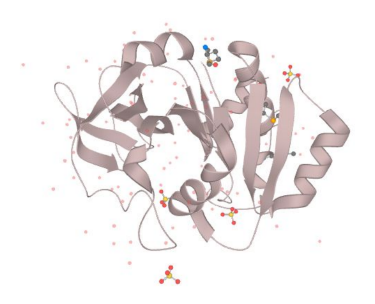
# **Characterizing Nucleotidyl Transferase**

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## Introduction

The aim of this project is to characterize a single domain, particularly Nucleotidyl Transferase from the organism Yersinia pseudotuberculosis - a bacterial species that most commonly causes foodborne illness. We have been assigned a domain sequence from which we build a sequence model to provide structural and functional characterization of the domain family. All the results represented in this report can be reconstructed by using this code available on <u>Github</u>.

#### **Domain Model Definition**

To find a good model that represents our assigned domain, we developed an automated process for building HMM and PSSM models based on an input multiple sequence alignment. Prior to running this automated procedure we took two steps. Firstly, using the InterPro API we gathered all reviewed proteins belonging to SwissProt along with the domain position within them and defined them as our ground truth. Then, we performed a BLAST search against UniProt and its clusters to retrieve homologous proteins in order to perform a multiple sequence alignment to detect conserved regions. We used the MSA to build our models upon it. We chose to work with UniRef50 afterwards because we obtained a lot of significant hits ( $e \le 10^{-20}$ ) and the UniRef cluster allowed for easier programmatic access.

### **Building and evaluating the models**

Once we obtained the homologous proteins, we wanted to have a broader choice for our models so we developed an easy way of building them using Python. Prior to building them, we performed MSA using T-Coffee, Muscle and ClustalOmega. The alignments produced were not very noisy but using Jalview we removed redundant sequences to gather more alignments. Having obtained sixteen alignments, we built an HMM and a PSSM model for each and we evaluated them with respect to our ground truth ([Table 1], [Table 2] in the Appendix). We realized that overall the alignments built by T-Coffee performed best, while varying redundancy thresholds do not produce better results.

The final model we chose to work with is an HMM model built upon an MSA produced by T-Coffee. We selected it since it produced the best results metrics-wise and overall HMMs are a generalization of the profile concept and take the positions of indels into account when modeling the relationship of the sequences as a "family", to represent the most probable reality.

## **Domain Family Characterization**

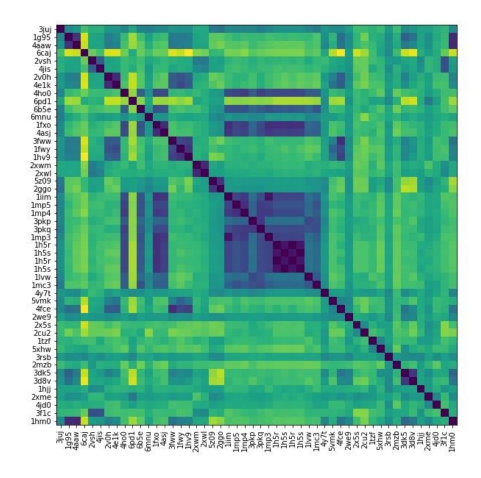
Having defined a model, we examine the structural and functional characteristics of the protein family. By identifying two datasets, one used for the structural and the other for the functional characterization, we were able to gain some insights regarding the family. The datasets constructed are:

- 1. **family\_structures**, obtained by using the HMM created previously to search sequence databases for homologous sequences on HMMER. Only the PDB structures with a minimum overlap of 80% were kept to build the dataset.
- 2. **family\_sequences**, which comprises all UniRef90 sequences matching the HMM. This was retrieved in .xml format.

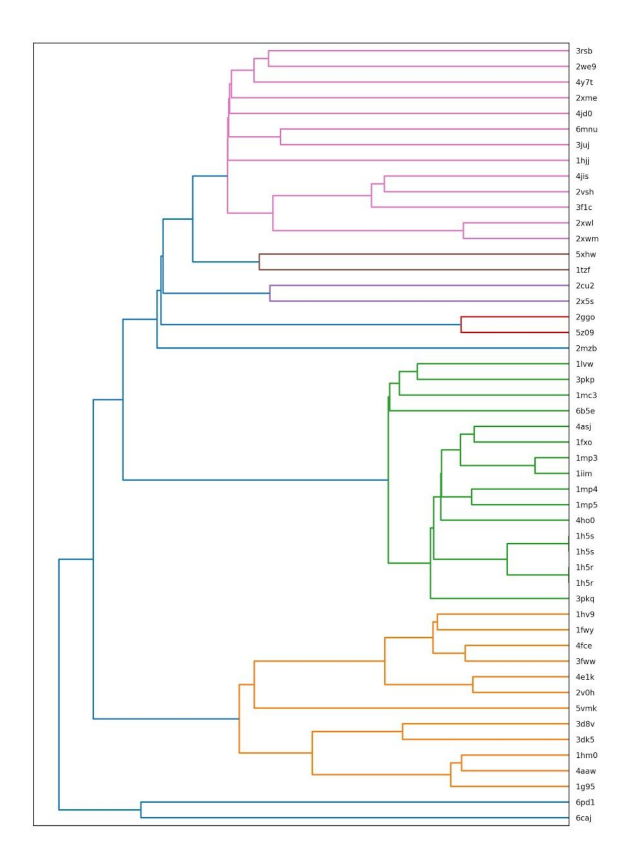
#### Structural Characterization

Homology modeling is based on the observation that related protein sequences adopt similar three-dimensional structure. It is the structure that determines the molecular function of the protein.

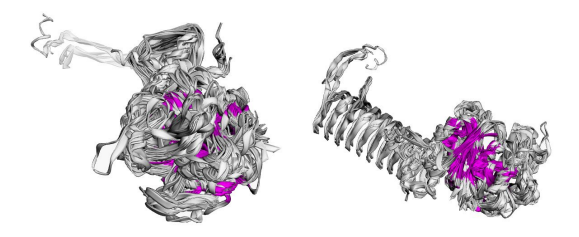
The first task after acquiring the family structures was to perform a pairwise structural alignment using TM-align, which uses a simple approach with gapless threading and secondary structure similarity. To interpret the results, we used RMSD to tell us how close two structures are. This value equals 0 for identical structures, and is very high if the two are significantly different. The results are plotted in a heatmap, purple indicating closer structures and yellow the opposite. We can clearly observe a cluster in the middle of the plot, indicating the similarity.



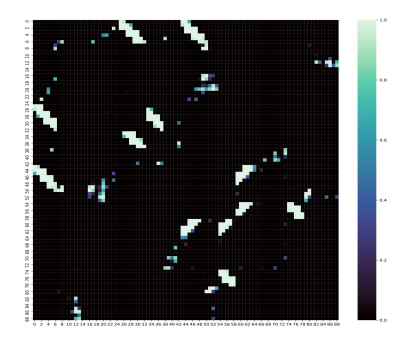
We also constructed a dendrogram to represent the hierarchical clustering of the RMSD values obtained. Analyzing it, we assume that the far down sequences are outliers that have a very high structural difference. By doing a box plot analysis, we confirmed that one of these should be removed from further investigation (6caj).



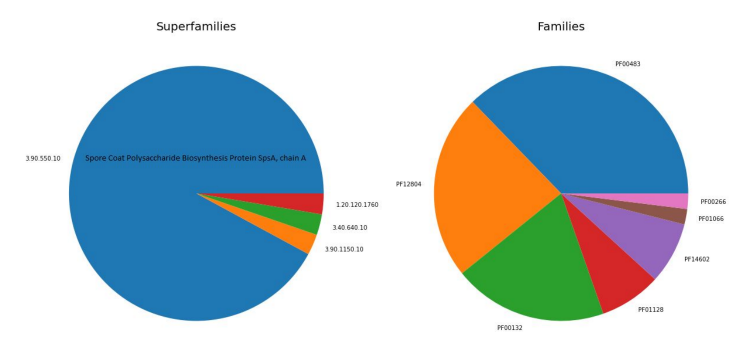
To identify the most conserved positions, we carried out a multiple structure alignment making use of mTM-align server and the PDB structures excluding outliers. We worked with the common core alignment (highlighted in magenta colour in the following figure) to spot long range conserved contacts.



Essentially, we built a distance matrix for every domain structure by setting a threshold of 12 sequence positions for the separation. Likewise, we produced a contact map examining only the areas separated no more than 8Å.



Finally, by using the PDB IDs from the family structures in the "Retrieve/ID mapping" service from UniProt and cross-referencing Gene3D and Pfam, we derived the CATH superfamilies as well as the Pfam families concerning the proteins we worked with. As a result, four superfamilies and seven families were identified. The most dominant superfamily is 3.90.550.10 and each of the structures was identified with it, whereas there are three families with a significant influence: PF00483 - which is the family of the input sequence assigned (Y. pseudotuberculosis); PF12804; and PF00132.



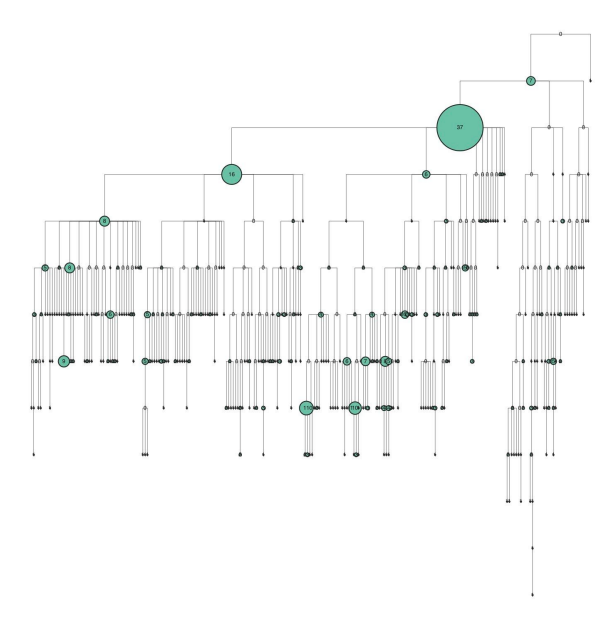
## Taxonomy

In this section, we address the evolutionary relationships among sequences. We make use of the family\_sequences database, comprising 664 clusters containing a total of 1179 protein sequences.

Each cluster is characterized by a taxonomic ID (i.e. the cluster identification), a representative member ID (that is the ID of the representative member of the cluster) and a taxonomic ID for each protein contained in the cluster.

In order to find a representative tree branch, we decided to consider the cluster ID and download the NCBI taxonomy database using the <a href="ete3">ete3</a> library. Once we found the lineage of all the proteins, we counted the abundance of each ID in the lineage and set the dimension of the respective node accordingly.

Finally, we were able to find the taxonomy tree of our family sequences. In particular, we chose to exploit the <u>toytree</u> library. There are 242 leaves and all the other nodes are internal. The most common nodes can be seen in [Table 3] in the <u>Appendix</u>. The taxonomy tree is shown in the following figure:



#### **Functional Characterization**

In order to provide a functional characterization of our domain, we finally analyzed the GO annotations of each retrieved sequence. For clarification, <u>Gene Ontology</u> is a structured, controlled vocabulary for the classification of gene function at the molecular and cellular level, divided in three separate sub-ontologies or GO types: biological process, molecular function and cellular component. The needed information was already contained in the family sequences dataset, so we simply extracted the annotations (entity/dbReference) as explained for the taxonomy lineage. For the same purposes, we needed the annotations from the SwissProt dataset.

It is important at this point to recall that, according to the true path rule, a gene annotated to a term is also implicitly annotated to each ancestor of that term in the GO graph, so the following statistics will be performed, for each sequence, on the set of all the annotations relative to the sequence itself and to its ancestors, in order to obtain more reliable results.

The next step was to perform a functional enrichment analysis: it consists of applying statistical tests to verify if genes of interest are more often associated with certain biological functions than what would be expected in a random set of genes. We chose Fisher's exact test as a reference, given that, especially in the case of strongly unbalanced datasets like ours, it usually outperforms the common alternatives such as fold increase, computed aside and clearly greater than 1, even if not in agreement with the F-test about the order of the first most significant annotations. The results are shown below, in an absolute scale and grouped by sub-ontology.

Note that, given both the huge dimension of the entire SwissProt with respect to the number our matching sequences and the good performances of our model, the resulting right-tailed p-values are extremely small, indicating that our model performs very well and that the found terms are strongly likely to be correct.

de	children	fold_inc	two_p	r_p	1_p	
						GOi
transferase activity, transferring phosphorus	597.0	14.005525	0.000004	0.000000e+00	1.0	GO:0016772
carbohydrate metabolic process	539.0	14.843562	0.000003	0.000000e+00	1.0	GO:0005975
anatomical structure morphogenesis	704.0	37.104949	0.000002	0.000000e+00	1.0	GO:0009653
transferase activity, transferring acyl groups	412.0	27.466950	0.000002	1.362233e-316	1.0	30:0016747
regulation of anatomical structure morphogenesis	393.0	26.136851	0.000002	5,894371e-309	1.0	30:0022603
transferase activity, transferring acyl groups	458.0	23.247296	0.000001	8.738476e-294	1.0	30:0016746
regulation of developmental process	1713.0	15.653365	0.000002	5.754133e-240	1.0	30:0050793
cellular lipid metabolic process	557.0	12.966483	0.000002	4.762986e-232	1.0	90:0044255
lipid metabolic process	672.0	11.679714	0.000003	1.883860e-217	1.0	30:0006629
carbohydrate derivative biosynthetic process	319.0	8.951599	0.000003	1.589272e-199	1.0	30:1901137
nucleobase-containing small molecule metabolic	642.0	8.558009	0.000003	4.438470e-194	1.0	30:0055086
regulation of biological quality	1506.0	10.133582	0.000002	7.156973e-184	1.0	30:0065008
developmental process	3212.0	9.637514	0.000003	1.637212e-178	1.0	30:0032502
transferase activity	2478.0	5.134042	0.000006	2.402559e-175	1.0	30:0016740
nucleobase-containing compound biosynthetic pr	457.0	7.784914	0.000004	1.882027e-171	1.0	0:0034654

The word cloud in the figure below contains the 15 most enriched GO terms descriptions, independently from their sub-ontology, which is nevertheless represented by their score-proportionally vanishing colours: in red the children of the molecular\_function node and in blue the ones under the biological process class.

regulation of cell shape
anatomical structure morphogenesis
external encapsulating structure organization
amino sugar biosynthetic process glycosaminoglycan metabolic process

UDP-N-acetylglucosamine diphosphorylase activity
peptidoglycan metabolic process
cellular carbohydrate biosynthetic process
polysaccharide metabolic process lipooligosaccharide metabolic process
lipid A biosynthetic process glycolipid biosynthetic process

It is easy to note that no green term shows up, which would be relative to the cellular\_component branch, for which the lower p-value is lower with respect to the other terms, but still significant in the first cases. Moreover, it is interesting that only one molecular\_function term shows up, revealing that with high probability (p-value < 1e<sup>308</sup>) our domain can be annotated with depth of 6 *UDP-N-acetylglucosamine diphosphorylase activity* term, GO:0003977, and so with all its ancestors.

Note that, with the p-value threshold alone, it may be hard to tell which are the most interesting hits, so a condition about the number of children of each term was added, in order to prefer, among the top-ranked results, the terms which are higher up in the main branches, as an assurance of the stability of the results. In this case the resulting word cloud is slightly more general and shows higher level terms of the two previously mentioned sub-ontologies, and in general most of them have as children the nodes in the previous word cloud, confirming we are on the right path.

```
transferase activity, transferring acyl groups other than amino-acyl groups cellular lipid metabolic process

anatomical structure morphogenesis

lipid metabolic process

regulation of developmental process
developmental process
transferase activity, transferring acyl groups
nucleobase-containing small molecule metabolic process
transferase activity, transferring phosphorus-containing groups
regulation of anatomical structure morphogenesis

carbohydrate derivative biosynthetic process
transferase activity
carbohydrate metabolic process
regulation of biological quality
```

Finally, the most significant enriched branches are shown below, coherently with the previous results.

depth	def	Sub-Ontology	
			GOi
3	regulation of cell shape	biological_process	GO:0008360
2	anatomical structure morphogenesis	biological_process	GO:0009653
3	cell morphogenesis	biological_process	GO:0000902
3	cell wall organization	biological_process	GO:0071555
3	transferase activity, transferring phosphorus	cellular_component	GO:0016772
3	cellular carbohydrate metabolic process	biological_process	GO:0044262
3	carbohydrate metabolic process	biological_process	GO:0005975
2	cell wall organization or biogenesis	biological_process	GO:0071554
3	transferase activity, transferring acyl groups	cellular_component	GO:0016746
3	regulation of developmental process	biological_process	GO:0050793
3	cellular lipid metabolic process	biological_process	GO:0044255
3	lipid metabolic process	biological_process	GO:0006629
3	nucleobase-containing small molecule metabolic	biological_process	GO:0055086
2	regulation of biological quality	biological_process	GO:0065008

## Summary

To summarize, using an automated process and by exploiting some of the available Multiple Sequence Alignment tools like T-Coffee, ClustalOmega and Muscle we developed a model that represents our domain precisely considering the important metrics. We then constructed the family structures and sequences databases to be able to provide additional insights into the structural and functional characteristics of our domain. After performing pairwise and multiple sequence structural alignments, we obtained the CATH superfamily 3.90.550.10 and several families of which the most dominant was our starting sequence's family - PF00483. For the functional part, we found out that it only covers two of the three sub-ontologies namely the biological processes (biosynthetic and metabolic processes) and molecular function (transferase activity).

## Appendix

 $Model\ Names: /models/model\_type/UR50\_Alignment-Software\_Redundancy-Threhsold$ 

## 1. Sequence Metrics

accuracy	precision	sensitivity	specificity	MMC	f1_score	model_name
0.998423	0.391760	0.965577	0.998456	0.614588	0.557377	/models/hmm/UR50_CO_R96
0.998338	0.380667	0.982788	0.998354	0.611158	0.548775	/models/pssm/UR50_CO_R96
0.995792	0.192281	0.969019	0.995820	0.430801	0.320889	/models/hmm/UR50_CO_R98
0.998320	0.377333	0.974182	0.998345	0.605810	0.543969	/models/pssm/UR50_CO_R98
0.996165	0.207290	0.969019	0.996193	0.447377	0.341523	/models/hmm/msa_modified_clustalomega
0.998341	0.381333	0.984509	0.998356	0.612227	0.549736	/models/pssm/msa_modified_clustalomega
0.998412	0.390278	0.967298	0.998444	0.613966	0.556160	/models/hmm/UR50_MU_R92
0.998341	0.381333	0.984509	0.998356	0.612227	0.549736	/models/pssm/UR50_MU_R92
0.998483	0.399563	0.944923	0.998538	0.614042	0.561637	/models/hmm/UR50_MU_R95
0.998310	0.375333	0.969019	0.998340	0.602601	0.541086	/models/pssm/UR50_MU_R95
0.998904	0.484007	0.989673	0.998913	0.691732	0.650085	/models/hmm/UR50_TC_R96
0.998324	0.378000	0.975904	0.998347	0.606879	0.544930	/models/pssm/UR50_TC_R96
0.998912	0.486005	0.986231	0.998926	0.691958	0.651136	/models/hmm/UR50_TC
0.998310	0.375333	0.969019	0.998340	0.602601	0.541086	/models/pssm/UR50_TC
0.998736	0.447012	0.965577	0.998770	0.656594	0.611111	/models/hmm/UR50_MU_R93
0.998352	0.383333	0.989673	0.998361	0.615436	0.552619	/models/pssm/UR50_MU_R93

## 2. Position Metrics

model_name	f1_score	ммс	specificity	sensitivity	precision
/models/hmm/UR50_CO_R96	0.700152	0.703797	0.999623	0.780827	0.634587
/models/pssm/UR50_CO_R96	0.821488	0.834378	0.999703	0.996851	0.698593
/models/hmm/UR50_CO_R98	0.659466	0.661030	0.999586	0.710022	0.615631
/models/pssm/UR50_CO_R98	0.821773	0.834724	0.999706	0.997639	0.698619
/models/hmm/msa_modified_clustalomega	0.566281	0.567959	0.999593	0.525077	0.614502
/models/pssm/msa_modified_clustalomega	0.825365	0.837667	0.999711	0.996303	0.704494
/models/hmm/UR50_MU_R92	0.486412	0.499033	0.999669	0.397448	0.626689
/models/pssm/UR50_MU_R92	0.828359	0.840395	0.999721	0.997348	0.708340
/models/hmm/UR50_MU_R95	0.477184	0.487562	0.999645	0.395862	0.600556
/models/pssm/UR50_MU_R95	0.965022	0.965420	0.999956	0.994358	0.937367
/models/hmm/UR50_TC_R96	0.748381	0.757001	0.999627	0.882171	0.649828
/models/pssm/UR50_TC_R96	0.839262	0.849587	0.999742	0.994664	0.725858
/models/hmm/UR50_TC	0.838641	0.839130	0.999837	0.871012	0.808590
/models/pssm/UR50_TC	0.988695	0.988705	0.999989	0.994448	0.983009
/models/hmm/UR50_MU_R93	0.484768	0.496073	0.999650	0.399782	0.615642
/models/pssm/UR50_MU_R93	0.962610	0.963053	0.999952	0.993567	0.933524

## 3. Nodes' Abundance

Organism	Abundance
Bacteria	37
Proteobacteria	16
Clostridium	11
Streptococcus	10
Shewanella	9
Gammaproteobacteria	8
Mycobacteriaceae	8
Pasteurellaceae	8
Bacillaceae	7
cellularorganisms	7
Streptomyces	6
Lactobacillaceae	6
Terrabacteriagroup	6
Vibrionaceae	6