centrifuge tube, labeled *Thawed PBMC*. **TIP: Insert tip into complete RPMI when pipetting, be careful to not create bubbles.**
6. Take 1 mL of the cell/complete RPMI mixture.
- 7. Pipette into original thawed cell vial, rinse inside the vial with the complete RPMI to recover additional thawed cells. **TIP: Insert tip into complete RPMI, be careful to not create bubbles.**
- 8. Draw up cell/complete RPMI mixture and pipette back into the 15 mL centrifuge tube. **TIP: Insert tip into complete RPMI and pipette gently up and down. Be careful to not create bubbles.**
9. Centrifuge cells for 10 minutes at 300 rcf.
10. While the cells are centrifuging, take the IL-2 (1 µg/mL) out from -20°C and thaw at room temperature.
- **CRITICAL: Use IL-2 aliquot that has been frozen at -20°C for less than a month. Do not use IL-2 that has been previously thawed.**
11. After cells are centrifuged, check for cell pellet.
- 12. Aspirate supernatant. **TIP: Be careful not to aspirate cell pellet.**

- a. Use pipette to remove last bit of supernatant.
13. Resuspend cell pellet in 1 mL of fresh complete RPMI.
 - a. Mix well to resuspend. **TIP: Make sure to pipette around the tube to ensure there are no clumps or bubbles.**
14. Slowly add additional complete RPMI to a final concentration of 1×10^6 cells/mL.
15. Mix thawed IL-2 thoroughly by carefully pipetting up and down.
16. Dilute 100 μ L of 1 μ g/mL IL-2 per 10 mL of cell suspension to a final concentration of 10 ng/mL IL-2.
- **CRITICAL: Discard thawed IL-2 aliquot if there is any volume remaining. IL-2 must only be thawed once.**
- 17. Mix with serological pipette. **TIP: Gently pipet up and down 3-5 times, be careful to not create bubbles.**
- 18. Transfer cell suspension to flask. For 3-5 mL of cell volume, transfer to a T25 flask and for 8-10 mL of cell volume, transfer to a T75 flask. **TIP: Slowly pipette down the side of the flask as to not create bubbles.**
- 19. Spread out cell suspension by rocking flask carefully to fully cover the bottom of the flask. **TIP: Be careful to not make bubbles.**
20. Move to incubator for overnight recovery at 37°C, 5% CO₂.

Chapter 4: Pre-Sample Enrichment

Materials Required

Complete RPMI (37°C)
15 mL Centrifuge Tube
Overnight Recovered Cells from Chapter 3 **or**
Fresh PBMC if Working with Fresh Samples
Lo-Bind Microcentrifuge Tube for Cell Count

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. Transfer cells from flask into 15 mL centrifuge tube.
- 2. Add complete RPMI to flask and rinse 5 times. **TIP: Make sure to spread out the complete RPMI to gather maximum number of cells.**
 - For T75 Flask add 3 mL
 - For T25 Flask add 2 mL
 - For 6 Well Plate add 1 mL
3. Transfer cell/complete RPMI mixture to the 15 mL centrifuge tube.
- 4. Mix well 5 times with 10 mL serological pipette. **TIP: Be careful not to create bubbles.**

- 5. Take a 10 μ L aliquot of your cells and transfer to a Lo-Bind Microcentrifuge Tube for cell counting.
CRITICAL: See Appendix D1 for cell counting instructions.
- 6. Centrifuge cells for 10 minutes at 300 rcf. While cells are centrifuging, use hemocytometer to count cells.
CRITICAL: See Appendix D1 for cell counting instructions.
- **CRITICAL: If cells are less than 80% viable, proceed to Appendix D2 Dead Cell Depletion Protocol using Ficoll.**
- 7. Proceed immediately to next chapter.

Chapter 5: CD8 Sample Enrichment

Materials Required

Complete RPMI (37°C)
 RoboSep Buffer (4°C)
 Miltenyi CD8 Microbeads, Human, 2 mL (4°C)
 MACS LS Column
 Prepared Cells from Chapter 4
 Enrichment Kit:
 MACS Metal Plate/Magnet Kit
 3 x 15 mL Centrifuge Tubes (*Discard, Flow Through, CD8 fraction*)
 Lo-Bind Microcentrifuge Tube for Post-Enrichment CD8

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. Remove the centrifuged cells and check for cell pellet.
- 2. Aspirate supernatant. **TIP: Be careful not to aspirate the cells.**
 - a. Use pipette to aspirate remaining supernatant.
- **CRITICAL: For every 1×10^7 cells, resuspend in 80 μ L RoboSep (4°C) and 20 μ L of CD8 beads (4°C).**
3. Add 80 μ L of cold RoboSep to 15 mL centrifuge tube containing 1×10^7 or fewer cells.
4. Vortex the Miltenyi CD8 Microbeads at a slow speed for 10 seconds.
5. Add 20 μ L of Miltenyi CD8 Microbeads and mix well by gently pipetting up and down 5 times.
- **TIP: Make sure to eliminate clumps so that beads are evenly distributed among cells. Be careful not to create bubbles.**
6. Incubate in refrigerator (4°C) for 15 minutes.
- 7. After 15 minutes, add 2 mL of cold RoboSep. **TIP: Not necessary to mix for this step.**
8. Centrifuge cells for 10 minutes at 300 rcf.
- **TIP: Keep RoboSep in refrigerator during enrichment process.**

- 9. Set up MACS sorting by setting metal plate in tissue culture hood and placing magnet on metal plate. Place LS column in magnet with wings facing out and align the 15 mL centrifuge tube labeled “Discard” under the LS column. **CRITICAL: LS Column should not touch the tubes.**
- 10. After cells are centrifuged, check for cell pellet and continue with MACS separation.
- 11. Aspirate RoboSep from cell pellet. **TIP: Since it is a small volume, use pipette for this step to prevent accidental aspiration of the cell pellet.**
- 12. For 1×10^8 or fewer cells, resuspend with 500 μ L of cold RoboSep.
 - a. Mix well to resuspend by gently pipetting up and down 5 times. **TIP: Make sure to pipette around the tube to ensure there are no clumps or bubbles.**
 - **CRITICAL: Be careful not to let column dry out. Do not add liquid when there is already liquid in the LS Column.**
- 13. Start with the LS column over the “Discard” tube, add 3 mL of cold RoboSep to LS Column. **CRITICAL: Be careful not to create bubbles or touch sides of LS column. Let all the RoboSep flow through before moving on to the next step. As a reminder, be careful to not let the column dry out.**
- 14. Unscrew and keep cap for “Flow Through” tube. **NOTE: This is in preparation for next step to ensure the column does not dry out during the transition.**
- 15. When the last drop falls through to the “Discard” tube, move the rack over so the LS column is over the “Flow Through” tube. **CRITICAL: Be careful not to let column dry. If there is one drop remaining that will not fall, move on to the next step.**
- 16. Increase volume of pipette to 800 μ L to ensure all 500 μ L of the cell suspension is drawn up.
- 17. Mix cell suspension by gently pipetting up and down 5 times. **NOTE: This ensures that the cells are evenly dispersed after sitting.**
- 18. Draw up all 500 μ L of cell suspension and pipette carefully into the center of the LS column without touching sides of the column.
- 19. Wash 3 times with 3 mL of cold RoboSep.
 - a. First wash: Rinse inside walls of cell suspension tube with 3 mL of cold RoboSep before transferring the mixture to LS Column. **NOTE: This is to retrieve any cells that have been left behind.**
 - i. Pipette all the mixture into LS Column after last drop passes through or does not fall from step 18. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
 - b. Second wash: Add 3 mL of RoboSep into LS Column after last drop passes through or does not fall. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
 - c. Third wash: Add 3 mL of RoboSep into LS Column after last drop passes through or does not fall. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
- 20. After the last drop of the third wash passes through or does not fall, remove the LS Column carefully from the magnet, and place carefully on the tube labeled for “CD8 fraction.”
- 21. Cap the “Flow Through” tube, this will be used for the CD4 enrichment. **Do not discard.**

- 22. Add 5 mL of cold RoboSep to the LS column. **CRITICAL: Be careful not to touch the sides.**
- 23. Take plunger, smoothly push down on the plunger to push the RoboSep buffer through the LS Column. **TIP: Lift up at the end of the plunging action so that the liquid does not splash back onto the LS Column tip.**
- 24. Set LS Column back on the “CD8 fraction” tube.
- **CRITICAL: Do not allow the plunger to interact with external contaminants. It will be used for one more step.**
- 25. Loosen up plunger. Remove plunger briefly from column and hold in one hand.
- 26. Add another 4 mL of cold RoboSep to the LS Column.
- 27. Take plunger, smoothly push down on the plunger to push the RoboSep buffer through the LS Column. **TIP: Lift up at the end of the plunging action so that the liquid does not splash back onto the LS Column tip.**
- 28. Discard LS Column and plunger.
- 29. Centrifuge “CD8 fraction” tube and “Flow Through” (CD8 Depleted PBMC Fraction) tube for 10 minutes at 300 rcf.
- 30. After cells are centrifuged, check for cell pellets.
- 31. Aspirate RoboSep buffer from “CD8 fraction” and “Flow Through” (CD8 Depleted PBMC Fraction) tubes. **TIP: Be careful to not aspirate cell pellet.**
- 32. Use pipette to aspirate the remaining supernatant from each tube. **TIP: Be careful to not aspirate cell pellet.**
- 33. Add 1 mL complete RPMI to “CD8 fraction” and resuspend cell pellet. **TIP: Make sure there are no clumps or bubbles.**
- 34. Add an additional 1 mL of complete RPMI and mix thoroughly by gently pipetting up and down 5 times. **TIP: Make sure there are no clumps or bubbles.**
- 35. Aliquot 10 μ L of the “CD8 fraction” into a Lo-Bind Microcentrifuge tube and proceed to cell count. **CRITICAL: See Appendix D1 for cell counting instructions.**
- 36. Move “CD8 fraction” tube to incubator until Cell Stimulation (Chapter 7).

Chapter 6: CD4 Sample Enrichment

Materials Required

Complete RPMI (37°C)
 RoboSep Buffer (4°C)
 Miltenyi CD4 Microbeads, Human, 2 mL (4°C)
 MACS LS Column
 CD8 Depleted Fraction or Overnight Recovered Cells from Chapter 4 if CD8 Depletion is Not Necessary
 Enrichment Kit:
 MACS Metal Plate/Magnet Kit
 3 x 15 mL Centrifuge Tubes (*Discard, Flow Through, CD4 fraction*)
 Lo-Bind Microcentrifuge Tube for Post-Enrichment CD4

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

- **CRITICAL: For every 1×10^7 cells, resuspend in 80 μ L RoboSep (4°C) and 20 μ L of CD4 beads (4°C).**
 1. Add 80 μ L of cold RoboSep to 15 mL centrifuge tube containing "CD8 depleted fraction" of 1×10^7 cells or fewer.
 2. Vortex the Miltenyi CD4 Microbeads at a slow speed for 10 seconds.
 3. Add 20 μ L of Miltenyi CD4 Microbeads and mix well by gently pipetting up and down 5 times.
- **TIP: Make sure to eliminate clumps so that beads are evenly distributed among cells. Be careful not to create bubbles.**
 4. Incubate in refrigerator (4°C) for 15 minutes.
- 5. After 15 minutes, add 2 mL of cold RoboSep. **TIP: Not necessary to mix for this step.**
- 6. Centrifuge cells for 10 minutes at 300 rcf.
- **TIP: Keep RoboSep in refrigerator during enrichment process.**
- 7. Set up MACS sorting by setting metal plate in tissue culture hood and placing magnet on metal plate. Place LS column in magnet with wings facing out and align the 15 mL centrifuge tube labeled "Discard" under the LS column. **CRITICAL: LS Column should not touch the tubes.**
- 8. After cells are centrifuged, check for cell pellet and continue with MACS separation.
- 9. Aspirate RoboSep from cell pellet. **TIP: Since it is a small volume, use pipette for this step to prevent accidental aspiration of the cell pellet.**
- 10. For 1×10^8 or fewer cells, resuspend with 500 μ L of cold RoboSep.
 - a. Mix well to resuspend by gently pipetting up and down 5 times. **TIP: Make sure to pipette around the tube to ensure there are no clumps or bubbles.**

- **CRITICAL: Be careful not to let column dry out. Do not add liquid when there is already liquid in the LS Column.**
- 11. Start with the LS column over the "Discard" tube, add 3 mL of cold RoboSep to LS Column. **CRITICAL: Be careful not to create bubbles or touch sides of LS column. Let all the RoboSep flow through before moving on to next step. As a reminder, be careful to not let the column dry out.**
- 12. Unscrew and keep cap for "Flow Through" tube. **NOTE: This is in preparation for next step to ensure the column does not dry out during the transition.**
- 13. When last drop falls through to "Discard" tube, move the rack over so the LS column is over the "Flow Through" tube. **CRITICAL: Be careful not to let column dry. If there is one drop remaining that will not fall, move on to the next step.**
- 14. Increase volume of pipette to 800 μ L to ensure all 500 μ L of the cell suspension is drawn up.
- 15. Mix cell suspension by gently pipetting up and down 5 times. **NOTE: This ensures that the cells are evenly dispersed after sitting.**
- 16. Draw up all 500 μ L of cell suspension and pipette carefully into the center of the LS column without touching sides of the column.
- 17. Wash 3 times with 3 mL of cold RoboSep.
 - a. First wash: Rinse inside walls of cell suspension tube with 3 mL of RoboSep before transferring the mixture to LS Column. **NOTE: This is to retrieve any cells that have been left behind.**
 - i. Pipette all of the mixture into LS Column after last drop passes through or does not fall from step 16. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
 - b. Second wash: Add 3 mL of RoboSep into LS Column after last drop passes through or does not fall. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
 - c. Third wash: Add 3 mL of RoboSep into LS Column after last drop passes through or does not fall. **CRITICAL: Be careful not to let LS Column dry out or allow pipette to touch sides.**
- 18. After the last drop of the third wash passes through or does not fall, remove the LS Column carefully from the magnet, and place carefully on the tube labeled for "CD4 fraction."
- 19. Cap the "Flow Through" tube and discard.
- 20. Add 5 mL of cold RoboSep to the LS column. **CRITICAL: Be careful not to touch the sides.**
- 21. Take plunger, smoothly push down on the plunger to push the RoboSep buffer through the LS Column. **TIP: Lift up at the end of the plunging action so that the liquid does not splash back onto the LS Column tip.**
- 22. Set LS Column back on the "CD4 fraction" tube.
- 23. Loosen up plunger. Remove plunger briefly from column and hold in one hand.
- **CRITICAL: Do not allow the plunger to interact with external contaminants. It will be used for one more step.**
- 24. Add another 4 mL of cold RoboSep to the LS Column.
- 25. Take plunger, smoothly push down on the plunger to push the RoboSep buffer through the LS Column. **TIP: Lift up at the end of the plunging action so that the liquid does not splash back onto the LS Column tip.**

26. Discard LS Column and plunger.
27. Centrifuge “CD4 fraction” tube for 10 minutes at 300 rcf.
28. After cells are centrifuged, check for cell pellet.
- 29. Aspirate RoboSep buffer from “CD4 fraction” tube. **TIP: Be careful to not aspirate cell pellet.**
- 30. Use pipette to aspirate the remaining supernatant from each tube. **TIP: Be careful to not aspirate cell pellet.**
- 31. Add 1 mL complete RPMI to “CD4 fraction” and resuspend cell pellet. **TIP: Make sure there are no clumps or bubbles.**
- 32. Add an additional 1 mL of complete RPMI and mix thoroughly by gently pipetting up and down 5 times. **TIP: Make sure there are no clumps or bubbles.**
- 33. Aliquot 10 μ L of the CD4 fraction into a Lo-Bind Microcentrifuge tube and proceed to cell count. **CRITICAL: See Appendix D1 for cell counting instructions.**
34. Move “CD4 fraction” tube to incubator until Cell Stimulation. (Chapter 7).

Chapter 7: Cell Stimulation

Materials Required

Complete RPMI (37°C) Anti-CD28 (4°C) Sterile 1X PBS (room temperature) Prepared anti-CD3 Coated 96 Well Plate from Chapter 2 15 mL Centrifuge Tube (<i>CD8 in complete RPMI</i>) Incubated CD8 Tube in complete RPMI Incubated CD4 Tube in complete RPMI 2 x Lo-Bind Microcentrifuge Tubes for Cell Count (label <i>CD8</i> , <i>CD4</i>)
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All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

- 1. Vortex anti-CD28 at a slow speed for 10 seconds. **TIP: Ensure contents are well suspended.**
- 2. Spin anti-CD28 in a mini centrifuge for 10 seconds. **TIP: Ensure that contents are all in the bottom of the vial.**
- 3. Prepare anti-CD28 and complete RPMI mixture by supplementing complete RPMI with 5 μ g/mL of anti-CD28. **CRITICAL: Volume is dependent on number of cells.**
 - a. Dilute 25 μ L of 1 mg/mL anti-CD28 into 5 mL of complete RPMI.
 - b. Use serological pipette to mix thoroughly.
4. Take CD8 and CD4 cells from incubator.

5. Centrifuge CD8 and CD4 cells for 10 minutes at 300 rcf.
6. After cells are centrifuged, check for cell pellet.
- 7. Aspirate supernatant with pipette. **TIP: Make sure to use a manual pipette to prevent accidental aspiration of cell pellet.**
8. Use pipette to mix the anti-CD28 and complete RPMI mixture to ensure it is evenly distributed.
- 9. Using the anti-CD28/complete RPMI mixture from step 8, resuspend CD8 cells to a cell concentration of 1×10^6 cells/mL. **TIP: Resuspend as thoroughly as possible, but gently.**
10. Repeat step using CD4 cells.
11. Prepare Stimulation Plate.
 - a. Remove previously prepared anti-CD3-coated 96 well plate from refrigerator and aspirate the 1X PBS. **TIP: Aspirate from the edge of well.**
 - b. Add 100 μ L of 1X PBS to each well to rinse.
 - c. Aspirate 1X PBS from each well. **TIP: Aspirate from the edge of the well**
- 12. Mix CD8 cells by pipetting up and down gently 5 times. Add 100 μ L of cell suspension to coated wells on plate. **CRITICAL: Be careful not to create bubbles. This will maximize even stimulation of cell suspension.**
13. Repeat step using CD4 cells.
14. Incubate plate for 24 hours at 37°C, 5% CO₂.

Chapter 8: Chip Thaw

Materials Required

IsoCode Chips in Vacuum Sealed Bag (-20°C)

Methods

- 1. Take vacuum sealed bag containing IsoCode chips from -20°C. **CRITICAL: Chips must stay sealed until Chip Loading (Chapter 10).**
2. Place on bench to thaw at ambient temperature 30 - 60 minutes prior to use.
3. While chips and samples thaw, prepare liquid reagents and attach all reagent tubes to IsoLight. Refer to the IsoLight System Guide for detailed instructions.

Chapter 9: Cell Staining

NOTE: Please read before proceeding with cell staining.

There are 3 options for cell staining depending on your experimental design:

1. Membrane staining (violet) is described in Chapter 9a.
2. Dual staining with membrane stain (violet) and CD45RO-specific surface marker stain (red) is achieved by following Chapter 9a and Chapter 9b sequentially.
3. CD4 and CD8- specific surface marker staining is described in Chapter 9c.

Cell Staining Reagents

Test Material	Catalog Number	Color
Membrane stain	STAIN-1001-1	Violet
Allophycocyanin anti-human CD45RO (APC-CD45RO)	STAIN-1004-1	Red
Alexa Fluor 647 anti-human CD4 (AF647-CD4)	STAIN-1002-1	Red
Alexa Fluor 647 anti-human CD8 (AF647-CD8)	STAIN-1003-1	Red

Chapter 9a: Cell Membrane Staining (Violet)

Materials Required

Anti-CD3/CD28 Stimulated CD8 & CD4 Cells in 96 Well Plate from Chapter 7
 2 x Lo-Bind Microcentrifuge Tubes (CD8, CD4)
 Sterile 1X PBS (room temperature)
 Complete RPMI (37°C)
 Membrane Stain (-20°C)
 Membrane Stain Diluent (DMSO) (-20°C)

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. Prepare membrane stain stock.
 - a. Thaw tube of membrane stain diluent (DMSO) at room temperature.
 - b. Spin tubes of membrane stain and membrane stain diluent (DMSO) in a mini centrifuge for 10 seconds to collect the contents at the bottom of the tubes.
 - c. Add 20 µL of membrane stain diluent (DMSO) directly to the tube of membrane stain. Pipet up and down 15 times gently to resuspend.

● **CRITICAL: Membrane stain must be prepared fresh. Discard remaining stain – do not store.**

- 2. Prepare stain master mix by diluting 2 μL of membrane stain into 1 mL of 1X PBS in a Lo-Bind microcentrifuge tube (1:500 final dilution). With the same pipette tip, pipette up and down 10 times to ensure all membrane stain has been released. Depending on sample number and cell count, additional tubes of stain master mix may need to be prepared. **CRITICAL: Failure to follow these steps will negatively impact cell counts.**
 - a. With a P1000 set to 500 μL , gently pipette the stain master mix up and down **15 times**.
 - b. **Gently vortex** the stain master mix for **5 seconds**.
 - c. **Ensure master mix is mixed well before adding stain to cells.**
3. Remove 96 well plate with CD8 and CD4 cells from incubator.
4. Mix CD8 cells by pipetting up and down. Transfer cells to a Lo-Bind microcentrifuge tube by using P100 pipette to draw up 100 μL at a time in a gentle, circular motion until well is empty. **NOTE: Pool wells if there are replicates.**
5. Repeat Step 4 using CD4 cells.
6. Centrifuge cells for 10 minutes at 300 rcf.
7. After cells are centrifuged, check for cell pellets.
- 8. Aspirate supernatant with a pipette.* **TIP: Be careful not to aspirate the cell pellets.**

*NOTE: Supernatants may be stored at -80°C for population assay.
9. Add 1mL of PBS to dilute any remaining media and mix by pipetting up and down.
10. Centrifuge cells for 10 minutes at 300 rcf.
11. After cells are centrifuged, check for cell pellets.
- 12. Aspirate supernatant with a pipette.* **TIP: Be careful not to aspirate the cell pellets.**
- 13. For every 1×10^6 cells, add 100 μL of **well mixed** stain master mix to each cell suspension tube. **CRITICAL: Pipet mix the cells 15 times. Be careful to not create bubbles. Gently remix master mix if it has been sitting for longer than a few minutes.**
14. Incubate for 5 minutes at 37°C in the dark.
- 15. Gently pipet mix the cell suspension **15 times**. **CRITICAL: Be careful to not create bubbles.**
16. Incubate for an additional 5 minutes at 37°C in the dark.
- 17. After incubation, add 5 times the volume of complete RPMI. **CRITICAL: Pipet mix the cells 15 times. Be careful to not create bubbles.**
18. Incubate for 10 minutes at 37°C in the dark.
- 19. Take 10 μL of cells to count. Count cells using a hemocytometer and determine percent of viable cells as described in Appendix D1. **TIP: Cell counting can be done while cells are incubating.**
20. Centrifuge stained cells for 10 minutes at 300 rcf.

21. After cells are centrifuged, check for cell pellets.
- 22. Aspirate supernatant with a pipette. **TIP: Be careful not to aspirate the cell pellets.**
NOTE: If dual staining with CD45RO, immediately proceed to Chapter 9b.
23. If only staining with membrane stain, resuspend the cell pellet with complete RPMI to a cell density of 1×10^6 cells/mL. Proceed to Chapter 10.

Chapter 9b: Surface Marker-Specific Staining, APC-CD45RO (Red)

Materials Required

Cells stained with Membrane Stain from Chapter 9a
APC anti-human CD45RO (4°C)
Complete RPMI (37°C)

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. After completing steps of Chapter 9a, Steps 1-22, resuspend the cell pellet with 16 μ L complete RPMI.
2. Add 4 μ L of APC anti-human CD45RO (1:5 final dilution). Mix gently by pipetting up and down.
3. Incubate for 20 minutes at room temperature in the dark.
- 4. After incubation, add 1 mL of complete RPMI to each sample tube. **TIP: Mix gently, be careful not to create bubbles.**
- 5. Take 10 μ L of cells to count. Count cells using a hemocytometer and determine percent of viable cells as described in Appendix D1. **TIP: Cell counting can be done while cells are incubating.**
6. Centrifuge stained cells for 10 minutes at 300 rcf.
7. After cells are centrifuged, check for cell pellets.
- 8. Aspirate supernatant with a pipette. **TIP: Be careful not to aspirate the cell pellets.**
9. Resuspend the cells with complete RPMI to a cell density of 1×10^6 cells/mL. Proceed to Chapter 10.

Chapter 9c: Surface Marker-Specific Staining AF647 anti-human CD4 (Red) and AF647 anti-human CD8 (Red)

Materials Required

Anti-CD3/CD28 Stimulated CD8 & CD4 Cells in 96 Well Plate from Chapter 7
 2 x Lo-Bind Microcentrifuge Tubes (CD8, CD4)
 Sterile 1X PBS (room temperature)
 Complete RPMI (37°C)
 AF647 anti-human CD4 (4°C)
 AF647 anti-human CD8 (4°C)

All the following steps should take place in a sterile tissue culture hood. Centrifugation steps should be performed at 21°C.

Methods

1. Remove 96 well plate with CD8 and CD4 cells from incubator.
2. Mix CD8 cells by pipetting up and down. Transfer cells to a Lo-Bind microcentrifuge tube by using P100 pipette to draw up 100 µL at a time in a gentle, circular motion until well is empty. **NOTE: Pool wells if there are replicates.**
3. Repeat Step 2 using CD4 cells.
4. Centrifuge cells for 10 minutes at 300 rcf.
5. After cells are centrifuged, check for cell pellets.
- 6. Aspirate supernatant with a pipette.* **TIP: Be careful not to aspirate the cell pellets.**
 *NOTE: Supernatants may be stored at -80°C for population assay.
7. Spin tubes of AF647 anti-human CD4 and AF647 anti-human CD8 in a mini centrifuge for 10 seconds to collect stain at the bottom of the tube.
8. Resuspend cell pellet with 18 µL complete RPMI.
- 9. Add 2 µL of AF647 anti-human CD8 to CD8+ samples (1:10 final dilution). Mix gently by pipetting up and down. **CRITICAL: Be sure to use appropriate stain for each cell subset.**
- 10. Add 2 µL of AF647 anti-human CD4 to CD4+ samples. Mix gently by pipetting up and down. **CRITICAL: Be sure to use appropriate stain for each cell subset.**
11. Incubate for 20 minutes at room temperature in the dark.
- 12. After incubation, add 1 mL of complete RPMI to each sample tube. **TIP: Mix gently, be careful not to create bubbles.**
- 13. Take 10 µL of cells to count. Count cells using a hemocytometer and determine percent of viable cells as described in Appendix D1. **TIP: Cell counting can be done while cells are incubating.**
14. Centrifuge stained cells for 10 minutes at 300 rcf.

15. After cells are centrifuged, check for cell pellets.
- 16. Aspirate supernatant with a pipette. **TIP: Be careful not to aspirate the cell pellets.**
17. Resuspend the cells with complete RPMI to a cell density of 1×10^6 cells/mL. Proceed to Chapter 10.

Chapter 10: Chip Loading

Materials Required (Pre-prepared)

Pre-Thawed IsoCode Chips in Vacuum Sealed Bag from Chapter 8 Stained CD8 Cells at 1×10^6 cells/mL Stained CD4 Cells at 1×10^6 cells/mL
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Methods

- 1. Remove IsoCode chips from vacuum sealed bag and place on a flat surface. **CRITICAL: Keep protective blue film on bottom of chip.**
- 2. Resuspend CD8 fraction by gently pipetting up and down 15 times. Immediately proceed to chip loading. Pipette 30 μ L of cell suspension into IsoCode chip. **CRITICAL: Be careful not to create bubbles. Insert pipette tip vertically into inlet port until tip lightly touches bottom, and slowly pipette 30 μ L into inlet port. Be careful not to eject second step of pipette—it will cause bubbles.**
- 3. Resuspend CD4 fraction by gently pipetting up and down 15 times. Immediately proceed to chip loading. Pipette 30 μ L of cell suspension into IsoCode chip. **CRITICAL: Be careful not to create bubbles. Insert pipette tip vertically into inlet port until tip lightly touches bottom, and slowly pipette 30 μ L into inlet port. Be careful not to eject second step of pipette—it will cause bubbles.**
4. Let IsoCode chips sit for one minute on a flat surface.
- 5. Check bottom of chip to ensure liquid has entered the chip. **TIP: If liquid has not flowed, tap IsoCode chip on flat surface lightly.**
6. When inserting IsoCode chip into IsoLight, make sure barcode is facing up and towards you with the magnet facing the IsoLight. Take the blue film off while inserting each IsoCode chip into the IsoLight.

NOTE: Please refer to the loading instructions of the IsoLight instrument for details.

D. Appendix

D1 Protocol: Cell Quantification & Viability

Materials Required

Hemocytometer 10 μ L aliquot of cells Trypan Blue

NOTE: Automated cell counters can be used in this protocol EXCEPT prior to loading cells on chip due to spectral overlap of the stains. Manual cell counting is required prior to loading on the chip.

- 1. Using a P10 pipette, add equal volume of Trypan blue solution to 10 μ L of sample. Mix gently to resuspend.
TIP: Make sure to pipette around the tube to ensure there are no clumps or bubbles.
- 2. Load onto hemocytometer. **CRITICAL: Be careful not to overfill or create bubbles.**
- 3. Count and record viable (clear) and dead cells (blue) of all four 16-square corners.
- **CRITICAL: If more than 200 cells/16 squares were counted, repeat count using a 1:5 or 1:10 dilution with 1X PBS or complete RPMI using a fresh sample aliquot.**
- 4. Calculate the concentration of cells as follows:
 - a. Concentration (cells/mL) = Average per square cell count $\times 10^4$ \times dilution factor
- 5. Calculate the number of cells as follows:
 - a. Number of cells = Cell concentration (cells/mL) from D.1.3 \times total volume of cell suspension (mL)
- 6. Calculate percent viable cells:
 - a. % Viable cells = $100 \times \text{number of viable cells} / [\text{number of viable cells} + \text{number of dead cells}]$

D2 Protocol: Dead Cell Removal Using Ficoll

Materials Required

Complete RPMI (37°C) Cells (Minimum 3×10^6) 2 x 15 mL Centrifuge Tubes Ficoll Paque
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- **CRITICAL: It is recommended to start this protocol with a minimum of 3×10^6 total cells.**
- 1. Carefully add 6 mL of Ficoll to the bottom of the required number of 15 mL centrifuge tube(s) prior to harvesting stimulation cultures.
- 2. Centrifuge cells for 10 minutes at 300 rcf.
- 3. Remove cells from centrifuge, check for cell pellet.
- 4. Aspirate supernatant. **TIP: Be careful not to aspirate cell pellet.**
 - a. Use pipette to aspirate remaining supernatant.
- 5. Resuspend the pellet(s) in 7 mL of complete RPMI. **TIP: Be careful not to create bubbles.**
- **CRITICAL: Do not use more than 1×10^7 cells of your suspension per Ficoll tube.**
- 6. Add the cell suspension(s) VERY SLOWLY to the tube(s) containing Ficoll. **CRITICAL: Place the tip of your pipette on the wall of the tube, close to the Ficoll layer. Add cell suspension VERY SLOWLY.**
- **CRITICAL: This step must be done carefully and slowly to avoid mixing of the layers.**
- 7. Centrifuge tubes for 20 minutes at 300 rcf without brake or acceleration.
- **CRITICAL: Turn acceleration and brakes off to preserve the density layers established during centrifugation.**
- 8. While cells centrifuge, prepare appropriate number of 15 mL centrifuge tube(s) containing 6 mL of complete RPMI.
- 9. Remove cells from centrifuge, check for cloudy layer which are the viable cells.
- 10. Aspirate a small volume of the supernatant. **CRITICAL: Be careful not to aspirate cloudy layer containing viable cells.**
- 11. Using a P1000 pipette, collect the viable cells by recovering the cloudy layer between Ficoll and complete RPMI media
- 12. Transfer cells into the 15 mL centrifuge tube(s) containing complete RPMI.
- 13. Aliquot 10 μ L of cell/complete RPMI mixture(s) into a Lo-Bind Microcentrifuge Tube(s) and proceed to cell count. **CRITICAL: See Appendix D1 for cell counting instructions.**

Troubleshooting & References

Please contact Support at 475-221-8402 & support@isoplexis.com with specific troubleshooting questions.

Problem	Possible Reason	Solution
Low quality cell count on chip <i>Cell Counting & Concentration related</i>	<ul style="list-style-type: none"> Recommended cell concentrations not used Issue with Cell Counting procedure Trypan Blue may have debris Poor cell removal from anti-CD3 plate 	<ul style="list-style-type: none"> Use recommended cell concentrations during overnight incubation (Chapter 7) Use appropriate dilutions recommended in Appendix D1. Do a recount if initial count does not seem accurate Quick spin Trypan Blue to pellet potential debris, remove aliquot from top of Trypan Blue. Start with fresh aliquot of Trypan Blue. Thoroughly mix cells in well with pipette prior to transferring to tube (refer to Step 9a.4 and 9a.5 or Step 9c.2 and 9c.3)
Low quality cell count on chip <i>Stain Process related</i>	<ul style="list-style-type: none"> Use of media other than the recommended media in protocol which could interact with cell stain Use of stains not recommended in protocol Recommended stain concentration, incubation time and/or incubation temperature not used Membrane stain was stored prior to use. 	<ul style="list-style-type: none"> Use complete RPMI media following recipe in Table 7 Use IsoPlexis provided validated stain (Table 4: Cell Staining Reagents) Follow staining steps as highlighted in Chapter 9 Use only freshly prepared membrane stain per Chapter 9.
Low quality cell count on chip <i>Technique Detail related</i>	<ul style="list-style-type: none"> Bubbles loaded onto chip, especially at Chip Loading Detection of potential artifacts such as debris, cell clumping, inefficient enrichment possibly due to: <ul style="list-style-type: none"> Pipetting wrong concentration Not fully inserting column into MACS separator Reagents not stored at recommended temperatures Recommended number of cells not loaded on chip 	<ul style="list-style-type: none"> Follow Critical step in 10.2 and 10.3 to avoid introduction of bubbles on chip Ensure use of a sterile space to reduce introduction of potential contaminants. Use dedicated pipettes, tips, and tubes for sterile work. Pipette up and down gently and throughout protocol to reduce clumps. Follow closely the Critical steps and tips in Chapters 5 and 6 (CD8 and CD4 Sample Enrichments) Load recommended number of cells (30,000 cells per chip) (Chapter 10)
Limited frequency of stimulated cells, i.e. those with cytokine signal, on chip <i>Viability related</i>	<ul style="list-style-type: none"> Leaving thawed cells in DMSO for an extended period Low viable cells due to low viability input sample and lack of utilization of Ficoll-Paque Decreased viability due to cell shock 	<ul style="list-style-type: none"> After thaw, quickly transfer cells from DMSO to complete RPMI to ensure viability of cells. Verify viability of cells is above 80% as stated in Chapter 4 to ensure protocol is being performed with the highest quality of cells. Use Ficoll Paque in Appendix D2 if viability is less than 80% Use reagents at recommended temperatures (i.e. always use warmed media [37°C])

<p>Limited frequency of stimulated cells, i.e. those with cytokine signal, on chip <i>Stimulation step related</i></p>	<ul style="list-style-type: none"> • Recommended anti-CD3/anti-CD28 stimulation concentration was not used • Recommended anti-CD3/anti-CD28 stimulation duration was not used 	<ul style="list-style-type: none"> • Use anti-CD3/anti-CD28 concentrations listed in Chapter 2 and Chapter 7 • Use anti-CD3/anti-CD28 timing listed in Chapter 7 • Use recommended vendor as listed in Table 3
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