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April 27, 2022

1 Lab: Hyper-Parameter Optimization with PCA

PCA is often applied as a pre-processing step with classifiers. When using PCA in this manner, one must select the number of PC components to use along with parameters in classifier. In this lab, we will demonstrate how to performing this *hyper-parameter optimization*. In doing the lab, you will learn to:

- Combine PCA with data scaling.
- Compute and visualize PC components
- Select the number of PCs with K-fold cross validation
- Implement the multi-stage classifier pipeline in sklearn
- Perform automatic parameter search using GridSearchCV in combination with a pipeline.

We first download the basic packages.

```
[2]: import numpy as np
import matplotlib
import matplotlib.pyplot as plt
```

1.1 Downloading the Data

We will use a very simple wine dataset, commonly used in teaching machine learning class. The problem is to classify the type of red wine from features of the wine such as the alchohol and other chemical components. There are three possible wine types.

```
[1]: from sklearn.datasets import load_wine
  from sklearn.model_selection import KFold
  data = load_wine()

# TODO 1: print the features names in data.feature_names and data.target_names
  print('features names: '+ str(data.feature_names))
  print('target_names: '+str(data.target_names))

features names: ['alcohol', 'malic_acid', 'ash', 'alcalinity_of_ash',
  'magnesium', 'total_phenols', 'flavanoids', 'nonflavanoid_phenols',
  'proanthocyanins', 'color_intensity', 'hue', 'od280/od315_of_diluted_wines',
  'proline']
  target_names: ['class_0' 'class_1' 'class_2']
```

Get the data matrix X from data.data and the target values y from data.target. Print the number of samples, number of features and number of classes.

1.2 Perform PCA for Visualization

Before performing PCA, you should scale the data matrix to remove the mean and normalize the variance of the different components. For this purpose, create a StandardScaling object scaling. Then fit the scaling with the entire data X. Transform the data and let Xs be the scaled data.

```
[5]: from sklearn.preprocessing import StandardScaler

# TODO 3:
# scaling = ...
# scaling.fit(...)
# Xs = ...
scaling = StandardScaler(with_mean=True, with_std=True)
scaling.fit(X)
Xs = scaling.transform(X)
```

Now, fit a PCA on the scaled data matrix Xs. You can use the sklearn PCA method. In order that we can visualize the results set n_components=2. Select svd_solver='randomized' and whiten=True. Use the pca.transform method to find, Z, the coefficients of Xs in the PCA basis.

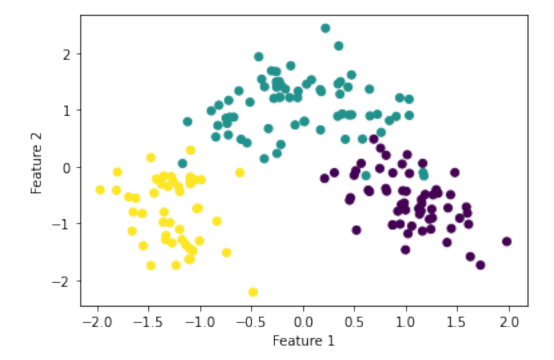
```
[6]: from sklearn.decomposition import PCA

# TODO 4:
# ncomp = 2
# pca = PCA(...)
# Z = ...
# Construct the PCA object
ncomp = 2
pca = PCA(n_components=ncomp,
svd_solver='randomized', whiten=True)
pca.fit(Xs)
Z = pca.transform(Xs)
```

In the transformed basis, each data sample is represented by a two dimensional vector, Z[i,0], Z[i,1]. Plot a scatter plot of the transformed data. Use different marker colors for the different classes. If you did everything, you should see that the classes are quite well separated with even two PCs.

```
[7]: # TODO 5:
    # plt.scatter(...)
    plt.scatter(Z[:,0],Z[:,1],c=y)
    plt.xlabel('Feature 1')
    plt.ylabel('Feature 2')
```

[7]: Text(0, 0.5, 'Feature 2')



Now, refit the scaled data Xs using n_components=nfeatures where nfeatures is the number of features. This is the maximum number of PCs. Get the singular values from pca.singular_values_ and plot the portion of variation as a function of the number of PCs. The PoV for using n PCs is:

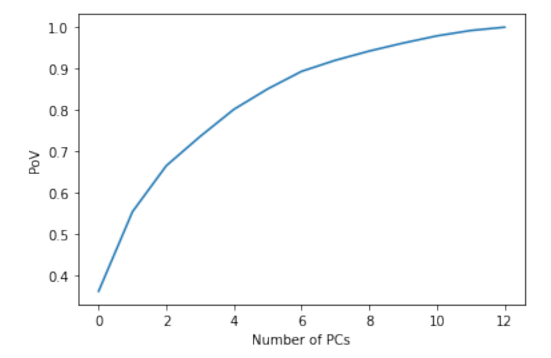
```
PoV[n] = \sum_{i=0}^{n-1} s[i]**2 / \sum_{i=0}^{d-1} s[i]**2
```

where s[i] is the i-th singular value and d is the number of features. You should see that the 4 PCs contains more than 70% of the variance.

```
[8]: # TODO 6:
    nfeatures = X.shape[1]
    pca1 = PCA(n_components=nfeatures,
```

```
svd_solver='randomized', whiten=True)
Z = pca1.fit_transform(Xs)
S=pca1.singular_values_
Ssquare=S**2
denom=np.sum(Ssquare)
numerator=np.cumsum(Ssquare)
PoV=numerator/denom
plt.plot(PoV)
plt.xlabel('Number of PCs')
plt.ylabel('PoV')
```

[8]: Text(0, 0.5, 'PoV')



1.3 Using PCA with Classification

We will now use data scaling and PCA as a pre-processing step for logistic classification. The number of PCs to use can be found with cross-validation. Complete the code below which tries different number of PCs components to use and measures the test accuracy for each value.

```
[9]: from sklearn.preprocessing import StandardScaler
  from sklearn.linear_model import LogisticRegression
  nfold = 5

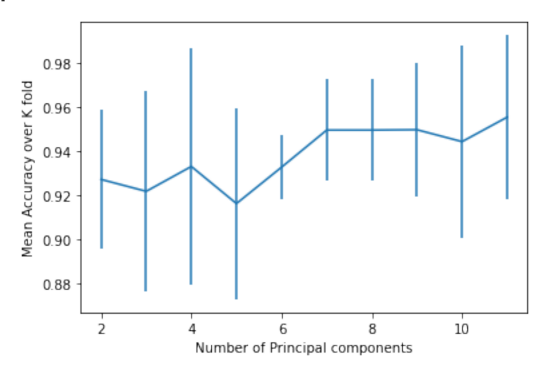
# Create a K-fold object
  kf = KFold(n_splits=nfold)
```

```
kf.get_n_splits(X)
# Number of PCs to try
ncomp_test = np.arange(2,12)
num_nc = len(ncomp_test)
# Accuracy: acc[icomp,ifold] is test accuracy when using `ncomp =_
→ncomp_test[icomp] in fold ifold.
acc = np.zeros((num_nc,nfold))
# ========
# TODO 7: (combined all)
# Loop over number of components to test
for icomp, ncomp in enumerate(ncomp_test):
    # Look over the folds
   for ifold, I in enumerate(kf.split(X)):
        Itr, Its = I
       Xtr = X[Itr]
       ytr = y[Itr]
       Xts = X[Its]
       yts = y[Its]
        scalingtr = StandardScaler(with_mean=True, with_std=True)
        scalingtr.fit(Xtr)
       Xtrs = scalingtr.transform(Xtr)
        pca = PCA(n_components=ncomp, svd_solver='randomized', whiten=True)
       pca.fit(Xtrs)
        Ztr = pca.transform(Xtrs)
        logreg = LogisticRegression(multi_class='auto',solver='lbfgs')
       logreg.fit(Ztr,ytr)
       Xtss = scalingtr.transform(Xts)
        Zts = pca.transform(Xtss)
        yhat=logreg.predict(Zts)
        acc[icomp, ifold] = np.mean(yhat == yts)
```

Use the plt.errorbar function to plot the mean accuracy with error bars corresponding to the standard error of the accuracy as a function of the number of components. Find the optimal number of PCs to use according to the normal rule and one SE rule. If you did it correctly, you should get an accuracy of around 96%.

```
# TODO:
         Optimal order with the normal rule
         Optimal order with one SE rule
acc_mean = np.mean(acc,axis=1)
acc_std = np.std(acc,axis=1)*np.sqrt(nfold)/np.sqrt(nfold-1)
plt.errorbar(ncomp_test, acc_mean, yerr=acc_std, fmt='-')
plt.xlabel('Number of Principal components')
plt.ylabel('Mean Accuracy over K fold')
imax = np.argmax(acc_mean)
print("The optimal number of PCs with the normal rule is{0:d}".
→format(ncomp_test[imax]))
acc_tgt = acc_mean[imax] - acc_std[imax]
I = np.where(acc_mean >= acc_tgt)[0]
iopt = I[0]
print("The optimal number of PC's with the one SE rule is {0:d}".
 →format(ncomp_test[iopt]))
```

The optimal number of PCs with the normal rule is 11 The optimal number of PC's with the one SE rule is 2



1.4 Hyper-Parameter Optimization with GridCV.

We will now try a more complex classifier – a support vector classifier with a radial basis function. When we use such a classifer, there will be a number of parameters to tune. When the number of parameters to tune becomes large, writing a loop over multiple parameters as we did above becomes

cumbersome. The sklearn package has a very nice routine, GridSearchCV to perform this sort of parameter search.

Before, we do this we need to create an estimator Pipeline. An estimator pipeline is a sequence of transformations followed by an estimator that will operate on the transformed data. Create the following pipeline:

- Create a StandardScaler() object called scaler for the first transformation
- Create a PCA() object called pca for the second transformation
- Create a SVC() object called svc for the final SVM classifier. Set the parameter kernel='rbf'.

Once you have the three steps defined, you can create the pipeline with the command:

```
pipe = Pipeline(steps=[('scaler', scaler), ('pca', pca), ('svc', svc)])
```

```
[13]: from sklearn.pipeline import Pipeline
  from sklearn.model_selection import GridSearchCV
  from sklearn.svm import SVC

# TODO 9:
# scaler = ...
# pca = ...
# svc = ...
# pipe = Pipeline(steps=[('scaler', scaler), ('pca', pca), ('svc', svc)])
  scaler = StandardScaler()
  pca = PCA()
  svc = SVC(kernel='rbf')
  pipe = Pipeline(steps=[('scaler', scaler), ('pca', pca), ('svc', svc)])
```

We next define all the parameters that we want to search over. Define the following arrays:

- ncomp_test: values from 3 to 10 representing number of PCs to test
- C_test: values of C in the SVC to test. Use 10^{-2}, 10^{-1}, ..., 10^{3}
- gam_test: values of gamma in the SVC to test. Use 10^{-3}, 10^{-2}, ..., 10^{1}

```
[27]: # TODO 10:
    # ncomp_test = ...
    # c_test = ...
    # gam_test = ...
    ncomp_test = np.arange(3,11)
    num_nc = len(ncomp_test)
    c_test = np.logspace(-2, 3, num = 6, base = 10)
    gam_test = np.logspace(-3, 1, num = 5, base=10)
```

Next, we create a dictionary params of the form:

```
params = {'pca_n_components': ncomp_test, 'svc_C' : c_test, ...}
```

Each key in the dictionary is the of the form estimator__param and the value is the values to be tested.

Finally, an object estimator = GridSearchCV(...) from pipe and params. Set cv=5, train_score=True and iid=False. Fit the estimator from the data X,y. Then the estimator will perform the cross-validation over all the parameters. This may take a minute since we are search over so many parameters.

```
[29]: # TODO 12:
    # estimator = GridSearchCV(...)
    # estimator.fit(...)
    estimator = GridSearchCV(pipe,params,cv=5, return_train_score=True)
    estimator.fit(X,y)
```

Print the best test score and best parameters. They are fields in **estimator**. If you did it correctly, it should be a little higher than the logistic regression (about 0.97 to 0.98 accuracy).

```
[30]: # TODO 13:
    print('The best test score is '+format(estimator.best_score_,".4f"))
    print('The best parameters are '+str(estimator.best_params_))
```

The best test score is 0.9778

The best parameters are {'pca_n_components': 5, 'svc_C': 1.0, 'svc_gamma': 0.1}

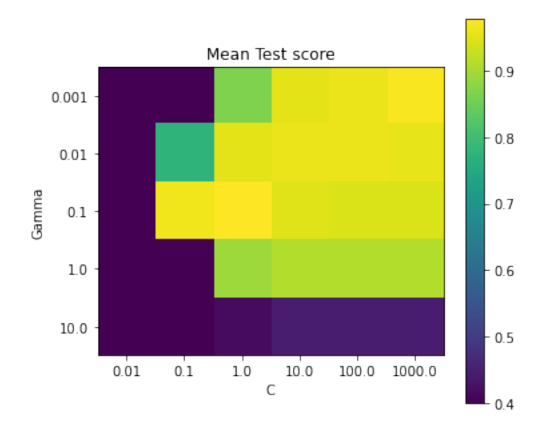
Finally, you can get the test score for all the parameter choices from

```
test_score = estimator.cv_results_['mean_test_score']
```

Use the imshow command to plot the mean test score over gamma and C for the value n_components=5.

```
[31]: # TODO 14:
    test_score = estimator.cv_results_['mean_test_score']
    print(test_score.shape)
    pc_n=estimator.cv_results_['param_pca__n_components'].data.astype(int)
    tsc=estimator.cv_results_['mean_test_score'][np.where(pc_n==5)]
```

```
print(estimator.cv_results_['param_svc_C'][np.where(pc_n==5)])
print(estimator.cv_results_['param_svc__gamma'][np.where(pc_n==5)])
print(tsc.shape)
print(tsc)
tsc=tsc.reshape(len(c_test),len(gam_test))
print(tsc)
print(c_test.astype(object))
print(gam_test.astype(object))
plt.figure(figsize=(6, 5))
plt.subplots_adjust(left=.2, right=0.95, bottom=0.15, top=0.95)
plt.imshow(tsc.T,interpolation='nearest')
plt.xlabel('C')
plt.ylabel('Gamma')
plt.colorbar()
plt.yticks(np.arange(len(gam_test)), gam_test)
plt.xticks(np.arange(len(c_test)), c_test)
plt.title('Mean Test score')
plt.show()
(240.)
10.0 10.0 10.0 10.0 100.0 100.0 100.0 100.0 100.0 1000.0 1000.0 1000.0
1000.0 1000.07
[0.001 0.01 0.1 1.0 10.0 0.001 0.01 0.1 1.0 10.0 0.001 0.01 0.1 1.0 10.0
0.001 0.01 0.1 1.0 10.0 0.001 0.01 0.1 1.0 10.0 0.001 0.01 0.1 1.0 10.0]
(30,)
[0.39904762 0.39904762 0.39904762 0.39904762 0.39904762 0.39904762
0.77603175 0.96619048 0.39904762 0.39904762 0.86587302 0.95507937
0.97777778 0.89365079 0.41603175 0.95507937 0.96095238 0.95
0.91047619 0.44412698 0.96111111 0.96095238 0.94428571 0.91047619
0.44412698 0.97206349 0.95555556 0.94428571 0.91047619 0.44412698]
[[0.39904762 0.39904762 0.39904762 0.39904762 0.39904762]
[0.39904762 0.77603175 0.96619048 0.39904762 0.39904762]
[0.86587302 0.95507937 0.97777778 0.89365079 0.41603175]
[0.95507937 0.96095238 0.95
                                 0.91047619 0.44412698]
[0.96111111 0.96095238 0.94428571 0.91047619 0.44412698]
[0.97206349 0.95555556 0.94428571 0.91047619 0.44412698]]
[0.01 0.1 1.0 10.0 100.0 1000.0]
[0.001 0.01 0.1 1.0 10.0]
```



[]: