

## Zoom In and Out: A Comprehensive Immunologic Evaluation of Human, Murine and Rat Samples

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### Introduction

Fluorochrome-conjugated antibodies applied in conventional FACS are widely used but have limited utility for high-parameter studies. Spectral cytometry overcomes those limitations since the emission spectrum of every fluorescence molecule is detected across a defined wavelength range. Employing spectrum cytometry for immunophenotyping at NUVISAN we can zoom in and out of the immune system of:

- Human
- Mouse
- Rat

with less sample material needed to extract complex information vital for immunological studies. Immune responses to cancer are highly influenced by the tumor microenvironment, where the delicate balance between suppressor and effector cells steers the prognosis after therapy. Proper establishment of multiparameter panels require accurate assessment of isolation and preservation protocols as practical considerations on epitope stability of immune cells vital for downstream assays and immunophenotyping.

We use spectral cytometry to map immune cell subsets and correlate them with tumor progression in human, mice and rats. Using unsupervised dimensionality reduction tools (e.g viSNE) we identify major immune subsets as well as analyze their expression of stimulatory and inhibitory molecules in tissues and periphery.

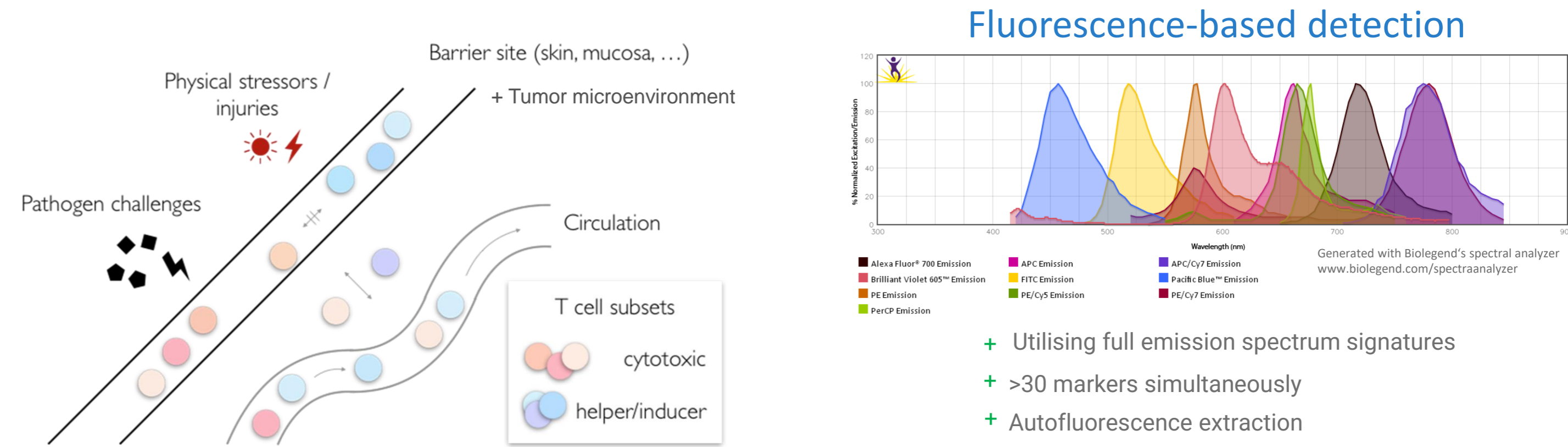


Fig. 1: Cytotoxic and helper-type circulating and tissue resident T cells at barrier sites as well as tumor microenvironment.

### Employing Spectral Cytometry for Immunophenotyping

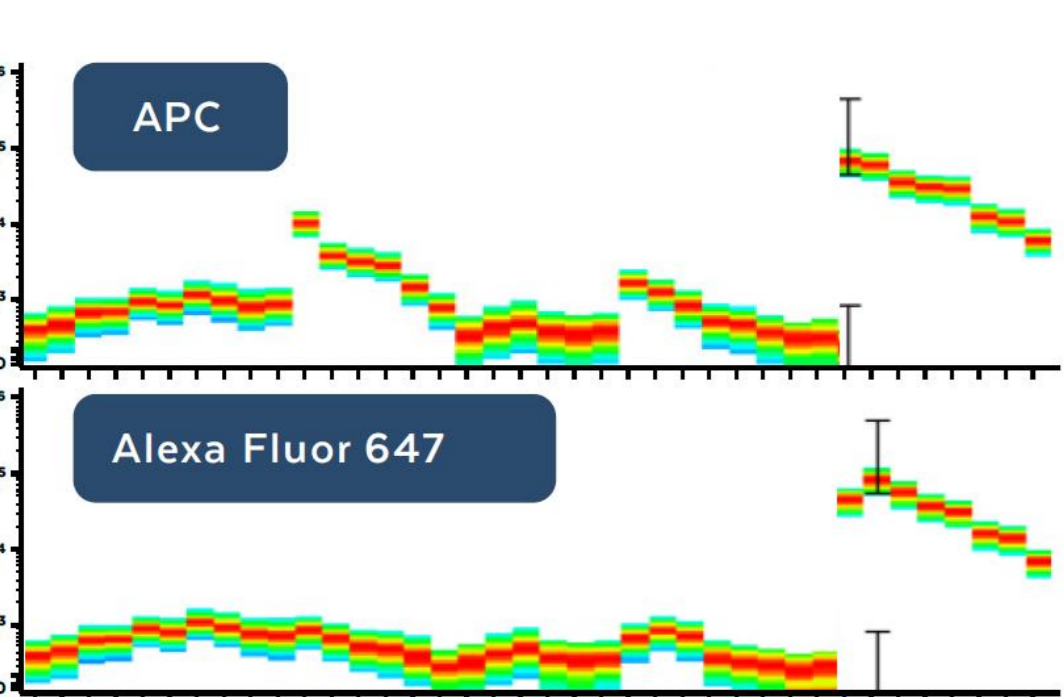


Fig. 2: Full spectrum plots from a 3-laser CYTEK Aurora show distinct signatures for APC and Alexa Fluor 647.

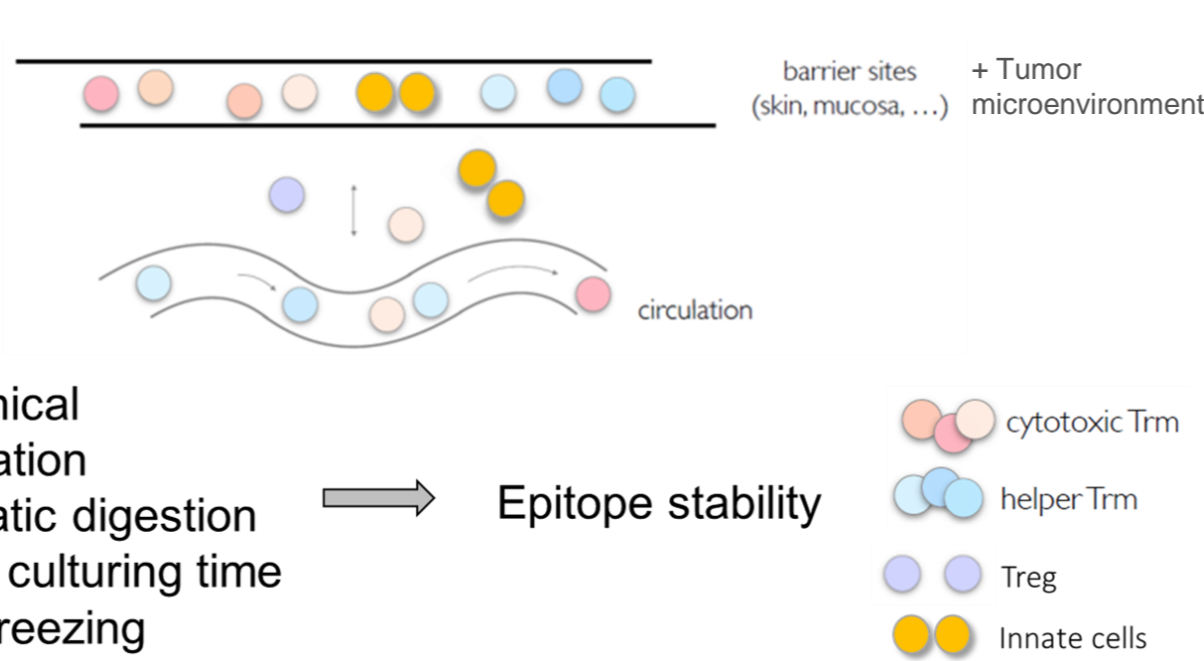


Fig. 3: Establishment of multiparameter panels require proper assessment of isolation and preservation protocols as practical considerations on epitope stability of immune cells vital for downstream assays and immunophenotyping.

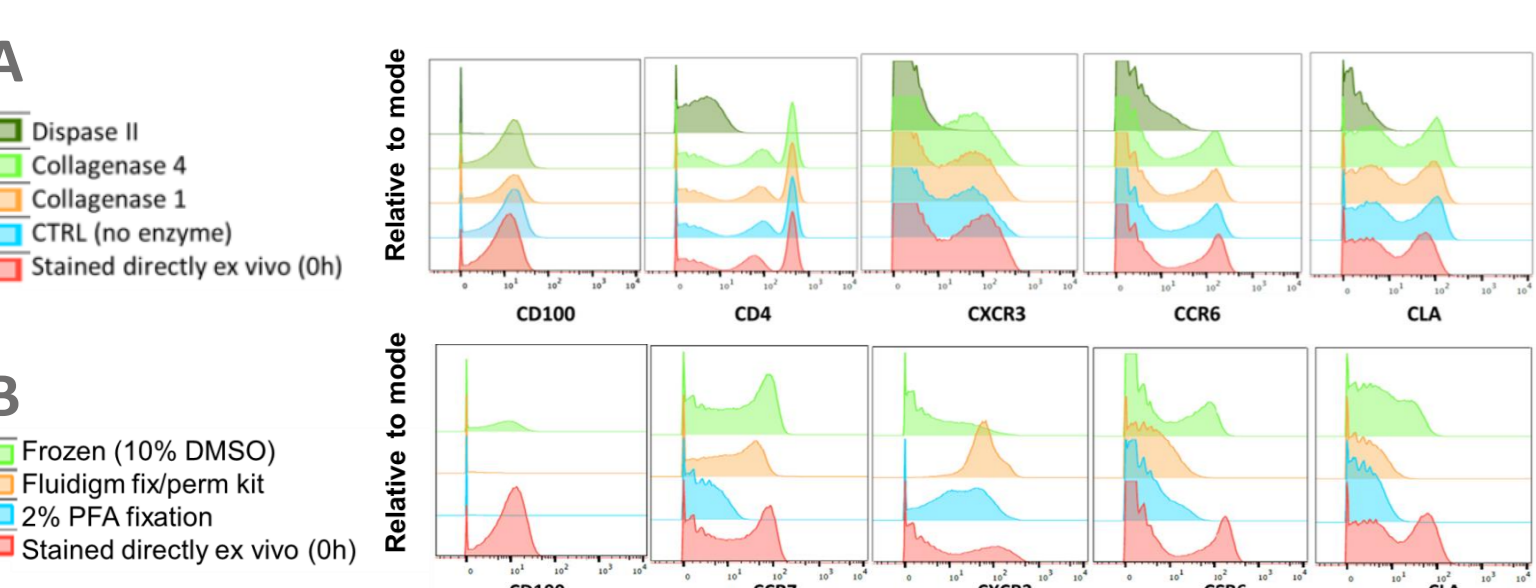


Fig. 4: A) Representative examples of effect of collagenase 1, 4 and Dispase II after 6 hours of digestion, 37 °C onto epitope recovery. B) Representative examples of effect of fixation and freezing onto epitope recovery. Human PBMCs, gated on live single CD45+ cells.

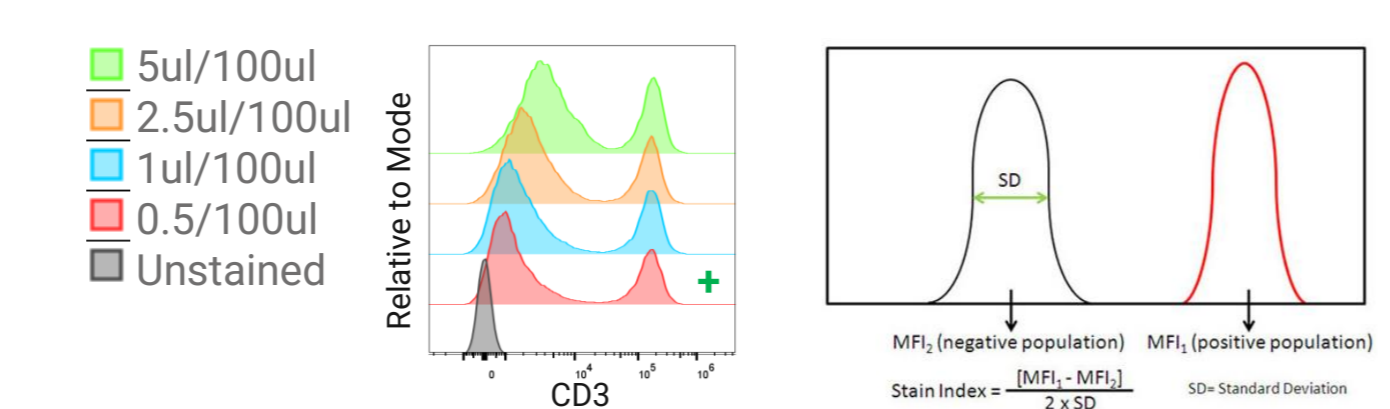


Fig. 5: Representative example of CD3 titration for the identification of the best stain index value.

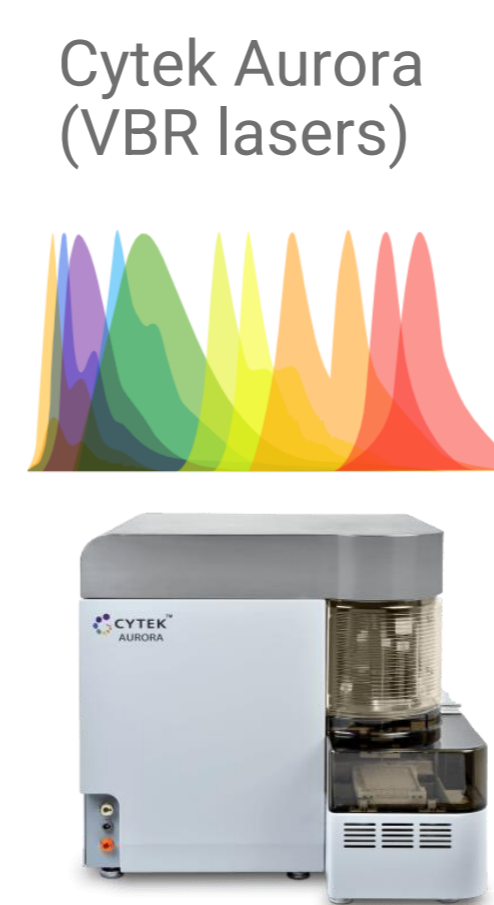
Off the shelf NUVISAN panels available

**Human Immunophenotyping Panel** (≥30 parameters) of B, T, NK, DC, ILC and monocyte subsets in PBMCs.

**Murine Immunophenotyping Panel** (≥ 28 parameters) of B, T, NK, DC, and monocyte subsets in Thymus, blood, spleen, BM, tumors.

**Rat Immunophenotyping Panel** (≥ 13 parameters) of B, T, NK, DC, and monocyte subsets in Thymus, blood, spleen, BM, tumors.

| human     | mouse      | rat       |
|-----------|------------|-----------|
| CD11c     | CD4        | CD4       |
| CD45RA    | CD19       | CD45RA    |
| CD3       | Iy6G       | CD8       |
| CD25      | CD69       | CD3       |
| IgD       | CD45       | CD62L     |
| CD95      | CD44       | Gr        |
| CD11b     | CD11b      | Ki67      |
| CD38      | CD62L      | CD161a    |
| CD57      | Nkp46      | CD45RC    |
| CD27      | CD137/41bb | Live/dead |
| CD123     | CD8        | CD28      |
| CD127     | CD45R/B220 | IgM       |
| HLADR     | CTLA4      | CD38      |
| CCR7      | F4/80      | CD45      |
| CD19      | CD3        |           |
| CD16      | Iy6C       |           |
| TCRgd     | TCRgd      |           |
| CD14      | CD11c      |           |
| CD8       | PD1        |           |
| CD1c      | CD25       |           |
| PD1       | SLAMF7     |           |
| CD56      | CD206      |           |
| CD45RA    | Live/dead  |           |
| CD28      | MHCII(A/E) |           |
| SLAMF7    | CD38       |           |
| CCR3      |            |           |
| CCR6      |            |           |
| cKIT      |            |           |
| IgM       |            |           |
| Live dead |            |           |



Advantage of Spectral flow cytometry:  
**Less sample material needed to extract more complex information!**

Establishments & optimisations of FACS panels are crucial steps in flow cytometry experiments, as they optimize sensitivity, specificity, and reproducibility, leading to improved data quality and reliability. Find out more at <https://www.nuvisan.com/home.html> to see how we are committed to supporting cutting-edge research endeavours. As part of our comprehensive suite of services, we take great pride in offering specialized assistance as well as expertise in immunology and Fluorescence-Activated Cell Sorting (FACS) experiments.



### Human immunophenotyping

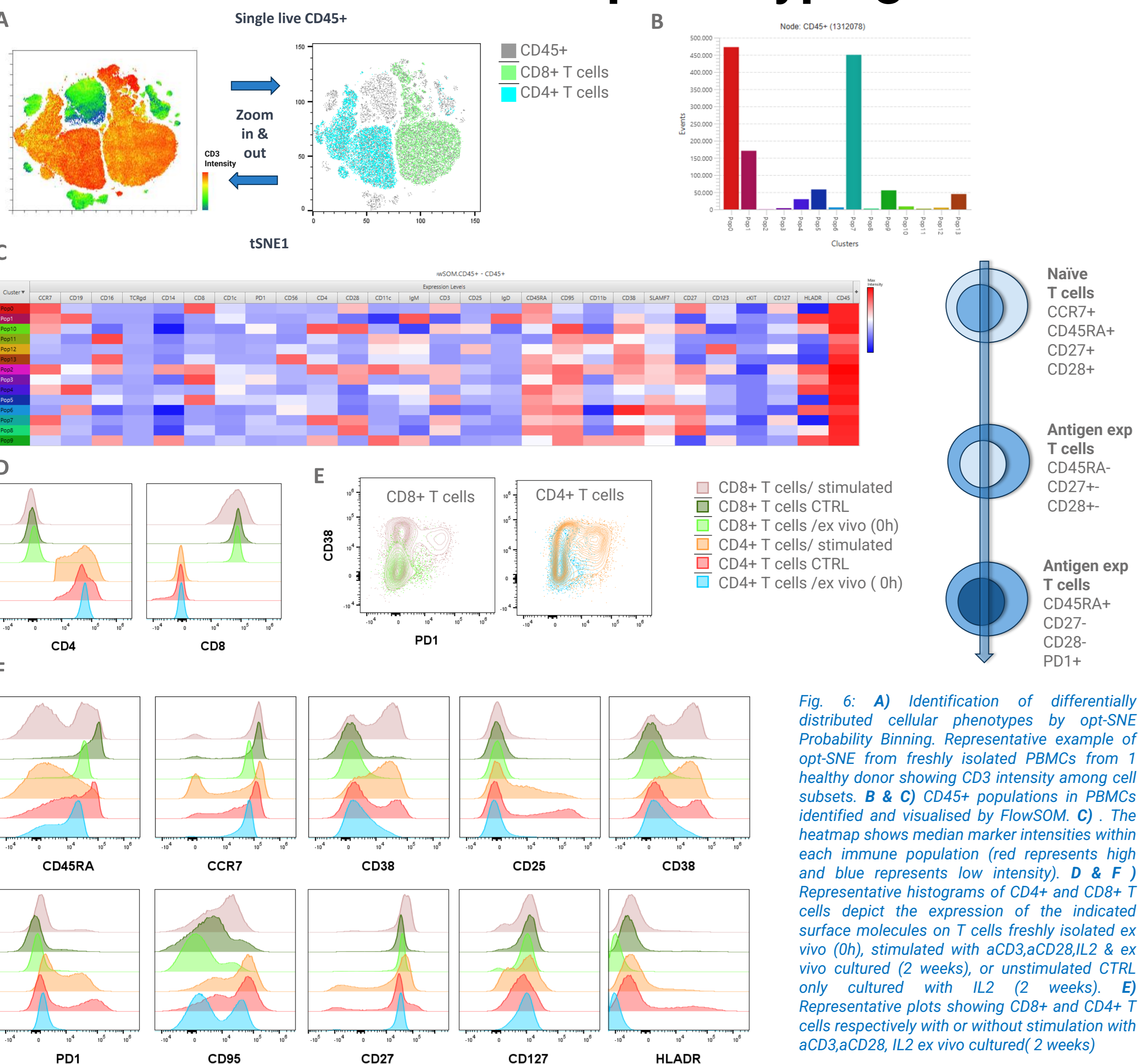


Fig. 6: A) Identification of differentially distributed cellular phenotypes by opt-SNE Probability Binning. Representative example of opt-SNE from freshly isolated PBMCs from 1 healthy donor showing CD3 intensity among cell subsets. B & C) CD45+ populations in PBMCs identified and visualised by FlowSOM. C) The heatmap shows median marker intensities within each immune population (red represents high and blue represents low intensity). D & E) Representative histograms of CD4+ and CD8+ T cells freshly isolated ex vivo (0h), stimulated with aCD3,aCD28,IL2 & ex vivo cultured (2 weeks), or unstimulated CTRL only cultured with IL2 (2 weeks). E) Representative plots showing CD8+ and CD4+ T cells respectively with or without stimulation with aCD3,aCD28, IL2 ex vivo cultured (2 weeks)

### Murine immunophenotyping

Experimental design 4 groups (aPD1 vs isotype control, aCTLA4 vs isotype control)

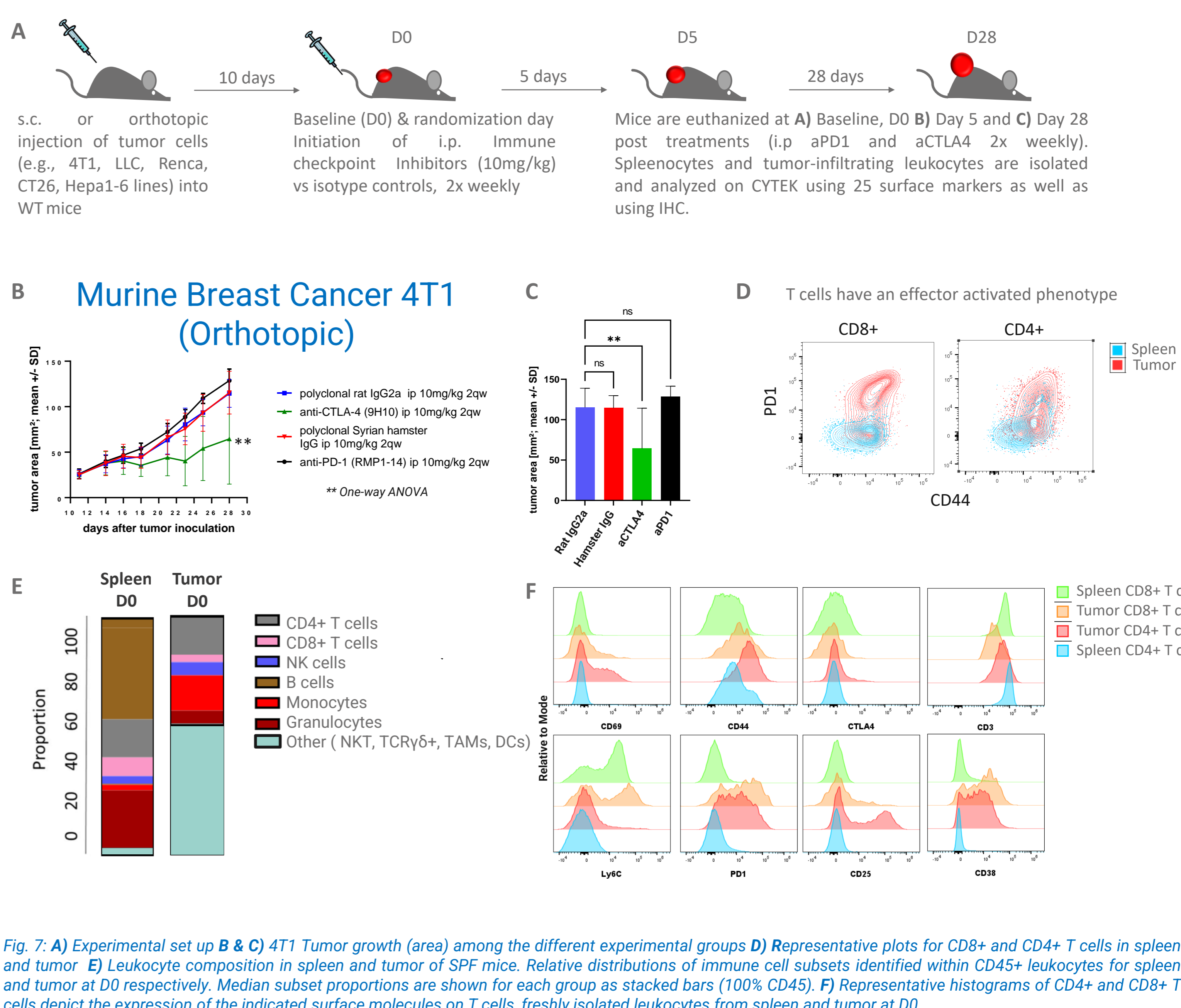


Fig. 7: A) Experimental set up B & C) 4T1 Tumor growth (area) among the different experimental groups D) Representative plots for CD8+ and CD4+ T cells in spleen and tumor E) Leukocyte composition in spleen and tumor of SPF mice. Relative distributions of immune cell subsets identified within CD45+ leukocytes for spleen and tumor at D0 respectively. Median subset proportions are shown for each group as stacked bars (100% CD45). F) Representative histograms of CD4+ and CD8+ T cells depict the expression of the indicated surface molecules on T cells, freshly isolated leukocytes from spleen and tumor at D0.