

Sample Preparation Guidelines for Cytek cFluor[®] Immunoprofiling Kit 14 Color RUO Kit

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Introduction

For anyone working with the Cytek cFluor[®] Immunoprofiling Kit 14 color RUO Kit to prepare and acquire whole blood or peripheral blood mononuclear cells (PBMCs), here is Cytek's recommended sample preparation instructions. If you are acquiring the samples on your Cytek Aurora or Cytek Northern Lights cytometer, here are 3 additional items to make your workflow easier:

- 1. Import the cFluor fluorescent tags to the fluorescent tag lists in your SpectroFlo[®] Library section. If you already have existing cFluor tags in your library, delete them or overwrite them with the tags in this list
- 2. Download the SpectroFlo Experiment Template for PBMCs or for whole blood and import it into SpectroFlo.
- 3. Download and read the Acquisition Protocol

Please note that this kit is designed for research use only and is not for use in diagnostic or therapeutic procedures.

Materials Needed

- PBMCs or Whole Blood collected in EDTA blood collection tubes
- ViaDye™ Red Fixable Viability Dye, Cytek Biosciences Cat: R7-60008
- Cytek Immunoprofiling Kit 14 color RUO kit, Cytek Biosciences, Cat: R7-40000
- PBS, pH 7.4, Corning, Cat: 21-040-CM
- Stain Buffer (BSA), BD Biosciences, Cat: 554657 or equivalent
- BD FACS Lysing Solution, BD Biosciences, Cat: 349202 or equivalent
- 12 mm x 75 mm, round bottom 5 mL polystyrene tubes, Corning Falcon, Cat: 352008, or equivalent
- 2 mL, V-bottom, 96 well deep well plates, Corning, Cat: 3960, or equivalent
- U-bottom, 96 well plates, Corning, Cat: 3365, or equivalent
- 4% paraformaldehyde in PBS, Santa Cruz Biotechnology, Cat: sc-281692, or equivalent
- UltraComp eBeads™ Plus Compensation Beads, Thermo Fisher Scientific, Cat: 01-3333-41

Thawing PBMCs

For the 14C kit, plan on using 5 x 10⁶ to 10 x 10⁶ cells for Reference Controls and for the Multicolor Sample.

- 1. Pre-warm ~50 mL RPMI (supplemented with 10% FBS, 1% Penicillin/Streptomycin) at 37°C for at least 30 minutes
- 2. Thaw PBMC vial quickly in 37°C water bath until the core is loose
- 3. Transfer the cells into a 50 mL conical tube
- 4. Add 1 mL of warm media to the empty cryovial. Set it aside
- 5. Drop-by-drop, slowly add 10 mL of warm media to the cells in the 50 mL conical tube while gently swirling the tube to mix

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- 6. Pour the contents of the cryovial from step (4) into the 50 mL conical tube
- 7. Add additional media to complete the final volume to 20 mL
- 8. Centrifuge at 200 x g, 8 minutes
- 9. Decant the supernatant and blot on paper towel or remove the supernatant by aspiration

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- 10. Gently resuspend the pellet in 2 mL of warm media by pipetting up and down using a serological pipet
- 11. Repeat steps (7)-(10)
- 12. Resuspend in proper volume of warm media for a final cell concentration of 2.5 x 106 cells/ml and verify by counting

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13. Loosen the cap on the 50 mL conical tube, place the cells in the cell culture incubator until ready to use

Preparing ViaDye Red Fixable Viability Dye

- 1. Completely thaw DMSO
- 2. Add 100 µL DMSO to the lyophilized ViaDye Red Fixable Viability Dye stock (=1 mM stock solution)
- 3. Vortex to mix thoroughly
- 4. Aliquot and freeze at -20°C until use
- 5. Thaw an aliquot of the stock solution at room temperature, protected from light, before each use. **NOTE:** Do not re-freeze or re-use the viability dye.
- 6. Dilute the stock solution at 1:500 in PBS (=2 μ M working solution)
- 7. Use the working solution at 5 μL per test

Protocol for Staining PBMCs in Tubes

Plan on using 2×10^5 cells for each reference control (14 fluorescence controls, 1 ViaDye Red Fixable Viability Dye control, and 1 unstained control), and 1 x 10⁶ cells for each multicolor sample.

Viability Reference Control

- 1. Label a 12 mm x 75 mm tube for the Viability Reference Control
- 2. Add 2×10^5 cells to the tube
- 3. Add PBS to complete the final volume to 3 mL
- 4. Centrifuge at 400 x g, 5 minutes at room temperature
- 5. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 6. Vortex thoroughly
- 7. Repeat steps (3)-(6) if the volume in step (2) is greater than 1 mL
- 8. Add 5 μ L of working solution ViaDye Red Fixable Viability Dye to the cell pellet
- 9. Vortex thoroughly
- 10. Incubate for 15 minutes at room temperature, protected from light
- 11. Add 3 mL of Stain Buffer
- 12. Centrifuge at 400 x g, 5 minutes at room temperature
- 13. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 14. Vortex thoroughly
- 15. Resuspend in 150 µL Stain Buffer



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16. Acquire at medium flow rate within 2 hours post staining if cells are not fixed **NOTE**: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Tubes" to fix the samples in 1% paraformaldehyde.

Single Stain Reference Controls

- 1. Label a 12 x 75 mm tube for each Single Stain Reference Control
- 2. Add 2×10^5 cells to each tube

NOTE: See Table for sample type recommendations for each marker.

- 3. Add Stain Buffer to complete the final volume to 3 mL
- 4. Centrifuge at 400 x g, 5 minutes at room temperature
- 5. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 6. Vortex thoroughly
- 7. Repeat steps (3)-(6) if the volume in step (2) is greater than 1 mL
- 8. Add 5 μL of appropriate monoclonal antibody to the cell pellet
- 9. Vortex thoroughly
- 10. Incubate for 20 minutes at room temperature, protected from light
- 11. Add 3 mL of Stain Buffer
- 12. Centrifuge at 400 x g, 5 minutes at room temperature
- 13. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 14. Vortex thoroughly

NOTE: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Tubes" to fix the samples in 1% paraformaldehyde.

- 15. Resuspend in 150 μL Stain Buffer
- 16. Acquire at medium flow rate within 2 hours post staining if cells are not fixed

Table 1. Sample Type Recommendations for optimal Reference Controls

Laser	Target	Fluorochrome	Recommended Sample Type
Violet	CD3	cFluor [®] V420	Cells or Beads
	CD14	cFluor [®] V450	Cells or Beads
	CD45	cFluor [®] V547	Cells Only
Blue	CD8	cFluor [®] B515	Cells or Beads
	CCR7	cFluor [®] BYG575	Cells or Beads
	lgD	cFluor [®] BYG667	Cells Only
	CD45RA	cFluor [®] B690	Cells Only
	CD19	cFluor [®] BYG710	Cells or Beads
	CD25	cFluor [®] BYG781	Cells or Beads
Red	CD127	cFluor [®] R659	Cells or Beads
	CD16	cFluor [®] R668	Cells Only
	CD56	cFluor [®] R720	Cells or Beads
	CD4	cFluor [®] R780	Cells or Beads
	CD27	cFluor [®] R840	Cells or Beads



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Multicolor Sample

- 1. Label a 12 x 75 mm tube for each Multicolor sample
- Prepare the antibody cocktail in a 1.5 mL tube. For one Multicolor sample, add 5 µL of all antibodies included in the kit, one-by-one, except for cFluor BYG575 CCR7 and cFluor BYG667 IgD

NOTE: cFluor BYG575 CCR7 and cFluor BYG667 IgD need to be added separately. DO NOT add these antibodies in the cocktail mix.

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NOTE: Prepare one extra test for the multicolor cocktail to account for any reagent loss in the process (ex. make multicolor cocktail for 6 tests if you have 5 multicolor samples to stain). Take 60 µL of the cocktail per multicolor sample and discard any leftover if not used within 10 days of cocktailing.

- 3. Add 1 x10⁶ cells to Multicolor Sample tube
- 4. Add PBS to complete the final volume to 3 mL
- 5. Centrifuge at 400 x g, 5 minutes at room temperature
- 6. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 7. Vortex thoroughly
- 8. Repeat steps (4)-(7) if the volume in step (3) is greater than ImL
- 9. Add 5 μL of working solution ViaDye Red Fixable Viability Dye to the cell pellet
- 10. Vortex thoroughly
- 11. Incubate for 15 minutes at room temperature, protected from light
- 12. Add 3 mL of Stain Buffer
- 13. Centrifuge at 400 x g, 5 minutes at room temperature
- 14. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 15. Vortex thoroughly
- 16. Add 5 μL of cFluor BYG667 lgD. Vortex thoroughly
- 17. Incubate for 10 minutes at room temperature, protected from light
- 18. Add 5 μL of cFluor BYG575 CCR7. Vortex thoroughly
- 19. Incubate for 10 minutes at room temperature, protected from light
- 20. Add 60 μ L/test of the antibody cocktail prepared in step (2)
- 21. Vortex thoroughly
- 22. Incubate for 20 minutes at room temperature, protected from light
- 23. Add 3 mL of Stain Buffer
- 24. Centrifuge at 400 x g, 5 minutes at room temperature
- 25. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 26. Vortex thoroughly

NOTE: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Tubes" to fix the samples in 1% paraformaldehyde.

- 27. Resuspend in 300 μL Stain Buffer
- 28. Acquire at medium flow rate within 2 hours post staining if cells are not fixed

NOTE: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Tubes" to fix the samples in 1% paraformaldehyde.



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Cell Fixation in Tubes

If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow these steps to fix the samples in 1% paraformaldehyde and acquire within 24 hours of fixation.

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- 1. Dilute 4% paraformaldehyde in PBS to make 1% paraformaldehyde solution
- 2. Pellet the cells by centrifugation at 400 x g, 5 minutes at room temperature, if the cells are in suspension
- 3. Decant supernatant and blot on paper towel or remove the supernatant by aspiration

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- 4. Add 300 μL of 1% paraformaldehyde to the cell pellet
- 5. Vortex thoroughly
- 6. Incubate for 20 minutes at room temperature, protected from light
- 7. Add 3 mL of Stain Buffer
- 8. Centrifuge at 400 x g, 5 minutes at room temperature
- 9. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 10. Vortex thoroughly
- 11. Resuspend in 150 μL Stain Buffer for Single Stain Reference Controls and 300 μL for Multicolor Samples
- 12. Store at 4°C and acquire within 24 hours post fixation

Protocol for Staining PBMCs in 96 well Plates

Plan on using 2×10^5 cells for each Single Stain Reference Control (14 fluorescence controls, 1 ViaDye Red Fixable Viability Dye control and 1 unstained control), and 1×10^6 cells for each Multicolor Sample. Prepare separate plates for Single Stain Reference Controls and Multicolor Samples.

Use a 96 deep well V-bottom plate (polystyrene or polypropylene) to prepare the cells and transfer the final sample to a 96 well U-bottom plate (polypropylene) for acquisition, if required.

Viability Reference Control

- 1. Using a 96 deep well V-bottom plate, add 2×10^5 cells to Viability Reference Control well
- 2. Add PBS to complete the final volume to 2 mL $\,$
- 3. Centrifuge at 400 x g, 5 minutes at room temperature
- 4. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 5. Resuspend well by pipetting up and down
- 6. Repeat steps (2)-(5) if the volume in step (1) is bigger than 1 mL
- 7. Add 5 μL of working solution ViaDye Red Fixable Viability Dye to the cell pellet
- 8. Mix well by pipetting up and down
- 9. Incubate for 15 minutes at room temperature, protected from light
- 10. Add Stain buffer to complete the final volume to 2 mL
- 11. Centrifuge at 400 x g, 5 minutes at room temperature
- 12. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 13. Resuspend in 200 µL Stain Buffer by pipetting up and down
- 14. Transfer the sample to 96 well U-bottom polypropylene plate, if required
- 15. Acquire at medium flow rate within 2 hours post staining if cells are not fixed





NOTE: If the samples need to be stored at 4°C for more than 1 hour prior to collecting data, follow the steps in "Cell Fixation in Plates" to fix the samples in 1% paraformaldehyde.

Single Stain Reference Control

- Using a 96 deep well V-bottom plate, Add 2 x 10⁵ cells to each Single Stain Reference Control well
 NOTE: See Table 1 for sample type recommendations for each marker.
- 2. Add Stain Buffer to complete the final volume to 2 mL
- 3. Centrifuge at 400 x g, 5 minutes at room temperature
- 4. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 5. Resuspend well by pipetting up and down
- 6. Repeat steps (2)-(5) if the volume in step (1) is greater than 1 mL
- 7. Add 5 μL of mAb to the cell pellet in each well
- 8. Mix well by pipetting up and down
- 9. Incubate for 20 minutes at room temperature, protected from light
- 10. Add Stain Buffer to complete the final volume to 2 mL per well
- 11. Centrifuge at 400 x g, 5 minutes at room temperature
- 12. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 13. Resuspend in 200 μL Stain Buffer by pipetting up and down
- 14. Transfer the sample to 96 well U-bottom polypropylene plate, if required
- 15. Acquire at medium flow rate within 2 hours post staining if cells are not fixed
 - **NOTE:** If the samples need to be stored at 4oC for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Plates" to fix the samples in 1% paraformaldehyde

Multicolor Sample

 Prepare antibody cocktail in a 1.5 mL tube. For one Multicolor sample, 5 µL of 12 antibodies one-by-one, except for cFluor BYG575 CCR7 and cFluor BYG667 IgD.

NOTE: cFluor BYG575 CCR7 and cFluor BYG667 IgD need to be added separately. DO NOT add these antibodies in the cocktail mix.

NOTE: Prepare one extra test for the multicolor cocktail to take in account for any reagent loss in the process (ex. make multicolor cocktail for 6 tests if you have 5 multicolor samples to stain). Take 60 μ L of the cocktail per multicolor sample and discard any leftover if not used within 10 days of cocktailing.

- 2. Using a 96 deep well V-bottom plate, add 1 x 10⁶ cells to Multicolor Sample wells
- 3. Add PBS to complete the final volume to 2 mL
- 4. Centrifuge at 400 x g, 5 minutes at room temperature
- 5. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 6. Resuspend well by pipetting up and down
- 7. Repeat steps (3)-(6) if the volume in step (2) is greater than 1 mL
- 8. Resuspend well by pipetting up and down
- 9. Add 5 μL of working solution ViaDye Red Fixable Viability Dye to the cell pellet
- 10. Mix well by pipetting up and down

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- 11. Incubate for 15 minutes at room temperature, protected from light
- 12. Add Stain Buffer to complete the final volume to 2 mL
- 13. Centrifuge at 400 x g, 5 minutes at room temperature
- 14. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 15. Resuspend well by pipetting up and down
- 16. Add 5 μL of cFluor BYG667 lgD
- 17. Mix well by pipetting up and down
- 18. Incubate for 10 minutes at room temperature, protected from light
- 19. Add 5 µl of cFluor BYG575 CCR7
- 20. Add 60 μ l /test of the antibody cocktail prepared in step (1)
- 21. Mix well by pipetting up and down
- 22.Incubate for 20 minutes at room temperature, protected from light
- 23. Add Stain Buffer to complete the final volume to 2 mL
- 24. Centrifuge at 400 x g, 5 minutes at room temperature
- 25. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 26. Resuspend in 200 μL Stain Buffer by pipetting up and down
- 27. Transfer the sample to 96 well U-bottom polypropylene plate, if required
- 28. Acquire at medium flow rate within 2 hours post staining if cells are not fixed

NOTE: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Plates" to fix the samples in 1% paraformaldehyde

Cell Fixation in Tubes

If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow these steps to fix the samples in 1% paraformaldehyde and acquire within 24 hours post fixation.

- 1. Dilute 4% paraformaldehyde in PBS to make 1% paraformaldehyde solution
- 2. Pellet the cells by centrifugation at 400 x g, 5 minutes at room temperature, if the cells are in suspension
- 3. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 4. Add 300 μL of 1% paraformaldehyde to cell pellet in each well
- 5. Mix well by pipetting up and down
- 6. Incubate for 20 minutes at room temperature, protected from light
- 7. Add Stain Buffer to complete the final volume to 2 mL
- 8. Centrifuge at 400 x g, 5 minutes at room temperature
- 9. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 10. Resuspend in 200 μL Stain Buffer
- 11. Transfer the sample to 96 well U-bottom polypropylene plate, if required
- 12. Store at 4°C and acquire within 24 hours post fixation



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Protocol for Staining in Whole Blood

Plan to use 100 µl of whole blood for each Reference Control and for each Multicolor Sample.

Single Stain Reference Control

- 1. Label a 12 x 75 mm tube for each Single Stain Reference Control
- 2. Add 100 μI of whole blood to each tube

NOTE: See Table for sample type recommendations for each marker.

- 3. Add 5 μ L of appropriate monoclonal antibody to the cell pellet
- 4. Vortex thoroughly
- 5. Incubate for 20 minutes at room temperature, protected from light
- 6. Add 2 mL of 1X FACS Lysing Solution, vortex well for 10 seconds
- 7. Incubate the samples in the dark for 10 minutes, at room temperature
- 8. Centrifuge at 400 x g, 5 minutes at room temperature
- 9. Decant and blot on paper towel or remove the supernatant by aspiration
- 10. Wash with 3 ml of Stain Buffer
- 11. Centrifuge at 400 x g, 5 minutes at room temperature
- 12. Decant and blot on paper towel or remove the supernatant by aspiration
- 13. Resuspend in 300 μ L Stain Buffer
- 14. Acquire at medium flow rate within 2 hours post staining
 - NOTE: The samples can be stored at 4°C for up to 24 hours post-staining.

Table 1. Sample Type Recommendations for optimal Reference Controls

Laser	Target	Fluorochrome	Recommended Sample Type
Violet	CD3	cFluor [®] V420	Cells or Beads
	CD14	cFluor® V450	Cells or Beads
	CD45	cFluor [®] V547	Cells Only
Blue	CD8	cFluor [®] B515	Cells or Beads
	CCR7	cFluor [®] BYG575	Cells or Beads
	lgD	cFluor [®] BYG667	Cells Only
	CD45RA	cFluor [®] B690	Cells Only
	CD19	cFluor [®] BYG710	Cells or Beads
	CD25	cFluor [®] BYG781	Cells or Beads
Red	CD127	cFluor [®] R659	Cells or Beads
	CD16	cFluor [®] R668	Cells Only
	CD56	cFluor [®] R720	Cells or Beads
	CD4	cFluor [®] R780	Cells or Beads
	CD27	cFluor [®] R840	Cells or Beads

Multicolor Sample

- 1. Label a 12 x 75 mm tube for each Multicolor sample
- Prepare antibody cocktail in a 1.5 mL tube. For one Multicolor sample, 5 µL of the 12 antibodies one-by-one, except for cFluor BYG575 CCR7 and cFluor BYG667 lgD

NOTE: cFluor BYG575 CCR7 and cFluor BYG667 IgD need to be added separately. DO NOT add these antibodies in the cocktail mix.



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NOTE: Prepare one extra test for the multicolor cocktail to take in account for any reagent loss in the process (ex. make multicolor cocktail for 6 tests if you have 5 multicolor samples to stain). Take 60 μ L of the cocktail per multicolor sample and discard any leftover if not used within 10 days of cocktailing.

- 3. Add 100 μl of whole blood to the Multicolor Sample tube
- 4. Add 5 μL of cFluor BYG667 lgD. Vortex thoroughly
- 5. Incubate for 10 minutes at room temperature, protected from light
- 6. Add 5 μL of cFluor BYG575 CCR7. Vortex thoroughly
- 7. Incubate for 10 minutes at room temperature, protected from light
- 8. Add 60 μ L/test of the antibody cocktail prepared in step (2)
- 9. Vortex thoroughly
- 10. Incubate for 20 minutes at room temperature, protected from light
- 11. Add 2 mL of 1X FACS Lysing Solution, vortex well for 10 seconds
- 12. Incubate the samples in the dark for 10 minutes, at room temperature
- 13. Centrifuge at 400 x g, 5 minutes at room temperature
- 14. Decant supernatant and blot on paper towel or remove the supernatant by aspiration

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- 15. Wash with 3 mL of Stain Buffer
- 16. Centrifuge at 400 x g, 5 minutes at room temperature
- 17. Decant supernatant and blot on paper towel or remove the supernatant by aspiration
- 18. Resuspend in 300 μL Stain Buffer
- 19. Acquire at medium flow rate within 2 hours post staining

NOTE: The samples can be stored at 4°C for up to 24 hours post-staining.

NOTE: If the samples need to be stored at 4°C for more than 2 hours prior to collecting data, follow the steps in "Cell Fixation in Tubes" to fix the samples in 1% paraformaldehyde

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