

# Approach to Bivariate Analysis of Data Acquired Using the Maxpar Direct Immune Profiling Assay

# Introduction

The Fluidigm Maxpar® Direct™ Immune Profiling Assay™ (Cat. No. 201325) was validated with Maxpar Pathsetter™ software (Cat. No. 401018). The software provides fast, reliable, and flexible automated analysis of FCS files from human PBMC and whole blood cells stained with the Maxpar Direct Immune Profiling Assay and acquired on a Helios™ mass cytometer. Maxpar Pathsetter automatically reports on 37 immune populations and is the recommended tool for reporting and analysis of results from the Maxpar Direct Immune Profiling Assay. Alternatively, bivariate gating can be used to define and enumerate these immune populations. The Maxpar Direct Immune Profiling Assay gating example, provided here, was developed using Cytobank, specifically for use with the Maxpar Direct Immune Profiling Assay. This gating strategy is based on the flow cytometry results data templates created by the Human Immunology Project Consortium (HIPC) [1], and on the Maxpar Direct Immune Profiling Assay probability state model used by Maxpar Pathsetter software. The gating example provides recommendations for gating immune populations in whole blood using markers available in the kit. The gating example can also be used for PBMC and is available as Fluidigm\_Maxpar Direct Immune Profiling Assay\_201325\_Gating Example\_v1.0.

# Symbols and Abbreviations in This Document

# **Symbols**

Purple squares denote navigation options within Cytobank.

Pink squares denote example populations in plots.

#### **Terms and Abbreviations**

Beads: EQ™ Four Element Calibration Beads (Cat. No. 201078)

DC: dendritic cell

MAIT cell: mucosal-associated invariant T cell

mDC: myeloid dendritic cell
NK cell: natural killer cell
NKT cell: natural killer T cell
pDC: plasmacytoid dendritic cell

Treg: regulatory T cell
Th: T helper cell

# **Accessing the Public Experiment**

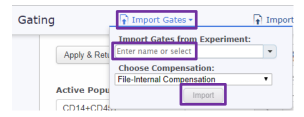
An example analysis for the panel kit, Fluidigm\_Maxpar Direct Immune Profiling Assay\_201325\_Gating Example\_v1.0, is available for reference at premium.cytobank.org.

# **Create a Clone of the Public Experiment**



The Public Experiment is read-only. To access the Gating tab, create a personal copy by cloning the experiment (available under the Actions tab).

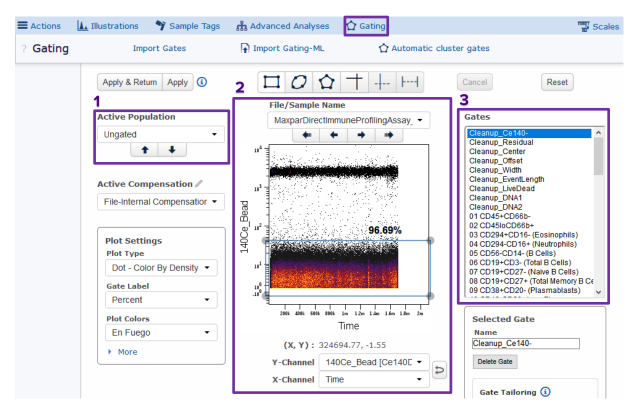
# **Apply Gating Strategy to New Data**



The gating strategy can be applied to new experiments by using the Import Gates function in the Gates tab. Enter the Cytobank experiment number **221569** in the dialog box and click **Import**. This applies the gating strategy of the selected Cytobank experiment to the files in the new experiment.

# **Overview of the Cytobank Gating Tab**

The Gating tab within Cytobank is used to create and adjust the gating strategies.



# Key features:

- 1 Active Population: Choose this population to view and adjust the gate.
- **2** File/Sample Name plot: View the gate parameters for the active population.
- **3** Gates: Choose a gate name to view and edit.

# **Defining Gates and Populations**

In Cytobank, the terms gate and population are not interchangeable.

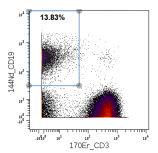


Figure 1. CD19+CD3- gate.

A gate is a selected region in the plot. Gates are defined on single parameters in histograms or two parameters in bivariate plots, for example, the CD19+CD3- gate (Figure 1). Gates can be rectangular, elliptical, or polygonal in shape.

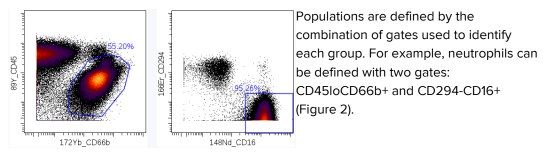
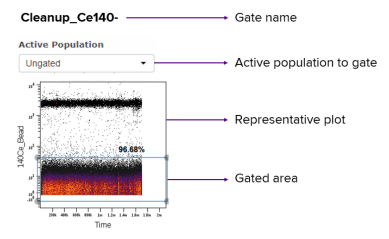


Figure 2. CD45loCD66b+ and CD294-CD16+ gates used to define the neutrophil population

# Using the Maxpar Direct Immune Profiling Assay Cell Gating Strategy

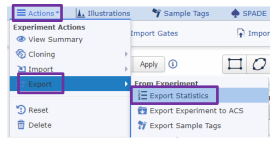
The steps for the cleanup and cell gating are outlined below with the gate name, Active Population, and a representative plot.



To adjust gates:

- 1 Select the gate name in the Gates box.
- 2 Select the **Active Population** from the drop-down menu.
- **3** Review the gate and modify to select the appropriate region if required.

# **Exporting Statistics from a Cytobank Experiment**



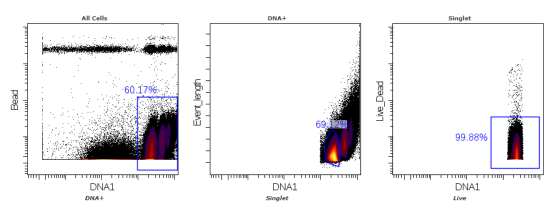
In addition to the gating strategy, statistics such as event counts and marker intensities can be exported using the Export Statistics tool.

A template for exporting event counts for each population is provided within the Public

Experiment (Fluidigm\_Maxpar Direct Immune Profiling Assay\_201325\_Gating Example\_v1.0). For more information on Cytobank features visit support.cytobank.org.

# **Data Cleanup**

Prior to cell gating using antibody targets (markers), a cleanup strategy is used to remove debris, normalization beads, doublets, and dead cells [2]. A common cleanup method used in mass cytometry is depicted below. DNA+ events are gated on DNA+Bead- (191Ir+140Ce-), then singlet events are gated using Event Length vs. DNA. The most abundant DNA+ Event\_Length (low/int) population is selected, followed by viable cell gating using a viability marker such as Cell-ID™ Intercalator-Rh (191Ir+103Rh-).

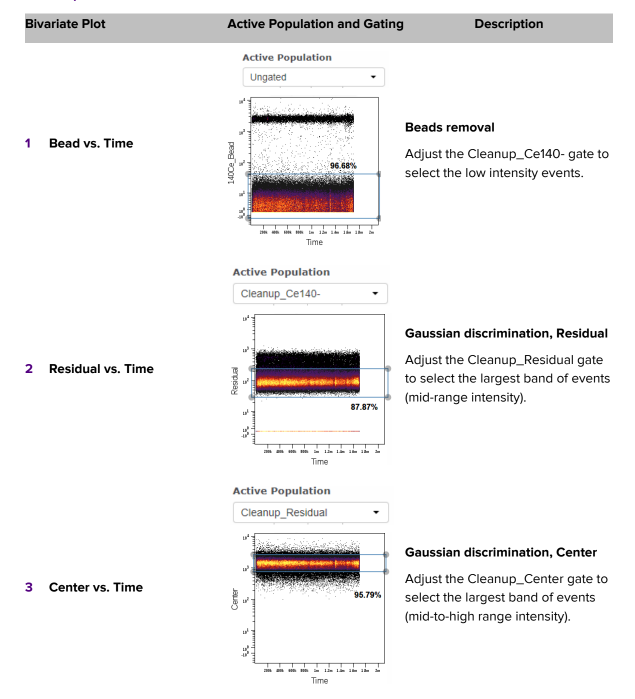


A new robust cleanup strategy was developed by Verity Software House and Fluidigm. This method has shown better aggregate and doublet removal than commonly applied gating strategies. In addition to the common parameters used (DNA, Bead, Event Length, Viability), this cleanup method uses Gaussian parameters for each event. The Gaussian discrimination (GD) channels (Center, Width, Offset, Residual) are recorded for each FCS file generated by Helios.

# **Cleanup Strategy**

Performing the automated cleanup routine using Maxpar Pathsetter software (Cat. No. 401018) is recommended. Below is an example of the manual cleanup strategy, which can be applied to an FCS file normalized in CyTOF® Software version 6.7.1016 (or higher). Each cleanup parameter is plotted against time. The gates are adjusted to remove aggregates, debris, beads, doublets, and dead cells.

# **Cleanup Gates**



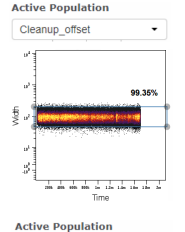
# 4 Offset vs. Time

# Cleanup\_Center 99.23% 101 101 2004 4004 6008 1006 Im 12m 12m 12m 12m 12m 2m Time

# Gaussian discrimination, Offset

Adjust the Cleanup\_Offset gate to select the largest band of events (mid-range intensity).

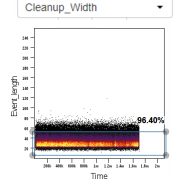
# 5 Width vs. Time



# Gaussian discrimination, Width

Adjust the Cleanup\_Width gate to select the largest band of events (mid-range intensity).

# 6 Event Length vs. Time

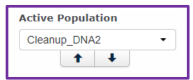


# **Event Length**

Adjust the Cleanup\_EventLength gate to select the largest band of events (low range intensity).

# **Bivariate Plot Active Population and Gating** Description **Active Population** Cleanup\_EventLength **Cell viability** Adjust the Cleanup\_LiveDead gate Live Dead vs. Time to select the largest band of events 99.85% (low range intensity). 200k 400k 600k 800k 1m 12m 1.4m 1.5m 1.5m 2m Time **Active Population** Cleanup\_LiveDead Nucleated (DNA-containing) cells, **94.06**% 1911r Adjust the Cleanup\_DNA1 (191lr) **DNA1 vs. Time** gate to select the largest band of events (mid-to-high range intensity). 600k 800k 1m 12m 1.4m 1.6m 1.8m 2m Time **Active Population** Cleanup\_DNA1 Nucleated (DNA-containing) cells, 1931r 97.96% Adjust the Cleanup\_DNA2 (193Ir) **DNA2** vs. Time 193Ir DNA2 gate to select the largest band of events (mid-to-high range intensity).

# **Global Parent Population**

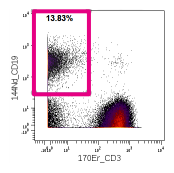


After each of the cleanup gates (Cleanup\_Ce140- to Cleanup\_DNA2) is applied, the Cleanup\_DNA2 population is considered to be the live singlet population.

This is the global parent population that will be used for subsequent immune cell gating.

# **Immune Cell Gating**

The ability to gate is dependent on the staining intensity of each marker and the resolution between positive and negative populations. Before analysis of critical samples, a preliminary pilot experiment should be performed on noncritical samples. Review pilot data for antibody staining quality. Evaluate pilot experiment marker intensities that are lower or higher than expected, which may affect the ability to identify populations.



The Maxpar Direct Immune Profiling Assay gating strategy is a manual method that uses bivariate plots to gate on positive and negative regions to identify different immune populations. Gates are adjusted based on the marker expression. For instance, B lymphocytes are gated as CD3-CD19+. The CD3 population is used as a negative population for CD19+ events. For some markers, a negative population may not be available in PBMC samples. View each gate in each file to ensure correct gating.

Many gates are used to identify multiple populations. The gates used to define each population, including intermediate populations, are listed in Appendix: Population Gating Tables. In the gating descriptions below, populations starting with a checkbox sign  $(\square)$  are considered end populations.

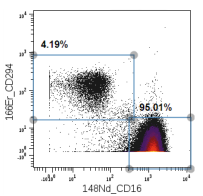
NOTE For more information, contact your local Fluidigm field application specialist.

# **Neutrophils and Eosinophils**

# **Bivariate Plot Active Population and Gating** Description **Active Population** Cleanup DNA2 Granulocyte exclusion CD66b vs. CD45 The CD45 vs. CD66b plot is used 42.64% to gate on granulocytes (CD45loCD66b+) and excludes granulocytes from subsequent CD45 CD45+ populations: 01 CD45+CD66b-(Lymphocytes, DCs, Monocytes) 02 CD45loCD66b+ (Granulocytes) 172Yb\_CD66b NOTE CD66b expression and granulocyte frequency are sensitive to sample preparation and sample type. PBMC preparations may have few granulocytes.

# **Active Population**

02 CD45loCD66b+ (Granul ▼



#### **☑** Eosinophils and neutrophils

From active population #02 CD45loCD66b+ (Granulocytes), the CD16 vs. CD294 plot is used to identify the two following granulocyte populations:

- 03 CD294+CD16-: Eosinophils
- 04 CD294-CD16+: Neutrophils

NOTE CD16 expression on granulocytes is sensitive to sample preparation and sample type. PBMC preparations may have reduced CD16 expression on granulocytes.

NOTE Neutrophils and eosinophils have been observed to nonspecifically bind to certain antibodies. Adding heparin to whole blood samples prior to staining (as followed in the Maxpar Direct Immune Profiling Assay Cell Staining and Data Acquisition User Guide, PN 400286) reduces nonspecific binding [3].

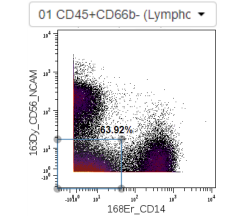
# 2 CD16 vs. CD294

# **B Cell Subsets**

1 CD14 vs. CD56

# Bivariate Plot Active Population and Gating Description

#### **Active Population**



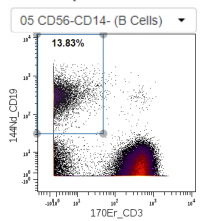
# NK cell and monocyte exclusion

Starting from active population #01 CD45+CD66b- (Lymphocytes, DCs, Monocytes), the CD56-CD14- gate is used to exclude natural killer cells and monocytes.

• 05 CD56-CD14- (B Cells)

2 CD3 vs. CD19

# **Active Population**

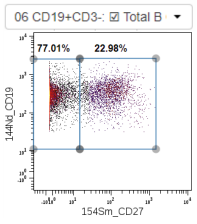


#### ☑ Total B cells

From active population #05 CD56-CD14- (B Cells), the CD19+CD3- gate excludes T cells and gates on total B cells.

• 06 CD19+CD3-: Total B Cells

# **Active Population**



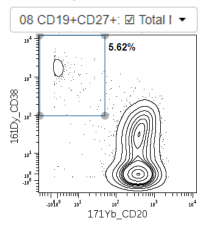
#### ☑ Naive and total memory B cells

From active population #06 CD19+CD3-: Total B Cells, the CD27 vs. CD19 plot includes two gates to identify naive B cells and total memory B cells:

- 07 CD19+CD27-: Naive B Cells
- 08 CD19+CD27+: Total Memory B Cells

3 CD27 vs. CD19

# **Active Population**



# ☑ Plasmablasts

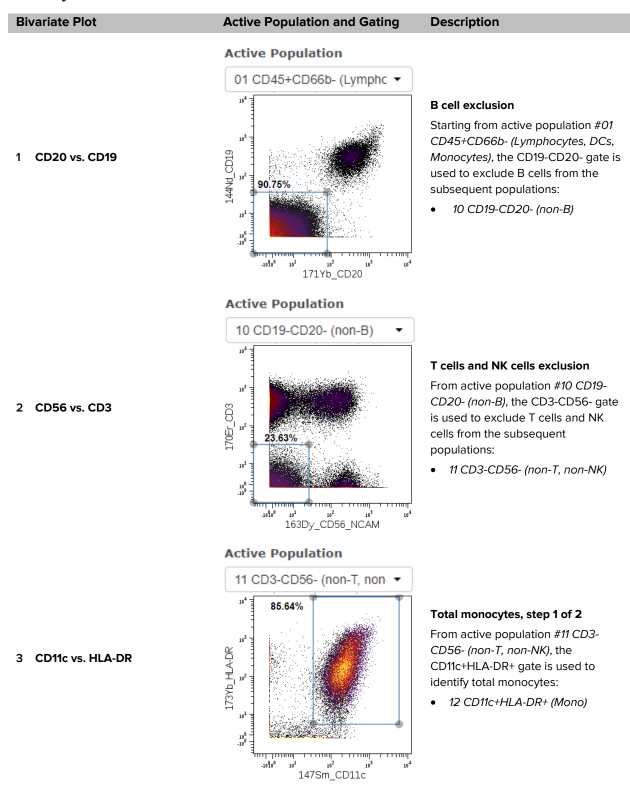
From active population #08 CD19+CD27+: Total Memory B Cells, the CD38+CD20- gate identifies the plasmablast population:

• 09 CD38+CD20-: Plasmablasts

**NOTE** For this gate it is recommended to use a contour plot to better identify rare plasmablast events.

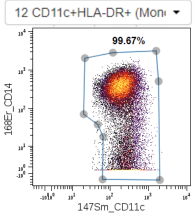
4 CD20 vs. CD38

# **Monocytes**



4 CD11c vs. CD14

# **Active Population**

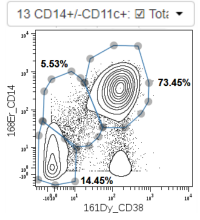


# ☑ Total monocytes step 2 of 2

From active population #12 CD11c+HLA-DR+ (Mono), the CD14+/-CD11c+ gate is used to confirm the presence of CD14 on total monocytes. CD14lo/-CD11clo/events are excluded using this gate:

13 CD14+/-CD11c+: Total Monocytes

# **Active Population**



# ☑ Monocytes, classical, transitional, and nonclassical

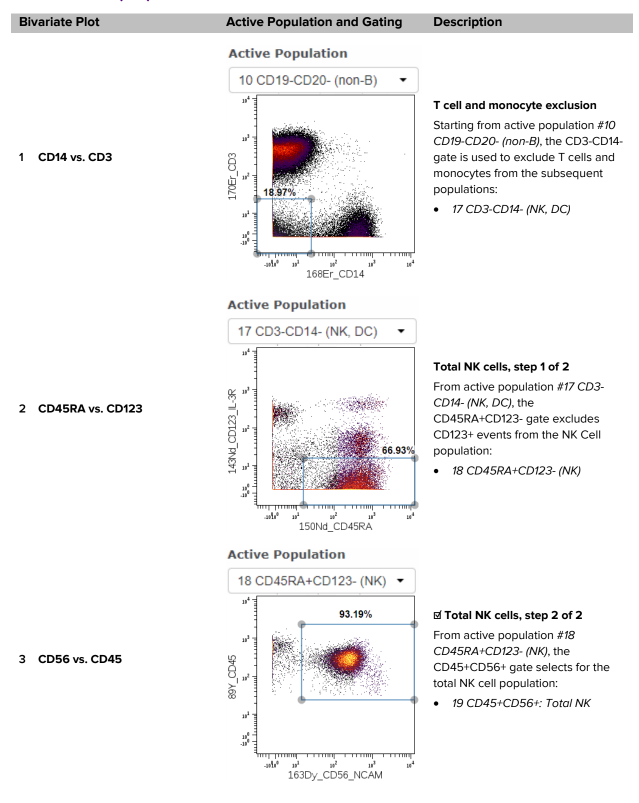
From active population #13 CD14+/-CD11c+: Total Monocytes, the CD14 vs. CD38 plot is used to distinguish between classical, transitional, and nonclassical monocytes:

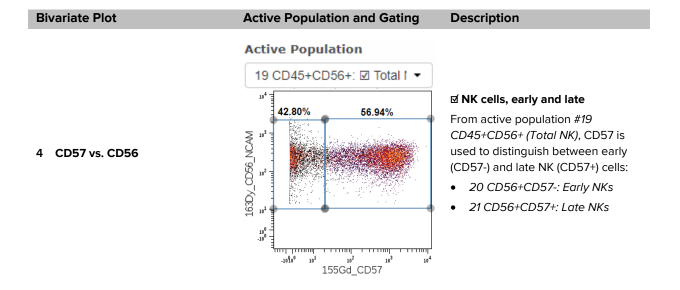
- 14 CD14+CD38+: Classical Monocytes
- 15 CD38lo/-CD14int: Transitional Monocytes
- 16 CD38-CD14-: Nonclassical Monocytes

NOTE For this gate, using a contour plot is recommended to better identify transitional monocytes events.

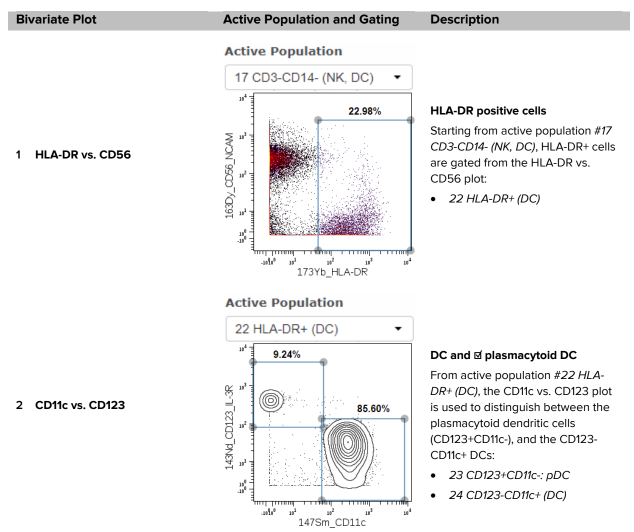
#### 5 CD14 vs. CD38

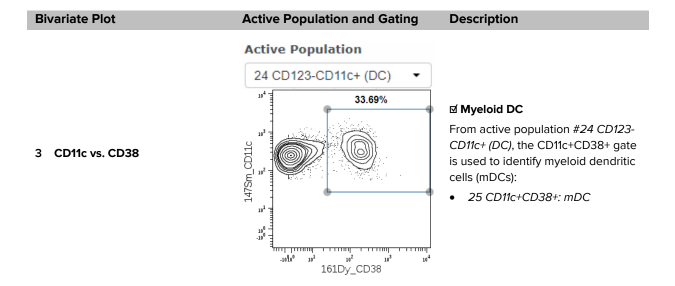
# **Natural Killer (NK) Cells**



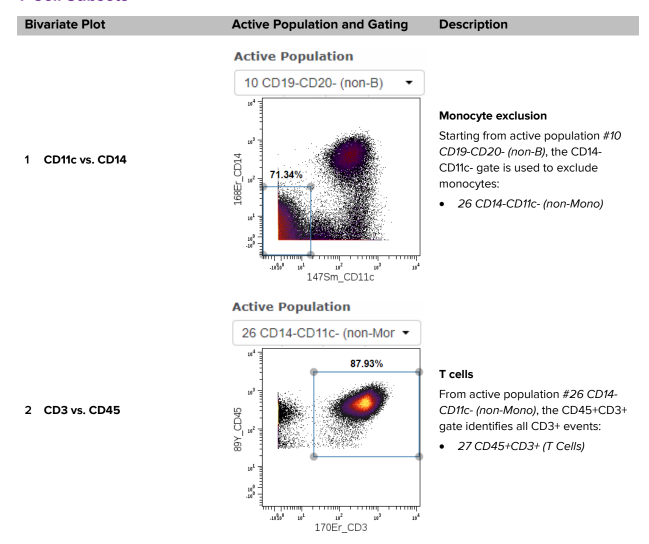


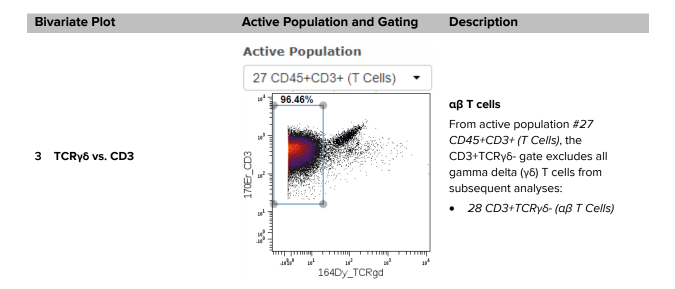
# **Dendritic Cells (DCs)**





# **T Cell Subsets**



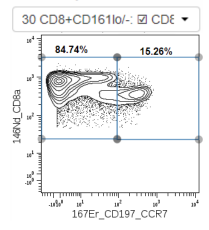


# CD8 $\alpha\beta$ T Cell Subsets

Bivariate Plot	Active Population and Gating	Description	
1 CD8a vs. CD4	Active Population  28 CD3+TCRγδ- (αβ T Cell ▼  10 <sup>4</sup> 41.09%  140Nd_CD8a	CD8 αβ T cells, step 1 of 2 From active population #28 CD3+TCRγδ- (αβ T Cells), the CD4- CD8+ gate identifies the CD8 T cell population:  • 29 CD4-CD8+ (CD8 αβ T Cells)	
2 CD161 vs. CD8a	Active Population  29 CD4-CD8+ (CD8 αβ T C ▼  81.91%  81.91%  1010  10	☑ CD8 αβ T cells, step 2 of 2  From active population #29 CD4- CD8+ (CD8 αβ T Cells), the CD161 vs. CD8a plot is used to gate out the CD8a+CD161hi cells from the subsequent CD8 populations:  • 30 CD8+CD161lo/-: CD8 αβ T Cells	

3 CCR7 vs. CD8a

# **Active Population**



# CD8 $\alpha\beta$ T cell stages, initial gate

From active population #30 CD8+CD161lo/-: CD8 a\(\beta\) T Cells, the CCR7 vs. CD8a plot is used to distinguish between the CCR7+ and CCR7- populations. These are intermediate subsets:

- 31 CD8+CCR7hi
- 34 CD8+CCR7lo/-

**NOTE** For this gate it is recommended to use a contour plot to better identify the transition between CCR7hi and CCR7lo/-, which typically occurs around 10<sup>2</sup>.

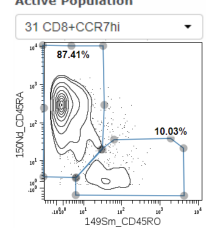
# 

From active population #31 CD8+CCR7hi, the CD45RO vs. CD45RA plot is used to identify naive (CD45RA+CD45RO-) and central memory (CD45RA-CD45RO+) CD8 T cells:

- 32 CD45RA+CD45RO-: CD8 Naive
- 33 CD45RA-CD45RO+: CD8 Central Memory

NOTE CD45RA+CD45RO- gate is used again in active population #39, and CD45RA-CD45RO+ gate is used again in active populations #40, #42, #46, and #50.

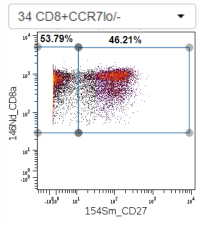
# Active Population



5 CD27 vs. CD8a

4 CD45RO vs. CD45RA

# **Active Population**



# 

From active population #34 CD8+CCR7/o/-, the CD27 vs. CD8a plot is used to gate on CD8 effector memory (CD27+) and CD8 terminal effector (CD27-) cells:

- 35 CD8+CD27+: CD8 Effector Memory
- 36 CD8+CD27-: CD8 Terminal Effector

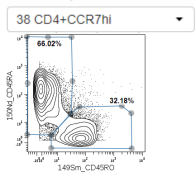
# CD4 aß T Cell Subsets

**Bivariate Plot Active Population and Gating** Description **Active Population** 28 CD3+TCRyδ- (αβ T Cell ▼ ☑ CD4 αβ T cells 56.73% From active population #28 10<sup>3</sup> CD3+TCRy $\delta$ - ( $\alpha\beta$  T Cells), the 1 CD8a vs. CD4 145Nd\_CD4 CD4+CD8- gate is used to identify CD4 T cells: 37 CD4+CD8-: CD4 αβ T Cells -10¶0<sup>0</sup> 146Nd\_CD8a **Active Population** CD4  $\alpha\beta$  T cell stages, initial gate 37 CD4+CD8-: ☑ CD4 αβ 1 ▼ The CCR7 vs. CD4 plot is used to distinguish between the CCR7hi and 104 CCR7lo/- populations. These are 61.85% 38.15% intermediate subsets: 103 38 CD4+CCR7hi 2 CCR7 vs. CD4 145Nd\_CD4 41 CD4+CCR7lo/-**NOTE** The transition between CCR7hi and CCR7lo/- typically occurs around 10<sup>2</sup> and is observed more easily as a contour plot. 167Er\_CD197\_CCR7

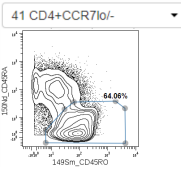
# **Active Population and Gating**

# Description

# **Active Population**

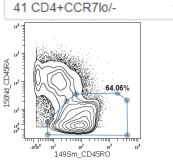


# **Active Population**

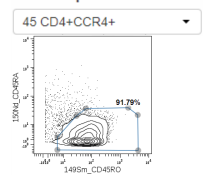


# 3 CD45RO vs. CD45RA

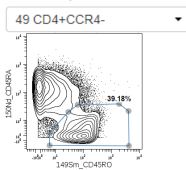
NOTE The CD45RA+CD45RO-gate was previously used for CD8 T cell subsets. The following are the same gates used in CD4 T cell populations.



#### **Active Population**



# **Active Population**



# ☑ CD4 αβ T cells, naive and central memory

From active population #38 CD4+CCR7hi, the CD45RO vs. CD45RA plot is used to identify naive (CD45RA+CD45RO-) and central memory (CD45RA-CD45RO+) CD4 T cells:

- 39 CD45RA+CD45RO-: CD4 Naive
- 40 CD45RA-CD45RO+: CD4 Central Memory

# CD4 $\alpha\beta$ T cells, effector memory and terminal effector

From active population #41 CD4+CCR7lo/-. CD45RA-CD45RO+ is used as an intermediate population to identify CD4 effector memory and terminal effector cells:

42 CD4+CCR7Io/-CD45RA-CD45RO+

# CD45RA-CD45RO+ selection for intermediate population of Treg cells

From active population #45 CD4+CCR4+, CD45RA-CD45RO+ is used as an intermediate population to identify T regulatory cells:

46 CD4+CCR4+CD45RA-CD45RO+

# CD45RA-CD45RO+ selection for intermediate population of Th1-like

From active population #49 CD4+CCR4-, CD45RA-CD45RO+ is used as an intermediate population to identify Th-1 like cells:

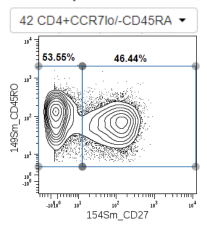
50 CD4+CCR4-CD45RA-CD45RO+

4 CD27 vs. CD45RO

# **Active Population and Gating**

# Description

# **Active Population**

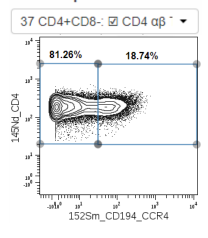


# 

From active population #42 CD4+CCR7lo/-CD45RA-CD45RO+, the CD27 vs. CD45RO plot it used to identify CD4 effector memory and terminal effector cells:

- 43 CD45RO+CD27+: CD4 Effector Memory
- 44 CD45RO+CD27-: CD4
   Terminal Effector

# **Active Population**



# CCR4 positive and negative selection for Treg and Th1-like intermediate populations

From active population #37 CD4+CD8-: CD4 a\beta T Cells, the CCR4 vs. CD4 plot is used to identify the intermediate population T regulatory cells (CD4+CCR4+).

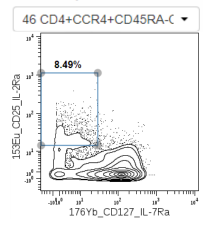
From active population #48 CD4+CXCR5-, the CCR4 vs. CD4 plot is used to identify the intermediate population Th1-like cells (CD4+CCR4-):

- 45 CD4+CCR4+
- 49 CD4+CCR4-

**NOTE** Active population #13 Total Monocytes can be used to identify the positive and negative threshold of CCR4.

5 CCR4 vs. CD4

# **Active Population**



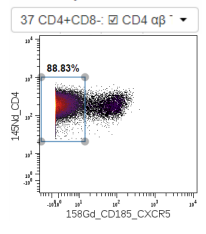
# ☑ Treg (T regulatory cells)

From active population #46 CD4+CCR4+CD45RA-CD45RO+, Treg cells are identified by gating on CD25hiCD127lo/- cells:

• 47 CD25hiCD127lo/-: Treg

# 6 CD127 vs. CD25

# **Active Population**



# CD4+CXCR5- selection for Th1-like, Th2-like, and Th-17 like intermediate populations

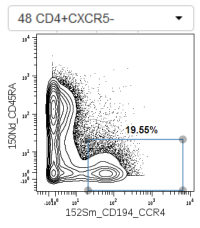
From active population #37 CD4+CD8-: CD4 a\beta T cells, CXCR5-cells are gated as an intermediate population to identify Th-1-like, Th2-like, and Th17-like cells:

48 CD4+CXCR5-

NOTE Chemokine receptor expression is sensitive to sample type, preparation, and treatment. For PBMC samples, the intensities of CXCR5 should be evaluated on noncritical samples.

# 7 CXCR5 vs. CD4

# **Active Population**



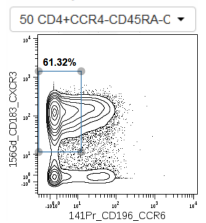
# CD45RA-CCR4+ selection for Th2like and Th17-like intermediate populations

From active population #48 CD4+CXCR5-, CD45RA-CCR4+ cells are gated as an intermediate population to identify Th2-like, and Th17-like cells:

52 CD45RA-CCR4+

# 8 CCR4 vs. CD45RA

# **Active Population**



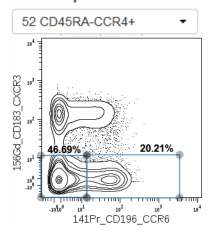
#### ☑ Th1-like

From active population #50 CD4+CCR4-CD45RA-CD45RO+, Th1like cells are gated as CXCR3+CCR6-:

51 CXCR3+CCR6-: Th1-like

# 9 CCR6 vs. CXCR3

# **Active Population**



# ☑ Th2-like and Th17-like

From active population #52 CD45RA-CCR4+, the CXCR3 vs. CCR6 plot is used to identify Th2-like and Th17-like cells:

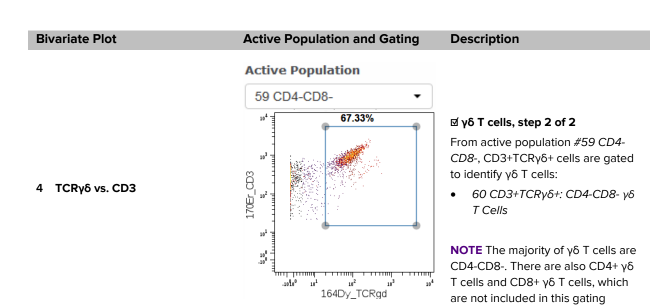
53 CXCR3-CCR6-: Th2-like

54 CXCR3-CCR6+: Th17-like

NOTE Chemokine receptor expression is sensitive to sample type, preparation, and treatment. For PBMC samples, the intensities of CXCR3 and CCR6 should be evaluated on noncritical samples to determine the ability to gate CD4 Th-like subsets.

# CD4- MAIT/NKT Cells and CD4-CD8- γδ T Cells

**Bivariate Plot Active Population and Gating** Description **Active Population** 28 CD3+TCRyδ- (αβ T Cell ▼ CD4 exclusion 30.71% From active population #28 CD3+TCRγδ- (αβ T Cells), CD3+CD4-1 CD4 vs. CD3 cells are gated as an intermediate 170Er CD3 population to identify CD4-MAIT/NKT cells: 55 CD3+CD4-145Nd CD4 **Active Population** 55 CD3+CD4-**☑ MAIT/NKT cells** From active population #55 25.11% CD3+CD4-, CD28+CD161hi cells are .60Gd CD28 gated to identify CD4- mucosal-2 CD161 vs. CD28 associated invariant T (MAIT)/natural killer T (NKT) cells: 56 CD28+CD161hi: CD4-MAIT/NKT NOTE MAIT and NKT cells can be 151Eu\_CD161 resolved with anti-TCR  $V\alpha7.2$ antibody. **Active Population** 27 CD45+CD3+ (T Cells) γδ T cells, step 1 of 2 From active population #27 CD45+CD3+ (T Cells), the CD4-CD8cells are gated as an intermediate 3 CD8a vs. CD4 [45Nd\_CD4 population to identify CD4-CD8-  $\gamma\delta$ T cells: 59 CD4-CD8-146Nd\_CD8a



strategy.

# **Basophils**

Bivariate Plot	Active Population and Gating	Description	
1 CD11c vs. HLA-DR	Active Population  11 CD3-CD56- (non-T, non ▼  10 1	Basophils, step 1 of 2 From active population #11 CD3- CD56- (non-T, non-NK), the CD11c- HLA-DR- gates on an intermediate population used to identify basophils:  57 HLA-DR-CD11c-	
2 CD294 vs. CD123	Active Population  57 HLA-DR-CD11c-  104  65.64%  65.64%	<ul> <li>☑ Basophils, step 2 of 2</li> <li>From active population #57 HLA-DR-CD11c-, basophils are gated as CD123+CD294+:</li> <li>● 58 CD123+CD294+: Basophils</li> <li>NOTE Basophils are a granulocyte population that can be found in PBMC preparations.</li> </ul>	

166Er\_CD294

# **Additional Gating Comments**

# **Monocyte Gating Strategy**

Classical, transitional, and nonclassical monocytes are more commonly gated using CD14 and CD16 (Figure 3) [4–6]. The CD38 marker is also expressed by monocytes [7–11]. The Maxpar Pathsetter software probability state immunophenotyping model produces more robust results using CD38 instead of CD16. Manual gating of total monocytes and monocyte subsets—classical, transitional, and nonclassical—using CD38 vs. CD14 were comparable to Pathsetter immunophenotyping model. Results from back-gating each monocyte subset from CD14 vs. CD38 to CD14 vs. CD16 appear similar to the common strategy (Figure 4). This document shows a CD14 and CD38 gating strategy. However, each user can determine a preferred gating strategy.

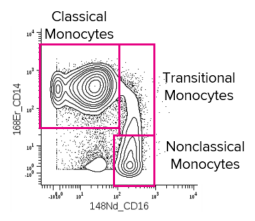


Figure 3. Common gating strategy for monocyte subsets using CD14 and CD16

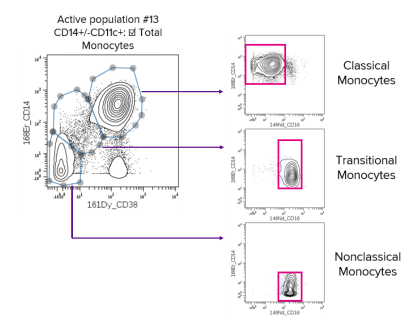


Figure 4. Back-gating the monocyte subsets from CD14 vs. CD38 to CD14 vs. CD16.

# **Further Gating Memory B Cells**

Memory B cells can be further divided into IgD+ Memory B cells and IgD- Memory B cells. This can be visualized by gating IgD vs. CD27 on total B cells (active population #06).

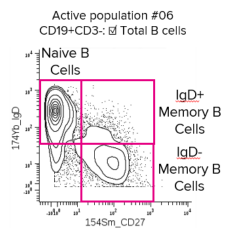


Figure 5. Gating strategy for identifying IgD+ and IgD- memory B cells.

# CD66b- Neutrophils

The Maxpar Pathsetter probability state immunophenotyping model identifies a population that has been labeled CD66b- Neutrophils. This population is identified as lineage negative (CD3-CD19-CD56-HLA-DR-CD123-CD45-CD66b-) phenotype with sample-dependent CD16lo/+ expression observed. Neutrophils have been shown to express different isoforms of CD66, such as CD66a, CD66b, CD66c, and CD66d [12,13], which may not be captured by CD66b in the panel. When comparing the spatial localization of CD66b- Neutrophils to CD66b+ Neutrophils in the Pathsetter Cen-se<sup>r™</sup> plot\*, these two populations cluster close together, suggesting that they may be related. With the addition of more markers to the panel, the CD66b- Neutrophil population may be more accurately defined.

<sup>\*</sup> Cen-se' (Cauchy-Enhanced Nearest-neighbor Stochastic Embedding) is an unsupervised nonlinear dimensionality reduction algorithm developed by Verity Software House, based on the original t-SNE (t-distributed stochastic neighbor embedding) algorithms commonly used for visualizing high-dimensional mass cytometry data [14].

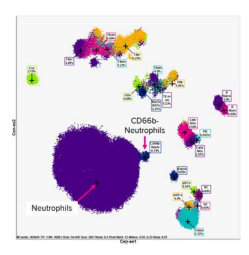


Figure 6. Cen-se' plot in Pathsetter showing the nearby clustering of the Neutrophils and CD66b-Neutrophils in a whole blood sample. The Neutrophil populations are not displayed in the Cen-se' plot of the default Pathsetter report.

# **Additional considerations**

- Marker intensities may vary across different samples. To adjust the gate for each specific file and/or population, use the Gate Tailoring box below the Gates box. For instance, the CD45RA vs. CD45RO gate can be adjusted for CD4 and CD8 T cells.
- Gates can be added and removed for each population in the Population Manager.
- Always check all gates for each file.
- For more information on Cytobank features go to support.cytobank.org.

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# **Appendix: Population Gating Tables**

Table 1. Gates used to distinguish cell populations stained by the Maxpar Direct Immune Profiling Assay (Cat. No. 201325). All listed cell populations should also include the cleanup gates described in the cleanup strategy.

		Population	Gate #	Gates	Population	Gate #	Gates
		Neutrophils	02	CD45loCD66b+	Facinantila	02	CD45loCD66b+
	es		04	CD294-CD16+	Eosinophils	03	CD294+CD16-
	Granulocytes		01	CD45+CD66b-			
			10	CD19-CD20-			
	anı	Basophils	11	CD3-CD56-			
	Gr		57	HLA-DR-CD11c-			
			58	CD123+CD294+			
			01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			26	CD14-CD11c-		26	CD14-CD11c-
		CD8 αβ T cells	27	CD45+CD3+	CD8 αβ T cells,	27	CD45+CD3+
		(Total;	28	CD3+TCRγδ-	Naïve	28	CD3+TCRγδ-
		CD161lo/-)	29	CD4-CD8+	raive	29	CD4-CD8+
			30	CD8+CD161lo/-		30	CD8+CD161lo/-
						31	CD8+CCR7hi
						32	CD45RA+CD45RO-
		CD8 αβ T cells, Central Memory	01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
Lymphocytes	S		26	CD14-CD11c-		26	CD14-CD11c-
\ <u>\f_2</u>	Cells		27	CD45+CD3+	CD8 αβ T cells,	27	CD45+CD3+
မို	-		28	CD3+TCRγδ-	Effector	28	CD3+TCRγδ-
E	ср8		29	CD4-CD8+	Memory	29	CD4-CD8+
\ \frac{1}{2}	O		30	CD8+CD161lo/-		30	CD8+CD161lo/-
			31	CD8+CCR7hi		34	CD8+CCR7lo/-
			33	CD45RA-CD45RO+		35	CD8+CD27+
		CD8 αβ T cells, Terminal Effector	01	CD45+CD66b-			
			10	CD19-CD20-			
			26	CD14-CD11c-			
			27	CD45+CD3+			
			28	CD3+TCRγδ-			
			29	CD4-CD8+			
			30	CD8+CD161lo/-			
			34	CD8+CCR7lo/-			
			36	CD8+CD27-			

		Population	Gate #	Gates	Population	Gate #	Gates
			01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			26	CD14-CD11c-		26	CD14-CD11c-
		CD4 αβ T cells (Total)	27	CD45+CD3+	CD4 αβ T cells,	27	CD45+CD3+
			28	CD3+TCRγδ-	Naïve	28	CD3+TCRγδ-
			37	CD4+CD8-		37	CD4+CD8-
						38	CD4+CCR7hi
						39	CD45RA+CD45RO-
			01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			26	CD14-CD11c-		26	CD14-CD11c-
		CD4 αβ T	27	CD45+CD3+	CD4 αβ T cells,	27	CD45+CD3+
		cells, Central	28	CD3+TCRγδ-	Effector	28	CD3+TCRγδ-
		Memory	37	CD4+CD8-	Memory	37	CD4+CD8-
			38	CD4+CCR7hi		41	CD4+CCR7lo/-
			40	CD45RA-CD45RO+		40	CD45RA-CD45RO+
						43	CD45RO+CD27+
			01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			26	CD14-CD11c-		26	CD14-CD11c-
es	<u>8</u>	CD4 αβ T cells, Terminal Effector	27	CD45+CD3+		27	CD45+CD3+
\ ₹	Sel		28	CD3+TCRγδ-	Treg	28	CD3+TCRγδ-
Lymphocytes	CD4 T Cells		37	CD4+CD8-		37	CD4+CD8-
Εď	<b>D4</b>		41	CD4+CCR7lo/-		45	CD4+CCR4+
Ę	บ		40	CD45RA-CD45RO+		46	CD45RA-CD45RO+
			44	CD45RO+CD27-		47	CD25hiCD127lo/-
		Th1-like	01	CD45+CD66b-	Th2-like	01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			26	CD14-CD11c-		26	CD14-CD11c-
			27	CD45+CD3+		27	CD45+CD3+
			28	CD3+TCRγδ-		28	CD3+TCRγδ-
			37	CD4+CD8-		37	CD4+CD8-
			48	CD4+CXCR5-		45	CD4+CXCR5-
			49	CD4+CCR4-		52	CD45RA-CCR4+
			50	CD45RA-CD45RO+		53	CXCR3-CCR6-
			51	CXCR3+CCR6-			
		Th17-like	01	CD45+CD66b-			
			10	CD19-CD20-			
			26	CD14-CD11c-			
			27	CD45+CD3+			
			28	CD3+TCRγδ-			
			37	CD4+CD8-			
			45	CD4+CXCR5-			
			52	CD45RA-CCR4+			
			54	CXCR3-CCR6+			

		Population	Gate #	Gates	Population	Gate #	Gates
			01	CD45+CD66b-		01	CD45+CD66b-
	ells		10	CD19-CD20-		10	CD19-CD20-
	ŭ	Gamma-delta T	26	CD14-CD11c-	MAIT/NKT CD4-	26	CD14-CD11c-
	Ē	cells, CD4-CD8-	27	CD45+CD3+	cells	27	CD45+CD3+
	Other T cells		59	CD4-CD8-		55	CD3+CD4-
	O		60	CD3+TCRγδ+		56	CD28+CD161hi
			01	CD45+CD66b-		01	CD45+CD66b-
		Total B cells	05	CD56-CD14-	Nation Deadle	05	CD56-CD14-
			06	CD19+CD3-	Naive B cells	06	CD19+CD3-
	<u>s</u>					07	CD19+CD27+
	cells		01	CD45+CD66b-		01	CD45+CD66b-
es	В	T	05	CD56-CD14-		05	CD56-CD14-
7		Total Memory B cells	06	CD19+CD3-	Plasmablasts	06	CD19+CD3-
ğ		Cells	08	CD19+CD27+		08	CD19+CD27+
Lymphocytes						09	CD38+CD20-
Ę			01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
		Total NK cells	17	CD3-CD14-	Early NK cells	17	CD3-CD14-
			18	CD45RA+CD123-	Early NK Cells	18	CD45RA+CD123-
	S		19	CD45+CD56+		19	CD45+CD56+
	ll es					20	CD56+CD57-
	NK cells	Late NK cells	01	CD45+CD66b-			
	_		10	CD19-CD20-			
			17	CD3-CD14-			
			18	CD45RA+CD123-			
			19	CD45+CD56+			
			21	CD56+CD57+			
		Total Monocytes	01	CD45+CD66b-		01	CD45+CD66b-
			10	CD19-CD20-		10	CD19-CD20-
			11	CD3-CD56-	Classical	11	CD3-CD56-
			12	CD11c+HLA-DR+	Monocytes	12	CD11c+HLA-DR+
	rtes		13	CD14+/-CD11c+		13	CD14+/-CD11c+
	Monocytes					14	CD38+CD14hi
	ouo	Transitional Monocytes	01	CD45+CD66b-		01	CD45+CD66b-
	Σ		10	CD19-CD20-		10	CD19-CD20-
			11	CD3-CD56-	Nonclassical	11	CD3-CD56-
			12	CD11c+HLA-DR+	Monocytes	12	CD11c+HLA-DR+
			13	CD14+/-CD11c+		13	CD14+/-CD11c+
			15	CD38lo/-CD14int		16	CD38-CD14-
	SIIs	Plasmacytoid Dendritic cells	01	CD45+CD66b-		01	CD45+CD66b-
	: ce		10	CD19-CD20-		10	CD19-CD20-
	itic		17	CD3-CD14-	Myeloid Dendritic cells	17	CD3-CD14-
	ndr		22	HLA-DR+	Denuntic cens	22	HLA-DR+
	Dendritic cells		23	CD123+CD11c-		24	CD123-CD11c+
	Δ					25	CD11c+CD38+

Table 2. Active cell populations identified in Cytobank for manual gating of the Maxpar Direct Immune Profiling Assay (Cat. No. 201325). Active populations that contain parentheses are descriptors of intermediate populations for which those gates are used. Populations ending with a descriptor following a colon and starting with a checkbox sign (🗹) are end populations.

01 CD45+CD66b- (Lymphocytes, DCs, Monocytes)	31 CD8+CCR7hi
02 CD45loCD66b+ (Granulocytes)	32 CD45RA+CD45RO-: ☑ CD8 Naive
03 CD294+CD16-: ☑ Eosinophils	33 CD45RA-CD45RO+: ☑ CD8 Central Memory
04 CD294-CD16+: ☑ Neutrophils	34 CD8+CCR7lo/-
05 CD56-CD14- (B cells)	35 CD8+CD27+: ☑ CD8 Effector Memory
06 CD19+CD3-: ☑ Total B cells	36 CD8+CD27-: ☑ CD8 Terminal Effector
07 CD19+CD27-: ☑ Naive B cells	37 CD4+CD8-: ☑ CD4 αβ T cells
08 CD19+CD27+: ☑ Total Memory B cells	38 CD4+CCR7hi
09 CD38+CD20-: ☑ Plasmablasts	39 CD45RA+CD45RO-: ☑ CD4 Naive
10 CD19-CD20- (non-B)	40 CD45RA-CD45RO+: ☑ CD4 Central Memory
11 CD3-CD56- (non-T, non-NK)	41 CD4+CCR7lo/-
12 CD11c+HLA-DR+ (Mono)	42 CD4+CCR7Io/-CD45RA-CD45RO+
13 CD14+/-CD11c+: ☑ Total Monocytes	43 CD45RO+CD27+: ☑ CD4 Effector Memory
14 CD14+CD38+: ☑ Classical Monocytes	44 CD45RO+CD27-: ☑ CD4 Terminal Effector
15 CD38lo/-CD14int: ☑ Transitional Monocytes	45 CD4+CCR4+
16 CD38-CD14-: ☑ Nonclassical Monocytes	46 CD4+CCR4+CD45RA-CD45RO+
17 CD3-CD14- (NK, DC)	47 CD25hiCD127lo/-: ☑ Treg
18 CD45RA+CD123- (NK)	48 CD4+CXCR5-
19 CD45+CD56+: ☑ Total NK	49 CD4+CCR4-
20 CD56+CD57-: ☑ Early NKs	50 CD4+CCR4-CD45RA-CD45RO+
21 CD56+CD57+: ☑ Late NKs	51 CXCR3+CCR6-: ☑ Th1-like
22 HLA-DR+ (DC)	52 CD45RA-CCR4+
23 CD123+CD11c-: ☑ pDC	53 CXCR3-CCR6-: ☑ Th2-like
24 CD123-CD11c+ (DC)	54 CXCR3-CCR6+: ☑ Th17-like
25 CD11c+CD38+: ☑ mDC	55 CD3+CD4-
26 CD14-CD11c- (non-Mono)	56 CD28+CD161hi: ☑ CD4- MAIT/NKT
27 CD45+CD3+ (T cells)	57 HLA-DR-CD11c-
28 CD3+TCRγδ- (αβ T cells)	58 CD123+CD294+: ☑ Basophils
29 CD4-CD8+ (CD8 αβ T cells)	59 CD4-CD8-
30 CD8+CD161lo/-: ☑ CD8 αβ T cells	60 CD3+TCRγδ+: ☑ CD4-CD8- γδ T cells

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