COMP5318 MACHINE LEARNING AND DATA MINING ASSIGNMENT 2 REPORT

OPTIMIZED CLASSIFICATION ON FOREST COVERTYPE

Based on kNN, Linear Regression and Random Forest Algorithms

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Abstract

This project re-classifies Covtype dataset using three classification algorithms: kNN, Linear Regression and Random Forest. Evaluating the performance of these three supervised algorithms and choose the most appropriate one fitting in solving similar datasets classification problems. Finally, kNN was considered as the best.

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INTRODUCTION

Classification aims to classify input data with common characteristics into the same categories as efficient as possible. These years, methods of classification are proposed increasingly. However, the performance of each method are quite different.

In this assignment, we have chosen the Covtype dataset as an object, a classical dataset with 581,012 instances of forests and 54 dimensions in judging the cover type, divided into seven labels. Since the dataset has been classified, we chose three supervised learning methods to re-classify in order that we can compare the performance of these three methods, and through deeper analyzing, we would choose one of them as recommended method. The three algorithms are: kNN, Linear Regression and Random Forest.

As the performance of an algorithm mainly includes time consumption and accuracy, the implementation has strengthened our understanding of each algorithms, including their efficiency and situations they fits for separately. Moreover, we got experience on comparing and choosing the appropriate methods when confronting with real problems, which is premise for rewriting and optimizing classical machine learning methods.

SPECIFICATION

The whole project was running and tested on ThinkPad T540P with i5-4200m@2.5GHz and 8G Ram.

The OS type is Ubuntu 16.04LTS.

PREVIOUS WORK

Before working on the problem, we have referred to several literature to find successful instances in dealing with similar dataset. This process helped us define the algorithms we would use for achieving the target time-saving and cost-effective in the process.

The characteristics of Covtype dataset are summarized as following:

1. Massive instances but less dimensions.

There are over 580000 instances shown in the dataset, however, only over 50 dimensions used for classification

- 2. Data in the dataset are discrete objects, not continuous.
- 3. With 7 labels given, the re-classify process should be a supervised learning process.
- 4. There are 7 cover types. Therefore, this is a multi-classification problem.

Based on the former dataset characteristics, following appropriate methods was found.

Method 1: VFDT (Very Fast Decision Tree learner), a decision-tree leaning system based on Hoeffding trees.

This algorithm is a representation of data stream classification technology. Since data today usually appears in data stream format, with the real-time, continuous(not actually continuous, just numerous), infinite, and non-reproducible four properties, and static classification cannot satisfy the real needs, classification for data stream is becoming more and more prevalent.

VFDT has the ability to incorporate and classify new information online in shorter training time by dividing the income data stream. It is powerful in dealing with large datasets. Moreover, as a ready-to-use model, VFDT can be used after the first few data have been trained, and its quality increase smoothly with time.

Method 2: Bagging and Boosting.

The two methods also came up from data stream classification wave and usually used as ensemble classification methods to generate advanced classifiers.

Bagging can be used to enhance the effect of classifier, which produces several replicate training sets by random sampling, then getting corresponding weak classifiers by training them separately, and integrating them finally.

Similar with bagging, boosting uses all weak classifiers to form a strong classifier. However, instances in boosting are all based with certain weight corresponding to the importance of each repetition. So adjusting the weights can create more accurate classifiers.

Method 3: Round Robin classification, a method based on separate-and-conquer rule algorithms.

It has attracted much attention in neural networks and SVM (support vector machines) communities. The basic idea is to transform multi-classification problems into binary classification problems. During the process, one classifier is applied for each pair of classes and ignoring all others when using only training examples for these two classes. Then the complexity is lower. Round Robin classification has been proved to get further improvement

by integrated with bagging algorithm mentioned above.

METHODS AND DESIGN

Preprocessing

The preprocessing in this project can be divided into two types: Preprocessing of the dataset itself in order to generate predicting set and training set; preprocessing for a defined method for improving the accuracy. This part will only introduce the former, and the special preprocessing for each algorithms themselves will be followed with algorithms implementation.

In the preprocessing, we used the origin dataset to form four subsets called train sample, train target, predict sample, predict target. It was completed by generate samples and targets.py.

Firstly, the whole dataset was separated into ten parts equally and randomly. Choose one subset from the ten and extracting the last column from the subset, as the last column is the label of instance.

Secondly, defining the remaining 53 columns of the chosen subset as predicting set, called predict sample in the procedure, and naming the last column as predict target individually. It is kept for testing the accuracy.

Thirdly, integrate the remaining nine subsets into one, then repeating the former process to get training set, called train sample and train target in the procedure.

Finally, repeating the process above all to form ten-folds.

Algorithm Selection

For supervised learning, we have learnt four methods: kNN(k-NearestNeighbor), Naive Bayes, Linear Regression, and SVM(Support Vector Machine). And there are several algorithms that we haven't learnt, such as Random Forest, Adaboost, SVR etc. In this assignment, we would like to choose three algorithms with superior diversity in theory, so that the comparison result would be more obvious. At the same time, time consumption is also an important index to consider.

In theory, SVM and Linear Regression have several similarities, and both requires highdimensional vector calculation with high accuracy also high time complexity. Naive Bayes and kNN are both low complexity relatively, however, Naive Bayes is better to deal with equal-weight-dimension dataset.

As a result, we chose kNN, Linear Regression and Random Forest as the comparison objects, for these three algorithms are not only diverse in performance, but can be completed in the same external open-source library: sklearn. As is known, variable libraries are possible to influence algorithm performance.

Algorithm Introduction

This part will introduce the algorithms separately and describe the significance of each parameter in details.

kNN

K-nearest neighbors algorithm (k-NN) is a non-parametric method used for classification. Representing instance x as $a_1(x)$, $a_s(x)$,..., $a_n(x)$

$$d(x_i, y_i) = \sqrt{\sum_{r=1}^{n} (a_r(x_r) - a_r(x_j))^2}$$

Where: $a_r(x)$ = the r.th attribute of x, $d(x_i, x_j)$ = distance between instance x_i and x_j for discrete dataset, it performs:

$$\hat{f}(x_q) \leftarrow argmax \sum_{i=1}^k \delta(v, f(x_i))$$

Where: v is an element of V set,V is finite set v_1 , v_2 , v_3 , ..., v_n select k nearest instances represented as x_1 , x_2 ,..., x_k . Then return $\hat{f}(x_q)$ In the Covtype dataset, the attribute was fixed as 53. kNN needs no special preprocessing. The dataset was directly classified through knn.py

Linear Regression

Logistic Regression can be used as a binary as well as multi-classification regression when the dependent variable is dichotomous. Using the thought of logistic regression, in our assignment, we can build up a linear model:

$$z_i = \vec{x}_i \cdot \vec{w}_i = w_0 + x_{i1} w_1 + x_{i2} w_2 + \dots + x_{i(D+1)} w_{(D+1)}$$

for i=1,2,... n, where:

w stands for the parameter of the learnt.

x stands for test data, namely forest instance here.

Equating the linear model to a probability p(x) with logistic transformation applied.

$$\log(\frac{p(\vec{x}_i)}{1 - p(\vec{x}_i)}) = \vec{x}_i \cdot \vec{w}_i = z_i$$

Therefore, we could derive:

$$p(\vec{x}_i \cdot \vec{w}_i) = \frac{1}{1 + e^{-z_i}}$$

Also, we can have loss function:

$$loss(\vec{w}) = -l(\vec{w}) = \sum_{i=1}^{N} \log(1 + e^{z} - y_i z_i)$$

Where y_i is 0 or 1 in logistic regression.

Based on the above process and applying gradient descent algorithm for each label, we can get a estimated weights vector w for every label. Using this vector, we can get the most probable label for a specific data.

In this assignment, we designed a specific preprocessing part for improving the performance of LR method, accordingly, we used K-Means in advance in order to achieve statistical outlier removal. Thus, the original dataset had been re-classified with KMeans, and formed 7 new clusters. Then LR worked on the new clustering dataset both to observe if there was an optimization on LR performance.

Random Forest

A Random Forest consists of a collection of simple tree predictors, each of which has the ability to produce a response when presented with a set of predictor values and can also be used to classify the final result. The optimal size of predictor variables is given by log(2M+1), where M is the number of inputs.

Given a set of simple trees and a set of random predictor variables, the Random Forest

method defines a margin function that measures the extent to which the average number of votes for the correct class exceeds the average vote for any other class present in the dependent variable. Given an ensemble of classifiers $h_1(x), h_2(x), ..., h_K(x)$, and with the training set drawn at random from the distribution of the random vector Y, X, define the margin function as:

$$mg(\vec{X}, Y) = av_k I(h_k(\vec{X}) = Y) - \max_{j \neq Y} av_k I(h_k(\vec{X}) = Y)$$

The error can be defines as: $PE^* = P_{X,Y}(mg(\vec{X}, Y) < 0)$

While implementation of Random Forest, we did similar preprocessing as LR methods, using KMeans to cluster the dataset in advance and then evaluating the performance.

EXPERIMENTS AND DISCUSSION

Performance

The index for evaluating the performance of each algorithms are: precision, recall, f1-score and support. They are calculated as follows:

Precision = TP/(TP + FP)

Recall = TP/(TP + FN)

$$F1 - Score = P * R/2(P + R), P = Precision, R = Recall$$

where: TP means actually true and predict positive; FP means actually false but predict positive; FN means actually false and predict negative.

Table 1-3 shows the performance of three algorithms using 10 folds cross validation.

In addition to precision, recall and f1-score, we derive a more detailed confusion matrix showing the differences between true data and our prediction. From this matrix, we could easily see the confused classifications. Rows are predictions and columns are real data. The matrix for each algorithm are shown in Table 4-6, which is derived from randomly selected training samples and prediction samples.

Table 1: kNN performance table

Folds	Precision	Recall	F1-score	Support
1	0.97	0.97	0.97	58102
2	0.97	0.97	0.97	58102
3	0.97	0.97	0.97	58102
4	0.97	0.97	0.97	58102
5	0.97	0.97	0.97	58102
6	0.97	0.97	0.97	58102
7	0.97	0.97	0.97	58102
8	0.97	0.97	0.97	58102
9	0.97	0.97	0.97	58102
10	0.97	0.97	0.97	58102

Table 2: Logistic Regression performance table

Folds	Precision	Recall	F1-score	Support
1	0.76	0.71	0.73	58102
2	0.73	0.71	0.73	58102
3	0.75	0.71	0.72	58102
4	0.75	0.71	0.73	58102
5	0.75	0.71	0.73	58102
6	0.75	0.71	0.72	58102
7	0.75	0.71	0.73	58102
8	0.75	0.71	0.73	58102
9	0.75	0.71	0.72	58102
10	0.75	0.71	0.73	58102

 Table 3: Random Forest performance table

Folds	Precision	Recall	F1-score	Support
1	0.95	0.94	0.94	58102
2	0.95	0.95	0.95	58102
3	0.95	0.95	0.95	58102
4	0.95	0.95	0.95	58102
5	0.94	0.94	0.94	58102
6	0.95	0.95	0.95	58102
7	0.95	0.95	0.95	58102
8	0.95	0.95	0.95	58102
9	0.95	0.95	0.95	58102
10	0.95	0.95	0.95	58102

Table 4: kNN Detailed Confusion Matrix

Predicted/True	1.0	2.0	3.0	4.0	5.0	6.0	7.0	All
1.0	20670	604	1	0	10	1	41	21327
2.0	532	27436	38	0	63	16	5	28090
3.0	0	43	3516	9	3	48	0	3619
4.0	0	0	47	214	0	16	0	277
5.0	11	81	11	0	867	6	0	976
6.0	1	44	62	9	3	1605	0	1724
7.0	40	7	0	0	1	0	2041	2089
All	21254	28215	3675	232	947	1692	2087	58102

Table 5: Logistic Regression Detailed Confusion Matrix

Predicted/True	1.0	2.0	3.0	4.0	6.0	7.0	All
1.0	14290	6238	5	0	0	355	20888
2.0	4973	22876	546	0	57	63	28515
3.0	5	417	3109	11	132	0	3674
4.0	0	0	194	39	11	0	244
5.0	22	876	57	0	10	0	965
6.0	1	625	1012	2	91	0	1731
7.0	1084	23	7	0	0	971	2085
All	20375	31055	4930	52	301	1389	58102

Table 6: Random Forest Detailed Confusion Matrix

Predicted/True	1.0	2.0	3.0	4.0	5.0	6.0	7.0	All
1.0	20198	1082	0	0	4	3	40	21327
2.0	1074	26890	44	3	42	28	9	28090
3.0	2	91	3432	9	3	82	0	3619
4.0	0	1	44	224	0	8	0	277
5.0	23	212	14	0	719	8	0	976
6.0	3	62	152	6	1	1500	0	1724
7.0	112	8	0	0	0	0	1969	2089
All	21412	28346	3686	242	769	1629	2018	58102

Table 7: Running Time

Name Running Time(seconds per folds on average)

kNN 8.22 Logistic Regression 183.11 Random Forest 13.1

Runtime

Table 7 shows the running time on average for each algorithm

Comparison and Discussion

Totally, from the results shown above, we found that for a single algorithm, each fold of ten has similar performance and running time, which means that the functions we used from external libraries are quite stable and scientific.

kNN algorithm performed excellent with the lowest time cost as well as the highest precise, running out the results within 10 seconds(almost all around 8 seconds) average in each fold, and the precision of all label are the same with the value 0.97, which is a considerable value. The result reflects the Knn algorithm's benefits:

- 1. Simply and effective
- 2. Training time complexity is O(n) (has linear relationship with the size of training data set)
- 3. Not sensitive to the outlier, since we haven't used outlier remove on it.

However, the value seemed too much for a classifier.

LR performed worst among these three algorithms, with around 3 minutes running each fold, and the accuracy was all around 0.7 for each label and each fold. Since we have known that there are only 53 dimensions in the dataset, the running speed was not ideal. However, to some extent, it is also a good result for a classifier.

Random Forest also performed perfect just next to kNN algorithm with high accuracy and short running time. Only in the fifth fold there was a slight fluctuation. The average running time of Random Forest algorithm among ten folds data was 13.1 seconds as Table 7 has shown, 5 seconds slower than kNN algorithm.

It seems that kNN performed best and should be chosen. However, evaluating an algo-

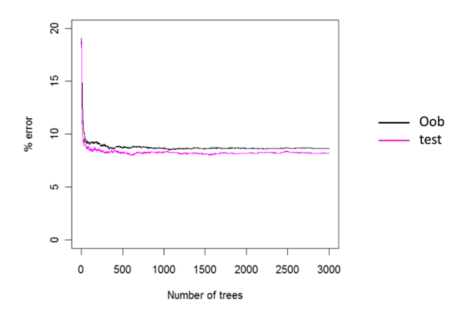
rithm for datasets should not only focus on one experiment result but also the stability of it.

As is widely known, Knn shows its weakness when the training data set is unbalance, for example the number of label 2 is much smaller than the other 6 labels. In this circumstance, when we try to classify label 2, we actually calculate the nearest label to label 2, not the label 2 itself. However, our dataset is relatively balance: there are certain amount of intersection and overlapping among datasets. So, the kNN has a good performance.

Moreover, kNN is weak at classifying large scale of data such as the data stream mentioned above. However, we have summarized that our dataset has fewer dimensions, and the scale of data and the complexity is not so large as other types if dataset. Therefore, kNN is appropriate.

For Random Forest, it has benefits that inherited many of the advantage of CART (Classification and Regression Trees) like:

- 1. Computationally simple and quick to fit, even for large data set.
- 2.No formal distributional assumption
- 3.Can balance the varieties for unbalanced data set.
- 4.No over-fitting in processing relatively big data set Just like the picture shows:



Thus, the RF may not look as efficient as KNN after running small dataset. However, as the more and more training data sets put into training, the performance of RF will gradually improved. That's the reason why RF fits for larger scale of dataset, but not the scale like

Covtype dataset.

In summary, we consider kNN is the best algorithm in dealing with this type

Reflection

After experiencing the whole process of machine learning, we find that we can do better in the following part:

As the description document explained, the dataset contains binary numbers and decimal numbers. With common sense, we should transform the different types of data into unified numbers before training the data set.

It seems that the different types of columns did not affect the result too much. Through searching relative researches, we make an explanation that when the data in high dimension space, thus the factor that different types of data will not affect each other or even the training model itself too much.

However, what if we do preprocessing with different types of columns separately? If we divide the original dataset into different parts, and how to integrate them again then clustering? We have not research about that.

CONCLUSIONS AND FUTURE WORK

In conclusion, we choose kNN as the recommended algorithm in dealing with the similar scale of dataset.

Then what can we do in the future? Here are two objectives:

1.Optimize Random Forest:

Take Boost algorithm into consideration to optimize the Random Forest method. Try to use Boosting to iterate the data set, and try to modify our new training model based on the last iteration.

2. Testing on other algorithms:

Although we have used three methods to analysis this work, we still want to use other methods like SVM, NB to training the data set. Future work will lie on analyzing the advantage of each method to determine a suitable method for this work.

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APPENDIX: HOW TO RUN

- 1. Generate 10 folds. Run *generate_samples_and_targets.py*. The generated file should be in *data* folder.
- 2. Run Python file for each algorithm. e.g. in command line, run *knn_with_orc.py*. The result should be in the *result* folder.