UCS2612 Machine Learning Laboratory

A9. Applications of dimensionality reduction techniques

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CSE-A

Aim

Develop a python program to perform dimensionality reduction using PCA and LDA. Visualize the features from the dataset and interpret the results obtained by the model using Matplotlib library.

Dataset:- http://www3.dsi.uminho.pt/pcortez/wine/winequality.zip

Import Libraries

```
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
from sklearn.metrics import roc_curve, roc_auc score
from sklearn.model selection import train test split
import seaborn as sns
import matplotlib.pyplot as plt
from sklearn.preprocessing import LabelEncoder
from sklearn.preprocessing import StandardScaler, MinMaxScaler
from sklearn.preprocessing import StandardScaler
from sklearn.ensemble import RandomForestClassifier,
GradientBoostingClassifier
from sklearn.metrics import accuracy score, classification report,
confusion matrix
import numpy as np
import matplotlib.pyplot as plt
```

Read dataset

```
# importing or loading the dataset
data = pd.read csv("C:/Users/ashwi/Downloads/ML
Lab/A9/winequality/winequality-red.csv", header=0, sep=";")
data.head()
   fixed acidity volatile acidity citric acid residual sugar
chlorides \
             7.4
                              0.70
                                            0.00
                                                             1.9
0.076
                              0.88
                                            0.00
             7.8
                                                             2.6
0.098
                              0.76
                                            0.04
             7.8
                                                             2.3
0.092
                              0.28
            11.2
                                            0.56
                                                             1.9
0.075
             7.4
                              0.70
                                                             1.9
                                            0.00
0.076
   free sulfur dioxide total sulfur dioxide density pH sulphates
0
                  11.0
                                         34.0 0.9978 3.51
                                                                    0.56
1
                  25.0
                                         67.0 0.9968 3.20
                                                                    0.68
```

2	15	.0		54.0	0.9970	3.26	0.65	
3	17	.0		60.0	0.9980	3.16	0.58	
4	11	.0		34.0	0.9978	3.51	0.56	
alcoho								
1 9. 2 9.	8 5							
3 9.	8 6							
4 9.								
data.desc							,	
	xed acidity 1599.000000	volatile ac 1599.		citric 1599.0		esidual s 1599.0	_	
mean std	8.319637 1.741096	0.527821 0.179060		0.270976 0.194801			2.538806 1.409928	
min 25%	4.600000 7.100000	0.	120000 390000	0.0	00000 90000	0.9	00000	
50%	50% 7.900000		520000	0.260000		2.2	2.200000	
75% max	9.200000 15.900000		640000 580000		20000 00000		00000	
		free sulfur	dioxide	total	sulfur	dioxide		
density count 15		1599	.000000		1599	.000000		
1599.0000 mean	0.087467	15	.874922		4 6	5.467792		
0.996747 std	0.047065	1.0	.460157		3.0	2.895324		
0.001887								
min 0.990070	0.012000		.000000			5.000000		
25% 0.995600	0.070000	7	.000000		22	2.000000		
50% 0.996750	0.079000	14	.000000		38	3.000000		
75% 0.997835	0.090000	21	.000000		62	2.000000		
max	0.611000	72	.000000		289	0.00000		
1.003690								
count 15	рН 599.000000	sulphates 1599.000000	1599.00		qual 1599.000	000		
mean std	3.311113 0.154386	0.658149 0.169507	10.42		5.636 0.807			

3.310000 0.620000 10.200000 6.000000 3.400000 0.730000 11.100000 6.000000 4.010000 2.000000 14.900000 8.000000
--

Data Pre-processing

!				
data.corr()				
	fixed acidity v	volatile acidity	citric acid \	
fixed acidity	1.000000	-0.256131	0.671703	
volatile acidity	-0.256131	1.00000	-0.552496	
citric acid	0.671703	-0.552496	1.000000	
residual sugar	0.114777	0.001918	0.143577	
chlorides	0.093705	0.061298	0.203823	
free sulfur dioxide	-0.153794	-0.010504	-0.060978	
total sulfur dioxide	-0.113181	0.076470	0.035533	
density	0.668047	0.022026	0.364947	
рН	-0.682978	0.234937	-0.541904	
sulphates	0.183006	-0.260987	0.312770	
alcohol	-0.061668	-0.202288	0.109903	
quality	0.124052	-0.390558	0.226373	
	residual sugar	chlorides free s	ulfur	
dioxide \				
fixed acidity	0.114777	0.093705	-0.153794	
volatile acidity	0.001918	0.061298	-0.010504	
volatile actualty	0.001916	0.001290	-0.010304	
citric acid	0.143577	0.203823	-0.060978	
residual sugar	1.000000	0.055610	0.187049	
-				
chlorides	0.055610	1.000000	0.005562	
free sulfur dioxide	0.187049	0.005562	1.000000	
	0.00000	0 045400	0 667666	
total sulfur dioxide	0.203028	0.047400	0.667666	
density	0.355283	0.200632	-0.021946	
density	0.333203	0.200032	-0.021940	
рН	-0.085652	-0.265026	0.070377	
E	0.00002	11200020	3.3.3377	
sulphates	0.005527	0.371260	0.051658	
-				
alcohol	0.042075	-0.221141	-0.069408	
quality	0.013732	-0.128907	-0.050656	

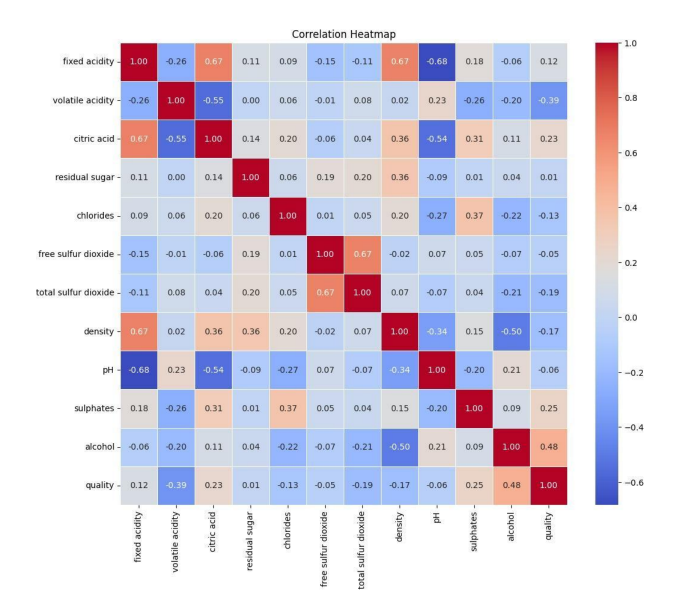
```
total sulfur dioxide density pH
sulphates \
                                -0.113181 0.668047 -0.682978
fixed acidity
0.183006
volatile acidity
                                 0.076470 0.022026 0.234937 -
0.260987
citric acid
                                 0.035533 0.364947 -0.541904
0.312770
                                 0.203028 0.355283 -0.085652
residual sugar
0.005527
chlorides
                                 0.047400 0.200632 -0.265026
0.371260
free sulfur dioxide
                                 0.667666 -0.021946 0.070377
0.051658
total sulfur dioxide
                                 1.000000 0.071269 -0.066495
0.042947
                                 0.071269 1.000000 -0.341699
density
0.148506
                                 -0.066495 -0.341699 1.000000 -
Нф
0.196648
sulphates
                                  0.042947 0.148506 -0.196648
1.000000
                                 -0.205654 -0.496180 0.205633
alcohol
0.093595
                                 -0.185100 -0.174919 -0.057731
quality
0.251397
                       alcohol quality
                     -0.061668 0.124052
fixed acidity
                   -0.202288 -0.390558
volatile acidity
citric acid
                     0.109903 0.226373
residual sugar
                     0.042075 0.013732
chlorides
                    -0.221141 -0.128907
free sulfur dioxide -0.069408 -0.050656
total sulfur dioxide -0.205654 -0.185100
density
                    -0.496180 -0.174919
Нф
                     0.205633 - 0.057731
                     0.093595 0.251397
sulphates
                     1.000000 0.476166
alcohol
quality
                    0.476166 1.000000
data.dropna(inplace=True)
scaler standard = StandardScaler()
data standardized = scaler standard.fit transform(data)
scaler normal = MinMaxScaler()
data normalized = scaler normal.fit transform(data)
```

```
data standardized = pd.DataFrame(data standardized,
columns=data.columns)
data normalized = pd.DataFrame(data normalized, columns=data.columns)
data standardized.head()
   fixed acidity volatile acidity citric acid residual sugar
chlorides \
      -0.528360
                         0.961877 -1.391472
                                                    -0.453218 -
0.243707
      -0.298547
                         1.967442 -1.391472
                                                      0.043416
0.223875
   -0.298547
                         1.297065 -1.186070
                                                     -0.169427
0.096353
      1.654856
                        -1.384443 1.484154
                                                     -0.453218 -
0.264960
      -0.528360
                         0.961877 - 1.391472
                                                     -0.453218 -
0.243707
   free sulfur dioxide total sulfur dioxide density pH
sulphates \
             -0.466193
                                  -0.379133 0.558274 1.288643 -
0.579207
                                   0.624363 0.028261 -0.719933
              0.872638
0.128950
                                   0.229047 0.134264 -0.331177 -
             -0.083669
0.048089
              0.107592
                                   0.411500 0.664277 -0.979104 -
0.461180
                                  -0.379133 0.558274 1.288643 -
             -0.466193
0.579207
   alcohol quality
0 -0.960246 -0.787823
1 -0.584777 -0.787823
2 -0.584777 -0.787823
3 -0.584777 0.450848
4 -0.960246 -0.787823
data normalized.head()
   fixed acidity volatile acidity citric acid residual sugar
chlorides \
       0.247788
                         0.397260
                                          0.00
                                                      0.068493
0.106845
       0.283186
                                          0.00
                                                      0.116438
                         0.520548
0.143573
      0.283186
                         0.438356
                                          0.04
                                                      0.095890
0.133556
      0.584071
                         0.109589
                                          0.56
                                                      0.068493
```

```
0.105175
4 0.247788
                        0.397260
                                        0.00 0.068493
0.106845
  free sulfur dioxide total sulfur dioxide density
sulphates \
             0.140845
                                  0.098940 0.567548 0.606299
0
0.137725
                                  0.215548 0.494126 0.362205
             0.338028
0.209581
                                  0.169611 0.508811 0.409449
             0.197183
0.191617
                                  0.190813 0.582232 0.330709
             0.225352
0.149701
                                  0.098940 0.567548 0.606299
             0.140845
0.137725
  alcohol quality
0 0.153846
                0.4
1 0.215385
                0.4
2 0.215385
                0.4
3 0.215385
                0.6
4 0.153846
                0.4
```

EDA

```
plt.figure(figsize=(12, 10))
sns.heatmap(data.corr(), annot=True, cmap='coolwarm', fmt='.2f',
linewidths=0.5)
plt.title('Correlation Heatmap')
plt.show()
```



Splitting the data into testing and training

```
# distributing the dataset into two components X and Y
X_red = data.iloc[:, 0:11].values
y_red = data.iloc[:, 11].values

X_train_red, X_test_red, y_train_red, y_test_red =
train_test_split(X_red, y_red, test_size=0.2, random_state=0)
```

Feature engineering

```
# performing preprocessing part
from sklearn.preprocessing import StandardScaler
sc = StandardScaler()

X_train_red = sc.fit_transform(X_train_red)
X_test_red = sc.transform(X_test_red)
```

Building PCA model

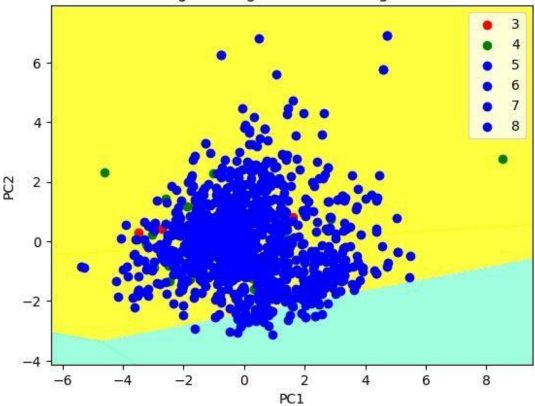
```
from sklearn.decomposition import PCA
PCa = PCA (n components = 2)
X train red = PCa.fit transform(X train red)
X test red = PCa.transform(X test red)
explained variance = PCa.explained variance ratio
from sklearn.linear model import LogisticRegression
classifier = LogisticRegression(random state = 0)
classifier.fit(X train red, y train red)
LogisticRegression(random state=0)
y pred red = classifier.predict(X test red)
from sklearn.metrics import confusion matrix
cm = confusion matrix(y test red, y pred red)
print(cm)
accuracy score (y test red, y pred red)
[[0 0 0 2 0 0]
 [0 0 4 7 0 0]
 [ 0 0 89 45 1 0]
 [ 0 0 55 81 6 0]
 [ 0 0 4 21 2 0]
[0 0 0 2 1 0]
0.5375
```

Visualisation of PCA model

```
# result through scatter plot
from matplotlib.colors import ListedColormap
```

```
X set, y set = X train red, y train red
X1, X2 = np.meshgrid(np.arange(start = X set[:, 0].min() - 1,
                     stop = X set[:, 0].max() + 1, step = 0.01),
                     np.arange(start = X \text{ set}[:, 1].min() - 1,
                     stop = X set[:, 1].max() + 1, step = 0.01))
plt.contourf(X1, X2, classifier.predict(np.array([X1.ravel(),
             X2.ravel()).T).reshape(X1.shape), alpha = 0.75,
             cmap = ListedColormap(('yellow', 'white', 'aquamarine')))
plt.xlim(X1.min(), X1.max())
plt.ylim(X2.min(), X2.max())
for i, j in enumerate(np.unique(y set)):
   plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
                c = ListedColormap(('red', 'green', 'blue'))(i), label
= j)
plt.title('Logistic Regression (Training set)')
plt.xlabel('PC1') # for Xlabel plt.ylabel('PC2')
# for Ylabel
plt.legend() # to show legend
# show scatter plot
plt.show()
C:\Users\nithi\AppData\Local\Temp\ipykernel 4776\4008791166.py:18:
UserWarning: *c* argument looks like a single numeric RGB or RGBA
sequence, which should be avoided as value-mapping will have
precedence in case its length matches with *x* & *y*. Please use the
*color* keyword-argument or provide a 2D array with a single row if
you intend to specify the same RGB or RGBA value for all points.
 plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
```

Logistic Regression (Training set)



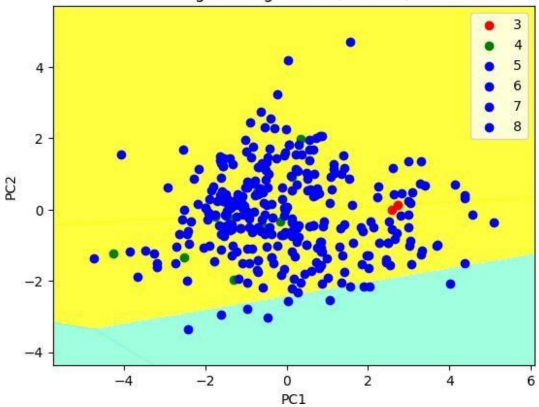
```
# Visualising the Test set results through scatter plot
from matplotlib.colors import ListedColormap
X set, y set = X test red, y test red
X1, X2 = np.meshgrid(np.arange(start = X_set[:, 0].min() - 1,
                     stop = X set[:, 0].max() + 1, step = 0.01),
                     np.arange(start = X set[:, 1].min() - 1,
                     stop = X set[:, 1].max() + 1, step = 0.01))
plt.contourf(X1, X2, classifier.predict(np.array([X1.ravel(),
             X2.ravel()]).T).reshape(X1.shape), alpha = 0.75,
             cmap = ListedColormap(('yellow', 'white', 'aquamarine')))
plt.xlim(X1.min(), X1.max())
plt.ylim(X2.min(), X2.max())
for i, j in enumerate(np.unique(y set)):
    plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
                c = ListedColormap(('red', 'green', 'blue'))(i), label
= j)
```

```
# title for scatter plot
plt.title('Logistic Regression (Test set)')
plt.xlabel('PC1') # for Xlabel
plt.ylabel('PC2') # for Ylabel
plt.legend()

# show scatter plot
plt.show()

C:\Users\nithi\AppData\Local\Temp\ipykernel_4776\3957206111.py:19:
UserWarning: *c* argument looks like a single numeric RGB or RGBA
sequence, which should be avoided as value-mapping will have
precedence in case its length matches with *x* & *y*. Please use the
*color* keyword-argument or provide a 2D array with a single row if
you intend to specify the same RGB or RGBA value for all points.
   plt.scatter(X_set[y_set == j, 0], X_set[y_set == j, 1],
```

Logistic Regression (Test set)



Building LDA model

 $\label{lem:constraint} from \ sklearn. discriminant_analysis \ import \ Linear Discriminant Analysis \ as \ LDA$

```
lda = LDA(n components = 2)
X train red = lda.fit transform(X train red, y train red)
X test red = lda.transform(X test red)
from sklearn.linear model import LogisticRegression
from sklearn.metrics import accuracy score
classifier = LogisticRegression(random state = 0)
classifier.fit(X train red, y train red)
LogisticRegression(random state=0)
y pred = classifier.predict(X test red)
print(y pred)
[5\ 5\ 6\ 6\ 7\ 5\ 6\ 5\ 6\ 6\ 6\ 6\ 5\ 5\ 7\ 6\ 6\ 5\ 6\ 6\ 6\ 5\ 5\ 5\ 5\ 5\ 6\ 6\ 6\ 5\ 5\ 5\ 5
5 6 5 6 5 6 6 6 6 5 6 5 5 5 6 6 5 6 6 6 6 6 5 6 5 6 6 6 5 6 5 6 6
6 7
5 7
5 6
5 6
5 5 5 5 5 6 6 5 6 5 5 5 5 5 5 5 5 6 5 7 6 6 6 7]
accuracy = accuracy score(y test red, y pred)
print("Accuracy of LDA model:", accuracy)
Accuracy of LDA model: 0.5375
```

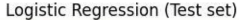
Visualisation of LDA model

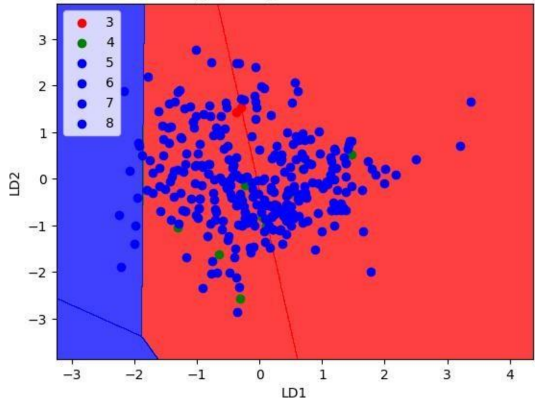
```
from matplotlib.colors import ListedColormap
X_set, y_set = X_test_red, y_test_red

X1, X2 = np.meshgrid(np.arange(start = X_set[:, 0].min() - 1, stop =
X_set[:, 0].max() + 1, step = 0.01), np.arange(start = X_set[:,
1].min() - 1, stop = X_set[:, 1].max() + 1, step = 0.01))

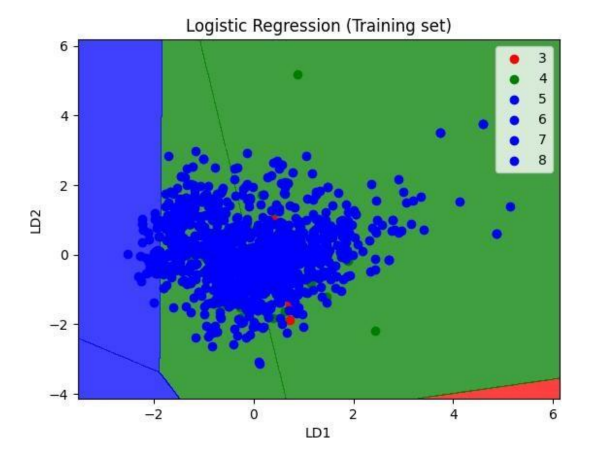
plt.contourf(X1, X2, classifier.predict(np.array([X1.ravel(), X2.ravel()]).T).reshape(X1.shape), alpha = 0.75, cmap = ListedColormap(('red', 'green', 'blue')))
```

```
plt.xlim(X1.min(), X1.max())
plt.ylim(X2.min(), X2.max())
for i, j in enumerate(np.unique(y set)):
    plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
                c = ListedColormap(('red', 'green', 'blue'))(i), label
= \dot{j})
plt.title('Logistic Regression (Test set)')
plt.xlabel('LD1')
plt.ylabel('LD2')
plt.legend()
plt.show()
C:\Users\nithi\AppData\Local\Temp\ipykernel 4776\531035640.py:12:
UserWarning: *c* argument looks like a single numeric RGB or RGBA
sequence, which should be avoided as value-mapping will have
precedence in case its length matches with *x* & *y*. Please use the
*color* keyword-argument or provide a 2D array with a single row if
you intend to specify the same RGB or RGBA value for all points.
  plt.scatter(X set[y set == j, 0], X_set[y_set == j, 1],
```





```
from matplotlib.colors import ListedColormap
X set, y set = X train red, y train red
X1, X2 = np.meshgrid(np.arange(start = X set[:, 0].min() - 1, stop =
X = [:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0].max() + 1, step = 0.01), np.arange(start = X set[:, 0
1].min() - 1, stop = X set[:, 1].max() + 1, step = 0.01))
plt.contourf(X1, X2, classifier.predict(np.array([X1.ravel(), X2.ravel()]
).T).reshape(X1.shape), alpha = 0.75, cmap = ListedColormap(('red',
'green', 'blue')))
plt.xlim(X1.min(), X1.max())
plt.ylim(X2.min(), X2.max())
for i, j in enumerate(np.unique(y set)):
          plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
                                           c = ListedColormap(('red', 'green', 'blue'))(i), label
= j)
plt.title('Logistic Regression (Training set)')
plt.xlabel('LD1')
plt.ylabel('LD2')
plt.legend()
plt.show()
C:\Users\nithi\AppData\Local\Temp\ipykernel 4776\171321940.py:12:
UserWarning: *c* argument looks like a single numeric RGB or RGBA
sequence, which should be avoided as value-mapping will have
precedence in case its length matches with *x* & *y*. Please use the
*color* keyword-argument or provide a 2D array with a single row if
you intend to specify the same RGB or RGBA value for all points.
    plt.scatter(X set[y set == j, 0], X set[y set == j, 1],
```



Inference

- 1. After applying PCA, you can analyze the principal components to understand which original features contribute the most to the variance in the data. You can also visualize the data in reduced dimensions to explore patterns or clusters.
- 2. After applying LDA, you can interpret the learned linear discriminants to understand how the classes are separated in the reduced-dimensional space. LDA provides insight into which features are most discriminative for class separation.

Learning Outcomes

- 1. Implementation of Pre-processing, EDA and feature selection.
- 2. Implementation of PCA nad LDA models and visualising it.
- 3. Displaying the confusion matrix.
- 4. Understanding the techniques of dimentionality reduction.

GITHUB LINK

https://github.com/KavinSiva13/ML-A4/tree/main/Assignment-9