Prediction of hypertension drug therapy response using K-NN imputation and SVM algorithm

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ABSTRACT

Hypertension is a degenerative disease but its healing takes a long time by consuming hypertension drugs until patient’s lifetime. The research is conducted to predict response of drug therapy using bioinformatics approach which is a blend of biological and informatics engineering methods. It is used medical record data of hypertensive patient in drug therapy which has an impact on genetic characteristics. The data is constructed as modelling for learning process. Then, it is implemented as a prediction whether the blood pressure is under control or not. However, the amount data have no values, then they are required to be applied preprocessing data. Therefore, this research is proposed K-Near Nearest Neighbor (K-NN) Imputation algorithm for refining data. After that, it is implemented using Support Vector Machine (SVM) algorithm for prediction. The experiment result is achieved the highest accuracy rate of 90% at the best parameter value \( \lambda = 0.9, \Sigma = 2, C = 0.1, \varepsilon = 0.001 \) in ten times iterations.

Keywords: Hypertension, K-NN imputation, Missing value, Prediction, SVM

1. INTRODUCTION

Hypertension is a degenerative disease but it is a serious problem in the world. Approximately, there are 65 million patients that are diagnosed hypertension in ratio 1:3 adults and it is 28% American’s prehypertension [1]. However, it is only 31% that the targeted blood pressure is achieved which the systolic blood pressure is less than 140 mmHg and or the diastolic blood pressure is less than 90 mmHg. In Indonesia, the prevalence of hypertension reaches 31.7%. Based on the previous research, it was found that 65.8% of patients in Harapan Kita polyclinic were detected having hypertension and only 39.3% reached the blood pressure target [2]. The 66% of hypertensive patients consume medicine on regular, and 60.7% of those have not reached the targeted blood pressure. As the previous research at the heart polyclinic of RSU dr. Saiful Anwar Malang shows that it is only 20.3% of hypertensive patients achieve the target of blood pressure [3]. These results show a high rate of uncontrolled hypertension.

Uncontrolled hypertension is a risk factor for cardiovascular events which is caused of the coronary heart disease and the cerebrovascular disease [4]. Based on the World Health Organization (WHO), uncontrolled hypertension is effect to 7 million deaths in each productive age and 64 million disabilities [5]. Therefore, the efforts to decrease blood pressure is addressed to achieve the targets by adequate therapy. It is a very important to reduce mortality and morbidity of the related hypertension diseases. The most causes of hypertension are multi-factorial, including the related activity of the renin angiotensin system, increasing sympathetic system, obesity, stress, excessive salt consumption and genetics. Angiotensin receptor blockers (ARBs) are powerful vasodilators for inhibitor of Renin Angiotensin Aldosterone System (RAAS) as well as Angiotensin Converting Enzyme inhibitors (ACEi). Therapy using ACEi and ARB has proven many clinical benefits and it is widely used in clinical practice [6]. Uncontrolled blood pressure may be related to...
adherence, choice of drug combinations, or genetic variant molecules which are involved in RAAS. Research in Japan showed that the renin C-5312T variant, which nucleotide substitution from C to T in the nucleotide sequence-5312, was an independent predictor of resistance for ARB users in Japan [7].

The other hand, many researchers have developed tools for diagnosis some diseases using bioinformatics approach including machine learning algorithm. The previous research has been conducted on pairwise DNA sequence alignment between hepatitis B virus (HBV) and hepatocarcinoma (HCC) using modified dynamic programming which improved performance of computation space and time [8]. Also, the related research have been conducted for classification of breast cancer using logistic regression, diabetic analytics using data mining approach. Beside that, classification of brain tumor image segmentation is implemented using hybrid strategy for clustering and segmentation method [9-11]. Furthermore, Support vector Machine (SVM) is one of machine learning algorithms which has high accuracy in medical research. This algorithm is applied to detect hypertention based on radial pulse wave and some risk factors obesity, stress, systolic and diastolic blood pressure, physical exercises, cigarette consumption and diet lifestyle [12-13]. However, the involved data of this research is incomplete. Several features have null values. Therefore, this study is purposed the K-NN Imputation and SVM algorithm to implement a prediction system based on the characteristics of genetic variation in hypertensive patients against drug therapy. The system is developed using patient data that treats hypertention drugs including polymorphime from the angiotensin and renin genes which have an important role in the cardiovascular system. The first step is preprocessing data to solve the missing value using KNN-Imputation, then it will be constructed the model in SVM method in order to predict the drug therapy respond for hypertensive patients.

2. HYPERTENSION PATHOLOGY

Hypertension is a complex pathophysiological disease. Systemic blood pressure regulation is multifactorial. It is basically the end result of cardiac autoregulation and peripheral vascular resistance (Figure 2.1) [14]. The renin system angiotensin aldosterone plays an important role in the regulation of blood pressure, electrolyte balance, and the pathogenesis of atherosclerosis [15-17]. In the initial phase is increasing RAAS activity which occurs by increasing production of angiotensin and or expression or activity of Renin (REN). The Renin catalyzes the breakdown of angiotensinogen (AGT) into angiotensin I. Then, it is catalyzed to angiotensin II with angiotensin converting enzyme (ACE). Furthermore, Angiotensin II increases blood pressure through strong vasoconstriction and sodium retention [15-16]. As an illustration it can be shown in Figure 1.

Figure 1. The renin system of angiotensin aldosterone (RAAS) in regulating blood pressure
Furthermore, based on the pathophysiological complexity of hypotension there are various anti-hypertensive therapies which aim to inhibit pathophysiology. There are many classes of antihypertensive drugs such as RAAS inhibitors consisting of ACEi and ARB, calcium channel blockers (CCB), diuretics, nervous system inhibitors sympathetic such as beta blockers, alpha blockers, and the details are shown in Table 1 [18].

| Type of Drug                          | Name of Drug       | Dosage                                      |
|---------------------------------------|--------------------|---------------------------------------------|
| ACE inhibitors                        |                    |                                             |
| Captopril                             |                    | 12.5-50 mg twice daily                      |
| Enalapril                             |                    | 5-40 mg once daily or in two equally divided doses |
| Fosinopril                            |                    | 10-40 mg once daily                         |
| Lisinopril                            |                    | 5-40 mg once daily                          |
| Perindopril erbumine                  |                    | 4-8 mg once daily                           |
| Perindopril arginine                  |                    | 5-10 mg once daily                          |
| Quinapril                             |                    | 5-40 mg once daily or in two equally divided doses |
| Ramipril                              |                    | 2.5-10 mg once daily or in two equally divided doses |
| Trandolapril                          |                    | 1-4 mg once daily                           |
| Calcium channel blockers-dihydropyridine |                    |                                             |
| Amlopipline                           |                    | 2.5-10 mg once daily                        |
| Felodipine                            |                    | 5-20 mg once daily (controlled release)     |
| Lercanidipine                         |                    | 10-20 mg once daily                         |
| Nifedipine                            |                    | 10-40 mg once daily (conventional)          |
| Calcium channel blockers-nondihydropyridine |                | 20-120 mg once daily (controlled release)  |
| Diltiazem                             |                    | 180-360 mg once daily (controlled release) |
| Verapamil                             |                    | 120-240 mg once daily (controlled release) |
| Angiotensin II receptor antagonists   |                    |                                             |
| Candesartan                           |                    | 8-16 mg once daily                          |
| Eprosartan                            |                    | 600-800 mg once daily                       |
| Irbesartan                            |                    | 150-300 mg once daily                       |
| Losartan                              |                    | 50-100 mg once daily                        |
| Telmisartan                           |                    | 20-80 mg once daily                         |
| Olmesartan                            |                    | 20-40 mg once daily                         |
| Thiazide diuretics                    |                    |                                             |
| Chlorthalolone                        |                    | 12.5-25 mg once daily                       |
| Hydrochlorothiazide                   |                    | 12.5-25 mg once daily                       |
| Indapamide                            |                    | 12.5-2.5 mg once daily                      |
| Beta-blockers                         |                    |                                             |
| Bisoprolol                            |                    | 1.25-10 mg once daily                       |
| Atenolol                              |                    | 25-100 mg once daily                        |
| Carvedilol                            |                    | 12.5-50 mg once daily                       |
| Metoprolol tartrate                   |                    | 100-400 mg twice daily                      |
| Metoprolol succine (controlled release)|                | 50-100 mg twice daily                       |
| Osxrenolol                            |                    | 12-190 mg daily                             |
| Other                                 |                    |                                             |
| Clonidine                             |                    | 40-160 mg twice daily                       |
| Hydralazine                           |                    | 50-300 μg twice daily                       |
| Methyldopa                            |                    | 12.5-100 mg twice daily                     |
| Moxonidine                            |                    | 125-500 mg twice daily                      |
| Prazosin                              |                    | 200-600 μg daily                            |
|                                      |                    | 0.5-10 mg twice daily                       |

3. RESEARCH METHOD

The system consists of two main stages, are preprocessing data using K-NN Imputation algorithm and prediction using SVM method as shown in Figure 2.
3.1. K-Nearest Neighbor (K-NN) Imputation

K-Nearest Neighbor (KNN) Imputation is a method to get the attribute value based on the similarity between new cases and old cases at the appropriate features. According to Olivas [19], it is technique of Machine Learning to handle missing values data by imputation under considering the most similar record values.

At the first stage, input data is taken from medical record of hypertensive patients. However, the number of ignorance data is required to fill values using this method which is based on the appropriate data similarity. By separately, the data is selected which has complete value and has incomplete values. The next process is calculating the euclidean distance of data value to be shorted the similarity. Finally, it is selected to the most frequency values.

3.2. SUPPORT VECTOR MACHINE (SVM)

The Support Vector Machine (SVM) method is a linear classification method by finding the best hyperplane that functions as a separator of two classes in input space. The basic concept of Support Vector Machine is linear classifiers, and then it is developed into non-linear classifiers by incorporating kernel tricks in high-dimensional space as in Figure 3.

In Figure 3, it is illustrated the SVM method. The thick black line in the middle is a hyperplane that separates data +1 and data -1, in this study the data to be used is positive review data and negative review data. The closest point to the hyperplane is called Support Vector. The distance between a support vector and a hyperplane is called a margin. A support vector is a point that intersects a small black line.

Basically, SVM method is a linear classifier that can only be used for linear data. Therefore, it is developed by adding a kernel trick in order to classify non-linear data. The classified data must be transformed to the vector space in high dimension. The kernel trick functions that can be used in non-linear SVM classifications are Polynomial, Gaussian (RBF) and Sigmoid. Each label is denoted y_i ∈ {-1, +1} for i = 1, 2, ..., n, where n is the amount of data. It is assumed that +1 and -1 classes can be completely separated from the hyperplane, which is defined:

\[ w \cdot x + b = 0 \]  \hspace{1cm} (1)

Data xi is included into -1 and it is as stated in the (2).

\[ w \cdot x_i + b \leq -1 \]  \hspace{1cm} (2)

Data xi is included into +1 and it is as stated in the (3)
\[ w \cdot x_i + b \geq +1 \]  

(3)

The largest margin is calculated by maximizing the distance between the nearest point and hyperplane.

\[ \frac{1}{\|w\|} \]  

(4)

In general, therefore the real problem is a non-linearly separable form, then two classes cannot be separated by a hyperplane completely. Therefore, SVM modification is needed by entering kernel functions. The non-linear SVM concept is to change the data \( x \) that is mapped by the function \( \Phi(x) \) to a higher dimensional vector space. This mapping aims to represent data in the new vector space. The learning process of SVM is finding support vectors by dot product data that has been transformed into the new space. The dot product value can be calculated without knowing the data transformation process \( \Phi \). The kernel function provides convenience in the SVM learning process to determine support vectors in non-linear data [20]. The kernel function can be formulated as (5)

\[ K(x_i, x_j) = \Phi(x_i) \cdot \Phi(x_j) \]  

(5)

In the SVM method, Radial Basis Function (RBF) is a kernel to be applied in this research as in (6)

\[ K(x_i \cdot x_j) = \exp\left(-\frac{\|x_i-x_j\|^2}{2\sigma^2}\right) \]  

(6)

The next step is to make predictions by implementing the Sequential Support Vector Machine method including: calculation of the Hessian matrix, iteration to reach the maximum in least error rate or Max (| \( \delta \alpha \) |) <\( \epsilon \). After that, the bias and similarities between the testing data and training data are calculated. As a result, it will be obtained the positive or negative classes as it shows in Figure 4. It is a flow diagram of the Support Vector Machine sequential process.

4. RESULT AND DISCUSSION

The data is taken from Syaiful Anwar Malang Hospital, at heart polyclinic. The feature details are as in Table 2 with balanced classes, which have the same total of data in each class. However, before testing with different data, validation tests are achieved with accuracy rate of 100%. This shows the system that is built is reliable.

| Feature            | Remark                                      |
|--------------------|---------------------------------------------|
| Code               | Identify of patient                         |
| Gender             | Male/ Female                                |
| Age                | Birth date                                  |
| Ethnic             | Javanese or others                          |
| Waist C            | Waist circumference                         |
| Hip C              | Hip Circumference                           |
| Weight             | Weight of patient                           |
| Height             | Height of patient                           |
| Smoking            | Active/ passive                             |
| Menopause          | Stop of menstruation                        |
| Hypertention       | Historical Hypertension                     |
| Ur                 | Ureum                                       |
| Cr                 | Creatine                                    |
| HDL                | High Density Lipoprotein                    |
| LDL                | Low Density Lipoprotein                     |
| TG                 | Trigliserida                                |
| Cholesterol        | Level of cholesterol total                  |
| Glycemia           | Level of blood sugar                        |
| AGT pre            | Level of Angiotensinogen at pre intervention (drug) |
| AGT post           | Level of Angiotensinogen at post intervention (drug) |
| DeltaAGT           | Difference between AGTPre and AGTPost       |
| Difference AGT     | up/ down                                    |
| Quartile AGT       | Quartile of Angiotensinogen level            |
| Q3AGTPost          | The last quartile of Angiotensinogen         |
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The interface of prediction system is shown at Figure 5. There are some input parameter for training and testing data including lambda, gamma, C, epsilon and the number of iteration.

![Figure 5. Implementation interface](image)

Then, based on the experiment result that the accuracy rate is achieved at 90%. It is applied at the best parameter value as in Table 3.

| Parameter  | Value |
|------------|-------|
| lambda     | 0.9   |
| sigma      | 2     |
| C          | 0.1   |
| Epsilon    | 0.001 |
| Number of iteration | 10   |

The accuracy rate of testing result is effect to many factors including preprocessing data of missing value using KNNI algorithm which have accuracy rate of 87%. Therefore, the next process for prediction is achieved of 90%.

5. CONCLUSION

Identification system of drug therapy response for hepatitis patients has been applied using a combination of K-Nearest Neighbor (K-NN) Imputation and Support Vector Machine (SVM) algorithm. The amount missing value data have been covered using K-NN Imputation based on the similarity measure of the attribute value. It was applied at the initial stage before implemented SVM algorithm for prediction. Overall, the accuracy result is achieved of 90%.

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