

The background is a dark blue gradient filled with numerous small, multi-colored dots in shades of red, orange, yellow, green, and light blue. A prominent, thick, wavy line of bright pink and magenta particles flows from the bottom left towards the top right, creating a sense of dynamic movement.

Elabscience®

Multi-Color Panel Design in Flow Cytometry

Elabscience Bionovation Inc.



About Us

A Reliable Research Partner in Life Science and Medicine

Elabscience® stands at the forefront of biotechnology innovation, expertly combining independent design, R&D, manufacturing, and sales to deliver premier reagents and services for cell detection research. Our diverse product portfolio includes advanced solutions for detecting membrane and intracellular proteins (Flow cytometry antibodies), secreted proteins (ELISA kits), cell glycolipid metabolic intermediates and inorganic salts (Metabolism Assays), and comprehensive assessments of cellular function and health (Cell Apoptosis Assay, Cell cycle Assay, Cell Proliferation /Cytotoxicity/Viability).

To keep pace with the rapid advancements in research, we are dedicated to the continuous development of cutting-edge antibody and protein reagents, ensuring that our products evolve to meet the latest scientific needs. Our commitment extends beyond cell detection to include sophisticated cell isolation and characterization, empowering researchers to tackle the most complex challenges in cell biology. We pride ourselves on maintaining stringent quality control for every product, enhancing the accuracy and reliability of your experimental results. Our relentless pursuit of excellence since 2009 has established our presence in over 150 countries and regions worldwide.

Elabscience® is dedicated to addressing the evolving challenges in life sciences and healthcare. With a focus on delivering competitive, innovative solutions and driven by an unwavering commitment to excellence, we strive to be your trusted research partner in life sciences and medicine.

23,000 +

Citations

1,000 +

Cooperation Units

150 +

Countries Worldwide

5

Technical Platforms

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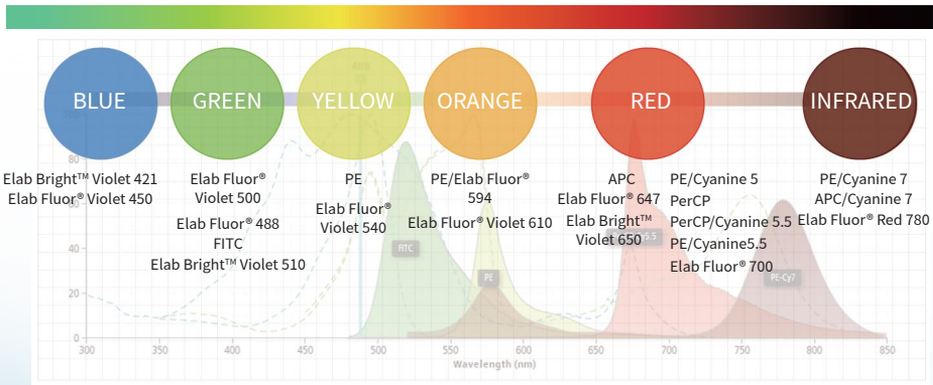
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01 Panel Design Principles



- Balance Antigen Density and Fluorochrome Brightness**
 High abundance antigen + Dim Fluorochrome.
 Low abundance antigen + Bright Fluorochrome.
- Avoid Spectral Overlap among Fluorochrome**
 Low abundance antigen can be detected in non-interference channel.
 High abundance antigen must be detected in channels that do not interfere with other channels.
- Minimize the Complexity of Analysis**
 Allow the spillover of mutually exclusive antigens.
 Allow the spillover of co-expressed antigens with highly abundance.
 Allow the spillover of offspring to their parents, but not the opposite.
- Use Tandem Fluorochromes Carefully**
 Tandem fluorochromes are necessary in multi-color panel design.
 Easily degraded when exposed to light or undergoing fixation.
 Follow protocols strictly to avoid tandem fluorochromes degradation.
- Cautions with Experiment Working Buffers**
 The acidic buffer or fixing step may destruct some dyes.
 eg: FITC is susceptible to low pH condition
 Fixation and extended storage lead to dye degradation.

02 Steps of Multi-Color Panel Design



■ STEP 01

Select the Target Markers

■ STEP 02

Check the Flow Cytometer Information

■ STEP 03

Check Fluorochrome Information

■ STEP 04

Pair Antigen with Fluorochrome

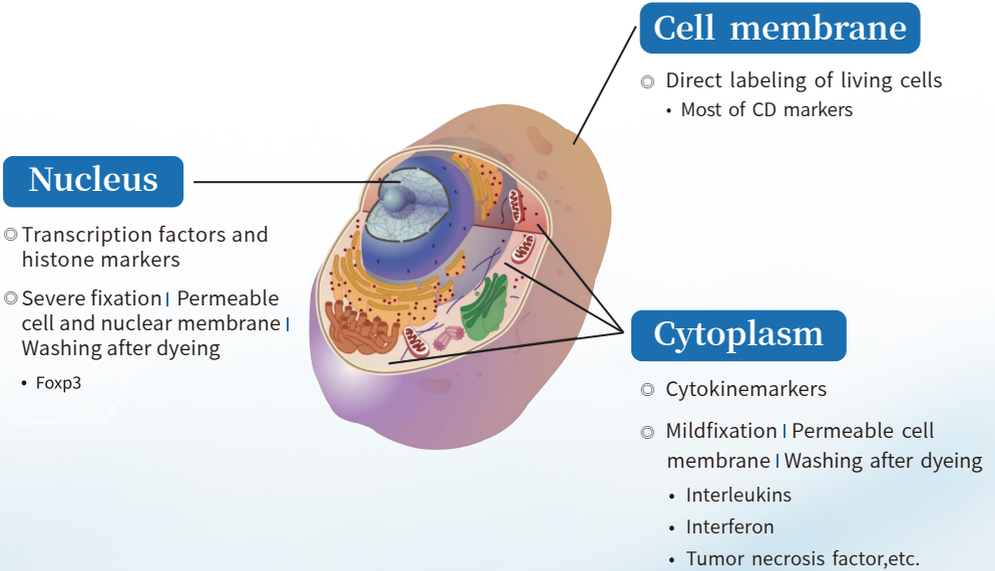
STEP 01 **Select the Target Markers**

Refer to Relevant Literature and Select the Target Markers

Human	Marker
B Cells	CD19
T Cells	CD3, CD4, CD8
Treg Cells	CD4, CD25, CD127
Th1/Th2/Th17 Cells	CD4, IFN- γ , IL-4, IL-17
Dendritic Cells	CD1c, CD83, CD141, CD209, MHC II
Natural Killer Cells	CD3 ⁺ , CD16, CD56
Macrophage	CD11b, CD68, CD163
Monocyte	CD14, CD16, CD64
Plasma Cells	CD138
Red Blood Cells	CD235a

Mouse	Marker
B Cells	CD19
T Cells	CD3, CD4, CD8
Treg Cells	CD4, CD25, Foxp3
Th1/Th2/Th17 Cells	CD4, IFN- γ , IL-4, IL-17
Dendritic Cells	CD11c, MHC II
Natural Killer Cells	CD3 ⁺ , CD49b (clone DX5) or NK1.1
Macrophage	F4/80, CD11b, CD80, CD86, CD206
Monocyte	CD11b, CD115, Gr-1, Ly-6C
Plasma Cells	CD138
Red Blood Cells	TER-119

Check the Marker Locations



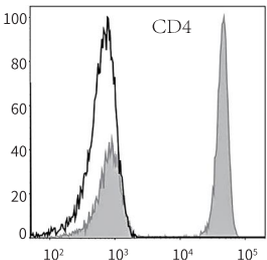
Classification of cell markers

- ☰ Generally speaking, most CD markers are located on the surface of cytomembrane. Cytokines, such as interleukins and interferon (IFN- α , IFN- β and IFN- γ), tumor necrosis factors (TNF- α , TNF- β) etc., are intracellular markers. And Foxp3 is the most popular intranuclear marker.
- ☰ For the intracellular and intranuclear markers, the cell needs to be fixed and broken before staining. If there is any intracellular or intranuclear maker, by conventional method, the first step is to stain the surface markers. Because "fixation" is easy to damage the tandem fluorescein, tandem dyes shall be not used in this step.

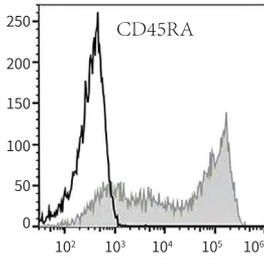
Check the Antigen Abundance

The antigen abundance can be roughly divided into three categories according to the expression of the corresponding antigen on/ in the cell types:

Easy to identify and antigens with obvious separation of negative-positive groups which can be easily distinguished. eg: CD3, CD4, CD19, etc.



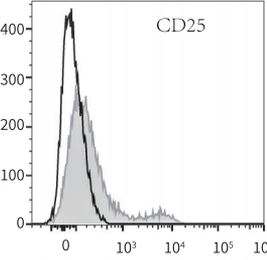
Negative and positive groups can be easily distinguished



Easy to identify, high abundance antigen expresses continuously. eg: CD27, CD28, CD45RA, CD45RO, etc.

High abundance antigen expresses continuously

Low abundance, active marker, or unknown but key markers. eg: CD25, STAT5, Foxp3, etc.

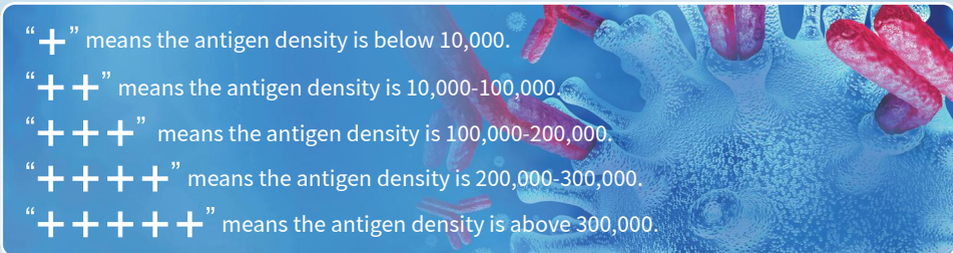


Low expression

Common Cell Surface Antigen Density

Cell Type	Marker	Density
Lymphocyte	CD3	++
	CD4	++
	CD8	++
	CD19	+
T cells	TCR	+++
	CD2	++
	CD3	+++
	CD5	++
	CD7	++
	CD45	++++
CD4+T cells	CD4	+++
	CD28	++
	CCR5	++
CD8+T Cells	CD8	++
	CD28	++
Monocyte	CD14	+++
	CD32	++
	CD64	++

Cell Type	Marker	Density
B cells	CD19	++
	CD20	+++
	CD21	++++
	CD22	++
	HLA-DR	+++
	CD11a	++
	CD40	+
Dendritic cells	CD86	++
	CD80	+
	CD11a	++
	CD40	++
NK Cells	CD80	+++
	CD86	++++
	CD56	++
Red blood cells	Glycophorin A	+++++
Neutrophils	CD14	+
	CD16	++++
Basic granulocyte	CD23	++



“+” means the antigen density is below 10,000.

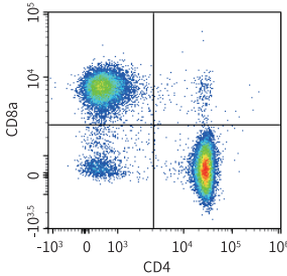
“++” means the antigen density is 10,000-100,000.

“+++” means the antigen density is 100,000-200,000.

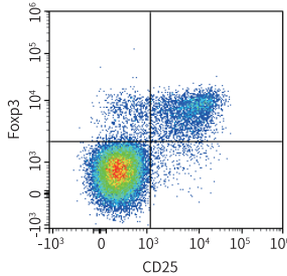
“++++” means the antigen density is 200,000-300,000.

“+++++” means the antigen density is above 300,000.

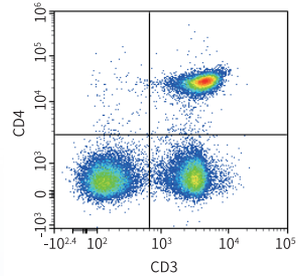
Check the Markers Interrelation



Mutual exclusion



Co-expression



Offspring and parents

The markers' relationship includes mutual exclusion, co-expression, offspring and parents, etc.

- ☞ Mutual exclusion means that two antigens will not be expressed on one cell at the same time, that is, if there is Protein A, there will be no Protein B, or if there is Protein B, there will be no Protein A. And mutually exclusive antigens allow fluorochrome spillover. eg: T cells are divided into CD4⁺ T cells and CD8⁺ T cells. CD4⁺ T cells express CD4 but not express CD8, and CD8⁺ T cells express CD8 rather than CD4.
- ☞ Antigen co-expression means that two antigens are expressed on the same cell. eg: Mouse Treg cells express CD25 and Foxp3 at the same time. Co-expressed but highly expressed antigens allow spillover.
- ☞ If the markers are offspring and the parents. Parents must be analyzed first. It means that the offspring antigen is analyzed on the basis of the parent antigen. eg: All T cells express CD3, and T cells are divided into CD4⁺ T cells and CD8⁺ T cells. In this case, CD3 is the parent, CD4 and CD8 are the offspring. Generally speaking, the spillover of offspring to parents is allowed, but spillover of parents to offspring is forbidden.

STEP 02 Check the Flow Cytometer Information

Channel and optional fluorochrome

Flow cytometer	Excitation	Detector (Filter)	Common fluorochrome
Take the flow cytometer with double laser as an example	488 nm	530/30	FITC, Elab Fluor® 488
		575/26	PE
		610/20	PE/TR, PE/Elab Fluor® 594
		695/40	PerCP/Cyanine5.5, PE/Cyanine5, PerCP
		780/60	PE/Cyanine7
	633 nm	660/20	APC, Elab Fluor® 647
		730/45	Elab Fluor® 700
		780/60	APC/Cyanine7, Elab Fluor® Red 780

Different manufacturers or different models have different configurations, even if the same model may have different configurations. When designing the panels, we must check the configuration of flow cytometer before we select appropriate fluorochrome. It is suggested to check the information as below:

- ① Excitation: there are several lasers can be used as excitation wavelength. The common flow cytometer lasers are 405nm, 488nm, 561nm, 633nm, etc.
- ② Detector: detectors are used to analysis emission wavelength.

STEP 03 Check Fluorochrome Information

- ☰ Check the fluorochrome excitation and emission wavelength, and confirm which fluorochrome can be used on the Flow cytometer according to the information of laser and detector.
- ☰ Check the relative brightness of the selected fluorochrome.
- ☰ Check the spillover among the fluorochrome.
- ☰ Check the characteristics of different fluorochrome, and select the appropriate fluorochrome according to the experimental purpose and requirements.

Fluorochrome Wavelength Information

Fluorochrome	Fluorochrome Emission Color	Excitation Laser Lines (nm)	Excitation Max(nm)	Emission Max(nm)
Elab Bright™ Violet 421	Blue	405	406	423
Elab Fluor® Violet 450	Blue	405	410	450
Elab Fluor® Violet 500	Green	405	410	501
Elab Bright™ Violet 510	Green	405	327,405	512
Elab Fluor® 488	Green	488	495	520
FITC	Green	488	490	530
Elab Fluor® Violet 540	Yellow	405	402	548
PE	Yellow	488/561	495, 565	575
Elab Fluor® Violet 610	Orange	405	421	613
PE/Elab Fluor® 594	Orange	488/561	495, 565	620
Elab Bright™ Violet 650	Red	405	407	646
APC	Red	633	650	660
Elab Fluor® 647	Red	633	650	670
PE/Cyanine5	Red	488/561	495, 565, 655	670
PerCP	Red	488	440, 480, 675	675
PerCP/Cyanine5.5	Red	488	440, 480, 675	675
PE/Cyanine5.5	Far Red	488/561	495, 565, 675	690
Elab Fluor® 700	Far Red	640	696	719
PE/Cyanine7	Infrared	488/561	495, 565, 755	775
Elab Fluor® Red 780	Infrared	633	625	765
APC/Cyanine7	Infrared	633	650, 760	780

Relative Brightness of Common Fluorochrome



Violet (405 nm)

- Elab Fluor® Violet 500
- Elab Fluor® Violet 450
- Elab Fluor® Violet 540
- Elab Fluor® Violet 610
- Elab Bright™ Violet 510
- Elab Bright™ Violet 421
- Elab Bright™ Violet 650

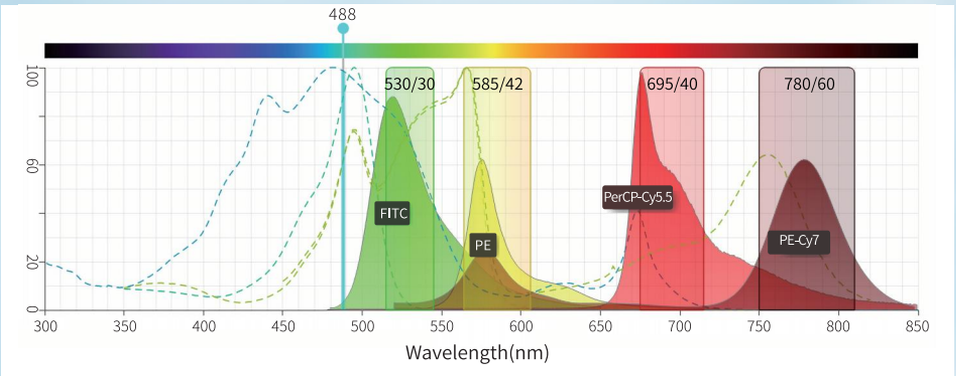
Blue (488 nm)

- PerCP
- Elab Fluor® 488
- FITC
- PerCP/Cyanine5.5
- PE
- PE/Cyanine5
- PE/Cyanine5.5
- PE/Elab Fluor® 594
- PE/Cyanine7

Red (640 nm)

- Elab Fluor® Red 780
- APC/Cyanine7
- APC
- Elab Fluor® 647
- Elab Fluor® 700

Overlap Information of Fluorochrome



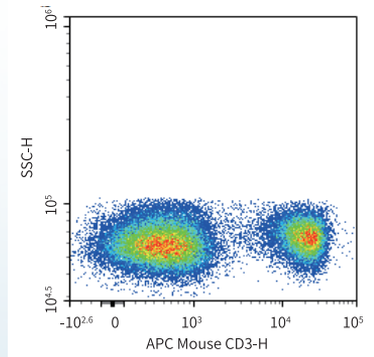
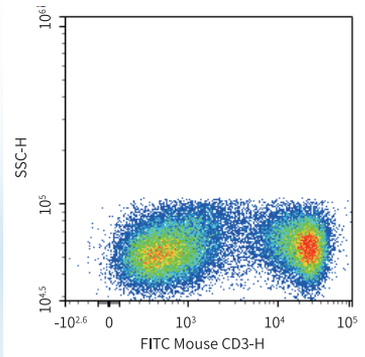
Fluorochrome Characteristics

Fluorochrome	Characteristics
Elab Bright™ Violet 421	Violet laser excitation, exhibits high brightness, excellent stability, and minimal spectral overlap with other fluorochromes.
Elab Fluor® Violet 450	A novel small-molecule fluorochromes that can replace Pacific Blue.
Elab Fluor® Violet 500	Violet laser excitation, exhibits a large Stokes shift, high brightness, good photostability and solubility, and is pH-insensitive.
Elab Bright™ Violet 510	Violet laser excitation, conjugated fluorochrome, high brightness, chemically stable, and photobleaching-resistant.
FITC	Easily affected by pH value. When the pH value decreases, the fluorescence intensity also decreases.
Elab Fluor® 488	Resistant to light and remains stable in a wide pH value (pH4-10).
Elab Fluor® Violet 540	Violet laser excitation, with a large Stokes shift, high stability, good water solubility, and moderate fluorescence intensity.
PE	High brightness, relatively stable.
Elab Fluor® Violet 610	Violet or yellow laser excitation, with a large Stokes shift, high brightness, photostability, good solubility, not affected by pH value.
PE/Elab Fluor® 594	The donor has a high molar extinction coefficient, resulting in stronger signal intensity for the tandem fluorochrome.
APC	High brightness, less stable than PE.
Elab Fluor® 647	Offers good fluorescence quantum yield and photostability; fluorescence is stable over pH 4-10.
Elab Bright™ Violet 650	Violet laser excitation, conjugated fluorochrome, high brightness (though slightly less bright than Elab Bright™ Violet 421), good stability, photobleaching-resistant and minimal spectral overlap with other fluorochromes.
PE/Cyanine 5	High brightness, easy to quench, not suitable to fixation, no matching with APC.
PerCP	Features a high extinction coefficient, high quantum yield, and a large Stokes shift.
PerCP/Cyanine5.5	Relatively stable (brightness and fixation) tandem fluorochrome.
PE/Cyanine5.5	Possesses a large Stokes shift, high fluorescence quantum yield, and good stability.
Elab Fluor® 700	Bright and stable; unaffected by pH changes in the range of 4-10 and exhibits good photostability.
PE/Cyanine7	High brightness, easy to quench, not suitable for fixation, no overlap with FITC, little interference and spillover with APC.
Elab Fluor® Red 780	Can replace APC/cyanine 7. Suitable for fixation and has less spillover to APC detector.
APC/Cyanine7	Weak brightness, not suitable for the analysis of low abundance antigens. Easy to quench and not suitable for fixation.

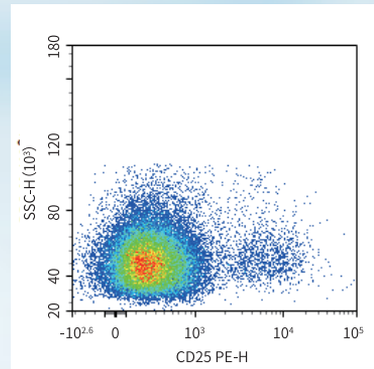
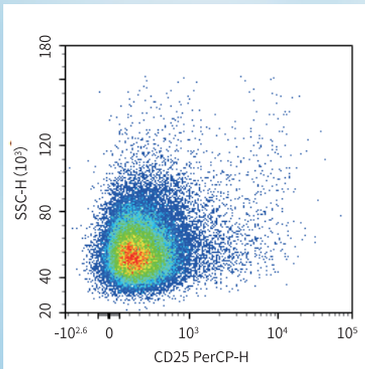
STEP 04 Pair Antigen with Fluorochrome

Balance Antigen Density and Fluorochrome Brightness

For high abundance antigen, weak or strong fluorochrome can be selected. As shown in the figure, high abundance antigen CD3 selects weak fluorochrome FITC or strong fluorochrome APC, in both situations, the results can be obviously observed.



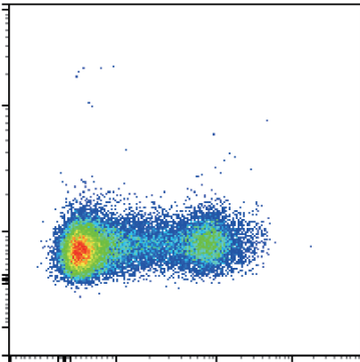
For low abundance antigen, strong fluorochrome must be selected. As shown in the figure, weak fluorochrome PerCP is selected by low abundance antigen CD25, leading to the inseparability of Negative-Positive cell groups. If strong fluorochrome PE is used, positive cell groups can be obvious to observe.



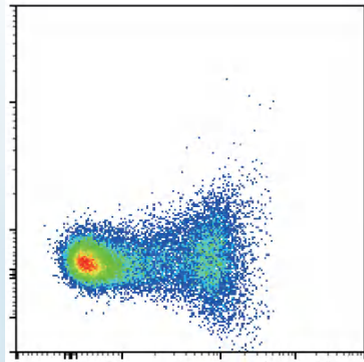
Avoid Spectral Overlap between Fluorochrome

Different fluorochrome may have spectral overlap. Try to use the fluorochrome combination with less spectral overlap in color matching, which can reduce the complexity of data analysis.

When the overlap occurs, fluorochrome compensation can only eliminate the background. For the reduced sensitivity of the disturbed detectors, it does not work.



No interference

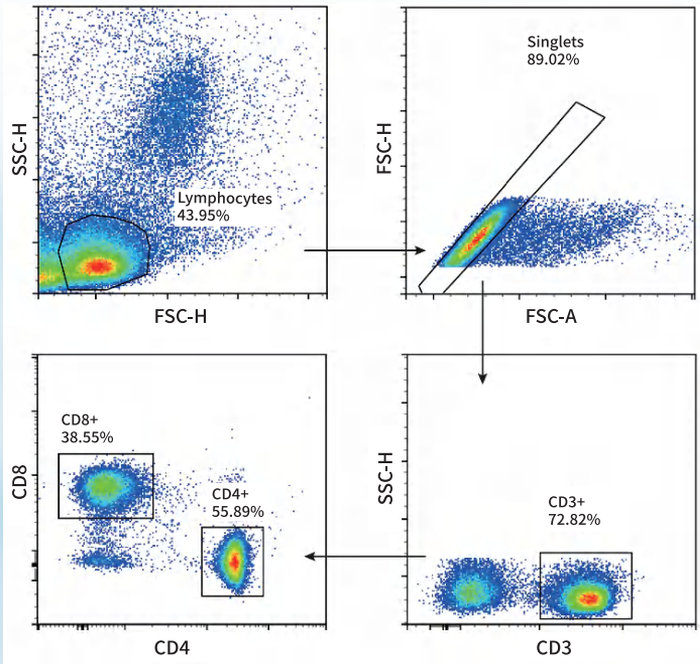


Serious interference

03 Cases of Multi-Color Panel Design

Case 1: Mouse Spleen T cells (3-color)

Marker	Fluorochrome	Clone No.	Cat. No.
CD3	Elab Fluor® Violet 450	17A2	E-AB-F1013Q
CD4	APC	GK1.5	E-AB-F1097E
CD8	Elab Fluor® Red 780	53-6.7	E-AB-F1104S

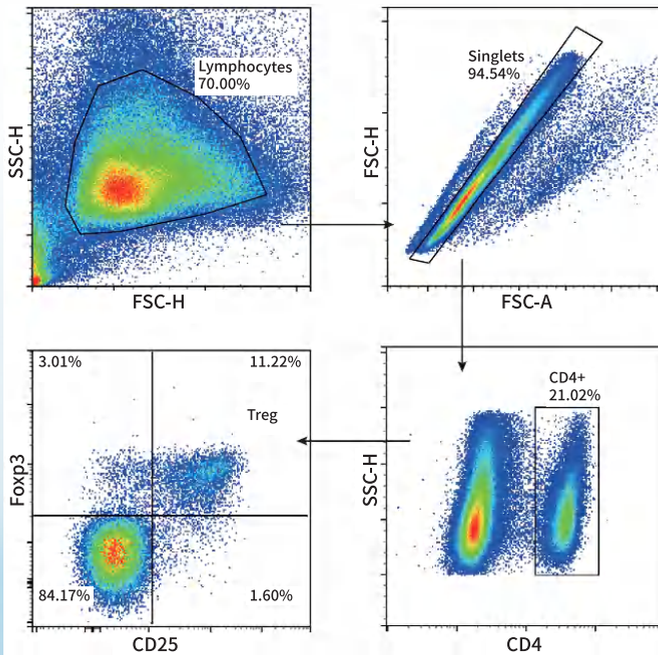


Tips:

- 1 Easy to distinguish the Negative-Positive cell groups, and there is no need for single staining tubes for compensation.
- 2 CD3/4/8 cells are easily distinguished, and generally speaking, isotype control is unnecessary.
- 3 The key factor of this experiment is the lysis of red blood cells. Excessive or insufficient lysis of red blood cells will lead to the unclear lymphocyte groups.

Case 2: Mouse Spleen Treg (3-color)

Marker	Fluorochrome	Clone No.	Cat. No.
CD4	FITC	GK1.5	E-AB-F1097C
CD25	APC	PC-61.5.3	E-AB-F1102E
Foxp3	PE	3G3	E-AB-F1238D

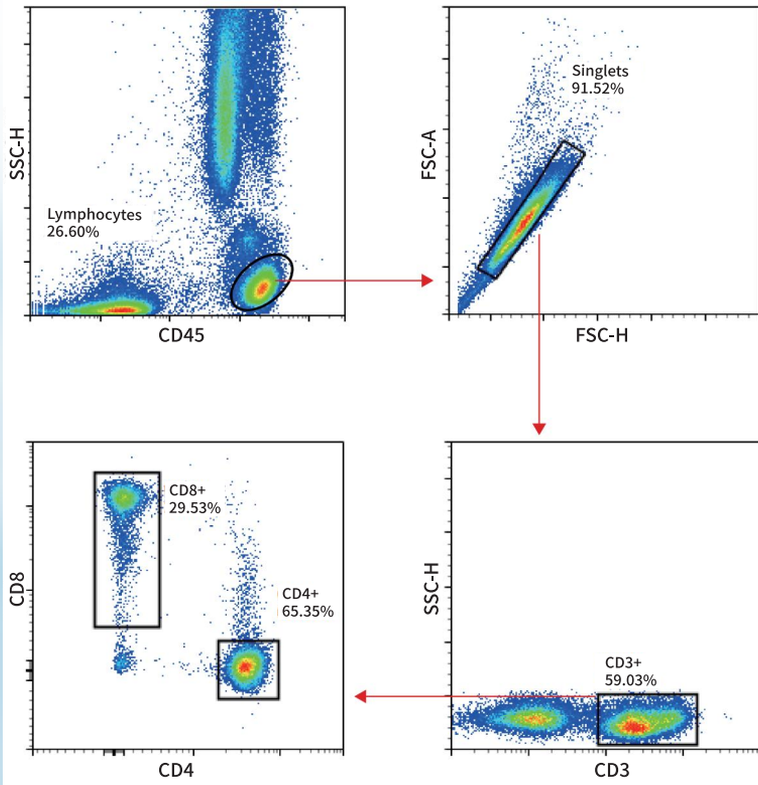


Tips:

- 1 Mouse Treg market is CD4⁺ CD25⁺ Fxp3⁺.
- 2 CD4⁺ cell group is obvious, and there is no need of isotype control. But CD25 and Fxp3 groups are not obvious, and isotype controls are needed.
- 3 There is fluorochrome spillover, and it is necessary to set single staining tubes for compensation.
- 4 Inappropriate use of Fixation/Permeabilization buffer may cause high background and unclear cell clustering. Please be careful.

Case 3: Human Peripheral Blood T Cells (4-color)

Marker	Fluorochrome	Clone No.	Cat. No.
CD45	Elab Fluor® Violet 450	HI30	E-AB-F1137Q
CD3	APC	OKT3	E-AB-F1001E
CD4	FITC	RPA-T4	E-AB-F1109C
CD8a	PE	OKT-8	E-AB-F1110D

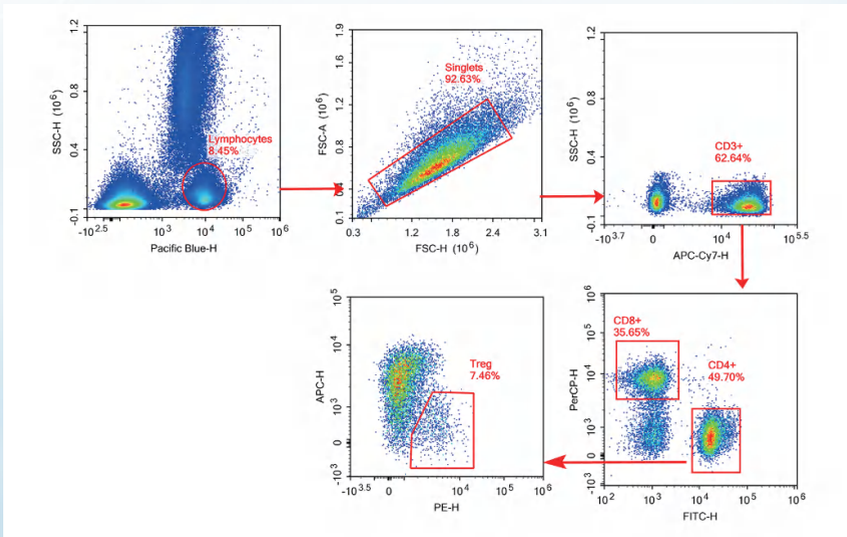


Tips:

- 1 For human peripheral blood T cells, it is suggested to use CD45, which can easily gate the lymphocyte group.
- 2 The cell groups are obvious, and there is no need to set single staining tubes for compensation.

Case 4: Human Peripheral Blood Treg (6-color)

Marker	Fluorochrome	Clone No.	Cat. No.
CD45	Elab Fluor® Violet 450	HI30	E-AB-F1137Q
CD3	Elab Fluor® Red 780	OKT3	E-AB-F1001S
CD4	FITC	RPA-T4	E-AB-F1109C
CD8a	PerCP/Cyanine5.5	OKT-8	E-AB-F1110J
CD25	PE	BC96	E-AB-F1194D
CD127	Elab Fluor® 647	A019D5	E-AB-F1152M

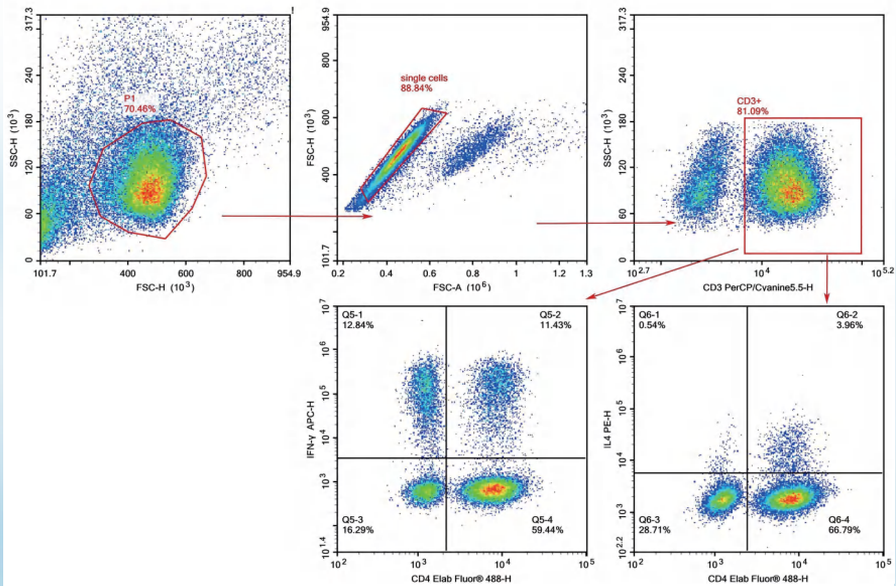


Tips:

- 1 Detecting human Treg by CD127 is no need of Fixation/Permeabilization step.
- 2 Gate the lymphocyte directly through CD45 and SSC, and then analyze the proportion of CD4⁺ CD25⁺ CD127^{-/low} cells. Treg cells account is about 3%-10% of lymphocytes in normal human peripheral blood.
- 3 It is suggested to set single staining tubes for compensation.

Case 5: Human Peripheral Blood Th1/Th2 (4-color)

Marker	Fluorescence	Clone No.	Cat. No.
CD3	PerCP/Cyanine5.5	UCHT1	E-AB-F1230J
CD4	Elab Fluor® 488	SK3	E-AB-F1352L
IFN-γ	APC	B27	E-AB-F1196E
IL-4	PE	MP4-25D2	E-AB-F1203D



Tips:

- 1 PMA stimulation can cause partial endocytosis of CD4 on the surface of human T cells, so we need to choose the CD4 clone SK3 with minimal impact on endocytosis.
- 2 Isotype Controls for IFN-γ and IL-4 are necessary, since the expression of cytokines is generally not high.
- 3 CD3⁺ CD4⁺ IFN-γ⁺ is Th1 cells, CD3⁺ CD4⁺ IL-4⁺ is Th2 cells.
- 4 The Permeabilization buffer may cause significant damage to cells, so it is recommended that the cell precipitates formed after centrifugation should be dispersed into cell suspensions before adding the Permeabilization buffer to reduce cell damage.

04 Data Analysis Services

You can also provide the original data of experimental results and logical relationship of markers to technical support. We can provide professional data analysis services for you.

**01**

Customer provides
basic information.



Estimate experimental
design and data integrity.

**02****03**

Data analysis.



Provide analysis results.

**04**



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