

# **CYTEK 20-Color AML Panel Instructions**

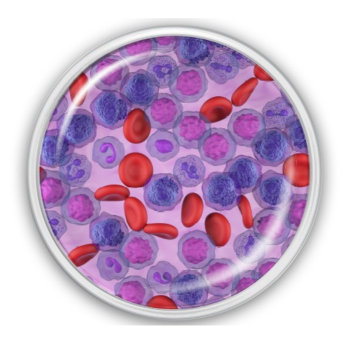
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**CYTEK 20-Color AML Panel** 



#### Introduction

For anyone working with the Cytek® 20-Color AML Panel to prepare and acquire bone marrow cells in Cytek® Northern LightTM or Aurora cytometer equipped with violet, blue and red lasers or higher, here are Cytek's recommended sample preparation procedures\*. These are 3 additional items to make your workflow easier:

- 1. Import the Cytek® 20-Color AML Panel Tags to the fluorescent tag lists in your SpectroFlo® Library section. If you already have existing tags in your library, delete them or overwrite them with the tags in this list.
- 2. Import experiment template "Cytek® 20-Color AML Panel Template" into SpectroFlo®.
- 3. Refer to Cytek® 20-Color AML Panel Acquisition Protocol for a step-by-step guide for sample acquisition and analysis in SpectroFlo®.
- Please note that this kit is designed for research use only and is not for use in diagnostic or therapeutic procedures. The following method has only been tested in bone marrow collected in the Heparin tube.
- For best results, resuspend cells in stain buffer after staining and analyze samples on Northern LightTM with 3 Lasers or Aurora within 2 hours post staining. Fixation with 1% paraformaldehyde following the procedure described in this protocol on page 4 can be performed if acquisition needs to be done at a later time, however, be aware of possible changes in the MFI for some antigens as well as quantitative differences compared to fresh samples in the enumeration of some populations.

## **Materials**

- Fresh Bone Marrow collected in Heparin tubes
- Cytek® 20-Color AML Panel, cFluor® Reagent Kit (19C) (P/N R7-40009) and CD19 Monoclonal Antibody (SJ25C1), Super Bright™ 780, eBioscience™ (P/N 78-0198-42)
- Cytek® RBC Lyse/Fix Solution 10X, R7-60010
- PBS, pH7.4, Corning 21-040-CM
- Cell strainer, Corning, 40 μm, 07-201-430
- Stain Buffer (BSA), BD Biosciences, 554657
- Paraformaldehyde solution 4% in PBS, Tonbo™, TNB-8222
- Cytek® FSPTM CompBeads, B7-10011

## **Sample Preparation**

#### Stain and lyse/fix Bone Marrow in Tubes

- Collect bone marrow into Heparin tubes\*
- Filter through a 40-μm cell strainer, and count cells using the hematology analyzer or flow cytometer, adjust cell conc. around 10 x 106 cells/mL in Stain Buffer

NOTE: If the cell count is > 20 x 106 cells/mL, dilute to 10 x 106 cells/mL in Stain Buffer.

 Plan on using ~400,000 cells for each Single Stain Reference Control (20 fluorescence, and 1 Unstained Control), and ~1 million cells for each Multicolor Sample

**NOTE:** For AML MRD evaluation, using  $\sim 10 \times 106$  cells for each Multicolor Sample (do not need to dilute the samples).

## Single Color Reference Controls

- 1. Label a 12 x 75 mm tube for each Single Stain Reference Control
- 2. Add ~50 µL of filtered bone marrow or 1 drop of Cytek® FSP™ CompBeads to each Single Stain Reference Control tube

**NOTE:** See Table 1 on page 4 for reference control type recommendations for each marker.

- 3. Add 5 µL of appropriate monoclonal antibody
- 4. Vortex thoroughly
- 5. Incubate for 20 minutes at room temperature, protected from light
- 6. For single stained cells add 2 mL 1X RBC Lyse/Fix solution

**NOTE:** Prepare 1X RBC Lyse/Fix Solution from 10X RBC Lyse/Fix solution with deionized water. For example, to make 50 mL add 5 mL of 10X RBC Lyse/Fix solution, and 45 mL of deionized water.

- 7. Vortex the tube briefly to mix
- 8. Incubate in the dark for 15 minutes at room temperature
- 9. Centrifuge at 530 x g, 5 minutes at room temperature
- 10. Decant and blot on paper towel
- 11. Vortex thoroughly to resuspend the cell pellet
- 12. Add 3 mL of stain buffer to the tube
- 13. Centrifuge at 530 x g, 5 minutes at room temperature
- 14. Decant and blot on paper towel
- 15. Vortex thoroughly to resuspend the cell pellet
- 16. For single stain beads, wash twice by adding 2 mL of stain buffer (or PBS contain 1% BSA), centrifuging (at 600 x g for 6 minutes), and aspirating the supernatant leaving approximately 50 μL of supernatant in the tube each time.
- 17. Resuspend in 300 μL Stain Buffer or go to step (1) in "Cell Fixation in Tubes" on page 3 to fix the cells or beads in 1% paraformaldehyde

**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 Tubes" on page 4 to fix the samples in 1% paraformaldehyde

18. Acquire at medium or high flow rate within 2 hours post staining if cells are not fixed

# **Multicolor Sample**

- 1. Label a 12 x 75 mm tube for each Multicolor sample
- 2. Prepare antibody cocktail according to the number of Multicolor samples. Add 5 μL per test of each antibody. **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 100 μL of the cocktail per multicolor sample and discard any leftover. Make antibody cocktails fresh each time before use and DO NOT re-use pre-made cocktails. Centrifuge the antibody cocktails at 8,000 -10,000 x g, 5 minutes at room temperature to avoid antibody aggregates. Take 100 μL supernatant per test.
- 3. Add ~100 µL of filtered bone marrow to Multicolor Sample tube
- 4. Vortex thoroughly
- 5. Incubate for 20 minutes at room temperature, protected from light
- 6. Add 2 mL 1X RBC Lyse/Fix solution
- 7. Vortex the tube briefly to mix
- 8. Incubate in the dark for 15 minutes at room temperature
- 9. Centrifuge at 530 x g, 5 minutes at room temperature
- 10. Decant and blot on paper towel
- 11. Vortex thoroughly to resuspend the cell pellet
- 12. Add 3 mL of stain buffer to the tube
- 13. Centrifuge at 530 x g, 5 minutes at room temperature
- 14. Decant and blot on paper towel
- 15. Vortex thoroughly to resuspend the cell pellet
- 16. Resuspend in 300 μL Stain Buffer or go to step (1) in "Cell Fixation in Tubes" on page 3 to fix the cells in 1% paraformaldehyde

**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 Tubes" on page 4 to fix the samples in 1% paraformaldehyde

17. Acquire at medium or high flow rate within 2 hours post staining if cells are not fixed

# **Cell Fixation in Tubes**

If the samples need to be stored at 4oC 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. Add 300 μL of 1% paraformaldehyde to cell pellet.
- 3. Vortex thoroughly.
- 4. Incubate for 20 minutes at room temperature, protected from light
- 5. Add 3 mL of Stain Buffer
- 6. Centrifuge at 400 x g, 5 minutes at room temperature
- 7. Decant and blot on paper towel
- 8. Vortex thoroughly
- 9. Resuspend in 300 µL Stain Buffer for Single Stain Reference Controls and 400 µL for Multicolor Samples
- 10. Store at 4oC and acquire within 24 hours post-fixation

Table 1. Reference Control Type Recommendations for Single Color Reference Controls

| Laser  | Target | Fluorochrome      | Recommended Control Type |
|--------|--------|-------------------|--------------------------|
| Violet | CD16   | cFluor® V420      | Cells or Beads           |
|        | CD14   | cFluor® V450      | Cells or Beads           |
|        | HLA-DR | cFluor® V505      | Cells or Beads           |
|        | CD4    | cFluor® V547      | Cells or Beads           |
|        | CD11b  | cFluor® V610      | Cells Only               |
|        | CD19   | Super Bright™ 780 | Cells or Beads           |
|        | CD7    | cFluor® B515      | Cells or Beads           |
|        | CD15   | cFluor® B548      | Cells or Beads           |
|        | CD34   | cFluor® BYG575    | Cells Only               |
|        | CD33   | cFluor® BYG610    | Cells Only               |
|        |        |                   |                          |

| Blue | CD71  | cFluor® BYG667 | Cells Only     |
|------|-------|----------------|----------------|
|      | CD38  | cFluor® B690   | Cells or Beads |
|      | CD117 | cFluor® BYG710 | Cells or Beads |
|      | CD56  | cFluor® BYG750 | Cells Only     |
|      | CD10  | cFluor® BYG781 | Cells or Beads |
| Red  | CD13  | cFluor® R659   | Cells or Beads |
|      | CD5   | cFluor® R685   | Cells Only     |
|      | CD123 | cFluor® R720   | Cells or Beads |
|      | CD64  | cFluor® R780   | Cells or Beads |
|      | CD45  | cFluor® R840   | Cells Only     |

NOTE: Recommendations are for use with Cytek® FSPTM CompBeads only.

# For Research Use Only. Not intended for use in diagnostic procedures.

cFluor® V547, cFluor® B515, cFluor® B548, cFluor® BYG610, cFluor® R685, cFluor® R720 and cFluor® R840 are equivalent to CF®405L, CF®488A, CF®514, PE-CF®596R, CF®660C, CF®700 and APC-CF®790T respectively, manufactured and provided by Biotium, Inc. under an Agreement between Biotium and Cytek (LICENSEE). The manufacture, use, sale, offer for sale, or import of the product is covered by one or more of the patents or pending applications owned or licensed by Biotium. The purchase of this product includes a limited, non-transferable immunity from suit under the foregoing patent claims for using only this amount of product for the

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DOC-00500 Rev A, Effective Date: 01/02/2023

#### **Documents / Resources**

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| CYTEK 20-Color AML Panel [pdf] Instructions          |
| 20-Color AML Panel, AML Panel, 20-Color Panel, Panel |
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## References

User Manual

Manuals+, Privacy Policy

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