

FACs Analysis ESLIM_018_001

Purpose

The characterisation of of main leukocyte subsets in peripheral blood cells from mice by flow cytometry.

Reliable values of frequencies of leukocyte clusters are very much dependent on the appropriate preparation, acquisition and gating of leukocytes. This method involves ammonium chloride erythrocyte lyses, thus preventing interference by larger amounts of erythrocytes. For all stainings, a monoclonal antibody to the cell surface glycoprotein CD45 is used to be able to create a CD45+ gate, allowing discrimination of leukocytes from debris, erythrocytes and thrombocytes. Furthermore, staining with propidium iodide (PI) gates out dead cells. Samples are acquired until a number of 30000 living CD45+ cells is reached for each sample (ref Adler and Busch 2007, submitted). Data are compensated analysed and using FlowJo software.

This protocol is part of the EMPReSSslim pipeline and is carried out on a cohort of 10 males and 10 females (age matched) at a time.

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1.0 Purpose:

- 1.1 The characterisation of of main leukocyte subsets in peripheral blood cells from mice by flow cytometry.

Reliable values of frequencies of leukocyte clusters are very much dependent on the appropriate preparation, acquisition and gating of leukocytes. This method involves ammonium chloride erythrocyte lyses, thus preventing interference by larger amounts of erythrocytes. For all stainings, a monoclonal antibody to the cell surface glycoprotein CD45 is used to be able to create a CD45+ gate, allowing discrimination of leukocytes from debris, erythrocytes and thrombocytes. Furthermore, staining with propidium iodide (PI) gates out dead cells. Samples are acquired until a number of 30000 living CD45+ cells is reached for each sample (ref Adler and Busch 2007, submitted). Data are compensated analysed and using FlowJo software.

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2.0 Scope:

- 2.1 Individuals who have been trained, and are competent in performing the procedures described herein must follow this procedure.
- 2.2 Any queries, comments or suggestions, either relating to this SOP in general or to a specific problem encountered during a procedure, should be addressed to the Project Leader.
- 2.3 Any deviances from this protocol must be reported to the Project Leader.

3.0 Safety Requirements:

- 3.1 General laboratory procedures should be followed, which include: no eating, no chewing gum, no drinking, and no applying of cosmetics in the work area. Laboratory coats must be worn at all times in the work area, unless the protocol

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4.0 Associated Documents:

- 4.1 ESLIM_028_001: Titration of antibody solutions for FACS analysis
- 4.2 ESLIM_018_001_Appendix_1: FACS analysis – Appendix 1: Definition of main lineages
- 4.3 ESLIM_024_001: Blood collection by retro-orbital puncture

5.0 Notes:

- 5.1 The selection of antibody-conjugate and the design of the staining panel has to be adapted to each laboratory, depending on the flow cytometer in the facility. It is possible to use a broader antibody-panel, as long as the gating strategy (appendix) is respected.

6.0 Quality Control:

- 6.1 Before the acquisition of the samples, run a sample of calibration beads, sphero Ultra Rainbow, 3 μm ϕ (spherotech Cat. URFP-30-2), in order to test the regular running of the flow cytometer (LSR II).

7.0 Equipment:

- 7.1 Microhematocrit tube
- 7.2 Centrifuge (enabling centrifugation of tubes/Microtiter plate, and temperature control)
- 7.3 7 colour Flow cytometer (LSR II)
Example of configuration in table 1:



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Table 1:

Laser	P	Longpass Dichroic Mirror	Bandpass Filter	Fluochrome	blood staining 1	blood staining 2
Blue: 488nm	G	-	488/10		Side scatter	Side scatter
	F	505LP	530/30	FITC		IgD
	E	550LP	575/26	PE		NK 1.1/USA 2
	D	600LP	610/20		Propidium Iodide	Propidium Iodide
	B	685LP	695/40	PerCPCy55	CD4	Gr1
	A	755LP	780/60	PECy7		CD19
Violet: 405nm	B	-	450/50	Pacific Blue	CD3	CD11b
Red: 633nm	C	-	660/20	APC	CD25	CD5
	B	710LP	730/45	Alexa Fluor 700	CD45	CD45
	A	755LP	780/60	APC- Alexa750	CD8a	

7.4 FlowJo software (for data analysis)

8.0 Supplies:

8.1 Eppendorf tubes



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- 8.2 Microtiter plate, round bottom
- 8.3 Heparin: Liquemin® N 25000 (Roche)
- 8.4 NH₄Cl-Tris (room temperature)
- 8.5 FACS buffer
1000ml PBS, 5g BSA, 660µl NaN₃ 30%, pH 7.45, filtered
- 8.6 Lysis buffer:
45ml sol.A + 5ml sol.B. –
sol.A: 1000ml aqua dest., 9.1 g (≈0.17M) NH₄Cl, filtered
sol.B: 1000ml aqua dest., 28.8g Tris-HCl, pH 7.5, filtered.
- 8.7 Propidium Iodide (PI) solution:
1:500 of stock solution
2mg/ml Propidium Iodide in PBS (stored at -20°C)
- 8.8 Fc-block-solution:
1:500 of stock solution (2mg/ml) in FACS buffer
- 8.10 Antibodies:
Prepare sufficient amount of antibody-mix within FACS buffer, having calculated the amount of antibodies to be used, based on titration (see EMPReSS SOP 6_001 Revision 0: Titration of antibody solutions for FACS analysis). Tables 2 and 3 describe the staining panels for EMPReSSslim pipeline.



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Table 2: Staining panel 1-mix:

Name	Stock	Ref
PerCpCy55-conjugated Rat anti-mouse CD4 (RM4-5)	1:1000 of 0.2 mg/ml	BD 5500954
PB-conjugated Rat anti mouse CD3 (17A2)	1:100 of 0.2 mg/ml	eBioscience 57-0032-82
APC-conjugated Rat anti-mouse CD25 (PC61)	1:100 of 0.2 mg/ml	BD 557192
APC-Alexa750-conjugated Rat anti-mouse CD8a (53-6.7)	1:400 of 0.1 mg/ml	Caltag MCD0827
Alexa Fluor 700-conjugated Rat anti-mouse CD45 (30F11)	1:2000 of 0.5 mg/ml	BioLegend 103128

Table 3: Staining panel 2-mix

Name	Stock	Ref
FITC-conjugated Rat anti-mouse IgD (11-26c.2a)	1:1000 of 0.5 mg/ml	BD 553439
PE-conjugated Rat anti-mouse NK-T/NK (U5A2-13)	1:200 of 0.2 mg/ml	BD 550082
PE-conjugated Rat anti-mouse NK 1.1 (PK 136) (Note: to be used only for C57BL/6 mice)	1:200 of 0.2 mg/ml	BD 553165
PerCpCy55-conjugated Rat anti-mouse Gr1 (RB6-8C5)	1:800 of 0.2 mg/ml	BD 552093
PECy7-conjugated Rat anti mouse CD19 (ID3)	1:1000 of 0.2 mg/ml	BD 552854
PB-conjugated Rat anti mouse CD11b (M1/70)	1:800 of 0.1 mg/ml	Caltag RM2828
APC-conjugated Rat anti-CD5 (53-7.3)	1:2000 of 1mg/ml	BD 550035
Alexa Fluor 700-conjugated	1:2000 of 0.5 mg/ml	BioLegend 103128

Rat anti-mouse (30F11)	CD45		
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8.13 Single Colour Control:

Rest of blood from all mice is pooled and used for single colour controls. Single colour controls (SCC) contain a single antibody and propidium iodide (PI), and are prepared simultaneously. Furthermore, one unstained sample (also containing PI) is prepared. In table 4 is the antibody used only for single colour controls in order to compensate PI-channel):

Table 4: PI channel compensation antibody

Name	Stock	Ref
PE-Alexa Fluor 610-conjugated Rat anti-mouse CD4 (RM4-5)	1:100 of 0.1 mg/ml	Caltag MCD0422

9.0 Procedure:

This protocol is carried out on a cohort of 20 mice (10 males and 10 females). It uses 7 colour flow cytometry with a 2 panel staining system.

- 9.1 Put 10 μ l of Heparin into an Eppendorf tube.
- 9.2 Collect 300 μ l of blood from the tail vein of the mouse (see EMPReSS SOP 3_011 Revision 0: Blood collection by tail venipuncture) and mix gently with the heparin. Repeat for all the mice in the cohort.
- 9.3 Centrifuge the samples (530RCF, 3 min, 10°C) and take 50-100 μ l of the plasma from each sample for further analysis.
- 9.4 Resuspend each cell pellet in 500 μ l NH₄Cl-Tris (room temperature) and filter the solutions through a nylon net into separate 1.5ml Eppendorf tubes.
- 9.5 Shake the tubes for 10 minutes at room temperature
- 9.6 For each mouse sample, pipette 200 μ l of the cell suspension into 2 wells of a U-bottomed 96 well plate.

bottomed 96-well plate.



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- 9.7 Pool the remaining cell suspensions and pipette 100 μ l into 18 wells for single colour controls (SCC) and 1 well for an unstained control.
- 9.8 Centrifuge the plate (530RCF, 3 min, 10°C). Discard the supernatants.
- 9.9 Resuspend the cells in 200 μ l NH₄Cl-Tris (RT) per well and shake the plate for 10 minutes at room temperature.
- 9.10 Centrifuge the plate (530RCF, 3 min, 10°C). Discard the supernatant.
- 9.11 Resuspend the cells in 200 μ l FACS buffer per well.
- 9.12 Centrifuge the plate (530RCF, 3 min, 10°C). Discard the supernatant.
- 9.13 Resuspend the cells in 50 μ l Fc-block-solution per well, and incubate on ice for 20 minutes.
- 9.14 Add 150 μ l FACS buffer to each well and centrifuge the plate (530RCF, 3 min, 10°C). Discard the supernatant.
- 9.15 Resuspend the cells in 50 μ l Antibody-Mix as follows:
 - i. In the 2 wells that originally contained 200 μ l cells use the Antibody mix for staining panel 1 and staining panel 2 respectively.
 - ii. In the 18 wells which originally contained 100 μ l pooled cells, use a different single antibody for each well. This is the SCC.
Note: to compensate the PI-channel antibody conjugated to PE-Alexa Fluor 610 is used.
 - iii. The 1 remaining well is resuspended in 50 μ l FACS buffer as an unstained control.
- 9.16 Protect from the light and incubate on ice for 15 minutes.
- 9.17 Add 50 μ l PI solution per well and then incubate in the dark, on ice, for a further 5 minutes.
- 9.18 Add 100 μ l FACS buffer to each well and centrifuge the plate (530RCF, 3 min,

10°C). Discard the supernatant.



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- 9.19 Wash the cells by adding 200 μ l FACS buffer per well and centrifuge the plate (530RCF, 3 min, 10°C). Discard the supernatant. Repeat this a wash further four times.
- 9.20 Resuspend the cells in 100 μ l FACS buffer and store at 4°C until data acquisition.
- 9.21 Define a “stopping gate” of living CD45 positive cells as 30 000 living cells. Data will be acquired until this number is reached. Storage gate is defined as ‘all events’.
- 9.22 Complete interpretation of the results, including compensation, using the FlowJo software.

10.0 Supporting information

11.0 History Review:

17 Aug 08: Track changes accepted and SOP numbers updated

Parameters and Metadata

T cell CD4+ percentage ESLIM_018_001_001 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: T_cell_CD4_percentage

T cell CD8+ percentage ESLIM_018_001_002 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: T_cell_CD8_percentage

T cell CD3+ percentage ESLIM_018_001_003 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: T_cell_CD3_percentage

CD25+ percentage of CD4+ T cells ESLIM_018_001_004 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: CD25_percentage_of_CD4_T_cells

B cell CD19+ percentage ESLIM_018_001_005 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: B_cell_CD19_percentage

Mature B cells IgD+ percentage ESLIM_018_001_006 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: Mature_B_cells_IgD_percentage

Granulocyte Gr1+ percentage ESLIM_018_001_007 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: Granulocyte_Gr1_percentage

NK cell percentage ESLIM_018_001_008 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: NK_cell_percentage

Monocyte gate ESLIM_018_001_009 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: %

Description: Monocyte_gate

Number of leukocytes ESLIM_018_001_010 | v1.0

simpleParameter

Req. Analysis: false

Req. Upload: true

Is Annotated: true

Unit Measured: count

Description: Number_of_leukocytes

Equipment name ESLIM_018_001_801 | v1.0

procedureMetadata

Req. Analysis: false

Req. Upload: true

Is Annotated: false

Description: Equipment_name

Equipment manufacturer ESLIM_018_001_802 | v1.0

procedureMetadata

Req. Analysis: true

Req. Upload: true

Is Annotated: false

Description: Equipment_manufacturer

Equipment model ESLIM_018_001_803 | v1.0

procedureMetadata

Req. Analysis: true

Req. Upload: true

Is Annotated: false

Description: Equipment_model

Method of blood collection ESLIM_018_001_804 | v1.0

procedureMetadata

Req. Analysis: true

Req. Upload: true

Is Annotated: false

Description: Method_of_blood_collection

EMPreSSID for blood collection SOP ESLIM_018_001_805 | v1.0

procedureMetadata

Req. Analysis: false

Req. Upload: false

Is Annotated: false

Description: EMPReSSID_for_blood_collection_SOP

Date/time of blood collection ESLIM_018_001_806 | v1.0

procedureMetadata

Req. Analysis: false

Req. Upload: true

Is Annotated: false

Description: DateTime_of_blood_collection

Sample status ESLIM_018_001_811 | v1.0

procedureMetadata

Req. Analysis: true

Req. Upload: true

Is Annotated: false

Description: Sample_status

Options: fresh, frozen,

Anaesthesia used for blood collection ESLIM_018_001_812 | v1.0

procedureMetadata

Req. Analysis: true

Req. Upload: true

Is Annotated: false

Description: Anaesthesia_used_for_blood_collection

Date of measurement ESLIM_018_001_813 | v1.0

procedureMetadata

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

Description: Date_of_measurement
