

Aquarray GmbH Hermann-von-Helmholtz-Platz 6 76344 Eggenstein - Leopoldshafen Germany

## Protocol

## Staining with Hoechst and Propidium Iodide (PI) with I-DOT Mini cell dispenser

Preparation of staining solution:

Hoechst 33343: ThermoScientific, #62248; final dilution 1:10000 (excitation 355nm laser) Propidium Iodide: ThermoScientific # BMS500PI; final dilution 1:3000 (excitation 535nm laser) Calcein: ThermoScientific #C1430; final dilution 1:2000 (excitation 945 nm laser)

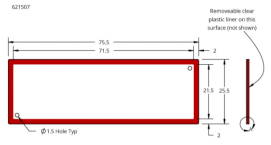
For one 672 DMA 33,6  $\mu L$  of staining solution (exact calculation) would be required for each slide.

Prepared dilutions have to be 4x more concentrated as final dilution: Hoechst 1:2500 PI 1:750 Calcein 1:500

Material: DMA with 672 spots SecureSeal<sup>™</sup> Hybridization Chamber (21,5x71,5x0,8 mm): Grace Bio-Labs (#621507)

Preparing a final volume 200 μL:

- 1. Add 0,08 μl Hoechst and 0,26 μL Propidium Iodide to 199 μL PBS.
- 2. Print 50 nL of staining solution on each droplet (150 nL culture medium containing cells) with protocol "staining with Hoechst-Calcein-PI on 672 DMA".
- 3. After addition of the staining solution put the DMA back the into Petri dish with humidified lids and incubate for 15 min in the cell incubator at 37°C.
- 4. Meanwhile prepare a SecureSeal chamber



(image taken from Grace Biolabs)



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- 5. Cut two small squares of a transparent adhesive tape to close the two holes ( $\emptyset$  1,5mm) on the left lower and right upper part.
- 6. Remove the DMA carefully from the Petri dish with humidified lids and put the sticky chamber quickly on the DMA to avoid drying of the cells/droplets. Ensure that the sticky chamber is well fixed on the DMA. Transfer to the microscope for imaging.