

Resource Summary Report

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OrthoMCL DB: Ortholog Groups of Protein Sequences

RRID:SCR_007839

Type: Tool

Proper Citation

OrthoMCL DB: Ortholog Groups of Protein Sequences (RRID:SCR_007839)

Resource Information

URL: <http://www.orthomcl.org/cgi-bin/OrthoMclWeb.cgi>

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Description: OrthoMCL is a genome-scale algorithm for grouping orthologous protein sequences. It provides not only groups shared by two or more species/genomes, but also groups representing species-specific gene expansion families. OrthoMCL starts with reciprocal best hits within each genome as putative in-paralog/recent paralog pairs and reciprocal best hits across any two genomes as putative ortholog pairs. Related proteins are interlinked in a similarity graph. Then MCL (Markov Clustering algorithm, Van Dongen 2000; www.micans.org/mcl) is invoked to split mega-clusters. This process is analogous to the manual review in COG construction. MCL clustering is based on weights between each pair of proteins, so to correct for differences in evolutionary distance the weights are normalized before running MCL.

Synonyms: OrthoMCL DB

Resource Type: data or information resource, database

Keywords: FASEB list

Resource Name: OrthoMCL DB: Ortholog Groups of Protein Sequences

Resource ID: SCR_007839

Alternate IDs: r3d100012462, nif-0000-03230

Alternate URLs: <https://doi.org/10.17616/R3FZ27>

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Record Last Update: 20260829T183008+0000

Ratings and Alerts

No rating or validation information has been found for OrthoMCL DB: Ortholog Groups of Protein Sequences.

No alerts have been found for OrthoMCL DB: Ortholog Groups of Protein Sequences.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 1051 mentions in open access literature.

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

Kwak Y, et al. (2025) A dual insect symbiont and plant pathogen improves insect host fitness under arginine limitation. *mBio*, 16(4), e0358824.

Aspinwall J, et al. (2025) Behavioral Manipulation of *Ixodes scapularis* by *Ehrlichia muris euclairensis*: Implications for Tick-Borne Disease Transmission. *bioRxiv : the preprint server for biology*.

Yang S, et al. (2025) Comparative Genomics Reveals Evidence of the Genome Reduction and Metabolic Potentials of *Alilineobacillus hadale* Isolated from Challenger Deep Sediment of the Mariana Trench. *Microorganisms*, 13(1).

Ros-Lucas A, et al. (2025) Identification of novel compounds against *Trypanosoma cruzi* using AlphaFold structures. *Computational and structural biotechnology journal*, 27, 1838.

Sun R, et al. (2025) Chromosome-level genome assembly of a high-yield Chinese soybean variety Mengdou1137 unlocks genetic potential of disease and lodging resistance. *TAG. Theoretical and applied genetics. Theoretische und angewandte Genetik*, 138(6), 119.

Bian C, et al. (2025) Chromosome-level genome assemblies of five *Sinocyclocheilus* species. *GigaByte (Hong Kong, China)*, 2025, gigabyte155.

Westaway JAF, et al. (2025) Genomic epidemiology of *Plasmodium knowlesi* reveals putative genetic drivers of adaptation in Malaysia. *PLoS neglected tropical diseases*, 19(3), e0012885.

Guo L, et al. (2025) Duckweed: a starch-hyperaccumulating plant under cultivation with a combination of nutrient limitation and elevated CO₂. *Frontiers in plant science*, 16, 1531849.

Chen N, et al. (2025) Genome-wide identification of the AAP gene family and expression analysis under tissue-specific expression in five legumes. *BMC genomics*, 26(1), 173.

Wang Y, et al. (2025) Genome-Driven Functional Validation of *Bacillus amyloliquefaciens* Strain MEPW12: A Multifunctional Endophyte for Sustainable Sweet Potato Cultivation. *Microorganisms*, 13(6).

Salamzade R, et al. (2025) zol and fai: large-scale targeted detection and evolutionary investigation of gene clusters. *Nucleic acids research*, 53(3).

Christel S, et al. (2025) Catabolic pathway acquisition by rhizosphere bacteria readily enables growth with a root exudate component but does not affect root colonization. *mBio*, 16(1), e0301624.

He Y, et al. (2025) Comparative Genomic Analysis of *Campylobacter* Plasmids Identified in Food Isolates. *Microorganisms*, 13(1).

Huang M, et al. (2025) Goose multi-omics database: A comprehensive multi-omics database for goose genomics. *Poultry science*, 104(3), 104842.

Prostak S, et al. (2025) Frog-killing chytrid fungi deploy different strategies to regulate intracellular pressure in cell types that have or lack a cell wall. *bioRxiv : the preprint server for biology*.

Gong L, et al. (2025) Haplotype-resolved genome assembly and genome-wide association study identifies the candidate gene closely related to sugar content and tuber yield in *Solanum tuberosum*. *Horticulture research*, 12(6), uhaf075.

Gu H, et al. (2025) Chromosome-level assemblies of the White bream *Parabramis pekinensis*. *Scientific data*, 12(1), 871.

Salvà-Serra F, et al. (2025) Description of *Pseudomonas imrae* sp. nov., carrying a novel class C β -lactamase gene variant, isolated from gut samples of Atlantic mackerel (*Scomber scombrus*). *Frontiers in microbiology*, 16, 1530878.

Wang Y, et al. (2025) Telomere-to-telomere genome of common bean (*Phaseolus vulgaris* L., YP4). *GigaScience*, 14.

Wang C, et al. (2025) Genomic-transcriptomic analysis identifies the Syrian hamster as a superior animal model for human diseases. *BMC genomics*, 26(1), 286.