# Immunophenotyping RBRCLA\_IMM\_001

## **Purpose**

This test differentiates immune cell sub-populations via flow cytometry.

**Description:** increased CD4-positive T cell number (MP:0008074), decreased CD4-positive T cell number (MP:0008075), etc..

## **Experimental Design**

- Minimum number of animals: 3M + 3F
- Age at test: Week 57
- Sex: We would expect the results of this test to show sexual dimorphism

## **Equipment**

## Equipment

- Scissors and forceps for biopsy
- Precision balance
- Calibrated single and multichannel pipettes
- Plate shaker
- Refrigerated centrifuge
- Flow Cytometer (capable of distinguishing a minimum of 8 colours per well)
- Tissue dissociator:
  - GentleMACS tissue dissociator OR
  - Equipment for manual dissociation
- Cell counter equipment:
  - Orflo Moxi-Z Cell counter OR
  - Coulter Vicell XR OR Life Technologies Attune® Flow Cytometer

#### **Supplies**

- 96-well V-bottomed plates (Falcon #353263)
- Petri dishes
- Dispensing troughs
- Extra long 10 µl pipette tips for antibody solutions
- (if using GentleMACS for dissociation) C Tubes. It is acceptable to re-use these once.
- 50ml Falcon tubes
- Cell strainers e.g. 70m cell strainers that fit 50ml Falcon tubes (BD Falcon, #352350) OR Nytex
- Cell counter recipients (i.e., slides/cassettes/etc. for cell counter)
- (if sample processing delayed) RPMI 1640

- (if sample processing on same day) HBSS (with phenol red)
- CS (calf serum)
- PBS with Mg2+, with Ca2+ (for enzyme buffer used for DNAse and Collagenase D digestions)
- PBS without Mg2+, without Ca2+ (for <u>FACS buffer</u> to be used in all steps subsequent to enzymatic digest)
- EDTA (final concentration 2mM)
- Digestion enzyme (Collagenase D from Roche, #11088858001) stock solution in enzyme buffer (see below), aliquoted and stored at -20°C
- DNAse I stock solution (Sigma, #DN25) in enzyme buffer (see below), aliquoted and stored at -20°C
- RBC lysis buffer (eBioscience #00-4300-54 or BD Biosciences #555899, both 10X from manufacturer)
- **HEPES** (pH 7.2)

## **Procedure**

This protocol requires several steps in the collection, preparation and analysis of the samples. Each one is detailed separately below.

## Reagent preparation

Note that two different PBS solutions are required for the protocol below, one with Ca2+ and with Mg2+, another without Ca2+ and without Mg2+.

- Collection buffer:
  - (*if spleens are to be processed on the same day*) HBSS with Ca2+/Mg2+ and phenol red (Life Technologies 14170161; check if it has phenol red) *OR*
  - (*if analysis will be delayed*) RPMI medium with 2% CS added.
- **FACS buffer** (for all steps subsequent to enzymatic digest; stable for up to 1 month in the fridge):
  - PBS 1X <u>without</u> Ca2+/Mg2+ *OR*
  - HBSS 1X <u>without</u> Ca2+/Mg2+
  - EDTA 2mM
  - 2% CS (v/v)
  - 10mM HEPES
- **Enzyme buffer** (for DNAse and Collagenase D digestions; Stable for up to 1 month in the fridge):
  - PBS with Ca2+ and Mg2+ OR
  - HBSS 1X with Ca2+/Mg2+
  - 2% CS (v/v):
  - 10mM HEPES
- RBC Lysis buffer: Prepare a 1X solution in ddH<sub>2</sub>0 from lysis buffer.
- **Stopping buffer** (require 300 µl per sample):
  - 1x PBS without Ca2+ and without Mg2+ or HBSS
  - 0.1 M EDTA (37.5 g/L)
- Antibody cocktails for Panels 1 & 2
  - Protect antibodies and prepared cocktails from direct light.
  - Mastermix concentration, storage temperature and stability to be determined after panels 1 and 2 have been finalised and tested.

- Each sample will require 50 μl (or up to 100 μl) of diluted 1X antibody cocktail.
- Antibody cocktails should be gently but thoroughly mixed or quickly vortexed to ensure homogeneity of the solutions.
- In order to eliminate aggregated antibodies from your mix, centrifuge each antibody cocktail for 8 min at 20,000xg and 8°C prior to staining cells.

## Read buffer / dead cell exclusion dye

- SytoxBlue at 1:10000 concentration in FACS buffer OR
- SytoxGreen at 1:20000 concentration in FACS buffer
- Zombie Near Infra-Red live dead from Biolegend at 1:2000 concentration
- Require 200 I per well (i.e. 400 I for each spleen).
- Enzyme cocktail (working solution): 3 ml per each spleen, containing final concentrations of:
  - DNAse I: 30 g
  - Collagenase D: 600 Mandl Units

**NOTE**: To top up to the 3ml use enzyme buffer; any intermediate dilutions of the enzyme stock solutions should be prepared with <u>enzyme buffer</u>.

## Other preparations on the day

- Bring RBC lysis buffer and stop solution to room temperature.
- Prepare wet ice box, label tubes, etc.

## Note all centrifuge steps are: 5 min, 400 x g at 8°C

## Spleen collection

- Collect the spleen from euthanized mice.
- Remove all fat from the spleen and weigh the organ on a petri dish (do not hydrate the organ before weighing it as this would lead to substantial errors in measurement).
- Place the spleen in a 1.5ml eppendorf tube with 1 mL of sample collection buffer on ice.
   Use:
  - (if spleens are to be processed on the same day) HBSS without calcium, without magnesium but with phenol red OR
  - (if analysis will be delayed) RPMI with 2% CS buffer.

## Spleen dissociation / digests

## If using a GentleMacs tissue dissociator:

- Add the spleen to a GentleMACS C tube containing 3 ml of 1X enzyme cocktail.
- Clip the tube on GentleMACS dissociator and run programme spleen\_2.
- Incubate cell suspension for 30 minutes with gentle mixing at least every 5 minutes. Register incubation temperature.
- Run programme spleen 3.
- Add 300 L of stopping buffer and mix by inversion to block enzymatic digestion and dissociate T cell-dendritic cell interactions.
- Filter cell suspension:
  - through 70 m Nylon mesh filter into a 50 mL Falcon tube OR

- directly from C-tubes pour splenocyte suspension through 30 mm CellTrics Partec filters (#04-0042-2316) into 15 ml tubes.
- (optional) Wash the GentleMACS C tube with 5ml <u>FACS buffer</u>, filter and pool with flow-through from previous step.
- Centrifuge for 5 minutes, 400 x g at 8°C and discard supernatant.
- Resuspend total splenocytes in 1 mL cold <u>FACS buffer</u> and keep on ice (this step is not required if counting is performed on the attune).

## **OR**, if performing manual digests:

- Place weighed spleen in 12x75mm tube containing 1ml of collagenase solution in 1X HBSS with Ca2+ and Mg2+ (0.17-0.2 Wünsch unit/ml)
- Mince into fine pieces using small scissors, place on ice until all samples are minced.
- Add 2ml collagenase (0.17-0.2 Wünsch unit/ml) to each tube and place in a 37°C water bath for 30 minutes.
- Tricturate (pipetting vigorously up and down using a 1 mL pipetman) the mixture to break up clumps.
- Spin at 500 x g in a swing bucket rotor for 5 min at 10°C. Decant the supernatant, rack the tubes or vortex to resuspend the pellet. Add 2ml <u>FACS buffer</u>, mix well by vortexing, take 10 µl for the counting step.
- Dilutions for counting: 2 serial 1:10 dilutions (10μl cells + 90μl <u>FACS buffer</u>, then 10μl of the 1:10 dilution + 90μl buffer.)
- Spin for 5min, 500 x g at 10°C, decant supernatant, blot the top of the tube, resuspend pellet at 1x10<sup>8</sup> cells/ml.

## Cell counting

- Perform a cell count on an aliquot of the re-suspended cells (adjust concentration according to the cell counter method used).
- Note down the cell count, correct for dilution and calculate the concentration in cells per μl.
- Cell count:
  - <u>If performed before RBC lysis</u>, pipette the volume containing approximately 4 million cells/well to a 96 well plate in horizontal fashion starting from A1 onwards for panel 1 staining.
  - <u>If performed after RBC lysis</u>, pipette the volume containing approximately 1-2 million cells/well to a 96 well plate in horizontal fashion starting from A1 onwards for panel 1 staining.
- Do the same for panel 2 staining in separate wells leaving a few empty rows between the panels to avoid cross contamination.
- Top up to final volume of 100 ml using <u>FACS buffer</u>, centrifuge, discard supernatant and keep plate on wet ice.

## Red blood cell lysis, blocking & staining

- Remove plate from ice and add 30 to 100 ml of 1X RBC lysis buffer (at room temperature) to each cell pellet from the previous step.
- Pipette up and down 2-3 times to break up the pellet and ensure complete lysis. Alternatively, vortex the edges of the plates, then pipet quickly once to ensure resuspension is ideal for optimal lysis.

• Incubate for 1 minute at room temperature and then return to ice and add 100 to 200 ml of <u>FACS buffer</u> (to stop lysis) to each well.

**Note**: Following RBC lysis, every centrifugation step can be performed at 2000rpm for 1 minute in a 96 well plate, which significantly speeds up the protocol. Do take care to resuspend the cells very well to prevent HTS clumping.

- Centrifuge, discard supernatant and resuspend in 200 ml <u>FACS buffer</u> (this step is not required if lysis was performed in 30 μl, since there will be enough volume left in the well for a bigger wash of 200 μl; saves time on a spin).
- Again centrifuge and discard supernatant and resuspend in 50 ml of 1:100 Fc block and incubate on ice for 10 min. Top up to 200 ml using <u>FACS buffer</u> after incubation.
- Take antibody (AB) cocktails from the fridge. In order to eliminate aggregated ABs from your mix before use, centrifuge each AB cocktail for 8 min at 20,000 x g and 4°C.
- Centrifuge plate, discard supernatant and resuspend in 50 to 100 ml 1X AB mix in appropriate wells for individual panels followed by incubation on ice and in the dark for 20 min.
- If using Sytox Blue/Sytox Green as live/dead discriminator:
  - Top up to 200 ml with <u>FACS buffer</u> after incubation. Centrifuge, discard supernatant and resuspend in 200 ml <u>FACS buffer</u>.
  - When ready to read plate, centrifuge again and discard supernatant. Resuspend the pellet in 200 ml of read buffer (Sytox Blue diluted 1:10000 in <u>FACS buffer</u>; Sytox Green diluted 1:20000 in <u>FACS buffer</u>).
- If using Zombie NIR dye as live/dead discriminator:
  - Add 200 ml of PBS (RT) to all samples
  - Spin at 2000 rpm for 1 minute 8°C
  - Add 100 ml/well of Zombie Near-IR Live/Dead dye (1/2000) made up in PBS incubate at room temperature for 10 mins, add 200 ml <u>FACS buffer</u>.

## **General Recommendations for Setting up Cytometer**

Set up the analyser to aim acquire 300,000 viable events (live cells) for each of Panels 1 and 2. 500,000 are recommended for panel 2 in order to increase robustness of myeloid population of low frequencies (macrophages, DCs).

## **Gating Panel 1**

Parameters	Gating steps			
Panel A live leukocyte count				Τ
T cells (panel A)	number of live leukocytes	CD5+	CD161-	$\top$
NKT cells (panel A)	number of live leukocytes	CD5+	CD161+	
NK cells (panel A)	number of live leukocytes	CD5-	CD161+	
Others	number of live leukocytes	CD5-	CD161-	П
CD4 T cells	number of live leukocytes	CD5+	CD161-	CD
CD8 T cells	number of live leukocytes	CD5+	CD161-	CD
DN T cells	number of live leukocytes	CD5+	CD161-	CD
DP T cells	number of live leukocytes	CD5+	CD161-	CD
CD4 NKT cells	number of live leukocytes	CD5+	CD161+	CD
CD8 NKT cells	number of live leukocytes	CD5+	CD161+	CD
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DN NKT cells	number of live leukocytes	CD5+	CD161+	CD
CD4 CD25+ T cells		number of CD5+	CD161-	CD
CD4 CD25- T cells		number of CD5+	CD161-	CD
CD8 CD25+ T cells		number of CD5+	CD161-	CD
CD8 CD25- T cells		number of CD5+	CD161-	CD
DN CD25+ T cells		number of CD5+	CD161-	CD
DN CD25- T cells		number of CD5+	CD161-	CD
CD4 CD25+ NKT cells		number of CD5+	CD161+	CD
CD4 CD25- NKT cells		number of CD5+	CD161+	CD
CD8 CD25+ NKT cells		number of CD5+	CD161+	CD
CD8 CD25- NKT cells		number of CD5+	CD161+	CD
DN CD25+ NKT cells		number of CD5+	CD161+	CD
DN CD25- NKT cells		number of CD5+	CD161+	CD
CD4 CD44+CD62L- T cells		number of CD5+	CD161-	CD
CD4 CD44+CD62L+ T cells		number of CD5+	CD161-	CD
CD4 CD44-CD62L+ T cells		number of CD5+	CD161-	CD
CD4 CD44-CD62L- T cells		number of CD5+	CD161-	CD
CD8 CD44+CD62L- T cells		number of CD5+	CD161-	CD
CD8 CD44+CD62L+ T cells		number of CD5+	CD161-	CD
CD8 CD44-CD62L+ T cells		number of CD5+	CD161-	CD
CD8 CD44-CD62L- T cells		number of CD5+	CD161-	CD
DN CD44+CD62L- T cells		number of CD5+	CD161-	CD
DN CD44+CD62L+ T cells		number of CD5+	CD161-	CD
DN CD44-CD62L+ T cells		number of CD5+	CD161-	CD
DN CD44-CD62L- T cells		number of CD5+	CD161-	CD
CD4 CD44+CD62L- NKT cells		number of CD5+	CD161+	CD
CD4 CD44+CD62L+ NKT cells		number of CD5+	CD161+	CD
CD4 CD44-CD62L+ NKT cells		number of CD5+	CD161+	CD
CD8 CD44+CD62L- NKT cells		number of CD5+	CD161+	CD
CD8 CD44+CD62L+ NKT cells		number of CD5+	CD161+	CD
CD8 CD44-CD62L+ NKT cells		number of CD5+	CD161+	CD
DN CD44+CD62L- NKT cells		number of CD5+	CD161+	CD
DN CD44+CD62L+ NKT cells		number of CD5+	CD161+	CD
DN CD44-CD62L+ NKT cells		number of CD5+	CD161+	CD

## Gating Panel B

	1				
Parameters	Gating steps				
Panel B live leukocyte count					
Neutrophils	Live	CD11b+	Ly6G+		
Monocytes	Not Granulocytes	CD11b+	Ly6C High		
Eosinophils	Not Monocytes	CD11b+	SSC-H High		
NK Cells (panel B)	Not Eosinophils	CD161+	CD19-	CD5-	
NK Subsets (Q1)	Not Eosinophils	CD161+	CD19-	CD5-	С
NK Subsets (Q2)	Not Eosinophils	CD161+	CD19-	CD5-	С
NK Subsets (Q3)	Not Eosinophils	CD161+	CD19-	CD5-	С

NK Subsets (Q4)	Not Eosinophils	CD161+	CD19-	CD5-	С
NKT Cells (panel B)	Not Eosinophils	CD161+	CD19-	CD5+	
NKT Subsets (Q1)	Not Eosinophils	CD161+	CD19-	CD5+	С
NKT Subsets (Q3)	Not Eosinophils	CD161+	CD19-	CD5+	С
T Cells (panel B)	Not Eosinophils	CD161-	CD5+		
T Subset	Not Eosinophils	CD161-	CD5+	Ly6C+	
B Cells	Not Eosinophils	MHCII+	CD19+		
B1B Cells	Not Eosinophils	MHCII+	CD19+	CD5+	
B2B Cells	Not Eosinophils	MHCII+	CD19+	CD5-	
Follicular B Cells	Not Eosinophils	MHCII+	CD19+	CD5-	С
pre-B Cells	Not Eosinophils	MHCII+	CD19+	CD5-	С
MZB	Not Eosinophils	MHCII+	CD19+	CD5-	С
cDCs	Not Eosinophils	MHCII+	CD19-	CD11c+	$\mathbf{L}$
cDCs CD11b Type	Not Eosinophils	MHCII+	CD19-	CD11c+	С
pDCs	Not Eosinophils	Not T Cells	Ly6C+	CD317+	$\mathbf{L}$
RP Macrophage (F4/80+)	Not Eosinophils	MHCII+	F4/80+		$\mathbf{L}$
or					$\mathbf{L}$
RP Macrophage (CD19-CD11c-)	Not Eosinophils	MHCII+	CD19-	CD11c-	$\prod$

## **Parameters and Metadata**

# Spleen weight RBRCLA\_IMM\_001\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: true Is Annotated: false

**Unit Measured:** g

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# Percentage of live gated events in Panel A RBRCLA\_IMM\_002\_001

| v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

Unit Measured: %		
T cells (panel A) RB simpleParameter	RCLA_IMM_003_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NKT cells (panel A) simpleParameter	RBRCLA_IMM_004_001   v	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NK cells (panel A) simpleParameter	RBRCLA_IMM_005_001   v1	.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
Others PRDCLA IMM O	06 001 1 1 0	

# **Others** RBRCLA\_IMM\_006\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

CD4 T cells RBRCLA_simpleParameter	IMM_007_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 T cells RBRCLA_simpleParameter	IMM_008_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
DN T cells RBRCLA_IM simpleParameter	1M_009_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: false
DP T cells RBRCLA_IM simpleParameter	IM_010_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: false

# CD4 NKT cells RBRCLA\_IMM\_011\_001 | v1.0

simpleParameter

Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 NKT cells RBR simpleParameter	CLA_IMM_012_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: false
DN NKT cells RBRC simpleParameter	LA_IMM_013_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD4 CD25+ T cells	RBRCLA_IMM_014_001   v	1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true

**CD4 CD25- T cells** RBRCLA\_IMM\_015\_001 | v1.0

Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 CD25+ T cells simpleParameter	RBRCLA_IMM_016_001   v	1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 CD25- T cells simpleParameter	RBRCLA_IMM_017_001   v1	.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
DN CD25+ T cells F simpleParameter	RBRCLA_IMM_018_001   v1.0	0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

## **DN CD25- T cells** RBRCLA\_IMM\_019\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

CD4 CD25+ NKT ce	EIIS RBRCLA_IMM_020_00	1   v1.0
	Req. Upload: false	
CD4 CD25- NKT ce	<b>IIS</b> RBRCLA_IMM_021_001	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 CD25+ NKT ce	<b>ells</b> RBRCLA_IMM_022_00 <sup>2</sup>	1   v1.0
	Req. Upload: false	Is Annotated: false
CD8 CD25- NKT ce	<b>IIS</b> RBRCLA_IMM_023_001	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

# DN CD25+ NKT cells RBRCLA\_IMM\_024\_001 | v1.0

simpleParameter

Req. Analysis: false		Is Annotated: true
DN CD25- NKT cell simpleParameter	S RBRCLA_IMM_025_001	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
	quired events in Pa	nel A RBRCLA_IMM_026_
001   v1.0 simpleParameter		
Req. Analysis: false	Req. Upload: false	Is Annotated: false
Total number of ac	quired events in Pa	nel B rbrcla_imm_027_
simpleParameter		
Req. Analysis: false	Req. Upload: false	Is Annotated: false

# CD4 CD44+CD62L- T cells RBRCLA\_IMM\_028\_001 | v1.0

simpleParameter

Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD4 CD44+CD62L simpleParameter	+ T cells rbrcla_imm_	_029_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD4 CD44-CD62LosimpleParameter	+ T cells rbrcla_imm_	030_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD4 CD44-CD62L-simpleParameter	- T cells rbrcla_imm_0	031_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

CD8 CD44+CD62L- T cells RBRCLA\_IMM\_032\_001 | v1.0

Req. Analysis: false	Req. Upload: false	
CD8 CD44+CD62L+simpleParameter	- T cells rbrcla_imm_0	033_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 CD44-CD62L+ simpleParameter	T cells rbrcla_imm_0	34_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
CD8 CD44-CD62L-simpleParameter	T cells rbrcla_imm_03	35_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

# DN CD44+CD62L- T cells RBRCLA\_IMM\_036\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

DN CD44+CD62L+ simpleParameter	T cells rbrcla_imm_03	37_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false
DN CD44-CD62L+ simpleParameter	T cells rbrcla_imm_03	8_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false
DN CD44-CD62L- simpleParameter	<b>Cells</b> RBRCLA_IMM_039	9_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false
CD4 CD44+CD62L simpleParameter	- NKT cells rbrcla_in	MM_040_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true

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## CD4 CD44+CD62L+ NKT cells RBRCLA\_IMM\_041\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

## CD4 CD44-CD62L+ NKT cells RBRCLA\_IMM\_042\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

## CD8 CD44+CD62L- NKT cells RBRCLA IMM 043 001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

## CD8 CD44+CD62L+ NKT cells RBRCLA\_IMM\_044\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

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CD8 CD44-CD62L+ NKT cells RBRCLA\_IMM\_045\_001 | v1.0

Req. Analysis: false	Req. Upload: false	Is Annotated: false
DN CD44 CDC2L		
simpleParameter	NKT cells rbrcla_imm	1_046_001   v1.0
	Req. Upload: false	
DN CD44+CD62L+ simpleParameter	NKT cells rbrcla_imi	M_047_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
DN CD44-CD62L+ I simpleParameter	NKT cells rbrcla_imm	1_048_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

# Percentage of live gated events in Panel B RBRCLA\_IMM\_049\_001 | v1.0

simpleParameter

Req. Analysis: false	Req. Upload: false	Is Annotated: false
Unit Measured: %		
Neutrophils RBRCLA_simpleParameter	_IMM_050_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
Monocytes RBRCLA_I simpleParameter	MM_051_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
Eosinophils RBRCLA simpleParameter	_IMM_052_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true

# NK Cells (panel B) RBRCLA\_IMM\_053\_001 | v1.0

simpleParameter

Req. Analysis: false	Req. Upload: false	Is Annotated: true
NK Subsets (Q1) RE simpleParameter	BRCLA_IMM_054_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NK Subsets (Q2) RE simpleParameter	BRCLA_IMM_055_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NK Subsets (Q3) REsimpleParameter	BRCLA_IMM_056_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NK Subsets (Q4) RE simpleParameter	BRCLA_IMM_057_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true

NKT Cells (panel EsimpleParameter	<b>3)</b> RBRCLA_IMM_058_001	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NKT Subsets (Q1) simpleParameter	RBRCLA_IMM_059_001   v <sup>2</sup>	1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
NKT Subsets (Q3) simpleParameter	RBRCLA_IMM_060_001   v <sup>2</sup>	1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
T Cells (panel B) R simpleParameter	BRCLA_IMM_061_001   v1.0	)
Req. Analysis: false	Req. Upload: false	Is Annotated: true

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# T Subset RBRCLA\_IMM\_062\_001 | v1.0 simpleParameter Req. Analysis: false Req. Upload: false Is Annotated: true B Cells RBRCLA\_IMM\_063\_001 | v1.0 simpleParameter Req. Analysis: false Req. Upload: false Is Annotated: true B1B Cells RBRCLA\_IMM\_064\_001 | v1.0 simpleParameter

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

## B2B Cells RBRCLA\_IMM\_065\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

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## Follicular B Cells RBRCLA\_IMM\_066\_001 | v1.0

Req. Analysis: false	Req. Upload: false	Is Annotated: true
Follicular B Cells (6 simpleParameter	CD21/35+) RBRCLA_IMI	M_067_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
Transitional B Cells simpleParameter	S RBRCLA_IMM_068_001	v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
Transitional B Cells simpleParameter	s (CD21/35 low) RBR	CLA_IMM_069_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
MZB RBRCLA IMM 070	001 Lv1 0	

**IVIZD** RBRCLA\_IMM\_070\_001 | V1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

MZB (CD21/35 high	<b>1)</b> RBRCLA_IMM_071_001	v1.0
	Req. Upload: false	
<b>CDCs</b> RBRCLA_IMM_072 simpleParameter	2_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true
cDCs CD11b Type simpleParameter	RBRCLA_IMM_073_001   v1	1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: true
pDCs RBRCLA_IMM_076 simpleParameter	4_001   v1.0	
Req. Analysis: false	Req. Upload: false	Is Annotated: true

## RP Macrophage (F4/80+) RBRCLA\_IMM\_075\_001 | v1.0

simpleParameter

Req. Analysis: false	Req. Upload: false	Is Annotated: true

# RP Macrophage (CD19- CD11c-) RBRCLA\_IMM\_076\_001 | v1.0

simpleParameter

Req. Analysis: false Req. Upload: false Is Annotated: true

## Equipment name RBRCLA\_IMM\_077\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: FACS, Fortessa\_1, LSR II, Flow cytometer,

.....

## Equipment manufacturer RBRCLA\_IMM\_078\_001 | v1.0

procedureMetadata

Req. Analysis: true Req. Upload: true Is Annotated: false

Options: BD Biosciences, Beckman Coulter,

## Equipment model RBRCLA\_IMM\_079\_001 | v1.0

procedureMetadata

Req. Analysis: true Req. Upload: true Is Annotated: false

Options: BD LSRFortessa Cell Analyzer, H47100123, Gallios, FACSAria III, BD LSR-II, CANTO-II,

CS&T Bead Iot RBRCLA\_IMM\_080\_001 | v1.0 procedureMetadata

**Is Annotated:** false

## Anesthesia RBRCLA\_IMM\_081\_001 | v1.0

Reg. Analysis: false Reg. Upload: true

procedureMetadata

Reg. Analysis: true Reg. Upload: true Is Annotated: false

**Options:** Injection narcosis with Sodium Pentobarbital (Somnopentyl), none, Injection narcosis with Ketamine (100mg/kg)/Xylazine (10mg/kg), Injection narcosis with Tribromoethanol (Avertin), Isoflurane, Injection narcosis with Medetomidine/Midazolam/Butorphanol,

## Cell digestion RBRCLA\_IMM\_082\_001 | v1.0

procedureMetadata

Req. Analysis: true Req. Upload: true Is Annotated: false **Options:** GentleMACS, manual, Cell digestion agent RBRCLA\_IMM\_083\_001 | v1.0 procedureMetadata Req. Analysis: false Req. Upload: true Is Annotated: false Options: Collagenase D, Collagenase II, Cell digestion agent manufacturer RBRCLA\_IMM\_084\_001 | v1.0 procedureMetadata Req. Analysis: false Req. Upload: true Is Annotated: false Options: Roche, Worthington, Gibco,

# Cell digestion agent catalog number RBRCLA\_IMM\_085\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: #11088858001, CLS2LS004176, 17101-015, Cell counting RBRCLA\_IMM\_086\_001 | v1.0 procedureMetadata Req. Analysis: false Req. Upload: true Is Annotated: false Options: pre-lysis, post-lysis, Cell counting equipment manufacturer RBRCLA\_IMM\_087\_001 | v1 .0 procedureMetadata Reg. Analysis: false Reg. Upload: true Is Annotated: false Options: Life Technologies, American Optical, Beckman Coulter, BD Biosciences,

## Cell counting equipment model RBRCLA\_IMM\_088\_001 | v1.0

Merck Millipore, Orflo, Nexcelom Bioscience, IntelliCyt,

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

<b>Options:</b> Countess Automated Cell Counter, Reichert Brightline, Gallios, BD LSR-II, Scepter, Attune, Moxi Z, 4468770, Cellometer Auto T4, iQue Screener PLUS,		
Cell counting equiposedureMetadata	oment name RBRCLA	_IMM_089_001   v1.0
Req. Analysis: false	Req. Upload: true	Is Annotated: false
Cell lysis buffer ma	anufacturer RBRCLA_I	MM_090_001   v1.0
Req. Analysis: false	Req. Upload: true	Is Annotated: false
Options: eBioscience, BD Pha	armLyse, Jax, JMC, LONZA,	
Cell lysis buffer car	talog number RBRCL	A_IMM_091_001   v1.0
Req. Analysis: false	Req. Upload: true	Is Annotated: false
<b>Options:</b> 00-4300-54, 555899	, home brew, 10-548E,	

# Date and time of sacrifice RBRCLA\_IMM\_092\_001 | v1.0

procedureMetadata

Req. Analysis: false	Req. Upload: true	Is Annotated: false
Date and time of sa procedureMetadata	ample preparation R	BRCLA_IMM_093_001   v1.0
Req. Analysis: false	Req. Upload: true	Is Annotated: false
Sample storage ter	mperature until ana	lysis (in Celsius) RBR
CLA_IMM_094_001   v1.0 procedureMetadata		
Req. Analysis: false	Req. Upload: true	Is Annotated: false
Unit Measured: C		
FCS repository referencedureMetadata	erence (URL/ID) RBR	CLA_IMM_095_001   v1.0
Req. Analysis: false	Req. Upload: false	Is Annotated: false

## Balanced salt solution type RBRCLA\_IMM\_096\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false Options: HBSS, PBS, Balanced salt solution manufacturer RBRCLA\_IMM\_097\_001 | v1.0 procedureMetadata Req. Analysis: false Req. Upload: true Is Annotated: false Options: Sigma, Life Technologies, Wisent, Wako, Gibco, Biochrom, Balanced salt solution catalog number RBRCLA\_IMM\_098\_001 | v1 .0 procedureMetadata Req. Analysis: false Req. Upload: true Is Annotated: false Options: D1408, H6136-1L, 041-20211, 14190-144, L 182-10, HBSS 1X 14170-088, 14175-095.

## RPMI manufacturer RBRCLA IMM 099 001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: Sigma, Life Technologies, Jax, Wako, Gibco, none used,

## RPMI catalog number RBRCLA\_IMM\_100\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: R8758, 11875-101, home brew, 189-02145, 31800-022, none used, 11875-093,

## DNAse I manufacturer RBRCLA\_IMM\_101\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: Sigma,

## DNAse I catalog number RBRCLA\_IMM\_102\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

## Dead cell exclusion dye RBRCLA\_IMM\_103\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: Sytox Blue, Sytox Green, Zombie NIR, DAPI, Propidium Iodide,

# Dead cell exclusion dye manufacturer RBRCLA\_IMM\_104\_001 | v1.

0

procedureMetadata

Reg. Analysis: false Reg. Upload: true Is Annotated: false

Options: Life Technologies, Biolegend, Sigma, home brew,

## Dead cell exclusion dye catalog number RBRCLA\_IMM\_105\_001 |

v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

**Options:** S34857, S-34860, 423106, D9542, S11348, home brew, R37606,

# Cell digestion temperature (in Celsius) RBRCLA\_IMM\_106\_001 | v1

.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: 37, RT,

## Panel A FCS file(s) RBRCLA\_IMM\_107\_001 | v1.0

seriesMediaParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

**Increments:** Minimum 1

## Panel B FCS file(s) RBRCLA\_IMM\_108\_001 | v1.0

seriesMediaParameter

Req. Analysis: false Req. Upload: false Is Annotated: false

Increments: Minimum 1

## Automated analysis RBRCLA\_IMM\_109\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: true Is Annotated: false

Options: Yes, No,

Collection buffer manufacturer RBRCLA\_IMM\_110\_001 | v1.0 procedureMetadata

Req. Analysis: false Req. Upload: false Is Annotated: false

Options: Life Technologies,

Collection buffer catalog number number RBRCLA\_IMM\_111\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: false Is Annotated: false

**Options:** 24020,

.....

Req. Analysis: false Req. Upload: false Is Annotated: false Options: Life Technologies, FACS buffer catalog number RBRCLA\_IMM\_113\_001 | v1.0 procedureMetadata Reg. Analysis: false Reg. Upload: false Is Annotated: false **Options:** 14175, Enzyme buffer manufacturer RBRCLA\_IMM\_114\_001 | v1.0 procedureMetadata Reg. Analysis: false Reg. Upload: false Is Annotated: false Options: Life Technologies,

# Enzyme buffer catalog number RBRCLA\_IMM\_115\_001 | v1.0

procedureMetadata

Req. Analysis: false Req. Upload: false Is Annotated: false

**Options:** 14025,