



Perfusion Live Microscopy Using Zeiss LSM 780 and Ibidi Perfusion Sets with SPY650 DNA Dye V.2

Emir Bora Akmeriç¹

¹MDC



protocol.

AG Gerhardt

Emir Bora Akmeriç

Step by step protocol for setting up live microscopy experiments with Ibidi perfusion sets

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https://protocols.io/view/perfusion-live-microscopy-using-zeiss-lsm-780-and-b2f3qbqn

Emir Bora Akmeriç

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Cell Seeding 1h

- 1 Check whether HUVECs in T25/T75 are confluent
- 2 Gelatinize 2 or 3 Ibidi 0.4 luer u-slides with 0.2% gelatin in water

5m

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3	Bring trypsin, PBS, media and FBS to to 37C inside cell culture incubator	25m
4	Trypsinize dish and count cells. A minimum of 500k cells are needed for 2 slides	15m
5	Seed slides with HUVECs at a density of 2.2 million/mL, with 100 uL volume	10m
6	Add 120 uL of EGM2 media with antibiotics 20 to 30 minutes after seeding.	5m
7	Put one pair of male luer couplers, 2 sets of syringes and 1 set of male extenders(for microscopy, there are some on my bench. Basically 2 tubes inside a pipette tip box) incubator for overnight degassing	
Live mi	croscopy prep 1h 10m	
8	In a 2 mL eppi, dilute 1:1000 spy650 DNA in CO2 independent medium(transparent,	10m
	a falcon in 4C. If not, there is also a stock you can add Gentamicin to)	silould be
9		and the gates
9	a falcon in 4C. If not, there is also a stock you can add Gentamicin to) Prepare one set of ibidi flow unit by adding 14 mL of CO2 independent medium. Externale coupling with the tubes in the extra coupling box. You should have slightly elon tubes in a tube>male adaptor>female joiner>extra tube>male coupler>luer female or	and the gates oupler
-	a falcon in 4C. If not, there is also a stock you can add Gentamicin to) Prepare one set of ibidi flow unit by adding 14 mL of CO2 independent medium. Extermale coupling with the tubes in the extra coupling box. You should have slightly elon tubes in a tube>male adaptor>female joiner>extra tube>male coupler>luer female corder Aspirate medium from the slides and add 250 uL of previous mix into each slide. Income	end the gates oupler ubate at

best way to do this is putting two pipette tip boxes on the bottom right part of the chamber and

	Continue with connecting the air tube and electric cable, there is a stage exit for such cables. Turn on microscope and incubator and open ZEN. Calibrate, pinch test and start running at 37C but without CO2	
13	Bring one slide and a plastic clamp to the microscopy room. While clamped carefully connect the slide to the unit. Dry couplings as usual with kimwipe	
14	Wipe both sides of the slide with isopropanol as well as the objective	
15	On ZEN, withdraw the imaging setup from an image in Anna/210709_Wt_flow 5m	
16	Turn on Definite Focus. Click Find Surface/Focus	
17	Check focus on Live and make necessary manual focusing adjustments and click store focus	
18	Check whether definite focus is on for focus strategy, enable definite focus for every tile scan	
19	Check that Tile scan is set to 3x3 and that a 250+ frame timed capture will be done, change autosave to stream	
20	Start experiment. If possible, check whether everything is in focus after the first 10 minutes	
ost microscopy(next day) 1d		
21	After 24 hours, dismantle the flow unit. You can bring the unit with the it to the 4C room and 1d	
	can take it apart and clean afterwards.	

then setting up the flow unit on top of these boxes, making sure that the setup is stable.