Notes

- PBMC = peripheral blood mononuclear cells.

- Principle of separation (source GE Healthcare’s booklet Ficoll Paque PLUS 1.077 g/mL): The separation of blood cells is performed by buoyant density centrifugation. The median buoyant densities of human monocytes, lymphocytes and thrombocytes (platelets) are 1.067-1.077, 1.073-1.077 and 1.04–1.08 g/mL, respectively. With the exception of basophils, polymorphonuclear leukocytes (PMNs) have a much greater buoyant density ($\rho > 1.082$ g/mL) than mononuclear cells. Hence, “on centrifugation, cells in the blood sample sediment towards the blood/Ficoll-Paque media interface, where they come in contact with the Ficoll PM400 present in Ficoll-Paque products. Red blood cells are efficiently aggregated by this agent at room temperature. Aggregation increases the rate of sedimentation of the red cells, which rapidly collect as a pellet at the bottom of the tube, where they are well separated from lymphocytes with yields:

- some lymphocytes with 

- platelet aggregation (PMNs) have 

- mononuclear cells. Granulocytes also sediment to the bottom of the Ficoll media layer. This process is facilitated by an increase in their densities caused by contact with the slightly hypertonic Ficoll-Paque media. Thus, on completion of centrifugation, both granulocytes and red blood cells are found at the bottom of the tube, beneath the Ficoll-Paque product. Lymphocytes, monocytes, and platelets are not dense enough to penetrate into the Ficoll Paque PLUS medium layer having a density of 1.077 g/mL. These cells therefore collect as a concentrated band at the interface between the original blood sample and the Ficoll-Paque product. This banding enables the mononuclear cells to be recovered with high yield in a small volume with little mixing with the Ficoll-Paque media. When Ficoll-Paque PREMIUM 1.073 is used (instead of Ficoll-Paque PLUS 1.077 g/mL), some lymphocytes with densities >1.073 will enter the Ficoll-Paque media layer and the resulting cell preparation will be enriched for lower density cells like mesenchymal stromal/stem cells and monocytes. Washing and centrifuging the harvested cells subsequently removes platelets, any contaminating FicollPaque media, and plasma. The resulting cell suspension then contains highly purified, viable lymphocytes, monocytes, mesenchymal stromal/stem cells, and is suitable for further studies.”

- Some critical steps are:
  - Perform the isolation and freezing of PBMCs less than 8 hours after the blood collection. Best results are obtained if the procedure is performed less than 2 hours after blood collection.
  - Dilute the blood with PBS prior to overlay: “When erythrocytes in whole blood are aggregated, some lymphocytes are trapped in the clumps and therefore sediment with the erythrocytes. This tendency to trap lymphocytes is reduced by diluting the blood. Dilution gives a better lymphocyte yield and reduces the size of the red cell clumps. Aggregation of erythrocytes is enhanced at higher temperatures (37°C), which consequently decreases the yield of lymphocytes. At lower temperatures (4°C), however, the rate of aggregation is decreased but the time of separation is increased, which also decreases the yield of lymphocytes. A compromise temperature of 18–20 °C gives optimal results” (source: GE Healthcare).
  - Minimize the time that the cells remain in a pellet or in contact with the Ficoll.
  - Wash the cells carefully after the Ficoll.
  - Do not stop during the cell freezing but add slowly the DMSO-containing solution to the cells (in most cases cells are first resuspended in FBS and DMSO-containing solution is then added dropwise). Most labs freeze cells either at 4°C or at room temperature.

- The expected PBMC yields from whole blood for healthy populations are (source: HIV AIDS Network coordination PBMC processing SOP):

<table>
<thead>
<tr>
<th>Population</th>
<th>Mononuclear Cell Yield Range (cells/mL blood)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>(0.8 to 3.2) x 10^6</td>
</tr>
<tr>
<td>Pediatric—less than 6 months</td>
<td>(3 to 10) x 10^6</td>
</tr>
<tr>
<td>Pediatric—6 mo. to 2 years</td>
<td>(2 to 9) x 10^6</td>
</tr>
<tr>
<td>Pediatric—2 to 5 years</td>
<td>(1 to 6) x 10^6</td>
</tr>
<tr>
<td>Pediatric—more than 5 years</td>
<td>(0.8 to 4) x 10^6</td>
</tr>
<tr>
<td>Pediatric—Unknown age</td>
<td>(1 to 10) x 10^6</td>
</tr>
</tbody>
</table>

- Buffy coats are isolated from whole blood at the blood bank by an initial centrifugation without a density gradient. This initial spin concentrates red blood cells at the bottom and plasma on top, the buffy coat forms the interface and contains most of the leukocytes and platelets from the whole blood (although it still looks like blood). Each unit (~470 mL) of whole blood yields a ~60-45 mL buffy coat, which contains ~200-800*10^6 mononuclear cells. At the Purpan EFS blood bank, the night shift team performs this spin on the day of the blood collection, stores the buffy coat at 20°C and makes it available to the CPTP on the next morning.

- PBMCs are usually frozen right away after the Ficoll isolation and used subsequently in functional assays. However, with functional tests on fresh or frozen cells, a resting period (typically overnight) is often recommended after Ficoll isolation or cell thawing.
Equipment, reagents and reagent preparation

- PBS without calcium and magnesium. Keep and use at room temperature.
- Density gradient media suitable for human blood cells: e.g. Ficoll-Paque PLUS (density 1.077 g/mL, GE Healthcare Pharmacia) or Pancoll human density 1.077 g/mL (PanBiotech ref P04-60500). Do not use Ficoll-Paque PREMIUM density 1.073 g/mL (gives greater enrichment in monocytes or mesenchymal stem cells) or 1.084 g/mL (adapted for rodent PBMC).
- DMSO Grade culture (e.g. Sigma ref D2650). After opening, undiluted DMSO is stable at room temperature (15 to 30°C) when protected from light and moisture, for 6 months. Reagent may be aliquoted in small amounts to help preserve sterility. Label aliquots with “DMSO,” the date opened/aliquoted, the expiration date (six months from opening) and tech initials. Protect aliquots from light.
- Heat-inactivated FBS thawed and stored at 2 to 8°C (stable for one calendar month).
- NALGENE® Mr. Frosty, 1°C/minute cryo-freezing container (can hold 18 cryovials). Mr Frosty requires isopropanol that needs to be replaced once a month (in a specific waste bottle) and level must be checked after each freeze-thaw cycle. Alternatively, some dry cryopreservation modules can be used such as the StrataCooler CryoLite preservation module (32 cryovial holder from Agilent Technologies) that cools down at a controlled rate of 0.4–0.6°C/minute. The freezing boxes must be left to dry between uses and cooled down at 4°C overnight before the PBMC isolation.
- 2 mL cryotubes (e.g. Biosigma CL2ARBIPS).

PBMC isolation from buffy coats or whole blood

1. Store blood tubes at room temperature prior to the isolation of PBMCs. Measure the usable whole blood volume within 0.5 mL. All steps below are performed at room temperature.

2. Blood dilution:
   2.1 Remix the blood by gently inverting the blood collection tube 6-8 times.
   2.2 Wipe off the blood tubes or buffy coat bags with 70% ethanol.
   2.3 Transfer the blood tubes or the buffy coat in a single recipient that can contain at least twice its volume.
   2.4 Dilute the blood with equal amount of PBS.
   2.5 Mix gently the blood and PBS.

3. Density Gradient Cell Separation:
   3.1 Invert the Ficoll-Paque PLUS or Pancoll density 1.077 g/mL (optimized for human cells as explained above) bottle several times to ensure thorough mixing
   3.2 Add Ficoll-Paque at the bottom of a new tube without touching the side of the tube since the reagent is toxic.
      3.2.1 Use 4 mL Ficoll-Paque PLUS for 6 to 10 mL diluted blood in a 15 mL conical tube.
      3.2.2 Use 20 mL Ficoll-Paque PLUS for 30 to 35 mL diluted blood in a 50 mL conical tube.
      3.2.3 Typically with buffy coats we use 4 x 50 mL conical tubes filled with 20 mL Ficoll + 30-35 mL diluted buffy coat
   3.3 Gently overlay the Ficoll with the diluted blood. Allow the diluted blood to flow down the side of the tube and pool on top of the density gradient media surface without breaking surface plane (see scheme below).

4. Gradient centrifugation: Centrifuge at room temperature at 760 xg (1900 rpm on Heraeus Megafuge 1.0R) for 20 min with the brakes OFF since the deceleration disrupts the density gradient. If the centrifuge starts shaking, stop immediately but gently; reweigh and balance the tubes.
Human PBMC isolation and freezing

**5. PBMC collection and washing**

5.1 Thorough washing of buffy coats = 5 or more washes is important in order to reduce platelet contamination. With whole blood, 2 to 3 washes are generally sufficient.

5.2 Label new sterile conical tube(s), same number and same volumes as for the density gradient cell separation, which will contain the PBMCs.

5.3 Remove as much plasma layer as possible.

5.4 Collect the mononuclear cells from the plasma/Ficoll interface with a disposable transfer pipet. Transfer the cells from each Ficoll tube into a new sterile conical tube. If several Ficoll tubes were used for the same donor, do not pool cells yet. While collecting the cells, be sure to aspirate plasma only and as little Ficoll as possible (typically 2 mL from a 15 mL tube and 8 mL from a 50 mL tube). Fill each tube with PBS.

5.5 Perform 3 (blood) or 5 washes (buffy coats):

5.5.1 Spin cells 350 x (1300 rpm on Heraeus Megafuge 1.0R) for 8 min. Do not forget to switch on again the centrifuge brakes.

5.5.2 Remove carefully the supernatant (the pellet is loose because of red blood cells) by pipetting off the supernatant without touching the pellet (do not even tilt the tube...). Suspend the cell pellet by gentle pipetting (do not use the p1000 with narrow tips but rather a p5 – 25 mL) in a low volume of PBS and fill the tube with additional PBS.

5.5.3 After the 2nd wash, tubes from the same donor can be pooled 2 by 2.

5.5.4 For increased removal of platelets, perform one wash with centrifugation at low speed i.e. 200 xg (900 rpm on Heraeus Megafuge 1.0R) for 10 min with the centrifuge brakes on. Do not perform this step if the blood volume was low.

6. Analysis of PBMC count and viability (can be done during the last wash):

6.1 Pipet off the supernatant and loosen the pellet by adding 0.5 to 1 mL PBS and gently resuspend cells with the 1 mL pipet. Add PBS to bring cells at approximately 5x10⁶ cells/ml (max 10.10⁶ cells/ml), knowing that each mL of blood will give a rough average of 1.5x10⁶ PBMCs or that a buffy coat contains 200 million to 1 billion PBMCs.

6.1.1 Ex: suspend PBMC a 50 mL whole blood collection into 15 mL PBS

6.1.2 Ex: suspend PBMCs from a 65 mL buffy coat into 50 mL PBS

6.2 Determine cell concentration and viability with the Cellometer 2000 (linearity: 2 to 12x10⁶ cells/mL)

6.2.1 Principle: Cells are stained with acridine orange, AO, and propidium iodide, PI. AO is a nuclear staining (nucleic acid binding) dye permeable to both live and dead cells. It stains all nucleated cells to generate green fluorescence. PI can only enter dead cells with compromised membranes. PI stains all dead nucleated cells to generate red fluorescence. Cells stained with both AO and PI fluoresce red due to quenching, so all live nucleated cells fluoresce green and all dead nucleated cells fluoresce red.

6.2.2 Protocol: Remove the 2 plastic covers of the counting slide (each slide has 2 counting chambers). Mix 20 microl cells with 20 microl AO/PI. Mix 5 times and load 20 microl
into the chamber. Adjust the focus (perform an image preview) and read immediately and only once the cell staining (perform a cell count using the “immune cells, low RBC” program). Record the live cell concentration and the percentage of live cells.

6.2.3 QOC (whole blood) should show a fresh PBMC yield above 8x10^5 PBMC/mL whole blood and viability above 95%. Long processing time and poor technique may adversely affect the viability.

**PBMC freezing (below 4°C freezing protocol), example for 2.10^7 cells/cryovial**

1. Thaw an aliquot of FBS and allow cooling at 2 to 8°C.
2. Put the NALGENE® Mr. Frosty or StrataCooler freezing boxes at 2 to 8°C overnight.
3. Prepare the 20% DMSO/FBS mixture and allow cooling at 2 to 8°C. The mixture can be stored at 2 to 8°C for 1 working day.
4. Hand-label the cryotubes or print cryolabels (e.g. Protocol /Participant code/PBMC/date yyyy-mm-dd).
5. Perform the steps below on an ice pan and do not stop during the procedure:
   5.1 Suspend PBMCs in cold FBS at 4x10^7 cells/mL: first add 0.5 to 1 mL FBS, mix/detach the cells by gentle pipetting with a p5-25 mL (do not use narrow tips) and then add the remaining volume of FBS.
   5.2 Add the same volume of cold FBS 20% DMSO dropwise SLOWLY 1 drop per second while swaying the tube to gently mix the cell suspension (final solution: FBS 10% DMSO).
   5.3 Mix gently 3 times by tube inversion.
   5.4 Using the p5 mL pipette, aliquot 1 mL cell suspension to each cryotube, firmly close the lid and put the tubes into the NALGENE® Mr. Frosty.
   5.5 If freezing boxes are not available, the following “low technology” method works equally well:
      5.5.1 Place the cryovials in a Styrofoam rack (e.g., racks for 15 ml conical tubes).
      5.5.2 Place a second Styrofoam container of the same type over the first one and tape the two containers together.
      5.5.3 Insert into a plastic bag leaving some air in the bag before taping it closed.
6. Place the freezing boxes or the Styrofoam container immediately into a -80°C freezer for 12 to 24 h (max 4 days), then transfer the cryovials into the liquid nitrogen tank.
7. Avoid any temperature increase during the transfer in the nitrogen tank and, in general, prior to the thawing of the cells.