

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



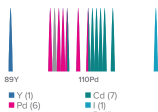
Reasons to choose CyTOF

# THE CYTOF

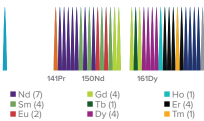
# Reasons to take your high-dimensional

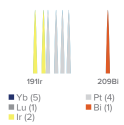
#### Precision data with discrete signals

Not impacted by spectral overlapping of fluorochromes and tissue autofluorescence



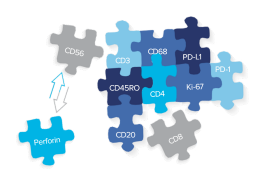
■ Rh (1)





#### 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.



#### Reproducible and comparable

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

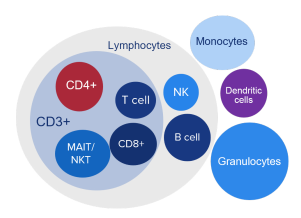


## research beyond the limits of fluorescence



#### Capture rare or unexpected cell populations

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





# 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





#### **Trusted by researchers**

The leading technology for high-parameter immune research





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

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



#### Risks of fluorescence

for high-parameter studies



**Missed** cell populations or **false positives** 



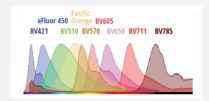
More iterations required in panel design



**Reduced sensitivity** where fluorescence overlap occurs



**Higher resource use** to compensate for spectral overlap



# From sample collection to high-dimensional insights in 3 days\*

### Flow cytometry

CD294

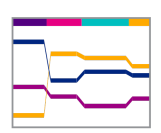
HLA-DR

Get started with the validated Maxpar® Direct<sup>™</sup> Immune Profiling Assay<sup>™</sup>.

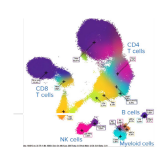
#### 1 tube 30 markers (<300 µL of whole blood)

#### CD4 CD38 CD11c CD45RA CCR6 CD14 CD45RO CCR7 CD20 CD66b

#### 5 min analysis



#### 37 cell populations



Surface profiles Cell proliferation **Apoptosis** Metabolism Phosphoproteins Cytokine production Transcription factors

### Minimal signal overlap

CHANNEL						3	ŧ		ě			ê		B			B					
89Y	1	0	٥	۰	0	0	0	0	0	0	۰	0	0	۰	0	0	۰	0	0	0	0	۰
142Nd	۰	1	0.02	٥	۰	۰	۰	۰	۰	۰	۰	۰	0.00	۰	۰	۰	۰	۰	۰	۰	۰	۰
14294		aa	1	0	0		0	0	0	0	0	0	0	۰	0	0	0	0	۰	0	0	۰
149Nd	۰	aa	0	1	0	0.01	۰	0	0	0	۰	۰	0	0	0	0	0	0	•	0	0	0
1475m		0	0	0	1	0.02	۰	0	0	0	0		۰	۰	0	۰	0	0	۰	۰	0	۰
146Nd	۰	۰	۰	0	۰	1	۰	0	0	0	0	0	0	0	0	0	0	0	•	0	0	0
1492m			۰	۰	0	0.01	1	۰	٥	۰	۰	۰	۰	۰	0	۰	۰	0	۰	٥	۰	۰
191Eu	0	0	٥	0	0	0	•	1	0.01	0.02	0	0	0	۰	0	0	0	0	0	0	0	0
D2Sm	0	0	0	0	0	0	0		1	0	0.01	0	0	۰	0	0	0	0	0	0	0	0
5 X u	0	0	۰	0	0	0	0	0.01	0	1	0	0	0	•	0	0	0	0	0	0	0	0
D49m		0	٥	0	0		0.01	0	0.08	0.02	1	0.02	0	0	0	0	0	0	0	0	0	0
155 Gd	0	۰	۰	0	0		۰	0		0	0.01	1		•	0	0	0	0.02	•	0	0	0
156 Gd		۰	۰	0	0	0	0	0	0	0	۰	0	1	0	0	0	0	0	0	002	0	
165Ho	0	0	0	0	0	0	0	0	0	0	0	0	۰	1	0	0	0	0	۰	0	۰	۰
1076	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1	0.07	0	0	0	0	0	0
1600		0	0	0	0		0	0	0	0	0	0		•	0.07	1		0	•	0	0	
10 9 Tm			۰	0	0		۰	۰	0	0	۰	0		•	0	0.02	1		•	0		
17176			۰	0	۰		0	0	0	0	۰	۰		•	0	0	•	1	000	000		
17 2/b			۰	0	0	0		0		0		0			0	0		0.01	1	0.10	aa	aa
17-4Yb			۰	0	۰	0	0	0	0	0	0	0	0		0		0	0	0.02	1	aa	aa
179Lu			۰	0	۰							0									1	۰
176Yb	-	0	0	0			-	-	-	-	-	0		-		-	-		-	-		1



The CyTOF flow cytometry image (far left) shows minimal spillover between metal channels when compared with the same panel from a competitor spectral flow cytometer (left).

Quantify and visualize 40-plus markers in a single run.

\* After panel and image analysis optimization

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.

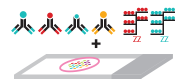
### **Tissue imaging**

Get started with our Maxpar IMC<sup>™</sup> Cell Segmentation Kits and ready-to-go high-plex panels.

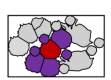
Simultaneous staining

One-step detection

High-plex data in minutes

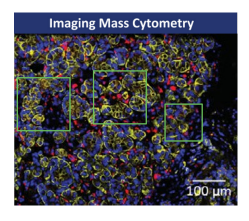


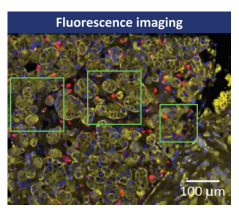




Tissue architecture
Protein modifications
Signaling pathway activation
Cell injury states
Cell proliferation
Transcriptional signatures

## **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

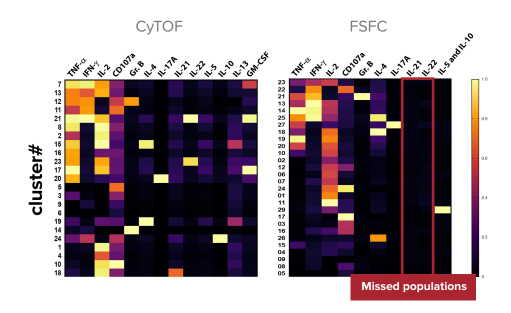
## See more with CyTOF

## Flow cytometry

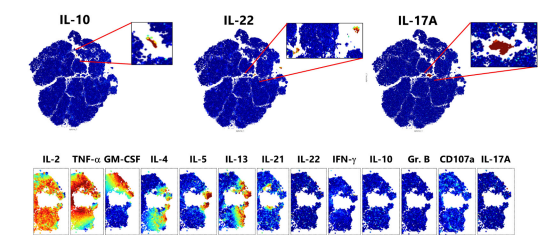
CyTOF detects more immune subpopulations in a single cell compared to full spectrum flow cytometry (FSFC).

# 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



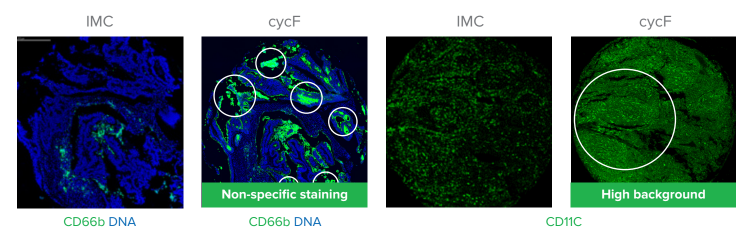
This data features findings from a comparison study using a 28-marker common panel including 12 cytokines. Data provided by Boston University. Application of opt-SNE to this CyTOF dataset highlights distinct combinations of effector functions from type 1, type 2 and type 17 lineages.

# See your true biology

## See clearly with IMC

## Tissue Imaging

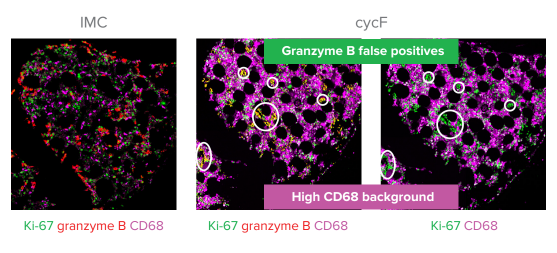
IMC allows highly specific staining without the challenges of autofluorescence or false positives inherent to cyclic fluorescence-based imaging (cycF).



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



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

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.

# The CyTOF Advantage.

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Over 2,000 peer-reviewed publications

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