From the PSVS Results Report, we could know that there are 20 models generated by CYANA. And main analyses that the report shows are Distance Violations per Model, Dihedral Angle Violations per Model, RPF Scores, RMSD, Ramachandran Plot Summary, Global Quality Scores and Visual tools.

1. Distance violations per model

Distance violations per model

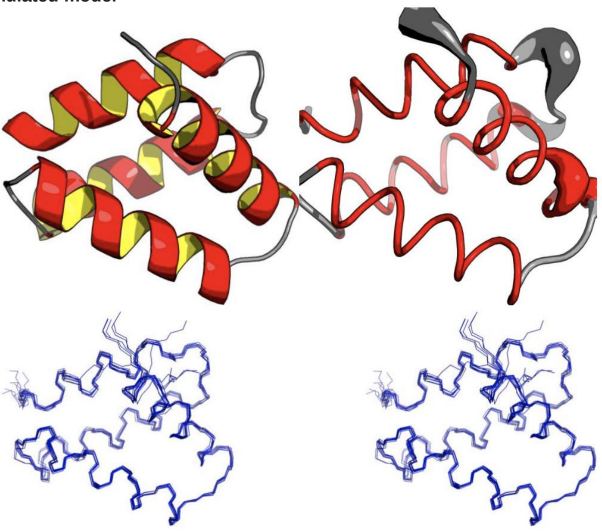
Calculated using sum over r^-6

$$0.1 - 0.2 \text{ Å} \quad 0.2 - 0.5 \text{ Å} > 0.5 \text{ Å}$$

 $0.2 \qquad 0.1 \qquad 0$

Experimental data determine the upper limit of distance. After calculation using sum of minus six square of limits (base values), PSVS will give number of distances (which are 0.1 - 0.2 Angstrom more, 0.2 - 0.5 Angstrom more and > 0.5 Angstrom more than base values) over number of models. Here, most cases are more than 0.5 Angstrom which are not good.

2. Simulated model



3. Several scores

RPF Scores

Recall Precision F-measure DP-score 0.919 0.972 0.945 0.834

RMSD All residues Ordered residues² Selected residues³
All backbone atoms 0.8 Å 0.3 Å 0.3 Å
All heavy atoms 1.1 Å 0.8 Å 0.8 Å

Ramachandran Plot Summary for selected residues³ from Procheck

Most favoured regions Additionally allowed regions Generously allowed regions Disallowed regions 93.9% 0.0% 0.0%

Ramachandran Plot Summary for selected residues³ from Richardson Lab's Molprobity

Most favoured regions Allowed regions Disallowed regions View plot View model summary 99.4% 0.6% 0%

Global quality scores

Program	Verify3D	ProsaII (-ve)	Procheck (phi-psi) ³	$Procheck (all)^3$	MolProbity Clashscore
-Raw score	0.32	1.05	0.30	-0.17	25.82
Z -score I	-2.25	1.65	1.49	-1.01	-2.91

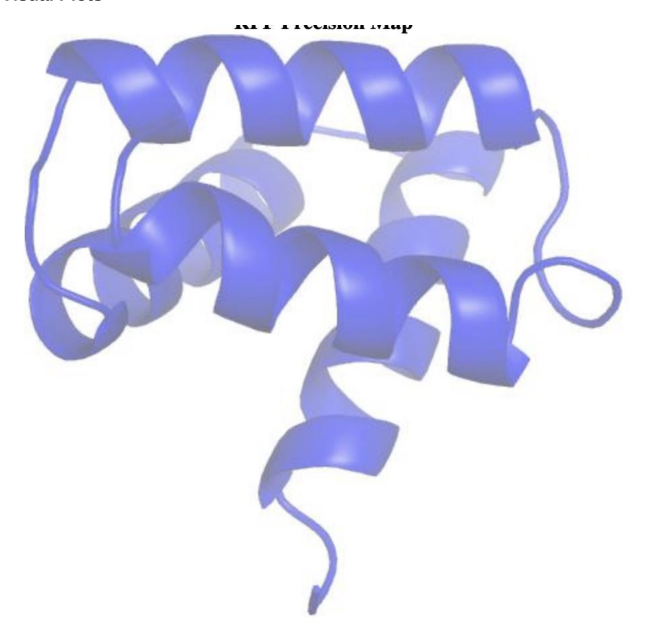
RPF Scores are scores to analyze the comparison between models and experiments data. Recall score is used to calculate how many contacts are matched with real contacts in data. Precision score is used to calculate how many real contacts that appear in expected contacts of models. The good result is, these two scores are comparable. F-measure score considers contacts in models and in data simultaneously. F-measure score is normalized into this range and turn out to be a new score which is called DP-score. DP-score is what we usually looks at to determine the quality of models. Generally speaking, DP-score > 0.7 and other three scores > 0.9 are expected. In this case, DP-score is 0.834 and other three scores are more than 0.9. So models are reliable. Using specific tools to determine quality of models are still necessary.

This part is calculation of RMSD of either backbone or heavy atoms. The reason that RMSD of all heavy atoms are more than that of backbone atoms is, heavy atoms also appear in side chains which may not converge. Since PSVS calculates mean values of RMSD of 20 models, RMSD will much low. The expectation of these two is, RMSD of all backbone atoms is less than 0.7~0.8 Angstrom and RMSD of all heavy atoms is less than 1.5 Angstrom.

It is good to see in Ramachandran Plot Summary, most favoured regions have 93.9% (from Procheck) and 99.4% (from Richardson Lab's Molprobity) and nothing in Disallowed regions.

There are 5 programs to calculate different quality scores. In the table shown above, raw scores are calculated directly from programs. Z-score is a measure of how many standard deviations below or above the population mean a raw score is. Therefore, the best Z-score is more than 0. However, since experimental data is not very perfect, the expected raw score is more than -0.5 and Z-score is more than -3. Generally, scores from MolProbity Clashscore (which is also calculated from Richardson Lab's Molprobity) and Procheck will be not so good if structures are from CYANA. In this case, we could see that all z-scores are good.

4. Visual Plots



It is RPF score of each residue. Bluer means higher score and redder means low score. In this case, light blue parts could be checked with data again to see if there is any improvement that could be done. The analysis of the plot below is similar.



