Prediction of RNA- and DNA-Binding Proteins Using Various Machine Learning Classifiers

Mehdi Poursheikhali Asghari and Parviz Abdolmaleki *

Department of Biophysics, Faculty of Biological Sciences, Tarbiat Modares University, Tehran, Iran

Abstract

Background: Nucleic acid-binding proteins play major roles in different biological processes, such as transcription, splicing and translation. Therefore, the nucleic acid-binding function prediction of proteins is a step toward full functional annotation of proteins. The aim of our research was the improvement of nucleic-acid binding function prediction.

Methods: In the current study, nine machine-learning algorithms were used to predict RNA- and DNA-binding proteins and also to discriminate between RNA-binding proteins and DNA-binding proteins. The electrostatic features were utilized for prediction of each function in corresponding adapted protein datasets. The leave-one-out cross-validation process was used to measure the performance of employed classifiers.

Results: Radial basis function classifier gave the best results in predicting RNA- and DNA-binding proteins in comparison with other classifiers applied. In discriminating between RNA- and DNA-binding proteins, multilayer perceptron classifier was the best one.

Conclusion: Our findings show that the prediction of nucleic acid-binding function based on these simple electrostatic features can be improved by applied classifiers. Moreover, a reasonable progress to distinguish between RNA- and DNA-binding proteins has been achieved.

Keywords: DNA-binding proteins, Machine-learning algorithms, RNA-binding proteins

Introduction

Protein-RNA interactions play a fundamental role in different bioprocesses, such as transcription, splicing and translation. RNA-Binding Proteins (RBPs) are pivotal ingredients of RNA splicing, export, stability, localization and translation. They organize all aspects of RNA biogenesis from maturation, surveillance, nucleocytoplasmic transfer to subcellular localization, translation and decomposition. Therefore, a full comprehension of a diverse variety of cellular processes requires the identification of RBPs.

So far, various computational methods have been developed for the identification of RBPs. Some of them have been based on sequence-derived features such as amino acid composition, dipeptide composition, composition-transition-distribution of seven physicochemical properties, evolutionary information in terms of position-specific scoring matrices and functional domain composition. The majority of sequence-based methods used the Support Vector Machine (SVM) algorithm for identifying RBPs. On the other hand, alternative methods have utilized the electrostatic features of the protein surface patches in order to identify RBPs. Similar to RBPs, DNA-Binding Proteins (DBPs) have been predicted by using two different approaches. In the structure-based approach, the structural alignment and threading-based methods as well as the electrostatic features of the surface patches of proteins have been utilized for identifying DBPs. The electrostatic features have been used in this approach since large positively charged surface patches of proteins usually participate in interaction with DNA molecule. The second approach is the prediction of DBPs based on the sequence information. Numerous machine-learning algorithms have been built based on different encoding schemes of the protein sequence to predict DBPs, in this approach.

Machine-learning algorithms are extensively used to predict the structure and function of proteins. Actual-
ly, machine learning presents one of the most robust approaches to constructing predictive models in settings where experimentally validated training data are available. At present, however, it is unclear whether the available experimental data regarding DNA-protein and RNA-protein interactions are sufficient for successfully training classifiers using machine learning algorithms. Against this fact, this study applies machine learning approaches to train electrostatic-based classifiers for predicting DBPs and RBPs.

This study was done using simple electrostatic features including charge, dipole and quadrupole moments for predicting RNA- and DNA-binding proteins by means of neural network method. Electrostatic interactions are among the most significant indicators to be considered when one will determine the function of proteins. It is now generally recognized that one must analyze the electrostatic forces in a protein to understand its function.

Here, this electrostatic-based approach was extended by applying 9 various machine learning classifiers for identifying RNA- and DNA-binding proteins and also for discriminating RBPs from DBPs. Our goal was to improve the classification accuracy in each of the comparisons. The results demonstrate that this approach can be used by other researchers in this field for more accurate nucleic acid-binding function prediction.

Materials and Methods

Datasets

The protein datasets of Ahmad and Sarai were used for our analysis. These datasets consisted of 160 RBP chains, 143 DBP chains and 2441 non nucleic acid-binding protein (Ctrl) chains.

Electrostatic features

The extracted electrostatic features of Ahmad and Sarai work were used in the current study. These features included the charge, dipole and quadrupole moments of protein chains. The detailed description of these features can be found at Ahmad S et al.

Machine learning algorithms

In this study, nine classification algorithms of Alternating Decision Tree (ADTree), K-nearest neighbor (K-NN), L1 Regularized Logistic Regression (L1 RLR), L2 Regularized Logistic Regression (L2 RLR), Multi-layer Perceptron classifier (MLPClassifier), Random Forest (RF), Radial Basis Function classifier (RBF-Classifier), RealAdaBoost and Sequential Minimal Optimization (SMO) algorithm were used to predict the nucleic acid-binding function of proteins (i.e., RNA- and DNA-binding) and also to differentiate between RBPs and DBPs. Waikato Environment for Knowledge Analysis (WEKA) package version 3.7.10, an ensemble of machine learning algorithms, was used to perform classifying tasks. Below, there is a brief explanation of nine employed classifiers and their corresponding parameter values:

ADTree: It generates an alternating decision tree. 20 boosting iterations were used in our experiments. All other parameters of the algorithm were set to default. Our classification was done with ADTree function of WEKA.

K-NN: It is a standard non-parametric classification method. The basic idea of the K-NN method is that a new case will be classified as the most frequent class among its K-Nearest Neighbors. K-NN was used with Euclidean distance and distance weighting (1/distance) and also 75 values of K ranging from K=1 to K=75 were examined and the best ones were selected in terms of the area under receiver operating characteristic curve (AUC) measure. All other parameters of the algorithm were set to default. IBk function of WEKA was used for classification.

L1 RLR and L2 RLR: Logistic regression is a well-established method for the classification or prediction of binary response function based on the various independent features. The regression utilizes an objective function and the number of its parameters is as large as the number of features. Usually the objective function contains also a regularization term. It penalizes model details of unnecessary complexity, focuses on the most concerned features, and thus avoids over-fitting of the data used for training (parameter optimization). The most common variants of regularization methods are L1 regularization, also known as Lasso and L2 regularization also known as ridge regression. As penalty term, the L1 regularization adds the sum of the absolute values of the model parameters to the objective function, whereas the L2 regularization adds the sum of squares of parameters.

LibLINEAR function of WEKA was utilized for doing prediction with these algorithms. The parameter cost (c) was changed from value 1 to 10 and the best one for our classifications was selected. The features were normalized and probability estimates for classification problems were generated. All other parameters of the algorithm were set to default.

MLPClassifier: It trains a multilayer perceptron with one hidden layer. MLPClassifier function was used and the number of units was changed in the hidden layer and the best architecture for our classifications was selected. All other parameters of the algorithm were set to default.

RF: It is an ensemble classifier method based on decision trees. After a large number of trees are generated, each tree in the forest gives a vote for a class and the most popular class among trees for a test instance presents the final classification. A few parameters influence the performance of RF models, such as the number of trees in the forest (ntree) and the number of variables considered at each split (mtry). RandomForest function was used in this study and 500 trees were grown in each experiment. For the number of variables randomly selected at each node, the default value that was equal to the square root of the feature...
dimension was used. Random forests were trained with a maximum depth of 30 trees.

RBFClassifier: It implements radial basis function networks for classification, trained in a fully supervised manner. RBFClassifier function was utilized and different numbers of base function were examined and the best one for classification was selected. All other parameters of the algorithm were set to default.

Real AdaBoost: It boosts a two-class classifier using the Real AdaBoost method. The default parameter values of RealAdaBoost function were used for our classifications.

SMO: It is an implementation of SVM algorithm that globally replaces all missing values and transforms nominal attributes into binary ones. It also normalizes all attributes by default. SMO is conceptually simple, easy to implement and faster in computation. Fitting logistic regression models to the outputs of the SMO could in addition provide probability estimates.

SMO function was used and linear, polynomial (of degree 2) and radial basis function kernels in our classifications were examined and the best ones were selected. The cost factor c was appropriately chosen during the training time. All other parameters of the SMO algorithm were set to default.

Leave-One-Out Cross-Validation (LOOCV)

The performance of our models trained on Ahmad and Sarai datasets was assessed using LOOCV. In this procedure, one sample was taken out of the whole dataset and was used as the test instance, and the remaining samples were used as training instances. Then, the prediction was made for the test sample. This process was repeated n times (n=total number of samples), and the final performance results were obtained by averaging over all the test results. The sample can be a protein sequence, a protein chain, a DNA sequence, and so on. In this study, the sample was the protein chain. There is not any biological reason for picking this strategy. This is a statistical procedure.

Performance measures

Various performance measures were used to evaluate the results. These include accuracy, precision, recall, f-measure and the area under Receiver Operating Characteristic (ROC) curve (known as AUC). Accuracy shows proximity of measurement of results to the true value. It can be calculated as $\frac{TP}{TP+FN}$, where TP refers to true and F refers to false, whereas P is positive class and N is negative class. Recall $\frac{TP}{TP+FN}$ relates to the classifier's ability to identify positive instances while precision $\frac{TP}{TP+FP}$ is the fraction of predicted instances as positive class that is correctly predicted. F-measure $\frac{(2*precision*recall)}{(precision+recall)}$ is the geometric mean of precision and recall. Also, models based on the ROC curve were evaluated which plot the true positive rate against false positive rate. The AUC value reported by an ROC curve is equal to the probability that a classifier will rank a randomly chosen positive instance higher than a randomly chosen negative one. The AUC is a standard non threshold-dependent index for performance evaluation.

ROC curves

For better comparison of applied machine-learning algorithms on 3 datasets, their corresponding ROC cur-
Comparison with other methods

The results of this research are comparable with the results of Ahmad and Sarai because their protein datasets and also their extracted features were used in this study. The difference between their work and the current study is utilizing nine machine-learning algorithms other than the neural network they employed.

Obtained results in this study show improvement in the performance measures, especially in the AUC measure, when comparing with their work. Ahmad and Sarai reached to the AUC value of 0.78 for RBPs versus ctrl dataset, while this measure improved by 0.07 by means of RBFClassifier algorithm in the current study. Likewise, the accuracy was improved by 0.03 and reached to the 0.94. Another algorithm which improved these measures was K-NN which raised the AUC and accuracy values up to 0.84 and 0.94, respectively. Among 9 machine-learning algorithms employ-
RNA-and DNA-Binding Proteins Prediction

Discussion

In this study, 9 machine-learning algorithms were used for discrimination between RBPs and ctrl chains, DBPs and ctrl chains and finally between RBPs and DBPs. The obtained results demonstrated that our selected classification algorithms can further improve predictions constructed with only 5 electrostatic features obtained from low-resolution protein structures. Then, these features show the capability for nucleic acid-binding function prediction and also for discriminating RNA-binding from DNA-binding function. Adding more informative features to the prediction process can improve the performance measures and hence increase the classification accuracy.

Two robust classifiers in 3 different comparisons were RBFClassifier and MLPClassifier. These two classifiers have architectures similar to neural network. Therefore, similar to Ahmad and Sarai work, neural network methods show significant robustness for prediction of nucleic acid-binding function with the help of electrostatic features.

The importance of this work is to support this idea that simple electrostatic features such as charge, dipole and quadruple moments are useful for identification of nucleic acid-binding function. These electrostatic features can be combined with other sequence or structure-based features for precise function prediction at the level of nucleic acid-binding. Also, the discrimination between RNA- and DNA-binding function is reinforced by these features as well as other suitable ones.

Given the relatively small sizes of the nucleic acid-binding proteins analyzed in this study, discrepancies in the results obtained using different classifiers to predict nucleic acid-binding proteins must be interpreted with caution. It will be important to evaluate these methods on larger, more complete datasets of experimentally validated nucleic acid-binding proteins as they become available.

Notes for practical implementation

The overall workflow for practical implementation of our method is shown in figure 4. Given the protein structure as the input, five electrostatic features are extracted from the query protein. Then, the RBFClassifier as the most powerful method in discriminating between RNA/DNA-binding and non-nucleic acid-binding proteins uses its trained structure to determine whether the query is a RNA/DNA-binding protein or not. If the protein function was predicted as nucleic acid-binding (or RNA/DNA-binding), then the classification continues to distinguish RNA-binding function from DNA-binding utilizing the trained structure of MLPClassifier, i.e. the best-selected classifier trained on the nucleic acid-binding proteins dataset.

Conclusion

Nine different machine-learning algorithms other than neural network with a higher capability for nucleic acid-binding function prediction have been introduced. These classifiers showed reasonable improvement that highlights their potential to be used by other researchers for nucleic acid-binding function prediction. It is hoped that the use of alternative sequence or structure-based features as well as the electrostatic features will reinforce nucleic acid-binding function prediction protocols.
Acknowledgement

We would like to thank Dr Ebrahim Barzegari Asadabadi for his useful comments.

References

15. Ma X, Guo J, Sun X. Sequence-based prediction of RNA-binding proteins using random forest with mini-
RNA-and DNA-Binding Proteins Prediction

110

Avicenna Journal of Medical Biotechnology, Vol. 11, No. 1, January-March 2019


