Computational fishing and structural analysis of MIPS protein from two important plant groups

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ABSTRACT. Myo Inositol 1-Phosphate Synthase (MIPS), which catalyzes the first step of inositol metabolism, has been reported from a diverse range of organism like bacteria to human including different groups of plants and animals. The present work is carried out to explore and analyze structural forms of the respective MIPS proteins from complete sequenced genome or proteome available on database of one representative from two important plant groups viz. bryophyte (Physcomitrella patens) and pteridophyte (Selaginella moellendorfii). Previously reported characteristic MIPS sequences was used to identify its homolog ones from those members under study. The explored sequences compared with a number of MIPS varieties from other plant members to study the conserveness or evolution of the protein/enzyme. ProtParam tool provided necessary theoretical physicochemical data of the predicted proteins, the three-dimensional structures were predicted through homology modelling with identified amino acid data. Structural evaluation and stereochemical analyses were performed using ProSA-web displaying Z-scores and Molprobity visualising Ramachandran plot.

1. INTRODUCTION

A rapid and fruitful development of structural and functional genomics and proteomics during the period of twentieth century’s revolution in molecular and computational biology unfolded the chapter of proper storage, retrieval and analysis of genomic proteomic or transcriptomic data for all time. And there are several options to get rid of a particular gene or protein or else like of interest which further can be analysed for different purpose. In very recent time computational determination of structural and functional aspects of a protein/gene derive an equal importance as like wet lab experimental outputs. For example, a number of computational tools are developed and evolved on regular basis to predict the secondary or quaternary structure of a protein or an enzyme and its validation that can be completed within a very short period of time.

Myo inositol 1-phosphate synthase (MIPS) is the key functional enzyme of myo-Inositol biosynthesis pathway. It belongs to NAD-5 super family and act as an NAD$^+$ binding oxidoreductase (Majumder et al 1997) and found to be present throughout a evolutionarily wide range of living organisms (Majumder et al, 2003). The distribution and evolution of MIPS genes from a probable ancestor to cyanobacteria, eubacteria and archaea, and ultimately to higher eukaryotes such as plants and animals, indicate that MIPS started evolving from different sources (Majumder et al., 2003) and that MIPS enzyme activity is distributed in evolutionary diverse phyla, from eubacteria, archaebacteria, cyanobacteria, algae, fungi to higher plants and animals. In plants the enzyme has been reported from many organisms belonging to algae, fungi, gymnosperm, angiosperm etc. Although a few previous work has been carried out from bryophyte (Chhetri et al 2006) and pteridophyte members (Chhetri et al 2005,2006,2008, Jha et al 2013) but still those does not reveal any sequence or structural data from any member of the mentioned groups and that made interested in the current investigation through In Silico method. A large number of computational tools are available for making predictions regarding the identification and structure prediction of proteins. When a whole genome data are reported from a bryophyte (Physcomitrella patens) and
pteridophyte (*Selaginella moellendorffii*), those were used to match and find out the MIPS sequence followed by the bioinformatics methodology known as Protein fishing previously followed by many researchers and in recent Basu et al. (2015).

The enzyme MIPS probably exists in homotetrameric form in native state as evidenced by crystallography (Norman et al., 2002; Stieglitz et al. 2004 etc). It involves in internal oxidoreduction and cyclization & needs NAD+ as a cofactor which is mostly reconverted. MIPS convert glucose 6-phosphate to L-myoinositol 1-phosphate in an NAD+ dependent manner. The NAD+ binding in eukaryotic MIPS involves a Rossman fold characterized by a GXGGXXG motif typical of an oxidoreductase (Kleiger and Eisenberg, 2002).

Crystal structure analysis of MIPS from *Saccharomyces cerevisiae* revealed that each monomer of the homo-tetrameric MIPS has three functionally important structural domains namely the NAD binding Rossman fold, the catalytic binding site and the core domain. Those studies also exemplified induced fit model for binding of the substrate with the catalytic domain of the enzyme. (Stein & Geiger, 2002).

2. METHODOLOGY

2.1 Subject Sequences: Two different MIPS variety sequences, one from the lower plant group green algae (*Chlamydomonas reinhardtii*, Accession No. EDP05930) and another from halophytic monocotyledon (*Porteresia coarctata*, Accession No. AAP74579) were downloaded from NCBI Genpept database (www.ncbi.nlm.nih.gov) and used for sequence homology searching in target plants proteome.

2.2 Protein fishing by local alignment: Those two MIPS amino acid sequences were served as query for BLASTp against whole Proteome of each of our two target plant i.e. *Physcomitrella patens* and *Selaginella moellendorffii* using the California Department of Energy, JGI’s database Phytozome v10 (http://phytozome.jgi.doe.gov/pz/portal.html#!search?show=BLAST). Working comparison matrix was BLOSUM62. The proteins with top hit on BLASTp or with significant similarity were fished out and reported as probable MIPS variety sequences of those target organisms.

2.3 Protein’s physicochemical properties: The theoretical physicochemical properties were computed using the Expasy’s Prot-Param (Gasteiger et al., 2005) (http://us.expasy.org/tools/protparam.html) prediction server (Table 1). The output parameters were theoretical isoelectric point (pI), molecular weight, total num-ber of positive and negative residues, extinction coefficient (Gill and Von Hippel, 1989), instability index (Guruprasad et al., 1990), aliphatic index (Ikai et al., 1980) and grand average hydrophathy (GRAVY) (Kyte and Doolo-ttle, 1982).

2.4 Multiple sequence alignment: To confirm the presence of previously reported conserved regions of the protein a multiple sequence alignment among the diverse variety of MIPS protein from different group of plants was laid down with the help of online server (http://multalin.toulouse.inra.fr/multalin).

2.5 Phylogenetic analysis of sequences: Phylogenetic tree was generated by the UPGMA method using MEGA 6 software (Fig. 4) followed by validation with 100 replicates of Bootstrapping.

2.6 Quaternary structure prediction: The explored sequence’s three dimensional structure prediction was performed using Modeller 9.12. In brief, this method aligns a query sequence to one or more template structures as determined by crystallization/X-ray diffraction, or NMR spectrometry. Suitable templates (PDB id 1LA2) were selected through pdbBLAST at NCBI. E-value and sequence similarity (Table 2) was carefully considered in this respect. As the protein exist in homotetrameric form in its native state so the top hit of homotetrameric structure report was
selected for modelling template. The NAD+ Ligand in the template structure have been transferred to the model because the ligand are annotated as biologically relevant to this particular protein or enzyme. Besides, to establish the model it should be in contact with Ligand and there should have no clash between the ligand and protein when interacting residues with the ligand are conserved between the template-target. Molecule visualisation was done with Chimera 1.10 (Pettersen et al 2004)

2.6 Evaluation of theoretical model: To check the stereochemical stability the psi and phi dihedral angles of each amino acid residues in the protein structures were determined with Ramachandran plots (Ramachandran et al., 1963) analysis from Molprobity (Chen et al 2010) server (http://molprobity.biochem.duke.edu/) . The predicted in silico models were also evaluated using the ProSA web server, ProSA calculates the overall quality score. The output provides a z-score for the model that indicates the overall model quality; this value was determined from the plot during prediction.

2.7 Submission to PMDB database: The models generated were successfully submitted in Protein model database (PMDB) (https://bioinformatics.cineca.it/PMDB/).

3. RESULT AND DISCUSSION:
Both the two way of BLASTp C. Reinhardtii and P. coarctata (Fig. 1) against the Selaginella moellendorffii proteome showed a single hit of resultant protein sequence. It was Gene 270269 which encode 511 residue long protein that found to be 72% identical having 89% positive similarity with myo Inositol 1-Phosphate synthase of C. reinhardtii. On the other hand this was 67.1% identical with 79% positive similarity in terms of P. coarctata belonged variety of mIPS.

Likewise when the same methodology was followed in case of Physcomitrella patens, interestingly we found two genes (Fig. 2) from two different chromosome actually showed almost equal strength of similarity. One is Phpat.004G003200 which have 66.7% identity & 78% positive similarity with both C. reinhardtii and P. Coarctata MIPS. Another is Phpat.003G121000 that have 66.8% identity & 77.3% positive similarity with both C. reinhardtii and P. coarctata.
Now the explored amino acid sequences can be considered for further analysis after they named MIPSsm and MIPSpp respectively. Computational tools provide researchers to understand physico-chemical and structural properties of proteins. The theoretical calculation of different useful parameters (Table 1) like mol. Wt., extinction coefficient, isoelectric point etc reveal an overview on the primary structure and stability of the proteins.

<table>
<thead>
<tr>
<th>Subject Protein</th>
<th>Length</th>
<th>negatively charged residues (Asp + Glu)</th>
<th>positively charged residues (Arg + Lys)</th>
<th>M. wt.</th>
<th>Extinction coefficients</th>
<th>Isoelectric point</th>
<th>Aliphatic index</th>
<th>Instability index:</th>
<th>Grand average of hydrophobicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIPSpp</td>
<td>511</td>
<td>64</td>
<td>55</td>
<td>56776.0</td>
<td>51590</td>
<td>5.41</td>
<td>88.88</td>
<td>32.55</td>
<td>-0.209</td>
</tr>
<tr>
<td>MIPSsm</td>
<td>511</td>
<td>63</td>
<td>55</td>
<td>56767.9</td>
<td>53205</td>
<td>5.73</td>
<td>87.32</td>
<td>38.79</td>
<td>-0.229</td>
</tr>
</tbody>
</table>

MIPSsm have a few more positively charged residues (Arg+Lys) and a greater instability index than the MIPSpp. Isoelectric point (pI) is the pH value at which the molecule carries no charges or the negative and positive charges are equal. Proteins are found to be stable and compact at their pI. The computed pI value (pI < 7) will be useful for developing buffer systems for purification by isoelectric focusing method.

**Conserved domains:** The Multiple Sequence Alignment of protein is a powerful tool for grouping proteins into families and allows subsequent analysis of evolutionary issues (Balasubramanian et al., 2012). The comparison of homologous protein sequences is the most effective means of identifying common active sites or binding domains. The multiple sequences alignment of amino acids with other group of plant members MIPS variety sequence (Fig. 3) obtained from database revealed some distinctly conserved domains that supports the earlier observation highlighted by Majumder et al, 2003 which are essential for functional property of the protein. One of them is the most important Rossmann fold sequence GXGGXXG type that functions as a NAD+ binding domain. The domain lies in between 75 to 85 no. residue in the MSA image (Fig. 3).
Fig. 3: Multiple sequence alignment of MIPS proteins among different plant, yeast and the subject sequences from Selaginella & Physcomitrella under study

**Phylogenetic analysis:** Various sequence information of gene or protein offers the possibility to investigate the gene function in vivo with parallel phylogenetic analyses of multigene families to address their evolution within and across species (Vincentz *et al.*, 2003). Within families, the protein structure and catalytic residues that determine the substrate specificity are generally conserved.

After confirmation of the explored proteins existence in NAD-5 super family with the help of CD search tool at NCBI, phylogenetic analysis of MIPS proteins among selected organisms (Fig. 4) also provided powerful information about ancestry of the protein. According to result the cyanobacteria Oscillatoria and Yeast forms an out-group and the members under current study are found to be in the same branch which suggests a common ancestor that conforms to the common plant taxonomical lineage.

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**Fig. 4:** Evolutionary relationships among taxa in respect to MIPS
The evolutionary history of the MIPS protein among different plants and yeast and cyanobacteria was inferred using the UPGMA method [Sneath and Sokal 1973]. The optimal tree with the sum of branch length = 7.32697491 is shown. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (100 replicates) are shown next to the branches [Felsenstein, 1985]. The evolutionary distances were computed using the Poisson correction method [Zuckerkandl and Pauling, 1965] and are in the units of the number of amino acid substitutions per site. The analysis involved 9 amino acid sequences. All positions containing gaps and missing data were eliminated. There were a total of 364 positions in the final dataset. [Data adopted from: MEGA6 software]

**Homology modelling:** Protein 3D structure is important in understanding protein interactions, function and their localization [Jones 2001]. Quaternary structure plays an important role in defining protein function by facilitating allostery and cooperativity in the regulation of ligand binding (Matthew et al., 2001). Homology modeling is the most common structure prediction method. In this study, the quaternary structures of *Physcomitrella* and *Selaginella* MIPS proteins were predicted by considering the biological assembly that provided the functional state of the protein. The NAD+ ligands also incorporated in time of model generation (Fig. 5) to fix the functionality of the protein with its structure. NAD+ characteristically bound to the GWGGNNNG domain which exists in an alpha-helix of each monomer.

<table>
<thead>
<tr>
<th>Model Protein</th>
<th>PMDB id</th>
<th>Template</th>
<th>Sequence similarity with template</th>
<th>Sequence identity with template</th>
<th>E value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIPSpp</td>
<td>PM0080022</td>
<td>1LA2</td>
<td>99</td>
<td>51</td>
<td>1e-168</td>
</tr>
<tr>
<td>MIPSsm</td>
<td>PM0080023</td>
<td>1LA2</td>
<td>99</td>
<td>53</td>
<td>3e-174</td>
</tr>
</tbody>
</table>

Table 2: Summarized values after PDB-BLAST for template selection

![Fig. 5](image-url): The quaternary structures of MIPSsm and MIPSpp generated by Modeller and visualized by Chimera (For comparison both two models are attached in a single background)
The Z-score value, a measure of model quality that predicts the total energy of the structure (Wiederstein and Sippl, 2007), was determined for the MIPSsm & MIPSpp with the help of PROSA server. Z-score also measures the deviation of total energy of the structure with respect to an energy distribution derived from random conformations. The calculated scores i.e. both -7.66 for MIPSpp and -7.4 for MIPSsm are well within the range of scores (Fig. 6) typically found for proteins of similar size.

The residue score plot (Fig. 7) shows local model quality by plotting energies as a function of amino acid sequence position $i$. In general, positive values correspond to problematic or erroneous parts of the input structure. A plot of single residue energies usually contains large fluctuations and is of limited value for model evaluation. Hence the plot is smoothed by calculating the average energy over each 40-residue fragment $s(i,i+39)$, which is then assigned to the 'central' residue of the fragment at position $i+19$ (thick line). A second line with a smaller window size of 10 residues is shown in the background of the plot (thin line) [adopted from ProSA web server]. The plots indicate that in both the cases the modelled structures are found to have good stability and overall quality.
Table 3: Stereo chemical properties of modelled proteins

<table>
<thead>
<tr>
<th>Model Protein</th>
<th>PMDB id</th>
<th>residue in favoured region</th>
<th>Outlier</th>
<th>Z-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIPSpp</td>
<td>PM0080022</td>
<td>93% (1888/2031)</td>
<td>35/2031</td>
<td>-7.6</td>
</tr>
<tr>
<td>MIPSsm</td>
<td>PM0080023</td>
<td>92.4% (1878/2032)</td>
<td>46/2032</td>
<td>-7.44</td>
</tr>
</tbody>
</table>

Fig 8: Screenshot of Molprobity Ramachandran Plot (Left- MIPSpp, Right- MIPSsm)

Per Residue geometry and the overall geometry of the protein structures was analyzed with Ramachandran plots (Ramachandran et al., 1963). In a polypeptide, the main chain N-Cα and Cα-Cb bonds are relatively free to rotate. These rotations are represented by the torsion angles φ and ψ respectively. Fig. 8 shows the quantitative evaluation of Physcomitrella and Selaginella MIPS structures generated using Ramachandran plots provided by the Molprobity web server. Both Physcomitrella and Selaginella MIPS had almost equal number of residues in favoured (>92%) regions. The prominence of residues in the right handed α-helix and β-sheet regions suggested that the structure was rigid with more right helices. Physcomitrella MIPS showed higher quality compared to the Selaginella MIPS protein in respect to having fewer residues in the outlier region and more residues in the allowed region (Table 3). Selaginella MIPS had 97.74% of residues in the allowed region. More residues were observed in the allowed region of Physcomitrella (98.28%). There were more outliers in the Selaginella MIPS (2.26%) than in Physcomitrella (1.72 %). These values were almost similar with the the template X ray crystallography structure present in Protein Data Bank. After validation 3D models were successfully submitted to PMDB with id PM0080022 and PM0080023.

4. CONCLUSION

When the enzyme myo-inositol 1-Phosphate synthase is crucially important in diversification evolution and comparison study, the void of sequence data from the previously mentioned important group can at least be mortally fulfilled by the referred sequence of current study. Furthermore it is well known that due to low resolution and high R factors of many crystal structures, homology modelling is now regarded as an important technique for not only in obtaining the 3D-structure of a putative protein but also for refining the existing low accuracy experimental
structures. The above analysis through homology models may provide an insight to simulations and related computational studies such as drug docking calculations and protein-drug or enzyme-target interactions to obtain useful insights into structure–function relationships. The partial phylogenetic analysis including the reported data and its positive inference finally adds a strong support in current method of study.

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Reference

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