PROTOCOL VitroGel[®] MSC



CAT NO. VHM03, VHM03S

RECOMMENDED MATERIALS AND REAGENTS

- VitroGel[®] MSC (Cat# VHM03)
- Cells
- Cell culture medium
- Conical tubes (15 mL or 50 mL)
- Micropipette; low retention pipette tips
- Centrifuge
- Cell culture plate

3D Cell Culture Protocol

- 1. Bring VitroGel MSC to room temperature or warm at 37°C.
- 2. Prepare the MSC suspension in the culture medium.
 - Recommended cell concentration > 0.8×10^6 cells/mL.
 - **Optional:** if culture medium contains critical supplement (e.g. 2% Human Platelet Lysate (HPL), prepare cell suspension with 3X supplement (e.g. 6% HPL).
- 3. Add 1mL VitroGel MSC to 500 µL cell suspension and gently pipette up and down 5-10 times to mix thoroughly. (Keep VitroGel and cell suspension at 2:1 v/v mixing ratio).
- 4. Transfer the hydrogel mixture to a well plate. Gently tilt/swirl the well plate to ensure there is an even covering on the bottom of each well. The recommended volume of hydrogel for well plates is list below.

	6 well plate	12 well plate	24 well plate	48 well plate	96 well plate
Volume per well	1200 μL	600 μL	300 μL	150 μL	50 μL

- 5. Wait 10-15 min at room temperature for a soft gel formation.
- (Note: During the hydrogel forming process, do not disrupt the hydrogel by tilting or shaking the well plate).
 6. Carefully add additional medium to cover the hydrogel. The recommended volume of cover medium for well plates is listed below

	6 well plate	12 well plate	24 well plate	48 well plate	96 well plate
Volume per well	1200 μL	600 μL	300 μL	150 μL	50 μL

Place the well plate in an incubator and change the cover medium every 48 hours.
 (Note: We recommend to only change 50-80% of the top medium without disturbing the hydrogel).



Hydrogel-Cell Bead Protocol

- 1. Bring VitroGel MSC to room temperature or warm at 37°C.
- 2. Prepare the MSC suspension in the culture medium.
 - Recommended cell concentration > 0.8×10^6 cells/mL.
 - <u>Optional:</u> if culture medium contains critical supplement (e.g. 2% Human Platelet Lysate (HPL)), prepare cell suspension with 3X supplement (e.g. 6% HPL).
- 3. Add 1mL VitroGel MSC to 500 µL cell suspension and gently pipette up and down 5-10 times to mix thoroughly. (Keep VitroGel and cell suspension at 2:1 v/v mixing ratio).
- 4. Add cell culture medium to the well plate. The recommended volume of hydrogel for well plates is list below.

	6 well plate	12 well plate	24 well plate	48 well plate	96 well plate
Volume per well	3000 µL	1500 μL	750 μL	300 μL	100 µL

5. Using a pipettor with a 100 μL tip, carefully pipette the hydrogel-cell mixture into the well plate as droplets. (roughly 5-10 droplets per 100μL of hydrogel-cell mixture.

Optional: control the final size of hydrogel-cell beads by adjusting the volume of the droplets: for small beads, 1-5µL per droplet, for large beads, 20-50µL per droplet).

<u>**Tip:**</u> Press the pipette plunger to create a droplet on the pipette tip, lower the pipette tip to release the droplet by contacting the surface of culture medium.

Place the well plate in an incubator and change the medium every 48-72 hours.
 (Note: We recommend to only change 50-80% of the top medium without disturbing the hydrogel beads).

2D Hydrogel Coating Protocol

- 1. Bring VitroGel MSC to room temperature or warm at 37°C.
- 2. Add 1mL VitroGel MSC to 500µL cell culture medium and gently pipette up and down 5-10 times to mix thoroughly.

(Note: Keep VitroGel and cell medium at 2:1 v/v mixing ratio.

Optional: If culture medium contains critical supplement (e.g. 2% Human Platelet Lysate (HPL)), prepare culture medium with 3X supplement (e.g. 6% HPL) to mix with VitroGel MSC to get 1X final concentration of supplement).

3. Transfer the hydrogel mixture to a well plate. Gently tilt/swirl the well plate to ensure there is an even covering on the bottom of each well. The recommended volume of hydrogel for well plates is listed below.

	6 well plate	12 well plate	24 well plate	48 well plate	96 well plate
Volume per well	1200 μL	600 μL	300 µL	150 μL	50 µL

4. Wait 10-15 min at room temperature for a soft gel formation.

(Note: During the hydrogel forming process, do not disrupt the hydrogel by tilting or shaking the well plate).
5. Carefully add medium with cells on top of hydrogel (Recommend cell concentration of 5 x 10⁵ cells/mL). The recommended volume of cell medium for well plates is listed below.

	6 well plate	12 well plate	24 well plate	48 well plate	96 well plate
Volume per well	1200 μL	600 μL	300 μL	150 μL	50 µL

6. Wait 10-15 min at room temperature for a soft gel formation.

(Note: During the hydrogel forming process, do not disrupt the hydrogel by tilting or shaking the well plate).



Protocol for Cell Recovery from VitroGel MSC

- For 3D cell culture and 2D hydrogel coating, refer to **Protocol-1** of the VitroGel Cell Recovery Solution Protocol.
- For hydrogel-cell bead culture, refer to Protocol-2 of the VitroGel Cell Recovery Solution Protocol."

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