

# Resource Summary Report

Generated by [SPARC SAWG](#) on Aug 9, 2026

## CORUM

RRID:SCR\_002254

Type: Tool

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### Proper Citation

CORUM (RRID:SCR\_002254)

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### Resource Information

**URL:** <http://mips.helmholtz-muenchen.de/genre/proj/corum>

**Proper Citation:** CORUM (RRID:SCR\_002254)

**Description:** Database of manually annotated protein complexes from mammalian organisms. Annotation includes protein complex function, localization, subunit composition, literature references and more. All information is obtained from individual experiments published in scientific articles, but data from high-throughput experiments is excluded.

The majority of protein complexes in CORUM originates from man (65%), followed by mouse (14%) and rat (14%)., THIS RESOURCE IS NO LONGER IN SERVICE.  
Documented on September 16,2025.

**Abbreviations:** CORUM

**Synonyms:** CORUM the Comprehensive Resource of Mammalian protein complexes, CORUM - the Comprehensive Resource of Mammalian protein complexes

**Resource Type:** database, data or information resource

**Defining Citation:** [PMID:19884131](#), [PMID:17965090](#)

**Keywords:** mammalian protein, protein, protein complex, protein function, FASEB list

**Funding:** BMBF 031U212C

**Availability:** THIS RESOURCE IS NO LONGER IN SERVICE

**Resource Name:** CORUM

**Resource ID:** SCR\_002254

**Alternate IDs:** nif-0000-02688, OMICS\_01904, r3d100011272

**Alternate URLs:** <http://mips.gsf.de/genre/proj/corum>

**Record Creation Time:** 20220129T080212+0000

**Record Last Update:** 20260808T190352+0000

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## Ratings and Alerts

No rating or validation information has been found for CORUM.

No alerts have been found for CORUM.

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## Data and Source Information

**Source:** [SciCrunch Registry](#)

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## Usage and Citation Metrics

We found 164 mentions in open access literature.

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

Rathore U, et al. (2026) Systematic discovery of pro- and anti-HIV host factors in primary human CD4+ T cells. *Cell*, 189(12), 3651.

Frommelt F, et al. (2025) The solute carrier superfamily interactome. *Molecular systems biology*, 21(6), 632.

Dang V, et al. (2025) VerteBrain reveals novel neural and non-neural protein assemblies conserved across vertebrate evolution. *bioRxiv : the preprint server for biology*.

Liu X, et al. (2025) Mealtime alters daily rhythm in nuclear O-GlcNAc proteome to regulate hepatic gene expression. *bioRxiv : the preprint server for biology*.

Pio-Lopez L, et al. (2025) Universal multilayer network embedding reveals a causal link between GABA neurotransmitter and cancer. *BMC bioinformatics*, 26(1), 149.

Le Minh G, et al. (2025) GATAD2B O-GlcNAcylation Regulates Breast Cancer Stem-like Potential and Drug Resistance. *Cells*, 14(6).

Aç?kgöz MA, et al. (2025) Secondary Metabolites and Biological Activities From *Achillea gypsicola* Hub-Mor. Under Foliar Applications of Chitosan and Salicylic Acid. *Food science & nutrition*, 13(5), e70225.

Soriaga LB, et al. (2025) Shared host genetic landscape of respiratory viral infection. *Proceedings of the National Academy of Sciences of the United States of America*, 122(20), e2414202122.

Salovska B, et al. (2025) A robust multiplex-DIA workflow profiles protein turnover regulations associated with cisplatin resistance and aneuploidy. *Nature communications*,

16(1), 5034.

Kenaston MW, et al. (2025) Yellow Fever Virus Interactomes Reveal Common and Divergent Strategies of Replication and Evolution for Mosquito-borne Flaviviruses. *bioRxiv* : the preprint server for biology.

Girault V, et al. (2025) Multi-proteomic profiling of the varicella-zoster virus-host interface reveals host susceptibilities to severe infection. *Nature microbiology*, 10(8), 2048.

Lu Y, et al. (2025) Decoding the Effects of Bexarotene treatment on brain of AD-like model mice: Single-Cell Transcriptomics and Chromatin Accessibility Analysis. *Research square*.

Gonzalez-Lozano MA, et al. (2025) EndoMAP.v1 charts the structural landscape of human early endosome complexes. *Nature*, 643(8070), 252.

Gonzalez-Lozano MA, et al. (2025) EndoMAP.v1, a Structural Protein Complex Landscape of Human Endosomes. *bioRxiv* : the preprint server for biology.

Buzzao D, et al. (2025) FunCoup 6: advancing functional association networks across species with directed links and improved user experience. *Nucleic acids research*, 53(D1), D658.

Wright SN, et al. (2025) State of the interactomes: an evaluation of molecular networks for generating biological insights. *Molecular systems biology*, 21(1), 1.

González JT, et al. (2025) Age-invariant genes: multi-tissue identification and characterization of murine reference genes. *Aging*, 17(1), 170.

Ibrahim IH, et al. (2025) Altered protein lipidation in cancer: A hidden link to sensory neuronal dysfunction. *Computational biology and chemistry*, 120(Pt 1), 108656.

Sweeney KM, et al. (2024) CRISPR screen for protein inclusion formation uncovers a role for SRRD in the regulation of intermediate filament dynamics and aggresome assembly. *PLoS genetics*, 20(2), e1011138.

Wang K, et al. (2024) Longitudinal molecular profiling elucidates immunometabolism dynamics in breast cancer. *Nature communications*, 15(1), 3837.