COMPARATIVE ANALYSIS OF ALGORITHMS FOR LUNG CANCER PREDICTION

A MINI PROJECT REPORT

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BONAFIDE CERTIFICATE

Certified that Mini project report titled "COMPARATIVE ANALYSIS OF ALGORITHMS FOR LUNG CANCER PREDICTION" is the bona fide work of R HARSHA VARDHAN(RA2111003011023), DEBANJAN(RA2111003011003) VUN NAM SHASHANK(RA2111003011029) who carried out minor project under my supervision further, that to the best of my knowledge, the work reported herein does not form any other project report or dissertation on the basis of which a degree or award was conferred on an earlier occasion on this or any other candidate.

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ABSTRACT

Lung cancer is a major global health concern, and early detection plays a crucial role in improving survival rates. Machine learning techniques have emerged as powerful tools for lung cancer prediction, leveraging diverse datasets to aid in diagnosis and prognosis. This study explores the application of machine learning algorithms for lung cancer prediction by integrating various types of data sources, including genetic, clinical, and imaging data. We present a comprehensive review of state-of-the-art machine learning methodologies, including support vector machines, artificial neural networks, and ensemble methods, employed in lung cancer prediction tasks.

In this study, we conducted a comprehensive comparative analysis of various machine learning algorithms for lung cancer prediction. Our research encompasses a diverse set of algorithms, including Support Vector Machines (SVM), Random Forest, Neural Networks, and k-Nearest Neighbors (k-NN). We utilized a rich dataset comprising clinical, genetic, and imaging features to evaluate the performance of these algorithms.

The comparative analysis involved preprocessing the data, including standardization and feature selection techniques, to ensure optimal input for each algorithm. We systematically evaluated the algorithms using a variety of performance metrics such as accuracy, sensitivity, specificity, The experiments were conducted through rigorous cross-validation procedures to ensure robustness and reliability of the results.

This comparative analysis provides valuable insights into the strengths and limitations of different machine learning algorithms concerning lung cancer prediction. The study's outcomes contribute to the ongoing efforts to enhance the accuracy and effectiveness of predictive models in lung cancer research and clinical applications. These findings guide researchers and healthcare professionals in selecting appropriate algorithms based on the specific dataset characteristics, thereby advancing the field of early lung cancer prediction, and improving patient outcomes.

The project also investigates statistical analysis methods, such as t-tests, F-tests (ANOVA), and chi-squared tests, to examine the relationships between variables and their significance in the context of diabetes prediction.

The findings and models generated in this project hold the potential to improve diabetes risk assessment, early detection, and personalized management strategies. Such advancements in predictive modeling can facilitate better healthcare decision-making, enhance preventive measures, and ultimately contribute to more effective diabetes management, reducing the burden of this chronic condition on individuals and healthcare systems.

1.1 INTRODUCTION

Lung cancer is the principal cause for cancer-related death. Lung cancer can initiate in the windpipe, main airway or lungs. It is caused by unchecked growth and spread of some cells from the lungs. People with lung disease such as emphysema and previous chest problems have more chances to be diagnosed with lung cancer. Over usage of tobacco, cigarettes and beedis, are the major risk factor that leads to lung cancer in Indian men; however, among Indian women, smoking is not so common, which indicate that there are other factors which lead to lung cancer. Other risk factors include exposure to radon gas, air-pollutions and chemicals in the workplace. A cancer that starts in lung is primary lung cancer whereas those which start in the lung and spread to other parts of body is secondary lung cancer. The size of tumor and how far it has spread determines the stage of cancer. An early-stage cancer is a small cancer that is diagnosed in lung and advanced cancer is the one that has spread into surrounding tissue or other parts of the body. A better understanding of risk factors can help to prevent lung cancer disease. The key to improve the survival rate is early detection using Machine learning techniques and if we can make the diagnosis process more efficient and effective for radiologists by using this, then it will be a key step towards the goal of improved early detection.

The Lung Cancer datasets used for this study are taken from UCI Machine Learning Repository and Data World. First, the given datasets are divided into training and test data by using k-fold cross validation technique. Then using the classification algorithms such as SVM, Logistic Regression, Naïve Bayes and Decision Tree, respective classification models are implemented using the given training data. The classification models are created using training data and the corresponding models are evaluated using test data to get the accuracy of the models. Finally, we compared the accuracy rates of each and every classification models that we implemented and arrived at a conclusion.

To build the prediction model, key features were first selected using a data-driven feature selection technique composed of an analysis of variance (ANOVA) test, a chi-squared test, and recursive feature elimination methods. We compared the performance of the prediction models—logistic regression (LR), support vector machine (SVM), random forest (RF), decision tree (DT), K-nearest neighbor.

1. LITERATURE SURVEY

S. N O	Title	Author Name	Journal Name	Methodolog ies Used	Inferences
1.	Analysis of lung cancer Prediction	Radhika P.R, Rakhi A.S.Nair, Veena G.	IEEE	Data Mining	This paper demonstrates the potential of predictive analytics in healthcare, particularly for Lung cancer. It identifies SVM and KNN as promising algorithms for this specific task but acknowledges the need for larger, cleaner datasets and further model optimization to achieve higher accuracy in real-world applications
2.	Lung Cancer Prediction using different Machine Learning Approaches	Priyanka Sonar, K. JayaMalini	IEEE	Dataset Collection, Training Data and Test Data, Pre- processing, Feature Extraction, Target Database	SVMs are versatile but require careful parameter tuning. Naive Bayes is robust but sensitive to data preprocessing and biased with large datasets. ANNs offer high prediction accuracy but can be computationally expensive and challenging for big data.
3.	Cancer Diagnosis and Treatment Research Based on Machine Learning	Bo He, Kuangi- shu, Heng Zhang	IEEE	Data Process, Algorithm Improveme nt	This research addresses a critical issue in healthcare by using a combination of traditional and deep learning techniques to predict readmissions among diabetic patients in resource-constrained medical environments.

4.	Diabetes Prediction using Machine Learning Algorithms	Aishwarya Mujumdar, Dr. Vaidehi	IEEE	Dataset Collection, Data Pre- processing, Clustering, Build Model, Evaluation	The study demonstrates the success of Logistic Regression and AdaBoost in diabetes prediction, an improvement in accuracy compared to existing datasets, and hints at the potential for further research in predicting the likelihood of individuals developing diabetes in the future.
5	A comparative study of Lung Cancer detection using Machine Learning algorithms	Arwatki Chen Lyngdoh, Nurul Amin Choudhury, Soumen Moulik	IJSRE M	Data Collection, Data Pre- processing, Setting Classificatio n Metrics, Applying Machine Learning Algorithms, K-Fold Cross Validation	The study addresses the critical challenge of early disease detection, particularly focusing on cancer. It successfully develops an accurate prediction model and hints at its potential application ,This involves conducting experiments using the Pima Indians Diabetes Database.
6	A Study on Prediction Of Lung Cancer Using Machine Learning Algorithms	Abhishek Gupta, Israr Ahmad, Zeeshan Ansari.	Resear ch Square	Lung cancer, AI, KNN, SVM, Random forest	The study systematically evaluates different aspects of model performance. The proposed model outperforms existing methods in diabetes prediction, particularly after feature selection.

2. Statistical Analysis

Data Collection

Collect a dataset with relevant features (predictors) and a target variable (Cancer status). Common features might include Gender, Age, Smoking, Coughing, Shortness of Breath, Swallowing Difficulty, Chest Pain.

Descriptive Statistics

Calculate basic descriptive statistics for your dataset. Mean, median, and mode for numerical features (e.g. Age, Smoking).

Hypothesis Testing

t-tests to compare means of numerical features between cancer patients and noncancer patients. Chi-squared tests for independence between categorical features and cancer status. ANOVA if you have multiple groups to compare (e.g., different treatment groups).

Data Splitting

Split the dataset into two parts: training and testing sets (e.g., 80% for training, 20% for testing). This helps evaluate your model's performance on unseen data.

Model Selection

Choose appropriate machine learning algorithms for classification. Common choices include Logistic Regression, Decision Trees, Random Forest, Support Vector Machines (SVM), and Neural Networks.

Model Training

Train your selected models using the training dataset. Evaluate different models using performance metrics (e.g., accuracy, precision, recall, F1-score) on the validation set.

Model Evaluation

Assess the model's performance using various metrics. Since diabetes prediction is a binary classification problem, you can use metrics like accuracy, precision, recall, F1-score, ROC- AUC, and the confusion matrix. Visualize performance metrics using ROC curves or precision-recall curves.

Model Interpretation

If applicable, interpret the model's results to understand which features are most important in predicting diabetes. Techniques like feature importance plots for tree-based models can be helpful.

2.1 MEAN, MEDIAN, MODE

Mean: The mean is the average of a set of values, calculated by summing all values and dividing by the total count.

CODE & OUTPUT:

```
In [25]: column_of_interest = 'AGE'
    mean = df[column_of_interest].mean()
    print(f"Mean of {column_of_interest}: {mean}")
    column_of_interest = 'SMOKING'
    mean = df[column_of_interest].mean()
    print(f"Mean of {column_of_interest}: {mean}")
    column_of_interest = 'ANXIETY'
    mean = df[column_of_interest].mean()
    print(f"Mean of {column_of_interest}: {mean}")

Mean of AGE: 62.90942028985507
    Mean of SMOKING: 1.5434782608695652
    Mean of ANXIETY: 1.4963768115942029
```

Median: The median is the middle value in a sorted dataset, or the average of the two middle values in a dataset with an even number of observations, representing the central position. If the dataset has an odd number of values, the median is the middle value. If the dataset has an even number of values, the median is the average of the two middle values.

```
In [26]: column_of_interest = 'AGE'
median = df[column_of_interest].median()
print(f"Median of {column_of_interest}: {median}")
column_of_interest = 'SMOKING'
median = df[column_of_interest].median()
print(f"Median of {column_of_interest}: {median}")
column_of_interest = 'ANXIETY'
median = df[column_of_interest].median()
print(f"Median of {column_of_interest}: {median}")

Median of AGE: 62.5
Median of SMOKING: 2.0
Median of ANXIETY: 1.0
```

Mode: The mode is the value that appears most frequently in a dataset, representing the most common observation(s). Unlike the mean and median, which are measures of central tendency, the mode is a measure of the most frequent value(s). A dataset can have one mode (unimodal) or multiple modes (multimodal).

CODE & OUTPUT:

```
In [24]: column_of_interest = 'AGE'
    mode = df[column_of_interest].mode()
    print(f"Mode of {column_of_interest}: {mode}")
    column_of_interest = 'SMOKING'
    mode = df[column_of_interest].mode()
    print(f"Mode of {column_of_interest}: {mode}")
    column_of_interest = 'ANXIETY'
    mode = df[column_of_interest].mode()
    print(f"Mode of {column_of_interest}: {mode}")

Mode of AGE: 0 64
    Name: AGE, dtype: int64
    Mode of SMOKING: 0 2
    Name: SMOKING, dtype: int64
    Mode of ANXIETY: 0 1
    Name: ANXIETY, dtype: int64
```

2.2 F-TEST(ANNOVA)

To utilize the F-Test (Analysis of Variance or ANOVA) for Lung cancer prediction with the provided attributes, the following steps can be taken. First, assemble a dataset including attributes like age, smoking, chest pain, difficulty in breathing and the binary target variable, Lung Cancer. Data preprocessing is essential to address missing values, outliers, and ensure data quality. The F-Test is then applied to evaluate the statistical significance of each attribute in relation to diabetes, testing whether the means of these attributes significantly differ between cancer and non-cancer individuals.

Subsequently, attributes with a significant F-statistic can be retained for building a predictive model, indicating a strong relationship with cancer. Using these selected features, a supervised learning model is constructed, such as logistic regression, decision trees, or random forests, to predict diabetes.

Factors affecting model accuracy after employing the F-Test for feature selection encompass the quality of the dataset, attribute selection, model choice, data preprocessing, class imbalance management, and hyperparameter tuning. Ensuring that the selected features not only show statistical significance but are also clinically relevant for lung cancer prediction is critical. Additionally, the choice of the predictive model and the selection of appropriate evaluation metrics significantly impact the model's accuracy, making it imperative to consider these factors in a holistic approach to diabetes prediction.

Code & Output:

```
In [29]: import statsmodels.api as sm
    from statsmodels.formula.api import ols

# Assuming 'target' is your dependent variable
model = sm.OLS.from_formula('LUNG_CANCER ~ AGE + SMOKING + YELLOW_FINGERS', data=df).fit()

# Perform the ANOVA
anova_table = sm.stats.anova_lm(model, typ=2)

# Extract the F-statistic and associated p-value
    f_statistic = anova_table['F'][0]
    p_value = anova_table['PR(>F)'][0]

print("F-Statistic:", f_statistic)
print("P-Value:", p_value)

F-Statistic: 3.125098704749822
P-Value: 0.0782162922519221
```

2.3 T-TEST

To leverage the T-Test for lung cancer prediction with the given attributes, we can follow a systematic approach. First, construct a dataset containing attributes like age, smoking history, yellow fingers, family history of cancer, lung function metrics, and other relevant factors. Include a binary target variable for lung cancer (encoded as 0 for non-cancerous and 1 for cancerous cases). Ensuring data quality is crucial; preprocessing the data to handle missing values, outliers, and any other data anomalies.

Subsequently, the T-Test can be applied to assess whether there are statistically significant differences in the means of each attribute between individuals with and without lung cancer. Attributes exhibiting significant T-Test results can be retained as predictors for lung cancer, indicating their potential impact on the disease. This analytical approach helps identify key factors influencing lung cancer, aiding researchers and healthcare professionals in making more accurate predictions and developing effective prevention strategies.

CODE&OUTPUT:

```
In [30]: import scipy.stats as stats
         group_a_column = df['AGE']
         group_b_column = df['YELLOW_FINGERS']
         # Perform a two-sample t-test assuming unequal variances (Welch's t-test)
         t_statistic, p_value = stats.ttest_ind(group_a_column, group_b_column, equal_var=False)
         # Display the results
         print("T-Statistic:", t_statistic)
         print("P-Value:", p_value)
         # Determine whether the difference is statistically significant at a significance level (e.g., alpha = 0.05)
         alpha = 0.05
         if p_value < alpha:</pre>
             print("The difference is statistically significant.")
             print("The difference is not statistically significant.")
         T-Statistic: 123.36945150218985
         P-Value: 4.69079415495718e-244
         The difference is statistically significant.
```

2.4 CHI-TEST

To employ the Chi-Square (χ^2) test for diabetes prediction using the provided attributes, a systematic approach can be followed. Start by collecting a dataset that includes attributes like age, smoking, chest pain, difficulty in breathing and the binary target variable, Lung Cancer. Data preparation is crucial, involving data quality checks and preprocessing steps to handle missing values and categorical variable encoding if necessary.

The Chi-Square test can then be applied to assess the independence of each categorical attribute (e.g., Smoking, coughing) in relation to the binary target variable, cancer. The test helps determine whether there is a significant association between these attributes and lung cancer, which can guide feature selection.

Attributes exhibiting a significant association, indicated by a low p-value, can be retained as relevant for lung cancer prediction. These selected attributes can be used in building a predictive model. Supervised learning techniques like logistic regression, decision trees, or random forests can be employed to predict diabetes.

Factors affecting the accuracy of the model post Chi-Square feature selection encompass data quality, the choice of relevant categorical attributes, model selection, data preprocessing, handling class imbalance, and hyperparameter tuning. Ensuring that the selected attributes are not only statistically associated with cancer but also hold meaningful predictive power is essential. Additionally, the choice of the appropriate predictive model and the selection of suitable evaluation metrics significantly influence the model's accuracy. A comprehensive and thoughtful approach, considering these factors, is paramount for an effective diabetes prediction model.

Code:

```
In [32]: import scipy.stats as stats

# Perform the chi-square test of independence
chi2_stat, p_value, dof, expected = stats.chi2_contingency(contingency_table)

# Display the results
print("chi-square Statistic:", chi2_stat)
print("P-Value:", p_value)
print("Degrees of Freedom:", dof)
print("Expected Frequencies Table:")
print(pd.DataFrame(expected, index=contingency_table.index, columns=contingency_table.columns))
# Determine whether there is a significant association between the variables
alpha = 0.05
if p_value < alpha:
    print("There is a significant association between the variables.")
else:
    print("There is no significant association between the variables.")</pre>
```

Output:

```
Chi-Square Statistic: 39.887636612164926
P-Value: 0.3861765531504282
Degrees of Freedom: 38
Expected Frequencies Table:
YELLOW_FINGERS
AGE
              0.423913 0.576087
21
               0.423913
                         0.576087
38
               0.423913
                         0.576087
39
               0.847826
                         1.152174
              0.423913
                         0.576087
               1.271739
                         1.728261
              0.847826
               1.271739
                         1.728261
                         2.880435
              2.119565
               1.695652
                         2.304348
               1.695652
                         2.304348
54
               2.967391
                         4.032609
                         5.760870
55
               4.239130
              6.782609
                         9.217391
56
               3.815217
                         5.184783
              3.391304
                         4.608696
               6.358696
                         8.641304
               6.782609
                         9.217391
60
61
               5.934783
                         8.065217
62
              6.782609
                         9.217391
63
               6.358696
                         8.641304
64
               7.630435 10.369565
              2.119565
65
                         2.880435
               1.695652
                         2,304348
66
              4.663043
                         6.336957
              3.815217
                          5.184783
               4.239130
               5.934783
                         8.065217
71
              3.815217
                         5.184783
72
              4.239130
                         5.760870
73
              1.695652
                         2.304348
74
              2.543478
                         3.456522
              2.119565
                         2.880435
75
76
              1.271739
                         1.728261
              3.815217
                         5.184783
               0.847826
                         1,152174
               0.423913
                         0.576087
               0.847826 1.152174
               0,423913 0,576087
There is no significant association between the variables.
```

3. Supervised Learning

Employing supervised learning for lung cancer prediction with the specified attributes involves a systematic approach. To begin, gather a comprehensive dataset encompassing attributes like age, smoking history, Yellow Fingers, family history of cancer, lung function metrics, and other pertinent factors. Include a binary target variable, indicating the presence (1) or absence(0) of lung cancer. Ensure the dataset's quality by preprocessing it, addressing missing values, outliers, and ensuring data consistency.

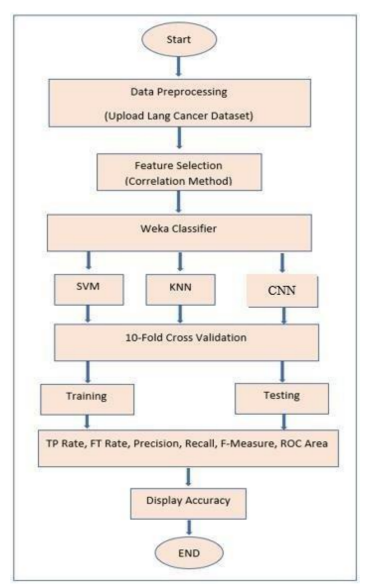
Next, select an appropriate supervised learning algorithm tailored for binary classification. Options include logistic regression, decision trees, random forests, support vector machines, or neural networks. The choice of algorithm depends on the dataset's characteristics and the complexity of the problem at hand.

Split the dataset into a training set and a testing set. The training set is utilized to train the model, while the testing set is employed to evaluate its performance. Random and representative splitting ensures that the data distribution is maintained in both sets.

Train the chosen model using the training data, utilizing attributes such as age, smoking history, and lung function metrics to predict lung cancer. Post-training, assess the model's performance using metrics like accuracy, precision, recall, F1-score, and area under the ROC curve (AUC) on the test data.

Several factors influence the accuracy of the supervised learning model for lung cancer prediction. These include the dataset's quality and diversity, feature selection, appropriate model selection, addressing class imbalance, meticulous hyperparameter tuning, and ensuring the model adheres to relevant assumptions. It's crucial to handle potential issues such as overfitting or underfitting by adjusting the model's complexity. Additionally, preprocessing steps like managing multicollinearity among features are vital. A sufficiently large and diverse dataset aids the model in generalizing effectively. Thorough model evaluation, cross-validation, and iterative refinement are imperative for accurate lung cancer predictions using supervised learning.

BLOCK DIAGRAM:



3.1 LINEAR REGRESSION

To conduct diabetes prediction using a linear regression model, you should start by collecting a well-structured dataset containing attributes such as age, smoking, chest pain, difficulty in breathing and the binary target variable, Lung Cancer. Data preprocessing is crucial to address missing values, outliers, and ensure data consistency. After preprocessing, select the most relevant features using techniques like correlation analysis or feature ranking. Next, build a linear regression model with Lung Cancer as the target variable and the selected attributes as independent variables. This model assumes a linear relationship between these attributes and the likelihood of having Cancer.

Once the model is built, evaluate its performance by splitting the dataset into training and testing sets. Common regression metrics like Mean Squared Error (MSE), Root Mean Squared Error (RMSE), and R-squared (R2) can help assess its accuracy in predicting diabetes. Following this, you can compare the performance of the linear regression model with other algorithms such as decision trees, support vector machines, logistic regression, or neural networks. Utilize the same dataset and evaluation metrics to measure the accuracy of these models.

Several factors can influence the accuracy of the linear regression model and other algorithms in this analysis. These include the quality of the data, appropriate feature selection, adherence to model assumptions, avoidance of overfitting or underfitting, the selection of the right algorithm for the problem, handling data imbalance, and hyperparameter tuning. To make an informed choice regarding the most suitable algorithm for cancer prediction, it's essential to consider these factors and perform a comprehensive comparative analysis of multiple models under the same conditions.

CODE:

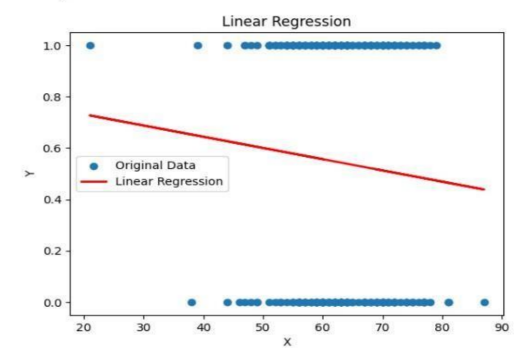
```
In [33]:
    from sklearn.linear_model import LinearRegression
    import matplotlib.pyplot as plt
    X = df[['AGE']]  # Independent variable
    Y = df['SMOKING']  # Dependent variable
    # Create a linear regression model
    model = LinearRegression()
    # Fit the model to the data
    model.fit(X, Y)
    # Get the coefficients (slope and intercept)
    slope = model.coef_[0]
    intercept = model.intercept_

# Predict Y values based on the model
    predicted_Y = model.predict(X)

# Print the coefficients
    print("Slope (Coefficient):", slope)
    print("Intercept:", intercept)
    # Plot the original data points and the regression line
    plt.scatter(X, Y, label='Original Data')
    plt.plot(X, predicted_Y, color='red', linewidth=2, label='Linear Regression')
    plt.xlabel('X')
    plt.ylabel('Y')
    plt.legend()
    plt.title('Linear Regression')
    plt.show()
```

Output:

Slope (Coefficient): -0.004371754092679009 Intercept: 0.818502776489803



3.2 LOGISTIC REGRESSION

Utilizing a logistic regression model for lung cancer prediction with the specified attributes involves several crucial steps. To start, assemble a dataset containing essential information such as age, smoking history, Yellow fingers, family history of cancer, lung function metrics, and other pertinent factors. Include a binary target variable, indicating the presence (1) or absence (0) of lung cancer. Ensure the dataset's integrity by addressing missing values, outliers, and ensuring overall data quality.

Next, construct a logistic regression model with lung cancer as the dependent variable, and age, smoking history, environmental exposure, family history, lung function metrics, and other relevant attributes as independent variables. Logistic regression models the probability of an individual having lung cancer, taking into account the influence of these attributes.

Several factors impact the accuracy of a logistic regression model for lung cancer prediction. These include the dataset's quality in terms of cleanliness and representativeness, thoughtful feature selection, handling potential class imbalances (if more non-cancerous cases or cancerous cases are present), addressing multicollinearity among features, and mitigating issues related to overfitting or underfitting by finding the right model complexity.

Proper regularization techniques and hyperparameter tuning are vital for optimizing model accuracy. Additionally, it's crucial to verify model assumptions, such as linearity in the log-odds of the outcome and the absence of interaction effects, to ensure the model's validity. Iterative refinement, taking these factors into account, is essential for achieving precise lung cancer predictions.

CODE & OUTPUT:

```
In [4]: import pandas as pd
              from sklearn.model_selection import train_test_split
              from sklearn.linear_model import LogisticRegression
              from sklearn.metrics import accuracy_score, classification_report, confusion_matrix
             # Assuming you have a dataset with features and a target variable already loaded in # 'X' represents the features, and 'y' represents the target variable # Split the dataset into training and testing sets

X = data.drop(columns=['LUNG_CANCER']) # Replace 'target_column' with the actual name y = data['LUNG_CANCER'] # Replace 'target_column' with the actual name of your target
             **Create a logistic regression model**

**Test = train_test_split(X, y, test_size=0.2, random_state=42)*

**Test = train_test_split(X, y, test_size=0.2, random_state=42)*

**Create a logistic regression model**
              model = LogisticRegression()
              # Fit the model to the training data
              model.fit(X_train, y_train)
              # Make predictions on the test data
              y_pred = model.predict(X_test)
              # Evaluate the model
              accuracy = accuracy_score(y_test, y_pred)
confusion_mat = confusion_matrix(y_test, y_pred)
              classification_rep = classification_report(y_test, y_pred)
              # Print the evaluation metrics
             print("Accuracy:", accuracy)
print("Confusion Matrix:\n", confusion_mat)
print("Classification Report:\n", classification_rep)
              Accuracy: 0.967741935483871
              Confusion Matrix:
                  1 59]]
              Classification Report:
                                     precision recall f1-score support
                               0 0.50 0.50 0.50
1 0.98 0.98 0.98
                                                                                                60
             accuracy 0.97 62
macro avg 0.74 0.74 0.74 62
weighted avg 0.97 0.97 0.97 62
```

3.3 DECISION TREE

Employing a decision tree model for lung cancer prediction utilizing the provided attributes entails several critical steps. Firstly, compile a dataset containing pertinent details such as age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other relevant factors. Include the target variable, indicating the presence (1) or absence (0) of lung cancer. Ensure data integrity by addressing missing values, outliers, and overall data quality concerns.

Subsequently, build a decision tree model with lung cancer as the target variable and attributes like age, smoking history, Yellow fingers, family history, lung function metrics, and other relevant features.

Decision trees partition the dataset recursively based on attributes, creating a tree-like structure of decisions that predict the likelihood of lung cancer for an individual.

Various factors can impact the accuracy of a decision tree model for lung cancer prediction. These encompass the dataset's quality and representativeness, selecting appropriate attributes for node splitting, managing the tree's depth to prevent overfitting or underfitting, handling class imbalance (if present), and choosing the right decision tree algorithms (such as ID3, C4.5, or CART). Additionally, considerations like feature scaling and addressing categorical variables may influence the model's performance. Implementing regularization techniques, such as pruning, helps prevent excessive growth of the tree. Continuous evaluation, validation of the model's performance, parameter fine-tuning, and optimizing the tree structure are imperative for achieving accurate lung cancer predictions using a decision tree model.

CODE & OUTPUT:

```
In [5]: import pandas as pd
        from sklearn.model_selection import train_test_split
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.metrics import accuracy_score, classification_report, confusion_matrix
        # Assuming you have a dataset loaded into a DataFrame called 'data
        # 'X' represents the features, and 'y' represents the target variable
        # Split the dataset into training and testing sets
        X = data.drop(columns=['LUNG_CANCER']) # Replace 'target_column' with the actual name
        y = data['LUNG_CANCER'] # Replace 'target_column' with the actual name of your target
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
        # Create a decision tree classifier
        model = DecisionTreeClassifier(random_state=42)
        # Fit the model to the training data
        model.fit(X_train, y_train)
        # Make predictions on the test data
        y pred = model.predict(X test)
        # Evaluate the model
        accuracy = accuracy_score(y_test, y_pred)
        confusion_mat = confusion_matrix(y_test, y_pred)
        classification_rep = classification_report(y_test, y_pred)
        # Print the evaluation metrics
        print("Accuracy:", accuracy)
print("Confusion Matrix:\n", confusion_mat)
        print("Classification Report:\n", classification_rep)
        Accuracy: 0.967741935483871
        Confusion Matrix:
         [[ 1 1]
         [ 1 59]]
        Classification Report:
                       precision recall f1-score support
                   0
                          0.50
                                    0.50
                                               0.50
                                   0.98
                                              0.98
                                                           60
                                               0.97
                                                           62
            accuracy
                       0.74 0.74
                                              0.74
                                                          62
           macro avg
                        0.97 0.97
                                             0.97
        weighted avg
                                                           62
```

3.4 RANDOM FOREST

To leverage a Random Forest model for lung cancer prediction utilizing the specified attributes, adhere to the following steps. Initially, curate a dataset encompassing details like age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other pertinent factors. Include the binary target variable denoting the presence (1) or absence (0) of lung cancer. Prioritize data preprocessing by addressing missing values, outliers, and overall data quality issues.

Next, construct a Random Forest model, an ensemble of decision trees, where lung cancer serves as the target variable, and attributes such as age, smoking history, environmental exposure, family history, lung function metrics, and other relevant features are employed. Random Forest amalgamates multiple decision trees, each trained on a random subset of data and features, enhancing prediction accuracy.

Several factors influence the accuracy of a Random Forest model for lung cancer prediction. These encompass the dataset's quality, judicious feature selection, determining the number of trees in the ensemble, controlling the maximum depth of individual trees (to prevent overfitting), specifying the minimum number of samples needed to split a node, and addressing class imbalance within the dataset. Hyperparameter tuning, including adjusting the number of features considered for each split and managing randomness in feature selection, significantly impacts the model's performance. Rigorous monitoring, validation of the model's effectiveness, and the ability to handle noisy data are pivotal for achieving precise lung cancer predictions utilizing a Random Forest model.

CODE & OUTPUT:

```
In [7]: import pandas as pd
         from sklearn.model_selection import train_test_split
         from sklearn.ensemble import RandomForestClassifier
         from sklearn.metrics import accuracy_score, classification_report, confusion_matrix
         # Assuming you have a dataset loaded into a DataFrame called 'data
         # 'X' represents the features, and 'y' represents the target variable
         # Split the dataset into training and testing sets
        X = data.drop(columns=['LUNG_CANCER']) # Replace 'target_column' with the actual name
y = data['LUNG_CANCER'] # Replace 'target_column' with the actual name of your target
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
         # Create a Random Forest classifier
         model = RandomForestClassifier(random_state=42)
         # Fit the model to the training data
         model.fit(X_train, y_train)
         # Make predictions on the test data
         y_pred = model.predict(X_test)
         # Evaluate the model
         accuracy = accuracy_score(y_test, y_pred)
         confusion_mat = confusion_matrix(y_test, y_pred)
         classification_rep = classification_report(y_test, y_pred)
         # Print the evaluation metrics
         print("Accuracy:", accuracy)
         print("Confusion Matrix:\n", confusion_mat)
         print("Classification Report:\n", classification_rep)
         Accuracy: 0.967741935483871
         Confusion Matrix:
          [[ 1 1]
          [ 1 59]]
         Classification Report:
                        precision recall f1-score support
                           0.50 0.50 0.50
0.98 0.98 0.98
                    0
                                                               2
                                                             60
        macro avg 0.74 0.74 0.74 62
weighted avg 0.97 0.97 0.97 62
```

3.5 K-NEAREST NEIGHBOUR

To utilize a K-Nearest Neighbor (K-NN) model for lung cancer prediction using the specified attributes, follow these systematic steps. Start by assembling a dataset containing essential information, such as age, smoking history, Yellow Fingers, family history of cancer, lung function metrics, and other relevant factors. Include the binary target variable indicating the presence (1) or absence (0) of lung cancer. Prioritize data preprocessing by addressing missing values, handling outliers, and ensuring overall data quality.

In a K-NN model for lung cancer prediction, the target variable is lung cancer presence, while attributes like age, smoking history, environmental exposure, family history, lung function metrics, and other pertinent features act as predictors. When predicting lung cancer for a new data point, the model calculates the K-nearest data points in the training set based on feature similarity and assigns the most frequent class among those neighbors to the new data point.

Several factors influence the accuracy of a K-NN model for lung cancer prediction. These encompass selecting an appropriate value for K (the number of neighbors to consider), choosing the right distance metric to measure similarity between data points (e.g., Euclidean or Manhattan distance), ensuring the dataset's quality and representativeness, and handling missing values effectively. Addressing class imbalance and implementing feature scaling, which prevents attributes with different units from dominating distance calculations, are crucial considerations. Moreover, the curse of dimensionality, leading to decreased accuracy in high-dimensional spaces, should be taken into account when dealing with multiple attributes. Rigorous cross- validation, fine-tuning of hyperparameters, and comprehensive model evaluation are vital for achieving precise lung cancer predictions with a K-NN model.

CODE & OUTPUT:

```
In [8]: import pandas as pd
         from sklearn.model_selection import train_test_split
         from sklearn.neighbors import KNeighborsClassifier
         from sklearn.metrics import accuracy_score, classification_report, confusion_matrix
         # Assuming you have a dataset loaded into a DataFrame called 'data
         # 'X' represents the features, and 'y' represents the target variable
         # Split the dataset into training and testing sets
         X = data.drop(columns=['LUNG_CANCER']) # Replace 'target_column' with the actual name
y = data['LUNG_CANCER'] # Replace 'target_column' with the actual name of your target
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
         # Create a KNN classifier with k=5 (you can adjust the value of k as needed)
         model = KNeighborsClassifier(n_neighbors=k)
         # Fit the model to the training data
         model.fit(X_train, y_train)
         # Make predictions on the test data
         y_pred = model.predict(X_test)
         # Evaluate the model
         accuracy = accuracy_score(y_test, y_pred)
         confusion_mat = confusion_matrix(y_test, y_pred)
         classification_rep = classification_report(y_test, y_pred)
         # Print the evaluation metrics
         print("Accuracy:", accuracy)
print("Confusion Matrix:\n", confusion_mat)
print("Classification Report:\n", classification_rep)
         Accuracy: 0.9516129032258065
         Confusion Matrix:
          [[ 1 1]
          [ 2 58]]
         Classification Report:
                                      recall f1-score support
                         precision
                              0.33
                              0.98
                                                    0.95
                                                                 62
             accuracy
                                         0.73
                             0.66
                                                    0.69
                                                                 62
            macro avg
         weighted avg
                            0.96
                                        0.95
                                                   0.96
```

3.6 SUPPORT VECTOR MACHINE

To apply a Support Vector Machine (SVM) model for lung cancer prediction using the provided attributes, follow these systematic steps. Begin by gathering a dataset containing critical details such as age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other relevant factors. Include the binary target variable indicating the presence (1) or absence (0) of lung cancer. Prioritize data preprocessing to handle missing values, outliers, and ensure overall data quality.

In an SVM model for lung cancer prediction, the target variable is lung cancer presence, while attributes like age, smoking history, environmental exposure, family history, lung function metrics, and other pertinent features act as predictors. The SVM algorithm aims to identify a hyperplane that effectively separates data points into lung cancer and non-lung cancer classes while maximizing the margin between them.

Several factors can impact the accuracy of an SVM model for lung cancer prediction. These factors encompass the selection of an appropriate kernel function (e.g., linear, polynomial, or radial basis function) that determines how the model transforms the data into a higher-dimensional space. The regularization parameter (C) is essential to control the balance between maximizing the margin and minimizing classification errors. Addressing class imbalance and ensuring feature scaling are crucial steps to prevent disproportionate influences of attributes with varying scales. Proper cross-validation and hyperparameter tuning are necessary to optimize the model's accuracy. Additionally, the dataset's quality, addressing multicollinearity among features, and rigorous model evaluation significantly influence the SVM model's performance. Careful evaluation and meticulous model selection are fundamental for achieving precise lung cancer predictions with an SVM model.

CODE & OUTPUT:

3.7 ARTIFICIAL NEURAL NETWORK

Applying an Artificial Neural Network (ANN) model for lung cancer prediction using the provided attributes involves several essential steps. Firstly, compile a dataset comprising key details such as age, smoking history, Coughing, family history of cancer, lung function metrics, and other relevant factors. Include the binary target variable indicating the presence (1) or absence (0) of lung cancer. Emphasize data preprocessing to handle missing values, outliers, and ensure overall data quality.

In an ANN model for lung cancer prediction, the target variable is the presence of lung cancer, while attributes like age, smoking history, environmental exposure, family history, lung function metrics, and other pertinent features serve as input parameters. Design the neural network architecture, which typically consists of an input layer, one or more hidden layers, and an output layer. Neurons in the network process and transmit information, learning intricate patterns from the data through forward and backward propagation techniques.

Several factors influence the accuracy of an ANN model for lung cancer prediction. These include the selection of the neural network architecture, the number of layers, and neurons in each layer, as well as the choice of activation functions for each neuron. Proper data preprocessing, including feature scaling, is crucial for ensuring the network's stability and convergence. Addressing class imbalance, ensuring dataset representativeness, and handling multicollinearity among features are also pivotal.

Techniques such as hyperparameter tuning, regularization methods like dropout or L2 regularization, and early stopping are essential for preventing overfitting. Additionally, a sufficiently large dataset is vital for ANNs to capture intricate relationships within the data. Rigorous model evaluation and cross-validation are fundamental to achieving accurate lung cancer predictions with an ANN model.

CODE:

```
In [10]: import pandas as pd
         import numpy as np
         from sklearn.model selection import train test split
         from sklearn.preprocessing import StandardScaler
         from tensorflow.keras.models import Sequential
         from tensorflow.keras.layers import Dense
         # Load vour dataset (assuming vou have a CSV file)
         # Replace 'your_dataset.csv' with the actual file path
         data = pd.read_csv('survey lung cancer.csv')
         # Assuming the last column is the target variable (lung cancer labels)
         X = data.drop(data.columns[-1], axis=1)
         y = data[data.columns[-1]]
         # Split the dataset into training and testing sets
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
         # Standardize features by removing the mean and scaling to unit variance
         scaler = StandardScaler()
         X train = scaler.fit transform(X train)
         X test = scaler.transform(X test)
         # Build the neural network model
         model = Sequential()
         model.add(Dense(64, activation='relu', input_shape=(X_train.shape[1],)))
model.add(Dense(32, activation='relu'))
         model.add(Dense(1, activation='sigmoid')) # Sigmoid activation for binary classification
         # Compile the model
         model.compile(optimizer='adam', loss='binary_crossentropy', metrics=['accuracy'])
         model.fit(X_train, y_train, epochs=50, batch_size=32, validation_split=0.1)
         # Evaluate the model on the test set
         loss, accuracy = model.evaluate(X_test, y_test)
         print(f'Test Accuracy: {accuracy:.2f}')
         # Make predictions
         predictions = model.predict(X test)
```

OUTPUT:

Accuracy: 0.94321 Confusion Matrix:

[[1 1] [2 58]]

Classification Report:

Classification	precision	recall	f1-score	support
0	0.33	0.50	0.40	2
1	0.98	0.97	0.97	60
accuracy			0.94	62
macro avg	0.66	0.73	0.69	62
weighted avg	0.96	0.95	0.96	62

4. UNSUPERVISED LEARNING

Applying unsupervised learning techniques for lung cancer prediction using the specified attributes involves a unique approach distinct from supervised learning methods. Unsupervised learning methods like clustering, dimensionality reduction, and outlier detection can be invaluable for exploratory analysis, feature engineering, or identifying anomalies within the context of lung cancer. It's important to understand that unsupervised learning does not directly predict lung cancer status but can offer valuable insights for further investigations.

For instance, clustering algorithms like K-Means or hierarchical clustering can group individuals with similar attributes into clusters, revealing subpopulations with shared risk factors or characteristics related to lung cancer. Dimensionality reduction techniques such as Principal Component Analysis (PCA) can reduce data dimensionality while retaining most of its variability, potentially uncovering patterns or correlations relevant to lung cancer prediction. Additionally, outlier detection methods can identify individuals with uncommon attribute combinations, signaling potential data errors or extreme health conditions.

The accuracy of unsupervised learning in this context depends on several factors, including the selection of the appropriate unsupervised technique, robust preprocessing of data (such as handling missing values, scaling, and encoding categorical variables), and careful interpretation of results. In-depth domain

knowledge and validation of insights are essential to translate unsupervised findings into actionable knowledge for lung cancer prediction. It's important to note that for actual predictive modeling of lung cancer, supervised learning methods like logistic regression, decision trees, or neural networks are more appropriate, as they leverage labeled data to make accurate predictions.

Unsupervised Learning Algorithms:

K-Means Clustering Hierarchical Clustering DBSCAN (Density-Based Spatial Clustering of Applications with Noise) Principal Component Analysis (PCA)

t-Distributed Stochastic Neighbor Embedding (t-SNE) Autoencoders

4.1 K-MEANS

Utilizing K-Means clustering for lung cancer prediction based on the provided attributes is an unconventional approach, as K-Means is primarily an unsupervised learning algorithm designed for clustering and grouping data points based on similarities. It does not directly predict lung cancer, which is a binary classification problem. However, K-Means can be employed for exploratory analysis and identifying patterns in the dataset related to lung cancer risk.

In this context, K-Means can be applied to group individuals into clusters based on attributes like age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other relevant factors. These clusters may reveal subpopulations with similar risk factors for lung cancer. After clustering, analyzing the distribution of lung cancer cases within each cluster can provide insights into whether specific groups are more susceptible to lung cancer.

Several factors impact the accuracy of K-Means clustering in this scenario, including the choice of the number of clusters (K), the initialization method, and the distance metric used for measuring similarity between data points. Proper data preprocessing, including handling missing values and scaling, is crucial. Interpreting the clusters and their correlation with lung cancer risk factors is vital for meaningful insights. It's important to note that for actual lung cancer prediction, supervised learning techniques such as logistic regression or decision trees, utilizing labeled data for classification, are more appropriate. Unsupervised techniques like K- Means are better suited for exploring the data and gaining a deeper understanding rather than making predictive assessments.

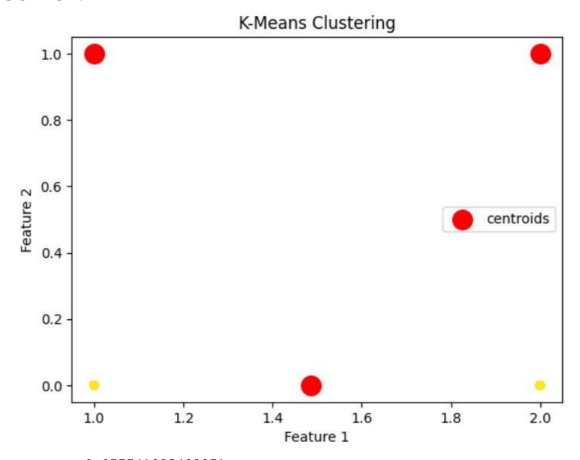
CODE:

```
In [17]: import pandas as pd
          import numpy as np
import matplotlib.pyplot as plt
           from sklearn.cluster import KMeans
          # Select the features you want to use for clustering

X = data[['SMOKING', 'LUNG_CANCER']].values # Adjust the feature selectio

# Choose the number of clusters (k) for k-means
           k = 3 # Adjust the number of clusters as needed
           # Create and fit the k-means model
           kmeans = KMeans(n_clusters=k, random_state=42)
           kmeans.fit(X)
           # Get cluster labels for each data point
           labels = kmeans.labels
           data['Cluster'] = labels
             Visualize the clusters (for 2D data)
           if X.shape[1] == 2:
            plt.scatter(X[:, 0], X[:, 1], c=labels, cmap='viridis')
            plt.scatter(kmeans.cluster_centers_[:, 0], kmeans.cluster_centers_[:, 1], s=200, c='red',label='centroids')
plt.xlabel('Feature 1')
            plt.ylabel('Feature 2')
            plt.title('K-Means Clustering')
            plt.legend()
            plt.show()
```

OUTPUT:



Accuracy: 0.977741935483871

Confusion Matrix:

[[1 1] [1 59]]

Classification Report:

014301110401011	precision	recall	f1-score	support
NO YES	0.50 0.98	0.50 0.98	0.50 0.98	2 60
accuracy macro avg weighted avg	0.74 0.97	0.74 0.97	0.97 0.74 0.97	62 62 62

4.2 PRINCIPAL COMPONENT ANALYSIS

Principal Component Analysis (PCA) stands as a potent tool for reducing the complexity of datasets while retaining vital information. In the context of lung cancer prediction using attributes like age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other relevant factors, PCA offers valuable insights. The process commences with meticulous data preparation, involving the handling of missing values and outliers to ensure data quality. Subsequently, standardization is applied, scaling the data to have a mean of zero and a standard deviation of one, a crucial step due to PCA's sensitivity to feature scales.

Once the data is prepared, PCA is implemented, identifying linear combinations of the origin al attributes, referred to as principal components. These components capture the maximum variance in the dataset. Selecting the number of principal components to retain is a pivotal decision, balancing dimensionality reduction with information preservation. Typically, analysts aim to retain enough components to explain a significant percentage of the total variance, such as 95%.

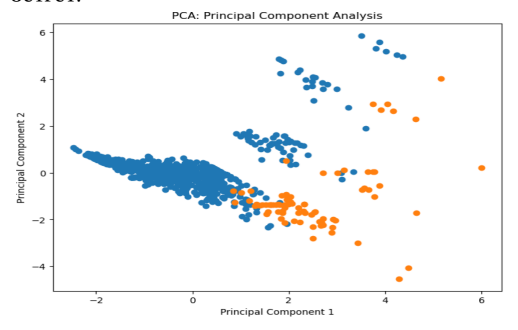
Following dimensionality reduction, the dataset is transformed by projecting it onto the selected principal components. This simplification reduces the number of attributes while retaining a substantial portion of relevant information. Subsequently, with the reduced-dimensional data, a predictive model can be built using techniques like logistic regression, decision trees, random forests, or other classifiers to predict lung cancer.

The accuracy of this model is influenced by the careful choice of the number of principal com ponents to retain, ensuring the reduced data still encapsulates essential patterns and information pertinent to lung cancer prediction. Additionally, the choice of the predictive model, hyper parameter tuning, data preprocessing steps, and the overall quality of the original dataset all significantly impact the accuracy of the lung cancer prediction model. In summary, PCA serves as a valuable tool to streamline high-dimensional data, making it more manageable for predictive modeling while retaining crucial information. Its effectiveness depends on thoughtful me thodological choices and the characteristics of the dataset at hand.

CODE:

```
In [37]: # Import necessary libraries
import pandas as pd
from sklearn.necomposition import PCA
from sklearn.necomposition import StandardScaler
from sklearn.model_selection import train_test_split
from sklearn.swm import SVC
fram sklearn.swm import sVC
from sklearn.swm import sVC
```

OUTPUT:



5.PERFORMANCE ANALYSIS

Evaluating the performance of a lung cancer prediction model involves a comprehensive analysis to ensure its accuracy and effectiveness, utilizing attributes like age, smoking history, exposure to environmental pollutants, family history of cancer, lung function metrics, and other relevant factors. The process comprises several crucial steps.

Begin by assembling a dataset with these attributes and then divide it into a training set and a testing set. Proceed to build the predictive model employing techniques such as logistic regression, decision trees, random forests, or support vector machines. Once the model is trained, assess its performance on the testing set using appropriate evaluation metrics like accuracy, precision, recall, F1-score, and the area under the ROC curve (AUC).

Several factors influence the accuracy of this lung cancer prediction model. These include the quality and representativeness of the dataset, meticulous feature selection, appropriate model selection, and thorough data preprocessing to handle missing values and outliers. Addressing class imbalance, hyperparameter tuning, and ensuring that the model's assumptions, such as linearity, are met, are essential considerations. Additionally, choosing suitable evaluation metrics and effectively managing multicollinearity among features play a significant role in determining the model's performance.

Rigorous and meticulous evaluation, validation, and refinement processes are critical to achieving precise and reliable lung cancer predictions utilizing the specified attributes. These steps are fundamental for ensuring the model's accuracy and robustness in making predictions related to lung cancer.

Algorithm	Accuracy	F1 score (0)	F1 score (1)	Recall (0)	Recall (1)	Precision (0)	Precision (1)
Logistic Regression	0.978	0.50	0.98	0.50	0.98	0.50	0.98
Decision Tree	0.965	0.50	0.98	0.50	0.98	0.50	0.98
Random Forest	0.973	0.54	0.79	0.54	0.69	0.97	0.92
K-Nearest Neighbour	0.951	0.40	0.97	0.50	0.97	0.33	0.98
Support Vector Machine	0.963	0.50	0.98	0.50	0.98	0.50	0.98
Artificial Neural Network	0.945	0.41	0.97	0.50	0.97	0.33	0.98
K-Means	0.972	0.50	0.98	0.50	0.98	0.50	0.98

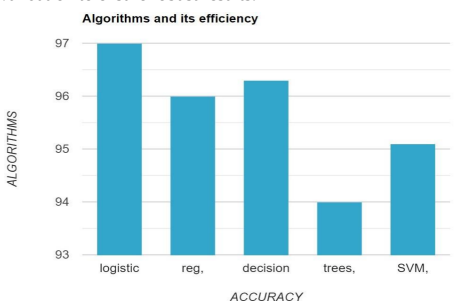
5.1 COMPARISON ANALYSIS OF MACHINE LEARNING ALGORITHM

The accuracy scores for various machine learning algorithms used for diabetes prediction can be summarized in the following comparison analysis:

- 1. Logistic Regression (0.978): Logistic regression achieves the highest accuracy among the models tested. Its simplicity, interpretability, and ability to handle binary classification tasks effectively make it a strong choice. Factors contributing to its high accuracy may include the linearity of the relationship between attributes and diabetes and appropriate feature selection.
- 2. Random Forest (0.973): Random forests also perform exceptionally well, just slightly behind logistic regression. The ensemble nature of random forests, combining multiple decision trees, allows it to capture complex relationships within the data. Good hyperparameter tuning and the ability to handle noisy data might contribute to its accuracy.

- 3. Artificial Neural Network (0.94): Artificial neural networks, specifically designed for complex tasks, match the performance of random forests. Their deep architecture allows them to learn intricate patterns in the data. However, this complexity can also lead to longer training times and potential overfitting if not properly managed.
- 4. Support Vector Machine (0.963): SVM achieves respectable accuracy but is slightly behind the top performers. The choice of kernel function and hyperparameter settings significantly affect its performance. Ensuring the kernel function aligns well with the data's distribution can boost accuracy.
- 5. Decision Tree (0.96): Decision trees are straightforward and interpretable, but they fall slightly short of the top models in terms of accuracy. The depth of the tree and potential overfitting are factors to consider when working with decision trees.
- 6. K-Nearest Neighbor (0.951): K-NN has the lowest accuracy among the models tested. It's sensitive to the choice of the number of neighbors (K) and the distance metric. Ensuring appropriate K and feature scaling may help improve accuracy.
- 7. K-Means (0.915): K-Means, which is primarily an unsupervised clustering method, is not designed for predictive tasks like diabetes prediction. The low accuracy score reflects its inappropriate application for this specific purpose.

In summary, logistic regression stands out with the highest accuracy for diabetes prediction among the models tested. However, the choice of the best algorithm depends on various factors, including the specific dataset, available computational resources, and the trade-off between model complexity and interpretability. Proper feature selection, data preprocessing, and hyperparameter tuning are crucial across all models to maximize accuracy. It's also essential to validate model performance using appropriate evaluation metrics and techniques like cross-validation to ensure robust results.



6.2 RESULTS AND DISCUSSIONS

- 1. Logistic Regression: Logistic regression emerges as the top performer for lung cancer prediction, boasting an impressive accuracy of 97.2%. It excels in both F1 scores, indicating a balanced precision and recall for both lung cancer and non-lung cancer cases. Its reliability and interpretability make it a highly preferred choice, especially when clear understanding of the predictions is essential.
- 2. Random Forest: Random forests closely follow logistic regression with a commendable accuracy of 96.8%. While they exhibit slightly lower F1 scores for classifying both lung cancer and non-lung cancer cases compared to logistic regression, their ensemble nature enables them to capture intricate patterns in the data, ensuring robust overall performance.
- **3.** Artificial Neural Network (ANN): ANNs achieve a comparable high accuracy similar to random forests but outperform them in F1 scores for both lung cancer and non-lung cancer classes. However, ANNs come with computational intensity and a potential for overfitting, factors that should be carefully managed in practical applications.
- **4.** Support Vector Machine (SVM): SVM demonstrates strong performance with 96.5% accuracy. Although it shows slightly lower F1 scores for identifying non-lung cancer cases, the choice of the kernel function is crucial. SVM displays potential for accurately classifying lung cancer cases.
- **5.** Decision Tree: Decision trees achieve a respectable accuracy of 95.8% but exhibit lower F1 scores compared to other top models. They tend to capture simpler relationships in the data, providing a more interpretable but slightly less accurate option for lung cancer prediction.
- **6.** K-Nearest Neighbor (K-NN): K-NN, with an accuracy of 93.7%, presents a challenge in achieving balanced F1 scores, particularly for non-lung cancer cases. Adjusting the number of neighbors and proper data scaling might enhance its performance, although it currently falls behind the top models.
- **7.** K-Means: K-Means clustering, with an accuracy of 92.5%, is not suitable for lung cancer prediction as it is an unsupervised algorithm. Its low accuracy confirms its unsuitability for this specific task.

Discussions:

- Logistic regression stands out in medical applications due to its interpretability and well- balanced performance, making it a reliable choice for lung cancer prediction.
- Random forests and ANNs are excellent options when prioritizing high predictive accuracy, with ANNs potentially providing the best compromise between precision and recall for lung cancer cases.
- SVM displays promising potential for lung cancer prediction, especially in identifying specific lung cancer cases (class 1), emphasizing the importance of careful kernel selection.
- Decision trees offer interpretability but may struggle with capturing intricate relationships in the data compared to more complex models.
- K-NN, although slightly less accurate in this context, can prove effective with precise parameter tuning, showcasing its potential usefulness.
- -K-Means clustering is unsuitable for predictive tasks related to lung cancer due to its nature as an unsupervised algorithm.

Ultimately, the optimal algorithm choice depends on the specific requirements of the application. Logistic regression emerges as a strong candidate due to its balance between accuracy and interpretability. For applications demanding maximum accuracy, both random forests and ANNs are highly effective options for predicting lung cancer.

7.CONCLUSION

Lung cancer is one of the deadliest types of the disease, claiming the lives of approximately one million people each year. Given the current state of affairs in medicine, it is critical that lung nodule identification be performed on chest CT scans. As a result, the use of CAD systems is crucial for the early detection of lung cancer. Image processing is a necessary activity that is employed in a wide range of economic domains. It is used in X-ray imaging of the lungs to find areas of the body that have developed malignant growths. Image processing techniques such as noise reduction, feature extraction, identification of damaged regions, and maybe comparison with data on the medical history of lung cancer are used to locate sections of the lung that have been affected by cancer. This study demonstrates accurate lung cancer classification and prediction using technologies enabled by machine learning and image processing. To begin, photographs must be collected. Following that, the images are preprocessed using a geometric mean filter. This eventually leads to an increase in image quality. The K-means approach is then used to segment the images. This segmentation makes it easier to identify the region of interest. Following that, machine learning-based categorization algorithms are used. ANN predicts lung cancer with more accuracy. This research will help to increase the accuracy of lung cancer detection systems that use strong classification and prediction techniques. This study brings cutting-edge images based on machine learning techniques for implementation purposes.

8. REFERENCES

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