

Homology modeling using SWISS-Model for Biochemistry

In 1 collection

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1 И

Works for me

This protocol is published without a DOI



ABSTRACT

Protocol for homology modeling proteins for use in Biochemistry I at James Madison University. Protocol guides students to use the SWISS-Model web server (citations below).

The protocol directs users to save data in OSF or the <u>Open Science Framework</u>. This is the preferred project management tool for the class and is required for JMU students using this for the course. Other users can use whichever system is preferred.

Citations for servers:

1. Waterhouse, A., Bertoni, M., Bienert, S., Studer, G., Tauriello, G., Gumienny, R., Heer, F. T., de Beer, T. A. P., Rempfer, C., Bordoli, L., Lepore, R., and Schwede, T. (2018) *SWISS-MODEL: homology modelling of protein structures and complexes*. Nucleic Acids Res. 46, W296–W303.

PROTOCOL CITATION

Michael Friedman, Chris Berndsen 2020. Homology modeling using SWISS-Model for Biochemistry I. **protocols.io**

https://protocols.io/view/homology-modeling-using-swiss-model-for-biochemist-bkmjku4n

COLLECTIONS (i)

B

Biochemistry I methods

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LAST MODIFIED

Sep 08, 2020

PROTOCOL INTEGER ID

41355

PARENT PROTOCOLS

Part of collection

Biochemistry I methods

GUIDELINES

This protocol guides students through homology modeling and analysis of the resulting model. This protocol uses the CRX DNA binding domain to generate the results thus the shown images and results will vary.

The protocol directs users to save data in OSF or the <u>Open Science Framework</u>. This is the preferred project management tool for the class and is required for JMU students using this for the course. Other users can use whichever system is preferred.

MATERIALS TEXT

SWISS-MODEL server: https://swissmodel.expasy.org/

Phyre² server: http://www.sbg.bio.ic.ac.uk/~phyre2/html/page.cgi?id=index

A sequence in FASTA format

Internet connection

Structure viewing program such as YASARA or UCSF Chimera

Open Science Framework account (JMU students only)

BEFORE STARTING

Gather your sequence in FASTA format (an example is shown below)

>seq_name	
MASDETEASETEAMDAET	

NCBI BLAST 10m

1 Navigate to NCBI BLAST (Basic Local Sequence Alignment Tool) and paste your sequence into the "Enter Query Sequence" box.



1.1 The standard settings for the search are shown in the table.

	Default Setting	What it
		does
Enter Query Sequence		
Query Subrange	(Blank)	Limits
		search to
		a part of
		the
		sequence.
		Can be
		useful if
		there are
		common
		motifs/do
		mains in
		the
		sequence.
Choose Search Set		

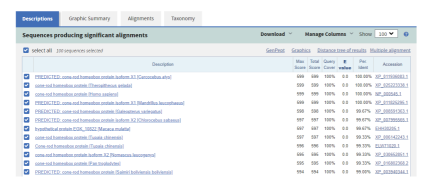
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- 1.2 Record any changes to the settings in Step 2.1 below:
- 1 3 Press BLAST and wait until the results return.

Thie search can take up to © 01:00:00 hour

Analysis of BLAST results to ID sequence

2 Results will be returned as shown as below:



2.1 Column definitions from the **Descriptions** tab of the results.

	Table column	What it tells
		you
Description		Tells you
		identify of
		matching
		sequence.
		Predicted or
		hypothetical in
		title indicates
		protein has not
		been verified.
Max Score		During
		alignment
		identities,
		similarities,
		and gaps are
		scored. This
		indicates the
		best score if
		the sequence
		was aligned
		multiple times
Total Score		If many
		disconnected
		parts matched
		this is the sum
		of the max
		scores for
		those

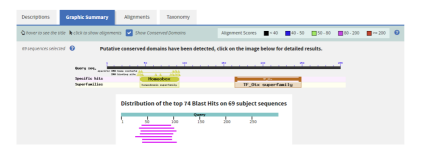
Query Cover	Indicates the percentate of the query sequence found in the match. 100% means all of the sequence was found.
E value	E(xpect) value tells you how many sequences that would rank higher if this was a random match. 0 or very small numbers are good.
Per. Ident	How much of the sequence was identical in sequence. Need >40% for good homology model.
Accession	The accession number for the sequence. Can be clicked to take you to the info card on that sequence.

2.2 Record your best 5 sequences and their statistics in the table below.

Sequence Description	Max Score	Total Score	Query Coverage	E value	Per Ident	Accession

3 In the **Graphic Summary** tab, you can view the domains in your sequence.

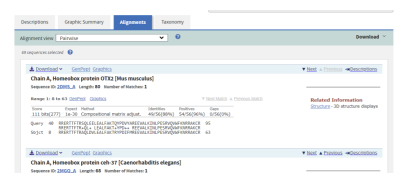
A **domain** is a part of the sequence with a known fold/shape/structure. A **motif** is a sequence that has a shape or function. Typically domains can fold on their on, while motifs are shorter pieces within domains.



3.1 Record any domains or motifs in the table below along with the approximate position within the sequence. This can help in the modeling and support the accuracy of your model later on.

Domain/Motif name	position (this should be a number/set of numbers)

4 In the Alignments tab, the actual sequence alignment (the data) are shown.



- 4.1 Each alignment shows the following key information:
 - Identities and their location within the sequence.
 - Positives and their location within the sequence.
 - Gaps and their location within the sequence.
 - The alignment: Your sequence is the top row, the matched sequence in the middle row (+ means similar), and the sequece from the database (called Sbjct).
 - Position number of the sequence match. These are the numbers at each end of the sequences.
- **4.2** Press the *Download* link to the top right of the alignment and select *Text* you will get a complete file of your results. Upload this to your OSF folder for this project and name the file:

BLAST_alignment_[Group_name]_[sequence_name].txt

Replace [Group_name] with your name/group name without the brackets. Replace [sequence_name] with the name of the sequence.

Indicate your OSF file location as a link within a note on this step.

THIS IS YOUR DATA FILE FOR THE SEARCH!

Analysis of BLAST results to ID potential modeling templates

5 and repeat search but limit the Database to Protein Data Bank proteins (pdb). This search will identify proteins of known structure that match your protein and can suggest if your modeling attempt will be successful. Record your sequence matches in the table.

5.1 Accession numbers here lead to the information on the structure which may help when using SWISS-MODEL. These accession numbers are the PDB ID numbers.

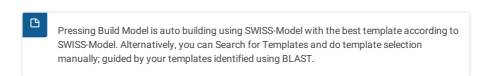
Sequence Description	Max Score	Total Score	Query Coverage	E value	Per Ident	Acce ssion

Table for recording results from PDB focused BLAST.

5.2 The top five structures here are potential **templates structures** which you can use to model your sequence. This means these structures are similar at the sequence level to your sequence and *potentially* will result in a similar structure to your sequence.

Start a New Modell	ing Project ⊚	Copy and paste your sequence	ce here	
Target Sequence(s): (Format must be FASTA.	Paste your target sequence(s) or UniP	rotKB AC here	Supported Inputs	Θ
Clustal, plain string, or a valid UniProtKB AC)	+ Upload Target Sequence File Validate		Label so you know what it be with minintext.	will
Project Title:	Untitled Project			
Email:	Optional			
	Search For Templates	Build Model	Last step! S	Select
	By using the SWISS MODEL server you agree to comply with	h the following terms of use and to cite the corresponding articles.		

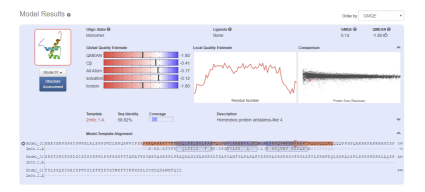
6.1 Follow the instructions on the image above to start the modeling by Build Model. Initial steps can take up to © 00:20:00



6.2 If you choose to do manual building via the Search for Templates tab. Record your templates below.

Template name (second column in table)		

7 Once model building (either manual or automated) is complete, a screen as shown below will appear.



7.1 Useful information from this screen

- GMQE for Global Model Quality Estimation is scored from zero to 1 and indicates model quality based on the alignment with numbers closer to 1 indicating a more reliable model.
- QMEAN indicates the model quality based on structural features and the quality of the chemistry such as torsion angles and solvation. A good model has a number that is more positive, although a good model can have a negative QMEAN score. Less than -4 and model has bad chemistry.
- Local Quality Estimate indicates model quality on a per residue basis and can indicate if there are sections of hte model that are problematic (such as the ends of the model in the report above)
- Model-Template alignment shows how well the template structure and the sequence align and
 what parts of the model were used. Blue colors means better alignment while red colors mean
 worse alignment and modeling. Secondary structure is also indicated with tubes for α-helix and
 arrows for β-sheet.
- 8 The grey Model button leads to a menu to download information.

Two key options:

- 1. PDB format results in just the homology model, which can be viewed in YASARA or Chimera
- 2. **Model Report** downloads a .zip with the PDB file model and an HTML based report of the model process including the statistics shown in Step 8.1.

Download both and upload both files to OSF.

Name the PDB file:

SWISS_model_[Group_name]_[sequence_name].pdb

Replace [Group_name] with your name/group name without the brackets. Replace [sequence_name] with the name of the sequence.

Name the zip file:

SWISS_data_[Group_name]_[sequence_name].zip

Replace [Group_name] with your name/group name without the brackets. Replace [sequence_name] with the name of the sequence.

THESE ARE YOUR DATA FILES FOR SWISS MODEL!

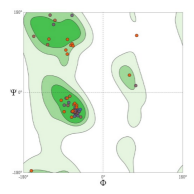
- 8.1 Indicate your OSF file location as a link within a note on this step.
- 9 The Structure Assessment button leads to a new page showing a basic geometric and chemical assessment of the model.
 - 9.1 The Ramachandran plot indicates if the **phi/psi angles** are appropriate for a protein structure and is interactive. The Phi angle is the **dihedral angle** for the rotation of the N-C α bond while the psi angle is for rotation around the C α -C bond of the amino acid backbone. The ideal angles for helices, sheets, and coils shown in green areas below are known to minimize steric clashes between atoms.

Use the camera tool to record the Ramachandran plot and upload it to your OSF.

Name the image file:

SWISS phipsi [Group name] [sequence name]

Replace [Group_name] with your name/group name without the brackets. Replace [sequence_name] with the name of the sequence. Add a note with the link to your file.



Ramachandran plot

- 9.2 The Molprobity Results are numerical scores based on the model and indicate what percentage of amino acids that fall in the ideal geometry category and have minimal clashes. The check boxes allow for visualization of the bad amino acids and can be useful to see if there are general model problems or localized issues. Localized issues can be fixed, general problems cannot.
- 9.3 Record your Molprobity numbers.

	Deviant amino acids
Molprobity Score	
Clash Score	
Ramachandran favored	
Ramachandran outliers	
Rotamer outliers	
C-beta deviations	
Bad bonds	
Bad angles	

10 Make sure you have recorded all the required data.

If you have completed the Phyre modeling. Save the record, export to PDF and upload this file to OSF in the Notebook files.

Ligand identification using SWISS-Model

- 11 Clues to functionality can be gleaned from comparing unknown or predicted structures to with previously characterized structures of known function and characteristics. These scans can be biased against novel proteins or proteins with similar structures but distinct functions, but for initial guesses can be powerful. These methods align the structure and/or amino acid sequence to a database of structures with known ligand binding sites and look for structures with the similarity in amino acid composition, position, and over 3-D similarity. The idea being that similar structures lead to similar functions.
- 12 Return back to your SWISS-Model search and note any models with ligands present. This is noted in the far right column of the templates page.



Identify the ligand by clicking on the ligand name. Record ligands in the top few hits in the table below.

I	Ligand name

- 14 Click the Name of the template to see where the ligands bind to the protein.
 - 14.1 Generally ligands are classified into nonfunctional binders, covalent, and non-covalent binders. The latter two categories are the most interesting. Hovering over the ligand name shows the molecule bound to the protein and left-clicking on the name zooms the structure to show the specifics of ligand binding, including the weak interactions between the amino acids and ligand.
- 15 Download the top two hits with ligands bound from the server and align it to your model in YASARA and record the RMSD value that YASARA returns to you.
 - A perfect match in RMSD is 0, while a poor match is one where the RMSD value is >3 Å, however a high RMSD value does not mean there are not regions of local similarity. A visual comparison is always helpful!
- Observe if there is any match in the ligand/substrate binding sites between your model and the template structures with ligands bound.
 - Does the ligand "fit" into the aligned sites? It will not be perfect, so look for how bumps could fit into holes or nearby holes! Weak interactions also should be analyzed.
 - 16.1 Record the ligand name and possible interactions between your model and the ligand below.

Ligand name	Source structure PDB ID	Interacting amino acids in the model structure (three letter code and amino acid number)