Yue_Mu_Final_Project

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1 STA663 Final Project

1.1 Background

- K-means clustering method remains the most popular clustering method and one of the top 10 algorithms in data mining. However, it is not a good clustering method in terms of efficiency or quality. Recent work has focused on improving the initialization procedure, which leads to k-means++ and kmeans|| algorithm.
- Basic K-means clustering uses the Lloyd algorithm, which starts with k arbitrary centers and performs an EM-type local search till convergence. However, it is very sensitive to initialization and takes many iterations to converge.
- K-means++ solves this problem by first choosing a center at random, and then sampling one point each iteration based on their probability distribution. It does spread out the centers pretty well, but it needs k passes over the data, which can be a problem when dataset is large.
- K-means||, also called scalable K-means++, oversamples by sampling each point independently with a larger probability, which is intuitively equivalent to updating the distribution much less frequently. It turns out to be more efficient than K-means++.

1.2 Algorithm

```
In [1]: from __future__ import division
    import os
    import sys
    import glob
    import matplotlib.pyplot as plt
    import matplotlib.cm as cm
    import numpy as np
    import pandas as pd
    import time
    %matplotlib inline
    %precision 4
    plt.style.use('ggplot')
```

1.2.1 K-means Algorithm (Simple)

```
In [2]: def initialization(data,k):
    """Return the initial set of cluster centers."""

#Generate k random indices between 0 and the number of rows in the dataset
    centers_index = np.random.choice(range(data.shape[0]), k, replace=False)
    return data[centers_index, :]

def kmeans1(centers,data,k):
```

```
"""Implement K-means clustering algorithm. Return converged cluster centers."""
            converge = False
            phi = []
            iterations = 0
            while (not converge) and (iterations < 1000):
                #Find out which cluster each data point is the closest to
                min_dist = np.zeros(data.shape[0])
                min_index = np.zeros(data.shape[0])
                for i,p in enumerate(data):
                    result = minimum_distance(p,centers)
                    min_dist[i] = result[0]
                    min_index[i] = result[1]
                #Calculate phi
                phi_val = np.sum(min_dist)
                phi.append(phi_val)
                #Calculate the new centers
                new_centers = np.empty(centers.shape)
                for i in range(0, k):
                    if data[min_index == i,:].shape[0] == 0:
                        new_centers[i] = centers[i]
                    else:
                        new_centers[i] = np.mean(data[min_index == i, :], axis=0)
                #Compare old centers with new centers to see if the algorithm has converged
                if np.array_equal(centers, new_centers):
                    converge = True
                else:
                    centers = new_centers
                iterations += 1
            return (iterations, phi, centers, min_index)
        def minimum_distance(p, centers):
            """Return the distance to the closest cluster center"""
           min_index = 0
           min_dist = sys.float_info.max
            for i, cc in enumerate(centers):
                d = np.sum((p - cc) ** 2)
                if min_dist > d:
                    min_dist = d
                    min_index = i
            return (min_dist,min_index)
1.2.2 K-means Algorithm (Vectorized)
In [3]: def kmeans2(centers,data,k):
            """Implement K-means clustering algorithm. Return converged cluster centers."""
            converge = False
```

```
iterations = 0
            while (not converge) and (iterations < 1000):
                #Find the Euclidean distance between a center and a data point
                data2 = data[:, np.newaxis, :]
                d2 = (data2 - centers) ** 2
                #Calculate the total distance to each center for each data point.
                distance = np.sum(d2, axis=2)
                #Find out which cluster each data point belongs to.
                min_index = np.zeros(distance.shape)
                min_index[range(distance.shape[0]), np.argmin(distance, axis=1)] = 1
                #Calculate phi
                phi_val = np.sum(distance[min_index == 1])
                phi.append(phi_val)
                #Calculate the new centers
                new_centers = np.empty(centers.shape)
                for i in range(0, k):
                    if data[min_index[:, i] == 1,:].shape[0] == 0:
                        new_centers[i] = centers[i]
                    else:
                        new_centers[i] = np.mean(data[min_index[:, i] == True, :], axis=0)
                #Compare old centers with new centers to see if the algorithm has converged
                if np.array_equal(centers, new_centers):
                    converge = True
                else:
                    centers = new_centers
                iterations += 1
            return (iterations, phi, centers, min_index)
1.2.3 K-means++ Algorithm (Simple)
In [4]: def kmeanspp1(data, k):
            """Implement the K-means++ algorithm. Return k initial cluster centers."""
            #Sample a point uniformly at random from the data
            centers = data[np.random.choice(range(data.shape[0]),1), :]
            #Iterate k-1 times through the dataset to select the initial cluster centers
            while centers.shape[0] < k:
                min_dist = np.zeros(data.shape[0])
                for i,p in enumerate(data):
                    min_dist[i] = minimum_distance(p,centers)[0]
                #Calculate cost of the data
                phi = np.sum(min_dist)
                #Calculate the probability distribution
                prob = min_dist/phi
                #Select the next centroid using the probability distribution calculated
                centers = np.vstack([centers, data[np.random.choice(range(data.shape[0]),1, p = prob),
            return centers
```

phi = []

```
1.2.4 K-means++ Algorithm (Vectorized)
```

```
In [5]: def kmeanspp2(data, k):
            """Implement the K-means++ algorithm. Return k initial cluster centers."""
            #Sample a point uniformly ar random from the data
            centers = data[np.random.choice(range(data.shape[0]),1), :]
            data2 = data[:, np.newaxis, :]
            #Iterate k-1 times through the dataset to select the initial cluster centers
            while centers.shape[0] < k:
                d2 = (data2 - centers) ** 2
                distance = np.sum(d2, axis=2)
                min_index = np.zeros(distance.shape)
                min_index[range(distance.shape[0]), np.argmin(distance, axis=1)] = 1
                min_dist=distance[min_index == 1]
                phi = np.sum(min_dist)
                #Calculate the probability distribution
                prob = min_dist/phi
                #Select the next centroid using the probability distribution calculated
                centers = np.vstack([centers, data[np.random.choice(range(data.shape[0]),1, p = prob),
            return centers
```

1.2.5 Scalable K-means++ Algorithm

```
In [6]: def scalablekmeanspp(data,k,l):
            """Implement the K-means++ algorithm. Return k initial cluster centers."""
            #Sample a point uniformly at random from the data
            centers = data[np.random.choice(range(data.shape[0]),1), :]
            data2 = data[:, np.newaxis, :]
            min_dist = np.zeros(data.shape[0])
            for i,p in enumerate(data):
                min_dist[i] = minimum_distance(p,centers)[0]
            phi = np.sum(min_dist)
            for i in range(int(round(np.log(phi)))):
                d2 = (data2 - centers) ** 2
                distance = np.sum(d2, axis=2)
                min_index = np.zeros(distance.shape)
                min_index[range(distance.shape[0]), np.argmin(distance, axis=1)] = 1
                min_dist=distance[min_index == 1]
                phi = np.sum(min_dist)
                for j,p in enumerate(data):
                    prob = l*min_dist[j]/phi
                    u=np.random.uniform(0,1)
                    if prob >= u:
                        centers = np.vstack([centers, p])
            #Now we have an initial set of cluster centers that is greater than k
            #Find number of points in dataset closer to each center than any other centers
            d2 = (data2 - centers) ** 2
            distance = np.sum(d2, axis=2)
            min_index = np.zeros(distance.shape)
            min_index[range(distance.shape[0]), np.argmin(distance, axis=1)] = 1
            weights = np.array([np.count_nonzero(min_index[:, j]) for j in range(centers.shape[0])]).re
            #Recluster the weighted points into k clusters using kmeans++
```

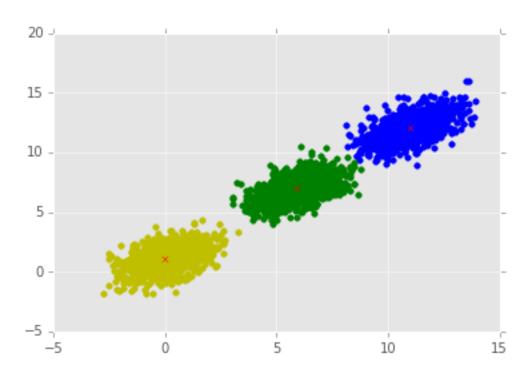
new_centers = centers[np.random.choice(range(centers.shape[0]),1), :]

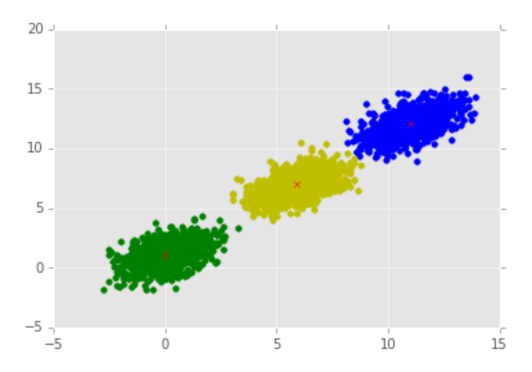
```
centers2 = centers
index = np.where(centers2==new_centers)[0]
centers2 = np.delete(centers2,index[0],axis=0)
weights = np.delete(weights,index[0])
while new_centers.shape[0] < k:
    #Calculate the probability distribution based on weights
    prob = weights/np.sum(weights)
    #Select the next centroid using the probability distribution calculated
    c = centers2[np.random.choice(range(centers2.shape[0]),1, p = prob.ravel()), :]
    index = np.where(centers2==c)[0]
    new_centers = np.vstack([new_centers, c])
    #Remove the selected center and its corresponding weight
    centers2 = np.delete(centers2,index[0],axis=0)
    weights = np.delete(weights,index[0])
#Implement kmeans clustering to obtain the new centers
new_centers2 = kmeans2(new_centers,centers,k)[2]
return new_centers2
```

1.3 Simulate Data

```
In [7]: def generate_data(n):
            """Return mixed data with cluster labels."""
            mean0 = [0, 1]
            cov0 = [[1, 0.5], [0.5, 1]]
            data0 = np.random.multivariate_normal(mean0, cov0, n)
            data0 = np.hstack((data0, np.ones((data0.shape[0],1))))
            mean1 = [6, 7]
            cov1 = [[1, 0.5], [0.5, 1]]
            data1 = np.random.multivariate_normal(mean1, cov1, n)
            data1 = np.hstack((data1, np.ones((data1.shape[0],1)) * 2))
            mean2 = [11, 12]
            cov2 = [[1, 0.5], [0.5, 1]]
            data2 = np.random.multivariate_normal(mean2, cov2, n)
            data2 = np.hstack((data2, np.ones((data2.shape[0],1)) * 3))
            data = np.vstack((data0, data1, data2))
            np.random.shuffle(data)
            print (data.shape)
            return data
        data1=generate_data(1000)[:,0:2]
(3000, 3)
In [8]: def generate_data(n):
            """Return mixed data with cluster labels."""
            mean0 = [0, 1]
            cov0 = [[1, 0.5], [0.5, 1]]
            data0 = np.random.multivariate_normal(mean0, cov0, n)
            data0 = np.hstack((data0, np.ones((data0.shape[0],1))))
```

```
mean1 = [5, 6]
            cov1 = [[1, 0.5], [0.5, 1]]
            data1 = np.random.multivariate_normal(mean1, cov1, n)
            data1 = np.hstack((data1, np.ones((data1.shape[0],1)) * 2))
            mean2 = [10, 11]
            cov2 = [[1, 0.5], [0.5, 1]]
            data2 = np.random.multivariate_normal(mean2, cov2, n)
            data2 = np.hstack((data2, np.ones((data2.shape[0],1)) * 3))
            mean3 = [15, 16]
            cov3 = [[1, 0.5], [0.5, 1]]
            data3 = np.random.multivariate_normal(mean2, cov2, n)
            data3 = np.hstack((data3, np.ones((data3.shape[0],1)) * 4))
            mean4 = [20, 21]
            cov4 = [[1, 0.5], [0.5, 1]]
            data4 = np.random.multivariate_normal(mean2, cov2, n)
            data4 = np.hstack((data4, np.ones((data4.shape[0],1)) * 5))
            data = np.vstack((data0, data1, data2, data3, data4))
            np.random.shuffle(data)
            print (data.shape)
            return data
        data2=generate_data(5000)[:,0:2]
(25000, 3)
1.4 Test
In [229]: k=3
          result = kmeans2(initialization(data1,k),data1,k)
          centers = result[2]
          min_index = result[3]
          #Plot clusters
          cols=iter(['b','g','y'])
          for i in range (k):
              plt.scatter(data1[min_index[:,i] == 1, :][:,0], data1[min_index[:,i] == 1, :][:,1], color
          plt.scatter(centers[:,0], centers[:,1], color='r', marker='x')
('Initial Centers:', array([[ 8.2972, 8.0749],
       [ 6.5101, 6.9435],
       [-1.2449, -0.4135]))
('Required', 4, 'iterations to converge.')
Out[229]: <matplotlib.collections.PathCollection at 0x7fe759310e10>
```

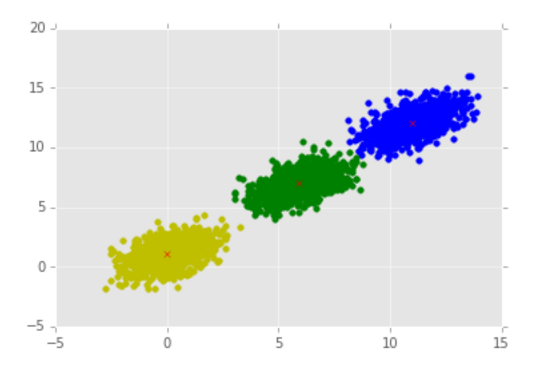




```
In [232]: result = kmeans2(scalablekmeanspp(data1,k,2),data1,k)
         centers = result[2]
         min_index = result[3]
         #Plot clusters
         cols=iter(['b','g','y'])
         for i in range (k):
             plt.scatter(data1[min_index[:,i] == 1, :][:,0], data1[min_index[:,i] == 1, :][:,1], color
         plt.scatter(centers[:,0], centers[:,1], color='r', marker='x')
('Initial Centers:', array([[ 12.5395, 12.3895],
       [ 10.388 , 12.0946],
       [ 6.6019, 7.9372]]))
('Required', 7, 'iterations to converge.')
('Initial Centers:', array([[ 11.1654, 12.1567],
       [ 5.877 , 6.9726],
       [ 0.4497, 1.459]]))
('Required', 3, 'iterations to converge.')
```

8

Out[232]: <matplotlib.collections.PathCollection at 0x7fe75923bf90>



1.5 Comparison of Algorithms

```
In [53]: np.random.seed(200)
         result = kmeans2(initialization(data1,3),data1,3)
         #Print number of iterations till convergence
         print result[0]
         #Print the clustering cost after initialization
         print result[1][0]
46372.4191656
In [55]: np.random.seed(200)
         result = kmeans2(kmeanspp2(data1,3),data1,3)
         print result[0]
         print result[1][0]
13765.1574859
In [54]: np.random.seed(200)
         result = kmeans2(scalablekmeanspp(data1,3,1.5),data1,3)
         print result[0]
         print result[1][0]
3
6396.64504607
In [66]: np.random.seed(140)
         result = kmeans2(initialization(data2,5),data2,5)
         print result[0]
         print result[1][0]
```

Based on the results above, we can see that K-means++ reduces number of Lloyd iterations by a great amount, and scalable K-means++ does that even more than K-means++, especially when the dataset is bigger and has more clusters. The efficiency of scalable K-means++ can also be reflected in the initial clustering cost. The smaller the initial clustering cost is, the better the initial centroids are, meaning that they are closer to the actual cluster centers. The initial clustering cost of scalable K-means++ is significantly lower than the other two algorithms.

1.6 Code Efficiency

Line #	Hits	Time	Per Hit	% Time	Line Contents
8					def kmeans1(centers,data,k):
9					"""Implement K-means clustering algorithm. I
10					
11	1	2	2.0	0.0	converge = False
12	1	2	2.0	0.0	phi = []
13	1	0	0.0	0.0	iterations = 0
14	4	11	2.8	0.0	while (not converge) and (iterations < 1000)
15					#Find out which cluster each data point
16	3	30	10.0	0.0	<pre>min_dist = np.zeros(data.shape[0])</pre>
17	3	10	3.3	0.0	<pre>min_index = np.zeros(data.shape[0])</pre>
18	9003	10552	1.2	2.5	<pre>for i,p in enumerate(data):</pre>
19	9000	381187	42.4	91.3	result = minimum_distance(p,centers

9000	14599	1.6	3.5	min_dist[i] = result[0]
9000	9386	1.0	2.2	min_index[i] = result[1]
				#Calculate phi
3	40	13.3	0.0	<pre>phi_val = np.sum(min_dist)</pre>
3	7	2.3	0.0	phi.append(phi_val)
				#Calculate the new centers
3	15	5.0	0.0	<pre>new_centers = np.empty(centers.shape)</pre>
12	32	2.7	0.0	<pre>for i in range(0, k):</pre>
9	514	57.1	0.1	if data[min_index == i,:].shape[0] =
				new_centers[i] = centers[i]
				else:
9	1051	116.8	0.3	new_centers[i] = np.mean(data[m:
				$ extit{\#Compare old centers with new centers t}$
3	109	36.3	0.0	<pre>if np.array_equal(centers, new_centers)</pre>
1	1	1.0	0.0	converge = True
				else:
2	1	0.5	0.0	<pre>centers = new_centers</pre>
3	3	1.0	0.0	iterations += 1
1	0	0.0	0.0	return (iterations, phi, centers, min_index
	9000 3 3 12 9 9 9 3 1 2 3	9000 9386 3 40 3 7 3 15 12 32 9 514 9 1051 3 109 1 1 2 1 3 3	9000 9386 1.0 3 40 13.3 3 7 2.3 3 15 5.0 12 32 2.7 9 514 57.1 9 1051 116.8 3 109 36.3 1 1 1.0 2 1 0.5 3 3 1.0	9000 9386 1.0 2.2 3 40 13.3 0.0 3 7 2.3 0.0 3 15 5.0 0.0 12 32 2.7 0.0 9 514 57.1 0.1 9 1051 116.8 0.3 3 109 36.3 0.0 1 1 1.0 0.0 2 1 0.5 0.0 3 3 1.0 0.0

We can see that majority of the time was spent on running the minimum_distance function because it requires running through the entire dataset. We can try to optimize this part of the algorithm using Cython.

```
In [9]: %load_ext cythonmagic
In [10]: %%cython -a
         import numpy as np
         cimport numpy as np
         cimport cython
         @cython.boundscheck(False)
         @cython.wraparound(False)
         def cminimum_distance(np.ndarray[np.float64_t, ndim=1] p, np.ndarray[np.float64_t, ndim=2] cen
             cdef int min_index = 0
             cdef float min_dist = 1.79769313486e+308
             cdef int i
             cdef np.ndarray[np.float64_t, ndim=1] cc
             for i, cc in enumerate(centers):
                 d = np.sum((p - cc) ** 2)
                 if min_dist > d:
                     min_dist = d
                     min_index = i
             return (min_dist,min_index)
Out[10]: <IPython.core.display.HTML at 0x7fa432851b90>
```

1.6.1 K-means Algorithm (Cython)

```
In [11]: def kmeans3(centers,data,k):
             """Implement K-means clustering algorithm. Return converged cluster centers."""
```

```
phi = []
             iterations = 0
             while (not converge) and (iterations < 1000):
                 #Find out which cluster each data point is the closest to
                 min_dist = np.zeros(data.shape[0])
                 min_index = np.zeros(data.shape[0])
                 for i,p in enumerate(data):
                     #Here use the cython version of the minimum_distance function
                     result = cminimum_distance(p,centers)
                     min_dist[i] = result[0]
                     min_index[i] = result[1]
                 #Calculate phi
                 phi_val = np.sum(min_dist)
                 phi.append(phi_val)
                 #Calculate the new centers
                 new_centers = np.empty(centers.shape)
                 for i in range(0, k):
                     if data[min_index == i,:].shape[0] == 0:
                         new_centers[i] = centers[i]
                     else:
                         new_centers[i] = np.mean(data[min_index == i, :], axis=0)
                 #Compare old centers with new centers to see if the algorithm has converged
                 if np.array_equal(centers, new_centers):
                     converge = True
                 else:
                     centers = new_centers
                 iterations += 1
             return (iterations, phi, centers, min_index)
1.6.2 K-means++ (Cython)
In [12]: def kmeanspp3(data, k):
             """Implement the K-means++ algorithm. Return k initial cluster centers."""
             #Sample a point uniformly at random from the data
             centers = data[np.random.choice(range(data.shape[0]),1), :]
             #Iterate k-1 times through the dataset to select the initial cluster centers
             while centers.shape[0] < k:</pre>
                 min_dist = np.zeros(data.shape[0])
                 for i,p in enumerate(data):
                     #Here use the cython version of the minimum_distance function
                     min_dist[i] = cminimum_distance(p,centers)[0]
                 #Calculate cost of the data
                 phi = np.sum(min_dist)
                 #Calculate the probability distribution
                 prob = min_dist/phi
                 #Select the next centroid using the probability distribution calculated
                 centers = np.vstack([centers, data[np.random.choice(range(data.shape[0]),1, p = prob),
```

converge = False

return centers

It seems that in this case vectorization increases the efficiency the most.

In []: