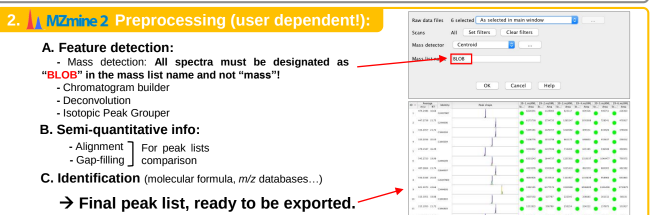
1. LC-MS² analysis and data conversion: conversion of proprietary files into universal .mzXML format using MSConvert (ProteoWizard).





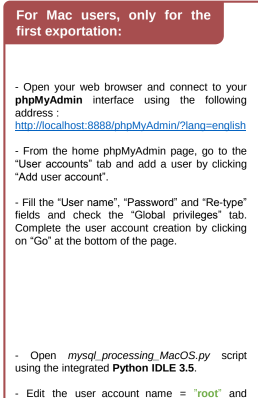
3. Exportation:



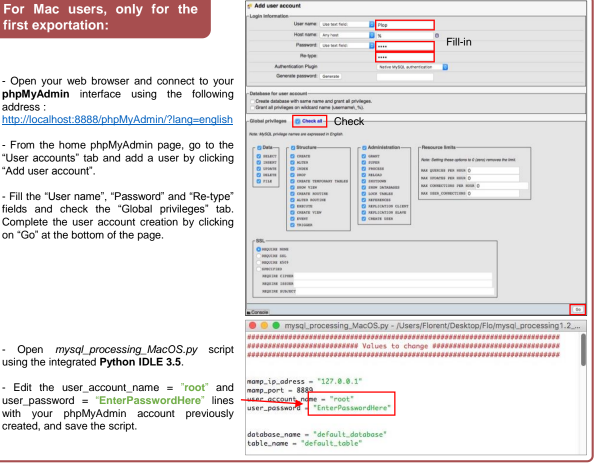
- Download Python 3.5 (or upper version) and Mamp 4.1.1 (Mac users) or WampServer 3.0.6 (Windows users).
- Download the 4 python scripts afforded at https://github.com/ElMouth/MZM2-MN
- Open **Mamp** and start the servers or simply open **WampServers**.

Only for first use: - Update the pip package for Python by launching the get-pip.py script via Python Launcher.

- Import the Pandas, XLWT, and PyMySQL libraries for Python by launching the installing_modules.py script with Python Launcher.



created, and save the script.

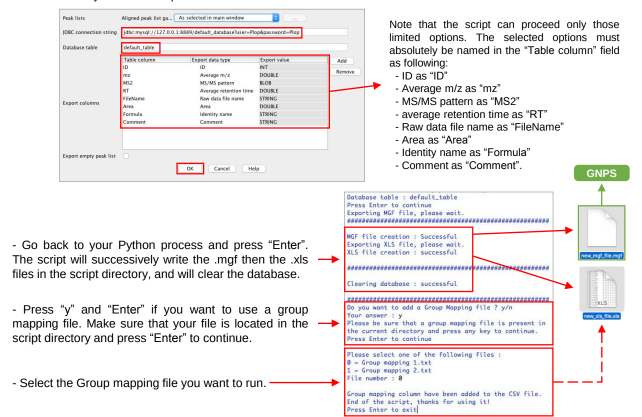


- With your MZmine 2 processed data in hand and the local server started, launch the mysql_processing.py (Mac or Win) script using the Python launcher (for Mac users Python IDLE (then press F5)).

- A new mySQL database
(default_database) and table
(default_table) are automatically
generated. Then, you are asked to
export your MZmine 2 data using the
given JDBC connection string and
Database table parameters:

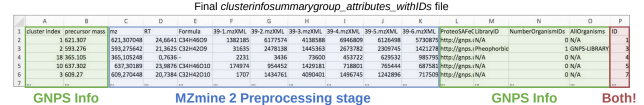
Type "copyright"
>>> WARNING: The
Visit http://www
Connection to Ma
Connection to Ma
Connection to Ma
Issue "copyright"
>>> WARNING: The
Visit http://www

- Go back to MZmine 2 and click in the "peak list methods" tab, "Import/Export" and "Export to SQL database".
- Fill the "JDBC connection string" and "Database table" fields using the parameters afforded in the script and select the data you want to export. Click "OK":



4. GNPS:

- Launch a GNPS job with the exported .mgf file. Make sure the MS-Clustering tab is **deactivated** and the Minimum cluster size set to 1.
- Download the clustered data from GNPS and open the *clusterinfosummarygroup_attributes_withIDs* file with Excel. Simply order this table by increasing RT (*RTMean* column), that corresponds in fact to you Mzmine *ID*. This column can be renamed *ID* and all other useless columns can be delete (all except *cluster index*, the newly renamed *ID*, *precursor mass*, *ProteoSAFeClusterLink*, *LibraryID*, *NumberOrganismIDs* and *AllOrganisms*).
- Open the .xls file from MZmine and order it by increasing *ID* value too. Copy this table and just append it to the *clusterinfosummarygroup_attributes_withIDs* file:



- Open Cytoscape and import a new network from the *networkedges_selfloop* file. Load the modified data table *clusterinfosummarygroup_attributes_withIDs* for accurate semi-quantitative pie chart drawing based on relative areas and molecular formula implementation.