

Liver-Chip R1 Co-Culture Protocol

November 07, 2024

EP-234 Rev. A

Liver-Chip R1 Co-Culture Protocol

Overview

Introduction This protocol describes the general steps for using the Emulate Liver-Chip R1 BioKit Co-Culture. For instructions on using the Liver-Chip **S1** BioKit Co-Culture, please see EP227.

CAUTION



Extreme consideration must be taken when using the Emulate Chip with a Zoë. The consumable type must be configured on the User Interface (UI) according to the chip being used (Chip-S1 or Chip-R1). This setting impacts the Prime, Regulate, and Flow parameters for the chip. A mismatch between the UI setting and the physical consumable could lead to instrument failure.

Chip-R1 can only be used with Pod-2. Chip-S1 can only be used with Pod-1.

To use Chip-R1 on Zoë-CM1, ensure the firmware is on v1.2.4 or above To use Chip-R1 on Zoë-CM2, ensure the firmware is on v1.4.5 or above*

Firmware updates can be performed through Utility Hub on the Emulate website.

*If your Zoë-CM2 is on v1.3.0 or below, please contact Emulate Support.

| | | | 윰 | ZOE 12345 |
|----------|----------|----------------|-----------|-------------|
| Prime C | Cycle | | | |
| Regula | te Cycle | Start | | |
| Flow | 🛱 Chip-R | 1 | Stretch | |
| LOCATION | MEDIA | RATE | PARAMETER | RATE |
| Тор | Liquid | $30 \ \mu L/h$ | Strain | 0 % |
| Bottom | Liquid | 30 µL/h | Frequency | O Hz |

Follow the below steps to select the Chip-R1 as the consumable type:

| Step | Action |
|------|--|
| 1 | Use the Dial to highlight "Chip-S1" or "Chip-R1" on the display. |
| 2 | Press the Dial Button to select the displayed chip type. |
| 3 | Rotate the Dial to toggle to "Chip-R1" |
| 4 | Press the Dial Button to select "Chip-R1" |

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Part I: Liver-Chip R1 BioKit Co-Culture

Overview

| Introduction | This part provides an overview of the Liver-Chip R1 BioKit Co-Culture as well as its key components, shipping information, and storage specifications. | | | |
|---------------|--|-------------------------------|--|--|
| Components | The Liver-Chip R1 BioKit Co cells listed in the table below | p-Culture includes the pre-qu | alified primary human liver | |
| | Category | Channel Location | Type of Cells | |
| | Parenchymal Epithelial Cells | Тор | Hepatocytes | |
| | Non-parenchymal cells (LSEC) | Bottom | Liver sinusoidal endothelial cells (LSECs) | |
| Cell Shipping | Cells are shipped in cryogenic storage vacuum flasks. | | | |
| Cell Storage | Cells must be stored in the vapor phase of liquid nitrogen until use. | | | |

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Part II: Experimental Overview

Overview

| Introduction | This part gives an overview of the experimental workflow. |
|---|--|
| Day -X: Reagent Preparation | Aliquot reagents (media supplements, ECM, Matrigel, etc.) |
| Day -5: Thaw LSECs | Prepare LSEC culture flask Thaw and plate LSECs |
| Day -2: (Optional) Passage LSECs | It is up to the user's discretion to perform additional passaging It may not be necessary if the LSECs are confluent enough on this day |
| Day -1: Chip Preparation | Prepare chipsPrepare ECM solutionCoat chips with ECM |
| Day 0: Hepatocytes to Chips, Hepatocyte Overlay | Prepare hepatocyte seeding medium Prepare chips Thaw hepatocytes Adjust cell density Seed hepatocytes to top channel Seed a well plate Prepare overlay medium Overlay hepatocytes |

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Overview, Continued

| Harvest LSECs Seed LSECs to bottom channel Gas equilibration of media Prime Pods Wash chips Chips to Pods | | Seed LSECs to bottom channel Gas equilibration of media Prime Pods Wash chips |
|--|--|--|
|--|--|--|

Pods to Zoë

Day 2+: Maintaining and Sampling

• Sampling and media replenishment

Part III: Equipment and Materials Required

Overview

| Introduction | Ensure all equipment, materials, and reagents are accessible prior to each experiment. Below are the catalog numbers for the specific materials needed. |
|--------------|---|
| Note | Exact catalog numbers are not provided for some required materials, as several brands and models are accepted. |

Required Equipment A list of equipment needed for this protocol is provided below:

| Equipment | Description | Supplier | Catalog Number |
|---|------------------------------|-----------|----------------|
| Liver-Chip R1 BioKit | Co-Culture 12- or | Emulate | BIO-LH-CO12R1 |
| Co-Culture | 24-pack | | BIO-LH-CO24R1 |
| Zoë-CM2 [®] Culture | 1 per 12 chips | Emulate | ZOE-CM2 |
| Module | | | |
| Orb-HM1 [®] Hub | 1 per 4 Zoës | Emulate | ORB-HM1 |
| Module | | | |
| Chip Cradle | Autoclaved, 1 per 6 chips | Emulate | CHIP-CRD |
| Steriflip [®] -HV Filters | Sterile, 0.45 µm | EMD | SE1M003M00 |
| | PVDF filter | Millipore | |
| Square Cell Culture Dish (120 x 120 mm) | Sterile, 1 per 6 chips | VWR | 82051-068 |
| Collagen type-1 | 24-well, flat-bottom, | Corning | 356408 |
| coated plates | TC-treated | | |
| Handheld vacuum aspirator | - | Corning | 4930 |
| Aspirating pipettes | 2-mL, polystyrene, | Corning / | 357558 |
| | individually wrapped | Falcon | |
| Aspirating tips | Sterile (autoclaved) | - | - |
| Serological Pipettes | 2-mL, 5-mL, 10-mL, | | |
| | and 25-mL low- | | |
| | endotoxin, sterile | | |
| Pipette | P20, P200, and | - | - |
| | P1000 | | |
| Pipette Tips | P20, P200, and | - | - |
| | P1000 sterile, filter, | | |
| | low-adhesion | | |

Required Equipment, Continued

Overview, Continued

| Equipment | Description | Supplier | Catalog Number |
|-----------------------|------------------------|----------|----------------|
| Conical tubes | 15-mL and 50-mL | - | - |
| | polypropylene, sterile | | |
| Eppendorf Tubes® | 15-mL, sterile | - | - |
| Parafilm [®] | - | - | - |
| Microscope (with | For bright-field | - | - |
| Camera) | imaging | | |
| Hemocytometer | - | - | - |
| Water bath (or | - | - | - |
| beads) | | | |
| Vacuum set-up | -70 kPa minimum | - | - |
| | achievable pressure | | |
| T25 Flask | - | - | - |
| T75 Flask | - | - | - |
| Ice bucket | - | - | - |
| 70% ethanol wipes | For surface | - | - |
| | sterilization | | |

Required Materials

A list of reagents, media, and supplements needed for this protocol in addition to the Emulate Liver-Chip R1 BioKit Co-Culture is provided below:

| Reagent | Description | Supplier | Catalog Number |
|-------------------------------|---------------------|----------|----------------|
| Dulbecco's PBS | 1X | Corning | 21-031-CV |
| (DPBS -/-) (without | | | |
| Ca2+, Mg2+) | | | |
| 10X DPBS (-/-) | 10X | Corning | 20-031-CV |
| (without Ca++, Mg++) | | | |
| Trypan blue | 0.4% solution | Sigma | 93595 |
| Percoll [®] Solution | 100% stock | Sigma | P4937 |
| | solution | | |
| Trypsin-EDTA | 0.05% trypsin | Sigma | T3924 |
| solution | | | |
| WEM Medium (+) | Williams' medium | Sigma | W4128 |
| | E with phenol red | | |
| | (+) | | |
| WEM Medium (-) | Williams' medium | Sigma | W1878 |
| | E no phenol red (-) | | |

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Overview, Continued

| Required |
|------------|
| Materials, |
| Continued |

| Reagent | Description | Supplier | Catalog Number |
|--------------------|-----------------|-----------------|----------------|
| CSC medium (Kit) | LSEC medium & | Cell Systems | 4Z3-500 |
| | supplements | - | |
| Culture boost™ | 50X supplement | Cell Systems | 4CB-500 |
| Attachment Factor™ | 1X | Cell Systems | 4Z0-210 |
| Matrigel® | LDEV-free | Corning | 354234 |
| Fibronectin | Bovine protein, | ThermoFisher | 33010-018 |
| | plasma | | |
| Collagen type I | Rat tail; HC | Corning | 354249 |
| Penicillin- | 10,000 U / mL; | Sigma | P4333 |
| streptomycin | 10 mg / mL | | |
| L-GlutaMax™ | 200 mM | ThermoFisher | 35050-061 |
| L-Ascorbic Acid | 100-mg powder | Sigma | 5960 |
| Dexamethasone | 100-mg powder | Sigma | D4902 |
| Fetal bovine serum | Sterile, heat- | Sigma | F4135 or F8317 |
| (FBS) | inactivated | | |
| ITS+ | Premix | Corning | 354352 |
| | supplement | | |
| Cell culture grade | For fibronectin | Corning | 25-055-CVC |
| water | solution | | |
| | preparation | | |
| Cell culture grade | For | Millipore Sigma | D2650 |
| DMSO | dexamethasone | | |

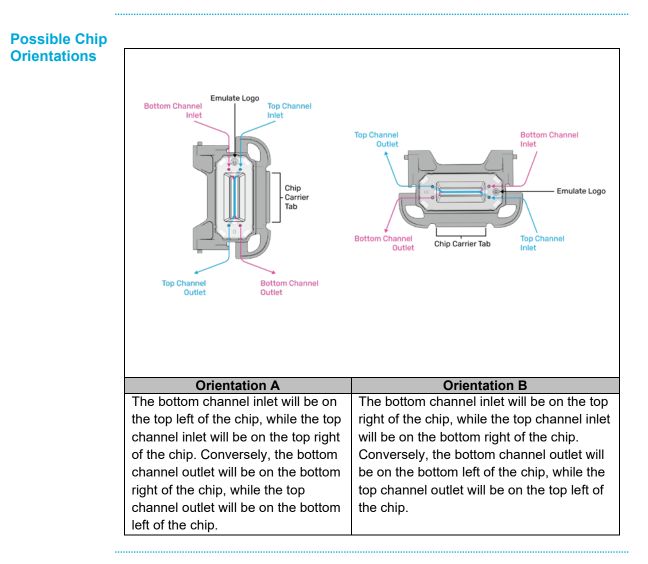
Part IV: Workstation Preparation and Chip Handling Techniques

Overview

| Aseptic Techniques | Always work with chips in a sterile environment, such as the biosafety cabinet (BSC). |
|-----------------------|---|
| | Before beginning the experiment, prepare, sanitize, and organize the work surface of the BSC. Arrange tips, media, and other necessary materials in the sterile field so they are easily within reach but not blocking the path of airflow. |
| | Use 70% ethanol to thoroughly sanitize all reagents and materials before placing them into the BSC. |
| | Always avoid touching the chip directly. Handle the chip carrier only by the sides, or by the tab, with gloves. |
| | Do not remove chips from the chip carrier until after the experiment. |
| Cell Storage | Always store cryopreserved cells in the vapor phase of liquid nitrogen. Never store them in dry ice nor in a -80°C freezer. Chronic temperature fluctuations can cause severe damage to cell membranes and the cytoskeletal components. |
| | |

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Chip Handling Techniques



Pipetting

While 50 μ L (top channel) and 20 μ L (bottom channel) are the standard volumes used throughout the protocol, there can be some flexibility in the actual volumes used:

| Channel | Volume Range |
|----------------|--------------|
| Top Channel | 35–50 μL |
| Bottom Channel | 15–20 μL |

These volumes allow for simple pipetting and a slight overfill to avoid bubbles or dry channels.

 \bullet All wash steps, unless otherwise stated, are performed using 200 μL of the specific wash solution.

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Chip Handling Techniques, Continued

| Channel and | The specific channel and membrane dime | ensions are outlined below: | | |
|-------------|--|-----------------------------------|--|--|
| Membrane | Top Channel | | | |
| Dimensions | Width x Height Dimensions | 1000 μm x 1000 μm | | |
| | Area | 25.66 mm ² | | |
| | Volume | 24.52 μL | | |
| | Imaging distance from the bottom of | 172 µm | | |
| | the chip to the top of the membrane | | | |
| | 950um | 1 | | |
| | 1000um 1000um 1050um | | | |
| | Bottom | n Channel | | |
| | Width x Height Dimensions | 1050 µm x 100 µm | | |
| | Area | 29.46 mm ² | | |
| | Volume | 2.97 μL | | |
| | | nbrane | | |
| | Pore diameter | 3.0 µm | | |
| | Pore spacing | Random distribution (track-etched | | |
| | | membrane) | | |
| | Porosity | 2.8% | | |
| | Thickness | 22 μm | | |
| | Co-Culture Region | | | |
| | Area | 16.56 mm ² | | |

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Chip Handling Techniques, Continued

Pipetting Solution into Channels

Follow the steps below for pipetting solution into the Organ-Chip when coating, washing, and seeding cells prior to attaching the chip to Zoë.

Note: Always introduce liquid to the endothelial channel before the epithelial channel.

| Step | Action |
|------|--|
| 1 | Take a P200 pipette with a sterile pipette tip and collect the solution to |
| | be added to the Organ-Chip. |
| 2 | Place the pipette tip perpendicular to the chip channel inlet, ensuring |
| | that the tip is securely in the port. |
| 3 | Steadily dispense the liquid through the channel. |

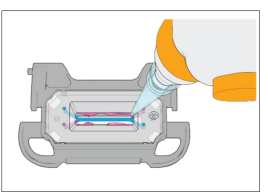
CAUTION



There is additional resistance in the bottom channel of Chip-R1 compared to Chip-S1. Therefore, when pipetting into the bottom channel, depress the plunger fully and hold the pipette at the port for several seconds to allow the full volume to be pipetted through the bottom channel.



When aspirating from the gasket window, take precaution NOT to make contact with the channel feature itself. The top gas exchanger film can easily be punctured by a pipette tip.



Note on Aspiration: Careful consideration must be given to aspiration steps during Chip-R1 seeding. Residual liquid over the outlet ports may be a contributing factor to the formation of cell distribution gradients when handling the chips post-seeding.

Suggestions for aspiration and avoiding gradients:

- 1. Aspirate excessive volume from the outlets with a vacuum aspirator.
- 2. Do not work directly over the port as this may cause the cell medium to aspirate from the channel itself.
- 3. Do not drag the aspirator tip across the top of the chip as this can cause scratches on the top layer.

Part V: Liver-Chip R1 Co-Culture Protocol

Protocol Overview

Introduction This section lists the basic steps for using Liver-Chip R1 BioKit Co-Culture in experiments.

Timeline

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| Day -X: Reagent Preparation | 15 |
| Day -5: Thawing Liver Sinusoidal Endothelial Cells (LSECs) | 17 |
| Day -1: Chip Preparation | 20 |
| Day 0: Hepatocytes to Chip & Matrigel Overlay | 25 |
| Day 1: LSECs to Chip, Chips to Pods, Pods to Zoë | 40 |
| Day 2+: Chip Maintenance and Sampling | 55 |

Day -X: Reagent Preparation

Aliquot Reagents

Introduction Aliquot reagents prior to use so the stock solutions do not undergo multiple freezethaw cycles.

Fibronectin (ECM)

| Reagent | Conc. [Stock] | Amount | Volume | Solvent |
|-------------|---------------|--------|--------|--------------------|
| Fibronectin | 1 mg / mL | 1 mg | 1 mL | Cell culture grade |
| | | | | water |

- Resuspend 1 mg fibronectin in 1 mL of cell culture grade water according to manufacturer's instructions.
- Create single-use volume aliquots and store them at -20°C. For 12 chips, this equates to 37.5 μL of Fibronectin for 1.5 mL of final ECM volume (see Pages 22 and 23 for calculations).

Matrigel (Overlay)

| Reagent | Volume |
|----------|----------------|
| Matrigel | Varies per lot |

The Matrigel bottle must be thawed overnight on ice, either in the back of the $2-6^{\circ}$ C refrigerator or in a cold room. Add water to ensure the ice is slushy, as the solution gels rapidly at temperatures above 10°C. For aliquoting, use pipette tips and tubes that have been prechilled to -20°C, maintain aliquots on slushy ice, and transfer the aliquots immediately to -20°C.

- Calculate and prepare the aliquot volume needed to achieve a concentration of 0.25 mg/mL in the overlay media.
- Store aliquots at -20°C.

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Aliquot Reagents, Continued

Culture Supplements

| Reagent | Conc. [Stock] | Volume | Solvent |
|-----------------|---------------|--------------------|--------------------|
| L-Ascorbic acid | 50 mg / mL | Calculate based on | Cell culture grade |
| | | amount purchased | water |
| Dexamethasone | 10 mM | Calculate based on | Cell culture grade |
| | | amount purchased | DMSO |
| Dexamethasone | 1 mM | Calculate based on | Cell culture grade |
| | | amount purchased | DMSO |

- Resuspend each supplement to the stock concentration in the table above.
- Aliquot each supplement to single-use volumes (i.e., 12 chips, see pages 26 and 37 for details).
- Please check the Complete Hepatocyte Seeding Medium and Complete Hepatocyte Maintenance Medium preparation tables in the appendix to adjust the aliquot size based on your experiment size.
- For L-ascorbic acid, prepare 160–200 µL aliquots.
- For 10 mM dexamethasone, prepare 20 µL aliquots.
- For 1 mM dexamethasone, prepare 10 µL aliquots.
- Store aliquots at -20°C.

Day -5: Thawing Liver Sinusoidal Endothelial Cells (LSECs)

| Goals | Thaw and expand LSEC culture media in flask | | |
|-----------------------|---|----------|--|
| Required Materials | Complete LSEC Culture Medium (at 37°C) 15 mL conical tube Attachment Factor™ T-75 flask Serological pipettes Pipettes and tips Aspirator Centrifuge 70% ethanol | | |
| Key Steps | Tonic | See Page | |

| Торіс | See Page |
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| Prepare LSEC Culture Media and Flask | 18 |
| Thaw and Plate LSECs | 19 |

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Prepare LSEC Culture Media and Flask

LSEC Culture Media

Base LSEC Culture Medium (500 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-------------|--------|---------------|---------------|---------|----------|
| CSC basal | 490 mL | - | - | Cell | 4Z3-500 |
| medium | | | | Systems | |
| Culture- | 5 mL | - | 1% | Cell | 4CB-500 |
| boost | | | | Systems | |
| Pen / Strep | 5 mL | - | 1% | Sigma | P4333 |

• Store the LSEC Culture Medium at 4°C.

• Use the LSEC Culture Medium within 30 days of preparation.

Complete LSEC Culture Medium (50 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-----------|--------|---------------|---------------|--------------|----------|
| Base LSEC | 45 mL | - | - | Recipe Above | - |
| Culture | | | | | |
| Medium | | | | | |
| FBS | 5 mL | - | 10% | Sigma | F4135 |

• Store the Complete LSEC Culture Medium at 4°C.

• Use the Complete LSEC Culture Medium within 7 days of preparation.

Prepare Flask

| Step | Action |
|------|--|
| | Warm a sufficient amount of Complete LSEC Culture Medium and |
| 1 | Attachment Factor™ to 37°C. 15 mL of medium is needed for |
| | thawing, and an additional 15 mL is needed for each flask. |
| 2 | Label the culture flask with the relevant information (e.g., cell type, |
| 2 | passage number, date, initials). |
| | Pipette Attachment Factor onto the growth surface of the flask until it |
| 3 | is fully covered. 5 mL of Attachment Factor is used for each T-75 |
| | flask. |
| 4 | Place the prepared flask into the 37°C incubator to coat the surface. |
| 4 | Maintain this temperature until the cells are plated for at least 5 min. |

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Thaw and Plate LSECs

Note on LSECs

LSECs are the only cells in this protocol that must be plated and expanded before being seeded in the chip. LSECs initial passage may vary per Liver-Chip R1 BioKit Co-Culture. Reach out to your dedicated Scientific Liaison or Emulate Support for further guidance based on the lot and passage of LSECs you received.

If needed, LSECs can be further expanded by passaging prior to chip seeding, but should not exceed passage 5 from the initial BioKit. Reach out to Emulate support (support@emulatebio.com) for any questions related to LSEC seeding density optimization.

Thawing and Maintaining Cells

| Step | Action |
|------|---|
| 1 | Immerse the vial(s) of cells in a 37°C water bath without submerging the cap. Closely observe and gently agitate the vials. Remove them from the water bath just before the last of the ice disappears. |
| 2 | Spray the vial(s) with 70% ethanol and wipe them dry before placing them into the BSC. |
| 3 | Immediately transfer the contents of the vial(s) into a sterile 15 mL conical tube containing 3 mL of warm Complete LSEC Culture Medium. |
| 4 | Rinse the vial(s) with 1 mL of Complete LSEC Culture Medium and collect the run-off in the 15 mL tube. |
| 5 | Bring the volume to 15 mL with Complete LSEC Culture Medium. |
| 6 | Centrifuge 200 x g for 5 min at room temperature. |
| 7 | Aspirate and discard the supernatant, leaving approximately 100 µL of medium covering the cell pellet. |
| 8 | Loosen the pellet by gently flicking the tube. |
| 9 | Re-suspend cells in 15 mL of Complete LSEC Culture Medium. |
| 10 | Aspirate and discard excess Attachment Factor from the T-75 flask that was prepared. Note: It is unnecessary to rinse or dry the flask prior to adding cells. |
| 11 | Add the LSEC suspension to the freshly coated T-75 flask. |
| 12 | Incubate overnight at 37°C and 5% CO ₂ . |
| 13 | Refresh the Complete LSEC Culture Medium every other day until the cells are seeded in the chip. |

Day -1: Chip Preparation

| Overview | | |
|-----------------------|---|--------------------|
| Goals | Coat the inner channels with a mixture of collagen I and fibron for cell attachment | ectin ECM proteins |
| Required Materials | Chip-R1 Rigid Chips (12 Chips per Zoë) 15 mL conical tubes DPBS (- / -) at room temperature DPBS (- / -) aliquot at 4°C Collagen I Fibronectin 70% ethanol Square Cell Culture Dish (120 x 120 mm) Ice and ice bucket Pipettes and filtered tips Aspirator and sterile tips | |
| Key Steps | Tania | Cao Dava |

| Торіс | See Page |
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| Prepare Chips | 21 |
| Prepare ECM Solution | 22 |
| Coat Chips with ECM | 24 |

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Prepare Chips

Steps

| Step | Action |
|------|---|
| 1 | Spray the chip packaging with 70% ethanol and bring it into the BSC. |
| 2 | Open the packaging, place the Chip Cradle in the dish, and then carefully insert 6 chips into the Chip Cradle (see Figure 1). Note: For ease of workflow, ensure the carrier's tab is pointing to the right and that all chips are facing the same direction within the dish. |
| | <image/> <image/> |
| | |
| 3 | Label each chip carrier tab with the corresponding chip's ID number. |

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Prepare ECM Solution

Before YouPrepare fresh ECM before each use by combining the individual ECM componentsBeginwith cold DPBS to reach the final working concentrations. The ECM solution will
coat both channels.

Needed Volumes

For human Liver-Chips, the ECM working concentrations are:

| Reagent | Concentration |
|-------------|---------------|
| Collagen I | 100 μg / mL |
| Fibronectin | 25 μg / mL |

Key Steps

| Step | Action |
|------|--|
| 1 | Bring a full ice bucket to the BSC. |
| 2 | Thaw one aliquot of fibronectin (1 mg / mL) on ice. Always maintain each ECM component and mixture on ice. |
| 3 | Calculate the volume of ECM solution needed to coat all chips. 1. Volume required per chip: ~100 μL 2. For every batch of 12 chips, prepare 1.5 mL of ECM solution: 12 chips x 100 μL / chip = 1.2 mL of ECM solution. 1.2 mL + extra 300 μL = 1.5 mL of ECM solution. |
| 4 | Combine the components to prepare the ECM working solution. |
| 5 | Keep the ECM solution on ice until it is used. |

| Ľ | Г | L | Е | | |
|---|---|---|---|--|--|
| | | | | | |

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Prepare ECM Solution, Continued

Example ECM Calculation

ECM Calculation Example for 12 chips:

SolutionConcentrationCollagen I stock concentration8.41 mg / mL (C1)Collagen I final concentration0.1 mg / mL (C2)Fibronectin stock concentration1 mg / mL (C1)Fibronectin final concentration0.025 mg / mL (C2)Stock VolumeCollagen I (X) or fibronectin (Y) (V1)Final volume of ECM solution1.5 mL (V2)

Collagen Calculation:

 $(8.41 \text{ mg} / \text{mL}) \times (\text{X mL}) = (0.1 \text{ mg} / \text{mL}) \times (1.5 \text{ mL})$

X = 17.83 μ L of collagen I stock solution

Fibronectin Calculation:

 $(1 \text{ mg} / \text{mL}) \times (Y \text{ mL}) = (0.025 \text{ mg} / \text{mL}) \times (1.5 \text{ mL})$

Y = 37.5 µL of fibronectin

DPBS Calculation

Volume DPBS =

(total volume of ECM needed) - (volume of collagen I) - (volume of fibronectin)

= 1500 μL – 17.83 μL – 37.5 μL

= 1444.67 µL of DPBS

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Coat Chips with ECM

Steps

| Step | |
|--------|--|
| 1 | Using a P200 pipette, draw 100 μ L of ECM solution. (Each chip will use 100 μ L.) |
| 2 | Carefully introduce ECM solution through the bottom channel inlet |
| 2 | until a small droplet forms on the outlet. |
| 2 | Without releasing the plunger, move the pipette containing the |
| 3 | remaining ECM solution to the top channel inlet. |
| 4 | Introduce ECM solution through the inlet, leaving small droplets of |
| 4 | excess ECM solution on both ports in both channels (see Figure 2). |
| | |
| 5 | Figure 2. Chip in chip carrier with small ECM droplets at ports. Wash any bubbles from the channel with the ECM solution. |
| 6 | Repeat steps 1–6 for each chip. |
| 7 | To prevent evaporation during incubation, fill the central reservoir with |
| 1 | 0.75–1 mL of DPBS (see Figure 3). Place the lid onto the dish. |
| | |
| Figure | 3. Pipette filling central reservoir of Chip Cradle with 0.75 mL DPBS. |
| | For best results, incubate the chips at 4°C overnight, then at 37°C for |
| 8 | at least 1 h the following day prior to seeding. |
| 0 | Note: Chips can be stored at 4°C for up to 2 days if kept moist. |
| | |

Day 0: Hepatocytes to Chip & Matrigel Overlay

Overview

Goals

• Thaw and seed hepatocytes in chip.

Required Materials

- Complete Hepatocyte Seeding Medium (at 37°C)
- Percoll solution (at room temperature)
- 10X DPBS (at room temperature)
- 1X DPBS (at room temperature)
- Serological pipettes
- Pipettes and filtered tips
- Aspirator and sterile tips
- 50 mL conical tubes
- Diluted trypan blue counting solution
- Hemocytometer
- 24-well collagen I-coated plate
- 70% ethanol
- Microscope
- Complete Hepatocyte Maintenance Medium (at 4°C)
- Complete Hepatocyte Maintenance Medium (at 37°C)
- Hepatocyte Overlay Medium (at 4°C)
- Matrigel aliquot (at 4°C in slushy ice)
- Ice bucket and ice

Key Steps

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| Thaw Hepatocytes | 29 |
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|--|----|----|--|

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Prepare Hepatocyte Seeding Media

Hepatocyte Seeding Media

Base Hepatocyte Seeding Medium (500-mL bottle)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|--------------|--------|---------------|---------------|--------|----------|
| WEM + | 490 mL | - | - | Sigma | W4128 |
| (with phenol | | | | | |
| red) | | | | | |
| Pen / Strep | 5 mL | - | 1% | Sigma | P4333 |
| L-GlutaMAX | 5 mL | - | 1% | Gibco | 35050-61 |

• Store the Base Hepatocyte Seeding Medium at 4°C.

• Use the Base Hepatocyte Seeding Medium within 30 days of preparation.

Complete Hepatocyte Seeding Medium (150 mL)

This amount is sufficient for 12 chips

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-----------------|---------|---------------|---------------|---------|----------|
| Base Hepatocyte | 140.835 | - | - | Recipe | - |
| Seeding Medium | mL | | | Above | |
| ITS+ premix | 1.5 mL | - | 1% | Corning | 354352 |
| Ascorbic Acid | 150 µL | 50 mg / mL | 0.05 mg / mL | Sigma | 5960 |
| Dexamethasone | 15µL | 10 mM | 1 µM | Sigma | D4092 |
| FBS | 7.5 mL | - | 5% | Sigma | F4135 |

• Store the Complete Hepatocyte Seeding Medium at 4°C.

• Use the Complete Hepatocyte Seeding Medium within 3 days of preparation.

90% Percoll Solution (20 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|------------------|--------|---------------|---------------|--------|----------|
| Percoll Solution | 18 mL | 100% | 90% | Sigma | P4937 |
| 10X DPBS (- / -) | 2 mL | 100% | 10% | - | - |

• Maintain the 90% Percoll Solution at room temperature prior to use.

• Prepare the Percoll Solution fresh for each use.

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Prepare Hepatocyte Seeding Media, Continued

Trypan Blue Cell Counting Solution

| Reagent | Volume | Source | Cat. No. |
|---------------------|--------|--------------|----------|
| Complete Hepatocyte | 40 µL | Recipe Above | - |
| Seeding Medium | | | |
| Trypan Blue | 5 µL | Sigma | 93595 |

• Maintain the Trypan Blue Cell Counting Solution at room temperature prior to use.

• Prepare the Trypan Blue Cell Counting Solution fresh for each use.

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Prepare Chips

Steps

| Step | Action |
|------|---|
| 1 | Transfer the ECM-coated chips from the incubator into the BSC. |
| 2 | Fully aspirate the ECM from both channels. |
| | Gently wash each channel with 200 µL of Complete Hepatocyte |
| 3 | Seeding Medium. Aspirate the medium outflow on each chip's |
| | surface, leaving medium in both channels. |
| | Repeat the wash with an additional 200 μ L of Complete Hepatocyte |
| 4 | Seeding Medium per channel, leaving the excess medium outflow that |
| | covers the inlet and outlet ports. |
| 5 | Cover the Square Cell Culture Dish (120 x 120 mm) and place the |
| 5 | chips into the incubator until the cells are ready for seeding. |

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Thaw Hepatocytes

| Before You Begin | Before thawing the cryopreserved hepatocytes, make sure all equipment is organized and ready for use. Also, be sure that all required reagents are prepared and have reached the appropriate temperature. |
|---------------------------|---|
| Tips for Thawing Cells | Up to three vials of cryopreserved hepatocytes can be thawed at the same time. Once they are thawed, the contents of each should be combined into one 50 mL conical tube (See Step 6 in "Steps" below") and processed as one sample. As the cells are thawing, it is critical to work as quickly but as gently as possible. This will help maximize cell recovery and minimize damage to the hepatocytes. Do not allow the cells to thaw at room temperature or on ice. Once the hepatocytes are thawed, dilute them in the Complete Hepatocyte Seeding Medium as soon as possible to prevent DMSO toxicity within the cryoprotectant. |
| Steps | |

| Step | Action |
|------|--|
| | Place 3 mL of warm Complete Hepatocyte Seeding Medium into a |
| 1 | sterile 50 mL conical tube. |
| 2 | Remove the required number of cryovials. |
| | Spray each cryovial with 70% ethanol and wipe it dry. Twist the cap a |
| 3 | quarter of a full turn to relieve any internal pressure, then re-tighten it. |
| | Immediately place the frozen vial in a 37°C water bath without |
| | submerging the cap. Rapidly thaw hepatocytes by gently swirling the |
| 4 | vials in the water bath until only a small ice pellet remains. This |
| | process should take only 60–90 seconds. Thawing any longer will |
| | decrease viability and cell yield. |
| | When one small ice pellet remains, immediately remove the vial from |
| 5 | the water bath, wipe it dry, spray it with 70% alcohol, and wipe it dry |
| | again before placing it into the BSC. |
| | Quickly transfer the contents of the vial into the 3 mL of Complete |
| 6 | Hepatocyte Seeding Medium in the sterile 50 mL conical tube |
| | prepared in Step 1. |
| 7 | Rinse the cryovial with 1 mL of Complete Hepatocyte Seeding |
| 1 | Medium and transfer it to the 50 mL conical tube. |
| | With gentle agitation and swirling, slowly add enough of the |
| 8 | Complete Hepatocyte Seeding Medium to bring the total volume to |
| | 35 mL. |
| 9 | Add 15 mL of 90% Percoll Solution, bringing the total volume to 50 |
| | mL. |
| 10 | Cap the tube tightly and slowly invert it three times to mix the cell |
| | solution. |
| 11 | Centrifuge the cells at 96 x g for 6 min at room temperature. |

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Thaw Hepatocytes, Continued

Steps, Continued

| Step | Action |
|------|--|
| 12 | Return the tube to the BSC. Carefully aspirate the supernatant, |
| 12 | leaving 3–5 mL. Ensure the pellet remains undisturbed. |
| 13 | Tilt and rotate the tube to gently re-suspend the cell pellet in the |
| 15 | remaining medium. |
| 14 | Gently add enough Complete Hepatocyte Seeding Medium to bring |
| 14 | the total volume to 50 mL. |
| 15 | Centrifuge the cells at 72 x g for 4 min at room temperature. |
| 16 | Return the tube to the BSC. Carefully aspirate the supernatant, |
| 10 | leaving 1–2 mL. Ensure the pellet remains undisturbed. |

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Adjust Cell Density

Overview Human hepatocytes must be seeded in the Liver-Chip at a density of 2.8 x 10⁶ cells / mL. It is essential to ensure the seeding density is accurate for viable, functional cells and long-term culture.

Note: This seeding density was optimized with Hepatocyte donor 2214423. The seeding density may vary and must be optimized based on the donor that is part of your BioKit. Reach out to Emulate support (support@emulatebio.com) for any questions related to hepatocyte seeding density optimization.

Steps

| Step | Action | |
|------|---|--|
| 1 | Tilt and rotate the tube to gently resuspend the cell pellet. | |
| 2 | Measure the total suspension volume using a 5 mL pipette. | |
| 3 | Confirm the cell pellet has disappeared, sufficiently rotate to homogenize the cell suspension and transfer 5 µL of the cell suspension to the Trypan Blue Cell Counting Solution, generating a 1:10 dilution. | |
| 4 | Mix the trypan blue solution thoroughly and count the cells using a hemocytometer. | |

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Cell Counting and Viability Assessment

Cell Counting and Viability Assessment

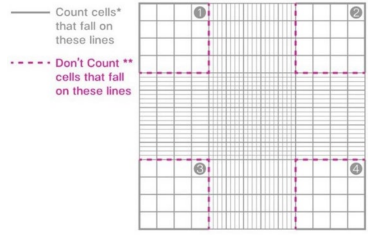


Figure 4. Example hemocytometer

1. Count both viable and non-viable cells in each quadrant of the hemocytometer (see Figure 4).

Live Cell Count, Dead Cell Count, Total Cell Count.

2. Calculate the percentage viability of the cell solution

Live Cells + Total Cells x 100 = % Viability

3. Calculate the viable cell concentration. The dilution factor is 10 when prepared in the trypan blue solution above.

Live Cell Count x (1 x 10 x 10⁴) ÷ 4 = Viable Cell Concentration (cells / mL)

- 4. Calculate the viable cell yield.
- 5. Viable Cell Concentration x Cell Suspension Volume = Viable Cell Yield (cells)

Viable Cell Yield + Desired Density = Reconstitution Volume

DilutingAfter calculating the Viable Cell Yield, dilute the hepatocytes with warm CompleteHepatocytesHepatocyte Seeding Medium to the required final cell density: 2.8 x 10⁶ cells / mL (if
using hepatocyte donor 2214423).

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Cell Counting and Viability Assessment, Continued

AdditionalIf the Viable Cell Concentration is less than 2.8 x 106 cells / mL (when using
hepatocyte donor 2214423):

| Action |
|---|
| Leave the hepatocyte cell suspension undisturbed at room |
| temperature for at least 5 mins. This will allow the cells to settle at the |
| bottom of the tube. |
| Gently remove enough from the top of the supernatant to decrease |
| total cell suspension volume. |
| Re-count the cell suspension and recalculate the appropriate seeding |
| density accordingly (Steps 5 and 6 on the next page). This will help |
| to avoid subjecting hepatocytes to mechanical stress with |
| centrifugation. |
| |

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Seed Hepatocytes to the Top Channel

Overview

Work with one chip at a time. After seeding the first chip, use a microscope to assess the cell density within the channel. Adjust the density of the cell suspension as necessary for the rest of the chips.

Steps

| Step | Action | | |
|--|--|--|--|
| 1 | Bring the square dish containing the prepared chips to the BSC. | | |
| 2 | Without touching the ports, carefully aspirate excess medium droplets | | |
| 2 | from the surface of one chip. | | |
| 3 | Very gently agitate the cell suspension before seeding each chip to | | |
| 5 | ensure a homogeneous cell suspension. | | |
| | Quickly and steadily pipette 40 μ L of the cell suspension into the top | | |
| 4 | channel inlet port while aspirating the outflow | | |
| | fluid from the surface. Do not directly touch the outlet port. | | |
| | Cover the dish and transfer it to the microscope to check the seeding | | |
| 5 | density within the chip (see Figure 5). | | |
| Ŭ | Note: At this stage, cells with optimal seeding density will form an | | |
| _ | even cell layer with ~half a cell radius between individual cells. Seeding density too low Seeding density optimal Seeding density too high | | |
| Immediatel seedin 24 hours seedin | g after | | |
| | Figure 5. Hepatocyte seeding density reference chart. | | |
| | If the seeding density is not optimal or cell distribution is not even, | | |
| 6 | return the chips to the BSC, and wash the channel with 200 μL of fresh medium twice. Do not aspirate the medium from the | | |
| U | channels. Adjust the cell density accordingly and repeat Steps 3–5 | | |
| | until the density in the channel is correct. | | |
| | | | |

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Seed Hepatocytes to the Top Channel, Continued

| Steps, | Step | Action | | |
|-----------|----------|--|--|--|
| Continued | | After confirming the correct cell density, seed cells in the remaining chips. | | |
| 7 | | Note: Minimize the amount of time the cells are outside the incubator | | |
| | | by seeding batches of no more than 12 chips at a time and by | | |
| | | immediately placing the batches into the incubator at 37°C. | | |
| | | Place the chips (with the DPBS-filled reservoir) in the incubator at | | |
| | 8 | 37°C for 4 h (see Figure 6 for examples of attachment). | | |
| | 0 | Note: Achieving the correct seeding density is essential for the | | |
| | | success of the chips. | | |
| | Figure 6 | The left image shows appropriate attachment. The right image shows appropriate attachment. | | |

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Seed a Well Plate

Overview It is recommended to always seed any remaining hepatocytes into a 24-well plate pre-coated with collagen I as a control for cell quality.

Key Steps

| Step | Action |
|------|---|
| 1 | Dilute the hepatocyte suspension with warm Complete Hepatocyte |
| | Seeding Medium to a final density of 0.8 x 10⁶ cells / mL . |
| 2 | Add 400, 500, and 600 μ L of the cell suspension to three separate |
| | wells of the 24-well plate. |
| 3 | Mix each well to ensure an even suspension and allow the cells to |
| | settle for 5 min on the microscope stage. After this, inspect the |
| | densities under a microscope. |
| 4 | Determine which of the three wells depicts the optimal seeding |
| | density. Then, using that well's volume, plate the remaining cells into |
| | individual wells until no cells remain. |
| 5 | Place the well plate onto an incubator shelf. To ensure even cell |
| | distribution across the bottoms of the culture wells, first move the |
| | plate in a figure-eight motion across the shelf at least three times. |
| | Then, move it in a crisscross pattern at least three times. Afterward, |
| | leave the plate undisturbed until the following day to allow the cells to |
| | fully attach. |

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Prepare Hepatocyte Maintenance Medium

Hepatocyte Maintenance Medium

Base Hepatocyte Maintenance Medium (500 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|--------------|--------|---------------|---------------|--------|----------|
| WEM (without | 490 mL | - | - | Sigma | W1818 |
| phenol red) | | | | | |
| Pen / Strep | 5 mL | 100X | 1% | Sigma | P4333 |
| L-GlutaMAX | 5 mL | 100X | 1% | Gibco | 35050-61 |

- Store the Base Hepatocyte Maintenance Medium at 4°C.
- Use the Base Hepatocyte Maintenance Medium within 30 days of preparation.

Complete Hepatocyte Maintenance Medium (50 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-----------------|--------|---------------|---------------|--------------|----------|
| Base Hepatocyte | 49.445 | - | - | Recipe above | - |
| Maintenance | mL | | | | |
| Medium | | | | | |
| ITS + premix | 500 µL | - | 1% | Sigma | 354352 |
| Ascorbic acid | 50 µL | 50 mg / mL | 0.05 mg / mL | Sigma | 5960 |
| Dexamethasone | 5 µL | 1 mM | 100 nM | Sigma | D4902 |

• Store Complete Hepatocyte Maintenance Medium at 4°C.

• Use the Complete Hepatocyte Maintenance Medium within 3 days of preparation.

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Prepare Overlay Medium

Before YouSlowly thaw the Matrigel aliquot on ice (which should be made slushy with water)Beginfor 30 min or until thawed. Keep the Matrigel aliquot in slushy ice at all times, as this
solution gels rapidly at temperatures above 10°C. To maintain an even coating, use
pre-chilled pipettes, tips, and tubes stored at -20°C prior to use as well as cold
medium during preparation and overlay.

Hepatocyte Overlay Medium

Prepare Hepatocyte Overlay Medium by adding Matrigel to the Complete Hepatocyte Maintenance Medium to achieve a final Matrigel concentration of 0.25 mg/mL.

Example calculation is below:

Hepatocyte Overlay Medium (20 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|---------------------|---------|---------------|---------------|--------------|----------|
| Complete Hepatocyte | 19.5 mL | - | - | Recipe above | - |
| Maintenance Medium | | | | | |
| Matrigel | 0.5 mL | 10 mg / mL | 0.25 mg / mL | Corning | 354234 |

• Keep both the Matrigel aliquot and the Hepatocyte Overlay Medium on slushy ice.

Steps for preparation:

| Step | Action |
|------|--|
| 1 | Prepare the Hepatocyte Overlay Medium by diluting ice-cold, thawed Matrigel into ice- cold Complete Hepatocyte Maintenance Medium using prechilled tips to a final concentration of 0.25 mg / mL, as directed above. |
| 2 | Gently mix the overlay medium well and keep the Hepatocyte Overlay Medium on ice at all times. |

Note: Introducing polymerized Matrigel to the channels of the chip can cause uneven distribution and may impact cell viability and chip performance. If any traces of solidified Matrigel are observed in the Hepatocyte Overlay Medium, discard the entire medium and prepare a fresh batch following the steps above, ensuring all reagents and equipment are kept at 4°C during this step.

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Overlay Hepatocytes

| Steps for | | |
|-------------------------|------|---|
| Overlaying | Step | Action |
| Hepatocytes in Chips | 1 | In the BSC, gently pipette 200 µL of warm Complete Hepatocyte Maintenance Medium through the top channel of each chip to remove cell debris from the hepatocyte monolayer. |
| | 2 | Aspirate the media outflow at the outlets, leaving the media within the channel. |
| | 3 | Incubate the chips for 1 h at 37°C. |
| | 4 | Using cold tips, gently pipette 200 µL of the cold Hepatocyte Overlay Medium to the top channel of each chip, leaving droplets covering both the inlet and outlet ports (See Figure 7). |
| | 5 | Figure 7. Chip with Hepatocyte Overlay Medium covering ports. |
| | 5 | Incubate chips overnight at 37°C. |
| | | |

Steps for Overlaying Hepatocytes in Well Plates

| Step | Action |
|------|--|
| 1 | In the BSC, vigorously swirl the 24-well plate of hepatocytes to |
| 1 | release any cell debris and unattached cells from the monolayer. |
| 2 | Aspirate the medium from each well. |
| 2 | Add 500 µL of cold Hepatocyte Overlay Medium to each well and |
| 3 | return the plate to the incubator to sit overnight. |

Day 1: LSECs to Chip, Chips to Pods, Pods to Zoë

Overview

Goals

- Seed LSECs in the chip
- De-gas and equilibrate media
- Connect chips to Pods
- Connect Pods to Zoë

Required Materials

- LSEC Seeding Medium (at 37°C)
- LSEC Seeding Medium (at 4°C)
- Complete Hepatocyte Maintenance Medium (at 37°C)
- Trypan blue counting solution
- Hemocytometer
- 15 mL conical tube
- Serological pipettes
- Pipettes and filtered tips
- Aspirator and sterile tips
- Ice bucket and ice
- 70% ethanol
- Microscope
- Installed and qualified Zoë
- Prepared chips
- Pods (sterile), 1 per chip
- Tray, 1 per 6 chips
- Steriflip[®] filtration unit: PVDF filter 0.45 µm (sterile)

Key Steps

| Торіс | See Page |
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| Wash Chips | 42 |
| Harvest LSECs | 43 |
| Seed LSECs to Bottom Channel | 44 |
| Prepare LSEC Maintenance Medium | 45 |
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| Prime Pods | 48 |
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| Pods to Zoë | 53 |

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Prepare LSEC Seeding Medium

Goals Seed LSECs into the bottom channel, in the LSEC Seeding Medium.

LSEC Seeding Medium

LSEC Seeding Medium (100 mL)

.....

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-------------------|--------|---------------|---------------|----------------|----------|
| Base LSEC Culture | 90 mL | - | - | Prepared above | - |
| Medium | | | | | |
| FBS | 10 mL | - | 10% | Sigma | F4135 |

- Maintain 35 mL at 37° C and 65 mL at 4° C
- Store the remaining LSEC Seeding Medium at 4°C.
- Use the LSEC Seeding Medium within 3 days of preparation.

Trypan Blue Cell Counting Solution (45 µL)

| Reagent | Volume | Source | Cat. No. |
|---------------------|--------|--------------|----------|
| LSEC Seeding Medium | 40 µL | Recipe Above | - |
| Trypan Blue | 5 µL | Sigma | 93595 |

- Maintain the Trypan Blue Cell Counting Solution at room temperature prior to use.
- Prepare three tubes of the Trypan Blue Cell Counting Solution.
- Always prepare the solution fresh before each use.

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Wash Chips

| Action |
|--|
| Gently pipette 200 µL of warm Complete Hepatocyte Maintenance |
| Medium into the top channel of each chip. Aspirate the outflow, |
| leaving the media in the channel. |
| Gently pipette 200 µL of warm LSEC Seeding Medium to the bottom |
| channel of each chip. Aspirate the outflow, leaving the media in the |
| channel. |
| Return the chips to incubator until the LSECs are ready for seeding. |
| - |

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Harvest LSECs

| Before You | To seed the bottom channel, LSECs in culture must be harvested and counted. |
|------------|--|
| Begin | LSECs should be adjusted to a final seeding density of 4.5×10^6 cells / mL. |

| Step | Action |
|------|---|
| 1 | Bring the culture flask containing LSECs from the incubator the BSC. |
| 2 | Aspirate culture media and add 15 mL of 1X DPBS to wash the |
| 2 | culture surface. Aspirate the DPBS wash. |
| 3 | Add 3 mL of trypsin-EDTA to the flask. Incubate for 2–3 min at 37°C. |
| | Tap the side of the flask gently. Inspect the culture under the |
| 4 | microscope to verify complete cell detachment from the culture |
| | surface. |
| 5 | Add 9 mL of warm LSEC Seeding Medium to the flask, and pipette |
| 0 | gently to mix while collecting all cells from the culture surface. |
| 6 | Transfer the contents of the flask (12 mL) into a sterile 15 mL conical |
| 0 | tube. |
| 7 | Add 3 mL of LSEC Seeding Medium the tube, bringing the total |
| · · | volume to 15 mL. |
| 8 | Centrifuge LSECs at 200 x g for 5 min at room temperature. |
| | Carefully aspirate the supernatant, leaving approximately 100 μ L of |
| 9 | medium above the cell pellet. |
| | Note: The cell pellet will be very small, so be sure to aspirate gently. |
| 10 | Loosen the cell pellet by flicking the tube gently. |
| 11 | Using a P1000 pipette, gently resuspend the cells by adding 100 μ L |
| | of cold LSEC Seeding Medium. |
| | Pipette gently to create a homogenous mixture, and transfer 5 μ L of |
| 12 | the cell suspension to the Trypan Blue Counting Solution. This will |
| | create a 1:10 dilution. |
| 13 | Mix the counting solution thoroughly. Count the cells using a |
| | hemocytometer (See "Cell Counting and Viability Assessment"). |
| 14 | Dilute the LSECs to 4.5 x 10 ⁶ cells / mL. |
| 15 | Keep the LSEC cell suspension on ice until you are ready to seed the |
| | chips. |

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| | | | | |

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Seed LSECs to Bottom Channel

Before YouWork with one chip at a time. After seeding the first chip, assess the cell density
within the channel using a microscope. Adjust the density of the cell suspension
accordingly for the next chips if necessary. After all six chips have been seeded,
immediately invert the chips.

| Step | Action |
|------|---|
| | Working with one chip, seed 15–20 μ L of the LSEC cell suspension |
| | into the bottom channel while aspirating the outflow. |
| | Assess the cell density within the channel using a microscope. Make |
| 1 | any necessary adjustments to the density of the cell suspension for |
| | the next chips. Seed remaining chips. Proceed to Step 2 immediately |
| | Note: There is additional resistance while pipetting the bottom |
| | channel in Chip-R1 compared to Chip-S1 (see Page 13). |
| | After seeding, aspirate the DPBS from the reservoir and invert the |
| 2 | Chip Cradle (see Figure 8). |
| 2 | Note: Each Chip Cradle can support up to six chips inside a Square |
| | Cell Culture Dish (120 x 120 mm). |
| | Place a small reservoir (15 mL conical tube cap containing sterile |
| 3 | DPBS) inside the Square Cell Culture Dish (120 x 120 mm) to provide |
| | humidity for the cells. Replace the dish lid. |
| | |
| | |

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Prepare LSEC Maintenance Medium

LSEC Maintenance Medium

LSEC Maintenance Medium (100 mL)

| Reagent | Volume | Conc. [Stock] | Conc. [Final] | Source | Cat. No. |
|-------------------|--------|---------------|---------------|--------|----------|
| Base LSEC Culture | 98 mL | - | - | - | - |
| Medium | | | | | |
| FBS | 2 mL | - | 2% | Sigma | F4135 |

- Store at 4°C.
- Use the LSEC Maintenance Medium within three days of preparation.

Note: If the seeding density is low, you can use 10% FBS for 24–48 h and then switch to 2% FBS.

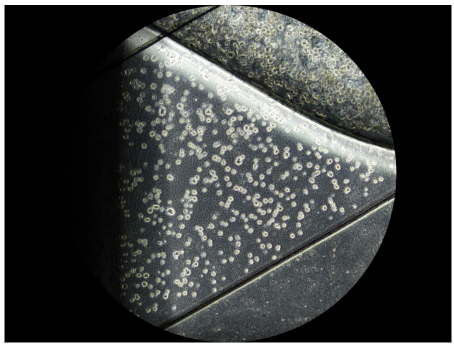


Figure 9. If seeding density is low, continued use of 10% FBS is recommended. If seeding density is optimal (above), culture cells with 2% FBS.

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Gas Equilibration of Media

CAUTION



The media equilibration step is imperative to successfully culture Organ-Chips. Omitting this step will cause bubbles to form in the chip, in the Pod, or both, which will in turn negatively impact cell viability and result in irregular flow

Before You Begin

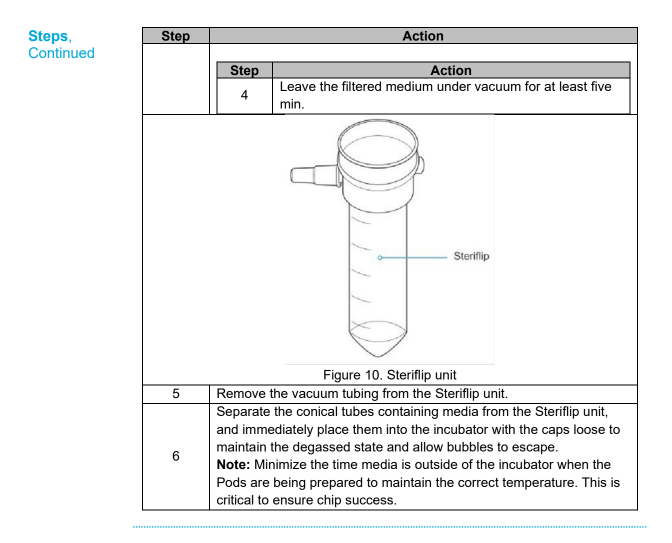
- Once the culture media for both chip channels have been warmed for at least 1 h, the gas equilibration step can be performed.
- Ensure the medium is outside of a warmed environment (such as an incubator or bath) for no longer than 10 min, as gas equilibrium can become compromised when medium is allowed to cool.
- If the vacuum pump is not located close to the water bath (e.g., inside the BSC), it is recommended to place some clean water warmed at 37°C inside the BSC to minimize cooling during the media equilibration step.

Steps

| Step | Action |
|------|--|
| 1 | east 3.3 mL of Complete Hepatocyte Maintenance Medium chip in a 50 mL conical tube. |
| 2 | east 3.3 mL of LSEC Maintenance Medium for each chip in the 50 mL conical tube. |
| 3 | th 50 mL conical tubes of media at 37°C in a water bath or n for at least 1 h. |
| 4 | Action Attach each conical tube containing warmed media to a Steriflip unit (See Figure 10). With the unit "right-side up" (medium in the bottom conical tube), apply vacuum for 10 seconds. Invert the Steriflip-connected tubes, and check that the medium begins to pass from the top conical tube to the bottom tube. Note: The vacuum must operate at a minimum of -70 kPa. At this pressure, it should take about 2 seconds for 10 mL of media to flow through the filter. If it takes longer, |
| | stop and refer to "Media take too long to pass through Steriflip" in the troubleshooting section. |

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Gas Equilibration of Media, Continued



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Prime Pods

CAUTION



Ensuring successful Pod priming is best practice for successful liquid-liquid connection between the chip and Pod. Failure to do so can cause bubbles to form in the chip, in the Pod, or both, which will in turn negatively impact cell viability and result in irregular flow.

Steps

| Step | Action |
|---------|---|
| 1 | Sanitize the exterior of the Pod packaging and trays with 70% |
| I | ethanol, wipe them, and transfer them to the BSC. |
| 2 | Open the Pod package and place the Pods into the trays. Orient the |
| Z | Pods with the reservoirs facing the back of the tray (See Figure 11). |
| Tray Ha | Pods Indle Figure 11. Chips and Pods inserted into a tray. |
| | Pipette 3 mL of pre-equilibrated, warm media to each inlet reservoir. |
| | In the top channel inlet reservoir, add Complete Hepatocyte |
| 3 | Maintenance Medium; in the bottom channel inlet reservoir, add |
| | LSEC Maintenance Medium. |
| 4 | Pipette 300 μ L of pre-equilibrated, warm media to each outlet |
| 4 | reservoir, directly over each outlet Via (see Figure 12). |

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Prime Pods, Continued

| Steps, | Step | Action |
|-----------|------|--|
| Continued | | Bottom Channel Top Channel |
| | | |
| | | Via |
| | | Figure 10: Tag view of a Dad |
| - | | Figure 12: Top view of a Pod. Bring trays with Pods to the incubator and slide them completely into |
| | 5 | Zoë with the tray handle facing outward. |
| | 6 | Before running the Prime Cycle, confirm the Zoë is set to the correct chip type. If the incorrect chip is active, use the dial to select the Chip-R1 consumable on the Zoë UI (See Page 2). |
| | | Run the Prime Cycle on Zoë. |
| | | |
| | | Step Action |
| | | 1 Use the Dial to highlight "Prime" on the display. |
| | 7 | 2 Press the Dial Button to select "Prime." |
| | 7 | 3 Rotate the Dial clockwise to highlight "Start." |
| | | 4 Press the Dial again to select "Start" and begin. |
| | | Note: Once "Start" is selected, there will be a sound as Zoë engages the Pods. |
| | | Close the incubator door and allow Zoë to prime the Pods; this |
| | 8 | process takes approximately one min. |
| | | Note: The status bar will read "Ready," if the Prime Cycle is complete. |
| | 9 | Remove the tray from Zoë and bring them to the BSC. |
| | 10 | Verify that the Pods were successfully primed. This is critical for |
| | .0 | success. |

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Prime Pods, Continued

| Pod Priming | Take out the tray and inspect the top of the Pods (See Figure 13) to verify the |
|--------------|--|
| Verification | presence of small media droplets through the Pod window at all four fluidic ports. |

| lf | Then |
|--|---|
| Droplets are not visible through the top window | Re-run the Prime Cycle on those Pods. |
| | If the issue persists, contact Emulate Support. |
| Any outlet port does not show a droplet, but the | Ensure Step 4 of "Priming Steps" has been |
| inlet port does. | performed correctly. |
| Any media escaped onto the tray (this | Clean the tray using a wipe sprayed with 70% |
| may occur more often by the outlet ports). | ethanol. |

Figure 13

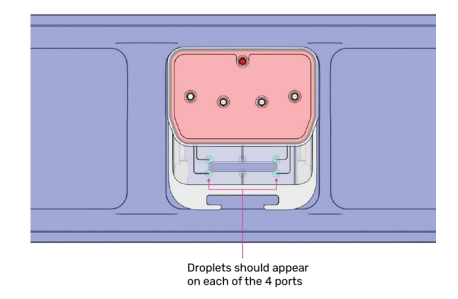


Figure 13. Top view of chip in Pod with fluidic posts covered with droplets.

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Wash Chips

| Action |
|--|
| Transfer the seeded chips in the Square Cell Culture Dish (120 x 120 |
| mm) from the incubator to the BSC. |
| Gently wash each top channel with warm, equilibrated Complete |
| Hepatocyte Maintenance Medium to remove any bubbles. |
| Place small droplets of equilibrated Complete Hepatocyte |
| Maintenance Medium on each chip's top channel inlet and outlet port. |
| Gently wash each chip's bottom channel with warm, equilibrated |
| LSEC Maintenance Medium to remove any possible bubbles as well |
| as to replace with de-gassed media. |
| Place small droplets of equilibrated LSEC Maintenance Medium on |
| each chip's bottom channel inlet and outlet ports. |
| |

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Chips to Pods

| Step | Action |
|------|---|
| | Hold one chip (in a chip carrier) in the dominant hand and one Pod in |
| 1 | the other hand. Slide the chip carrier into the tracks on the underside |
| | of the Pod until the chip carrier has fully seated. |
| 0 | Place a thumb on the chip carrier tab. Gently, but firmly, press the tab |
| 2 | in and up until it engages with the Pod. |
| | Aspirate any excess media on the chip surface through the Pod |
| | window. |
| | |
| | Note: This is a crucial step as media can easily fall into the recess of |
| 3 | the Chip-R1 gasket during connection or at any point when leaving |
| | droplets over the ports (i.e., during ECM coating). DO NOT drag the |
| | aspirator tip across the top of the chip as this can cause scratches on |
| | the top layer. Instead, aspirate against the corners and edges of the |
| | chip gasket window. |
| | Figure 14: Aspirating excess media from chip gasket window. |
| | Place the Pod and connected chip onto the tray. Additionally, clean all |
| 4 | excess media from the trays and bottoms of Pods using a wipe |
| | sprayed with 70% ethanol. |
| 5 | Repeat Steps 1–4 for each Pod and chip carrier. |
| 0 | Confirm that there is sufficient media in each Pod inlet and outlet |
| 6 | reservoir. Also Ensure that the Pod lids are flat and secure. |
| -7 | Bring the tray to the incubator until all samples are connected to keep |
| 7 | the equilibrated media warm. |
| | · · |

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Pods to Zoë

Steps

| Step | Actio | on | | | |
|------|---|-------------------------------------|--|--|--|
| | Immediately place the trays holding | | | | |
| 1 | prevent media from cooling and losing its gas equilibration. | | | | |
| | Program the appropriate Organ-Chip culture conditions on Zoë. | | | | |
| 2 | These conditions will start as soon as the Regulate Cycle is complete | | | | |
| 2 | For the Liver-Chip R1, set the flow ra | ate 30 μ L / h for both top and | | | |
| | bottom channels. | | | | |
| | Run the Regulate Cycle: | | | | |
| | | Action | | | |
| | 1 Using the Dial, highlight the | | | | |
| | | elect "Regulate," and rotate the | | | |
| | Dial clockwise to "Start." | | | | |
| | 3 Press the Dial again to se | - | | | |
| | Regulate Cycle (see Figu | | | | |
| | | ed, there will be a sound as Zoë | | | |
| | engages the Pods. | | | | |
| | | 군 ZOE 123456 | | | |
| | Prime Cycle | | | | |
| | Regulate Cycle Start | | | | |
| 3 | Flow 🗁 Chip-R1 | Stretch | | | |
| | LOCATION MEDIA RATE | PARAMETER RATE | | | |
| | Top Liquid 30 µL/h | Strain 0 % | | | |
| | Bottom Liquid 30 µL/h | Frequency 0 Hz | | | |
| | | | | | |
| | | | | | |
| | | U | | | |
| | Figure 15. Zoë UI showing Regulate Selection | | | | |
| | 4 Make sure the "Activation | " button is glowing blue. | | | |
| | | | | | |
| | | | | | |

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Pods to Zoë, Continued

| Steps, | Step | | | Action | | |
|-----------|------|--|--|---------------------|-------------------|------------------|
| Continued | • | The Reg | ulate Cycle takes | s 2 h to finish. Th | ere is a prog | ress bar at the |
| | | top of the | e UI screen (see | Figure 16). | | |
| | | Step | | Actio | n | |
| | | | | | 융 | ZOE 123456 |
| | | Р | rime Cycle | | | |
| | | R | egulate Cycle | Start | | |
| | | F | low 🗟 Chip-R1 | | Stretch | |
| | | L | DCATION MEDIA F | ATE | PARAMETER | RATE |
| | | Т | op Liquid | 30 µL/h | Strain | 0 % |
| | | в | ottom Liquid | 30 µL/h | Frequency | O Hz |
| | | | | | | |
| | | | | | | |
| | | | | | | (j) |
| | | Figur | ra 16. 7aë III.aar | oon obouring Doo | nulata Cuala | |
| | | - | | | | in progress. |
| | | | | • | | ause Zoë by |
| | 4 | | • • • | - | • • | - |
| | | · - | | | | |
| | | Remove the Pod lids. Gently aspirate media from all four Pod | | | | |
| | 5 | | - | • | till covers th | e reservoir Vias |
| | | | | | annol cnoci | fic worm |
| | | | | | | |
| | 6 | 6 | | | | |
| | | - | | | | |
| | | | | | • | |
| | 7 | • | , , | • | , Zoë will be | gin flow at the |
| | | - | Prime Cycle Start Flow Chip-R1 Stretch Ioocation MEDIA Rate Ioocation Bottom Liquid 30 m/h Ioocation Ioocation Bottom Liquid 30 m/h Ioocation Ioocation Ioocation Bottom Liquid 30 m/h Ioocation Ioocation Ioocation Ioocation Figure 16. Zoë UI screen showing Regulate Cycle in progress. Frequency 0 m/h Ioocation Ioocatiocation Ioocation | accord follow | | |
| | | | | liate Cycle, nowe | ever, il it is ne | ecessary, ioliow |
| | | 11030 310 | ,ps. | | | |
| | | Step | | Actio | n | |
| | | 1 | - | | - | |
| | Note | 2 | | | - | and rotate the |
| | | | | | | weit 4 min for |
| | | | | - | | |
| | | | • | | | |
| | | | | | | |
| | | | | 5 | | |

Day 2+: Chip Maintenance and Sampling

Overview

Goals

- Maintain chips in Zoë
- Cell culture inspection
- Collect samples for analysis

Required Materials

Chips in PodsCell culture media

CAUTION



Failure to replenish media on time will cause air to be introduced into the Pod and chip, which ultimately results in cell death and may cause damage to Zoë Culture Module. Do not fill reservoir past 4 mL of the total volume during replenishment.

Steps

| Step | Action | | | | |
|------|---|--|--|--|--|
| 1 | Pause Zoë by pressing the silver "Activation" button. | | | | |
| 2 | Remove the trays and place them into the BSC. | | | | |
| 3 | Visually inspect each chip for bubbles. | | | | |
| 4 | Using a microscope, assess the morphology and viability of cells in the chips. Capture representative images at 10X or 20X magnification at the following locations (see Figure 17): | | | | |
| | Inlet Junction | | | | |
| | Center of Channel | | | | |
| | Outlet Junction | | | | |
| | Outlet Junctions Center Inlet Junctions Image: Control of the second | | | | |
| 5 | 5 Remove Pod lids and collect effluent from the Pod outlet reservoirs while not disturbing the Pod reservoir Vias. | | | | |

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Overview, Continued

| Steps, | Step | |
|-----------|------|----------------------------------|
| Continued | | Gently aspirate any mediun |
| | 6 | thin liquid film still covers th |
| | | introduced into them. |
| | | Refill the Pod reservoirs wit |
| | 7 | medium. Then, perform a V |
| | 1 | in the reservoir directly over |

| 6 | Gently aspirate any medium not collected for analysis, ensuring that a thin liquid film still covers the reservoir Vias so that no air is introduced into them. |
|---|--|
| 7 | Refill the Pod reservoirs with the appropriate fresh cell culture medium. Then, perform a Via wash by pipetting 200 μ L of the medium in the reservoir directly over the top of the Via to dislodge any bubbles. |
| 8 | Place the Pod lids back on and return the trays to Zoë. |
| 9 | Press the silver "Activation" button to resume pre-set Organ-Chip culture conditions. Zoë will engage when the "Activation" button glows blue. |

Action

Part VI: Troubleshooting

Troubleshooting

Troubleshooting Table

| Issue | Section | Step | Recommendation | |
|-----------------|---------------------|---------------------|---|--|
| Bubbles are | Any step related to | Any step related to | Wash the channel with the appropriate | |
| present in | chip handling, | chip handling, | solution until all bubbles have been | |
| channel | such as ECM | such as ECM | removed. If bubbles persist, it may be | |
| | coating and cell | coating and cell | helpful to aspirate the channel dry and | |
| | seeding. | seeding. | slowly re-introduce solution. | |
| Bubbles in the | Any step related to | Any step related to | Dislodge bubbles using pipette tip or | |
| ports upon | chip handling, | chip handling, | aspirate the channels and reintroduce | |
| introduction of | such as ECM | such as ECM | appropriate media. | |
| media into the | coating, and cell | coating and cell | | |
| chip | seeding. | seeding. | | |
| Media takes | Chips to Pods and | Equilibration of | Vacuum pressure is not reaching | |
| too long to | Pods to Zoë | Media | -70kPa. Find an alternate vacuum | |
| pass through | | | source with the appropriate pressure. | |
| Steriflip | | | | |
| Pods do not | Chips to Pods and | Prime Pods | If Pods do not prime on the first | |
| prime | Pods to Zoë | | attempt, ensure that medium covers all | |
| | | | Pods Vias, and run the Prime Cycle | |
| | | | again. If the problems persist, record | |
| | | | the Pod lot number, and replace it with | |
| | | | a new Pod. | |
| Screen is | Chips to Pods and | Any step related to | Power off Zoë and turn it on again. If | |
| frozen or | Pods to Zoë; | Organ-Chip | the problem persists, contact our | |
| unresponsive | Maintaining and | culture on Zoë | support team. | |
| | Sampling | | | |
| Pods are stuck | Maintaining and | Any step related to | The Pod lid is not secured. Try | |
| in Zoë | Sampling | Organ-Chip | wiggling the tray to the right and left as | |
| | | culture on Zoë | you slide it out while keeping it level. If | |
| | | | the problem persists, contact our | |
| | | | support team. | |
| Pods not | Maintaining and | Maintenance and | There is inherent variability with Zoë; | |
| flowing | Sampling | Regulate | however, large fluctuations and major | |
| properly or | | | flow issues primarily result from | |
| evenly/ | | | bubbles. To remove bubbles and allow | |
| Bubbles | | | for flow, remove the chip from the Pod, | |
| observed in | | | flush the chip with media, re-prime the | |
| chip | | | Pod with media, and reconnect the | |
| | | | chip to the Pod. | |

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Troubleshooting, Continued

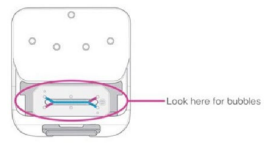
Potential Root If there is a high failure rate due to bubbles, or if bubbles are persistent, despite Causes of performing the above mitigation step (See Figure 18), check for the following: **Bubbles**

| lf | Then | |
|--|---|--|
| Medium is not sufficiently equilibrated | Be sure to follow media preparations steps in | |
| before adding to Pods | section "Gas Equilibration of Media." | |
| Vacuum for Steriflip too weak | Ensure that media passes through the Steriflip in | |
| | ~10 seconds. | |
| Incorrect Steriflip used | Confirm the correct Steriflip unit is being used | |
| | (Millipore SE1M003M00). | |
| Medium not warmed correctly before Steriflip | Be sure to follow the media preparation steps in | |
| step | the section "Gas Equilibration of Media". | |
| Insufficient priming occurring | Disconnect the chip and re-prime the Pod. | |
| | Ensure media droplets are present in all ports. | |

Figure 18

Images of Bubbles in an Organ-Chip

Images of Bubbles in Chip







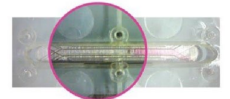
Small bubbles in Top Channel



Both Channels dry



Bubble plug in Bottom Channel



Bubble plug in Top Channel



Bubble plug in Top Seeding

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