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## iGEM Calibration Protocol - Flow Cytometry Cell Size

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In devel.



#### **ABSTRACT**

Conversion from forward scatter to Eµm is not a linear function, so data cannot be converted simply by multiplying with a scaling factor. We thus recommend use of software tools for processing data with size calibration.

This protocol can be combined with bead-based fluorescence calibration.

### MATERIALS

NAME ~	CATALOG # V	VENDOR V	CAS NUMBER $\vee$ RRID $\vee$
Size calibration beads	PPS-6K	SpheroTech	

#### MATERIALS TEXT

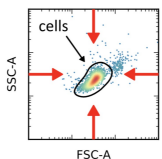
- Experimental samples: transformed per your desired experimental procedure
- Flow cytometer with a forward scatter channel (typically a 488nm excitation and a 488nm/10nm emission filter). Measurements on this channel will be converted to equivalent μm diameter (Εμm).
- TASBE Flow Analytics Analysis software

## BEFORE STARTING

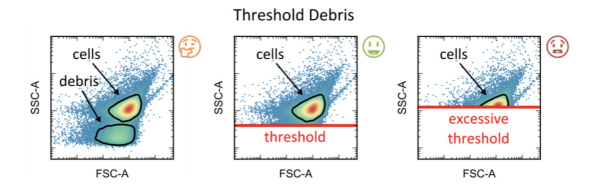
Read through this entire protocol carefully before you start your experiment and prepare any materials you may need. See the "Results" section for an example of a completed data analysis spreadsheet.

- 1 Prepare and culture experimental samples according to your desired experimental procedure.
- Prepare experimental samples as needed for running through your flow cytometer.
- 3 Follow SpheroTech directions for preparation for PPS-6K bead sample.
- ▲ Measure all samples in flow cytometer
  - Using the bead sample, adjust FSC channel voltage so that the top bead peak is below detector saturation.
  - Using the negative process control sample, adjust forward-scatter and side-scatter voltages to place the strong cell peak as close to the center of the detector range as possible (see example illustrated below).

# Center FSC and SSC



• Instrument gating should be set to ensure that no cell events are discarded (see example illustrated below).



- Collect at least 10,000 events per sample.
- 5 Compute the Eµm/a.u. scaling function from bead sample using TASBE Flow Analytics to create a TASBE color model.
- 6 Convert experimental data to Eµm using <u>TASBE Flow Analytics</u> analysis functions.

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