UCD Genome Center	SOP Standard Operating Procedure [Under Development]	Page 1 of 2
date: 06/02/2011	Extraction of Mammalian Tissue Samples: Liver	Code No.: liver 060220101

Issued: 06/02/2011 Valid From: Not yet valid	Validity Area: UC Davis Genome Center, Metabolomics Core and Research Laboratories
Responsible:	
This SOP supersedes: SOP 05-28-2008	Approved:

# **Extraction of Mammalian Tissue Samples: Liver**

#### 1. References:

• Fiehn O, Kind T (2006) Metabolite profiling in blood plasma. In: Metabolomics: Methods and Protocols. Weckwerth W (ed.), Humana Press, Totowa NJ (in press)

#### 2.Starting material:

• Liver sample: weigh 4mg per sample into 2mL Eppendorf tubes.

## 3. Equipment:

- Centrifuge (Eppendorf 5415 D)
- Calibrated pipettes 1-200μl and 100-1000μl
- Eppendorf tubes 2mL, clear (Cat. No. 022363204)
- Centrifuge tubes 50mL, polypropylene
- Eppendorff Tabletop Centrifuge (Proteomics core Lab.)
- ThermoElectron Neslab RTE 740 cooling bath at −20°C
- MiniVortexer (VWR)
- Orbital Mixing Chilling/Heating Plate (Torrey Pines Scientific Instruments)
- Speed vacuum concentration system (Labconco Centrivap cold trap)
- Turex mini homogenizer

## 4. Chemicals

- Acetonitrile, LCMS grade (JT Baker; Cat. No.9829-02)
- Isopropanol, HPLC grade (JT Baker; Cat. No. 9095-02)
- Methanol
- Acetone
- Crushed ice
- $18 \text{ M}\Omega$  pure water (Millipore)
- Nitrogen line with pipette tip
- pH paper 5-10 (EMD Chem. Inc.)

### 5. Procedure

Preparation of extraction mix and material before experiment:

- 1. Switch on bath to pre-cool at  $-20^{\circ}$ C ( $\pm 2^{\circ}$ C validity temperature range)
- 2. Check pH of acetonitrile and isopropanol (pH7) using wetted pH paper

UCD Genome Center	SOP Standard Operating Procedure [Under Development]	Page 2 of 2
date: 06/02/2011	Extraction of Mammalian Tissue Samples: Liver	Code No.: liver 060220101

- 3. Make the extraction solution by mixing acetonitrile, isopropanol and water in proportions 3:3:2
- 4. De-gas the extraction solution for 5 min with nitrogen. Make sure that the nitrogen line was flushed out of air before using it for degassing the extraction solvent solution

### Sample Preparation

- 1. Weigh 4mg tissue sample in to a 2mL Eppendorf tube.
- 2. Add 1mL extraction solvent to the tissue sample and homogenize for 45 seconds ensuring that sample resembles a powder. In between samples, clean the homogenizer in solutions of methanol, acetone, water, and the extraction solvent in the order listed.
- 3. Vortex samples for 10 seconds, then 5 minutes on 4°C shaker.
- 4. Centrifuge the samples for 2 minutes at 14,000 rcf. Aliquot  $500\mu$ L supernatant for analysis, and  $500\mu$ L for a backup. Store backup aliquots in the -20°C freezer.
- 5. Evaporate one 500μl analysis aliquot in the Labconco Centrivap cold trap concentrator to complete dryness (typically overnight).
- 6. The dried aliquot is then re-suspended with 500µl 50% acetonitrile (degassed as given)
- 7. Centrifuge for 2 minutes at 14,000 rcf using the centrifuge Eppendorf 5415.
- 8. Remove supernatant to a new Eppendorf tube.
- 9. Evaporate the supernatant to dryness in the Labconco Centrivap cold trap concentrator.
- 10. Submit to derivatization.

The residue should contain membrane lipids because these are supposedly not soluble enough to be found in the 50% acetonitrile solution. Therefore, this 'membrane residue' is now taken for membrane lipidomic fingerprinting using the nanomate LTQ ion trap mass spectrometer. Likely, a good solvent to redissolve the membrane lipids is e.g. 75% isopropanol (degassed as given above). If the 'analysis' aliquot is to be used for semi lipophilic compounds such as tyrosine pathway intermediates (incl. dopamine, serotonine etc, i.e. polar aromatic compounds), then these are supposedly to be found together with the 'GCTOF' aliquot. We can assume that this mixture is still too complex for Agilent chipLCMS. Therefore, in order to develop and validate target analysis for such aromatic compounds, we should use some sort of Solid Phase purification. We re-suspend the dried 'GCTOF' aliquot in 300 user (degassed as before) to take out sugars, aliphatic amino acids, hydroxyl acids and similar logP compounds. The residue should contain our target aromatics. We could also try to adjust pH by using low concentration acetate or phosphate buffer. The residue could then be taken up in 50% acetonitrile and used for GCTOF and Agilent chipMS experiments. The other aliquot should be checked how much of our target compounds would actually be found in the 'sugar' fraction.

#### 6. Problems

To prevent contamination disposable material is used. Control pH from extraction mix.

### 7. Quality assurance

For each sequence of sample extractions, perform one blank negative control extraction by applying the total procedure (i.e. all materials and plastic ware) without biological sample.

#### 8. Disposal of waste

Collect all chemicals in appropriate bottles and follow the disposal rules.

UCD Genome Center	SOP Standard Operating Procedure [Under Development]	Page 3 of 2
date: 06/02/2011	Extraction of Mammalian Tissue Samples: Liver	Code No.: liver 060220101