

HOW TO COUNT CELLS AND DETERMINE PERCENT VIABILITY USING THE TRYPAN BLUE EXCLUSION METHOD

MATERIALS

- 0.4% Trypan Blue
- Neubauer Hemacytometer and cover slip
- Dulbecco's Phosphate buffered saline (DPBS) optional

When you receive an AllCells fresh product, please count the cells immediately upon receipt. Cryopreserved products should be counted immediately after thawing before any further manipulation. AllCells is not liable for any cell loss during subsequent processing or manipulation.

FRESH CELL PRODUCTS

1. Proper aseptic technique and personal protective equipment should always be used when handling cell products.
2. Mix the cell suspension thoroughly and to disperse any cell clumps to ensure even distribution of the cells.
3. For fresh cell products shipped in a transfer bag, use both hands to pick up the transfer bag by the top and bottom edges and invert the bag side to side (approximately 1 second for each rotation) for 5 total rotations to mix.
 - a. Wipe the transfer line with an alcohol swab. Make sure the line is clamped before cutting the line with sterilized scissors.
 - b. After the line has been cut, unclamp the line and dispense a 2-3mL aliquot of the cell product into a sterile tube.
 - c. Mix the cells in the tube by inverting 3-5 times or if cell lumps are present, pipette up and down at least 5 times with a P1000 pipetman.

- d. Immediately transfer 50-100µL aliquot of the cell product into a fresh tube for dilution prior to counting.
- e. Dilute the test sample with the appropriate volume of DPBS to target approximately 2M cells/mL and mix thoroughly.

Example Dilution:

Our apheresis product (LP, FR, Full) contains approximately 1×10^{10} total nucleated cells in about 250mL. The estimated cell concentration is 4×10^7 cells/mL (1×10^{10} cells/250mL).

A 20-fold dilution (50µL sample + 950µL DPBS) will result in a test sample of approximately 2×10^6 cells/mL. The final dilution factor is 40 (20 x 2) after 1:1 dilution with 0.4% trypan blue for counting.

- f. Proceed to Step 5.
4. For all other containers, mix the cells by gently pipetting 25-50% of the cell suspension up and down at least 5 times using an appropriately sized pipettor or serological pipette.

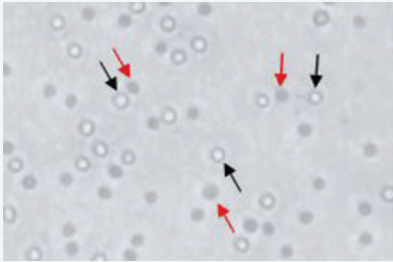


Figure 1:
Differentiation of live and dead cells using trypan blue dye exclusion. Black arrows indicate live cells and red arrows indicate dead cells stained with dye.

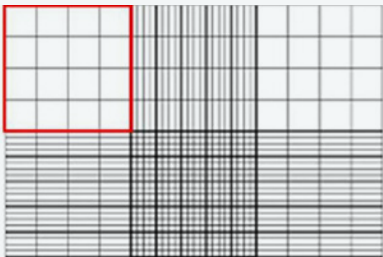


Figure 2:
Hemocytometer with Neubauer gradients (from http://www.hausser-scientific.com/products/hausser_bright_line.html). A counting quadrant is highlighted in red outline.

- a. For whole fresh bone marrow product shipped in a bottle, a 4-10 fold dilution of the test sample with DPBS before addition of trypan blue and counting is recommended.
 - b. Based on our target cell number and fill sizes, no additional dilution for our other fresh cell products is needed; however, an optional additional dilution of test samples from our 25, 100 and 300M fill sizes with DPBS before proceeding to Step 5 may facilitate counting.
5. Immediately take a 10 μ L aliquot of the cell product from the vial/tube after mixing.
 6. Combine 10 μ L of the diluted/undiluted test sample with 10 μ L 0.4% trypan blue. The test sample should be counted within 5 minutes after mixing with trypan blue.
 7. Load the hemacytometer with approximately 10 μ L of the trypan blue-labeled sample. Make sure the entire chamber is filled with the test sample.
 8. Under bright field, dead cells appear light to dark blue while viable cells appear clear to autofluorescent (Figure 1).
 9. Count all the viable cells in a quadrant (Figure 2). To achieve better accuracy, ≥ 100 viable cells should be counted. More than one quadrant may need to be counted to achieve ≥ 100 viable cells counted.
 10. Count the dead cells in the same quadrants.
 11. Calculations to determine the cell concentration and percent viability.

$$\text{Cell Concentrations (cells/mL)} = \frac{\# \text{ total viable cells counted}}{\# \text{ quadrants counted}} \times \text{final dilution factor} \times 10^4^*$$

$$\text{Cell Viability (\%)} \times 100 = \frac{\# \text{ viable cells counted}}{\# \text{ viable cells} + \# \text{ dead cells counted}} \times 100$$

CRYOPRESERVED CELL PRODUCTS

1. Follow the AllCells procedure for thawing cells.
2. After the cell product is completely thawed, invert the vial 5 times to mix the cells and immediately transfer a 20 μ L aliquot of the cell product into a microcentrifuge tube.
3. For product unit sizes containing 1M cells or less, follow Steps 5-11 as instructed for counting Fresh Cell products.
4. If the unit size of the cell product is >1M cells, dilute the test sample by slowly adding (around one drop per second) an appropriate volume of cold (2-8 $^{\circ}$ C) DPBS to target approximately 2M cells/mL concentration.

Example Dilution:

Our LP, CR, MNC, 10M product contains no less than 10×10^6 total viable nucleated cells in 1.8mL.

The estimated cell concentration is $\geq 5.6 \times 10^6$ cells/mL ($\geq 10 \times 10^6$ cells/1.8mL). A 3-fold dilution (20 μ L sample + 40 μ L DPBS) will result in a test sample of approximately $\geq 1.87 \times 10^6$ cells/mL. The final dilution factor is 6 (3 x 2) after 1:1 dilution with 0.4% trypan blue for counting.

5. Gently mix the diluted test sample and follow Steps 5-11 as instructed for counting Fresh Cell products.

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