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Instructions for Analyzing FTICR-MS Data acquired by the 21T (with Thermo Velos Pro)

Required Programs and Files

Formularity (<https://omics.pnl.gov/software/formularity>)

- Database file (.bin)
- Requires Windows (not compatible with MacOS/Linux)

Provided calibration file (.ref)

Provided R scripts

Procedure Overview

This protocol will outline how to generate an FTICR-MS data report. In brief, the provided .txt files contain peak lists which were generated using NOMSI software (PNNL, Richland, WA). These peak lists will be loaded into Formularity, a program capable of calibrating peaks, combining samples, and assigning chemical formulas, to create an unprocessed ICR report and an accompanying .log file. In order to detect samples that may have calibrated poorly, the provided “Processing_Formularity_Log.R” script can be used to generate a list of suspicious samples.

Detailed Procedure

1. Download Formularity from <https://omics.pnl.gov/software/formularity>. While on the website, be sure to also download the “CIA_DB_2016_11_21.bin” database file.
 - a. Downloading Formularity will also provide documentation and some example data.
 - b. For further details regarding this program, please refer to *Tolić et al, 2017 - Anal. Chem.*
2. After downloading Formularity, ensure that the settings mirror those in Figure 1 (also listed below)
 - a. Under the Spectra files section:
 - i. Charge: 1
 - ii. Ionization: proton_detachment
 - b. Under the Calibration section, drag and drop the "**Hawkes_neg.ref**" (doi: 10.1002/lom3.10364) file into the box specified “Drop calibration file” and keep the default settings (listed below if changed).
 - i. Select “quadratic” from the Regression drop-down menu
 - ii. Start tolerance, ppm: 4
 - iii. End tolerance, ppm: 0.1
 - iv. Rel. factor: 1000000
 - c. Under the peak filters section, ensure:
 - i. Min S/N: 2
 - ii. Min rel. abund.: 0
 - iii. Max rel. abund.: 1
 - d. Under the “CIA formula finder” tab:

- i. Drag and drop the WHOI_CIA_DB_2016_11_21.bin file into the box specified “Drop DB files”. Settings should be default
 - ii. “Alignment” needs to be checked
 - iii. Alignment tolerance, ppm: 0.5
 - iv. Formula tolerance, ppm: 0.3
 - v. DB mass limit: 500
 - vi. Formula score: min(N+S+P) & The lowest error
 - vii. Max relationship gaps: 5
 - viii. Error: AMU – 0.00002
 - ix. Within the “Formula building blocks” box, ensure that CH₂, CH₄O-1, H₂, and O are checked.
 - x. User-defined filter: O>0 AND N<=4 AND S<4 AND P<=2
 - xi. Ensure that “Use relationship”, “Use formula filters”, and “CIA” (top right in the Spectra Files section) are all checked.
3. With the correct settings, you can now drag and drop the provided .txt files into the box labeled “Drop Spectra Files” in the upper right of Formularity.
 - a. The box will turn red and Formularity will begin generating the report, which can take up to 30-45 minutes depending on sample number, and the program might seem unresponsive.
4. Once the program finishes running, two files will be generated (Report.csv and a log file) in the folder containing the .xml files.
 - a. We recommend renaming the report to match your dataset name; this will prevent it from being overwritten if you run Formularity again.
5. Next, edit the "**Processing_Formularity_Log_v2.R**" script to include your dataset name and directory containing the log file and run it through either R, RStudio, or command line.
 - a. This will generate two files: one with all calibration results and another specifying poorly calibrated samples. **We recommend removing these samples.**
6. You now have a finalized FTICR-MS Report! We recommend using the R package “ftmsRanalysis” (<https://github.com/EMSL-Computing/ftmsRanalysis>) to further process the data (i.e., filter peaks by mass, calculate molecular properties, classify compounds, etc.).

Columns in FTICR output are as follows:

- Mass - Measured ionic mass (m/z)
- Atoms in the analyzed molecule; identified by an algorithm in Kujawinski and Behn 2006 and modified as described in Tfaily et al. 2017
 - C - Carbon
 - H - Hydrogen
 - O - Oxygen
 - N - Nitrogen
 - C13 – Carbon 13
 - S - Sulfur

- P - Phosphorus
- Na - Sodium
- El_Comp - Function group(s) that are known.
- NeutralMass - Zero-charge or neutral molecular mass. Calculated from observed ionic mass by balancing charge with protons or electrons.
- Error_ppm - Parts per million error calculated as [expected molecular mass - measured molecular mass]/expected molecular mass * 1e6.
- Candidates - Possible known molecules the analyzed molecule could be. Assigned by checking against a known database. Value of -1 when molecule was not checked against database.
- All remaining columns identified by sample name. These are from the spectra for each sample where the numerical value is the peak intensity at the particular molecular mass.

Formularity

Spectra files
 Charge: 1 Ionization: proton_detachment ☒ CIA ☐ IPA
 Adduct: Result: M-p Drop Spectra Files

Calibration
 Drop calibration file: Hawkes_neg.ref
 Regression: linear Start tolerance, ppm: 5.000000 End tolerance, ppm: 0.500000
 Rel. factor: 1000000.0000

Peak filters
 Min S/N: 7.000 Min rel. abund.: 0.000 Max rel. abund.: 1.000

CIA formula finding IPA formula finding Error plot CIA DB inspector File convertor CIA filter inspector About

Formula assignment
 Alignment ☒ Alignment tolerance, ppm: 0.500
 Loaded: WHOI_CIA_DB_2016_11_21.bin
 Formula tolerance, ppm: 0.500000
 DB mass limit: 500.000
 Formula score: min(N+S+P) & The lowest error
 Use formula filters ☒
 Use relationship ☒ Max relationship gaps: 2 Error: AMU 0.00002000
 Formula building blocks
☒ CH2 ☐ CH4O-1 ☒ H2 ☐ C2H4O ☐ CO2 ☐ C2H2O ☒ O
 Special filter: None
 User-defined filter: O>0 AND (N+S+P)<6 AND S<3 AND P<2
 Use default advanced ☒ Advanced ...

Save parameters Load parameters Load CIA parameters

Figure 1: Default Formularity settings.