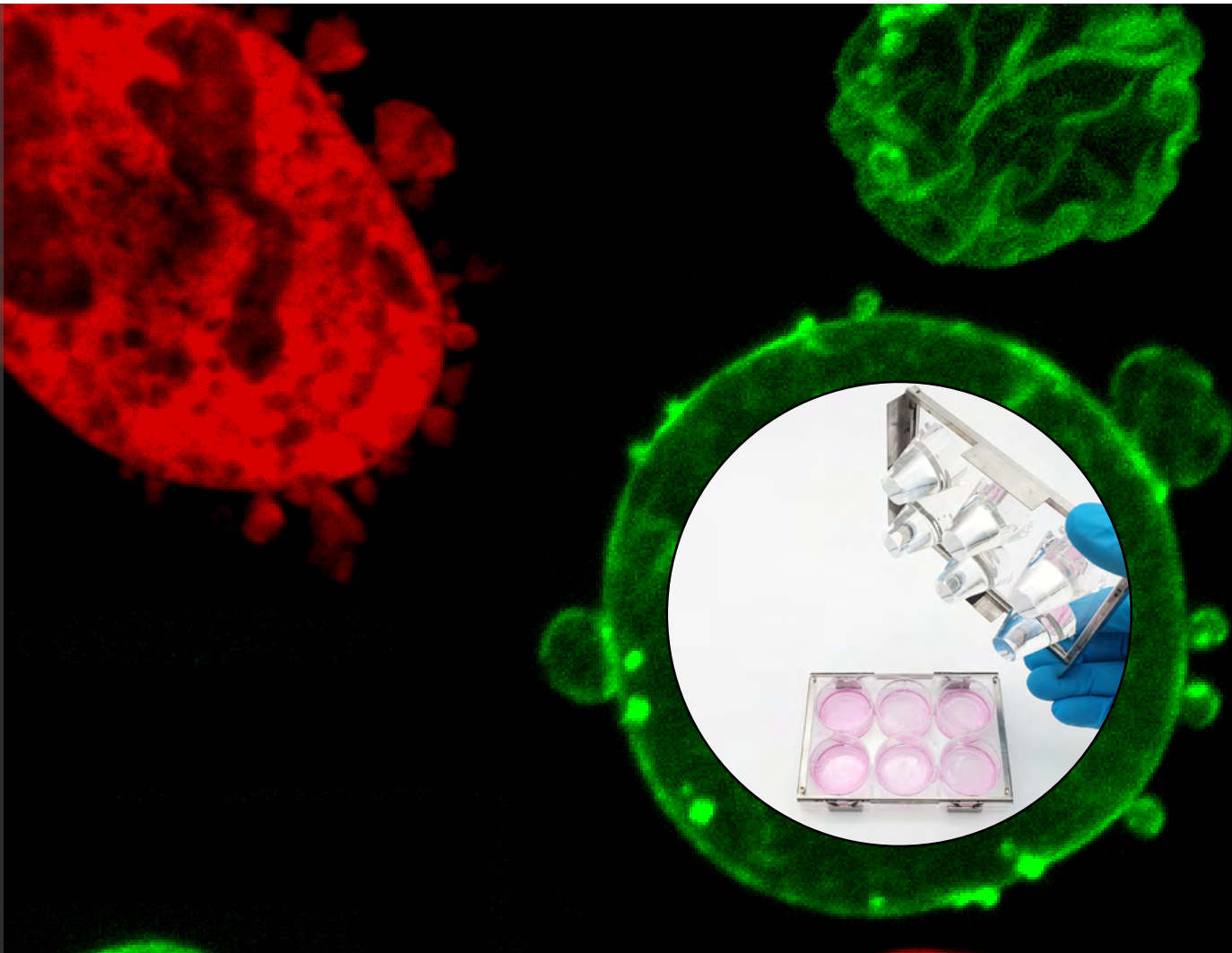


4Dcell static 6-well cell confiner

User protocol- CSOW 620

USER GUIDE



How to use 4Dcell static 6-well confiner 'CSOW 620'

Support

A video is available online to understand each step to handle perfectly your CSOW 620:
www.4dcell.com/cell-culture-systems/cell-confinement/static-cell-confiner

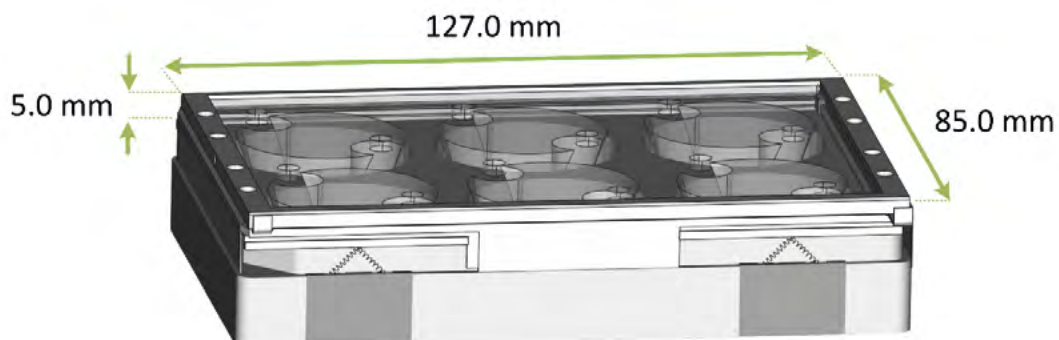
Material Included

4DCell CSOW 620
Confinement slides/coverslips with PDMS micropillars (12 units)
Confinement PDMS pistons (12 units)
Small blade
Tape
Caps for closing the holes on the lid

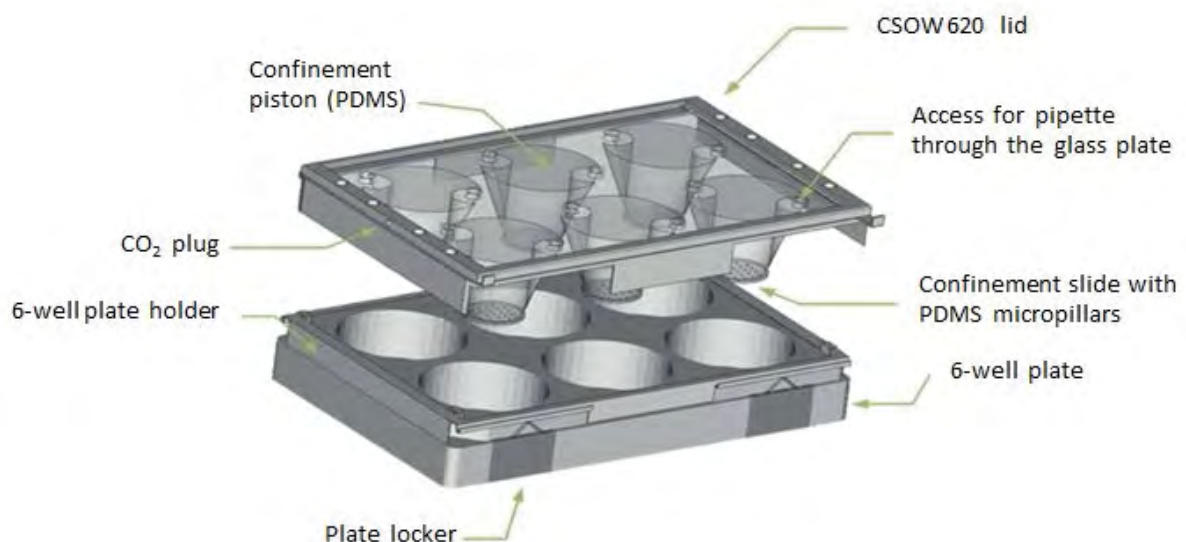
Other recommended material

70% ethanol
Tweezers
6-well plates (Glass Bottom MatTek)

CSOW 620



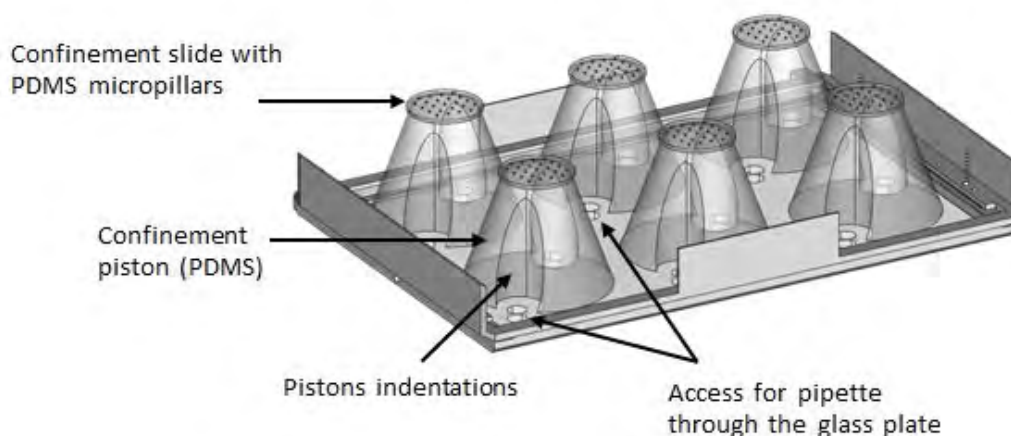
CSOW 620 parts



A. Handling and assembling the confiner

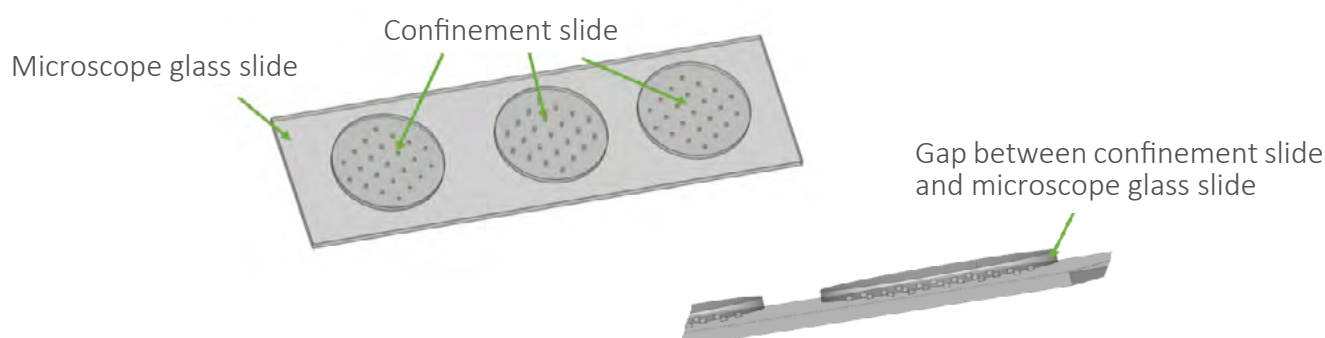
Note: The different items of the confiner kit are shipped in sterile bags. To avoid contamination, only manipulate the items using aseptic techniques in a laminar flow hood.

1. Clean the two components of the CSOW 620 confiner, the lid (top) and holder (bottom), using absorbant paper and 70 % ethanol.
2. Clean the pistons using 70% ethanol. If the PDMS pistons have particles of dust these can hinder the good attachment to the glass and interfere with imaging. In this case, remove the dust particles with the tape provided and rinse once again with 70% ethanol.
3. Attach the pistons to the CSOW 620 lid making sure the indentations of the pistons are aligned with the holes in the glass plate.

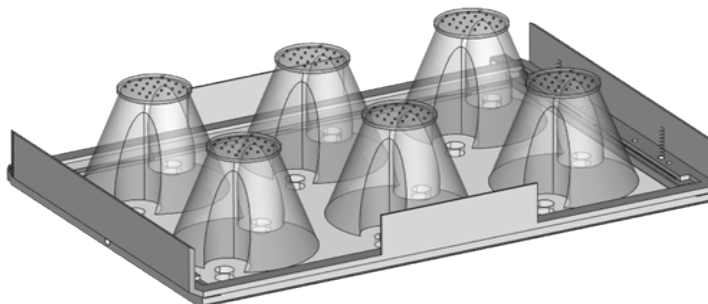


The PDMS is usually sticky, enabling it to be attached to glass very easily. The holes in the lid (access for pipette through the glass plate) can be used to pipette media, aspirate it, etc. The holes can be closed with the white caps provided in the kit. If you find it difficult to remove the caps, please use tweezers to do so.

4. Confinement slides are individually shipped according to confinement height with the pdms pillars facing down. You can use a blade to gently remove the confinement slide (be careful not to break it). Place the blade in the small gap between the confinement slide and the microscope glass slide. Afterwards, tilt the blade so that the confinement slide detaches from the microscope glass slide.

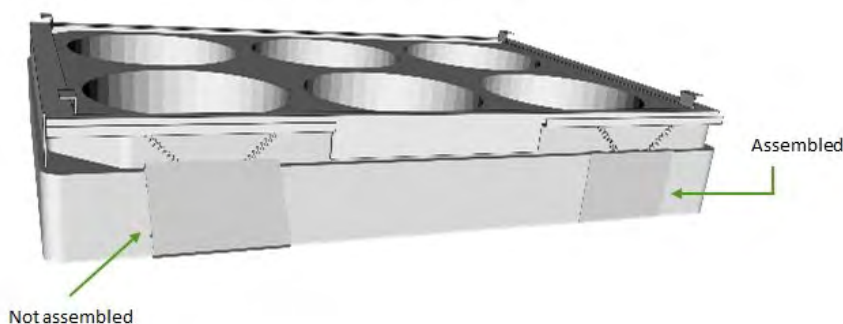


5. Place the confinement slide on the PDMS piston with the pillars facing up as depicted in the figure below. Make sure the slides are well centered on the pistons to avoid breaking the slides once in contact with the inner circle on the bottom of the plate. Avoid creating air bubbles between the PDMS piston and the glass coverslip.

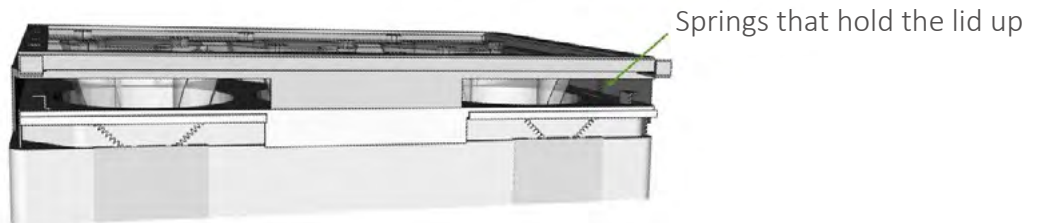


Tip for cell culture (section B of this protocol, page 6): Incubate the pistons and the confinement slides in culture medium to equilibrate the PDMS for at least one hour. To do this you can put the CSOW 620 lid containing the confinement pistons + the slides on a regular 6-well plate. The springs attached to the inside of the lid will prevent the confinement slides from touching the bottom of the plate. Note that PDMS absorbs small hydrophobic molecules from the medium. Therefore, if drugs are used in the experiment, these drugs should to be included in the medium in this incubation step. You don't need to lock the confiner at this stage (resting position).

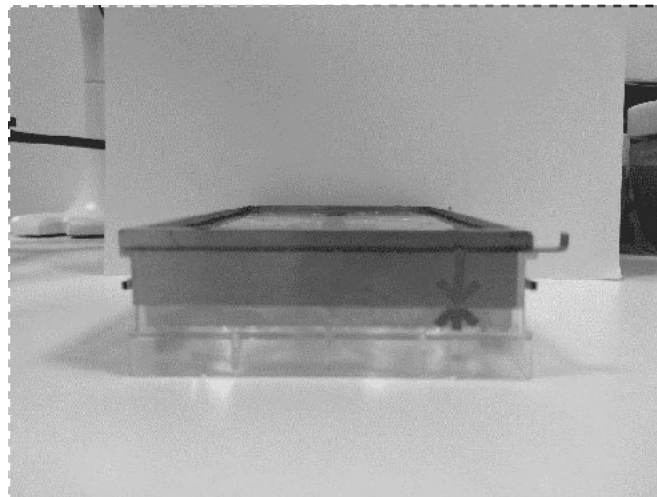
6. To assemble the bottom part of the confiner to the 6-well plate, gently pull the metal pads down so that they grab on to the bottom of the plate. The pads should be pulled one by one, starting with one pad from one side, then the next pad from the diagonally opposite side, then the last two in the same manner (never pull the two pads from the same side one after the other).



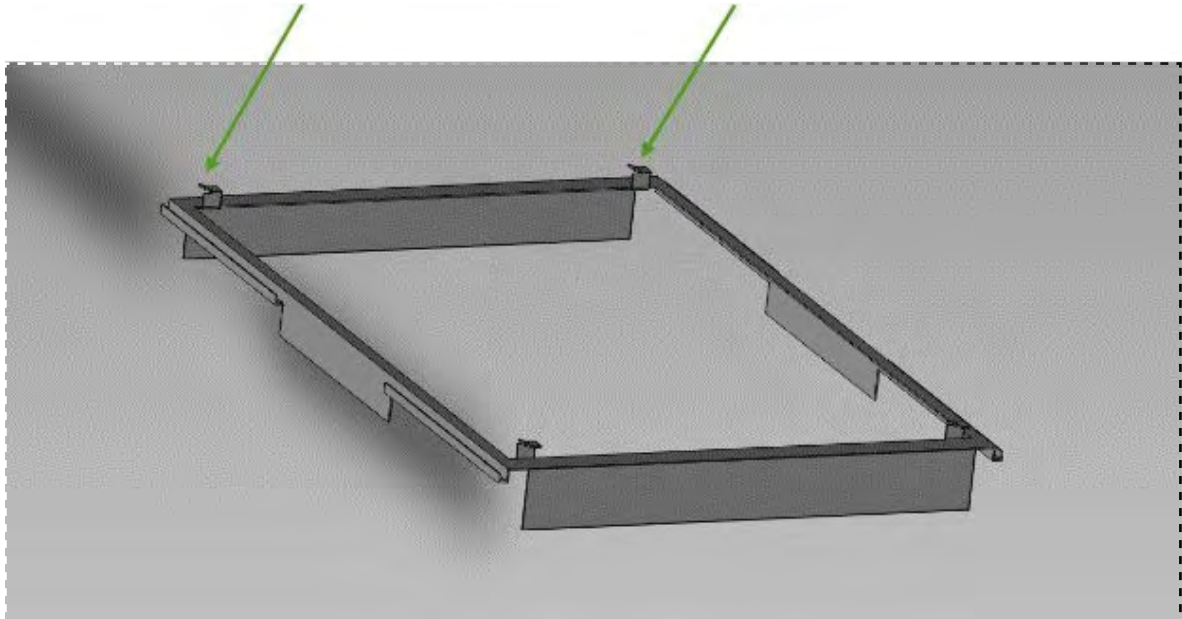
7. Place the lid on the top of the assembled holder. Allow it to be in the resting position, in which the springs hold the lid up.



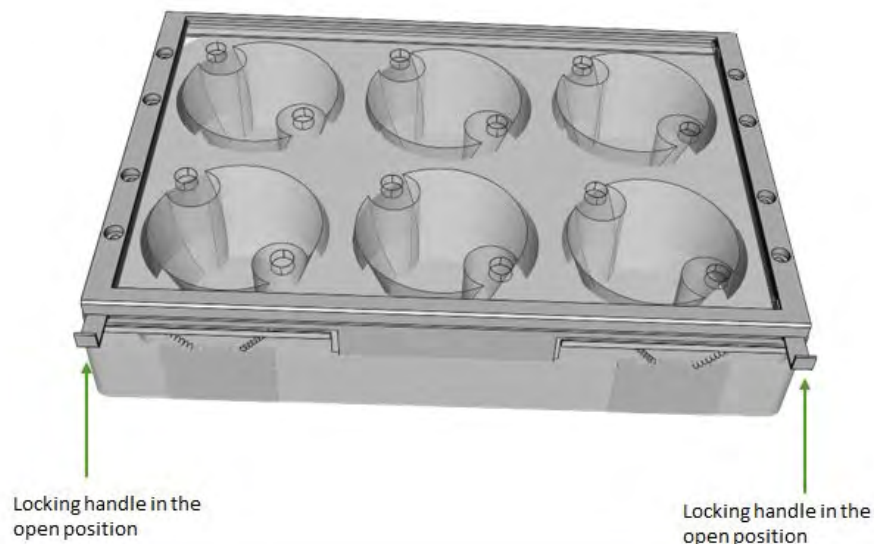
Be sure to align the marks:



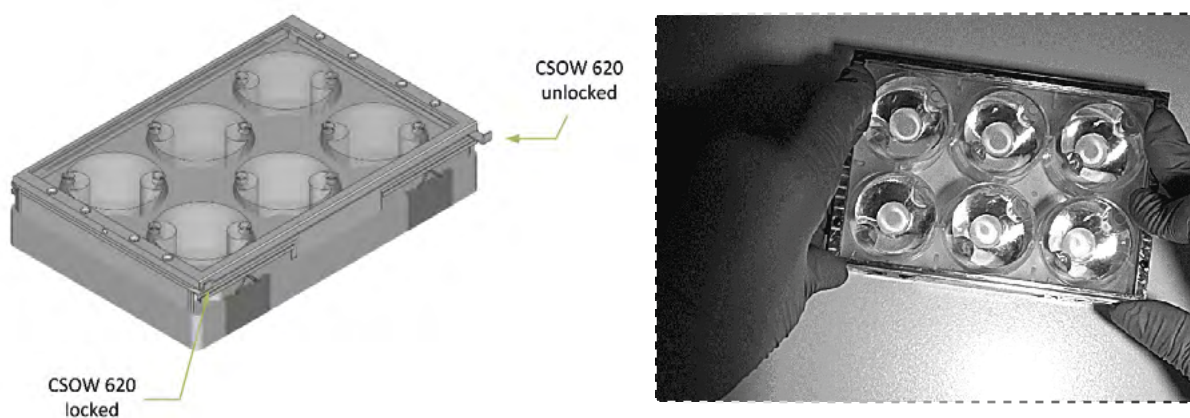
The tilted part of the plate holder should be facing the front when you assemble the lid and the holder.



Please be careful because the locking only works if the plate is assembled in the correct orientation. Make sure that the locking handles are open:



8. To lock the confiner, press the lid down on both left and right sides (as shown below, right image) and simultaneously lock the two handles by pushing them in. If you feel resistance, make sure the plate is assembled in the correct orientation as described in step 7.



9. To unlock the confiner, gently pull the locking handles out and let the lid come up. If it does not come up naturally, gently move it upwards.

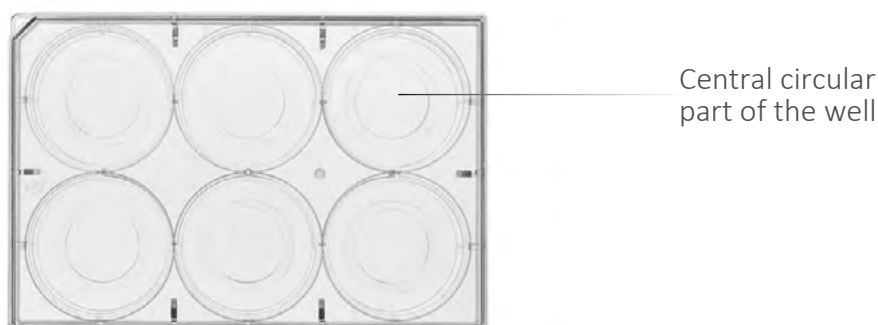
10. If you have issues removing the holding pads from the 6-well plate (when needing to change the plate), you can use tweezers to help you remove the metal pads.

B. Handling and assembling the confiner with cells

11. Make sure the confiner with the pistons are properly cleaned (steps 1 to 2).

12. Proceed to steps 4 to 6.

13. To ensure cell adherence, the 6-well plate can be coated with fibronectin or other ECM matrix of your choice(1). 500 μ L of fibronectine is enough to cover the central circular part of the well. Incubate for 1 hour at room temperature.



14. Remove the fibronectin solution.

15. Ressuspend the cells at the desired density in 500 μ L of cell culture medium and pipette into the wells. Let the cells adhere to the substrate (the timing depends on the cell type)*.

16. Place the confiner lid on top (after incubation in another 6-well plate for 1 hour at 37°C, 5% CO₂, as described in step 7)**.

17. To confine the cells, press the lid down on both left and right sides (as shown above in step 8) and simultaneously lock the two handles. If you feel resistance, make sure the plate is assembled in the correct orientation as described in step 7.

18. Add 2 mL of medium through the holes on the lid that give access to the bottom of the plate.

19. To release the confinement, gently pull the locking handles out and let the lid come up. If it does not come up naturally, gently move it upwards (see images on step 8).

**We recommend a volume of 500 μ L of cell suspension if you want to retrieve the cells for gene analysis after the confinement. Alternatively, if you don't need to remove the cells for later analysis, you can resuspend them in a bigger volume of medium. In that case, the volume will take the whole well and not only the central circular part.*

***Incubate the pistons and the confinement slides in culture medium to equilibrate the PDMS for at least one hour. To do this you can put the CSOW 620 lid containing the confinement pistons + the slides on a regular 6-well plate. The springs attached to the inside of the lid will prevent the confinement slides from touching the bottom of the plate.*

Note that PDMS absorbs small hydrophobic molecules from the medium. Therefore, if drugs are used in the experiment, these drugs should to be included in the medium in this incubation step. You don't need to lock the confiner at this stage (resting position).

(1) With the glass bottom MatTek 6-well plates, 500 μ L is enough volume to coat the inner circle of the well. References: Liu *et al.*, 2015, Cell 160, 659–672