

VISVESVARAYA TECHNOLOGICAL UNIVERSITY
“JnanaSangama”, Belgaum -590014, Karnataka.



LAB REPORT
On

MACHINE LEARNING

Submitted by

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1BM21CS086

**in partial fulfillment for the award of the degree of
BACHELOR OF ENGINEERING
in
COMPUTER SCIENCE AND ENGINEERING**



**B.M.S. COLLEGE OF ENGINEERING
(Autonomous Institution under VTU)
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**B. M. S. College of Engineering,
Bull Temple Road, Bangalore 560019
(Affiliated To Visvesvaraya Technological University, Belgaum)
Department of Computer Science and Engineering**

CERTIFICATE



This is to certify that the Lab work entitled "**MACHINE LEARNING**" carried out by **KANJIKA SINGH (1BM21CS086)**, who is bonafide student of **B. M. S. College of Engineering**. It is in partial fulfillment for the award of **Bachelor of Engineering in Computer Science and Engineering** of the Visvesvaraya Technological University, Belgaum during the year 2023-24. The Lab report has been approved as it satisfies the academic requirements in respect of Machine Learning Lab - **(22CS6PCMAL)**work prescribed for the said degree.

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Course outcomes:

CO1	Apply machine learning techniques in computing systems
CO2	Evaluate the model using metrics
CO3	Design a model using machine learning to solve a problem
CO4	Conduct experiments to solve real-world problems using appropriate machine learning techniques

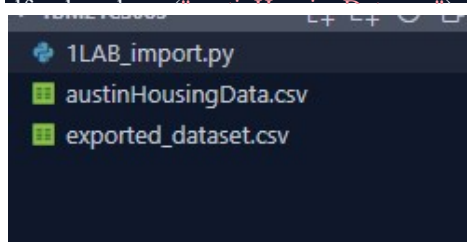
1. Write a python program to import and export data using Pandas library functions

Importing

```
import pandas as pd
data = pd.read_csv('content/austinHousing.csv')
data.head()
```

Output:

```
PS C:\Users\student\Documents\1BM21CS083> python -u "c:\Users\student\Documents\1BM21CS083\1LAB_import.py"
      zpid      city    streetAddress zipcode ... numOfBathrooms numOfBedrooms numOfStories   homeImage
0  111373431  pflugerville  14424 Lake Victor Dr  78660 ...           3.0              4                 2  111373431_ffce26843283d3365c11d81b8e6bcd6f-p_f...
1  120900430  pflugerville   1104 Strickling Dr  78660 ...           2.0              4                 1  120900430_8255c127be8dcf0a1a18b7563d987088-p_f...
2  2084491383  pflugerville  1408 Fort Dessau Rd  78660 ...           2.0              3                 1  2084491383_a2ad649e1a7a098111dcea084a11c855-p...
3  120901374  pflugerville   1025 Strickling Dr  78660 ...           2.0              3                 1  120901374_b469367a619da85b1f5ceb69b675d88e-p_f...
4  60134862  pflugerville  15005 Donna Jane Loop  78660 ...           3.0              3                 2  60134862_b1a48a3df3f11e005bb913873e98ce2-p_f.jpg
[5 rows x 47 columns]
PS C:\Users\student\Documents\1BM21CS083>
```



Exporting

```
url = "archive.ics.uci.edu/ml/machine-learning-databases/iris/iris.data"
col_names = ["sepal-length", "sepal-width", "petal length",
             "petal-width", "class"]
iris_data = pd.read_csv(url, names=col_names)
iris_data.head()
```

```
import pandas as pd
url = "https://archive.ics.uci.edu/ml/machine-learning-databases/iris/iris.data"

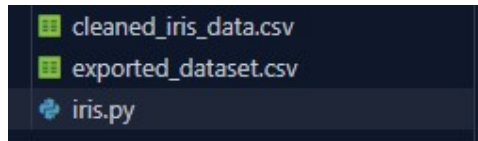
col_names = ["sepal_length_in_cm",
            "sepal_width_in_cm",
            "petal_length_in_cm",
            "petal_width_in_cm",
            "class"]
iris_data = pd.read_csv(url, names=col_names)

print(iris_data.head())

iris_data.to_csv("cleaned_iris_data.csv")
```

Output:

```
PS C:\Users\student\Documents\1BM21CS083> python -u "c:\Users\student\Documents\1BM21CS083\1lab\iris.py"
   sepal_length_in_cm  sepal_width_in_cm  petal_length_in_cm  petal_width_in_cm      class
0           5.1          3.5            1.4            0.2  Iris-setosa
1           4.9          3.0            1.4            0.2  Iris-setosa
2           4.7          3.2            1.3            0.2  Iris-setosa
3           4.6          3.1            1.5            0.2  Iris-setosa
4           5.0          3.6            1.4            0.2  Iris-setosa
PS C:\Users\student\Documents\1BM21CS083>
```



2. Demonstrate various data pre-processing techniques for a given dataset

3.

Algorithm

- 1) Import dataset using pandas
- 2) Perform dataset.shape() to analyse shape of dataset
- 3) use isnull() function from pandas to analyse missing values.
- 4) drop or fill missing values according to your usecase. Example dropna() and ffill().
- 5) use libraries like Scikit learn etc to perform more preprocessing if required

```
%matplotlib inline
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
import sklearn

dataset = pd.read_csv("Data.csv")
df = pd.DataFrame(dataset)
print(df.head())

      Country   Age   Salary Purchased
0    France  44.0  72000.0      No
1     Spain  27.0  48000.0     Yes
2  Germany  30.0  54000.0      No
3     Spain  38.0  61000.0      No
4  Germany  40.0       NaN     Yes

X = df.iloc[:, :-1].values
y = df.iloc[:, -1].values

print(X)

[['France' 44.0 72000.0]
 ['Spain' 27.0 48000.0]
 ['Germany' 30.0 54000.0]
 ['Spain' 38.0 61000.0]
 ['Germany' 40.0 nan]
 ['France' 35.0 58000.0]
 ['Spain' nan 52000.0]
 ['France' 48.0 79000.0]
 ['Germany' 50.0 83000.0]
 ['France' 37.0 67000.0]]

print(y)

['No' 'Yes' 'No' 'No' 'Yes' 'Yes' 'No' 'Yes' 'No' 'Yes']
```

```
df.isnull().sum()
```

```
Country      0  
Age         1  
Salary       1  
Purchased    0  
dtype: int64
```

```
df1 = df.copy()
```

```
# summarize the shape of the raw data  
print("Before:",df1.shape)  
  
# drop rows with missing values  
df1.dropna(inplace=True)  
  
# summarize the shape of the data with missing rows removed  
print("After:",df1.shape)
```

```
Before: (10, 4)  
After: (8, 4)
```

```
: df2
```

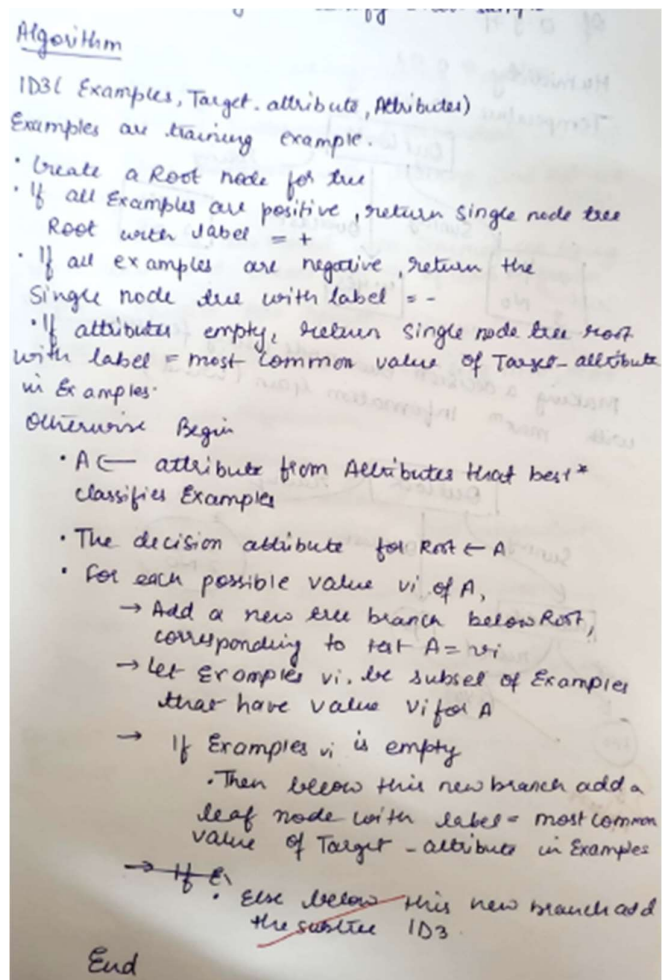
```
:   Country  Age   Salary Purchased  
0   France  44.0  72000.0     No  
1   Spain   27.0  48000.0    Yes  
2   Germany 30.0  54000.0     No  
3   Spain   38.0  61000.0     No  
4   Germany 40.0     NaN      Yes  
5   France  35.0  58000.0    Yes  
6   Spain   NaN   52000.0     No  
7   France  48.0  79000.0    Yes  
8   Germany 50.0  83000.0     No  
9   France  37.0  67000.0    Yes
```

```
: pd.get_dummies(df2)
```

```
:   Age   Salary Country_France Country_Germany Country_Spain Purchased_No Purchased_Yes  
0  44.0  72000.0        True        False       False      True      False  
1  27.0  48000.0       False       False       True      False      True  
2  30.0  54000.0       False       True       False      True      False  
3  38.0  61000.0       False       False       True      True      False  
4  40.0    NaN        False       True       False      False      True  
5  35.0  58000.0        True       False       False      False      True  
6  NaN   52000.0       False       False       True      True      False  
7  48.0  79000.0        True       False       False      False      True  
8  50.0  83000.0       False       True       False      True      False  
9  37.0  67000.0        True       False       False      False      True
```

4.

Decision Tree Algorithm



Code and output:

```

✓ [6] import pandas as pd
import numpy as np
import math
# Reading the dataset (Tennis-dataset)
data = pd.read_csv('PlayTennis.csv')

```

```

✓ [2] data.head()

```

	outlook	temp	humidity	windy	play
0	sunny	hot	high	False	no
1	sunny	hot	high	True	no
2	overcast	hot	high	False	yes
3	rainy	mild	high	False	yes
4	rainy	cool	normal	False	yes
5	rainy	cool	normal	True	no
6	overcast	cool	normal	True	yes
7	sunny	mild	high	False	no
8	sunny	cool	normal	False	yes
9	rainy	mild	normal	False	yes
10	sunny	mild	normal	True	yes
11	overcast	mild	high	True	yes
12	overcast	hot	normal	False	yes
13	rainy	mild	high	True	no

```

✓ [3] def find_entropy(data):
    """
        Returns the entropy of the class or features
        formula: - Σ P(X)logP(X)
    """
    entropy = 0
    for i in range(data.nunique()):
        x = data.value_counts()[i]/data.shape[0]
        entropy += (- x * math.log(x,2))
    return round(entropy,3)
def information_gain(data, data_):
    """
        Returns the information gain of the features
    """
    info = 0
    for i in range(data_.nunique()):
        df = data[data_ == data_.unique()[i]]
        w_avg = df.shape[0]/data.shape[0]
        entropy = find_entropy(df.play)
        x = w_avg * entropy
        info += x
    ig = find_entropy(data.play) - info
    return round(ig, 3)
def entropy_and_infogain(datax, feature):
    """
        Grouping features with the same class and computing their
        entropy and information gain for splitting
    """
    for i in range(data[feature].nunique()):
        df = datax[datax[feature]==data[feature].unique()[i]]
        if df.shape[0] < 1:
            continue
        display(df[[feature, 'play']].style.applymap(highlight)\n            .set_properties(subset=[feature, 'play'], **{'width': '80px'})\n            .set_table_styles([{'selector': 'th', 'props': [(['background-color', 'lightgray'],\n                ('border', '1px solid gray'),\n                ('font-weight', 'bold')]),\n                {'selector': 'td', 'props': [(['border', '1px solid gray'])]},\n                {'selector': 'tr:hover', 'props': [(['background-color', 'white'],\n                    ('border', '1.5px solid black')])]}]))\n
        print(f'Entropy of {feature} - {data[feature].unique()[i]} = {find_entropy(df.play)}')
        print(f'Information Gain for {feature} = {information_gain(datax, data[feature])}')

```

```
✓ [4] print(f'Entropy of the entire dataset: {find_entropy(data.play)}')  
Entropy of the entire dataset: 0.94
```

```
✓ [5] entropy_and_infogain(data, 'outlook')
```

	outlook	play
0	sunny	no
1	sunny	no
7	sunny	no
8	sunny	yes
10	sunny	yes

Entropy of outlook - sunny = 0.971

	outlook	play
2	overcast	yes
6	overcast	yes
11	overcast	yes
12	overcast	yes

Entropy of outlook - overcast = 0.0

	outlook	play
3	rainy	yes
4	rainy	yes
5	rainy	no
9	rainy	yes
13	rainy	no

Entropy of outlook - rainy = 0.971

Information Gain for outlook = 0.246

```
✓ [10] entropy_and_infogain(data, 'temp')
```

	temp	play
0	hot	no
1	hot	no
2	hot	yes
12	hot	yes

Entropy of temp - hot = 1.0

	temp	play
3	mild	yes
7	mild	no
9	mild	yes
10	mild	yes
11	mild	yes
13	mild	no

Entropy of temp - mild = 0.918

	temp	play
4	cool	yes
5	cool	no
6	cool	yes
8	cool	yes

Entropy of temp - cool = 0.811

Information Gain for temp = 0.029

```
▶ entropy_and_infogain(data, 'humidity')
```

	humidity	play
0	high	no
1	high	no
2	high	yes
3	high	yes
7	high	no
11	high	yes
13	high	no

Entropy of humidity - high = 0.985

	humidity	play
4	normal	yes
5	normal	no
6	normal	yes
8	normal	yes
9	normal	yes
10	normal	yes
12	normal	yes

Entropy of humidity - normal = 0.592

Information Gain for humidity = 0.151

```
✓ [12] entropy_and_infogain(data, 'windy')
```

	windy	play
0	False	no
2	False	yes
3	False	yes
4	False	yes
7	False	no
8	False	yes
9	False	yes
12	False	yes

Entropy of windy - False = 0.811

	windy	play
1	True	no
5	True	no
6	True	yes
10	True	yes
11	True	yes
13	True	no

Entropy of windy - True = 1.0

Information Gain for windy = 0.048

```

✓ 0s  sunny = data[data['outlook'] == 'sunny']
sunny.style.applymap(highlight)\n.set_properties(subset=data.columns, **{'width': '100px'})\n.set_table_styles([{'selector': 'th', 'props': [('background-color', 'lightgray'), ('border', '1px solid gray'), ('font-weight', 'bold')]},\n{'selector': 'tr:hover', 'props': [('background-color', 'white'), ('border', '1.5px solid black')]}])

```

	outlook	temp	humidity	windy	play
0	sunny	hot	high	False	no
1	sunny	hot	high	True	no
7	sunny	mild	high	False	no
8	sunny	cool	normal	False	yes
10	sunny	mild	normal	True	yes

```

✓ 0s  [14] print(f'Entropy of the Sunny dataset: {find_entropy(sunny.play)}')

```

Entropy of the Sunny dataset: 0.971

```

✓ 0s  [15] entropy_and_infogain(sunny, 'temp')

```

	temp	play
0	hot	no
1	hot	no

Entropy of temp - hot = 0.0

	temp	play
7	mild	no
10	mild	yes

Entropy of temp - mild = 1.0

	temp	play
8	cool	yes

Entropy of temp - cool = 0.0
Information Gain for temp = 0.571

```

✓ 0s  [16] entropy_and_infogain(sunny, 'humidity')

```

	humidity	play
0	high	no
1	high	no
7	high	no

Entropy of humidity - high = 0.0

	humidity	play
8	normal	yes
10	normal	yes

Entropy of humidity - normal = 0.0
Information Gain for humidity = 0.971

```

✓ 0s  [17] entropy_and_infogain(sunny, 'windy')

```

	windy	play
0	False	no
7	False	no
8	False	yes

Entropy of windy - False = 0.918

	windy	play
1	True	no
10	True	yes

Entropy of windy - True = 1.0
Information Gain for windy = 0.02

```

✓ [18] rainy = data[data['outlook'] == 'rainy']
rainy.style.applymap(highlight)\n    .set_properties(subset=data.columns, **{'width': '100px'})\n    .set_table_styles([{('selector': 'th', 'props': [(('background-color', 'lightgray'), ('border', '1px solid gray'),\n        ('font-weight', 'bold'))],\n        ('selector': 'tr:hover', 'props': [(('background-color', 'white'), ('border', '1.5px solid black'))])}])

```

	outlook	temp	humidity	windy	play
3	rainy	mild	high	False	yes
4	rainy	cool	normal	False	yes
5	rainy	cool	normal	True	no
9	rainy	mild	normal	False	yes
13	rainy	mild	high	True	no

```

✓ [19] print(f'Entropy of the Rainy dataset: {find_entropy(rainy.play)}')

```

Entropy of the Rainy dataset: 0.971

```

✓ [20] entropy_and_infogain(rainy, 'temp')

```

	temp	play
3	mild	yes
9	mild	yes
13	mild	no

Entropy of temp - mild = 0.918

	temp	play
4	cool	yes
5	cool	no

Entropy of temp - cool = 1.0

Information Gain for temp = 0.02

```

✓ [21] entropy_and_infogain(rainy, 'humidity')

```

	humidity	play
3	high	yes
13	high	no

Entropy of humidity - high = 1.0

	humidity	play
4	normal	yes
5	normal	no
9	normal	yes

Entropy of humidity - normal = 0.918

Information Gain for humidity = 0.02

```

✓ [22] entropy_and_infogain(rainy, 'windy')

```

	windy	play
3	False	yes
4	False	yes
9	False	yes

Entropy of windy - False = 0.0

	windy	play
5	True	no
13	True	no

Entropy of windy - True = 0.0

Information Gain for windy = 0.971

5. Build KNN Classification model for a given dataset.:

KNN algorithm

Step1: Select optimal value of K

Step2: calculate Euclidean distance

Step3: find nearest neighbor

Step4: vote for classification (majority)

voting) distance (x, x_i) = $\sqrt{\sum_j d_{ij}^2} = \sqrt{(x_j - x_{ij})^2}$

Code and output:

```
✓  [1] import numpy as np
    import pandas as pd
    import matplotlib.pyplot as plt
    import plotly.express as px
    import seaborn as sns

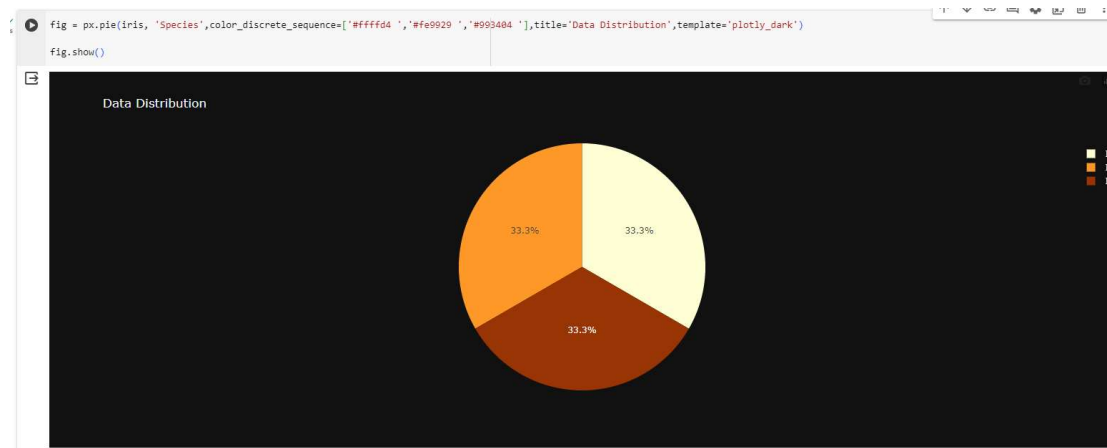
✓  [3] iris = pd.read_csv("Iris.csv") #Load Data
    iris.drop('Id',inplace=True,axis=1) #Drop Id column

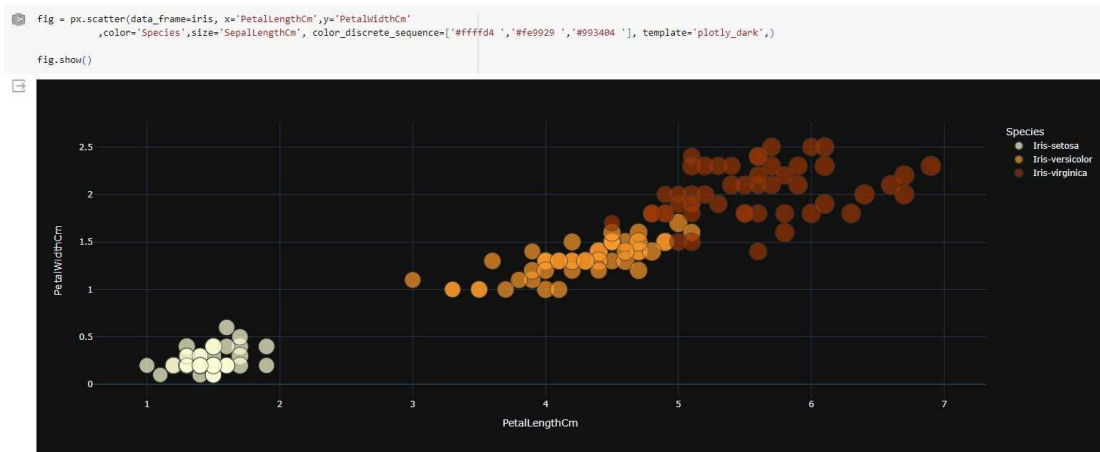
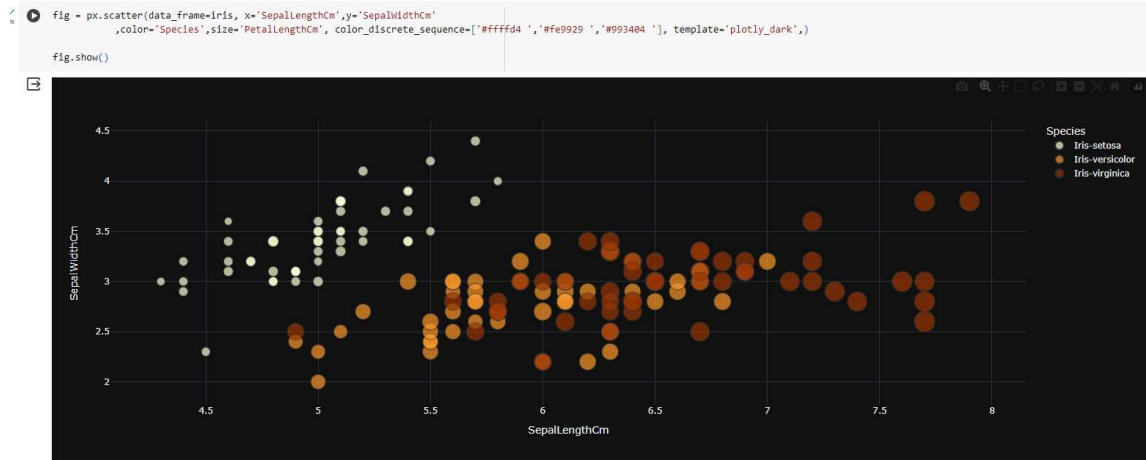
✓  [4] iris.head()
```

	SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm	Species	
0	5.1	3.5	1.4	0.2	Iris-setosa	
1	4.9	3.0	1.4	0.2	Iris-setosa	
2	4.7	3.2	1.3	0.2	Iris-setosa	
3	4.6	3.1	1.5	0.2	Iris-setosa	
4	5.0	3.6	1.4	0.2	Iris-setosa	

Next steps: View recommended plots

```
✓  [5] X = iris.iloc[:, :-1] #Set our training data
    y = iris.iloc[:, -1] #Set training labels
```





```
ls  class KNN:  
'''  
    K-Nearest Neighbors (KNN) classification algorithm  
  
    Parameters:  
    -----  
    n_neighbors : int, optional (default=5)  
        Number of neighbors to use in the majority vote.  
  
    Methods:  
    -----  
    fit(X_train, y_train):  
        Stores the values of X_train and y_train.  
  
    predict(X):  
        Predicts the class labels for each example in X.  
  
    """  
    def __init__(self, n_neighbors=5):  
        self.n_neighbors = n_neighbors  
  
    def euclidean_distance(self, x1, x2):  
        """  
            Calculate the Euclidean distance between two data points.  
  
        Parameters:  
        -----  
        x1 : numpy.ndarray, shape (n_features,)  
            A data point in the dataset.  
  
        x2 : numpy.ndarray, shape (n_features,)  
            A data point in the dataset.  
  
        Returns:  
        -----  
        distance : float  
            The Euclidean distance between x1 and x2.  
        """  
        return np.linalg.norm(x1 - x2)
```

```
def fit(self, X_train, y_train):
    """
    Stores the values of X_train and y_train.

    Parameters:
    -----
    X_train : numpy.ndarray, shape (n_samples, n_features)
        The training dataset.

    y_train : numpy.ndarray, shape (n_samples,)
        The target labels.
    """
    self.X_train = X_train
    self.y_train = y_train

def predict(self, X):
    """
    Predicts the class labels for each example in X.

    Parameters:
    -----
    X : numpy.ndarray, shape (n_samples, n_features)
        The test dataset.

    Returns:
    -----
    predictions : numpy.ndarray, shape (n_samples,)
        The predicted class labels for each example in X.
    """
    # Create empty array to store the predictions
    predictions = []
    # Loop over X examples
    for x in X:
        # Get prediction using the prediction helper function
        prediction = self._predict(x)
        # Append the prediction to the predictions list
        predictions.append(prediction)
    return np.array(predictions)
```

```

def _predict(self, x):
    """
    Predicts the class label for a single example.

    Parameters:
    -----
    x : numpy.ndarray, shape (n_features,)
        A data point in the test dataset.

    Returns:
    -----
    most_occuring_value : int
        The predicted class label for x.
    """
    # Create empty array to store distances
    distances = []
    # Loop over all training examples and compute the distance between x and all the training examples
    for x_train in self.X_train:
        distance = self.euclidean_distance(x, x_train)
        distances.append(distance)
    distances = np.array(distances)

    # Sort by ascendingly distance and return indices of the given n neighbours
    n_neighbors_idxs = np.argsort(distances)[: self.n_neighbors]

    # Get labels of n-neighbour indexes
    labels = self.y_train[n_neighbors_idxs]
    labels = list(labels)
    # Get the most frequent class in the array
    most_occuring_value = max(labels, key=labels.count)
    return most_occuring_value

```

```

[1] def train_test_split(X, y, random_state=42, test_size=0.2):
    """
    Splits the data into training and testing sets.

    Parameters:
    X (numpy.ndarray): Features array of shape (n_samples, n_features).
    y (numpy.ndarray): Target array of shape (n_samples,).
    random_state (int): Seed for the random number generator. Default is 42.
    test_size (float): Proportion of samples to include in the test set. Default is 0.2.

    Returns:
    Tuple[numpy.ndarray]: A tuple containing X_train, X_test, y_train, y_test.
    """
    # Get number of samples
    n_samples = X.shape[0]

    # Set the seed for the random number generator
    np.random.seed(random_state)

    # Shuffle the indices
    shuffled_indices = np.random.permutation(np.arange(n_samples))

    # Determine the size of the test set
    test_size = int(n_samples * test_size)

    # Split the indices into test and train
    test_indices = shuffled_indices[:test_size]
    train_indices = shuffled_indices[test_size:]

    # Split the features and target arrays into test and train
    X_train, X_test = X[train_indices], X[test_indices]
    y_train, y_test = y[train_indices], y[test_indices]

    return X_train, X_test, y_train, y_test

```

```
[12] X_train, X_test, y_train, y_test = train_test_split(X.values, y.values, test_size = 0.2, random_state=42) #
```

```

[13] model = KNN(7)
model.fit(X_train, y_train)

[14] def compute_accuracy(y_true, y_pred):
    """
    Computes the accuracy of a classification model.

    Parameters:
    y_true (numpy array): A numpy array of true labels for each data point.
    y_pred (numpy array): A numpy array of predicted labels for each data point.

    Returns:
    float: The accuracy of the model, expressed as a percentage.
    """
    y_true = y_true.flatten()
    total_samples = len(y_true)
    correct_predictions = np.sum(y_true == y_pred)
    return (correct_predictions / total_samples)

[15] predictions = model.predict(X_test)
accuracy = compute_accuracy(y_test, predictions)
print(f" our model got accuracy score of : {accuracy}")

our model got accuracy score of : 0.9666666666666666

[16] from sklearn.neighbors import KNeighborsClassifier
skmodel = KNeighborsClassifier(n_neighbors=7)
skmodel.fit(X_train, y_train)

    KNeighborsClassifier
KNeighborsClassifier(n_neighbors=7)

[17] sk_predictions = skmodel.predict(X_test)
sk_accuracy = compute_accuracy(y_test, sk_predictions)
print(f" sklearn-model got accuracy score of : {sk_accuracy}")

sklearn-model got accuracy score of : 0.9666666666666667
```

6.

Linear Regression

→ Linear Regression

Training phase:

1. Input: Training data with one predictor (x) and one target variable (y)

2. compute mean

3. estimate coefficients

$$\beta = \frac{\text{cov}(x, y)}{\text{var}(x)}$$

$$\beta_0 = \bar{y} - \beta \bar{x}$$

Prediction phase:

1. Input: New x_{test}

2. Compute prediction: use $\hat{y}_{\text{pred}} = \beta_0 + \beta x_{\text{test}}$ to predict \hat{y}_{pred}

Output:

~~model.predict([123])~~

→ array([107.82727115])

Code and output:

```
[1] import math
import numpy as np
import pandas as pd
import plotly.express as px
import pickle

[2] # Load the training and test datasets
train_data = pd.read_csv('train.csv')
test_data = pd.read_csv('test.csv')

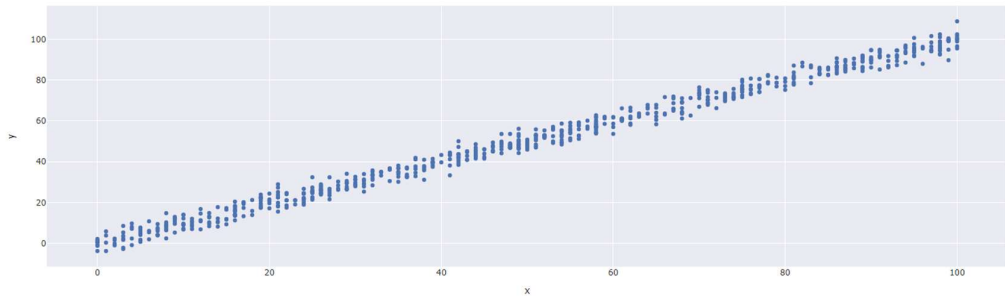
# Remove rows with missing values
train_data = train_data.dropna()
test_data = test_data.dropna()

[3] train_data.head()
```

	x	y
0	24.0	21.549452
1	50.0	47.464463
2	15.0	17.218656
3	38.0	36.586398
4	87.0	87.288984

Next steps: [View recommended plots](#)

```
[4] px.scatter(x=train_data['x'], y=train_data['y'], template='seaborn')
```



```
[5] # Set training data and target
X_train = train_data['x'].values
y_train = train_data['y'].values

# Set testing data and target
X_test = test_data['x'].values
y_test = test_data['y'].values
```

```

[6] """
Standardizes the input data using mean and standard deviation.

Parameters:
    X_train (numpy.ndarray): Training data.
    X_test (numpy.ndarray): Testing data.

Returns:
    Tuple of standardized training and testing data.
"""

# Calculate the mean and standard deviation using the training data
mean = np.mean(X_train, axis=0)
std = np.std(X_train, axis=0)

# Standardize the data
X_train = (X_train - mean) / std
X_test = (X_test - mean) / std

return X_train, X_test

X_train, X_test = standardize_data(X_train, X_test)

[7] X_train = np.expand_dims(X_train, axis=-1)
X_test = np.expand_dims(X_test, axis=-1)

[8] class LinearRegression:
    """
    Linear Regression Model with Gradient Descent

    Linear regression is a supervised machine learning algorithm used for modeling the relationship
    between a dependent variable (target) and one or more independent variables (features) by fitting
    a linear equation to the observed data.

    This class implements a linear regression model using gradient descent optimization for training.
    It provides methods for model initialization, training, prediction, and model persistence.

    Parameters:
        learning_rate (float): The learning rate used in gradient descent.
        convergence_tol (float, optional): The tolerance for convergence (stopping criterion). Defaults to 1e-6.

    Attributes:

        ATTRIBUTES:
            W (numpy.ndarray): Coefficients (weights) for the linear regression model.
            b (float): Intercept (bias) for the linear regression model.

    Methods:

        initialize_parameters(n_features): Initialize model parameters.
        forward(X): Compute the forward pass of the linear regression model.
        compute_cost(predictions): Compute the mean squared error cost.
        backward(predictions): Compute gradients for model parameters.
        fit(X, y, iterations, plot_cost=True): Fit the linear regression model to training data.
        predict(X): Predict target values for new input data.
        save_model(filename=None): Save the trained model to a file using pickle.
        load_model(filename): Load a trained model from a file using pickle.

    Examples:
        >>> from linear_regression import LinearRegression
        >>> model = LinearRegression(learning_rate=0.01)
        >>> model.fit(X_train, y_train, iterations=1000)
        >>> predictions = model.predict(X_test)
        """

        def __init__(self, learning_rate, convergence_tol=1e-6):
            self.learning_rate = learning_rate
            self.convergence_tol = convergence_tol
            self.W = None
            self.b = None

        def initialize_parameters(self, n_features):
            """
            Initialize model parameters.

            Parameters:
                n_features (int): The number of features in the input data.
            """
            self.W = np.random.randn(n_features) * 0.01
            self.b = 0

        def forward(self, X):
            """
            Compute the forward pass of the linear regression model.

            Parameters:
                X (numpy.ndarray): Input data of shape (m, n_features).
            """

```

```
[8]     Returns:
          numpy.ndarray: Predictions of shape (m,).
        """
        return np.dot(X, self.W) + self.b

    def compute_cost(self, predictions):
        """
        Compute the mean squared error cost.

        Parameters:
            predictions (numpy.ndarray): Predictions of shape (m,).

        Returns:
            float: Mean squared error cost.
        """
        m = len(predictions)
        cost = np.sum(np.square(predictions - self.y)) / (2 * m)
        return cost

    def backward(self, predictions):
        """
        Compute gradients for model parameters.

        Parameters:
            predictions (numpy.ndarray): Predictions of shape (m,).

        Updates:
            numpy.ndarray: Gradient of W.
            float: Gradient of b.
        """
        m = len(predictions)
        self.dW = np.dot(predictions - self.y, self.X) / m
        self.db = np.sum(predictions - self.y) / m
def fit(self, X, y, iterations, plot_cost=True):
    """
    Fit the linear regression model to the training data.

    Parameters:
        X (numpy.ndarray): Training input data of shape (m, n_features).
        y (numpy.ndarray): Training labels of shape (m,).
        iterations (int): The number of iterations for gradient descent.
        plot_cost (bool, optional): Whether to plot the cost during training. Defaults to True.
    
```

```
[8]
    Raises:
        AssertionError: If input data and labels are not NumPy arrays or have mismatched shapes.

    Plots:
        Plotly line chart showing cost vs. iteration (if plot_cost is True).
    """
    assert isinstance(X, np.ndarray), "X must be a NumPy array"
    assert isinstance(y, np.ndarray), "y must be a NumPy array"
    assert X.shape[0] == y.shape[0], "X and y must have the same number of samples"
    assert iterations > 0, "Iterations must be greater than 0"

    self.X = X
    self.y = y
    self.initialize_parameters(X.shape[1])
    costs = []

    for i in range(iterations):
        predictions = self.forward(X)
        cost = self.compute_cost(predictions)
        self.backward(predictions)
        self.W -= self.learning_rate * self.dW
        self.b -= self.learning_rate * self.db
        costs.append(cost)

        if i % 100 == 0:
            print(f'Iteration: {i}, Cost: {cost}')

        if i > 0 and abs(costs[-1] - costs[-2]) < self.convergence_tol:
            print(f'Converged after {i} iterations.')
            break

    if plot_cost:
        fig = px.line(y=costs, title="Cost vs Iteration", template="plotly_dark")
        fig.update_layout(
            title_font_color="#41BEE9",
            xaxis=dict(color="#41BEE9", title="Iterations"),
            yaxis=dict(color="#41BEE9", title="Cost")
        )
        fig.show()
    
```

```

        fig.show()
    def predict(self, X):
        """
        Predict target values for new input data.

        Parameters:
            X (numpy.ndarray): Input data of shape (m, n_features).

        Returns:
            numpy.ndarray: Predicted target values of shape (m,).
        """
        return self.forward(X)

    def save_model(self, filename=None):
        """
        Save the trained model to a file using pickle.

        Parameters:
            filename (str): The name of the file to save the model to.
        """
        model_data = {
            'learning_rate': self.learning_rate,
            'convergence_tol': self.convergence_tol,
            'W': self.W,
            'b': self.b
        }

        with open(filename, 'wb') as file:
            pickle.dump(model_data, file)

    @classmethod
    def load_model(cls, filename):
        """
        Load a trained model from a file using pickle.

        Parameters:
            filename (str): The name of the file to load the model from.
        """
        with open(filename, 'rb') as file:
            pickle.dump(model_data, file)

    @classmethod
    def load_model(cls, filename):
        """
        Load a trained model from a file using pickle.

        Parameters:
            filename (str): The name of the file to load the model from.

        Returns:
            LinearRegression: An instance of the LinearRegression class with loaded parameters.
        """
        with open(filename, 'rb') as file:
            model_data = pickle.load(file)

        # Create a new instance of the class and initialize it with the loaded parameters
        loaded_model = cls(model_data['learning_rate'], model_data['convergence_tol'])
        loaded_model.W = model_data['W']
        loaded_model.b = model_data['b']

        return loaded_model

9] lr = LinearRegression(0.01)
lr.fit(X_train, y_train, 10000)

Iteration: 0, Cost: 1670.0184887161677
Iteration: 100, Cost: 227.15535101517312
Iteration: 200, Cost: 33.84101696145528
Iteration: 300, Cost: 7.9408253395546575
Iteration: 400, Cost: 4.4787260872934835
Iteration: 500, Cost: 4.005803317750673
Iteration: 600, Cost: 3.943513116253261
Iteration: 700, Cost: 3.9351674953098015
Iteration: 800, Cost: 3.9340493517293096
Converged after 863 iterations.

```

```

Iteration: 800, Cost: 3.9340493517293096
Converged after 803 Iterations.

Cost vs Iteration

[10] lr.save_model('model.pkl')

[11] model = LinearRegression.load_model("model.pkl")

[12] class RegressionMetrics:
    @staticmethod
    def mean_squared_error(y_true, y_pred):
        """
        Calculate the Mean Squared Error (MSE).

        Args:
            y_true (numpy.ndarray): The true target values.
            y_pred (numpy.ndarray): The predicted target values.

        Returns:
            float: The Mean Squared Error.
        """
        assert len(y_true) == len(y_pred), "Input arrays must have the same length."
        mse = np.mean((y_true - y_pred) ** 2)
        return mse

    @staticmethod
    def root_mean_squared_error(y_true, y_pred):
        """
        Calculate the Root Mean Squared Error (RMSE).

        Args:
            y_true (numpy.ndarray): The true target values.
            y_pred (numpy.ndarray): The predicted target values.

        Returns:
            float: The Root Mean Squared Error.
        """
        assert len(y_true) == len(y_pred), "Input arrays must have the same length."
        mse = RegressionMetrics.mean_squared_error(y_true, y_pred)
        rmse = np.sqrt(mse)
        return rmse

    @staticmethod
    def r_squared(y_true, y_pred):
        """
        Calculate the R-squared (R^2) coefficient of determination.

        Args:
            y_true (numpy.ndarray): The true target values.
            y_pred (numpy.ndarray): The predicted target values.
        """

```

```
[12]     @staticmethod
[12]     def r_squared(y_true, y_pred):
[12]         """
[12]             Calculate the R-squared ( $R^2$ ) coefficient of determination.
[12]
[12]             Args:
[12]                 y_true (numpy.ndarray): The true target values.
[12]                 y_pred (numpy.ndarray): The predicted target values.
[12]
[12]             Returns:
[12]                 float: The R-squared ( $R^2$ ) value.
[12]             """
[12]             assert len(y_true) == len(y_pred), "Input arrays must have the same length."
[12]             mean_y = np.mean(y_true)
[12]             ss_total = np.sum((y_true - mean_y) ** 2)
[12]             ss_residual = np.sum((y_true - y_pred) ** 2)
[12]             r2 = 1 - (ss_residual / ss_total)
[12]             return r2
```

```
[13] y_pred = model.predict(X_test)
[13] mse_value = RegressionMetrics.mean_squared_error(y_test, y_pred)
[13] rmse_value = RegressionMetrics.root_mean_squared_error(y_test, y_pred)
[13] r_squared_value = RegressionMetrics.r_squared(y_test, y_pred)

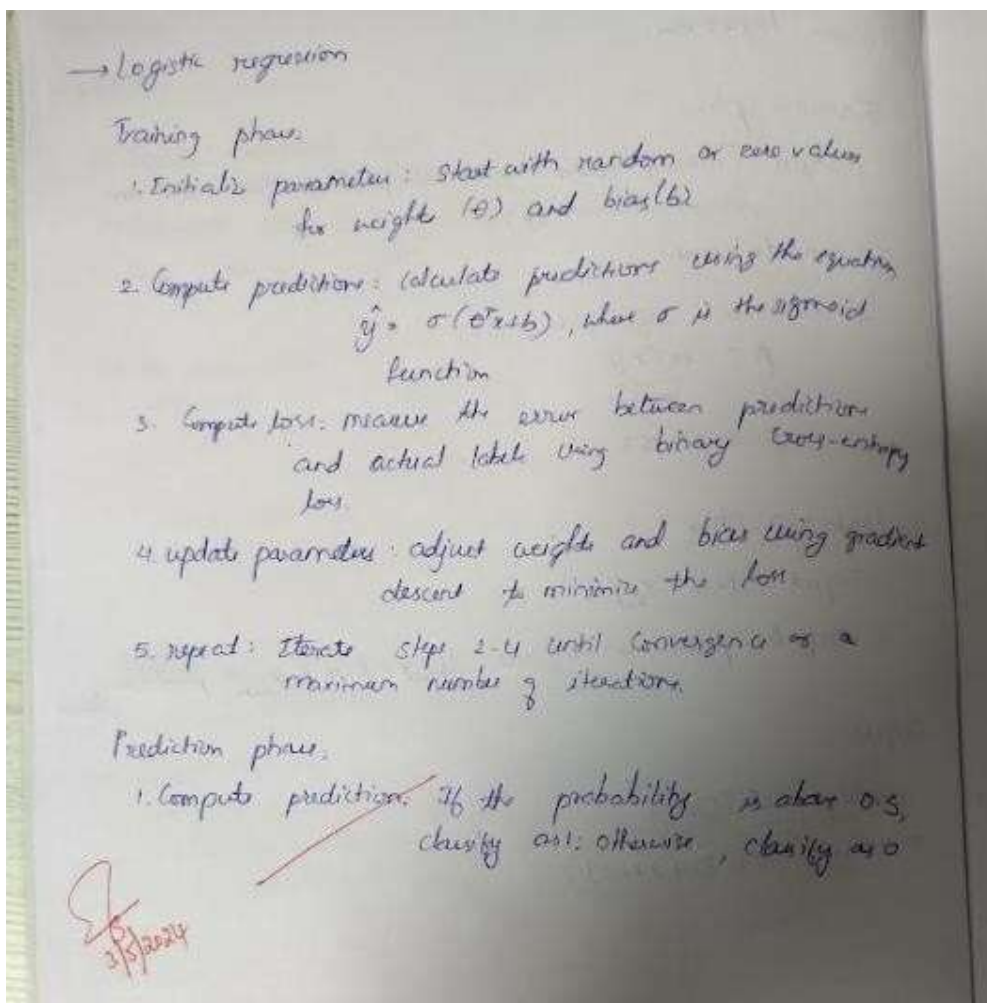
print(f"Mean Squared Error (MSE): {mse_value}")
print(f"Root Mean Squared Error (RMSE): {rmse_value}")
print(f"R-squared (Coefficient of Determination): {r_squared_value}")

Mean Squared Error (MSE): 9.44266965025894
Root Mean Squared Error (RMSE): 3.07289271701095
R-squared (Coefficient of Determination): 0.9887898724670081
```

```
▶ model.predict([[2]])
array([107.82727115])
```

7.

Logistic Regression



```

✓ [3] import math
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
import plotly.express as px
import pprint
import pickle

✓ [4] df = pd.read_csv('breast-cancer.csv')

✓ [5] df.head()

   id diagnosis radius_mean texture_mean perimeter_mean area_mean smoothness_mean compactness_mean concavity_mean concave points_mean ... radius_worst texture_worst perimeter_worst are
0  842302      M     17.99    10.38    122.80   1001.0    0.11840    0.27760    0.3001    0.14710 ...     25.38    17.33    184.60
1  842517      M     20.57    17.77    132.90   1326.0    0.08474    0.07864    0.0869    0.07017 ...     24.99    23.41    158.80
2  84300903    M     19.69    21.25    130.00   1203.0    0.10960    0.15990    0.1974    0.12790 ...     23.57    25.53    152.50
3  84348301    M     11.42    20.38    77.58    386.1     0.14250    0.28390    0.2414    0.10520 ...     14.91    26.50    98.87
4  84358402    M     20.29    14.34    135.10   1297.0    0.10030    0.13280    0.1980    0.10430 ...     22.54    16.67    152.20
5 rows × 32 columns

```

```

✓ [6] df.drop('id', axis=1, inplace=True) #drop redundant columns

✓ [7] df['diagnosis'] = (df['diagnosis'] == 'M').astype(int) #encode the label into 1/0

✓ [8] corr = df.corr()

```



```
[12]: # Get the absolute value of the correlation
cor_target = abs(corr["diagnosis"])

# Select highly correlated features (threshold = 0.2)
relevant_features = cor_target[cor_target>0.2]

# Collect the names of the features
names = [index for index, value in relevant_features.items()]

# Drop the target variable from the results
names.remove('diagnosis')

# Display the results
pprint.pprint(names)

[ 'radius_mean',
  'texture_mean',
  'perimeter_mean',
  'area_mean',
  'smoothness_mean',
  'compactness_mean',
  'concavity_mean',
  'concave points_mean',
  'symmetry_mean',
  'radius_se',
  'perimeter_se',
  'area_se',
  'compactness_se',
  'concavity_se',
  'concave points_se',
  'radius_worst',
  'texture_worst',
  'perimeter_worst',
  'area_worst',
  'smoothness_worst',
  'compactness_worst',
  'concavity_worst',
  'concave points_worst',
  'symmetry_worst',
  'fractal_dimension_worst']
```

```
[13] X = df[names].values
y = df['diagnosis'].values

[14] def train_test_split(X, y, random_state=42, test_size=0.2):
    """
    Splits the data into training and testing sets.

    Parameters:
        X (numpy.ndarray): Features array of shape (n_samples, n_features).
        y (numpy.ndarray): Target array of shape (n_samples,).
        random_state (int): Seed for the random number generator. Default is 42.
        test_size (float): Proportion of samples to include in the test set. Default is 0.2.

    Returns:
        Tuple[numpy.ndarray]: A tuple containing X_train, X_test, y_train, y_test.
    """
    # Get number of samples
    n_samples = X.shape[0]

    # Set the seed for the random number generator
    np.random.seed(random_state)

    # Shuffle the indices
    shuffled_indices = np.random.permutation(np.arange(n_samples))

    # Determine the size of the test set
    test_size = int(n_samples * test_size)

    # Split the indices into test and train
    test_indices = shuffled_indices[:test_size]
    train_indices = shuffled_indices[test_size:]

    # Split the features and target arrays into test and train
    X_train, X_test = X[train_indices], X[test_indices]
    y_train, y_test = y[train_indices], y[test_indices]

    return X_train, X_test, y_train, y_test
```

```

' [15] X_train, X_test, y_train, y_test = train_test_split(X,y)

' [16] def standardize_data(X_train, X_test):
    """
    Standardizes the input data using mean and standard deviation.

    Parameters:
        X_train (numpy.ndarray): Training data.
        X_test (numpy.ndarray): Testing data.

    Returns:
        Tuple of standardized training and testing data.
    """
    # Calculate the mean and standard deviation using the training data
    mean = np.mean(X_train, axis=0)
    std = np.std(X_train, axis=0)

    # Standardize the data
    X_train = (X_train - mean) / std
    X_test = (X_test - mean) / std

    return X_train, X_test

X_train, X_test = standardize_data(X_train, X_test)

' [17] def sigmoid(z):
    """
    Compute the sigmoid function for a given input.

    The sigmoid function is a mathematical function used in logistic regression and neural networks
    to map any real-valued number to a value between 0 and 1.

    Parameters:
        z (float or numpy.ndarray): The input value(s) for which to compute the sigmoid.

    Returns:
        float or numpy.ndarray: The sigmoid of the input value(s).

    Example:
        >>> sigmoid(0)
        0.5
    """
    # Compute the sigmoid function using the formula: 1 / (1 + e^(-z)).
    sigmoid_result = 1 / (1 + np.exp(-z))

    # Return the computed sigmoid value.
    return sigmoid_result

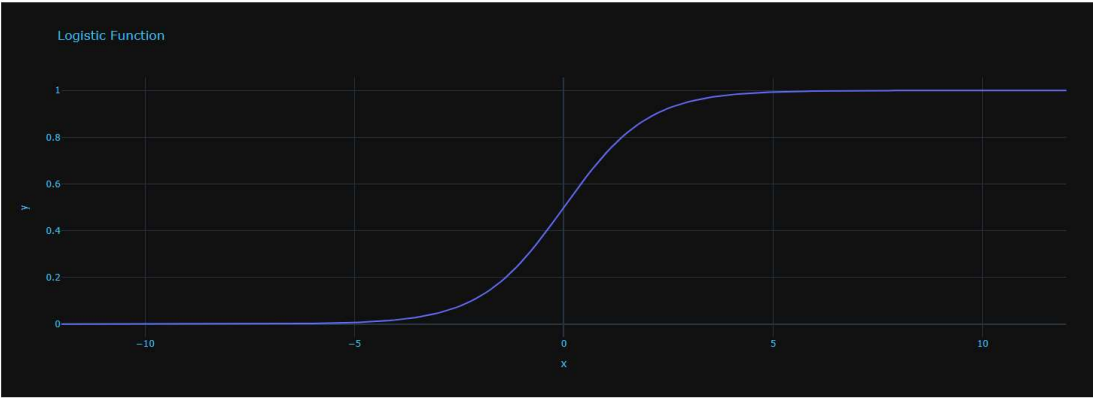
```

```

[1] z = np.linspace(-12, 12, 200)

fig = px.line(x=z, y=sigmoid(z), title='Logistic Function', template="plotly_dark")
fig.update_layout(
    title_font_color="#41BEE9",
    xaxis=dict(color="#41BEE9"),
    yaxis=dict(color="#41BEE9")
)
fig.show()

```



```
[19] class LogisticRegression:
    """
    Logistic Regression model.

    Parameters:
        learning_rate (float): Learning rate for the model.

    Methods:
        initialize_parameter(): Initializes the parameters of the model.
        sigmoid(z): Computes the sigmoid activation function for given input z.
        forward(X): Computes forward propagation for given input X.
    """

    def __init__(self, learning_rate=0.0001):
        np.random.seed(1)
        self.learning_rate = learning_rate
```

```
    def initialize_parameter(self):
        """
        Initializes the parameters of the model.
        """
        self.W = np.zeros(self.X.shape[1])
        self.b = 0.0

    def forward(self, X):
        """
        Computes forward propagation for given input X.

        Parameters:
            X (numpy.ndarray): Input array.

        Returns:
            numpy.ndarray: Output array.
        """
        Z = np.matmul(X, self.W) + self.b
        A = sigmoid(Z)
        return A

    def compute_cost(self, predictions):
        """
        Computes the cost function for given predictions.

        Parameters:
            predictions (numpy.ndarray): Predictions of the model.

        Returns:
            float: Cost of the model.
        """
        m = self.X.shape[0] # number of training examples
        # compute the cost
        cost = np.sum((-np.log(predictions + 1e-8) * self.y) + (-np.log(1 - predictions + 1e-8)) * (
            1 - self.y)) # we are adding small value epsilon to avoid log of 0
```

```
[19]     """
    # number of training examples
    m = self.X.shape[0]
    # compute the cost
    cost = np.sum((-np.log(predictions + 1e-8) * self.y) + (-np.log(1 - predictions + 1e-8)) * (1 - self.y))
    # we are adding small value epsilon to avoid log of 0
    cost = cost / m
    return cost
def compute_gradient(self, predictions):
    """
    Computes the gradients for the model using given predictions.

    Parameters:
        predictions (numpy.ndarray): Predictions of the model.
    """
    # get training shape
    m = self.X.shape[0]

    # compute gradients
    self.dW = np.matmul(self.X.T, (predictions - self.y))
    self.dW = np.array([np.mean(grad) for grad in self.dW])

    self.db = np.sum(np.subtract(predictions, self.y))

    # scale gradients
    self.dW = self.dW * 1 / m
    self.db = self.db * 1 / m

def fit(self, X, y, iterations, plot_cost=True):
    """
    Trains the model on given input X and labels y for specified iterations.

    Parameters:
        X (numpy.ndarray): Input features array of shape (n_samples, n )
        y (numpy.ndarray): Labels array of shape (n_samples, 1)
        iterations (int): Number of iterations for training.
        plot_cost (bool): Whether to plot cost over iterations or not.

    Returns:
        None.
    """
    self.X = X
```

```

19]     self.X = X
        self.y = y

        self.initialize_parameter()

        costs = []
        for i in range(iterations):
            # forward propagation
            predictions = self.forward(self.X)

            # compute cost
            cost = self.compute_cost(predictions)
            costs.append(cost)

            # compute gradients
            self.compute_gradient(predictions)

            # update parameters
            self.W = self.W - self.learning_rate * self.dW
            self.b = self.b - self.learning_rate * self.db

            # print cost every 100 iterations
            if i % 1000 == 0:
                print("Cost after iteration {}: {}".format(i, cost))

        if plot_cost:
            fig = px.line(y=costs,title="Cost vs Iteration",template="plotly_dark")
            fig.update_layout(
                title_font_color="#41BEE9",
                xaxis=dict(color="#41BEE9",title="Iterations"),
                yaxis=dict(color="#41BEE9",title="cost")
            )
            fig.show()
    def predict(self, X):
        """
        Predicts the labels for given input X.

        Parameters:
        X (numpy.ndarray): Input features array.

        Returns:
        numpy.ndarray: Predicted labels.
        """

```

```
[19]     predictions = self.forward(X)
             return np.round(predictions)

     def save_model(self, filename=None):
        """
        Save the trained model to a file using pickle.

        Parameters:
            filename (str): The name of the file to save the model to.
        """
        model_data = {
            'learning_rate': self.learning_rate,
            'W': self.W,
            'b': self.b
        }

        with open(filename, 'wb') as file:
            pickle.dump(model_data, file)

    @classmethod
    def load_model(cls, filename):
        """
        Load a trained model from a file using pickle.

        Parameters:
            filename (str): The name of the file to load the model from.

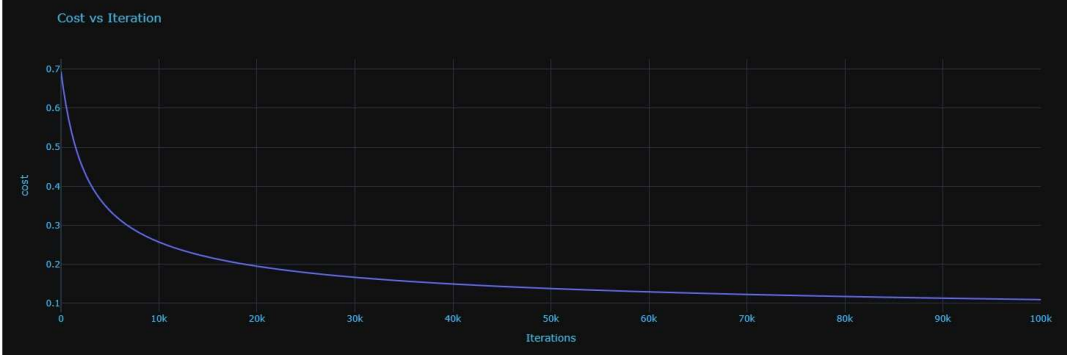
        Returns:
            LogisticRegression: An instance of the LogisticRegression class with loaded parameters.
        """
        with open(filename, 'rb') as file:
            model_data = pickle.load(file)

        # Create a new instance of the class and initialize it with the loaded parameters
        loaded_model = cls(model_data['learning_rate'])
        loaded_model.W = model_data['W']
        loaded_model.b = model_data['b']

        return loaded_model
```

```
[⑩] lg = LogisticRegression()
lg.fit(X_train, y_train, 100000)
```

Cost after iteration 0: 0.6931471606598454
Cost after iteration 10000: 0.2570770370558246
Cost after iteration 20000: 0.10520178673689726
Cost after iteration 30000: 0.16685820756163852
Cost after iteration 40000: 0.14978939548676498
Cost after iteration 50000: 0.1381876134031554
Cost after iteration 60000: 0.1296814121248933
Cost after iteration 70000: 0.123114403998813
Cost after iteration 80000: 0.11785163788790082
Cost after iteration 90000: 0.113513771386002



```

[22] lg.save_model("model.pkl")

[23] class ClassificationMetrics:
    @staticmethod
    def accuracy(y_true, y_pred):
        """
        Computes the accuracy of a classification model.

        Parameters:
        y_true (numpy array): A numpy array of true labels for each data point.
        y_pred (numpy array): A numpy array of predicted labels for each data point.

        Returns:
        float: The accuracy of the model, expressed as a percentage.
        """
        y_true = y_true.flatten()
        total_samples = len(y_true)
        correct_predictions = np.sum(y_true == y_pred)
        return (correct_predictions / total_samples)

    @staticmethod
    def precision(y_true, y_pred):
        """
        Computes the precision of a classification model.

        Parameters:
        y_true (numpy array): A numpy array of true labels for each data point.
        y_pred (numpy array): A numpy array of predicted labels for each data point.

        Returns:
        float: The precision of the model, which measures the proportion of true positive predictions
        out of all positive predictions made by the model.
        """
        true_positives = np.sum((y_true == 1) & (y_pred == 1))
        false_positives = np.sum((y_true == 0) & (y_pred == 1))
        return true_positives / (true_positives + false_positives)

```

```

[23]     @staticmethod
    def recall(y_true, y_pred):
        """
        Computes the recall (sensitivity) of a classification model.

        Parameters:
        y_true (numpy array): A numpy array of true labels for each data point.
        y_pred (numpy array): A numpy array of predicted labels for each data point.

        Returns:
        float: The recall of the model, which measures the proportion of true positive predictions
        out of all actual positive instances in the dataset.
        """
        true_positives = np.sum((y_true == 1) & (y_pred == 1))
        false_negatives = np.sum((y_true == 1) & (y_pred == 0))
        return true_positives / (true_positives + false_negatives)

    @staticmethod
    def f1_score(y_true, y_pred):
        """
        Computes the F1-score of a classification model.

        Parameters:
        y_true (numpy array): A numpy array of true labels for each data point.
        y_pred (numpy array): A numpy array of predicted labels for each data point.

        Returns:
        float: The F1-score of the model, which is the harmonic mean of precision and recall.
        """
        precision_value = ClassificationMetrics.precision(y_true, y_pred)
        recall_value = ClassificationMetrics.recall(y_true, y_pred)
        return 2 * (precision_value * recall_value) / (precision_value + recall_value)

```

```
[24] model = LogisticRegression.load_model("model.pkl")
```

```
✓ [24] model = LogisticRegression.load_model("model.pkl")  
0s  
✓ [25] y_pred = model.predict(X_test)  
accuracy = ClassificationMetrics.accuracy(y_test, y_pred)  
precision = ClassificationMetrics.precision(y_test, y_pred)  
recall = ClassificationMetrics.recall(y_test, y_pred)  
f1_score = ClassificationMetrics.f1_score(y_test, y_pred)  
  
print(f"Accuracy: {accuracy:.2%}")  
print(f"Precision: {precision:.2%}")  
print(f"Recall: {recall:.2%}")  
print(f"F1-Score: {f1_score:.2%}")  
  
Accuracy: 98.23%  
Precision: 100.00%  
Recall: 95.24%  
F1-Score: 97.56%
```

8. Build Support vector machine model for a given dataset

Algorithm:

SVM

- ① define kernel function
Eg. $K(x_1, x_2) = x_1 \cdot x_2$
- ② solve quadratic programming
- ③ compute weight and bias
- ④ identify support vectors
- ⑤ Make predictions

Output

```
Model = SVC()  
model.fit(X_train, y_train)  
predictions = model.predict(X_test)  
accuracy = (y_test, predictions)
```

10.9823088

arrays.

```

✓ [1] from google.colab import drive
drive.mount('/content/drive')

↳ Mounted at /content/drive

✓ [2] import seaborn as sns
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import plotly.express as px

✓ [3] df = pd.read_csv('/content/drive/MyDrive/breast-cancer.csv')
df.head()

→ id diagnosis radius_mean texture_mean perimeter_mean area_mean smoothness_mean compactness_mean concavity_mean concave
   0 842302 M 17.99 10.38 122.80 1001.0 0.11840 0.27760 0.3001 0.14710 ...
   1 842517 M 20.57 17.77 132.90 1326.0 0.08474 0.07864 0.0869 0.07017 ...
   2 84300903 M 19.69 21.25 130.00 1203.0 0.10960 0.15990 0.1974 0.12790 ...
   3 84348301 M 11.42 20.38 77.58 386.1 0.14250 0.28390 0.2414 0.10520 ...
   4 84358402 M 20.29 14.34 135.10 1297.0 0.10030 0.13280 0.1980 0.10430 ...

5 rows × 32 columns

```



```

✓ [5] df.drop('id', axis=1, inplace=True) #drop redundant columns

✓ [6] df.describe().T

→ count mean std min 25% 50% 75% max
  radius_mean 569.0 14.127292 3.524049 6.981000 11.700000 13.370000 15.780000 28.11000
  texture_mean 569.0 19.289649 4.301036 9.710000 16.170000 18.840000 21.800000 39.28000
  perimeter_mean 569.0 91.969033 24.298981 43.790000 75.170000 86.240000 104.100000 188.50000
  area_mean 569.0 654.889104 351.914129 143.500000 420.300000 551.100000 782.700000 2501.00000
  smoothness_mean 569.0 0.096360 0.014064 0.052630 0.086370 0.095870 0.105300 0.16340
  compactness_mean 569.0 0.104341 0.052813 0.019380 0.064920 0.092630 0.130400 0.34540
  concavity_mean 569.0 0.088799 0.079720 0.000000 0.029560 0.061540 0.130700 0.42680
  concave points_mean 569.0 0.048919 0.038803 0.000000 0.020310 0.033500 0.074000 0.20120
  symmetry_mean 569.0 0.181162 0.027414 0.106000 0.161900 0.179200 0.195700 0.30400
  fractal_dimension_mean 569.0 0.062798 0.007060 0.049960 0.057700 0.061540 0.066120 0.09744
  radius_se 569.0 0.405172 0.277313 0.111500 0.232400 0.324200 0.478900 2.87300
  texture_se 569.0 1.216853 0.551648 0.360200 0.833900 1.108000 1.474000 4.88500
  perimeter_se 569.0 2.866059 2.021855 0.757000 1.606000 2.287000 3.357000 21.98000
  area_se 569.0 40.337079 45.491006 6.802000 17.850000 24.530000 45.190000 542.20000
  smoothness_se 569.0 0.007041 0.003003 0.001713 0.005169 0.006380 0.008146 0.03113
  compactness_se 569.0 0.025478 0.017000 0.002552 0.013020 0.020450 0.023450 0.12540

```

```

    [7] df['diagnosis'] = (df['diagnosis'] == 'M').astype(int) #encode the label into 1/0

    [8] corr = df.corr()

    [10] # Get the absolute value of the correlation
        cor_target = abs(corr["diagnosis"])

        # Select highly correlated features (threshold = 0.2)
        relevant_features = cor_target[cor_target>0.2]

        # Collect the names of the features
        names = [index for index, value in relevant_features.items()]

        # Drop the target variable from the results
        names.remove('diagnosis')

        # Display the results
        print(names)

    [11] X = df[names].values
        y = df['diagnosis']

    [12] def scale(X):
        """
        Standardizes the data in the array X.

        Parameters:
        X (numpy.ndarray): Features array of shape (n_samples, n_features).

        Returns:
        numpy.ndarray: The standardized features array.
        """
        # Calculate the mean and standard deviation of each feature
        mean = np.mean(X, axis=0)
        std = np.std(X, axis=0)

        # Standardize the data
        X = (X - mean) / std
        return X

    [13] X = scale(X)

    [14] def train_test_split(X, y, random_state=42, test_size=0.2):
        """
        Splits the data into training and testing sets.

        Parameters:
        X (numpy.ndarray): Features array of shape (n_samples, n_features).
        y (numpy.ndarray): Target array of shape (n_samples,).
        random_state (int): Seed for the random number generator. Default is 42.
        test_size (float): Proportion of samples to include in the test set. Default is 0.2.

        Returns:
        Tuple[numpy.ndarray]: A tuple containing X_train, X_test, y_train, y_test.
        """
        # Get number of samples
        n_samples = X.shape[0]

        # Set the seed for the random number generator
        np.random.seed(random_state)

        # Shuffle the indices
        shuffled_indices = np.random.permutation(np.arange(n_samples))

        # Determine the size of the test set
        test_size = int(n_samples * test_size)

        # Split the indices into test and train
        test_indices = shuffled_indices[:test_size]
        train_indices = shuffled_indices[test_size:]

        # Split the features and target arrays into test and train
        X_train, X_test = X[train_indices], X[test_indices]
        y_train, y_test = y[train_indices], y[test_indices]

        return X_train, X_test, y_train, y_test

    [15] X_train, X_test, y_train, y_test = train_test_split(X, y, test_size = 0.2, random_state=42) #split the data into traing and validation

```

```

[18] class SVM:
    """
    A Support Vector Machine (SVM) implementation using gradient descent.

    Parameters:
    -----
    iterations : int, default=1000
        The number of iterations for gradient descent.
    lr : float, default=0.01
        The learning rate for gradient descent.
    lambda : float, default=0.01
        The regularization parameter.

    Attributes:
    -----
    lambda : float
        The regularization parameter.
    iterations : int
        The number of iterations for gradient descent.
    lr : float
        The learning rate for gradient descent.
    w : numpy array
        The weights.
    b : float
        The bias.

    Methods:
    -----
    initialize_parameters(X)
        Initializes the weights and bias.
    gradient_descent(X, y)
        Updates the weights and bias using gradient descent.
        """
    def update_parameters(dw, db):
        """Updates the weights and bias."""
    def fit(X, y):
        """Fits the SVM to the data."""
    def predict(X):
        """Predicts the labels for the given data.

    """

    def __init__(self, iterations=1000, lr=0.01, lambda=0.01):
        """
        Initializes the SVM model.

        Parameters:
        -----
        iterations : int, default=1000
            The number of iterations for gradient descent.
        lr : float, default=0.01
            The learning rate for gradient descent.
        lambda : float, default=0.01
            The regularization parameter.
        """
        self.lambda = lambda
        self.iterations = iterations
        self.lr = lr
        self.w = None
        self.b = None
    def initialize_parameters(self, X):
        """
        Initializes the weights and bias.

        Parameters:
        -----

```

```

✓ [18]      X : numpy array
             The input data.
"""
m, n = X.shape
self.w = np.zeros(n)
self.b = 0

def gradient_descent(self, X, y):
    """
    Updates the weights and bias using gradient descent.

    Parameters:
    -----
    X : numpy array
        The input data.
    y : numpy array
        The target values.
    """
    y_ = np.where(y <= 0, -1, 1)
    for i, x in enumerate(X):
        if y_[i] * (np.dot(x, self.w) - self.b) >= 1:
            dw = 2 * self.lambdab * self.w
            db = 0
        else:
            dw = 2 * self.lambdab * self.w - np.dot(x, y_[i])
            db = y_[i]
        self.update_parameters(dw, db)

def update_parameters(self, dw, db):
    """
    Updates the weights and bias.

    Parameters:
    -----
    dw : numpy array
        The change in weights.
    db : float
        The change in bias.
    """
    self.w = self.w - self.lr * dw
    self.b = self.b - self.lr * db

```

```

def fit(self, X, y):
    """
    Fits the SVM to the data.

    Parameters:
    -----
    X : numpy array
        The input data.
    y : numpy array
        The target values.
    """
    self.initialize_parameters(X)
    for i in range(self.iterations):
        self.gradient_descent(X, y)

def predict(self, X):
    """
    Predicts the class labels for the test data.

    Parameters
    -----
    X : array-like, shape (n_samples, n_features)
        The input data.

    Returns
    -----
    y : numpy array
        The predicted class labels.
    """

```

```

[18]     Returns
-----
y_pred : array-like, shape (n_samples,)
    The predicted class labels.

"""
# get the outputs
output = np.dot(X, self.w) - self.b
# get the signs of the labels depending on if it's greater/less than zero
label_signs = np.sign(output)
#set predictions to 0 if they are less than or equal to -1 else set them to 1
predictions = np.where(label_signs <= -1, 0, 1)
return predictions

[19] def accuracy(y_true, y_pred):
"""
Computes the accuracy of a classification model.

Parameters:
-----
y_true (numpy array): A numpy array of true labels for each data point.
y_pred (numpy array): A numpy array of predicted labels for each data point.

Returns:
-----
float: The accuracy of the model
"""
total_samples = len(y_true)
correct_predictions = np.sum(y_true == y_pred)
return (correct_predictions / total_samples)

[20] model = SVM()
model.fit(X_train,y_train)
predictions = model.predict(X_test)

accuracy(y_test, predictions)

→ 0.9823008849557522

[28] model.predict([-0.47069438, -0.16048584, -0.44810956, -0.49199876, 0.23411429,
0.02765051, -0.10984741, -0.27623152, 0.41394897, -0.03274296,
-0.18269561, -0.22105292, -0.35591235, -0.16192949, -0.23133322,
-0.26903951, -0.16890536, -0.33393537, -0.35629925, 0.4485028 ,
-0.10474068, -0.02441212, -0.19956318, 0.18320441, 0.19695794])

→ array(0)

```

9. Build k-Means algorithm to cluster a set of data stored in a .CSV file.

Algorithm:

K Means clustering algorithm

- ① Select number K to decide the number of clusters
- ② Select random K points or centroids
- ③ Assign each point to closest centroid for predefined cluster
- ④ Calculate the variance and place a new centroid of each cluster.
- ⑤ Repeat 3, 4, reassign the centroid
- ⑥ If any reassignment occurs, goto step 4, else goto finish.
- ⑦ The model is ready.

```

88 [1] from google.colab import drive
drive.mount('/content/drive')

→ Mounted at /content/drive

1s [2] import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import plotly.express as px
import seaborn as sns
import plotly.graph_objects as go

1s [3] iris = pd.read_csv("/content/drive/MyDrive/Iris.csv") #Load Data
iris.drop('Id',inplace=True,axis=1) #Drop Id column

1s [4] X = iris.iloc[:, :-1] #Set our training data
y = iris.iloc[:, -1] #We'll use this just for visualization as clustering doesn't require labels

1s [5] class Kmeans:
    """
        K-Means clustering algorithm implementation.

    Parameters:
        K (int): Number of clusters

    Attributes:
        K (int): Number of clusters
        centroids (numpy.ndarray): Array containing the centroids of each cluster

    Methods:
        __init__(self, K): Initializes the Kmeans instance with the specified number of clusters.
        initialize_centroids(self, X): Initializes the centroids for each cluster by selecting K random points from the dataset.
        assign_points_centroids(self, X): Assigns each point in the dataset to the nearest centroid.
        compute_mean(self, X, points): Computes the mean of the points assigned to each centroid.
        fit(self, X, iterations=10): Clusters the dataset using the K-Means algorithm.
    """

    def __init__(self, K):
        assert K > 0, "K should be a positive integer."
        self.K = K

    def initialize_centroids(self, X):
        assert X.shape[0] >= self.K, "Number of data points should be greater than or equal to K."

        randomized_X = np.random.permutation(X.shape[0])
        centroid_idx = randomized_X[:self.K] # get the indices for the centroids
        self.centroids = X[centroid_idx] # assign the centroids to the selected points

    def assign_points_centroids(self, X):
        """
            Assign each point in the dataset to the nearest centroid.

        Parameters:
            X (numpy.ndarray): dataset to cluster

        Returns:
            numpy.ndarray: array containing the index of the centroid for each point
        """
        X = np.expand_dims(X, axis=1) # expand dimensions to match shape of centroids
        distance = np.linalg.norm((X - self.centroids), axis=-1) # calculate Euclidean distance between each point and each centroid
        points = np.argmin(distance, axis=1) # assign each point to the closest centroid
        assert len(points) == X.shape[0], "Number of assigned points should equal the number of data points."

```

```

✓ [5]     points = np.argmin(distance, axis=1) # assign each point to the closest centroid
0s      assert len(points) == X.shape[0], "Number of assigned points should equal the number of data points."
return points

def compute_mean(self, X, points):
    """
    Compute the mean of the points assigned to each centroid.

    Parameters:
    X (numpy.ndarray): dataset to cluster
    points (numpy.ndarray): array containing the index of the centroid for each point

    Returns:
    numpy.ndarray: array containing the new centroids for each cluster
    """
    centroids = np.zeros((self.K, X.shape[1])) # initialize array to store centroids
    for i in range(self.K):
        centroid_mean = X[points == i].mean(axis=0) # calculate mean of the points assigned to the current centroid
        centroids[i] = centroid_mean # assign the new centroid to the mean of its points
    return centroids

def fit(self, X, iterations=10):
    """
    Cluster the dataset using the K-Means algorithm.

    Parameters:
    X (numpy.ndarray): dataset to cluster
    iterations (int): number of iterations to perform (default=10)

    Returns:
    numpy.ndarray: array containing the final centroids for each cluster
    numpy.ndarray: array containing the index of the centroid for each point
    """
    self.initialize_centroids(X) # initialize the centroids
    ...
    ...


✓ [5]     """
0s      self.initialize_centroids(X) # initialize the centroids
for i in range(iterations):
    points = self.assign_points_centroids(X) # assign each point to the nearest centroid
    self.centroids = self.compute_mean(X, points) # compute the new centroids based on the mean of their points

    # Assertions for debugging and validation
    assert len(self.centroids) == self.K, "Number of centroids should equal K."
    assert X.shape[1] == self.centroids.shape[1], "Dimensionality of centroids should match input data."
    assert max(points) < self.K, "Cluster index should be less than K."
    assert min(points) >= 0, "Cluster index should be non-negative."

return self.centroids, points

```

```

✓ [6]  X = X.values

✓ [7]  kmeans = Kmeans(3)

centroids, points = kmeans.fit(X, 1000)

✓ [8]  fig = go.Figure()
fig.add_trace(go.Scatter(
    x=X[points == 0, 0], y=X[points == 0, 1],
    mode='markers', marker_color='#DB4CB2', name='Iris-setosa'
))
fig.add_trace(go.Scatter(
    x=X[points == 1, 0], y=X[points == 1, 1],
    mode='markers', marker_color='#c9e9f6', name='Iris-versicolour'
))

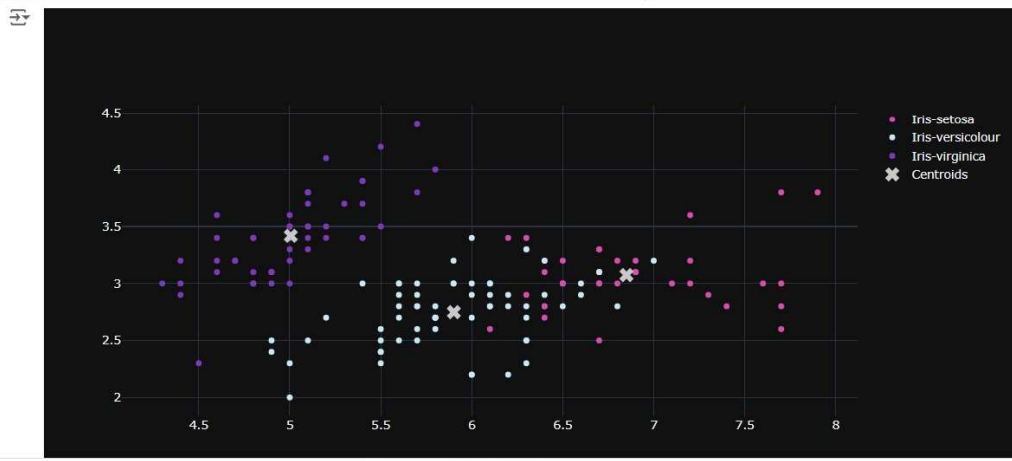
```

```

[8] ))
fig.add_trace(go.Scatter(
    x=X[points == 2, 0], y=X[points == 2, 1],
    mode='markers',marker_color="#7D3AC1",name='Iris-virginica'
))

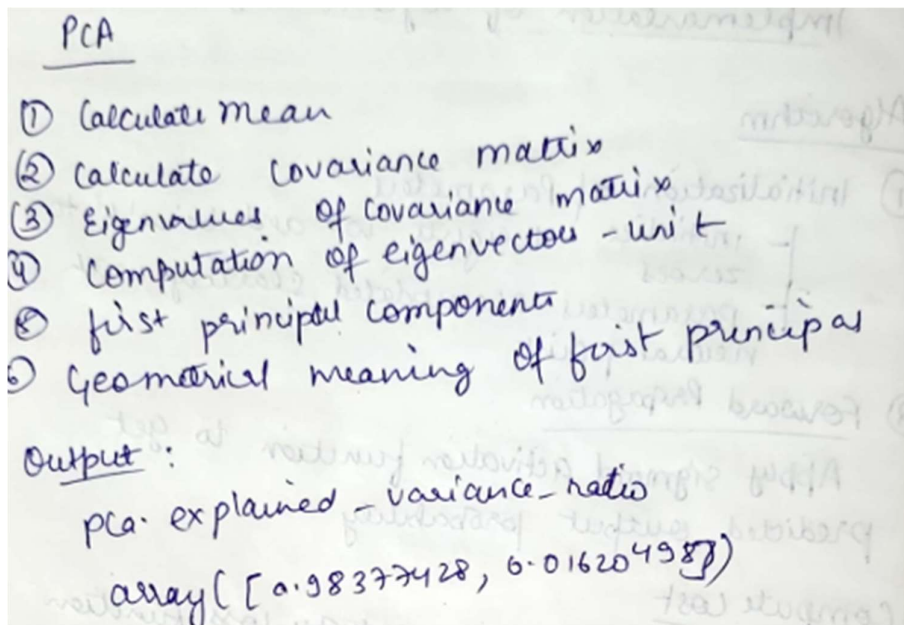
fig.add_trace(go.Scatter(
    x=centroids[:, 0], y=centroids[:,1],
    mode='markers',marker_color="#CAC9CD",marker_symbol=4,marker_size=13,name='Centroids'
))
fig.update_layout(template='plotly_dark',width=1000, height=500,)


```



10. Implement Dimensionality reduction using Principle Component Analysis

(PCA)Algorithm:



+ Code + Text

✓ RAM Disk

```

✓ [1] from google.colab import drive
drive.mount('/content/drive')

↳ Mounted at /content/drive

✓ [2] import seaborn as sns
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import plotly.express as px
import plotly.graph_objects as go
from plotly.subplots import make_subplots

✓ [3] df = pd.read_csv('/content/drive/MyDrive/breast-cancer.csv')
df.head()

```

	id	diagnosis	radius_mean	texture_mean	perimeter_mean	area_mean	smoothness_mean	compactness_mean	concavity_mean	concave_points_mean	... radius_worst	
0	842302	M	17.99	10.38	122.80	1001.0	0.11840	0.27760	0.3001	0.14710	...	25.38
1	842517	M	20.57	17.77	132.90	1326.0	0.08474	0.07864	0.0869	0.07017	...	24.99
2	84300903	M	19.69	21.25	130.00	1203.0	0.10960	0.15990	0.1974	0.12790	...	23.57
3	84348301	M	11.42	20.38	77.58	386.1	0.14250	0.28390	0.2414	0.10520	...	14.91
4	84358402	M	20.29	14.34	135.10	1297.0	0.10030	0.13280	0.1980	0.10430	...	22.54

5 rows x 32 columns

```
[4] df.drop('id', axis=1, inplace=True) #drop redundant columns

[5] df['diagnosis'] = (df['diagnosis'] == 'M').astype(int) #encode the label into 1/0

[6] corr = df.corr()

[7] # Get the absolute value of the correlation
cor_target = abs(corr["diagnosis"])

# Select highly correlated features (threshold = 0.2)
relevant_features = cor_target[cor_target>0.2]

# Collect the names of the features
names = [index for index, value in relevant_features.items()]

# Drop the target variable from the results
names.remove('diagnosis')

# Display the results
print(names)

[8] ['radius_mean', 'texture_mean', 'perimeter_mean', 'area_mean', 'smoothness_mean', 'compactness_mean', 'concavity_mean', 'concave p

[9] X = df[names].values

[10] X = df[names].values

[11] class PCA:
    """
    Principal Component Analysis (PCA) class for dimensionality reduction.
    """

    def __init__(self, n_components):
        """
        Constructor method that initializes the PCA object with the number of components to retain.

        Args:
        - n_components (int): Number of principal components to retain.
        """
        self.n_components = n_components
    def fit(self, X):
        """
        Fits the PCA model to the input data and computes the principal components.

        Args:
        - X (numpy.ndarray): Input data matrix with shape (n_samples, n_features).
        """
        # Compute the mean of the input data along each feature dimension.
        mean = np.mean(X, axis=0)

        # Subtract the mean from the input data to center it around zero.
        X = X - mean

        # Compute the covariance matrix of the centered input data.
        cov = np.cov(X.T)
```

```

✓ [11] # Compute the covariance matrix of the centered input data.
cov = np.cov(X.T)

# Compute the eigenvectors and eigenvalues of the covariance matrix.
eigenvalues, eigenvectors = np.linalg.eigh(cov)
# Reverse the order of the eigenvalues and eigenvectors.
eigenvalues = eigenvalues[::-1]
eigenvectors = eigenvectors[:, ::-1]

# Keep only the first n_components eigenvectors as the principal components.
self.components = eigenvectors[:, :self.n_components]

# Compute the explained variance ratio for each principal component.
# Compute the total variance of the input data
total_variance = np.sum(np.var(X, axis=0))

# Compute the variance explained by each principal component
self.explained_variances = eigenvalues[:self.n_components]

# Compute the explained variance ratio for each principal component
self.explained_variance_ratio_ = self.explained_variances / total_variance
def transform(self, X):
    """
    Transforms the input data by projecting it onto the principal components.

    Args:
    - X (numpy.ndarray): Input data matrix with shape (n_samples, n_features).

    Returns:
    - transformed_data (numpy.ndarray): Transformed data matrix with shape (n_samples, n_components).
    """
    # Center the input data around zero using the mean computed during the fit step.
    X = X - np.mean(X, axis=0)

```



```

✓ [11] # Project the centered input data onto the principal components.
Ctrl+M B     transformed_data = np.dot(X, self.components)

return transformed_data

def fit_transform(self, X):
    """
    Fits the PCA model to the input data and computes the principal components then
    transforms the input data by projecting it onto the principal components.

    Args:
    - X (numpy.ndarray): Input data matrix with shape (n_samples, n_features).
    """
    self.fit(X)
    transformed_data = self.transform(X)
    return transformed_data

```



```

✓ [12] pca = PCA(2)

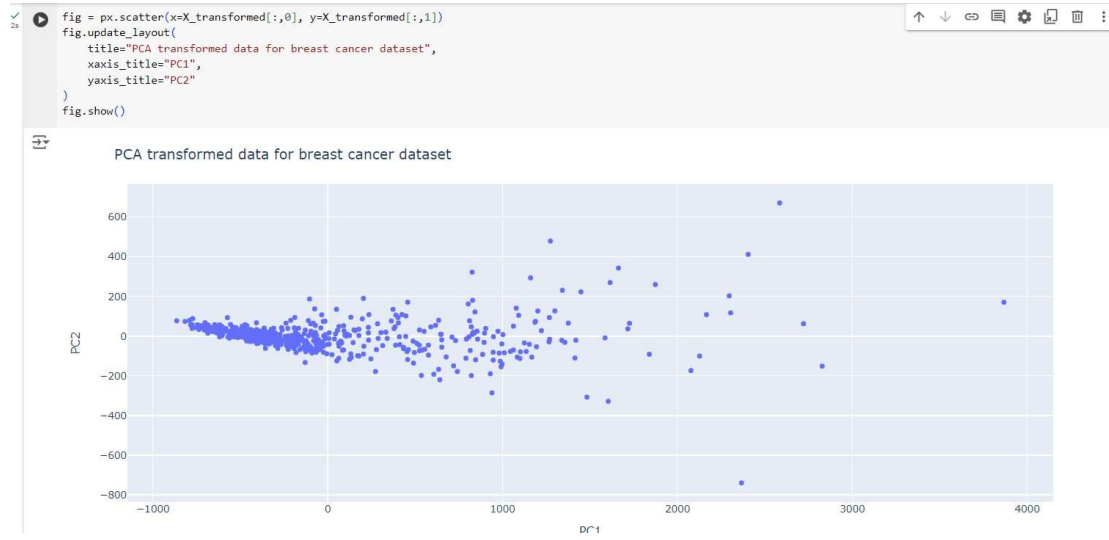
✓ [13] pca.fit(X)

✓ [14] pca.explained_variance_ratio_
→ array([0.98377428, 0.01620498])

✓ [15] X_transformed = pca.transform(X)

✓ [16] X_transformed[:, 1].shape
→ (569,)

```



11. Build Artificial Neural Network model with back propagation on a given dataset

Algorithm:

Build ANN with backpropagation

- Create feed forward network with n inputs, n hidden units, nout outputs.
- Initialize all network weights to small random numbers.
- Until the termination condition is met,
Do
 - for each (\vec{x}, \vec{t}) in training examples
 - Propagate input forward
 - Propagate error backward
 - for each hidden unit h, calculate error
 - update weight

```

✓ 0s 1 import numpy as np
   from sklearn.model_selection import train_test_split

   db = np.loadtxt("./content/duke-breast-cancer.txt")
   print("Database raw shape (%s,%s)" % np.shape(db))

   ↵ Database raw shape (86,7130)

✓ 0s 2 [3] np.random.shuffle(db)
   y = db[:, 0]
   x = np.delete(db, [0], axis=1)
   x_train, x_test, y_train, y_test = train_test_split(x, y, test_size=0.1)
   print(np.shape(x_train),np.shape(x_test))

   ↵ (77, 7129) (9, 7129)

✓ 0s 3 [4] hidden_layer = np.zeros(72)
   weights = np.random.random((len(x[0]), 72))
   output_layer = np.zeros(2)
   hidden_weights = np.random.random((72, 2))

✓ 0s 4 [5] def sum_function(weights, index_locked_col, x):
   result = 0
   for i in range(0, len(x)):
       result += x[i] * weights[i][index_locked_col]
   return result

✓ 0s 5 [6] def activate_layer(layer, weights, x):
   for i in range(0, len(layer)):
       layer[i] = 1.7159 * np.tanh(2.0 * sum_function(weights, i, x) / 3.0)

✓ 0s 6 [7] def soft_max(layer):
   soft_max_output_layer = np.zeros(len(layer))
   for i in range(0, len(layer)):
       denominator = 0
       for j in range(0, len(layer)):
           denominator += np.exp(layer[j] - np.max(layer))
       soft_max_output_layer[i] = np.exp(layer[i] - np.max(layer)) / denominator
   return soft_max_output_layer

✓ 0s 7 [8] def recalculate_weights(learning_rate, weights, gradient, activation):
   for i in range(0, len(weights)):
       for j in range(0, len(weights[i])):
           weights[i][j] = (learning_rate * gradient[j] * activation[i]) + weights[i][j]

✓ 0s 8 [9] def back_propagation(hidden_layer, output_layer, one_hot_encoding, learning_rate, x):
   output_derivative = np.zeros(2)
   output_gradient = np.zeros(2)
   for i in range(0, len(output_layer)):
       output_derivative[i] = (1.0 - output_layer[i]) * output_layer[i]
   for i in range(0, len(output_layer)):
       output_gradient[i] = output_derivative[i] * (one_hot_encoding[i] - output_layer[i])
   hidden_derivative = np.zeros(72)
   hidden_gradient = np.zeros(72)
   for i in range(0, len(hidden_layer)):
       hidden_derivative[i] = (1.0 - hidden_layer[i]) * (1.0 + hidden_layer[i])
   for i in range(0, len(hidden_layer)):
       sum_ = 0
       for j in range(0, len(output_gradient)):
           sum_ += output_gradient[j] * hidden_weights[i][j]
       hidden_gradient[i] = sum_ * hidden_derivative[i]
   recalculate_weights(learning_rate, hidden_weights, output_gradient, hidden_layer)
   recalculate_weights(learning_rate, weights, hidden_gradient, x)

```

```

✓ [10] one_hot_encoding = np.zeros((2,2))
    for i in range(0, len(one_hot_encoding)):
        one_hot_encoding[i][i] = 1
    training_correct_answers = 0
    for i in range(0, len(x_train)):
        activate_layer(hidden_layer, weights, x_train[i])
        activate_layer(output_layer, hidden_weights, hidden_layer)
        output_layer = soft_max(output_layer)
        training_correct_answers += 1 if y_train[i] == np.argmax(output_layer) else 0
        back_propagation(hidden_layer, output_layer, one_hot_encoding[int(y_train[i])], -1, x_train[i])
    print("MLP Correct answers while learning: %s / %s (Accuracy = %s) on %s database." % (training_correct_answers, len(x_train),
                                                                                           training_correct_answers/len(x_train), "Duke breast cancer"))

→ MLP Correct answers while learning: 44 / 77 (Accuracy = 0.5714285714285714) on Duke breast cancer database.

✓ [11] testing_correct_answers = 0
    for i in range(0, len(x_test)):
        activate_layer(hidden_layer, weights, x_test[i])
        activate_layer(output_layer, hidden_weights, hidden_layer)
        output_layer = soft_max(output_layer)
        testing_correct_answers += 1 if y_test[i] == np.argmax(output_layer) else 0
    print("MLP Correct answers while testing: %s / %s (Accuracy = %s) on %s database." % (testing_correct_answers, len(x_test),
                                                                                           testing_correct_answers/len(x_test), "Duke breast cancer"))

→ MLP Correct answers while testing: 8 / 9 (Accuracy = 0.8888888888888888) on Duke breast cancer database

```

11a. Implement Random forest ensemble method on a given dataset.

Algorithm:

Random Forest:

- ① Import libraries
- ② Load and prepare dataset
- ③ Train the data before that do `train test split`.
- ④ Initialize Random forest regressor.
- ⑤ Train model.
- ⑥ Make predictions
- ⑦ Evaluate using MSE (mean square error)

Output

$$\text{Accuracy} = 0.93$$

```

[17] from google.colab import drive
drive.mount('/content/drive')

Mounted at /content/drive

[18] import math
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
import plotly.express as px

[19] iris = pd.read_csv("/content/drive/MyDrive/Iris.csv") #Load Data
iris.drop('Id',inplace=True,axis=1) #Drop Id column

[20] iris.head().style.background_gradient(cmap =sns.light_palette("seagreen", as_cmap=True)
)

SepallengthCm SepalWidthCm PetallengthCm PetalWidthCm Species
0 5.100000 3.500000 1.400000 0.200000 Iris-setosa
1 4.900000 3.000000 1.400000 0.200000 Iris-setosa
2 4.700000 3.200000 1.300000 0.200000 Iris-setosa
3 4.600000 3.100000 1.500000 0.200000 Iris-setosa
4 5.000000 3.600000 1.400000 0.200000 Iris-setosa

[21] X_df = iris.iloc[:, :-1] #Set our training dataframe
y_df = iris.iloc[:, -1] # Set our training labels dataframe

[22] fig = px.pie(iris, 'Species',color_discrete_sequence=['#3dec84','#009688','#2E8B57'],title='Data Distribution',template='plotly')
fig.show()

Data Distribution

```

The pie chart illustrates the distribution of Iris species. It consists of three equal-sized segments, each representing one-third of the total data. The legend indicates the colors for each species: Iris-setosa (light green), Iris-versicolor (dark green), and Iris-virginica (medium green).

Species	Percentage
Iris-setosa	33.3%
Iris-versicolor	33.3%
Iris-virginica	33.3%

```

[23] iris['Species'] = iris['Species'].astype("category")
codes = iris['Species'].cat.codes

```

```

Ds [23] iris['Species'] = iris['Species'].astype("category")
      codes = iris['Species'].cat.codes

Ds [24] def train_test_split(X, y, random_state=42, test_size=0.2):
      """
      Splits the data into training and testing sets.

      Parameters:
          X (numpy.ndarray): Features array of shape (n_samples, n_features).
          y (numpy.ndarray): Target array of shape (n_samples,).
          random_state (int): Seed for the random number generator. Default is 42.
          test_size (float): Proportion of samples to include in the test set. Default is 0.2.

      Returns:
          Tuple[numpy.ndarray]: A tuple containing X_train, X_test, y_train, y_test.
      """
      # Get number of samples
      n_samples = X.shape[0]

      # Set the seed for the random number generator
      np.random.seed(random_state)

      # Shuffle the indices
      shuffled_indices = np.random.permutation(np.arange(n_samples))

      # Determine the size of the test set
      test_size = int(n_samples * test_size)

      # Split the indices into test and train
      test_indices = shuffled_indices[:test_size]
      train_indices = shuffled_indices[test_size:]

      # Split the features and target arrays into test and train
      X_train, X_test = X[train_indices], X[test_indices]
      y_train, y_test = y[train_indices], y[test_indices]

      return X_train, X_test, y_train, y_test

Ds [25] X = iris.iloc[:, :-1].values
      y = iris.iloc[:, -1].values.reshape(-1,1)
      X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=.2, random_state=41)

Ds [26] from sklearn.tree import DecisionTreeClassifier
      m = DecisionTreeClassifier()

Ds [27] class RandomForest:
      """
      A random forest classifier.

      Parameters
      -----
      n_trees : int, default=7
          The number of trees in the random forest.
      max_depth : int, default=7
          The maximum depth of each decision tree in the random forest.
      min_samples : int, default=2
          The minimum number of samples required to split an internal node
          of each decision tree in the random forest.

      Attributes
      -----
      n_trees : int
          The number of trees in the random forest.
      max_depth : int
          The maximum depth of each decision tree in the random forest.
      min_samples : int
          The minimum number of samples required to split an internal node
          of each decision tree in the random forest.
      trees : list of DecisionTreeClassifier
          The decision trees in the random forest.
      """

      def __init__(self, n_trees=7, max_depth=7, min_samples=2):
          """
          Initialize the random forest classifier.
      """

```

```

[27]     The decision trees in the random forest.

def __init__(self, n_trees=7, max_depth=7, min_samples=2):
    """
    Initialize the random forest classifier.

    Parameters
    -----
    n_trees : int, default=7
        The number of trees in the random forest.
    max_depth : int, default=7
        The maximum depth of each decision tree in the random forest.
    min_samples : int, default=2
        The minimum number of samples required to split an internal node
        of each decision tree in the random forest.
    """

    self.n_trees = n_trees
    self.max_depth = max_depth
    self.min_samples = min_samples
    self.trees = []

def fit(self, X, y):
    """
    Build a random forest classifier from the training set (X, y).

    Parameters
    -----
    X : array-like of shape (n_samples, n_features)
        The training input samples.
    y : array-like of shape (n_samples,)
        The target values.

    Returns
    -----
    self : object
        Returns self.
    """

    # Create an empty list to store the trees.
    self.trees = []
    # Concatenate X and y into a single dataset.
    dataset = np.concatenate((X, y.reshape(-1, 1)), axis=1)
    # Loop over the number of trees.

    # Create an empty list to store the trees.
    self.trees = []
    # Concatenate X and y into a single dataset.
    dataset = np.concatenate((X, y.reshape(-1, 1)), axis=1)
    # Loop over the number of trees.
    for _ in range(self.n_trees):
        # Create a decision tree instance.
        tree = DecisionTreeClassifier(max_depth=self.max_depth, min_samples_split=self.min_samples)
        # Sample from the dataset with replacement (bootstrapping).
        dataset_sample = self.bootstrap_samples(dataset)
        # Get the X and y samples from the dataset sample.
        X_sample, y_sample = dataset_sample[:, :-1], dataset_sample[:, -1]
        # Fit the tree to the X and y samples.
        tree.fit(X_sample, y_sample)
        # Store the tree in the list of trees.
        self.trees.append(tree)

    return self

def bootstrap_samples(self, dataset):
    """
    Bootstrap the dataset by sampling from it with replacement.

    Parameters
    -----
    dataset : array-like of shape (n_samples, n_features + 1)
        The dataset to bootstrap.

    Returns
    -----
    dataset_sample : array-like of shape (n_samples, n_features + 1)
        The bootstrapped dataset sample.
    """

    # Get the number of samples in the dataset.
    n_samples = dataset.shape[0]
    # Generate random indices to index into the dataset with replacement.
    np.random.seed(1)
    indices = np.random.choice(n_samples, n_samples, replace=True)
    # Return the bootstrapped dataset sample using the generated indices.
    dataset_sample = dataset[indices]
    return dataset_sample

def most_common_label(self, v):

```

```
[27]     def most_common_label(self, y):
        """
        Return the most common label in an array of labels.

        Parameters
        -----
        y : array-like of shape (n_samples,)
            The array of labels.

        Returns
        -----
        most_occurring_value : int or float
            The most common label in the array.
        """
        y = list(y)
        # get the highest present class in the array
        most_occurring_value = max(y, key=y.count)
        return most_occurring_value

    def predict(self, X):
        """
        Predict class for X.

        Parameters
        -----
        X : array-like of shape (n_samples, n_features)
            The input samples.

        Returns
        -----
        majority_predictions : array-like of shape (n_samples,)
            The predicted classes.
        """
        #get prediction from each tree in the tree list on the test data
        predictions = np.array([tree.predict(X) for tree in self.trees])
        # get prediction for the same sample from all trees for each sample in the test data
        preds = np.swapaxes(predictions, 0, 1)
        #get the most voted value by the trees and store it in the final predictions array
        majority_predictions = np.array([self.most_common_label(pred) for pred in preds])
        return majority_predictions
```

```
[28] def accuracy(y_true, y_pred):
    """
    Computes the accuracy of a classification model.

    Parameters:
    y_true (numpy array): A numpy array of true labels for each data point.
    y_pred (numpy array): A numpy array of predicted labels for each data point.

    Returns:
    float: The accuracy of the model, expressed as a percentage.
    """
    y_true = y_true.flatten()
    total_samples = len(y_true)
    correct_predictions = np.sum(y_true == y_pred)
    return (correct_predictions / total_samples)
```

```
[35] from sklearn.preprocessing import LabelEncoder

label_encoder = LabelEncoder()
y_train_encoded = label_encoder.fit_transform(y_train)
y_test_encoded = label_encoder.transform(y_test)

/usr/local/lib/python3.10/dist-packages/sklearn/preprocessing/_label.py:116: DataConversionWarning:
A column-vector y was passed when a 1d array was expected. Please change the shape of y to (n_samples, ), for example using ravel().
/usr/local/lib/python3.10/dist-packages/sklearn/preprocessing/_label.py:134: DataConversionWarning:
A column-vector y was passed when a 1d array was expected. Please change the shape of y to (n_samples, ), for example using ravel().
```

```
[39] from sklearn.preprocessing import LabelEncoder  
  
label_encoder = LabelEncoder()  
y_train_encoded = label_encoder.fit_transform(y_train.ravel())  
y_test_encoded = label_encoder.transform(y_test.ravel())  
model = RandomForest(10, 10, 2)  
model.fit(X_train, y_train_encoded)  
  
predictions = model.predict(X_test)  
accuracy(y_test_encoded, predictions)
```

→ 0.9333333333333333

```
▶ from sklearn.tree import DecisionTreeClassifier  
  
# Create and train the decision tree model  
dt = DecisionTreeClassifier()  
dt.fit(X_train, y_train_encoded)  
  
# Make predictions on the test data  
predictions = dt.predict(X_test)  
  
# Calculate accuracy  
accuracy(y_test_encoded, predictions)
```

→ 0.9

11b. Implement Boosting ensemble method on a given dataset.

Algorithm:

31/05/2024AdaBoost Algorithm

- ① Import libraries
- ② Load & prepare data
- ③ Initialize AdaBoost model - (learning_rate, n_estimators)
- ④ Make the model train
- ⑤ Make predictions
- ⑥ Evaluate model on metrics like Mean Absolute Error.

Output

accuracy: 0.9467

S. Ichchou

```
[15] # Compute error rate, alpha and w
def compute_error(y, y_pred, w_i):
    ...
    Calculate the error rate of a weak classifier m. Arguments:
    y: actual target value
    y_pred: predicted value by weak classifier
    w_i: individual weights for each observation

    Note that all arrays should be the same length
    ...
    return (sum(w_i * (np.not_equal(y, y_pred)).astype(int)))/sum(w_i)

def compute_alpha(error):
    ...
    Calculate the weight of a weak classifier m in the majority vote of the final classifier. This is called
    alpha in chapter 10.1 of The Elements of Statistical Learning. Arguments:
    error: error rate from weak classifier m
    ...
    return np.log((1 - error) / error)

def update_weights(w_i, alpha, y, y_pred):
    ...
    Update individual weights w_i after a boosting iteration. Arguments:
    w_i: individual weights for each observation
    y: actual target value
    y_pred: predicted value by weak classifier
    alpha: weight of weak classifier used to estimate y_pred
    ...
    return w_i * np.exp(alpha * (np.not_equal(y, y_pred)).astype(int))
```

```

[16] # Define AdaBoost class
class AdaBoost:

    def __init__(self):
        self.alphas = []
        self.G_M = []
        self.M = None
        self.training_errors = []
        self.prediction_errors = []

    def fit(self, X, y, M = 100):
        ...
        Fit model. Arguments:
        X: independent variables - array-like matrix
        y: target variable - array-like vector
        M: number of boosting rounds. Default is 100 - integer
        ...

        # Clear before calling
        self.alphas = []
        self.training_errors = []
        self.M = M

        # Iterate over M weak classifiers
        for m in range(0, M):

            # Set weights for current boosting iteration
            if m == 0:
                w_i = np.ones(len(y)) * 1 / len(y) # At m = 0, weights are all the same and equal to 1 / N
            else:
                # (d) Update w_i
                w_i = update_weights(w_i, alpha_m, y, y_pred)

            # (a) Fit weak classifier and predict labels
            G_m = DecisionTreeClassifier(max_depth = 1)      # Stump: Two terminal-node classification tree
            G_m.fit(X, y, sample_weight = w_i)
            y_pred = G_m.predict(X)

            self.G_M.append(G_m) # Save to list of weak classifiers

            # (b) Compute error
            error_m = compute_error(y, y_pred, w_i)
            w_i = update_weights(w_i, alpha_m, y, y_pred)

            # (a) Fit weak classifier and predict labels
            G_m = DecisionTreeClassifier(max_depth = 1)      # Stump: Two terminal-node classification tree
            G_m.fit(X, y, sample_weight = w_i)
            y_pred = G_m.predict(X)

            self.G_M.append(G_m) # Save to list of weak classifiers

            # (b) Compute error
            error_m = compute_error(y, y_pred, w_i)
            self.training_errors.append(error_m)

            # (c) Compute alpha
            alpha_m = compute_alpha(error_m)
            self.alphas.append(alpha_m)

            assert len(self.G_M) == len(self.alphas)

    def predict(self, X):
        ...
        Predict using fitted model. Arguments:
        X: independent variables - array-like
        ...

        # Initialise dataframe with weak predictions for each observation
        weak_preds = pd.DataFrame(index = range(len(X)), columns = range(self.M))

        # Predict class label for each weak classifier, weighted by alpha_m
        for m in range(self.M):
            y_pred_m = self.G_M[m].predict(X) * self.alphas[m]
            weak_preds.iloc[:,m] = y_pred_m

        # Calculate final predictions
        y_pred = (1 * np.sign(weak_preds.T.sum())).astype(int)

        return y_pred

```

```
[17] import pandas as pd
     import numpy as np
     from sklearn.model_selection import train_test_split
     from sklearn.tree import DecisionTreeClassifier

    # Dataset
    df = pd.read_csv('/content/spambase.data', header = None)

    # Column names
    names = pd.read_csv('/content/spambase.names', sep = ':', skiprows=range(0, 33), header = None)
    col_names = list(names[0])
    col_names.append('Spam')

    # Rename df columns
    df.columns = col_names

    # Convert classes in target variable to {-1, 1}
    df['Spam'] = df['Spam'] * 2 - 1

    # Train - test split
    X_train, X_test, y_train, y_test = train_test_split(df.drop(columns = 'Spam').values,
                                                       df['Spam'].values,
                                                       train_size = 3065,
                                                       random_state = 2)
```

```
▶ # Fit model
ab = AdaBoost()
ab.fit(X_train, y_train, M = 400)

# Predict on test set
y_pred = ab.predict(X_test)

from sklearn.metrics import accuracy_score

# Calculate accuracy
accuracy = accuracy_score(y_test, y_pred)
print("Accuracy:", accuracy)
```

```
→ Accuracy: 0.9440104166666666
```