

Resource Summary Report

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ENZYME

RRID:SCR_002487

Type: Tool

Proper Citation

ENZYME (RRID:SCR_002487)

Resource Information

URL: <http://www.expasy.org/enzyme/>

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Description: A repository of information relative to the nomenclature of enzymes. It is primarily based on the recommendations of the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (IUBMB) and it describes each type of characterized enzyme for which an EC (Enzyme Commission) number has been provided. These include * EC number * Recommended name * Alternative names (if any) * Catalytic activity * Cofactors (if any) * Pointers to the Swiss-Prot protein sequence entry(ies) that correspond to the enzyme (if any) * Pointers to human disease(s) associated with a deficiency of the enzyme (if any) We believe that the ENZYME database can be useful to anybody working with enzymes and that it can be of help in the development of computer programs involved with the manipulation of metabolic pathways. Available services include downloading ENZYME by FTP as well as report forms for a new ENZYME entry or for an error/update in an existing entry.

Abbreviations: ENZYME

Synonyms: Enzyme nomenclature database

Resource Type: service resource, database, storage service resource, data repository, data or information resource

Defining Citation: [PMID:10592255](#)

Keywords: enzyme commission, enzyme name, nomenclature, enzyme categories, enzyme classification, enzyme nomenclature, enzyme reaction categories, gold standard

Availability: Free, Available for download, Freely available

Resource Name: ENZYME

Resource ID: SCR_002487

Alternate IDs: nif-0000-02803

Alternate URLs: <http://www.expasy.ch/enzyme/>

Record Creation Time: 20220129T080213+0000

Record Last Update: 20260728T043134+0000

Ratings and Alerts

No rating or validation information has been found for ENZYME.

No alerts have been found for ENZYME.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 97 mentions in open access literature.

Listed below are recent publications. The full list is available at [nidm-terms](#) .

Duan C, et al. (2025) FGeneBERT: function-driven pre-trained gene language model for metagenomics. Briefings in bioinformatics, 26(2).

Yang WH, et al. (2025) Ribonuclease 1 Induces T-Cell Dysfunction and Impairs CD8+ T-Cell Cytotoxicity to Benefit Tumor Growth through Hijacking STAT1. Advanced science (Weinheim, Baden-Wurttemberg, Germany), 12(13), e2404961.

Baltsavia I, et al. (2025) MjCyc: Rediscovering the pathway-genome landscape of the first sequenced archaeon, Methanocaldococcus (Methanococcus) jannaschii. iScience, 28(1), 111546.

Esembaeva MA, et al. (2025) Evaluation of Genome-Scale Model Reconstruction Strategies for Lentilactobacillus kefir DH5 and Deciphering Its Metabolic Network. Metabolites, 15(12).

Passin V, et al. (2025) Depletion of macrophages and osteoclast precursors mitigates iron overload-mediated bone loss. IUBMB life, 77(1), e2928.

Levé M, et al. (2025) Metabolomics and metagenomics in mice reveal the role of the gut microbiota in tryptophan metabolism. iScience, 28(11), 113751.

Guerrero-Egido G, et al. (2024) bacLIFE: a user-friendly computational workflow for genome analysis and prediction of lifestyle-associated genes in bacteria. Nature communications, 15(1), 2072.

Cattin-Ortolá J, et al. (2024) Cargo selective vesicle tethering: The structural basis for binding of specific cargo proteins by the Golgi tether component TBC1D23. *Science advances*, 10(13), eadl0608.

Ashour DJ, et al. (2023) Zasp52 strengthens whole embryo tissue integrity through supracellular actomyosin networks. *Development (Cambridge, England)*, 150(7).

Alp M, et al. (2023) Drug screening of α -amylase inhibitors as candidates for treating diabetes. *Journal of cellular and molecular medicine*, 27(15), 2249.

Pavic K, et al. (2023) Structural mechanism for inhibition of PP2A-B56 γ and oncogenicity by CIP2A. *Nature communications*, 14(1), 1143.

Ramírez-Palacios C, et al. (2023) Computational prediction of α -transaminase selectivity by deep learning analysis of molecular dynamics trajectories. *QRB discovery*, 4, e1.

Val-Torregrosa B, et al. (2022) Phosphate-induced resistance to pathogen infection in *Arabidopsis*. *The Plant journal : for cell and molecular biology*, 110(2), 452.

Sardar P, et al. (2022) De novo metatranscriptomic exploration of gene function in the millipede holobiont. *Scientific reports*, 12(1), 16173.

Nursimulu N, et al. (2022) Architect: A tool for aiding the reconstruction of high-quality metabolic models through improved enzyme annotation. *PLoS computational biology*, 18(9), e1010452.

Wang B, et al. (2022) MiR-630 suppresses non-small cell lung cancer by targeting vimentin. *Journal of clinical laboratory analysis*, 36(9), e24536.

Cattin-Ortolá J, et al. (2021) Sequences in the cytoplasmic tail of SARS-CoV-2 Spike facilitate expression at the cell surface and syncytia formation. *Nature communications*, 12(1), 5333.

Guo Y, et al. (2021) TMT-based quantitative proteomic analysis reveals the spleen regulatory network of dexamethasone-induced immune suppression in chicks. *Journal of proteomics*, 248, 104353.

Cheng H, et al. (2021) Full-Length Transcriptome of *Thalassiosira weissflogii* as a Reference Resource and Mining of Chitin-Related Genes. *Marine drugs*, 19(7).

Magalhães P, et al. (2021) Urinary fetuin-A peptides as a new marker for impaired kidney function in patients with type 2 diabetes. *Clinical kidney journal*, 14(1), 269.