

ProtacMLA: PROTAC Mutation and Ligand analysis tool

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ABOUT

- ProtacMLA is an online web-based tool that helps users extract, visualize, and analyse the Proteolysis targeting chimeras (PROTACs).
- This online freely accessible web tool helps to extract the PDB format file of PROTAC or the PROTAC complex and analyze the PROTAC region.
- This tool helps to:
 - Extract and analyse ligands acting as the E3 ligase binding ligand and the linker
 - Mutate the amino acid region that acts like the target protein-binding region of the PROTAC molecule
- The tool also enlists databases and tools that are useful in the PROTAC and protein study.

Homepage

The screenshot shows the ProtacMLA homepage with a blue header and a white main content area. The header includes the logo and navigation links. The main content area is divided into several sections: a PDB Viewer section at the top right, a Mutation section in the middle left, a Ligand analysis section in the middle right, and a Databases & Tools section at the bottom. The Mutation and Ligand analysis sections contain input fields and buttons for file uploads and analysis. The Databases & Tools section lists various resources for protein and ligand research.

ProtacMLA - PROTAC Mutation & Ligand Analysis TUTORIAL

Structure Details

Load a PDB to view details...

Ligands

No ligands detected yet...

Show Ligands

Viewer Options

Enter PDB ID (e.g., 6lu7) Load PDB from RCSB or Upload Local File: Choose File | No file chosen

Peptide Mutation TUTORIAL

Generate ready-to-use UCSF Chimera Python scripts with desired mutation and optional minimization.

1) Structural Inputs

PDB ID Local PDB structure file path

Output structure location Desired Output Format

PROTAC Ligand analysis TUTORIAL

Enter a PDB ID or upload a .pdb file. The tool finds non-polymer ligands and enriches them via the RCSB ChemComp API.

PDB ID Load by ID

We download the .pdb file from RCSB and analyze it in your browser.

Or upload a .pdb file | No file chosen Analyze Upload

We read only in your browser; nothing is uploaded anywhere else.

Exclusions (common non-ligands):

We ignore standard amino acids, water, and typical ions/solvents by default. You can add/remove three-letter codes below (comma-separated).

ALA, ARG, ASN, ASP, CYS, GLN, GLU, GLY, HIS, ILE, LEU, LYS, MET, PHE, PRO, SER, THR, TRP, TYR, VAL, SEC, PHE, HDX, IAT, DDD, NA, CL, MG, ZN, CA, SO4, PO4, HEH, NAG, MAN, BNA, PEG, MPO, GOL

[Open PBEE Binding Affinity Prediction Colab](#)

Databases & Tools for PROTAC and Protein Research

Databases

- PROTAC-DB 3.0**: Manually curated database for PROTAC molecules with degradation capacity, binding affinities and cellular activities.
- RCSB PDB**: Archive of 3D structures of large biomolecules (Proteins, DNA, RNA) and their complexes.
- PROTACpedia**: Manually curated collaborative and comprehensive resource for PROTACs and their complexes.
- PDBbind+**: Comprehensive collection of experimental Binding affinity data for biomolecules recorded in RCSB PDB.
- AlphaFold DB**: Collection of over 200 million predicted protein structures using AlphaFold.
- BindingDB**: Binding affinity database focusing on protein complexes considered as drug targets.
- PubChem**: Open chemistry database for small and large molecules.
- ZINC15**: A free database of commercially available compounds for virtual screening.
- DrugBank**: Comprehensive database of approved & experimental drugs.

Tools

- DeepPROTACs**: A deep neural network model that predicts degradation capacity of PROTAC molecule based on structure of given target protein and E3 ligase.
- DIFFPROTACs**: Transformers based model that learns and generates new PROTAC linkers based on given ligands.
- SwissADME**: Website that allows to compute physicochemical descriptors and predict ADME parameters, pharmacokinetic properties, druglike nature and medicinal chemistry friendliness of small molecules.
- PlayMolecule**: A virtual environment for drug discovery where simulation, AI and data are integrated.
- ChemProp**: A machine learning package for chemical property prediction using D-MPNN architecture.
- DeepChem**: Open-source ML toolchain that democratizes the use of Deep-learning in drug discovery, material science, quantum chemistry and biology.
- ProteinPlus**: A webserver for structure-based molecule modeling, provides functionalities for structure quality assessment, hydrogen placement, search alternative conformations.

PDB Viewer section

Mutation section

Database and Tools section

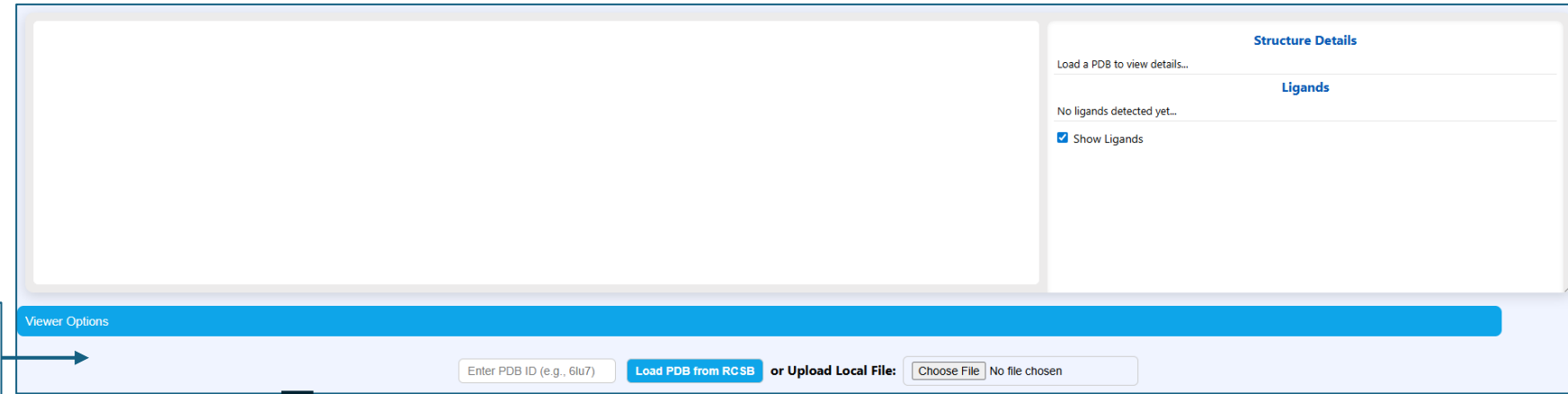
PBEE protein-peptide binding affinity predictor

Ligand analysis section

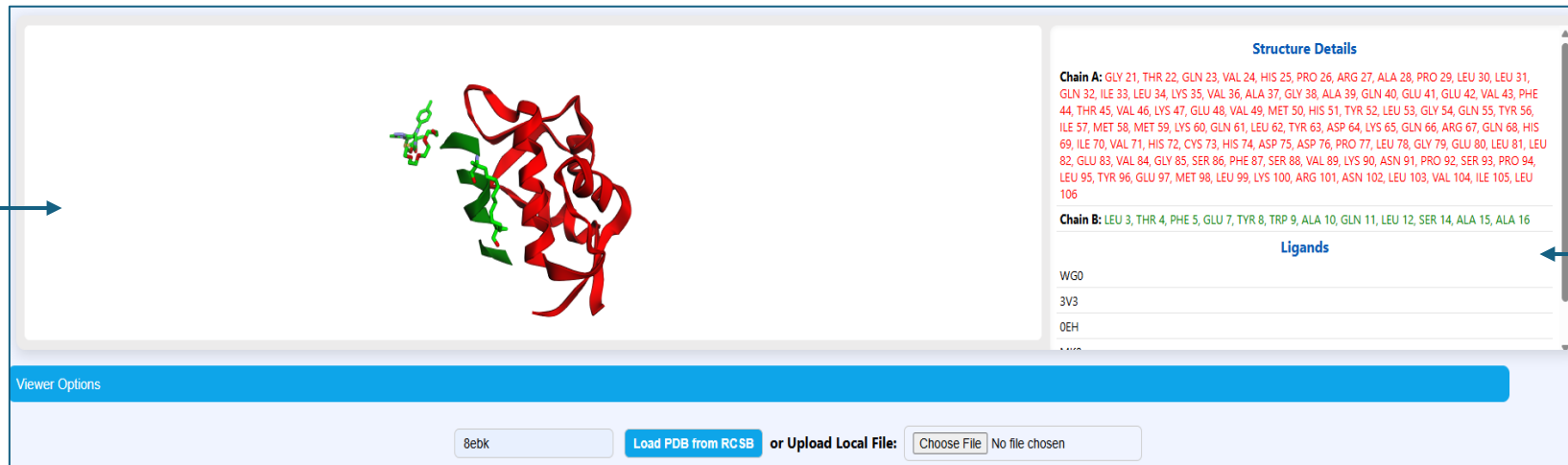
PDB structure viewer section

This section helps users visualize 3D structures along with their residues, chains, and involved ligands, allowing them to identify their target complex.

The viewer is designed in a way that it excludes common ions, water molecules, and uncommon ligands.



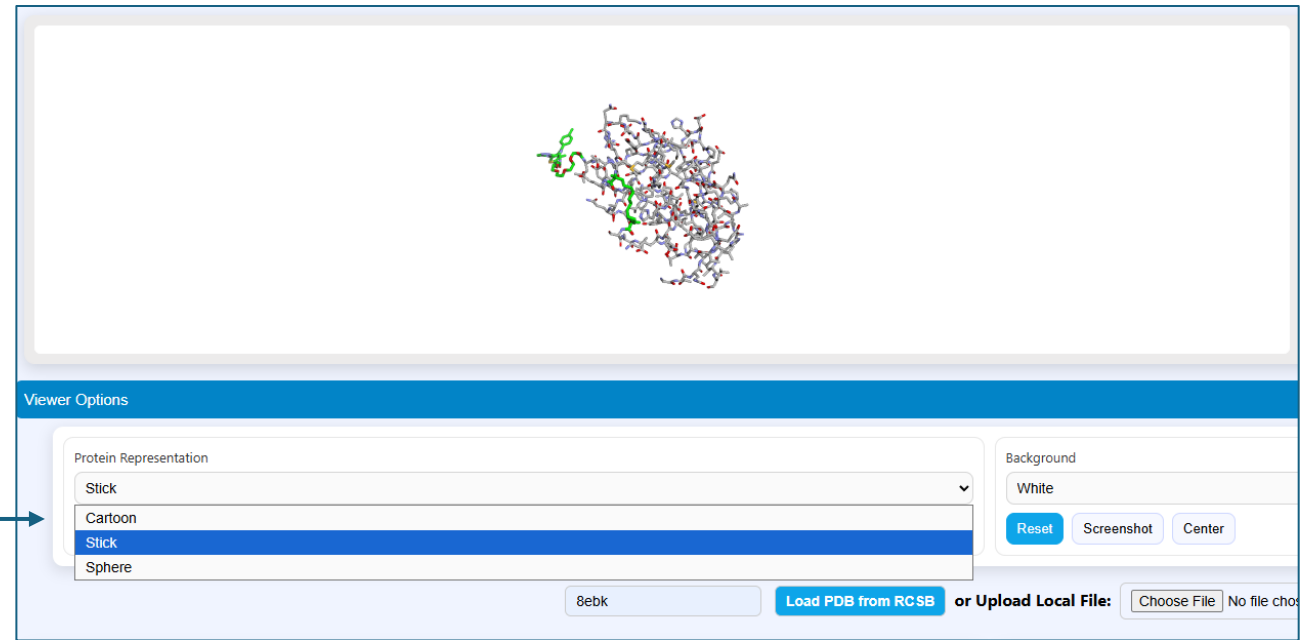
Users may enter the PDB ID or upload their PDB structure from the local system.



The 3D structure is loaded, which can be visualized here

Details of the structure, including their residues, chain, and involved ligands, may be found here

Using the 'Viewer option', the user may change the structure representation to cartoon/stick/sphere for easy visualization of the structure components.



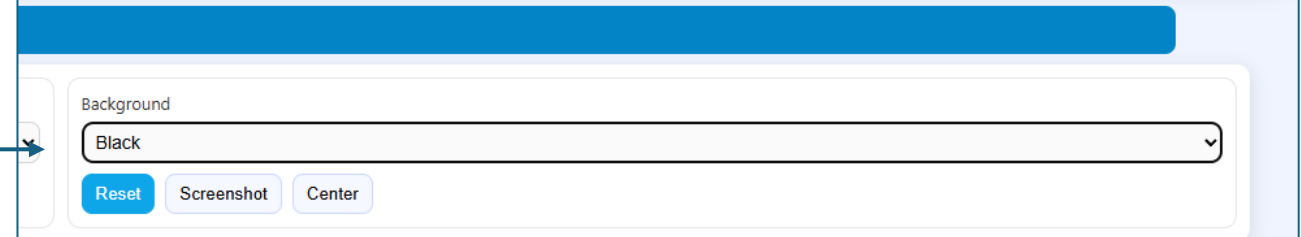
The screenshot shows a protein structure in stick representation within a viewer window. Below the window is a 'Viewer Options' panel. The 'Protein Representation' dropdown menu is open, showing options: Stick, Cartoon, Stick (highlighted), and Sphere. To the right, the 'Background' dropdown is set to 'White'. Below these are 'Reset', 'Screenshot', and 'Center' buttons. At the bottom, there is a 'Load PDB from RCSB' button and an 'Upload Local File' section with a 'Choose File' button and 'No file chosen' text.

Using the 'Viewer option', the user may also change the background to white/black/soft blue to increase the visibility of small components.



The screenshot shows the same protein structure in stick representation, but now on a black background. The 'Viewer Options' panel is partially visible, showing the 'Background' dropdown set to 'Black'. To the right, there is a 'Structure Details' section with two chains of amino acid sequences. Chain A is highlighted in red and Chain B in green. Below that is a 'Ligands' section with a list of ligand IDs: WGO, 3V3, 0EH, and 1V2.

The options for Reset (reset to default settings), Screenshot (save structure image in PNG format), and Center (move the structure to the center of the view window) are also included.



This is a close-up of the 'Viewer Options' panel. The 'Background' dropdown menu is open, showing 'Black' as the selected option. Below the dropdown are three buttons: 'Reset', 'Screenshot', and 'Center'.

Databases and Tools for PROTAC and Protein Research

This section of the tool is designed to help users access various available resources for computational drug discovery and bioinformatics using PROTAC.

The included databases and tools range from several PROTAC-specific resources to general ones.

Databases



PROTAC-DB 3.0

RCSB PDB

PROTACpedia

PDBbind+

AlphaFold DB

BindingDB

PubChem

ZINC15

DrugBank

Tools



DeepPROTACs

DiffPROTACs

SwissADME

PlayMolecule

ChemProp

DeepChem

ProteinPlus

PROTAC-DB 3.0

- **Purpose:** A manually curated database specifically for PROteolysis-TArgeting Chimeras (PROTACs). It's a specialized resource for researchers working on targeted protein degradation.
- **Key Features:** It provides detailed information on PROTAC molecules, including their chemical structures, and crucial biological data like degradation capacity (DC50 and Dmax), binding affinities (IC50, Ki, Kd), and cellular activities. The updated version, 3.0, has expanded its entries and now includes pharmacokinetic data, which is essential for assessing the druggability of these molecules. The database also categorizes the components of a PROTAC: the warhead (the part that binds to the target protein), the E3 ligase ligand, and the linker.

RCSB PDB (Protein Data Bank)

- **Purpose:** The central global archive for 3D structures of large biomolecules. It serves as a fundamental resource for structural biologists, providing a detailed view of the atomic coordinates of proteins, DNA, RNA, and their complexes.
- **Key Features:** PDB entries are derived from experimental methods like X-ray crystallography, NMR spectroscopy, and cryo-electron microscopy. The database is a go-to source for understanding molecular function, interactions, and for structure-based drug design. It includes extensive annotations, visualizations, and tools for searching and analyzing structures.

PROTACpedia

- **Purpose:** A collaborative, high-quality, and freely accessible resource for PROTACs. Unlike commercial databases, it's designed to be a community-driven platform.
- **Key Features:** It provides a manually curated collection of data on PROTAC molecules. The collaborative nature of the platform means that registered users can contribute new data, helping to expand the resource. This makes it a dynamic and up-to-date source of information for the PROTAC research community.

PDBbind+

- **Purpose:** A specialized database that bridges structural and energetic information. It's a comprehensive collection of experimentally determined binding affinity data for protein-ligand complexes that have a corresponding 3D structure in the RCSB PDB.
- **Key Features:** It is an invaluable resource for developing and validating computational methods in drug discovery, such as docking and scoring functions. The database provides a "refined set" of high-quality data, which is a standard benchmark for testing the performance of molecular modeling algorithms.

AlphaFold DB

- **Purpose:** A collection of over 200 million predicted protein structures generated by Google DeepMind's AlphaFold AI system. It dramatically expands the number of available protein structures beyond those determined experimentally.
- **Key Features:** For many organisms, it provides a comprehensive predicted proteome. Each structure comes with a per-residue confidence score (pLDDT), allowing users to assess the reliability of the prediction for different regions of the protein. This database is accelerating research by providing structural models for proteins that have not yet been experimentally characterized.

BindingDB

- **Purpose:** A publicly accessible database focused on the binding affinities of small molecules to proteins, particularly those considered to be drug targets.
- **Key Features:** It contains millions of data points, including Ki, Kd, IC50, and EC50 values. This data is critical for medicinal chemists and computational modelers for developing Structure-Activity Relationships (SAR), training machine learning models, and validating docking methods. It also provides links to related information in other databases like RCSB PDB and PubChem.

PubChem

- **Purpose:** An open chemistry database maintained by the National Institutes of Health (NIH). It's a foundational resource for a vast array of chemical information.
- **Key Features:** It includes a massive collection of information on chemical structures, physical properties, biological activities, and more. Data is contributed by a wide range of sources, including government agencies and vendors. It's a go-to resource for chemical searching, whether by structure, name, or other identifiers.

ZINC15

- **Purpose:** A free and comprehensive database of commercially available chemical compounds. It is optimized for virtual screening.
- **Key Features:** ZINC15 contains over 230 million purchasable compounds, all in "ready-to-dock," 3D formats. This is a crucial feature for computational chemists who need to quickly prepare large libraries of molecules for docking simulations. It also provides various filters based on molecular properties like molecular weight and LogP, making it easy to create focused screening libraries.

DrugBank

- **Purpose:** A unique bioinformatics and cheminformatics resource that combines detailed drug information with comprehensive drug target data.
- **Key Features:** It includes data on approved and experimental drugs, providing their chemical, pharmacological, and pharmaceutical details. It also offers a wealth of information on the drugs' protein targets, including sequences and pathways. This database is a powerful tool for linking drugs to their mechanisms of action and for drug repurposing studies.

TOOLS

DeepPROTACs

- **Purpose:** A deep neural network model designed to predict the degradation capacity of a PROTAC molecule.
- **Key Features:** It takes the 3D structures of the target protein and the E3 ligase as input, along with the PROTAC molecule's structure. The model uses a combination of Graph Convolutional Networks (GCNs) and other neural network architectures to predict whether a given PROTAC will effectively degrade its target. This tool helps in the early-stage rational design of PROTACs, as it can filter out potentially ineffective molecules before synthesis.

PlayMolecule

- **Purpose:** A virtual environment for drug discovery that integrates simulations, AI, and data. It provides a comprehensive platform for various computational workflows.
- **Key Features:** It offers a suite of applications for tasks like protein and small molecule preparation, virtual screening, binding mode analysis, and relative binding affinity predictions. It can be accessed via a graphical user interface, a Python API, or the command line, providing flexibility for different user needs. It's designed to streamline complex drug discovery processes.

DiffPROTACs

- **Purpose:** A generative AI model based on diffusion and transformer architectures for designing new PROTAC linkers.
- **Key Features:** This tool addresses a major challenge in PROTAC design: the linker. It can learn the properties of existing linkers and generate novel ones that connect a given target ligand and E3 ligase ligand. It's a powerful approach for exploring the vast chemical space of potential linkers to find ones that optimize the properties of the PROTAC molecule.

ChemProp

- **Purpose:** An open-source machine learning package for chemical property prediction, specifically using Directed Message Passing Neural Networks (D-MPNNs).
- **Key Features:** This is a coding-focused tool for researchers who want to build their own predictive models. It simplifies the process of training and using powerful graph neural networks on molecular data. It can predict a wide range of properties for single molecules, reactions, and multi-molecule systems, making it highly versatile for cheminformatics tasks.

ProteinPlus

- **Purpose:** A web server for structure-based molecular modeling, focusing on supporting life scientists who work with protein structures.
- **Key Features:** It provides a suite of functionalities for preparing and analyzing protein structures, especially in the context of protein-ligand interactions. Key tools include structure quality assessment, hydrogen placement, binding site prediction (DoGSiteScorer), and the generation of aligned protein structure ensembles. It also features tools for 2D ligand interaction diagrams and finding alternative conformations.

SwissADME

- **Purpose:** A web-based tool for computing physicochemical descriptors and predicting Absorption, Distribution, Metabolism, and Excretion (ADME) parameters for small molecules.
- **Key Features:** It helps researchers quickly assess the "druglike" nature of their compounds. It predicts properties such as water solubility, gastrointestinal absorption, blood-brain barrier penetration, and adherence to rules like Lipinski's Rule of Five. This is an essential tool for early-stage drug discovery to filter out molecules with poor pharmacokinetic properties.

DeepChem

- **Purpose:** An open-source machine learning toolchain that aims to democratize the use of deep learning in drug discovery, materials science, quantum chemistry, and biology.
- **Key Features:** It provides high-level APIs for building and deploying deep learning models on molecular datasets. It includes a wide variety of models and data featurizers, making it easy to get started with complex machine learning tasks. It's a foundational library for researchers who want to apply AI to scientific problems.