

Analysis Protein Drugs Design Program with Quality and Structure Prediction

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ABSTRACT

A complete lack of hair will result when the "hairless protein" associated with the hairless gene, which is required for hair growth, ceases to function. The coordinates of this gene are 22027873-22045326 on chromosome 8. The JmjC domain superfamily includes the hairless gene, which also contributes to the process of histone demethylation. The domain sequence is 212 amino acids long and spans residue positions 946 to 1157 in the hairless protein, which has 1189 residues. JmjC domains have been discovered in more than 100 eukaryotic and bacterial sequences based on considerable sequence similarity, including the human hairless gene, which is mutated in people with alopecia universalis. To homology model the JmjC domain in the hairless protein, we have tried the bioinformatics method. NCBI-BLASTP, EBIClustalW, SMART, 3D-PSSM, DeepView/Swiss-PDB Viewer, PyMOL, and WhatCheck are the tools and programs utilized in this work. Using the template crystal structure of the likely antibiotic production protein from *Thermus thermophilus* HB8, the structure of the JmjC domain is predicted. Modeled domain structure's minimized energy value was -3394.570 KJ/mol. The simulated domain structure was validated using the WHAT IF-Proteins Model Check tool.

Keywords: development, design, protein

I. INTRODUCTION

The maintenance of hair development depends greatly on the mammalian hairless (*hr*) gene. The hairless protein 1 protein, which has 1189 amino acids, is encoded by the *hr* gene. Shrivastava, et al. 320 International Journal of Bioinformatics and Biological Science: v.1 n.3&4 Sept-Dec 2013 Although the *hr* gene was discovered around 75 years ago, the biochemical function and structure of the protein it encodes have not yet been anticipated. Alopecia universalis congenita (ALUNC) and atrichia with papular lesions are both conditions caused by *hr* gene defects. (APL). APL is an autosomal recessive condition that is defined by papillary lesions throughout the majority of the body and almost complete absence of hair, while ALUNC is a rare autosomal recessive form of hair loss that is characterized by hair follicles without hair. Congenital hair loss, whether total or partial, can occur alone or in conjunction with other abnormalities. Most isolated congenital alopecia families, according to reports, follow an autosomal recessive pattern of inheritance. The gene for autosomal recessive congenital alopecia has previously been linked to a 15 cM area on chromosome 8 p21–22 3 in a large inbred Pakistani family where affected individuals exhibit universal congenital alopecia, which is the lack of any hair at all.

Families with congenital atrichia have been discovered to have recessive mutations in the *hr* gene, which was originally located near to the MU interval 2. A rare autosomal recessive variant of complete alopecia known as atrichia with papular lesions (APL) causes widespread body-wide papular lesions made up of keratin-filled cysts as well as hair loss that begins shortly after birth. The pathophysiology of this condition has been linked to mutations in the *hr* gene, a putative single zinc finger transcription factor 4. A significant development in functional proteomics is the comparative homology modeling and 3D structure prediction of the JmjC domain in hairless proteins. In order to align the target sequence of the hairless protein with the template sequence and structure, the study's goals were to (i) identify a template or parent structure related to the target sequence, (ii) find secondary structures for the target and template sequences, and (iii) identify all well-conserved portions of the alignment between these two. (iii) Using the alignment of the target and template, one may determine the backbone coordinates for the target sequence. Using our understanding of the factors that determine protein structure, we can (i) build the loop for target sequence segments for which coordinates cannot be determined from the template due to insertions and deletions in the alignment (typically in loop regions of the protein), and (ii) build side chains determined by the target sequence onto the backbone model created from the template structure and loop construction.

II. MATERIALS AND METHODS

Finding protein structures related to the target sequence and using some of them as templates was the first step in comparative modeling of hairless proteins. For additional investigation, protein sequences from various species were downloaded from NCBI (found at www.ncbi.nlm.nih.gov) and saved in the Fasta file format.

The target sequence of the hairless protein was searched against sequence databases like PIR, TrEMBL/Swiss-Prot, and structure databases like PDB and SCOP 6. It was submitted to the NCBI-BLAST server to retrieve the closest relevant homologs. The hairless protein's amino acid sequence was acquired from SWISS-PROT 7. As a reference sequence 8, the Swiss-Prot sequence with Primary (citable) accession O43593 was utilized. A basic local alignment (NCBI-BLASTP; <http://www.ncbi.nlm.nih.gov/blast/Blast.cgi>) 9 search was conducted to compare the sequence of the hairless protein with a library or database of sequences, identify library sequences that resembled the sequence of the hairless protein, and find templates by selecting the PDB database as the target database search. To identify and annotate genetically mobile domains and to analyze the domain structures found in hairless protein, the Simple Modular Architecture Research Tool (SMART; <http://smart.embl-heidelberg.de/>) 10 was employed. SMART results and the JmjC domain sequence were used as templates for multiple sequence alignment using the ClustalW program (<http://www.ebi.ac.uk/Tools/clustalw/index.html>) 11.

To get the best possible global sequence alignment, ClustalW alignment was once more manually altered and realigned using ClustalW with the default values for Gap Opening, Gap Extension Penalty, and DNA weight matrix. Then, using this multiple sequence alignment file 5, phylogenetic trees were constructed. Once the best templates have been chosen, the sequences of the templates and the target were aligned using ClustalW in order to acquire the best alignment while modeling where appropriate. The target sequence for this alignment was the JmjC domain in the hairless protein. To find protein fold recognition, the threading approach of the three-dimensional protein secondary structure modeling procedure (3D-PSSM; <http://www.sbg.bio.ic.ac.uk/3dpssm/index2.html>) 12 was used. Software called DeepView (Swiss-PdbViewer) is employed to model a protein 13. The DeepView sequence alignment option was chosen, and gaps were manually added using the multiple sequence alignment-ClustalW output and the 3DPSSM alignment of the template with the query protein, which is the hairless protein.

The alignment feature in DeepView was used to fit the superimposed models of the JmjC domain and the antibiotics synthesis protein from *Thermus thermophilus* HB8 (PDB ID: 1V70 Chain A) for model creation. A web-based integrated service devoted to protein structure homology modeling is called the SWISS-MODEL Workspace. The generated project file was uploaded to the SWISSMODEL automated homology model-building server 15 for model calculation. It is developed for modeling, visualization, and analysis of biological systems such as proteins, nucleic acids, lipid bilayer assemblies, etc. Using the WHATIF-Protein Model Check tool 16, a stereochemical quality check was conducted on the provided model.

A few problematic side chain conformations were found and fixed. DeepView was used to minimize energy consumption on the final structure, and the procedure was repeated. The protein evaluation server WHAT IF (<http://swift.cmbi.ru.nl/servers/html/index.html>) protein evaluation server, which includes several other protein evaluation tools, was employed in the quality assessment of projected domain structure. In order to evaluate and identify abnormal bond angles between heavy atoms in amino acids, we used the Ramachandran plot.

III. RESULTS AND DISCUSSION

The amino acid sequence in Fasta format was retrieved from the SwissProt accession number of the hairless protein for the BLAST search. Results from BLASTP (Figure 1a, 1b) demonstrated that bit scores and anticipated alignment values were very subpar and did not demonstrate a meaningful alignment. Because the alignment bit scores were so low, it was clear that no useful template structure was available to predict the structure of the full hairless protein.

The JmjC domain was discovered using SMART tool in the hairless protein. The result showed that the domain was present in the hairless protein sequence from residue position 946 to 1157 at an E-value of 3.10×10^{-40} (Figure 2a).

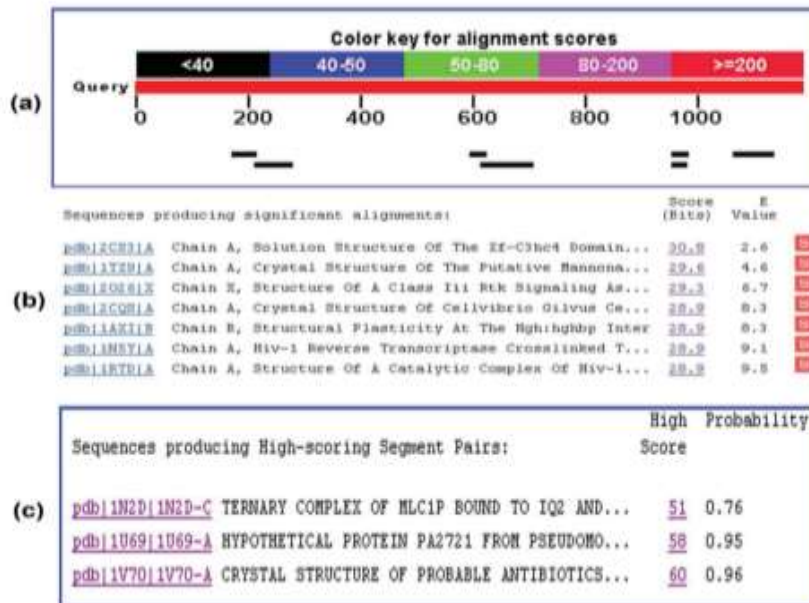


Figure 1

In Shrivastava, et al. 322 International Journal of Bioinformatics and Biological Science: v.1 n.3&4 Sept. 1: (a) Distribution of seven BLAST hits on the hairless protein query sequence; the black color bar graphic indicates that the sequence length of the Blast hit is fewer than 40 residues. The BLAST hit's bit score (b), with the lowest value suggesting a higher number of gaps and substitutions related to each matched sequence. (c) SMART-BLAST hit, which indicates a higher likelihood of aligning the JmjC domain with three of the templates based on the score value of the hit.

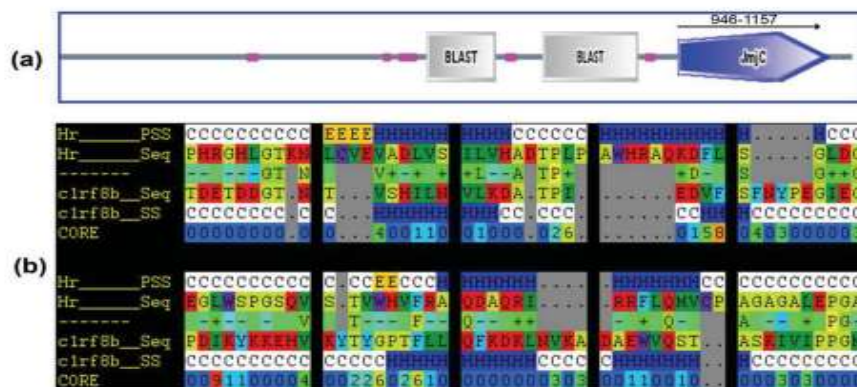


Figure 2

Human Hairless Protein Homology Modeling International Journal of Bioinformatics and Biological Science: v. 1 n. 3&4 Sept.-Dec. 2013 323 Fig. 2: (a) SMART Domain Prediction; the JmjC domain is present in the hairless protein's query sequence at residue positions 946 to 1157. (b) 3D-PSSM result; alignment of JmjC domain with template sequence of Saccharomyces cerevisiae eIF4E translation initiation factor (PDB ID: 1RF8 Chain B). The crystal structures of the PA2721 protein from Pseudomonas aeruginosa PAO1 (PDB ID: 1U69 Chain A) and the probable antibiotics synthesis protein from Thermus thermophilus HB8 (PDB ID: 1V70 Chain A) were produced as a result of SMART-BLAST of the JmjC domain (Figure 1c). Only 6% of the two chosen templates with the JmjC domain showed similarity in ClustalW-multiple sequence alignment (Figure 3a), making multiple template-based structure prediction impossible. Using antibiotic synthesis protein (1V70) as a template, pairwise alignment of the JmjC domain sequence revealed a 25% similarity. (Figure 3b). The PA2721 protein's alignment with the JmjC domain, however, only revealed 4% similarity. (Figure 3c). These alignment results suggested that choosing the template structure of the antibiotic manufacturing protein for the JmjC domain 3D structure prediction might have biological importance. For modeling the JmjC domain of the hairless protein, the yeast translation initiation factor eIF4E from Saccharomyces cerevisiae (PDB ID: 1RF8 Chain B) was chosen from the 3D-PSSM results with a 22% similarity (Figure 2b). Instead of producing a secondary structure-based template for the full region of the JmjC domain, the chosen eIF4E template had an incorrect alignment with the JmjC domain (Figure 2b).

Finally, we moved on with the crystal structure of a protein from *Thermus thermophilus* HB8 that is likely to synthesize antibiotics in order to anticipate the structure of the JmjC domain.

With the aid of the ClustalW alignment output, the sequence alignment of the template with the JmjC domain was manually performed in the DeepView alignment option. The outcome is depicted in Figure 2. The structure was then aligned and superimposed, and it was sent to the Swiss-model server for modeling and assigning coordinates to the JmjC domain structure. Due to the aligned and superimposed structure that was provided to the Swiss-model by Shrivastava, et al. 324 International Journal of Bioinformatics and Biological Science: v.1 n.3&4 Sept-Dec 2013 Fig. 3: (a) ClustalW-Multiple sequence alignment of the template sequence with the JmjC domain; (b) Pairwise alignment of the JmjC domain with one of the template antibiotics synthesis proteins (PDB ID: 1V70; chain A); and (c) Pairwise alignment of the JmjC domain with another one of the PA2721 protein from *Pseudomonas aeruginosa* PAO1 (PDB ID: 1U69 Chain A). The symbols "*" and ":" denote identical, "." denotes semi-conserved substitution in alignment, and "." denotes conserved substitution. Human Hairless Protein Homology Modeling International Journal of Bioinformatics and Biological Science: v. 1 n. 3&4 Sept.-Dec. 2013 325 Fig. 4: DeepView-Alignment was used to align the sequences of the JmjC domain and the antibiotic production protein (PDB ID: 1V70; chain A). SER160 and LEU153, two amino acid residues, were found in the Ramachandran plot's forbidden zone (Figure 3), as previously mentioned 16.

IV. STRUCTURE PREDICTION

For the creation of the loop in LEU153, the residues ARG151 and GLU155 were chosen as anchor amino acids, whereas GLY158 and TRP162 were chosen for the development of the loop in SER160. After creating loops, the anticipated JmjC domain structure's computed minimized energy was -4680.609 KJ/mol.



Figure 3

All of the amino acid residues are present in the permitted region of Ramachandran plot 16 in the final modelled structure, which was validated by Ramachandran plot (Figure 5b). The WHATIF -Protein Model Check observed results showed that the predicted structure's Ramachandran Z-score was -2.555. This score accurately depicted all residues' backbone conformations in relation to the Ramachandran plot's known-allowed regions. It was anticipated that the RMS Zvalue will fall between -4.0 and 4.0. The RMS Z-score for bond lengths was 0.494, and all bond angles were consistent with accepted bond angles when utilizing a 4 sigma tolerance. The protein structures were well-refined, as indicated by the backbone conformation Z-score of -0.520. Figure 6 depicts the anticipated structure of the JmjC domain of the human hairless protein. Using PyMOL, a free and versatile molecular graphics software package 17, the resulting molecular models' structures were represented. The beta-meander motif, a straightforward super secondary protein shape made up of successive anti-parallel β -strands connected by hairpin loops, is used to represent the JmjC domain structure. This JmjC domain structure uses an eight hairpin loops, one alpha helix, nine beta strands compact fold. We have provided a solid molecular model of the JmjC domain of the human hairless protein in this study, giving the biological functions of this protein a structural underpinning. *Thermus thermophilus* HB8 potential antibiotics synthesis protein crystal structures that were stored in the Protein Data Bank were used to construct a homology model of the JmjC domain. It has an energy minimization of -4680.609 KJ/mol. The most significant finding of our modeling results, however, is that the homology model Shrivastava, et al. 326 International Journal of Bioinformatics and Biological Science: v.1 n.3&4 Sept-Dec 2013 Fig. 3: Two amino acid residues, SER160 and LEU153, are present in the forbidden area in the model provided by the SWISS-MODEL server in (a) Ramachandran plot for the JmjC domain before loop building, and they are absent in (b) Ramachandran plot for the validated JmjC domain following loop formation. International Journal of Bioinformatics and Biological Science: v. 1 n. 3&4 Sept.-Dec. 2013 327 Fig. 2: (a) Modeled representation of the JmjC domain structure; the expected structure takes the shape of a Beta-meander motif, in which two antiparallel strands are connected by a hairpin loop that is frequently made up of one or more glycine or proline residues.

CONCLUSION

This structure is shown as a ribbon model (colored by secondary structure in the JmjC illustration, with the helix colored pink, the sheets colored green, and the loops colored light grey and cyan). (HelixPink; Sheets-Yellow; loops: Light grey and cyan). To build the structure representation, PyMOL was employed. the human JmjC domain as a molecular target in alopecia universalis, is of sufficient quality to use as a structural model for upcoming structure-based inhibitor creation initiatives.

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