# Classification Systems

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# March 22, 2019

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# 1 Classification Systems

In this practical, you are asked to compare the prediction error of:

- 1. The Naive Bayes Classifier
- 2. LDA
- 3. QDA
- 4. Nearest Shrunken Centroids Classifier

On the Breast Cancer dataset provided in the previous notebooks, and the Prostate cancer dataset attached. The details about this last dataset are found in the reference:

Singh, D., Febbo, P., Ross, K., Jackson, D., Manola, J., Ladd, C., Tamayo, P., Renshaw, A., D'Amico, A., Richie, J., Lander, E., Loda, M., Kantoff, P., Golub, T., & Sellers, W. (2002). Gene expression correlates of clinical prostate cancer behavior. Cancer Cell, 1, 203–209.

This dataset is in CSV format and the last column contains the class label. The task of interest is to discriminate between normal and tumor tissue samples.

Importantly:

Use a random split of 2 / 3 of the data for training and 1 / 3 for testing each classifier. Any hyper-parameter of each method should be tuned using a grid-search guided by an inner cross-validation procedure that uses only training data. To reduce the variance of the estimates, report average error results over 20 different partitions of the data into training and testing as described above. Submit a notebook showing the code and the results obtained. Give some comments about the results and respond to these questions:

What method performs best on each dataset? What method is more flexible? What method is more robust to over-fitting?

```
Script 1.0.1 (python)
import warnings
warnings.filterwarnings("ignore")
3 %matplotlib inline
4 import numpy as np
5 import pandas as pd
  import matplotlib.pyplot as plt
7 import matplotlib.lines as mlines
8 import matplotlib as mpl
9 from matplotlib import colors
import seaborn as sns; sns.set()
import scipy.stats as stats
12 import scipy as sp
13 from scipy import linalg
14 from sklearn.naive_bayes import GaussianNB
from sklearn.discriminant_analysis import LinearDiscriminantAnalysis
from sklearn.discriminant_analysis import QuadraticDiscriminantAnalysis
17 from sklearn.naive_bayes import GaussianNB
18 from sklearn.neighbors import NearestCentroid
19 from sklearn.decomposition import PCA
20 from sklearn.pipeline import Pipeline
21 from sklearn.model_selection import train_test_split, RepeatedStratifiedKFold, GridSearchCV
22 from sklearn import preprocessing
from sklearn.metrics import accuracy_score, make_scorer, confusion_matrix
```

#### 1.1 Methods

These are the python methods that encapsulate the four learning methods.

#### 1.1.1 Implementation details

#### **Quadratic Discriminant Analysis**

Before training the classifier we have chosen a good value for the corresponding regularization hyper-parameter with a grid-search guided by cross-validation.

The regularization parameter regularizes the covariance matrix estimate as

$$(1 - \lambda) \cdot \Sigma + \lambda \cdot I$$

#### **Nearest Centroids**

Before training the classifier we have chosen a good value for the shrinkage threshold hyper-parameter with a grid-search guided by cross-validation.

This procedure leads to a reduction in the number of features, by zeroing all deltas that exceed the threshold.

They take the form:

$$\mu_{kj}=m_j+\Delta_{kj}\,,$$

where  $\Delta_{ki}$  is the shrunken component

#### Selecting the best parameter value

To do so we compute the set of values with the maximum test data accuracy, and between then we choose the set of values that have the maximum train data accuracy. From this set we choose the lowest value.

```
Script 1.1.1 (python)
   def get_component_number(df_data, desired_variance=99.0, scaling=False):
2
       Obtain the number of components that explains a %desired_variance
3
       Args:
4
           df_data (dataframe): dataframe of features in cols and samples in rows
5
           desired_variance (float): desired explained variance
           scaling (boolean): True if pre-scaling is needed prior to compute PCA
7
       Returns:
8
           int: number of components to maintain to have a explained variance >=
9
       desired_variance
           float: variance explained for the number of components returned
10
11
           numpy array: cumulative variance by number of components retained
12
13
       if scaling:
           df_data_2 = preprocessing.StandardScaler().fit_transform(df_data)
14
15
       else:
           df_{data_2} = df_{data}
16
       # project the data into this new PCA space
17
       pca = PCA().fit(df_data_2)
18
19
       desired_variance = desired_variance/100.0
       explained_variance = np.cumsum(pca.explained_variance_ratio_)
20
       component_number = 0
21
       for cumulative_variance in explained_variance:
22
23
           component_number += 1
           if cumulative_variance >= desired_variance:
24
```

```
25
       return component_number, cumulative_variance, explained_variance
26
27
28
  def create_datasets_from_file(data_file, header, random_state, label_pos,
29
                                  label_value, features_ini, features_fin=None,
30
                                  with_dim_red=False, retained_variance=99.0):
31
       """Create training and test sets from file
32
33
           Args:
34
               data_file (string): Name of the data file (csv) of samples a features
35
               header (string): None or position of the header (pandas read_csv parameter)
36
               random_state (int): Seed for the random split (as needed for sklearn
37
       train_test_split)
               label_pos (int): Column of the labels in data_file
38
               label_value (int): Value of the label to asign internal '1' value
39
               features_ini (int): First column of features in data_file
40
               features_fin (int): Last column + 1 of features in data_file. If None, last
41
      column of file.
               with_dim_red (bool): If True, it performs a dimensionality reduction by PCA
42
               retained_variance (float): If dimensionality reduction, variance to retain
43
44
45
           Returns:
               (np.array): train set scaled
46
47
                (np.array): test set scaled
               (np.array): class labels for the train set
48
               (np.array): class labels for the test set
49
50
51
       data = pd.read_csv(data_file, header = header)
52
       if features_fin == None:
53
           X = data.values[:, features_ini:].astype(np.float)
54
       else:
55
           X = data.values[ :, features_ini:features_fin].astype(np.float)
56
       y = (data.values[:, label_pos] == label_value).astype(np.int)
57
58
       # Split dataset between training and test
59
       x_train, x_test, y_train, y_test = train_test_split(X, y,
60
                                                             test_size=1.0/3,
61
                                                              → random_state=random_state)
       # Data standardization
62
       scaler = preprocessing.StandardScaler().fit(x_train)
63
       x_train_scaled = scaler.transform(x_train)
64
       x_test_scaled = scaler.transform(x_test)
65
       # Check standardization
66
67
       for i in range (1, np.size(x_train_scaled,1)):
           assert round(np.var(x_train_scaled[:,0]),3) == round(np.var(x_train_scaled[:,i]),3),\
68
           "Warning: revise data standardization"
69
70
       if with_dim_red:
71
72
           desired_variance = retained_variance
73
           component_number, _, _ =\
```

```
get_component_number(x_train_scaled, desired_variance, scaling=None)
74
            print("Features reduced to", component_number)
75
76
            pca = PCA(n_components = component_number)
            pca.fit(x_train_scaled)
77
78
            x_train_scaled = pca.transform(x_train_scaled)
            x_test_scaled = pca.transform(x_test_scaled)
79
80
81
       return x_train_scaled, x_test_scaled, y_train, y_test
82
   def prediction_accuracy(x_train, x_test, y_train, y_test, method_func, method_param,
83
       param_value):
        """Estimate parameter given training and test sets:
84
85
            Args:
                x_train (np.array): train set
86
                x_{test} (np.array): test set
87
                y_train (np.array): class labels for the train set
88
                y\_test (np.array): class labels for the test set
89
                method_func (string) : name of the learning method
90
91
                param (string): name of learning method parameter
                param_value (float): value of parameter to try
92
            Returns:
93
                float: best parameter value to use in prediction
94
95
        11 11 11
96
97
       if method_param != "" :
            params = {method_param : param_value}
98
       else:
99
            params ={}
100
       method = globals()[method_func](**params)
101
102
        # Training
103
       method.fit(x_train, y_train)
104
105
        # Prediction
106
107
       y_pred = method.predict(x_test)
       conf = confusion_matrix(y_test, y_pred)
108
       TN = conf[0][0]
109
       TP = conf[1][1]
110
       FP = conf[0][1]
111
       FN = conf[1][0]
112
113
       print(conf)
       print('Predicion accuracy is: %f' % ((TP + TN) / (TN + TP + FP + FN)))
114
       print('True postive rate is: %f' % (TP / (TP + FN)))
115
       print('True negative rate is: %f\n' % (TN / (TN + FP)))
116
117
118
   def estimate_parameter(x_train, x_test, y_train, y_test, method_func, param, param_values):
        """Estimate parameter given training and test sets:
119
120
            Args:
                x_train (np.array): train set
121
                x_test (np.array): test set
122
123
                y_train (np.array): class labels for the train set
124
                y_test (np.array): class labels for the test set
```

```
method_func (string) : name of the learning method
125
                param (string): name of learning method parameter
126
                param_values (list of float): list of parameter values to try
127
            Returns:
128
                (float): best parameter value to use in prediction
129
130
        11 11 11
131
        # Pipeline for estimate the regularization parameter
       pipeline = Pipeline([ ('method', globals()[method_func]()) ])
133
134
        # Construct the grid the hyperparameter candidate shronk theshold
135
       param_grid = { 'method__' + param : param_values }
136
137
138
        # Evaluating
       skfold = RepeatedStratifiedKFold(n_splits=10, n_repeats=1, random_state=0)
139
       gridcv = GridSearchCV(pipeline, cv=skfold, n_jobs=1, param_grid=param_grid,\
140
                scoring=make_scorer(accuracy_score))
141
       result = gridcv.fit(x_train, y_train)
142
143
        # Accuracies
144
       accuracies = gridcv.cv_results_['mean_test_score']
145
       std_accuracies = gridcv.cv_results_['std_test_score']
146
147
       test_accuracies = np.ones(len(param_values))
148
149
       for i in range(len(param_values)):
150
           method_params = {param : param_values[ i ]}
151
           method = globals()[method_func](**method_params)
152
153
            method.fit(x_train, y_train)
            test_accuracies[ i ] = accuracy_score(method.predict(x_test), y_test)
154
155
       max_test_accuracy = max(test_accuracies)
156
157
       # Obtain best_param_value as max
158
       best_param_value = 0
159
       best_train_accuracy = 0
160
       for i in range(len(param_values)):
161
            if test_accuracies[ i ] == max_test_accuracy:
162
                if accuracies[i] > best_train_accuracy:
163
164
                    best_train_accuracy = accuracies[i]
165
                    best_param_value = param_values[i]
166
        #best_param_value = param_values[ np.argmax(accuracies) ]
167
        # Plot
168
       plt.figure(figsize=(15, 10))
169
170
       plt.grid()
171
       line1, = plt.plot(param_values, accuracies, 'o-', color="g")
       line2, = plt.plot(param_values, test_accuracies, 'x-', color="r")
172
       plt.fill_between(param_values, accuracies - std_accuracies / np.sqrt(10), \
173
            accuracies + std_accuracies / np.sqrt(10), alpha=0.1, color="g")
174
       plt.title("Different hyper-parameter " + param + " values for " + method_func)
175
176
       plt.xlabel('Hyper-parameter')
```

```
plt.xticks(np.round(np.array(param_values), 2))
177
        plt.ylabel('Classification Accuracy')
178
        plt.ylim((min(min(accuracies), min(test_accuracies)) - 0.1,
179
                  min(1.02, max(max(accuracies), max(test_accuracies)) + 0.1)))
180
181
        plt.xlim((min(param_values), max(param_values)))
182
        legend_handles = [ mlines.Line2D([], [], color='g', marker='o', \
183
                                   markersize=15, label='CV-estimate'), \
184
185
                        mlines.Line2D([], [], color='r', marker='x', \
                                   markersize=15, label='Test set estimate')]
186
        plt.legend(handles=legend_handles, loc = 3)
187
        plt.show()
188
189
190
        print("Best param value:", best_param_value)
        return best_param_value
191
192
   def learn_dataset(data_file, header, random_state, label_pos,
193
                      label_value, features_ini, features_fin=None,
194
                      with_dim_red=False, retained_variance=99.0):
195
        """Learn data sets from file, methods:
196
                1. The Naive Bayes Classifier
197
                2. LDA
198
                3. QDA
199
                4. Nearest Shrunken Centroids Classifier
200
201
                data_file (string): Name of the data file (csv) of samples a features
202
203
                header (string): None or position of the header (pandas read_csv parameter)
                random_state (int): Seed for the random split of sets (as needed for sklearn
204
        train_test_split)
                label_pos (int): Column of the labels in data_file
205
                label_value (int): Value of the label to asign internal '1' value
206
                features_ini (int): First column of features in data_file
207
                features_fin (int): Last column + 1 of features in data_file. If None, last
208
        column of file
                with_dim_red (bool): If True, it performs a dimensionality reduction by PCA
209
                retained_variance (float): If dimensionality reduction, variance to retai
210
211
212
213
        X_train_scaled, X_test_scaled, y_train, y_test = \
            create_datasets_from_file(data_file, header, random_state,
214
215
                                       label_pos, label_value, features_ini,

→ features_fin=features_fin,
                                       with_dim_red=with_dim_red,
216

→ retained_variance=retained_variance)
        print(X_train_scaled.shape)
217
218
        # Naive Bayes accuracy
219
        prediction_accuracy(X_train_scaled, X_test_scaled, y_train, y_test, "GaussianNB", "", "")
220
221
        # LDA accuracy
222
        prediction_accuracy(X_train_scaled, X_test_scaled, y_train, y_test,
223
        → "LinearDiscriminantAnalysis", "", "")
```

```
224
225
        # QDA estimate reg parameter
226
       param_values = np.linspace(0, 1, 10).tolist()
227
       best_param_value = estimate_parameter(X_train_scaled, X_test_scaled, y_train, y_test,\
228
                           "QuadraticDiscriminantAnalysis", "reg_param", param_values)
229
       # QDA accuracy
       # Best parameter reg value according CV estimate
230
       prediction_accuracy(X_train_scaled, X_test_scaled, y_train, y_test, \
231
                            "QuadraticDiscriminantAnalysis", "reg_param", best_param_value)
232
233
       # Centroids
234
235
       # Best parameter shrink_threshold value according CV estimate
       param_values = np.linspace(0, 8, 20).tolist()
236
       best_param_value = estimate_parameter(X_train_scaled, X_test_scaled, y_train, y_test,\
237
                           "NearestCentroid", "shrink_threshold", param_values)
238
        # Centroids accuracy
239
       prediction_accuracy(X_train_scaled, X_test_scaled, y_train, y_test, "NearestCentroid",
240
        → "shrink_threshold", best_param_value)
```

#### 1.2 Breast cancer

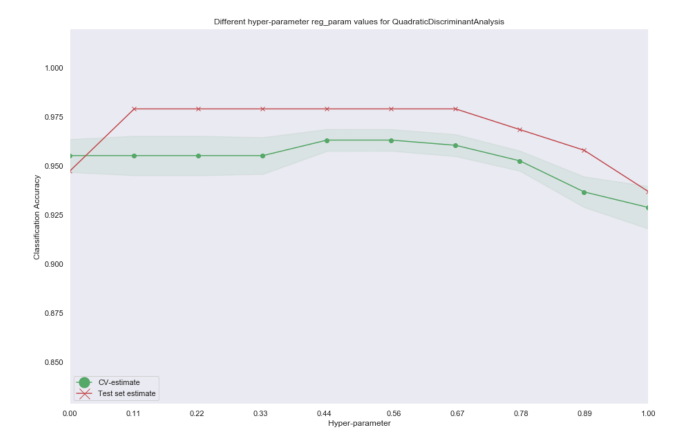
```
Script 1.2.1 (python)

1  # Breast Cancer
2  data_file = './data/wdbc.csv'
3  learn_dataset(data_file, None, 1, 1, "B", 2, features_fin = None, with_dim_red = False)
```

```
Output

(379, 30)
[[ 61     5]
     [ 7 117]]
Predicion accuracy is: 0.936842
True postive rate is: 0.943548
True negative rate is: 0.924242

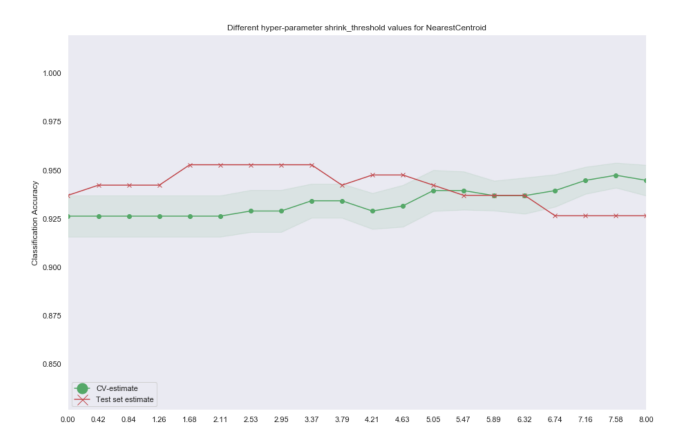
[[ 60     6]
     [ 1 123]]
Predicion accuracy is: 0.963158
True postive rate is: 0.991935
True negative rate is: 0.909091
```



## Output

[[ 62 4] [ 0 124]]

Predicion accuracy is: 0.978947 True postive rate is: 1.000000 True negative rate is: 0.939394



3.37

3.79 4.21

Hyper-parameter

4.63

5.05 5.47 7.16

### Output

Best param value: 3.3684210526315788

1.68

[[ 60 6] [ 3 121]]

Predicion accuracy is: 0.952632 True postive rate is: 0.975806 True negative rate is: 0.909091

#### 1.3 Prostate cancer

# Script 1.3.1 (python) 1 # Prostate Cancer data\_file = './data/prostate.csv' 3 learn\_dataset(data\_file, 0, 1, -1, 1, 0, -1, with\_dim\_red = True)

### Output

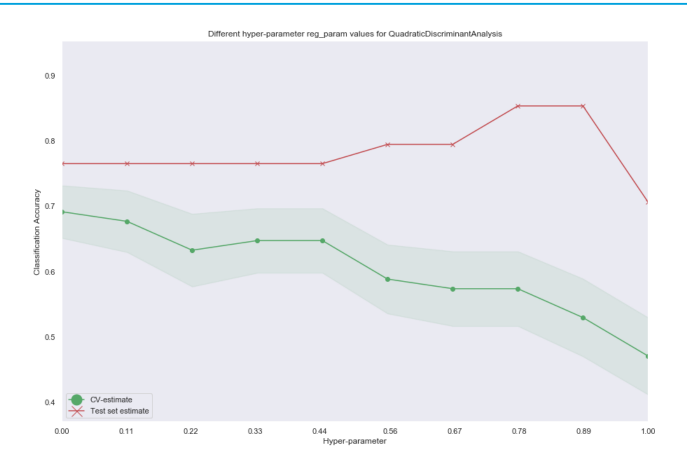
Features reduced to 63 (68, 63)

[[11 5] [ 1 17]]

Predicion accuracy is: 0.823529 True postive rate is: 0.944444 True negative rate is: 0.687500

[[16 0] [ 3 15]]

Predicion accuracy is: 0.911765 True postive rate is: 0.833333 True negative rate is: 1.000000

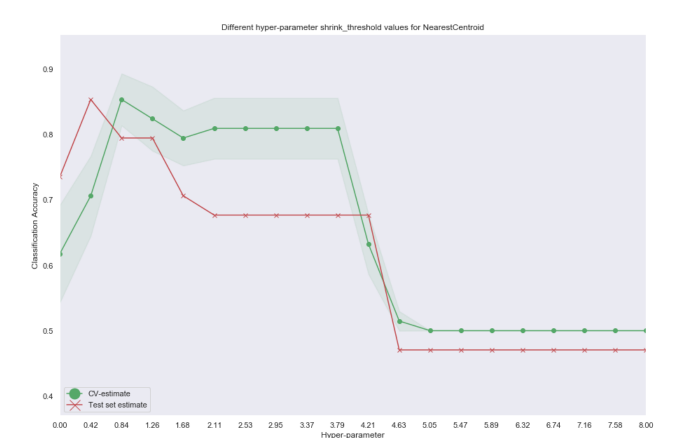


### Output

Best param value: 0.777777777777777

[[12 4] [ 1 17]]

Predicion accuracy is: 0.852941 True postive rate is: 0.944444 True negative rate is: 0.750000



### Output

Best param value: 0.42105263157894735

[[14 2] [ 3 15]]

Predicion accuracy is: 0.852941 True postive rate is: 0.833333 True negative rate is: 0.875000

### 1.4 Conclusions

We observe that **QDA** performs very poorly in the prostate dataset, given the high dimensionality of this dataset, which do not ease the accurate computation of the covariance matrices. Perhaps if we perform previously a dimensionality reduction by PCA, we'll improve this result.

**NSC** performs in this case much better due to the reduced number of parameters and the feature selection properties of this classifier and more consistently between both cases (prostate and breast).