Structural descriptor database: a new tool for sequence-based functional site prediction
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Abstract

Background: The Structural Descriptor Database (SDDB) is a web-based tool that predicts the function of proteins and functional site positions based on the structural properties of related protein families. Structural alignments and functional residues of a known protein set (defined as the training set) are used to build special Hidden Markov Models (HMM) called HMM descriptors. SDDB uses previously calculated and stored HMM descriptors for predicting active sites, binding residues, and protein function. The database integrates biologically relevant data filtered from several databases such as PDB, PDBSUM, CSA and SCOP. It accepts queries in fasta format and predicts functional residue positions, protein-ligand interactions, and protein function, based on the SCOP database.

Results: To assess the SDDB performance, we used different data sets. The Trypsion-like Serine protease data set assessed how well SDDB predicts functional sites when curated data is available. The SCOP family data set was used to analyze SDDB performance by using training data extracted from PDBSUM (binding sites) and from CSA (active sites). The ATP-binding experiment was used to compare our approach with the most current method. For all evaluations, significant improvements were obtained with SDDB.

Conclusion: SDDB performed better when trusty training data was available. SDDB worked better in predicting active sites rather than binding sites because the former are more conserved than the latter. Nevertheless, by using our prediction method we obtained results with precision above 70%.

Background

In the post-genomic era, functional identification of proteins and their interactions became an essential step for the understanding of the molecular machinery of the cell. Advances in structural genomics and proteomics [1-3] are producing a huge amount of information. Structural information is useful for inferring the protein biochemical function because often this function is strongly correlated with the protein 3D structure [4-6]. Moreover, structural properties can be useful for detecting a set of special amino acids that is strongly related to function. These amino acids are the protein functional residues, e.g. active site and binding site residues. Functional residues are the key players in protein-protein interactions and in protein-ligand interactions. Some recent studies have sought to find the common maximal set of functional res-

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idues for related proteins [7-14]. Those studies aimed at finding ligand functional sites may have a major impact on drug discovery development [15].

Protein-ligand interactions can be solved by assessing the co-crystallized ligand structure. Nowadays, there are several applications for this purpose. One such application is LigBase [16], a 3D ligand-binding database derived from the Protein Data Bank (PDB) [17]. In LigBase, a ligand-binding site is an amino acid with at least one atom within 5 Å from a ligand atom. There are many other applications based on the same principle, including PDBSUM [18], BindingDB [12], and PLD [13]. On the other hand, protein-protein interactions are hard to solve because large protein complexes are often more difficult to crystallize than protein-ligand complexes [19-22]. However, when functional residues of a protein family are known, they can be used to predict the function of related proteins. Identifying these residues from amino acid sequences alone would be useful for the understanding of the interactions of proteins with no 3D structure defined.

In order to predict functional sites in proteins with unknown function, the PROSITE database [23] uses a method based on regular expressions [24] (ScanProSite [25]) to detect functional site patterns and profiles within proteins with unknown function. In addition, Henschel and colleagues [22] presented a method based on hidden Markov models (HMM) [26-28] to predict functional sites in both protein-ligand and protein-protein interactions. Their research compiled sequential segments that constituted structural features of an interaction site into a hidden Markov model, namely the HMM descriptor. In their study, each Multiple Sequence Alignment (MSA) that represents a protein family generates fragments according to the position of functional residues, and for each fragment an HMM is built. Segmented HMMs are merged into one final HMM by their linking with insert states. This final HMM is called the HMM descriptor. Collection of HMM descriptors can be used to screen sequence databases by predicting functional site positions. Henschel and colleagues [22] focused on the protein-protein functional site predictions, but they also evaluated ligand-binding descriptors for the ATP-binding.

Herein, we present a method based on hidden Markov models (HMM) that uses functional sites of known ligand-protein interactions to predict functional sites in query sequences. Our method was based on the study done by Henschel et al [22], with some differences. First, we built the HMM from full MSAs rather than from functional sites of MSAs segments. Second, HMMs for the prediction of functional residues were trained with both multiple sequence alignments based on structural alignments and with column alignments which corresponded to functional residue positions. Third, our approach focused on predictions of protein-ligand functional sites rather than predictions of protein-protein functional sites. The ligand-binding residues are likely to be important in many applications such as in designing small molecule inhibitors for drug discovery. Finally, we developed a web-accessible database, called Structural Descriptor Database (SDDDB) that uses our prediction method and allow the detection of functional sites, biological activity, and of ligand interactions in query sequences.

The SDDDB was built with information from several databases, namely the PDB [17], SCOP [29], PDB-SUM [18], and CSA [30]. Each family in the SCOP database was associated with a library of HMM descriptors. We selected 1,902 SCOP families and 70,215 PDB entries. Functional sites were obtained from PDBSUM (binding-ligand residues) and from CSA (active-ligand residues). Structural alignments were derived from the 3DCOFFEE aligner [31]. We used the HMMER package [32] for building HMM descriptors. We also built a user friendly website to turn both the database and the functional site prediction method available to anyone interested at http://www.sddb.lncc.br. In this website, query sequences can be scored by using a library of HMM descriptors. The SCOP classification, functional residue positions, and ligand binding interactions are considered in order to indicate the sequence most likely function. Also, SDDDB contains links to all related databases, which allows the user to have immediate access to additional information on the protein of interest.

Results and discussion

In order to validate our prediction method, we performed experiments using three different data sets. We started by assessing one specific family, namely the Trypsion-like Serine proteases. Next, we assessed the remaining SCOP families. Finally, we compared our results with those on the ATP-binding study done by Henschel and colleagues [22]. In the first experiment, the functional residue positions were defined from protease study [33]. In the second and third experiments, the functional residue positions were extracted from the PDBSUM and the CSA databases. In the first and second experiments we wanted to assess our prediction method when both data was available, curated data supplied by expert knowledge and automated data calculated from computational methods, as done previously by others [34-36].

Trypsion-like Serine proteases data set

In our first experiment, we only considered SCOP sequences from the Trypsion-like Serine protease family. We selected a total of 39 proteins. Sequences were separated into 8 groups, according to ligand specificity. These groups are Chymotrypsin, Elastase, Granzyme, Thrombin,
Urokinase, Trypsin, Plasminogen Activator, and a hybrid group formed by Procarboxypeptidase A-S6 subunit III, Granzyme K, and Neuropsin. Table 1 shows for each group the PDB identify, chain, start residue and end residue that refer to the PDB coordinate file for each protein selected. Also, table 1 shows alignment identity for each group.

We performed cross validation by applying the leaving one out method [37], in each group. First, training set sequences were aligned using the 3DCOFFEE tool [31]. Next, for each alignment, positions of binding site alignment and of active site alignment were defined, according to [33]. HMM descriptors were built using both alignments and alignment-columns, which corresponded to

| Table 1: Groups of Trypsion-like Serine protease. |
|-----------------------------------------------|
| Group name | PDB-Ids | Chain | Start Residue | End Residue | Alignment identity |
|------------|---------|-------|---------------|-------------|--------------------|
| Chymotrypsin | 1gmc    | A     | 1             | 245         | 52%                |
|            | 1cho    | E     | 1             | 245         |                    |
|            | 1azz    | A     | 16            | 246         |                    |
|            | 1eq9    | A     | 16            | 244         |                    |
| Kallikrein | 1ao5    | A     | 16            | 246         | 32%                |
|            | 1op8    | A     | 16            | 246         |                    |
|            | 1aut    | C     | 16            | 243         |                    |
|            | 1bgv    | A     | 16            | 245G        |                    |
|            | 1lo6    | A     | 16            | 243         |                    |
|            | 1eax    | A     | 16            | 244         |                    |
|            | 1m9u    | A     | 16            | 242         |                    |
| Granzyme   | 1euf    | A     | 16            | 243         | 62%                |
|            | 1iau    | A     | 16            | 244         |                    |
|            | 1fi8    | A     | 16            | 244         |                    |
| Thrombin   | 1bth    | H     | 16            | 242         | 47%                |
|            | 1id5    | H     | 16            | 244         |                    |
|            | 1etr    | H     | 16            | 247         |                    |
|            | 1fis    | A     | 16            | 244         |                    |
|            | 1h8d    | H     | 16            | 246         |                    |
|            | 1fxy    | A     | 16            | 246         |                    |
|            | 1gvk    | B     | 16            | 245         |                    |
|            | 1fpk    | C     | 16            | 245         |                    |
|            | 1rfn    | A     | 16            | 245         |                    |
| Urokinase  | 1fiw    | A     | 16            | 254         | 43%                |
|            | 1fiz    | A     | 16            | 257         |                    |
|            | 1fv9    | A     | 1             | 244         |                    |
| Trypsin    | 1d6r    | A     | 16            | 245         | 73%                |
|            | 1mct    | A     | 16            | 245         |                    |
|            | 1slu    | B     | 16            | 245         |                    |
|            | 3tgi    | E     | 16            | 245         |                    |
|            | 1h4w    | A     | 16            | 246         |                    |
|            | 1h8b    | A     | 16            | 245         |                    |
|            | 1trn    | A     | 16            | 246         |                    |
| Plasminogen activator | 1a5i | A | 1A | 244 | 46% |
|            | 1bda    | A     | 1A            | 244         |                    |
|            | 1gvz    | A     | 16            | 246         |                    |
| Hybrid group | 1fon    | A     | 9             | 240         | 29%                |
|            | 1mza    | A     | 14            | 248         |                    |
|            | 1npm    | A     | 16            | 243         |                    |

Group name, PDB coordinates details, such as PDB-ld, chain, start residue, and end residue of each protein used in first experiment. In addition, the later column shows the identity of alignment of each protein group.
the residues of the functional sites. These HMM descriptors were used for screening and predicting functional residue positions in the test-sequence set.

Table 2 reports precision and recall values for each group, for e-values > 1. The Trypsion-like Serine protease experiment achieved an average of 99.61% of precision and 99.62% of recall, when predicting the active site. Regarding the binding site predictions, this experiment achieved an average of 98% of precision and 97% of recall. These results show that active site residues are more conserved than binding site residues. Therefore, HMM descriptors performed better when active sites were assessed. Also, we assessed protein function prediction by classifying test-sequence according to the SCOP database. As a result, 100% of the proteins were correctly classified into the SCOP class, fold, super family, and family. Additionally, 97.1% was classified into its correct domain.

**SCOP data set**

The second experiment was performed considering the SCOP families selected for this study (see Method section for election criterion), with the exception of the Trypsion-like Serine protease family. A total of 1,901 families were selected. Similarly as in the first experiment, we performed cross validation by applying the leave one out test [37]. Sequences in each family were separated into groups according to their ligand interactions. Training sequences were aligned using 3DCOFFEE [31]. Binding and active site alignment positions were obtained by running an algorithm that recognizes functional ligand residues in alignments, as explained in Methods. Next, HMM descriptors were built using both alignment and functional ligand positions. Finally, each HMM descriptor was used for screening, predicting, and classifying a test-sequence.

The SCOP families experiment achieved precision of 84% and recall of 70.8%, in predicting the active site. For binding site prediction, precision and recall were 78% and 57%, respectively. As observed in the first experiment, our methodology obtained better results in predicting active sites rather than in predicting binding sites. Indeed, active site residues are more conserved than binding residues. Therefore, we applied different levels of tolerance when defining an alignment column into a binding or active site. For instance, we set as binding site columns those that presented more than 50% of their binding residues aligned, whereas for active sites columns we required 80% of active residues aligned. These parameter settings narrowed the active site predictions because they decreased the number of false positives that could be incorporated in the training data, which in turn increases precision. The recall also increases because fewer cases of false negatives are incorporated in the active site column.

We assessed the classification with respect to the SCOP database, and found that all proteins were correctly classified into the SCOP class, fold, super family, and family. Around 96% of the proteins were classified into their correct domain.

**ATP-binding data set**

We compared our results with those obtained by Hen schel and colleagues [22]. For that, we assessed HMM descriptors performance for the particular study of the ATP-binding site. We considered 94 SCOP families in which a ATP-ligand or ADP-ligand was present, totaling 10,520 sequences. Notice that their tests were performed on SWISS-PROT [38] database, and they used 36,774 sequences, while we used the SCOP database, a dataset smaller than Swiss-Prot. Unfortunately, direct compari-

| Group name          | AS Precision | AS Recall | BS Precision | BS Recall |
|---------------------|--------------|-----------|--------------|-----------|
| Chymotrypsin        | 97           | 98.8      | 97.8         | 96.8      |
| Kallikrein          | 100          | 100       | 99           | 98.6      |
| Granzyme            | 100          | 99.2      | 100          | 98.9      |
| Thrombin            | 99.9         | 100       | 98           | 97.2      |
| Urokinase           | 100          | 100       | 97.9         | 97.1      |
| Trypsin             | 100          | 100       | 100          | 100       |
| Plasminogen activator| 100         | 99.8      | 98.5         | 97.3      |
| hybrid group        | 100          | 99.2      | 96.1         | 96.3      |

Precison and recall for each group in Trypsion-like Serine protease data set

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sons are not possible because their source code is not available. For this experiment we assessed precision-recall for e-values < 1. Table 3 shows our results and those of Henschel and colleagues’. Their original results can be found in table 1[22]. They achieved a precision of 89.43%, whereas we achieved 86.39%. However, our results had a recall rate of 55.72%, whereas theirs was 25.17%. This indicates that our prediction method has produced fewer false negatives. While our method detected more false positives, it also detected more true positives.

Assessing HMM Descriptors performance
To evaluate the performance of HMM descriptors, we built precision-recall curves for our first and second experiments. Figure 1-a shows precision-recall curves for active site predictions, whereas figure 1-b reports precision-recall curves for binding site predictions. Table 4 shows protein function classification for both experiments. We noted that the first experiment performed better than the second one in all cases. We compared the experiment performance for active site prediction only, and we observed that the area under the graphic is greater for the Trypsion-like Serine protease experiment than for the SCOP family experiment (figure 1-a). Also, figures 1-b shows greater areas for the Trypsion-like Serine protease experiment. Results in the first experiment were expected because the training data set is a curated data, as they were collected from the literature. We believe they are more reliable than computationally-calculated functional residues. This significant difference in the performance of these experiments may be due to separation into groups according to ligand specificity. Proteins from the same family can present different functional sites when interacting with the same ligand, i.e., the type of amino acids involved in interaction might not be the same ones. Thus, these proteins present different ligand specificity. Naturally, the alignment of proteins with similar ligand specificity keeps the functional sites aligned. As a result, HMM descriptors derived from these alignments perform better than those built by using alignments that contain all the proteins that interact with the same ligand, which explains the better results obtained in the first experiment. In any way, HMM Descriptor performed better achieved precision above 70% and recall above 50%, in both cases.

Table 3: Matches for ATP-binding data set.

|          | Precision | Recall |
|----------|-----------|--------|
| Henschel et al | 89.43  | 25.17  |
| SDDB     | 86.39    | 55.72  |

Precision and recall for both Henschel and colleagues prediction method, and our approach (SDDB).

Conclusion
With this study we aimed to develop a web-based tool that allowed the use of a computational method for the prediction of functional sites and protein function. For that, we created a prediction method based on the study of Henschel and colleagues [22] in which they developed a method for predicting functional sites in proteins with unknown function. Our web-based tool does two main tasks: it screens the SCOP database for the prediction of the protein function, and it points out the positions of its functional residues. Likewise the study of Henschel and colleagues, our prediction method is based on the hidden Markov model descriptors and structural properties. However, our HMMs were built using information from structural alignments, ligand binding sites, and active sites.

Our web-accessible database is a user friendly interface that allows users to analyze their own data. After setting the parameters, users can submit their protein sequences. Each protein is then analyzed and results are sent by email. Each sequence submitted is scanned against a library of HMM descriptors and results are displayed in two levels. In the first level of results, it shows all predicted functional residues and protein interactions with its ligands. In the second, it shows the predicted functional residues for each interacting ligand, individually. HMMs and alignments are available for download in the result page.

To validate our prediction method, we performed experiments that achieved significant precision-recall ranges. Our approach is accurate in determining protein-ligand interactions and interaction-residues positions. Therefore, SDDB is useful for functional annotation and for predicting functional residues. Knowing the positions of functional residues may provide insights into the biological activity of the protein and reveal new targets for drug development.

Methods
The web-based tool was developed in Java version 1.5 and is now served through a Tomcat apache web server running on a Linux operating system. The web-site was built with servlets, JSP, and java scripts technologies. To store our data, we built a relational database manager using Postgres, version 8.2. The SDDB was populated with data extracted from public databases such as SCOP, version 1.71, PDB (downloaded in May, 2007), PDBSUM (downloaded in May, 2007) and CSA version 2.2.5. PDB is a repository for 3D structural data of proteins widely used in structural proteomic analysis. SCOP is a manually inspected database of protein folds that is particularly interesting to our study because it describes structural and evolutionary relationships between proteins, including all entries in PDB. PDBSUM is a database that provides, for each PDB file, a summary of the key structural features,
Comparison of Trypsion-like Serine protease and SCOP descriptors through precision-recall curves

**Figure 1**

Comparison of Trypsion-like Serine protease and SCOP descriptors through precision-recall curves. SDDB performance for all data sets, as measured by precision-recall curves. Each point in the graphic corresponds to a different e-value cutoff. A – Active site prediction. B – Binding site prediction.
including protein-ligand interaction and binding sites positions. Finally, CSA catalogs catalytic residues in enzymes of 3D structures. Based on the family level of SCOP, we extracted and classified the majority of protein-ligand interactions found in the PDB. HMMs were built from structural alignment derived from 3DCOFFEE, using HMMER package. This stage is the so-called training phase (see below). The Inference phase is the step in which query sequences are submitted to SDDB tool.

**Training Phase**

In this phase data was fed into the SDDB to be later used for inference. Initially, we used other databases to obtain the necessary data such as SCOP families, PDB structure files, ligands, and functional ligand residues. Next, structural alignments were built from the SCOP protein group using 3DCOFFEE. Finally, HMMER was performed to build HMM descriptors from both structural alignments and from functional ligand residues.

**Extracting data from external databases**

Initially, we selected all proteins in each SCOP family, except for those proteins without either ligand interactions or shared ligand interactions. Also, we removed the proteins whose size $T$ was not in the range $[M - M/2, M + M/2]$, where $M$ is the average size of the sequences in its respective SCOP family. This step is necessary because some of the SCOP family sequences greatly vary in size, which results in alignments with many gaps. As a result, HMMs built from a gapped alignment do not recognize homologous proteins well. With this initial data processing, we obtained 1,902 families with 70,215 sequences.

Next, selected sequences were separated in groups according to the SCOP family level. For each sequence, we extracted from PDB its three-dimensional structure required for the structural alignment. From PDBSUM, we extracted the set of ligands that interact with each sequence and for each ligand we also extracted its protein binding sites. Figure 2 shows a LIGPLOT [39] scheme of the binding ligand residues (downloaded from PDBSUM) of the interaction between E. coli's DNA ligase (PDB id 1a0i) [40] and the ATP (Adenosine-5'-triphosphate) ligand. The active ligand residues of the interaction between SCOP protein and PDB ligand were extracted from the CSA database.

**Building Structural Groups**

Let $f$ be an arbitrary SCOP family. All proteins of $f$ make up the core structural group of this family, namely $SG_f$ (structural group of $f$). The alignment of $SG_f$ proteins was called $A_f$, and it was used as the starting point for the construction of the HMM descriptor that represents the $f$ called by HMM$_f$. A family $f$ is made up of a set of proteins $P = \{p_1,...,p_Q\}$, where $Q$ is the number of sequences within an $f$. Each protein $p_i \in P$ interacts with a set of ligands $L = \{L_1,...,L_n\}$ downloaded from the PDBSUM. Each family $f$ was divided into groups of proteins that interact with the same ligand. Then, a group was created only by proteins of $f$ that interact with ligand $L_i$. This group was called $SG_fL_i$ (structural group formed by a subset of $f$ that interacts with $L_i$). Each $SG_fL_i$ was defined as the side structural group. The alignment of $SG_fL_i$ proteins was called $A_fL_i$. This alignment was the starting point for the construction of the HMM descriptors, namely HMM$_fL_i$. An HMM descriptor was built using both the MSA ($A_f$ or $A_fL_i$) and the alignment-columns, which corresponded to the residues of the functional sites (see below). Figure 3-a shows how each SCOP family was segmented in structural groups and how HMM descriptors were created from these groups.

**Algorithm for recognizing functional ligand residues in alignments**

The building of HMM descriptors required both the entire MSA and the columns referring to the amino acid alignment of the functional sites. Each protein $p$ within the alignment has a set of amino acids that form, in regard to its interaction with a specific ligand, its functional sites, i.e., the ligand determines the set of functional sites of that protein. Because each protein $p$ in $A_f$ has a set of functional sites associated to the $L_i$ interaction, we directly applied the algorithm for the identification of the functional sites. On the other hand, the $A_f$ alignment requires knowing the cumulative sites for all involved proteins, before applying the algorithm for the identification of functional sites. The cumulative sites of a protein $p$ were calculated by combining sets of functional sites of the protein $p$ with each of its ligands. Thus, if a protein $p$ has interactions with ligands $L = \{L_1,...,L_j\}$, a set called $pL_k$ includes the positions of the functional sites that are associated to the interaction between $p$ and $L_k$, where $k \leq j$. As a result, the cumulative sites of a protein $p$ that has the set of ligands $L$ is created by $pL = pL_1 \cup ... \cup pL_j$. 

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**Table 4: SCOP classification.**

| Class | Fold | Super Family | Family | Domain |
|-------|------|--------------|--------|--------|
| Trypsion-like Serine protease | 100  | 100          | 100    | 100    | 97.1 |
| SCOP | 100  | 100          | 100    | 100    | 96.2 |

SCOP classification for Trypsion-like Serine protease and SCOP data sets.
After identifying the cumulative functional sites, we applied the algorithm described below. Each column in an MSA is individually assessed. Whenever 50% of the sequences contained a binding residue in that column, the column was marked as a binding site of the alignment. If 80% of the sequences contained an active residue at that column, then this column was marked as an active site of the alignment. Therefore, for each alignment \( A_f \), there is an array \( B_f \) with all elected columns of \( A_f \), that represent functional site positions in \( A_f \). Therefore, \( B_f \) contained functional sites of proteins of \( f \) and \( B_fLi \) contained functional sites of a subset of proteins of \( f \) that interacts with the \( L_i \) ligand.

**HMM construction**

The HMM descriptors were built using an HMMER package. Given a \( SG_f \) of a arbitrary SCOP family denoted by \( f \), we built a HMM from \( A_f \) and \( B_f \). We also built an HMM for each alignment in \( SG_fL \), where \( L \) is the set of ligands that interact with proteins of the \( f \). Next, we built an HMM for each protein in \( A_f \) and \( A_fLi \), as follows. Each alignment con-
contains a $P = \{p_1, \ldots, p_n\}$ protein subset, where each $p_i$ is a protein of the $f$, since $i \leq n \leq Q$. For each $p_i$, an HMM was built from its alignment with the consensus-sequence of the HMM that contains $p_i$. The consensus-sequence is obtained from the alignment used to build the HMM, considering the most frequent amino acids in each column. These HMMs, called HMM classifications, were built to aid the SCOP classification of proteins of unknown function. Figure 3-b shows how to build HMM classifications for a hypothetical family $f_i$. First, the consensus-sequence (called consensus-sequence $f_i$) is extracted from HMM $f_i$. Next, each protein $p_i$ in $SG_{f_i}$, where $SG_{f_i}$ is the structural group of $f_i$ (see Building Structural Groups section), is aligned with consensus-sequence $f_i$ through hmmalign program of HMMER package. This alignment

**Figure 3**

**Creating HMM descriptors from SCOP families.** Each SCOP family was segmented in structural groups and HMM descriptors were created from these groups. A – Image shows the building of HMM descriptors for a hypothetical family, namely $f_i$. First, $A_{f_i}$ is built by aligning all proteins of $SG_{f_i}$ structural group. Next, HMM $f_i$ is built from both $A_{f_i}$ alignment and functional site positions of $A_{f_i}$, called $B_{f_i}$. Finally, $f_i$ is divided into groups of proteins that interact with the same ligand, and HMM $f_i L_i$ are built in the same way, since $L$ is the number of ligand and $1 \leq i \leq N$. B – In order, for building HMM classifications, the consensus-sequence $f_i$ and each protein in $SG_{f_i}$ are aligned by producing $A_{\alpha} P_{\alpha}$ where $1 \leq i \leq Q$. The building of HMM $f_i p_i$ classifier is based on $A_{\alpha} P_{\alpha}$.
is called \(A_p \rho p\), where \(\rho p \in SG_{f_i} = \{p_1, ..., p_Q\}\) and \(Q\) is the number of sequences in \(f_i\). Each \(A_p \rho p\) results in the building of \(\text{HMM}_f \rho p_i\).

**Inference Phase**

After establishing and characterizing the training phase, the web-tool can then be used for predicting functional sites and protein function of user query sequences. Users can submit sequences in fasta format to the web site. Each sequence \(p\) is submitted to the \(\text{HMM}_f \) where \(f\) represents the families selected by the user. Let \(\theta_{f_i}\) be e-value assigned to scoring of \(p\) by \(\text{HMM}_f \). If \(\theta_{f_i} < \theta_{f_i} \rho p\) is accepted, i.e., it is likely that \(p\) is a member of the \(f\). The \(\theta_j\) is the threshold necessary in order for \(p\) to belong to \(f\). When \(p\) is accepted, the cumulative functional site positions and the \(\rho\) protein function are predicted, followed by \(p\) scoring by all \(\text{HMM}_f L\), where \(L_i \in L\) and \(L\) is the set of all ligands that interact with proteins of \(f\). Similarly, if \(\theta_{f_i} L_{1, i} < \theta_{\rho \pi_i}\), then \(p\) is accepted, where \(\theta_{\rho \pi_i}\) is the e-value assigned to scoring of \(p\) by \(\text{HMM}_f \rho \pi_i\) and \(\theta_{\rho \pi_i}\) is the threshold for considering \(p\) interacting with \(L_i\). When \(p\) is accepted, the functional sites of interaction between \(p\) and \(L_i\) are predicted.

**Predicting binding and active ligand residues**

The alignment-columns containing functional sites were mapped to the HMM states, i.e., either match or insert states, as shown in figure 4. The figure shows a partial alignment of proteins for the globin family, in which the column labeled \(B_3\) represents a binding site position in the alignment mapped to the match state \(M_2\). Similarly, the column labeled \(A_3\), which represents an active site position in the alignment, mapped to the match state \(M_3\). Note that the column labeled \(B_3\) mapped an insert state \(I_{1, 3}\). Hence all columns in the alignment were represented by one HMM state. When a protein \(p\) is scored by an HMM, the Viterbi algorithm [41] provides both \(Pr[p|\text{HMM}_f\] and \(\pi\), where \(\pi\) is the best path of \(p\) through HMM states. If \(\pi\) is given by \(M_1 M_2 I_3 ... M_n\), then it means that \(a_1\) was recognized by the match state \(M_1\), \(a_2\) was recognized by the \(M_2\), \(a_3\) was recognized by the insert state \(I_3\), until \(a_n\) was recognized by \(M_n\). Therefore, if \(M_2\) is a state that represents a binding site, it is likely that \(a_2\) in \(p\) is a binding site residue.

**Annotating query sequences**

After submitting a \(p\) protein to an HMM descriptor and verifying that its e-value is smaller than \(\theta_{f_i}\) it is necessary to determine the following parameters for the SCOP classification: class, fold, super family, family, and domain. We annotated \(p\) by comparing it with all HMM classifications (see section HMM construction) and by associating to \(p\) the same classification of \(\rho p\), where \(p\) is the protein most similar to \(p\), and which belongs to the HMM classification. For instance, suppose that some protein \(p\) has been scored by \(\text{HMM}_f\) as in figure 3-b. Also, consider that \(p\) has been classified with e-value \(\theta_{f_i} p\), where \(\theta_{f_i} p\leq \theta_{f_i}\). Next, \(p\) is scored by each \(\text{HMM}_f p_i\), where \(p_i \in \{p_1, ..., p_Q\}\) and \(Q\) is the number of sequences in \(f_i\). Each \(\text{HMM}_f p_i\) assigns a \(\theta_{f_i} p_i\) e-value to \(p\). Suppose \(\theta_{f_i} p_i<\theta_{f_i} p_i<...<\theta_{f_i} p_i\), thus \(p\) receives the SCOP classification of \(p_i\).

**Authors’ contributions**

JSB performed the studies and participated in writing the manuscript. JHF participated in the design of the study and ATRV participated in the design and coordination of the study, and in writing the manuscript. All authors read and approved the final manuscript.

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**Figure 4**

Mapping functional residues to HMM states. Each column in the globin alignment maps to either match or insert state, including the columns that represent functional sites. The columns labeled by \( A_s \) represents active site positions, whereas \( B_s \) columns represents binding site positions. \( M, I, \) and \( D \) represent match, insert and delete states in HMM architecture, respectively. In this illustration, \( A_{s1} \) mapped to \( M_8 \) state, and the \( B_{s1}, B_{s2} \) and \( B_{s3} \) columns mapped to \( M_2, M_{13} \) and \( I_{15} \) states, respectively.
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