Data Mining in Dermatological Diagnosis: A Method for Severity Prediction

Manjusha K. K  
Research Scholar  
Karpagam University  
Coimbatore, India

Sankaranarayanan K  
Dean (PG Studies)  
Sri Ramakrishna Institute of Technology  
Coimbatore, India

Seena P  
Asst. Professor  
Dept.of Dermatology  
Govt. Medical College  
Kottayam, India

ABSTRACT
Medical diagnosis is an important but complicated task that should be performed with great accuracy and efficiency. In dermatology there are a lot of diseases which shows similar pattern in appearance and symptoms. So diagnosis of a single patient can differ significantly if he is examined by different physicians. So we need a standard format for predicting such a disease. Today, automated medical analysis help doctors to diagnose and predict diseases, at a very fast pace. This study goes through different dermatological diseases with similar symptoms which may even prove fatal if left unattended at exact time. Medical dataset used for this work contain 230 instances with 22 attributes. In this paper we have experimented on data gathered from the southern part of Kerala, India. For better prediction calculation of accuracy of mining algorithm is important. Two data mining classification algorithm Naive Bayes’(NB) and J48 was used for data analysis. Weka Open Source Software, a build in software tool for data mining was used. The GUI developed in Java, reveals the chances of different dermatological disease and also finds out the probabilities of occurrence of each disease.

Keywords
Data mining, Dermatology, Diagnosis, Weka.

1. INTRODUCTION
Data Mining is the process of extracting hidden knowledge from data. Classification algorithms always find a useful rules or classes from large amount of data. Data mining application includes several fields like banking, insurance and Crime detection including health care. Medical Industry faces many problems due to the increase of types of diseases and their specific management. In addition, the amount of data generated by healthcare transactions is too large, diverse and complex to be analyzed by traditional methods. The application of data mining on medical data can foreground new, useful and potentially lifesaving knowledge. Data mining in medical analysis helps to increase diagnostic accuracy, reduce treatment cost and save human resources [1]. Knowledge discovery in medical databases is a well-defined process and data mining is an essential step. Data mining is, in short, “Knowledge mining from data”. Data mining is the process of analyzing data from different views and summarizing it into useful information. Classification algorithms find a set of rules to represent data into classes. It includes two steps; the first step tries to find a model for the class attribute as a function of other variables of the datasets. In the second step the related class of each record is determined by applying formerly designed model on the new and unseen dataset [2]. A popular algorithm based on probability theory is Naive Bayes’ algorithms. A predictive model algorithm for classification task is induction of decision trees.

2. BACKGROUND
The large growth of medical databases available in technologically advanced countries has motivated medical researchers in those countries to use data mining for knowledge discovery from these databases. With the steady increase in the volume of stored data, data mining techniques assume an increasingly important role in arriving at patterns and extracting knowledge to provide better patient care and effective diagnostic capabilities. This makes it difficult to analyze the data in order to make important decision regarding patient health. So, it becomes essential to generate a powerful tool for analyzing and extracting important information from this complex data and derive a vital knowledge from it for future reference and research. The analysis of health data can provide a great boost to healthcare by enhancing the performance of patient management tasks. Data mining technologies can provide benefits to healthcare organization for grouping the patients having similar type of diseases or health issues so that healthcare organizations can prescribe the most effective treatments [3]. Data mining applications can be developed to evaluate the effectiveness of medical treatments. By comparing and contrasting causes, symptoms and courses of treatments, data mining can deliver an analysis of the most effective courses of action. For example, the outcomes of patient groups treated with different drug regimens for the same disease or conditions can be compared to determine which treatments work best and are most cost effective. Data mining is vital in the most critical sector of healthcare both in developed and developing countries. Considering the immense benefit it has ushered in, one can envisage the tremendous scope for furthering studies in this field.

2.1 Decision tree
Decision trees are one of the most powerful tool in data mining and knowledge discovery. It has been used in analysis of large and complex bulk of data in order to discover useful patterns. The basic decision tree algorithm is called ID3 (Iterative Dichotomizer3). ID3 can handle only discrete values, but the successor C4.5 can handle numeric values. Classification and Regression Trees (CART) approach is suited for analysis of categorical and continuous datasets. J48 is the implementation of ID3 algorithm developed by the WEKA [4]. It can handle different types of data like numeric, nominal, textual data and can also process incorrect or missing values. J48 can be implemented in data mining packages in different platforms and easy to understand
because of its presentation. J48 show high performance with small effort.

2.2 Naive Bayes

In medical data mining, Naive Bayes classification plays an important role. It is a probabilistic classification based on the Bayes theorem.

The probability of data record $X$ having the class label $C_i$ is

$$P(X|C_i) P(C_i)$$

$$P(C_i|X) = \frac{P(X|C_i) P(C_i)}{P(X)}$$

The class label $C_i$ with largest conditional probability value determines the category of the data record.

It is very practical when the dimensionality of the inputs is high. The word “Naive” implies the independence between all attributes. Naive Bayes (NB) is a machine-learning method that has been used for over 50 years in biomedical informatics [5]. It requires only small amount of training data to estimate the parameter which is very useful for health care applications [6]. Naive Bayes computes conditional probabilities of the classes given with the instance and select the class with highest posterior [7]. Regardless of this simplified assumption and naive design, naive bayes classifier works well in many complex real world situations. Bayes classification is outperformed by current approaches, like boosted trees or random forests.

2.3 Dermatological Diseases

This research work was directed towards the prediction and analysis of some commonly seen skin diseases and symptoms focusing on the relationship of symptoms to find important factors and rules that affect skin disorders. The selected conditions have certain common features and this part of the work discusses the eight types of dermatological conditions namely rubella, Kawasaki disease, scarlet fever, fifth disease (erythema infectiosum), no vaccination subitum, (exantheme subitum or roseola infantum), measles, chickenpox and entrovirus.

3. RELATED WORKS

Chang et al [8] conducted five experiments focusing on six major skin diseases and used decision tree of data mining combining with neural network classification methods to construct best predictive model in dermatology. The study predicted and analysed six commonly seen skin diseases namely psoriasis, seborrhic dermatitis, lichen planus, pityriasis rosea, chronic dermatitis and pityriasis rubra pularis. All classification technology could predict the disease with considerable accuracy with neural network model having the highest level of accuracy of 92.62%.

Zeon et al [9] developed a disease prediction system, DOCAID, for predicting typhoid, malaria, jaundice, tuberculosis and gastroenteritis based on patient symptoms and complaints employing Naive Bayes Classifier algorithm [9]. An accuracy of 91% accuracy in predicting the diseases were reported by the authors.

Theodorali et. al [10] developed prediction model for predicting the final outcome in patients suffering from severe injuries after accident. The analysis included a comparison of data mining techniques using classification, clustering and association algorithms. Using this analysis they obtained results in terms of sensitivity, specificity, positive predictive value and negative predictive value and compared the results between different prediction models.

4. METHODOLOGY

The research work is structured into 3 stages as represented in figure 1. The first stage includes data collection & pre processing and producing training data and analyzing variables. In the second stage we use Weka tool to check the accuracy of the models. The third stage presents explanation of the prediction model

4.1 Data collection

Data was collected from various tertiary health care centres in Kottayam and Alappuzha districts of Kerala. The study was developed on the basis of the survey and the questionnaires are prepared and filled out by skin specialist in Kottayam Medical College. After filtering and correcting missing values we got 230 skin disease data and this data is used for prediction of eight skin diseases which show similar symptoms. Among all, 44 pieces of data were chickenpox, 28 rubella, 29 Kawasaki disease, 31 scarlet fever, 41 measles, 27 fifth disease, 12 entrovirus and 18 no vaccination subitum. In this study 21 factors exist and all are clinical attributes. The attributes are temperature (>90°F or <90°F), level of fever (90-95, 95-102, >102), duration of fever (number days fever started), type of exantheme (Maculopapules, vesicular, maculopapular rash), progressive exantheme (slowly or quickly), painful exantheme (Y,N), type of exantheme (kopik spot, pharyngitis, blisters, aarbeiting), localization (face, neck, cheek, body), conjunctivitis, seasons (winter, summer, autumn, spring), age (baby, young, elder ), prior contact, patient medication (medicine A, medicine B, medicine C), whether patient is vaccinated or not, recent journey etc.

4.2 Data analysis software

Weka (“Waikato Environment for Knowledge Analysis”) is a popular suite of machine learning software written in Java, developed at University of Waikato, New Zealand [11]. For analysing data we are using Weka 3.7.9. It contains a
collection of algorithms for data analysis and predictive modelling, together with GUI for easy access to this functionality. Weka supports several standard data mining tasks like data pre-processing, classification, clustering, association rules, visualization and feature selection. It also offer different test option like Cross validation, using training set, test set, percentage split etc. We have used Naive bayes method, J48 decision tree to perform the mining and classification process.

4.3 Data pre-processing
The database of this study contains a total of 230 skin disease cases. After data pre-processing all data was divided into two sections: They were the training data set with 180 and test data set with 50 respectively to verify the accuracy of predictive model. The data is analyzed and implemented in WEKA tool. Data mining finds out the valuable information hidden in huge volumes of data. Weka tool is a collection of machine learning algorithms for data mining techniques, written in Java. It consists of data pre-processing, classification, regression, association rules, clustering and visualization tools. We have used Naive bayes method and J48 decision tree to perform the mining and classification process. We have used 10 folds cross validation to minimize any bias in the process and improve the efficiency of the process.

5. RESULT AND DISCUSSION
The Graphical User Interface developed in Java Apache-Tomcat-5.5.35 and the results when the diagnosis on the basis of imported input has been shown in the figure 2.

Fig 2: Graphical user interface

This software gave accurate results, with the consultation of doctors. The doctors can input symptoms into the software and get the most probable disease from the eight diseases. The data was compared using weka software. The results of the experimentation are shown in figure 3.

Fig 3: Experimentation results

The results showed that all the eight diseases depend on all attributes. The attributes fever, season, vaccination, headache and aarbeitong play an important role for the prediction. 197 patients value depends on fever. Vaccination is also an important factor, because most of the disease depends on whether the person is vaccinated or not.

The data set used for the experiment contains 230 instances with 22 attributes and eight class attributes to test and substantiate the difference among classification algorithms. The classification algorithm includes Naive Bayes, J48. After analysing data with WEKA tool results shows that the highest correctly classified instances is 228 (99.13%) by decision tree and naive bayes shows 201 (87.4%) correctly classified instances respectively (figure 4).

Fig 4: Simulation result of data mining algorithm

The time taken is also an important parameter when we are comparing results. Naive Bayes classification requires only 0.01s and J48 requires 0.03s.

The work was also targeted in comparing the performance of the various algorithms with dataset as given in figure 5. The result showed that Naive bayes produced less precision and true Positive rate as compared with J48 algorithm. J48 is more efficient in all parameter like TP-rate, FP-rate, Precision, Recall and ROC area.
Doctors to clear their confusion when predicting diseases using J48. Thus, we conclude that this software helps the best to predict the demographic dependence of the disease. The data for the system predicts eight different skin diseases and can help only; we can extend it with other parameters. This expert system also helps to save the time and expense of patients. This can be extended to predict other type of diseases or we can use the same dataset with other data mining techniques. However computer aided diagnosis must be regarded only as one form of supportive measure. Medical practitioner’s responsibility and general patient medical care must be given the most priority.

7. REFERENCES

6. CONCLUSION
A computer aided model has been developed for the analysis of different disease. In this software we are using 22 attributes only; we can extend it with other parameters. This expert system predicts eight different skin diseases and can help doctors to predict the disease confidence. The data for the study was collected from a limited region and future study can be done by collecting data from a wide region which will help to predict the demographic dependence of the disease. The best prediction model was obtained with algorithm developed using J48. Thus, we conclude that this software helps the doctors to clear their confusion when predicting diseases with similar symptoms and helps to take better decision. The expert system also helps to save the time and expense of patients. This can be extended to predict other type of diseases or we can use the same dataset with other data mining techniques. However computer aided diagnosis must be regarded only as one form of supportive measure. Medical practitioner’s responsibility and general patient medical care must be given the most priority.

Fig 5: Performance study of data mining algorithms
Confusion matrices produced by J48 and Naive Bayes are given in table 1 and table 2 respectively.

Table 1. Confusion matrix of J48

<table>
<thead>
<tr>
<th>A</th>
<th>b</th>
<th>C</th>
<th>D</th>
<th>e</th>
<th>F</th>
<th>g</th>
<th>h</th>
<th>Classified as</th>
</tr>
</thead>
<tbody>
<tr>
<td>28</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>a=rubella</td>
</tr>
<tr>
<td>0</td>
<td>29</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>b=kawasaki</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>31</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>c=scarlet</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>41</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>d=measles</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>27</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>e=fifthdisease</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>44</td>
<td>0</td>
<td>0</td>
<td>f=chickenpox</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>12</td>
<td>0</td>
<td>2</td>
<td>g=entrovirus</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>16</td>
<td>h=novaccination</td>
</tr>
</tbody>
</table>

Table 2. Confusion matrix of naïve bayes

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>c</th>
<th>d</th>
<th>e</th>
<th>F</th>
<th>g</th>
<th>h</th>
<th>Classified as</th>
</tr>
</thead>
<tbody>
<tr>
<td>28</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>a=rubella</td>
</tr>
<tr>
<td>0</td>
<td>29</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>b=kawasaki</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>31</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>c=scarlet</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>0</td>
<td>39</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>d=measles</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>9</td>
<td>7</td>
<td>11</td>
<td>0</td>
<td>0</td>
<td>e=fifthdisease</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>43</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>f=chickenpox</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>6</td>
<td>3</td>
<td>3</td>
<td>g=entrovirus</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>18</td>
<td>0</td>
<td>h=novaccination</td>
</tr>
</tbody>
</table>