

# Ready-To-Use 3D Human Blood Brain Barrier Spheroid Kit

SP3D-HBBBS Cat. #SP3D-8738

## **Product Description**

The blood brain barrier (BBB) is a specialized capillary bed that separates the brain from the circulatory system and protects the brain from most pathogens [1]. Endothelial tight junctions supported by pericytes and astrocytes are primarily responsible for the highly selective nature of the BBB, restricting the passage of numerous solutes, most antibodies, and some antibiotics [2]. As such, efforts to understand the mechanisms underlying BBB integrity have been critical to developing techniques that are able to penetrate the BBB to deliver therapeutic or diagnostic molecules to the brain. Due to the complexities of the BBB, it is difficult to study in a 2-dimensional *in vitro* system, which inherently lacks multiple aspects of the physiological microenvironment. ScienCell<sup>TM</sup>'s ready-to-use 3D human blood brain barrier spheroid kit offers cryopreserved 3D multicellular BBB spheroids comprised of human umbilical vein endothelial cells, brain vascular pericytes, and astrocytes at a 1:1:1 ratio, recapitulating the intracellular interactions at BBB. A highly unique feature of this kit is that researchers can achieve the functional and homogenous 3D BBB spheroids in 24-48 hours after thawing (Fig. 1-3), without encountering the long and complex procedures involved in 3D cell culture.

**Kit Components (Included)** 

3D Cell Culture Components							
Cat #	# of vials	Product Name	Quantity	Storage			
SP-8738	1	Human Blood Brain Barrier Spheroids	$1 \times 10^4$	Liquid			
		(SP-HBBB)	spheroids	nitrogen			
3D-8701	1	3D-BBB Spheroid Medium – basal	200 mL	2-8 °C			
		(3D-BBBSpM)					
3D-8752	1	3D-BBB Spheroid Supplement	2 mL	-20 °C			
		(3D-BBBSpS)					
0010	1	Fetal Bovine Serum (FBS)	10 mL	-20 °C			
0583	1	Penicillin/Streptomycin Solution (P/S)	2 mL	-20 °C			
0343 (or) 0353	1	Ultra-Low Binding Culture Plates 1		RT			
(or) 0383		(24-, 48-, or 96- well plate)					

### **Quality Control**

SP3D-HBBBS is tested for the formation of functional and uniform 3D BBB spheroids according to the included protocol. All components are negative for bacterial and fungal contamination.

#### **Product Use**

SP3D-HBBBS are for research use only. It is not approved for human or animal use, or application in clinical or *in vitro* diagnostic procedures.

# **Shipping**

SP-8738, 3D-8752, 0010, and 0583 are shipped on dry ice. 3D-8701, and [0343 (or) 0353 (or) 0383] are shipped at room temperature.

## References

- [1] Bernacki J, Dobrowolska A, Nierwiñska K, Maecki A. (2008) "Physiology and pharmacological role of the blood-brain barrier." *Pharmacological Reports*. 60: 600-622.
- [2] Daneman R, Zhou L, Kebede A, Barres B. (2010) "Pericytes are required for blood-brain barrier integrity during embryogenesis." *Nature*. 468(7323): 562-566.

#### **Procedure:**

### Step I: Preparing the complete 3D spheroid medium

- 1. Thaw 3D-BBB spheroid supplement (3D-BBBSpS; Cat. #3D-8752), fetal bovine serum (FBS; Cat. #0010), and penicillin/streptomycin solution (P/S solution; Cat. #0583) at 37°C. Mix 3D-BBSpS, FBS and P/S solution into the 3D-BBB spheroid medium (3D-BBBSpM; Cat. #3D-8701) by gently swirling the medium bottle around.
  - a. 3D-BBBSpM medium is **viscous** and optimized for homogenous spheroid formation.
  - b. Warm the complete 3D-BBBSpM medium only to **room temperature** before use.
  - c. When stored in the dark at 4°C, the complete medium is stable for one month.

### Step II: Thawing and maintaining the ready-to-use 3D spheroids

- 2. One frozen vial contains  $\ge 1 \times 10^4$  spheroids, which is sufficient for plating into **half of a multiwell plate** (e.g. 24-, 48-, and 96-well ultra-low binding plate).
- 3. Place the frozen vial in a 37°C water bath. Hold and rotate the vial gently until the contents completely thaw. Promptly remove the vial from the water bath, wipe it down with 70% ethanol, and transfer it to the sterile field.
- 4. Carefully remove the cap without touching the interior threads. GENTLY pipette spheroid suspension up and down **one time** to disperse potential spheroid aggregates.
- 5. Gently transfer the spheroid suspension into a fresh 50 mL conical tube.
- 6. Add 12 mL of 3D culture medium to the above 50 mL conical tube.
- 7. Resuspend spheroids in 3D culture medium by gently pipetting up and down for ~ 5-7 times using a serological pipette.

Note: 3D culture medium has a high viscosity; thus, pipetting slowly is important to avoid the formation of bubbles.

8. Aliquot the suggested volumes (see **Table A, column 2**) of spheroid suspension into each well of the included ultra-low binding plate (24-, or 48- or 96-well plate).

123Plate formatsVolume per wellTotal number of wells24-well~ 1000 μL12 wells48-well~ 500 μL24 wells96-well~ 250 μL48 wells

Table A: An Example of Suggested Medium Volumes Per Well

- 9. Incubate spheroids at 37°C in a 5 % CO<sub>2</sub> incubator.
- 10. For best results, do not disturb the culture for at least 16 hours after the culture has been initiated.

11. Next day, change 60-70 % of the top layer of the medium using a pipette by hand to remove the residual DMSO. (Do not use a vacuum aspirator). After 1<sup>st</sup> medium change, change 60-70% of the top layer of the medium every 3-4 days.

Note: Spheroids are situated at the bottom of the well <u>due to the viscosity of the 3D culture medium</u>. Thus, centrifugation of the plates is not necessary, and spheroid loss will not occur by changing 60-70 % of the top layer of the medium by pipetting.

12. Monitor the health of spheroids every day under the microscope. Human BBB spheroids are recovered and ready for your experiment after 2-3 days post thawing (see Figure 1).

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Figure 1 – At 100x magnification, brightfield images of ready-to-use 3D human BBB spheroids at 48 hours after thawing.

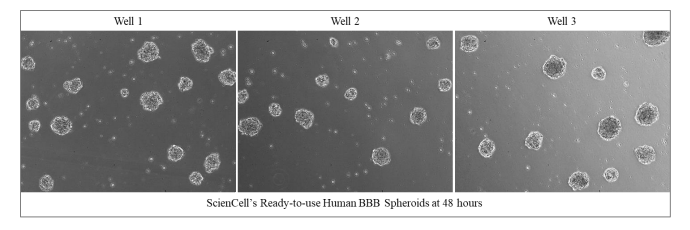


Figure 2 – Day 7; Top – Endothelial cell marker VWF (green), and astrocyte marker GFAP (red). Bottom – Endothelial cell marker CD31 (red) and pericyte marker PDGF receptor  $\beta$  (green) (taken at 200x magnification).

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Figure 3 – Day 7; Expression of the tight junction marker ZO1 (red) on the surface on the human BBB spheroids (taken at 200x magnification).

