**BD Phosflow™ Protocol for Human Whole Blood Samples**

## Protocol I (Detergent Method)

### Reagents Required

<table>
<thead>
<tr>
<th>Full Name</th>
<th>Short Name</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellular stimuli</td>
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<td></td>
</tr>
<tr>
<td>BD Phosflow™ Lyse/Fix Buffer, 5X</td>
<td>Lyse/Fix Buffer</td>
<td>558049</td>
</tr>
<tr>
<td>Distilled water</td>
<td></td>
<td></td>
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<tr>
<td>Phosphate buffered saline containing CaCl₂ and MgCl₂, 1X</td>
<td>PBS</td>
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<tr>
<td>BD Phosflow™ Perm/Wash Buffer I, 10X</td>
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<td>BD Phosflow™ fluorochrome-conjugated antibodies to phosphoproteins</td>
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<td></td>
</tr>
<tr>
<td>BD™ fluorochrome-conjugated antibodies to cell surface antigens (optional)</td>
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</tbody>
</table>

### Procedural Notes

- Methods and kinetics of activation vary for each phosphorylated cell signaling molecule. Select appropriate stimuli and stimulation times before beginning the protocol. See the [Suggested Stimulation Conditions for Phosphoprotein Detection](#) chart for more information.
- Intracellular phosphoproteins and cell surface antigens can be stained simultaneously. However, if there is difficulty resolving surface marker stains, surface staining can be performed before fixation or between fixation and permeabilization. Refer to the [Tested Surface Markers](#) chart, the BD FACSelect™ Buffer Compatibility Resource, and BD Phosflow™ Alternative Protocol 1: Fix–Stain–Perm or BD Phosflow™ Alternative Protocol 2: Stain–Fix–Perm for more information.
- Strict adherence to time and temperature recommendations for fixation, permeabilization, and staining is necessary for optimal resolution of phosphoprotein and cell surface marker stains.
- Be sure to remove the majority of the supernatant after each centrifugation step. High residual volumes of supernatant will dilute buffers in subsequent steps, which could result in poor staining.

### Reagent Preparation

- Prepare 1X Lyse/Fix Buffer according to the Technical Data Sheet (TDS) instructions by diluting in distilled or deionized water. Warm to 37°C for 15 to 30 minutes prior to use.
- Prepare 1X Perm/Wash Buffer I according to the TDS instructions by diluting in distilled water. Use at room temperature.

### Procedure

1. Collect whole blood in the presence of anticoagulant (EDTA or sodium heparin). EDTA is preferred for preserving light scatter properties when PMA is used as an activator. Each tube to be stained will require approximately 200 µL of whole blood.
2. Treat the cells with appropriate stimuli, and incubate at 37°C for an appropriate length of time (1 to 30 minutes; see Procedural Note). An untreated control sample should be set up in parallel.
3. After the stimulation period, fix the cells immediately by adding 10 volumes of pre-warmed Lyse/Fix Buffer to the samples. Mix well by inverting 5 to 10 times or by vortexing. Samples should be mixed promptly and thoroughly to ensure complete erythrocyte lysis.
4. Incubate the cells at 37°C for 10 to 12 minutes.
5. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
6. Vortex to disrupt the cell pellet.
7. Wash the cells:
   a. Add a volume of PBS equivalent to the volume of Lyse/Fix Buffer used.
   b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
   c. Vortex to disrupt the cell pellet.
8. Permeabilize the cells by adding 1 mL of Perm/Wash Buffer I for 1–10 x 10⁶ cells (minimum 1 mL). Mix gently and incubate for 15 to 30 minutes at room temperature.
9. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
10. Vortex to disrupt the cell pellet.
11. Wash the cells:
    a. Add a volume of Perm/Wash Buffer I equivalent to the volume used for permeabilization.
    b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.
12. Resuspend the cells in Perm/Wash Buffer I at a final concentration of 5–10 x 10⁶ cells/mL.
13. Transfer 100 µL of the cell suspension (0.5–1 x 10⁶ cells) to each 12 x 75-mm BD Falcon™ tube and add the recommended volume of BD Phosflow antibody.
14. Mix and incubate at room temperature for 60 minutes protected from light.
15. Wash the cells:
    a. Add at least 3 mL of Perm/Wash Buffer I.
    b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.
16. Resuspend the cells in approximately 500 µL of Perm/Wash Buffer I prior to flow cytometric analysis.
BD Phosflow™ Protocol for Human Whole Blood Samples

Protocol II and III (Mild or Harsh Alcohol Method)

Reagents Required

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<tr>
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<td>Cellular stimuli</td>
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</tr>
<tr>
<td>BD Phosflow™ Lyse/Fix Buffer, 5X</td>
<td>Lyse/Fix Buffer</td>
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<td>Distilled water</td>
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<td>BD Phosflow™ Perm Buffer III*</td>
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<tr>
<td>BD™ fluorochrome-conjugated antibodies to cell surface antigens (optional)</td>
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*Select either Perm Buffer II or III based on the surface markers and phosphospecific antibodies used. See the Tested Surface Markers chart and the BD FACSelect™ Buffer Compatibility Resource for more information.

Procedural Notes

- Methods and kinetics of activation vary for each phosphorylated cell signaling molecule. Select appropriate stimuli and stimulation times before beginning the protocol. See the Suggested Stimulation Conditions for Phosphoprotein Detection chart for more information.

- Intracellular phosphoproteins and cell surface antigens can be stained simultaneously. However, if there is difficulty resolving surface marker stains, surface staining can be performed before fixation or between fixation and permeabilization. Refer to the Tested Surface Markers chart, the BD FACSelect™ Buffer Compatibility Resource, and BD Phosflow™ Alternative Protocol 1: Fix–Stain–Perm or BD Phosflow™ Alternative Protocol 2: Stain–Fix–Perm for more information.

- Strict adherence to time and temperature recommendations for fixation, permeabilization, and staining is necessary for optimal resolution of phosphoprotein and cell surface marker stains.

- Be sure to remove the majority of the supernatant after each centrifugation step. High residual volumes of supernatant will dilute buffers in subsequent steps, which could result in poor staining.

Reagent Preparation

- Prepare 1X Lyse/Fix Buffer according to the TDS instructions by diluting in distilled or deionized water. Warm to 37°C for 15 to 30 minutes prior to use.

- Ensure that Perm Buffer II or III is chilled to between -20°C and 4°C.

Procedure

1. Collect whole blood in the presence of anticoagulant (EDTA or sodium heparin). EDTA is preferred for preserving light scatter properties when PMA is used as an activator. Each tube to be stained will require approximately 200 µL of whole blood.

2. Treat the cells with appropriate stimuli, and incubate at 37°C for an appropriate length of time (1 to 30 minutes; see Procedural Note). An untreated control sample should be set up in parallel.
3. After the stimulation period, fix the cells immediately by adding 10 volumes of pre-warmed Lyse/Fix Buffer to the samples. Mix well by inverting 5 to 10 times or by vortexing. Samples should be mixed promptly and thoroughly to ensure complete erythrocyte lysis.
4. Incubate the cells at 37°C for 10 to 12 minutes.
5. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
6. Vortex to disrupt the cell pellet.
7. Wash the cells:
   a. Add a volume of PBS equivalent to the volume of Lyse/Fix Buffer used.
   b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
   c. Vortex to disrupt the cell pellet. Insufficient cell resuspension prior to permeabilization may lead to cell clumping.
8. Permeabilize the cells by adding 1 mL of pre-chilled Perm Buffer II or III for 1–10 x 10⁶ cells (minimum 1 mL). Vortex to mix and incubate for 30 minutes on ice.
9. Wash the cells:
   a. Before pelleting the cells, add at least 3 mL of Stain Buffer for every 1 mL of Perm Buffer used.
   b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
   c. Vortex to disrupt the cell pellet.
10. Wash the cells two additional times:
    a. Add a volume of Stain Buffer equivalent to that used in Step 9.
    b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.
    d. Repeat steps a–c.
11. Resuspend the cells in Stain Buffer at a final concentration of 5–10 x 10⁶ cells/mL.
12. Transfer 100 µL of the cell suspension (0.5–1 x 10⁶ cells) to each 12 x 75-mm BD Falcon tube and add the recommended volume of BD Phosflow antibody.
13. Mix and incubate at room temperature for 60 minutes protected from light.
14. Wash the cells:
    a. Add at least 3 mL of Stain Buffer.
    b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.
15. Resuspend the cells in approximately 500 µL of Stain Buffer prior to flow cytometric analysis.
**Protocol IV (Harsh Detergent Method)**

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**Procedural Notes**

- Methods and kinetics of activation vary for each phosphorylated cell signaling molecule. Select appropriate stimuli and stimulation times before beginning the protocol. See the Suggested Stimulation Conditions for Phosphoprotein Detection chart for more information.

- Intracellular phosphoproteins and cell surface antigens can be stained simultaneously. However, if there is difficulty resolving surface marker stains, surface staining can be performed before fixation or between fixation and permeabilization. Refer to the Tested Surface Markers chart, the BD FACSelect™ Buffer Compatibility Resource, and BD Phosflow™ Alternative Protocol 1: Fix–Stain–Perm or BD Phosflow™ Alternative Protocol 2: Stain–Fix–Perm for more information.

- Permeabilization with Perm Buffer IV might result in decreased cell recovery. For maximal cell recovery, avoid aspirating the supernatant during all post-permeabilization washes. Instead, decant the supernatant and gently blot the tube edge on an absorbent surface to minimize residual volume.

- Longer permeabilization time or using a ratio of cell to buffer volume outside the recommended ratio might result in increased cell loss and poorer fluorescent surface marker and/or phosphoprotein-specific antibody staining and detection.

- Perm Buffer IV may be used at a 1X or 0.5X concentration. The 1X concentration might result in increased cell loss and decreased ability to stain certain cell surface markers, but it provides optimal resolution of certain intracellular phosphoprotein stains. The 0.5X concentration results in less optimal staining of some intracellular phosphoproteins, but it may provide improved cell recovery and improved compatibility for staining cell surface CD markers. See the BD FACSelect™ Buffer Compatibility Resource for more information.

- Strict adherence to time and temperature recommendations for fixation, permeabilization, and staining is necessary for optimal resolution of phosphoprotein and cell surface marker stains.

- Be sure to remove the majority of the supernatant after each centrifugation step. High residual volumes of supernatant will dilute buffers in subsequent steps, which could result in poor staining.

**Reagent Preparation**

- Prepare 1X Lyse/Fix Buffer according to the TDS instructions by diluting in distilled or deionized water. Warm to 37°C for 15 to 30 minutes prior to use.

- Prepare 1X or 0.5X Perm Buffer IV according to the TDS instructions by diluting in 1X PBS. Use at room temperature. See Procedural Notes and the BD FACSelect™ Buffer Compatibility Resource for information on choosing between 1X and 0.5X Perm Buffer IV.
BD Phosflow™ Protocol for Human Whole Blood Samples

Procedure

1. Collect whole blood in the presence of anticoagulant (EDTA or sodium heparin). EDTA is preferred for preserving light scatter properties when PMA is used as an activator. Each tube to be stained will require approximately 200 µL of whole blood.

2. Treat the cells with appropriate stimuli, and incubate at 37°C for an appropriate length of time (1 to 30 minutes; see Procedural Note). An untreated control sample should be set up in parallel.

3. After the stimulation period, fix the cells immediately by adding 10 volumes of pre-warmed Lyse/Fix Buffer to the samples. Mix well by inverting 5 to 10 times or by vortexing. Samples should be mixed promptly and thoroughly to ensure complete erythrocyte lysis.

4. Incubate the cells at 37°C for 10 to 12 minutes.

5. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.

6. Vortex to disrupt the cell pellet.

7. Wash the cells:
   a. Add a volume of PBS equivalent to the volume of Lyse/Fix Buffer used.
   b. Centrifuge at 600g for 6 to 8 minutes and remove the supernatant, leaving no greater than 50 µL of residual volume.
   c. Vortex to disrupt the cell pellet.

8. Permeabilize the cells by slowly adding 1 mL of Perm Buffer IV drop by drop for 0.5–2.0 x 10^6 cells (minimum 1 mL). Vortex to mix and incubate for 15 to 20 minutes at room temperature.

9. Centrifuge at 600g for 6 to 8 minutes. Decant the supernatant and gently blot the tube edge on an absorbent surface, leaving no greater than 50 µL of residual volume.

10. Vortex to disrupt the cell pellet.

11. Wash the cells twice:
    a. Add at least 3 mL of Stain Buffer.
    b. Centrifuge at 600g for 6 to 8 minutes. Decant the supernatant and gently blot the tube edge on an absorbent surface, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.
    d. Repeat steps a–c.

12. Resuspend the cells in Stain Buffer at a final concentration of 5–10 x 10^6 cells/mL.

13. Transfer 100 µL of the cell suspension (0.5–1 x 10^6 cells) to each 12 x 75-mm BD Falcon tube and add the recommended volume of BD Phosflow antibody.

14. Mix and incubate at room temperature for 60 minutes protected from light.

15. Wash the cells:
    a. Add at least 3 mL of Stain Buffer.
    b. Centrifuge at 600g for 6 to 8 minutes. Decant the supernatant and gently blot the tube edge on an absorbent surface, leaving no greater than 50 µL of residual volume.
    c. Vortex to disrupt the cell pellet.

16. Resuspend the cells in approximately 500 µL of Stain Buffer prior to flow cytometric analysis.