

# Resource Summary Report

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## PyRosetta

RRID:SCR\_018541

Type: Tool

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### Proper Citation

PyRosetta (RRID:SCR\_018541)

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### Resource Information

**URL:** <http://www.pyrosetta.org/>

**Proper Citation:** PyRosetta (RRID:SCR\_018541)

**Description:** Interactive Python based interface to Rosetta molecular modeling suite. Stand alone Python based implementation of Rosetta molecular modeling package that allows users to write custom structure prediction and design algorithms using major Rosetta sampling and scoring functions.

**Synonyms:** Python Rosetta

**Resource Type:** software application, standalone software, software resource

**Defining Citation:** [PMID:20061306](#)

**Keywords:** Molecular modeling, custom structure prediction, design algorithm, energy function, scoring function, bio.tools

**Funding:** NIGMS R01 GM73151;  
NIGMS R01 GM078221;  
NSF 0846324

**Availability:** Free, Freely available

**Resource Name:** PyRosetta

**Resource ID:** SCR\_018541

**Alternate IDs:** biotools:pyrosetta

**Alternate URLs:** <https://bio.tools/pyrosetta>

**Record Creation Time:** 20220129T080340+0000

**Record Last Update:** 20260821T043040+0000

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## Ratings and Alerts

No rating or validation information has been found for PyRosetta.

No alerts have been found for PyRosetta.

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## Data and Source Information

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

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## Usage and Citation Metrics

We found 25 mentions in open access literature.

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

Ludwig MS, et al. (2026) Extracellular 70???kDa heat shock protein in blood plasma binds insulin and modulates glycaemic control in vivo. *Cell stress & chaperones*, 31(4), 100180.

Dishner IT, et al. (2026) Circularity in Sequence-Controlled Copolyamides Enabled by Regioselective Enzymatic Hydrolysis. *Journal of the American Chemical Society*, 148(17), 18285.

Vangaru S, et al. (2025) To pack or not to pack: revisiting protein side-chain packing in the post-AlphaFold era. *bioRxiv : the preprint server for biology*.

Vangaru S, et al. (2025) To pack or not to pack: revisiting protein side-chain packing in the post-AlphaFold era. *Briefings in bioinformatics*, 26(3).

Riveros Maidana RLB, et al. (2025) Benchmarking Machine Learning Models for HIV-1 Protease Inhibitor Resistance Prediction: Impact of Data Set Construction and Feature Representation. *Journal of chemical information and modeling*, 65(19), 10037.

Roel-Touris J, et al. (2024) The structural landscape of the immunoglobulin fold by large-scale de novo design. *Protein science : a publication of the Protein Society*, 33(4), e4936.

Sazhnyev Y, et al. (2024) Somatic Mutations within Myocilin due to Aging May Be a Potential Risk Factor for Glaucoma. *Genes*, 15(2).

Andrianov GV, et al. (2024) vScreenML v2.0: Improved Machine Learning Classification for Reducing False Positives in Structure-Based Virtual Screening. *International journal of molecular sciences*, 25(22).

Richards LS, et al. (2023) Cryo-EM Structure of a Human LECT2 Amyloid Fibril Reveals a Network of Polar Ladders at its Core. *bioRxiv : the preprint server for biology*.

Custodio JM, et al. (2023) Structural and physical features that distinguish tumor-controlling from inactive cancer neoepitopes. *Proceedings of the National Academy of Sciences of the United States of America*, 120(51), e2312057120.

Bhardwaj G, et al. (2022) Accurate de novo design of membrane-traversing macrocycles. *Cell*, 185(19), 3520.

Tubiana J, et al. (2022) Reduced B cell antigenicity of Omicron lowers host serologic response. *Cell reports*, 41(3), 111512.

Hickerson BT, et al. (2022) Host receptor-targeted therapeutic approach to counter pathogenic New World mammarenavirus infections. *Nature communications*, 13(1), 558.

Koehler Leman J, et al. (2021) Ensuring scientific reproducibility in bio-macromolecular modeling via extensive, automated benchmarks. *Nature communications*, 12(1), 6947.

Milighetti M, et al. (2021) Predicting T Cell Receptor Antigen Specificity From Structural Features Derived From Homology Models of Receptor-Peptide-Major Histocompatibility Complexes. *Frontiers in physiology*, 12, 730908.

Maguire JB, et al. (2021) XENet: Using a new graph convolution to accelerate the timeline for protein design on quantum computers. *PLoS computational biology*, 17(9), e1009037.

Pan X, et al. (2021) De novo protein fold families expand the designable ligand binding site space. *PLoS computational biology*, 17(11), e1009620.

Julian AT, et al. (2021) 3DFI: a pipeline to infer protein function using structural homology. *Bioinformatics advances*, 1(1).

Zheng Y, et al. (2020) Histone Deacetylase HDA9 and WRKY53 Transcription Factor Are Mutual Antagonists in Regulation of Plant Stress Response. *Molecular plant*, 13(4), 598.

Yang J, et al. (2020) Predicting the viability of beta-lactamase: How folding and binding free energies correlate with beta-lactamase fitness. *PloS one*, 15(5), e0233509.