



Translating the complex immune system into disease insights with **CyTOF** technology

Don't miss the unexpected.



Reasons to choose CyTOF

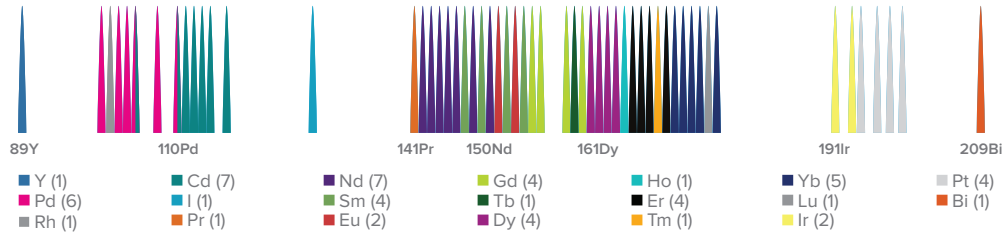
THE CYTOF

Reasons to take your high-dimensional

1

Precision data with discrete signals

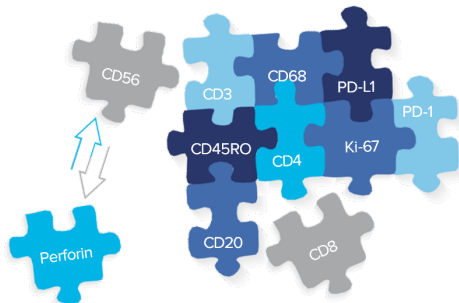
Not impacted by spectral overlapping of fluorochromes and tissue autofluorescence



2

Easy panel design to complete experiments quicker

Large number of available antibodies without overlap simplifies panel design and expansion



Start with ready-to-go panels and easily swap markers in and out.

3

Reproducible and comparable

Stained samples can be frozen, stored and shipped to support longitudinal studies and multi-site workflows.



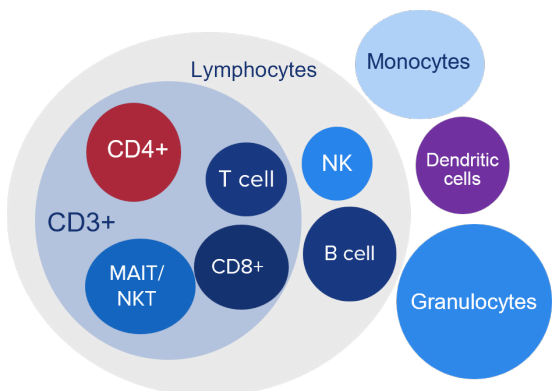
ADVANTAGE

research beyond the limits of fluorescence

4

Capture rare or unexpected cell populations

Unbiased, high-dimensional profiling of 40-plus markers to uncover diverse immune subpopulations



Cytometry by time-of-flight (CyTOF® technology)

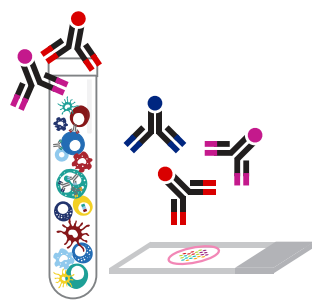
Applies purified heavy metal labels, not normally found in biological systems, instead of fluorophores



5

Minimal sample required, saving on limited clinical research material

Simultaneous staining and detection from a single tube or tissue scan, without multiple staining controls or time-consuming cyclic protocols



Risks of fluorescence for high-parameter studies



Missed cell populations or false positives



More iterations required in panel design



Reduced sensitivity where fluorescence overlap occurs



Higher resource use to compensate for spectral overlap

6

Trusted by researchers

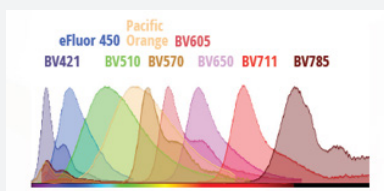
The leading technology for high-parameter immune research



>2,000 publications



>200 clinical trials

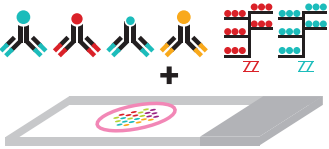


Whether you are analyzing suspension or tissue samples, time-of-flight (TOF) technology combined with Maxpar[®] reagents enables a streamlined end-to-end workflow to complete high-parameter experiments faster than fluorescence-based detection.

Tissue imaging

Get started with our Maxpar IMC[™] Cell Segmentation Kits and ready-to-go high-plex panels.

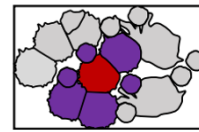
Simultaneous staining



One-step detection

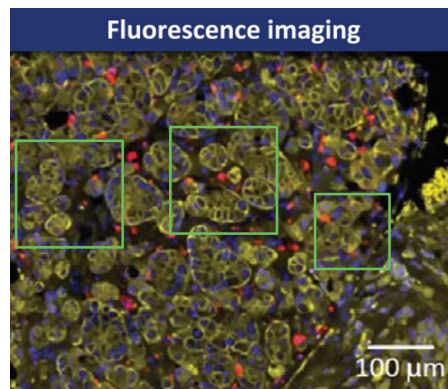
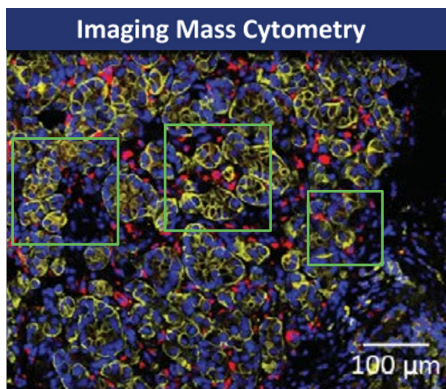


High-plex data in minutes



- Tissue architecture
- Protein modifications
- Signaling pathway activation
- Cell injury states
- Cell proliferation
- Transcriptional signatures

Clear spatial imaging



The Imaging Mass Cytometry[™] image (far left) shows many well-defined red signals from CD68 that are indistinct or missing from the fluorescence image (left).

Without compromise.

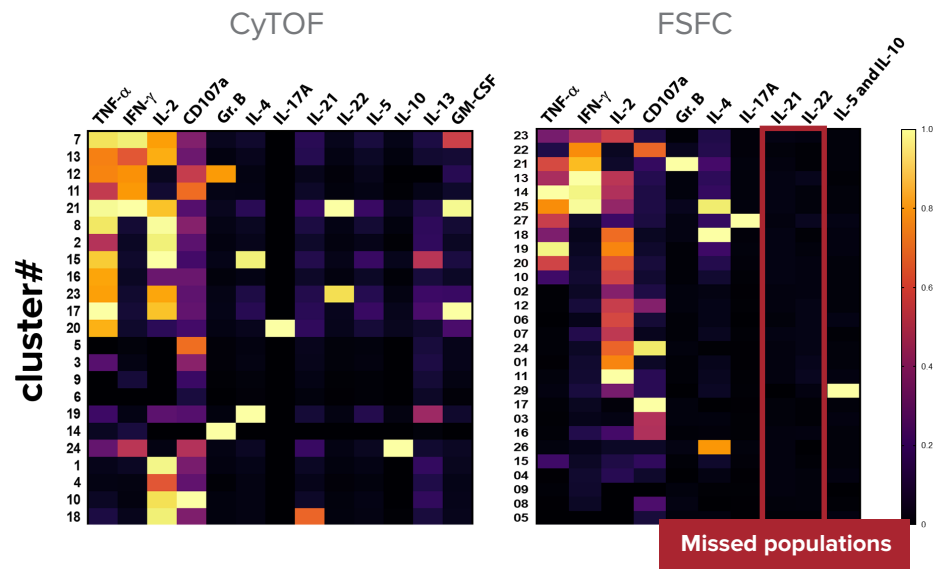
Results you can trust, reproduce and publish

See more with CyTOF Flow cytometry

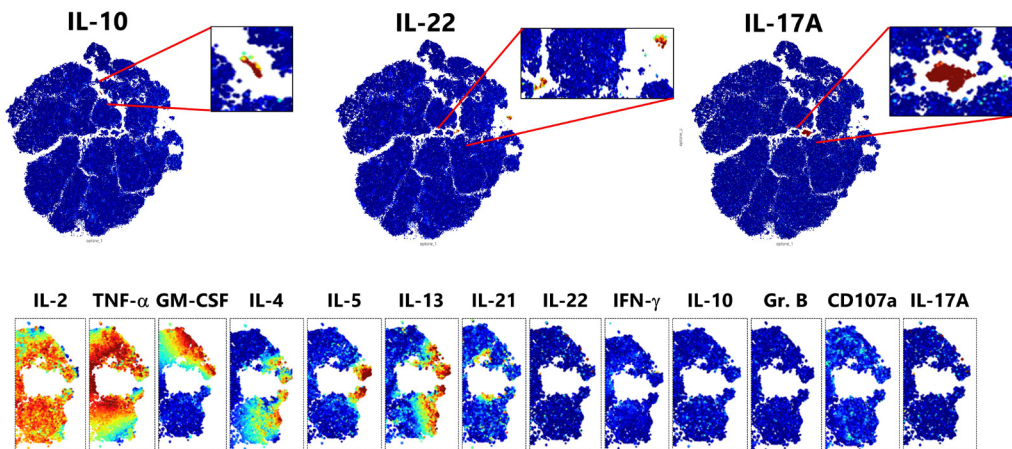
CyTOF detects more immune subpopulations in a single cell compared to full spectrum flow cytometry (FSFC).

Detect more cell populations with greater functional diversity.

The heat map on the left shows detection of IL-21 and IL-22, not present in the spectral data. Additionally, IL-5 and IL-10 were detected by CyTOF in independent channels.



Clear resolution and detection of low-frequency immune cells



Application of opt-SNE to this CyTOF dataset highlights distinct combinations of effector functions from type 1, type 2 and type 17 lineages.

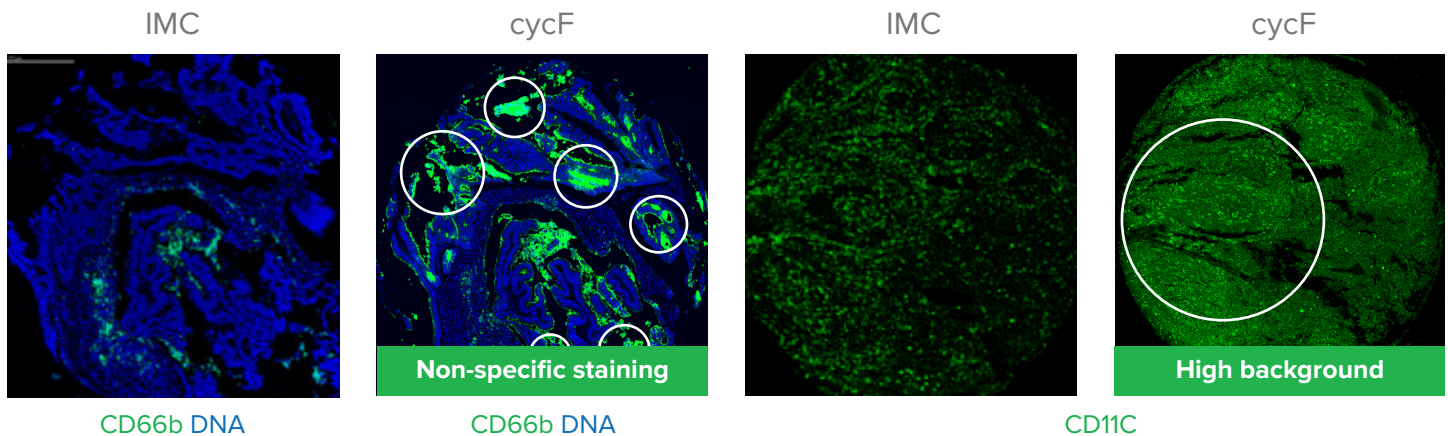
This data features findings from a comparison study using a 28-marker common panel including 12 cytokines. Data provided by Boston University.

See your true biology

See clearly with IMC

Tissue Imaging

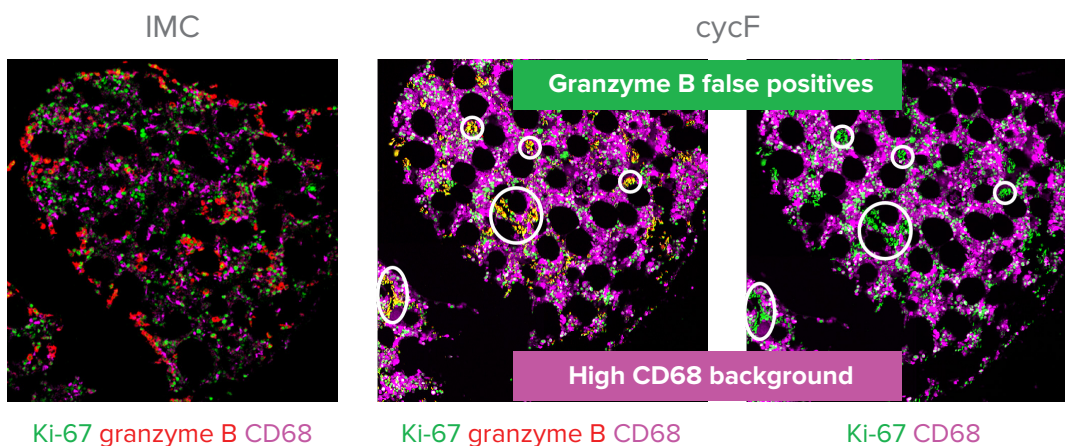
IMC allows highly specific staining without the challenges of autofluorescence or false positives inherent to cyclic fluorescence-based imaging (cycF).



The cycF image (right) shows non-specific false-positive CD66b signal in colon adenocarcinoma.

CD11c signal is obscured by autofluorescence with cycF (right) in lymph node tissue.

Image highly-autofluorescent tissue types without challenge



In bone marrow, cycF data (right) shows granzyme B signal co-localized to the nucleus with Ki-67. The normal cytoplasmic or membranous localization of granzyme B can only be seen with IMC (left). CD68 signal is clear with IMC but obscured with cycF.

Data generated from a comparison study with multiplex cyclic fluorescence (cycF) using a 27-marker common panel. Cyclic fluorescence data provided by Georgetown University.

The CyTOF Advantage.

A trusted technology used in

Over 200 clinical trials

Over 2,000 peer-reviewed publications

For **any** phase
of research

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