

Structural bioinformatics

RCSB protein Data Bank: exploring protein 3D similarities via comprehensive structural alignments

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Abstract

Motivation: Tools for pairwise alignments between 3D structures of proteins are of fundamental importance for structural biology and bioinformatics, enabling visual exploration of evolutionary and functional relationships. However, the absence of a user-friendly, browser-based tool for creating alignments and visualizing them at both 1D sequence and 3D structural levels makes this process unnecessarily cumbersome.

Results: We introduce a novel pairwise structure alignment tool (rcsb.org/alignment) that seamlessly integrates into the RCSB Protein Data Bank (RCSB PDB) research-focused RCSB.org web portal. Our tool and its underlying application programming interface (alignment.rcsb.org) empowers users to align several protein chains with a reference structure by providing access to established alignment algorithms (FATCAT, CE, TM-align, or Smith–Waterman 3D). The user-friendly interface simplifies parameter setup and input selection. Within seconds, our tool enables visualization of results in both sequence (1D) and structural (3D) perspectives through the RCSB PDB RCSB.org Sequence Annotations viewer and Mol* 3D viewer, respectively. Users can effortlessly compare structures deposited in the PDB archive alongside more than a million incorporated Computed Structure Models coming from the ModelArchive and AlphaFold DB. Moreover, this tool can be used to align custom structure data by providing a link/URL or uploading atomic coordinate files directly. Importantly, alignment results can be bookmarked and shared with collaborators. By bridging the gap between 1D sequence and 3D structures of proteins, our tool facilitates deeper understanding of complex evolutionary relationships among proteins through comprehensive sequence and structural analyses.

Availability and implementation: The alignment tool is part of the RCSB PDB research-focused RCSB.org web portal and available at rcsb.org/alignment. Programmatic access is available via alignment.rcsb.org. Frontend code has been published at github.com/rcsb/rcsb-pecos-app. Visualization is powered by the open-source Mol* viewer (github.com/molstar/molstar and github.com/molstar/rcsb-molstar) plus the Sequence Annotations in 3D Viewer (github.com/rcsb/rcsb-saguaro-3d).

1 Introduction

Pairwise alignments of 3D structures of proteins are invaluable for understanding evolutionary relationships between and among proteins, to infer functional annotations from well-studied protein structures, and for analyses of Computed Structure Models (CSMs), e.g. from AlphaFold DB (Varadi 2024), ModelArchive (Schwede 2009), or ESM Metagenomic Atlas (Lin 2023). Moreover, the success of machine learning-based protein structure prediction underlines the relevance of these tools as they can be used to transfer ligand binding information from experimentally-determined structures to CSMs or align multiple individual experimentally-determined structures of domains to the CSM of the full-length polypeptide chain.

In this work, we present the RCSB Protein Data Bank (RCSB PDB) research-focused RCSB.org alignment tool, which provides an intuitive graphical user interface (rcsb.org/alignment)

plus an application programming interface (API, alignment.rcsb.org). These entry points empower users to align multiple 3D structures with one reference structure in a pairwise manner. These tools succeed a previously provided RCSB.org structure alignment capabilities (Prlić 2010). Uniquely, our new tool distinguishes itself from comparable solutions (Li 2014, Wiederstein and Sippl 2020, Holm et al. 2023, van Kempen 2023, Prochazka 2024, Zurkowski 2024) by offering a powerful range of parameterizable alignment methods. Users can explore these alignments, both at the 1D sequence level and in 3D, ensuring a seamless and comprehensive experience.

2 Results

The RCSB.org alignment tool (rcsb.org/alignment) facilitates 3D superposition of multiple protein structures to one

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reference structure and integrates with other RCSB.org web portal APIs (Rose 2021, Burley 2022, 2023, Bittrich 2023).

2.1 Required input and options

Individual structures can be selected for alignment by typing in the corresponding PDB entry identifier, by pasting a URL to a file [mmCIF, BinaryCIF (Sehna 2020)], or PDB formats are all supported), or by uploading an externally-provided atomic coordinates file (in various file formats, see Section 2.2). One polypeptide chain must be selected for each entry. Optionally, a sequence range can be included to align only a specific set of residues (and ignore the others). For the alignment, users can choose from a range of established methods, including jFATCAT, jCE, jCE-CP, Smith–Waterman 3D (Lafita 2019), and TM-align (Zhang and Skolnick 2005). Algorithm-specific parameters can also be specified. jFATCAT supports both rigid and flexible alignments: rigid alignments will optimize the superposition of two protein chains in their entirety, whereas flexible alignments can fragment chains into smaller sections and align fragment pairs, while allowing for flexibility in regions connecting fragments. Flexible alignments are particularly useful for partially disordered proteins or those with flexible hinges such as calmodulin or the catalytic domains of protein kinases.

2.2 File upload and URL support

Users can align structures from arbitrary URLs in any of the supported formats: mmCIF, BinaryCIF (Sehna 2020), plus the legacy PDB format.

Local files can serve as input by selecting the “File Upload” input mode. Files are processed by a dedicated API at user-upload.rcsb.org. The API endpoint responds to file uploads with a unique token, which can be used to retrieve the file and initially restricts file access to the original uploader but can be shared freely. The access token grants access to the file for 90 days. Uploaded files can be referenced by their URLs.

2.3 Result overview

Figure 1 showcases the output of a structure alignment. Chain A of 1rpj (D-allose binding protein) serves as reference, chain A of 2oen (distant, sugar-binding homolog) and chain A of AF-P39265-F1 [CSM of D-allose binding protein with UniProt identifier P39265, (Varadi 2024)] were aligned to this reference structure.

For all input structures (Fig. 1A) except the first listed or reference structure, scores of their 3D structure (Fig. 1B) and 1D sequence alignment (Fig. 1C) with respect to the reference are reported. An interactive combination of a 1D sequence view (bottom-left panel) and 3D structure view (bottom-right panel) allows users to interrogate the alignment. The blue boxes on the left side of the sequence tracks (Fig. 1D) summarize the content of each aligned structure by providing a break-down of the aligned polymer chain (1st box), other polymer chains (2nd box), and non-polymer ligands (3rd box). These boxes can be used to toggle the representation of the corresponding item. Hovering over 1D sequence positions (Fig. 1E) or amino acid residues in 3D (Fig. 1F) will highlight the corresponding residue(s), both at the 1D sequence level and in 3D. Sequence features are represented by the RCSB Sequence Annotations Viewer (Segura 2020, 2022). 3D visualization is powered by the Mol* 3D viewer (Sehna 2021).

2.4 Exporting and sharing alignments

Several result files are offered for download (Fig. 1G): Transformed mmCIF files can be reopened in a 3D viewer that will show all structures superimposed, sequence alignments are available as FASTA format file, and 3D transformation matrices are offered in JSON format.

Alignment results can be bookmarked and shared with other users by obtaining a URL that includes all inputs and parameters needed to recreate an alignment.

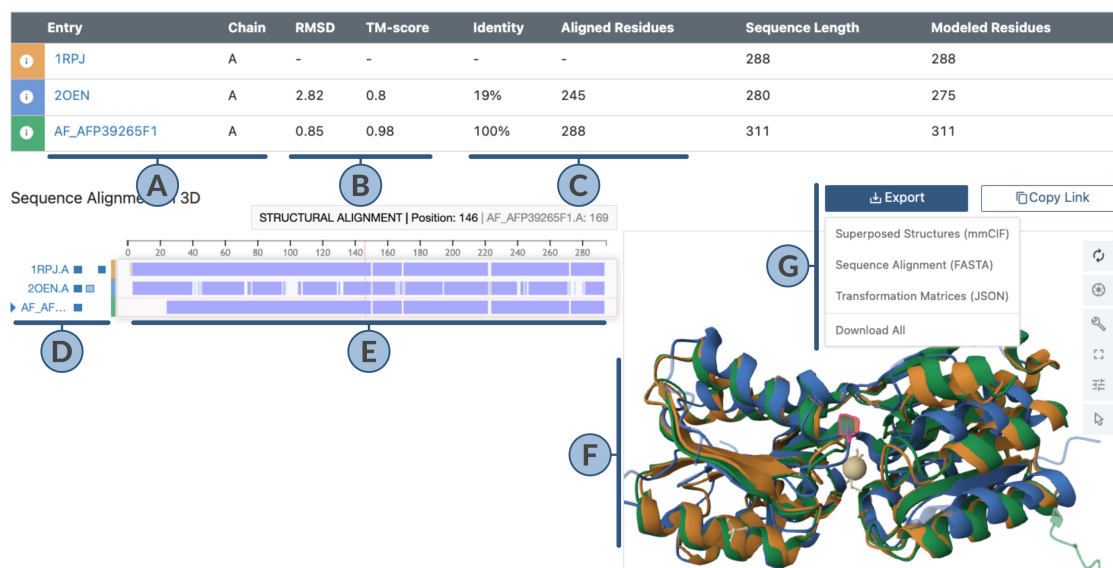


Figure 1. Output of the pairwise alignment application. (A) Summary of aligned chains. (B) 3D Structure-based scores (Root-Mean-Square-Deviation or RMSD in Ångström Units; TM-score, range 0.00–1.00). (C) Sequence-based scores (% identity, number of aligned residues). (D) Toggle visibility of aligned chain, all polymers, and all ligands via 1st, 2nd, and 3rd box buttons, respectively. (E) Structurally aligned sequence regions in a darker shade, regions aligned only at sequence level in a lighter shade, non-aligned gaps are left empty. (F) 3D visualization of aligned structures. (G) Tools to export and share alignment.

2.5 Structure alignment API

The Pairwise Structure Alignment tool is built atop a public API (alignment.rcsb.org), which provides a convenient way to compute structure alignments in a pairwise manner. Time required for computation depends on structure size and the alignment method used. Because some comparisons can take considerable time, alignment jobs run asynchronously. Each user request is assigned a unique identifier in the form of a ticket. This ticket serves as a key to track the progress of the alignment job. Users can check the status of their ticket, allowing them to monitor the processing stages until the job reaches completion.

3 Conclusion

The RCSB PDB research-focused RCSB.org web portal Pairwise Structure Alignment application is a novel tool that allows users to align multiple 3D protein structures to one reference protein chain. Selection from a number of well-established alignment methods is permitted and can be parameterized freely. Alignments of experimentally-determined structures from the PDB archive, CSMs from AlphaFold DB and the ModelArchive are integrated, and data from public URLs plus uploading of user-supplied atomic coordinates files are all supported. Moreover, structure data can be referenced using UniProt accession numbers, AlphaFold DB identifiers, or ESM Metagenomic Atlas identifiers. A combination of the RCSB Sequence Annotations Viewer (Segura 2020, 2022) and the Mol* 3D viewer (Sehna 2021) enables interactive interrogation of alignments by users, both at the 1D sequence level and in 3D. Resulting alignments can be saved in different formats and can be bookmarked and shared via URL.

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Conflict of interest

None declared.

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Applications Note