

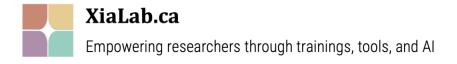
Spectra Processing, Compound Annotation, Functional Insight and Causal Analysis using MetaboAnalyst 6.0

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Schedule

Part I: 2:15 pm – 4:15 pm

- **2:15 2:30:** General introduction
- 2:30 3:00: Untargeted metabolomics
 - ✓ LC-MS & MS/MS spectral processing
 - √ From peaks to functions
- **3:05 3:25**: Live demo
- **3:25 4:15:** Hands on practice

Part II: 4:30 p.m. – 6:30 p.m.

- **4:30 5:15**: Background
 - ✓ Statistical analysis
 - √ Causal analysis
- 5:15 5:35: Live demo
- **5:40 6:20**: Hands on practice
- **6:20 6:30**: Summary & discussion

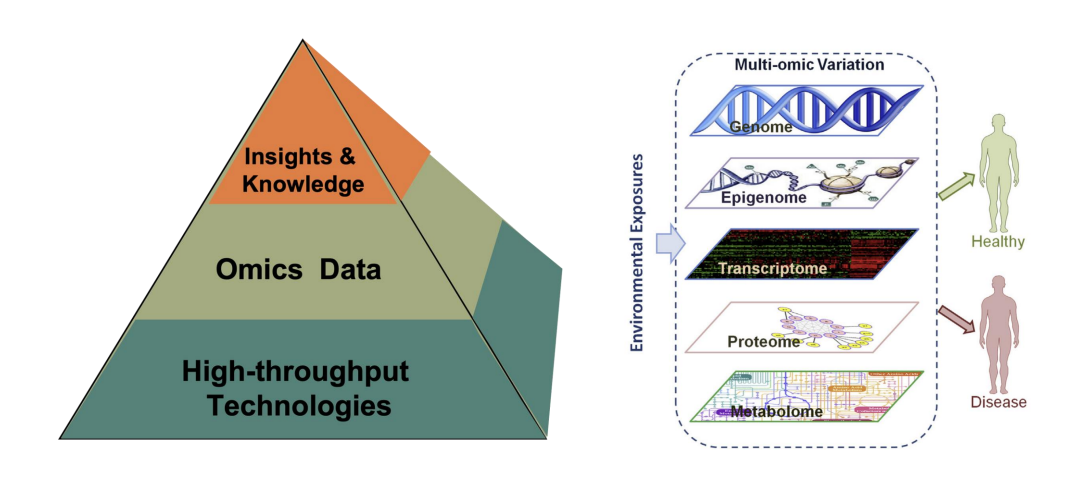
Github Repository

https://github.com/xia-lab/Metabolomics_2024

- Slides (in PDF format);
- Example data;
- Reference literatures;
- Contact information.

General Intro to Omics Data Science

Omics & multi-omics era



Two distinct challenges

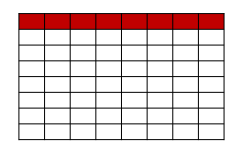
Size challenge (raw data)

- ➤ Raw reads, spectra, images
- ➤ Large (GB ~TB)
- ➤ Large storage and computing resources

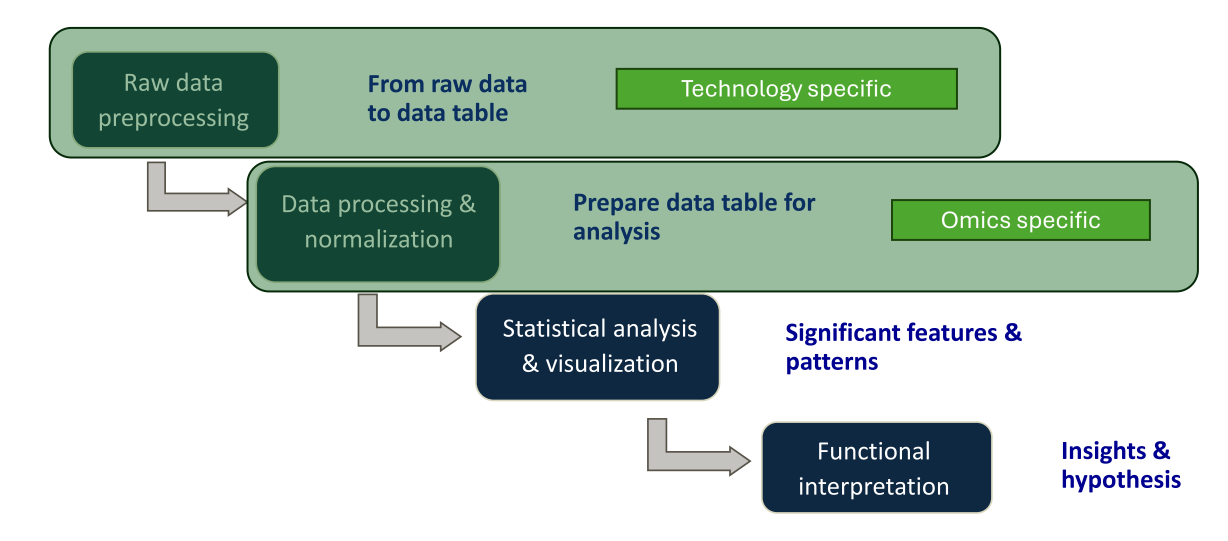
Complexity challenge (feature table)

- > Feature table (abundance, intensities)
- ➤Small (100s KB ~ MB)
- ➤ High-dimensional, missing values
- ➤ Data analysis starts here

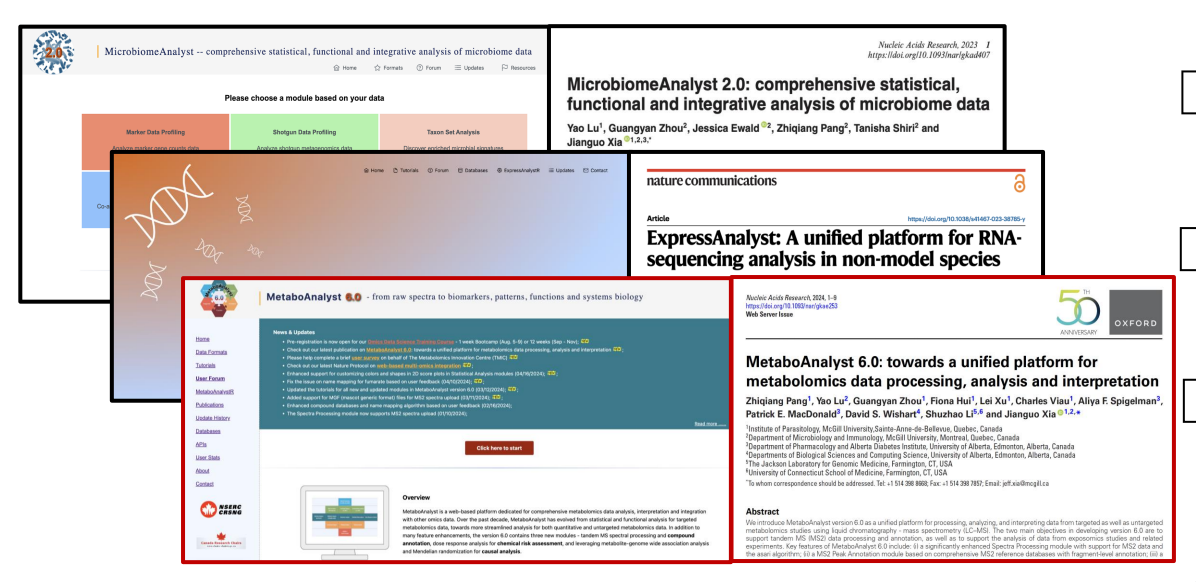




General workflow for single omics



Unified frameworks for single omics analysis



Omics Data Science Training Course

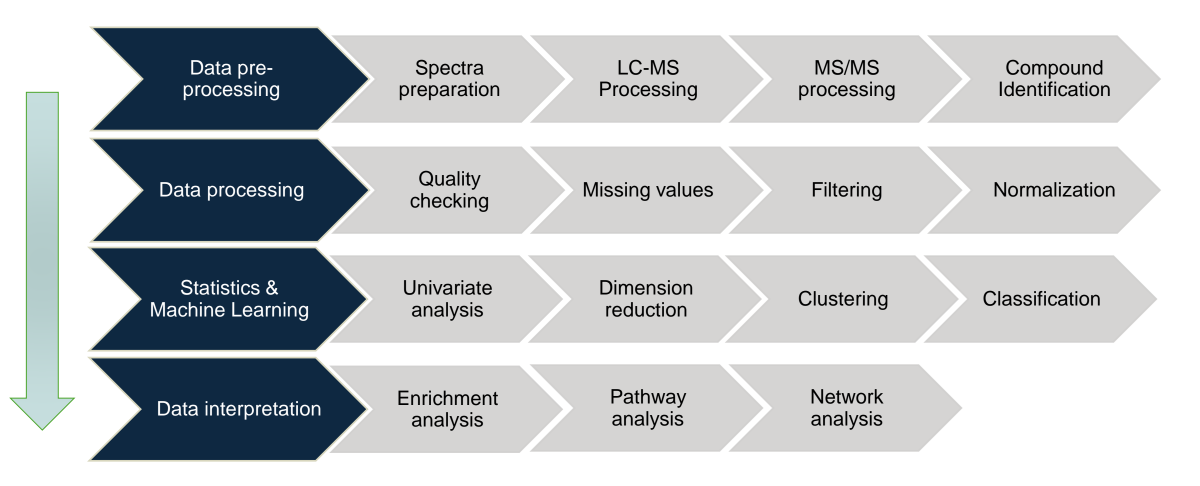
- Summer Bootcamp
 - O Aug. 5 9, **9:30 16:30**
- Regular Session
 - Saturday morning 9:30 12:00, Sept. Nov.

Topics	Tools	1. Nature Communications (2023) 2. Current Protocol (2023)		
Transcriptomics	ExpressAnalyst			
Biological Networks	NetworkAnalyst miRNet	3. Methods in Molecular Biology (2022) 4. Nucleic Acids Research (2020) 5. Nucleic Acids Research (2019)		
Metabolomics	MetaboAnalyst	Nucleic Acids Research (2024) Nature Protocols (2022) Current Protocols in Bioinformatics (2019)		
Microbiomics	2.0 MicrobiomeAnalyst	9. Nucleic Acids Research (2023) 10. Nature Protocols (2020)		
Multi-omics	OmicsNet OmicsAnalyst	11. Nature Protocol (2024) 12. Nucleic Acids Research(2022) 13. Nucleic Acids Research (2021)		
Special Topics	Biomarker analysis & meta-analysis	14. Metabolomics (2013) 15. Nature Protocols (2015)		
	RNAseq processing for non-model species	16. Genome Research (2021)		
	LC-MS and MS/MS spectra processing	17. Nature Communications (2024)		

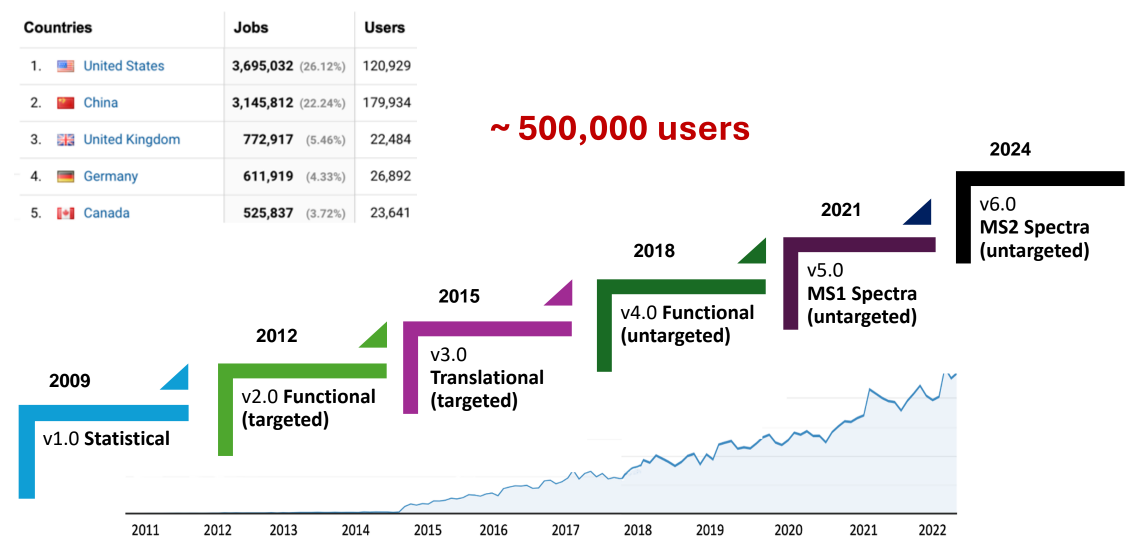
Key Publications

- Liu, P., Ewald J., Legrand E., Jeon, Y.S., Sangiovanni, J., Hacariz, O., Pang, Z., Zhou, G., Head, J., Basu, N., and Xia, J. @ (2023) "A unified platform for RNA-seq analysis in non-model species", <u>Nature Communications</u> 14, 2995.
- Ewald J., Zhou, G., Lu, Y., and Xia, J.@ (2023) "Using ExpressAnalyst for Comprehensive Gene Expression Analysis in Model and Non-Model Organisms" <u>Current Protocols</u> 3 (11), e922
- Chang, L., Xia, J.@ (2022) "MicroRNA Regulatory Network Analysis Using miRNet 2.0" Chapter Transcription Factor Regulatory Networks, 185-204, Methods in Molecular Biology, Humana Press, New York, NY.
- Chang, L., Zhou, G., Soufan, O. and Xia, J.@ (2020) "miRNet 2.0: network-based visual analytics for miRNA functional analysis and systems biology" <u>Nucleic Acids Research</u> (doi: 10.1093/nar/gkaa467)
- Zhou, G., Soufan, O., Ewald, J., Hancock, REW, Basu, N. and Xia, J@. (2019) "NetworkAnalyst 3.0: a visual analytics platform for comprehensive gene expression profiling and meta-analysis" Nucleic Acids Research 47 W234-241
- Pang, Z., Lu, Y., Zhou, G., Hui, F., Xu, L., Viau, C., Spigelman, A., MacDonald, P., Wishart, D., Li, S., and Xia, J. @ (2024)
 "MetaboAnalyst 6.0: towards a unified platform for metabolomics data processing, analysis and interpretation" <u>Nucleic Acids Research</u> (doi: 10.1093/nar/gkae253)
- Pang, Z., Zhou, G., Ewald, J., Chang, L., Hacariz, O., Basu, N., Xia, J.@ (2022) "Using MetaboAnalyst 5.0 for LC-HRMS spectra processing, multi-omics integration and covariate adjustment of global metabolomics data" <u>Nature Protocols</u> (doi: 10.1038/s41596-022-00710-w).
- Chong, J., Wishart, D.S. and Xia, J.@ (2019) "Using MetaboAnalyst 4.0 for Comprehensive and Integrative Metabolomics Data Analysis". <u>Current Protocols in Bioinformatics</u> 68, e86 (doi: 10.1002/cpbi.86)
- Lu, Y., Zhou, G., Ewald, J., Pang, Z., Shiri, T., and Xia, J.@ (2023) "MicrobiomeAnalyst 2.0: comprehensive statistical, functional and integrative analysis of microbiome data" Nucleic Acids Research (doi: 10.1093/nar/gkad407).
- Chong, J., Liu, P. and Xia, J@. (2020) "Using MicrobiomeAnalyst for comprehensive statistical, functional and meta-analysis of microbiome data" <u>Nature Protocols</u> (doi: 10.1038/s41596-019-0264-1)
- Ewald J., Zhou, G., Lu, Y., Kolic, J., Ellis, C., Johnson, J., Macdonald P., and Xia, J. @ (2024) "Web-based Multi-omics Integration Using the Analyst Software Suite", Nature Protocols (doi: 10.1038/s41596-023-00950-4
- Zhou, G., Pang Z., Lu, Y., Ewald, J., Xia, J.@ (2022) "OmicsNet 2.0: a web-based platform for multi-omics integration and network visual analytics" <u>Nucleic Acids Research</u> (doi:10.1093/nar/gkac376).
- Zhou, G., Ewald, J., and Xia, J.@ (2021) "OmicsAnalyst: a comprehensive web-based platform for visual analytics of multi-omics data" <u>Nucleic Acids Research</u> (10.1093/nar/gkab394)
- Xia, J., Broadhurst, D., Wilson, M. and Wishart, D. (2013) "Translational biomarker discovery in clinical metabolomics: an introductory tutorial". Metabolomics (doi: 10.1007/s11306-012-0482-9)
- Xia, J@., Gill, E., and Hancock, R.E.W@. (2015) "NetworkAnalyst for Statistical, Visual and Network-based Approaches for Metaanalysis of Expression Data" <u>Nature Protocols</u> 10 (6), 823-844
- Liu, P., Ewald, J., Galvez, J., Head, J., Crump, D., Basu, N. and Xia, J.@ (2021) "Ultrafast functional profiling of RNA-seq data for nonmodel organisms" <u>Genome Research</u> (doi: 10.1101/gr.269894.120)
- Pang, Z., Xu, L., Viau, C., Lu Y., Salavati, R., Basu, N., and Xia, J.@ (2024) "MetaboAnalystR 4.0: a unified LC-MS workflow for global metabolomics" Nature Communications (doi:10.1038/s41467-024-48009-6)

Common Tasks for Metabolomics



MetaboAnalyst roadmap



MetaboAnalystR



MetaboAnalystR 3.0: Towards an Optimized Workflow for Global Metabolomics;

MetaboAnalystR 2.0: From Raw Spectra to Biological Insights; MetaboAnalystR 4.0: a unified LC-MS/MS workflow for global metabolomics

MetaboAnalystR
1.0: flexible and reproducible analysis of metabolomics data;

MetaboAnalyst 6.0 Modules

Input Data Type	Available Modules (click on a module to proceed, or scroll down to explore a total of 18 modules including utilities)				
LC-MS Spectra (mzML, mzXML or mzData)			Spectra Processing [LC-MS w/wo MS2]		
MS Peaks (peak list or intensity table)		Peak Annotation [MS2-DDA/DIA]	Functional Analysis [LC-MS]	Functional Meta-analysis [LC-MS]	
Generic Format (.csv or .txt table files)	Statistical Analysis [one factor]	Statistical Analysis [metadata table]	Biomarker Analysis	Statistical Meta-analysis	Dose Response Analysis
Annotated Features (metabolite list or table)		Enrichment Analysis	Pathway Analysis	Network Analysis	
Link to Genomics & Phenotypes (metabolite list)			Causal Analysis [Mendelian randomization]		

Resources for this workshop

Slides & materials:

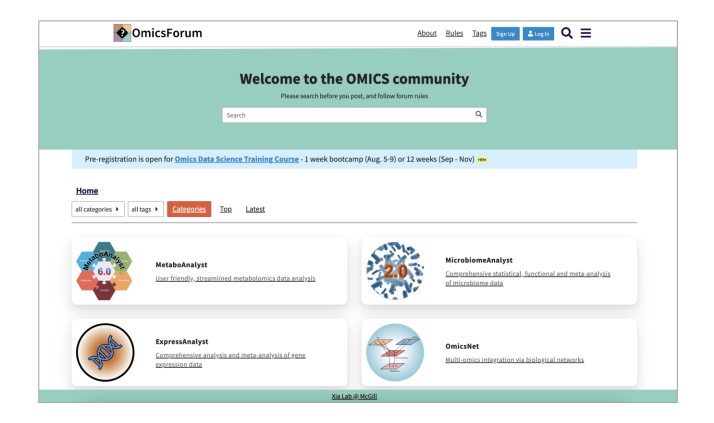
https://github.com/xia-lab/Metabolomics 2024

MetaboAnalyst servers:

- https://www.metaboanalyst.ca
- ➤ https://new.metaboanalyst.ca
- https://dev.metaboanalyst.ca

User Forum:

https://omicsforum.ca



Cautions & recommendations

- **➤ Do not open multiple tabs!**
 - > Results will overwrite each other!
- > Form a group and share computers
 - Make new friends
 - Help each other & reduce stress
- ➤ Page display may be **slow** due to bandwidth limitation
 - ➤ Be patient
 - ➤ Reduce bandwidth consumption by forming group