



Sep 22, 2021

# MAIT Cell Intracellular Cytokine Staining

In 1 collection

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dx.doi.org/10.17504/protocols.io.bmg2k3ye

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ABSTRACT

This is part 3.5 of the "Study of MAIT Cell Activation in Viral Infections In Vivo" collection of protocols.

**Collection Abstract:** MAIT cells are abundant, highly evolutionarily conserved innate-like lymphocytes expressing a semi-invariant T cell receptor (TCR), which recognizes microbially derived small intermediate molecules from the riboflavin biosynthetic pathway. However, in addition to their TCR-mediated functions they can also be activated in a TCR-independent manner via cytokines including IL-12, -15, -18, and type I interferon. Emerging data suggest that they are expanded and activated by a range of viral infections, and significantly that they can contribute to a protective anti-viral response. Here we describe methods used to investigate these anti-viral functions in vivo in murine models. To overcome the technical challenge that MAIT cells are rare in specific pathogen-free laboratory mice, we describe how pulmonary MAIT cells can be expanded using intranasal bacterial infection or a combination of synthetic MAIT cell antigen and TLR agonists. We also describe protocols for adoptive transfer of MAIT cells, methods for lung homogenization for plaque assays, and surface and intracellular cytokine staining to determine MAIT cell activation.

ATTACHMENTS

Study of MAIT Cell
Activation in Viral Infections
In Vivo.pdf

DOI

dx.doi.org/10.17504/protocols.io.bmg2k3ye

**EXTERNAL LINK** 

https://link.springer.com/protocol/10.1007/978-1-0716-0207-2\_17

PROTOCOL CITATION

Timothy S C Hinks, Bonnie van Wilgenburg, Huimeng Wang, Liyen Loh, Marios Koutsakos, Katherine Kedzierska, Alexandra J. Corbett, Zhenjun Chen 2021. MAIT Cell Intracellular Cytokine Staining.

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https://dx.doi.org/10.17504/protocols.io.bmg2k3ye

COLLECTIONS (1)



Study of MAIT Cell Activation in Viral Infections In Vivo

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Citation: Timothy S C Hinks, Bonnie van Wilgenburg, Huimeng Wang, Liyen Loh, Marios Koutsakos, Katherine Kedzierska, Alexandra J. Corbett, Zhenjun Chen

#### **KEYWORDS**

Virus, MAIT cell, Flow cytometry, MR1-tetramer, Infection, Mouse

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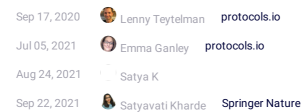
#### CREATED

Sep 17, 2020

LAST MODIFIED

Sep 22, 2021

#### OWNERSHIP HISTORY



#### PROTOCOL INTEGER ID

42234

### PARENT PROTOCOLS

Part of collection

Study of MAIT Cell Activation in Viral Infections In Vivo

## MATERIALS TEXT

For materials, please refer to the Guidelines section of the "Study of MAIT Cell Activation in Viral Infections In Vivo" collection.

## SAFETY WARNINGS

Personal protective equipment (PPE) should be worn at all times (gloves, lab coat, & eye protection) (see Notes 3 and 4).

For hazard information and safety warnings, please refer to the SDS (Safety Data Sheet).

# To analyze MAIT cell frequencies and function during viral infection.

- Prewarm collagenase media and shaking incubator to § 37 °C.
- Mice should be euthanized (e.g., using a rising concentration of CO<sub>2</sub> with a second method to confirm death).



Open the diaphragm by cutting the rib cage to expose both the heart and lungs. Gently perfuse the right ventricle with **B mL** – **10 mL ice-cold RPMI** to remove circulating blood. Perfuse using a 10-mL syringe and a 26-G needle.

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Proper perfusion will result in lung inflation and a color change to pink/ white.

- 4 Remove lungs (see Note 18) using scissors to cut through the hilum and place into a 24-well plate containing ice-cold RPMI to transfer organs to the laboratory.
- 5 Chop lungs into very small pieces (see Note 9).
- 6 Place lung tissue into a 1-mL Eppendorf tube containing 0.5 mL/lung of pre-warmed collagenase/DNase medium. This should also contain, **□**0.5 μl Brefeldin A (1:1000) (final concentration [M]3.0 μg/mL).
- 7

Incubate tubes on their sides in a shaking incubator at ₹ 37 °C , at № 100 rpm - № 180 rpm , for ७ 01:30:00 .

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After © 01:30:00 pour digested tissue through a 70- $\mu$ m cell strainer and force through into Petri dish with the plunger from a 1-mL syringe. Rinse residual cells into a total of  $\blacksquare$ 10 mL FACS wash in 10 mL falcon tubes at  $\blacksquare$ 8 Room temperature. Centrifuge at 9400 x g, Room temperature, 00:05:00.

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Resuspend in 2 mL per lung (see Note 19) of pre-warmed TAC lysis buffer at § 37 °C. Vortex well, then place in a pre-warmed water bath at § 37 °C. After © 00:05:00 neutralize by adding an equal volume of FACS buffer.

Centrifuge at \$\text{ 400 x g, Room temperature , 00:05:00}.

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Numbers of lung cells can be estimated using a hemocytometer or spectrophotometer (see Notes 20 and 21).

- 11 Transfer 100 μl containing 0.5–1 million cells to a 96-well U- or V-bottom plate format or into FACS tubes for staining, passing them through a 40-μm mesh (see Note 22).
  - (a)  $1 \times 100 \,\mu\text{L}$  into a plate for surface stain (steps 16-22).
  - (b)  $2 \times 100 \,\mu$ L (unstimulated and stimulated) to a second plate (see Note 23), and include a no Brefeldin control (steps 12, 13, and 21–30).

In vitro stimulation phase:

- 12 Keep the cells for the surface stain § On ice, while setting up PMA/Ionomycin stimulation to induce production of cytokines of interest.
  - (a) PMA final concentration: [M]20 ng/mL .
  - (b) Ionomycin [M]1 μg/mL.
  - (c) 1000x stock Brefeldin A (final concentration [M]3.0 µg/mL).

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Incubate for (303:00:00 at § 37 °C with [M]5 % CO2.

### Surface staining:

- 14 During stimulation phase perform surface staining for extracellular markers (see Note 24).
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Add 20 µl 2.4G2 (anti-CD16/32) containing

**3.2** μl MR1-6-FP tetramer (no fluorochrome conjugate) to block non-specific binding. Incubate for **00:15:00** dark, **8 Room temperature**.

17

Add surface cocktail (Table 2) using a cocktail made up in  $\Box 10~\mu I$  FACS buffer . Pipette carefully to mix. Stain for  $\odot 00:20:00-\odot 00:30:00$  at & Room temperature.

- 18 For single color controls use splenocytes or compensation beads.
- 19 🕲 🍂

Wash cells *twice* with  $\Box 2$  mL FACS buffer, centrifuging at  $\textcircled{3}400 \times g$ , 00:05:00 (or if using plate format wash three times with  $\Box 200 \mu I$  FACS buffer centrifuging for 000:02:00 at  $\textcircled{3}400 \times g$ ).

- 20 Resuspend cells in 100 μl FACS buffer (see Note 25). To enable estimation of absolute cell numbers, add a known number of calibration beads.
  - 20.1 (a) Vortex calibration beads hard. Dilute (1:10) counting beads in PBS before using. To each sample 
    25 μl of these diluted beads was added, and an additional 10 μl of beads were saved to be counted with a hemocytometer, giving a count of X in a large square, i.e., X × 10<sup>4</sup> beads/mL (which is X x 10 beads/μL, or X × 10 × 25 beads/sample). Typically add a total of 25,000 beads per sample.
  - 20.2 (b) When samples have been acquired on flow cytometer, these calibration beads can be detected using their FSC/SSC profile and the absolute number of cells of interest can be estimated using the

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following approach. Total number of MAIT cells per sample = Number of MAIT cells counted on flow cytometer × Number of beads added/Number of beads counted/proportion of total lung cell suspension actually used for staining.

# Intracellular staining:

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After 3-h stimulation, continue processing the cells for intracellular staining. Resuspend into FACS tube with +  $\blacksquare 1$  mL PBS . Centrifuge at  $\textcircled{3}400 \times g$ , 00:05:00 . (Alternatively, if in 96-well format resuspend in  $\blacksquare 100 \ \mu l$  PBS , centrifuge at  $\textcircled{3}400 \times g$ , 00:02:00 and repeat.)

22 Resuspend in 20 µl PBS with 0.4 µl Zombie Yellow for 00:15:00.

23

Add  $\blacksquare 20~\mu l$  2.4G2 (anti CD16/32) SN containing  $\blacksquare 0.2~\mu l$  unlabeled MR1-6-FP tetramer to block non-specific tetramer staining. Incubate for  $\circlearrowleft 00:15:00~\text{dark}$ , & Room temperature .

- 24 Add surface cocktail (Table 3) using a cocktail made up in  $\Box 10~\mu l$  FACS buffer . Pipette carefully to mix. Stain for  $\odot 00:20:00 \odot 00:30:00$  at & Room temperature .
- For single color controls use splenocytes or compensation beads. These may be available from being made up earlier in the protocol.

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Wash cells *once* with □1 mL FACS buffer, centrifuging at ⊕400 x g, 00:05:00 (or if using plate format wash *twice* with □200 μI FACS buffer centrifuging for ⊕00:02:00 at ⊕400 x g).

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Resuspend in  $\Box 200~\mu l$  commercially available Fixation/Permeabilizationsolution and incubate for  $\odot 00:30:00$  & On ice .

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Wash with **□2** mL Perm Wash (diluted 1:9 with FACS buffer). Centrifuge at **⊕400** x g, 00:05:00 (or if using plate format wash *twice* with **□200** µl Perm Wash centrifuging for **⊕0:02:00** at **⊕400** x g).

29

Resuspend in 50 µl Perm Wash containing intracellular cocktail (Table 4) and pipette carefully to mix. Incubate

for **© 00:45:00** or leave to stain **© Overnight** .

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Analyze cells on flow cytometer.