

Instructions µ-Slide 8 well



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 8 well is an array of 8 square fields where cells can be cultivated and subsequently investigated with microscopical methods. It is intended for optimization of experimental parameters like antibody dilution, seeding density or most effective drug concentration.

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 plastic bottom, 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	

Geometry

The μ -Slide 8 well provides standard slide format according to ISO 8037/1.

Geometry of μ–Slide 8 well		
Number of wells	8	
Dimensions of wells ($w \times l \times h$) in mm	$9.4\times10.7\times6.8$	
Growth area per well	1.0 cm^2	
Recommended filling volume per well	300 µl	
Total height with lid	8 mm	
Bottom matches coverslip	No. 1.5	

μ-Slide Surfaces

Depending on the type of cells and the special application you are using, you will need μ -Slides with different surfaces. If you do not require any special adhesion molecules

for your application, the best choice will be ibiTreat, a tissue culture treated surface.

We provide precoated μ –Slides with adhesion substrates like Collagen IV, Fibronectin, Poly–L–Lysin, and Poly–D–Lysin. Such adhesion substrates have been shown to stimulate the adhesion and growth of various cell lines in μ –Slides. Only high–quality substrates are used 1 .

The uncoated μ –Slide is manufactured from hydrophobic plastic. For the cultivation of most cell lines, it is indispensable to treat the uncoated μ –Slide with biopolymers, which mediate cell adhesion and growth.

Coating your µ-Slide 8 well

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.

- Prepare your coating solution according to the manufacturer's specifications or reference.
- Apply 300 µl per well and leave at room temperature for at least 30 minutes.
- Aspirate the solution and wash with ultra-pure water. Let dry at room temperature.

Seeding cells

- Trypsinize and count cells as usual. Dilute the cell suspension to the desired concentration. Depending on your cell type, application of a $5-11 \times 10^4$ cells/ml suspension should result in a confluent layer within 2–3 days.
- Apply 300 μl cell suspension into each well of the μ–Slide. Avoid shaking as this will result in inhomogeneous distribution of the cells.

¹Collagen IV, BD Cat.-Nr. 35 6233, Fibronectin, BD Cat.-Nr. 354008, Poly-L-Lysin, Sigma Cat.-Nr. P4832, Poly-D-Lysin, BD Cat.-Nr. 354210



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• Cover reservoirs with the supplied lid. Incubate at 37°C and 5 % CO₂ as usual.

Undemanding cells can be left in their seeding medium for up to three days and grow to confluence there. However, best results might be achieved when the medium is changed every 1–2 days. Carefully aspirate the old medium and replace it by 300 μ l/well fresh medium.

Slide on an inverted microscope. You can use any fixative of your choice. The $\mu\text{-Slide}$ 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.

Tip:

As you may know from 96 well plates, the bent meniscus at the air–liquid interphase in small open wells destroys the phase contrast effect of your microscope image. To avoid this problem, we recommend using our channel Slides such as the μ –Slides I and VI or the open μ –Slide 18 well flat.

Preparation for Cell Microscopy

To analyze your cells, no special preparations are necessary. Cells can be observed live, or fixed directly in the µ–

Immersion Oil

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

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

μ-Slide 8 well family

The μ -Slide 8 well family is available with different surfaces. See table below for choosing your μ -Slide 8 well.



Ordering Number	Treatment or Coating	Characteristics
80826	ibiTreat, sterile	hydrophilic, tissue culture treated
80822	Collagen IV, sterile	protein coating
80823	Fibronectin, sterile*	protein coating
80824	Poly-L-Lysine, sterile	biopolymer coating
80825	Poly-D-Lysine, sterile*	biopolymer coating
80821	uncoated, sterile	hydrophobic

^{*} available on request only



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Selected References

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