

Identifying neutrophils in immunocompetent mice infected with Mycobacterium abscessus

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INTRODUCTION

preexisting lung conditions such as cystic fibrosis or chronic obstructive pulmonary disease. Regardless of the therapy, Mycobacterium abscessus (Mabs), a bacteria prevalent in the environment targets patients with weakened immune system or Mabs can develop high levels of resistance to antibiotics, preventing a complete clearance of the infection.

NTM infection, however, is still controversial. While neutrophils could restrict the growth of NTM, they are also associated with the aggravation of clinical symptoms, such as bronchitis to this infection. These cells play a key role as the first line of defense against pathogens in the respiratory tract and are investigations have highlighted the critical role of neutrophils in predominant in the bronchial lumen. The role of neutrophils on Macrophages are crucial reservoir of Mabs; however, recent

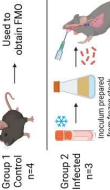
The goal of this mini-project is to identify neutrophil populations in uninfected and M abcessus-infected C57BL/6 mice by using flow-cytometry, while learning to use the Aurora Cytex flow cytometer and FlowJo software for data analysis.

OBJECTIVES

- 1. Identify neutrophil populations in the lungs of C57BL/6 mice under uninfected and Mycobacterium abscessus-infected conditions using flow cytometry.
- Compare neutrophil and macrophage profiles between infected and uninfected mice to assess immune cell recruitment and activation during M. abscessus infection.

EXPERIMENTAL STRATEGY

CDIIc NINFECTED



Mice were anesthetized and rom frozen stock

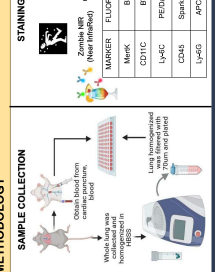
1x10^6 CFU/mouse was

Figure 1. Mice were humanely euthanized following approved ethical guidelines

ACKNOWLEDGEMENTS

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 - Anne Chenchar for providing some of the mice used on these experiments
 - 3. All members of the MIP730 class for their feedback

METHODOLOGY



STAINING PROTOCOL

Hamster IgG nonspecific binding of Abs CLONE to Fc receptors on cells. Rat FC Block. reduce FLUOROPHORE BV605

READING

Cytex aurora 4 lasers: Compensation beads Biolegend 424601 V, B,Y/G, R

ANALYSIS

FLOWJO Version10.4

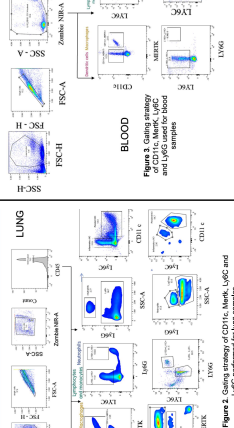
Rat IgG2b, k Rat IgG2a, k

Spark Blue 550 PE/Dazzle 594

APC/fire 750

Rat IgG2c,k

GATING STRATEGY



power of the study, affecting the accuracy of the Small sample size significantly reduced the statistical

> this study provided valuable insights for improving the experiment, including the addition of markers to enable a more in-depth characterization of the cell populations involved in the

of sample processing, including the considerable time required for data analysis. However,

The aim of this study was to understand the use of flow cytometry and the analysis of the data obtained through FlowJo, which was highly successful in highlighting the challenges

Ly6G performed for lung samples

MERTK

MERTK

CONCLUSIONS

LIMITATIONS

- > Markers selected for this experiment were insufficient characterize the intended cell populations in the samples and to optimize a flow cytometry panel, such as CD11b, CD62L for neutrophil activation and CxCR2 (neutrophil aging)
- ▶ Limited experience and exposure to this type of experiment may have influenced data quality and contributed to variability between the samples.

during infection, however, other studies have shown that M. abscessus infection leads to

an upregulation of neutrophil migration to the lungs.

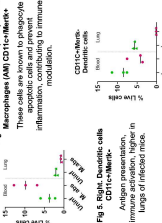
The characterization of the neutrophil population was obstructed by the extremely low presence of live cells in the sample. The first explanation could be an error on laboratory processing of the sample, a second explanation could include a depletion of neutrophils

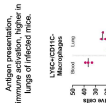
infection mechanism in both mouse lung and blood.

RESULTS

CD11C+/Mertk+ Dendritic cells

Fig 4. Left. mDendritic cells (mDC)/Alveolar







Inflammatory macrophages

High expression of Ly6C+

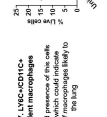
LY6C+/CD11C+

Macrophages

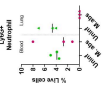
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Fig 6. Left. LY6C+/CD11C-





 Γ X ϵ G



 Γ X ϵ G

Species (ROS), NETs formation.

Neutrophils

Fig LY6G+

infected animals due to lower



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