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# 1 Project Name: Kmeans Clustering Implementation

```
In [1]: from kmeans import *

from sklearn.ensemble import RandomForestClassifier
from sklearn.cluster import SpectralClustering
from sklearn.datasets import make_blobs
from sklearn.datasets import load_breast_cancer
from sklearn.metrics import confusion_matrix
from sklearn.metrics import accuracy_score

import numpy as np
import numpy as np
import matplotlib.pyplot as plt
from PIL import Image
from collections import Counter
```

# 2 Blob Clustering

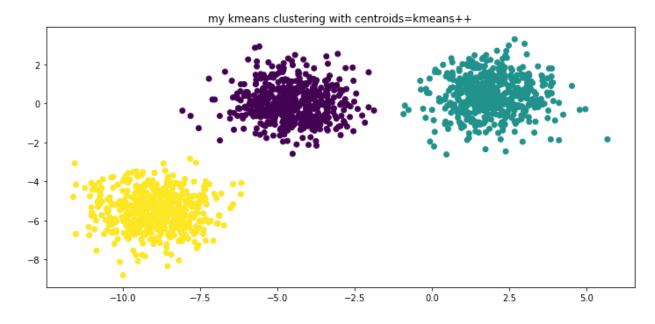
- From the first graph with 2 subplots, we can see that kmeans++ method give better output than centroids=None method.
- From the second graph, we can see that kmeans++ method can really make sparse initial centroids. That's where it works better than normal method.

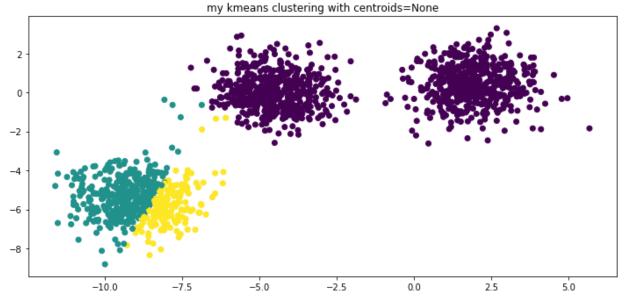
#### 2.1 kmeans

```
In [2]: # kmeans++ has sparse initial centroids
k = 3
X, y = make_blobs(n_samples=1500, random_state=170)
```

```
In [3]: # my prediction
        results = []
        for c in ['kmeans++', None]:
            centroids, clusters = kmeans(X=X,k=k,centroids=c)
            y_pred = np.zeros([len(X),1])
            for cluster_number in range(k):
                y pred[clusters[cluster_number]] = cluster_number
            y pred = y_pred.reshape(1,-1)[0]
            results.append(y_pred)
        plt.figure(figsize=(12, 12))
        plt.subplot(211)
        plt.scatter(X[:, 0], X[:, 1], c= results[0])
        plt.title('my kmeans clustering with centroids=kmeans++')
        plt.subplot(212)
        plt.scatter(X[:, 0], X[:, 1], c= results[1])
        plt.title('my kmeans clustering with centroids=None')
```

Out[3]: Text(0.5, 1.0, 'my kmeans clustering with centroids=None')

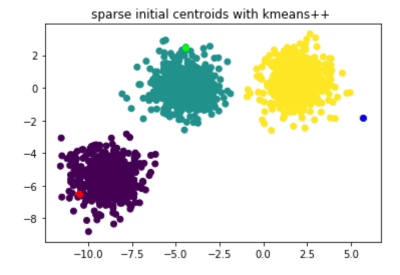




#### 2.2 kmeans++

```
In [4]: # kmeans++ initialization gives sparse initial centroids
    centroids = init_centroids(X=X,k=k) # init sparse centroids, not randomly i
    plt.scatter(X[:, 0], X[:, 1], c= y)
    plt.scatter(centroids[0,0],centroids[0,1],c='red')
    plt.scatter(centroids[1,0],centroids[1,1],c='blue')
    plt.scatter(centroids[2,0],centroids[2,1],c='lime')
    plt.title('sparse initial centroids with kmeans++ ')
```

```
Out[4]: Text(0.5, 1.0, 'sparse initial centroids with kmeans++ ')
```



## 3 Binary Prediction

- In this kind of case, kmeans clustering can be used for classification, normally with good accuracy.
- Here, it's important to link our cluster number with its correct group, we should flip\_over predictions when necessary.

```
In [5]: cancer = load_breast_cancer()
X = cancer.data
y = cancer.target
print('{:.2%} observations has y_true=1'.format(y[y==1].shape[0]/y.shape[0]
# y=0 cancer, y=1 benign
```

62.74% observations has y\_true=1

#### 3.1 Mehod 1 to match groups with labels

- The Basic idea, is to identify each cluster's size.
- Sort values in y by its counts, then associate clusters in y\_pred with labels by finding its rank in value counts.
- To be more specific, in this case, the bigger cluster would be assigned lable=1.
- Pros: can deal with multi\_classification problems.

• Cons: when clusters' size are similar, it may fail to find its correct label.

```
In [6]: def sort by counts(y):
            # return a list of tuples, like [(1, 357), (0, 212)]
            return sorted(Counter(y).items(),key=lambda x:x[1],reverse=True)
        def flip over if necessary(y,y pred):
            sorted_y = sort_by_counts(y)
            sorted_y_pred = sort_by_counts(y_pred.reshape(1,-1)[0])
            # similar distribution, same order in counts
            if [item for item,count in sorted_y] == [item for item,count in sorted_
                return y pred
            else:
                y_pred_new = y_pred
                for i in range(len(sorted y)):
                    y pred new[clusters[int(sorted y pred[i][0])]] = sorted y[i][0]
                return y pred new
        for i in range(1,11):
            print('Test Number {} '.format(i))
            k = 2
            centroids, clusters = kmeans(X, k)
            y_pred = np.zeros([len(X), 1])
            for cluster_number in range(k):
                y pred[clusters[cluster number]] = cluster number
            accuracy_before = accuracy_score(y, y pred.reshape(1, -1)[0])
            print('Accuracy before flip over : \n {:.4f}'.format(accuracy before))
            # flip over according to the value counts distribution
            y_pred = flip_over_if_necessary(y,y_pred)
            accuracy = accuracy_score(y, y_pred.reshape(1, -1)[0])
            print('Accuracy : \n {:.4f}'.format(accuracy))
            if accuracy<0.5:</pre>
                print('Failed !!! Because predicted clusters have similar sizes !!!
            print('Confusion Matrix : \n {} \n'.format(confusion_matrix(y, y_pred))
        Test Number 1
        Accuracy before flip_over :
         0.8225
        Accuracy:
         0.1775
        Failed !!! Because predicted clusters have similar sizes !!!
        Confusion Matrix:
         [[ 6 206]
         [262 95]]
        Test Number 2
        Accuracy before flip_over :
         0.1019
        Accuracy:
         0.8981
        Confusion Matrix:
         [[160 52]
            6 351]]
        Test Number 3
```

```
Accuracy before flip_over :
 0.8699
Accuracy:
 0.8699
Confusion Matrix:
 [[199 13]
 [ 61 296]]
Test Number 4
Accuracy before flip_over :
 0.8576
Accuracy :
 0.8576
Confusion Matrix :
 [[132 80]
 [ 1 356]]
Test Number 5
Accuracy before flip_over :
0.1951
Accuracy :
0.8049
Confusion Matrix:
 [[101 111]
 [ 0 357]]
Test Number 6
Accuracy before flip_over :
0.2531
Accuracy :
 0.2531
Failed !!! Because predicted clusters have similar sizes !!!
Confusion Matrix:
 [[ 5 207]
 [218 139]]
Test Number 7
Accuracy before flip over :
0.1775
Accuracy :
0.1775
Failed !!! Because predicted clusters have similar sizes !!!
Confusion Matrix:
 [[ 6 206]
 [262 95]]
Test Number 8
Accuracy before flip over :
 0.1265
Accuracy :
 0.8735
Confusion Matrix :
 [[143 69]
 [ 3 354]]
Test Number 9
Accuracy before flip over :
```

```
0.0914
Accuracy:
0.9086
Confusion Matrix:
[[170 42]
[ 10 347]]

Test Number 10
Accuracy before flip_over:
0.1054
Accuracy:
0.8946
Confusion Matrix:
[[177 35]
[ 25 332]]
```

### 3.2 Mehod 2 to match groups with labels

- The basic idea, is to investigate the performace of clustering.
- To be more specific, in this case, if accuracy<0.5, we flip\_over the prediction, and recalculate the metrics.
- In this case, method 2 works better than method 1, because 62.74% observations has y\_true=1. When the clusters' size don't differ that much, method 1 would fail to implement flip\_over, even though it's really necessary.

```
In [7]: accuracies_kmeans = []
        for i in range(1, 6):
            print('Test Number {} '.format(i))
            k = 2
            centroids, clusters = kmeans(X, k)
            y_pred = np.zeros([len(X), 1])
            for cluster_number in range(k):
                y_pred[clusters[cluster_number]] = cluster_number
            accuracy_before = accuracy_score(y, y_pred.reshape(1, -1)[0])
            if accuracy_before<0.5: # simple-judge</pre>
                print('Accuracy before flip_over : \n {:.4f}'.format(accuracy_befor
                y pred = np.where(y pred==1,0,1) # flip it
            accuracy = accuracy_score(y, y_pred.reshape(1, -1)[0])
            accuracies_kmeans.append(accuracy)
            print('Accuracy : \n {:.4f}'.format(accuracy))
            print('Confusion Matrix : \n {} \n'.format(confusion matrix(y, y pred))
        print('In Average, Accuracy_kmeans = {:.4f}'.format(np.mean(accuracies_kmea
        Test Number 1
        Accuracy:
         0.8717
        Confusion Matrix :
         [[199 13]
         [ 60 297]]
        Test Number 2
        Accuracy:
         0.9033
        Confusion Matrix:
         [[174 38]
         [ 17 340]]
        Test Number 3
        Accuracy before flip_over :
         0.0914
        Accuracy:
         0.9086
        Confusion Matrix:
         [[168 44]
         [ 8 349]]
        Test Number 4
        Accuracy:
         0.8225
        Confusion Matrix:
         [[206
                 61
         [ 95 262]]
        Test Number 5
        Accuracy before flip over :
         0.1037
        Accuracy:
```

```
0.8963
Confusion Matrix :
  [[178   34]
  [ 25  332]]
In Average, Accuracy_kmeans = 0.8805
```

# 4 Image Compression

## 4.1 Greyscale image

• set the points that has similar grey\_value to a single (centroid) value.

```
In [8]: image_path = 'north-africa-1940s-grey.png'
img = np.array(Image.open(image_path))
# the greyscale value given by img[i][j]
```

In [9]: display(Image.fromarray(img))



```
In [10]: X = img.reshape(-1,1)
# Number of cluster k : how many grey_values are chosen.
k = 4
centroids, clusters = kmeans(X,k,centroids='kmeans++')
# get the four centroids, many points grey_value are around these centroids
centroids = centroids.astype(np.uint8)

y_pred = np.zeros([len(X), 1])
# convert all the grey_values to the 4 centroids
for cluster_number in range(k):
    y_pred[clusters[cluster_number]] = centroids[cluster_number]
```

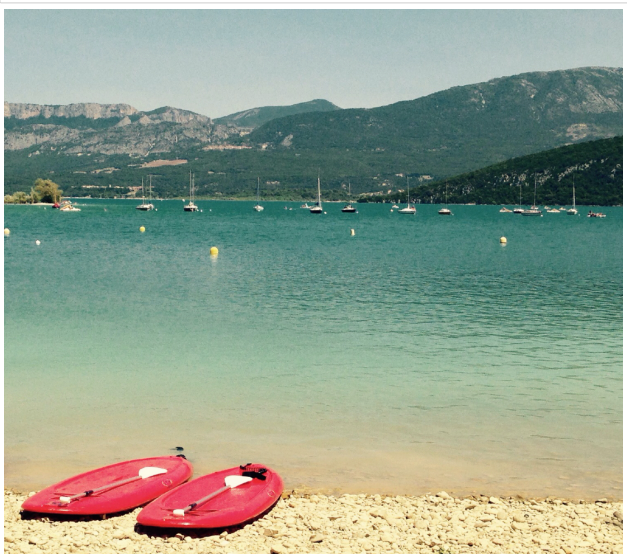
```
In [11]: y_pred = y_pred.reshape(img.shape)
grey_img = y_pred.astype(np.uint8)
display(Image.fromarray(grey_img))
```



### 4.2 Color image

• set the points that have similar color to a single (centroid) color.

```
In [16]: image_path_2 = 'lake.jpeg'
img_2 = np.array(Image.open(image_path_2))
display(Image.fromarray(img_2))
```



```
In [17]: X = img_2.reshape(-1,1,3) # 3 for red, greeb, blue
k = 32

centroids, clusters = kmeans(X,k,centroids= 'kmeans++')
centroids = centroids.astype(np.uint8)
y_pred = np.zeros([len(X), 3])
# convert all the colors to the 32 colors
for cluster_number in range(k):
    y_pred[clusters[cluster_number]] = centroids[cluster_number]
```



# 5 Advanced: Spectral Clustering

```
In [19]: cancer = load_breast_cancer()
X = cancer.data
y = cancer.target
```

```
In [20]: accuracies_spectral = []
         for i in range(1,6): # different random state in rf for each i
             print('Test Number {}'.format(i))
             rf= RandomForestClassifier() # different random state
             rf.fit(X,y)
             S = similarity_matrix(X,rf)
             cluster = SpectralClustering(n clusters=2, affinity='precomputed')
             y pred = cluster.fit predict(S) # pass similarity matrix not X
             if accuracy_score(y,y_pred)<0.5: # might needs to filp over</pre>
                 y pred = np.where(y pred==1,0,1)
             accuracy = accuracy score(y, y pred)
             accuracies_spectral.append(accuracy)
             print('Accuracy : \n {:.4f}'.format(accuracy))
             print('Confusion Matrix : \n {} \n'.format(confusion matrix(y, y pred))
         print('In Average, Accuracy spectral = {:.4f}'.format(np.mean(accuracies_sp
         Test Number 1
         Accuracy:
          0.9701
         Confusion Matrix:
          [[201 11]
          [ 6 351]]
         Test Number 2
         Accuracy:
          0.9684
         Confusion Matrix:
          [[201 11]
          [ 7 350]]
         Test Number 3
         Accuracy:
          0.9772
         Confusion Matrix:
          [[204 8]
          [ 5 352]]
         Test Number 4
         Accuracy:
          0.9719
         Confusion Matrix:
          [[202 10]
          [ 6 351]]
         Test Number 5
         Accuracy:
          0.9789
         Confusion Matrix:
          [[205 7]
          [ 5 352]]
         In Average, Accuracy spectral = 0.9733
```

```
In [21]: print('In Average, Accuracy_kmeans = {:.4f}'.format(np.mean(accuracies_kmeal print('In Average, Accuracy_spectral = {:.4f}'.format(np.mean(accuracies_spectral = Accuracy_kmeans = 0.8805)
In Average, Accuracy_kmeans = 0.8805
In Average, Accuracy_spectral = 0.9733
```

 Spectral\_Clustering has higher average accuracy than Kmeans\_Clustering in the cancer\_or\_not classification problem.

# 6 Summary

- I implemented kmeans clustering (both kmeans and kmeans++ algorithm) from scratch.
- I achieved these applications with my own kmeans clustering code: blob clustering, binary prediction, image compression, and spectral clustering.