

Document # MXV.SYS.005		Effective: 30JUL2021
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Title: Moxi V – System Check Bead Procedure

# MATERIALS REQUIRED

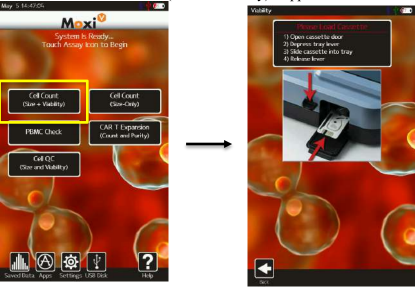
- Moxi V (Gemini Bio-Products Cat #MXV102)
- System Check Beads, Orange/Red Channel (Gemini Bio-Products Cat# MXA028)
- Cassettes: Type S+ (Gemini Bio-Products Cat# MXC030)

# **PREPARATION**

- 1. Ultrasonicate Both Check beads for 60 seconds at room temperature (22.5°C)
- 2. Vortex the beads at the highest speed setting for 30 seconds.
- 3. If the "Fluorescent Gain" setting has been changed, change it back to "Default" (Go to "Settings" from the Home screen. Touch "Fluorescence Gain" field to toggle)

# **RUNNING THE TESTS**

- 1. Mix beads by slowly inverting the bottle 30x initially (5x between runs, if repeats are being performed)
- 2. Turn Moxi V unit on and select "Cell Counts (Size + Viability)" App



- 3. Insert a cassette and wait for the laser cassette alignment to complete.
- 4. When the alignment completes (and prompted "Enter the Sample" at the top left black bar), pipette 60μL bead 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 beads will begin to settle) and the test will automatically run.

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- 1. The results will be automatically gated by the system if the "Auto Gate" setting is on. If not, or to fine-tune the gate locations, follow the steps above (and described below).
  - a. Touch the blue histogram overlay icon at the bottom right of plot, to overlay the size (red, x- axis) and fluorescence (blue, y-axis) histograms
  - b. Adjust the size gates (blue vertical markers) to place it in the "valley" (minima) of the red histogram, to the left of the bead peaks (see image above/middle).
  - c. Toggle to enable fluorescent gating by touching the red gate icon (top right of scatter plot in the image above/right). Place the fluorescent gate just below the upper (Bright) bead population (below the upper peak on the blue histogram)
- 2. With this gating, you can record all relevant stats
  - a. The total concentration is listed at the top of the scatter plot, between the blue/size gates.
  - b. Dark Bead concentration and size is listed in the "Live Cells" black box to the lower/left of the scatter plot.
  - c. Bright Bead concentration and size is listed in the Dead Cells" black box to the lower/right of the scatter plot.
- 3. Verify the reported concentration and size to the expected values on the bottle. The margin of error should be 15% or less for both concentration and size. *Note: For exact lot size and concentration data, please email tech support@orflo.com with the bead lot number to obtain a COA for that lot.*

## FACTORS THAT CAN AFFECT RESULTS

- 1. Temperature of the bead solution or environment can slightly affect the reported diameter. Bead solution and environment should be within 20-25°C.
- 2. Microbial contamination of the bead solution can also affect the reported diameter. Avoid contamination by using a clean pipette tip to aliquot the solution. Store the beads in 2-8°C when not in use.
- 3. Improper mixing can affect the reported concentration. Slow inversion of the bottle after sonicating/vortexing is key to ensuring single-bead suspension.

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Document	#	Revision #	Reason
MXV.SYS.005		002	Updated company information from Orflo to Gemini Bio Simplified protocol instructions, updated system photos, revised CV acceptance criteria

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