A protein classification engine based on stochastic finite state automata

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Abstract: Accurate protein classification is one of the major challenges in modern bioinformatics. Motifs that exist in the protein chain can make such a classification possible. A plethora of algorithms to address this problem have been proposed by both the artificial intelligence and the pattern recognition communities. In this paper, a data mining methodology for classification rules induction is proposed. Initially, expert – based protein families are processed to create a new hybrid set of families. Then, a prefix tree acceptor is created from the motifs in the protein chains, and subsequently transformed into a stochastic finite state automaton using the ALERGIA algorithm. Finally, an algorithm is presented for the extraction of classification rules from the automaton.

Keywords: protein classification, data mining, bioinformatics, motifs, finite state automata

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1. Protein Classification

Protein classification is currently one of the most interesting problems in computational biology. The biological action of proteins is traditionally identified by time consuming and expensive in-vitro experiments. However, recent developments in bioinformatics have enabled the use of computational tools and techniques towards this end [1]. Clustering algorithms [2], artificial neural networks [3], decision trees [4, 5] and statistical models [6] are few of the methods currently employed. Motifs that can be found in a protein chain have provided a higher level of abstraction to the problem, since protein properties are mainly defined by them. Motifs can be either patterns, which are short aminoacid chains with a specific order, or profiles, which are computational representations of multiple sequence alignments derived by the use of hidden Markov models.

On the other hand, proteins can be assorted into families, each family containing proteins with similar functions. Those families can either be expert–based, meaning that they have been experimentally specified and their significance is biologically meaningful, or computer–generated, meaning that they have been created with the use of unsupervised protein classification algorithms. In the latter case the major disadvantage is that the protein classes will not necessarily have any biological meaning or significance. In the former, expert – based classes are often overlapping, thus adding to the complexity of the classification algorithms.

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2. Core Engine

In [7] a technique for extraction of classification rules using finite state automata was introduced, the classes being expert-based protein families. The technique is outlined in Figure 1. First, the training set is constructed from a set of known proteins. The next step is the creation of the Prefix Tree Acceptor (PTA) using the protein chains of the train set. Using the ALERGIA algorithm [8], the PTA is converted to an equivalent Stochastic Finite State Automaton (SFSA), in which every transition is associated with a probability. Since the SFSA is a generalized representation of the protein structure, information can be obtained directly from it. As a result, certain probabilities can be calculated, such as the probability of a protein chain to contain a certain motif or a specific subset of motifs. Using these probabilities, rules can be extracted to better describe the form of the protein chains. In this paper we extend this technique by introducing both a hybrid form of classification and a new algorithm for the extraction of classification rules from the automata.

Figure 1: FSA based technique for protein classification rules induction

3. Methodology Outline

The proposed methodology for rule induction consists of three sequential parts: a) Preprocessing, b) Model Creation and c) Rule Extraction.

During the Preprocessing phase, all the protein classes in the training set are recombined to create Virtual Classes. These constitute a hybrid form of the original partitioning, where the overlapping areas between classes are assigned to new classes, as shown in Figure 2. In this case, the classes that arise still maintain their biological meaning, but they also represent more closely the proteins they contain. On an algorithmic level, this step creates disjoint sets of proteins, which can be independently processed for classification rules.

For each ensuing Virtual Class, during the Model Creation phase, a SFSA is created using the algorithm described in [7], modified as follows: before the construction of the PTA, the motif list is
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transformed in order to move the most frequently appearing motifs at the beginning of each list. This transformation ensures that the order of appearance of the motifs in the lists will have no impact on the classification rules.

Figure 2: Virtual protein classes

The final step in the methodology is the extraction of classification rules from the SFSA. This is accomplished using the Max_Probability_Path algorithm, presented in Box 1. This algorithm is a modification of the shortest path: it finds the maximum probability path from the starting vertex to any other vertex in the SFSA. However, only the paths to the terminating vertices are important due to the fact that they define the conditions necessary for a protein to belong to a class. For each vertex the algorithm stores its predecessor, thus allowing tracing the path from the starting vertex to any other vertex in the SFSA.

Box 1: Max_Probability_Path algorithm

The concept behind the algorithm is that the probability of any path from the starting node is equal to the product of the probabilities of the edges that comprise the path. A classification rule is derived from a path simply by reading the edge “types”, i.e. the motifs that exist in the path. The probability of the path is a quantitative measure of the interest of the rule: the higher the probability of the path the stronger the rule will be.
An overview of the complete methodology is presented in Figure 3.

![Figure 3: Methodology Outline](image)

### References


