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# 🌐 Nuclei isolation from fresh frozen human colon tissue for 10X Genomics Multi-omics (ATAC + GEX) assay

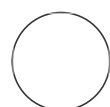
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## DISCLAIMER

This protocol is still being optimized! At the moment, we are testing if we can achieve better results without using the collagenase/dispase as dispase is a proteinase.

The buffers are based on the user guides from 10X Genomics.

## ABSTRACT

Nuclei isolation protocol used for sn multi-omics assay on fresh frozen human colon tissue (full thickness and muscularis layer only).

## MATERIALS

### BUFFERS

Buffers based on 10X Genomics user guides.

Collagenase/Dispase 10269638001 from Sigma

A	B	C	D	E	F
Collagenase Lysis Buffer	Stock conc		Final conc		V (µl)
Tris-HCl pH 7.4	1000	mM	10	mM	10,5
NaCl	5000	mM	10,0	mM	2,1
MgCl <sub>2</sub>	1000	mM	3	mM	3,2
DTT	1000	mM	1	mM	1,1

OPEN ACCESS

DOI:

[dx.doi.org/10.17504/protocols.io.5jyl8pn17g2w/v1](https://dx.doi.org/10.17504/protocols.io.5jyl8pn17g2w/v1)

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**Protocol status:** In development

It works, but we are still developing and optimizing this protocol

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**PROTOCOL integer ID:** 83804

**Keywords:** ASAPCRN

A	B	C	D	E	F
RNAseIn	40	U/uL	1	U/uL	26,3
Dispase/collagenase	0,1	mg/mL	0,003	mg/mL	31,5
NF H2O					975,5
Total Volume					1050,0
Wash Buffer 1	Stock conc		Final conc		V (µl)
Tris-HCl (pH 7.4)	1000	mM	10	mM	28,4
NaCl	5000	mM	10	mM	5,7
MgCl2	1000	mM	3	mM	8,5
BSA	10	%	1	%	283,5
Tween-20	10	%	0,1	%	28,4
DTT	1000	mM	1	mM	2,8
RNase in plus (promega)	40	U/ul	1	U/ul	70,9
Water					2406,9
Total Volume					2835,0
0.1x Lysis	Stock conc		Final conc		V (µl)
Tris-HCl (pH 7.4)	1000	mM	10	mM	5,3
NaCl	5000	mM	10	mM	1,1
MgCl2	1000	mM	3	mM	1,6
Tween-20	10	%	0,01	%	0,5
NP-40	10	%	0,01	%	0,5
Digitonin	0,5	%	0,002	%	2,1
BSA	10	%	1	%	52,5
DTT	1000	mM	1	mM	0,5
Rnase Inhibitor Protector (SIGMA)	40	U/ul	1	U/ul	13,1
Water					447,8

A	B	C	D	E	F
Total Volume					525,0
Wash Buffer 2	Stock conc		Final conc		V (μl)
Tris-HCl (pH 7.4)	1000	mM	10	mM	10,5
NaCl	5000	mM	10	mM	2,1
MgCl <sub>2</sub>	1000	mM	3	mM	3,2
BSA	10	%	1	%	105,0
Tween-20	10	%	0,1	%	10,5
DTT	1000	mM	1	mM	1,1
Rnase Inhibitor Protector (SIGMA)	40	U/ul	1	U/ul	26,3
Water					891,5
Total Volume					1050,0
1X Diluted Nuclei buffer	Stock conc		Final conc		V(μl)
Nuclei buffer	20	X	1	X	10
DTT	100	mM	1	mM	2
Rnase Inhibitor Protector (SIGMA)	40	U/ul	1	U/ul	5
Water					183,0
Total Volume					200

Instruments and equipment:

- Kai Medical 2 mm Biopsy punch BP-20F
- WPI Noyes scissors 12 cm S/S 15 mm blades
- VWR®, Disposable Pestles and Cordless Motor for Pellet Mixing
- PluriSelect 40 μm, 20 μm filters
- Flowmi 40 μm Cell strainer for 1000P
- Luna FL Cell Counter
- Swing out rotor centrifuge

## Sample preparation

Place the sample preparation instruments on dry ice: biopsy punch, 1.5 mL EP tube, tweezers,

- 1 petri dish.
- 2 Using 2 mm biopsy punch, aliquot 1.5-2 tissue pieces per patient. Place the pieces in a fresh 1.5 mL EP tube placed on dry ice. Repeat with a fresh biopsy punch for each multiplexed patient.

## Experiment preparation

- 3 Prepare all buffers fresh and on wet ice. Add the the detergents, DTT and RNase inhibitor just before use.
- 4 Place all the plastics and filters on wet ice.

## Tissue homogenization

- 5 **Perform all steps on wet ice.** Add 500  $\mu$ L Collagenase Lysis Buffer to the tube with multiplexed patient samples. Let thaw slightly.
- 6 Cut the tissue with scissors until there are no more pieces visible. **Time** = around 4 min (depending on the amount and the characteristics of the tissue).
  - 6.1 After cutting, place the scissors in a fresh 1.5 mL EP tube placed on ice.
- 7 Further homogenize the tissue using an automatic plastic pellet pestle, for 45 s.

- 7.1** Place the pellet pestle in the EP tube instead of the scissors.
- 8** Mince the tissue for 1 min.
- 9** Wash the scissors using 250  $\mu$ L Collagenase Lysis Buffer (while adding it to the tissue).
- 10** Homogenize the tissue with the used pellet pestle x15 (by hand).
- 11** Wash the pestle with 250 $\mu$ L Collagenase Lysis Buffer and add it to the tissue.
- 12** Gently pipet up and down 10X with P1000.
- 13** Centrifugate for 5 min at 500g at 4°C, remove the supernatant and resuspend in 1 mL Wash Buffer 1.
- 14** With P1000, transfer the sample onto 40 and 20  $\mu$ m stacked strainers placed on 25 mL EP tube.

- 15 Wash the 1.5 mL EP tube with additional 1 mL Wash buffer 1, transfer onto the top filter.
- 16 Centrifugate the 25 mL EP tube for 5 min at 500g, 4°C.
- 16.1 Gently remove the supernatant.
- 17 Resuspend the pellet in 700 µL of Wash Buffer 1 and filtrate through Flowmi filter into a fresh DNA LoBind 1.5 mL EP tube.
- 18 Centrifugate for 5 min at 500g at 4°C
- 18.1 Gently remove the supernatant.

## Nuclei permeabilization

- 19 Resuspend the pellet in 500 µL 0.1x Lysis Buffer, pipette mix X5 with P1000.
- 20 Incubate on ice for 1 min (tissue quality dependent).

**21** Add 1000  $\mu$ L of Wash Buffer 2 and pipette mix 5X with P1000.

**22** Centrifugate for 5 min at 500 g at 5°C.

**22.1** Very gently remove the supernatant.

## Nuclei resuspension and counting

**23** Resuspend the pellet in 25-50  $\mu$ L 1X Diluted Nuclei Buffer.

**24** Cell count the nuclei and proceed to the 10X Genomics Chromium Next GEM Single Cell Multiome ATAC + Gene Expression assay.