DISULFIND: a disulfide bonding state and cysteine connectivity prediction server

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ABSTRACT

DISULFIND is a server for predicting the disulfide bonding state of cysteines and their disulfide connectivity starting from sequence alone. Optionally, disulfide connectivity can be predicted from sequence and a bonding state assignment given as input. The output is a simple visualization of the assigned bonding state (with confidence degrees) and the most likely connectivity patterns. The server is available at http://disulfind.dsi.unifi.it/.

INTRODUCTION

Disulfide bridges play a major role in the stabilization of the folding process and, consequently, in studies related to structural and functional properties of specific proteins. In addition, knowledge about the disulfide bonding state of cysteines may help the experimental structure determination process and may be useful in other genomic annotation tasks.

DISULFIND uses a combination of machine learning algorithms to predict intrachain bridges from sequence alone. Similar to many other tools of this kind, it solves the prediction problem in two steps. First, the disulfide bonding state of each cysteine is predicted by a binary classifier; second, cysteines that are known to participate in the formation of bridges are paired to obtain a connectivity pattern.

RELATED WORKS

Early work on bonding state employed representations based on local-window multiple alignment profiles and neural networks for discrimination (1,2). Mucchielli–Giorgi *et al.* (3) introduced the idea of adding a global descriptor to improve prediction accuracy. Ceroni *et al.* (4) proposed a method based on a combination of string and vector kernels in conjunction with support vector machines (SVMs). Song *et al.* (5)

applied a linear discriminant using dipeptides as features. Martelli *et al.* (6) suggested the use of hidden Markov models to refine local predictions obtained via neural networks. SVMs are also used in the method presented in (7).

Prediction of connectivity patterns was pioneered in (8) with a method based on weighted graph matching, implemented in the prediction server DCON. Vullo and Frasconi (9) introduced the use of multiple alignment profiles by means of recursive neural networks (RNNs). In this approach, (that still underpins DISULFIND) a global score is assigned to an entire connectivity pattern. In the DAG RNN approach described in (7,10), the probability for a disulfide bond is computed for each pair of cysteines. The associated DIpro server (which also predicts bonding state) is described in (11). Taskar et al. (12) formulated disulfide connectivity as a structured-output prediction problem and solved it using a generalized large-margin machine. Ferrè and Clote (13) proposed a feedforward neural-network architecture with hidden units associated with cysteine pairs and inputs encoding secondary structure; the method is behind the prediction server DiANNA (14). Zhao et al. (15) confirmed that the profile of distances between bonded cysteines is an important feature for prediction of connectivity patterns. This idea has been further exploited in conjunction with SVMs to develop the method behind the prediction server PreCys (16). Finally, CysView (17) is a server that predict patterns by comparison of a query sequence to annotated data bases.

MATERIALS AND METHODS

Multiple alignment profiles

Prediction of protein structural properties is typically more accurate when incorporating evolutionary information encoded in multiple alignment profiles. Profiles are used in DISULFIND both in bonding state and connectivity prediction. They are calculated by using one iteration of the

The authors wish it to be known that, in their opinion, the first three authors should be regarded as joint first authors.

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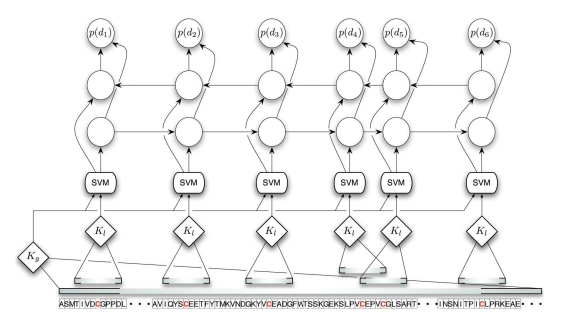


Figure 1. Architecture of the bonding state predictor. The lower level provides independent cysteine predictions based on a local kernel k_I on local attributes, and a global kernel k_g on the entire sequence. The upper level is a BRNN (represented here schematically by its graphical model) that outputs a disulfide-bonding probability $p(d_i)$ for each cysteine, based on all SVM predictions.

PSI-BLAST program run on Swiss-Prot and TrEMBL using the BLOSUM62 matrix and an E-value cutoff of 0.005.

Prediction of disulfide bonding state

DISULFIND employs an SVM binary classifier to predict the bonding state of each cysteine, followed by a refinement stage that classifies all the cysteines in a chain in a collective fashion (18), that is, by deciding the overall bonding state assignment of an entire chain rather than making several independent predictions (one for each cysteine). The overall architecture is shown in Figure 1. The SVM receives as input both local and global features [see also (3)]. Local features consist of a window of position specific conservations derived from multiple alignment, centered around the target residue. Global features (amino acid composition, chain length, number of cysteines and average cysteine conservation) provide information about the bonding class of the entire chain (all cysteines bonded, none or mix), which is strongly correlated with the subcellular compartment where the protein resides (reducing versus oxidizing environments).

The refinement stage is motivated by the observation that single cysteines are not independently sampled. Linkage occurs between pairs forming a disulfide bridge but also among sets of cysteines that coordinate a metal ion. A second source of linkage is due to the fact that bonding state is very often a global property of the protein chain and not a local property of individual cysteines (2,3). The effects of correlation are mitigated in two ways. First, we trained a bidirectional recurrent neural network (BRNN) (19) to predict a globally correct sequence of bonding state assignments, given a (possibly incorrect) sequence of locally calculated predictions. At each cysteine position i, the BRNN output is computed using the logistic function and can be therefore interpreted as the conditional probability $p(d_i)$ that the

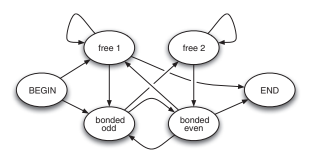


Figure 2. Finite state automaton used in the final stage of bonding state prediction.

cysteine is disulfide-bonded given the input sequence. A position-specific prediction confidence is then defined as

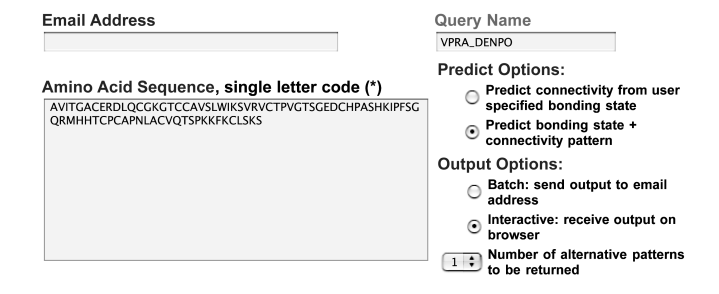
$$c_i = 2(\max\{p(d_i), 1 - p(d_i)\} - 0.5).$$

Second, we enforce the number of bonded cysteines to be even (interchain bridges are ignored) using a finite state automaton (shown in Figure 2). Given the sequence of bonding state probabilities (computed by the BRNN), the most likely sequence of bonding states is obtained by running a Viterbi algorithm. Similar ideas (but using a hidden Markov models rather than an automaton) were presented in (6).

Prediction of disulfide connectivity

We assume in this subsection that disulfide-bonding state of cysteines is given (either entered manually by the user or predicted using the method described above). The method used in DISULFIND is fully detailed in (9) and briefly summarized

A connectivity pattern can be conveniently represented as an undirected graph whose vertices are cysteines and edges



Before submitting your first query please read the following note

Submit Query Reset Fields

Figure 3. Screenshot of the DISULFIND input form.

are disulfide bridges. The problem thus consists of mapping an input sequence with annotated cysteines into an output graphs representing disulfide connectivity. This structured output prediction problem can be cast in the traditional supervised learning setting by introducing a regression problem defined as follows. The input is formed by the annotated sequence and a candidate connectivity pattern. The target is a real valued score, defined as the fraction of correctly assigned bridges. During training the target score is known and we use it to train a recursive neural network in regression mode. Prediction is carried out by running the trained network on all possible connectivity patterns and choosing the one yielding maximum score. The number of possible disulfide patterns connecting 2B cysteines is (2B-1)!! where the double factorial n!! is defined as the product of all odd integers that are less or equal to n.

In order to limit computational efforts, DISULFIND can assign at most five disulfide bridges (in this case the number of candidates to be evaluated is 945). Two remarks are relevant to this limitation. First, chains with more than five bridges are rare (no more than 10% of the Swiss-Prot chains annotated with disulfide bridges). Second, the prediction accuracy is already lowfor chains having five bridges because of a limited number of available training examples; hence prediction of patterns with six or more bridges would be very inaccurate.

IMPLEMENTATION

DISULFINDis available both as a standalone service at http://disulfind.dsi.unifi.it/ and as part of PredictProtein (20).

The current version (DISULFIND 1.1, released in February 2006), incorporates some improvements in the presentation interface.

Interface

The input to the predictor is entered via an HTTP form using the SEND method. The main fields (see Figure 3) are the following.

Email address The address were results will be sent if the email output option is selected.

Query name An optional field that allows to label the sequence with an user provided identifier.

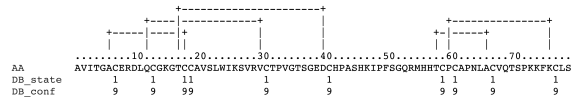
Amino acid sequence The protein sequence using standard amino acid one-letter codes. Spaces and newlines are automatically stripped.

Predict options In its normal behavior, DISULFIND predicts both bonding state and connectivity. If the bonding state is known in advance, users may check the corresponding option in the user interface and after the form is submitted they will be presented a screen where the bonding state of each cysteine can be manually assigned. In this case only predicted connectivity will be returned.

Output options There are two possible output operation modes. In email mode, after the form is submitted, a job is scheduled in the server and results are returned in ASCII format to the indicated email address. In browser mode, results are returned to the HTTP client (see Figure 4).

Alternatives By default DISULFIND only returns the most likely connectivity pattern. By setting the number of alternatives to an integer k in the range (1,3), the k best ranking patterns will be returned.

Results for VPRA DENPO



```
80
DB_state
DB_conf
DB bond
          bond(7,19)
DB bond
          bond(13,31)
DB bond
          bond(18,41)
          bond(59,67)
DB_bond
DB bond
          bond(61,77)
```

Conn_conf 0.441431

Figure 4. DISULFIND output.

The output presented to the user consists of the original sequence annotated with predictions as shown in Figure 4. The following items are returned in the output screen:

AA The original amino acid sequence;

DB state Predicted disulfide bonding state (1 = disulfide)bonded, 0 = not disulfide bonded;

DB conf Confidence of disulfide bonding state prediction (0 = low to 9 = high); a red color means that the Viterbi aligner overruled the SVM predition for that residue in order to achieve a consistent prediction at the chain level (i.e. an even number of disulfide bonded cysteines, as interchain bonds are ignored);

Conn conf Confidence of connectivity assignment given the predicted disulfide bonding state. The confidence in this case is the predicted score associated with the connectivity pattern, i.e. the fraction of correctly assigned bridges—see details in (9). Although the score is a number in (0,1), it should not be confused with the probability that the pattern is correct.

The above output is repeated if multiple alternative patterns are requested. Since all the alternatives share the same bonding state prediction, fields DB state and DB conf are only shown in the output presentation of the most likely pattern.

Performance

Under regular load conditions, a query can be answered in about 30-60 s. CPU time depends on the sequence length and the number of disulfide bridges. Most CPU time is used by PSI-BLAST for calculation of multiple alignment profiles. The 20-fold cross validation performance of the bonding state prediction stage is reported in Table 1. In order to assess the significance of the confidence score (see Equation 1), we report in Figure 5 the accuracy and rejection rate of the

Table 1. DISULFIND bonding state predictor: experimental results on a 20-fold cross validation procedure (PDB Select July 2005)

Method	Q_2	Q_p
Loc29	86 ± 1	73 ± 2
BRNN Loc29+f	88 ± 1	82 ± 2
BRNN Loc29+f FSA	88 ± 1	83 ± 2

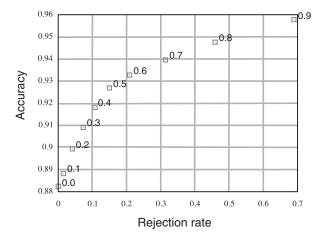


Figure 5. Accuracy versus rejection rate of the abstaining bonding state predictor for different confidence cutoff values (the rejection rate is the fraction of cysteines that are predicted at a confidence level below the cutoff value shown at the right of each point in the curve).

bonding state classifier that abstains when the confidence is lower or equal to a given cutoff. It can be seen, for example, that accuracy improves to $Q_2 = 92.7\%$ at a rejection rate of 15.0% for a confidence cutoff of 0.5.

Table 2. Leave-one-out validation results of disulphide connectity prediction

Number of bridges	Number of chains	Q_P	Q_c
2	156	75.0	75.0
3	146	46.6	55.7
4	99	50.5	63.4
5	45	17.8	42.7
All	446	54.5	60.2

Concerning disulfide connectivity, leave-one-out estimates of prediction accuracy on a set of 446 Swiss-Prot Sequences (9) are reported in Table 2 [note that results reported in (9) were based on a 4-fold cross validation]. Q_p is the fraction of correctly assigned patterns, while Q_c is the fraction of correctly predicted bridges. If multiple alternative are selected, the probability that a correct pattern is included increases. Results obtained considering the top k = 3 configurations are $Q_p = 66.3$, $Q_c = 69.5$.

Statistics

DISULFIND has served a total of over 7000 tasks from almost 50 national domains since April 2003 and is currently serving an average of 60 queries per week. Hundreds of queries per month have been served via PredictProtein since July 2004.

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Conflict of interest statement. None declared.

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