Regularized maximum correntropy machine

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Abstract

In this paper we investigate the usage of regularized correntropy framework for learning of classifiers from noisy labels. The class label predictors learned by minimizing transitional loss functions are sensitive to the noisy and outlying labels of training samples, because the transitional loss functions are equally applied to all the samples. To solve this problem, we propose to learn the class label predictors by maximizing the correntropy between the predicted labels and the true labels of the training samples, under the regularized Maximum Correntropy Criteria (MCC) framework. Moreover, we regularize the predictor parameter to control the complexity of the predictor. The learning problem is formulated by an objective function considering the parameter regularization and MCC simultaneously. By optimizing the objective function alternately, we develop a novel predictor learning algorithm. The experiments on two challenging pattern classification tasks show that it significantly outperforms the machines with transitional loss functions.

Keywords: Pattern classification, Label noise, Maximum Correntropy criteria, Regularization

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1. Introduction

The classification machine design has been a basic problem in the pattern recognition field. It tries to learn an effective predictor to map the feature vector of a sample to its class label [1, 2, 3, 4, 5, 6, 7, 8, 9]. We study the supervised multi-class learning problem with $L$ classes. Suppose we have a training set denoted as $D = \{(x_i, y_i)\}, i = 1, \cdots, N$, where $x_i = [x_{i1}, \cdots, x_{iD}]^T \in \mathbb{R}^D$ is the $D$ dimensional feature vector of the $i$-th training sample, and $y_i \in \{1, \cdots, L\}$ is the class label of $i$-th training sample. Moreover, we also denote the label indicator matrix as $Y = [Y_{li}] \in \mathbb{R}^{L \times N}$, and $Y_{li} = 1$ if $y_i = l$, and $-1$ otherwise. We try to learn $L$ class label predictors $\{f_l(x)\}, l = 1, \cdots, L$ for the multi-class learning problem, where $f_l(x)$ is the predictor for the $l$-th class and $\theta$ is its parameter. Given a sample $x_i$, the output of the $l$-th predictor is denoted as $f_l(x_i)$, and we further denote the prediction result matrix as $F_\theta = [F_{\theta li}] \in \mathbb{R}^{L \times N}$, and $F_{\theta li} = f_l(x_i)$. To make the prediction as precise as possible, the target of predictor learning is to learn parameter $\theta$, so that the difference between true class labels of the training samples in $Y$ and the prediction results in $F_\theta$ could be minimized, while keeping the complexity of the predictor as low as possible. To measure how well the prediction results fit the true class label indicator, several loss functions $L(F_\theta, Y)$ could be considered to compare the prediction results in $F_\theta$ against the true class labels of the training samples in $Y$, such as the 0-1 loss function, the square loss function, the hinge loss function, and the logistic loss function. We summarize various loss functions in Table 1.

These loss functions introduced in Table 1 have been used widely in various learning problems. One common feature of these loss functions is that a sample-wise loss function is applied to each training sample equally and then the losses of all the samples are summed up to obtain the final overall loss. The sample-wise loss functions are of exactly the same form with the same parameter (if they have parameters). The basic assumption behind this loss function is that the training samples are of the same importance. However, due to the limitation of the sampling technology and noises occurred during the sampling procedure,
Table 1: Various empirical loss functions for predictor learning

| Title       | Formula of $L(F_{\theta}, Y)$                                                                 | Notes                                                                 |
|-------------|-----------------------------------------------------------------------------------------------|----------------------------------------------------------------------|
| 0-1 Loss    | $\sum_{i} \mathbb{I}[F_{\theta Y_{i}} < 0]$, where $\mathbb{I}(\cdot)$ is the indicator function and $\mathbb{I}(\cdot) = 1$ if $\cdot$ is true, 0 otherwise. | The 0-1 loss function is NP-hard to optimize, non-smooth and non-convex. |
| Square Loss | $\sum_{i} ||F_{\theta Y_{i}} - Y_{i}||_{2}^{2}$                                                 | The square loss function is a convex upper bound on the 0-1 loss. It is smooth and convex, thus easy to optimize. |
| Hinge Loss  | $\sum_{i} \min(1 - F_{\theta Y_{i}} Y_{i}, 0)$                                               | The hinge loss function is not smooth but subgradient descent can be used to optimize it. It is the most common loss function in SVM. |
| Logistic Loss | $\sum_{i} \ln(1 + e^{-F_{\theta Y_{i}} Y_{i}})$                                           | This loss function is also smooth and convex, and is usually used in regression problem. |

there are some noisy and outlying samples in real-world applications. If we use the transitional loss functions listed in Table 1, the noisy and outlying training samples will play more important roles even than the good samples. Thus the predictors learned by minimizing the transitional loss functions are not robust to the noisy and outlying training samples, and could bring a high error rate when applied to the prediction of test samples.

Recently, regularized correntropy framework has been proposed for robust pattern recognition problems [10, 11, 12, 13]. In [14], He et al. argued that the classical mean square error (MSE) criterion is sensitive to outliers, and introduced the correntropy to improve the robustness of the presentation. Moreover,
the $l_1$ regularization scheme is imposed on the correntropy to learn robust and sparse representations. Inspired by their work, we propose to use the regularized correntropy as a criterion to compare the prediction results and the true class labels. We use correntropy to compare the predicted labels and the true labels, instead of comparing the feature of test sample and its reconstruction from the training samples in He et al.’s work. Moreover, an $l_2$ norm regularization is introduced to control the complexity of the predictor. In this way, the predictor learned by maximizing the correntropy between prediction results and the true labels will be robust to the noisy and outlying training samples. The proposed classification Machine Maximizing the Regularized CorrEntropy, which is called RegMaxCEM, is supposed to be less sensitive to outlining samples than those with transitional loss functions. Yang et al. [15] also proposed to use correntropy to compare predicted class labels and true labels. However, in their framework, the target is to learn the class labels of the unlabeled samples in a transductive semi-supervised manner, while we try to learn the parameters for the class label predictor in a supervised manner.

The rest of this paper is structured as follows: In Section 2, we propose the regularized maximum correntropy machine by constructing an objective function based on the maximum correntropy criterion (MCC) and developing an expectation maximization (EM) based alternative algorithm for its optimization. In Section 3, the proposed methods are validated by conducting extensive experiments on two challenging pattern classification tasks. Finally, we give the conclusion in Section 4.

2. Regularized Maximum Correntropy Machine

In this section we will introduce the classification machine maximizing the correntropy between the predicted class labels and the true class labels, while keeping the solution as simple as possible.
2.1. Objective Function

To design the predictors $f^l_\theta(x)$, we first represent the data sample $x$ as $\bar{x}$ in the linear space and the kernel space as:

\[
\bar{x} = \begin{cases} 
  x, & \text{(linear)}, \\
  K(\cdot, x), & \text{(kernel)},
\end{cases}
\]  

(1)

where $K(\cdot, x) = [K(x_1, x), \cdots, K(x_N, x)]^T \in \mathbb{R}^N$ and $K(x_i, x_j)$ is a kernel function between $x_i$ and $x_j$. Then a linear predictor $f^l_\theta(x)$ will be designed to predict whether the sample belongs to the $l$-th class as

\[
f^l_\theta(x) = w_l^T \bar{x} + b_l, \ l = 1, \cdots, L,
\]  

(2)

where $\theta = \{(w_l, b_l)\}_{l=1}^L$ is the parameters of the predictors, $w_l \in \mathbb{R}^D$ is the linear coefficient vector and $b_l \in \mathbb{R}$ is a bias term for the $l$-th predictor. The target of predictor designing is to find the optimal parameters to have the prediction result $f^l_\theta(x_i)$ of the $i$-th sample to fit its true class label indicator $Y_{li}$ as well as possible, while keeping the solution as simple as possible. To this end, we consider the following two problems simultaneously when designing the objective function:

Prediction Accuracy Criterion based on Correntropy To consider the prediction accuracy, we could learn the predictor parameters by minimizing a loss function listed in Table 1 as

\[
\min_{\theta} L(F_\theta, Y)
\]  

(3)

However, as we mentioned in Section 1, all these loss functions are applied to all the training samples equally, which is not robust to the noisy samples and outlying samples. To handle this problem, instead of minimizing a loss function to learn the predictor, we use the MCC [10] framework to learn the predictor by maximizing the correntropy between the predicted results and the true labels.
Remark 1: In previous studies, it has been demonstrated that the MCC is robust to outliers, for example, see [10]. Based on this, we assume that the predictors developed by MCC should also be insensitive to outliers.

Correntropy is a generalized similarity measure between two arbitrary random variables $A$ and $B$. However, the joint probability density function of $A$ and $B$ is usually unknown, and only a finite number of samples of them are available as $\{(a_i, b_i)\}_{i=1}^{d}$. It leads to the following sample estimator of correntropy:

$$V(A, B) = \frac{1}{d} \sum_{i=1}^{d} g_{\sigma}(a_i - b_i), \quad (4)$$

where $g_{\sigma}(a_i - b_i) = \exp\left(-\frac{(a_i - b_i)^2}{2\sigma^2}\right)$ is a Gaussian kernel function, and $\sigma$ is a kernel width parameter. For a learning system, MCC is defined as

$$\max_{\vartheta} \frac{1}{d} \sum_{i=1}^{d} g_{\sigma}(a_i - b_i) \quad (5)$$

where $\vartheta$ is the parameter to be optimized in the criterion so that $B$ is as correlated to $A$ as possible.

Remark 2: $\vartheta$ is usually a parameter to define $B$, but not the kernel function parameter $\sigma$. In the learning system, we try to learn $\vartheta$ so that with the learned $\vartheta$, $B$ is correlated to $A$. For example, in this case, $A$ is the true class label matrix while $B$ is the predicted class label matrix, and $\vartheta$ is the predictor parameter to define $B$.

To adapt the MCC framework to the predictor learning problem, we let $A$ be the prediction result matrix $F_{\theta}$ parameterized by $\theta$, and $B$ be the true class label matrix $Y$, and we want to find the predictor parameter $\theta$ such that $F_{\theta}$ becomes as correlated to $Y$ as possible under the MCC framework. Then, the following correntropy-based predictor learning model will be obtained:
\[
\max_\theta V(F_\theta, Y),
\]
\[
V(F_\theta, Y) = \frac{1}{L \times N} \sum_{l=1}^{L} \sum_{i=1}^{N} g_{\sigma}(F_{\theta li} - Y_{li}).
\] (6)
Note that in [10], MCC is used to measure the similarity between a test sample and its sparse linear representation of training samples, while in this work it is used to measure the similarity between the predicted class label and its true label. Also note that the dependence on \(\sigma\) in (6) and later (8), (11) relies on the dependence of the kernel function \(g_{\sigma}(\cdot)\). In our experiments, the \(\sigma\) value is calculated as
\[
\sigma = \frac{1}{2 \times L \times N} \sum_{l=1}^{L} \sum_{i=1}^{N} \|F_{\theta li} - Y_{li}\|_2^2
\] following [10].

**Predictor Regularization** To control the complexity of the \(l\)-th predictor independently, we introduce the \(l^2\)-based regularizer \(\|w_l\|^2\) to the coefficient vector \(w_l\) of the \(l\)-th predictor. We assume that the predictors of different classes are equally important, and the following regularizer is introduced for multi-class learning problem:
\[
\min_{w_l} \frac{1}{L} \sum_{l=1}^{L} \|w_l\|^2
\] (7)

**Remark 3:** The \(l^2\) norm is also used by support vector regression as a measure of model complexity. However, in support vector classification, this regularization term is either obtained by a “maximal margin” regularization or obtained by a “maximal robustness” regularization for certain type of feature noises [16]. Thus our \(l^2\) norm regularization term can also be regarded as a term to seek maximal margin or robustness.

**Remark 4:** The \(l_2\)-regularization is used in comparison to the \(l_1\)-regularization in our model. Using \(l_1\)-regularization we can seek the sparsity of the predictor coefficient vector, but it cannot guarantee the minimal model complexity, maximal margin or maximal robustness like the \(l_2\)-regularization, thus we choose to use the \(l_2\)-regularization. In the future, we will ex-
plore the usage of $l_1$-regularization to see if the prediction results can be improved.

By substituting $\theta = \{w_l, b_l\}_{l=1}^L$, $F_{\theta l_l} = f_{\theta l_l}^l(x_i)$, and combining both the predictor regularization term in (7) and the prediction accuracy criterion term based on correntropy in (6), we obtain the following maximization problem for the maximum correntropy machine:

$$\max_{(w_l, b_l)_{l=1}^L} \frac{1}{L \times N} \sum_{l=1}^L \sum_{i=1}^N g_{\sigma}(f_{\theta l_l}^l(x_i) - Y_{l_i}) - \alpha \frac{1}{L} \sum_{l=1}^L ||w_l||^2 \tag{8}$$

where $\alpha$ is a tradeoff parameter. This optimization problem is based on correntropy using a Gaussian kernel function $g_{\sigma}(x)$. It treats the prediction of individual training samples of individual classes differently. By this way, we can give more emphasis on samples with correctly predicted class labels, while those noisy or outlying training samples will have small contributions to the correntropy. In fact, when the regularizer term is introduced, (8) is a case of the regularized correntropy framework [14].

2.2. Optimization

Due to the nonlinear attribute of the kernel function $g_{\sigma}(x)$ in the objective function in (8), direct optimization is difficult. An attribute of the kernel function $g_{\sigma}(x)$ is that its derivative is also the same kernel function, and if we set its derivative to zero to seek the optimization of the objective, it is not easy to obtain a close form solution. However, according to the property of the convex conjugate function, we have:

**Proposition 1** There exists a convex conjugate function $\varphi$ of $g_{\sigma}(x)$ such that

$$g_{\sigma}(x) = \max_p (p||x||^2 - \varphi(p)) \tag{9}$$

and for a fixed $x$, the maximum is reached at $p = -g_{\sigma}(x)$. This proposition is taken from [17], which is further derived from the theory of convex conjugated functions. It is further discussed and used in many applications such as [10, 14].
By substituting (9) into (8), we have the augmented optimization problem in an enlarged parameter space

$$\max_{(w_l, b_l)_{l=1}^L, P} \frac{1}{L \times N} \sum_{l=1}^L \sum_{i=1}^N \left[ P_{li} \| f_{w_l, b_l}^l(x_i) - Y_{li} \|^2 - \varphi(P_{li}) \right] - \alpha \frac{1}{L} \sum_{l=1}^L \| w_l \|^2$$

$$= \frac{1}{L \times N} \sum_{l=1}^L \sum_{i=1}^N \left[ P_{li} \| w_l^\top \tilde{x}_i + b_l - Y_{li} \|^2 - \varphi(P_{li}) \right] - \alpha \frac{1}{L} \sum_{l=1}^L \| w_l \|^2,$$

(10)

where $P = [P_{li}] \in \mathbb{R}^{N \times L}$ are the auxiliary variable matrix. To optimize (10), we adapt the EM framework to solve $P$ and $\{(w_l, b_l)\}_{l=1}^L$ alternately.

2.2.1. Expectation Step

In the expectation step of the EM algorithm, we calculated the auxiliary variable matrix $P$ by fixing $\theta$. Obviously, according to Proposition 1, the maximum of (10) can be reached at

$$P = -g_\sigma(F_\theta - Y),$$

$$P_{li} = -g_\sigma(w_l^\top \tilde{x}_i + b_l - Y_{li}).$$

(11)

Note that $g_\sigma(X)$ is the element-wise Gaussian function. With fixed predictor parameters, the auxiliary variable $-P_{li}$ can be regarded as confidence of prediction result of the $i$-th training sample regarding to the $l$-th class. The better the $l$-th prediction result of the $i$-th sample fits the true label $Y_{li}$, the larger the $-P_{li}$ will be.

Remark 5: It is interesting to see if there is any relation between the auxiliary variables in $P$ and the slack variables in SVM, since both can be viewed as measures of classification losses. The slack variables in SVM are the upper boundaries of hinge losses of the training samples, while the auxiliary variables in $P$ are a dissimilarity measure between the predicted labels and the true labels under the framework of the MCC rule, which is also a loss function. Meanwhile, the auxiliary variables in $P$ also play a role of weights of different training samples as in (10), so that the learning can be robust to the noisy labels, but the auxiliary variables in SVM do not have such functions.
Remark 6: In the expectation step, we actually solve an alternative optimization of solving $P$ while fixing $\{(w_l, b_l)\}_{l=1}^{L}$. However, according to Proposition 1, the solution for this optimization problem is in the form of (11), which can be calculated directly and makes it an expectation step of the EM algorithm.

2.2.2. Maximization Step

In the maximization step of the EM algorithm, we solve the predictor parameters $\{(w_l, b_l)\}_{l=1}^{L}$ while fixing $P$. The optimization problem in (10) turns out to be

$$\max_{\{(w_l, b_l)\}_{l=1}^{L}} \frac{1}{L \times N} \sum_{l=1}^{L} \sum_{i=1}^{N} [P_{li}||w_l^T \bar{x}_i + b_l - Y_{li}||^2 - \varphi(P_{li})] - \frac{1}{L} \sum_{l=1}^{L} ||w_l||^2. \tag{12}$$

Noticing $P_{li} < 0$ and removing terms irrelevant to $w_l$ and $b_l$, the maximization problem in (12) can be reformulated as the following dual minimization problem:

$$\min_{\{(w_1, b_1), \ldots, w_L, b_L\}} O(w_1, b_1, \ldots, w_L, b_L),$$

$$O(w_1, b_1, \ldots, w_L, b_L) = \frac{1}{L \times N} \sum_{l=1}^{L} \sum_{i=1}^{N} (-P_{li}||w_l^T \bar{x}_i + b_l - Y_{li}||^2) + \alpha \sum_{l=1}^{L} ||w_l||^2. \tag{13}$$

To simplify the notations, we define a vector $u_l = [u_{l1}, \ldots, u_{lN}]^\top \in \mathbb{R}^N$ so that $u_{li}^2 = -\frac{1}{N} P_{li}$. With $u_l$, the objective function in (13) can be rewritten as

$$O(w_1, b_1, \ldots, w_L, b_L) = \frac{1}{L} \sum_{l=1}^{L} [\|(w_l^T \bar{x}_i + b_l - Y_{li})\|^2 + \alpha ||w_l||^2]$$

$$= \frac{1}{L} \sum_{l=1}^{L} [(w_l^T \bar{X}_l + b_l u_l^T - \bar{Y}_l)(w_l^T \bar{X}_l + b_l u_l^T - \bar{Y}_l)^\top + \alpha u_l^\top w_l], \tag{14}$$

where $\bar{X}_l = [u_{l1} \bar{x}_1, \ldots, u_{lN} \bar{x}_N] \in \mathbb{R}^{D \times N}$ is the matrix containing all the training sample feature vectors weighted by $u_l$, and $\bar{Y}_l = [u_{l1} Y_{l1}, \ldots, u_{lN} Y_{lN}] \in \mathbb{R}^N$ is the $l$-th row of $Y$ weighted by $u_l$. 




Obviously, the optimization problem in (13) is a linear least squares problem. Analytical solution for (13) could be obtained easily. By setting the derivative of $O(w_1, b_1, \ldots, w_L, b_L)$ with regard to $b_l$ to zero, we have

$$\frac{\partial O(w_1, b_1, \ldots, w_L, b_L)}{\partial b_l} = \frac{1}{2L} (w_l^\top X_l + b_l u_l^\top - Y_l) 1_N = 0$$

$$\Rightarrow b_l = \frac{(Y_l - w_l^\top X_l) 1_N}{u_l^\top 1_N} = y_l - w_l^\top x_l,$$

where $y_l = \frac{Y_l 1_N}{u_l^\top 1_N}$ and $x_l = \frac{X_l 1_N}{u_l^\top 1_N}$. By substituting (15) to $O(w_1, b_1, \ldots, w_L, b_L)$, we have

$$O(w_1, \ldots, w_L) = \frac{1}{L} \sum_{l=1}^{L} \{ [w_l^\top (X_l - x_l u_l^\top)] [w_l^\top (X_l - x_l u_l^\top)]^\top - 2[w_l^\top (Y_l - y_l u_l^\top)]^\top + \alpha w_l^\top w_l \}$$

(16)

By setting the derivative of $O(w_1, \ldots, w_L)$ with regard to $w_l$ to zero, we have the optimal solution $w_l^*$

$$\frac{\partial O(w_1, \ldots, w_L)}{\partial w_l} = \frac{1}{2L} [2(X_l - x_l u_l^\top)(X_l - x_l u_l^\top)^\top w_l - 2(X_l - x_l u_l^\top)(Y_l - y_l u_l^\top)^\top + 2\alpha w_l] = 0$$

$$\Rightarrow w_l^* = [(X_l - x_l u_l^\top)(X_l - x_l u_l^\top)^\top + \alpha I]^{-1} (X_l - x_l u_l^\top)(Y_l - y_l u_l^\top)^\top,$$

(17)

where $I$ is an $D \times D$ identity matrix. Then we substitute $w_l^*$ into (15), and we will have the optimal solution of $b_l^*$,

$$b_l^* = y_l - w_l^\top x_l$$

(18)

2.3. Algorithm

Algorithm 1 summarizes the predictor parameter learning procedure of RegMaxCEM. The E-step and the M-step will be repeated for $T$ times.

3. Experiments

In the experiments, we will evaluate the proposed classification method on two challenging pattern classification tasks — bacteria identification [18] and
Algorithm 1 RegMaxCEM Learning Algorithm.

**Input:** Training set: \( \mathcal{D} = \{(x_i, y_i)\}_{i=1}^N \);

Initialize the auxiliary variable matrix \( P^0 = -1_{L \times N} \);

Represent each sample \( x_i \) as \( \widetilde{x}_i \) as in (1);

for \( t = 1, \cdots, T \) do

**Maximization-Step:** Update the predictor parameters \( \theta^t = \{(w^T_l, b^T_l)\}_{l=1}^L \) as in (17) and (18) by fixing \( P^{t-1} \).

**Expectation-Step:** Update the auxiliary variable matrix \( P^t \) as in (11) by fixing the predictor parameters \( \theta^t \).

end for

**Output:** Predictor parameters \( \theta^T = \{(w^T_l, b^T_l)\}_{l=1}^L \).

3.1. Experiment I: Bacteria Identification

3.1.1. Dataset and Setup

High-precision identification of bacteria is quite important for the diagnosis of cancers and bacterial infections. Recently, ensemble aptamers (ENSaptamers), which utilizes a small set of nonspecific DNA sequences, has been proposed to provide an effective platform for the detection of bacteria [18]. ENSaptamers is a sensor array with seven sensors, and each sensor is designed using a DNA element.

For the experiment, we collected in total 66 samples of 6 different bacteria, including S.typhimurium, S.flexneri, E.coli (CAU 0111), S.sonnei, S.typhi and E.coli (ATCC 25922). The number of samples for each bacteria varies from 9 to 13. Given an unknown bacteria sample with its fluorescence response patterns of ENSaptamer, the task is to identify which bacteria it is. To this end the seven fluorescence response patterns of ENSaptamer against the sample will be used to construct the 7-dimensional feature vector, and then the sample will be classified into one of the known bacteria using the RegMaxCEM predictor.

To conduct the experiment, we randomly split the entire dataset into two
non-overlapping subsets — the training set and the test set. 33 samples were used as training sample in the training set, while the remaining 33 ones as test samples. The predictor parameters of RegMaxCEM were trained using the feature vectors and class labels of the training samples. Then the class labels of the test samples were predicted by the trained predictor, and compared to their true labels to calculate the classification accuracy. The random split process (training/test) was repeated for ten times and the accuracies over these ten splits were reported as classification performance.

3.1.2. Results

We compare our proposed method against classifiers based on other loss functions, including square loss, hinge loss and logistic loss. The 0-1 loss is the simplest loss function, but difficult to optimize, thus is not compared in the experiment. The boxplots of accuracies of different methods using both linear and kernel representations are illuminated in Figure 1. As shown in Figure 1, predictor produced by maximizing the correntropy yields improvements over other loss functions. Given the extremely small variation of classification accuracies over the ten splits, though the improvements of the accuracies are not large in absolute terms (around 0.1), they are consistent and significant. To verify whether the improvements are statistically significant, we performed the paired t-tests to the accuracies of the proposed method and other compared methods. The null hypothesis of the T-test is that the accuracies of the proposed method and the compared methods come from distributions with equal means. The $P$ values of the t-tests are reported as measurements of statistically significance. A low $P$ value implies that the difference between the proposed method and the compared methods are statistically significant. The $P$ values are reported in Table 2. As we can see from the table, all the improvements archived by RegMaxCEM, for both linear representation and kernel representation, are statistically significant at the 0.05 significance level. This is not surprising: There are some noisy and outlying samples in the training set, which have been utilized by the methods with square loss, hinge loss or logistic loss as equally as other
samples, thus they bring some bias to the predictor. However, the RegMaxCEM
has the potential of filtering these samples, which can result in reliable learning
of predictors in practice. It is also interesting to notice that the square loss,
hinge loss and logistic loss have archived very similar classification accuracies.
Though they used different loss functions, these loss functions are applied to
the training samples equally.

![Boxplots of accuracies of bacteria identification.](image)

(a) Linear representation

(b) Kernel representation

Figure 1: Boxplots of accuracies of bacteria identification.

3.2. Experiment II: DNA-Binding Site Prediction

It is very important to predict the DNA-binding sites in proteins for under-
standing the molecular mechanisms of protein-DNA interaction. In this experi-




Table 2: P values of paired T-tests on accuracies of ten splits of RegMaxCEM and compared methods on bacteria identification.

| Linear representation | Kernel representation |
|-----------------------|-----------------------|
| Compared methods      | P values              | Compared methods | P values |
| Square Loss           | 0.0266                | Square Loss      | 0.0118   |
| Hinge Loss            | 0.0243                | Hinge Loss       | 0.0224   |
| Logistic Loss         | 0.0115                | Logistic Loss    | 0.0095   |

3.2.1. Dataset and Setup

The PDNA-62 database for DNA-binding site prediction has been used in this experiment. This database contains 8,163 sites in proteins in total. Among these sites, 1,215 of them are DNA-binding sites, while the remaining 6,948 sites are non-binding sites. We select 1,000 DNA-binding sites and 5,000 non-binding sites from the PDNA-62 database to construct our database for the experiment. Given a candidate site, the goal of DNA-binding site prediction is to predict whether it is a DNA-binding site or not. To this end, the evolutionary information, solvent accessible surface area and the protein backbone structure features were extracted from the site, and then combined to construct the feature vector. The feature vector was further inputted into the classifier to distinguish DNA-binding sites from the non-binding sites [19].

To conduct the experiment, we employed the 10-fold cross validation. The database was split into 10 non-overlapping folds randomly, one of which was used as the test set, while the rest 9 of them were used as the training set. The procedure was repeated for 10 times so that each fold was used as the test set once.

The prediction performance was measured by the receiver operating characteristic (ROC) and recall-precision curves. The usage of ROC curve is mainly due to the imbalanced classes. The ROC curve is created by plotting false posi-
tive rate (FPR) against true positive rate (TPR), while recall-precision curve is obtained by plotting recall against precision. The FPR, TPR, recall and precision are defined as:

\[
\text{FPR} = \frac{FP}{FP + TN}, \quad \text{TPR} = \frac{TP}{TP + FN},
\]

\[
\text{recall} = \frac{TP}{TP + FN}, \quad \text{precision} = \frac{TP}{TP + FP},
\]

(19)

where \(TP\) is the number of DNA-binding sites predicted correctly, \(FP\) is the number of non-binding sites predicted as DNA-binding sites wrongly, \(TN\) is the number of non-binding sites predicted correctly, while \(FN\) is the number of DNA-binding sites predicted as non-binding sites wrongly. For a better predictor, its ROC curve should be closer to the top left corner of the figure, while the recall-precision curve should be closer to the top right corner. Besides the two curves, the area under the ROC curve (AUC) is also used as a single measurement of the prediction. A better predictor will have a larger AUC value.

3.2.2. Results

The ROC and recall-precision curves of the proposed method and compared methods are reported in Figure 2. The predictors using linear and kernel representations are both illuminated. The AUC values of the ROC curves are reported in Table 3 as well. Overall the proposed method clearly outperform the other methods significantly, although there is some variability in prediction performance over different representation types. From Table 3, we could see that the accuracy of the predictor is slightly increased by using the kernel representation instead of the linear representation. The regularized correntropy based predictors gives much better results than other methods on both representations. An interesting result from the DNA-binding prediction on this dataset is that the predictor with the hinge loss function outperforms other two methods.
Figure 2: ROC and recall-precision curves on DNA-Binding site prediction experiment using both linear and kernel representations.
Table 3: AUC values of ROC curves on DNA-Binding site prediction experiment.

| Methods       | Linear representation AUC | Kernel representation AUC |
|---------------|---------------------------|----------------------------|
| RegMaxCEM     | 0.9226                    | 0.9344                     |
| Square Loss   | 0.8768                    | 0.8891                     |
| Hinge Loss    | 0.8908                    | 0.8961                     |
| Logistic Loss | 0.8747                    | 0.8776                     |

4. Conclusion and Future Work

In this paper, we present a novel regularized predictor learning model for multi-class pattern recognition problems. The predictor are learned by maximizing the correntropy between the prediction results and the true class labels. By applying the MCC rule, we could treat different training samples differently, so that the noisy and outlying training samples have less impact on the learning of predictors. Compared with the existing predictor models with various loss functions, it is robust to the noisy and outlying training samples. The experiments on bacteria identification and DNA-binding site prediction show that a good predictor may benefit much from a well designed loss function based on MCC. The proposed method outperformed the predictor with other popularly used loss functions. In the future, we will investigate if the regularized maximum correntropy framework can be used to regularize ranking score learning [20, 21], data representation [22, 23, 24, 25, 26, 27, 28, 29] Moreover, we also plan to extend the proposed regularized correntropy based classifier for bioinformatics problems [30, 31, 32, 33, 34], wireless sensor network [35], hardware fault detection [36, 37, 38, 39, 40], and large scale data analysis on distributed big data platforms [41, 42, 43, 44, 45].

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