

VERSION 2

MAR 19, 2024

OPEN BACCESS



DOI:

dx.doi.org/10.17504/protocols.io.x 54v92e8ml3e/v2

Protocol Citation: Wenxin Zhao, Sheng Zhong 2024. Protocol for Preparing Brain Samples for MUSIC. **protocols.io** https://dx.doi.org/10.17504/protoc ols.io.x54v92e8ml3e/v2Version created by Wenxin Zhao

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

Protocol for Preparing Brain Samples for MUSIC V.2

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Human BioMolecular Atlas Program (HuBMAP) Method Development Community

MUSIC



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DISCLAIMER

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The protocols.io team notes that research involving animals and humans must be conducted according to internationally-accepted standards and should always have prior approval from an Institutional Ethics Committee or Board.

ABSTRACT

Here states the detailed procedure to prepare brain samples for MUSIC study.



Created: Mar 19, 2024

Last Modified: Mar 19, 2024

PROTOCOL integer ID: 96945

Tissue pulverization and crosslinking

- 1 Cut a portion of post-mortem human brain frontal cortex sample collected from The Brain and Body Donation Program (BBDP) at Banner Sun Health Research Institute on dry ice with heavy razor blades, and collect 50 mg of the sample in a 1.5 mL LoBind tube.
- 2 Thaw the 50 mg of brain sample on ice, and chop the tissue into smaller pieces by pestle. Store the rest of the sample at -80°C.
- 3 Incubate the sample with 10 mL of 2 mM disuccinimidyl glutarate (DSG) in 1X PBS in a 15 mL LoBind tube at room temperature for 45 min with gentle rotation.
- 4 Wash once with 10 mL of 1X PBS by centrifugation at 1,000 x g for 4 min.
- Resuspend the sample in 15 mL of 1X PBS containing 3% formaldehyde, and incubate for 10 min with a gentle rotation.
- **6** Quench the crosslinking reaction by the addition of 5 mL of 1.25 M glycine followed by an incubation of 5 min with a rotation.

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7 Centrifuge the sample at 1,000 x g for 4 min, and wash the sample twice with ice-cold 1X PBS containing 0.3% BSA (wt/vol).

Nuclei isolation

- **8** Use Chromium Nuclei Isolation kit (10X genomics, 1000494) to isolate nuclei from crosslinked cortex samples.
- **9** Transfer 50 mg frozen tissue into pre-chilled sample dissociation tube.
- Add 400 μ L of Lysis Buffer to Sample Dissociation Tube. Dissociate tissue with plastic pestle until homogeneous.

	A	В
	Component	Volume (µL)
	Lysis Reagent	1000
Г	Reducing Agent B	1
Г	Sufactant A	10
	Total Volume	1011

Lysis Buffer

- Add 600 μ L of lysis buffer into the tube, and mix 10 times by pipetting. Incubate on ice for 10 min.
- 12 Equally load the solution into two nuclei isolation column, and centrifuge the tubes at 16,000 x g for 20 sec at 4°C.

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- Vortex the flowthrough in the collection tube that contains nuclei for 10 sec at 3,200 rpm or max speed to resuspend nuclei.
- 14 Centrifuge the collection tubes for 3 min at 500 x g at 4°C to pellet nuclei. Carefully discard the supernatant.
- 15 Resuspend the nuclei in 500 μL of Debris Removal Buffer provided by the kit by pipetting 15 times.

	A	В
Г	Component	Volume (µL)
Г	Debris Removal Reagent	500
Г	Reducing Agent B	0.5
	Total Volume	500.5

Debris Removal Buffer

- 16 Centrifuge the nuclei at 700 x g for 10 min at 4°C. Carefully discard the supernatant.
- 17 Resuspend the nuclei in 1 mL of Wash and Resuspension Buffer.

A	В
Component	Volume (µL)
1X PBS	1750
10% BSA	200
RNase Inhibitor (40X)	50
Total Volume	2000

Wash and Resuspension Buffer

18 Centrifuge the nuclei at 500 x g for 5 min at 4°C. Carefully discard the supernatant.