

Tube Formation Assays With the μ -Plate Angiogenesis 96 Well

1. Introduction

The proven well-in-a-well format of the μ -Slide Angiogenesis is also available in a 96 well version. The special geometry provides a meniscus-free surface consisting of both gel and overlaid medium, which allows for high quality phase contrast imaging, and a homogeneous cell distribution.

This Application Note describes the handling of the μ -Plate Angiogenesis 96 Well with Matrigel™ and a multichannel pipette.

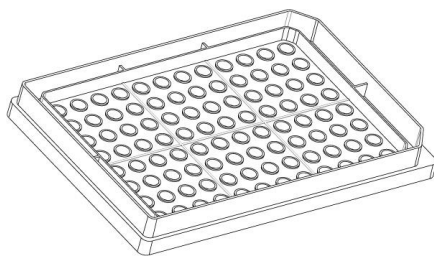


Fig. 1: μ -Plate Angiogenesis 96 Well

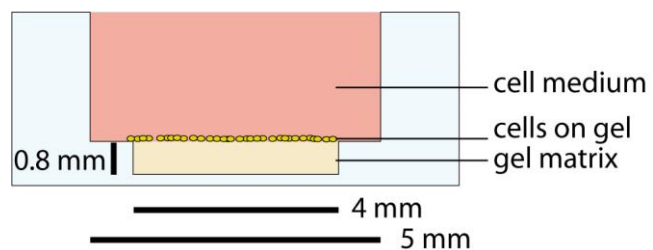


Fig. 2: Cross section of a single well

2. Material

For a Tube Formation Assay with the μ -Plate Angiogenesis 96 Well, the following material is needed:

Cells:	HUVEC (PromoCell, C-12200, C-12203)	1 x 10 ⁴ per well
Medium:	Endothelial Cell Growth Medium (PromoCell, C-22010)	70 μ l per well
Gel matrix:	Corning® Matrigel® Growth Factor Reduced, (#356231)	10 μ l per well
Plate:	μ -Plate Angiogenesis 96 Well, ibiTreat (ibidi, 89646)	1 plate
Detach reagent:	Accutase (PromoCell, C-41310)	8 ml per T75 flask

Further required equipment:

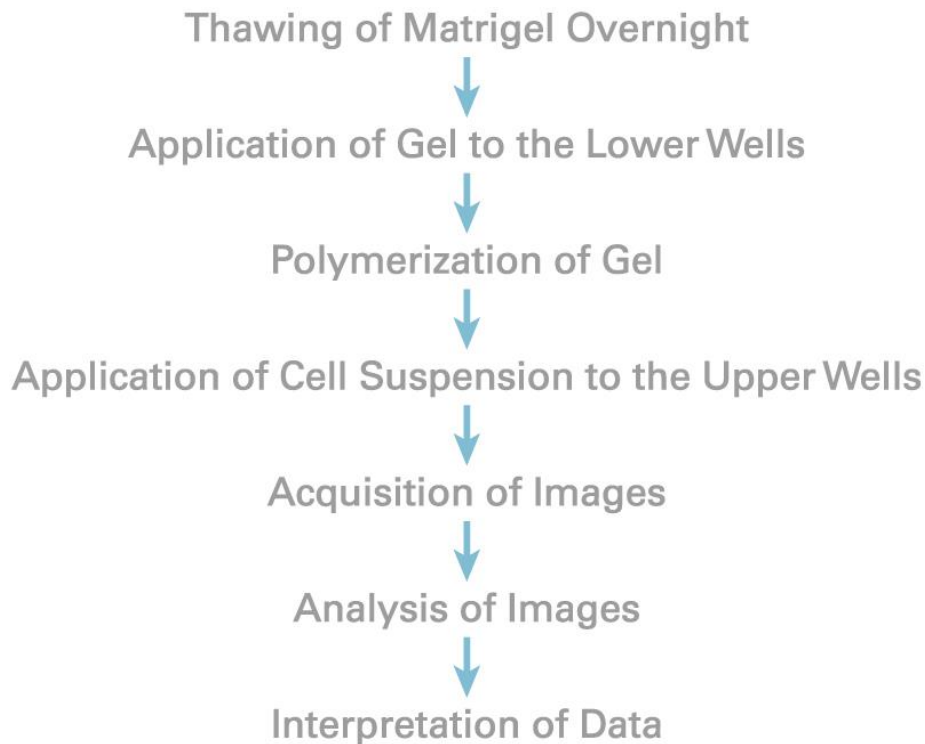
- Scale paper
- Cooling rack or crushed ice
- PCR stripe to prepare 8 aliquots for the multi-channel pipette
- Multi-channel pipette with tips

Please note: The volume of the medium and gel that is mentioned above is for one well only. You need to calculate the total amount of your required volume by multiplying it with the number of wells that will be used. Also, pipetting with the multi-channel pipette will require some additional liquid to ensure the correct uptake for all of the wells.

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3. Work Flow

The protocol steps are the same as in Application Note 19 "Tube Formation in μ -Slide Angiogenesis":



4. Preparation of Gel and Plate

Gel Application

To use the gel with a multi-channel pipette, it must be distributed to the 8 small tubes of a PCR-stripe (alternatively in one column of wells of a 96-well plate with conical bottom), and cooled on ice.

Add at least 10 μ l more gel than you actually need to fill each of the wells.

Example: For pipetting the whole μ -Plate Angiogenesis 96 well (12 columns), add 130 μ l to each of the 8 wells.



Fig.3: PCR-stripe with Matrigel on ice

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Follow these steps:

- 1) The day before seeding the cells, place the Matrigel™ on ice in the refrigerator at 4°C. The gel can slowly thaw overnight.

Note: Always use precooled pipette tips (4°C) for pipetting the gel!

- 2) Place the PCR-strip (or 96-well plate) and the vial with the thawed gel on ice or in any appropriately cooled stage. Transfer the required gel volumes to the wells of the PCR-stripe with a cooled pipette tip.
- 3) Remove the μ -Plate Angiogenesis 96 Well from the sterile packaging and pull off the protection foil (see Figure 4).
- 4) Adjust your multi-channel pipette to 10 μ l and take up the gel (see Figure 5).
- 5) When applying the gel to the wells of the μ -Plate Angiogenesis 96 Well, hold the pipette upright to prohibit the liquid from flowing into the edge of the upper well (see Figure 6).

Pipetting Tips

To avoid air bubbles, make three uptake and push-out movements (10 μ l) with the pipette, while leaving the tip in the gel. Then transfer 10 μ l aliquots to the wells.

Due to the high viscosity of Matrigel™, it might be necessary to adjust the pipet volume to slightly more or less than 10 μ l (see [Application Note 19 "Tube Formation Assays in \$\mu\$ -Slide Angiogenesis"](#)).

To control the right amount of gel, observe the gel in a phase contrast microscope with a low magnification (e.g., 4x).

Be sure to finish the whole pipetting procedure in a few minutes, so as to avoid evaporation effects.

Gelation

After filling all the required wells with Matrigel™, cover the μ -Plate Angiogenesis with the supplied lid and place it in the incubator to allow for the polymerization of the gel (30–60 minutes).

To help minimize evaporation, the side reservoirs can be filled with sterile water. Fill only half of the volume of the reservoirs to avoid any overflow of the liquid into the wells.

During the polymerization of the gel, prepare the cell suspension.

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Fig.4: Pull off the protection foil.

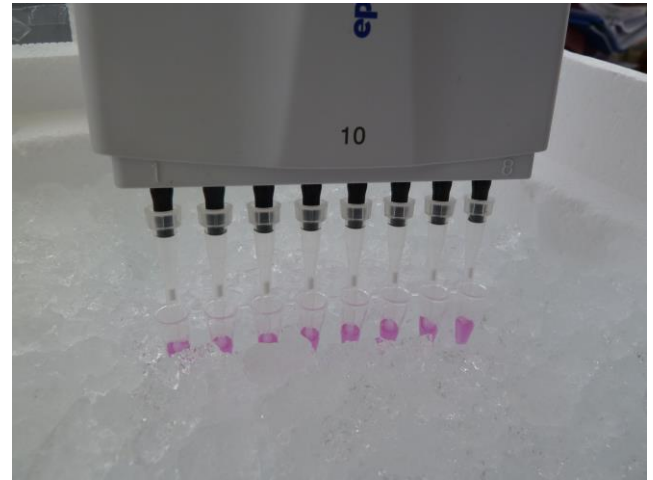


Fig.5: Take in 10 µl of gel from the PCR-stripe.

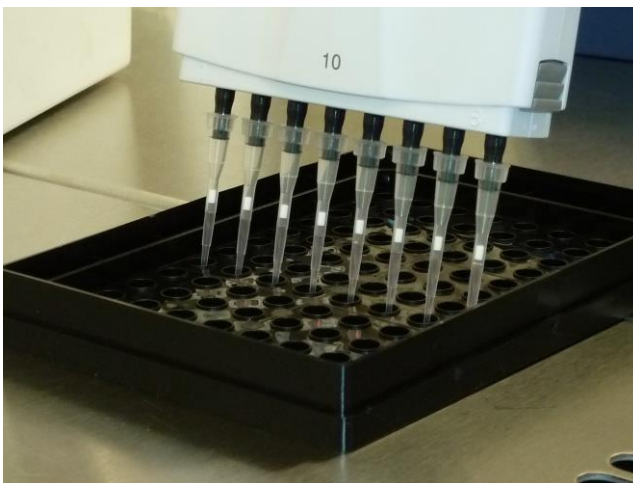


Fig.6: Apply 10 µl of Matrigel™ to the lower well.

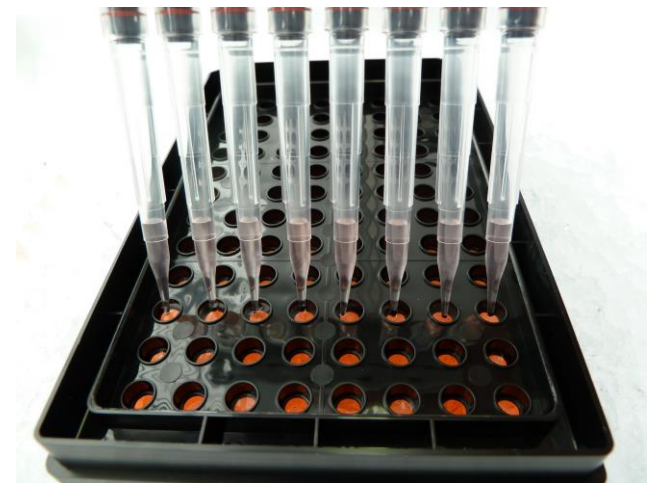


Fig.7: Apply 70 µl of cell suspension to the upper well.

5. Seeding Cells

The number of cells seeded onto the surface of the gel is a crucial parameter to obtain reliable results. The cell type and size determine the number of cells that are needed. For the best results, optimize the cell seeding number before starting an experimental series.

Follow these steps:

- 1) For a final cell number of 10,000 cells per well, adjust a cell suspension of 1.4×10^5 cells/ml. Then mix thoroughly.
- 2) Take the µ-Plate from the incubator.
- 3) Apply 70 µl of cell suspension to each upper well. Keep the pipet tips upright and take care not to touch the gel with the pipet tip.
- 4) Close the µ-Plate Angiogenesis with the lid. The plate is now ready for observation.

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- 5) After a few minutes, the cells will have sunk down to the gel surface and will be lying in one plane. However, due to the geometry of the wells, some cells will also be found on the plastic margins (not only on the gel).

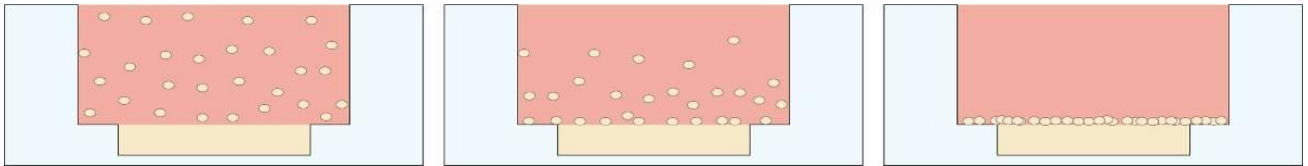


Fig.8: Sinking process of cells. After some minutes the cells will have fallen to the gel surface.

6. Observation on the Microscope

The network formation on top of the gel surface will follow a certain development characteristic, depending on your cell type. When observing the total tube length, for example, the parameters will rise to a maximum value during the first few hours after seeding, and then slowly decline in the following hours, until reaching a base value.

Immediately after seeding the cells, position the plate on an inverted microscope that is equipped with an incubation chamber. Program the positions you want to observe on your imaging system, and then start a time-lapse recording. For HUVEC, we recommend using a small magnification (4x or 10x) and a time interval of one hour in between the single images. Note: It is possible that cells will migrate into the gel and change the focal plane. Therefore, it is recommended to use a software autofocus program to get sharp pictures over an elapsed amount of time.

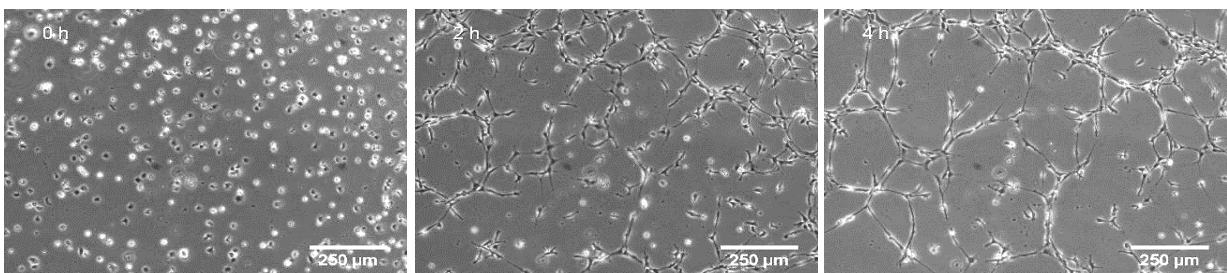


Fig.9: Time lapse pictures with a 10x magnification at 0, 2, and 4 hours.

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7. Data Analysis

For reproducible results and a fast and objective data analysis, we recommend using the automated [Tube Formation ACAS Image Analysis](#) system. Using this web-based platform, you can conveniently upload your images and your analysis results will be ready for download within minutes.

In ACAS, the images are automatically analyzed considering a variety of parameters, such as loop count, average loop area, average tube length (per loop), total tube length and branch count.

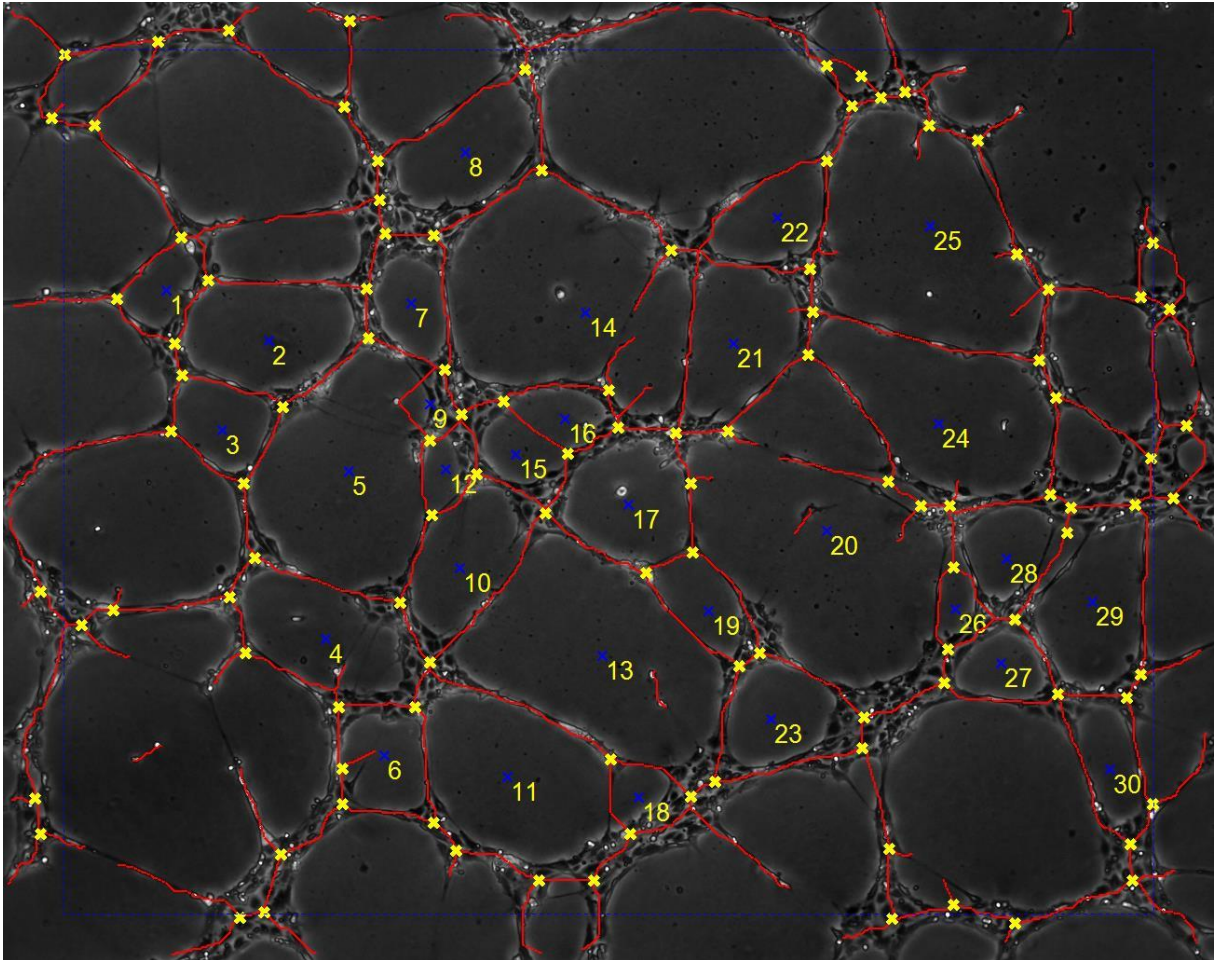


Fig.10: Data analysis with the ACAS Tube Formation Module. The tubes are shown in red and the branching points are shown as yellow marks. The numbers indicate a closed loop of tubes with the blue mark as the center of the loop.