

INTERNATIONAL JOURNAL IN PHARMACEUTICAL SCIENCES



Journal Homepage: https://www.ijpsjournal.com

Review Article

A Step Toward the Future: Using Machine Learning to Detect Leukemia

ABSTRACT

Nakul Magotra

Chandigarh University, Gharuan, Punjab

ARTICLE INFO Received: 17 July 2023

Accepted: 19 July 2023 Published: 27 July 2023 Keywords: automatic leukemia detection, acute lymphoblastic leukemia, lymphocyte image segmentation, machine learning DOI: 10.5281/zenodo.8187907

INTRODUCTION

The identification of malignant neoplastic illness may involve microscopic examination of peripheral blood smears. Nevertheless, this type of detailed microscopic evaluation takes a lot of time, is fundamentally subjective, and is controlled by the clinical knowledge and expertise of the hematopathologist. An affordable laptop power assisted technology for measuring peripheral blood samples must be created in order to get over these problems. In this study, methods for machine-controlled detection and subclassification of acute lymphocytic leukemia (ALL) using image processing and machine

learning techniques are proposed. The machinecontrolled unwellness recognition approach heavily relies on the choice of the appropriate segmentation theme. Hence, fresh techniques are envisaged to divide the traditional images of mature white cells and malignant lymph cells into their component morphological regions.

The segmentation problem is self-addressed within the supervised framework in order to make the planned schemes viable from a practical and realtime stand point. The segmentation problem is developed as picture element classification, picture element bunch, and picture element

*Corresponding Author: Nakul Magotra Address: Chandigarh University, Gharuan, Punjab

Email : nakulsharma1322@gmail.com

Relevant conflicts of interest/financial disclosures: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.



Leukemia is a cancer of the bone marrow, a spongy tissue that secretes into the bones and serves as the site for the production of blood cells. One of the most prevalent kinds of leukemia in adults is acute myeloid leukemia (AML). Leukemia has non-specific signs and symptoms that are also similar to those of other interpersonal illnesses. The only way to accurately diagnose leukemia is by manually examining a stained blood smear or bone marrow aspirate under the microscope. However, this approach takes more time and is less precise. This paper describes a method for the automatic recognition and classification of AML in blood smears. Classification techniques include decision trees, logistic regression, support vector machines, and naive bayes. labeling problems separately in these projected strategies, which include Gaussian Naive Bayes, Logistic Regression, Decision Tree Classifier, Random Forest Classifier, and SVM field modeling. The effectiveness of machinecontrolled classifier systems is evaluated against manual results supplied by a panel of hematopathologists using a thorough validation procedure. It is observed that conventional and malignant lymphocytes have different physical characteristics.

An Affordable technology is proposed to mechanically recognize lymphoblasts and observe dead peripheral blood samples. Metameric nucleus and protoplasm sections of the white cell images were used to derive morphological, textural, and color options. In comparison to other individual models, an ensemble of Random Forest classifiers demonstrates the highest classification accuracy of 96.5%. These techniques include inexpensive categorization, nucleus and protoplasm feature extraction, and lymph cell picture segmentation. An enhanced theme is also planned, and the results are related to use of the flow cytometer, to subtype malignant neoplastic disease blast images supporting cell lineages. By using this theme, it will be possible to establish whether blast cells are myeloid or derived from body fluid. The extracted alternatives from the leukemic blast images are mapped onto one of the two teams using an ensemble of call trees. To evaluate each model's performance, numerous studies and tests are carried out. Performance metrics, such as the precision and efficacy of the anticipated machinecontrolled systems based on routine diagnostic procedures.

A. LEUKEMIA

Leukemia is a collection of diverse blood-related malignancies with varying etiologies, pathophysiology, prognoses, and therapeutic responses (Bain, 2010). In today's world, leukemia is seen as a significant problem because it can afflict anyone, including children, adults, and occasionally newborns as young as 12 months. Leukemia is one of the top 15 most prevalent types of cancer in adults, according to a World Health Organization report, while it is the most common disease in children (Kampen,2012). The next sections will explain the blood cell lineage, different forms of leukemia, diagnostic techniques now in use, available treatments, and prognostic factors in an effort to better understand leukemia.

B. LEUKEMIA TYPES

Your doctor can identify the type of leukemia you have with the use of lab tests. The course of treatment varies for each kind of leukemia. Chronic and Acute Leukemias The name "leukemia" refers to how swiftly the disease progresses and worsens:

Acute: Typically, acute leukemia progresses quickly. Malignant neoplastic illness cells are growing rapidly, and these abnormal cells don't function like regular white blood cells do. A bone marrow examination could reveal low quantities of healthy blood cells and a high concentration of malignant neoplastic disease cells. Blood cancer patients may have extreme fatigue, bruising easily, and frequent infections.

Chronic: The progression of chronic leukemia can be gradual. Malignant neoplastic illness cells function almost identically to regular white blood cells. The first indication of illness may not be a physical illness; instead, it may be abnormal results from a routine biopsy. The biopsy, for instance, can reveal a high concentration of malignant neoplastic illness cells. Malignant neoplastic disease cells may eventually replace healthy blood cells if they are not treated.

C. Myeloid and Lymphoid Leukemia's

The afflicted kind of white blood cell is sometimes used to name leukemias:

Myeloid: Myeloid, myelogenous, or leukaemia are all terms used to describe leukemia that originates in myeloid cells.



Lymphoid: Lymphoid, lymphoblastic, or lymphocytic leukemia is a type of leukemia that develops in lymphoid cells. The lymph nodes, which swell, may become a collection point for lymphoid leukemia cells.

The Four Most Common Leukemia

Types Myeloid cells are affected by acute myeloid leukemia (AML), which spreads swiftly. Blast cells from leukemia gather in the bone marrow and blood. In 2013, an estimated 15,000 People will receive an AML diagnosis. The majority (about 8,000) will be 65 years of age or older, and about 870 children and teenagers can contract this sickness.

Rapid growth is a feature of acute lymphoblastic leukemia (ALL), which affects bodily fluid cells. Blast cells from leukemia typically gather in the blood and bone marrow. More than a dozen People have been given the diagnosis of beat 2013. The majority (almost 3,600) are children and teenagers. Myeloid cells are affected by chronic myeloid leukemia (CML), which often develops slowly at first. The variety of white blood cells has increased, according to blood testing. Moreover, the bone marrow contains a very small variety of leukemic blast cells. CML will be identified in about 6,000 Americans in 2013. Just approximately 107 children and teenagers can have this illness, and almost 0.5 (about 2,900) will be 65 years of age or older.

Lymphoid cells are affected by chronic lymphocytic leukemia (CLL), which normally develops slowly. The variety of white blood cells has increased, according to blood testing. The traditional blood corpuscle makes the aberrant cells function almost identically. In 2013, there were 16,000 CLL diagnoses in Americans. Around 10,700 are 65 years of age or older. Almost nobody is affected by this ailment. Around 6,000 additional instances of other, less prevalent types of blood cancer are expected in 2013.

PROPOSED EXPERIMENTAL METHOD

A. Logistic Regression

In essence, a supervised classification algorithmic rule is what logistic regression is. The target variable (or output), y, will only accept distinct values for the provided set of alternatives (or inputs), X, during a classification problem. Contrary to popular assumption, a regression model could include supply regression. To determine the likelihood that a given data input belongs to the class designated as "1," the program creates a regression model. Similar to how linear regression presumes that the data follows a linear function, logistic regression uses the sigmoid function to model the data.



Only when a call threshold is taken into consideration does logistic regression transform into a classification technique. Setting the threshold value, which depends on the classification problem itself, is a crucial aspect of logistic regression.

The exactness and recall values have a negative impact on the judgment for "the price | the worth" of the threshold value. Although it would be ideal for each exactness and memory to be identical, this is rarely the case. To determine the threshold in the event of a Precision-Recall trade-off, we consider the following factors: -

1. Low Precision/High Recall: We choose a call price that contains a low accuracy of exactitude or a high value of Recall in applications where we want to reduce the number of false negatives without actually reducing the amount of false positives. As an illustration, in a very cancer

diagnostic application, we frequently classify any affected patient as unaffected without paying much attention to whether the patient has already received a de jure cancer diagnosis. This may be because additional medical illnesses can identify the absence of cancer, but they cannot detect the presence of the disease in a candidate who has previously been rejected.

2. High Precision/Low Recall: In cases where we want to reduce the number of false positives without actually reducing the number of false negatives, we often choose a call that has a high precision value or a low recall value. For instance, if we are inclined to categorize customers based on whether or not they will react positively or negatively to a made-to-order advertisement, we would like to be absolutely certain that the consumer will react positively to the promotion because, in the event of a negative reaction, the consumer may lose out on a potential sale.

B. Gaussian Naive Bayes

A statistical classification method based on the Bayes Theorem is called naive Bayes. One of the only supervised learning algorithms is this one. The algorithmic rule that is quick, accurate, and dependable is the Naive Thomas Bayes classifier. For extremely large data sets, naive Thomas Bayes classifiers perform quickly and accurately. The naive Thomas Bayes classifier makes the assumption that the influence of a particular feature in a particularly class is dependent on a number of variables. A loan applicant, for instance, may or may not be interesting depending on their financial success, history of prior loans and group actions, age, and placement. These options are still considered separately even though they are interconnected. This assumption is regarded as naive because it makes calculation easier.

$$P(h|D) = \frac{P(D|h)P(h)}{P(D)}$$

1. P(h): the likelihood that hypotheses h being correct (regardless of the data). commonly known as the preceding probability of h.

2. P(D): is short for probability of the data (regardless of the hypothesis). It's commonly referred to as the prior likelihood.

3. P(h|D): is the likelihood that hypothesis h is true given the facts D. Often, this is referred to as posterior probability.

4. P(**D**|**h**): the likelihood of knowing d if the premise h were correct. Often, this is referred to as posterior probability.

C. Decision Tree Classifier

A decision tree is a tree structure that resembles a flowchart where each leaf node symbolizes the outcome and each interior node represents a feature (or attribute). In an excessive call tree, the root node at the top is recognized. It gains the ability to divide data according to attribute value. It divides the tree in a judgment call partitioning method that is extremely algorithmic. You benefit from greater cognitive process using this flowchart-like layout. It is essentially a graphic representation of a flow chart design that matches human level thinking. Decision trees are easy to understand because of this, and an interpretive response will result in the loss of a possible customer transaction.



A white box form of ML algorithm is the decision tree. Internal decision-making logic is shared, but this is not the case with black box algorithms like neural networks. When compared to the neural network algorithm, it trains more quickly. The



amount of records and number of attributes in the given data determine the temporal complexity of decision trees. The decision tree is a nonparametric or distribution-free strategy that does not rely on the assumptions of a probability distribution. High dimensional data can be handled by decision trees accurately.

D. Random Forest Classifier

A supervised learning algorithm is random forests. Both classification and regression can be done with it. This approach is also the simplest and most adaptable. Trees comprise a forest. Using randomly chosen data samples, random forests build decision trees, extract a forecast from each tree, and then use voting to determine which answer is best. It also offers a pretty accurate indication of how important convenience is. A supervised machine learning approach based on ensemble learning is known as random forest. A type of learning called ensemble learning involves combining the same algorithm or other algorithms to produce more accurate predictive models. The random forest algorithm creates a forest of trees by combining various algorithms of the same type, or many decision trees, hence the term "random forest". The random forest approach can be applied to both classification and regression tasks.



E. Support Vector Machines

Support vector classification (SVC) and support vector regression (SVR), often known as SVMs, are supervised learning algorithms that can be applied to classification and regression issues, respectively. It is only used for small datasets because processing them takes too long. We will concentrate on SVC in this set.

The fundamental assumption to make in this case is that the hyperplane is more likely to correctly classify points in its field or classes as there are more SV points from the hyperplane. Because the position of the hyperplane will change if the vector positions change, SV points are crucial in determining the hyperplane. This hyperplane is also referred to as a margin planning hyperplane technically. Finding a hyperplane that divides characteristics into distinct domains is the foundation of SVM.

• Linear Kernel

When data can be split linearly, or along the same line, the linear kernel is employed. It is one of the most often utilized kernels. It is typically utilized when a specific data set contains a lot of properties. Text classification is one of the instances where there are too many features because each alphabet introduces a new feature. So, in Text Classification, we primarily use linear kernel.

• Radial basis function kernel(RBF)/ Gaussian Kernel:

Another well-liked kernel technique that is more frequently employed in SVM models is the Gaussian RBF (Radial Basis Function). A function whose value changes depending on how far it is from the origin or another point is the RBF kernel. The format of the Gaussian Kernel is as follows:

$$K(X_1, X_2) = exponent(-\gamma ||X_1 - X_2||^2)$$

||X1 - X2|| = Euclidean distance between X1 & X2

We determine the dot product (similarity) of X1 & X2 using the distance in the initial space.

> Confusion Matrix:

A confusion matrix, also known as an error matrix, is a standard table structure that permits visualization of the performance of an



algorithm, particularly one that deals with statistical classification problems.

In learning unexpectedly, a supervised learning one is typically referred to as a matching matrix. In the matrix, each row denotes a hypothetical class, whereas each column denotes occurrences in the actual class (or vice versa). The name refers to how simple it is to determine whether the system is conflating the two classes (ie usually confusing each other).

It is a particular kind of contingency table that has the same set of "classes" in both of its dimensions—"real" and "approximate"—and two dimensions (a combination of each class and dimension in the dimension table is a variable).

		Prediction		
		0	1	
tual	0	TN	FP	
Act	1	FN	ТР	

- False positive (FP) = A test result that incorrectly suggests the presence of a specific ailment or trait
- True positive (TP) = The proportion of true positivity that is successfully detected is measured by true positive (TP), which is also known as true positive rate or probability of detection in various fields.
- True Negative (TN) = The proportion of true negatives that are accurately detected is measured by True Negative (TN), also known as Specificity (also known as real negative rate).
- False Negative (FN) = False Negative (FN) results are tests that show a condition does not exist when in fact it does. An example

would be a test result that shows a person does not have cancer when in fact they do.

- Steps to the work Process:
 - 1. To make writing programs easier, we import packages and libraries in the first step.
 - 2. After loading the data, I will output the first seven rows of information.
 - 3. Examine the information and determine how many rows and columns are present.
 - 4. Produce graphs and encode hierarchical data to count counts.
 - 5. Produce a pair plot to show the association.
 - Divide the data set into a feature data set, also known as the independent data set (X), and a target data set, also known as the dependent data set, to begin setting the data for the model (Y).
 - 7. Rescale and redistribute the data.
 - 8. Provide a function that can store a variety of classification models, including logistic regression, decision trees, and random forests.
 - 9. To determine if each patient has cancer, build a model that incorporates all models and examine the accuracy scores on each model's training data.
 - 10. Display the confusion matrices and model precision on test data.
 - 11. Use the model that performed best on the test data by accuracy and metric.

RESULTS OF EXPERIMENTS AND DISCUSSION

The 569 blood samples that made up the dataset for this study were. In order to obtain the sample dataset of blood cancer for AML, which comprises of 569 and 33 different parameters on which the analysis is performed, we entered the Kaggle platform.

• Load the data

	id	diagnosis	radius_mean	texture_me:
0	842302	М	17.99	10.38
1	842517	Μ	20.57	17.77
2	84300903	М	19.69	21.25
3	84348301	Μ	11.42	20.38
4	84358402	Μ	20.29	14.34
5	843786	М	12.45	15.70
6	844359	М	18.25	19.98

A sampling of the data's first seven rows





Malignant (cancerous) and benign (noncancerous) diagnosis chart

• Construct a pair plot



• Create a heat map to illustrate the relationship.



• Display the accuracy and confusion matrices for each model on the test data

```
[[86 4]
[ 4 49]]
Model[0] Testing Accuracy = "0.9440559440559441!"
[[89 1]
[ 5 48]]
Model[1] Testing Accuracy = "0.958041958041958!"
[[87 3]
[ 2 51]]
Model[2] Testing Accuracy = "0.965034965034965!"
[[88 2]
[ 3 50]]
Model[3] Testing Accuracy = "0.965034965034965!"
[[85 5]
[ 6 47]]
Model[4] Testing Accuracy = "0.9230769230769231!"
[[84 6]
[ 1 52]]
Model[5] Testing Accuracy = "0.951048951048951!"
[[87 3]
[ 2 51]]
Model[6] Testing Accuracy = "0.965034965034965!"
```



• Demonstrate more methods for obtaining classification accuracy and other metrics

	precision	recall	f1-score	support		
0	0.95	0 99	0.97	9.9		
1	0.95	0.91	0.94	53		
-	0120	0.171	0121			
accuracy			0.96	143		
macro avg	0.96	0.95	0.95	143		
weighted avg	0.96	0.96	0.96	143		
0.958841958841958						
Model 2						
	precision	recall	f1-score	support		
0	0.98	0.97	0.97	90		
1	0.94	0.96	0.95	53		
accuracy			0.97	143		
macro avg	0.96	0.96	0.96	143		
weighted avg	0.97	0.97	0.97	143		
0.96503496503	4965					
Hadal 3						
MODEL 3			<i>(</i>			
	precision	Lecall	+1-score	support		
	0.07		0.07			
0	0.9/	0.96	0.97	50		
1	0.90	0.94	0.95	53		
24410241			0.07	145		
accuracy			0.97	143		
macro avg	0.96	0.96	0.96	143		
weighted avg	0.96	0.97	6.96	143		
0.96503496503	4965					
Nodel 4						
novea -						
	nnecicion	recall.	£1_55000	support		
	precision	recall	f1-score	support		
a	precision 0.93	recall	f1-score	support		
e 1	0.93	0.94	6.94	support 90		
0 1	0.93 0.90	recall 0.94 0.89	6.94 0.90	support 90 53		
0 1	0.93 0.90	recall 0.94 0.89	f1-score 0.94 0.90	90 53		
e 1 accuracy	precision 0.93 0.90	recall 0.94 0.89	f1-score 0.94 0.90 0.92 0.92	support 90 53 143 143		
accuracy macro avg	0.93 0.99 0.92 0.92	0.94 0.89 0.92	f1-score 0.94 0.90 0.92 0.92 0.92	support 90 53 143 143		
e 1 accuracy macro avg weighted avg	precision 0.93 0.90 0.92 0.92	recall 0.94 0.89 0.92 0.92	f1-score 0.94 0.90 0.92 0.92 0.92	90 53 143 143 143		
0 accuracy macro avg weighted avg 0.92307692307	0.93 0.99 0.92 0.92	0.94 0.89 0.92 0.92	f1-score 0.94 0.90 0.92 0.92 0.92	90 53 143 143 143		
0 accuracy macro avg weighted avg 0.92307692307	0.93 0.99 0.90 0.92 0.92	0.94 0.89 0.92 0.92	f1-score 0.94 0.90 0.92 0.92 0.92	support 90 53 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5	0.93 0.90 0.92 0.92 0.92	0.94 0.89 0.92 0.92	f1-score 0.94 0.90 0.92 0.92 0.92	90 53 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5	0.92 0.92 0.92 0.92 0.92 0.92	0.94 0.99 0.92 0.92 recall	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score	support 90 53 143 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5	precision 0.93 0.90 0.92 0.92 0.92 0.92 precision	0.94 0.89 0.92 0.92 recall	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score	support 90 53 143 143 143 143 support		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0	precision 0.93 0.90 0.92 0.92 0.92 precision 0.99	recall 0.94 0.92 0.92 recall 0.93	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96	support 90 53 143 143 143 143 support 90		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1	precision 0.93 0.90 0.92 0.92 0.92 0.92 0.92 precision 0.99 0.90	recall 0.94 0.92 0.92 recall 0.93 0.98	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94	support 90 53 143 143 143 143 support 90 53		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1	precision 0.93 0.90 0.92 0.92 0.92 0.92 precision 0.99 0.99	recall 0.94 0.92 0.92 recall 0.93 0.98	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94	support 90 53 143 143 143 143 support 90 53		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy	precision 0.93 0.90 0.92 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99	recall 0.94 0.92 0.92 recall 0.93 0.98	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95	support 90 53 143 143 143 143 support 90 53 143		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg	precision 0.93 0.90 0.92 0.92 0.92 0.92 precision 0.99 0.99 0.90 0.94	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95	support 90 53 143 143 143 143 support 90 53 143 143		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg	precision 0.93 0.90 0.92 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99 0.99	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95	support 90 53 143 143 143 143 53 53 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg	precision 0.93 0.90 0.92 0.92 0.92 0.92 precision 0.99 0.90 0.94 0.94	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95	f1-score 0.94 0.92 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95	support 90 53 143 143 143 143 143 53 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104	precision 0.93 0.90 0.92 0.99 0.95 0.55 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95	support 90 53 143 143 143 143 support 90 53 143 143 143 143		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6	precision 0.93 0.90 0.92 0.99 0.95 0.55 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95	support 90 53 143 143 143 143 53 53 143 143 143		
e 1 accuracy weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6	precision 0.93 0.90 0.92 0.99 0.95 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.95 recall	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95 0.95 1.95 0.95	support 90 53 143 143 143 143 53 143 143 143 143 143 5000000000000000000000000000000000000		
e accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6	precision 0.93 0.90 0.92 0.99 0.95 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95 recall	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.95 0.95 0.95 0.95 0.95 1-score	support 90 53 143 143 143 143 53 143 143 143 143 143		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 1 accuracy macro avg weighted avg 0.95104895104 Model 6	precision 0.93 0.90 0.92 0.99 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 00 0.98 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95 recall 0.97	f1-score 0.94 0.92 0.92 0.92 f1-score 0.96 0.95 0.95 0.95 1-score 0.95 0.95 0.95 0.95	support 90 53 143 143 143 143 53 143 143 143 143 143 143 90		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6 0 1	precision 0.93 0.90 0.92 0.95 0.99 0	recall 0.94 0.92 0.92 recall 0.93 0.96 0.95 recall 0.95 recall 0.97 0.96	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.95 0.95 0.95 f1-score 0.97 0.95	support 90 53 143 143 143 143 143 143 143 143 143 14		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6 0 1	precision 0.93 0.90 0.92 0.99 0.98	recall 0.94 0.92 0.92 recall 0.93 0.96 0.95 recall 0.95	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95 f1-score 0.95 0.95 0.95	support 90 53 143 143 143 143 143 143 143 143 143 14		
e 1 accuracy weighted avg 0.92307692307 Model 5 0 1 accuracy weighted avg 0.95104895104 Model 6 1 accuracy 1 accuracy	precision 0.93 0.90 0.92 0.99 0.95 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95 recall 0.97 0.96	f1-score 0.94 0.90 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95 f1-score 0.95	support 90 53 143 143 143 143 143 143 143 143 143 14		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 accuracy macro avg weighted avg 0.95104895104 Model 6 1 accuracy macro avg	precision 0.93 0.90 0.92 0.99 0.95 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95 recall 0.97 0.96 0.96 0.96 0.96 0.96 0.96	f1-score 0.94 0.92 0.92 0.92 f1-score 0.96 0.95 0.95 0.95 f1-score 0.97 0.95 0.95 0.97 0.96	support 90 53 143 143 143 143 53 143 143 143 143 143 143 143		
e 1 accuracy macro avg weighted avg 0.92307692307 Model 5 0 1 accuracy macro avg weighted avg 0.95104895104 Model 6 1 accuracy macro avg weighted avg	precision 0.93 0.92 0.99 0	recall 0.94 0.92 0.92 recall 0.93 0.98 0.96 0.95 recall 0.97 0.96 0.97 0.96 0.97	f1-score 0.94 0.92 0.92 0.92 f1-score 0.96 0.94 0.95 0.95 0.95 f1-score 0.97 0.95 0.97 0.96 0.97 0.96 0.97 0.96 0.97 0.96 0.97 0.95 0.97 0.95 0.97 0.95 0.97 0.95	support 90 53 143 143 143 143 143 143 143 143 143 14		

0.965034965034965

• Publish the Random Forest Classifier model's prediction along with the actual values

Table 1: Accuracy Table

Method	Accuracy Percentage
Logistic Regression	94.4%
K Nearest Neighbour	95.8%
Support Vector Machine	96.5%
(Linear Classifier)	
Support Vector Machine	96.5%
(RBF Classifier)	
Gaussian Naive Bayes	92.3%
Decision Tree Classifier	95.1%
Random Forest	96.5%
Classifier	

CONCLUSION

Create the prediction or classification based on the test data and display both the classification or prediction made by the Random Forest Classifier model and the patient's actual values, which indicate whether or not they have cancer. I've noticed that the model incorrectly diagnosed some patients as having cancer when they didn't, and it incorrectly classified individuals who actually did have cancer as not having cancer. Even though this model is good, I want it to be even better when it comes to other people's lives. I want it to be as accurate as possible and as good as, if not better than, doctors. So, each of the models needs to be fine-tuned a little bit further.

REFERENCES

 Cristianini, N., and J. Shawe-Taylor. "An Introduction to support vector machines and other kernel-based learning methods" New York: Cambridge University Press, 2000.

- 2. Vapnik, V. N. "The Ature of Statistical Learning Theory" New York: Springer, 1995.
- A. Madabhushi, "Digital pathology image analysis: opportunities and challenges," Imaging in Medicine, vol. 1, no. 1, pp. 7– 10, 2009.
- A. N. Esgiar, R. N. G. Naguib, B. S. Sharif, M. K. Bennett, and A. Murray, "Fractal analysis in the detection of colonic cancer images," IEEE Transactions on Information Technology in Biomedicine, vol. 6, no. 1, pp. 54–58, 2002.
- L. Yang, O. Tuzel, P. Meer, and D. J. Foran, "Automatic image analysis of histopathology specimens using concave vertex graph," in Medical Image Computing and Computer-Assisted Intervention—MICCAI 2008, pp. 833–841, Springer, Berlin, Germany, 2008.
- 6. R. C. Gonzalez, Digital Image Processing, Pearson Education India, 2009.

- S. Liao, M. W. K. Law, and A. C. S. Chung, "Dominant local binary patterns for texture classification," IEEE Transactions on Image Processing, vol. 18, no. 5, pp. 1107–1118, 2009.
- J. C. Caicedo, A. Cruz, and F. A. Gonzalez, "Histopathology image classification using a bag of features and kernel functions," in Artificial Intelligence in Medicine, vol. 5651 of Lecture Notes in Computer Science, pp. 126–135, Springer, Berlin, Germany, 2009.
- H. S. Wu, J. Barba, and J. Gil, "Iterative thresholding for segmentation of cells from noisy images" Journal of Microscopy, vol.197, no. 3, pp. 296–304, 2000.

HOW TO CITE: Nakul Magotra, A Step Toward the Future: Using Machine Learning to Detect Leukemia, Int. J. in Pharm. Sci., 2023, Vol 1, Issue 7, 317-325. https://doi.org/10.5281/zenodo.8187907

