

PRD004 Version 7 - 02/14 PROTOCOL

Catalog #201103A (500 µL)

Cell-ID[™] Intercalator-103Rh 500 µM



WARNING! CHEMICAL HAZARD. Before handling any chemicals, refer to the Material Safety Data Sheet (MSDS) provided by the manufacturer, and observe all relevant precautions.



NOTICE: HIGH CONCENTRATION. Cell-ID Intercalator-103Rh 500 μ M is a highly concentrated metal intercalator solution and must be diluted in accordance with this protocol to avoid early failure of the detector.

Description

Cell-ID Intercalator-103Rh is a cationic nucleic acid intercalator that contains natural abundance Rhodium (¹⁰³Rh) and is used in CyTOF® analysis for either discrimination of dead cells from live cells (if staining is done before cell fixation), or to discriminate single nucleated cells from doublets (if staining is done post-fixation). It is a live cell membrane-impermeable dye.



Note: While dilutions of the 500 μ M stock solution are suggested in the protocols below, the concentration can be titrated for individual cell types and experiments for optimal staining. It is suggested not to exceed 4 μ M intercalator concentration in the staining solution for cell singlet discrimination protocols.

Live/Dead Staining Protocol

- 1 Add Cell-ID Intercalator-103Rh directly into cell culture flask at a dilution of 1:500 and return cells to incubator for 15 minutes.
- 2 If desired count viable cells with Trypan Blue to confirm % viable observed with Cell-ID Intercalator-103Rh staining.
- Proceed to stain cells as usual with MaxPar® metal-conjugated antibodies and, following fixation, with Cell-ID Intercalator-Ir (Fluidigm Cat. 201192A or 201192B) to identify dead cells.

Notes

Cells with high Intercalator-Rh mean values in ¹⁰³Rh versus ¹⁹³Ir plot are considered dead since the membrane impermeable intercalator-Rh would have accumulated only in cells with a compromised membrane, and these cells should be excluded in gating strategies.

• For live/dead staining of thawed cells including PBMC, rest thawed cells for at least one hour before proceeding with addition of Intercalator-Rh in step 1.

Cell Singlet Discrimination—Staining Protocol A

- 1 Before intercalating, cells must be fixed.
 - If fixed with methanol, wash cells with PBS (without Ca²⁺ or Mg²⁺) before proceeding.
 - Cells may be used directly if fixed with formaldehyde (3.7%, 30min, RT).
- 2 Dilute Cell-ID Intercalator-103Rh 1:500 with PBS (without Ca²⁺ or Mg²⁺).
- 3 Use 0.5mL of working solution per 1x10⁶ cells/ tube.
- 4 Incubate 15-20 mins at room temperature.
- 5 Wash cells with 2 mL PBS (without Ca^{2+} or Mg^{2+}) per tube. Repeat once.

Cell Singlet Discrimination—Staining Protocol B (for use with the MaxPar® Cell Surface Staining Protocol)

- 1 After cell staining is complete, prepare 1 ml of cell intercalation solution for each sample by diluting Cell-ID Intercalator-103Rh 1:1000 into MaxPar® Fix and Perm Buffer (Fluidigm Cat. 201067) and mix by vortexing.
- 2 Add 1 ml of the intercalation solution prepared in step 1 to each tube and gently vortex. Incubate for 1 hour at room temperature or leave overnight at 4 °C.



Note: Cells can be left at 4 °C in the intercalation solution up to 48 hours.

- 3 Wash cells by adding 2 ml of MaxPar® Cell Staining Buffer (Fluidigm Cat. 201068), centrifuge and discard supernatant by aspiration.
- 4 Repeat for a total of two washes with MaxPar Cell Staining Buffer.
- 5 Wash cells with 2 ml of MaxPar® Water (Fluidigm Cat. 201069), centrifuge and discard supernatant by aspiration.
- 6 Leave cells pelleted until ready to run on CyTOF. Immediately prior to CyTOF data acquisition, adjust cell concentration to $2.5-5 \times 10^5/\text{ml}$ with MaxPar Water and filter cells into cell strainer cap tubes.
- 7 Acquire data on CyTOF.

Cell Singlet Discrimination—Staining Protocol B (for use with the MaxPar® Cell Surface Staining Protocol)

Technical Support

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