Comparative Protein Modelling

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Abstract: comparative protein modelling is predicting the 3D-structure of the given protein sequence by selecting a similar template sequence and modelling the sequence like that of the matched template sequence. We have different techniques to implement this time of modelling, in this project we are using Modeller tool to model our query protein based on the template protein. We also automated the process of protein modelling with modeler by suggesting a sample pipeline which saves time and space related issues and finally we evaluated our results using online tools such as TM-Score and Molprobity for obtaining different scores and results. For visual analysis, we have used chimera and Rasmol.

Motivation: proteins are very important functional units of our body. They are also used as chemicals, enzymes, medical drugs, and other hormones. They are building block of our body; dynamic maintenance of life is carried out by protein. All protein functions depend on the way it is structured i.e. on physical and chemical properties. Common thing of protein functionality is dependent on the type of amino acids that are grouped together to form a protein. So, to understand the functionality of the protein we need to understand its structure, for which we need to build a model of structure for the protein. With this motivation, we have built 10 of our taken query sequences to models and analyzed the properties of them under different conditions.

Results: With the given 10 proteins; 5 from CASP 11 and 5 from CASP 10 we have obtained better accuracy of TM-Scores for 6 proteins comparing with the CASP results. Results for each of the protein are explained separately in the results section. We have also implemented a sample pipeline for automation of the process of modelling using Modeller. Implemented few refinement steps on some proteins where the accuracy has been increased compared to the previous results before refinement.

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1 Introduction

The term comparative protein modelling or homology modelling or template based modelling are referred as the same where our own main goal is to model a protein 3D structure using the know templates. Here the templates are those with similar sequence to that of our unknown query protein. Based on the known protein i.e. our template we will model the query protein using the properties and alignment of the known template. The template contains sufficient information of spatial arrangement of residues and internal structure which helps in predicting our model. Comparative modelling of protein sequence is more reliable technique to obtain better results of protein structure modelling. Protein are one of most important functional units of our body, they do most of the work in cells and they are required for structure, function, and regulation of body's tissues and organs. The three-dimensional structure of the protein determines the functionality of the protein. The four levels of proteins i.e. the primary structure which is a sequence of amino acid residues determine the peptide chain. In the secondary structure, hydrogen bonds between the amino acids creates alpha helix, which is a spiral or coiled molecule and pleated sheet that looks like ribbon with regular peaks and valleys as a part of the fabric. The tertiary structure is for overall shape of the protein which are either globular or fibrous. Quaternary structures describe the proteins ap-

Our method of protein modelling starts with taking a query sequence. Query sequence is the one which we want to model as a three-dimensional structure for the protein then later identifying template and build model using the Modeller tool and we validate our results using varies validation techniques available online and do structural analysis of the proteins using visualization tools.

2 Methods

The whole process if protein modelling is described as follows:

- Select the required protein and get the sequence.
- Search for template in BLAST, PDB and other protein databases accordingly as required.
- Find the better matching sequence for the query sequence, this is our template. We may have single or multiple templates. Align the template with our query.
- Prepare our files in PIR format. Download the PDB formats for the templates. Save PIR formatted query file as .ali extension.
- Start the Modeller by giving input the query .ali file and other PDB files, according to the log file generated give inputs to the Modeller.
- vi. Validate the chosen best model and determine the accuracy.

To align the given sequences, we have used multiple sequence alignment techniques for multiple templates selected, some of them include T-Coffee and Clustal-Omega. At some cases, we have chosen only one template, where we have used Needleman-Wunch algorithm to align the sequences and converted all of them into PIR format. The pipeline that we have implemented asks for sequence and automatically converts them into PIR

format and saves query in .ali file and rest template sequences as .pir extension. The PDBs are automatically download once given the template IDs after script 1 execution. The program that we wrote has BioPython packages and uses NCBI pdb API call to download the PDB files given the template IDs. Alternatively, we can use REST API to download the pdb files. The REST API is of XML format.

For validation, we are using different tools available online and for visualization purpose we have used Chimera as our tool. The various online techniques that we have used to measure the accuracy of our model include TM-Score, Molprobity scores, and RMSD score.

All the steps are cleared explained clearly below where we have mentioned each of options that we have chosen for protein modelling.

The software distribution that we used for this project is described as below:

- a. Test cases i.e. Protein queries are taken from CASP website. CASP10 and CASP 11 are chosen to select 10 proteins which include:
 - CASP11 targets T0856, T0843, T0806, T0837, T0792 and
 - CASP10 targets T0757, T0666, T0678, T0651, T0694
- b. For template identification: BLAST, PDB, and SWISS-Model.
- Sequence alignment: Needleman-Wunch algorithm, and T-Coffee
- d. Software for protein modelling used is Modeller 9.17.
- Various languages used are python 2.7 and JAVA (Needleman-Wunch algorithm)
- f. Protein visualization software's: Chimera, and Rasmol
- g. Project Management and version control: GitHub

Process automation

The whole process of comparative protein modelling is as follows, Initially we find the templates and select the best templates matching our query sequence, then we align them and start the process or modelling by giving all the files as needed to the Modeller and based on the output obtained we decide whether to refine the model or end the process. But for running Modeller we need to provide certain inputs at each step. To ease the process, we have automated the system using python scripts. For this purpose we have implemented a sample pipeline where the process of using Modeller is made lot easier compared to the standard approach. Instead changing inputs in the script files every time, user needs to just enter the

```
Administrator. Command Prompt - python start.py

Picrosoft Windows (Upresion 6.1.7681)
Copyright 607 2889 Hisrosoft Copporation. All rights reserved.
C:\windows.pysten222.ed C:\Program Files\Modeller9.17\bin\Test-Pipeline
C:\Windows.pysten222.ed C:\Program Files\Modeller9.17\bin\Test-Pipeline
C:\Program Files\Modeller9.17\bin\Test-Pipeline
D:\Windows.pysten222.ed C:\Program Files\Modeller9.17\bin\Test-Pipeline
ESWISUREEYPD-IDSEURN ILISHANGANGITISSIKSEYRKLIGNPFPLH---DNUIDPLLTIPNUTAECSESGKRIF

NIKAN
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Fig 1:start.py script; User entering query sequence

sequences initially and template IDs later for each of the script to run. The script automatically converts the given aligned query sequence into PIR format as saves it as .ali extension file. At each script execution, based on the output generated by the script and after evaluating the results, we given the template ID as input to the next script and the process is the same till the end. Sample screenshot for the pipeline is as below, where the program asks for user input query sequence.

When the user enters the aligned sequence, the script process the sequence and ask for user to confirm by pressing enter, once the user clicks enter the first script of Modeller starts running. Now for each script, once the execution is complete the console ask for user input. From script 2 users' needs to enter just the template IDs and model ID for the last one accordingly after evaluating the log results generated by the Modeller. The PDB files required for the template can be automatically downloaded when we use BioPython libraries available for python. So, taking these dependencies we can download the PDB files directly. Another option is we can use REST API calls provided by PDB website where are already available in XML format. So, we use them to download the PDB files required for running script2. Similarly, the pipeline process is continued for each of the proteins.

3 Results

With the given ten CASP sequences, we divided the sequences among ourselves in the group and completed the process individually later we validated our results by exchanging the proteins we modelled among ourselves. For template search, we have used BLAST as our primary source and we played around with different option available with BLAST to match the best template that we can obtain from our resources. For most of the results that are obtained we have used PSI-BLAST option in BLAST and searched for template in PDB database. Each of the protein results are explained below separately and all the results obtained are discussed individually at first and then comparison is made with CASP results to show the accuracies of our protein model that we have generated.

3.1 CASP 10 and CASP 11

For the process of modelling the proteins, we have chosen 5 query sequences for CASP 10: T0757, T0666, T0678, T0651, T0694 and CASP 11: T0856, T0843, T0806, T0837, and T0792 respectively we have followed the same procedure as discussed in the pipeline. Initially we have taken the query sequence and searched for template in BLAST with option as pdb database and PSI-BLAST searching techniques. Depending the results obtained and percentage of identity between the chosen templates and query sequence we have used either of pair wise sequence alignment with Needleman-Wunch algorithm and T-Coffee for multiple sequence alignment between query and templates which are explained clearly for each of the protein below with results obtained for each of the protein. For pair-wise sequence alignment we have implemented a JAVA program from GitHub and for multiple sequence alignment we have used T-Coffee and Clustal-Omega accordingly when required and for validation and refinement purpose.

3.1.1 Protein modelling using Modeller

We have taken 5 protein query sequences from CASP 10 i.e. T0757, T0666, T0678, T0651, and T0694 respectively and from CASP 11 i.e. T0856, T0843, T0806, T0837, and T0792 as our query sequence individually and the models are selected based on the scores of DOPE, Molpdf, GA341 where Molpdf is Modeller objective function, DOPE is discrete optimized protein energy and GA341 as a percentage sequence identity between the template and model as a parameter. The results obtained are as follows:

• T0757:

we have taken the query sequence from CASP and used BLAST to obtain the template sequence, although we got couple of matching sequences but as result we took 4gak as out template and performed pair-wise sequence alignment between each of the query and template. Initially we tried with multiple sequence alignment but the TM-Score obtained for the protein sequence alignment is around 0.8, but when we tried pair-wise alignment with selected template we obtained TM-Score as 0.9629. The result of the Modeller scores is as follows:

Filename	Molpdf	DOPE score	GA341 score
Model1.pdb	1250.1788	33 -29520.337	89 1.00000
Model2.pdb	1209.114	l <mark>62 -29355.9</mark> 3	359 1.00000
Model3.pdb	1170.828	37 -29699.65	625 1.00000
Model4pdb	1374.1108	34 -29354.617	19 1.00000
Model5.pdb	1231.565	80 -29481.416	502 1.00000
Model6.pdb	1167.898	32 -29369.69	531 1.00000

We have chosen model3 for our result which has better TM-Score compared to other models. We have also considered Model3 as a part of refinement which has better Molprobity score 2.02 compared to Model2 which has score of 2.43, so as a refinement we have reconsidered Model3 as our solution even though it has high DOPE score but least Molpdf score.

• T0666:

For T0666 we have started the analysis and obtained multiple templates, after analysis we have chosen 3napA and 3ux4A as our resultant templates and performed multiple sequence alignment with query. The diagonalization matrix obtained as a comparison is as follows:

3napAA	.@23u	x4AA@
3napAA@2	264	2
3ux4AA@3	1	180

Based on the results which are bit confusing to choose the better template, we have performed modelling taking both the templates separately and after obtaining results we have chosen best template and model based on the TM-Score obtained for each of them respectively. Finally, we have chosen 3ux4A as the best matching template which and the results for model structures are as follows:

>> Summary of successfully produced models:

Filename Molpdf DOPE score GA341 sc	ore
Model1.pdb 1022.91730 -23121.25391 1.00000	
Model2.pdb 1053.24976 -23116.05078 1.00000	
Model3.pdb 950.99738 -23461.31836 0.99997	
Model4.pdb 1070.12378 -23245.58594 0.99999	
Model5.pdb 1003.10345 -23420.93164 0.99994	

Based on scores, we have choosen model2 as our best model which as average of scores as well comparatively and obtained a TM-Score of 0.9042 and Molprobity score as 2.92.

T0678:

We have performed pair-wise sequence alignment with the query protein taking 4epz as our template. The results obtained are better comparing to multiple sequence alignment that we have performed initially, later with refinement we have chosen pair-wise sequence alignment as best choice for this protein based on our template models. The results obtained for models are follows:

>> Summary of successfully produced models:

Filename	Molpo	df DOPE score	GA341 score
Model1.pdb	643.56921	-18270.39844	1.00000
Model2.pdb	613.73462	-18336.48047	1.00000
Model3.pdb	1147.81177	-17304.24219	1.00000
Model4.pdb	595.51093	-18572.18359	1.00000
Model5.pdb	590.30353	-18337.23242	1.00000

We fixed model2 as our best model based on the results of TM-Score of 0.9222 and Molprobity score as 2.19.

• T0651:

We have performed multiple sequence alignment for this protein later we have analyzed our results after final decision of template and model selection, here we noticed multiple-sequence alignment performed better by few points compared to pair-wise sequence alignment. The results obtained for models of chosen template are as follows:

>> Summary of successfully produced models:

Filename	Molpdf DOPE score	GA341 score
Model1.pdb	1590.79980 -30183.49805	1.00000
Model2.pdb	1579.90295 -30485.20117	1.00000
Model3.pdb	1718.90918 -30320.98438	1.00000
Model4.pdb	1441.80750 -30527.74219	1.00000
Model5.pdb	1662.43384 -30278.29883	1.00000
Model6.pdb	1435.20447 -30606.22852	1.00000

We have finally decided to take model6 as our result model based on taking average scores of DOPE and Molpdf. The TM-Score obtained is 0.97 and Molprobity score of 2.18.

• T0694:

Based on the results of BLAST, we have performed multiple sequence alignment of templates and query using T-Coffee and the results obtained for models are as follows:

>> Summary of successfully produced models:

Filename	Molpdf	DOPE score	GA341 score
		20552 50545	4 00000
Model1.pdb	1435.19836 -	38772.60547	1.00000
Model2.pdb	1422.11194 -	38440.36719	1.00000
Model3.pdb	1332.25159	-38742.54688	1.00000
Model4.pdb	1465.64661 -	38483.46484	1.00000
Model5.pdb	1380.60278 -	38905.53125	1.00000

We have chosen Model3 as our resultant model taking average of scores obtained and we have achieved a TM-Score of 0.9576 and Molprobity score of 2.21 which is comparatively better solution based on analysis of CASP results which we will discuss later in the paper.

T0843:

We have used multiple sequence alignment of different templates with query sequence and obtained better Molprobity score than CASP predictions available on the website. The TM-Score is also appeared to be more reliable and overall, we scored almost equal TM-Score and better Molprobity score compared with top appeared results. The models generated has the scores as below:

>> Summary of successfully produced models:

Filename	Molpd	f DOPE score	GA341 score
Model1.pdb	1987.00964	-45485.44531	1.00000
Model2.pdb	1966.13806	-45068.55859	1.00000
Model3.pdb	1899.45654	-45152.75781	1.00000
Model4.pdb	1810.79456	-45319.01563	1.00000
Model5.pdb	1907.92151	-45049.98828	1.00000

based on average of the results we have chosen model3 as our best model and it has better Molprobity compared to others.

• T0806:

We have used pair-wise sequence alignment for this model and obtained best TM-Score compared with the CASP results. Initially we tried with multiple sequence alignment but comparatively pair-wise sequence alignment performed better and the results obtained are as follows:

>> Summary of successfully produced models:

Filename	Molpd	If DOPE score	GA341 score
Model1.pdb	1111.07874	-31527.54883	1.00000
Model2.pdb	1288.70447	-31395.63867	1.00000
Model3.pdb	1436.59229	-31155.95898	1.00000
Model4.pdb	1199.31531	-31481.81055	1.00000
Model5.pdb	1460.71851	-30915.53516	1.00000
Model6.pdb	1187.11096	-31542.72852	1.00000

We have choosen Molpdf score as evaluating factor and based on the result as model1 performed better we made further analysis with TM-Score and Molprobity as an analysis factors and determined model1 as our best results which outperformed on CASP results obtaining TM-Score as 0.9682.

• T0837:

Specifically, to this protein, results can be improved and lot of refinement can be performed. The template that we identified with different means didn't yield any better results for us. We have tried using many techniques, like pair-wise, multiple-sequence alignment with T-Coffee and Clustal-Omega. Although TM-Score improved after identification of better template and pair-wise sequence alignment, still it is in between random and accepted model. Here we show TM-Score and Molprobity score than DOPE scores as they are important for validation and we've obtained the scores as follows:

TM-Score: 0.234 Molprobity: 3.18

• T0792

This protein has the similar case as above, but we have obtained fair results upon refinement. Initially the TM-Score was as low as 0.12. Later with refinement and proper identification of templates the score improved to 0.4172 and for few of the matching sequence i.e. templates didn't show any TM-Scores as there were no matching residues in modeled result and template. The various TM-Scores before refinement and after refinement are as follows:

Before refinement

Templates 54a9, 5cd7, 5a49 has no TM-Scores; template 4obm has TM-Score of 0.12

After refinement

2i6e gave me a score of 0.16 and 3s93 has score of 0.4172.

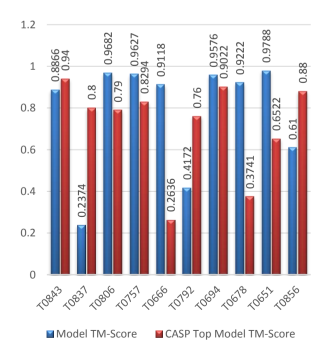
We have performed different techniques with 3s93 which is best template available, out of all the techniques multiple sequence alignment with templates that are selected after refinement gave better results.

3.2 Evaluation

For analyzing the accuracy of our data, we have compared our results with CASP results and based on the TM-Score results that we have obtained, 4 of our proteins outperformed the CASP results and are as shown below in Fig.2. TM-score obtained for the proteins T0806, T0757, T0666, T0694, T0678 and T0651 outperformed compared to that of CASP results and

Fig2: TM-Score comparison with best CASP models

TM-Score Comparision



When we consider protein T0843 it almost nearly modelled to that of CASP result and it has better Molprobity results compared to that of CASP result so overall, we can say it is well modelled comparing with CASP results. Few of the results are obtained upon refinement of the original modelling which was clearly explained in the paper above and the data table for the above graph is as shown below in Table 1.

Table1: Comparison of TM-scores with CASP best models

Protein	TM-Score	CASP best TM-Score
T0843	0.8866	0.94
T0837	0.2374	0.8
T0806	0.9682	0.79
T0757	0.9627	0.8294
T0666	0.9118	0.2636
T0792	0.4172	0.76
T0694	0.9576	0.9022
T0678	0.9222	0.3751
T0651	0.9788	0.6522
T0856	0.61	0.88

and the result comparison with CASP with molporbity scores are viewed in below table along with the type of alignment that we have used for our protein modelling is given in the below table 2.

Here, for CASP 10 sequence comparison the website hasn't provided the information on TM-scores but the scores for Molprobity are alone provided. So, we have considered the native structure to determine the TM-Score of CASP top predicted model with best Molprobity score. Here the native structure are the templates considered for modelling the protein, hence it is reliable to calculate the TM-Score with native structures. The native templates are taken from CASP website and used for calculation of TM score for the one with best Molprobity score. For the proteins from CASP 10, i.e. T0757, T0666, T0678, T0651, and T0694 the TM-score results are compared with the result of model with best Molprobity score in the CASP website and choosing native structure as template.

In the below table 2, we have made our comparisons with CASP Molprobity scores, the given CASP-Molprobity scores are the best scores chosen from the list of scores available after filtering them. The table also provides information on the type of sequence alignment that we have used for kind of protein modelling.

Here we want to specify the sequence alignment to analyze the scores obtained with sequence alignment, and we have noticed the scores either increase or decrease or remains the same. Hence it is hard to decide about following either of sequence or multiple template alignment.

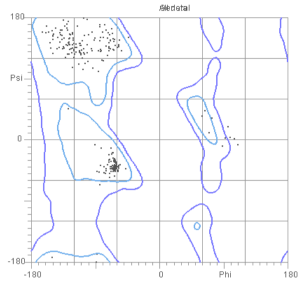
Table2: Molprobity and sequence alignment use; comparing Molprobity scores with best CASP models

<u>Protein</u>	Alignment Used	Molprobity Score	CASP-Molprobity Score
T0843	Multiple	2.42	3.02
T0837	Pair-wise	3.18	1.58
T0806	Pair-wise	2.04	1.37
T0757	Pair-wise	2.02(5 th in the CASP list)	0.83
T0666	Multiple	2.92	1.23
T0792	Multiple	3.02	1.18
T0694	Multiple	2.21	0.7
T0651	Multiple sequence	2.18 (13 th in CASP List)	1.27
T0678	Pair-wise	2.19	0.74
T0856	Pair-wise	2.82	1.66

We have also considered the Ramachandran plots to evaluate our data, here in this paper we want to discuss plots of two proteins using the plot graphs, one protein being the best results we obtained and the other is bad model that we have predicted. We have taken protein T0757 to validate our results as best model. T0757 has TM-Score of 0.9627, Molprobity score of 2.02 which is 5th place in the CASP list and it has favored Ramachandran plots of 98.37% (>98% is considered as our goral). We have used online Molprobity tool to validate our model for which the above results are obtained. The Ramachandran plots for generated model is as follows, initially we have taken favored regions of the plot and most of the points are in favored region except few, later we considered the allowed regions which shows very few of them being outliers and most of them being very near to the allowed region. This can be viewed in below

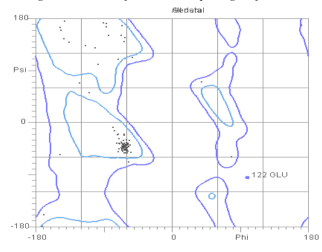
figure 3 below and with accepted regions, number of outliers were reduced.

Fig3. Ramachandran favored and accepted regions for T0757



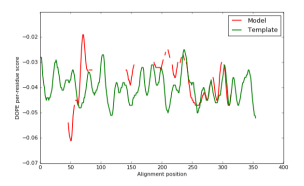
Next, we have taken the model which was generated by Modeller but with bad results compared to the CASP top model results for this we have considered the protein T0837 having a TM-Score of 0.2374 and Molprobity score as 3.18. The TM-Score specify it is almost a randomly generated model and it has Ramachandran favored plots as 90%, in the graph we identified how scattered the points in the above region of -180 and psi also, we can noticed that 122 GLU is an outlier and far away from the favored region and we can notice many points being away from favored region. Now when we consider the accepted regions in the graph as shown in fig4, still 122 GLU is far away but few points now are considered as accepted regions hence we have about 90% of favored regions as the result.

Fig4. Ramachandran favored and accepted regions for T0837



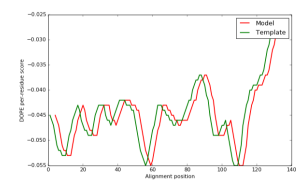
We also considered the plot generation of Modeller which finds the problem with modelled structures to identify the alignment problem with respective to DOPE scores. As T0837 has very less TM-Scores which might be due to less number of residue matches, we plotted the model vs template graph for T0837 and obtained the graph as shown in the figure 5.

Fig.5: Model vs Template plot for T0837



Here we can see that alignment positions comparing with model and selected template aren't matching properly and the in the above graph the red line represents the model and green line represents the template. The red line was plotting down at the beginning showing the varying match with the template. This tell us that the alignment is not proper and can be improved a lot. Also template selection is one of the important tasks of modelling, while selecting templates we need to make sure to select the one with least e values. Similarly when we consider a best modelled protein, where we have taken T0757 as our example, the graph obtained comparing modelled structural sequence alignment and template is as shown in fig 6

Fig6. Model vs Template plot for T0757



Here we can notice how well the template and model are aligned together, this shows the similar alignment positions of template and model and hence the score of TM-Score is high due to high number of matching residues. The e value of selected template is also less and alignment position comparing with DOPE score is pretty much interesting that they are almost aligned similarly which shows it to be a very good model.

3.3 Protein View using Chimera

Sample visualization for protein structure modelled is as shown in the Figure 7

Fig7: Protein view of T0806 using chimera-Ribbon view

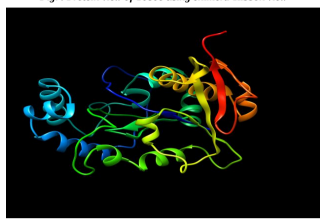
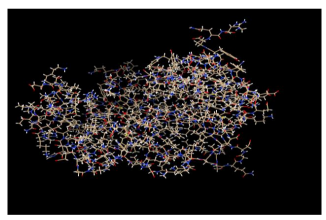


Fig8: Protein view of T0806 Chimera-All atom view



Which was represented in Ribbon view and Figure 8 as all atom view using chimera protein visualization software. We can also perform visual analysis using this software where we can see the bonds present and performed many structural and different kinds of visual analysis such as seeing the position of atoms, types of atoms and structural view of the model that was generated etc.

3.4 Case Studies

- We had a problem with protein identification initially as we have chosen pBlast as our option in BLAST search and later we have refined our results and identified better templates using PSI-BLAST option.
- We faced a lot of challenge with choosing the best of modelled templates, mainly because of scores obtained with Modeller as they are uneven and hard to select the correct one. For

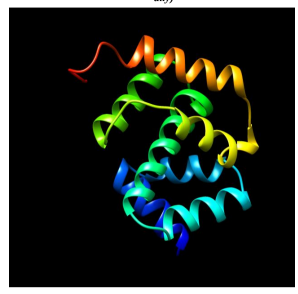
example, the scores obtained for T0651 protein are as shown in fig4. As we can see that obtained Molpdf, DOPE score and GA341 score for protein in the above Fig9. Are uneven so for best protein selection we have chosen the model with average scores, if there are bad results, we have chosen the one with best scores for each of category and tried with those models.

Fig9: T0651 Script 4 results using Modeller

> Distribution of short non-bonded contacts: DISTANCE1: 0.00 2.10 2.20 2.30 2.40 2.50 2.60 2.70 2.80 2.90 3.00 3.10 3.20 3.30 3.40 DISTANCE2: 2.10 2.20 2.30 2.40 2.50 2.60 2.70 2.80 2.90 3.00 3.10 3.20 3.30 3.40 3.50 FREQUENCY: 0 33 63 222 192 281 273 290 344 358 383 << end of ENERGY. >> Summary of successfully produced models: DOPE score Filename molpdf GA341 score asea1.B99990001.pdb 2208.54395 -18450.60156 0.02578 asea1.B99990002.pdb 1936.98352 -18376.74609 0.01414 qseq1.B99990003.pdb 2510.62158 -17843.83203 0.00694 asea1.899990004.pdb 0.04183 2116.11377 -18231.61719 qseq1.B99990005.pdb 2779.90063 -17003.68555 0.04731 qseq1.B99990006.pdb 2255.66235 -18772.46484 0.01694 Total CPU time [seconds]

- There are some case studies that we identified while carrying on the process such as, the sequence alignment didn't play a major role in obtaining an accurate template, but templates play a major and most of the role in protein modelling and in obtaining best model. The sequence alignment has very minimal role in modelling accurate 3D-structure of a protein. For example, we have considered the sequence T0792 for which
- We tried the sequence alignment of query with templates i.e. with both pair-wise and multiple sequence alignment along and other option with no sequence alignment for query with any of the templates. For a surprise, still the final obtained scores are the same for both cases. So, we concluded that major part of protein modelling depends on the template that we choose.
- Visual interpretation of protein doesn't always yield better results, and we can't assume that a better structured modelled has better scores. For example, if we visualize the below protein in fig10, it looks like a very well structured protein but when we validated the results, the TM-Score obtained for above protein model is just 0.2287. Hence visual interpretation based on structural analysis may completely fail for protein model.

Fig10: Ribbon View of a well-structured modelled (Visually)



- We also noticed that geometry of the structure plays very important roles in structuring a protein modelling, we can improve the score of protein model by refinement of the best model obtained.
- Another biggest confusion we have is with sequence alignment, at some cases we noticed that sequence alignment doesn't really play a major role. But some other cases the results were changed based on the sequence alignment that we have considered. For some cases with pair-wise sequence alignment we obtained better results, for some other cases with multiple sequence alignment. At time, we didn't find any difference between with and without alignment as both yielded same results for some of the test cases. Hence it is hard to identify the best practice but with results obtained, it is to be noted that pair-wise sequence is bit better compared to multiple but we have aligned only after identifying the template, that's a bad idea here.

Future analysis

We can perform lot more refinement to obtain better proteins. In the future, we want to identify better template by implementing machine learning techniques or by selecting wide range of databases, as we noticed that template plays a major key role in modelling the protein structure. The refinement process can be concentrated more towards the structural analysis of the protein than the sequence alignment, as the bon angles, bond lengths and other properties plays a key role in modelling of protein. Finally, we want to implement a better pipeline with visuals and easy to use integration on the front end for our pipeline there by automating the whole process of modelling and concentrating heavily on the data

analysis and data mining part of protein modelling. This helps a lot in time saving and implementation process.

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