**Protocol Capture**

**Intro**

Here I will describe how to run and analyze the results from restrained multistate design (R-MSD). R-MSD is used for multistate design, to minimize the energy of multiple protein complexes simultaneously. This is useful in many different contexts, for example if you want to design a single sequence that can bind multiple proteins with high affinity, or a sequence that stabilizes a protein in multiple conformations. In my benchmark cases my goal is to design the sequence of proteins that have multiple binding partners to increase affinity for all binding partners. I show an example case where I take a protein, FYN kinase, that has been crystallized with two binding partners, and design a new sequence that has low energy in complex with its two partners.

R-MSD is run completely within RosettaScripts, as just a combination of movers written specifically for the purpose of multistate design. The benefit of this is that you can add any other movers that are available within RosettaScripts to your protocol, i.e. backbone minimization, rigid body docking, atom pair constraints, etc. **All Rosetta commands were run with version 8641cc1735a37dff08c3f1857bbe3035908f7f04.** All analysis scripts are available for download at https://github.com/sevya/msd\_analysis\_scripts. Note: scripts provided in the analysis directory are dependent upon each other, and when moved from this directory may not function properly.

**PDB Preparation**

First I downloaded the PDBs from the RCSB and manually inspected them to remove all but one asymmetric unit. In this specific case, the PDB IDs I’m interested in are 1AVZ and 1M27. You can do this manually or with the clean\_pdbs.py script, located in /path/to/Rosetta/tools/protein\_tools/scripts/clean\_pdb.py. This script will download the specific chains of your structure and remove all non-proteinogenic molecules, which makes the structure easier to work with in Rosetta. The syntax for this is:

clean\_pdb.py 1avz ABC

clean\_pdb.py 1m27 ABC

In this case chain C in both 1avz and 1m27 is the FYN kinase that I want to design. However 1m27 has an extra leading valine residue at the N-terminus that is not present in 1avz. To simplify my protocol I removed this residue in PyMol before proceeding - this residue can also be removed using a text editor. Next, I reorder the chains to put the one protein common to both structures, FYN, as the first chain, rename the chains as A, B, and C, and renumber each chain starting from one. This makes my protocol slightly easier since both of my inputs are in the same format. The renumbering can be done by hand or with the script reorder\_pdb\_chains.py, which takes as input your desired chain order, desired chain ids, and the input and output PDBs. This script simply moves the order of the chains to the desired order and renames the chains, while also renumbering residues from 1 to N. Note that this does not change the coordinates of any atoms, just the order in the PDB file and the chain identifier. The command for this is:

reorder\_pdb\_chains.py --new\_chain\_order C,A,B --new\_chain\_ids A,B,C 1avz.pdb 1avz\_renum.pdb

reorder\_pdb\_chains.py --new\_chain\_order C,A,B --new\_chain\_ids A,B,C 1m27.pdb 1m27\_renum.pdb

Next I create 50 relaxed models from these starting PDBs, using the following commands, XML scripts and flags. Of the 50 relaxed models I selected the lowest energy model for the design process. The flags I use control the memory usage when Rosetta is building side chain rotamers (linmem\_ig), the number of extra rotamers to include in the library (ex1/2, use\_input\_sc), the number of models to make (nstruct), and a designation to include all side chain atoms (fullatom). For more information on Rosetta relax and available options see <https://www.rosettacommons.org/docs/latest/prepare-pdb-for-rosetta-with-relax.html>. Below are the commands used to create relaxed models.

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgccrelease @relax.flags -s 1avz\_renum.pdb -parser:protocol relax.xml

relax.xml:

<ROSETTASCRIPTS>

<SCOREFXNS>

</SCOREFXNS>

<TASKOPERATIONS>

<InitializeFromCommandline name=ifcl />

<RestrictToRepacking name=rtr />

</TASKOPERATIONS>

<FILTERS>

</FILTERS>

<MOVERS>

<FastRelax name=relax repeats=8 task\_operations=ifcl,rtr min\_type=lbfgs\_armijo\_nonmonotone/>

</MOVERS>

<APPLY\_TO\_POSE>

</APPLY\_TO\_POSE>

<PROTOCOLS>

<Add mover=relax />

</PROTOCOLS>

</ROSETTASCRIPTS>

relax.flags:

-database /path/to/Rosetta/main/database

-linmem\_ig 10

-ex1

-ex2

-in:file:fullatom

-out:file:fullatom

-use\_input\_sc

-nstruct 50

**Input files**

Once I have my processed PDBs I can generate the files needed for R-MSD. First I need to generate residue files (resfiles) specifying the designable and repackable residues for both of my complexes. Residues that are designable can be substituted with any other amino acid, whereas ones that are repackable can only be substituted with different rotational isomers (rotamers) of the same amino acid. For more information on resfile syntax and logic see <https://www.rosettacommons.org/manuals/archive/rosetta3.5_user_guide/d1/d97/resfiles.html>. In this case I want to design all residues on chain A that are at the interface of chain A and chains B+C. Since the two complexes have different binding partners they may have overlapping but not identical interface residues – in this case I only want to design interface residues common to both complexes. In addition I want to repack any residues on chains B+C that are at the interface. I need a residue file for each complex that specifies which residues are to be designed and repacked. The number of designable residues needs to be the same, but repackable residues can be unique to each complex. To get these residue files I use the following script and flags:

generate\_interface\_files.py --side1 A --side2 BC --design-side 1 --repack --output 1avz 1avz\_relaxed.pdb

generate\_interface\_files.py --side1 A --side2 BC --design-side 1 --repack --output 1m27 1m27\_relaxed.pdb

This identifies all residues at the interface between chains A and B+C, specifies side 1 as the one with designable residues, and signals to repack any residues at the opposing side of the interface. It also specifies a name for my output file, which will be followed by the extension .resfile. After generating my residue files I need to manually remove residues on the A chain that are at the interface of one complex but not the other, to make sure I end up with the same number of designable residues. The resfiles I used in my protocol are shown below:

1avz.resfile:

NATRO

start

12 A ALLAA EX 1 EX 2

13 A ALLAA EX 1 EX 2

14 A ALLAA EX 1 EX 2

15 A ALLAA EX 1 EX 2

16 A ALLAA EX 1 EX 2

35 A ALLAA EX 1 EX 2

48 A ALLAA EX 1 EX 2

1 C NATAA EX 1 EX 2

2 C NATAA EX 1 EX 2

3 C NATAA EX 1 EX 2

4 C NATAA EX 1 EX 2

5 C NATAA EX 1 EX 2

6 C NATAA EX 1 EX 2

7 C NATAA EX 1 EX 2

10 C NATAA EX 1 EX 2

12 C NATAA EX 1 EX 2

13 C NATAA EX 1 EX 2

16 C NATAA EX 1 EX 2

17 C NATAA EX 1 EX 2

20 C NATAA EX 1 EX 2

47 C NATAA EX 1 EX 2

48 C NATAA EX 1 EX 2

49 C NATAA EX 1 EX 2

50 C NATAA EX 1 EX 2

1m27.resfile:

NATRO

start

12 A ALLAA EX 1 EX 2

13 A ALLAA EX 1 EX 2

14 A ALLAA EX 1 EX 2

15 A ALLAA EX 1 EX 2

16 A ALLAA EX 1 EX 2

35 A ALLAA EX 1 EX 2

48 A ALLAA EX 1 EX 2

61 B NATAA EX 1 EX 2

63 B NATAA EX 1 EX 2

75 B NATAA EX 1 EX 2

76 B NATAA EX 1 EX 2

77 B NATAA EX 1 EX 2

78 B NATAA EX 1 EX 2

79 B NATAA EX 1 EX 2

82 B NATAA EX 1 EX 2

83 B NATAA EX 1 EX 2

85 B NATAA EX 1 EX 2

86 B NATAA EX 1 EX 2

**R-MSD Script**

Now that I have my input files I am ready to run R-MSD. The following script contains the R-MSD fixed backbone protocol:

<ROSETTASCRIPTS>

<SCOREFXNS>

<tal weights=talaris2013.wts >

<Reweight scoretype=res\_type\_constraint weight=1.0 />

</tal>

</SCOREFXNS>

<TASKOPERATIONS>

<InitializeFromCommandline name=ifcl />

<RestrictToRepacking name=rtr />

</TASKOPERATIONS>

<MOVERS>

<PackRotamersMover name=design scorefxn=tal task\_operations=ifcl />

<MSDMover name=msd1 design\_mover=design constraint\_weight=0.5 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd2 design\_mover=design constraint\_weight=1 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd3 design\_mover=design constraint\_weight=1.5 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd4 design\_mover=design constraint\_weight=2 resfiles=1avz.resfile,1m27.resfile debug=1 />

<FindConsensusSequence name=finish scorefxn=tal resfiles=1avz.resfile,1m27.resfile />

</MOVERS>

<FILTERS>

</FILTERS>

<APPLY\_TO\_POSE>

</APPLY\_TO\_POSE>

<PROTOCOLS>

<Add mover=msd1 />

<Add mover=msd2 />

<Add mover=msd3 />

<Add mover=msd4 />

<Add mover=finish />

</PROTOCOLS>

</ROSETTASCRIPTS>

In this case my design mover is a PackRotamersMover, which is given to each MSDMover as a submover. Note that I never actually call the design mover – it is called within the MSDMover. The four MSDMovers also specify a weight for residue constraints, which are ramped throughout the protocol, and a debug flag for extra output messages. The resfiles tag uses the files generated in the previous step to tell the MSDMover which residues should be linked together in multistate design. The resfiles don’t need to have designable residues at the same positions (i.e. position 1 on protein 1 can correspond to a position 10 on protein 2), but they must have the same number of total designable residues. **Note: R-MSD matches resfiles to structures by input order. You have to make sure that PDBs are specified on the command line in the same order as resfiles in your XML file.** FindConsensusSequence is a protocol to make sure that I end up with the same sequence on all states at the end of the protocol. It checks at each position specified in the resfiles if the two input PDBs have a different amino acid, and if they do it places each of the candidate amino acids onto all states, packs rotamers and checks the sum of energy across states. Whichever of the candidates results in the lowest energy across all states is accepted as the final identity.

I also need to setup a flags file that specifies Rosetta options – the following are the flags I used in my benchmark:

-in:file:fullatom

-out:file:fullatom

-database /path/to/Rosetta/main/database/

-linmem\_ig 10

-ex1

-ex2

-nstruct 100

-run:msd\_job\_dist

-run:msd\_randomize

The only flags specific to my protocol are the last two. Run:msd\_job\_dist is needed for the JobDistributor to be able to give multiple input poses to a mover at the same time, which is needed for multistate design. This protocol will fail and throw an error message without that flag. Run:msd\_randomize randomizes the order of input poses before applying each mover. This is not completely necessary for multistate design but is recommended, the reason being that there is slightly different behavior depending on the order in which PDBs are input. By randomizing the order before you keep this from biasing your results. More information on the other flags can be found at https://www.rosettacommons.org/docs/wiki/full-options-list.

**Running R-MSD**

Now that all my setup is done I can run the protocol. I’ll run it with the following command line:

/path/to/Rosetta/main/source/bin/rosetta\_scripts.default.linuxgccrelease @msd.flags -s 1avz.pdb 1m27.pdb -parser:protocol fix\_bb.xml -scorefile fix\_bb.fasc

This will generate 100 fixed backbone designs using R-MSD. For my backbone minimized designs I used all the same options, input files, and commands, with the only difference being my xml:

<ROSETTASCRIPTS>

<SCOREFXNS>

<tal weights=talaris2013.wts >

<Reweight scoretype=res\_type\_constraint weight=1.0 />

</tal>

</SCOREFXNS>

<TASKOPERATIONS>

<InitializeFromCommandline name=ifcl />

<RestrictToRepacking name=rtr />

<RestrictToInterfaceVector name=rtiv chain1\_num=1 chain2\_num=2,3 CB\_dist\_cutoff=10.0 nearby\_atom\_cutoff=5.5 vector\_angle\_cutoff=75 vector\_dist\_cutoff=9.0 />

</TASKOPERATIONS>

<MOVERS>

<PackRotamersMover name=design scorefxn=tal task\_operations=ifcl />

<MSDMover name=msd1 design\_mover=design constraint\_weight=0.5 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd2 design\_mover=design constraint\_weight=1 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd3 design\_mover=design constraint\_weight=1.5 resfiles=1avz.resfile,1m27.resfile debug=1 />

<MSDMover name=msd4 design\_mover=design constraint\_weight=2 resfiles=1avz.resfile,1m27.resfile debug=1/>

<FindConsensusSequence name=finish scorefxn=tal resfiles=1avz.resfile,1m27.resfile />

<TaskAwareMinMover name=min tolerance=0.001 task\_operations=rtiv type=lbfgs\_armijo\_nonmonotone chi=1 bb=1 jump=1 scorefxn=talaris2013 />

<FastRelax name=relax scorefxn=talaris2013 task\_operations=ifcl,rtr,rtiv repeats=1 />

</MOVERS>

<FILTERS>

</FILTERS>

<APPLY\_TO\_POSE>

</APPLY\_TO\_POSE>

<PROTOCOLS>

<Add mover=msd1 />

<Add mover=min />

<Add mover=msd2 />

<Add mover=min />

<Add mover=msd3 />

<Add mover=min />

<Add mover=msd4 />

<Add mover=min />

<Add mover=finish />

<Add mover=relax />

</PROTOCOLS>

</ROSETTASCRIPTS>

**MPI\_MSD File Preparation**

To run MPI\_MSD I used the same designable and repackable residues, and reformatted them for this application. Full documentation on the MPI\_MSD application is available at <https://www.rosettacommons.org/docs/latest/mpi-msd.html>. Briefly, I will describe the necessary input files. I need an entity resfile that specifies the residues that I am designing (FYN.entres), a correspondence file that maps my designable residues to an index (FYN.corr), and secondary resfiles that specify which additional residues are to be repacked (1avz.2res, 1m27.2res). The residues I include in these files are derived from the interface residues I used in R-MSD. In addition I need a fitness file (fitness.daf), and state files for each input pdb (1avz.state, 1m27.state).

The contents of my input files are shown below:

FYN.entres:

7

ALLAA EX 1 EX 2

start

1 A ALLAA EX 1 EX 2

2 A ALLAA EX 1 EX 2

3 A ALLAA EX 1 EX 2

4 A ALLAA EX 1 EX 2

5 A ALLAA EX 1 EX 2

6 A ALLAA EX 1 EX 2

7 A ALLAA EX 1 EX 2

FYN.corr:

1 12 A

2 13 A

3 14 A

4 15 A

5 16 A

6 35 A

7 48 A

1avz.2res:

NATRO EX 1 EX 2

start

1 C NATAA EX 1 EX 2

2 C NATAA EX 1 EX 2

3 C NATAA EX 1 EX 2

4 C NATAA EX 1 EX 2

5 C NATAA EX 1 EX 2

6 C NATAA EX 1 EX 2

7 C NATAA EX 1 EX 2

10 C NATAA EX 1 EX 2

12 C NATAA EX 1 EX 2

13 C NATAA EX 1 EX 2

16 C NATAA EX 1 EX 2

17 C NATAA EX 1 EX 2

20 C NATAA EX 1 EX 2

47 C NATAA EX 1 EX 2

48 C NATAA EX 1 EX 2

49 C NATAA EX 1 EX 2

50 C NATAA EX 1 EX 2

1m27.2res:

NATRO EX 1 EX 2

start

61 B NATAA EX 1 EX 2

63 B NATAA EX 1 EX 2

75 B NATAA EX 1 EX 2

76 B NATAA EX 1 EX 2

77 B NATAA EX 1 EX 2

78 B NATAA EX 1 EX 2

79 B NATAA EX 1 EX 2

82 B NATAA EX 1 EX 2

83 B NATAA EX 1 EX 2

85 B NATAA EX 1 EX 2

86 B NATAA EX 1 EX 2

fitness.daf:

STATE\_VECTOR avz 1avz.state

STATE\_VECTOR m27 1m27.state

FITNESS vmin(avz) + vmin(m27)

1avz.state:

1avz\_relax\_2\_0010.pdb FYN\_min.corr 1avz.2res

1m27.state:

1m27\_relax\_2\_0003.pdb FYN\_min.corr 1m27.2res

**Running MPI\_MSD**

To run MPI\_MSD I use the following command:

mpiexec -n 12 /path/to/Rosetta/main/source/bin/mpi\_msd.default.linuxgccrelease -database /path/to/Rosetta/main/database/ -entity\_resfile FYN.entres -fitness\_file fitness.daf -ms::pop\_size 100 -ms::generations 105 -ms::numresults 100 -no\_his\_his\_pairE -ms::fraction\_by\_recombination .02 -msd::double\_lazy\_ig\_mem\_limit 100 -ex1 -ex2

This runs the application on 12 processors and generates 100 output files.

**Design analysis**

To do my design analysis I first sorted by the fitness of all of my designs, which is the sum of energy of my input proteins. I analyzed the top ten designs for each of these three methods for fitness, sequence recovery, and similarity to evolutionary sequence profile. After identifying my top ten designs I used the Weblogo server to generate sequence logos, and the deep\_analysis script as a wrapper to calculate amino acid frequencies at each position and make my sequence logo. Deep analysis takes in a resfile to know which residues should be compared - however, just a note that you should make a separate resfile for only designable residues (FYN\_analysis.resfile), otherwise it will show a sequence logo for designable and repackable residues. The contents of this resfile are shown below:

FYN\_analysis.resfile:

start

12 A ALLAA EX 1 EX 2

13 A ALLAA EX 1 EX 2

14 A ALLAA EX 1 EX 2

15 A ALLAA EX 1 EX 2

16 A ALLAA EX 1 EX 2

35 A ALLAA EX 1 EX 2

48 A ALLAA EX 1 EX 2

Note: deep analysis does not link residues between complexes like R-MSD does. It’s most useful to analyze each input complex separately. However, since the result of my design run will be two complexes with exactly the same sequence at all designable positions, it’s really only necessary to analyze sequences from one of the complexes. An example command for this script is the following:

deep\_analysis --prefix 1avz\_fixbb\_ --native 1avz\_renum.pdb --stack\_width 30 --seq --format png --labels sequence\_numbers --res FYN\_analysis.resfile -s d \*pdb --path /path/to/weblogo

This will output a sequence logo, as well as a .tab file that contains all amino acid frequencies at all positions. From this file you can convert amino acid frequencies into a bitscore (which is equal to pi x log2(20 x pi) ) and calculate the native sequence recovery (defined as the bitscore of the native amino acid divided by total bitscore at a position).

**Evolutionary Sequence Profiles**

To generate an evolutionary sequence profile for each protein I used PSIBlast, with the following command:

psiblast -query fyn.fasta -db non\_redundant\_database.db -num\_iterations 2 -out out.txt -out\_pssm fyn\_pssm.txt -out\_ascii\_pssm fyn\_ascii\_pssm.txt

The ASCII PSSM contains amino acid frequencies for all positions in my FASTA file. I filtered by 1) residues that were specified in my resfile, and 2) residues that had a mutation in the top ten models produced by any of the three design protocols. I then compared this evolutionary PSSM to the design PSSM for each design method. To do this I calculated a squared difference matrix between the two PSSMs, and summed the difference over all amino acids at a given position. At each position, I subtracted this value from 2 and normalized by a factor of 2 to yield a percent similarity score. I then averaged the percent similarity over all positions to generate an overall percent similarity score.