



Extraction of Metabolome From Tissue/Organ For HILIC LC-MS

**For batches of samples larger than 50, please contact our Metabolomics Facility regarding experimental design*

If interested in **NAD(P)(H) metabolites, please contact us as the prep will change*

If these are **material-limiting samples, please contact us as the prep will change*

1. Freshly collect the tissue/organ into a tube. **Note:** Do not rinse with PBS, if rinse is needed, quickly use saline.
2. Snap freeze in liquid nitrogen and store at -80°C until analysis.
3. For extraction of metabolites, section ~20-25 mg of frozen tissue/organ, and record the wet weight.
4. Add 300 uL of ice-cold **methanol**/water 80:20 (vol/vol). **Note:** This volume may depend on tissue type/level of necrosis, etc. Please notify the Core if you optimize this volume!
5. Homogenize the tissue using the blue pestle/manual homogenizer (Sigma Z359947). **Note:** Use methanol/water 50:50 (vol/vol) to clean the probe between samples.
6. Subject to three freeze-thaw cycles between liquid nitrogen and 37 °C.
7. Vortex rigorously for 1 min, and then centrifuge at ~20,160 xg for 15 min in a refrigerated centrifuge.
8. Transfer the metabolite-containing supernatant into a new tube.
9. Perform protein quantitation on the supernatant (on the 80% methanol portion, NOT the pelleted protein). Standard curve calculations below:

[BSA]	Volume of 2.0mg/mL stock BSA	Volume of 80% methanol
2.0	500uL	0uL
1.0	250uL	250uL
0.5	125uL	375uL
0.25	62.5uL	437.5uL
0.125	31.2uL	468.7uL
0.0625	15.6uL	484.4uL
0	0uL	500uL



NOTE: Protein measurements will be on residual protein in the supernatant; most protein will crash out during the extraction and pellet. We recommend running the Pierce BCA kit with the widest range curve using 80% methanol as the diluent for the standard curve. Each extract should be run in triplicate with at least one 10x dilution factor to ensure your samples are in the linear range of the assay.

10. Transfer a volume that is equivalent to 10 ug of protein to a new Eppendorf tube.
11. Dry sample in SpeedVac.
12. Resuspend sample in 100 uL **acetonitrile**/water 80:20 (vol/vol) for a final concentration of [0.1mg/mL] of protein (of the supernatant).
13. Vortex rigorously for 1 min, and then centrifuge at ~20,160 *xg* for 15 min in a refrigerated centrifuge. **NOTE:** This second spin down is CRITICAL to preventing instrument contamination.
14. Prepare pooled sample(s) for each biological condition for MS2 data acquisition. From the LC-MS vials in Step 14, take an aliquot of each sample into a new LC-MS vial (with insert) to make the final volume of the pooled sample to be at least 45uL. **NOTE:** If your samples are isotopically labeled, you do NOT need to make a pooled sample, but you WILL need an unlabeled (a 0 time-point) sample.
15. Transfer the metabolite-containing supernatant to an LC-MS vial (with insert) and cap. **NOTE:** The Core will provide all LC-MS vials.