

Protocol for

PicoWells In **Chamber Slide**

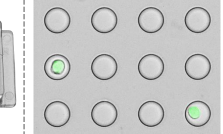
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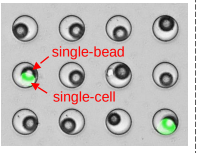
Plasma Treatment

Take the lid off and put the Pipette 1-2 mL of 30 mg/mL Place a Disk Magnet at the Add cold cell lysis buffer onto Remove the Disk Magnet. Place a PCR tube containing PicoWells In Chamber Slide in a BSA/PBS solution into each well; bottom of the picowells. Add the surface of picowells; Place a Cylinder Magnet on PBS on top of a strong Cube |plasma cleaner. Treat the surface | wash away unabsorbed BSA | the barcoded magnetic bead | incubate for 10 min; add fresh | the surface of picowells, and | Magnet. Insert the Cylinder of picowells with plasma for 1-2¹ after 1 h of incubation; add cell $\frac{1}{2}$ suspension ($\geq 10^6/mL$) and settle $\frac{1}{2}$ lysis buffer (optional). The Disk $\frac{1}{2}$ move the Cylinder Magnet Containing magnetic min to change the surface suspension properties of PDMS picowells down for 10 min (~10% picowells to capture single increase being hydrophobic from hydrophilic. wash away uncaptured cells.

Single-Cell Capture



Single-Bead Capture



Single-Cell Lysis

Beads Collection

(10⁵/mL); settle down for 10 min to allow most Magnet at the bottom will across the entire surface of beads with captured mRNA to picowells will have single cells); beads. Gently wash away the magnetic beads within uncaptured beads.

stability the picowells.

of picowells to collect mRNA.

Beads Transfer



the 1 into the PBS: stronaly stir up and the | magnetic beads with captured | down to transfer beads to the PCR tube.

Applications

- Single-Cell/Bead Isolation
- Single-Cell/Bead Imaging
- Sinale-Cell RNA-Sequencing

The Disk Magnet, Cylinder Magnet, and Cube Magnet need to be ordered separately.

