

JAN 16, 2024

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snPATHO-seq

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1 more workspace ↓



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ABSTRACT

Formalin-fixed paraffin-embedded (FFPE) samples are valuable but under-utilised in single-cell omics research due to their low DNA and RNA quality. Leveraging recent single-cell genomic technology advances, we introduce snPATHO-seq: a versatile method to derive high-quality single-nucleus transcriptomic data from FFPE samples.

MATERIALS

Reagents and consumables

- Ethanol
- Xvlene
- Nuclease Free water
- 1x Phosphate Buffer Saline (PBS, Ca2+ and Mg2+ free)
- Liberase TM or TH (Roche)
- Collagenase D or P (Roche)
- Hyaluronidase (CAS 37326-33-3, Calbiochem)
- RPMI1640 (Gibco)
- EZ Lysis Buffer (Sigma)
- 10% BSA (MACS® BSA Stock Solution, Miltenyi)
- Glycerol 50%
- RNAse Inhibitor (RiboLock from Thermo or RNA Protector from Roche)
- (Optional) 4',6-Diamidino-2-Phenylindole, Dihydrochloride (DAPI, ThermoFisher)
- 40 um and 70 um pluriStrainer filters (pluriSelect) (MACS SmartStrainers are also possible)
- 25 G needle

Equipment

- Thermomixer with adjustable shaking (Eppendorf)
- Swinging bucket refrigerated centrifuge

DOI: dx.doi.org/10.17504/protocol s.io.8epv5x58dq1b/v1

Protocol Citation: Tony Wang, Michael Roach, Jasmine Plummer, Luciano G Martelotto 2024. snPATHOseq. protocols.io

https://dx.doi.org/10.17504/p rotocols.io.8epv5x58dq1b/v1

MANUSCRIPT CITATION: https://doi.org/10.1101/2022. 08.23.505054

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Protocol status: Working We use this protocol and it's working

Created: Jan 10, 2024

Last Modified: Jan 16, 2024

PROTOCOL integer ID:

93229

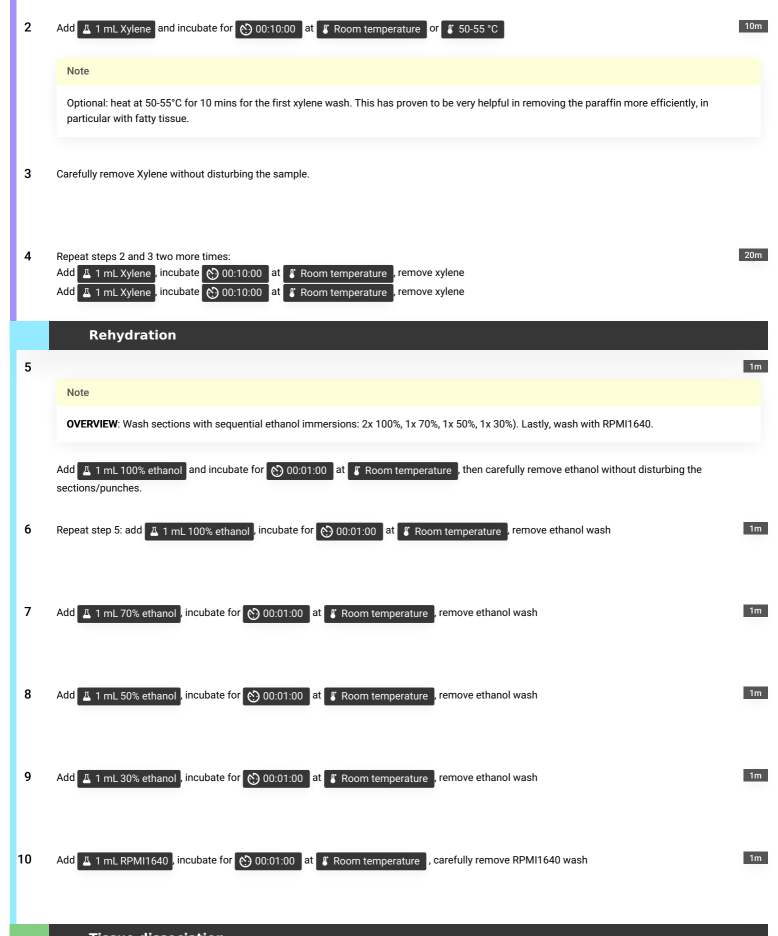
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Paraffin removal

Cut 1 or 2 approximately 25 µm-thick sections (punches are also possible) and transfer to a 1.5 mL Eppendorf tube.

Note

Store dry at 4°C if not used immediately. To keep it dry store in a container with silica beads. You can use the cylinder with silica beads that comes with 10x Genomics chips.



Prepare ☐ 1 mL Dissociation Solution

- [M] 0.25-1 mg/mL Liberase TM(*)
- [M] 0.25-1 mg/mL 10x Collagenase D(**)
- м 0.25-1 mg/mL Hyaluronidase
- [м] 1 U/uL RNAse Inhibitor

Add this 🗓 1 mL Dissociation Solution to the sections/punches.

Note

- (*) Liberase TH is very good enzyme too and can be used alone at 1-2.5 mg/mL.
- (**) Collagenase P (Roche) works fine as well.

IMPORTANT: I strongly recommend testing what concentration of enzymes works best for your tissue of interest. In our experience, Liberase TH alone at 1 mg/mL or the trio at 1 mg/mL Liberase TM + 1 mg/mL Collagenase D + 0.5 mg/mL Hyaluronidase works well with most of the samples we tested.

- Pro tip: Add 200 μL of the enzymatic cocktail and mince using a pestle for at least 10 strokes, then complete to 1 mL with the rest of the cocktail mix.
- Optional Pro tip: Before digestion, add 100 μL of digestion mix and homogenize the sample using a douncer/pestle by stroking 10-20 times. This helps in the digestion step. Then top up with the rest of the digestion mix.

12 Digest tissue for 45-60 mins(*) at 37°C in a Thermomixer at 800 RPM.

45m

(5) 800 rpm, 37°C, 00:45:00 in Thermomixer

Note

(*) Some blocks require longer digestion time. Inspect visually and help dissociation by pipetting up and down with a P1000 pipette.

IMPORTANT: Dissociation does not need to be complete; the objective here is to loosen up the material to facilitate the nuclei release. Dissociation completeness varies from block to block. Tissue does not need to be fully digested.

Lysing the cells

No

13

3111

OVERVIEW: Wash with lysis buffer. Resuspend and homogenize in small volume of lysis buffer. Add rest of lysis buffer and homogenise several times.

Add 🗸 400 µL Ez Lysis Buffer to the sample and mix by inverting 5 times, then centrifuge 😝 850 rcf, 4°C, 00:05:00

- Prepare 2 mL Lysis Solution as follows:

 - [M] 2 % (v/v) BSA
 - [M] 1 U/uL RNAse Inhibitor
- 15 Remove supernatant and add Δ 250 μL Lysis Solution (from step 14)

