

ARTIFICIAL INTELLIGENCE BASED IMPLEMENTATION APPROACH FOR PREDICTION OF BREAST CANCER MASSES

Abstract

Breast cancer is the common disease women face in this digital era. Based on the shape, size and density of the mammograms, benign and cancerous masses can be differentiated. Applications of machine learning in breast cancer are explored by focusing on predicting the possibility of a person having breast cancer. A few models are implemented in this chapter and a hybrid model named Voter Model is also implemented to have a better result. On an average the Voter model produces the results with an accuracy of 99.7%.

Keywords: Machine Learning, Breast Cancer Prediction, Voter Model

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I. INTRODUCTION

According to a breast cancer report [1], breast cancer stands as the most prevalent cancer affecting women in the United States. In recent years, there has been a noticeable rise in the incidence rate of breast cancer in developing nations. Numerous research studies have concentrated on differentiating between benign and malignant masses [13]. In this article, we introduce a hybrid prediction model called Voter Model, explicitly designed for forecasting breast cancer based on statistical data. Additionally, we present statistical projections for breast cancer occurrences in the United States for the year 2021. These projections indicate an expected 268,600 new cases of invasive breast cancer and 62,930 new cases of non-invasive breast cancer among women in the U.S. It's worth emphasizing that breast cancer mortality rates among women in the U.S. surpass those of most other types of cancer, except for lung cancer. A recent analysis also disclosed that on a global scale, a woman is diagnosed with breast cancer approximately every 3 minutes, and a woman loses her life to breast cancer about every 13 minutes.

In this chapter, machine learning concepts are used to predict Breast Cancer, using the dataset from The University of Wisconsin Breast Cancer Diagnosis Dataset (WBCD)[2] Section 2 describes the breast cancer dataset used and the features of a cancer cell taken into consideration. Section 3 talks about the algorithms use in breast cancer prediction. Section 4 gives an elaborate discussion on the factors taken into consideration while building various models. This Section 4 also gives the accuracies obtained at the end. Followed by this section are the references. The section 5 is the survey based on deep learning based algorithm

II. DATASET DESCRIPTION

The data collected so far can be classified into two groups: benign and malignant cases; 569 total cases, 357 classified as benign and 212 as malignant. The UC Irvine Machine Learning Repository is where the data utilized was located. The characteristics are calculated from a digital picture of a breast mass that was sampled using a fine needle aspiration (FNA). They characterize the characteristics of the visible cell nuclei in the picture. When creating the models, the following aspects were considered. Area, smoothness, smoothness (local variation in radius lengths), compactness, concavity (severity of concave regions of the contour), radius (mean of distances from center to points on the perimeter), texture (standard deviation of gray-scale values), perimeter, and area. Number of concaves, areas along the contour, symmetry, and fractal dimension.

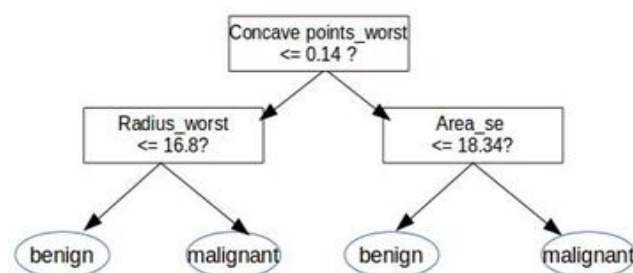


Figure 1: Decision Tree Classifier

III.MACHINE LEARNING BASED IMPLEMENTATION DETAILS

Python libraries such as Scikit-learn, Matplotlib, Keras were used for the models that were built.

- 1. Decision Tree Model:** A decision tree [3] is structured like a flowchart. Within this arrangement, every internal node functions as a "test" applied to an attribute, each branch illustrates the result of that test, and each leaf node represents a class label, which is the decision reached after considering all attributes. The paths from the tree's root to a leaf node define the criteria for classification. In Figure 1, you can see a portion of a decision tree classifier tailored for breast cancer. The test results provide answers to specific questions, and depending on these answers, a path within the tree is chosen. The final leaf node encountered indicates the class to which the sample belongs, classifying it as either benign or malignant.
- 2. Random Forest:** A random forest [4] consists of multiple decision trees, which collectively improve prediction reliability and accuracy. In the process of growing these trees, the Random Forest deliberately introduces an element of unpredictability into the model. When it comes to selecting features for splitting nodes, it chooses the best feature from a random subset of attributes rather than solely relying on the most significant one. This approach promotes greater diversity within the model. Decision trees are susceptible to overfitting [5], but Random Forest addresses this issue by generating random subsets of features and constructing smaller trees using them. In Figure 2, you can see three decision trees. Tree 1 and Tree 3 classify a test sample as malignant, whereas Tree 2 classifies it as benign. The Random Forest makes its prediction based on majority votes, ultimately predicting the sample as malignant.

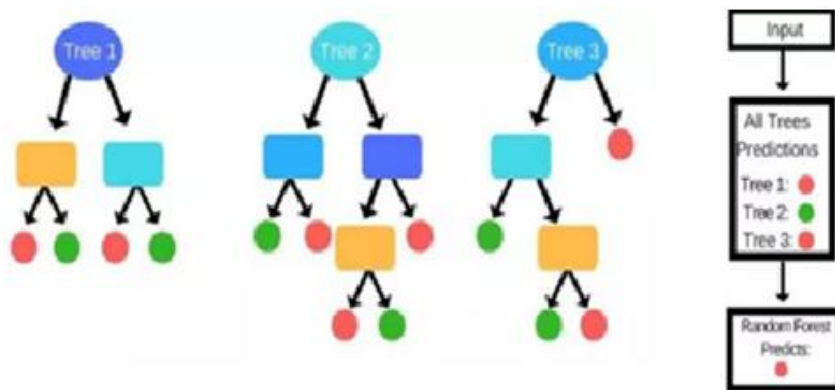


Figure 2: Random Forest Classifier

A model that fits the training data too well is referred to as over-fitting. This happens when a model learns every nuance and blip in the training data to the point where it performs horribly on untrained data. This is because the model learns ideas from the noise or random oscillations in the training data. However, fresh data might not support these ideas, and the model loses its ability to generalize.

- 3. Extra Tree:** Extra tree classifiers obtained by randomizing the random forest further. Instead of employing a bootstrap sample, each tree is trained using the whole learning sample, and the top-down splitting in the tree learner is random. A random cut-point is chosen rather than determining the locally optimum cut-point for each characteristic under consideration. The split that produces the greatest score is then selected to split the node out of all the randomly produced splits.
- 4. Support Vector Machine:** In the context of the support vector machine [6], every data point is represented as a point in an n-dimensional space, where 'n' corresponds to the number of features, and each feature's value corresponds to a specific coordinate in this space. The classification process involves the identification of a hyperplane that effectively separates the two classes. Specifically, the linear kernel seeks to establish a linear hyperplane for the purpose of classifying the samples.

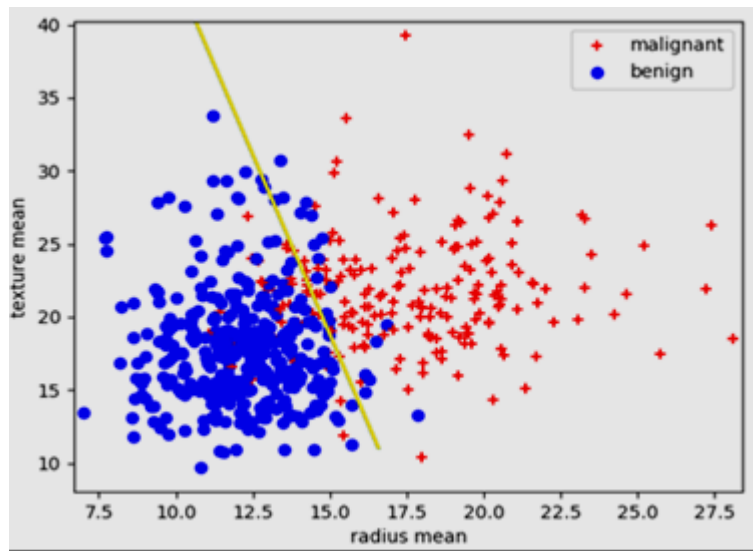


Figure 3: Support Vector Machine Classifier

Figure 3 shows an SVM classifier trained with features radius and texture. The red '+' represent malignant samples, while the blue circles represent benign samples. SVM classifier identifies the best hyperplane that classifies the data into their classes. This is represented by the yellow line. The model should consider accuracy as well as aim to maximize margin from samples to prevent overfitting.

- 5. Logistic Regression:** To make predictions, logistic regression utilizes an equation with linear components and independent predictors. The expected value can vary from negative infinity to positive infinity. The logistic regression outcome represents a binary class variable, where 0 corresponds to "no," and 1 corresponds to "yes." The result of the linear equation is compressed within the confined range of [0, 1]. The feature mean radius utilized in breast cancer prediction is represented by the X axis in Figure 4. The sigmoid function, which is employed to compress the anticipated value between 0 and 1, is shown by the blue curve. The samples above this curve are cancerous, whereas all those below it are benign.

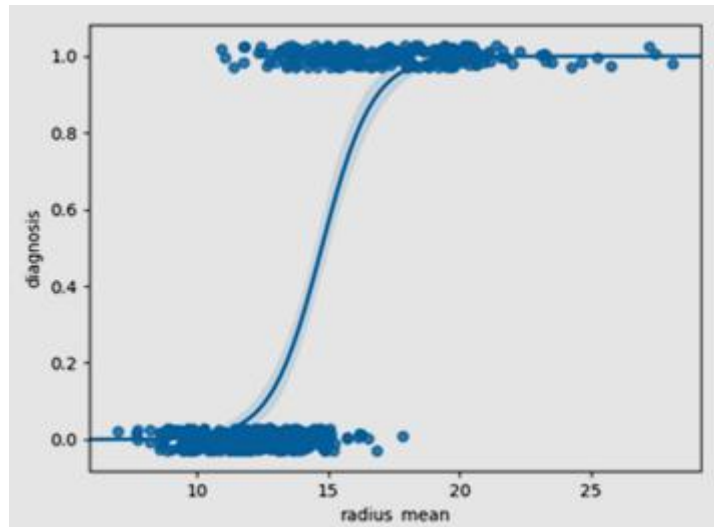


Figure 4: Logistic Regression Classifier

- 6. Naïve Bayes:** The Naive Bayes method, as described in reference [7], is a classification approach that relies on Bayes' Theorem [8]. It operates under the assumption that predictors are independent of each other. The Naive Bayes classifier assumes that the existence of a specific feature within a class has no connection to the existence of any other feature, which is why it is called "Naive.". $P(C_k | X)$ shows probability of a sample belonging to class malignant or benign. $P(C_k)$ represents this probability, while $P(X)$ represents the probability of a feature occurring, eg. mean symmetry being ≤ 0.15 .

$$\text{Equation 1: } P(C_k | X) = \frac{P(C_k)P(X|C_k)}{P(X)}$$

Equation 1 is the equation given by Bayes theorem.

- $P(C_k|X)$: Refers to the posterior probability of class C_k (the target) given the predictor X (attributes).
 - $P(C_k)$: Signifies the prior probability of class C_k , representing its initial probability before taking into account specific predictor variables.
 - $P(X | C_k)$: Denotes the likelihood, which represents the probability of observing predictor X given that class C_k is true.
 - $P(X)$: Represents the prior probability of predictor X , without considering any specific class information.
- 7. Artificial Neural Networks:** The neural network [9] captures information from the outcomes of previous data between cases. During training, the network is provided the results of previous cases as input along with the features. The neural network has an advantage over other methods in that it is also able to take features of all cases involved as inputs. Therefore, it can draw on the outcomes of previous training examples. The neural network used for the dataset under consideration is also shown here.

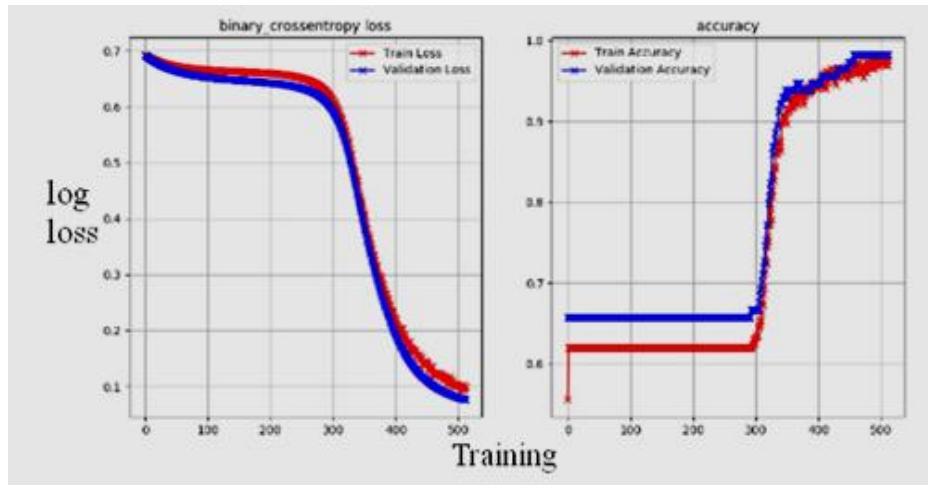


Figure 5: ANN classifier

Figure 5 describes how the loss (which should ideally be 0) decreases with each training iteration. The train and validation accuracy show a drastic increase as training progresses.

8. **Voter Model:** The No Free Lunch Theorem [10] states that any one algorithm that searches for an optimal cost or fitness solution is not universally superior to any other algorithm. In essence, different algorithms prove to be more effective for different data sets. Thus, instead of relying on a single algorithm completely, VoterModel algorithm relies equally on all of them.

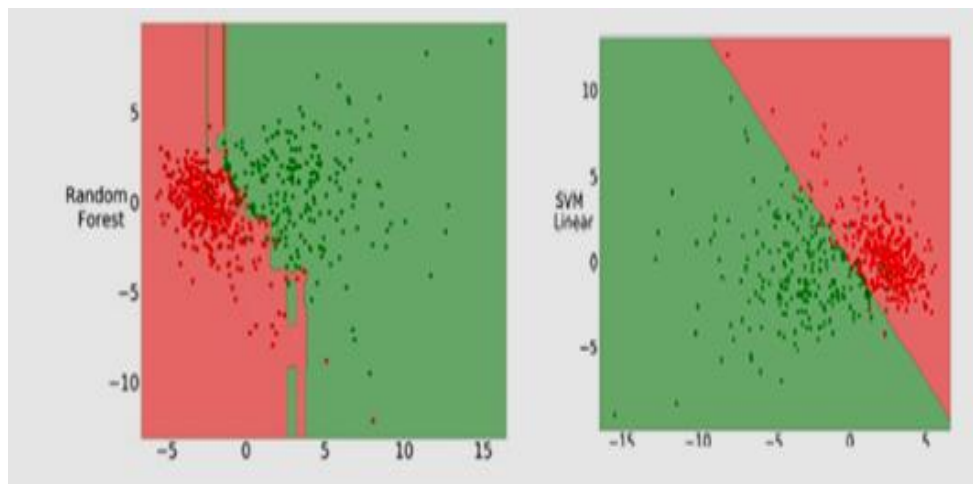


Figure 6: Fitting Curves of Random Forest and SVM Classifier

This can be explained by the Figure 6, which show the fitting curves of Random Forest and SVM classifiers. The features were reduced into 2 columns. The points (which indicates the samples) and curves (which indicates the boundaries) were plotted on the graph. Over fitting of boundaries to accommodate the points is clearly visible.

Voter Model Algorithm

Initialize votes for “benign” and “malignant” to 0
Train the data with the models under consideration.
Use the trained model to classify test data as “benign” or “malignant.”
If prediction is “malignant”:
 Increment votes of “malignant” by 1
Else:
 Increment votes of “benign” by 1
If “malignant” has higher vote count:
 Test data is considered as “malignant”
Else: Test data is considered as “benign”

Voter Model considers any machine learning model. Every model vote whether a test data is to be classified as benign or malignant. Based on most votes, a sample is classified as either benign or malignant using the model proposed. This can reduce over fitting as it prevents complete dependence on a single classifier. This has been proven based on the accuracy achieved by this model in comparison to the other models considered earlier.

IV. DISCUSSION

In the support vector machine model, proper parameter selection plays an important role in obtaining a correct classification. The linear kernel function is used to separate both the classes. Gamma should not be too high, as this can cause over-fitting.

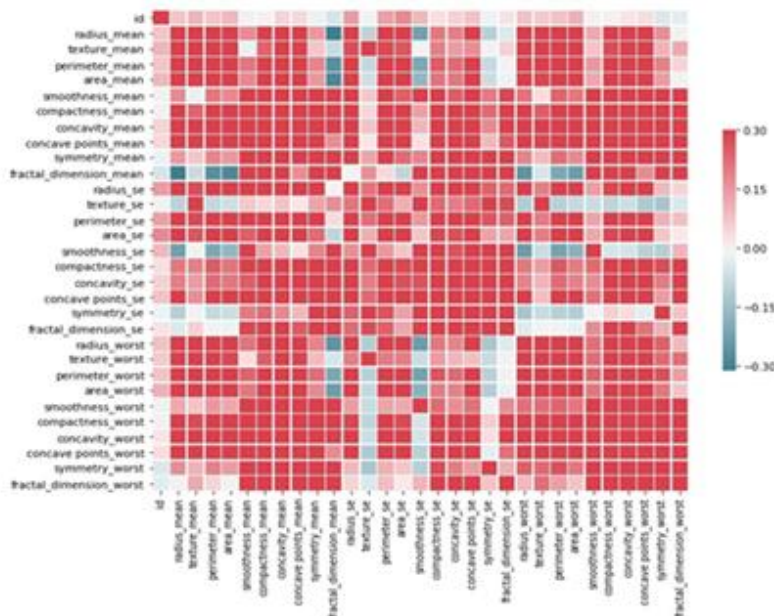


Figure 7: Correlation Matrix

The distribution of the remaining probability between the wrong classifications has no bearing on the value. A classification model whose output is a probability value between 0 and 1 is used to calculate cross-entropy loss, which employs a log function to gauge performance. As the anticipated probability becomes closer to the real label, cross-entropy loss goes down. For instance, it is poor and causes a significant loss to forecast a chance of 0.015 when the actual observation label is 1.

As seen in Fig. 7, the data was projected into a correlation matrix, a table that displays correlation coefficients between variables. The correlation between two variables is displayed in each cell of the table, which aids in choosing the features that may be utilized to train the model.

The correlation matrix helps to determine the correlated features, some of which are seen listed below.

Some positively correlated features identified are:

- Perimeter Mean and Radius Worst
- Area Mean and Radius Worst
- Texture Mean and Texture Worst
- Area Mean and Area Worst

From the correlation matrix, it was understood that radius, area, and perimeter essentially contain redundant information, which describes the physical appearance of a cell. Since area and perimeter are derived from radius, it is safe to discard both those columns. All the ‘worst’ columns can be discarded since they are a subset of the ‘mean’ columns.

For the random forest classifier and extra tree classifier, both the criteria- namely, Gini, as given by Equation 2, and entropy impurities, given by Equation 3, were implemented. Although both are often interchangeably used, for the Wisconsin Breast Cancer Diagnosis Dataset considered, entropy shows slightly better results. Gini prevents miscalculation, while entropy is used for exploratory analysis and can handle missing values. Entropy is apt for attributes that occur in classes.

Gini impurity:

$$\text{Equation 2: } \text{Gini}(E) = 1 - \sum_{j=1}^c p_j^2$$

where, P_j is the fraction of items labeled as class j .

Entropy:

$$\text{Equation 3: } H(E) = -\sum_{j=1}^c p_j \log p_j$$

where C is the number of classes

Table 1: Accuracies Obtained

Model	Columns	
	Criterion	Accuracy
Random Forest	Entropy	0.991
Random Forest	Gini	0.982
Extra Tree	Entropy	0.991
Extra Tree	Gini	0.982
Support Vector Machine	Linear Kernel	0.973
Logistic Regression	-	0.964
Naïve Bayes	-	0.956
Artificial Neural Network	-	0.999
Voter Model	-	0.997

Table 1 shows the accuracies obtained for various models, based on the different criteria considered.

V. SURVEY ON DEEP LEARNING METHODS

Utilizing the Xception deep learning model, [16] Yadavendra et al were able to attain exceptional results, with precision, recall, and F1 measures all reaching a commendable 0.90 under the same testing conditions. As a result, it was evident that the Xception method stands out as the superior choice among the various methods considered for classifying breast cancer tumors, demonstrating consistently high performance across these critical evaluation criteria. This signifies its robustness and effectiveness in accurately identifying and classifying such tumors, making it a preferred option for this task.

[17] Zheng, J et al introduced an innovative approach to breast cancer detection and early diagnosis by combining deep learning with the AdaBoost algorithm. They utilized the AdaBoost algorithm to create an ensemble classifier for the final prediction function. The results from evaluation tests demonstrated that proposed method exhibited superior predictive capabilities compared to other classifiers, with the deep-learning classifier standing out. Their analysis underscored the significant potential for rapid generalization and an efficiency boost in result prediction, driven by the neural network's automatic result derivation. Leveraging insights from the Convolutional Neural Network deep learning model, their DLA-EABA method contributed to enhancing system performance. They customized deep learning techniques to suit the unique attributes of each dataset, resulting in a tailored model for each one. The DLA-EABS method they put forth demonstrated remarkable accuracy in detecting breast cancer masses and subsequently improving patient survival rates. When benchmarked against existing methods, their approach consistently outperformed them in terms of performance.

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