

plt.xlabel('First principal component')
plt.ylabel('Second principal component')
plt.title('Fruits Dataset PCA (n\_components = 2)');

#### Manifold learning methods

### Multidimensional scaling (MDS) on the fruit dataset

### Multidimensional scaling (MDS) on the breast cancer dataset

(This example is not covered in the lecture video, but is included here so you can compare it to the results from PCA.)

```
In []: from sklearn.preprocessing import StandardScaler
    from sklearn.manifold import MDS
    from sklearn.datasets import load_breast_cancer

    cancer = load_breast_cancer()
    (X_cancer, y_cancer) = load_breast_cancer(return_X_y = True)

# each feature should be centered (zero mean) and with unit variance
    X_normalized = StandardScaler().fit(X_cancer).transform(X_cancer)

mds = MDS(n_components = 2)

X_mds = mds.fit_transform(X_normalized)

from adspy_shared_utilities import plot_labelled_scatter
    plot_labelled_scatter(X_mds, y_cancer, ['malignant', 'benign'])

plt.xlabel('First MDS dimension')
    plt.ylabel('Second MDS dimension')
    plt.ylabel('Second MDS dimension')
    plt.title('Breast Cancer Dataset MDS (n_components = 2)');

<IPython.core.display.Javascript object>
```

### t-SNE on the fruit dataset

(This example from the lecture video is included so that you can see how some dimensionality reduction methods may be less successful on some datasets.

Here, it doesn't work as well at finding structure in the small fruits dataset, compared to other methods like MDS.)

### t-SNE on the breast cancer dataset

Although not shown in the lecture video, this example is included for comparison, showing the results of running t-SNE on the breast cancer dataset. See the reading "How to Use t-SNE effectively" for further details on how the visualizations from t-SNE are affected by specific parameter settings.

# Clustering

<IPython.core.display.Javascript object>

### K-means

This example from the lecture video creates an artificial dataset with make\_blobs, then applies k-means to find 3 clusters, and plots the points in each cluster identified by a corresponding color.

```
In []:
    from sklearn.datasets import make_blobs
    from sklearn.cluster import KMeans
    from adspy_shared_utilities import plot_labelled_scatter

X, y = make_blobs(random_state = 10)

kmeans = KMeans(n_clusters = 3)
kmeans.fit(X)

plot_labelled_scatter(X, kmeans.labels_, ['Cluster 1', 'Cluster 2', 'Cluster 3'])
```

<IPython.core.display.Javascript object>

Example showing k-means used to find 4 clusters in the fruits dataset. Note that in general, it's important to scale the individual features before applying k-means clustering.

<IPython.core.display.Javascript object>

### Agglomerative clustering

### Creating a dendrogram (using scipy)

This dendrogram plot is based on the dataset created in the previous step with make\_blobs, but for clarity, only 10 samples have been selected for this example, as plotted here:

And here's the dendrogram corresponding to agglomerative clustering of the 10 points above using Ward's method. The index 0..9 of the points corresponds to the index of the points in the X array above. For example, point 0 (5.69, -9.47) and point 9 (5.43, -9.76) are the closest two points and are clustered first.

```
In []: from scipy.cluster.hierarchy import ward, dendrogram
plt.figure()
dendrogram(ward(X))
plt.show()
```

<IPython.core.display.Javascript object>

## **DBSCAN** clustering