



Oct 03, 2019

Frozen Tissue Nuclei Extraction

Carly Martin¹, Ashley Feirrer¹, Evan Macosko¹¹Broad Institute[dx.doi.org/10.17504/protocols.io.2frg6m6](https://doi.org/10.17504/protocols.io.2frg6m6) **Velina Kozareva** 

ABSTRACT

Protocol for extraction of nuclei from frozen tissue in preparation for single-nuclei sequencing (droplet-based/10X). This protocol is based strongly on a similar extraction protocol from the [McCarroll lab](#).

THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION

Welch JD, Kozareva V, Ferreira A, Vanderburg C, Martin C, Macosko EZ. Single-Cell Multi-omic Integration Compares and Contrasts Features of Brain Cell Identity. Cell. 2019 Jun 6.

190318_Nuclei_Extraction
_Protocol.pdf

1 Make dissociation buffer, 50 mLs per sample:

Dissociation Media (Carter et al. 2009)

	MW	final concentration (mM)	for 1L (in g or ml)	for 2L (in g or ml)
Na ₂ SO ₄	142.04	82	11.65	23.3
K ₂ SO ₄	174.26	30	5.23	10.46
glucose	180.2	10	1.81	3.62
HEPES	238.3	10	2.39	4.78
MgCl ₂ 6H ₂ O	203.31	5	5 ml (1M stock)	10 ml (1M stock)

2 **Make extraction buffer**, 3 mLs per sample:

Dissociation Buffer + 1% Kollidon VA64 + 1% Triton X100 + 1:40 RNase-inhibitor

3 **Chill all buffers to 4C**, and keep all buffers on ice when in use.

4 **Prepare equipment:**

Set out and chill the following equipment/supplies. The instructions below are for one sample.

- Cold block dissecting tray, kept at -20C
- Chill clean razor blades and glass slides on the cold dissecting tray
- Chill 3 mL syringe with 26 1/2 gauge needle, at 4C, one syringe and needle per sample
- Place a 12-well tissue-culture plate, well-bottoms colored with marker, on ice
 - * Color the underside of the culture plate with dark marker so you can see your sample dissociate
- Place 2 mL chilled extraction buffer in the first well of 12-well plate
- Place ~50 mL chilled DB in a falcon tube on ice
- Locate DAPI (1:1000) (<https://www.thermofisher.com/order/catalog/product/D1306>)
- Locate a 100um filter for falcon tubes, place in 4C
(<https://us.vwr.com/store/product/14796270/vwr-cell-strainers-dnase-rnase-free-non-pyrogenic-sterile>)
- Locate 50 µm eppendorf tube filter, place in 4C
(<https://us.sysmex-flowcytometry.com/consumables/celltrics-filters/sterile-single-pack-celltrics-filters/1445/sterile-single-pack-celltrics-filters-50/box?number=04-00-4-2327>)
- Place 2 50 mL tubes on ice
- Locate a c-Chip FR hemocytometer

5 **Locate supplies for FACS:**

- 96 well cold block, chilled to -20C:
(https://www.daigger.com/eppendorf-pcr-coolers-14616-group?gclid=EAlaIQobChMI2rKM7aDN4AIVDRgMCh2jkwRMEAQYBSABEgKeA_D_BwE)
- 200uL 5% BSA-DB
- 20uL collection buffer: DB with 1:40 RNase-inhibitor
 - * Prepare the collection buffer right before beginning FACS session

6 Begin extraction. Perform all steps with cold buffers on ice. Nuclei should never leave the ice or 4C environment.

Chop up tissue on the chilled dissection tray. If your sample is a very small piece (less than 3mm in any dimension) you can forgo this step and place sample immediately into the extraction buffer

Place the frozen sample on the glass slide and shave the frosty/freezer-burned tissue face you want to sample from off the sample and discard. This is damaged and not worth extracting. Then, moving rapidly but carefully, shave off “enough” sample to work with.

Rapidly mince the shaved sample with the 2 chilled razor blades that were used to shave the tissue block and scrape into the first well of a 6-well plate that contains 2 mL of cold extraction buffer. Pipette up and down 20 times.

7 Incubate in detergent with mechanical trituration:

Commence 10-min incubation, pipetting up and down 20x every ~2.5 min with a 1000 uL pipette set to 1000uL.

8 Perform mechanical dissociation:

Pull the sample into the chilled needle/syringe and express into the same well for a final mechanical dissociation. If large chunks remain, repeat this step, pulling the sample into the same syringe and expressing it into the same well.

9 Filter out large chunks of white matter:

Wet the 100-μ filter with 1000 uL cold DB then run sample through the filter into one of the chilled 50 mL falcon tubes. Rinse the filter with 1000uL of DB into the falcon tube.

10 Large volume wash:

Add chilled DB to the extracted filtered nuclei to a total volume of just under 30 mL.

11 **Split sample:**

Divide these diluted nuclei among 2 50 mL conical tubes (~15 mL per tube).

12 **Spin tubes** at 500 rcf in a swinging bucket centrifuge for 10 minutes at 4C.

13 **Remove supernatant:**

Carefully aspirate ~13 mLs wash buffer with a serological pipette. To keep from disturbing pellet, gently aspirate the rest of the supernatant with a P1000 pipette. Leave about 500-300uL of sample in the bottom of the tube. Pipette sample to resuspend nuclei, being careful not to introduce bubbles. Pool the appropriate sample pairs back together.

14 **Filter nuclei clumps:**

Wet the 50um eppendorf tube filter with cold DB, pass the ~0.5-1 mL of washed sample through the filter into an epi tube.

15 **Stain DNA:**

Add 1ul of DAPI to 1ml of nuclei. Remove 20ul and place into FR hemocytometer. Check your nuclei concentration and nuclei quality. Some debris is ok.

16 **FACS enrichment for singlets on singlet DAPI peak:**

We collect FACS'ed samples in PCR tubes held in a chilled (-20C) 96-well cold block to keep the collection volume as minimal as possible: (https://www.daigger.com/eppendorf-pcr-coolers-14616-group?gclid=EAlaIQobChMI2rKM7aDN4AIVDRgMCh2jkwRMEAQYBSABEgKeA_D_BwE)

We FACS on a Sony SH800 sorter, using a 70um chip, with these settings:

Sample Stop Condition

None

Recording Setting

Type Event Count
Value 100,000

Instrument Setting

Laser	Threshold
405nm On	Channel FSC
488nm On	Value 1.00%
561nm On	
638nm On	

Sensor Gain

FSC	4	BSC	32.0%
FL1	40.0%	FL2	38.5%
FL3	31.0%	FL4	32.0%
FL5	32.5%	FL6	37.5%

Sample Pressure 5

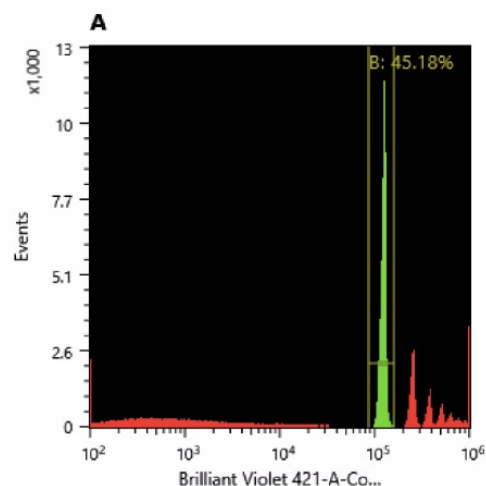
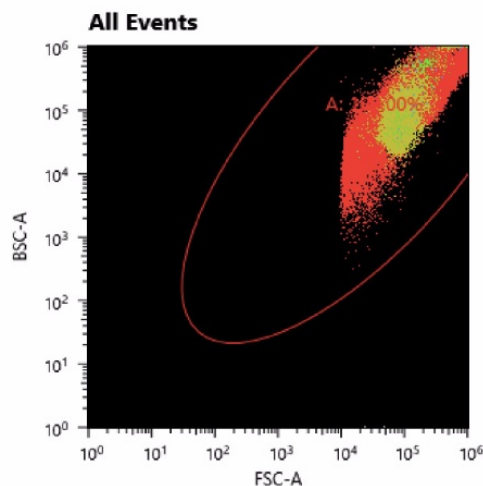
AD Advanced Setting

Forward Window Extension 50
Back Window Extension

Auto Parameters

Droplet Clock	50,700 Hz
Droplet Drive	46.88
Sort Delay	34
Sort Phase	200 deg
Charge	50.0 %
Deflection Left	1,291
Deflection Right	1,103
Enabled Control Breakoff	<input checked="" type="checkbox"/>

We gate on the singlet DAPI peak:



17 Prepare the collection PCR tubes:

- Fill the PCR tube with 5% BSA-DB, vortex, and remove BSA. *This coats the sides of the tube so that if nuclei hit the side of the tube they will fall to the bottom and not dry out.*
- Wash tube with 200 uL DB
- Make collection buffer, add 20uL collection buffer (DB with 1:40 RNase-inhibitor) to the PCR tube.

FACS into the tube using a 70um chip with 1%FSC threshold on the DAPI peak.

18 Calculate nuclei concentration:

After FACS, use a pipette to determine the volume in each tube.

Make a 1:10 dilution of your sample: combine 18ul chilled DB with 2ul of nuclei a PCR tube to make a 1:10 dilution. Mix and put into FR hemocytometer.

Visualize with the fluorescent scope, getting images of brightfield and DAPI-excited nuclei. Notice that debris is gone.

Count all 16 large squares to get the most accurate concentration estimate. Calculate the average of the 16 squares. Multiply that number by 10 (accounts for 1:10 dilution) then multiply by 5 (FC hemocytometer factor). This number is your final concentration in nuclei/uL.

19 Prep for 10X sequencing:

The maximum nuclei you can input is 17000 (which comes out to 10000 after loss). For 10x v3 the input nuclei volume is 46.6, which means the maximum concentration you can input is 364 n/uL.



This is an open access protocol distributed under the terms of the [Creative Commons Attribution License](https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited