Study on the Docking Analysis of Homocitrate Synthase Responsible for Hydrogen Production in *Nostoc Punctiforme ATCC29133*

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Abstract  In the case of Nitrogenase-based Hydrogen production, inactivation of the uptake hydrogenase (Hup) leads to significant increase in hydrogen production activity but it lasts only for few hours under the combined nitrogen atmosphere supplied by active hydrogenase. The catalytic FeMo cofactor of nitrogenase binds homocitrate, which is required for efficient nitrogen fixation. Hence blocking of Homocitrate synthase will be an efficient strategy for the significant hydrogen production. *Nostoc punctiforme* ATCC29133, a nitrogen fixing cyanobacterium that has the genes encoding only for the homocitrate synthase and not for uptake hydrogenase. Since the structure of homocitrate synthase is not yet elucidated, it was taken for modeling using Modeler9v5. The inhibitors for homocitrate synthase were identified from literature and were retrieved from Pubchem. The inhibitors identified were 1,10-phenanthroline, 2,2’-Dipyridyl, Hydroxyllysine, Iodoacetic acid, Oxalylglycine, Pipocelic acid, Thialysine, and oxalate were docked with the homocitrate synthase using Argus lab to compare their activities. However, among these eight inhibitors screened, 1,10-phenanthroline and Oxalylglycine were found to have best docking score and proved efficient in terms of both binding affinity and strong hydrogen bond interactions.

Keywords  Nitrogenase, Hydrogenase, Homocitrate Synthase, *Nostoc Punctiforme ATCC29133*, Modeling, Docking, And Hydrogen Production

1. Introduction

1.1. Role of Cyanobacteria in Fuel Production

Molecular hydrogen, also called the fuel of the future mainly due to its high conversion efficiency, recyclability, and nonpolluting nature, is mostly controlled by either photosynthetic or fermentative organisms and is produced by more environment friendly and less energy-intensive processes as compared to thermo-chemical and electrochemical processes. Cyanobacteria, (blue-green bacteria or blue-green algae) belonging to cyanophyceae, are a large and widespread group of photoautotrophic microorganisms, that fix atmospheric dinitrogen (N₂) into ammonia (NH₃), a form in which the nitrogen is further available for biological reactions[1]. Although quite uniform in nutritional and metabolic respects, cyanobacteria are a morphologically diverse group with unicellular, filamentous, and colonial forms[2]. It possess several enzymes that are involved in hydrogen metabolism: nitrogenase(s) catalyze the production of hydrogen (H₂) concomitantly with the reduction of nitrogen to ammonia; an uptake hydrogenase that catalyzes the consumption of hydrogen produced by the nitrogenase; and a bidirectional hydrogenase, which has the capacity to both uptake and produce hydrogen (Fig.1)[3-4]. In Nitrogenase-based hydrogen production, inactivation of uptake hydrogenase (Hup) leads to significant increase in hydrogen production activity[5]. However, the high-level-activity of the Hup mutants lasts only a few hours under air, a circumstance which seems to be caused by sufficient amount of combined nitrogen supplied by active nitrogenase.

While the uptake hydrogenase is present in all nitrogen fixing strains tested so far, the distribution of bidirectional enzyme is not universal (may be present in both nitrogen-fixing and non-nitrogen-fixing cyanobacteria). The molecular masses indicated for the uptake hydrogenase subunits are mean values calculated from the deduced amino acid sequences of *Anabaena* strain PCC 7120, *Nostoc* strain PCC 73102, and *A. variabilis* ATCC 29413, while the values for the subunits of the bidirectional enzyme are based on data exclusively from *A. variabilis* ATCC 29413 (Tamagnini *et al*., 2002).

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Fi gure 1. Enzymes and genes directly involved in hydrogen metabolism in cyanobacteria

1.2. Bioche mistry of the Hydrogen Producti on

A metallo-cluster called iron-molybdenum cofactor (Fe-Mo cofactor) is believed to provide the substrate binding and reduction site for biological nitrogen fixation. FeMo cofactor is contained within the nitrogenase MoFe protein, and X-ray crystallographic analysis has revealed that it consists of a metal sulfur core[Fe7S9Mo] and one molecule of (R)-homocitrate. The core is constructed from [MoFe3S3] and [Fe4S3] subfragments that are connected by three inorganic sulfide bridges located between pairs of Fe atoms from opposing fragments. The catalytic FeMo cofactor of nitrogenase binds homocitrate, which is required for efficient nitrogen fixation[6-10]. Homocitrate[(R)-2-hydroxy-1,2,4-butane tricarboxylic acid] is coordinated to the Mo atom through its 2-hydroxy and 2-carboxyl groups[11-13]. Although it is not yet known how the substrate interacts with FeMo cofactor during turnover, the presence of six coordinately unsaturated Fe atoms, as well as the attachment of homocitrate to the Mo atom, has invited speculation about the nature of substrate binding[6]. It has also been proposed that the carboxylate group coordinated to the Mo atom might serve as a leaving group in a mechanism that activates Mo to provide a substrate coordination site[14].

1.3. Role of Homocitrate Synthase

Homocitrate is synthesized by the nifV'-encoded enzyme Homocitrate synthase, which catalyses the condensation of acetyl coenzyme A and 2-oxoglutarate to give homocitrate and CoA. The systematic name of Homocitrate synthase (EC 2.3.3.14) is acetyl-CoA:2-oxoglutarate C-acetyltransferase (thioester-hydrolysing, carboxymethyl forming) while the other names in common use include 2-hydroxybutane-1,2,4-tricarboxy late 2-oxoglutarate-lyase, CoA-acetylating, acetyl-coenzyme A:2-ketoglutarate C-acetyl transferase, and homocitrate synthetase[15]. In fungi, such as yeast, and in extremely thermophilic bacteria, homocitrate is used as an intermediate in lysine biosynthesis through the α-aminoacidic acid pathway[16-17].

This enzyme was found to be a Zn-containing metalloenzyme and belongs to the family of transferases, specifically those acetyltransferases that convert acyl groups into alkyl groups on transfer. The R-carboxylate and R-oxo groups of R-ketoglutarate are perhaps required for optimal binding to coordinate to the active site Zn[18]. However, its activity can be inhibited by some of the metal-chelating and sulphydryl binding agents such as 1,10-phenanthroline, Oxalylglycine, 2,2'-Dipyridyl, Oxalate and Iodoacetic acid[19-20]. Moreover, in the in vivo condition, the activity of Homocitrate synthase was inhibited by Lysine and its analogues such as Hydroxylysine and Thialysine, which had a heterotrophic effect on 2-ketoglutaric binding sites. Perhaps, a second class of affectors found, included 2-aminoacidic acid, piperolic acid and dipicolinic acid, which affected the S-acetyl coenzyme A binding sites in Saccharomyces lipolytica[21].

1.4. Criterion for Choosing the Organism

Nostoc punctiforme ATCC29133 (Nostocaceae) is a nitrogen-fixing cyanobacterium that grows autotrophically with CO2 as the carbon source, utilizing an oxygen-producing photosynthetic mechanism for the generation of ATP and reductant. Interestingly, in this organism, heterocyst occupies 3-10% of total cells, which differentiate in response to the lack of nitrogen in the environment and acts as the site of nitrogen fixation. A latest release of Cyanobase (http://genome.kazusa.or.jp/cyanobase/NPUN), a genome database for Cyanobacteria, has revealed Nostoc punctiforme ATCC29133 to have the genes encoding solitarily for Homocitrate synthase and not for Uptake hydrogenase. And moreover, the Kyoto Encyclopedia of Genes and Genomes metabolic pathway database also suggested that lysine biosynthesis in Nostoc sp. to proceed by the diaminopimelic acid pathway, which does not involve homocitrate instead of α-aminoacidic acid pathway, which does involve homocitrate[5]. Hence blocking of Homocitrate synthase would be an efficient strategy for the significant hydrogen production[17].

1.5. Objective

With this perspective, the present study was aimed to work on cyanobacterial enzyme alteration to enhance the production of Hydrogen. Since, comparative modeling is a useful
tool in bioinformatics to predict the three dimensional structure of an unknown protein and also aids to understand protein function, the present study was aimed in this direction to model and predict the structure of Homocitrate synthase using Modeller 9v5 for the sequence (Accession No: B2J701) of Nostoc punctiforme ATCC29133, which was obtained from Uniprot database. Moreover, the interaction between Homocitrate synthase and its potential inhibitors such as 1,10-phenanthroline, 2,2’-Dipyridyl, Hydroxylsine, lodoacetic acid, Oxaoylglycine, Pipocolic acid, Thialysine, and Oxalate was also investigated Insilico by the process of docking.

2. Methodology

2.1. Retrieval of Target Protein Sequence

The protein sequence of homocitrate synthase from Nostoc punctiforme ATCC29133 was obtained from the Uniprot database (Accession No: B2J701) (http://www.uniprot.org/uniprot/B2J701). Since the three dimensional structure could not be ascertained in PDB (http://www.rcsb.org/uniprot/B2J701), an attempt was made to determine the 3D structure.

2.2. Template Identification

The NCBI BLAST was used to identify the template for modeling the three dimensional structure of Homocitrate synthase of Nostoc punctiforme ATCC29133, which revealed no suitable template identity. Therefore multiple templates (3BLE_A, 2CW6_A, 3BG9_B) were identified using 3D-JIGSAW (http://bmmc.cancerresearchuk.org/~3djigsaw/).

2.3. Model Generation and Validation

The three dimensional structure of Homocitrate synthase was predicted using MODELLER9V5 (http://www.salilab.org/modeller/) where the script "align2d.py" was employed to perform an alignment between the target and template sequence and the script "model-default.py" was employed to obtain a rough 3D model which was subjected to energy minimization using the steepest descent technique and Conjugate gradient technique to eliminate bad contacts between protein atoms through Swiss-Pdb Viewer (http://expasy.org/spdbv/). Perhaps, the backbone conformation of the rough model was inspected using the Phi/Psi Ramachandran plot obtained in the PROCHECK server (http://nihserver.mbi.ucla.edu/structures/).

2.4. Active Site Prediction

After obtaining the final model, the possible binding sites of Homocitrate synthase was searched using Q-site Finder (http://www.modelling.leeds.ac.uk/qsitefinder/) which works by binding the hydrophobic (CH3) probes to the protein, and finding clusters of probes with the most favorable binding energy. The higher cavity cluster was considered and the residues around the cluster were identified as the binding residues using Pymol. Thus, ten binding sites were obtained for Homocitrate synthase and since, the amino acid present in the first active site formed the binding pocket forming a hollow sphere arranged accordingly to the binding of Zinc metal was chosen to study interactions at this site with the chosen ligands.

2.5. Protein-Ligand Interaction Studies

Based on the previous studies[18-21] in microorganisms such as Saccharomyces cerevisiae and S. lipolytica, the following eight compounds viz., 1,10-phenanthroline; 2,2’-Dipyridyl; Hydroxylsine; lodoacetic acid; Oxaoylglycine; Pipocolic acid; Thialysine; and Oxalate were found to effectively inhibit Homocitrate synthase, of which 1,10-phenanthroline; 2,2’-Dipyridyl; Oxaoylglycine and Oxalate were metal chelating agents; while lodoacetic acid belongs to sulphydryl-binding agent; Hydroxylsine and Thialysine belongs to lysine analogues; and Pipocolic acid - an affector were chosen as the inhibitory ligands. The 3D structure of the inhibitors along with the Pubchem ID and Molecular weights (http://www.ncbi.nlm.nih.gov/Structure/index.shtml) were determined (Table 1) in order to dock with homocitrate synthase in Argus lab 4.0 (http://www.arguslab.com/downloads.htm), with the grid resolution of 0.40 Å where flexible ligand docking was done when the ligand was described as a torsion tree and the grids were constructed to overlay the binding site. Ligands root node (group of bonded atoms that do not have rotatable bonds) was placed on a search point in the binding site and a set of diverse and energetically favourable rotations were created. For each rotation, torsions in breadth-first order were constructed and those poses that survive the torsion search was scored. The N-lowest energy poses were retained and the final set of poses that underwent coarse minimization, re-clustering, and ranking was selected. The file of the receptor and ligand were uploaded in pdb format and mol format respectively. Docking precision was set to ‘Regular precision’ and ‘flexible ligand docking’ mode was employed for each docking run. The best ligand-receptor structure from the docked structure was chosen based on binding affinity and strong hydrogen bond interactions.

3. Results

3.1. Homology Modeling of Homocitrate Synthase

Homocitrate synthase of Nostoc punctiforme ATCC29133 constituted 377 aminoacids with the molecular weight of 41 kDa. The absence of three dimensional structures in PDB prompted to construct the 3D model, which provided valuable insight into molecular function and also enabled the analyses of its interactions with suitable inhibitors. Among the four conformations generated, the one with the least modeller objective function value was considered to be thermodynamically stable and chosen for further refinement and validation (Fig 2A). The stereochemistry of the constructed model was subjected to energy minimization to assess the quality of the structure. Ramachandran plot for the model showed 84.9% of the residues in the core region, 14.4% residues in the allowed regions and 0.6%, that is two residues in the disallowed region (Fig 3).
| S.NO | INHIBITOR NAME   | STRUCTURE | PUBCHEM ID | MOLECULAR WEIGHT [g/mol] |
|------|------------------|-----------|------------|--------------------------|
| 1    | 1,10-phenanthroline | ![Image](image1) | CID 1318   | 180.20532                |
| 2    | 2,2'-Dipyridyl    | ![Image](image2) | CID 1474   | 156.18392                |
| 3    | Hydroxlysine      | ![Image](image3) | CID 3032849 | 162.18696                |
| 4    | Iodoacetic acid   | ![Image](image4) | CID 5240   | 185.94849                |
| 5    | Oxalylglycine     | ![Image](image5) | SID 3881592 | 147.0862                 |
| 6    | Pipecolic acid    | ![Image](image6) | CID 849    | 129.15704                |
| 7    | Thialysine        | ![Image](image7) | CID 43056  | 211.12098                |
| 8    | Oxalate           | ![Image](image8) | SID 8145248 | 88.019                   |
Lakshmi P. T. V. et al. Study on the Docking Analysis of Homocitrate Synthase Responsible for Hydrogen Production in Nostoc punctiforme ATCC29133

Figure 2. Modeled structure of Homocitrate synthase represented with active site regions. A. Cartoon display of helices is shown in red color, sheets in yellow and coils in green. B. Ten number of active sites were identified and reported, of which the encircled site indicates the highly conserved active site region, which was chosen for ligand interaction.

X-axis shows Phi degrees range between -180 to +180 degrees.
Y-axis shows Psi degrees range between -180 to +180 degrees.
A, B, L are the most favoured regions shown in red color
a, b, l, p are the additional allowed regions shown in yellow color
~a, ~b, ~l, ~p are the generously allowed regions shown in pale yellow color.
White color shows the disallowed region.
Val 147 and Asn 189 are the two aminoacids present in disallowed regions.

Figure 3. Ramachandran plot of Homocitrate synthase built using Modeller9V5

3.2. Active Site Prediction
Among the ten binding sites obtained from the Pocket finder (Fig 2B), the highly conserved site was identified using Q-Site finder. The binding pocket containing 26 residues such as ASN7, ASP8, THR9, ARG12, ASP13, GLU41, LEU42, GLU43, ILE46, LEU70, TRP72, HIS92, ALA94, GLY 134, GLY135, GLU136, ASP137, ARG160, ARG162, CYS164, THR 166, HIS193, HIS195, ASN217, THR218 and ASN229 were conserved and therefore considered for docking analysis.

3.3. Docking of Homocitrate Synthase with Potential Inhibitors
Eight final docked conformations obtained for the different inhibitors were evaluated individually based on the number of hydrogen bonds formed; bond distance between atomic co-ordinates of the active site and inhibitor and the binding affinity (Fig 4). The docking energy showed that the inhibitors possessed the best binding affinity to the protein. Although the active site constituted about 26 residues, the ligands interacted with only 14 residues at THR9, ARG12, ASP13, GLU43, HIS92, GLU136, GLU225, ARG160, ARG162, HIS193, HIS195, ASN217, GLU225 and ASN229 respectively. However, among the chosen inhibitors, the best dock was performed by 1,10-Phenanthroline, which showed the energy of -7.67492Kcal/mol with two hydrogen bond interactions to the residue GLU at position 136 (Fig 4A) and Oxalylglycine which formed 6 hydrogen bonds (Fig 4E) but with slightly higher binding energy(-6.46383 Kcal/mol), than 1,10-Phenanthroline, with good bond distances. Iodoacetic acid and Oxalate may have least inhibition on protein due to the higher binding values when compared to other ligands and perhaps also showed more number of hydrogen bonds such as 5 and 12 respectively with higher bond lengths (Fig 4D and 4H). The hydrogen bond interactions between the eight inhibitors and Homocitrate synthase along with their bond distances and binding affinities are shown in Table 2.
Figure 4. Docking of ligands with homocitrate synthase
Proteins are shown in lines, ligand in stick mode and hydrogen bond as dotted lines. -A. Homocitrate synthase in complex with 1,10-Phenanthroline, B. Homocitrate synthase in complex with 2,2'-dipyridyl, C. Homocitrate synthase in complex with Hydroxyllysine, D. Homocitrate synthase in complex with Iodoacetic acid, E. Homocitrate synthase in complex with Oxalylglycine, F. Homocitrate synthase in complex with Pipecolic acid, G. Homocitrate synthase in complex with Thialysine, H. Homocitrate synthase in complex with Oxalate.

4. Discussions

The present study demonstrates the presence of genes for Homocitrate synthase and absence of genes for uptake hydrogenase in *Nostoc punctiforme* ATCC29133. Hence in this condition, *Nostoc sp.* produces hydrogen in excess than the other species having the genes for uptake hydrogenase. However, in spite of the production of enhanced hydrogen, the combined nitrogen atmosphere created by the active nitrogenase leads to the decrease in the net hydrogen production[5] reported that disruption of homocitrate synthase gene could lead to decrease in nitrogen fixation activity and increase in hydrogen production activity in *Klebsiella pneumoniae*. Homocitrate synthase produces homocitrate that binds to the FeMo cofactor of the nitrogenase, which is required for the efficient nitrogen fixation. Similarly, also in *Nostoc sp.*, Homocitrate synthase functions in the biosynthesis of homocitrate that bind with the FeMo cofactor of nitrogenase but not with lysine synthesis through the alpha-aminoacidic acid pathway as reported by the same co-workers[5]. In this way, homocitrate synthase, a Zn containing metalloenzyme that catalyses the condensation of acetyl CoA and alpha-ketoglutarate to yield homocitrate and CoA[18] if inhibited could probably improve the chances of enhancing the hydrogen production.

Hence, the activity of Homocitrate synthase was inhibited by metal-chelating reagents (1,10-phenanthroline, Oxalylglycine, Oxalate and 2,2'-Dipyridyl) and sulfhydryl binding reagent, Iodoacetic acid. In some of the yeast species, Homocitrate synthase is also believed to involve in biosynthetic pathway of lysine, in which Homocitrate synthase is inhibited by the lysine through feedback mechanism[21]. Since, some of the lysine analogues viz., Hydroxyllysine and Thialysine have the complete inhibitory action on homocitrate synthase and in particular on alpha-ketoglutarate binding sites and that certain effector molecule viz., pipecolic acid also affected the co-operativity of S-acetyl coenzyme A binding sites[21], they were used as small molecules to examine their interaction on the active site of Homocitrate synthase.

On analyzing the results obtained from the docking of metal chelating agents, was revealed to exhibit two hydrogen bonds and only one hydrogen bond respectively by 1,10-phenanthroline and Dipyridyl respectively. But in the case of Oxalate, it formed 12 hydrogen bonds in which some of the bonds showed higher bond length (more than 3Å) that
lead to the weaker interactions. In the case of Sulphydryl binding agent Iodoacetic acid also formed more number of hydrogen bonds (five) like Oxalate, which exhibited the least effect on protein due to its higher docking energy[22]. In the case of Lysine analogues, Hydroxylsine and Thialysine formed four and three hydrogen bonds respectively, but at the same time had a good inhibition for protein. The affector, Pipecolic acid formed only one hydrogen bond with the binding energy of -6.49758 Kcal/mol, which in spite of good binding energy, could not be considered as best because of least inhibitory action on protein due to its highest docking energy and the least number of hydrogen bond formed. However, based on the comparison of the analysis and with relative to binding energies, 1,10-Phenanthroline could be identified as the best potent ligand with very least binding energy of -7.67492 Kcal/mol than other inhibitors. Since Homocitrate synthase is a Zn-containing metalloenzyme, Zn metal is supposed to bind to the amino acids such as cysteine, Histidine, Aspartic acid, Glutamic acid, Asparagine, and Glutamine (given according to priority)[23-24]. Hence according to the least binding energy and the amino acids binding to the zinc metal, 1,10-Phenanthroline was revealed to be an effective interactor of the protein Homocitrate synthase. But according to the number of hydrogen bonds formed and the length of the hydrogen bonds and the amino acids binding to the zinc metal, Oxalylglycine is considered to be the best since it formed six hydrogen bonds of which four were bound to the appropriate aminoacids such as GLU43, HIS92, HIS193 and HIS195 but with slightly higher energy (-6.46383 Kcal/mol) than 1, 10-Phenanthroline with good bond lengths (less than three).

5. Conclusions

Since Homocitrate synthase is a Zn-containing metalloenzyme, it was shown to be effectively inhibited by the metal chelating agents, of which 1,10-Phenanthroline and Oxalylglycine showed the higher effect of inhibition when compared to other inhibitors under investigation. Both 1,10-Phenanthroline and Oxalylglycine showed very good binding energy and interactions with the active site residues compared to the other inhibitors. But basically any insilico analysis is either partial or incomplete if unproved under experimental (in vitro) conditions. Hence, in the direction of understanding the inhibitory mechanism, the work should be taken up to be implemented in wet lab which is our fore-front research (proposal applied to funding agency).

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**Webography**

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