

JAN 31, 2023

dx.doi.org/10.17504/protocol s.io.4r3l276d3g1y/v1

Protocol Citation: Elizabeth

https://dx.doi.org/10.17504/p rotocols.io.4r3l276d3g1y/v1

Neumann, Jamie Allen, Jeff

Spraggins 2023. Fixation

Protocol for Fresh Frozen Tissue Samples (post-MALDI).

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# ( Fixation Protocol for Fresh Frozen Tissue Samples (post-MALDI)

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#### **ABSTRACT**

## Scope:

To provide a method for removal of MALDI matrix, fixation and preparation of tissue sections for Cell DIVE.

## **GUIDELINES**

#### Special Notes:

- 1. Make sure no tissue drying, particularly after matrix removal.
- 2. Glass petri dishes can be used for more fragile tissues.
- 3. This protocol can be used on Cell DIVE fresh frozen samples.

## **MATERIALS**

#### Materials:

- 1. ITO slides with tissue from MALDI experiment
- 2. Coplin staining dish, Fisher S17495
- 3. Glass petri dish, Fisher 08-747C
- 4. Tabletop shaker

#### Reagents:

- 1. 4% paraformaldehyde in PBS, Fisher AAJ61899AK
- 2. Phosphate Buffered Solution (PBS), Fisher 70-011-044
- 3. Bovine Serum Albumin, Fisher BP671-10
- 4. Normal Donkey Serum, Jackson ImmunoResearch 017-000-121
- 5. Tween 20, Sigma P9416
- 6. Milliq ddH<sub>2</sub>O

## Solutions:

1. 1x PBS

100mL 10x PBS 900mL ddH<sub>2</sub>O

1000 mL total solution

2. Block Solution:

2.5 grams of BSA

5 mL of reconstituted donkey serum (reconstituted in 10mL of ddH<sub>2</sub>0)

0.25 mL of Tween 20 44.75 mL of 1X PBS

50 mL total block solution

# SAFETY WARNINGS

Paraformaldehyde should be used inside a chemical fume hood. For research use only.

1 After the tissue has been imaged via MALDI IMS, the slide should be placed in a coplin jar or glass petri dish with a solution of room temperature 4% paraformaldehyde in PBS (\*must be room temperature).

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working

Created: Jan 05, 2023

Last Modified: Jan 31, 2023

We use this protocol and it's

PROTOCOL integer ID: 74843

Keywords: Cell DIVE, MALDI IMS, fixation, matrix removal

2	Place jar/dish on a shaker for 5 minutes.
3	Discard the PFA (the MALDI IMS matrix will come off the slide in this step).

- 4 Add fresh 4% PFA in PBS and shake for 5 more minutes (for a total of 10 minutes in PFA).
  - Washing with PFA solution will fix and rehydrate the tissue as the matrix is removed. This preserves the tissue and antigens, and significantly reduces artifacts.
- Add fresh PBS for 3 minutes on shaker.
  Repeat this wash two more times for a total of 3 times for 3 minutes each.
  - pH should be neutral
- 6 Incubate the slides in 50 mg/mL bovie serum albumin (BSA), 10% (v/v) host-derived serum (depends on which antibodies they use), and 0.5% (v/v) Tween-20 in PBS for 30 minutes.
- **6.1** Be very careful. Tissue will be very fragile.
  - This permeabilizes the tissue slowly.
  - The Tween cannot be substituted for triton X, as triton X is too harsh and will degrade the tissue.
  - 30 minutes works for most. Can do up to 2 hours before the tissue gets very damaged/adherence issues/weird background.
- 7 Add fresh PBS for 3 minutes on shaker.
  Repeat this wash two more times for a total of 3 times for 3 minutes each.