# **Resource Summary Report**

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# Resource for Biocomputing Visualization and Informatics

RRID:SCR\_001374

Type: Tool

# **Proper Citation**

Resource for Biocomputing Visualization and Informatics (RRID:SCR\_001374)

#### **Resource Information**

URL: http://www.cgl.ucsf.edu/

Proper Citation: Resource for Biocomputing Visualization and Informatics

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**Description:** Biomedical technology resource center that develops software and web-based resources for the visualization and analysis of molecular structure, and related data, at scales ranging from the atomic to the supramolecular. They create tools for handling and integrating diverse types of biomolecular data, including atomic-resolution coordinates, density maps, sequences, annotations, and networks. Their primary efforts are in the visualization and analysis of structures of molecules and molecular assemblies, enzyme sequence-structure-function relationships, and network representations of protein similarity, binding interactions, and biological pathways. They provide technologies to enable identifying the molecular bases of disease and phenotypic variation, annotating proteins of unknown function, identifying targets for drug development, designing drugs, and engineering proteins with new functions. RBVI distributes software tools, including the popular UCSF Chimera visualization and analysis package, develops and hosts the Structure-Function Linkage Database, and provides access to state-of-the-art computational resources in support of research projects in these areas.

**Abbreviations: RBVI** 

Resource Type: biomedical technology resource center, training resource

**Keywords:** training resource, molecular modeling, software, molecular graphics, visualization, modeling, molecular structure, analysis, computation, computing and informatics technology center, FASEB list

Funding: NIGMS

Resource Name: Resource for Biocomputing Visualization and Informatics

Resource ID: SCR\_001374

Alternate IDs: nlx\_152531

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## Ratings and Alerts

No rating or validation information has been found for Resource for Biocomputing Visualization and Informatics.

No alerts have been found for Resource for Biocomputing Visualization and Informatics.

#### Data and Source Information

Source: SciCrunch Registry

# Usage and Citation Metrics

We found 151 mentions in open access literature.

**Listed below are recent publications.** The full list is available at <u>NIF</u>.

Kaewjiw N, et al. (2025) Domperidone inhibits dengue virus infection by targeting the viral envelope protein and nonstructural protein 1. Scientific reports, 15(1), 3817.

Hu Z, et al. (2024) Emodin alleviates cholestatic liver injury by modulating Sirt1/Fxr signaling pathways. Scientific reports, 14(1), 16756.

Yu Y, et al. (2024) Nuclear pore protein POM121 regulates subcellular localization and transcriptional activity of PPAR?. Cell death & disease, 15(1), 7.

Boonkua S, et al. (2024) Development of chimeric MrNV virus-like particles capable of binding to SARS-CoV-2-susceptible cells and reducing infection by pseudovirus variants. Scientific reports, 14(1), 31431.

Simbulan AM, et al. (2024) Immunoinformatics-guided approach for designing a panproteome multi-epitope subunit vaccine against African swine fever virus. Scientific reports, 14(1), 1354.

Manu P, et al. (2024) Computational Mutagenesis and Inhibition of Staphylococcus aureus AgrA LytTR Domain Using Phenazine Scaffolds: Insight From a Biophysical Study. BioMed research international, 2024, 8843954.

Mise K, et al. (2024) NDUFS4 regulates cristae remodeling in diabetic kidney disease. Nature communications, 15(1), 1965.

Munguía-Robledo S, et al. (2023) Lysine Methyltransferase EhPKMT2 Is Involved in the In Vitro Virulence of Entamoeba histolytica. Pathogens (Basel, Switzerland), 12(3).

Wang Y, et al. (2022) Cryo-EM analysis of Ebola virus nucleocapsid-like assembly. STAR protocols, 3(1), 101030.

Sadeghi M, et al. (2022) Inhibitory effect of flavonoid glycosides on digestive enzymes: In silico, in vitro, and in vivo studies. International journal of biological macromolecules, 217, 714.

Dechkla M, et al. (2022) Cry4Aa and Cry4Ba Mosquito-Active Toxins Utilize Different Domains in Binding to a Particular Culex ALP Isoform: A Functional Toxin Receptor Implicating Differential Actions on Target Larvae. Toxins, 14(10).

Nguyen T, et al. (2021) Direct IgG epitope mapping on bacterial AB toxins by cryo-EM. STAR protocols, 2(4), 100852.

Ahn C, et al. (2021) Mechanisms of typhoid toxin neutralization by antibodies targeting glycan receptor binding and nuclease subunits. iScience, 24(5), 102454.

Pickford P, et al. (2021) Partial agonism improves the anti-hyperglycaemic efficacy of an oxyntomodulin-derived GLP-1R/GCGR co-agonist. Molecular metabolism, 51, 101242.

Schmauder L, et al. (2021) Nematode CDC-37 and DNJ-13 form complexes and can interact with HSP-90. Scientific reports, 11(1), 21346.

Grataitong K, et al. (2021) Chimeric virus-like particles (VLPs) designed from shrimp nodavirus (MrNV) capsid protein specifically target EGFR-positive human colorectal cancer cells. Scientific reports, 11(1), 16579.

Ghodgaonkar-Steger M, et al. (2020) C-Terminal Motifs of the MTW1 Complex Cooperatively Stabilize Outer Kinetochore Assembly in Budding Yeast. Cell reports, 32(13), 108190.

Fonseca AM, et al. (2019) VAR2CSA Serology to Detect Plasmodium falciparum Transmission Patterns in Pregnancy. Emerging infectious diseases, 25(10), 1851.

Ge J, et al. (2018) Structure of mouse protocadherin 15 of the stereocilia tip link in complex

with LHFPL5. eLife, 7.

Su Z, et al. (2018) Electron Cryo-microscopy Structure of Ebola Virus Nucleoprotein Reveals a Mechanism for Nucleocapsid-like Assembly. Cell, 172(5), 966.