Resource Summary Report

Generated by NIF on May 17, 2025

Data independent acquisition proteomics workbench

RRID:SCR 021862

Type: Tool

Proper Citation

Data independent acquisition proteomics workbench (RRID:SCR_021862)

Resource Information

URL: https://usegalaxy.eu

Proper Citation: Data independent acquisition proteomics workbench (RRID:SCR_021862)

Description: Software tools for analysis of data independent acqusition type proteomics data are implemented in Galaxy framework. Enables complete DIA data analysis including spectral library generation based on DDA type data, analysis of DIA data in raw or mzml format, and differntial statistical analysis. All tools are based on open-source software such as diypasef, OpenSwath, swath2stats and MSstats.

Resource Type: software resource, software toolkit

Defining Citation: DOI:10.1101/2021.07.21.453197

Keywords: data independent acquisition, proteomics data, Galaxy framework,

Funding:

Availability: Free, Freely available

Resource Name: Data independent acquisition proteomics workbench

Resource ID: SCR_021862

Record Creation Time: 20220129T080357+0000

Record Last Update: 20250513T062212+0000

Ratings and Alerts

No rating or validation information has been found for Data independent acquisition proteomics workbench.

No alerts have been found for Data independent acquisition proteomics workbench.

Data and Source Information

Source: SciCrunch Registry

Usage and Citation Metrics

We found 192 mentions in open access literature.

Listed below are recent publications. The full list is available at NIF.

Lengyel M, et al. (2025) Zymogen granule protein 16B (ZG16B) is a druggable epigenetic target to modulate the mammary extracellular matrix. Cancer science, 116(1), 81.

Casella V, et al. (2025) Novel Insights into the Nobilamide Family from a Deep-Sea Bacillus: Chemical Diversity, Biosynthesis and Antimicrobial Activity Towards Multidrug-Resistant Bacteria. Marine drugs, 23(1).

Kunz L, et al. (2025) Avirulence depletion assay: Combining R gene-mediated selection with bulk sequencing for rapid avirulence gene identification in wheat powdery mildew. PLoS pathogens, 21(1), e1012799.

Banar M, et al. (2025) A novel broad-spectrum bacteriophage cocktail against methicillinresistant Staphylococcus aureus: Isolation, characterization, and therapeutic potential in a mastitis mouse model. PloS one, 20(1), e0316157.

Galvis J, et al. (2025) Using DIMet for Differential Analysis of Labeled Metabolomics Data: A Step-by-step Guide Showcasing the Glioblastoma Metabolism. Bio-protocol, 15(2), e5168.

Baei B, et al. (2025) Pharmacophore modeling and QSAR analysis of anti-HBV flavonols. PloS one, 20(1), e0316765.

Hermawaty D, et al. (2025) De novo transcriptome assembly and analysis during agarwood induction in Gyrinops versteegii Gilg. seedling. Scientific reports, 15(1), 2977.

Shaikh MA, et al. (2025) StCDF1: A 'jack of all trades' clock output with a central role in regulating potato nitrate reduction activity. The New phytologist, 245(1), 282.

Petroll R, et al. (2025) Enhanced sensitivity of TAPscan v4 enables comprehensive analysis of streptophyte transcription factor evolution. The Plant journal: for cell and molecular biology, 121(1), e17184.

Strateva TV, et al. (2025) First Detection and Genomic Characterization of Linezolid-

Resistant Enterococcus faecalis Clinical Isolates in Bulgaria. Microorganisms, 13(1).

Zhang N, et al. (2025) Deciphering the molecular logic of WOX5 function in the root stem cell organizer. The EMBO journal, 44(1), 281.

Dawson RA, et al. (2025) Carbon monoxide-oxidising Pseudomonadota on volcanic deposits. Environmental microbiome, 20(1), 12.

Fraccalvieri R, et al. (2025) Isolation and Characterization of Colistin-Resistant Enterobacteriaceae from Foods in Two Italian Regions in the South of Italy. Microorganisms, 13(1).

Tetzlaff S, et al. (2024) Small RNAs from mitochondrial genome recombination sites are incorporated into T. gondii mitoribosomes. eLife, 13.

Thompson RM, et al. (2024) Draft genome sequences of two Micromonospora strains isolated from the root nodules of Alnus glutinosa. Microbiology resource announcements, 13(3), e0113123.

Martín Del Pico E, et al. (2024) FAIRsoft-a practical implementation of FAIR principles for research software. Bioinformatics (Oxford, England), 40(8).

Vieira Da Cruz A, et al. (2024) Pyridylpiperazine efflux pump inhibitor boosts in vivo antibiotic efficacy against K. pneumoniae. EMBO molecular medicine, 16(1), 93.

Sageman-Furnas K, et al. (2024) Detailing Early Shoot Growth Arrest in Kro-0 x BG-5 Hybrids of Arabidopsis thaliana. Plant & cell physiology, 65(3), 420.

Nugroho A, et al. (2024) Transcriptome dataset of gall-rust infected Sengon (Falcataria falcata) seedlings using long-read PCR-cDNA sequencing. Data in brief, 52, 109919.

Beerling DJ, et al. (2024) Enhanced weathering in the US Corn Belt delivers carbon removal with agronomic benefits. Proceedings of the National Academy of Sciences of the United States of America, 121(9), e2319436121.