



Article Ensemble Learning or Deep Learning? Application to Default Risk Analysis

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Received: 19 January 2018; Accepted: 28 February 2018; Published: 5 March 2018

Abstract: Proper credit-risk management is essential for lending institutions, as substantial losses can be incurred when borrowers default. Consequently, statistical methods that can measure and analyze credit risk objectively are becoming increasingly important. This study analyzes default payment data and compares the prediction accuracy and classification ability of three ensemble-learning methods—specifically, bagging, random forest, and boosting—with those of various neural-network methods, each of which has a different activation function. The results obtained indicate that the classification ability of boosting is superior to other machine-learning methods including neural networks. It is also found that the performance of neural-network models depends on the choice of activation function, the number of middle layers, and the inclusion of dropout.

Keywords: credit risk; ensemble learning; deep learning; bagging; random forest; boosting; deep neural network

1. Introduction

Credit-risk management is essential for financial institutions whose core business is lending. Thus, accurate consumer or corporation credit assessment is of utmost importance because significant losses can be incurred by financial institutions when borrowers default. To control their losses from uncollectable accounts, financial institutions therefore need to properly assess borrowers' credit risks. Consequently, they endeavor to collate borrower data, and various statistical methods have been developed to measure and analyze credit risk objectively.

Because of its academic and practical importance, much research has been conducted on this issue. For example, Boguslauskas and Mileris (2009) analyzed credit risk using Lithuanian data for 50 cases of successful enterprises and 50 cases of bankrupted enterprises. Their results indicated that artificial neural networks are an efficient method to estimate the credit risk.

Angelini, Tollo, and Roli (Angelini et al. 2008) presented the application of an artificial neural network for credit-risk assessment using the data of 76 small businesses from a bank in Italy. They used two neural architectures to classify borrowers into two distinct classes: in bonis and default. One is a feedforward neural network and is composed of an input layer, two hidden layers and an output layer. The other is a four-layer feedforward neural network with ad hoc connections and input neurons grouped in sets of three. Their results indicate that neural networks successfully identify the in bonis/default tendency of a borrower.

Khshman (2009) developed a system of credit-risk evaluation using a neural network and applied the system to Australian credit data (690 cases; 307 creditworthy instances and 383 non-creditworthy instances). He compared the performance of the single-hidden layer neural network (SHNN) model and double-hidden layer network (DHNN). His experimental results indicated that the system with

SHNN outperformed the system with DHNN for credit-risk evaluation, and thus the SHNN neural system was recommended for the automatic processing of credit applications.

Yeh and Lien (2009) compared the predictive accuracy of probability of default among six data-mining methods (specifically, K-nearest neighbor classifier, logistic regression, discriminant analysis, naive Bayesian classifier, artificial neural networks, and classification trees) using customers' default payments data in Taiwan. Their experimental results indicated that only artificial neural networks can accurately estimate default probability.

Khashman (2010) employed neural-network models for credit-risk evaluation with German credit data comprising 1000 cases: 700 instances of creditworthy applicants and 300 instances where applicants were not creditworthy.¹ The results obtained indicated that the accuracy rates for the training data and test data were 99.25% and 73.17%, respectively. In this data, however, if one always predicts that a case is creditworthy, then the accuracy rate naturally converges to 70%. Thus, the results imply that there is only a 3.17% gain for the prediction accuracy of test data using neural network models.

Gante et al. (2015) also used German credit data and compared 12 neural-network models to assess credit risk. Their results indicated that a neural network with 20 input neurons, 10 hidden neurons, and one output neuron is a suitable neural network model for use in a credit risk evaluation system.

Khemakhem and Boujelbene (2015) compared the prediction of a neural network with that of discriminant analysis using 86 Tunisian client companies of a Tunisian commercial bank over three years. Their results indicated that a neural network outperforms discriminant analysis in predicting credit risk.

As is pointed out by Oreski et al. (2012), the majority of studies have shown that neural networks are more accurate, flexible and robust than conventional statistical methods for the assessment of credit risk.

In this study, we use 11 machine-learning methods to predict the default risk based on clients' attributes, and compare their prediction accuracy. Specifically, we employ three ensemble learning methods—bagging, random forest, and boosting—and eight neural network methods with different activation functions. The performance of each method is compared in terms of their ability to predict the default risk using multiple indicators (accuracy, rate of prediction, results, receiver operating characteristic (ROC) curve, area under the curve (AUC), and F-score).²

The results obtained indicate that the classification ability of boosting is superior to other machine-learning methods including neural networks. It is also found that the performance of neural-network models depends on the choice of activation function and the number of middle layers.

The remainder of this paper is organized as follows. Section 2 explains the data employed and the experimental design. Section 3 discusses the empirical results obtained. Section 4 presents concluding remarks.

2. Data and Experimental Design

2.1. Machine-Learning Techniques

Three ensemble-learning algorithms are employed in this study: bagging, random forest, and boosting. Bagging, developed by Breiman (1996), is a machine-learning method that uses bootstrapping to create multiple training datasets from given datasets. The classification results generated using the data are arranged and combined to improve the prediction accuracy. Because the bootstrap samples are mutually independent, learning can be carried out in parallel.

¹ The German credit dataset is publicly available at UCI Machine Learning data repository, https://archive.ics.uci.edu/ml/datasets/statlog+(german+credit+data).

² Lantz (2015) provides good explanation for machine learning methods.

Random forest, also proposed by Breiman (2001), is similar to bagging. It is a machine-learning method in which the classification results generated from multiple training datasets are arranged and combined to improve the prediction accuracy. However, whereas bagging uses all input variables to create each decision tree, random forest uses subsets that are random samplings of variables to create each decision tree. This means that random forest is better suited than bagging for the analysis of high-dimensional data.

Boosting is also a machine-learning method. Whereas bagging and random forest employ independent learning, boosting employs sequential learning (Schapire 1999; Shapire and Freund 2012). In boosting, on the basis of supervised learning, weights are successively adjusted, and multiple learning results are sought. These results are then combined and integrated to improve overall accuracy. The most widely used boosting algorithm is AdaBoost, proposed by Freund and Schapire (1996).

A neural network (NN) is a network structure comprising multiple connected units. It consists of an input layer, middle layer(s), and an output layer. The neural network configuration is determined by the manner in which the units are connected; different configurations enable a network to have different functions and characteristics. The feed-forward neural network is the most frequently used neural-network model and is configured by the hierarchical connection of multiple units. When the number of middle layers is greater than or equal to two, the network is called a deep neural network (DNN).

The activation function in a neural network is very important, as it expresses the functional relationship between the input and output in each unit. In this study, we employed two types of activation functions: Tanh and rectified linear unit (ReLU). These functions are defined as follows:

$$Tanh: f(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}}$$
$$ReLU: f(x) = \max(0, x)$$

The Tanh function compresses a real-valued number into the range [-1, 1]. Its activations saturate, and its output is zero-centered. The ReLU function is an alternative activation function in neural networks.³ One of its major benefits is the reduced likelihood of the gradient vanishing.

Although DNNs are powerful machine-learning tools, they are susceptible to overfitting. This is addressed using a technique called dropout, in which units are randomly dropped (along with their incoming and outgoing connections) in the network. This prevents units from overly co-adapting (Srivastava et al. 2014).

Thus, we use the following 11 methods to compare performance:

- 1. Bagging.
- 2. Random forest.
- 3. Boosting.
- 4. Neural network (activation function is Tanh).
- 5. Neural network (activation function is ReLU).
- 6. Neural network (activation function is Tanh with Dropout).
- 7. Neural network (activation function is ReLU with Dropout).
- 8. Deep neural network (activation function is Tanh).
- 9. Deep neural network (activation function is ReLU).
- 10. Deep neural network (activation function is Tanh with Dropout).
- 11. Deep neural network (activation function is ReLU with Dropout).

³ See LeCun et al. (2015).

2.2. Data

The payment data in Taiwan used by Yeh and Lien (2009) are employed in this study. The data are available as a default credit card client's dataset in the UCI Machine Learning Repository. In the dataset used by Yeh and Lien (2009), the number of observations is 25,000, in which 5529 observations are default payments. However, the current dataset in the UCI Machine Learning Repository has a total number of 30,000 observations, in which 6636 observations are default payments. Following Yeh and Lien (2009), we used default payment (No = 0, Yes = 1) as the explained variable and the following 23 variables as explanatory variables:

X1: Amount of given credit (NT dollar).

X2: Gender (1 = male; 2 = female).

X3: Education (1 = graduate school; 2 = university; 3 = high school; 4 = others).

X4: Marital status (1 = married; 2 = single; 3 = others).

X5: Age (year).

X6–X11: History of past payment tracked via past monthly payment records (-1 = payment on time; 1 = payment delay for one month; 2 = payment delay for two months; ...; 8 = payment delay for eight months; 9 = payment delay for nine months and above).

X6: Repayment status in September 2005.

X7: Repayment status in August 2005.

X8: Repayment status in July 2005.

X9: Repayment status in June 2005.

X10: Repayment status in May 2005.

X11: Repayment status in April 2005.

X12: Amount on bill statement in September 2005 (NT dollar).

X13: Amount on bill statement in August 2005 (NT dollar).

X14: Amount on bill statement in July 2005 (NT dollar).

X15: Amount on bill statement in June 2005 (NT dollar).

X16: Amount on bill statement in May 2005 (NT dollar).

X17: Amount on bill statement in April 2005 (NT dollar).

X18: Amount of previous payment in September 2005 (NT dollar).

X19: Amount of previous payment in August 2005 (NT dollar).

X20: Amount of previous payment in July 2005 (NT dollar).

X21: Amount of previous payment in June 2005 (NT dollar).

X22: Amount of previous payment in May 2005 (NT dollar).

X23: Amount of previous payment in April 2005 (NT dollar).

Because of the high proportions of no-default observations (77.88%), the accuracy rate inevitably remains at virtually 78% when all observations are used for analysis. It is difficult to understand the merit of using machine learning if we use all data. Thus, in this study we extracted 6636 observations randomly from all no-default observations to ensure that no-default and default observations are equal, thereby preventing distortion. As regards the ratio of training to test datasets, this study uses two cases, i.e., 90% to 10% and 75% to 25%.⁴

It is well known that data normalization can improve performance. Classifiers are required to calculate the objective function, which is the mean squared error between the predicted value

⁴ There are two typical ways to implement machine learning. One is to use training data, validation data, and test data, and the other is to use training data and test data. In the first approach, the result of the test is randomly determined and we cannot obtain robust results. Also, it is not advisable to divide the small sample into three pieces. Thus, we use the second approach in this study. We repeat the test results over 100 times to obtain robust results.

and the observation. If some of the features have a broad range of values, the mean squared error may be governed by these particular features and objective functions may not work properly. Thus, it is desirable to normalize the range of all features so that each feature equally contributes to the cost function (Aksoy and Haralick 2001). Sola and Sevilla (1997) point out that data normalization prior to neural network training enables researchers to speed up the calculations and to obtain good results. Jayalakshmi and Santhakumaran (2011) point out that statistical normalization techniques enhance the reliability of feed-forward backpropagation neural networks and the performance of the data-classification model.

Following Khashman (2010), we normalize the data based on the following formula:

$$z_i = \frac{x_i - x_{\min}}{x_{\max} - x_{\min}}$$

where z_i is normalized data, x_i is each dataset, x_{min} is the minimum value of x_i , and x_{max} is the maximum value of x_i . This method rescales the range of features to between 0 and 1. We analyze both normalized and original data in order to evaluate the robustness of our experimental results.

2.3. Performance Evaluation

We use accuracy to evaluate the performance of each machine-learning method. In our two-class problem, the confusion matrix (Table 1) gives us a summary of prediction results on a classification problem as follows:

Table 1. Confusion matrix

		Actual Class		
		Event	No-Event	
Predicted Class	Event No-Event	TP (True Positive) FN (False Negative)	FP (False Positive) TN (True Negative)	

Note that "true positive" indicates the case for correctly predicted event values; "false positive" indicates the case for incorrectly predicted event values; "true negative" indicates the case for correctly predicted no-event values: and "false negative" indicates the case for incorrectly predicted no-event values. Then, prediction accuracy rate is defined by,

prediction accuracy rate =
$$\frac{TP + TN}{TP + FP + FN + TN}$$

Furthermore, we repeat the experiments 100 times and calculate the average and standard deviation of the accuracy rate for each dataset.⁵

Next, we analyzed the classification ability of each method by examining the ROC curve and the AUC value. When considering whether a model is appropriate, it is not sufficient to rely solely on accuracy rate. The ratio of correctly identified instances in the given class is called the true positive rate. The ratio of incorrectly identified instances in the given class is called the false positive rate. When the false positive rate is plotted on the horizontal axis and the true positive rate on the vertical axis, the combination of these produces an ROC curve. A good model is one that shows a high true positive rate value and low false positive value. The AUC refers to the area under the ROC curve. A perfectly random prediction yields an AUC of 0.5. In other words, the ROC curve is a straight line connecting the origin (0, 0) and the point (1, 1).

⁵ We used set. seed(50) to remove the difference caused by random numbers in drawing the ROC curve and calculating the AUC.

We also report the F-score of each case, which is defined as follows:

$$F - score = \frac{2 \times recall \times precision}{recall + precision}$$

where *recall* is equal to TP/(TP + FN) and *precision* is equal to TP/(TP +FP). Thus, the F-score is the harmonic average of recall and precision.

3. Results

We implement the experiments using R—specifically, the "ipred" package for bagging, "randomForest" for random forest, "ada" package for boosting (adaboost algorithm), and "h2o" package for NN and DNN. Furthermore, we analyze the prediction accuracy rate of each method for two cases i.e., original and normalized data. Then, we examine the classification ability of each method based on the ROC curve, AUC value, and F-score.

Table 2a,b report the results obtained using the original data. The tables show that boosting has the best performance and yields higher than 70% prediction accuracy rate on average, with a small standard deviation for both training and test data. None of the neural network models exceed a 70% average accuracy rate for test data. Furthermore, they have relatively large standard deviation for test data. Thus, it is clear that boosting achieves a higher accuracy prediction than neural networks. The prediction accuracy rate for test data is less than 60% for bagging and random forest. In addition, the difference of ratios between training and test data (90%:10% or 75%:25%) does not have an obvious influence on the results of our analysis.⁶

Table 3a,b summarize the results obtained using normalized data. The tables show that boosting has the highest accuracy rate on test data, which is similar to the results obtained for the original data case. The average accuracy rate for boosting is more than 70% and it has the smallest standard deviation for both training and test data. None of the neural network models has an average prediction accuracy rate exceeding 70% for test data. Furthermore, they have relatively large standard deviation for test data. The prediction accuracy rate of bagging and random forest does not reach 60% on average for test date, which is similar to the case for the original data. In addition, the difference of ratios between training and test data (90%:10% or 75%:25%) does not have a major influence on the result, which is similar to the case with the original data. Our comparison of the results of the original data with the results of the normalized data reveals no significant difference in prediction accuracy rate.

Figures 1–11 display ROC curves with AUC and F-score for the case using normalized data and the ratio between the training and test data of 75% to 25%. In each figure, sensitivity (vertical axis) corresponds to the true positive ratio, whereas 1—specificity (horizontal axis) corresponds to the false positive ratio. The graphs indicate that the ROC curve for boosting and neural network models have desirable properties except for the case for the Tanh activation function with dropout.

The AUC values and F-score are also shown for each figure. It is found that the highest AUC value is obtained for boosting (0.769). The highest F-score is also obtained for boosting (0.744). Thus, the classification ability of boosting is superior to other machine-learning methods. This may be because boosting employs sequential learning of weights.

It is also found that the AUC value and F-score of NN are better than those of DNN when Tanh is used as an activation function. However, this result is not apparent when ReLU is used as an activation function. It is interesting to see the results of neural-network models with respect to the influence of dropout in terms of AUC value and F-score. When Tanh is used as an activation function, NN (DNN) outperforms NN (DNN) with dropout. On the other hand, when ReLU is used as an activation function, NN (DNN) with dropout outperform NN (DNN). Thus the performance of neural networks

⁶ The number of units in the middle layers of NN and DNN is determined based on the Bayesian optimization method. (See Appendix A for details.)

may be sensitive to the model setting i.e., the number of middle layers, the type of activation function, and inclusion of dropout.

		0		ng and test data i		
Method		Accuracy Ratio of	Training Data	Accuracy Ratio of Test Data		
			Average (%)	Standard Deviation	Average (%)	Standard Deviation
	Bagging		80.13	0.003	55.98	0.008
	Boosting		71.66	0.003	71.06	0.008
Random Forest		69.59	0.544	58.50	0.844	
Method		Accuracy Ratio of Training Data		Accuracy Ra	Accuracy Ratio of Test Data	
Model	Activation Function	Middle Layer	Average (%)	Standard Deviation	Average (%)	Standard Deviation
DNN	Tanh	2	70.66	0.721	68.93	0.972
NN	Tanh	1	71.01	0.569	69.59	0.778
DNN	Tanh with Dropout	2	58.47	3.566	58.46	3.404
NN	Tanh with Dropout	1	67.27	1.237	67.14	1.341
DNN	ReLU	2	69.57	0.707	68.61	0.863
NN	ReLU	1	68.81	0.708	68.30	1.008
DNN	ReLU with Dropout	2	69.97	0.903	69.01	0.956
NN	ReLU with Dropout	1	70.12	0.637	69.48	0.881
	(b) (Driginal Data	a: the Ratio of Trainin	ng and Test Data	is 90% to 10%	
Method		Accuracy Ratio of Training Data		Accuracy Ratio of Test Data		
	Wiethou		Average (%)	Standard Deviation	Average (%)	Standard Deviation
	Bagging		79.58	0.003	56.23	0.015
	Boosting		71.57	0.003	70.88	0.011
	Random Forest		68.55	0.453	58.77	1.331
	Method		Accuracy Ratio of	Accuracy Ratio of Training Data		tio of Test Data
Model	Activation Function	Middle Layer	Average (%)	Standard Deviation	Average (%)	Standard Deviation
DNN	Tanh	2	69.64	0.683	69.31	1.325
NN	Tanh	1	70.49	0.550	69.61	1.312
DNN	Tanh with Dropout	2	57.29	3.681	57.27	4.117
NN	Tanh with Dropout	1	66.37	1.619	66.25	1.951
DNN	ReLU	2	69.49	0.695	68.76	1.408
NN	ReLU	1	69.16	0.728	68.54	1.261
DNN	ReLU with Dropout	2	69.74	0.796	68.84	1.438
NN	ReLU with Dropout	1	70.26	0.573	69.55	1.210

Table 2. Prediction accuracy of	each method for original data.
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Method		Accuracy Ratio of	Training Data	Accuracy Ratio of Test Data		
	Method		Average (%)	Standard Deviation	Average (%)	Standard Deviation
	Bagging		80.12	0.003	56.15	0.008
	Boosting		71.66	0.004	70.95	0.007
Random Forest		69.67	0.565	58.39	0.880	
	Method		Accuracy Ratio of Training Data		Accuracy Ratio of Test Data	
Model	Activation Function	Middle Layer	Average (%)	Standard Deviation	Average (%)	Standard Deviation
DNN	Tanh	2	71.14	0.732	68.75	0.912
NN	Tanh	1	70.64	0.652	69.42	0.763
DNN	Tanh with Dropout	2	57.00	4.324	56.69	4.485
NN	Tanh with Dropout	1	68.09	0.641	68.01	0.904
DNN	ReLU	2	70.37	0.627	69.35	0.856
NN	ReLU	1	70.92	0.615	69.37	0.943
DNN	ReLU with Dropout	2	70.00	0.811	68.96	0.946
NN	ReLU with Dropout	1	70.25	0.692	69.56	0.813
	(b) N	lormalized d	lata: the ratio of train	ing and test data	a is 90% to 10%	
Method		Accuracy Ratio of Training Data		Accuracy Ra	tio of Test Data	
			Average (%)	Standard Deviation	Average (%)	Standard Deviation
	Bagging		79.54	0.003	56.28	0.013
	Boosting		71.50	0.003	70.80	0.012
	Random Forest		68.66	0.475	58.83	1.368
	Method		Accuracy Ratio of Training Data		Accuracy Ratio of Test Data	
Model	Activation Function	Middle Layer	Average (%)	Standard Deviation	Average (%)	Standard Deviation
DNN	Tanh	2	70.18	0.698	69.35	1.382
NN	Tanh	1	70.52	0.594	69.51	1.309
DNN	Tanh with Dropout	2	58.04	5.134	58.14	5.016
NN	Tanh with Dropout	1	67.33	1.285	67.13	1.787
DNN	ReLU	2	71.41	0.710	69.17	1.334
NN	ReLU	1	69.55	0.772	68.97	1.426
	ReLU with	2	69.76	0.785	69.13	1.426
DNN	Dropout ReLU with					

 Table 3. Prediction accuracy of each method for normalized data.

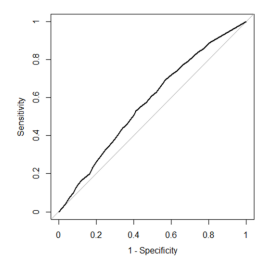


Figure 1. Receiver operating characteristic (ROC) curve for bagging. (Area under the curve (AUC) = 0.575, F-score = 0.520).

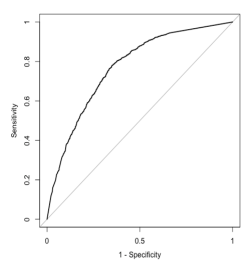


Figure 2. ROC curve for boosting. (AUC = 0.769, F-score = 0.744).

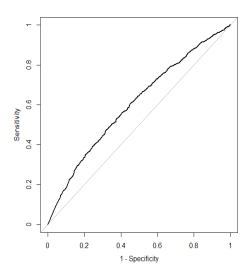


Figure 3. ROC curve for random forest. (AUC = 0.605, F-score = 0.714).

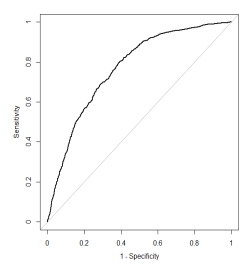


Figure 4. ROC curve for deep neural network (DNN) (Tanh). (AUC = 0.753, F-score = 0.721).

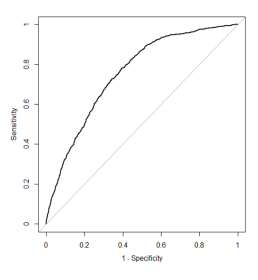


Figure 5. ROC curve for neural network (NN) (Tanh). (AUC = 0.768, F-score = 0.741).

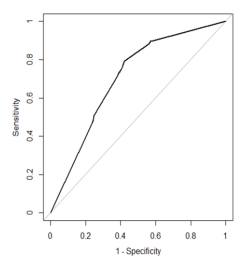


Figure 6. ROC curve for DNN (Tanh w/Dropout). (AUC = 0.600, F-score = 0.620).

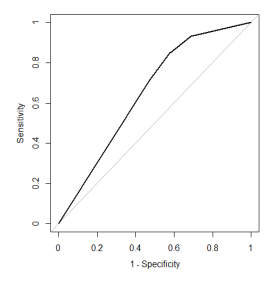


Figure 7. ROC curve for NN (Tanh w/Dropout). (AUC = 0.704, F-score = 0.717).

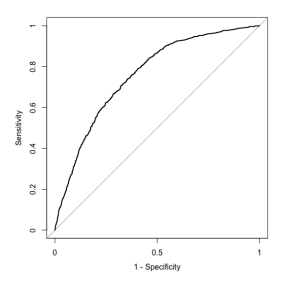


Figure 8. ROC curve for DNN (ReLU). (AUC = 0.751, F-score = 0.734).

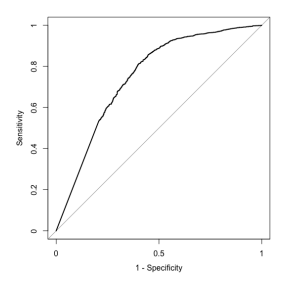


Figure 9. ROC curve for NN (ReLU). (AUC = 0.757, F-score = 0.727).

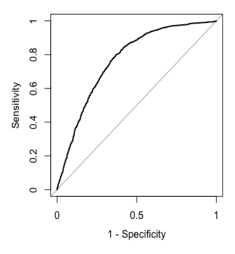


Figure 10. ROC curve for DNN (ReLU w/Dropout). (AUC = 0.765, F-score = 0.735).

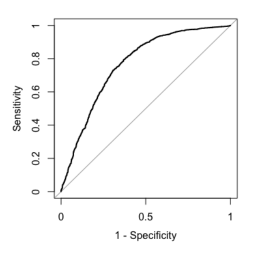


Figure 11. ROC curve for NN (ReLU w/Dropout). (AUC = 0.767, F-score = 0.730).

4. Conclusions

In this study, we analyzed default payment data in Taiwan and compared the prediction accuracy and classification ability of three ensemble-learning methods: bagging, random forest, and boosting, with those of various neural-network methods using two different activation functions. Our main results can be summarized as follows:

- (1) The classification ability of boosting is superior to other machine-learning methods.
- (2) The prediction accuracy rate, AUC value, and F-score of NN are better than those of DNN when Tanh is used as an activation function. However, this result is not apparent when ReLU is used as an activation function.
- (3) NN (DNN) outperforms NN (DNN) with dropout when Tanh is used as an activation function in terms of AUC value and F-score. However, NN (DNN) with dropout outperforms NN (DNN) when ReLU is used as an activation function in terms of AUC value and F-score.

The usability of deep learning has recently been the focus of much attention. Oreski et al. (2012) point out that the majority of studies show that neural networks are more accurate, flexible, and robust than conventional statistical methods when assessing credit risk. However, our results indicate that boosting outperforms the neural network in terms of prediction accuracy, AUC, and F-score. It is also well known that it is not easy to choose appropriate hyper-parameters for neural networks. Thus, neural networks are not always a panacea, especially for relatively small samples. Given this, it is

worthwhile to make effective use of other methods such as boosting. Our future work will be to apply a similar analysis to different data in order to check the robustness of our results.

Acknowledgments: We are grateful to the three anonymous referees for their helpful comments and suggestions. An early version of this paper was read at the Workshop of Big Data and Machine Learning. We are grateful to Zheng Zhang and Xiao Jing Cai for helpful comments and suggestions. This research was supported by a grant-in-aid from The Nihon Hoseigakkai Foundation.

Author Contributions: Shigeyuki Hamori conceived and designed the experiments; Minami Kawai, Takahiro Kume, Yuji Murakami and Chikara Watanabe performed the experiments, analyzed the data, and contributed reagents/materials/analysis tools; and Shigeyuki Hamori, Minami Kawai, Takahiro Kume, Yuji Murakami and Chikara Watanabe wrote the paper.

Conflicts of Interest: The authors declare no conflicts of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

Appendix Results of Bayesian Optimization

Method	Data	Ratio of Training and Test Data (%)	Input Layer	Middle Layer	Output Layer
Tanh	Original	75:25	23	7	2
Tanh	Original	90:10	23	5	2
Tanh with Dropout	Original	75:25	23	14	2
Tanh with Dropout	Original	90:10	23	12	2
ReLU	Original	75:25	23	3	2
ReLU	Original	90:10	23	7	2
ReLU with Dropout	Original	75:25	23	14	2
ReLU with Dropout	Original	90:10	23	19	2
Tanh	Normalized	75:25	23	5	2
Tanh	Normalized	90:10	23	5	2
Tanh with Dropout	Normalized	75:25	23	5	2
Tanh with Dropout	Normalized	90:10	23	10	2
ReLU	Normalized	75:25	23	11	2
ReLU	Normalized	90:10	23	4	2
ReLU with Dropout	Normalized	75:25	23	16	2
ReLU with Dropout	Normalized	90:10	23	12	2

Table A1.	Number	of units in	n middle la	yer for NN.
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Table A2. Number of units in middle layers for DNN.

Method	Data	Ratio of Training and Test Data (%)	Input Layer	Middle Layer 1	Middle Layer 2	Output Layer
Tanh	Original	75:25	23	5	17	2
Tanh	Original	90:10	23	2	9	2
Tanh with Dropout	Original	75:25	23	9	7	2
Tanh with Dropout	Original	90:10	23	3	11	2
ReLU	Original	75:25	23	4	6	2
ReLU	Original	90:10	23	4	9	2
ReLU with Dropout	Original	75:25	23	13	9	2
ReLU with Dropout	Original	90:10	23	5	20	2
Tanh	Normalized	75:25	23	6	17	2
Tanh	Normalized	90:10	23	4	3	2
Tanh with Dropout	Normalized	75:25	23	9	4	2
Tanh with Dropout	Normalized	90:10	23	3	18	2
ReLU	Normalized	75:25	23	4	6	2
ReLU	Normalized	90:10	23	10	7	2
ReLU with Dropout	Normalized	75:25	23	16	9	2
ReLU with Dropout	Normalized	90:10	23	5	21	2

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