Modelling and Functional Characterization of the Pyridoxamine Kinase/Phosphomethylpyrimidine Kinase Domain Family

Marco Uderzo

Department of Mathematics, University of Padua marco.uderzo@studenti.unipd.it ID: 2096998

Tanner Graves

Department of Mathematics, University of Padua tanneraaron.graves@studenti.unipd.it ID: 2073559

Claudio Palmeri

Department of Mathematics, University of Padua claudio.palmeri@studenti.unipd.it ID: 2062671

Abstract

This project aims to build a sequence model and provide a comprehensive functional characterization of the Pyridoxamine Kinase/Phosphomethylpyrimidine Kinase domain family. The models' accuracy is benchmarked against Pfam annotations in the SwissProt database. Furthermore, the project delves into the functional and structural properties of the domain family, involving taxonomic lineage analysis, Gene Ontology (GO) annotations assessment, and motif searching. (Include findings in the abstract)

1 Introduction

1.1 Protein Domains

In molecular biology, a protein domain represents a conserved part of a protein's sequence and three-dimensional structure, capable of evolving, functioning, and existing independently from the rest of the protein chain. These domains, each forming a stable and compact three-dimensional structure, are essential components in proteins, often occurring in various combinations across different proteins. Domains are fundamental in molecular evolution, serving as versatile building blocks that can be rearranged to form proteins with diverse functions. This adaptability and independence make them crucial in understanding protein structure and function.

1.2 Pyridoxamine Kinase (CHECK FOR ERRORS)

Pyridoxamine Kinase/Phosphomethylpyrimidine Kinase family is a group of enzymes that play key roles in various biochemical pathways, particularly in the metabolism of vitamins and coenzymes. This family includes two distinct but related enzymes:

- Pyridoxamine Kinase: This enzyme is involved in the vitamin B6 metabolism pathway. Vitamin B6 exists in different forms, including pyridoxamine, pyridoxal, and pyridoxine. Pyridoxamine kinase specifically catalyzes the phosphorylation of pyridoxamine, converting it into pyridoxamine 5'-phosphate. This is an important step in the salvage pathway of vitamin B6, which is crucial for its recycling and maintenance within the cell.
- Phosphomethylpyrimidine Kinase: This enzyme is a part of the thiamine (vitamin B1) biosynthetic pathway. It catalyzes the phosphorylation of hydroxymethylpyrimidine (HMP) to hydroxymethylpyrimidine phosphate. This step is essential in the synthesis of thiamine pyrophosphate (TPP), an active form of vitamin B1. TPP is a vital coenzyme in several enzymatic reactions, particularly those involved in carbohydrate metabolism.

Both these enzymes, due to their roles in vitamin metabolism, are crucial for maintaining cellular health and function. Disruptions in these pathways can lead to vitamin deficiencies, affecting numerous biological processes.

1.3 Objective of the Study

This project aims to build a sequence model and provide a comprehensive functional characterization of the Pyridoxamine Kinase/Phosphomethylpyrimidine Kinase domain family. (Write a small preamble of the goals of this project, even if it is similar to what is written in the abstract).

2 Methods and Results

2.1 Model Building

Firstly, we investigated the target family to model, having Pfam domain PF08543, and verified that the provided representative A0A0J9X285 protein sequence is indeed characteristic of the protein family. This was done by retrieving the seed alignment used to generate the HMM defining the Pfam family from *InterPro*, and aligning the representative query sequence to the seed alignment using *JalView*.

The query spans the length of the seed alignment and the gaps opened in the query correspond to low occupancy regions in the seed alignment. This bolsters our confidence that performing a homology search with our query sequence will be able to return sequences belonging to the PF08543 family. This was done by performing a Position-Specific Iterated BLAST (PSI-BLAST) search on *SwissProt*. The results were downloaded as a .fasta file and opened in *JalView*, where we added our query sequence as a reference. This FASTA was then submitted to *NCBI* (hosted clustal from JalView to return our MSA. ???)

The query sequence overlapped the primary conserved regions of the MSA, and the majority of positions outside of the query had very low occupancy, consisting of sequences that were unusually long. The query bounds for the MSA are observed to be reasonable bounds to trim the MSA, so poistions outside this range were trimmed from it.

Sequences that opened gaps more than a couple residues long were investigated by referencing the BLAST hit corresponding to that sequence. Many of these instances were from Eukariotes - which is atypical for this family - and were of reasonable quality. Since it is useful to include this information, no sequences reported by BLAST were discarded.

The MSA was finalized by removing the query sequence, and it was then processed to generate a Position-Specific Scoring Matrix (PSSM) using the command line PSI-BLAST tool, with the SwissProt database as the reference. Finally, the HMM was build using the hmmer hmmbuild command.

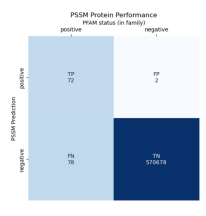
2.2 Model Evaluation

Using hmmsearch, and starting from the PSI-BLAST hits, we generated the predictions, extracting the list of alignments from the HMM search results, in order to understand the sequence alignments that relate to the Pyridoxamine Kinase/Phosphomethylpyrimidine Kinase domain family. After defining our ground truth by finding all proteins in *SwissProt* with the PF08543 domain, we proceeded to evaluating the PSSM model.

2.2.1 PSSM Protein-Level Performance Evaluation

The protein-level performances of the PSSM model are shown in the table below:

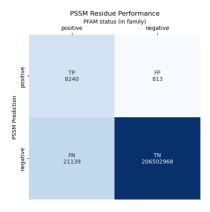
Metric	Value
Precision	0.894
Recall	0.227
F1-Score	0.361
Balanced Accuracy	0.613
MCC	0.45



2.2.2 PSSM Residue-Level Performance Evaluation

The residue-level performances of the PSSM model are shown in the table below:

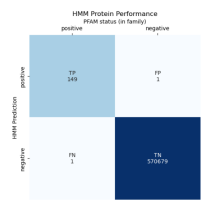
Metric	Value
Precision	0.91
Recall	0.28
F1-Score	0.429
Balanced Accuracy	0.64
MCC	0.505



2.2.3 HMM Protein-Level Performance Evaluation

The protein-level performances of the HMM model are shown in the table below:

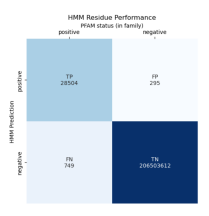
Value
0.993
0.993
0.993
0.996
0.993



2.2.4 HMM Residue-Level Performance Evaluation

The residue-level performances of the HMM model are shown in the table below:

Metric	Value
Precision	0.989
Recall	0.974
F1-Score	0.982
Balanced Accuracy	0.987
MCC	0.982

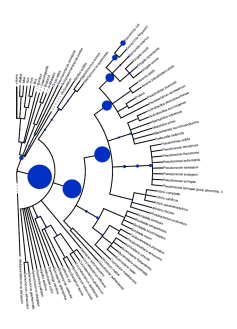


It is worth noting that the HMM model's performance is robust with respect to the starting alignment. (but are our models comparable with the corresponding Pfam?).

2.3 Taxonomy

To construct the taxonomic tree, we assembled the lineage data derived from the SwissProt database, corresponding to the protein family under investigation. The lineages were used to generate a comprehensive taxonomic hierarchy, which was enriched with node-specific information, including taxonomic names and the frequency of each taxon's occurrence within our data.

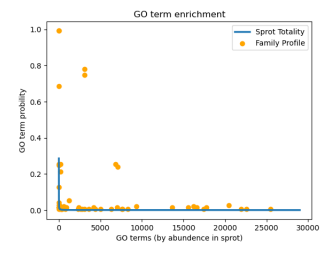
The taxonomic tree is shown below. In order to view it in full resolution, we refer to the corresponding <code>TaxonomyTree.pdf</code> file that can be found in the supplementary material.



2.4 Functional Enrichment with Gene Ontology Annotation

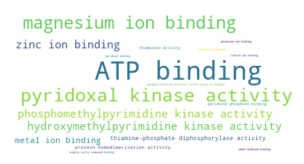
We performed Functional Enrichment Analysis using *Gene Ontology* (GO) annotations by extracting the *molecular function*, *cellular component*, and *biological process* data.

In order to visualize which GO Terms are characteristic of our family, we can plot the enrichment (probability) of observing a GO Term over both our model family and the totality of *SwissProt*. Selecting the terms with the highest odds, or the ratio of probability that the term is observed in the family and all of *SwissProt* gives us clues about which terms are most characteristic.

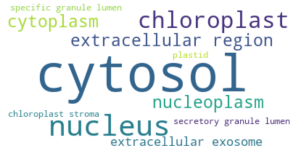


By using Fisher's Exact Test, we realized that the p-value is extremely close to zero (maybe add p=...) for terms with high odds, indicating that they are indeed characteristic of our family. However, as a consequence of how sparse GO Labels are for a sequence, when compared (to the abundance of our limited amout of terms (56) present in our family. ?? -; check this).

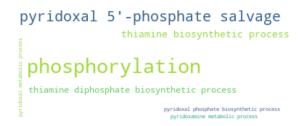
Below, we plot a word cloud of the Enriched Terms for each aspect:



Molecular Function Terms



Cellular Component Terms



Biological Process Terms

Most enriched branches:

- 2.5 Motifs
- 3 Results
- 4 Discussion
- 5 References

List and number all bibliographical references in 9-point Times, single-spaced, at the end of your paper. When referenced in the text, enclose the citation number in square brackets, for example. Where appropriate, include the name(s) of editors of referenced books.