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## What is wrong with my flow cytometry data?

Hints, tricks and pitfalls

**Webinar 5**  
**06.06.2023**

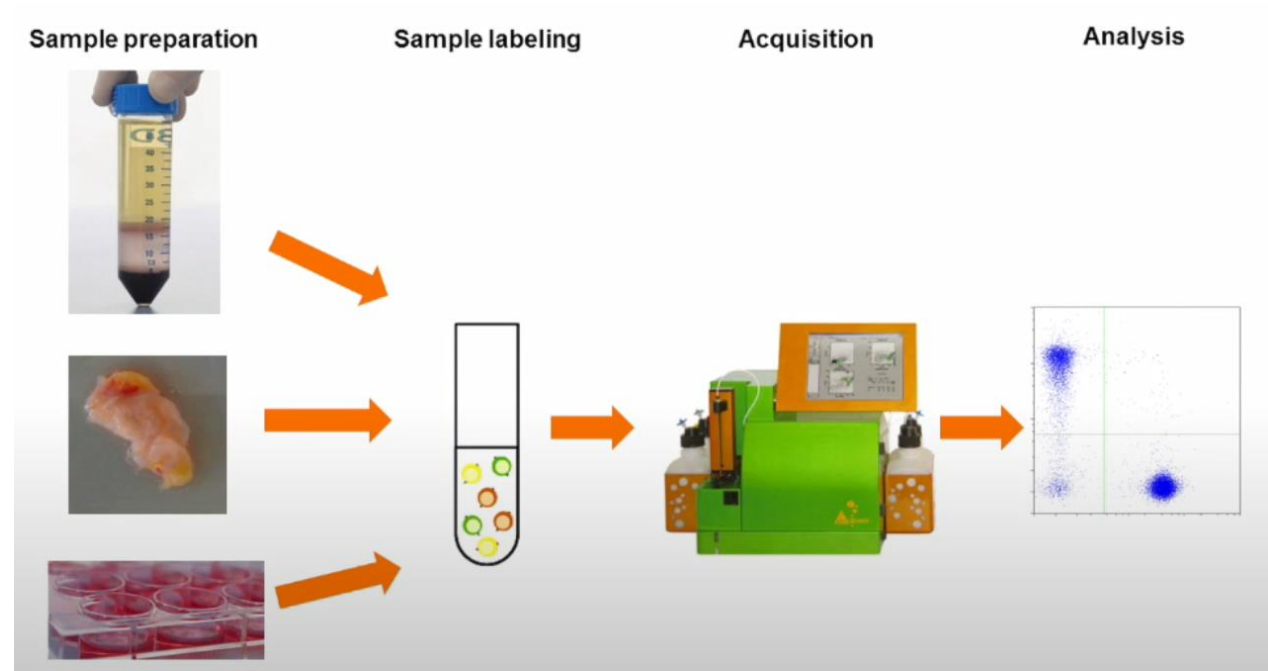
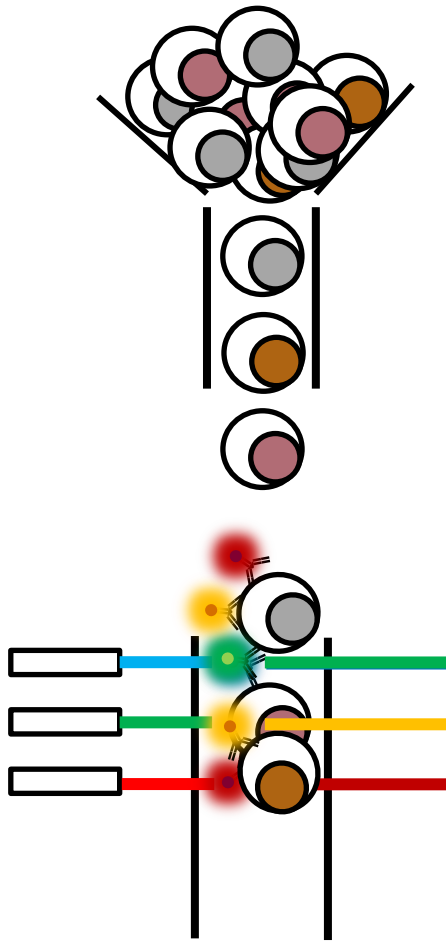
# The alpha and omega of preparing cells for flow cytometry



# What we need to talk about

1. Single cells
2. Preparation: blood, cells, tissues
3. Mechanical vs. enzymatic disintegration of tissues or cell layers
4. Doublets and dead cell exclusion
5. Fixation
6. Surface staining
7. Intracellular staining

# All about **single cells**...



<https://www.youtube.com/watch?v=mpOIRpIZhPg>

Viability and yield max!

# The A and O of **sample preparation**

“ better sample preparation leads to better results  
and easy data analysis



“ standard procedure:

- sampling: blood, tissues, freeze/thaw, cell harvest

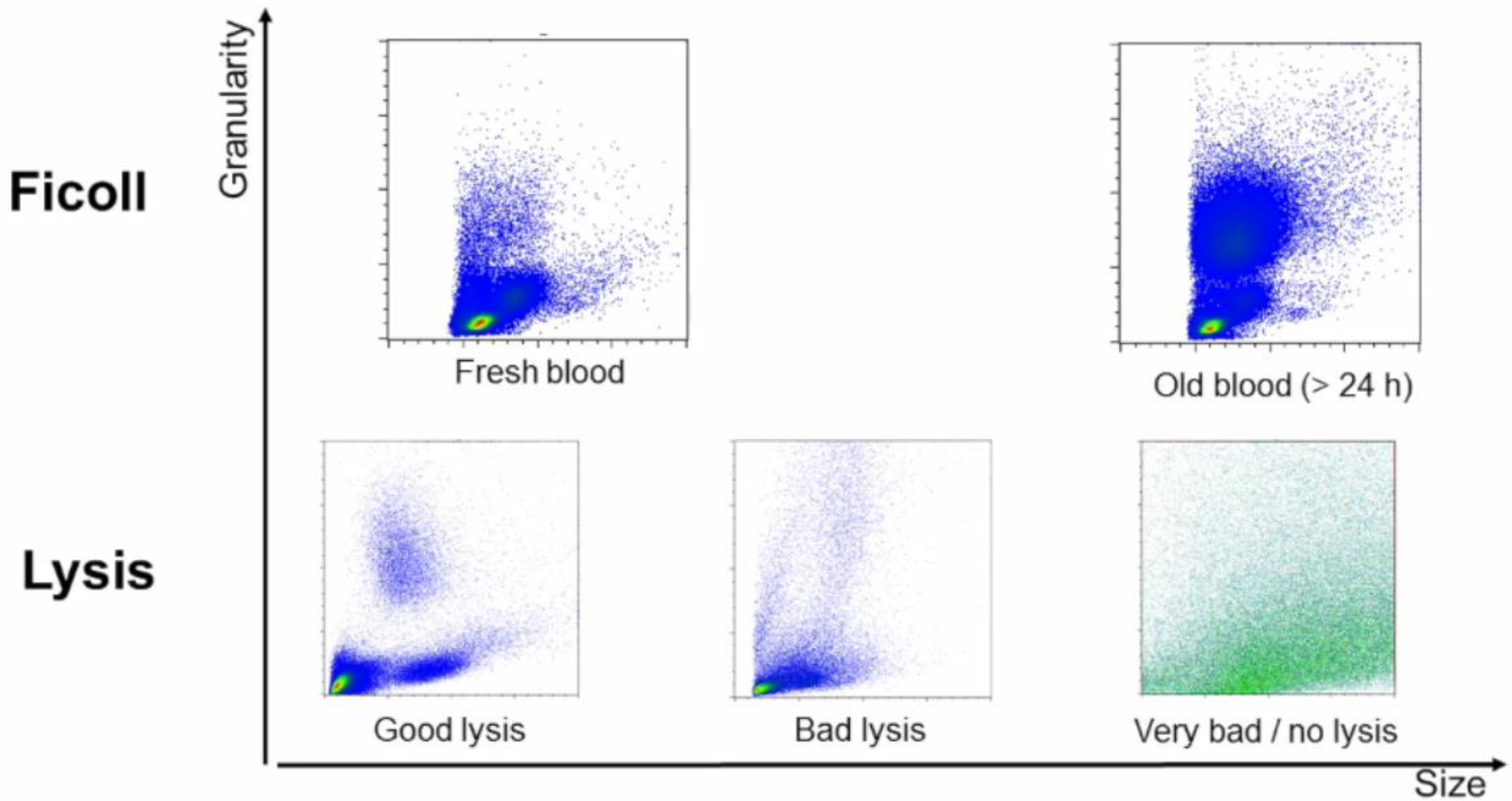
à single cell suspension

- wash with cell stain buffer (1X PBS + 1-2% BSA + 2-5 mM EDTA + 2mM  $\text{NaN}_3$ )

- stain and measure

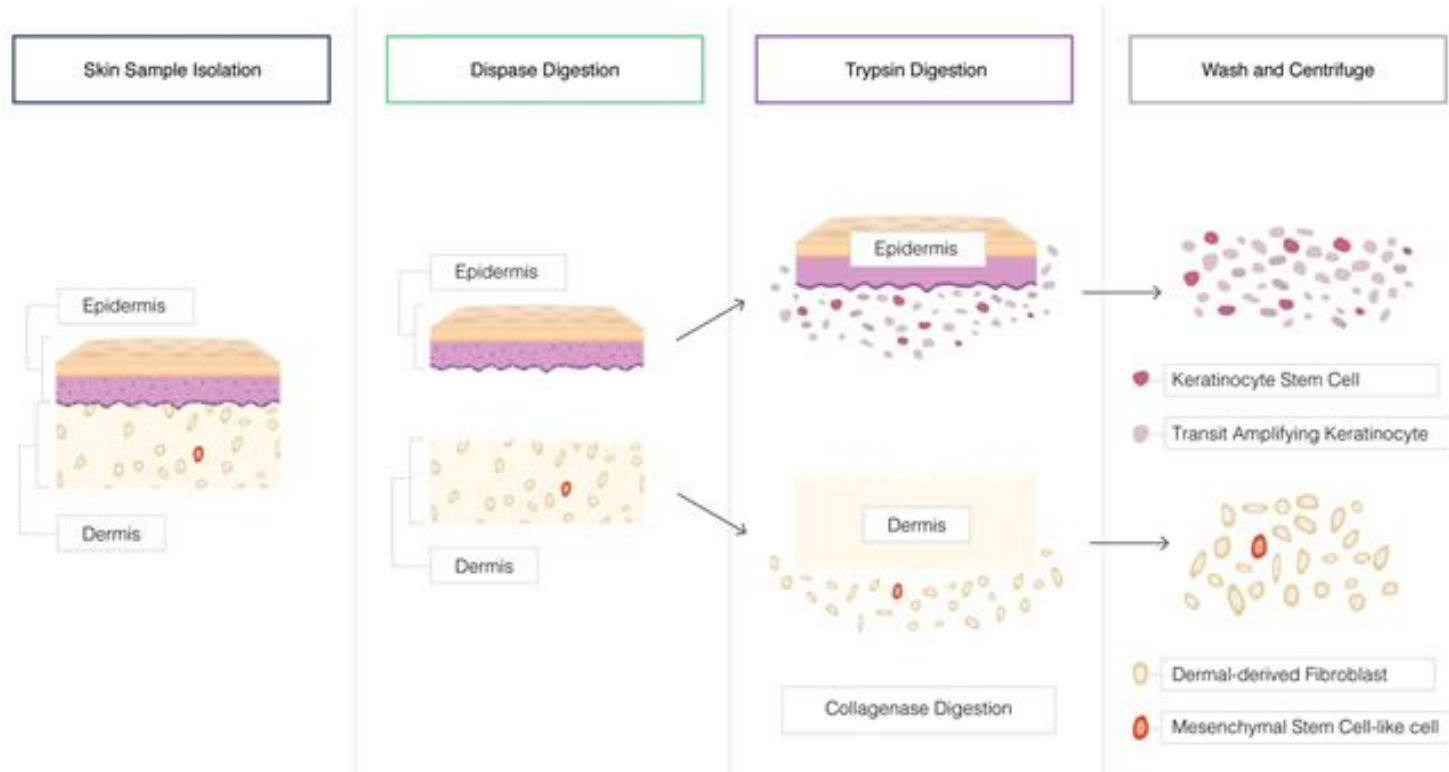
“ changing BSA and/or EDTA conc. might reduce doublets

# Sample prep: **Blood**



<https://www.youtube.com/watch?v=mpOIRpIzhPg>

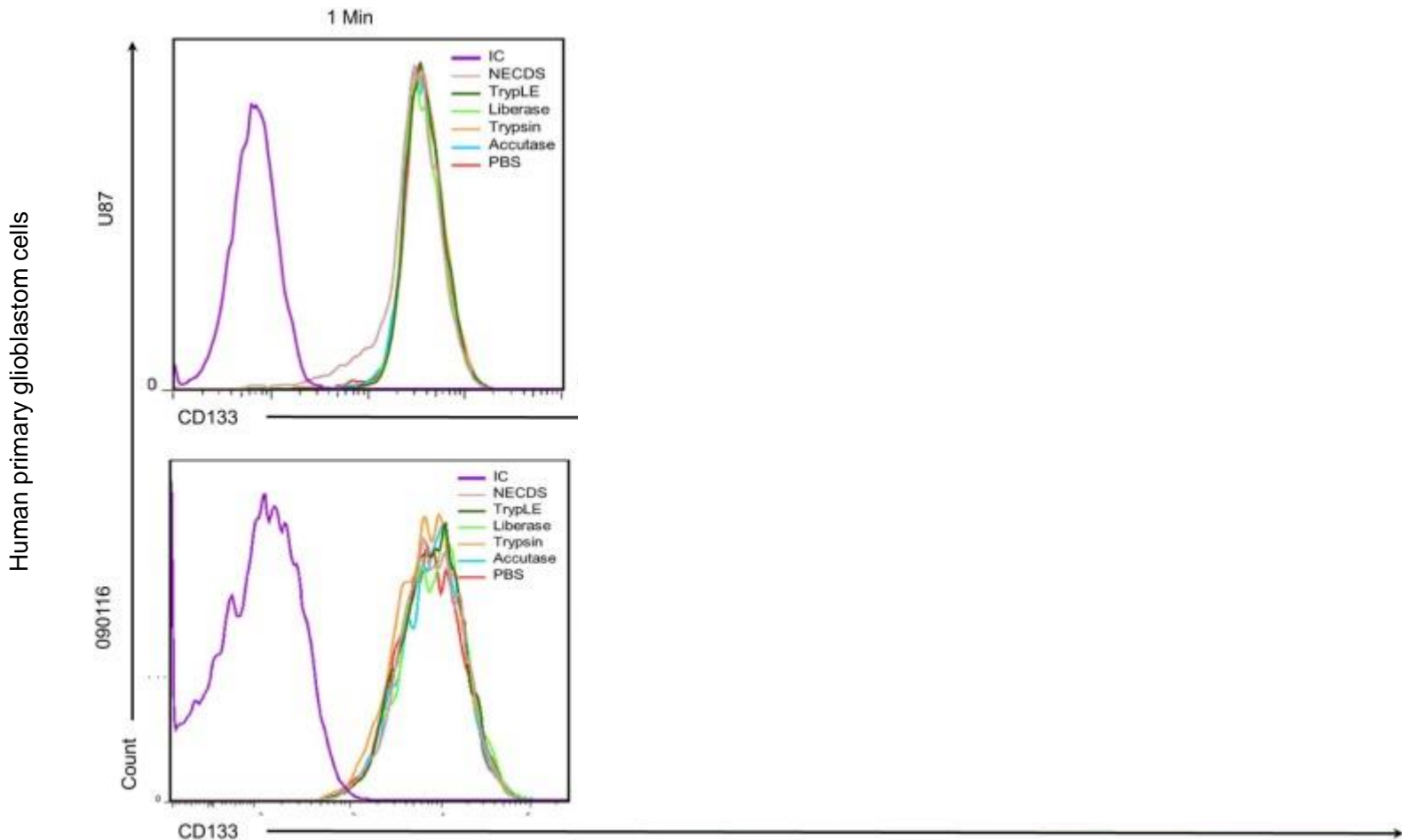
# Sample prep: **Tissues**



<https://www.flowmetric.com/cytometry-blog/sample-preparation-for-flow-cytometry>

- “ complex enzymatic digestion needed
- “ combination of enzymatic and mechanical protocols, include DNase
- “ goal: single cells with > 50% vitality and intact epitopes

# Sample prep: Enzymatic digestion



NECDS (non-enzymatic cell dissociation solution): detachment of primary cells, EDTA based

TrypLE (Gibco): animal origin-free, recombinant enzymes for dissociating adherent mammalian cells à substitutes trypsin, superior performances

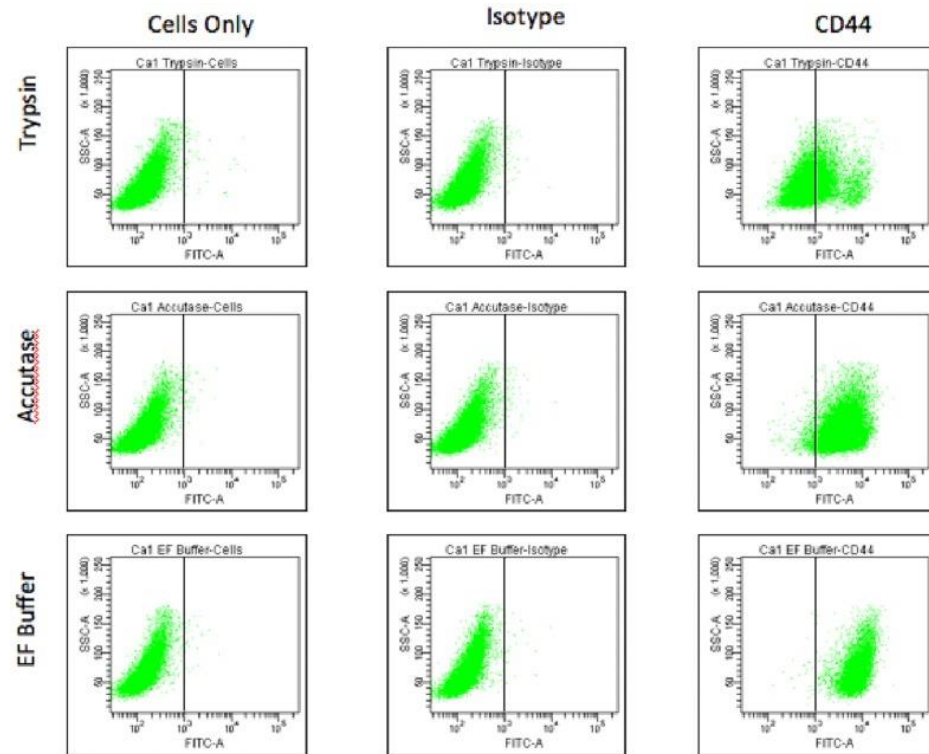
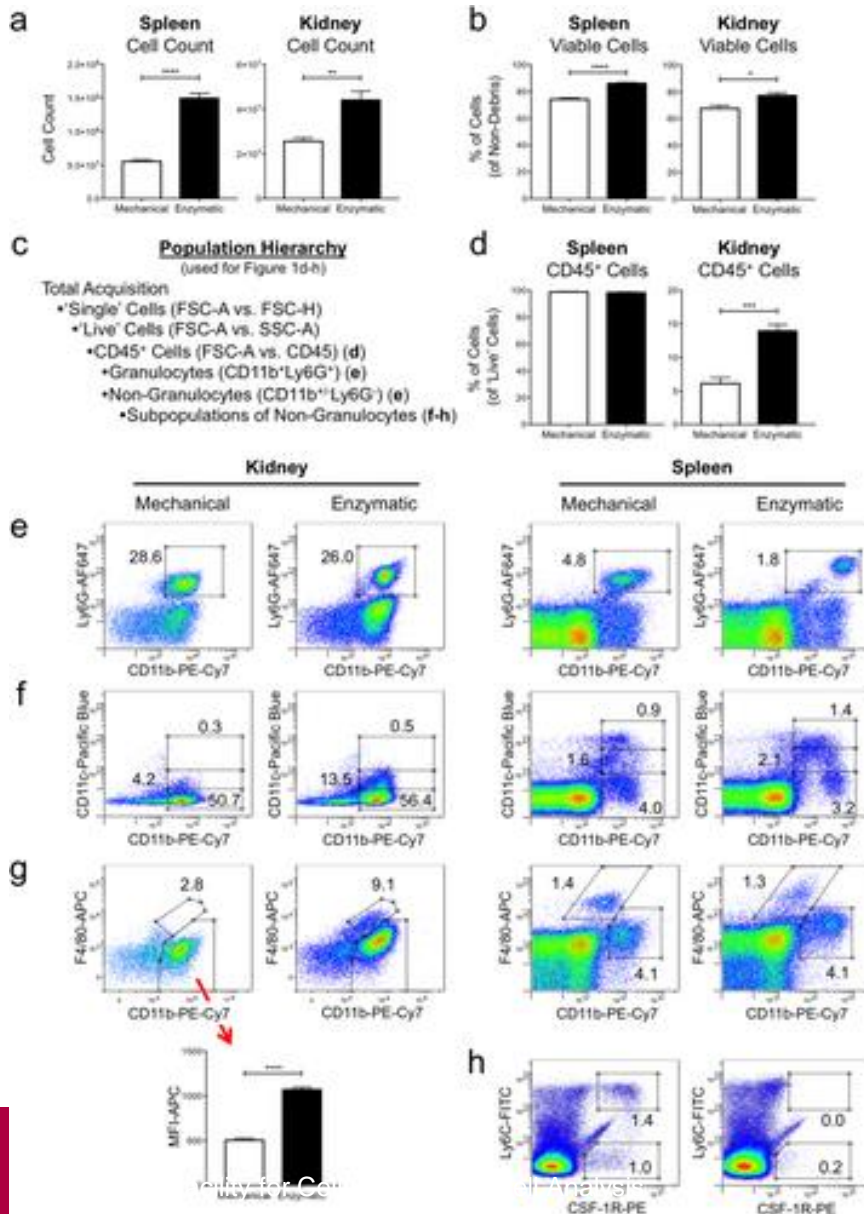
Liberase (Merck): Kollagenase I / II + Thermolysin (Protease)

Accutase: animal origin-free, proteolytisch und kollagenolytisch

CD133 . prominin-1 HSC,CSC marker



# Sample prep: Enzyme vs. scissors

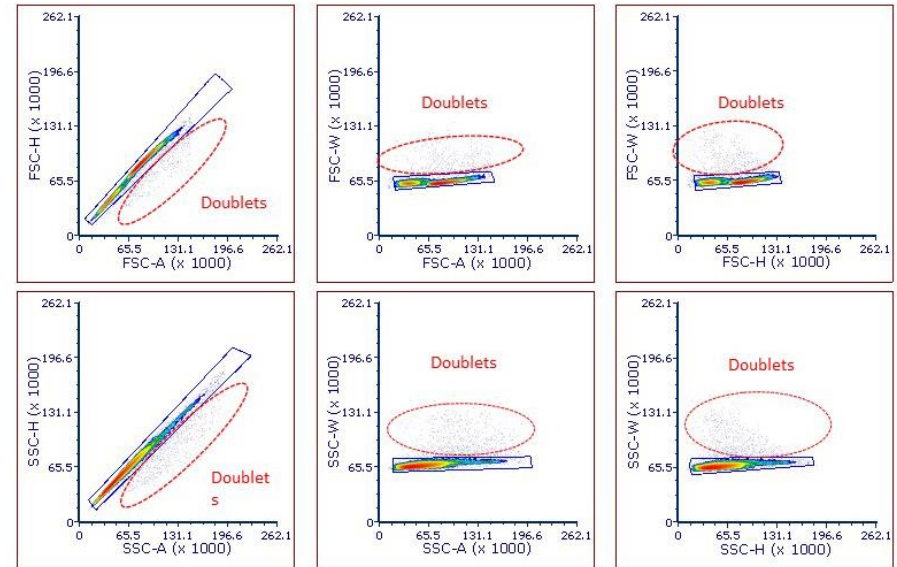


EF buffer = enzyme-free cell dissociation buffer (Invitrogen #13151-014)

# Sample prep: dead/ doublet cell ex

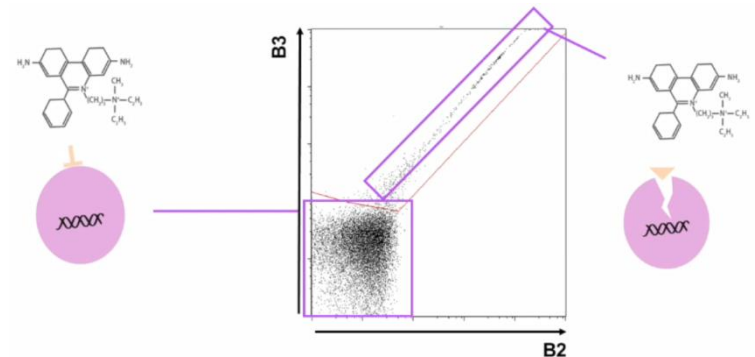
## Doublet cells

- “ wrong buffer composition
  - “ clumps do clog the cytometer
  - “ doublets cause false-positive signals
- à resuspend, filter (30-70 nm), vortex



## Dead cells

- “ density gradient:
  - “ high cell loss  $Q_A \in \tilde{A} D$
  - “ toxic
  - “ increase of autofluorescence
- à DNase treatment



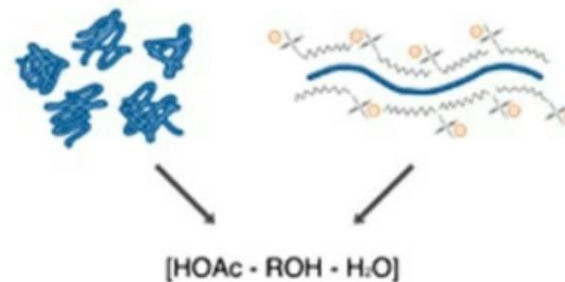
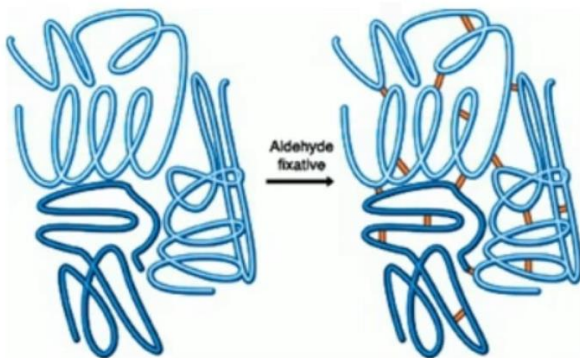
# Sample prep: Fixation

- “ fixation BEFORE staining - epitope alteration
- “ fixation AFTER staining - fluorophore instability
  - GFP, cyanine- and protein based fluorophors: signal loss after fixation (APC-Cy7 < APC-H7, PerCP, FITC, etc)
  - but some phospho-epitopes needs methanol demasking

## • Fixation

### • Formaldehyde

- Create bonds between lysine residues to cross link proteins
- Used between 0.5% and 4%
- Stable for 1-2 weeks but should be kept in PBS after initial fixation



### • Alcohols

- Usually 70% ethanol or methanol
- Precipitating/denaturing fixatives
- Dissolve lipids so permeabilize at the same time
- Coagulates proteins so may hide epitope
- Stable for months at 4°C or -20°C

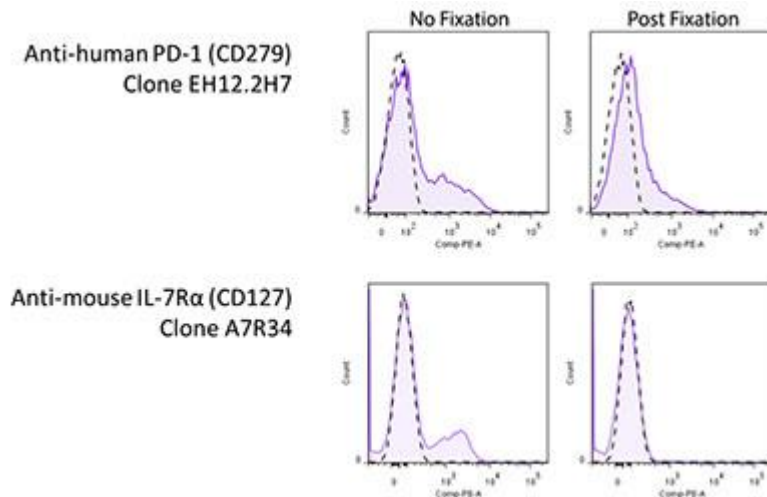
<https://www.youtube.com/watch?v=oeqiCa3nEwc>

# Sample prep: **Surface staining**

- “ coagulant (e.g. EDTA, Heparin) interfere with some epitopes
- “ check enzyme activity and whether targets are snibbled off%
- “ Ficoll and RBC Lysis buffer also interfere with surface epitopes
- “ blocking of Fc receptors when working with PBMCs or any other cell types expressing FcR
  - à exception: Fc-staining on monocytes, Ig-staining on B cells
- “ in case of fixation: check tandem fixability
- “ stimulation reduces or increases expression? à PMA/Iono causes internalization
- “ no azide in functional assays à ROS detection, Ca<sup>2+</sup>-flux

# Sample prep: Intracellular staining

- “ also block intracellularly
- “ dilute antibodies in saponin-based buffer (reversible)
- “ use bright fluorophors since expression is often weak
- “ use (FMO) controls



<https://www.biolegend.com/en-us/blog/fix-now-fix-later-considerations-for-the-use-of-paraformaldehyde-fixation-in-flow-cytometry>

- “ wash the cells adequately after each antibody incubation step (unbound antibodies are trapped in the cells in the case of intracellular staining)
- “ perform all protocol steps at 4°C
- “ permeabilize on ice

# Sample prep: **Take home message**

- “ the better the preparation and protocol of your samples the better your results/analysis is
- “ coagulants, Ficoll, lysis buffer, enzymes, fixation buffer interfere with epitope access and reduce signal of flurophors
- “ in multi-colour panels optimization towards high signal, low doublet rate, high viability is the goal!

Thank you for your attention.

See you next month: **4th July**

Next topic  
**When to use which buffer system**

## Enzymes for tissue dissociation

Enzyme	Type	Source	Target	
Trypsin	serine protease	bovine/porcine pancreas	predominantly at c-terminal side of AA Lys or Arg	relatively harsh
Papain	cysteine protease	<i>Carica papaya</i>	cleaves basic AAs	relatively harsh, but less digestive than Trypsin
Collagenase	protease, includes also phospholipases or neuraminidases	<i>Clostridium histolyticum</i>	peptide bonds in collagen; Different types available, depending on supplier Roche: A, B, D, H, P Sigma-Aldrich: I-IV, Worthington: Type 1-4	different collagenase types with different levels of unspecific activities
Liberase Blendzymes	blend of purified collagenases with neutral proteases	<i>Clostridium histolyticum</i> and <i>Bacillus polymyxa</i> or <i>B. thermoproteolyticus</i>	Supplied by Roche, more defined activity and composition	developed for pancreatic islet transplantation research
Dispase	protease	<i>Bacillus polymyxa</i>	fibronectin, collagen IV (collagen I)	separation of dermis and epidermis
DNase	nuclease DNase I: neutral DNase II; functions optimally at acidic pH	bovine pancreas	DNA	avoid cell clumping

<https://www.youtube.com/watch?v=mpOIRplZhPg>