Software Defect Prediction Using Variant based Ensemble Learning and Feature Selection Techniques

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Abstract: Testing is considered as one of the expensive activities in software development process. Fixing the defects during testing process can increase the cost as well as the completion time of the project. Cost of testing process can be reduced by identifying the defective modules during the development (before testing) stage. This process is known as “Software Defect Prediction”, which has been widely focused by many researchers in the last two decades. This research proposes a classification framework for the prediction of defective modules using variant based ensemble learning and feature selection techniques. Variant selection activity identifies the best optimized versions of classification techniques so that their ensemble can achieve high performance whereas feature selection is performed to get rid of such features which do not participate in classification and become the cause of lower performance. The proposed framework is implemented on four cleaned NASA datasets from MDP repository and evaluated by using three performance measures, including: F-measure, Accuracy, and MCC. According to results, the proposed framework outperformed 10 widely used supervised classification techniques, including: “Naïve Bayes (NB), Multi-Layer Perceptron (MLP), Radial Basis Function (RBF), Support Vector Machine (SVM), K Nearest Neighbor (KNN), kStar (K*), One Rule (OneR), PART, Decision Tree (DT), and Random Forest (RF)”.

Index Terms: Software Defect Prediction, Feature Selection, Classifier Variant, Ensemble Learning, Machine Learning Techniques

1. Introduction

Software testing is an important activity of quality assurance process, which ensures the delivery of qualitative product without any defects. As compared to other activities of software development life cycle, testing is the most expensive one as it consumes more resources than others [8,9,10]. This activity makes sure that all of the developed modules are bug free [6,7]. To bring down the overall development cost by keeping the quality intact, is a major issue for software development industry. The prediction of defective modules on the basis of historical development data can resolve this issue. In this procedure, the software modules which are more likely to be defective are identified before the testing activity. Due to which only those software modules are tested which are predicted as defective instead of all. With this approach, the cost of testing activity can be significantly reduced by keeping the quality intact [10,11,12]. The prediction of software defects is a binary classification problem as we have to identify that the particular module is defective or non-defective. Many researchers have focused on machine learning techniques to solve the binary classification problems such as: Network Intrusion Detection [21,22], Sentiment Analysis [13,14,15,16,17,18], Rainfall Prediction [19,20], and Software Defect Prediction [1,2,3,4,5]. The process of software defect prediction has been focused by many researchers in the last decade however improving the prediction accuracy has always been the main concern. This research proposes a classification framework for software defect prediction using variant based ensemble learning and feature selection techniques. There are five stages of the proposed framework: 1) Dataset Selection, 2) Variant Selection, 3) Pre-processing & Feature Selection, 4) Classification, and 5) Reflection of Results. The proposed framework is implemented on four of the cleaned datasets from NASA MDP repository including: JM1, KC1, PC4 and PC5. Moreover, three accuracy measure including: F measure, Accuracy and MCC are used for performance analysis. The performance of the proposed framework is compared with various supervised classifiers from a published research [1], which have used the same datasets and accuracy measures for performance analysis. The classifiers include: “Naïve Bayes (NB), Multi-Layer Perceptron (MLP), Radial Basis Function (RBF), Support Vector Machine (SVM), K
Nearest Neighbor (KNN), kStar (K*), One Rule (OneR), PART, Decision Tree (DT), and Random Forest (RF)”.

Results show that the proposed framework performed better than all of other classifiers from [1] in all of the three accuracy measures.

2. Related Work

Many researchers have focused on the use of machine learning techniques in order to predict the software defects before the testing stage by using the historical data (data of previously developed modules). The historical data consists of various software metrics collected during the development. This section discusses some recent studies, conducted on software defect prediction. All of the researchers which are going to be discussed in this section used cleaned version of datasets (D’”) from NASA MDP repository. Researchers in [1] performed a detailed performance analysis of various supervised classification techniques on software defect prediction. The techniques include: “Naïve Bayes (NB), Multi-Layer Perceptron (MLP), Radial Basis Function (RBF), Support Vector Machine (SVM), K Nearest Neighbor (KNN), kStar (K*), One Rule (OneR), PART, Decision Tree (DT), and Random Forest (RF)”.

The default parameters of the classification techniques are used and performance was evaluated by using six measures including: “Precision, Recall, F-Measure, Accuracy, MCC, and ROC Area”. The authors presented the results as a baseline for other studies so that any improvement in the accuracy by any proposed technique can be verified by comparing with the widely used base classifiers. Researchers in [2] presented a framework for software defect prediction. Feature selection and ensemble learning techniques are used to improve the performance of prediction. Two different dimensions are implemented in the framework, in one dimension feature selection is used and in second dimension feature selection activity is not performed. This process is done to identify the effect of feature selection process on the performance. Performance evaluation is performed through six measures such as: “Precision, Recall, F-Measure, Accuracy, MCC, and ROC Area”.

The results obtained from both of the dimensions are compared with each other. The results are also compared with other well-known and widely used classification techniques from [1] which have used the same datasets and performance measures. Researchers in [3] discussed the issue of imbalanced datasets and used three well known and widely used resampling techniques to resolve this problem during the process of software defect prediction. The used techniques include: “Random Under Sampling”, “Random Over Sampling” and “Synthetic Minority Oversampling Technique (SMOTE)”.

For classification, various widely used machine learning techniques are used and effects of the resampling techniques on the performance is analyzed by using four measures such as: “F-measure, Accuracy, MCC and ROC Area”. Researchers in [4] presented a framework to predict the defects at early stages of software development. To increase the performance of prediction, the researchers incorporated feature selection and ensemble learning techniques and for performance evaluation four measures are used: F-measure, Accuracy, MCC and ROC. Various feature selection techniques are used in the experiment. In the results, all feature selections techniques are compared with each other in all of the used performance measures. Then a detailed comparison is performed with various widely used supervised classification techniques from research [1]. Researchers in [5] proposed a classification framework to predict the defective modules before the testing. They used a multi filter feature section technique with an aggregated method and Artificial Neural Network (MLP). The framework is implemented with two different dimensions: first with oversampling technique and second without oversampling technique. The purpose behind introducing the oversampling technique in the framework is to analyze the effect of resampling on the performance of proposed classification framework. Performance evaluation is performed by using four accuracy measures: F-measure, Accuracy, MCC and ROC.

3. Materials and Methods

This paper contributes by proposing a classification framework for software defect prediction using variant based ensemble learning and feature selection techniques. The proposed framework (Figure 1) consists of five stages: 1) Dataset Selection, 2) Variants Selection, 3) Data Preprocessing & Feature selection, 4) Classification, 5) Reflection of results. WEKA tool is used for the implementation of proposed classification framework.
Selection of appropriate dataset is the first stage of the proposed framework. In this study we have selected four widely used cleaned datasets from NASA MDP repository including: JM1, KC1, PC4 and PC5 (Table 1). These datasets reflect the module wise metrics of software systems of NASA (Table 2) and their testing results that either these modules are defective or not (Fig. 2.). Each dataset consists of various records whereas each record represents a module in the form of an attribute set. The attributes are the software metrics which are generated during the development (Table 2).

Table 1. NASA Cleaned Datasets (D") [23]

| Dataset | Attributes | Modules | Defective | Non-Defective | Defective (%) |
|---------|------------|---------|-----------|---------------|---------------|
| JM1     | 22         | 7,720   | 1,612     | 6,108         | 20.8          |
| KC1     | 22         | 1,162   | 294       | 868           | 25.3          |
| PC4     | 38         | 1,270   | 176       | 1094          | 13.8          |
| PC5     | 39         | 1,694   | 458       | 1236          | 27.0          |
Each of the used datasets contains various independent attributes, and only one dependent attribute. The dependent attribute is the one which is going to be predicted, also known as target class and the independent attributes are those which are used to predict the dependent attribute. The independent attributes of the used datasets are mentioned in Table 2. The target class (dependent attribute) can contain only one from either of two values: ‘Y’ or ‘N’. ‘Y’ shows that the module is defective and ‘N’ reflects that the module is non-defective (Fig 2). Researchers in [23], provided two versions of cleaned NASA datasets: DS’ and DS’’. The instances in DS’ included duplicate and inconsistent values however DS’’ contains non-duplicate and consistent instances. These datasets are currently available at [24]. In this research we have used the DS’’ version of NASA datasets which is already been used by many researchers [1,2,3,4,5,25,26,27].

Table 2. Independent Attributes of Datasets

| Sr. # | Attributes                | JM1 | KC1 | PC4 | PC5 |
|-------|--------------------------|-----|-----|-----|-----|
| 1     | LOC_BLANK                | ✓   | ✓   | ✓   | ✓   |
| 2     | BRANCH_COUNT             | ✓   | ✓   | ✓   | ✓   |
| 3     | CALL_PAIRS               |     |     | ✓   | ✓   |
| 4     | LOC_CODE_AND_COMMENT     | ✓   | ✓   | ✓   | ✓   |
| 5     | LOC_COMMENTS             | ✓   | ✓   | ✓   | ✓   |
| 6     | CONDITION_COUNT          |     |     | ✓   | ✓   |
| 7     | CYCLOMATIC_COMPLEXITY    | ✓   | ✓   | ✓   | ✓   |
| 8     | CYCLOMATIC_DENSITY       | ✓   | ✓   | ✓   | ✓   |
| 9     | DECISION_COUNT           |     |     | ✓   | ✓   |
| 10    | DECISION_DENSITY         |     |     | ✓   | ✓   |
| 11    | DESIGN_COMPLEXITY        | ✓   | ✓   | ✓   | ✓   |
| 12    | DESIGN_DENSITY           | ✓   | ✓   | ✓   | ✓   |
| 13    | EDGE_COUNT               | ✓   | ✓   | ✓   | ✓   |
| 14    | ESSENTIAL_COMPLEXITY     | ✓   | ✓   | ✓   | ✓   |
| 15    | ESSENTIAL_DENSITY        | ✓   | ✓   | ✓   | ✓   |
| 16    | LOC_EXECUTABLE           | ✓   | ✓   | ✓   | ✓   |
| 17    | PARAMETER_COUNT          | ✓   | ✓   | ✓   | ✓   |
| 18    | GLOBAL_DATA_COMPLEXITY   | ✓   | ✓   | ✓   | ✓   |
| 19    | GLOBAL_DATA_DENSITY      | ✓   | ✓   | ✓   | ✓   |
| 20    | HALSTEAD_COMPUTATION     | ✓   | ✓   | ✓   | ✓   |
| 21    | HALSTEAD_DIFFICULTY      | ✓   | ✓   | ✓   | ✓   |
| 22    | HALSTEAD_EFFORT          | ✓   | ✓   | ✓   | ✓   |
| 23    | HALSTEAD_ERROR_EST        | ✓   | ✓   | ✓   | ✓   |
| 24    | HALSTEAD_LENGTH          | ✓   | ✓   | ✓   | ✓   |
Second stage of the framework deals with the selection of best variants from different classifiers (Fig.3). Six classifiers are optimized (tuned) to create the variants. The classifiers include: Support Vector Machine (SVM), Decision Tree (DT), K-Nearest Neighbor (KNN), Naive Bayes (NB), Random Forest (RF), and Multi-layer Perceptron (MLP). First, these classifiers are applied on all the datasets to predict the defective modules with default parameters (without tuning), and then different variants from each of these classifiers are created by optimizing their parameters. All of these variants are then used to predict the defective modules and those variants are selected for ensembles which have higher accuracy on all of the datasets. Only one variant is selected from one family (family included base classifier and its variants), suppose if four different variants of MLP are created then only one variant would be selected which performs higher than its own base classifier (with default parameters). If more than one variants perform better than base classifier then the one with the highest performance will be selected. Multiple variants are created of each base classifier. The first variant of each classifier is the base classifier itself with its default parameters. The accuracy of each of the later variants is compared with the first variant (base classifier) as well as with other variants within the family on each of the dataset. If any of the variant within the family cannot perform well than base classifier then no variant will be selected from that family.
Two variants of Naive Bayes (NB) are created (Table 3) by optimizing two parameters (“UseKernelEstimator” and “UseSupervisedDiscretization”). In first variant the default parameters are used, in which both the parameters are not selected whereas in later variants both the parameters are selected one by one.

Table 3. Variants of Naive Bayes

| Optimized parameter | NB-1 (Default) | NB-2 | NB-3 |
|---------------------|----------------|------|------|
| UseKernelEstimator / UseSupervisedDiscretization | None-selected | UseKernelEstimator | UseSupervisedDiscretization |

For Support Vector Machine (SVM), three variants are created (Table 4). The value of complexity parameter “C” is considered for optimizing, which defines hyper-plan and controls the misclassification. In the case of small value of C, a hyper-plan with large margin is defined which results in high misclassification rate in training data. If the value of C is large then it defines a hyper-plan with small margin which gives better results and classify maximum points correctly.

Table 4. Variants of Support Vector Machine

| Optimized parameter | SVM-1 (Default) | SVM-2 | SVM-3 | SVM-4 |
|---------------------|----------------|-------|-------|-------|
| C                   | 1              | 10    | 25    | 50    |

In Multi-layer Perceptron (MLP), the parameter “h” is optimized, which defines the two things 1) no of hidden layers, and 2) no of neurons in each hidden layer. The default value is “a” which defines one hidden layer, and the no of neurons in the hidden layer are decided by the following formula:

$$a = \frac{(attributes + Classes)}{2}$$

Only one hidden layer is used in all variants of MLP with different no of neurons as shown in Table 5.

Table 5. Variants of Multi-Layer Perceptron

| Optimized parameter | MLP-1 (Default) | MLP-2 | MLP-3 | MLP-4 | MLP-5 |
|---------------------|----------------|-------|-------|-------|-------|
| h                   | a              | 3     | 5     | 7     | 9     |

In Random Forest (RF), the parameter of “max depth” is optimized. By default RF has value 0 however two variants are created by changing its number (Table 6).

Table 6. Variants of Random forest

| Optimized parameter | RF-1 (Default) | RF-2 | RF-3 |
|---------------------|----------------|------|------|
| MaxDepth            | 0              | 10   | 15   |

During the optimization of K-Nearest Neighbor, three variants are created by optimizing the number of neighbors (Table 7).

Table 7. Variants of K-nearest Neighbours

| Optimized parameter | KNN-1 (Default) | KNN-2 | KNN-3 | KNN-4 |
|---------------------|----------------|-------|-------|-------|
| No of Neighbours    | 1              | 3     | 5     | 7     |

From Decision Tree (DT), 4 variants are created by tuning the parameter of confidence factor which controls the pruning (Table 8).
Table 8. Variants of Decision Tree

| Optimized Parameter | DT-1 (Default) | DT-2 | DT-3 | DT-4 | DT-5 |
|---------------------|----------------|------|------|------|------|
| Confidence Factor   | 0.25           | 0.20 | 0.15 | 0.10 | 0.05 |

All variants including the base classifiers (with default parameters) are used to classify the datasets with training-testing split ratio of 70:30. After the performance analysis on accuracy measure only three variants are selected: SVM-4, RF-3 and KNN-4, as these variants showed higher performance than the base classifiers as well as other variants within their family. The detailed results (accuracy of the selected variants) will be discussed in next section.

Third stage of proposed framework deals with two activities Data preprocessing and Feature Selection. In Data preprocessing, two tasks are performed: Resampling [31, 32] and Randomization. Resampling is performed to resolve the issue of class imbalance in the datasets as this issue can compromise the accuracy of proposed classification framework [2,3,4,5]. To perform this task, the buildin function of WEKA is used (weka.filters.supervised.instance.Resample). On the other hand, the randomization technique shuffles the instances of datasets. This process is also performed by using a builtin function of WEKA tool (weka.filters.unsupervised.instance.Randomize).

Feature Selection activity is applied on all the datasets in order to select only those features which highly participate in the classification process. As, it has been proved now from the studies [2], [5] that those features should be removed from the dataset which do not participate in classification process as these features may reduce the performance. In this research, feature selection is performed by “Cfs Subset Evaluator” [28, 29, 30] with BestFirst search method, whereas full dataset is given for training. In this approach, three directions are used for feature subset selection: Forward, Backward, and Bi-Directional. In result, three subsets are generated for each dataset. To choose the best subset, those attributes are selected which are common in each direction as shown in Table 9.

Table 9. Feature Subsets

| BestFirst (Direction) | JM1        | KC1        | PC4        | PC5        |
|-----------------------|------------|------------|------------|------------|
|                       | 1,3,4,6,7  | 1,3,8,9,10,14,19,20 | 4,8,17,.26 | 3,4,10,.13,17,.19,20,.21,35 |
| Forward               | .8,9,11,2  | 9,21        | 1,3,8,9,10,14,19,20 | 4,8,17,.26 | 3,4,13,.17,19,.20,25,.35 |
| Backward              | .8,9,16,2  | .21        | 1,3,8,9,10,14,19,20 | 4,8,17,.26 | 3,4,13,.17,19,.20,21,.35 |
| Bi-Directional        | .8,9,16,2  | .21        | 1,3,8,9,10,14,19,20 | 4,8,17,.26 | 3,4,13,.17,19,.20,35   |
| Common Features       | 1,3,4,6,7  | .8,9,20,21 | 1,3,8,9,10,14,19,20 | 4,8,17,.26 | 3,4,13,.17,19,.20,35   |

Classification is the fourth stage of the proposed framework. In this stage the ensembles are created by using three variants which are selected in second stage of framework. Purpose of ensemble learning is to combine the power of multiple classifiers in one classification model. Many ensemble techniques are available today such as Bagging, Boosting, Voting and Stacking etc. In this research, voting is selected to create ensembles. In this technique multiple sub models are created and each models performs its own prediction and then those predictions are combined on the basis of voting. Three variants selected in second stage including: SVM-4, RF-3 and KNN-4 are used to create various ensembles by using Voting technique. Ensemble with all possible combinations of selected variants are used one by one for classification using the subsets selected in third stage, and finally one ensemble having the combination of RF-3 and KNN-4 is chosen as it performed highest among all.
4. Results and Discussion

Fifth and the last stage of the proposed framework deals with the presentation and analysis of results. Three measures are used for the performance evaluation of proposed framework including: F-measure, Accuracy, and MCC. All of these performance measures are based on confusion matrix (Fig. 4).

![Confusion Matrix](image)

A confusion matrix is consists of four parameters: TP, FP, FN, and TN. These parameters are explained below:

- **TP** (True Positive): “Instances which are actually positive and also classified as positive”.
- **FP** (False Positive): “Instances which are actually negative but classified as positive”.
- **FN** (False Negative): “Instances which are actually positive but classified as negative”.
- **TN** (True Negative): “Instances which are actually negative and also classified as negative”.

All of three performance measures: F-measure, Accuracy, and MCC are calculated by using the parameters of confusion matrix. The brief description along with the calculation formula of each of the used performance measure is given below:

‘F-measure’ is one of the widely used measures to analyze the performance of classification techniques. To calculate this measure, two measures named ‘Precision’ and ‘Recall’ have to be calculated as F-measure is the average of these both measures.

**Precision** represents the ratio between True Positive instances and the instances which are classified as positive (True Positive + False Positive) as shown below:

$$\text{Precision} = \frac{TP}{(TP + FP)}$$

(1)

**Recall** represents the ratio between True Positive instances and the instances which are actually positive (True Positive + False Negatives) as shown below:

$$\text{Recall} = \frac{TP}{(TP + FN)}$$

(2)

And finally F-measure is the average of Precision and Recall, as shown below”

$$\text{F-measure} = \frac{\text{Precision} \times \text{Recall} + 2}{\text{Precision} + \text{Recall}}$$

(3)

**Accuracy** represents the ratio of the instances which are classified correctly and the total number of classified instances, as shown below:

$$\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN}$$

(4)

**MCC** represents the ratio of the observed classifications and the predicted classifications.

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

(5)
First the results of the variants from stage two are discussed. Multiple variants are created in each of the classifier family (base classifier family) and then those variants are used to predict the defect prone software modules by using all of the used datasets. Three variants from 6 families of the classifiers are selected as only those three performed with highest accuracies by outperforming the base classifiers of their families. The performance of selected variants including SVM-4, RF-3, and KNN-4 are reflected in the tables below (Table 10).

| Dataset | JM1  | KC1  | PC4  | PC5  |
|---------|------|------|------|------|
|         | Default Classifier | Variant | Default Classifier | Variant | Default Classifier | Variant | Default Classifier | Variant |
| SVM-4   | 79.188 | 79.4041 | 75.3582 | 76.7908 | 88.189 | 90.5512 | 74.2126 | 74.8031 |
| RF-3    | 80.1813 | 80.6131 | 77.937 | 79.0831 | 87.4016 | 87.4016 | 75.9843 | 76.7717 |
| KNN-4   | 73.9637 | 80.0086 | 69.341 | 77.9370 | 85.8268 | 85.8268 | 73.0315 | 76.5748 |

The final results of the proposed framework in terms of F-Measure, Accuracy and MCC are reflected and compared (Table 11 to Table 14) with the results of a published paper [1]. That paper [1] has used 10 widely used supervised classification models on the same version of cleaned NASA MDP datasets, which are used for experiments in this research. The classifiers used in the published papers include: “Naïve Bayes (NB), Multi-Layer Perceptron (MLP), Radial Basis Function (RBF), Support Vector Machine (SVM), K Nearest Neighbor (KNN), kStar (K*), One Rule (OneR), PART, Decision Tree (DT), and Random Forest (RF)”.

In this research, the performance measures are only discussed for the defective class (‘Y’) as we are predicting the defective modules not the non-defective modules. Results show that the proposed framework outperformed all 10 classifiers from published paper in all three performance measures on all of the used datasets. The results from published paper show the symbol ‘?’ in few places which indicates that the score in that measure cannot be calculated due to the issue of imbalance data. The proposed framework also solved this issue besides the overall high performance.

| Classifier | F-Measure | Accuracy | MCC |
|------------|-----------|----------|-----|
| NB         | 0.318     | 79.835   | 0.251 |
| RBF        | 0.181     | 80.397   | 0.215 |
| SVM        | ?         | 79.188   | ?    |
| kNN        | 0.348     | 73.963   | 0.186 |
| kStar      | 0.355     | 75.993   | 0.212 |
| OneR       | 0.216     | 77.158   | 0.126 |
| PART       | 0.037     | 79.490   | 0.104 |
| DT         | 0.348     | 79.101   | 0.252 |
| RF         | 0.284     | 80.181   | 0.244 |
| MLP        | 0.146     | 80.354   | 0.206 |
| Proposed Framework | 0.507 | 84.974 | 0.488 |

The results of JM1 dataset are shown in Table 11. It can be seen that the proposed framework outperformed in F-Measure, Accuracy, and in MCC with the scores of 0.507, 84.974, and 0.488 respectively.

| Classifier | F-Measure | Accuracy | MCC |
|------------|-----------|----------|-----|
| NB         | 0.400     | 74.212   | 0.250 |
| RBF        | 0.362     | 78.796   | 0.347 |
| SVM        | 0.085     | 75.358   | 0.151 |
| kNN        | 0.395     | 69.341   | 0.190 |
| kStar      | 0.419     | 72.206   | 0.238 |
| OneR       | 0.256     | 73.352   | 0.147 |
| PART       | 0.255     | 76.504   | 0.239 |
| DT         | 0.430     | 75.644   | 0.291 |
| RF         | 0.454     | 77.937   | 0.346 |
| MLP        | 0.358     | 77.363   | 0.296 |
| Proposed Framework | 0.548 | 83.9542 | 0.482 |
KC1 results are reflected in Table 12. It can be seen that the proposed framework shows high performance in all of the three measures with the scores of 0.548, 83.9542 and 0.482 respectively.

Table 13. PC4 Results

| Classifier | F-Measure | Accuracy | MCC    |
|------------|-----------|----------|--------|
| NB         | 0.404     | 86.089   | 0.334  |
| RBF        | 0.250     | 87.401   | 0.279  |
| SVM        | 0.286     | 88.189   | 0.342  |
| kNN        | 0.438     | 85.826   | 0.359  |
| kStar      | 0.330     | 81.889   | 0.225  |
| OneR       | 0.361     | 87.926   | 0.352  |
| PART       | 0.481     | 85.301   | 0.396  |
| DT         | 0.583     | 86.876   | 0.514  |
| RF         | 0.532     | 90.288   | 0.516  |
| MLP        | 0.562     | 89.763   | 0.515  |
| Proposed Framework | 0.68 | 91.8635 | 0.649 |

Table 13 reflects the results of PC4 dataset. It can be observed that the proposed framework performed better with highest performance in F-Measure, Accuracy, and in MCC with the score of 0.68, 91.8635, and 0.649 respectively.

Table 14. PC5 Results

| Classifier | F-Measure | Accuracy | MCC    |
|------------|-----------|----------|--------|
| NB         | 0.269     | 75.393   | 0.245  |
| RBF        | 0.235     | 75.590   | 0.251  |
| SVM        | 0.097     | 74.212   | 0.173  |
| kNN        | 0.498     | 73.031   | 0.314  |
| kStar      | 0.431     | 69.881   | 0.227  |
| OneR       | 0.387     | 71.259   | 0.209  |
| PART       | 0.355     | 75.787   | 0.274  |
| DT         | 0.531     | 75.000   | 0.361  |
| RF         | 0.450     | 75.984   | 0.322  |
| MLP        | 0.299     | 74.212   | 0.216  |
| Proposed Framework | 0.75 | 87.7953 | 0.669 |

PC5 results are shown in Table 14. It can be observed that the proposed framework outperformed in F-Measure, Accuracy, and in MCC with the score of 0.75, 87.7953 and 0.669 respectively.

5. Conclusion

In this paper, the researchers proposed a classification framework for the prediction of defect prone software modules in order to reduce the cost of testing process in software development life cycle. The key activities performed by the researchers in order to improve the performance include: feature selection and variant based ensemble classification. Feature selection is performed to eliminate those features which do not participate in the classification process and even reduce the performance of framework besides the high processing cost. In the process of variant selection, first the variants are created by optimizing six base classifiers, including: Support Vector Machine (SVM), Decision Tree (DT), K-Nearest Neighbor (KNN), Naive Bayes (NB), Random Forest (RF), and Multi-layer Perceptron (MLP). From the variants of each base classifier (classifier family), one variant is selected which performed higher than all other variants of its family including the base classifier. Three variants are selected in this research due to high performance including SVM-4, RF-3 and KNN-4. These variants are then integrated by using an ensemble technique: “Voting” with all possible combinations. One combination of variants: RF-3 and KNN-4 outperformed all the other combinations and is selected for classification in the proposed framework. The results of the framework are compared with the results of various widely used supervised classifiers from published paper, which have used the same datasets and performance measures for performance analysis. The comparative analysis have reflected the fact that the proposed framework outperformed all other classification techniques from the published paper and also resolved the issue of class imbalance. However it is suggested for future work to optimize more classifiers with extensive set of parameters so that more variants can be selected for ensemble learning.

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