r	Make sure to watch the video lecture and theory presentation for a full overview of PCA! Remember that PCA is just a transformation of your data and attempts to find out what features explain the most variance in your data. For example:
]:[import matplotlib.pyplot as plt import pandas as pd import numpy as np
	import seaborn as sns %matplotlib inline The Data
	Let's work with the cancer data set again since it had so many features. from sklearn.datasets import load_breast_cancer
]:[]:[<pre>cancer = load_breast_cancer()</pre>
	<pre>cancer.keys() dict_keys(['data', 'target', 'frame', 'target_names', 'DESCR', 'feature_names', 'filename']) print(cancer['DESCR'])</pre>
	breast_cancer_dataset: Breast cancer wisconsin (diagnostic) dataset
	Data Set Characteristics: :Number of Instances: 569 :Number of Attributes: 30 numeric, predictive attributes and the class
	:Attribute Information: - radius (mean of distances from center to points on the perimeter) - texture (standard deviation of gray-scale values) - perimeter - area - smoothness (local variation in radius lengths)
	- compactness (perimeter^2 / area - 1.0) - concavity (severity of concave portions of the contour) - concave points (number of concave portions of the contour) - symmetry - fractal dimension ("coastline approximation" - 1)
	The mean, standard error, and "worst" or largest (mean of the three worst/largest values) of these features were computed for each image, resulting in 30 features. For instance, field 0 is Mean Radius, field 10 is Radius SE, field 20 is Worst Radius. - class: - WDBC-Malignant
	- WDBC-Benign :Summary Statistics: ===================================
	radius (mean): 6.981 28.11 texture (mean): 9.71 39.28 perimeter (mean): 43.79 188.5 area (mean): 143.5 2501.0 smoothness (mean): 0.053 0.163
	compactness (mean): 0.019 0.345 concavity (mean): 0.0 0.427 concave points (mean): 0.0 0.201 symmetry (mean): 0.106 0.304 fractal dimension (mean): 0.05 0.097 radius (standard error): 0.112 2.873 texture (standard error): 0.36 4.885
	texture (standard error): 0.36 4.885 perimeter (standard error): 0.757 21.98 area (standard error): 6.802 542.2 smoothness (standard error): 0.002 0.031 compactness (standard error): 0.002 0.135 concavity (standard error): 0.0 0.396 concave points (standard error): 0.0 0.053
	symmetry (standard error): 0.008 0.079 fractal dimension (standard error): 0.001 0.03 radius (worst): 7.93 36.04 texture (worst): 12.02 49.54 perimeter (worst): 50.41 251.2 area (worst): 185.2 4254.0
	smoothness (worst): 0.071 0.223 compactness (worst): 0.027 1.058 concavity (worst): 0.0 1.252 concave points (worst): 0.0 0.291 symmetry (worst): 0.156 0.664 fractal dimension (worst): 0.055 0.208
	:Missing Attribute Values: None :Class Distribution: 212 - Malignant, 357 - Benign
	:Creator: Dr. William H. Wolberg, W. Nick Street, Olvi L. Mangasarian :Donor: Nick Street :Date: November, 1995 This is a copy of UCI ML Breast Cancer Wisconsin (Diagnostic) datasets.
	https://goo.gl/U2Uwz2 Features are computed from a digitized image of a fine needle aspirate (FNA) of a breast mass. They describe characteristics of the cell nuclei present in the image.
	Separating plane described above was obtained using Multisurface Method-Tree (MSM-T) [K. P. Bennett, "Decision Tree Construction Via Linear Programming." Proceedings of the 4th Midwest Artificial Intelligence and Cognitive Science Society, pp. 97-101, 1992], a classification method which uses linear programming to construct a decision tree. Relevant features
	were selected using an exhaustive search in the space of 1-4 features and 1-3 separating planes. The actual linear program used to obtain the separating plane in the 3-dimensional space is that described in: [K. P. Bennett and O. L. Mangasarian: "Robust Linear
	Programming Discrimination of Two Linearly Inseparable Sets", Optimization Methods and Software 1, 1992, 23-34]. This database is also available through the UW CS ftp server: ftp ftp.cs.wisc.edu cd math-prog/cpo-dataset/machine-learn/WDBC/
	 topic:: References W.N. Street, W.H. Wolberg and O.L. Mangasarian. Nuclear feature extraction for breast tumor diagnosis. IS&T/SPIE 1993 International Symposium on Electronic Imaging: Science and Technology, volume 1905, pages 861-870,
	San Jose, CA, 1993. - O.L. Mangasarian, W.N. Street and W.H. Wolberg. Breast cancer diagnosis and prognosis via linear programming. Operations Research, 43(4), pages 570-577, July-August 1995. - W.H. Wolberg, W.N. Street, and O.L. Mangasarian. Machine learning techniques to diagnose breast cancer from fine-needle aspirates. Cancer Letters 77 (1994)
]:[<pre>df = pd.DataFrame(cancer['data'],columns=cancer['feature_names']) #(['DESCR', 'data', 'feature_names', 'target_names', 'target'])</pre>
]:]:	mean mean mean mean mean mean mean mean
	0 17.99 10.38 122.80 1001.0 0.11840 0.27760 0.3001 0.14710 0.2419 0.07871 25.38 17.33 184.60 2019.0 0.1622 0.6656 0.7119 0.266 1 20.57 17.77 132.90 1326.0 0.08474 0.07864 0.0869 0.07017 0.1812 0.05667 24.99 23.41 158.80 1956.0 0.1238 0.1866 0.2416 0.18 2 19.69 21.25 130.00 1203.0 0.10960 0.15990 0.12790 0.2069 0.05999 23.57 25.53 152.50 1709.0 0.1444 0.4245 0.4504 0.24
	3 11.42 20.38 77.58 386.1 0.14250 0.28390 0.2414 0.10520 0.2597 0.09744 14.91 26.50 98.87 567.7 0.2098 0.8663 0.6869 0.25 4 20.29 14.34 135.10 1297.0 0.10030 0.13280 0.1980 0.10430 0.1809 0.05883 22.54 16.67 152.20 1575.0 0.1374 0.2050 0.4000 0.16 5 rows × 30 columns
A	PCA Visualization As we've noticed before it is difficult to visualize high dimensional data, we can use PCA to find the first two principal components, and visualize the data in this new, two-dimensional space, with a single scatter-plot. Before we do this though, we'll need to scale our data so that each feature has a single unit variance.
]:[single scatter-plot. Before we do this though, we'll need to scale our data so that each feature has a single unit variance. from sklearn.preprocessing import StandardScaler
	<pre>scaler = StandardScaler() scaler.fit(df) StandardScaler()</pre>
	scaled_data = scaler.transform(df) PCA with Scikit Learn uses a very similar process to other preprocessing functions that come with SciKit Learn. We instantiate a PCA object, find the principal components using the fit method, the apply the rotation and dimensionality reduction by calling transform().
]:[We can also specify how many components we want to keep when creating the PCA object. from sklearn.decomposition import PCA
]: [<pre>pca = PCA(n_components=2) pca.fit(scaled_data)</pre>
_	PCA(n_components=2) Now we can transform this data to its first 2 principal components. x_pca = pca.transform(scaled_data)
]:]:]:	scaled_data.shape (569, 30)
]:[x_pca.shape (569, 2)
]:	Great! We've reduced 30 dimensions to just 2! Let's plot these two dimensions out! plt.figure(figsize=(8,6)) plt.scatter(x_pca[:,0],x_pca[:,1],c=cancer['target'],cmap='plasma')
]:	plt.xlabel('First principal component') plt.ylabel('Second Principal Component') Text(0, 0.5, 'Second Principal Component')
	12.5 - 10.0 - 10.7 -
	Second Principal Components 2.5 - 0.0 - 0
	-5.0
	-7.5 - 0 5 10 15 First principal component Clearly by using these two components we can easily separate these two classes.
١	Interpreting the components we can easily separate these two classes. Unterpreting the components Unfortunately, with this great power of dimensionality reduction, comes the cost of being able to easily understand what these components represent.
	The components correspond to combinations of the original features, the components themselves are stored as an attribute of the fitted PCA object: pca.components_
]:	array([[0.21890244, 0.10372458, 0.22753729, 0.22099499, 0.14258969,
	[-0.23385713, -0.05970609, -0.21518136, -0.23107671, 0.18611302, 0.15189161, 0.06016536, -0.0347675, 0.19034877, 0.36657547, -0.10555215, 0.08997968, -0.08945723, -0.15229263, 0.20443045, 0.2327159, 0.19720728, 0.13032156, 0.183848, 0.28009203, -0.21986638, -0.0454673, -0.19987843, -0.21935186, 0.17230435, 0.14359317, 0.09796411, -0.00825724, 0.14188335, 0.27533947]])
]: [n this numpy matrix array, each row represents a principal component, and each column relates back to the original features. we can visualize this relationship with a heatmap: df_comp = pd.DataFrame(pca.components_,columns=cancer['feature_names'])
]: []:	<pre>plt.figure(figsize=(12,6)) sns.heatmap(df_comp,cmap='plasma',) </pre>
	-0.3
	-0.2 -0.1
	-0.0 0.1
	0.2
	mean te mean perir mean perir mean smooth mean compact mean comcave p mean symr mean concave p mean symr ean fractal dime worst symr worst concave points symmetry worst concave points symmetry worst servir worst symr orst fractal dime
	E Se Transfer No
	This heatmap and the color bar basically represent the correlation between the various feature and the principal component itself. Conclusion
	This heatmap and the color bar basically represent the correlation between the various feature and the principal component itself. Conclusion Hopefully this information is useful to you when dealing with high dimensional data!