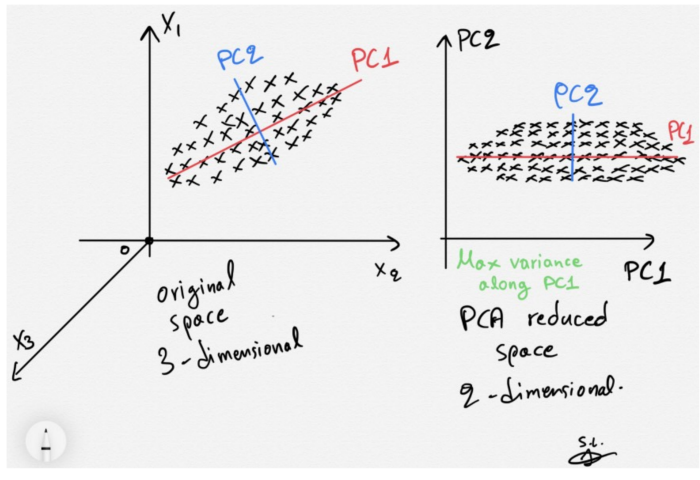
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PCA clearly explained —When, Why, How to use it and feature importance: A guide in Python

In this post I explain what PCA is, when and why to use it and how to implement it in Python using scikit-learn. Also, I explain how to get the feature importance after a PCA analysis.



Handmade sketch made by the author.

1. Introduction & Background

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Principal Components Analysis (PCA) is a well-known unsupervised dimensionality reduction technique that constructs relevant features/variables through linear (linear PCA) or non-linear (kernel PCA) combinations of the original variables (features). In this post, we will only focus on the famous and widely used linear PCA method.

The construction of relevant features is achieved by **linearly transforming correlated variables** into a smaller number of **uncorrelated** variables. This is done by **projecting** (dot product) the original data into the **reduced PCA space** using the eigenvectors of the covariance/correlation matrix aka the principal components (PCs).

The **resulting projected data** are essentially **linear combinations** of the **original** data **capturing most of the variance in the data** (Jolliffe 2002).

In summary, PCA is an **orthogonal transformation** of the data into a series of **uncorrelated** data living in the reduced PCA space such that the first component explains the most variance in the data with each subsequent component explaining less.

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2. When/Why to use PCA

• PCA technique is particularly useful in processing data where multicolinearity exists between the features/variables.



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Correlation vs

Collinearity vs





Multicollinearity





- PCA can be used when the dimensions of the input features are high (e.g. a lot of variables).
- PCA can be also used for **denoising** and **data compression**.

3. Core of the PCA method

Let \mathbf{x} be a matrix containing the original data with shape [n_samples, n_features].

Briefly, the PCA analysis consists of the following steps:

- First, the original input variables stored in x are z-scored such each original variable (column of x) has zero mean and unit standard deviation.
- The next step involves the construction and <u>eigendecomposition</u> of the **covariance** matrix **c**x= (1/n)x'x (in case of z-scored data the covariance is equal to the correlation matrix since the standard deviation of all features is 1).
- **Eigenvalues** are then **sorted** in a **decreasing** order representing decreasing variance in the data (the eigenvalues are equal to the variance I will prove this below using Python in Paragraph 6).
- Finally, the **projection** (transformation) of the **original normalized data** onto the **reduced PCA space** is obtained by **multiplying** (dot product) **the originally normalized data** by the **leading eigenvectors** of the covariance matrix i.e. the PCs.
- The new **reduced** PCA space **maximizes** the **variance** of the **original** data. To **visualize** the projected data as well as the contribution of the original variables, in a joint plot, we can use the **biplot**.

4. The maximum number of meaningful components

There is an upper bound of the meaningful components that can be extracted using PCA. This is related to the <u>rank</u> of the <u>covariance/correlation</u> matrix (cx). Having a data matrix x with shape [n_samples, n_features/n_variables], the <u>covariance/correlation</u> matrix would be [n_features, n_features] with <u>maximum rank</u> equal to min(n_samples, n_features).

Thus, we can have at **most** min(n_samples, n_features) **meaningful** PC **components/dimensions** due to the **maximum** <u>rank</u> of the covariance/correlation matrix.

5. Python example using scikit-learn and the Iris dataset

```
import numpy as np
import matplotlib.pyplot as plt
from sklearn import datasets
from sklearn.decomposition import PCA
import pandas as pd
from sklearn.preprocessing import StandardScaler
plt.style.use('ggplot')
# Load the data
iris = datasets.load_iris()
X = iris.data
y = iris.target
# Z-score the features
scaler = StandardScaler()
scaler.fit(X)
X = scaler.transform(X)
# The PCA model
pca = PCA(n_components=2) # estimate only 2 PCs
X_new = pca.fit_transform(X) # project the original data into the PCA
space
```

Let's plot the data before and **after** the **PCA** transform and also **color** code each point (sample) using the corresponding **class of the flower** (y) .

```
fig, axes = plt.subplots(1,2)
axes[0].scatter(X[:,0], X[:,1], c=y)
axes[0].set_xlabel('x1')
```

```
axes[0].set_ylabel('x2')
axes[0].set_title('Before PCA')

axes[1].scatter(X_new[:,0], X_new[:,1], c=y)
axes[1].set_xlabel('PC1')
axes[1].set_ylabel('PC2')
axes[1].set_title('After PCA')
plt.show()
```



PCA output of the above code.

We can see that in the PCA space, the **variance** is **maximized** along **PC1** (explains 73% of the variance) and **PC2** (explains 22% of the variance). Together, they explain 95%.

```
print(pca.explained_variance_ratio_)
# array([0.72962445, 0.22850762])
```

6. Proof of eigenvalues of original covariance matrix being equal to the variances of the reduced space

Mathematical formulation & proof

Assuming that the original input variables stored in x are z-scored such each original variable (column of x) has zero mean and unit standard deviation, we have:

- The covariance of the original space is: $\mathbf{C}_{\mathbf{X}} = \frac{1}{n}\mathbf{X}^T\mathbf{X}$
- PCA: Eigendecomposition of the covariance of the original space is:

$$\mathbf{C}_{\mathbf{X}} = \frac{1}{n} \mathbf{X}^T \mathbf{X} = \mathbf{U} \mathbf{\Lambda} \mathbf{U}^T$$

- Let $\mathbf{Y} = \mathbf{X}\mathbf{U}$ be the projected data (U stores the eigenvectors as columns)
- The covariance of the reduced PCA space is:

$$egin{aligned} \mathbf{C}_{\mathbf{Y}} &= rac{1}{n} \mathbf{Y}^T \mathbf{Y} \ &= rac{1}{n} (\mathbf{X} \mathbf{U}) (\mathbf{X} \mathbf{U})^T \ &= rac{1}{n} \mathbf{U}^T \mathbf{X}^T \mathbf{X} \mathbf{U} \ &= \mathbf{U}^T igg(rac{1}{n} \mathbf{X}^T \mathbf{X} igg) \mathbf{U} \ \mathbf{C}_{\mathbf{Y}} &= \mathbf{U}^T \mathbf{C}_{\mathbf{X}} \mathbf{U} \end{aligned}$$

-
$$\mathbf{C}_{\mathbf{Y}} = \mathbf{U}^T \mathbf{U} \mathbf{\Lambda} \mathbf{U}^T \mathbf{U} = \mathbf{\Lambda}$$
 which is diagonal.

Latex code written by the author.

A matrix above stores the **eigenvalues** of the **covariance** matrix of the original space/dataset.

Verify using Python

The **maximum variance proof** can be also seen by estimating the **covariance** matrix of the **reduced space**:

```
np.cov(X_new.T)

array([[2.93808505e+00, 4.83198016e-16],
        [4.83198016e-16, 9.20164904e-01]])
```

We observe that these values (on the diagonal we have the variances) are **equal** to the **actual eigenvalues** of the covariance stored in

```
pca.explained_variance_
array([2.93808505, 0.9201649 ])
```

7. Feature importance

pca.explained_variance_:

The **importance** of each **feature** is reflected by the **magnitude** of the **corresponding values in the eigenvectors** (higher magnitude — higher importance).

Let's find the most important features:

```
print(abs( pca.components_ ))
[[0.52106591 0.26934744 0.5804131 0.56485654]
[0.37741762 0.92329566 0.02449161 0.06694199]]
```

Here, pca.components_ has shape [n_components, n_features] Thus, by looking at the PC1 (first Principal Component) which is the first row

```
[[0.52106591 0.26934744 0.5804131 0.56485654]
```

we can conclude that **feature 1**, **3 and 4** are the **most important** for **PC1**. Similarly, we can state that **feature 2** and then **1** are the **most important** for **PC2**.

To sum up, we look at the absolute values of the eigenvectors' components corresponding to the **k** largest eigenvalues. In sklearn the components are sorted by explained variance. The larger they are these absolute values, the more a specific feature contributes to that principal component.

8. The biplot

The **biplot** is the best way to visualize **all-in-one** following a **PCA** analysis.

There is an implementation in $\underline{\mathbf{R}}$ but there is no standard implementation in **python** so I decided to write my **own function** for that:

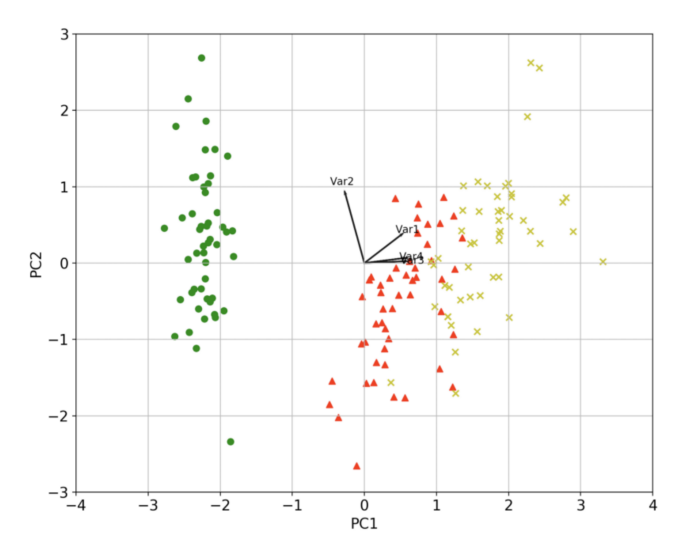
```
def biplot(score, coeff , y):
    Author: Serafeim Loukas, serafeim.loukas@epfl.ch
    Inputs:
       score: the projected data
       coeff: the eigenvectors (PCs)
      y: the class labels
    xs = score[:,0] # projection on PC1
    ys = score[:,1] # projection on PC2
    n = coeff.shape[0] # number of variables
    plt.figure(figsize=(10,8), dpi=100)
    classes = np.unique(y)
    colors = ['g','r','y']
    markers=['o','^','x']
    for s,l in enumerate(classes):
        plt.scatter(xs[y==1], ys[y==1], c = colors[s],
marker=markers[s]) # color based on group
    for i in range(n):
        #plot as arrows the variable scores (each variable has a score
for PC1 and one for PC2)
        plt.arrow(0, 0, coeff[i, 0], coeff[i, 1], color = 'k', alpha = 'k'
0.9, linestyle = '-', linewidth = 1.5, overhang=0.2)
        plt.text(coeff[i,0]* 1.15, coeff[i,1] * 1.15, "Var"+str(i+1),
color = 'k', ha = 'center', va = 'center', fontsize=10)
    plt.xlabel("PC{}".format(1), size=14)
    plt.ylabel("PC{}".format(2), size=14)
```

```
limx= int(xs.max()) + 1
limy= int(ys.max()) + 1
plt.xlim([-limx,limx])
plt.ylim([-limy,limy])
plt.grid()
plt.tick_params(axis='both', which='both', labelsize=14)
```

Call the function (make sure to run first the initial blocks of code where we load the iris data and perform the PCA analysis):

```
import matplotlib as mpl
mpl.rcParams.update(mpl.rcParamsDefault) # reset ggplot style

# Call the biplot function for only the first 2 PCs
biplot(X_new[:,0:2], np.transpose(pca.components_[0:2, :]), y)
plt.show()
```



The PCA biplot using my custom function.

We can again verify **visually** that **a**) the variance is maximized and **b**) that **feature 1**, **3 and 4** are the **most important** for **PC1**. Similarly, **feature 2** and then **1** are the **most important** for **PC2**.

Furthermore, **arrows** (variables/features) that point into the **same direction** indicate **correlation** between the variables that they represent whereas, the arrows heading in **opposite directions** indicate a **contrast** between the variables they represent.

Verify the above using **code**:

```
# Var 3 and Var 4 are extremely positively correlated
np.corrcoef(X[:,2], X[:,3])[1,0]
0.9628654314027957

# Var 2and Var 3 are negatively correlated
np.corrcoef(X[:,1], X[:,2])[1,0]
-0.42844010433054014
```

That's all folks! Hope you liked this article!

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References

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