Mass Spectrometry-based Multi-Omics: Combined Studies of Proteomics, Metabolomics, and/or Lipidomics

This manuscript (<u>permalink</u>) was automatically generated from <u>jessegmeyerlab/2022-multi-omics-review@b5c6d72</u> on January 6, 2022.

Authors

•	Yuming Jiang
	Department of Something, University of Whatever; Department of Whatever, University of Something

- Quinn Dickinson

Department of Something, University of Whatever; Department of Whatever, University of Something

- Amanda Momenzadeh
 - © 0000-0002-8614-0690 · © amandamomenzadeh
 Department of Biochemistry, Medical College of Wisconsin

Department of Biochemistry, Medical College of Wisconsin · Funded by Grant R21 AG074234; Grant R35 GM142502

Abstract

Studies that integrate unbiased measurements across at least two omics layers are often referred to as "multiomics". Measurable "omes" include the genome, transcriptome, proteome, and metabolome. Any combination of omic measures can be referred to as multiomics; for the scope of this review we focus on research combining proteomics and metabolomics. Mass spectrometry is the leading technique for analysis of the proteome and the metabolome. Due to improvements in sample preparation and data collection, more studies are incorporating both mass spectrometry-based proteomics and metabolomics. In this review, we discuss the perceived value of multiomics, advances in sample preparation and data collection, the current state of multiomic data integration, and clinical examples of multiomic analysis. Finally, we explore major barriers preventing democratization of mass spectrometry based multiomics to the same level as nucleic acid analysis, and we suggest solutions to break these barriers.

Introduction

A major goal of biomedical research is understanding how changes in biomolecule compositions of cells and tissues lead to disease phenotypes. The genome serves as a library of possible transcripts, some of which are instructions for proteins, and proteins act on metabolites. The genome thus indirectly determines the set of possible cellular phenotypic states, and the exact cellular state at any given time depends on interaction between the genome with endogenous and exogenous environmental cues. To understand how the genome connects to phenotype, measurement of the genome must be accompanied by measurement of downstream layers of the central dogma of biochemistry (Figure 1?).

Most RNAs encode for proteins, and aside from noncoding RNA and ribozymes, most of the potential functions encoded in a genome are carried out by proteins. Although extensive research has nearly commoditized RNA measurment or transcriptomics [1], multiple studies have found poor correlation between proteins and their corresponding mRNA [2]. Therefore, functional understanding of a biological state demands quantitations of all proteins that are present.

Large scale study of the proteins in a biological system is known as proteomics [3]. Proteomics originally meant large scale studies of protein quantities [4], but it has grown to include large scale studies of protein structure[5] [6] [7], protein-small molecule interactions[8], and protein-protein interactions [9].. Proteomics is increasingly applied to clinical studies to gain understanding of disease mechansims in humans [10].

Many argue that quantities of proteins are not sufficient to enable understanding and dissection of biological systems. The presence of a protein does not necessarily equal the presence of that protein's function. Protein functions are regulated in many ways, including allosteric feedback by metabolites and post-translational modifications such as phosphorylation, acylation, or proteolysis [11]. Roughly 2,700 cellular proteins are predicted to be enzymes [12], many of which act on metabolites that are intermediates required for cell proliferation or energy production.

The large scale measurement of the cellular metabolite pools is known as metabolomics [13], which is sometimes further separated into polar metabolomics and lipidomics [14]. Due to the chemical heterogeneity of metabolites, metabolomics studies use a variety of different separation and mass spectrometry (MS) techniques, including both gas chromatography (GC) [15] and liquid chromatography (LC)[16]. The field of metabolomics is trending toward more use of LC-MS over NMR [17].

Measurement of multiple omic layers is know as multiomics. MS measures peptides (or intact proteins [18]) for proteomic analysis, but MS also measures metabolites for metabolomic analysis. Increasingly studies use MS to measure both proteomes and metabolomes from the same sample. Although multiomics can refer to measuring at least any two omes, throughout this review we focus on multiomic measurements between proteomes and metabolomes.

Integrating data from multiomics should provide more information about the cellular state than the sum of each dataset. Multiomic data integration methods have been developed with different classes of goals. More work is needed to develop methods that take advantage of multiomic data to discover new biological insights into how systems work.

Previous reviews have covered related topics. A recent review surveyed multiomic studies in mitochondria [19] with a focus on sample preparation, qualitity control, methods to collect MS data, and linking measured molecules to phenotypes of respirometry. Another review gives an overview of

each omic method, considerations before performing a multiomic method, and multiomic integration [20]. Misra *et al.* reviewed multiomics with a focus on multiomic integration tools [21]. A metabolomics-centered review was recently published that discusses multiomic integration [22].

In this review we focus on the following topics with a focus on MS-based multiomics studies that combine proteomics and metabolomics:

- sample prepartion methods
- data collection methods
- data integration
- applications to model organisms
- applications to clinical studies

List of Planned Figures:

1. overview of how omic layers are related showing different 'flavors' of each omic analysis

genomics: transcription factor binding (chip-seq), long range structure (ATAC seq?), etc transcriptomics: microarrays, RNA-seq, long read nanopore or pacbio proteomics: interactomics, structural proteomics (thermal proteome profiling or CETSA), PTMomics metabolomics: polar metabolomics, lipidomics,

- 2. Workflow for typical MS-based multi-omics. from sample preparation, extraction to data collection, data intergration and processing.
- 3. Table of clinical multiomics studies
- 4. Figure showing concepts in multiomics integration?

5.

Sample Preparation for Multi-Omic Analysis

The basics: As with any methodology, the very first but also a crucial step for analytical success is proper sample preparation. This is particularly vital for mass spectrometry-based multi-omic analysis, the selected sample preparation strategy is a key determinant for information that will be obtained. Given the great diversity and complexity of biological samples being tested in proteomics and metabolomics, even subtle differences in sample preparation methods can have profound effects on the types of molecules being extracted, which further affect the retention time, signal stability and ionization efficiency. For example, analysis of different types of biomolecules requires specific sample extraction procedures. The most efficient extraction protocols often only cover a restricted type of biomolecules due to their different physicochemical properties. After any necessary dissection or collection, biological tissue samples require a step to

1, Sample preparation for proteomics

Protein preparation for MS analysis can be accomplished by many methods, so it is important to understand the steps leading to analysis. While intact proteins are typically studied by gel electrophoresis, the most common mass spectrometry workflows for complex protein samples analyze peptides, which are easier than proteins to fractionate by LC. Peptides also ionize and fragment more efficiently than whole proteins, resulting in spectra that are easier to interpret for protein identification. Peptide preparation involves reduction and alkylation of cysteines, digestion of

the sample into peptides, desalting and concentration of the peptides and final analysis of these peptides by ionization (e.g., ESI) plus orbitrap-based MS.

- 2, Sample preparation for metabolomics
- 2.1 non-targeted metabolomics

[23]

- 2.2 targeted metabolomics
- 2.3 lipidomics

[24]

3, Integrative sample prepatation for multi-omics

In the context of multi-omics analyses, being able to perform multiple measurements on the same sample can also decrease experimental variation. In this section, we will review several integrative sample preparation technical advances that increased the capabilities of multiomic analysis. One important goal is to decrease the variation between different samples, ultimately to .

<u>25</u>

<u> 26</u>

- -"An Integrated, High-Throughput Strategy for Multiomic Systems Level Analysis" [27]
- -"Detergent-Free Simultaneous Sample Preparation Method for Proteomics and Metabolomics" [28]
- -"MPLEx: a Robust and Universal Protocol for Single-Sample Integrative Proteomic, Metabolomic, and Lipidomic Analyses" [29]
- -"Single-platform 'multi-omic' profiling: unified mass spectrometry and computational workflows for integrative proteomics–metabolomics analysis" [30]
- -"Three-in-One Simultaneous Extraction of Proteins, Metabolites and Lipids for Multi-Omics" [31]

-"

New developments of mass spectrometry-based methods for multi-omics

- 4.1 proteomics
- 4.1.1 Traditional standard methods for proteomics

(Remember to mention here)

4.1.2 Direct infusion methods for proteome analysis (high-throughput methods)

For current proteomic analysis methods, time-consuming chromatographic separation (typically requiring 30–60 min per sample or even longer) is required to protect the coverage, repeatiability, robustness and quantification ability. However, with the rapid application of multiomics results in drug development, biomarker discovery studies and clinical diagnosis. High-throughput methods is highly desirable to boost these fields forward. [32]

(as a high-throughput method, MALDI based proteome analysis should be mentioned here, for example, the application of MALDI for identification of species of bacteria and fungas through their specific peptides)

4.2 metabolomics

To accurately and reliably interpret data derived from metabolomics and lipidomics studies, enormous mass spectrometry based methods were developed during the past decades. (remember to mention the application of MALDI for metabolites analysis, although the drawback of MALDI-tof is obvious.(

Drawbacks: 1,the background of organic matrix in the low molecular weight region 2,the obtained information of MALDI is still very limited, no more than 300 identified metabolites, and also quantification is difficult.

3,as a non-consistent ion source, currently TOF is the typical mass analyser for MALDI, which still suffers from relative low resolution. FTICR can connect MALDI)

4.2.2 Direct infusion mass spectrometry methods for high-throughput analysis of metabolites.()

(direct infusion and so called flow injection MS. do not know the differences, seems saying the same thing.) In typical metabolomic analysis workflow, MS is typically coupled with liquid or gas chromatography, enables separating metabolites within a complex sample and further increases the overall sensitivity by minimizing the ion suppression effect. However, this approach also bring for metabolomics analysis is that chromatographic separation is time-consuming (20–60 min per sample), preventing its application for high-throughput metabolomic screens of large scale samples that required for biomarker discovery studies and clinial analysis.

<u>[24]</u>

<u>[24]</u>

4.4 Integrated Strategy for Mass Spectrometry-Based Multiomics

Mass spectrometry (MS) serves as the centerpiece technology for proteome and metabolome analysis. To gain a better understanding of the multifaceted networks of myriad actions in complex organisms, integration of different multiomic layers is increasingly explored such as joint methods of different omics.

"Multiomics Method Enabled by Sequential Metabolomics and Proteomics for Human Pluripotent Stem-Cell-Derived Cardiomyocytes" [doi?]

[33]

[34, =pdf]

Multi-Omic Data Integration

Argonaut data integration manuscript - [36]

"Integration strategies of multi-omics data for machine learning analysis" [37]

Split strategies into five categories Early integration: Place all omics datasets into single matrix tends to focus on largest dataset, missing data from other omic datasets [38] simple, easy implementation, allows ML models to learn interactions between layers Mixed integration: Map each dataset into new representation befrooe combining Intermediate integration: Transform datasets into common and omic-specific representations Late Integration: analyze each omic layer separately and combine final predictions hierachical integration: base integration of datasets on regulatory relationships between omic layers

"Multi-omics data integration considerations and study design for biological systems and disease" [39]

mixomics R tools [40/journal.pcbi.1005752]

"Multi-omics Data Integration, Interpretation, and Its Application" [41]

Mass Spectrometry-based Multi-Omics Applied to Model Organisms

"Multiomics Method Enabled by Sequential Metabolomics and Proteomics for Human Pluripotent Stem-Cell-Derived Cardiomyocytes" [42]:

Multi-omics Reveal Specific Targets of the RNA-Binding Protein Puf3p and Its Orchestration of Mitochondrial Biogenesis [43]

Mitochondrial protein functions elucidated by multiomic mass spectrometry [44]

Multi-omic mitoprotease profiling reveals role for oct1p [45]

"An integrative systems genetic analysis of mammalian lipid metabolism" [46] * proteomics and lipidomics of mouse liver across 107 genetically different strains *

Clinical applications of multi-omics (proteomics, metabolomics, and/or lipidomics)

Literature currently focused on proteogenomics (integration of genome, transcriptome, and proteome) "Clinical multi-omics strategies for the effective cancer management" [47]

Holistic approach has the ability to improve prognostics and predictive accuracy of disease phenotypes to aid in better treatment and prevention "Multi-omics Data Integration, Interpretation, and Its Application" [48]

Approaches in cancer research:

- "Integration of Proteomics and Metabolomics Revealed Metabolite-Protein Networks in ACTH-Secreting Pituitary Adenoma" [47]: gas chromatography-mass spectrometry (GC-MS) for metabolomics, plus nano liquid chromatography tandem-mass spectrometry(nanoLC-MS/MS) proteomics. Using metabolomic and proteomic data to identify signaling pathways important in metabolic regulation of tumorigenesis. Reveal biomarkers for disease diagnosis, monitoring and therapeutic targets. Looking for changes in tumor on the pituitary gland (adrenocorticotropic hormone-secreting pituitary adenomas (ACTH-PA)). Downregulated glycolysis and fatty acid synthesis. Myc signaling pathway significantly participated in the metabolic changes and tumorigenesis of ACTH-PA.
- "L-Arginine Modulates T Cell Metabolism and Enhances Survival and Anti-tumor Activity" [49]: investigation of proteomic and metabolic adaptations underlying CD4+ T cell activation using MS. Proteins were analyzed by liquid chromatography-coupled mass spectrometry (LC-MS). In parallel, polar metabolites were analyzed by non-targeted flow-injection metabolomics (semi-quantitative method that allows rapid and deep profiling of metabolites. limitations: isobaric compounds cannot be discriminated + possible in-source degradation). Findings: increased L-arginine levels has effects on T cell activation, differentiation, and function.
- "Concurrent lipidomics and proteomics on malignant plasma cells from multiple myeloma patients:
 Probing the lipid metabolome" [50] comparing targeted and untargeted lipidomics and proteomics
 between relapsed and newly diagnosed MM patients
- "Proteometabolomics of bladder cancer: Current and future prospects" [doi:10.3233/CBM-150479?]: simultaneous proteome and metabolome data from urine and blood for urinary bladder cancer patient surveillance to aid in early detection of bladder cancer
- "Grade-Dependent Metabolic Reprogramming in Kidney Cancer Revealed by Combined Proteomics and Metabolomics Analysis" [51]

Approaches in COVID-19 research:

- "Proteomic and Metabolomic Characterization of COVID-19 Patient Sera" [52]: proteomic and metabolomic profiling of serum from 46 COVID-19 patients. Stable isotope labeled proteomics TMTpro (16plex) and and ultra performance liquid chromatography/tandem mass spectrometry (UPLC-MS/MS) untargeted metabolomics to analyze samples. Dysregulation of macrophage, platelet degranulation, complement system, and metabolic suppression in severe COVID-19 patients, useful for selection of biomarkers for severity evaluation. Implemented machine learning model using expression levels of 22 serum proteins and 7 metabolites with overall accuracy of 93.5% in the training set. *Large-Scale Multi-omic Analysis of COVID-19 Severity: [53]: investigation of proteins, lipids, and metabolites associated with COVID-19 status and severity in pts admitted with moderate to severe respiratory issues presumably related to SARS-CoV-2 infection.
- "Metabolomic/lipidomic profiling of COVID-19 and individual response to tocilizumab" [54]
- "Multi-Omics Resolves a Sharp Disease-State Shift between Mild and Moderate COVID-19" [55]
- "Multi-Omics integration analysis of respiratory specimen characterizes baseline molecular determinants associated with COVID-19 diagnosis" [56]
- "Longitudinal multi-omics analysis identifies responses of megakaryocytes, erythroid cells and plasmablasts as hallmarks of severe COVID-19 trajectories" [57]

Misc disease states:

• "Integration of metabolomics and proteomics in multiple sclerosis: From biomarkers discovery to personalized medicine" [58]: majority of biomarkers for MS are not sensitive and/or specific enough to be used for population screening. From the 188 proposed biomarker candidates for MS found in CSF, only 10 (5%) have been successfully verified, while 20 have been falsified. This is likely due to the inter-individual variation. Integrating omics approaches may help with this pitfall; proteomics in combination with the other "omics" may synergize to proide a more precise

- spectrum of information capable of better characterizing a phenotype. When proteins alone, metabolites alone and proteins and metabolites together were uploaded into the ingenuity pathway analysis (IPA) tool, number of matched molecules for "proteins and metabolites" is higher than the sum of molecules matched in the separate searches.
- "High-throughput mediation analysis of human proteome and metabolome identifies mediators of post-bariatric surgical diabetes control"[59] patients with type 2 diabetes randomized to RYGB vs nonsurgical diabetes/weight management; fasting plasma proteome and metabolome were assayed up to 3 years. Plasma proteome profiling was performed using the high-throughput DNA aptamer-based SOMAscan assay platform. Plasma metabolomics were profiled using a commercial semi-quantitative mass spectrometry-based platform (Metabolon, Inc.) using UPLC-MS/MS. Main mediator of improved glycemic control was GHR, which was reduced 3 months after RYGB. GH signaling = diabetogenic effects.

References

1. RNA-Seq: a revolutionary tool for transcriptomics

Zhong Wang, Mark Gerstein, Michael Snyder

Nature Reviews Genetics (2009-01) https://doi.org/crrz3v

DOI: https://doi.org/10.1038/nrg2484

2. Insights into the regulation of protein abundance from proteomic and transcriptomic analyses

Christine Vogel, Edward M Marcotte

Nature Reviews Genetics (2012-04) https://doi.org/bg9g

DOI: https://doi.org/10.1038/nrg3185

3. Mass-spectrometric exploration of proteome structure and function

Ruedi Aebersold, Matthias Mann

Nature (2016-09) https://doi.org/f83zqm

DOI: https://doi.org/10.1038/nature19949

4. Quantitative proteomics: challenges and opportunities in basic and applied research

Olga T Schubert, Hannes L Röst, Ben C Collins, George Rosenberger, Ruedi Aebersold

Nature Protocols (2017-07) https://doi.org/gbhfhm

DOI: 10.1038/nprot.2017.040 · PMID: 28569762

5. Measuring protein structural changes on a proteome-wide scale using limited proteolysis-coupled mass spectrometry

Simone Schopper, Abdullah Kahraman, Pascal Leuenberger, Yuehan Feng, Ilaria Piazza, Oliver Müller, Paul J Boersema, Paola Picotti

Nature Protocols (2017-11) https://doi.org/gcf7zn

DOI: <u>10.1038/nprot.2017.100</u> · PMID: <u>29072706</u>

6. Unlocking the secrets to protein-protein interface drug targets using structural mass spectrometry techniques

Angela Dailing, Alessandra Luchini, Lance Liotta

Expert Review of Proteomics (2015-09-03) https://doi.org/f74gnh

DOI: 10.1586/14789450.2015.1079487 · PMID: 26400464 · PMCID: PMC4858316

7. Fast photochemical oxidation of proteins (FPOP): A powerful mass spectrometry-based structural proteomics tool

Danté T Johnson, Luciano H Di Stefano, Lisa M Jones

Journal of Biological Chemistry (2019-08) https://doi.org/ghzz8z

DOI: <u>10.1074/jbc.rev119.006218</u> · PMID: <u>31262727</u> · PMCID: <u>PMC6690683</u>

8. Drug Target Identification in Tissues by Thermal Proteome Profiling

André Mateus, Nils Kurzawa, Jessica Perrin, Giovanna Bergamini, Mikhail M Savitski *Annual Review of Pharmacology and Toxicology* (2022-01-06) https://doi.org/gmwmqn
DOI: 10.1146/annurev-pharmtox-052120-013205 · PMID: 34499524

- 9. https://doi.org/10.1016/j.coisb.2017.04.015
- 10. Clinical applications of quantitative proteomics using targeted and untargeted dataindependent acquisition techniques.

Jesse G Meyer, Birgit Schilling

Expert review of proteomics (2017-05) https://www.ncbi.nlm.nih.gov/pubmed/28436239

DOI: <u>10.1080/14789450.2017.1322904</u> · PMID: <u>28436239</u> · PMCID: <u>PMC5671767</u>

11. Post-translational modification: nature's escape from genetic imprisonment and the basis for dynamic information encoding: Information encoding by post-translational modification

Sudhakaran Prabakaran, Guy Lippens, Hanno Steen, Jeremy Gunawardena Wiley Interdisciplinary Reviews: Systems Biology and Medicine (2012-11) https://doi.org/f39h4k DOI: 10.1002/wsbm.1185 · PMID: 22899623 · PMCID: PMCID: PMC3473174

12. [No title found]

Pedro Romero, Jonathan Wagg, Michelle L Green, Dale Kaiser, Markus Krummenacker, Peter D Karp

Genome Biology (2004) https://doi.org/fmkcgb

DOI: <u>10.1186/gb-2004-6-1-r2</u> · PMID: <u>15642094</u> · PMCID: <u>PMC549063</u>

13. Mass spectrometry-based metabolomics in health and medical science: a systematic review

Xi-wu Zhang, Qiu-han Li, Zuo-di Xu, Jin-jin Dou *RSC Advances* (2020) https://doi.org/gnxx6f
DOI: https://doi.org/10.1039/c9ra08985c

14. Mass spectrometry-based shotgun lipidomics – a critical review from the technical point of view

Fong-Fu Hsu

Analytical and Bioanalytical Chemistry (2018-10) https://doi.org/gd86z9

DOI: https://doi.org/10.1007/s00216-018-1252-y

15. Metabolomics by Gas Chromatography-Mass Spectrometry: Combined Targeted and Untargeted Profiling.

Oliver Fiehn

Current protocols in molecular biology (2016-04-01)

https://www.ncbi.nlm.nih.gov/pubmed/27038389

DOI: 10.1002/0471142727.mb3004s114 · PMID: 27038389 · PMCID: PMC4829120

16. Separation strategies for untargeted metabolomics

Gary | Patti

Journal of Separation Science (2011-12) https://doi.org/b5zd3w

DOI: 10.1002/jssc.201100532 · PMID: 21972197

17. Novel technologies for metabolomics: More for less

Paul Miggiels, Bert Wouters, Gerard JP van Westen, Anne-Charlotte Dubbelman, Thomas Hankemeier

TrAC Trends in Analytical Chemistry (2019-11) https://doi.org/gnx98m

DOI: <u>10.1016/j.trac.2018.11.021</u>

18. High-throughput quantitative top-down proteomics

Kellye A Cupp-Sutton, Si Wu

Molecular Omics (2020) https://doi.org/gnx98p

DOI: <u>10.1039/c9mo00154a</u> · PMID: <u>31932818</u> · PMCID: <u>PMC7529119</u>

19. Mass-spectrometric multi-omics linked to function – State-of-the-art investigations of mitochondria in systems medicine

TrAC Trends in Analytical Chemistry

(2019-10-01) https://www.sciencedirect.com/science/article/pii/S0165993619303668

DOI: 10.1016/j.trac.2019.115635

20. Multi-omics approaches to disease

Yehudit Hasin, Marcus Seldin, Aldons Lusis

Genome Biology (2017-05-05) https://doi.org/10.1186/s13059-017-1215-1

DOI: 10.1186/s13059-017-1215-1

21. Integrated omics: tools, advances and future approaches

Biswapriya B Misra, Carl Langefeld, Michael Olivier, Laura A Cox *Journal of Molecular Endocrinology* (2019-01) https://doi.org/gfx8hg

DOI: https://doi.org/10.1530/jme-18-0055

22. Multi-omics integration in biomedical research – A metabolomics-centric review

Maria A Wörheide, Jan Krumsiek, Gabi Kastenmüller, Matthias Arnold

Analytica Chimica Acta (2021-01) https://doi.org/gmsz75

DOI: https://doi.org/10.1016/j.aca.2020.10.038

23. Development of a plasma pseudotargeted metabolomics method based on ultra-high-performance liquid chromatography-mass spectrometry

Fujian Zheng, Xinjie Zhao, Zhongda Zeng, Lichao Wang, Wangjie Lv, Qingqing Wang, Guowang Xu

Nature Protocols (2020-08) https://www.nature.com/articles/s41596-020-0341-5

DOI: <u>10.1038/s41596-020-0341-5</u>

24. A complete workflow for high-resolution spectral-stitching nanoelectrospray direct-infusion mass-spectrometry-based metabolomics and lipidomics

Andrew D Southam, Ralf JM Weber, Jasper Engel, Martin R Jones, Mark R Viant *Nature Protocols* (2017-02) https://www.nature.com/articles/nprot.2016.156

DOI: 10.1038/nprot.2016.156

25. Multiomic analysis of a dried single-drop plasma sample using an integrated mass spectrometry approach

Weina Gao, Qiaoyun Zhang, Yiran Su, Peiwu Huang, Xue Lu, Qinyue Gong, Wendong Chen, Ruilian Xu, Ruijun Tian

Analyst (2020-10-12) https://pubs.rsc.org/en/content/articlelanding/2020/an/d0an01149e

DOI: 10.1039/d0an01149e

26. https://doi.org/10.3389/fgene.2021.635971

27. An Integrated, High-Throughput Strategy for Multiomic Systems Level Analysis

Danielle B Gutierrez, Randi L Gant-Branum, Carrie E Romer, Melissa A Farrow, Jamie L Allen, Nikesh Dahal, Yuan-Wei Nei, Simona G Codreanu, Ashley T Jordan, Lauren D Palmer, ... Richard M Caprioli

Journal of Proteome Research (2018-10-05) https://doi.org/gfhjtv

DOI: <u>10.1021/acs.jproteome.8b00302</u> · PMID: <u>30114907</u> · PMCID: <u>PMC6469931</u>

28. Detergent-Free Simultaneous Sample Preparation Method for Proteomics and Metabolomics

Alexandre Zougman, John P Wilson, Lee D Roberts, Rosamonde E Banks *Journal of Proteome Research* (2020-07-02) https://doi.org/10.1021/acs.jproteome.9b00662

29. MPLEx: a Robust and Universal Protocol for Single-Sample Integrative Proteomic, Metabolomic, and Lipidomic Analyses

Ernesto S Nakayasu, Carrie D Nicora, Amy C Sims, Kristin E Burnum-Johnson, Young-Mo Kim, Jennifer E Kyle, Melissa M Matzke, Anil K Shukla, Rosalie K Chu, Athena A Schepmoes, ... Thomas O Metz

mSystems (2016-06-28) https://doi.org/gk7gvg

DOI: https://doi.org/10.1128/msystems.00043-16

30. Single-platform 'multi-omic' profiling: unified mass spectrometry and computational workflows for integrative proteomics-metabolomics analysis

Benjamin C Blum, Fatemeh Mousavi, Andrew Emili

Molecular Omics (2018) https://doi.org/ghcv4s

DOI: <u>10.1039/c8mo00136g</u> · PMID: <u>30211418</u>

31. Three-in-One Simultaneous Extraction of Proteins, Metabolites and Lipids for Multi-Omics

Jianing Kang, Lisa David, Yangyang Li, Jing Cang, Sixue Chen

Frontiers in Genetics (2021-04-15) https://doi.org/gnxfsb

DOI: https://doi.org/10.3389/fgene.2021.635971

32. Quantitative shotgun proteome analysis by direct infusion

Jesse G Meyer, Natalie M Niemi, David J Pagliarini, Joshua J Coon

Nature Methods (2020-12) https://www.nature.com/articles/s41592-020-00999-z

DOI: 10.1038/s41592-020-00999-z

33. An Integrated Strategy for Mass Spectrometry-Based Multiomics Analysis of Single Cells

Yuanyuan Li, Hang Li, Yuping Xie, Shuo Chen, Ritian Qin, Hangyan Dong, Yongliang Yu, Jianhua Wang, Xiaohong Qian, Weijie Qin

Analytical Chemistry (2021-10-13) https://pubs.acs.org/doi/abs/10.1021/acs.analchem.0c05209

DOI: 10.1021/acs.analchem.0c05209

34. Multi-Omic Single-Shot Technology for Integrated Proteome and Lipidome Analysis

Yuchen He, Edrees H Rashan, Vanessa Linke, Evgenia Shishkova, Alexander S Hebert, Adam Jochem, Michael S Westphall, David J Pagliarini, Katherine A Overmyer, Joshua J Coon *Analytical Chemistry* (2021-02-22) https://pubs.acs.org/doi/abs/10.1021/acs.analchem.0c04764
DOI: 10.1021/acs.analchem.0c04764

35. An Integrated Strategy for Mass Spectrometry-Based Multiomics Analysis of Single Cells

Yuanyuan Li, Hang Li, Yuping Xie, Shuo Chen, Ritian Qin, Hangyan Dong, Yongliang Yu, Jianhua Wang, Xiaohong Qian, Weijie Qin

Analytical Chemistry (2021-10-13) https://pubs.acs.org/doi/abs/10.1021/acs.analchem.0c05209

DOI: <u>10.1021/acs.analchem.0c05209</u>

36. https://doi.org/10.1016/j.patter.2020.100122

37. Integration strategies of multi-omics data for machine learning analysis.

Milan Picard, Marie-Pier Scott-Boyer, Antoine Bodein, Olivier Périn, Arnaud Droit *Computational and structural biotechnology journal* (2021-06-22)

https://www.ncbi.nlm.nih.gov/pubmed/34285775

DOI: <u>10.1016/j.csbj.2021.06.030</u> · PMID: <u>34285775</u> · PMCID: <u>PMC8258788</u>

38. Integration of clinical chemistry, expression, and metabolite data leads to better toxicological class separation.

Jeppe S Spicker, Søren Brunak, Klaus S Frederiksen, Henrik Toft

Toxicological sciences: an official journal of the Society of Toxicology (2008-01-04)

https://www.ncbi.nlm.nih.gov/pubmed/18178960

DOI: 10.1093/toxsci/kfn001 · PMID: 18178960

39. Multi-omics data integration considerations and study design for biological systems and disease.

Stefan Graw, Kevin Chappell, Charity L Washam, Allen Gies, Jordan Bird, Michael S Robeson, Stephanie D Byrum

Molecular omics (2021-04-19) https://www.ncbi.nlm.nih.gov/pubmed/33347526 DOI: 10.1039/d0mo00041h · PMID: 33347526 · PMCID: PMC8058243

40. https://journals.plos.org/ploscompbiol/article?id

41. Multi-omics Data Integration, Interpretation, and Its Application

Indhupriya Subramanian, Srikant Verma, Shiva Kumar, Abhay Jere, Krishanpal Anamika Bioinformatics and Biology Insights (2020-01) https://doi.org/ggn3c8 DOI: https://doi.org/10.1177/1177932219899051

42. Multiomics Method Enabled by Sequential Metabolomics and Proteomics for Human **Pluripotent Stem-Cell-Derived Cardiomyocytes**

Elizabeth F Bayne, Aaron D Simmons, David S Roberts, Yanlong Zhu, Timothy J Aballo, Benjamin Wancewicz, Sean P Palecek, Ying Ge

Journal of Proteome Research (2021-10-01)

https://pubs.acs.org/doi/10.1021/acs.jproteome.1c00611

DOI: <u>10.1021/acs.jproteome.1c00611</u>

Multi-omics Reveal Specific Targets of the RNA-Binding Protein Puf3p and Its 43. **Orchestration of Mitochondrial Biogenesis.**

Christopher P Lapointe, Jonathan A Stefely, Adam Jochem, Paul D Hutchins, Gary M Wilson, Nicholas W Kwiecien, Joshua J Coon, Marvin Wickens, David J Pagliarini Cell systems (2017-12-13) https://www.ncbi.nlm.nih.gov/pubmed/29248374 DOI: <u>10.1016/j.cels.2017.11.012</u> · PMID: <u>29248374</u> · PMCID: <u>PMC5799006</u>

44. Mitochondrial protein functions elucidated by multi-omic mass spectrometry profiling. Jonathan A Stefely, Nicholas W Kwiecien, Elyse C Freiberger, Alicia L Richards, Adam Jochem, Matthew JP Rush, Arne Ulbrich, Kyle P Robinson, Paul D Hutchins, Mike T Veling, ... Joshua J Coon

Nature biotechnology (2016-09-26) https://www.ncbi.nlm.nih.gov/pubmed/27669165 DOI: 10.1038/nbt.3683 · PMID: 27669165 · PMCID: PMC5101133

45. Multi-omic Mitoprotease Profiling Defines a Role for Oct1p in Coenzyme Q Production.

Mike T Veling, Andrew G Reidenbach, Elyse C Freiberger, Nicholas W Kwiecien, Paul D Hutchins, Michael J Drahnak, Adam Jochem, Arne Ulbrich, Matthew JP Rush, Jason D Russell, ... David J **Pagliarini**

Molecular cell (2017-12-07) https://www.ncbi.nlm.nih.gov/pubmed/29220658 DOI: 10.1016/j.molcel.2017.11.023 · PMID: 29220658 · PMCID: PMC5730362

An integrative systems genetic analysis of mammalian lipid metabolism. 46.

Benjamin L Parker, Anna C Calkin, Marcus M Seldin, Michael F Keating, Elizabeth J Tarling, Pengyi Yang, Sarah C Moody, Yingying Liu, Eser J Zerenturk, Elise J Needham, ... Brian G Drew Nature (2019-02-27) https://www.ncbi.nlm.nih.gov/pubmed/30814737 DOI: <u>10.1038/s41586-019-0984-y</u> · PMID: <u>30814737</u> · PMCID: <u>PMC6656374</u>

https://doi.org/10.3389/fendo.2018.00678 47.

48. Multi-omics Data Integration, Interpretation, and Its Application:

Indhupriya Subramanian, Srikant Verma, Shiva Kumar, Abhay Jere, Krishanpal Anamika Bioinformatics and Biology Insights (2020-01-31)

https://journals.sagepub.com/doi/10.1177/1177932219899051

DOI: 10.1177/1177932219899051

L-Arginine Modulates T Cell Metabolism and Enhances Survival and Anti-tumor Activity 49. Roger Geiger, Jan C Rieckmann, Tobias Wolf, Camilla Basso, Yuehan Feng, Tobias Fuhrer, Maria Kogadeeva, Paola Picotti, Felix Meissner, Matthias Mann, ... Antonio Lanzavecchia

Cell (2016-10) https://doi.org/f88qkm

DOI: http://dx.doi.org/10.1016/j.cell.2016.09.031

50. Concurrent lipidomics and proteomics on malignant plasma cells from multiple myeloma patients: Probing the lipid metabolome

Ahmed Mohamed, Joel Collins, Hui Jiang, Jeffrey Molendijk, Thomas Stoll, Federico Torta, Markus R Wenk, Robert J Bird, Paula Marlton, Peter Mollee, ... Michelle M Hill *PLOS ONE* (2020-01-08) https://journals.plos.org/plosone/article?

id=10.1371/journal.pone.0227455

DOI: <u>10.1371/journal.pone.0227455</u>

51. Grade-Dependent Metabolic Reprogramming in Kidney Cancer Revealed by Combined Proteomics and Metabolomics Analysis

Hiromi I Wettersten, AAri Hakimi, Dexter Morin, Cristina Bianchi, Megan E Johnstone, Dallas R Donohoe, Josephine F Trott, Omran Abu Aboud, Steven Stirdivant, Bruce Neri, ... Robert H Weiss *Cancer Research* (2015-06-15) https://doi.org/f7h8zh

DOI: 10.1158/0008-5472.can-14-1703 · PMID: 25952651 · PMCID: PMC4470795

- 52. https://doi.org/10.1016/j.cell.2020.05.032
- 53. https://doi.org/10.1016/j.cels.2020.10.003
- 54. **Metabolomic/lipidomic profiling of COVID-19 and individual response to tocilizumab**Gaia Meoni, Veronica Ghini, Laura Maggi, Alessia Vignoli, Alessio Mazzoni, Lorenzo Salvati, Manuela Capone, Anna Vanni, Leonardo Tenori, Paolo Fontanari, ... Paola Turano *PLOS Pathogens* (2021-02-01) https://journals.plos.org/plospathogens/article?id=10.1371/journal.ppat.1009243
 DOI: 10.1371/journal.ppat.1009243
- 55. https://doi.org/10.1016/j.cell.2020.10.037
- 56. Multi-Omics integration analysis of respiratory specimen characterizes baseline molecular determinants associated with COVID-19 diagnosis

Jaswinder Singh Maras, Shvetank Sharma, Adil Bhat, Reshu Aggrawal, Ekta Gupta, Shiv K Sarin (2020-07-07) https://www.medrxiv.org/content/10.1101/2020.07.06.20147082v1

57. Longitudinal multi-omics analysis identifies responses of megakaryocytes, erythroid cells and plasmablasts as hallmarks of severe COVID-19 trajectories

Joana P Bernardes, Neha Mishra, Florian Tran, Thomas Bahmer, Lena Best, Johanna I Blase, Dora Bordoni, Jeanette Franzenburg, Ulf Geisen, Jonathan Josephs-Spaulding, ... HCA Lung Biological Network and the Deutsche COVID-19 Omics Initiative (DeCOI) (2020-09-18) https://www.medrxiv.org/content/10.1101/2020.09.11.20187369v1

58. Integration of metabolomics and proteomics in multiple sclerosis: From biomarkers discovery to personalized medicine

Piero Del Boccio, Claudia Rossi, Maria di Ioia, Ilaria Cicalini, Paolo Sacchetta, Damiana Pieragostino

PROTEOMICS – Clinical Applications (2016-04-01)

https://onlinelibrary.wiley.com/doi/10.1002/prca.201500083

DOI: 10.1002/prca.201500083

59. High-throughput mediation analysis of human proteome and metabolome identifies mediators of post-bariatric surgical diabetes control

Jonathan M Dreyfuss, Yixing Yuchi, Xuehong Dong, Vissarion Efthymiou, Hui Pan, Donald C Simonson, Ashley Vernon, Florencia Halperin, Pratik Aryal, Anish Konkar, ... Mary Elizabeth Patti *Nature Communications* (2021-11-29) https://www.nature.com/articles/s41467-021-27289-2

DOI: 10.1038/s41467-021-27289-2