



The ibidi product family is comprised of a variety of μ -Slides and μ -Dishes, which have all been designed for high–end microscopic analysis of fixed or living cells. The high optical quality of the material is similar to that of glass, so you can perform all kinds of fluorescence experiments with uncompromised resolution and choice of wavelength.

The μ -Slide Angiogenesis is a cell culture product for angiogenesis assays and direct cell culture. Cells can be grown on gel matrices, e.g. collagen gels, hyaluronic gels or BD MatrigelTM (Becton–Dickinson) or directly on the ibidi Standard Bottom.

Material

ibidi μ -Slides, μ -Dishes, and μ -Plates are made of a plastic that has the highest optical quality. The bottom material exhibits extremely low birefringence and autofluorescence, similar to that of glass. Also, it is not possible to detach the bottom from the upper part. The μ -Slides, μ -Dishes, and μ -Plates are not autoclavable, since they are only temperature–stable up to 80°C/175°F. Please note that gas exchange between the medium and incubator's atmosphere occurs partially through the polymer coverslip, which should not be covered.

Optical Properties ibidi Standard Bottom		
Refractive index n _D (589 nm)	1.52	
Abbe number	56	
Thickness	No. 1.5 (180 μm)	
Material	microscopy plastic/ polymer coverslip	

Please note! The ibidi standard bottom is compatible with certain types of immersion oil only. A list of suitable oils can be found on page 3.

Surfaces

The μ -Slide Angiogenesis is available with ibiTreat and the uncoated surface. The ibiTreat surface is a physical treatment and optimized for adhesion of most cell types. Many cell lines as well as primary cells were tested for good cell growth. Uncoated is a very hydrophobic surface and allows no direct cell growth.

A specific coating of the μ -Slide Angiogenesis can be done yourself following the procedure in section "Coating".

Geometry

The μ -Slide Angiogenesis provides standard slide format according to ISO 8037/1. The well to well distance of 9 mm (like 96 well plates) allows using multichannel pipettes.

Geometry of the µ–Slide Angiogenesis			
Number of wells	15		
Volume inner well	10 µl		
Diameter inner well	4 mm		
Depth inner well	0.8 mm		
Volume upper well	50 µl		
Diameter upper well	5 mm		
Growth area inner well	0.125 cm^2		
Coating area using 10 µl	0.23 cm^2		
Bottom	ibidi Standard Bottom		

Shipping and Storage

The μ -Slides, μ -Dishes and μ -Plates are sterilized and welded in a gas-permeable packaging. The shelf life under proper storage conditions (in a dry place, no direct sunlight) is listed in the following table.

Conditions			
Shipping conditions Storage conditions	Ambient RT (15-25°C)		
Shelf Life of Different Surfaces			
ibiTreat, glass bottom, ESS Collagen, Poly-Lysine	36 months 18 months		
Fibronectin	4 months		

Instructions



Coating

The uncoated μ -Slide must be coated to promote cell adhesion. If you want to establish a certain coating to match your needs, we recommend testing your coating procedure on both uncoated and ibiTreat μ -Slides, since we have observed that some biomolecules adhere differently to hydrophobic and hydrophilic plastic surfaces.

In tube formation assays the μ -Slide Angiogenesis is coated with a 0.8 mm thick layer of gel matrix.

- 1. Prepare your gel matrix according to the manufacturer's protocol or reference.
- 2. Fill the inner well with 10 µl liquid gel. Avoid air bubbles.
- 3. Let the gel polymerize under appropriate conditions.
- 4. Use as soon as possible.
- 5. If storage is needed fill sterile water around the wells to generate a humidified environment to hinder evaporation.

Non-gel based coatings are also possible. Please use $10 \ \mu$ l coating solution and calculate with an area to be coated of 0.23 cm² per well. Further information about coatings is provided in Application Note 08 "Cell culture coating".

Seeding Cells

In a tube formation assay cells are seeded on top of the polymerized gel matrix:



- 1. Trypsinize and count cells as usual. Dilute the cell suspension to the desired concentration. Depending on your cell type, we recommend $1-3 \times 10^5$ cells/ml.
- 2. Apply 50 µl of the cell suspension into the upper well. Do not touch the gel matrix with the pipet tip.
- 3. Cover the μ -Slide Angiogenesis with the supplied lid. Incubate at 37°C and 5 % CO₂ as usual.
- 4. Conduct your experiment.
- 5. Depending on the cell type, medium exchange is necessary every 1–2 days. Carefully aspirate the old medium and replace it by $50 \,\mu$ l fresh medium.

For a detailed protocol please refer to Application Note 19 "Tube Formation" and Application Note 5 "Tube Formation in μ -Plate Angiogenesis 96 well".

Further information about the optimization of experimental parameters and data analysis is provided in Application Note 27 "Tube Formation - Data Analysis".

Tip:

Air bubbles in the gel can be reduced by equilibrating the μ -Slide Angiogenesis before usage inside the incubator overnight.

Tip:

For less evaporation the space in-between the wells can be filled with sterile water or agarose. Add agarose to water or buffer solution (e.g. 0.1 g to 10 ml water). Melt agarose solution using a microwave or boiling water bath and allow the solution to cool to \sim 50°C.

Tip:

In case bent gel surfaces are created, increase or decrease the amount of gel used, until you get flat and even gels.

Experimental Setups

Alternatively, the μ -Slide Angiogenesis can be used for the following assays:

• Culture cells without a gel matrix directly in the minor wells.



• Fill the inner well with cells suspended inside a gel matrix. After gel polymerization, add 50 µl cell–free medium to fill the upper well.



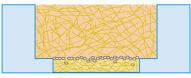
• Fill the inner well with a gel matrix and culture pieces of tissue or cell spheroids on it.

Instructions





• Sandwich Cell Culture: Fill the inner well with a gel matrix. Seed cells on top of the polymerized gel and imbed the cells with 50 µl gel in the upper well.



• Fill the inner well with a low volume of the gel matrix, e.g. 8 µl. Seed cells on top of the polymerized gel. If necessary gently shake the slide to make the cells slide into the middle of the well.



• Fill the inner well with fibroblasts suspended inside a gel matrix. Seed cells on top of the polymerized gel. Overlay the cell layer with medium and incubate for invasion of the cells into the gel matrix.



Preparation for Cell Microscopy

When gel matrices are used the optical quality and the use of high magnification objective lenses might be restricted. Without any gel cells can be observed live or fixed directly in the wells on an inverted microscope. You can use any fixative of your choice. The plastic material is compatible with a variety of chemicals, e.g. Acetone or Methanol. Further specifications can be found at www.ibidi.com. Due to the thin bottom of only 180 µm, high resolution microscopy is possible.

Immersion Oil

When using oil immersion objectives, use only the immersion oils specified in the table. The use of a nonrecommended oil could lead to the damage of the plastic material and the objective.

Company	Product	Ordering Number
ibidi	Immersion Oil	(ibidi) 50101
Zeiss	Immersol 518 F	(Zeiss) 444960
Zeiss	Immersol W 2010	(Zeiss) 444969
Leica	Immersion liquid	(Leica) 11513859

References

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J. Searle, M. Mockel, S. Gwosc, S. A. Datwyler, F. Qadri, G. I. Albert, F. Holert, A. Isbruch, L. Klug, and D. N. Muller. Heparin Strongly Induces Soluble fms-Like Tyrosine Kinase 1 Release In Vivo and In Vitro. *Arteriosclerosis, Thrombosis, and Vascular Biology*, 2011. doi: 10.1161ATVBAHA111.237784.

E. Vo, D. Hanjaya-Putra, Y. Zha, S. Kusuma, and S. Gerecht. Smooth-Muscle-Like Cells Derived from Human Embryonic Stem Cells Support and Augment Cord-Like Structures In Vitro. *Stem Cell Reviews and Reports*, 2010. doi: 10.1007/s12015-010-9144-3.



µ–Slide Angiogenesis Family

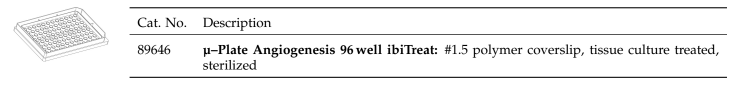
The μ -Slide Angiogenesis family is available with different surfaces and formats. See table below for choosing your μ -Slide and μ -Plate Angiogenesis, respectively.

µ–Slide Angiogenesis

	Cat. No.	Description
	81506	μ–Slide Angiogenesis ibiTreat: #1.5 polymer coverslip, tissue culture treated, sterilized
6999	81501	µ–Slide Angiogenesis uncoated: #1.5 polymer coverslip, hydrophobic, sterilized
*	81531	µ–Slide Angiogenesis Microdissection: PEN-membrane*, sterilized

* The PEN foil does not fit to standard cover slip thickness.

μ–Plate Angiogenesis 96 well



For research use only!

Further technical specifications can be found at www.ibidi.com. For questions and suggestions please contact us by e-mail *info@ibidi.de* or by telephone +49 (0)89/520 4617 0. All products are developed and produced in Germany. © ibidi GmbH, Am Klopferspitz 19, 82152 Martinsried, Germany.