# CORONARY HEART DISEASE USING SUPPORT VECTOR MACHINE

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#### **Abstract**

The preference of SVM kernel function with optimal features that flexibly applied for dynamic dataset is a new challenge. The restriction of technology and infrastructure support for diagnosing the bioinformatics at rural area is a major concern for developing countries towards excellent health services. Therefore, this study aimed at evaluating the utilization of Support Vector Machine (SVM) in classifying patients of coronary heart disease with Unstable Angina Pectoris (UAP), Non-Segment (ST) Elevation Myocardial Infarction (NSTEMI) and ST-Elevation Myocardial Infarction (STEMI) classes. So far, 280 samples were experimented with 17 attributes by considering four types of dataset, which include the original, reduced, pre-processing and K-Nearest Neighbours (k-NN). To evaluate the optimal parameter pairs in terms of accuracy and processing time for the above dataset types, 10-folds cross-validation and percentage split were carried out on Polynomial and Radial Basis Function (RBF) kernels. Waikato Environment for Knowledge Analysis (WEKA) tool for 10-folds reveals the optimum accuracy of 100% for polynomial kernel and 98.9% for RBF. Also, the percentage split of 70:30 affirms 100% accuracy with 0.06 seconds of processing time as the ideal values of Polynomial kernel test. Meanwhile, RBF exhibits 80:20 split for 100% accuracy with 0.08 seconds in dataset pre-processing. In a nutshell, SVM enhances the data precision and recall as well as minimizes the error possibility for the greatest classification of coronary heart disease patients in Polynomial and RBF kernel than other classifier such as Neural Network (NN). Therefore, the application of SVM improves the accuracy of coronary heart disease diagnostics.

Keywords: Neural network, K-nearest neighbours, Data mining, Support vector machine, Coronary heart diseases.

#### 1. Introduction

Data mining provides various manipulation services to achieve the prediction, classification, clustering, mapping, and anomalous detection of data. The utilization of this technique in various disciplines has evolved and shown a significant contribution to the field of knowledge, including medicine, finance, industry, technology, and even molecular biology as well as bioinformatics. With an emphasis on classification, the advent of methods in disaggregation data improves its usefulness and maneuverability in interpreting information, for examples Nijssen and Fromont [1] studied the optimal constraint of Decision tree method induction in pattern mining; Network and Tree-based methods were applied for data mining modeling in the corrosion of concrete sewer [2]; k-NN for scholarship recipient cases [3]; Multilayer Perceptron (MPL) for data mining in healthcare operations [4]; Naive Bayes approach in classifying the analysis of students' performance [5], Artificial Neural Network as a validation tool of Loud Haul Dump (LHD) machine performance characteristics [6], Neural Network in designating the water cycle problems [7] and the utilization of SVM in data mining [8].

Recently, the enforcement of the above methods in analysing the complex bioinformatics data was put into practice. Big data opportunities bring unprecedented potential and challenges in data mining and biological analysis systems in a cost-efficient manner [9]. Also, big data technology ensures that the biologist generates large amount of facts and measurement of genomic sequences, images of physiological structures, measuring the messenger Ribonucleic Acid (mRNA) and protein expression, transcription factor binding, and metabolite concentration with limitation of programming skills [10]. In addition, Majhi et al [11] utilized bioinformatics techniques to identify the early stages of diseases such as metabolic and urea cycle disorders, inborn errors and path-aligners through genetics analysing processes and proteomics reports, which are therefore compared with health care data. Furthermore, Dashtban and Balafar [12] found the significance of data mining as artificial intelligent tools in classifying the microarray cancer data.

The adoption of machine learning algorithms in bioinformatics accomplished the reduction of complex data and allocated the feature selection of biomarkers in raw data. Serra et al [13] verified the successful employment of machine learning techniques as well as clustering, classification, embedding techniques and networkbased approaches in addressing bioinformatics problems which include gene expression clustering, patient classification, brain network analysis, and identification of biomarkers. In addition, this technology's ability to capture biomedical data has reformed machine learning into a sophisticated way to solve the complexities of big data. The number of heterogeneity modalities in biological and neurobiological phenomena insists on the multi-view of intelligent data integration from several resources. In addition, multi-view learning and data integration offers greater statistical power analysis [14]. In the process of improvising classification parameters, especially in predicting bioinformatics data, a high level of precision is required to produce the best and most effective classifier tool. The classification techniques that involve data mining, as well as machine learning, reduce computational time and improve categorization precision in determining the optimum values as clarifying in the case of unknown protein sequence classification [15].

SVM is a classification method that produces a fairly high degree of accuracy and is commonly used compared with the conventional decision tree, ANN [16, 17] and other classifiers [18]. Furthermore, Sivagami [19] compared SVM, Multilayer Perception (MLP), One R, and Decision Tree J48 methods in the classification of breast cancer. The results showed that SVM with kernel type RBF provided the highest accuracy rate of 95%, 91% in polynomial type, and 90% in linear type. One R exhibited 83%, 80% in J48 and 74.1% in MLP, which is the lowest performance. The comparison of SVM and Left Anterior Descending (LDA) for the classification of Coronary heart showed accuracy at 96.86% and 78.18% respectively [20]. Furthermore, Mo and Xu [21] attempted to improve the performance of SVM based on the hybrid kernel function using the optimization of the Particle Swarm Optimization (PSO) algorithm in heart disease diagnosis. Meanwhile, the accuracy of SVM in the early diagnosis of a heart condition by modifying the kernel width using trial and error approach significantly increase by 18.2% [22]. This showed that the kernel function on SVM provides the opportunities in enhancing the accuracy. Unfortunately, some difficulties in choosing the SVM kernel function were encountered [23], as well as flexibility in dataset changing [24], selecting optimal features, and time-consumption [25].

Exploring the various advantages and problems in the SVM method, this study aimed at identifying Coronary Heart Disease (CHD) problems in Indonesia cases. This is interesting since most hospitals are equipped with standard technological equipment in terms of features, functions, and images in interpreting a disease, especially in rural areas. Simultaneously, medical doctors are required to provide a proper diagnostic and ensure appropriate treatment for patients. Furthermore, taking advantage of SVM in Bioinformatics, 280 CHD patients at Central Hospital in Pekanbaru, Riau were classified into three class outputs namely UAP, NSTEMI, and STEMI. By emphasizing on the kernel function impact [22, 23], the classification of Polynomial and RBF kernels using one-against-one multiclass SVM was analysed and investigated to explore the optimality of SVM accuracy. Therefore, this classification aids the diagnosis of CHD patients at an early stage flawlessly, to begin further treatment and also in clinical decision making [26].

The organization of this study begins with an introduction that explains the background, previous reviews on the SVM method, the objectives, the research work, and implications. Furthermore, detailed data, instruments, and step processes are elucidated in the research method. The output of Knowledge Discovery and Data mining (KDD) and SVM analysis as well as and SVM evaluation are deliberated in the research result and discussion. Finally, the conclusion is given as a resume and suggestion is made for future studies.

# 2. Research Method

Related to data resources, CHD patients in Central Hospital taking from 2011 to 2014 with the specification on 134, 64, and 82 cases of UAP, NSTEMI, and STEMI respectively. In selecting data that limits the patient's age beyond 25 years, 17 attributes were exploited and they were defined based on the reviews of previous researches [27-33] as presented in Table 1.

Table 1. Numbers of attributes

Code	Attributes
1	Age
2	gender
3	family history
4	heart history
5	history of diabetes mellitus
6	history of hypertension
7	history of cholesterol
8	obesity
9	systolic blood pressure
10	diastolic blood pressure
11	LDL levels
12	HDL levels
13	total cholesterol levels
14	triglyceride levels
15	blood levels glucose
16	elevation
17	cardiac enzymes

Following the employment of SVM in KDD [34].

# Step 1: Pre-processing

This step is to reduce data, therefore there is no missing value. The activity begins with data selection from CHD and then performed as an effort to feature subset selection by ignoring the irrelevant attributes CHD risk factors and missing values. In view of this, k-NN with the Euclidian distance calculation is performed in Eq. (1)

$$dist = \sqrt{\sum_{k=1}^{n} (pk - qk)^2} \tag{1}$$

where n is number of attributes, pk and qk values are the -k attribute.

# Step 2: Transformation

This step is to produce seventeen attributes and using k-NN and it is driven by discretizing the attributes with an equal width approach. The Equal width is one of the unsupervised discretizations of continuous features to obtain a better precision rate in dealing with data manipulation with high cardinality attributes [35] and its outputs become an input to the classification.

## Step 3: Classification using SVM

Subsequently, the core process of data mining, which is the one-against-one SVM multiclass method is defined with a value of d, sigma  $\sigma$ , and C as explained in Table 2.

Table 2. The define of SVM value

d	sigma (ơ)	C
1	1	0.01
2	2	0.02
3	3	0.03
4	4	0.04
5	5	0.05
		0.06
		0.07
		0.08
		0.09
		0.1
		0.2
		0.3
		0.4
		0.5
		0.6
		0.7
		0.6
		0.9
		1

The variable d is specified as the degree of the polynomial, the value of C is a constant that allows to trade off the influence of the higher and lower-order terms and this is a consideration for varying C values between 0.01 and 1. The selection values of d, and  $\sigma$  impact the performance accuracy, while C is selected based on the C function as a constraint, therefore, a greater value of C implies more penalty for classification errors. Meanwhile, the values of  $\sigma$  provide a good fit or an overfit to the data, when  $\sigma$  is large compared to the distance between the classes, it results in an overly flat discriminant surface. However, a smaller  $\sigma$  value compared to the distance between classes result in an over-fit [36]. A good choice for  $\sigma$  will be comparable to the distance between the closest members of the two classes. Furthermore, the highest accuracy of parameter pairs during the training session was found at C and  $\sigma$  for kernel RBF as well as C and d for the polynomial kernel. To process the data, WEKA 3.7.10, which is a powerful tool in data mining [37] and machine learning [38] was adopted.

# Step 4: Evaluation using SVM

The evaluation process was carried out to ensure the performance of the classification methods in the SVM with two kernel trick types on polynomial and RBF. The value of accuracy and time in the building model is thoroughly investigated to achieve the superlative one. Also, the 10-folds validation and confusion matrix with percentage splits on the portion of training data compare to test data in 40:60, 50:50, 60:40, 70:30, and 80:20 is applied to support the assessment process. However, there are no specific rules in the distribution of training-data and test-data, therefore, a large number of the former will represent the diversity of the data [39]. Furthermore, to calibrate the testing procedure and the overcoming of various issues related to percentage splits in defining the best *C* and parameter values, 10-folds validation was exploited. Also, the test simulation took place in four stages, viz the original dataset (with missing values), the reduced

(no missing values), the k-NN (with Euclidian distance calculation), and the Preprocessing (with KDD formation). Therefore, the success rate of classification, the determination of accuracy, error rate, precision, and recall values are performed based on the confusion matrix as depicted in Eqs. (2-5) [40] given by,

$$Accuracy = \frac{TP + TN}{P + N} \times 100\% \tag{2}$$

$$Error-rate = \frac{FP+FN}{P+N} \times 100\%$$
 (3)

$$Precision = \frac{TP}{TP + FP} \tag{4}$$

$$Recall = \frac{TP}{TP + FN} \tag{5}$$

TP (*True Positive*) = The amount of correctly classified data (*Actual class* (yes), *Predicted class* (yes)).

TN (*True Negative*) = The amount of correctly classified data (*Actual class* (no), *Predicted class* (no)).

FN (False Negative) = The amount of incorrectly classified data (Actual class (yes), Predicted class (no)).

FP (False Positive) = The amount of incorrectly classified data (Actual class (no), Predicted class (yes)).

P = Total of TP and FN N = Total of FP and TN

To scrutinize the performance of SVM with other classifiers, the calculation of confusion matrix in NN Multilayer perceptron is measured by considering the values of accuracy, error rate, precision, and recall. This was adopted because research has steadily built on the accuracy and efficiency of data mining using NN and SVM for medical prediction and classification tasks [41, 42]. NN methods were extensively adopted in classifying problems and as one of the most active research and application areas. Furthermore, SVM and NN have been used with high accuracy in classification with relatively small sample data [43, 44].

#### 3. The Results Result and Discussion

#### 3.1. The Result of KDD analysis

#### 3.1.1. Pre-processing data analysis

The data were manually selected from the medical record of 280 CHD patients at Central Hospital by paying special attention to the feature related to attributes and missing value treatments. The diversity of data based on the feature is shown in Table 1 and missing value consideration in Fig. 1. Table 3 explains that the increasing numbers of training data from 40%, 50%, 60%, 70%, and 80% are directly proportional to the diversity of data in accordance with seventeen attributes and three classes (UAP=1, NSTEMI=2, and STEMI=3). Consequently, the new pattern tested data is recognized easily. Also, Fig. 1 describes the transformation of pre-processing activity before and after manipulating the missing values by referring to k-NN distance calculation in Eq. (1). The missing values in the dataset

at number 28 column 11, 12, and 14 is replaced by 93, 57, and 84 respectively as well as the missing values at dataset number 69, and 71.

**Training Data Composition Feature** 40% 80% Classes Classes Area Area 37-44 Age (1) 25-31 45-51 32-37 52-58 38-43 59-65 44-49 66-72 50-55 ... 73-79 56-61 80-86 62-67 68-73 74-79 80-86 Gender (2) M M F Cardiac Norm Norm

Table 3. Data diversity according to the feature.

		•	Cart	mac		1	1011	11	55	U	C	•••	•	NOTH	1 .	105	U	U
	]	Enz	zym	es (1	<b>(7</b> )													
						]	Higl	1	0	31	2	6		High		0	68	51
No	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	Case
27	68	F	Yes	No	Yes	Yes	No	No	144	93	160.5	31.1	207	77	381	Yes	High	STEMI
28	63	М	No	Yes	Yes	Yes	No	No	140	90	?	?	212	?	262	Yes	High	STEMI
31	56	М	No	No	No	Yes	No	No	150	90	138.6	38.4	198	105	83	Yes	High	STEMI
69	68	М	No	No	No	No	No	No	100	70	?	?	164	241	76	Yes	High	STEMI
71	54	М	No	No	No	Yes	No	No	120	90	139.1	70	226	?	88	Yes	High	STEMI
			•					•			•			•				•
No	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	Case
27	68	F	Yes	No	Yes	Yes	No	No	144	93	160.5	31.1	207	77	381	Yes	High	STEMI
28	63	М	No	Yes	Yes	Yes	No	No	140	90	93	57	212	84	262	Yes	High	STEMI
31	56	М	No	No	No	Yes	No	No	150	90	138.6	38.4	198	105	83	Yes	High	STEMI
69	68	М	No	No	No	No	No	No	100	70	90.3	20.1	164	241	76	Yes	High	STEMI
71	54	м	No	No	No	Yes	No	No	120	90	139.1	70	226	140	88	Yes	High	STEMI

Fig. 1. Pre-processing with missing value.

# 3.1.2. Transformation data analysis

The medical records of CHD patients were collected in a variety of formats. Consequently, the discretization with the equal width approach was applied in expressing the standard range values from 0 to 1 as in Eq. (6).

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Series of range= the highest area – the lowest area

The number of categories

(6)

The discretization of attributes is depicted in Tables 4 and 5. Table 4 defines the values of attribute 1 for age discretization, attribute 9 for systolic blood pressure (BP), attribute 10 for diastolic blood pressure, attribute 11 for LDL, attribute 12 for HDL, attribute 13 for Total cholesterol, attribute 14 for Triglyceride, and attribute 15 for a glucose level. The rest of the attributes (2,3,4,5,6,7,8,16, and 17) were categorized into two series and discretized into 0 value for "No" and 1 for "Yes" as shown in Table 5. This discretization value will be the format for SVM input. The sample of format SVM input is described in Fig. 2.

#### 3.1.3. SVM mining analysis

To investigate the implication of pre-processing against SVM, the analysis is conducted by comparing the accuracy within dataset changes in the original data (without missing values), the reduced (with missing values), k-NN (with distance calculation), and pre-processing (KDD formatted). This was executed through the selection of the best parameters for 10-fold cross-validation in Table 6 and percentage split in Table 7 for four scenarios dataset. The graphical views of performances are shown in Figs. 3, 4, and 5.

The execution of 10-folds cross-validation in Table 6 explained that the preprocessing dataset improved accuracy level up to 100% and 98.9% in kernel polynomial and RBF, respectively with the superior parameters at C=0.02 and d=2, C=0.8 and  $\sigma=1$ , respectively. Similarly, Table 7 shows that the preprocessing dataset with the percentage split treatment also provided a significant growth of accuracy in polynomial and RBF kernel. Moreover, the execution time in model development considerably impacts the performance of pre-processing both in Polynomial and RBF kernel at the data composition of 70:30 and 80:20, respectively. Figures 3, 4, and 5 explained that the pre-processing dataset increases its performance in terms of time (s) and accuracy for Polynomial and RBF kernel.

Table 4. Attribute discretization

Age discr	etization (1)	Systolic TD di	scretization (Sis) (9)
Age (years)	Discretization	Systolic BP (mmHg	
25 < U <35	0	Sis<120	Optimal (0)
$35 \leq U < 45$	0.2	120< Sis <130	Normal (0.2)
45 < U <55	0.4	130< Sis <140	Normal Height (0.4)
55 < U <65	0.6	140< Sis <150	Low hypertension
_			(0.6)
$65 \le U < 75$	0.8	150< Sis <160	Moderate
			hypertension (0.8)
U ≥ 85	1	Sis >160	Severe hypertension
			(1)
	ias) discretization 10)	Discretization of	LDL (LDL) levels (11)
Diastolic BP	Discretization	LDL levels (mg/	Discretization
(mmHg)		dL)	
Dias<80	Optimal (0)	LDL<100	Optimal (0)
80 <u>&lt;</u> Dias<85	Normal (0.2)	100< LDL <130	Approaching optimal (0.25)
85 <u>&lt;</u> Dias<90	Normal Height (0.4)	130< LDL <160	Borderline high (0.5)
90 <u>&lt;</u> Dias<100	Low hypertension (0.6)	160< LDL <190	High (0.75)
100 <u>&lt;</u> Dias<110	Moderate hypertension (0.8)	LDL >190	Very high (1)
	(0.0)		
<b>Dias</b> >110	Severe		
_	Severe hypertension (1)		
Discretization o	Severe hypertension (1) of HDL(HDL) (12)	Discretization of to	tal cholesterol (Chol)(13)
Discretization of HDL levels	Severe hypertension (1)	Chol	tal cholesterol (Chol)(13) Discretization
Discretization o	Severe hypertension (1) of HDL(HDL) (12)		
Discretization of HDL levels	Severe hypertension (1) of HDL(HDL) (12)	Chol levels (mg	
Discretization of HDL levels (mg / dL)	Severe hypertension (1) of HDL(HDL) (12) Discretization	Chol levels (mg / dL)	Desirable (expected to be safe) (0) Borderline (must be
Discretization of HDL levels (mg / dL)  HDL<40	Severe hypertension (1)  ff HDL(HDL) (12)  Discretization  Low (0)	Chol levels (mg / dL) Chol <200	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control)
Discretization of HDL levels (mg / dL)  HDL<40	Severe hypertension (1)  ff HDL(HDL) (12)  Discretization  Low (0)	Chol levels (mg / dL) Chol <200	Desirable (expected to be safe) (0) Borderline (must be
Discretization of HDL levels (mg / dL)  HDL<40  40≤ HDL <60  HDL≥60	Severe hypertension (1)  f HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)
Discretization of HDL levels (mg / dL)  HDL<40  40≤ HDL <60  HDL≥60	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1)	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)
Discretization of HDL levels (mg / dL)  HDL<40  40≤ HDL <60  HDL≥60  Triglyceride d	Severe hypertension (1)  If HDL(HDL) (12) Discretization  Low (0)  Normal (0.5)  High (1) iscretization (14)	Chol levels (mg / dL) Chol <200  200 ≤ Chol <240  Chol ≥240  Glucose Level di	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)
Discretization of HDL levels (mg / dL)  HDL<40  40≤ HDL <60  HDL>60  Triglyceride d Triglyceride levels (mg / dL)	Severe hypertension (1)  f HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1) iscretization (14) Discretization	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)	Discretization  Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization
Discretization of HDL levels (mg / dL)  HDL<40  40≤ HDL <60  Triglyceride d Triglyceride levels (mg / dL)  trig <150	Severe hypertension (1)  f HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1) iscretization (14)  Discretization  Normal (0)	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)  Glu<40	Discretization  Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization  Optimal (0)
Discretization of HDL levels (mg / dL)  HDL<40  40 ≤ HDL <60  HDL ≥60  Triglyceride d Triglyceride levels (mg / dL)  trig <150  150 ≤ trig <200	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1) iscretization (14)  Discretization  Normal (0)  Borderline high (0.33)	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization  Optimal (0)  Normal (0.2)
Discretization of HDL levels (mg / dL)  HDL<40  40 ≤ HDL <60  Triglyceride d Triglyceride levels (mg / dL)  trig <150  150 ≤ trig <200  200 ≤ trig <500	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1)  iscretization (14)  Discretization  Normal (0)  Borderline high	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)  Glu<40 40≤ Glu<60  60≤ Glu<125	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization  Optimal (0)  Normal (0.2)  Normal Height (0.4)
Discretization of HDL levels (mg / dL)  HDL<40  40 ≤ HDL <60  HDL ≥60  Triglyceride d Triglyceride levels (mg / dL)  trig <150  150 ≤ trig <200	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1) iscretization (14)  Discretization  Normal (0)  Borderline high (0.33)	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)  Glu<40 40≤ Glu <60  60≤ Glu <125 125≤ Glu <145	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization  Optimal (0)  Normal (0.2)  Normal Height (0.4)  Low hypertension (0.6)
Discretization of HDL levels (mg / dL)  HDL<40  40 ≤ HDL <60  Triglyceride d Triglyceride levels (mg / dL)  trig <150  150 ≤ trig <200  200 ≤ trig <500	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1)  iscretization (14)  Discretization  Normal (0)  Borderline high	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)  Glu<40 40≤ Glu <125 125≤ Glu <145 145≤ Glu	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  scretization (Glu) (15)  Discretization  Optimal (0)  Normal (0.2)  Normal Height (0.4)  Low hypertension (0.6)  Moderate
Discretization of HDL levels (mg / dL)  HDL<40  40 ≤ HDL <60  Triglyceride d Triglyceride levels (mg / dL)  trig <150  150 ≤ trig <200  200 ≤ trig <500	Severe hypertension (1)  If HDL(HDL) (12)  Discretization  Low (0)  Normal (0.5)  High (1)  iscretization (14)  Discretization  Normal (0)  Borderline high	Chol levels (mg / dL) Chol <200  200≤ Chol <240  Chol ≥240  Glucose Level di Glucose Levels (mg/dL)  Glu<40 40≤ Glu <60  60≤ Glu <125 125≤ Glu <145 145≤ Glu <200	Desirable (expected to be safe) (0)  Borderline (must be aware- begin to control) (0.5)  High (1)  iscretization (Glu) (15)  Discretization  Optimal (0)  Normal (0.2)  Normal Height (0.4)  Low hypertension (0.6)

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Table 5. Attributes with two series discretization

Attributes	Discretiza	tion
Gender (2)	Male	1
	Female	0
Family History (3)	None	0
	Yes	1
Heart History (4)	None	0
	Yes	1
DM History (5)	None	0
	Yes	1
<b>Hypertension History (6)</b>	None	0
	Yes	1
Cholesterol History (7)	None	0
	Yes	1
Obesity (8)	None	0
	Yes	1
Elevation (16)	None	0
	Yes	1
Cardiac Enzymes (17)	None	0
	Yes	1

	Attributes discretization																	
No	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	Cases
1	0,4	1	0	0	0	0	0	0	0,4	0,2	0	0	0	0	0,4	0	0	UAP
2	0,4	1	0	0	0	0	0	0	0,8	0,8	0	0	0	0	0,4	0	0	UAP
3	0,4	0	0	1	0	0	0	0	0,2	0,2	0	0,5	0	0	0,4	0	0	UAP
4	0,6	1	0	1	0	0	0	0	0,2	0,2	0,25	0,5	0	0	0,8	0	0	UAP
5	0,6	1	0	0	1	0	0	0	1	0,2	0,25	0,5	0	0	0,4	0	0	UAP
6	0,6	1	0	1	1	1	1	0	0,2	0,2	0.5	1	0,5	0	0,4	0	1	NSTEMI
7	0,6	1	0	0	0	0	0	0	0,6	0,2	0	0	0	0,33	0,8	0	0	UAP
8	0,4	0	0	1	0	0	0	0	0,6	0,8	0,25	1	0,5	0	0,4	0	0	UAP
9	0,6	1	0	0	1	1	0	0	0,6	0,2	0.75	0	0,5	0	0,4	0	0	UAP
10	0,4	0	0	1	1	1	0	0	0,4	0,2	0,25	0,5	0,5	0,66	0,4	0	0	UAP
11	0,2	1	0	1	1	1	0	0	0,2	0	0,25	0	0	0	0,4	0	0	UAP
12	0,6	1	0	1	1	1	0	0	1	0,8	0,25	0	0	0,33	0,4	0	0	UAP
13	0,4	0	0	0	0	0	0	0	0	0	1	0	1	0	0,4	0	0	UAP
14	0,4	1	0	1	1	1	0	0	0,6	0,6	0	0	0	0	0,4	0	0	UAP
15	0,4	1	0	1	1	1	0	0	0,6	0.8	0.5	0	0,5	0,66	0,4	0	0	UAP
16	0,4	0	0	0	1	1	0	0	0,6	0,8	0,25	0	0	0	0,8	0	1	NSTEMI
17	0,8	0	0	0	0	0	0	0	1	0,8	0,25	0	0,5	0	0,8	1	1	STEMI
18	0,4	0	1	0	1	1	0	0	0,6	0,6	0	0	0	0	0,4	1	1	STEMI
19	0,6	1	0	0	0	0	0	0	0	0	1	0	1	0	0,4	1	1	STEMI
20	0,4	1	0	1	1	1	0	0	0.4	0,8	0,5	0	0	0	0,4	1	1	STEMI

Fig. 2. The sample of SVM input

Table 6. The accuracy of the best parameter - 10-fold cross validation.

Kernel		Polyn	omial	RBF				
Parameters	$\mathbf{C}$	$d/\sigma$	Accuracy	$\mathbf{C}$	$d/\sigma$	Accuracy		
Original Dataset	0.03	1	100%	0.01	1	47.9%		
Reduced Dataset	0.03	1	100%	0.01	1	48.1%		
k-NN Dataset	0.03	1	100%	0.01	1	47.9%		
<b>Pre-processing Dataset</b>	0.02	2	100%	0.8	1	98.9%		

Table 7. The accuracy of the best parameter pairs -percentage split.

Kernel		Polynoi	mial	RBF			
Parameters	DC	T (s)	Accuracy	DC	T(s)	Accuracy	
<b>Original Dataset</b>	70:30	27.49	100%	40:60	0.06	49.4%	
Reduced Dataset	70:30	21.37	100%	70:30	0.13	53.8%	
k-NN Dataset	80:20	24.55	100%	40:60	0.08	49.4%	
Pre-processing	70:30	0.06	100%	80:20	0.08	100%	
Dataset							

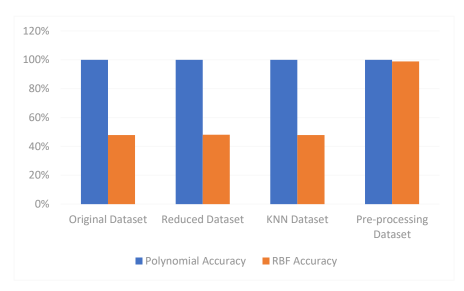


Fig 3. Dataset performance based on accuracy - 10-fold cross validation

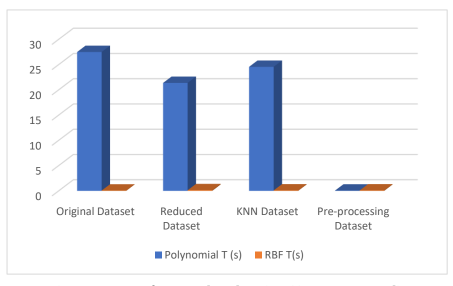


Fig 4. Dataset performance based on time (s) - percentage split

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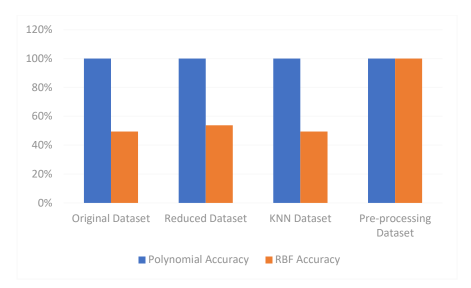


Fig 5. Dataset performance based on accuracy - percentage split

# 3.2. Testing

To evaluate the classification of CHD patient's dataset in SVM, the testing procedure was undertaken according to the Test Option Supplied on the Confusion Matrix formula [39]. The pre-processing dataset was put in place on 20% of tested data at C = 0.02 and d = 2 in the polynomial kernel and the values of C and  $\sigma$  are 0.8 and 1 respectively, in the RBF. In addition, the resemblance of SVM with another classifier, namely Multilayer perceptron Neural Network (NN) is operated to deeply observe the effectiveness of SVM. The confusion matrix for the above dataset of SVM and NN was explained in Table 8. This table showed that the classification in the pre-processing dataset for SVM is more accurate compared to NN, especially for RBF kernel. By comparing the values for error rate, precision and recall between polynomial kernel and RBF based on the confusion matrix computation as a side of SVM and NN, Fig. 6 is obtained. The figure showed that SVM for Polynomial kernel has 100% accuracy, "0" for error rate, and "1" for precision, and recall. Meanwhile, RBF kernel discharged from 51.79% into 100% accuracy, 0.48 into 1 for error rate, undefined into 1 for precision, and 0.52 into 1 for recall. Also, NN for polynomial kernel achieved 89% accuracy, "0.11" for error rate, and "0.89" for precision and recall.

Table o.	Comusic	ni Mati ix io	1 S v Ivi aliu i	111-1 Uly	iioiiiai aiiu	KDI.			
		SVM: Dat	aset Pre-pro	cessing					
		Polynomi	al		RBF				
Class		Prediction (	Class	Prediction Class					
	UAP	NSTEM	I STEMI	UAP	<b>STEMI</b>				
UAP	29	0	0	29	0	0			
NSTEMI	0	13	0	0	13	0			
<b>STEMI</b>	0	0	14	0	0	14			
Accuracy		100%			100%				
Error rate		0			0				
Precision		1			1				
Recall		1			1				
		NN-Mult	ilayer Perce	ptron					
Cla	ass	Pre	diction Class	<u> </u>					
	UAP	NSTEMI	STEMI						
UAP	18	0	1						
NSTEMI	2	18	1						
<b>STEMI</b>	0	2	14						
Accuracy	89%								
Error	0.11								

Table 8. Confusion Matrix for SVM and NN-Polynomial and RBF.

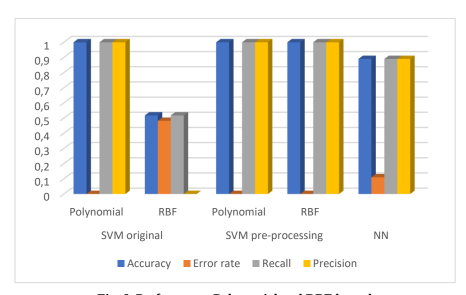


Fig. 6. Performance Polynomial and RBF kernel

## 3.3 Discussion

rate Precision

Recall

0.89

0.89

This result reveals that the pre-processing dataset in SVM provides significant values on the accuracy, error rate, precision, and recall, even though it exceeds NN capacity. As studied by [42], the SVM approach gives better predictive capability

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than other models, including NN. This, of course, has far-reaching implications in the medical context that require increasing sensitivity, specificity (the ability to predict the absence of the condition when it is not present) as well as discriminatory power of the classifier as key features to consider when comparing classifiers and diagnostic methods [45]. In the reviews on kernel type, the simulation presented that SVM polynomial is more reliable on the dataset changes compare to RBF. Consequently, the pre-processing prescription on SVM-RBF will undoubtedly boost RBF performance. Furthermore, selecting the specific kernel is an important research issue for kernel-based learning in the data mining area and the problem of SVM kernels is found in fitting the appropriate parameter values [46]. This investigation revealed that the SVM polynomial kernel mediates the accuracy and efficiency of the diagnostic results based on the parameters defined in CHD.

#### 4. Conclusions

This study successfully employed the SVM method in classifying the CHD patient's dataset. The simulation of the original, reduced, k-NN, and pre-processing datasets have shown the potential differences between polynomial and RBF kernel in terms of accuracy and processing time. The analysis of 10-folds cross-validation and percentage splits revealed the optimal pairs of parameters and data composition for polynomial and RBF kernel. Furthermore, the confusion matrix presented evidence that the pre-processing dataset delivered greater values in accuracy, precision, error rate, recall, and time model consumption than others. A comparative analysis between SVM and NN has shown the efficiency and accuracy of SVM in accelerating the unsurpassed classification of the CHD dataset with minimal errors. Therefore, this classification practically aids the doctors in suggesting medical assistance and taking a curative action. This result methodically answered the difficulties in choosing the SVM kernel function, which is flexible in changing the data set, optimal functionality, and time-consumption with high performance. Nevertheless, integrating SVM with other methods is a new solution to increase SVM performance for future work.

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Saturday, October 17, 2020

# **Letter of Acceptance**

# **To Whom It May Concern:**

This is to certify that

OKFALISA, LESTARI HANDAYANI, DINDA JUWITA P, MUHAMMAD AFFANDES, S.S.M.FAUZI3, SAKTIOTO

have submitted the paper entitled

# CORONARY HEART DISEASE USING SUPPORT VECTOR MACHINE

The paper has been reviewed and accepted for publication tentatively in

Volume 16, Issue 2 (April 2021).

The online publication of this paper is subject to a payment of USD300.

Best regards



Associate Professor Dr. Abdulkareem Sh. Mahdi Al-Obaidi, CEng MlMechE Executive Editor, Journal of Engineering Science & Technology http://jestec.taylors.edu.my/