# Detection of Chronic Kidney Disease using Machine Learning Algorithms with Least Number of Predictors

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Abstract-Chronic kidney disease (CKD) is one of the most critical health problems due to its increasing prevalence. In this paper, we aim to test the ability of machine learning algorithms for the prediction of chronic kidney disease using the smallest subset of features. Several statistical tests have been done to remove redundant features such as the ANOVA test, the Pearson's correlation, and the Cramer's V test. Logistic regression, support vector machines, random forest, and gradient boosting algorithms have been trained and tested using 10-fold cross-validation. We achieve an accuracy of 99.1 according to F1measure from Gradient Boosting classifier. Also, we found that hemoglobin has higher importance for both random forest and Gradient boosting in detecting CKD. Finally, our results are among the highest compared to previous studies but with less number of features reached so far. Hence, we can detect CKD at only \$26.65 by performing three simple tests.

Keywords—Chronic Kidney Disease (CKD); Random Forest (RF); Gradient Boosting (GB); Logistic Regression (LR); Support Vector Machines (SVM); Machine Learning (ML); prediction

# I. INTRODUCTION

Chronic kidney disease (CKD) is a significant public health problem worldwide, especially for low and mediumincome countries. Chronic kidney disease (CKD) means that the kidney does not work as expected and cannot correctly filter blood. About 10% of the population worldwide suffers from (CKD), and millions die each year because they cannot get affordable treatment, with the number increasing in the elderly. According to the Global Burden Disease 2010 study conducted by the International Society of Nephrology, chronic kidney disease (CKD) has been raised as an important cause of mortality worldwide with the number of deaths increasing by 82.3% in the last two decades [1, 2]. Also, the number of patients reaching end-stage renal disease (ESRD) is increasing, which requires kidney transplantation or dialysis to save patients' lives [1, 3, 4].

CKD, in its early stages, has no symptoms; testing may be the only way to find out if the patient has kidney disease. Early detection of CKD in its initial stages can help the patient get effective treatment and then prohibit the progression to ESRD [1]. It is argued that every year, a person that has one of the CKD risk factors, such as a family history of kidney failure, hypertension, or diabetes, get checked. The sooner they know about having this disease, the sooner they can get treatment. To raise awareness and to encourage those who are most susceptible to the disease to perform the tests periodically, we hope that the disease can be detected with the least possible tests and at low cost. So, the objective of this research is to provide an effective model to predict the CKD by least number of predictors.

In this paper, Section II reviews various research works that target the diagnosis of CKD using different intelligent techniques. Section III presents the dataset source and description. Section IV presents the methodology used for the prediction, including the data preprocessing steps and the modeling stage. Section V shows the results of the experiment and discusses the performance of ML algorithms in detecting CKD. Finally, Section VI includes the conclusion and future work of this work.

## II. LITERATURE REVIEW

# A. Related Work

In recent years, few studies have been done on the classification or diagnosis of chronic kidney disease. In 2013, T. Di Noia et al. [5], presented a software tool that used the artificial neural network ANN to classify patient status, which is likely to lead to end-stage renal disease (ESRD). The classifiers were trained using the data collected at the University of Bari over a 38-year period, and the evaluation was done based on precision, recall, and F-measure. The presented software tool has been made available as both an Android mobile application and online web application.

Using data from Electronic Health Records (EHR) in 2014, H. S. Chase et al. [6] identified two groups of patients in stage 3: 117 progressor patients (eGFR declined >3 ml/min/1.73m<sup>2</sup>/year) and 364 non-progressor patients (eGFR declined <1 ml/min/1.73m<sup>2</sup>). Where GFR is a glomerular filtration rate that commonly used to detect CKD. Based on initial lab data recorded, the authors used Naïve Bayes and Logistic Regression classifiers to develop a predictive model for progression from stage 3 to stage 4. They compared the metabolic complications between the two groups and found that phosphate values were significantly higher, but bicarbonate, hemoglobin, calcium, and albumin values were significantly lower in progressors compared to nonprogressors, even if initial eGFR values were similar. Finally, they found that the probability of progression in patients classified as progressors was 81% (73%-86%) and nonprogressors was 17% (13% - 23%).

Later in 2016, K. A. Padmanaban and G. Parthiban [7] aimed in their work to detect chronic kidney disease for diabetic patients using machine learning methods. In their research, they used 600 clinical records collected from a leading Chennai-based diabetes research center. The authors have tested the dataset using the decision tree and Naïve Bayes methods for classification using the WEKA tool. They concluded that the decision tree algorithm outweighs the Naïve Bayes with an accuracy of 91%.

A. Salekin and J. Stankovic [8] evaluated three classifiers: random forest, K-nearest neighbors, and neural network to detect the CKD. They used a dataset with 400 patients form UCI with 24 attributes. By using the wrapper method, a feature reduction analysis has been performed to find the attributes that detect this disease with high accuracy. By considering: albumin, specific gravity, diabetes mellitus, hemoglobin, and hypertension as features, they can predict the CKD with .98 F1 and 0.11 RMSE.

In the study carried out by W. Gunarathne, K. Perera, and K. Kahandawaarachchi [9], Microsoft Azore has been used to predict the patient status of CKD. By considering 14 attributes out of 25, they compared four different algorithms, which were Multiclass Decision Forest, Multiclass Decision Jungle, Multiclass Decision Regression, and Multiclass Neural Network. After comparison, they found that Multiclass Decision Forest performed the best with 99.1% accuracy.

H. Polat, H. D. Mehr, and A. Cetin [10] in their research used SVM algorithm along with two feature selection methods: filter and wrapper to reduce the dimensionality of the CKD dataset with two different evaluations for each method. For the wrapper approach, the ClassifierSubsetEval Greedy Stepwise search with the engine and WrapperSubsetEval with the Best First search engine were used. For the Filter approach, *CfsSubsetEval* with the Greedy Stepwise search engine and FilterSubsetEval with the Best First search engine were used. However, the best accuracy was 98.5% with 13 features using FilterSubsetEval with the Best First search engine using the SVM algorithm without mentioning which features were used.

P. Yildirim [11] studied the effect of sampling algorithms in predicting chronic kidney disease. The experiment was done by comparing the effect of the three sampling algorithms: Resample, SMOTE, and Spread Sup Sample on the prediction by multilayer perceptron classification algorithm. The study showed that sampling algorithms could improve the classification algorithm performance, and the resample method has a higher accuracy among the sampling algorithms. On the other hand, Spread Sub Sample was better in terms of execution time. A. J. Aljaaf et al. [12] examined in their study the ability of four machine learning (ML) models for early prediction of CKD, which were: support vector machine (SVM), classification and regression tree (CART), logistic regression (LR), and multilayer perceptron neural network (MLP). By using the CKD dataset from UCI and seven features out of 24, they compared the performance of these ML models. The results showed that the MLP model had the highest AUC and sensitivity. It was also noticeable that logistic regression almost had the same performance as MLP but with the advantage of the simplicity of the LR algorithm. Therefore, in our study, we can use the LR algorithm as a start or a benchmark and then use more complex algorithms.

Lastly in 2019, J. Xiao *et al.* [13] in their study established and compared nine ML models, including LR, Elastic Net, ridge regression lasso regression SVM, RF, XGBoost, knearest neighbor and neural network to predict the progression of CKD. They used available clinical features from 551 CKD follow-up patients. They conclude that linear models have the overall predictive power with an average AUC above 0.87 and precision above 0.8 and 0.8, respectively

## B. Dataset Concern

The dataset used in this study is a small dataset with small imbalance issue as will be described in Dataset section. Therefore, there are some concerns related to this dataset, which are an overfitting or generalization problem, imbalance, and the noise of the data. P. Yang et al. [14] in their review concluded that ensemble technique has the advantage of alleviating the problem of small size data by incorporating and averaging over multiple classifiers to reduce the probability of overfitting. Also, Deng et al. [15] found in their prediction of protein-protein interaction sites that the ensemble method can handle the imbalance problem and improve the prediction performance. Another survey by M. Fatima and M. Pasha [16] found that SVM provided improved accuracy to predict heart disease with the advantage of overfitting and noise [17].

#### III. DATASET

The dataset that supports this research is based on CKD patients collected from Apollo Hospital, India in 2015 taken over a two-month period. The data is available in the University of California, Irvine (UCI) data repository named Chronic\_Kidney\_Disease DataSet [18]. These data consisting of 400 observations suffer from missing and noisy value. The data includes 250 records of patients with CKD and 150 records of persons without CKD. Therefore, the percentage of each class is 62.5% with CKD and 37.5% without CKD. The ages of these observations are varied from 2 to 90 years old. It can be seen from Table I that the CKD dataset has 24 features including 11 numeric features and 13 nominal features, and the 25<sup>th</sup> feature indicates the classification or state of CKD.

Name	Description	Type: unit/ values		
Age (age)	Patient's age	Numeric: years		
Blood pressure (bp)	Blood pressure of the patient	Numeric: mm/Hg		
Specific gravity (sg)	The ratio of the density of urine	Nominal: 1.005, 1.010, 1.015, 1.020, 1.025		
Albumin (al)	Albumin level in the blood	Nominal: 0,1,2,3,4,5		
Sugar (su)	Sugar level of the patient	Nominal: 0,1,2,3,4,5		
Red blood cells (rbc)	Patients' red blood cells count	Nominal: normal, abnormal		
Pus cell (pc)	pus cell count of patient	Nominal: normal, abnormal		
Pus cell clumps (pcc)	Presence of pus cell clumps in the blood	Nominal: present, not present		
Bacteria (ba)	Presence of bacteria in the blood	Nominal: present, not present		
Blood glucose (bgr)	blood glucose random count	Numeric: mgs/dl		
Blood urea (bu)	blood urea level of the patient	Numeric: mgs/dl		
Serum creatinine (sc)	serum creatinine level in the blood	Numeric: mgs/dl		
Sodium (sod)	sodium level in the blood	Numeric: mEq/L		
Potassium (pot)	potassium level in the blood	Numeric: mEq/L		
Hemoglobin (hemo)	hemoglobin level in the blood	Numeric: gms		
Packed cell volume (pcv)	packed cell volume in the blood	Numeric		
White blood cell count (wc)	white blood cell count of the patient	Numeric: cells/cumm		
Red blood cell count (rc)	red blood cell count of the patient	Numeric millions/cmm		
Hypertension (htn)	Does the patient has hypertension on not	Nominal: yes, no		
Diabetes mellitus (dm)	Does the patient has diabetes or not	Nominal: yes, no		
Coronary artery disease (cad)	Does the patient has coronary artery disease or not	Nominal: yes, no		
Appetite (appet)	Patient's appetite	Nominal: good, poor		
Pedal Edema (pe)	Does patient has pedal edema or not	Nominal: yes, no		
Anemia (ane)	Does patient has anemia or not	Nominal: yes, no		
Class	Does the patient has kidney disease or not	Nominal: CKD, not CKD		

TABLE. I.	DESCRIPTION OF (	CKD DATASET
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IV. METHODOLOGY

# A. Data Preprocessing

Today's real-world datasets are susceptible to missing, noisy, redundant, and inconsistent data, especially clinical datasets. Working with low-quality data leads to low-quality results. Therefore, the first step in every machine learning application is to explore the dataset and understand its characteristics in order to make it ready for the modeling stage. This process is commonly known as data preprocessing.

1) Outliers: Outliers are extreme values located far away from the feature central tendency. Invalid outliers occur due to data entry errors, which are referred to as a noise in the data [19]. Medical data cannot be treated as other data in dealing with outliers since these outliers could be legitimate (valid) or important. For this reason, each outlier detected in the CKD dataset is checked to know if it is realistic or not. In this study, the extreme data points that go beyond the acceptable range medically have been treated as missing data and then modified as will be described in the missing data section. Box plots have been used to detect outliers in the CKD dataset, as Fig. 1 shows, there are some outliers detected for blood glucose random that reached 500 mg/dl. However, as mentioned in [20], the highest blood glucose level recorded in 2008 for a surviving patient reached 2,656 mg/dl. So, these outliers are legitimate and we should not change them.

In contrast, for potassium and sodium, three extreme data points are unacceptable. The highest potassium level observed was 7.6 mEq/L [21]. This means that a potassium level with 39 and 47, as shown in Fig. 2 is impossible and usually due to a mistake. Similarly, with sodium, as Fig. 3 shows, one extreme data point was detected, which is 4.5. Normally, sodium level should be between 135 and 145 mEq/L, and if it is less than 135, then the patient suffers from hyponatremia [22]. For this reason, a value of 4.5 is unacceptable or impossible.

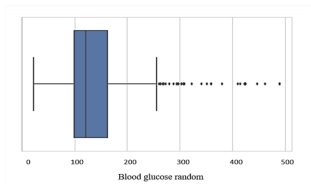


Fig. 1. Box Plot for Blood Glucose Random.

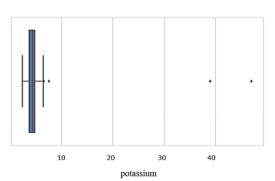


Fig. 2. Box Plot for Potassium.

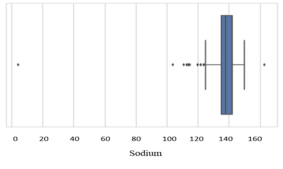


Fig. 3. Box Plot for Sodium.

2) Missing Values: In real-world datasets, missing data is a very common issue, especially in the medical area. Usually, every patient record and every attribute contains some missing values [23]. However, the chronic kidney disease dataset as shown in Fig. 4 has 96% of its variables having missing values; 60.75% (243) cases have at least one missing value, and 10% of all values are missing. There are different percentages of missing values for each variable, starting from 0.3% and reaching 38%, as shown in Table II.

Researchers in [9] used single imputation, such as mean and median, to impute the CKD dataset. However, according to Little's test [24], the missing values in CKD dataset are not missing completely at random (MCAR) with p-value <0.005. Therefore, single imputation cannot be used for handling missing values.

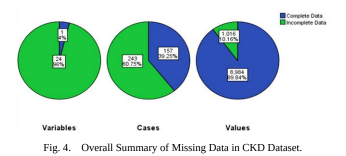
In this study, multiple imputations (MI) for replacing missing values in the CKD dataset. In multiple imputations (MI), missing values in the dataset are replaced m times, where m is usually a small number (from 3 to 10). We apply MI to produce five imputed datasets. The imputation process was based on linear regression for predicting continuous variables and logistic regression for categorical variables. Finally, we choose a dataset that has the nearest means and standard deviations for its variables to the original dataset.

*3) Data Reduction:* Data reduction means to reduce the number of features while maintaining a good analytical result. For this purpose, feature selection and features associations or correlation have been studied to remove redundant information.

a) Feature Associations: Pearson's correlation, Cramer's V, and ANOVA tests have been used to find relationships between variables. As shown in Fig. 5, and Fig. 6, there is a strong relationship between packed cell volume and hemoglobin and between hemoglobin and red cell count with the correlation coefficient of 0.89 and 0.79 respectively. Moreover, according to the ANOVA test, as shown in Table III, anemia also associated with PCV with pvalue <0.001 (2.16e<sup>-30</sup>). Another positive relationship was detected with a correlation coefficient of 0.68 between blood urea and serum creatinine.

TABLE. II. MISSING VALUES INFORMATION FOR EACH VARIABLE

	Missing		Valid	
Attribute Name	Number	Percent	Number	
Red blood cells	152	38.0%	248	
Red blood cell count	131	32.8%	269	
White blood cell count	106	26.5%	294	
Potassium	90	22.5%	310	
Sodium	88	22.0%	312	
Packed cell volume	71	17.8%	329	
Pus cell	65	16.3%	335	
Hemoglobin	52	13.0%	348	
Sugar	49	12.3%	351	
Specific gravity	47	11.8%	353	
Albumin	46	11.5%	354	
Blood glucose	44	11.0%	356	
Blood urea	19	4.8%	381	
Serum creatinine	18	4.5%	382	
Blood pressure	12	3.0%	388	
Age	9	2.3%	391	
Bacteria	4	1.0%	396	
Pus cell clumps	4	1.0%	396	
Coronary artery disease	2	0.5%	398	
Diabetes mellitus	2	0.5%	398	
Hypertension	2	0.5%	398	
Anemia	1	0.3%	399	
Pedal Edema	1	0.3%	399	
Appetite	1	0.3%	399	



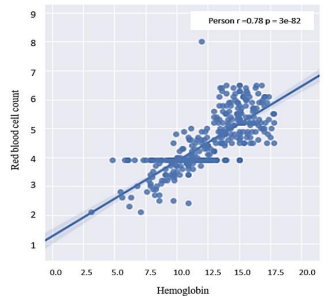


Fig. 5. Scatter Plot between Hemoglobin and Red Blood Cell Count.

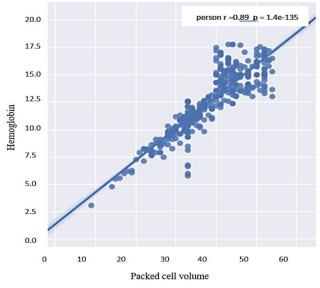


Fig. 6. Scatter Plot of PCV and Hemoglobin.

Feature 1 Categorical	Feature 2 Numeric	P- value
Diabetes mellitus	Blood glucose random	$1.29e^{-26}$
Sugar level	Blood glucose random	$2.40e^{-49}$
Hypertension	Blood pressure	$6.13e^{-0.8}$
Anemia	Packed cell volume	$2.16e^{-30}$
Red blood cell	Red blood cell count	$2.36e^{-13}$
Pus cell	White blood cell count	0.00147

TABLE. III. ANOVA TEST RESULTS

Since hemoglobin and serum creatinine have a stronger influence on the class attribute than their associated attributes, we decide to maintain them and remove the others as redundant attributes. Table IV shows the correlation between both numeric and nominal attribute and the class attribute.

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	Numeric variable	Correlation coefficient		Nominal variable	Correlation coefficient		
	Age	0.220		Albumin	0.730		
	Blood pressure	0.296		Red blood cell	0.540		
	Blood glucose random	0.399		Pus cell	0.420		
=	Blood urea	0.385	n	Pus cell clumps	0.214		
Pearson's correlation	Serum creatinine	0.361	correlation	Bacteria	0.120		
	Sodium	0.432	cor	Hypertension	0.590		
	Potassium	0.070	Cramer's V	Diabetes Mellitus	0.544		
	Haemoglobin	0.75	Cran	Coronary artery disease	0.236		
	Packed cell volume	0.72		Appetite	0.393		
	White blood	0.222		Pedal edema	0.365		
	cell count 0.222			Anemia	0.325		
	Red blood cell			Sugar level	0.432		
	count	0.666		Specific gravity	0.687		

TABLE. IV. CORRELATION BETWEEN ALL VARIABLES AND CLASS VARIABLE

Diabetes mellitus, sugar level, and blood glucose random almost measure the same thing, which is "sugar". The result proves the association by having p-value <0.001 (1.29  $e^{-26}$ ) when testing the correlation between them, and 0.55% coefficient between sugar level and diabetes according to Cramer's V test. The same procedure was applied to other associated features. In the end, nine features have been removed as redundant features. These features are blood glucose random, blood pressure, packed cell volume, red blood cell count, red blood cell (nominal), anaemia, sugar level, pus cell, and blood urea.

b) Feature Selection: The process of selecting the most discriminating features in a given dataset is known as feature selection. This process is enhancing the model's performance, reducing overfitting, and reducing the cost of building a model. Filter feature selection methods [25] selects features that have a stronger relationship with the outcome variable independent to the learning model. Therefore, use a measure or test independent to the learning algorithm to assess a subset of features. In this study, mutual information measure has been used as a feature selection method. Mutual information [25] measures the dependence of any kind of relationships between random variables.

4) Data transformation: In data transformation, data is transformed into appropriate forms for mining purposes [26]. Data transformation includes normalization, which is the process of scaling the attributes' values to fall within a small specific range [26]. It is usually applied before feature selection and modeling stages because different scales of attributes complicate the comparison of attributes and influence the ability of algorithms to learn [23]. However, in this study min-max normalization has been applied on numeric data types. Another data transformation has been done on categorical variables. This is because some ML algorithms cannot handle categorical variables, especially in regression problems. Therefore, categorical variables with n values are dummied in LR and SVM classifiers by converting each of them into n-1 dummy variables [27].

## B. Modeling

In the modeling stage, four machine learning algorithms have been applied to the dataset to assess their ability to detect CKD. These algorithms are logistic regression (LR), support vector machines (SVM), random forest (RF), and gradient boosting (GB).

1) Logistic regression: Logistic regression [28], also called logit model or logistic model, is a widely used model to analyze the relationship between multiple independent variables and one categorical dependent variable with the equation of the form:

$$\log\left[\frac{p}{1-p}\right] = a + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_i x_i \tag{1}$$

Where *p* is the probability of interest outcome, *a* is an intercept,  $\beta_1, \ldots, \beta_i$  are  $\beta$  coefficients associated with each variable *x*, and  $x_1, \ldots, x_i$  are the values of the predictor variables.

2) SVM: Support Vector Machines (SVM) [29] is a supervised learning model that is commonly used in classification problems. The idea of the SVM algorithm is to figure the optimal hyperplane that ideally separates all objects of one class from those objects of another class with the largest margins between these two classes. The objects that are far from the boundary are discarded from the calculation, while other data points that are located on the boundary will be maintained and determined as "support vectors" to get satisfactory computational efficiency [29]. The SVM algorithm has different kernel functions: radial basis function (RBF), linear, sigmoid, and polynomial. In this study, radial basis function has been chosen based on nested cross-validation results.

3) Ensemble method: Ensemble method [30] is a strategy for improving predictor or classifier accuracy. Ensemble method uses a combination of models to create an improved composite model to improve the performance. The main idea behind the ensemble technique is to group multiple "weak learners" to come up with a "strong learner". Two popular techniques for constructing ensembles are bagging and boosting. Both boosting and bagging can be used for prediction as well as classification [26, 30]. Bagging is an ensemble technique where many independent predictors or learners are built and their results are combined using the majority vote, whereas in boosting, the predictors or learners are made sequentially not independently. This sequential method because each classifier "pays more attention" to the training tuples that were misclassified by the previous classifier through assigning weights for each of them [26]. Random forest algorithm is an example of the "bagging" technique, whereas the gradient boosting algorithm is an example of the "boosting" technique. Fig. 7 shows the bagging and boosting structure in selecting samples for training.

a) Random Forest: Random forest (RF) is a bagging ensemble approach proposed by Breiman [31] that based on a machine learning mechanism called "decision tree". In a random forest, the "weak learners" in ensemble terms are decision trees [8, 32, 33]. Random forest imposes the diversity of each tree separately by selecting a random feature. After generating a large number of trees, they vote for the most common class. The random forest algorithm can deal with unbalanced data, it is robust against overfitting, and its runtimes are quite a bit faster [8, 31].

b) Gradient Boosting: Gradient boosting (GB) is an ensemble boosting technique that starts with "regression tree" as "weak learners". In general, the GB model adds an additive model to minimize the loss function by using a stage-wise sampling strategy. The loss function measures the amount at which the expected value deviates from the real value. Stagewise fashion put more emphasis on samples that are difficult to predict or misclassified. Unlike random forest, in GB, samples that are misclassified have a higher chance of being selected in training data [34]. GB reduces bias and variance and often provides higher accuracy, but the parameters should be tuned carefully to avoid overfitting. Therefore, nested cross-validation has been applied.

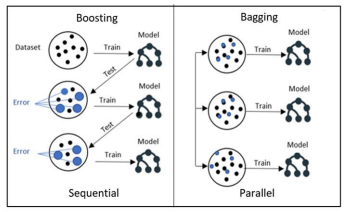


Fig. 7. Bagging and Boosting Structure.

#### V. RESULTS AND DISCUSSION

The result of each classifier has been evaluated using different evaluation metrics and validated against overfitting using 10-fold cross-validation. The nested cross-validation approach also has been applied for the purpose of tuning the models' parameters. The experiments are conducted using Python 3.3 programming language through the Jupyter Notebook web application. Several libraries from *Sciket-learn* [35] have been used, which is a free software for the machine learning library in Python. The evaluation measures considered in this study are accuracy using F1-measure, sensitivity, specificity, and area under the curve (AUC).

Each model generates different outputs depending on the different values of its parameters. By using nested cross-validation, the best performance for LR was with C=1000 and penalty=L2 with an accuracy of 98.9% using F1 measure. For the SVM model different values of "C", "gamma", and "kernel" have been tested. The best performance for SVM was with C=1 and gamma=3, and kernel = "RBF" (radial basis function) with an accuracy of 97.9% using F1 measure. For both RF and GB, the best results were with a number of trees=50 and Max\_depth=2, with an accuracy of 98.0% in RF and 99.1% in GB using F1 measure.

The experimental results of each model in terms of accuracy, F1-measure, precision, sensitivity, specificity, AUC are listed in Table V whereas the training and testing accuracies based on 10-fold cross-validation are listed in Table VI.

From the evaluation results, as Fig. 8 shows, all models have an excellent performance against detecting CKD with an accuracy > 97% using hemoglobin, specific gravity, and albumin features. By focusing on specificity and sensitivity, it is seen that all models also have the same specificity of 99.3% except RF (96.6), which means that all models were accurate in identifying the negative or healthy subjects. On the other hand, the highest sensitivity was obtained using the RF algorithm at 99.6%, which represents the percentage of correctly identified CKD patients.

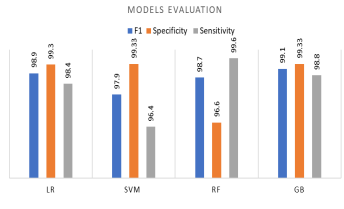


Fig. 8. Models Evaluation in Predicting CKD.

TABLE. V.	THE PREDICTIVE PERFORMANCE OF ML MODELS	

Classifier	Accura cy	F1	Precisi on	Sensiti vity	Specificit y	AUC
Logistic regressio n	98.75%	98.9 %	99.5 %	98.4 %	99.33 %	99.7 %
Support victor machines	97.5%	97.9 %	99.5%	96.4 %	99.33%	99.9 %
Random forest	98.5%	98.7%	98.0%	99.6%	96.6%	99.5%
Gradient boosting	99.0%	99.1 %	99.5%	98.8 %	99.33%	99.9%

 TABLE. VI.
 THE TRAIN AND TEST ACCURACIES OF ML MODELS

	LR	SVM	RF	GB
Train	99.4 %	97.52%	98.5%	99.7%
Test	98.75 %	97.5%	98.5%	99.0%

Hence, we achieve the highest detection performance with the GB model. This performance is higher than the performance achieved by [12] using a multilayer perceptron algorithm (MLP), seven features, and single-point split with 98.4% F1- measure. Also, higher performance Compared to study [8], were 98.0% F1-measure have been achieved using RF and five features. According to study [8], which also estimated the cost of each of 24 tests in the CKD dataset, performing these three features for detecting CKD would cost only \$26.65 while using all features will cost around \$451.36.

Since the higher results were achieved using RF, and GB algorithm, we also investigate the importance of the features in each of them. As shown in Fig. 9, haemoglobin has the highest score, whereas Albumin has the lowest score in both RF and GB. Looking at RF, the degree of importance is convergent for all variables, approximately from 0.29 to 0.44. Whereas in GB, there is a significant difference between the degree of importance of haemoglobin (0.77) and other features. Then, according to our result, we conclude that haemoglobin has played an essential role in detecting CKD.

However, this research is subject to some limitations related to the dataset used. First, the size of the dataset is considered to be small (400 instances), which may influence the reliability of the results. Second, difficulty finding is another dataset that has the same features in order to compare the results of the datasets.



Fig. 9. Importance of Features in RF and GB Models.

#### VI. CONCLUSION AND FUTURE WORK

This work examines the ability to detect CKD using machine learning algorithms while considering the least number of tests or features. We approach this aim by applying four machine learning classifiers: logistic regression, SVM, random forest, and gradient boosting on a small dataset of 400 records. In order to reduce the number of features and remove redundancy, the association between variables have been studied. A filter feature selection method has been applied to the remaining attributes and found that there are haemoglobin, albumin, and specific gravity have the most impact to predict the CKD. The classifiers have been trained, tested, and validated using 10-fold cross-validation. Higher performance was achieved with the gradient boosting algorithm by F1-measure (99.1 %), sensitivity (98.8%), and specificity (99.3%). This result is the highest among previous studies with less number of features and hence less cost. Therefore, we conclude that CKD can be detected with only three features. Also, we found that hemoglobin has the highest contribution in detecting CKD, whereas albumin has the lowest using RF and GB models.

Since the data used in this research is small, in the future, we aim to validate our results by using big dataset or compare the results using another dataset that contains the same features. Also, in order to help in reducing the prevalence of CKD, we plan to predict if a person with CKD risk factors such as diabetes, hypertension, and family history of kidney failure will have CKD in the future or not by using appropriate dataset.

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