

Pertaining to comparison of algorithms for Breast Cancer Diagnosis

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Abstract: In this paper an attempt is made to study the applicability of machine learning algorithms that try to predict the breast cancer type based on the data set. The goal of ML Algorithms is to predict and diagnosis to have a trained learning algorithms that gives the gene expression levels from cancer patient. The aim of this study is to compare different classification learning algorithms significantly to predict a benign from malignant cancer in Wisconsin breast cancer dataset. This study used the Wisconsin breast cancer dataset to compare five different learning algorithms, Bayesian Network, Naïve Bayes, Decision trees J4.8, ADTree, and Multi-layer Neural Network along with t-test for the best algorithm in terms of prediction accuracy. The experiment, has shown that Bayesian Network is significantly better than the other algorithms. Requirements are Mat Lab, Weka Tool, Java and Relational Database.

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I. INTRODUCTION

The diseases which cost so many lives, diagnostic computer-based applications are used widely. Robotics are playing a very important role in operating rooms. Also, the expert systems are presented in the intensive treatment rooms. In turn, using another aspect of Artificial Intelligence for breast cancer diagnosis is not unworthy. It is reported that breast cancer disease is the second most common cancer that affects women, and was the prevalent cancer in the world by the year of 2002[21]. Macmillan Cancer Support in London reports that, in the UK, breast cancer affects a significant number of Arab women and a small number of Arab men. This cancer is a very common type of cancer among women and the second highest cause of cancer death. In the United States, about one in eight women over their lifetime has a risk of developing breast cancer [1]. Breast cancer begins with the uncontrolled division of one cell inside the breast and results in a visible mass, called a tumor. The tumor can be either benign or malignant. The accurate diagnosis in determining whether the tumor is benign or malignant can result in saving lives. Therefore, the need for precise classification within the clinic is a cause of great concern for specialists and doctors. This importance of Artificial Intelligence has been motivated for the last 25 years, when scientists began to realize the complexity of taking certain decisions to treat particular diseases. The use of machine learning and data mining as tools in medical diagnosis becomes very effective and one of the critical diseases in medicine where the classification task plays a very essential role is the diagnosis of breast cancer. Therefore, machine learning techniques can help doctors to make an accurate diagnosis for breast cancer and make the correct classification of being benign or malignant tumor. There is no doubt that evaluation of data taken from the patient and decisions of doctors and specialists are the most important factors in the diagnosis, but expert systems and artificial intelligence techniques such as machine learning for classification tasks, also help doctors and specialists in a great deal

The aim of this paper is to investigate different machine learning techniques. It will use several algorithms and apply them on *Wisconsin breast cancer dataset*. Also focus on five machine learning techniques; Bayesian Network, Naïve Bayes, ADTree, J48, and Multilayer Neural Network (back-propagation) and analyze their results by study of these various algorithm.

The following sections give a brief introduction for the algorithms used in the experiment, and then discussion about the result and the analysis. Finally, frame the future work in the conclusion section.

II. BAYESIAN NETWORK

Bayesian Networks are graphical structures that give a chance to represent and reason about an uncertain domain. Bayesian Networks can be viewed as the merger between the probability theory and the graphical Theory. Bayesian Networks describe the probability distribution governing a set of variables by specifying a set of conditional independence assumptions along with a set of conditional probabilities. The nodes in the network represent variables(either discrete or continuous), and arcs represent the dependencies between these variables. The important concept in Bayesian Network is the conditional independence (Figure 1).

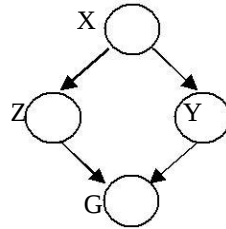


Figure.1 A simple structure of Bayesian Network.

The structure in Figure.1 shows that the probability of G is conditionally independent from X, given the probability of Z, Y and the relationship can be given in the following equation:

$$P(G|X,Z,Y) = P(G|Z,Y)$$

Each node is asserted to be conditionally independent of its non-descendants, given its immediate parents. This in fact, reduces the complexity of learning the target function. The joint probability distribution of all nodes in the network can be describe as follows:

$$P(y_1, y_2, y_3, \dots, y_n) = \prod_{i=1}^n P(y_i | \text{Parents}(y_i))$$

III. NAÏVE BAYES

Naïve Bayes are applied into learning tasks where each instance x is described by a conjunction of attribute values and where the target function $f(x)$ can take on any value from some finite set V. A set of training examples of the target function is provided, and a new instance is presented, described by the tuple of attribute values $\langle a_1, a_2, a_3, \dots, a_n \rangle$. The learner is asked to predict the target value for the new instance. The Bayesian approach used to classify the new instance is to assign the most predictable target value, given the attribute values $\langle a_1, a_2, a_3, \dots, a_n \rangle$ that describe the instances.

$$V_{NB} = \text{argmax} \prod_{i=1}^n P(a_i | V_j) P(V_j)$$

Naïve Bays can be considered as a special case of Bayesian Network (Figure.2).

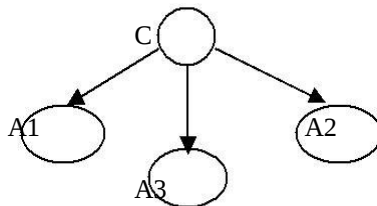


Figure.2 Naïve Bayes as a special case of Bayesian Network.

In Figure.2, each variable(A_i) is conditionally independent from other variables, given its class(C). However, the assumption in the Naïve Bayes has more constraining than the global assumption in the Bayesian Network, where in Bayesian Network the variables are governed by specifying a set of conditional independence assumptions along with a set of conditional probabilities.

IV. NEURAL NETWORK-BASED ALGORITHMS (NN)

A neural network can be defined as a model of reasoning based on the human brain [15]. Mainly, our brain consists of an intensive set of nerve cells connected to each other. The brain in humans contains about 10 billion and 60 trillion connections (synapses). When the brain uses multiple neurons in parallel, it can accomplish jobs much faster than the fastest computers nowadays.

Typically, the neuron consist of soma which represents the cell body, a number of fibers called dendrites, an axon which is a single long fiber , and synapses that represents connections between cells(Figure.3).

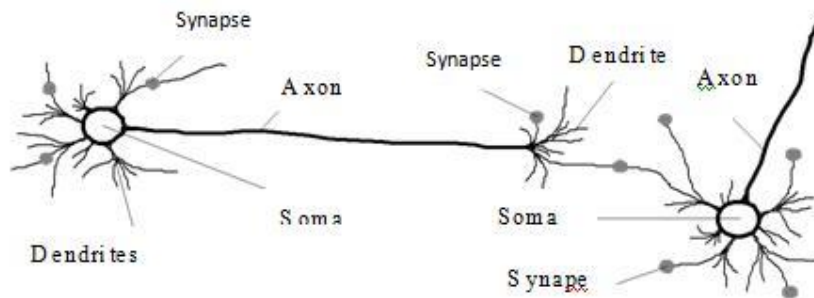


Fig.3. Biological Neural Network[15].

Similarly, an artificial Intelligence (ANN) is an interconnected group of artificial neurons that are based on a mathematical model to process information through neurons. ANN is sometimes called an adaptive system because it has the ability to change its structure based on the information being processed in the network. ANN's structure has input layers, hidden layers, and output layers (Figure.4).

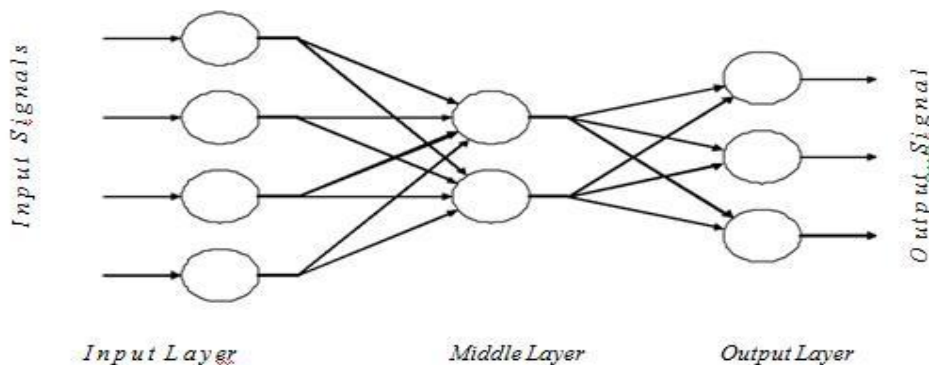


Fig.4. Artificial Biological Neural Network [15].

The figure above shows that the neurons are connected by links, and each link has a numerical weight associated with it. The weights express the strength of each neuron input and the learning is achieved by adjusting the weights through the links.

4.1. The Perceptron

The perceptron is considered as the simplest form of neural networks, and is based on the McCulloch and Pitts neuron model [15]. It has just a single neuron with adjustable synaptic weights and a hard limiter (Figure.5).

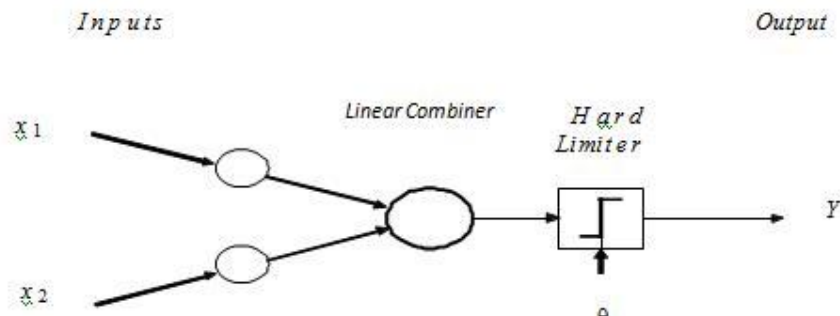


Fig.5. Single-layer two-input perceptron [15].

4.2. Multilayer neural network

A multilayer neural network is a feed-forward neural network (Figure.6). It has an input layer, one or more hidden layers, and output layer. The most popular algorithm is back-propagation which was first proposed in 1969 [15].

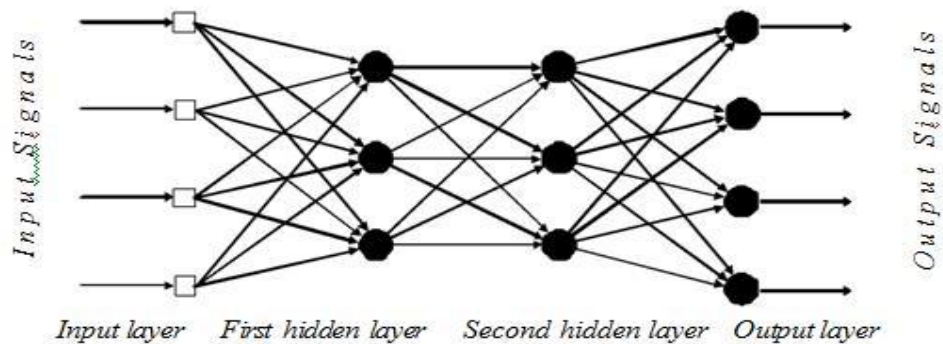


Fig6. Multilayer perception with two hidden layers [15].

Learning in multilayer network follows the same scheme as for a perceptron. A training data is presented to the network, and then the network computes its output pattern. If there is an error (difference between actual and desired output patterns), the weights in this case are adjusted to reduce the error by propagate the error backward through the output layer and hidden layers and then repeat the process again until reach the certain criteria.(The sum of squared error >- sufficient small number).

V. DECISION TREES ALGORITHMS

The trees are among the most popular inductive inference Decision algorithms. The learned in the decision trees is represented by a tree. Also, after the function decision tree is constructed the IF...THEN statements can be inferred easily.

5.1. ID3

The ID3 technique to build decision trees (with just nominal attributes) is based on the information theory and attempts to minimize the expected number of comparisons. The concept used to quantify information is *entropy*. Entropy is used to measure the amount of uncertainty, surprise or randomness in a set of data. Therefore, the ID3 finds the attribute that has the highest gain information or the lowest Entropy. Certainly, when all data in a set belongs to a single class, there is no uncertainty. In this case, the entropy is zero. $Entropy(S) = -p^+ \log_2(p^+) - p^- \log_2(p^-)$.

The goal of decision trees is to iteratively partition the given dataset into subsets, where all elements in each final subset belong to the same class[6].

5.2. C4.5

The decision tree algorithm C4.5 was designed by Quinlan [19]. It is designed to improve ID3 algorithm in different ways such as:

- Nominal and numeric data: the C4.5 can be used to classify either nominal or numeric attributes.

- Missing data: When the decision tree is built, missing data is simply ignored. To classify a record with a missing attribute value, the value for that item can be predicted based on what is known about the attribute values for the other records.

5.3. Alternating Decision Tree (ADTree)

The ADTree considered as another semantic for representing decision is trees[9]. ADTree, each decision node is replaced by two nodes: a prediction node (represented by an ellipse), and splitter node (represented by a rectangle). The decision tree in ADTree algorithm is identical while the prediction node is associated with a real valued number. As it is stated in the decision tree, an instance is mapped into a path along with the tree from the root to one of the leaves. However, unlike decision trees, the classification in ADTree that is associated with the path is not the label of the leaf. Instead, it is the sign of the sum of the prediction along the path.

VI. THE EXPERIMENT WITH WEKA TOOL

The Weka tool is a collection of machine learning algorithms, and data preprocessing tools [24]. It provides a complete implementation for data sets being used. Weka tool has a variety of methods for transforming data sets, such as the algorithms for discretisation and filtering. Generally, the Weka tool has the methods for all standard machine learning algorithms (regression, classification, clustering, association rules, and attribute selection, etc.) [24]. The data sets in Weka take the form of a single relation table in the ARFF format (Attribute- Relation File Format). The Weka tool has the ability to prepare the input data, evaluate learning methods statistically, and visualizing the input data with its result (output) during learning process.

6.1. 1 Preparing datasets in Weka

The standard way of representing datasets in Weka tool is called an ARFF (Figure.7).

```
@relation <name>
@attribute <attribute-name><type>
@data
```

Fig7. ARFF structure.

6.1.2 ARRF Structure

6.1.2.1 Header:

The header section of an ARFF file is very simple and merely defines the name of the dataset along with set of attributes and their associated types [20]. The following tags are considered when the file is constructed:

6.1.2.1.1 @ Relation:

The relation<name> tag is declared at the beginning of the file, where <name> is referred to the relation name you intend to use.

6.1.2.1.2 @ Attribute:

Attributes are defined as @attribute <attribute-name><type>, and can take either numeric, nominal, or string values.

6.1.2.1.3 @Data [<data-format>].

The instances in @data tag are written one per line with values for each attribute in turn, separated by commas. If a value is missing, it is represented by a single question mark "?". The class attribute is always written as a last attribute.

6.2 The experiments with dataset

6.2.1 Wisconsin Breast Cancer data set

This dataset was obtained from the University of Wisconsin Hospitals, Madison from Dr.William H. Wolberg, and formalized into ARFF file, Figure 8.

```
@relation Wisconsin Breast Cancer data
set @attribute SampleCodeNumber
numeric
@attribute ClumpThickness numeric
@attribute UniformityofCellSize numeric
@attribute UniformityofCellShape numeric
@attribute MarginalAdhesion numeric
@attribute SingleEpithelialCellSize
numeric
@attribute BareNuclei numeric
@attribute NormalNucleoli numeric
@attribute Mitoses numeric
@attribute Class {2, 4}
@data
1000025,5,1,1,1,2,1,3,1,1,2
1002945,5,4,4,5,7,10,3,2,1,2
1015425,3,1,1,1,2,2,3,1,1,2
```

Fig.8. Part of Wisconsin breast cancer dataset in ARFF format.

Wisconsin Breast Cancer data set consists of (10) features. All are numeric type: Clump Thickness, Uniformity of Cell Size, Uniformity of Shape, Cell Marginal Adhesion, Single Epithelial Cell Size, Bare Nuclei, Bland Normal Mitoses, Chromatin, Nucleoli. In addition, the class attribute has two values that the classification relies on: *Benign cancer* represented by (2), and *Malignant cancer* represented by (4). The dataset has 699 instances [Benign (458 instances), Malignant (241 instances)]. The data set has 16 instances with missing data (unavailable data) represented by "?".

VII. 7THE RESULT AND THE ANALYSIS

The aim of this study is to compare different learning algorithms to classify a benign cancer from malignant cancer in Wisconsin breast cancer dataset. One way to do the comparison is to estimate the error using any estimation procedure such as cross-validation for each learning algorithm and choose the algorithm whose estimate is smaller. This is still reasonable, if one algorithm has a lower estimated error than others in one dataset, then this algorithm can be consider as the model for this dataset. However, the difference between algorithms in this way might not be the significant difference. It is important to estimate the difference in errors significantly. The significant test called t-test (Student's t test) can be used to determine whether one learning algorithm outperforms another on a particular learning task.

7.1 The K-fold cross-validated paired t test

t-test is the most popular test among machine learning researchers and this the one in Weka tool. The dataset is divided into k disjoint sets of equal size used $T_1, T_2, T_3, \dots, T_n$ [Dietterich, 1996] where the size is at least 30. It then trains and tests the learning algorithm k times. In each time use T_i as a test set and the union of the other, where $T_j, j \neq i$, as training sets.

To begin by specify the significant level (confidence equals to 0.05). The experiment as shown in figure 9, clarifies that the Bayesian Network is significantly better among the other algorithms. It is notable that the difference between Bayesian Network and Naïve Bayes is not significantly and this is acceptable since Naïve Bayes is just a special case of Bayesian network, where the constraints on the assumptions in Bayesian Network are more global than Naïve Bayes (see section 2.1).

Confidence: 0.05 (two tailed)					
Dataset (1) bayes.Ba (2) bayes (3) funct (4) trees (5) trees.					
BreastCancer	(100)	97.20	 	96.11	95.58 * 95.49 * 94.92 *
	(v/*)	 	(0/1/0)	(0/0/1)	(0/0/1)
Key:					
(1) bayes.BayesNet '-D -Q bayes.net.search.local.K2 --P 1 -S BAYES					
-E bayes.net.estimate.SimpleEstimator --A 0.5' <u>7.4603744325877594E17</u>					
(2) bayes.NaiveBayes " <u>5.9952312017856973E18</u>					
(3) functions.MultilayerPerceptron '-L 0.3 -M 0.2 -N 500 -V 0 -S 0 -E					
20 -H a' <u>-1.18324858260785264E17</u>					
(4) trees.ADTree '-B 10 -E -3' <u>-1.53226483716769075E18</u>					
(5) trees.J48 '-C 0.25 -M 2' <u>-2.17733168393644448E17.</u>					

Fig.9. The result of the experiment.

The asterisks in Figure.9 means that the result is worse than the baseline (Bayesian Network in this case). For example, the function multilayer neural network is significantly worse than Bayesian Network. Figure.10 shows the Bayesian network that has been learned.

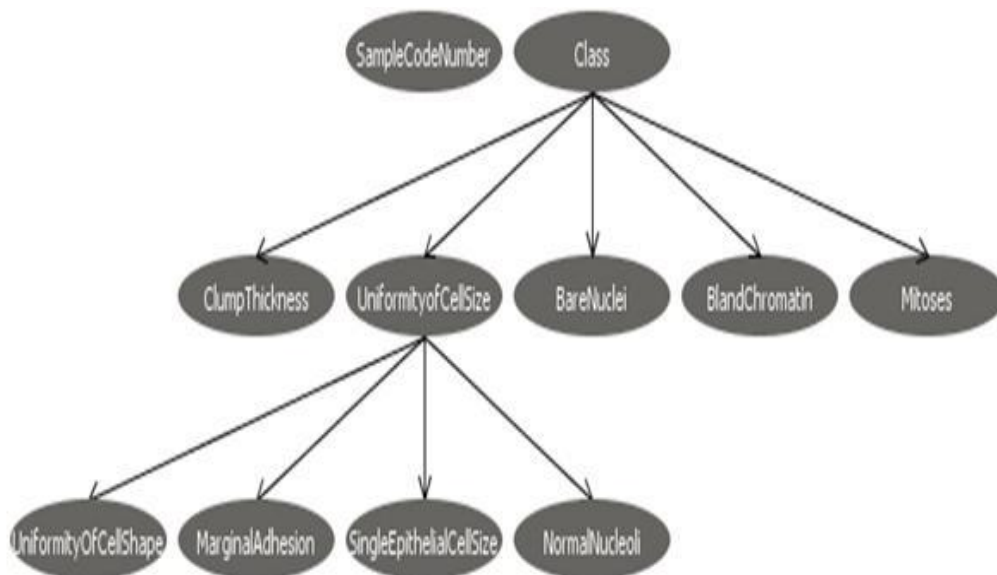


Fig.10 the Bayesian Network for Wisconsin breast cancer.

VIII. CONCLUSION AND FUTURE WORK

In this paper investigated five popular algorithms on one of the most important domains in medicine; namely breast cancer. This paper used Wisconsin breast cancer dataset that contains important risk factors. These factors usually are used to diagnose the breast cancer in labs and give a reliable result. This paper shown that using machine learning can help in cancer detection. However, it is also believe that the diagnosis of breast cancer can be more accurate if it is conducted in the genomic level. This will help in understanding for example the abnormality of genes and which are down-expressed/up-expressed. Starting from this motivation, also will work in the future in the genes-regularity networks using gene expression profiles for breast cancer. The consideration will be biased to Bayesian Networks and their interpretation to causal networks as it has not been investigated intensively in the literature.

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