cswHMM: A NOVEL CONTEXT SWITCHING HIDDEN MARKOV MODEL FOR BIOLOGICAL SEQUENCE ANALYSIS

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Keywords: Bioinformatics, Data-mining, Hidden markov models.

Abstract: In this work we created a sequence model that goes beyond simple linear patterns to model a specific type of higher-order relationship possible in biological sequences. Particularly, we seek models that can account for partially overlaid and interleaved patterns in biological sequences. Our proposed context-switching model (cswHMM) is designed as a variable-order hidden Markov model (HMM) with a specific structure that allows switching control between two or more sub-models. An important feature of our model is the ability of its sub-models to store their last active state, so when each sub-model resumes control it can continue uninterrupted. This is a fundamental variation on the closely related jumping HMMs. A combination of as few as two simple linear HMMs can describe sequences with complicated mixed dependencies. Tests of this approach suggest that a combination of HMMs for protein sequence analysis, such as pattern mining based HMMs or profile HMMs, with the context-switching approach can improve the descriptive ability and performance of the models.

1 INTRODUCTION

Biological sequences, especially protein sequences, code for 3-D objects, where even a sequentially distant position can fold into mutual proximity and vice versa, consecutive amino acids can have contrasting functions. As such, protein sequences contain interleaved gapped patterns of varying size, as well as long-distance interactions and complex dependencies that often need to be accounted for. That makes creating a good statistical model of a biological sequence extremely difficult.

Many approaches to representations of sequences by specialized data structures and models have been tried and are still valid, such as suffix trees (Bejerano, 2001), regular expressions (Nicolas, 2004) or motif descriptions (Bailey, 2009). Perhaps the most successful and widely known techniques are based on hidden Markov models (Karpplus, 1998), a class of stochastic models working with the probability of individual symbols in sequences. The most commonly used models are the profile hidden Markov models (pHMM) (Eddy, 1998) where each hidden state represents a specific position in the biological sequence. Models where states correspond to a group of sequence positions also exist. Such models have been used for transmembrane protein topology prediction (Viklund, 2004) or gene finding. (Pachter, 2001) Lately, more complex variations appeared such as jumping profile hidden Markov models (jpHMM) (Schultz, 2006) or the gapped pattern-mining-based hidden Markov model VOGUE (Zaki, 2010). VOGUE adds the ability of modelling gapped patterns and its biggest contribution is the idea of combing data mining with data modelling. We have adapted this data mining technique as a part of the parameter learning process of our model. jpHMM improve upon simple HMMs by allowing jumping between different sub-models. That allows jpHMM to better model alternating contexts in sequences. Strict adherence to the markovian property causes the history of visited states to be completely forgotten after every jump to a different sub-model.

To address this shortcoming, we propose a new type of hidden Markov model that is a combination of jumping profile HMMs and variable order HMMs. Same as jpHMM, our model consists of distinct sub-models, each representing a different context in the sequence. Similarly to jpHMM, our model can arbitrarily switch between these sub-models. Our model adds a context memory by remembering the last active state of each sub-model,
so that these can continue the computation uninterruptedly after resuming control. It means that each sub-model can represent different arbitrarily-gapped context in the sequence and cswHMM can combine them together. That is why cswHMM has a unique ability to analyze sequences that contain mixed signals with interleaving patterns.

We envision applications of cswHMM in several areas of biological sequence analysis. For example, proteins from a single family or superfamily often share a common amino acid core, but differ in the surface amino acids, often located in loops of varying lengths. While the composition of the core is driven by the necessity for predictable protein folding, the external parts may be determined by possible binding partners, the surrounding environment or a required enzymatic activity. The two kinds of sequences alternate in most currently known protein folds (Fernandez-Fuentes, 2010), therefore modeling such sequences with cswHMM seems natural. Another candidate for the application of cswHMM may be found in gene prediction. The position of genes in nucleotide sequences is often predicted using HMMs. In the case of eukaryotic genes, part of the HMM predicts exons interleaved with introns. Because the exons contain 3-nucleotide codons which must remain in-frame even between two exons separated by an intron, successful prediction of precise exon/intron boundaries requires a memory mechanism identical to the one proposed in cswHMMs (Majoros, 2009). Membrane proteins often form interfaces between two environments separated by the membrane, the parts of the protein that form one or the other part of such interface are commonly interleaved in the primary sequence (Krogh, 2001), again giving a possibility to model such sequences with cswHMM better than with alternative models. The other broader goal is to identify new gapped motifs in protein sequences and the possibility of their combination. This will allow us to contribute to a better understanding of protein sequence composition (Ganapathiraju, 2005).

The rest of the paper is organized as follows. In Section 2 we describe in detail the general design of cswHMM and its formal definition. In the end of the section we show a possible way of unsupervised learning of the cswHMM based on mined gapped patterns. In Section 3 we present two applications of our model to protein sequence analysis. Finally, Section 4 concludes our work and presents possible future research.

2 METHODOLOGY

HMM is the mathematical basis for our model since cswHMM is a case of a HMM with special topology. To summarize and unify the notification we will briefly describe the formal definition of HMM.

2.1 HMM Definition

HMMs are defined by a set of discrete hidden states $S = \{q_1, ..., q_n\}$ a transition function $T(q_x, q_y) = P(q_y|q_x)$ providing $P(q_y|q_x)$ the probability of transition from state $q_x$ to state $q_y$; by its output density probability function $O(q_x) = P(o|q_x)$ where $P(o|q_x)$ is a random variable over the set of possible observations determining probability of these observations. For applications in sequence analysis the set of possible observations is usually equal to sequence alphabet and it is a discrete variable. The last thing to define in HMM is a vector of prior probabilities. In the following text we omit definitions of these, since they are irrelevant to our model and can always be replaced by an additional initial state and a corresponding addition to the transition function.

2.2 cswHMM Definition

As explained in Section 1, the basic idea of cswHMM is to combine different sub-models so they can transit from one to another while keeping the last active state in all other sub-models in a specialized memory. A classical hidden Markov model is memoryless, but we can encode the information about the last active state in the topology of the HMM.

![Figure 1: a) Transition diagram of two HMMs we want to join. b) Transition diagram of joined cswHMM.](image)

To join two HMM sub-models with a set of states X and Y, we create a model with two sets of states, each composed of the Cartesian product of X and Y. Each state of cswHMM can be identified by an
ordered triplet $[i, q_x, q_y]$ where $i \in \{1,2\}$ is the number of the currently active sub-model. $q_x \in X$ and $q_y \in Y$ are the last active states in the respective sub-models. Figure 1 illustrates how two simple HMMs are joined to cswHMM. In the next paragraph we present the formal definition of cswHMM for joining of $n$ sub-models. We define the emission and transition probability functions of such model.

Suppose we have a set of HMMs $\Lambda = \{H_1, \ldots, H_n\}$, where each $H_i$ has states $s_i = \{q_{i1}, \ldots, q_{im}\}$, transition probability function $t_i(q_{ij}, q_{ik}) = P(q_{ik} | q_{ij})$, and output probability density function $o_i(o, q_{ij}) = P(o | q_{ij})$, where $j, k \in \{1, \ldots, m\}$ and $o$ are possible observation. Let there be an index set $I = \{1, \ldots, n\}$. Then cswHMM created as a combination of individual HMMs from $\Lambda$ would have states $S$ according to Equation 1. Equation 2 defines the emission probability function $O$ and the transition probability function $T$ of such cswHMM is defined by Equations 3-5.

$$S = I \times s_1 \times s_2 \times \ldots \times s_n = \bigcup_{i=1}^{m} \bigcup_{j=1}^{n} \{q_{i1}, \ldots, q_{in}\}$$

$$O(q_{i1}, \ldots, q_{in}) = a_i(q_{i_1})$$

$$T(q_{i1}, \ldots, q_{in}, q_{i_{j-1}}, q_{i_{j+1}}, \ldots, q_{in}) = t_i(q_{i_{j-1}}, q_{i_{j+1}}, \ldots, q_{i_{j+1}})$$

$$T(q_{i1}, \ldots, q_{in}, q_{i_{j-1}}, q_{i_{j+1}}, \ldots, q_{in}; i \neq l) = t_l(q_{i1}, q_{i_{l_1}}) * SW(i, l)$$

otherwise $T(x, y) = 0$

Where $SW(i, l)$ in the equation (4) is the probability of switching from state $j$ of sub-model $H_l$ to sub-model $H_i$. Essentially, parameter $SW(i, l)$ reflects the frequency of switching between contexts. Since we may not know the mapping of contexts on the data until the end of the learning procedure, estimation of this parameter is problematic and has to be done iteratively during the learning process or we must use some known properties of the data.

With this topology the cswHMM can switch between its underlying sub-models without losing knowledge of the continuity in the sub-models. That makes cswHMM a powerful tool for modelling data with interleaving patterns.

### 2.3 State Space Reduction

The drawback of such a general model is its complexity, since cswHMM composed of $n$ HMM, each with $m$ states $n * m^m$ has states. Since the algorithms for inference of HMM have time complexity $O = (N^2 * L)$ where $N$ is the number of states and $L$ is the length of analyzed sequence, the time complexity of inference for such a cswHMM would be $O = (n^2 * m^m * L)$. It is therefore necessary to build cswHMMs from a small number of relatively small sub-models in order to keep the models computationally feasible.

In real-world problems it is not usually necessary to combine all states of every sub-model, since in a linear sequence the number of directly neighbouring and possibly interleaved contexts is limited. The following methods can be used to lower the state complexity of cswHMM. If there are set of states of individual sub-model which do not switch to other sub-models, we can model such sub-models as a hierarchical HMM, where the “non-switching” set of states will be encapsulated in one higher-level state. The switching will take part only between these higher-level states.

The second method to lower state complexity is to combine only the sub-models that might interfere in the sequence. If we had two sub-models that we know will not mix in the sequence, we can create two separate cswHMMs, each with only one such model and then connect these cswHMM in parallel via a general state. This method allows us to create one complex model of a protein family that is made of many relatively small cswHMMs. Thus, the overall state complexity of the model is computationally feasible.

To conclude this section, let us emphasize that cswHMM may itself be a deep and complex model, but it is always a result of combining simpler HMMs. As such, its overall quality always depends on the sub-models we use.

### 2.4 Learning cswHMM

As mentioned before, the combination of models and the lack of precise knowledge about switching between them makes it difficult to learn model parameters directly from data. For HMMs this is commonly done by some type of EM algorithm. Inability to directly separate training data into sub-models requires a different approach. A possible
solution is to use data mining to estimate the separation of data into sub-models and a subsequent data modelling step based on the mined patterns. A similar technique of combining data mining and data modelling has been used in VOGUE (Zaki, 2010), we therefore used its backbone as a basis for our algorithm. Figure 2 shows the general scheme of the necessary algorithms. The main difference in our approach is the separation of patterns using clustering to obtain more models for different contexts. Next, we will describe individual parts of our algorithm in detail.

We used pattern mining algorithm VGS used as a base for the VOGUE model, which itself is a modification of cSPADE (Zaki, 2001), a method for constrained sequence mining. VGS finds all patterns of length at most $k$, with the maximum gap $g$ between each two elements that have minimal frequency $f$ in the sequence. Variables $k, g, f$ are parameters of the algorithm set by the user. VGS starts with patterns of lengths $k = 1$ and then incrementally couples those with enough positions in mutual proximity (defined by $g$) to reach frequency $f$. As an output of pattern mining we have a set of relevant patterns, each consisting of its ID and a list of positions of patterns occurring in the sequence.

In the second step of our algorithm we compute a similarity matrix in order to cluster mined patterns in the different sets representing individual contexts in the sequences. The logic behind “similarity” of two patterns is that if they frequently overlap and extend each other it is likely that they belong to the same context and therefore to the same pattern set. On the other hand patterns which frequently interleave and mix without coinciding with each other are more likely to belong to different contexts. We therefore define a similarity function $s(p_x, p_y) \rightarrow [0,1]$ in Equation 7 where $p_x$ and $p_y$ are patterns we want to compare.

$$s(p_x, p_y) = \left(\frac{1}{1 + \max(0, c_1 + \cdots + c_n)}\right)^\pi$$

Here $n$ is the number of instances of $p_x$ and $p_y$ overlapping in the sequence. $c_1, \ldots, c_n$ are respective distance coefficients of these collisions and $\pi$ is a coefficient determining the steepness of the function. Distance coefficients $c_i$ are defined by Equation 7.

$$c_i = \frac{w_a(n_x + n_y)}{2} - p \cdot w_p$$

where $n_x$ is the number of elements of pattern $p_x$ that lie in some gap of pattern $p_y$. Similarly, $n_y$ denotes the same for elements of $p_y$ in some gap of $p_x$. $p$ is the number of elements of patterns that coincide and $w_p$ and $w_a$ are weights set by the user.

For the clustering itself we used direct clustering as implemented in the clustering tool CLUTO (Karypis, 2002), since direct clustering is the best clustering method for small number of clusters. Each cluster represents one set of patterns that in turn represent one context in the sequence.

Individual HMMs which are the bases for our cswHMM are created from each pattern set similarly to the VOGUE algorithm, except we don’t add the gap states in the individual HMMs. Instead we add an additional HMM consisting of a single gap state, representing the parts of sequence not covered by any other HMMs. cswHMM is consequently created by combining these HMMs as described in Section 2.

Once we obtain an estimation of the appropriate cswHMM, we can separate data into sub-models and fine-tune our model using common EM methods, such as the popular Baum-Welsch algorithm.

3 VALIDATION AND POSSIBLE APPLICATIONS

3.1 PROSITE Classification

With this type of unsupervised learning we tested the performance of cswHMM as applied to the problem of protein classification. To compare our model with other existing HMMs, we chose the same dataset and type of experiment as the authors of the VOGUE (Zaki, 2010) model.

From the dataset of protein families PROSITE (Nicolas, 2004), ten families were chosen as a
testing dataset. Each family was divided into a training and testing set. Precisely, 75% of family members were used as training data for individual models, while remaining 25% of each family was compiled into a classification testing set.

Table 1: Accuracy of protein classification for individual protein families from PROSITE and overall accuracy.

<table>
<thead>
<tr>
<th>Class</th>
<th>cswHMM</th>
<th>VOGUE</th>
<th>HMMER</th>
<th>HMM</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDOC00662</td>
<td>81,82</td>
<td>81,82</td>
<td>72,73</td>
<td>27,27</td>
</tr>
<tr>
<td>PDOC00670</td>
<td>85,71</td>
<td>80,36</td>
<td>73,21</td>
<td>71,4</td>
</tr>
<tr>
<td>PDOC00561</td>
<td>90,48</td>
<td>95,24</td>
<td>42,86</td>
<td>61,9</td>
</tr>
<tr>
<td>PDOC00064</td>
<td>85,71</td>
<td>85,71</td>
<td>85,71</td>
<td>85,71</td>
</tr>
<tr>
<td>PDOC00154</td>
<td>71,88</td>
<td>71,88</td>
<td>71,88</td>
<td>59,38</td>
</tr>
<tr>
<td>PDOC00224</td>
<td>91,67</td>
<td>87,5</td>
<td>100</td>
<td>79,17</td>
</tr>
<tr>
<td>PDOC00271</td>
<td>91,89</td>
<td>89,19</td>
<td>100</td>
<td>64,86</td>
</tr>
<tr>
<td>PDOC00343</td>
<td>92,85</td>
<td>89,29</td>
<td>96,43</td>
<td>71,43</td>
</tr>
<tr>
<td>PDOC00397</td>
<td>80</td>
<td>100</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>PDOC00443</td>
<td>85,71</td>
<td>100</td>
<td>85,71</td>
<td>85,71</td>
</tr>
<tr>
<td>Average</td>
<td>86,38</td>
<td>85,11</td>
<td>80,43</td>
<td>67,66</td>
</tr>
</tbody>
</table>

We trained the cswHMM with different values of parameters of maximal pattern length, maximal gap length, minimal pattern frequency, similarity function coefficients and the switching probability. The models that gave the best performance were chosen. The optimal value of maximal pattern length was found to be 6 to 7. The maximal gap length was found not to significantly influence the results if higher than 5. The average probability of switching between sub-models was 0.86. This means that most of the modelled patterns were gapped.

Table 1 shows the comparison of results of different models on individual datasets and the overall probability of prediction. cswHMM is comparable with other methods and in overall probability it even slightly surpass them, but this type of application should not be the primary function of cswHMM. We use it more as a validation of our proposed model whose main purpose is to improve analyses of sequences with mixed contexts and to identify those contexts. To do so, we currently develop a tool to analyze individual sub-models and their performance during sequence analysis.

3.2 Loop Modelling

We analyzed eight arbitrarily selected protein families defined in the Pfam database (Finn, 2010), using the alignment of seed sequences of each family to determine conserved (protein core) and variable (loops) regions. In each alignment we identified possible loop positions as positions where more than 30% of aligned sequences had a gap. Amino acids at these positions were used to create a database of short sequences that represent the possible loops.

Table 2: Logarithmic probabilities for combined HMM, classic profile HMM and their comparison.

<table>
<thead>
<tr>
<th>Pfam code</th>
<th>cswHMM</th>
<th>pHMM</th>
<th>difference</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF00078</td>
<td>-3.738</td>
<td>-3.670</td>
<td>0.07</td>
<td>1.84</td>
</tr>
<tr>
<td>PF00024</td>
<td>-3.856</td>
<td>-3.756</td>
<td>0.10</td>
<td>2.66</td>
</tr>
<tr>
<td>PF00117</td>
<td>-3.722</td>
<td>-3.709</td>
<td>0.01</td>
<td>0.34</td>
</tr>
<tr>
<td>PF00171</td>
<td>-3.834</td>
<td>-3.798</td>
<td>0.04</td>
<td>0.95</td>
</tr>
<tr>
<td>PF00227</td>
<td>-3.699</td>
<td>-3.685</td>
<td>0.01</td>
<td>0.39</td>
</tr>
<tr>
<td>PF00246</td>
<td>-3.786</td>
<td>-3.695</td>
<td>0.09</td>
<td>2.46</td>
</tr>
<tr>
<td>PF03129</td>
<td>-3.856</td>
<td>-3.752</td>
<td>0.10</td>
<td>2.77</td>
</tr>
<tr>
<td>PF01436</td>
<td>-3.864</td>
<td>-3.812</td>
<td>0.05</td>
<td>1.36</td>
</tr>
<tr>
<td>Average</td>
<td>-3.794</td>
<td>-3.735</td>
<td>0.060</td>
<td>1.60</td>
</tr>
</tbody>
</table>

We trained a four-state HMM on this database to create a simple loop model for each family. Subsequently, we used the core profile HMM of each family and combined it with the corresponding loop model to create a simple cswHMM. We computed logarithmic probabilities of generating individual family without insertion states sequences with the new model and compared them with the logarithmic probabilities for classical pHMM. The results are shown in Table 2 and show slight improvement of generating probability with cswHMM.

The loop modelling experiment has shown that the cswHMM approach may find use in protein sequence analysis where blocks of amino acids interfacing with different environments (protein core, membrane or cytoplasm) are interspersed with contrasting blocks. Numerical treatment corresponding to the proposed cswHMM provided an average 1.6% improvement in sequence description, consistently better for all studied families (Table 2). Models used within the cswHMM framework could possibly represent different building blocks, where at least one of the block types follows a predetermined order of amino acids and where mixing of the blocks is variable or optional.

4 CONCLUSIONS AND FUTURE WORK

In this paper we have presented a new model that is a type of variable-order hidden Markov model with ability to analyze mixed contexts in sequences. We
also introduced one possible method for unsupervised learning of cswhHMM based on mined gapped patterns and showed two possible applications of such model to protein sequence analysis.

In our future work we plan to apply our cswhHMM to identify other mixed contexts in biological sequences. The possible application area is twofold. We can use our model to combine existing models of sequences that are known to have non-uniformly mixed contexts and in that way increase the description ability of those models. Some examples of such sequences are in Section 1. The other way of application of cswhHMMs is to use unsupervised learning over different sets of sequences and try to find new, currently unknown common gapped patterns and their possible combinations. In that way we could reveal new rules governing biological sequence composition.

ACKNOWLEDGEMENTS

This work has been supported by the Grant Agency of the Czech Republic grant GD204/08/H054.

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