Recursive NMF: Efficient Label Tree Learning for Large Multi-Class Problems

Lei Liu+, Prakash Mandayam Comar+, Sabyasachi Saha*, Pang-Ning Tan+, Antonio Nucci*
+Dept. of Computer Science, Michigan State University, East Lansing, MI 48824, USA
+{liulei1, mandayam, ptan}@msu.edu
*{saha, anucci}@narus.com

Abstract

Many object recognition or concept identification tasks require accurate detection of large number of classes. These applications present enormous challenges to traditional classification methods, which are mostly designed for solving problems with small number of classes. In this paper, we develop a method called recursive non-negative matrix factorization (RNMF) for building a hierarchical label tree over set of classes. The internal nodes of the tree employ linear classifiers to propagate a data instance to its corresponding leaf node, where one or more one class support vector machine (SVM) classifiers is applied to accurately predict its class. Our experiment results show that the proposed method achieves significant gain in test efficiency and comparable accuracy to some of the more expensive label tree learning methods.

1 Introduction

Classification with large number of classes is an important but challenging problem with many potential applications including malware detection [3], visual object recognition [4], and text categorization [6]. Existing approaches for multi-class classification employ a collection of binary classifiers, organized either in a flat (non-hierarchical) or a hierarchical structure. Non-hierarchical approaches, including one-versus-rest [10], one-versus-one [7], and error-correcting output coding (ECOC) [8], require invocation of all the binary classifiers during testing to determine its class. The efficiency of such methods during testing is therefore proportional to the number of binary classifiers that must be invoked during testing.

Hierarchical approaches [9, 2, 1, 4] help to improve test efficiency by organizing the classifiers in a tree or DAG-like structure. Their runtime complexity during testing is proportional to the depth of the hierarchy. For example, the Decision Directed Acyclic Graph (DDAG) [9] approach arranges the one-versus-one classifiers in a rooted binary DAG in order to reduce the runtime complexity for testing from $O(k(k-1)/2)$ to $O(k)$. In fact, it is possible to achieve sublinear complexity by training classifiers to discriminate more than two classes at each node of the hierarchical structure. Such a strategy was adopted by label tree learning methods such as [1, 4], which employ a linear classifier at each node to assign data instances from multiple classes to their corresponding child nodes. Only the binary classifiers located along the path from the root node to one of the leaf nodes in the hierarchical structure will be invoked to determine the class label of a test instance.

The hierarchical approaches also vary in terms of how their tree or DAG structure and the associated classes at each node are induced from training data. For example, Bengio et al. [1] developed their label embedding tree approach by training $k$ one-vs-rest classifiers to obtain an initial confusion matrix, which is then used as an affinity matrix for applying spectral clustering to partition the classes into smaller subgroups. Limitations of this method, which include the additional training time needed for constructing the initial confusion matrix and the hard partitions generated by the nodes of the tree, have been well-documented [4]. Deng et al. [4] presented an alternative strategy that simultaneously learns the class partitioning and the weights of the associated linear classifiers by optimizing a joint objective function. However, the approach requires solving an integer programming problem, which is NP-hard. The
authors proposed a relaxation of the problem to a linear program in order to find a polynomial time solution. This paper presents a novel label tree learning approach to speed up model testing in large multi-class problems. Unlike previous work, the leaf nodes of the tree contain multiple one-class SVM models to distinguish instances from different classes. This helps to shorten the depth of the tree without significantly degrading its accuracy. Our label tree learning algorithm partitions the classes and learns the weights of their classifiers simultaneously. It performs a soft partitioning of the classes by minimizing an information-theoretic loss function, which can be cast into a regularized non-negative matrix factorization problem. The regularization term is unique in that it minimizes the entropy of the node partitions. Our experimental results suggest that the induced tree achieves significant improvement in terms of test efficiency and comparable accuracy as other more complex tree learning methods.

2 Tree Learning with Recursive NMF

Let \( D = \{(x_1, y_1), (x_2, y_2), \ldots, (x_n, y_n)\} \) denote the training instances sampled from a distribution \( P \) over \( X \times Y \), where \( X = \mathbb{R}^d \) is the \( d \)-dimensional feature space and \( Y = \{1, 2, \ldots, k\} \) is the set of \( k \) distinct classes. The goal of multi-class classification is to learn a target function \( f \) that minimizes the classification error of any instance drawn from \( P \), i.e., \( \mathbb{P}(x, y) \sim P[f(x) \neq y] \). A typical approach for multi-class learning is to reduce the problem into multiple binary classification tasks. However, when the number of classes is large, applying all the binary classifiers to determine the predicted class becomes computationally infeasible.

This paper presents a novel and efficient label-tree learning algorithm for solving large multi-class problems. A label tree \([1]\) is a structure \( T = (V, E, \Phi, \Lambda) \), where \( V \) is the set of nodes, \( E \) is the set of directed links connecting each node to its children, \( \Phi \) is the set of classification functions and \( \Lambda \) is the set of class labels associated with the nodes. The label tree framework proposed in this study is shown in Figure 1. The framework recursively applies a regularized non-negative matrix factorization approach to learn the discriminant function at each internal node in the tree. The leaf nodes employ one or more 1-class SVM classifiers to identify the class label of a test instance.

Let \( X \) be an \( n \times d \) data matrix containing the feature vector of the training instances and \( Y \) be an \( n \times k \) matrix that represents the class label information, i.e., \( Y_{ij} = 1 \) if the \( i^{th} \) instance belongs to the \( j^{th} \) class, and zero otherwise. Let \( p \) be the branching factor of the tree. At the root node \( v_0 \), our goal is to learn a partition membership matrix \( L \) of size \( n \times p \) where \( n \) is the number of training instances assigned to the node and the associated feature weight matrix \( W \), which is a \( d \times p \) matrix that indicates the importance of each feature in defining the given partition. This is accomplished by minimizing the following objective function:

\[
\min_{L, W} D(X \parallel LW^T) + \lambda H(L^TY)
\]

\[
\text{s.t.} \quad L^T1_p = 1_n
\]

where \( D(X \mid LW^T) \) is the Kullback-Leibler divergence between the input data matrix \( X \) and the product of its latent factors \( LW^T \). The regularizer term \( H(L^TY) \) corresponds to the entropy of the class distribution of each partition. The regularizer is needed to avoid splitting instances from the same class into too many partitions. Since the entropy definition of the partitions requires the rows of matrix \( L^TY \) to be properly normalized (i.e., sum up to 1), i.e.,

\[
H(M) = -\sum_i M_{i+} \sum_j M_{ij} \log \frac{M_{ij}}{M_{i+}}
\]

where \( M_{i+} = \sum_j M_{ij} \) and \( M_{++} = \sum_{ij} M_{ij} \), the normalization is achieved by the constraint \( L^TY^T1_p = L^T1_p = 1_n \) since \( Y^T1_p = 1_p \). \( \lambda \) is a parameter provided by the user that controls the trade off between optimal partition of \( X \) and class purity of each partition. Thus, the Lagrange formulation of (1) is given by:

\[
\mathcal{L} = \sum_{i,j} [X_{ij} \log \frac{X_{ij}}{(LW^T)_{ij}} - X_{ij} + (LW^T)_{ij}]
\]

\[
- \lambda \sum_{ij} (L^TY)_{ij} \log (L^TY)_{ij}
\]

\[
- \sum_s \mu_s (\sum_m L_{sm}^T - 1)
\]

The objective function is minimized by taking its partial derivative with respect to the parameters \( W \) and \( L \) and

\(^1\lambda \) is set to the default value 1 in our experiments
setting them to zero:

$$\frac{\partial L}{\partial W_{pq}} = -\sum_i X_{ip} L_{iq} \in (LW_T)_{ip} + \sum_i L_{iq} = 0 \quad (2)$$

$$\frac{\partial L}{\partial L_{pq}} = -\sum_j X_{pj} W_{jq} \in (LW_T)_{pj} - \lambda \sum_j Y_{pj} \log(L^T Y)_{qj} - \lambda \sum_j Y_{pj} - \mu_q = 0 \quad (3)$$

By summing up over $p$ in Equation (3), we have

$$\mu_q = -\frac{1}{n} \sum_j X_{pj} W_{jq} \in (LW_T)_{pj} + \sum_j W_{jq} - \lambda \sum_j \frac{Y_{pj}}{n} \log(L^T Y)_{qj} - \lambda \quad (4)$$

Replacing (4) back into (3), the partial derivative can be rewritten as follow:

$$\frac{\partial L}{\partial L_{pq}} = -\sum_j X_{pj} W_{jq} \in (LW_T)_{pj} + \frac{1}{n} \sum_j X_{pj} W_{jq} \in (LW_T)_{pj} - \lambda \sum_j Y_{pj} \log(L^T Y)_{qj} + \lambda \sum_j \frac{Y_{pj}}{n} \log(L^T Y)_{qj} \quad (5)$$

The objective function can be solved using a gradient descent approach. Following the approach used in [5], the gradient descent formula can be transformed into a multiplicative update formula as follows:

$$W_{pq} = W_{pq} \frac{(X^T L)_{pq}}{(W L^T L)_{pq}}, \quad L_{pq} = L_{pq} \frac{\alpha}{\beta} \quad (6)$$

in which,

$$\alpha = \sum_j X_{pj} W_{jq} \in (LW_T)_{pj} + \lambda \sum_j Y_{pj} \log(L^T Y)_{qj}$$

$$\beta = \frac{1}{n} \sum_j X_{pj} W_{jq} \in (LW_T)_{pj} + \lambda \sum_j \frac{Y_{pj}}{n} \log(L^T Y)_{qj}$$

Algorithm 1 summarizes the key steps of our proposed recursive Non-Negative Matrix Factorization (RNMF) framework for label tree learning. We recursively apply Algorithm 1 to split the nodes in the tree until one of the following stopping criteria is met: (1) stop if all instances in a partition belong to the same class, or (2) stop if the node contains less than minleaf training instances, or (3) stop if there is only one class containing more than minclass instances\(^2\).

Instead of learning a separate classifier at each internal node, we assign data instances to their respective partitions as follows. Given a test instance $x_{test}$, we determine the partition it should be assigned to by computing the following $1 \times p$ vector: $\pi = x^T_{test} W_v (W^T_v W_v)^{-1}$ and choose the partition $j$ that has the largest magnitude, i.e., $j = \arg \max_k \pi_k$. The assignment procedure is repeated until the test instance reaches a leaf node, where we apply the 1-class SVM models associated with the node to predict its class label. The multi-class prediction using multiple 1-class SVM is described in our previous work [3].

### 3 Experimental Evaluation

We evaluated the performance of our algorithm on Wikipedia data from October 9, 2009. The original data contains about 3.6 million articles categorized into 336K categories. We choose the largest 214 categories, which contain 24,378 articles. We use averaged F1 score for all classes as our performance measure. The results reported in this study are obtained from 5-fold cross validation. We compared the performance of RNMF against two state-of-the-art label tree learning algorithms: DDAG [9] and the confusion matrix approach (CM) [1]. The results are summarized in Table 1. The results show that RNMF outperforms the confusion matrix based label embedding tree approach both in terms of their F1 score and test efficiency. Furthermore, the depth of our tree is also shorter. When compared against DAGSVM, which requires building a quadratic number

\(^2\)We set minleaf = 15 and minclass = 5 for our experiments.
of 1-vs-1 classifiers, our F1-score is slightly worse but our test time is at least 8 times faster.

Note that the performance of RNMF depends on the branching factor $p$ and the depth of the tree. If the tree has only a single node (i.e., a flat model consisting of $k$ one-class SVM models), our F1 score and test times are comparable to that of DAGSVM (as shown in Figures 2 and 3). As the depth of the tree increases, the test time improves significantly at the expense of decreasing F1 values. In fact, if we terminate the tree growing procedure at depth equals to 3, the F1 score is around 0.51 but the speedup is almost 3 times faster.

4 Conclusion

This paper presents a novel learning algorithm for large multi-class problems called Recursive NMF. The algorithm creates soft partitions of the classes by solving a regularized non-negative matrix factorization problem and uses a set of 1-class SVM models to predict the classes at the leaf nodes of the tree. Our experimental results on Wikipedia data suggests that RNMF outperforms a state-of-the-art label embedding tree algorithm [1] both in terms of its F1 score and test time. It also achieves comparable F1-score but higher testing efficiency compared to DDAG [9].

5 Acknowledgement

This research was conducted by the first two authors at Narus Corporation. Prakash Mandayam Comar and Pang-Ning Tan’s research are supported in part by ONR grant number N00014-09-1-0663.

References