## ORIGINAL ARTICLE

# An ensemble of reduced alphabets with protein encoding based on grouped weight for predicting DNA-binding proteins

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**Abstract** It is well known in the literature that an ensemble of classifiers obtains good performance with respect to that obtained by a stand-alone method. Hence, it is very important to develop ensemble methods well suited for bioinformatics data. In this work, we propose to combine the feature extraction method based on grouped weight with a set of amino-acid alphabets obtained by a Genetic Algorithm. The proposed method is applied for predicting DNA-binding proteins. As classifiers, the linear support vector machine and the radial basis function support vector machine are tested. As performance indicators, the accuracy and Matthews's correlation coefficient are reported. Matthews's correlation coefficient obtained by our ensemble method is  $\approx 0.97$  when the jackknife cross-validation is used. This result outperforms the performance obtained in the literature using the same dataset where the features are extracted directly from the amino-acid sequence.

**Keywords** Multi-classifier · Amino-acid alphabets · Support vector machine · DNA-binding proteins · Ensemble classifier

#### Introduction

In this paper, we deal with the DNA-binding proteins (Ahmad and Sarai 2005; Hwang et al. 2007; Kuznetsov et al. 2007; Lander et al. 2001; Ofran et al. 2007; Wang and Brown 2006; Yan et al. 2006) prediction problem proposing an ensemble (Nanni and Lumini 2006) of support vector machines (Cristianini and Shawe-Taylor 2000). Our results

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are very interesting; the fusion outperforms the best-published results on the same dataset when the features are extracted directly from the amino-acid sequence. We present a new multi-classifier construction based on the combination between the encoding method based on grouped weight (Zhang et al. 2006c) and a set of reduced alphabets obtained by a Genetic Algorithm (Nanni and Lumini 2008a).

Notice that the encoding method based on grouped weight obtain performance lower than that obtained by the pseudo amino acid composition, but when the encoding method based on grouped weight is combined with a set of reduced alphabets, the performance drastically increases.

In the literature, it is well known that the fusion among classifiers (Chou 2000) is a feasible method for improving the performance of a stand-alone method. Several ensemble methods are published in the bioinformatics literature: protein secondary structure prediction (Riis and Krogh, 1996); protein fold pattern prediction (Shen and Chou 2006); protein subcellular localization prediction (Shen and Chou 2006, 2007a, b, c, d); membrane protein type prediction (Chou and Shen 2007c); signal peptide prediction (Chou and Shen 2007e); peptide classification (Nanni and Lumini 2006a, b). Recently, based on the approach by fusing many individual classifiers, a powerful web-server package, called Cell-PLoc, was established (Chou and Shen 2008) for predicting the subcellular localization of proteins in various different organisms.

To extract features directly from the amino-acid sequence, the most known method is the pseudo amino acid composition. A pseudo amino acid composition generator (Shen and Chou e) was established at the website http://chou.med.harvard.edu/bioinf/PseAA/, originally introduced by Chou for protein subcellular localization (Chou 2001) and for enzyme functional class (Chou 2005). In the last year, a huge number of pseudo amino acid composition approaches have been proposed (Aguero-Chapin et al. 2006;



Caballero et al. 2007; Cai and Chou 2006; Chen et al. 2006a; Chen et al. 2006b; Chen and Li 2007a, b; Chou and Shen 2006a, b, 2007a, b, c, e; Diao et al. 2007; Ding et al. 2007; Du and Li 2006; Fang et al. 2007; Gao et al. 2005; Gonzalez-Diaz et al. 2006, 2007a, b, c; Kurgan et al. 2007; Li and Li 2007; Lin and Li 2007a, b; Liu et al. 2005a, b; Mondal et al. 2006; Mundra et al. 2007; Pan et al. 2003; Pu et al. 2007; Shen and Chou 2005a, b, 2006, 2007a, b, c, d, f, g, h; Shen et al. 2006, 2007; Shi et al. 2007; Wang et al. 2004, 2006; Xiao and Chou 2007; Xiao et al. 2006a, b; Zhang and Ding 2007; Zhang et al. 2006a, b, 2007; Zhou et al. 2007).

The encoding method based on grouped weight (Zhang et al. 2006c) is based on the division of the 20 amino acid residues into four different classes; in the original paper, this division is performed considering the hydrophobicity and charged character, and then to extract a set of features from different subsequences of each protein. To improve the method, we apply the method for building reduced alphabets proposed in Nanni and Lumini (2008b) for peptide classification. A different support vector machine (Cristianini and Shawe-Taylor 2000) is trained on each feature set (each extracted from a different alphabet). Finally, this pool of classifiers is combined by the vote rule (Duda et al. 2000).

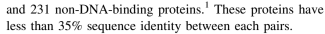
Notice that we use the same reduced alphabets proposed in Nanni and Lumini (2008b); these reduced alphabets are obtained to maximize the area under the receiver operating characteristic curve in two problems: HIV-protease, and recognition of T-cell epitopes. In these two datasets, the ensemble of classifiers obtains an average area under the receiver operating characteristic curve of 0.984.

Notice that the proteins of the problem of this paper are not considered for obtaining the reduced alphabets.

Several methods are proposed in the literature to deal with the DNA-binding proteins: Jones et al. (2003), Shanahan et al. (2004), Keil et al. (2004), and Tsuchiya et al. 2004) propose systems based on chemical or physical properties of amino acids; Ahmad and Sarai (2005), and Ahmad et al. (2004) propose systems based on neural network classifiers; Kuznetsov et al. (2006) propose a system based on structure information and sequence alignment profiles; Bhardwaj et al. (2005) propose a system based on electrostatic potentials and the amino acid composition of the protein; Fang et al. (2007) propose a system based on the Chou's pseudo-amino acid composition; and Nanni and Lumini (2008b) propose to combine the dipeptide composition of the protein with the features extracted from the gene ontology database (Chou and Shen 2007a, d).

## Materials and methods

We have used exactly the same datasets used in Fang et al. (2007). This dataset contains 118 DNA-binding proteins



The proposed method combined 11 classifiers where each classifier is trained using the encoding method based on grouped weight combined with a different reduced alphabet. Finally, these classifiers are combined by vote rule (Kittler et al. 1998). Notice that our ensemble is built by 11 classifiers since we have used K = 5 (as in Nanni and Lumini 2008b) different alphabets for each value of the size S = 4 of the reduced alphabets and for a given value of N (N = 1 or N = 2). Moreover, the baseline implementation based on the alphabets of Zhang et al. (2006c) also belongs to the ensemble.

In this work, an alternative way for the construction of reduced alphabets is applied; it is based on a Genetic Algorithm for grouping amino-acids (Nanni and Lumini 2008b). The objective function of the Genetic Algorithm is the maximization of the area under the receiver operating characteristic curve (Fawcett 2004) for a given classification problem using the N-gram feature extraction. K (K = 5) different alphabets are created for each value of the size S of the reduced alphabets and for a given value of N. The ith reduced alphabet is built considering the previous reduced alphabets of the same size S and of the same value of N. Simply, for the calculation of the objective function of the ith iteration of GA, the scores obtained by the ith reduced alphabet are combined by the mean rule with the scores obtained by the previous i-1 reduced alphabets. The mean rule (Eq. 1) selects as final score [ $score(\mathbf{s},c)$ ] the mean score of a pool of K classifiers:

$$score(\mathbf{s}, c) = \frac{1}{K} \sum_{i=1...K} sim_j(\mathbf{s}, c)$$
 (1)

Where:  $sim_j(s,c)$  is the similarity of the pattern (protein) **s** to the class c, obtained by the jth classifier; K is the number of combined classifier.

In this work, we use the ten alphabets obtained with: S = 4, N = 1; S = 4, N = 2. The area under the receiver operating characteristic curve, as in Nanni and Lumini (2008b), is maximized for two problems of peptide classification: HIV-protease; and recognition of T-cell epitopes.

The reduce alphabets are:

```
C1 = 'GPSV'; C2 = 'ACDFNW'; C3 = 'EKR';

C4 = 'HILMQTY'

C1 = 'QRSW'; C2 = 'AFHIKLMPV'; C3 = 'DENTY';

C4 = 'CG'

C1 = 'MNRTW'; C2 = 'ACHILPV'; C3 = 'DEFQ';

C4 = 'GKSY'

C1 = []; C2 = 'FHMNRSVWY'; C3 = 'ACEGIKLP';

C4 = 'DOT'
```



<sup>&</sup>lt;sup>1</sup> 107 DNA-binding proteins and 196 non-DNA-binding proteins have a hit in the GO database.

```
C1 = 'CDES'; C2 = 'AGHQTY'; C3 = 'FKVW';

C4 = 'ILMNPR'

C1 = 'HKR'; C2 = 'AGIMNQVWY';

C3 = 'CDEFPST'; C4 = 'L'

C1 = 'CMPS'; C2 = 'ADEFGHKLTVY'; C3 = 'IQ';

C4 = 'NRW'

C1 = 'AIPRSV'; C2 = 'CGHLNQW'; C3 = 'DEY';

C4 = 'FKMT'

C1 = 'FGHL'; C2 = 'ADKNPW';

C3 = 'CEIMQRSTVY'; C4 = '[]'

C1 = 'AKMWY'; C2 = 'FHILRST'; C3 = 'CDNQ';

C4 = 'EGPV'
```

Now, we describe the method used in this paper for the encoding method based on grouped weight; the original paper divided the 20 amino acid residues into four different classes (Zhang et al. 2006c): C1 = `GAVLIMPFW'; C2 = `QNSTYC'; C3 = `DE'; C4 = `HKR'. Since the amino acid residues are divided into four different classes, we can partition the amino acid residues into the following disjoint groups: C1 + C2 versus C3 + C4, or C1 + C3 versus C2 + C4, and C1 + C4 versus C2 + C3.

Given a protein  $\mathbf{p}$  we calculate three binary sequences (Eq. 2):

$$\begin{aligned} \mathbf{H1}_{\mathbf{p}}(j) &= 1 \text{ if } \mathbf{p}(j) \in C1 + C2 \text{ else } \mathbf{p}(j) = 0 \\ \mathbf{H2}_{\mathbf{p}}(j) &= 1 \text{ if } \mathbf{p}(j) \in C1 + C3 \text{ else } \mathbf{p}(j) = 0 \\ \mathbf{H3}_{\mathbf{p}}(j) &= 1 \text{ if } \mathbf{p}(j) \in C1 + C4 \text{ else } \mathbf{p}(j) = 0 \end{aligned} \tag{2}$$

We partition each binary sequence into L pieces of subsequence. In our implementation, the jth feature (j = 1, ..., L) is given by sum $(\mathbf{H1}_p(1:j \times S) = 1)/(j \times S)$ , where S = length of the protein/L and the function sum $(\mathbf{x})$  gives the number of 1 in the vector  $\mathbf{x}$ . We create L features for  $\mathbf{H1}$ ,  $\mathbf{H2}$ ,  $\mathbf{H3}$  and then we concatenate these three vectors.

For building the ensemble, we simply use the values of C1, C2, C3, and C4 obtained by the Genetic Algorithm.

Notice that to avoid any doubt in the implementation of the feature extraction method, we report the Matlab code in the "Appendix".

## Results and discussion

As performance indicators, we use Matthew's correlation coefficient (MCC) (Fang et al. 2007) and the accuracy (see Eq. 3).

where TP(i) is the number of correctly predicted DNA-binding proteins (true positives); TN(i), FP(i) and FN(i) are the numbers of true negatives, false positives, and false negatives, respectively.

As testing protocol, the rigorous jackknife test (Feng 2002, Chou and Zhang 1995) is used since it is demonstrated (Chou and Shen 2007d) that it is the best method for comparing different classifiers in a given dataset. For this reason, the jackknife test is widely adopted in the literature (Chen et al. 2006a, b, 2007; Chou and Shen 2006a; Diao et al. 2008; Ding et al. 2007; Du and Li 2006; Fang et al. 2007; Gao et al. 2005; Guo et al. 2006; Kedarisetti et al. 2006; Li and Li 2007; Lin and Li 2007a, b; Liu et al. 2007; Mondal et al. 2006; Niu et al. 2006; Shen and Chou 2007g; Shen et al. 2007; Wang et al. 2007; Sun and Huang 2006; Tan et al. 2007; Wang et al. 2005; Wen et al. 2006; Xiao and Chou 2007; Xiao et al. 2005a, b, 2006a; Zhang and Ding 2007; Zhang et al. 2006a, 2007; Zhou 1998; Zhou and Doctor 2003; Zhou et al. 2007).

We have used three different classifiers: linear support vector machine (LSVM); radial basis function support vector machine with C=1 and Gamma = 1 (RSVM1); radial basis function support vector machine with C=0.1 and Gamma = 1,000 (RSVM2). In Fig. 1, we report the performance of these three classifiers, using only the baseline encoding method based on grouped weight method, varying the value of L. In Fig. 1, we also report the performance of our ensemble (ENS) of radial basis function support vector machine<sup>2</sup> (with cost of the constrain violation = 0.1 and parameters of the radial based kernel = 1,000) classifiers. It is clear that the proposed method outperforms the stand-alone methods.

In Table 1, we report the performance of each standalone classifiers that build the ensemble (L = 10).

In Table 2, we report the MCC of other state-of-the-art methods, from Nanni and Lumini (2008a) and Fang et al. (2007), for predicting DNA-binding proteins.

From the analysis of the experimental results, the following observations may be made:

- 1. The encoding method based on grouped weight permits to obtain a reliable method.
- The proposed ensemble permits to improve the performance of the baseline encoding method based on grouped weight; the proposed ensemble of support vector machines obtains a Matthew's correlation

$$MCC(i) = \frac{TP(i) \times TN(i) - FP(i) \times FN(i)}{\sqrt{(TP(i) + FP(i)) \times (TP(i) + FN(i)) \times (TN(i) + FN(i)) \times (TN(i) + FP(i))}}$$
(3)

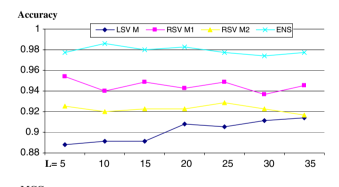
<sup>&</sup>lt;sup>2</sup> Implemented as in the OSU svm matlab toolbox

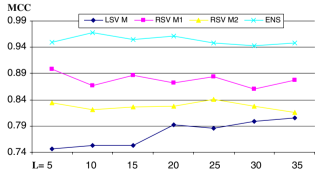
- coefficient of 0.968 when it is trained by 11 different set of reduced alphabets.
- The tested fusion permits to outperform several other state-of-the-art methods.
- 4. Our results show that Chou's pseudo amino-acid composition outperforms the encoding method based on grouped weight, but we show that the performance of the encoding method based on grouped weight increases if our method to build an ensemble of classifiers is used.

### **Conclusions**

We investigated how to build an ensemble of classifiers for the encoding method based on grouped weight. For building the ensemble, we used the same reduced alphabets proposed in Nanni and Lumini (2008b); these reduced alphabets are obtained to maximize the area under the receiver operating characteristic curve in two problems: HIV-protease, and recognition of T-cell epitopes.

The proposed ensemble of 11 support vector machines obtains a Matthew's correlation coefficient of 0.968;





 $\label{eq:Fig. 1} \textbf{Fig. 1} \ \ \textbf{Accuracy} \ \ \textbf{and} \ \ \textbf{MCC} \ \ \textbf{of three stand-alone} \ \ \textbf{methods} \ \ \textbf{and of the} \\ \textbf{proposed ensemble}$ 

Table 1 Performance of the classifiers that build the ensemble

0.8674
0.8972
0.9101
0.8733
0.872
0.7096
0.8592
0.886
0.8512
0.8568
0.8793

<sup>&</sup>lt;sup>a</sup> The set used in Zhang et al. (2006c)

**Table 2** Performance, in the same dataset, of other state-of-the-art methods for predicting DNA-binding proteins

Features	Classifier	MCC
Onthology-based	Stand-alone LSVM	0.928
Onthology-based	Random subspace LSVM	0.935
2-gram	Stand-alone LSVM	0.848
2-gram	Random subspace	0.942
Chou's pseudo amino-acid composition		0.924
Proposed ensemble		0.968

moreover, the validity of the novel approach is proved by the performance improvements obtained with respect to other state-of-the-art methods in the tested problem.

As a future study, we want to combine methods trained using different feature extraction methods, e.g., we can combine an ensemble builds using the onthology-based features, an ensemble based on the 2-gram composition, an ensemble based on the Chou's pseudo amino-acid composition, and the ensemble proposed in this work.

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## Appendix: Matlab code

The following function implements the base feature extraction method as detailed in "Materials and methods":



```
function F=featuresEBGW(ALFABETO,P,L)
C1=ALFABETO{1};
C2=ALFABETO{2};
C3=ALFABETO{3};
C4=ALFABETO{4};
uno=strcat(C1,C2);
due=strcat(C1,C3);
tre=strcat(C1,C4);
H=[];
for j=1:length(P)
  if sum(P(j)==uno)
H(j)=1;
  else
    H(j)=0;
  end
end
S=length(P)/L;
for j=1:L
  F(t)=sum(H(1:j*S)==1)/(j*S);
  t=t+1;
end
H=[];
for j=1:length(P)
if sum(P(j)==due)
    H(j)=1;
  else
    H(j)=0;
  end
end
for j=1:L
  F(t)=sum(H(1:j*S)==1)/(j*S);
  t=t+1;
end
H=[];
for j=1:length(P)
  if sum(P(j)==tre)
    H(j)=1;
  else
    H(j)=0;
  end
end
for j=1:L
  F(t)=sum(H(1:j*S)==1)/(j*S);
  t=t+1;
end
```



## The following rows implements the main of the system:

```
ALFABETO{1}='GAVLIMPFW';
ALFABETO{2}='QNSTYC';
ALFABETO(3)='DE':
ALFABETO{4}='HKR'
load c:\AlfabetiUniti.mat Alfabeto
% Alfabeto stores the reduced sets obtained in (Nanni and Lumini, 2008a)
       if NK==1
         ALF=ALFABETO;
       elseif NK>1 & NK<7
         ALF=Alfabeto{3}{NK-1};
       else
         ALF=Alfabeto{4}{NK-6};
       end
       F=[];
       for i=1:size(SEQ,2)
         F(i,:)=featuresEBGW(ALF,SEQ{i},L);
       res=F·
       for op=1:size(SEQ,2)
         TR = res; yTR = y; TR(op,:) = []; yTR(op) = [];
         TE=res(op,:);
         massimo=max(TR)+0.00001;
         minimo=min(TR);
         training=[];
         testing=[];
         training=TR;
         testing=TE;
         for i=1:size(TR,2)
           training (1: size (TR, 1), i) = double (TR (1: size (TR, 1), i) - minimo(i)) / (massimo(i) - minimo(i)); \\
           testing(1:size(TE,1),i)=double(TE(1:size(TE,1),i)-minimo(i))/(massimo(i)-minimo(i));
         end
         tra=[];
         TR=training;
         TE=testing
         [AlphaY, SVs, Bias, Parameters, nSV, nLabel] = rbfSVC(double(TR)', double(yTR),0.1,1000);
         [ER, Decision(op), Ns, ConfMatrix, lab(op)]=SVMTest(double(TE)', double(y(op)), AlphaY, SVs, Bias, Parameters, nSV,
nLabel);
       end
       VALUE(1)=sum(lab==y);
       %calcoloM MCC
       for i=1:3
         VALUE(i+1)=CalcoloMCC(lab,y,i)
       end
       labP(tt,:)=lab;
      tt=tt+1
    end
end
% vote rule
Q=[];
for i=1:size(SEQ,2)
  for j=1:max(y)
      Q(i,j)=sum(labP(:,i)==j);
   end
end
[a,b]=max(Q');
```

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