

Moxi Cyte Single Color Viability Reagent (MXA055) User's Guide

(Rev - 20250206)

FOR RESEARCH USE ONLY For Moxi V and Moxi GO II Systems



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Overview

The Moxi Cyte Single Color Viability Reagent (MXA055) is a Propidium Iodide (PI) – based staining reagent has been specially formulated for use with ORFLO's Moxi V or Moxi GO II instruments to provide rapid assessment of the viability of a cell sample. The assay is based on the general membrane-integrity/dye-exclusion method of assessing cell viability. PI is a cell membrane-impermeant, DNA-intercalcating dye that increases fluorescence over 50x upon binding to DNA. For healthy cells, the dye cannot penetrate the cell membrane and therefore cannot bind to its DNA and fluoresce. As a result, no fluorescence signal is recorded as this cell is counted. However, as cells lose membrane integrity through necrosis or apoptosis, this PI-based reagent permeates the cell and a strong fluorescence signal is generated.

Components

• 111 test (15ml) vial of ORFLO Moxi Cyte Single Color Viability Reagent (MXA055)

Storage and Handling

- Store the kit at 2-8°C
- Use product aseptically
- Minimize light exposure.



Protocol

- 1. Prepare cells into a single-cell suspension free of large (>30μm) particulate.
 - a. Notes:
 - i. Primary harvests (e.g. blood or tissue extractions):
 - 1. If large extracellular debris is present, pass sample through a $40\mu m$ cell strainer to remove it.
 - 2. If large amounts of RBCs are present in the sample, an appropriate RBC lysis method must be applied to get proper counts (RBCs will mask the signal of the cell population of interest due to their extremely high concentration levels).
 - ii. Cell aggregation/Clustering Clusters and aggregates can be dissociated through pipette trituration (repeatedly, e.g. 10x, pull a substantial fraction of the cell sample in and out of a pipette tip). If a dissociation treatment was used, perform this step while cells are still suspended in the dissociation reagent.
- 2. Mix cell suspension and Moxi Cyte Single Color Viability Reagent (invert several times to mix) according to the following table:

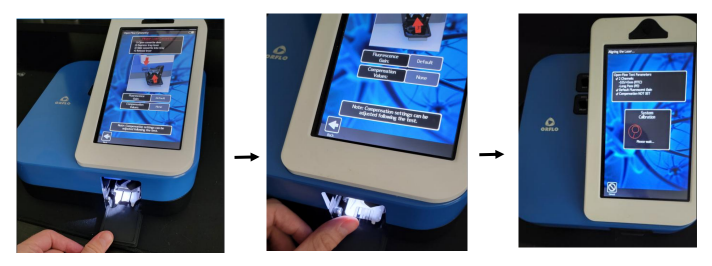
Concentration Of Original Cell Suspension	Moxi Cyte Viability Reagent Volume	Cell Suspension Volume	Dilution Factor
1×10^5 to 3×10^6 cells/ml	135 μL	15 μL	10x
3×10^6 to 6×10^6 cells/ml	190 μL	10 μL	20x
6 x 10 ⁶ to 1 x 10 ⁷ cells/ml	290 μL	10 μL	30x
>1 x 10 ⁷ cells/ml	490 μL	10 μL	50x

- 3. Incubate for 5 minutes in the dark at room temperature (25°C).
- 4. Analyze with the Moxi V or Moxi GO II using the "Cell QC" or "Cell Count (Size + Viability)" app within 20 minutes of initial mixing/staining.

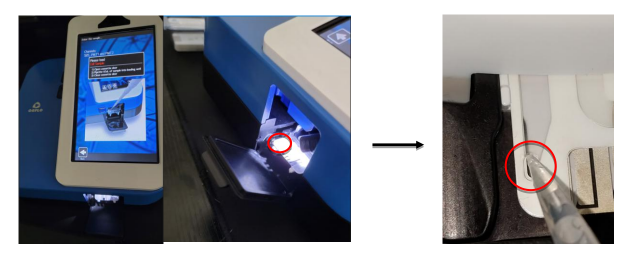


Running a Test (Note: For detailed Moxi V and Moxi GO II operation instructions and images, please see the latest User Guides at https://www.orflo.com/user-manuals/ as well as virtual demos for the Moxi V and Moxi GO II)

- 1. Prepare stained cell sample as described above.
- 2. Turn unit on and select the "Cell QC" or "Cell Count (Size + Viability)" app.
- 3. Open the door fully, insert an unused cassette, close the door, and wait for the laser cassette alignment to complete.



4. When the alignment completes (and prompted to "Enter the Sample" at the top left black bar), open the door to the first stop (45° angle) and pipette $60\mu L$ of the sample into the loading well in one fluid motion.



5. Immediately close the door (try not to delay longer than a second or two or the sample will begin to settle). Once the door is closed, the test will automatically run.



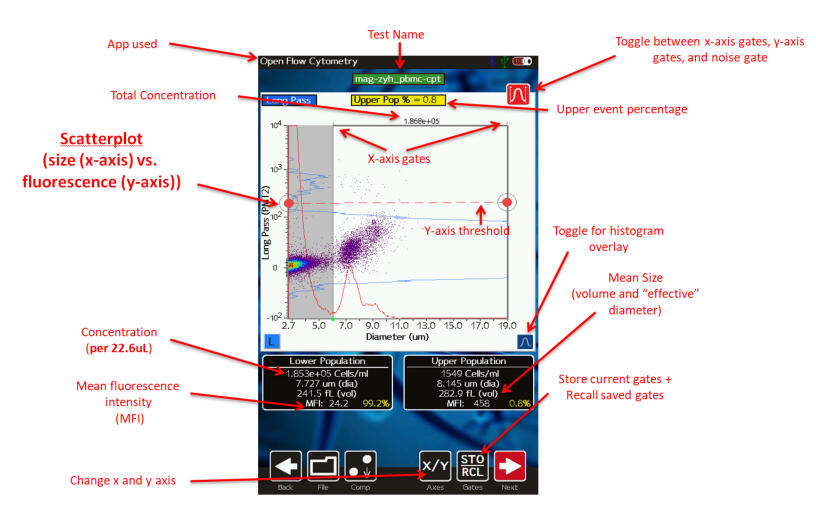
<u>Analyzing Results</u> (Note: For detailed Moxi V and Moxi GO II operation instructions and images, please see the latest User Guides at https://www.orflo.com/user-manuals/ as well as virtual demos for the Moxi V and Moxi V">Moxi V"<

- 1. Results are initially displayed as a Scatter/Dot Plot, Log[PMT V] vs. Diameter (μ m), with cell population gates set from the prior test run.
- 2. Refer to the image below for adjustments on the post-run data analysis screen

Moxi V: **Test Name** Toggle between x-axis gates, y-App used axis gates, and noise gate **Via-**034 Live Cell % = 26.7 **Total Concentration** Viability Percentage 3.743e+05 **Scatterplot** X-axis gates Dead (size (x-axis) vs. fluorescence (y-axis)) Toggle for histogram Y-axis threshold overlay Live **Excluded events** from noise gate Mean Size (shaded grey) (volume and "effective" diameter) 6.0 7.0 8.0 9.0 10.0 11.0 Diameter (um) Concentration (per 22.6uL) Live Cells Dead Cells 2.744e+05 Cells/ml 982e+04 Cells/m 4.906 um (dia) 61.83 fL (vol) MFI: 431 4.376 um (dia) Store current gates + Mean fluorescence Recall saved gates intensity (MFI) Save test results Change X-axis zoom Change x and y axis



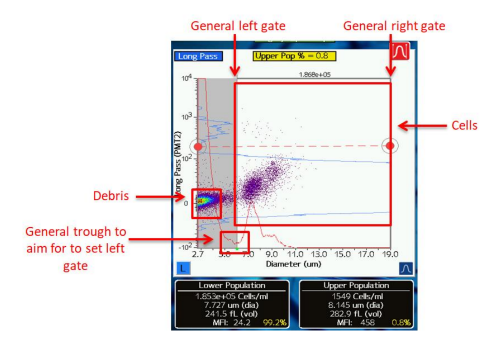
Moxi GO II:



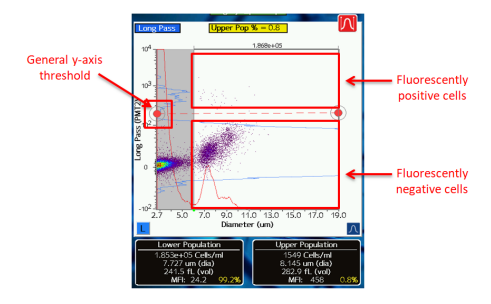
- 3. If it is difficult to view the cell population, touch the rescale icon to "zoom in" until you can clearly see the cells. This is only necessary for smaller particles, e.g. isolated nuclei.
- 4. Make sure to press the histogram overlay toggle (blue button at the bottom right of the scatterplot), as this will allow for significantly easier and more intuitive sample gating.



5. Adjust the size gates to include the cell region by dragging the blue markers to define the appropriate size region (image below). For general gating, the left x-axis gate should be set at the trough between the debris peak (\sim 3µm) and the cells. The right x-axis gate should be set all the way to the far right to include all cells. If auto-gating is enabled, the system will do this gating automatically. Note: The left marker can be angled if needed by toggling the left gate pivot (touch the red or green dot at the bottom of the gate to toggle).



6. Touch the red gate icon (top right of scatter plot, indicated in the left image below) to enable fluorescent gating (y-axis). For general gating for viability, set the gate at the beginning of the base of the non-fluorescent population (meaning it's including only clearly living cells) by using the histogram overlay.





- 7. Drag the red markers to position the dashed red live/dead (fluorescence) gating marker between the live and dead clusters. If auto-gating is enabled, the system will do this gating automatically.
- 8. *Sample/Test information* (refer to image at top of page 4 or 5 depending on instrument):
 - a. *Total Cell Concentration*: listed above the scatter/dot plot.
 - b. *Upper Population Percentage*: listed in the yellow box above the scatter/dot plot.
 - c. Lower Cell Population Concentration: Listed in the black box at the bottom-left of the scatter/dot plot.
 - d. *Lower Cell Population Mean Diameter*: Listed in the black box at the bottom-left of the scatter/dot plot.
 - e. *Upper Cell Population Concentration*: Listed in the black box at the bottom-right of the scatter/dot plot.
 - f. *Upper Cell Population Mean Diameter*: Listed in the black box at the bottom-right of the scatter/dot plot.
 - g. *Mean Fluorescence Intensity (MFI)*: Listed in the black box at the bottom for each cluster (Live/Dead).

Troubleshooting

- No/poor fluorescent separation:
 - a. Check total cell concentration Increase or decrease dilution factor (see table above) to achieve optimal concentrations.
 - b. Reagent may have expired or degraded Optimal kit storage conditions are **2-8°C in the dark**. Always use a clean pipette tip to avoid contamination of the reagent. If you suspect the reagent integrity may have been compromised, re-order new reagent.
- <u>Incomplete test or poor fluid flow:</u>
 - a. Over-concentration samples can result in clogging. Increase dilution factor (see table on Page 2) to achieve optimal concentrations.
 - b. Remove large particle/particulate by straining sample with a cell strainer (e.g. 40 μ m).
 - c. Break apart cell clusters/aggregates with pipette trituration.
- Cell population not visible/discernible:
 - a. Re-scale the x-axis after the test to better view smaller-diameter cell populations.
 - b. Ensure that the minimum concentration (>1 x 10^5 cells/ml) of cells are present in the sample.
 - c. Reduce potentially interfering cell size populations (e.g. peripheral cells, RBCs) via cell lysis, gradient centrifugation, or pre-plating techniques.