Basic Flow Cytometry Staining Protocol

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Date: 12/20/17
Modified: 5/15/19

Materials
- Cells to be stained
- FACS buffer – can be cell dependent
  - 1X Ca/Mg2+ free PBS
  - Either FBS (1-10%) or BSA (0.1-1%)
  - Optional: EDTA (0.5-5mM) if you have very sticky cells
  - Commonly used: 1% FBS in 1X PBS (consider filtering FBS to remove debris)
- Antibodies
- Live/dead stain (this protocol is for unfixable stains such as DAPI, PI, and the Sytox dyes)
- 5 mL flow tubes: Falcon #352008

Procedure
1. Prepare single cell suspension
2. lyse RBCs (optional, if needed)
3. Count cells
4. Adjust cell concentration to 0.1-5 million cells per 50 µL FACS buffer
   a. Number of cells needed depends on your experiment. General starting point is often 1 million cells.
5. Pipette 50 µL cells into Eppendorf or 5 mL flow tubes for staining
   a. Need one tube per sample and single stain controls
6. Add Fc block
   a. 1-5 µL per sample following manufacturer’s protocol
      i. Anytime you need to pipette 1 µL it’s easier to make a 1:10 dilution in FACS buffer and add 10 µL to each sample
7. Make sure samples are mixed by gently vortexing or pipetting up and down
8. Incubate at 4°C for 10-15 minutes
9. Add antibody cocktail – 50 µL total volume per tube
   a. For example, if you are adding 2 µL of antibody per tube, you would first combine 2 µL antibody and 48 µL FACS buffer, then add the 50 µL of antibody cocktail to the 50 µL of cells for a total staining volume of 100 µL
   b. For samples: make master mix (example below of 10X master mix) of all antibodies so that appropriate amount is in 50 µL (when added to cells, final staining volume is 100-110 µL, calculate for 100µL final volume)
      i. Always add an extra 1-2 to your master mix to account for pipetting error, so a 10x master mix will be good for 8-9 samples
10. Mix well and incubate for 30 minutes in the dark at 4°C (cover tubes in foil if needed)
11. Wash cells by adding FACS buffer
   a. 1 mL if staining in Eppendorf tubes
   b. 2 mL if staining in flow tubes
12. Centrifuge
13. Aspirate, pipette, or dump tubes to remove supernatant  
   a. Be careful to not disturb the pellet!
14. Resuspend in FACS buffer  
   a. Volume depends on the number of cells  
   b. Generally no less than 350 µL
15. If cells were stained in Eppendorf tubes, transfer to 5 mL FACS tubes
16. A few minutes before running samples on the cytometer, add live/dead stain (see manufacturers protocol)