Prediction of Disulfide Bonding Pattern Based on Support Vector Machine with Parameters Tuned by Multiple Trajectory Search

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Abstract: - The prediction of the location of disulfide bridges helps solving the protein folding problem. Most of previous works on disulfide connectivity pattern prediction use the prior knowledge of the bonding state of cysteines. In this study an effective method is proposed to predict disulfide connectivity pattern without the prior knowledge of cysteines’ bonding state. To the best of our knowledge, without the prior knowledge of the bonding state of cysteines, the best accuracy rate reported in the literature for the prediction of the overall disulfide connectivity pattern ($Q_p$) and that of disulfide bridge prediction ($Q_c$) are 48% and 51% respectively for the dataset SPX. In this study, the cystein position difference, the cystein index difference, the predicted secondary structure of protein and the PSSM score are used as features. The support vector machine (SVM) is trained to compute the connectivity probabilities of cysteine pairs. An evolutionary algorithm called the multiple trajectory search (MTS) is integrated with the SVM training to tune the parameters for the SVM and the window sizes for the predicted secondary structure and the PSSM. The maximum weight perfect matching algorithm is then used to find the disulfide connectivity pattern. Testing our method on the same dataset SPX, the accuracy rates are 54.5% and 60% for disulfide connectivity pattern prediction and disulfide bridge prediction when the bonding state of cysteines is not known in advance.

Key-Words: - Disulfide bonding pattern, SVM, multiple trajectory search

1 Introduction

Disulfide bonds play an important structural role in stabilizing protein conformations. The prediction of disulfide bonding pattern helps to a certain degree the prediction of the three-dimensional protein structure and hence its function because disulfide bonds impose geometrical constraints on the protein backbones. Some recent research works do have shown the close relation between the disulfide bonding patterns and the protein structures \cite{1, 2}. As shown in Fig. 1, the structures of (a) the tick anticoagulant peptide (1TAP), a protease inhibitor, and (b) the bovine pancreatic trypsin inhibitor (1QLQ), a serine protease inhibitor. Their sequence identity is only 18.2\%, the absence of significant sequence identity between 1TAP and 1QLQ was noted by Antuch et al. \cite{3} and PSI-BLAST searches in the SwissProt/TrEMBL and NR database were unsuccessful in identifying the similarity between these two proteins. In disulfide-bonding base classification, these two proteins have the same disulfide-bonding connectivity pattern (1-6, 2-3, 4-5), and all of which are classified in the BPTI-like superfamily in SCOP \cite{4}.

In the realm of the disulfide bond prediction, two problems are addressed. The first is the prediction of the disulfide bonding states and the second is the prediction of the disulfide bonding pattern. Recently, significant progress has been made in the prediction of the disulfide bonding states. Several methods based on statistical analysis \cite{5}, neural networks \cite{6, 7}, or support vector machines \cite{8} had been proposed. They are quite effective in predicting the bonding state of cysteines with the accuracy rates around 81\%-90\%.
Recently, several methods were proposed for the prediction of the disulfide bonding pattern. The first method was presented by Fariselli and Casadio [9]. They reduced disulfide connectivity to the graph matching problem in which vertices are oxidized cysteines and edges are labeled by the strength of interaction (contact potential) in the associated pair of cysteines. The Monte Carlo simulated annealing method is used to find the optimal values of contact potentials and finally the disulfide bridges are located by finding the maximum weight perfect matching. Fariselli et al. [10] improved their previous results by using neural networks to predict the cysteine pairwise interactions. Vullo and Frasconi [11] developed an ad-hoc recursive neural network for scoring labeled undirected graphs that represent the connectivity pattern and they improved the accuracy rate of bonding pattern prediction significantly from 34% to 44%. Cheng et al. [12] improved the prediction accuracy by using two-dimensional recursive neural networks to predict connectivity probabilities between cysteine pairs. Ferrè and Clote [13] designed the diresidue neural network to predict connectivity probabilities between cysteine pairs. They also used secondary structure information and diresidue frequencies in their training. Tsai et al. [14] used the support vector machine to predict connectivity probabilities between cysteine pairs. The features used in training the support vector machine are local sequence profiles and the linear distance of cysteines. All above mentioned methods are based on the reduction of the connectivity pattern prediction to the maximum weight perfect matching problem. The following four methods are not based on this reduction. Chen and Hwang [15] used the support vector machine to predict the bonding pattern directly. The features they used in training the support vector machine are the coupling between the local sequence environments of cysteine pairs, the cysteine separations, and the amino acid content. Zhao et al. [16] used a simple feature called cysteine separations profiles (CSP) to predict the connectivity patterns. Chen et al. [17] proposed a two-level model. Lu et al. [18] obtained the accuracy of 73.9% by using GA to optimize feature selection for the SVM. Song et al. [19] obtained the accuracy of 74.4% by using multiple sequence vectors and secondary structure. This accuracy rate is the best one found in the literature. Rubinstein et al. [20] analyzes the correlated mutation patterns in multiple sequence alignments to predict the disulfide bond connectivity. All these methods except Cheng et al. [12] and Ferrè et al. [13] assume that the bonding states are known. The method proposed by Cheng et al. [12] and Ferrè et al. [13] can be applied whether the bonding states are known or not.

**1.1 Support vector machine**

Support vector machine (SVM) is a supervised learning method used for classification [21]. It is believed to be superior to traditional statistical and neural network classifiers. However, it is critical to determine suitable combination of SVM parameters regarding classification performance. A special property of SVM is that it can simultaneously minimize the empirical classification error and maximize the geometric margin. It is a useful technique for data classification and regression and has become an important tool for machine learning and data mining. It is a powerful methodology for
solving problems in nonlinear classification, function estimation and density estimation which has also led to many applications, such as the image interpretation, data mining and other biotechnological fields [22-24]. SVM is generally used for data which can be classified into two clusters, however, classification of multiple clusters can also be easily expanded [25].

In general, SVM has better performance when competed with existing methods, such as neural networks and decision trees [26-28]. Recently, application of SVM in medicine has grown rapidly. For examples, it has been applied in prediction of RNA-binding sites in proteins [29], protein secondary structure prediction [30], remote sensing image classification [34], and electrical energy consumption forecasting [33], and pattern recognition and data classification [35].

The goal of support vector machine (SVM) is to separate multiple clusters by constructing a set of separating hyperplanes with greatest margin to the boundary of each cluster. For a two-class classification example, let us view the input data as a vector

\[ g(x) = w^T x + w_0 = 0 \]  

(1)

The distance of a point from a hyperplane is given by

\[ z = \frac{g(x)}{||w||} \]  

(2)

As shown in Fig. 2, the values of \( w \) and \( w_0 \) in Eq. (1) are scaled so that the values of \( g(x) \) at the nearest points in class 1 and class 2 equal to 1 and -1 respectively. Therefore, finding the hyperplane becomes a nonlinear quadratic optimization problem, which can be formulated as:

\[
\text{Minimize } J(w) = \frac{||w||^2}{2} \\
\text{Subject to } y_i(w^T x_i + w_0) \geq 1, \quad i = 1, 2, ..., N
\]  

(3)

The above minimizer must satisfy Karush-Kuhn-Tucker (KKT) condition, and it can be solved by considering Lagrangian duality. The problem can be stated equivalently by its Wolfe dual representation form:

\[
\text{Maximize } L(w, w_0, \lambda) = \frac{w^T w}{2} - \sum_{i=1}^{N} \lambda_i [y_i(w^T x_i + w_0) - 1] \\
\text{Subject to } w = \sum_{i=1}^{N} \lambda_i y_i x_i, \quad \sum_{i=1}^{N} \lambda_i y_i = 0, \quad \lambda \geq 0
\]  

(4)

where \( L(w, w_0, \lambda) \) is the Lagrangian function and \( \lambda \) is the vector of Lagrangian multipliers. By comparing Eqs. (3) and (4), it is noted that the first two constraints in Eq. (4) become equality constraints and this makes the problem easier to be solved. After a little bit algebra manipulation, Eq. (4) becomes

\[
\max_k \left( \sum_{i=1}^{N} \lambda_i y_i \right) - \frac{1}{2} \sum_{i,j} \lambda_i \lambda_j y_i y_j x_i^T x_j
\]  

(5)

Subject to \( \sum_{i=1}^{N} \lambda_i y_i = 0 \) with \( \lambda \geq 0 \)

As soon as the Lagrangian multipliers are obtained by maximizing the above equation, the optimal hyperplane can be obtained from

\[
w = \sum_{i:\lambda_i \neq 0} \lambda_i y_i x_i
\]  

in Eq. (4).

Once the optimal hyperplane is obtained, classification of a sample is performed based on the sign of the following equation:

\[ g(x) = w^T x + w_0 = \sum_{i=1}^{N} \lambda_i y_i x_i^T x + w_0 \]  

(6)

where \( N_s \) is the number of support vectors. For a vector \( x \in R^l \) in the original feature space, assume that there exists a mapping \( \phi \) from \( x \in R^l \) to \( y = \phi(x) \in R^k \), where \( k \) is usually much higher than \( l \). Then it is always true that

\[ \sum_{i} \phi_i(x) \phi_i(z) = K(x, z) \]  

(7)

where \( \phi_i(x) \) is the \( i \)th component of the mapping and the kernel function \( K(x, z) \) is a symmetric function satisfying the following condition.
\[ \int K(x,z)g(z)dxdz \geq 0, \text{ and } \int g(x)^2 dx \leq \infty \] (8)

For a nonlinear classifier, various kernels, including polynomial, radial basis function, and hyperbolic tangent, as shown in Eq. (9) can be used for mapping the original sample space into a new Euclidian space in which Mercer’s conditions are satisfied. The linear classifier can then be designed for classification.

\[ K(x,z) = (x^T z + 1)^q, \quad q > 0 \] (9a)
\[ K(x,z) = \exp\left(-\frac{\|x - z\|}{\sigma^2}\right) \] (9b)
\[ K(x,z) = \tanh(\beta x^T z + \gamma) \] (9c)

n-fold cross-validation of the SVM model is achieved by dividing the dataset into n folds. When some fold is reserved for testing, the other n-1 folds are used for training the model.

1.2 Multiple Trajectory Search

The multiple trajectory search (MTS) had been presented for large scale global optimization [38]. The MTS had also been used to solve the multi-objective optimization problems and obtained satisfactory results [39]. It uses multiple agents to search the solution space concurrently. Each agent does an iterated local search using one of three candidate local search methods. By choosing a local search method that best fits the landscape of a solution’s neighborhood, an agent may find its way to a local optimum or the global optimum.

1.2.1 Orthogonal Array and Simulated Orthogonal Array

The concept of orthogonal arrays which are used in experimental design methods is briefly introduced. Suppose in an experiment, there are k factors and each factor has q levels. In order to find the best setting of each factor’s level, qk experiments must be done. Very often, it is not possible or cost effective to test all qk combinations. It is desirable to sample a small but representative sample of combinations for testing. The orthogonal arrays were developed for this purpose. In an experiment that has k factors and each factor has q levels, an orthogonal array OA(n,k,q,t) is an array with n rows and k columns which is a representative sample of n testing experiments that satisfies the following three conditions. (1) For the factor in any column, every level occurs the same number of times. (2) For the t factors in any t columns, every combination of q levels occurs the same number of times. (3) The selected combinations are uniformly distributed over the whole space of all the possible combinations. In the notation OA(n,k,q,t), n is the number of experiments, k is the number of factors, q is the number of levels of each factor and t is called the strength.

The orthogonal arrays exist for only some specific n’s and k’s. So it is not appropriate to use the OA in some applications. Tseng et al. [38] proposed the simulated OA (SOA). The SOA satisfies only the first of the above mentioned three conditions, but it is easy to construct an SOA of almost any size. Suppose there are k factors and each factor has q levels, an m×k simulated orthogonal array SOA\textsubscript{m×k} with m being a multiple of q can be generated as follows. For each column of SOA\textsubscript{m×k}, a random permutation of 0, 1, ..., q-1 is generated and denoted as sequence C. Then the elements in C are picked one by one sequentially and filled in a randomly chosen empty entry of the column. If all elements in C were picked, the process picks elements again from the beginning of C. So in every column of SOA\textsubscript{m×k}, each of q elements will appear the same number of times (condition 1).

1.2.2 The Multiple Trajectory Search

The MTS generates M initial solutions by utilizing the simulated orthogonal array SOA\textsubscript{m×N}, where the number of factors corresponds to the dimension N and the number of levels of each factor is taken to be M. So each of 0, 1, ..., M-1 will appear once in every column. Using SOA tends to make these M initial solutions uniformly distributed over the feasible solution space. The initial search range for local search methods is set to half of the difference between the upper bound and the lower bound. Afterwards, local search methods will change the search range.

The MTS consists of iterations of local searches until the maximum number of function evaluations is reached. In the first iteration, the MTS conducts local searches on all of M initial solutions. But in the following iterations, only some better solutions are chosen as foreground solutions and the MTS conducts local searches on these solutions. Three local search methods are provided for the MTS. The MTS will first test the performance of three local search methods and then choose the one that performs best, that is, the one that best fits the landscape of the neighborhood of the solution, to do the search. After conducting the search on foreground solutions, the MTS applies Local Search 1 to the current best solution trying to improve the current best solution. Before the end of an iteration, some better solutions are chosen as the foreground solutions.
Evaluate function values of $X_i$'s 

Multiple Trajectory Search
/*Generate $M$ initial solutions*/
Build simulated orthogonal array $SOA_{MN}$
For $i = 1$ to $M$
  For $j = 1$ to $N$
    $X_{ij} = (i + (j-1)*SOA[i,j]) / (M-1)$
  End For
End For
End While
Evaluate function values of $X_i$'s
For $i = 1$ to $M$
  Enable[i] ← TRUE
  Improve[i] ← TRUE
  SearchRange $X_i$ = $(UPPER\_BOUND-LOWER\_BOUND) / (2^{j-1})$
End For

While (#ofEvaluation ≤ predefined_max_evaluation)
  For $i = 1$ to $M$
    If Enable[i] = TRUE
      Then
        Grade $X_i$ = 0
        $LS_1$ TestGrade ← 0
        $LS_2$ TestGrade ← 0
        $LS_3$ TestGrade ← 0
        For $j = 1$ to #ofLocalSearch
          $LS_1$ TestGrade ← $LS_1$ TestGrade +
            LocalSearch1($X_i$, SearchRange $X_i$)
          $LS_2$ TestGrade ← $LS_2$ TestGrade +
            LocalSearch2($X_i$, SearchRange $X_i$)
          $LS_3$ TestGrade ← $LS_3$ TestGrade +
            LocalSearch3($X_i$, SearchRange $X_i$)
        End For
        Choose the one with the best TestGrade and
        let it be LocalSearch $K$ * K may be 1, 2, or 3 */
    For $j = 1$ to #ofLocalSearch
      Grade $X_j$ ← Grade $X_j$ +
        LocalSearch$K$($X_j$, SearchRange $X_j$)
    End For
  End If
End For
For $i = 1$ to #ofLocalSearchBest
  LocalSearch1(BestSolution, SearchRangeBestSolution)
End For
For $i = 1$ to $M$
  Enable[i] ← FALSE
End For
Choose #ofForeground $X_i$'s whose Grade $X_i$ are best among the $M$ solutions and set their corresponding Enable[i] to TRUE
End While

In the MTS, three local search methods are used for searching different landscape of the neighborhood of a solution. Local Search 1 searches along one dimension from the first dimension to the last dimension. Local Search 2 is similar to Local Search 1 except that it searches along a direction derived from about one-fourth of dimensions. In both local search methods, the search range (SR) will be cut to one-half until it is less than $1 \times 10^{-15}$ if the previous local search does not make improvement. In Local Search 1, on the dimension concerning the search, the solution’s coordinate of this dimension is first subtracted by SR to see if the objective function value is improved. If it is, the search proceeds to consider the next dimension. If it is not, the solution is restored and then the solution’s coordinate of this dimension is added by $0.5 \times$SR, again to see if the objective function value is improved. If it is, the search proceeds to consider the next dimension. If it is not, the solution is restored and the search proceeds to consider the next dimension. Local Search 1 and Local Search 2 are listed in the following.

Function LocalSearch1($X_i$, SR)
If Improve[k] = FALSE
  Then
    SR = SR / 2
    If SR < 1e-15
      Then
        SR = (UPPER_BOUND-LOWER_BOUND) * 0.4
      End If
    End If
  End If
End If
If $X_i$ is better than current best solution
  Then
    grade = grade + BONUS1
    Update current best solution
  End If
If function value of $X_i$ is the same
  Then restore $X_i$ to its original value
Else
  If function value of $X_i$ degenerates
    Then
      restore $X_i$ to its original value
      $X_{ij} = X_{ij} - 0.5 \times$ SR
    End If
    If $X_i$ is better than current best solution
      Then
        grade = grade + BONUS1
        Update current best solution
    End If
    If function value of $X_i$ has not been improved
      Then
        restore $X_i$ to its original value
      Else
        grade = grade + BONUS2
        Improve[k] = TRUE
      End If
    Else
      grade = grade + BONUS2
      Improve[k] = TRUE
    End If
  End If
End If
End If
End For
return grade
Local Search 3 is different from Local Search 1 and Local Search 2. Local Search 3 considers three small movements along each dimension and heuristically determines the movement of the solution along each dimension. In Local Search 3, although the search is along each dimension from the first dimension to the last dimension, the evaluation of the objective function value is done after searching all the dimensions, and the solution will be moved to the new position only if the objective function has been improved at this evaluation. Local Search 3 is described in the following.

Function LocalSearch3(X, SR)
For i = 1 to N
    If Y_i is better than current best solution
        Then grade ← grade + BONUS1
            Update current best solution
    End If
    If grade has not been improved
        Then restore X_i to its original value
    Else
        grade ← grade + BONUS2
    Improve[k] ← TRUE
    End If
End For

2 Materials and Methods
2.1 Dataset
In order to compare the prediction accuracy rates with previously reported method by Cheng et al. [12], the same dataset SPX used by them was employed in the experiment. In SPX, all proteins were extracted from the PDB on May 17, 2004 that contain at least one intrachain disulfide bond and all
proteins that contain less than 12 amino acids were removed. Furthermore, to reduce overrepresentation of particular protein families, Cheng et al. used the UniqueProt, a protein redundancy reduction tool based on the HSSP distance [40], to choose 1018 proteins by setting the HSSP cut-off distance to 10. In SPX, the protein sequences were randomly divided into 10 subsets with roughly equal size for 10-fold cross-validation experiment.

2.2 Methodology
Cheng et al. [12] predicted the bonding state of cysteines first, and then they predicted the disulfide bond pattern for the predicted oxidized cysteines. Our method, instead of predicting bonding state first, directly predicts the bonding probability of all pairs of cysteines. Our method uses the cystein position difference, the cystein index difference, the predicted secondary structure of the protein and the PSSM score as features. The SVM is trained to compute the connectivity probabilities of all the cysteine pairs. The MTS [38] is used to evolve the parameters C and γ for SVM and window sizes for the predicted secondary structure and the PSSM. The maximum weight perfect matching algorithm is then used to find the disulfide connectivity pattern without the prior knowledge of the bonding state of cysteines.

2.3 Features
(1) NCPD (Normalized Cysteine Position Difference): Let \( \{c_1, c_2, ..., c_n\} \) be the positions of the cysteines in ascending order. The normalized cysteine position difference between \( c_i \) and \( c_j \) is defined as \( \frac{|c_i - c_j|}{(c_{max} - c_{min})} \).

(2) NCID (Normalized Cysteine Index Difference): Let \( \{c_1, c_2, ..., c_n\} \) be the positions of the cysteines in ascending order. The normalized cystein ordering index between \( c_i \) and \( c_j \) is defined as \( \frac{|i-j|}{n} \).

(3) PSSM: PSI-BLAST [41] is used to obtain the local sequence profiles. The output file contains four parts. The first part is the position-specific scoring matrix (PSSM). The PSSM score is used as one of features.

(4) PSS (Predicted Secondary Structure): The predicted secondary structure obtained by applying Jones’ prediction method [42] is used as a feature. In practice, the PSIPRED program is used to predict the secondary structure information.

2.4 Construction of Prediction Model Based on SVM
It is noted that SVM is superior to traditional statistical and neural network classifiers in many applications. However, it is critical to determine proper combination of SVM parameters (\( C \) and \( \gamma \)) in order to achieve good classification performance. The SVM implementation used in this work is LIBSVM [43]. Since the prediction rate is highly influenced by the value of the parameters \( C \) and \( \gamma \), the multiple trajectory search [38] is used for finding good settings of parameter values for the SVM and the window sizes for the PSS and the PSSM.

2.5 Multiple Trajectory Search for Selecting SVM Parameters and Window Sizes
The multiple trajectory search can find optimal or near-optimal solution within an acceptable time, and is faster than the dynamic programming or the branch-and-bound strategy. Previously, some research works applied the evolutionary algorithms to select features in the first phase and then used the selected features to train the SVM in the second phase. In this study, the MTS and the SVM training are tightly integrated. Since the values of parameters \( C \) and \( \gamma \) for the SVM are critical to classification accuracy of the SVM, selecting proper values of \( C \) and \( \gamma \) becomes an important task. Traditionally, the regular grid search strategy was used to perform the parameter value selection. However, it is very time-consuming. In this work, the MTS is integrated with the SVM training to select not only the value of parameters \( C \) and \( \gamma \) but also the window sizes of the PSS and the PSSM. As shown in Fig. 3, a chromosome is coded as \( S_i = (C_i, \gamma_i, S_{i1}, S_{i2}) \) where \( C_i \) and \( \gamma_i \) are the log values of the parameters \( C \) and \( \gamma \), and \( S_{i1}, S_{i2} \) are the window sizes for the PSS and the PSSM respectively. The fitness function is defined as the accuracy of the SVM on disulfide connectivity prediction. The flowchart of the integration of the MTS and SVM training is shown in Fig. 4.

<table>
<thead>
<tr>
<th>log_2 C</th>
<th>log_2 ( \gamma )</th>
<th>( S_1 )</th>
<th>( S_2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM Parameters</td>
<td>Window size for the PSS and the PSSM</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 3. Chromosome of the proposed method.
2.6 Maximum weight perfect matching

When a test protein is given, the connectivity probability of each cysteine pair will be computed by the trained SVM. A complete graph is then constructed with all cysteines as nodes and the weight associated with each edge is the disulfide connectivity probability of the pair of cysteines that are incident to this edge. The Gabow’s algorithm [44] is then applied to find the maximum weight perfect matching. Because the Gabow’s algorithm can only be applied with integer edge labels, the disulfide connectivity probability of the pair of cysteines is multiplied by 10000 and then truncated into an integer to represent the weight associated with the edge. This matching represents the prediction result of the disulfide connectivity pattern.

We set a probability threshold, when the disulfide connectivity probability of two cysteines is greater than the probability threshold, then there is a bond between the two cysteines, otherwise there is no bond between the two cysteines. For dataset SPX with 10-fold cross-validation, the results of parameter value selection by the proposed method are as follows: \( \log_2 C = 7.4 \) and \( \log_2 \gamma = -4.6 \) for SVM, the window sizes are 1 and 23 for the PSS and the PSSM. Moreover, the probability threshold is set to 0.22 for a bond to be existed between two cysteins. This value is determined empirically.

3 Experiment Results

In order to evaluate the performance of the prediction, two accuracy indices \( Q_P \) and \( Q_C \) are used:

\[
Q_P = \frac{C_P}{T_P} \quad \text{and} \quad Q_C = \frac{C_C}{T_C}
\]

where \( C_P \) is the number of proteins whose bonding patterns are correctly predicted; \( T_P \) is the total number of proteins in the test set; \( C_C \) is number of disulfide bridges that are correctly predicted and \( T_C \) is the total number of disulfide bridges in test proteins.
Tables 1 and 2 show the results for bridge classification and the prediction of the disulfide bonding pattern of the dataset SPX. For bridge classification, Cheng et al. [12] listed sensitivity and specificity instead of accuracy. From the definition of sensitivity and specificity, in general the value of accuracy lies between them. From Table 1, it is noted that the overall sensitivity is 52% for Cheng’s method and the prediction accuracy ($Q_c$) is 60% for our method. There is an increase of prediction accuracy from 52% to 60%.

As for the disulfide connectivity prediction, Cheng’s method with true secondary structure (SS) and solvent accessibility (SA) information in the inputs has the accuracy rate 51%. And Cheng’s method with predicted secondary structure (PSS) and predicted solvent accessibility (PSA) information in the inputs has the prediction accuracy 48% only. In this study, using predicted secondary structure in the inputs, the prediction accuracy is 54.5%. There is an increase of prediction accuracy from 48% to 54.5%.

Table 1. Bridge classification result for dataset SPX without the prior knowledge of the bonding state of cysteines.

<table>
<thead>
<tr>
<th># of bonds</th>
<th>Cheng et al. (2006)</th>
<th>This work</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sensitivity</td>
<td>Specificity</td>
</tr>
<tr>
<td>1</td>
<td>71%</td>
<td>48%</td>
</tr>
<tr>
<td>2</td>
<td>59%</td>
<td>60%</td>
</tr>
<tr>
<td>3</td>
<td>55%</td>
<td>61%</td>
</tr>
<tr>
<td>4</td>
<td>44%</td>
<td>48%</td>
</tr>
<tr>
<td>5</td>
<td>32%</td>
<td>35%</td>
</tr>
<tr>
<td>6</td>
<td>32%</td>
<td>36%</td>
</tr>
<tr>
<td>7</td>
<td>29%</td>
<td>32%</td>
</tr>
<tr>
<td>8</td>
<td>20%</td>
<td>22%</td>
</tr>
<tr>
<td>9</td>
<td>44%</td>
<td>52%</td>
</tr>
<tr>
<td>10</td>
<td>33%</td>
<td>36%</td>
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<td>17</td>
<td>53%</td>
<td>60%</td>
</tr>
<tr>
<td>25</td>
<td>32%</td>
<td>53%</td>
</tr>
<tr>
<td>26</td>
<td>31%</td>
<td>51%</td>
</tr>
<tr>
<td>All</td>
<td>52%</td>
<td>51%</td>
</tr>
</tbody>
</table>

Table 2. Disulfide connectivity prediction result for dataset SPX without the prior knowledge of the bonding state of cysteines.

<table>
<thead>
<tr>
<th># of bonds</th>
<th>Cheng et al. (2006)</th>
<th>This work</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$Q_p$ with SS and SA</td>
<td>$Q_p$ with PSS and PSA</td>
</tr>
<tr>
<td>1</td>
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<td>All</td>
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4 Conclusion

Recently, the progress in the prediction of the oxidation states of cysteines in protein sequence is significant. But for the prediction of the bonding pattern of cysteines, much research effort is still needed to improve the prediction accuracy. To these authors’ knowledge, all previous approaches except those presented by Ferrè et al. [13] and Cheng et al. [12] assume that the oxidation states of cysteines were known in advance. To practically solve the prediction of the bonding pattern of cysteines, this assumption eventually should be removed. In this work, without the prior knowledge of the oxidation states of cysteines, by integrating the MTS and the SVM training to tune parameters of SVM and the window sizes of the PSS and the PSSM, the proposed method achieves the accuracy of 54.5% on the bonding pattern prediction, which improves the accuracy of 51% obtained by Cheng et al. [12].

References.


