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Quick-start for Untargeted Metabolomics Analysis in GNPS

Tiago F. Leao¹, Chase M. Clark², Anelize Bauermeister¹, Emmanuel O. Elijah¹, Emily Gentry¹, Makhai Husband¹, Michelli Faria de Oliveira³, Nuno Bandeira^{1,4,5,6,*}, Mingxun Wang^{1,4,*}, Pieter C. Dorrestein^{1,7,8,*}

¹–Collaborative Mass Spectrometry Innovation Center, Skaggs School of Pharmacy and Pharmaceutical Sciences, University of California San Diego, La Jolla, California, USA.

²–Department of Medicinal Chemistry and Pharmacognosy, College of Pharmacy, University of Illinois at Chicago, Chicago, Illinois, USA.

³–Department of Medicine, University of California San Diego, La Jolla, California, USA.

⁴–Center for Computational Mass Spectrometry, University of California San Diego, La Jolla, California, USA.

⁵–Department of Computer Science and Engineering, University of California San Diego, La Jolla, California, USA.

⁶–Skaggs School of Pharmacy and Pharmaceutical Sciences, University of California San Diego, La Jolla, California, USA.

⁷–Departments of Pharmacology and Pediatrics, University of California San Diego, La Jolla, California, USA.

⁸–Center for Microbiome Innovation, University of California San Diego, La Jolla, California, USA.

To the editor - with this correspondence we aim to introduce Global Natural Product Social Molecular Networking (GNPS) to your readership and to highlight the quick-start interface we have recently developed. GNPS is a chemistry-focused, community-curated, small molecule mass spectrometry analysis ecosystem for untargeted metabolomics data.¹ GNPS was designed for the analysis and capture of mass spectrometry knowledge and data by the community. These capabilities are possible due to the deep integration of GNPS with the metabolomics data repository MassIVE. Co-localizing metabolomics data, knowledgebases, and computational infrastructure allows for maximal reuse by the community in the spirit of open data. As of February 2021, GNPS/MassIVE has grown to 1,800 public data sets (>490,000 mass spectrometry files, >1.2 billion tandem mass spectra) with on average over

*To whom correspondence should be addressed: pdorrestein@health.ucsd.edu, miw023@ucsd.edu, bandeira@eng.ucsd.edu).

Author contributions

M.W., N.B. and P.C.D. designed the project. M.W. and C.M.C. develop the code for the quick-start website. T.F.L. and A.B. tested the website and prepared the examples from public data. M.W. and A.B. prepared the online documentation. M.H., M.F.O., E.O.E. and E.G. collected the untargeted LC-MS/MS data published at MassIVE. T.F.L., M.W., N.B. and P.C.D. wrote this correspondence and all authors reviewed/edited the text and figure.

Competing Interests

P.C.D. is a scientific advisor to Sirenas, Galileo and Cybele and co-founder and scientific advisor to Omata and Enveda with approval by the University of California San Diego. M.W. is a co-founder of Omata Labs LLC.

300,000 accesses to the GNPS analysis ecosystem per month by users from more than 160 countries.

The GNPS ecosystem has been used in different fields, including natural products, exposomics, nutrition science, forensics, marine science, environmental science and clinical metabolomics (28.67% of all public files with metadata are associated with a disease condition and 21.36% of the human files are associated with a disease). Approximately 50 analytical tools are connected to the GNPS ecosystem/platform. These tools are accessible without having to convert or move raw data out of the platform. Uniquely, the tools at GNPS enable users to directly match their data to all public MS/MS reference libraries for annotation¹, and perform molecular networking, which enables the discovery of related metabolites (including transformations due to metabolism) without needing an isotopic label (Fig. 1a–f).^{1–3} The processed data tables and results from GNPS jobs can be exported into other downstream analysis infrastructures such as Cytoscape⁴, Metaboanalyst⁵ or QIIME2⁶, which provide interactive network, statistical, machine learning or multivariate analysis and visualization capabilities (Fig. 1g). In addition, all public data deposited at GNPS/MassIVE can be searched using MS/MS spectra (MASST)⁷ or using controlled vocabularies or annotations (ReDU).⁸ MASST and ReDU are allowing researchers to leverage the information in the entire GNPS/MassIVE repository to provide a global context to their data. Finally, all tools within GNPS aim to improve data analysis reproducibility by facilitating easy replication with identical parameters and the sharing of data analysis job results as URL links (Fig. 1).

However, GNPS is admittedly not very easy to use. Users must learn to convert data to open formats, how to transfer files via ftp clients into GNPS, and how to submit jobs before they can visualize the data as a molecular network (Fig. 1a–f) or as an interactive PCoA plot (Fig. 1g). We thus developed an easier-to-use quick start infrastructure (<https://gnps-quickstart.ucsd.edu>) for data conversion, validation of library batch files, public data set creation, classic and Feature-Based Molecular Networking (FBMN) for up to 50 files (87% of GNPS's usage, Fig. 1h). These improvements aim to make the infrastructure more accessible to a wider scientific community, including users interested in the study of cellular metabolic pathways or lipids. Complementing the quick-start, we have created documentation for all GNPS' tools (<https://ccms-ucsd.github.io/GNPSDocumentation/>).

Data conversion and upload is the first step to use GNPS, a pain point for many beginner users of mass spectrometry. Our online data conversion quick-start infrastructure, leveraging ProteoWizard⁹, enables the conversion of LC-MS/MS files from most vendor formats to the open data format .mzML, the preferred format for GNPS analysis. Once converted, the data can be used in two other quick-start modules: Classical¹ and FBMN³. Using Classical Molecular Networking, users can drag-and-drop files into up to three possible cohorts and select the type of instrument used (high vs low resolution). That is all that is necessary for users to run their first Classical GNPS Molecular Networking workflow. For FBMN, users can drag-and-drop a quantification table and a MS/MS spectral file (e.g. .mgf) that is generated according to the FBMN instructions (pre-processing instructions at <https://ccms-ucsd.github.io/GNPSDocumentation/featurebasedmolecularnetworking/>). Another GNPS quick-start module enables data deposition into MassIVE (<https://massive.ucsd.edu/>). The

last quick-start is to help build the knowledgebase in GNPS by allowing the community to deposit public MS/MS libraries. This quick-start allows validation of the library batch files to ensure compatibility with the GNPS ecosystem before uploading MS/MS reference spectra to the GNPS spectral database. None of the quick-start pages require login to run jobs. However, if the user creates an account, it can store jobs and results in the user space; otherwise, the jobs without a user account will be cleared periodically.

The inherent richness of untargeted metabolomics data makes analysis difficult. GNPS, a resource for metabolomics, lipidomics and natural products communities, has been developed for and by these communities to put forth a set of tools that help to tackle the complexities of untargeted LC-MS/MS data analysis. As a community resource, GNPS offers many innovative solutions/tools for data analysis that are continually growing due to community contributions. We hope the ever-expanding capabilities of GNPS can be made more accessible and reach a broader audience with the introduction here of the GNPS quick-start set of tools and interfaces.

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Code availability

The GNPS quick-start page can be accessed at <https://gnps-quickstart.ucsd.edu> and the github repository for the code is available at https://github.com/mwang87/GNPS_quickstart. We used the public MassIVE datasets MSV000085256 (DOI 10.25345/C5D410) and MSV000083437 (DOI 10.25345/C51G8P).

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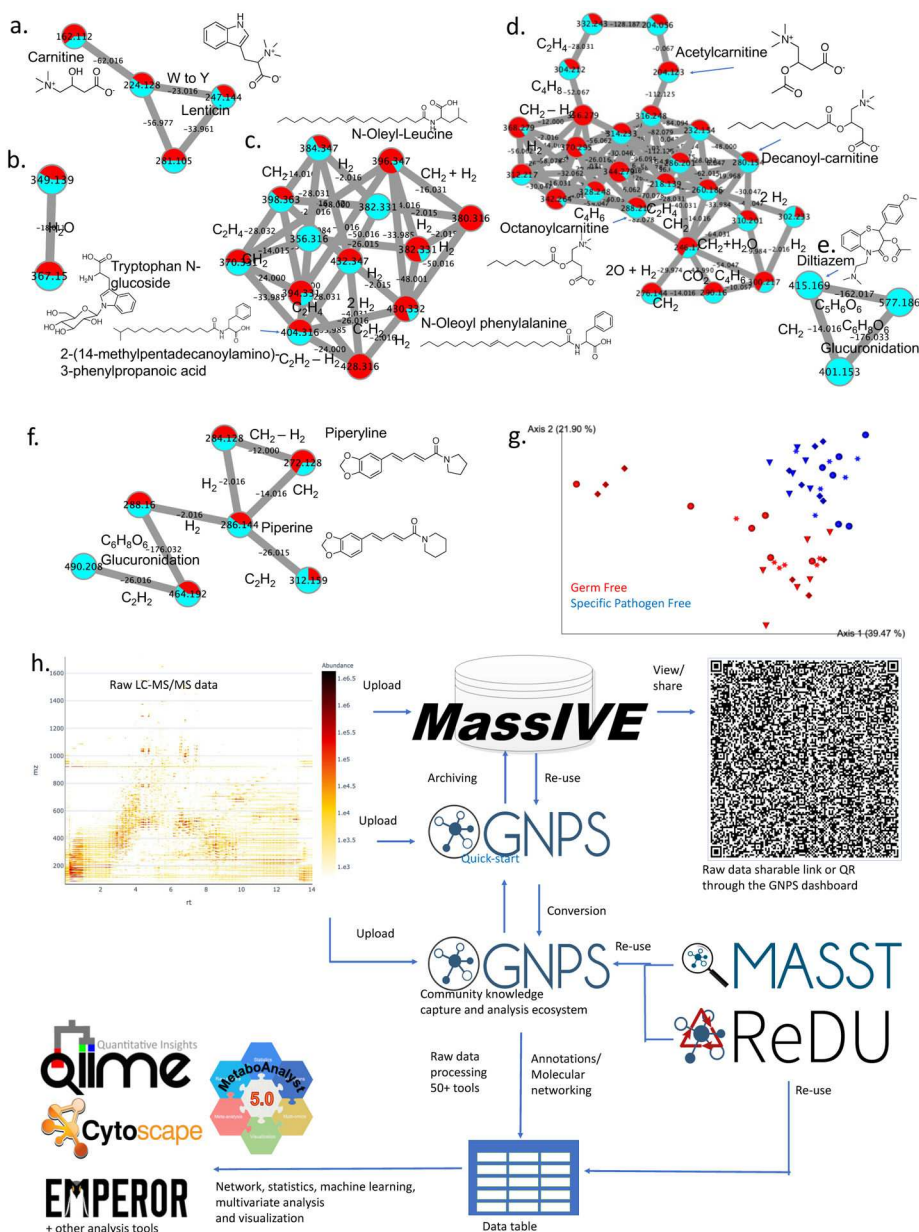


Fig. 1 |. The quick-start interface of GNPS can be used for Molecular Networking and multivariate analysis.

Representative molecular families obtained via Classical MS/MS Molecular Networking of plasma from an Alzheimer's disease (AD) (public MassIVE data set MSV000085256, DOI 10.25345/C5D410 and complete analysis job can be accessed at <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=ab00b95c6f2149bf92973c1a6bf19be6>). The molecular network is visualized with Cytoscape.⁴ Representative molecular families (sub-networks) reveal metabolites (a,b), lipids (c,d), drugs (e) or food-derived molecules (f) and their natural analogs (e.g. CH=CH, CH₂) and metabolites from phase I (e.g. reduction, demethylation or oxidation) and/or Phase II metabolism (e.g. glucuronidation). The families are annotated based on spectral matches against MS/MS reference libraries.¹ Blue are the relative levels in normal controls while red are the levels observed in the AD patients (relative levels

approximated via spectral count). **g**, PCoA visualization of the binary Jaccard similarity of the data from multiple duodenum sections associated with germ free and specific pathogen free mice (public MassIVE data set MSV000083437, DOI 10.25345/C51G8P) using Qiime 2⁶ and Emperor¹⁰ for visualization (An interactive link to the PCoA can be found here https://view.qiime2.org/visualization/?type=html&src=https%3A%2F%2Fcors.bridged.cc%2Fhttps%3A%2F%2Fgnps.ucsd.edu%2FProteoSAFe%2FDownloadResultFile%3Ftask%3D65a02c388b224f5dbd8311343731a8f2%26file%3Dqiime2_output%2Fqiime2_emperor.qzv%26block%3Dmain, n=4 mice for each condition, 6 sections each, each mouse is indicated by color and symbol). **h**, overview of the GNPS analysis ecosystem, including the new GNPS quick-start, analysis ecosystem and how it interfaces with the MassIVE repository, the GNPS search engines ReDU and MASST and the processing of data tables and external data table analysis and visualization tools. The QR code leads to the GNPS dashboard to visualize raw data.