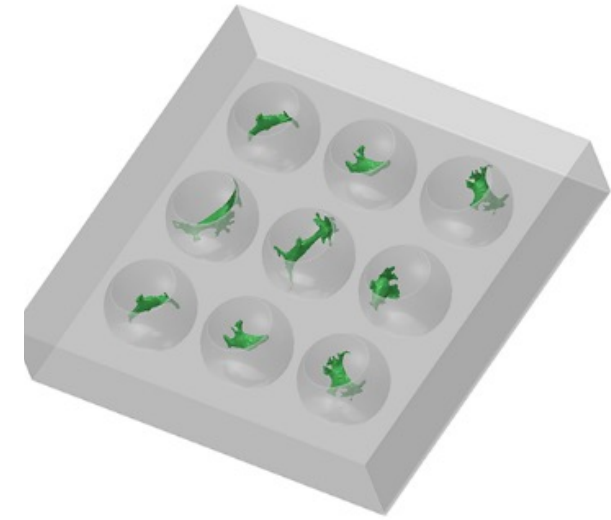


# Spherical Microwell Arrays for Mesenchymal Stem Cell Cultures



KUO, Hsu-Ting

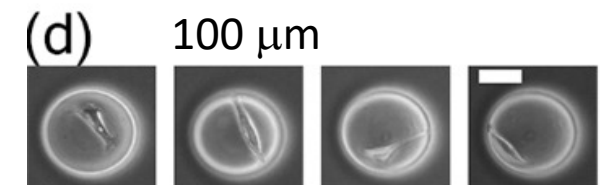
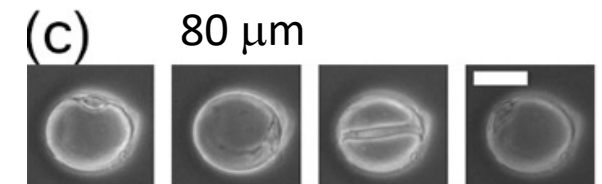
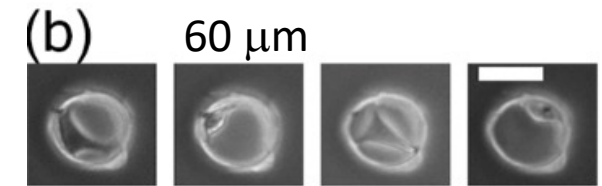
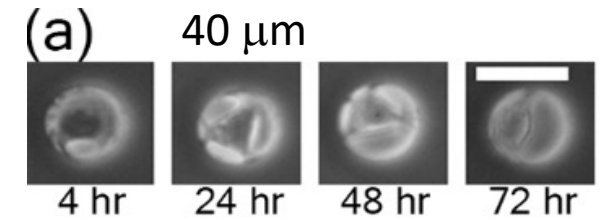
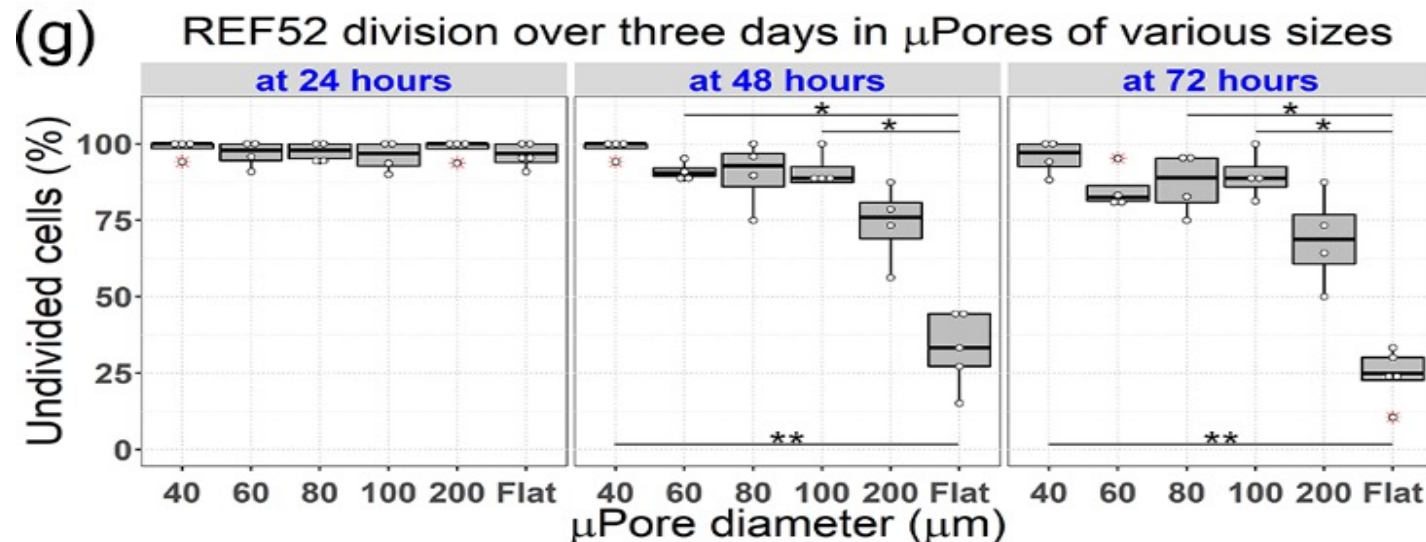
Supervisor: Keng-hui Lin

Institute of physics

# Motivation

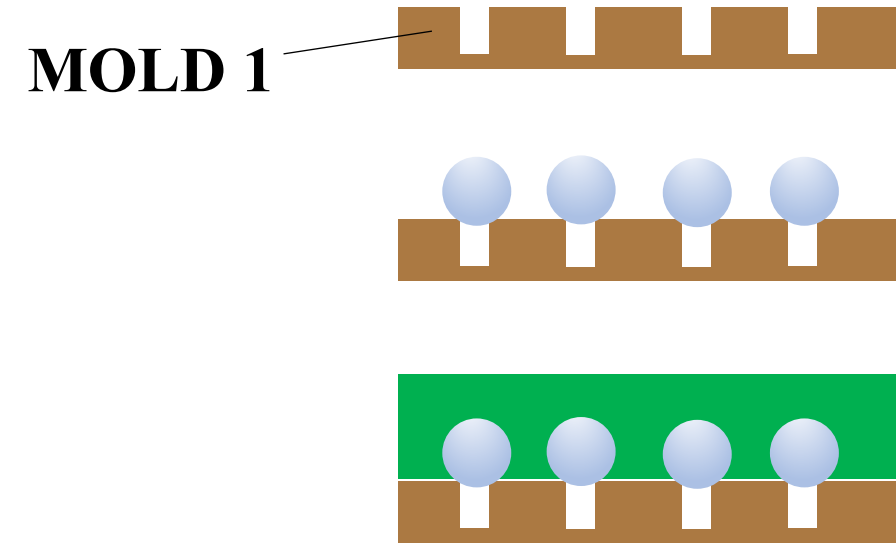
In Lin's lab, the former postdoc, Jonny Huang, developed a spherical microwell arrays as a novel 3D culture method.

Data shows that the self-renewal ability of REF52 cell decreases under 3D confinement



Rat embryonic fibroblast REF52

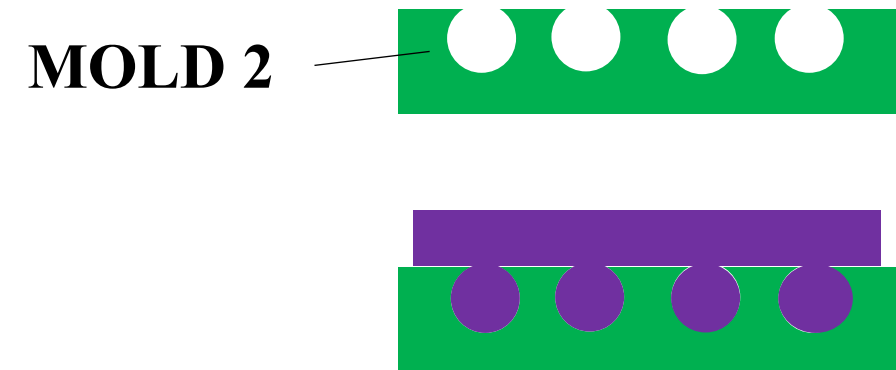
# Original protocol



**Step 1:** Create an array of micrometric cylindrical holes on PDMS

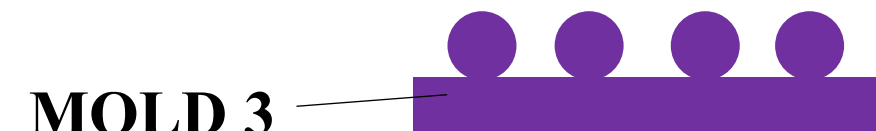
**Step 2:** Manually placed glass microspheres one by one onto the cylindrical holes

**Step 3:** Pour fast curing silicone on PDMS

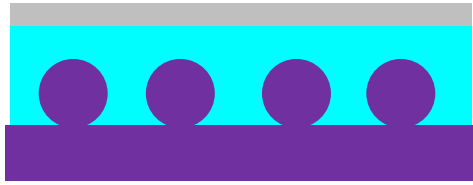


**Step 4:** Peel off the microspheres

**Step 5:** The second silicone template is applied with an off-the-shelf two-part epoxy resin and evenly appended to a glass-slide



→ **Step 6:** Obtain monolithic epoxy microsphere mold



**Step 7:** Using the mold to make PA gel



**Step 8:** Sterilization process (UV light)

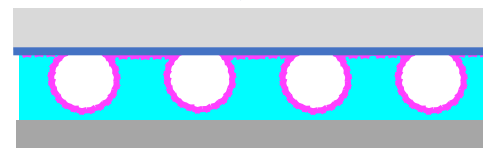


**Step 9:** Protein coating on the PA gel

 ECM proteins

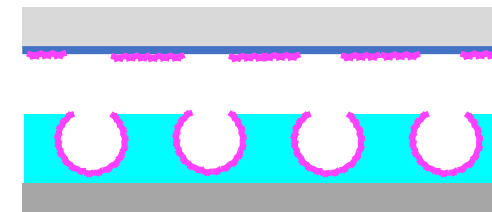
**Step 10:** Peel off extra protein

Pressure 



aldehyde-coated slide

overnight



top ECM removal

# Drawback of the Original Protocol



TIME CONSUMING:  
AROUND 3 DAYS



10 STEPS  
3 DIFFERENT MOLDS  
INVOLVED



LABOR CONSUMING

# Objective

Our goal is to:

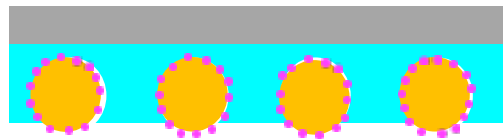
- **Simplify** the protocol of making 3D cell cultured platform
- Observe the change in self-renewal ability of **mesenchymal stem cells (MSC)** under 3D confinement.



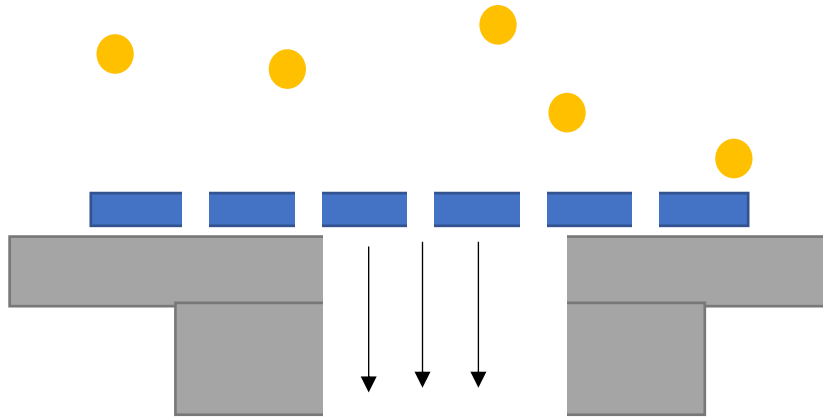
# ➤ Simplifying steps of making the 3D cell cultured platform

1. Simplify the procedure of beads placing
2. Avoid the step of peeling off extra protein

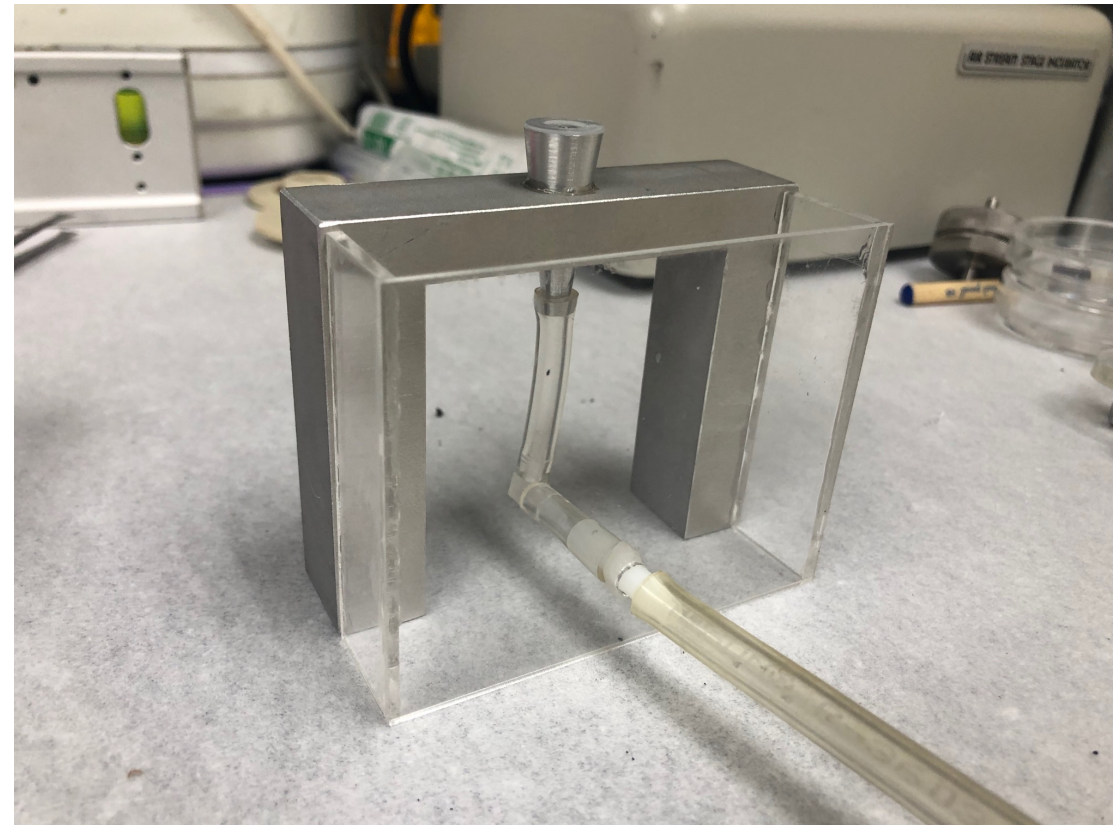
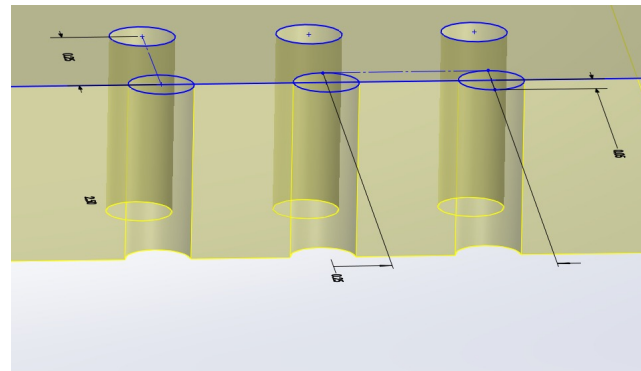
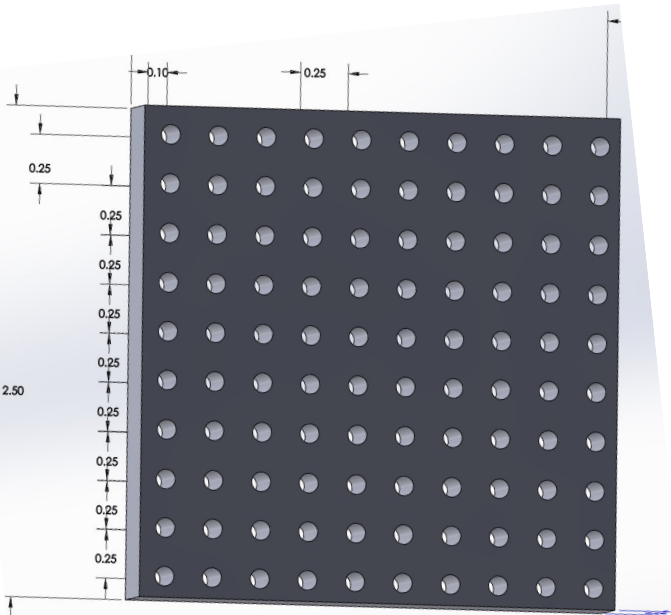
 Microspheres coating with ECM proteins



# Placing beads on an array of through holes



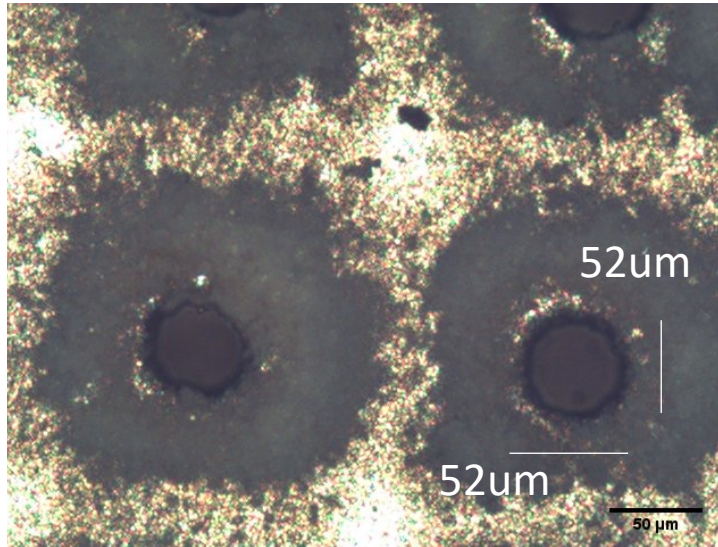
To achieve this purpose, I designed a chuck to hold the hole arrays.



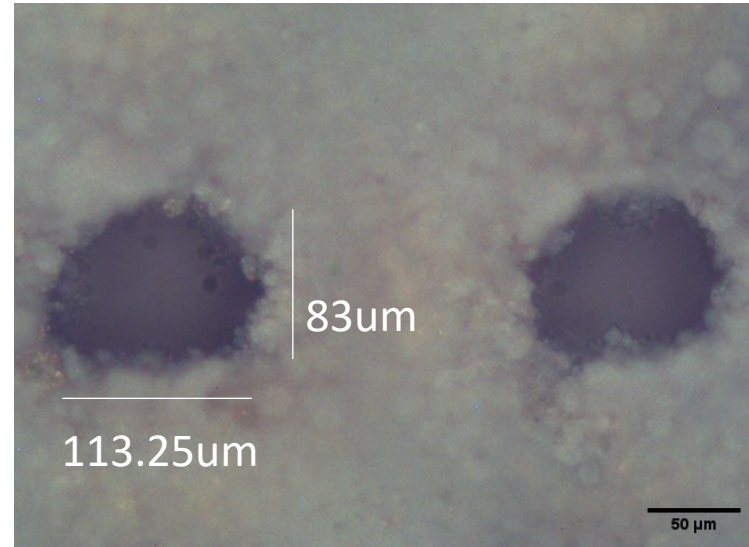


# Making Hole Arrays

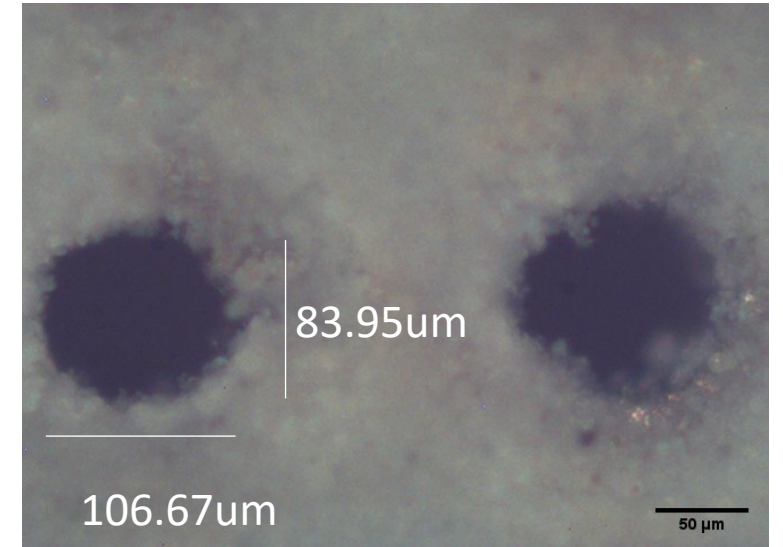
Custom-made by laser hole drilling.



2'' si wafer  
Thickness: 275um



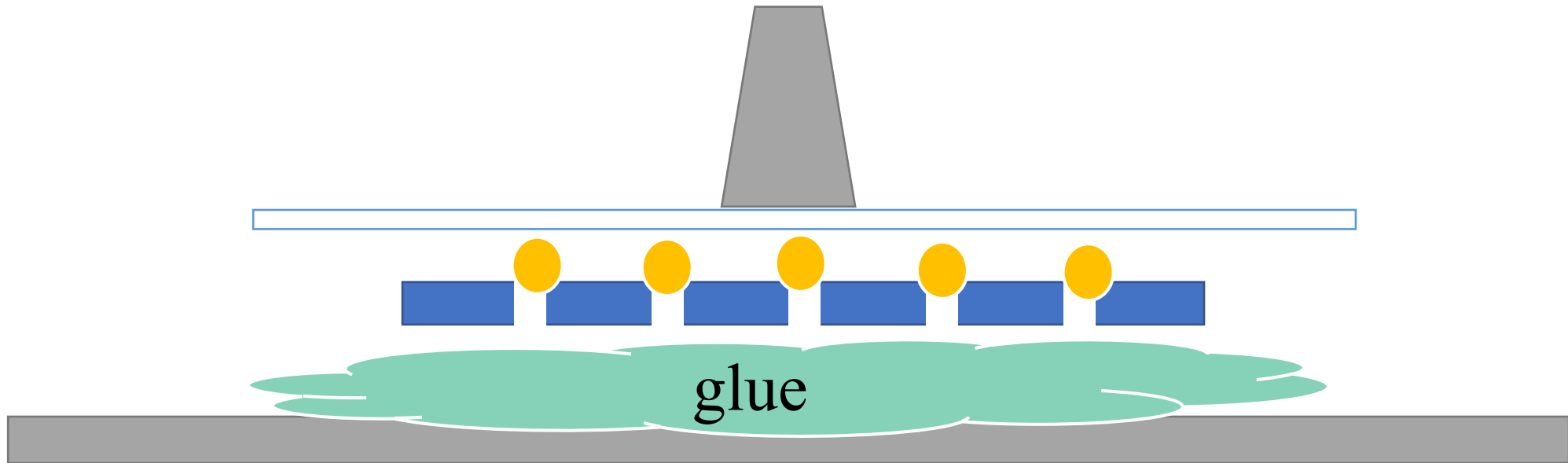
4'' si wafer  
Thickness: 400um



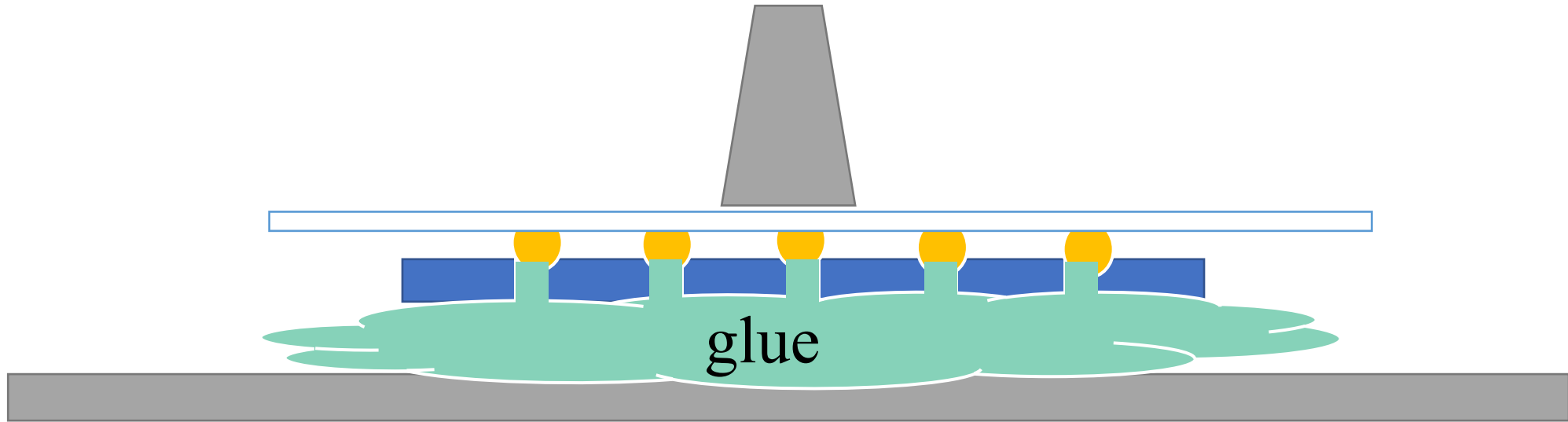
5'' si wafer  
Thickness: 500um

**Problem:** Non-spherical holes. The vacuum leaks when the spherical spheres sit on top.

Solution:  
Use glue to keep beads in place

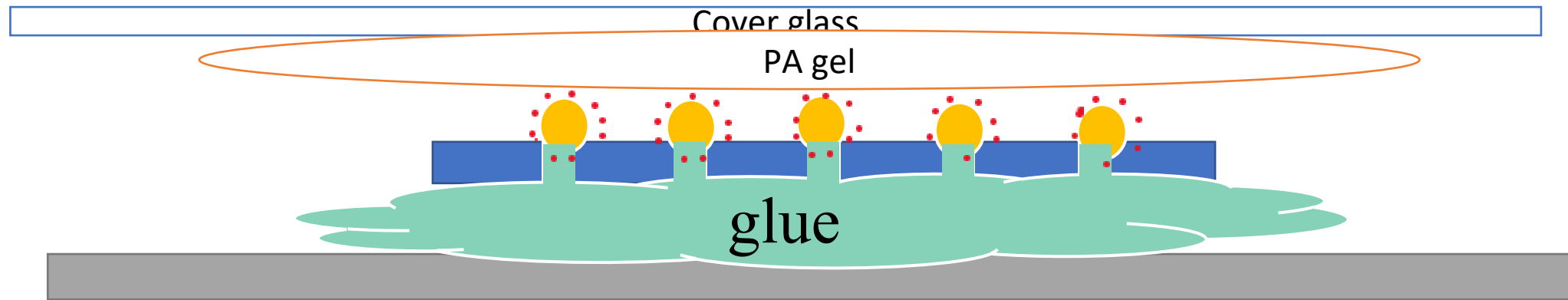


Solution:  
Use glue to keep beads in place



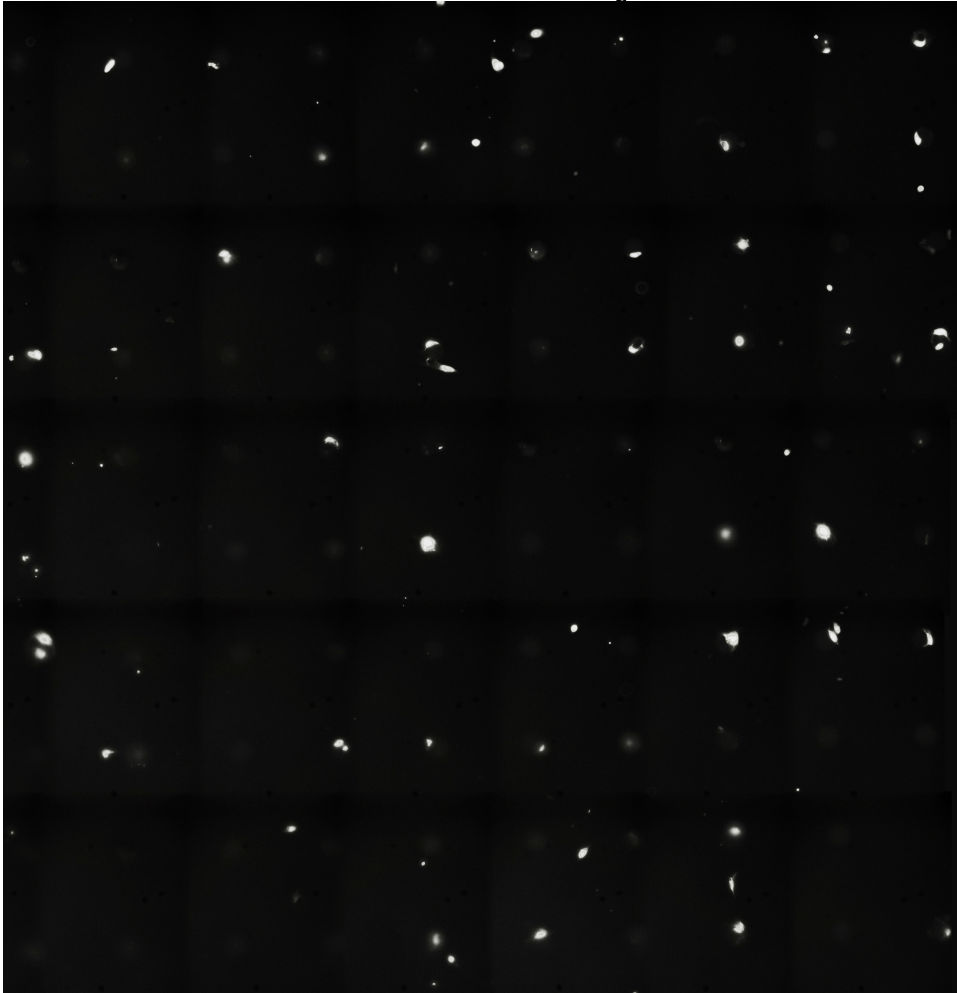
# ➤ Avoid the step of peeling off protein

1. Coat protein on beads beforehand



# MSC grown in 60 mm pores

2019/08/15 Day0



Single cell in pores undivided :  
N: 38 out of 38 → 100%

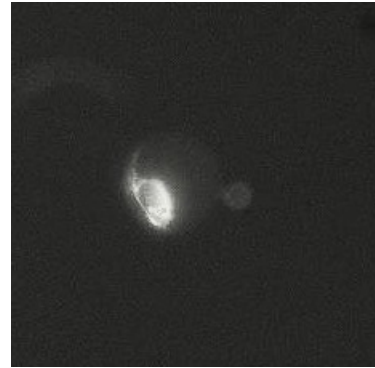
2019/08/19 Day3



20x TRITC Fluorescent dye: CellTracker

# MSC grown in 60 mm pores

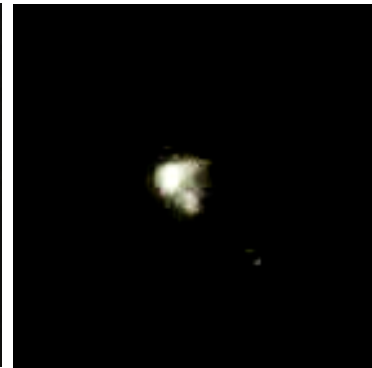
(a)



Day 0

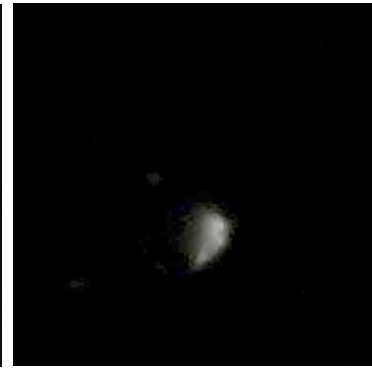
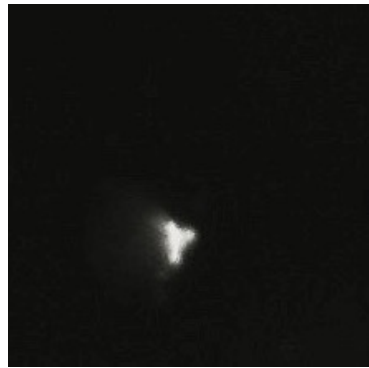
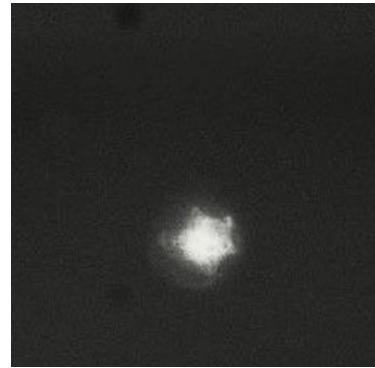


Day 3

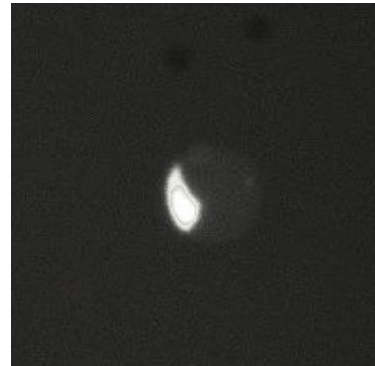


Day 5

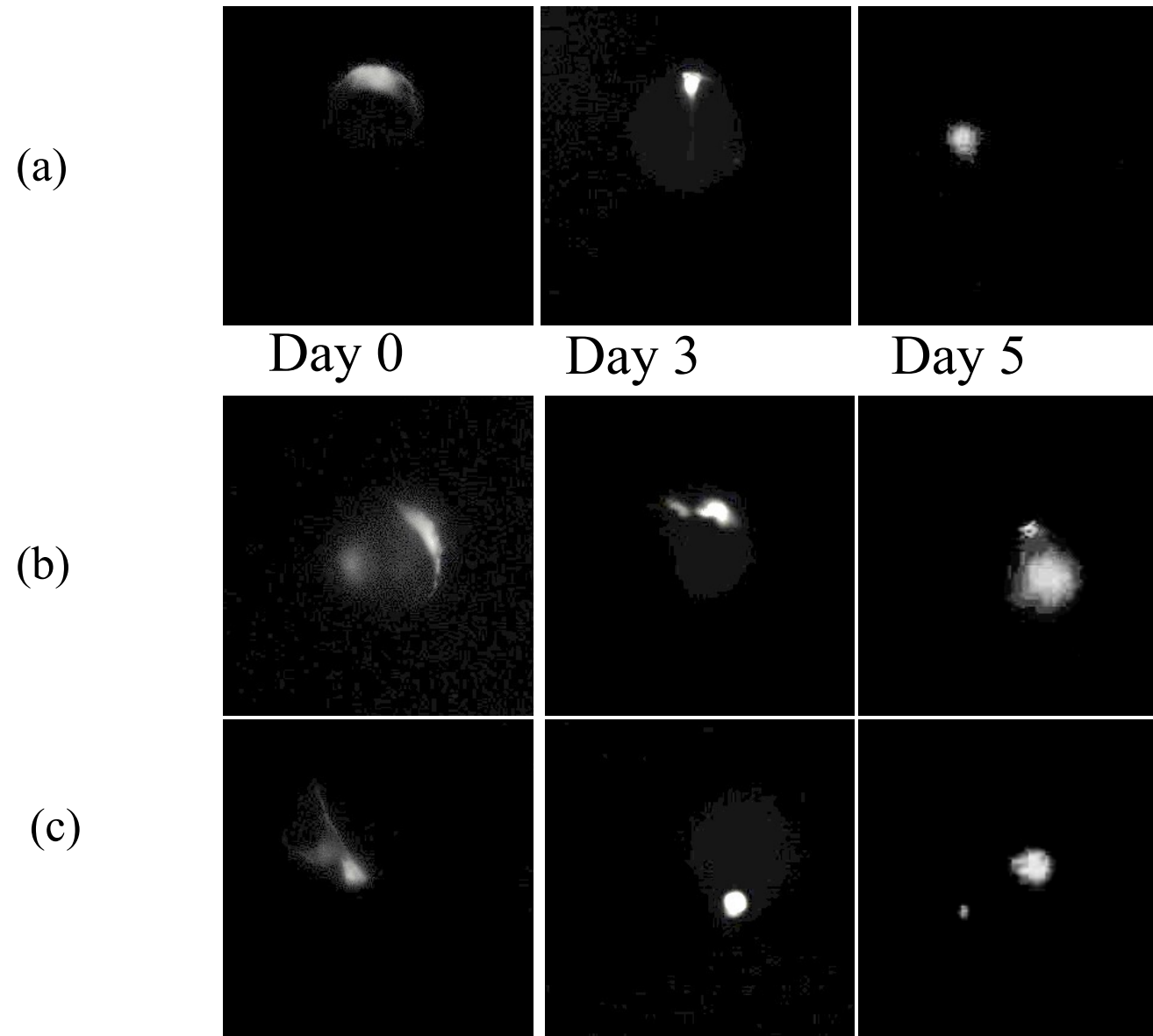
(b)



(c)



# MSC grown in 100 $\mu\text{m}$ pores



# Conclusion



It is possible to use the new protocol to replace the old method.



We also observe the cell cycle arrest of mesenchymal stem cells in spherical microwell arrays.



Questions?

Thank you!