

Cell Sorting

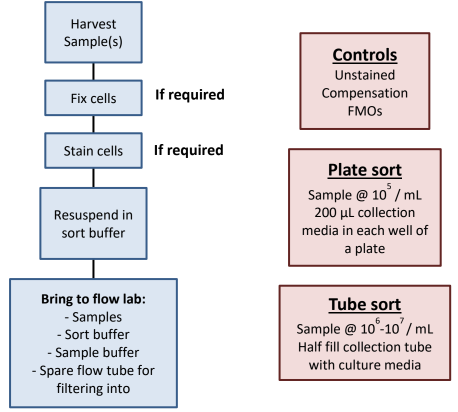
How it works?

You bring us sort sample(s) and appropriate controls and we sort the desired population within your sample.

Astrios: 5-lasers 355,405,488,561,630 nm; sort into tubes or plates; sort microparticles.

Remember, book your sort and submit a sort-form at least 2 days before sort-day https://dfcf.acls.analytical.unsw.edu.au/

Flow chart



Sort buffers

Basic	Sticky cell	Low viability	
1x PBS	1x Hanks BSS	1x Hanks BSS	
2 mM EDTA	5 mM EDTA	2 mM EDTA	
up to 2% serum	25 mM HEPES, pH 7	25 mM HEPES, pH 7	
	up to 2% serum	10U DNAse	
		up to 2% serum	

Next page, gotchas

	has <mark>/!\</mark>							
	There are a number of critical steps for successful cell sorts							
•	Cells MUST be filtered							
	 Even if your cells don't need it. 							
	 Consequence: Machine clogs, you take a big hit to your yield. Lose sort time for t rest of your samples. 							
	Bring correct controls							
	• Action required: Common controls: unstained, compensation, FMO							
	• Consequence : We will have to guess which population to sort if we can even							
	proceed with the sort.							
	Over confluent cells							
	 Action required: Keep your cells <70% confluent (adhesive), 8x10⁵ / mL (suspensic 							
	• Consequence: single-cell suspension will form aggregates faster, causing machine							
	blockages even if filtered. Cells will be starving and not behave as expected (unless							
	you always starve your cells?).							
Low viability samples								
	 Action required: Add DNAse to sample 							
	 Consequence: DNA released by dead cells causes clumping 							
	Low yield							
	• Main Causes:							
	 Population of interest was not as represented as expected 							
	 Total cell count was lower than expected 							
	 Clumped cells – which those events were aborted 							
	 Machine capability (up to 40% loss (aria), 25% loss (astrios). This includes 							
	cells exploding on impact in the tube							
	• Solutions:							
	Low % of desired cells; optimise transfection – don't just follow someone							
	else's protocol, establish the best conditions for your work.							
	 <u>Cell loss</u>; try to identify cause of loss (during washes, column), or increase 							
	input amount if possible							
	 <u>Clumped cells</u>; avoid over-confluence, we can filter the sample just before 							
	the sort to reduce cells re-adhering, use DNAse to break up low-viability							
	 samples <u>Exploding cells</u>; Astrios delivers higher yields for plate sorting 							
	- <u>Exploding cens</u> , Astrios delivers nigher yields for plate soluting							

Sort time; as predictable as pachinko.

	0.1	1.0	10.0	30.0	% cells of interest in	n sample
The sweet spot for a	2,000			event rate / sec		
reasonable sort time is	1666.7	166.7	16.7	5.6	minutes for 200k	
to sort a population with greater than 1%	5,000					
	666.7	66.7	6.7	2.2		
sample representation.	9,000					
	370.4	37.0	3.7	1.2		
- Well, sort at 9000/s all th - This is only possible if you		pristine.	We usual	lly run at 5	5000 events / s.	