## **Protocol capture**

#### Introduction

The following is a protocol capture of designing cyclic peptides based on antibody CDRH3 loops. It will review the design of CDRH3 loop from anti-influenza antibody C05.

All Rosetta commands for this publication were run with version 3e41de71be009712db5ba0f3b0cd1080a1603181, from March 2016.

All materials from this protocol capture can be downloaded from <a href="https://github.com/sevya/cyclic\_peptide\_protocol\_capture">https://github.com/sevya/cyclic\_peptide\_protocol\_capture</a>

## **Dependencies:**

Several scripts require the use of Python 2.7 as well as the Biopython package (<a href="https://github.com/biopython/biopython.github.io/">https://github.com/biopython/biopython.github.io/</a>). We recommend installing all necessary packages before beginning this protocol. Note that all analysis scripts will only function properly if they are in the correct directory as provided.

## **Structure preparation**

First, the C05 Fab structure (PDB ID 4fnl) was downloaded from the Protein DataBank (PDB; www.rcsb.org) and manually processed in PyMol. All waters were removed from the structure and non-protein residues were also removed. All residues except for one copy of the CDRH3 from chain H (residues 93-102, sequence AKHMSMQQVVSAGWERADLVGDAFDV) were deleted. The loop was then saved to a PDB file with the following command:

save CO5\_H3.pdb

The CDRH3 peptide was then renumbered using a python script to convert the numbering to start from 1 and ignore insertion codes. The renumbering script was run with the following command:

/path/to/Rosetta/tools/protein\_tools/scripts/pdb\_renumber.py
C05\_H3.pdb C05\_H3\_renum.pdb

Next the PeptideStubMover functionality in Rosetta was used to add a cysteine to the Nand C-termini of the CDRH3 peptide. The following command will add these cysteines:

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgc crelease -parser:protocol add\_disulfide.xml -s C05\_H3\_renum.pdb - out:prefix disulfide\_ -out:no\_nstruct\_label -extra\_res\_fa CYD.params

#### **GeneralizedKIC Loop Closure**

Now the peptide is ready for loop modeling simulations. We used Generalized Kinematic Closure (GeneralizedKIC) to close the loop and perturb the  $\varphi$  and  $\psi$  angles. Full of the GeneralizedKIC protocol documentation can be found https://www.rosettacommons.org/docs/latest/scripting\_documentation/RosettaScripts/co mposite\_protocols/generalized\_kic/GeneralizedKIC. Briefly, we first declare a bond between the cysteines at residues 1 and 28, and set the degrees of freedom to be used in loop modeling. We set residue 13 at the tip of the CDRH3 loop to be the anchor point of loop modeling, and the remaining residue to be mobile degrees of freedom. We next add a perturber that will modify the closed loop by perturbing the φ and ψ angles of all residues in the loop by a value drawn from Gaussian distribution centered at 15 degrees. Finally we add a single round of ROSETTA relax to add side chains and refine the structure before evaluating the energy. The GeneralizedKIC protocol will generate 20 solutions after loop closure, perturbation, and relaxation, and the lowest energy solution is reported as the final decoy. This entire protocol is repeated to generate 1,000 final output decoys. To create the models output directory and run the GeneralizedKIC protocol use the following command:

mkdir models

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgc
crelease @close\_relax.flags -s disulfide\_C05\_H3\_renum.pdb

# **Analysis**

We will use a python script to analyze the folded peptide models by calculating ROSETTA score and  $C\alpha$  RMSD for each of the models. To do so we will run the following command:

```
python
/path/to/Rosetta/tools/protein_tools/scripts/score_vs_rmsd.py --
table native_sc_vs_rmsd.tsv --native disulfide_C05_H3_renum.pdb
--CA --term total models/disulfide*pdb
```

```
python plot_score_vs_rmsd.py native_sc_vs_rmsd.tsv
```

This will make a plot of score vs RMSD for all models. In addition it will give a funnel discrimination score that is used to assess how well models converge on the native conformation. This score is derived from Conway et al.<sup>1</sup>. Overall a lower score (more negative) indicates that the structures are converging well.

### Peptide sequence redesign

Next we want to see if we can redesign the peptide for greater stability and convergence on the active conformation. As an example we will take the lowest RMSD structure and run fixed backbone ROSETTADESIGN to optimize the sequence. In a production run we

<sup>&</sup>lt;sup>1</sup> Patrick Conway et al., "Relaxation of Backbone Bond Geometry Improves Protein Energy Landscape Modeling.," *Protein Science* 23, no. 1 (January 2014): 47–55, doi:10.1002/pro.2389.

recommend to design more than one structure – in the manuscript we redesigned all peptides under 2 Å.

```
mkdir redesign/
sort -nk3 native_sc_vs_rmsd.tsv | head -2
```

Next copy the lowest energy model into the redesign folder. We provide an example structure for the purpose of this protocol capture. We will run 10 iterations of fixed backbone design and use the lowest scoring design. The resfile we use to guide design will allow all residues to be mutated to anything except for cysteine (ALLAAxC) and will disallow design on the N- and C-termini cysteines.

```
cp models/disulfide_C05_H3_renum_close_relax_0050.pdb redesign/
```

/path/to/Rosetta/main/source/bin/fixbb.default.linuxgccrelease s disulfide\_C05\_H3\_renum\_close\_relax\_0050.pdb -nstruct 10 out:prefix redesign\_ -ex1 -use\_input\_sc -resfile
redesign.resfile

```
redesign.resfile:
NATAA
start
2 - 27 A ALLAAXC
```

After making the peptide designs we will analyze the lowest scoring design to see if it improves the folding funnel. We will first use python scripts to convert the sequence from the PDB into a fasta file, then create a resfile from the fasta file to mutate our folding template.

/path/to/Rosetta/tools/protein\_tools/scripts/get\_fasta\_from\_pdb.py redesign\_disulfide\_C05\_H3\_renum\_close\_relax\_0050\_0009.pdb A redesign.fasta

python fasta\_to\_resfile.py redesign.fasta

This will create a resfile called redesign\_disulfide\_C05\_H3\_renum\_close\_relax\_0050\_0009.resfile that we will use to mutate our peptide template. Navigate back to the starting directory, create the template for folding and run the folding simulations:

cd ..

/path/to/Rosetta/main/source/bin/fixbb.default.linuxgccrelease - s disulfide\_C05\_H3\_renum.pdb -out:prefix d1\_ -resfile redesign/redesign\_disulfide\_C05\_H3\_renum\_close\_relax\_0050\_0009.r esfile -out:no\_nstruct\_label -use\_input\_sc

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgc crelease @close\_relax.flags -s d1\_disulfide\_C05\_H3\_renum.pdb After the design folding decoys are finished we can analyze the score and RMSD and compare to the native sequence:

```
python
/path/to/Rosetta/tools/protein_tools/scripts/score_vs_rmsd.py --
table d1_sc_vs_rmsd.tsv --native disulfide_C05_H3_renum.pdb --CA
--term total models/d1*pdb

python plot_score_vs_rmsd.py native_sc_vs_rmsd.tsv
d1_sc_vs_rmsd.tsv
```

# **Binding affinity measurement**

In the design process to this point we haven't accounted for the effect that a mutation may have on binding to the antigen. After we have identified a candidate peptide we next want to make sure that the mutations that stabilize the peptide do not interfere with an interaction hotspot. To do this we will model the redesigned sequence in the context of the antibody-antigen interface. We will thread the redesigned sequence over the co-crystal structure, perform a subtle refinement using ROSETTA relax with backbone constraints, and measure the binding energy.

First make a new subdirectory called ddg\_measure to place all the new files and navigate to this directory. Next we need to download and prepare the structure of the C05 co-crystal structure (PDB ID 4fp8). Download the structure and use PyMol to delete all chains except for A+H. Remove all waters and non-protein residues from the structure. Then remove all residues on the antibody except for the CDRH3 (residues 93-102, sequence AKHMSMQQVVSAGWERADLVGDAFDV). Save this structure as C05\_H3\_Ag.pdb

Next we will renumber the structure to make sure the numbering is uniform. Repeat the same command from earlier in the protocol to renumber the structure:

```
/path/to/Rosetta/tools/protein_tools/scripts/pdb_renumber.py
CO5_H3_Ag_renum.pdb
```

Once the structure is prepared we will run the relaxation on both the native sequence and redesigned sequence. As an example the sequence of one of the peptides provided in this manuscript (d1) is provided, along with a native resfile to measure the binding energy of the wild-type loop. Run the constrained relaxation with these two resfiles to create the models:

```
mkdir models
```

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgc crelease @relax.flags -out:suffix \_rlx\_d1 -parser:script\_vars resfile=d1 This will generate ten models for each of the sequences. To analyze the results use the following command to output the score and binding energy in the bound state:

```
sort -nk2 relax.fasc | grep rlx_native | head -1 | awk '{print $2" "$6}'
sort -nk2 relax.fasc | grep rlx_d1 | head -1 | awk '{print $2" "$6}'
```