CSE 847 Home Assignment 5

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1 Clustering: K-Means

1. Elaborate the relationship between k-means and spectral relaxation of k-means. Is it possible that we obtain exact k-means solution using spectral relaxed k-means?

Response: To discuss this issue, the rows of data matrix are considered to be the samples and columns are considered to be the features. Following the objective of k-means to reduce the sum-of-squared error (SSE), we can represent the same objective in a different format by considering each cluster as a set Π_j where j represent the cluster label and Π_j contains all the data points in the j^{th} cluster. The SSE can then be represented as:

$$SSE(\Pi_j) = \sum_{v \in \Pi_j} ||x_v - c_j||^2$$

where c_j is the cluster center of the corresponding cluster. The entire SSE can be found after summing over all the clusters.

After some transformations in matrix form, the objective function can be represented as:

$$SSE = \sum_{j=1}^{k} (trace(X_j^T X_j) - \frac{e^T}{\sqrt{n_j}} X_j^T X_j \frac{e}{\sqrt{n_j}})$$
 (1)

This expression could be further simplified as:

$$SSE = trace(X_i^T X_i) - trace(Y^T X_i^T X_i Y)$$
(2)

$$\text{where } Y = \left\{ \begin{array}{cccc} \frac{e}{\sqrt(n_1)} & 0 & . & . & 0 \\ 0 & \frac{e}{\sqrt(n_1)} & . & . & 0 \\ & & . & & \\ & & . & & \\ 0 & 0 & . & . & \frac{e}{\sqrt(n_K)} \end{array} \right\} \text{ which is an } (n \times k) \text{ orthogonal matrix.}$$

The goal of the approach then becomes to minimize SSE by maximizing $trace(Y^T X_j^T X_j Y)$. The spectral relaxation of k-means is inspired from the fact that instead of using this specific expression for Y, it is possible to use any arbitrary orthogonal matrix for Y. This leads to the relaxed maximization problem:

$$max_{Y^TY=I_k}trace(Y^TX_j^TX_jY)$$
(3)

The first k vectors in the left singular matrix of X can produce the Y^* that maximizes this expression. Finally, the clusters can be found using regular k-means on Y^* . That's how spectral version of k-means works. As evident from this discussion, we can see that both k-means and spectral k-means are trying to minimize the same error function, but spectral k-means is first trying to project the dataset to a lower dimensional space which makes it easier to capture the complex clustering structures.

The spectral-relaxed k-means become completely equivalent to k-means when the expression for Y becomes equal to the matrix mentioned in Equation 2.

- 2. Implementation of k-means. Submit all the source code to D2L along with a short report on your observation.
 - Implement the k-means in Matlab using the alternating procedure introduced in the class (you will not get the credit if you use the build-in kmeans function in Matlab).
 - Implement the spectral relaxation of k-means. Create a random dataset and compare the k-means and spectral relaxed k-means.

Response: In the previous part, the main concepts behind the spectral k-means was discussed. In both the cases, the regular k-means is used as the underlying clustering algorithm. It works using Block Coordinate Descend (BCD) minimizing one variable at a time by keeping the remaining ones fixed. The implementation of the algorithms was combined into a single code block as shown below. The spectral variant can be just used by passing true value for the 'spectral' parameter for kmeans. The code of the k-means implementation is also posted at: K-Means.

```
1 data = importdata('Data/Mall_Customers.csv');
2 data = data.data;
3 \, data = data(:,2:5);
4 \text{ num\_clusters} = [3:10];
5 \text{ SSE} = \mathbf{zeros}(1, \mathbf{size}(\text{num\_clusters}, 2));
6 for i=1:size(num\_clusters, 2)
       [cluster_assignments, cluster_centers] = kmeans_cluster(data,
7
          num_clusters(i));
      SSE(i) = compute_SSE(data, cluster_assignments);
8
      show_plot(data, cluster_assignments);
9
      saveas (gcf, streat ('Images/KMeans/Cluster_Arrangement_', int2str(
10
          num_clusters(i)),'.jpg'));
11 end
12 fig = figure; plot(num_clusters, SSE); xlabel('K'); ylabel('SSE'); title('
      Variation of SSE for different values of K');
13 saveas (gcf, strcat ('Images/KMeans/SSE_Convergence.jpg'));
14
15
16 function [cluster_assignments, cluster_centers] = kmeans_cluster(raw_data, k
      , spectral)
17 %
         Code to perform k-means clustering
18 %
         INPUTS:
                       = (n * m) matrix where n is the number of samples and m
19 %
             data
```

```
is\ the\ number\ of\ features
20 %
21 %
                           integer number of intended clusters
22 %
              spectral = boolean value representing spectral k-means if true,
23 %
                            else standard k-means
24 %
25 %
         OUTPUTS:
26 %
              cluster\_assignments = labels assigned to the each samples in
27 %
                                       [1,k]
28 %
              cluster\_centers
                                    = final clsuter centers found in the process
29 %
                                       of k-means
30
       % Assigning default values
31
       if nargin < 3
32
           if ~exist('spectral')
33
               spectral=false;
34
           end
35
       end
36
37
       if spectral
38
           \% for spectral relaxation, map the data samples to k-dimensional
39
           % feature space
40
           [U, \tilde{a}, \tilde{a}] = svd(raw_data);
41
           projection = U(:, 1:k);
42
           rand_mat = rand(k,k);
43
           orth_mat = orth(rand_mat);
44
           data = projection * orth_mat;
45
       else
46
           data = raw_data;
47
       end
48
49
       [\text{num\_samples}, ~\tilde{}] = \text{size}(\text{data});
50
       cluster_assignments = zeros(num_samples, 1);
51
       temp = randperm(num_samples);
52
       cluster\_center\_idx = temp(1:k);
53
       cluster_centers = data(cluster_center_idx , :);
54
       change = inf;
55
       count_iter = 0;
56
57
58
       while (change = 0)
           \% change represents the number cluster assignments that got changed
59
60
           % in the current iteration
           count_iter = count_iter + 1;
61
           prev_assignments = cluster_assignments;
62
63
           for cur_idx=1:num_samples
64
                \min_{-dist} = \inf;
65
                \min_{-idx} = -1;
66
```

```
67
                % for each sample, find the cluster center which is at min
68
                % distance
69
                for cluster_idx = 1:k
70
                    cur_dist = norm(data(cur_idx ,:) - cluster_centers(
71
                        cluster_idx ,:));
                    if(cur_dist < min_dist)</pre>
72
                         min_dist = cur_dist;
73
                         \min_{i} dx = cluster_{i} dx;
74
                    end
75
                end
76
                cluster_assignments(cur_idx,1) = min_idx;
77
           end
78
79
           for cluster_idx = 1:k
80
                % get the mean of each cluster
81
                cluster_centers(cluster_idx,:) = mean(data(cluster_assignments
82
                   = cluster_idx,:));
           end
83
84
           change = sum(prev_assignments ~= cluster_assignments);
85
86 %
              fprintf('Number of changes in iter %d: %d \ ', count_iter, change);
              show_plot(raw_data, cluster_assignments); % plot the clusters
87 %
       end
88
89
       SSE = compute_SSE(raw_data, cluster_assignments);
90
       fprintf('Final SSE for k=\%d: \%f\n', k, SSE);
91
92
93 end
94
95
96 function [SSE] = compute_SSE(data, cluster_assignments)
97 % Function to compute Sum of Squared Error
98 % INPUTS:
99 %
       data = the \ dataset \ used \ for \ clustering
100 %
       cluster\_assignments = labels for each sample in the data
101 %
       cluster\_centers = the centers found for each cluster
102 %
103 % OUTPUT:
       SSE = final sum of squared errors for the cluster config.
104 %
105
       num_clusters = size(unique(cluster_assignments),1);
106
       SSE = 0;
107
       for cluster_no = 1:num_clusters
108
            cluster_center = mean(data(cluster_assignments == cluster_no ,:));
109
           SSE = SSE + norm(data(cluster_assignments=cluster_no ,:)-
110
               cluster_center)^2;
```

```
end
111
112 end
113
114 function [] = show_plot(data, labels)
115\%
       Function to plot the cluster config.
116 %
       INPUTS:
117 %
            data = dataset used for clustering
118 %
            labels = the cluster label assigned to each sample
119 %
120 %
       OUTPUT:
121 %
            A plot representing the cluster config.
122
       k = size(unique(labels), 1);
123
       [~, num_features] = size(data);
124
125
126
       if num_features>2
127
            pcs = pca(data);
128
            reduced_data = data * pcs(:, 1:2);
129
       else
130
            reduced_data = data;
131
       end
132
133
       figure;
134
       hold on;
135
       gscatter(reduced_data(:,1),reduced_data(:,2),labels);
136
       title (strcat ('KMeans Clustering Arrangement for K=', int2str(k)));
137
       hold off;
138
       pause(2);
139
140 end
```

As further experimentation, the mall customer segmentation dataset available at Kaggle: Mall Customer has been used to test the two procedures. It has been observed that the K-means implementation is able to efficiently segregate the data into different clusters. The distribution of the final clusters for both K-Means and Spectral K-Means for varying values of k are displayed in Figure 1. The corresponding SSE values are further plotted in Figure 2. Although the SSE scores are decreasing as the value of k is increased, it does not mean the clustering is moving towards better labelling. But, it depends on the original number of categories present in the data. So, the value of k plays a crucial role over here. If it is selected appropriately, it can produce high quality clustering of the entire dataset. On the other hand, spectral k-means projects the original dataset to a low-dimension feature space. So, it is really hard to visualize the data in the original feature space. Even though the original feature space representation of the cluster arrangement is provided in Figure 1, it is somewhat deceptive because it is not the feature space where the clustering happened.

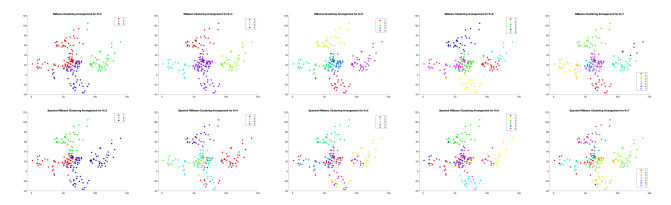


Figure 1: Final Cluster Assignments for KMeans and Spectral KMeans for varying K-values

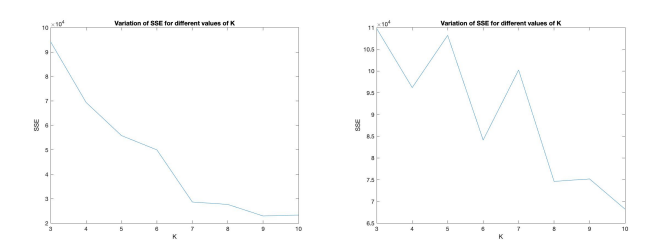
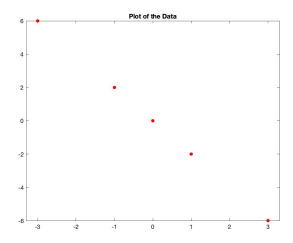


Figure 2: Plot of the SSE scores for different values of k

2 Principle Component Analysis

- 1. Suppose we have the following data points in 2d space (0, 0), (-1, 2), (-3, 6), (1, -2), (3, -6).
 - Draw them on a 2-d plot, each data point being a dot.
 - What is the first principle component? Given 1-2 sentences justification. You do not need to run Matlab to get the answer.
 - What is the second principle component? Given 1-2 sentences justification. You do not need to run Matlab to get the answer.

Response: The plot of the data in 2d space is shown in the following figure:



From the data, it can be observed that the mean of the data is (0,0). So, it is already a zero-mean design matrix. The process to find the principal components for the data matrix (let's say X) is to get the right singular vectors for X.

$$X = U \times \Sigma \times V^T$$
 [from SVD]

The principal components of X are just the vectors present in V.

SVD of X gives us the following matrix for V:

$$V = \begin{bmatrix} -0.4472 & 0.8944 \\ 0.8944 & 0.4472 \end{bmatrix}$$

The first principal component of the data is: $[-0.4472, 0.8944]^T$ The second principal component of the data is: $[0.8944, 0.4472]^T$

- 2. **Experiment**: We apply data pre-processing techniques to a collection of handwritten digit images from the USPS dataset (data in Matlab format: USPS.mat)1. You can load the whole dataset into Matlab by load USPS.mat. The matrix A contains all the images of size 16 by 16. Each of the 3000 rows in A corresponds to the image of one handwrit- ten digit (between 0 and 9). To visualize a particular image, such as the second one, first you need to convert the vector representation of the image to the matrix representation by A2 = reshape(A(2,:), 16, 16), and then use imshow(A2') for visualization. Implement Principal Component Analysis (PCA) using SVD and apply to the data using p = 10,50,100,200 principal components. Reconstruct images using the selected principal components from part 1.
 - Show the source code links for parts 1 and 2 to your github account.
 - The total reconstruction error for p = 10, 50, 100, 200.
 - A subset (the first two) of the reconstructed images for p = 10, 50, 100, 200.

Note: The USPS dataset is available at USPS Dataset. The image size is 16 by 16, thus the data dimensionality of the original dataset is 256. We used a subset of 3000 images in this homework.

Response: The steps used to find the principal components using SVD are:

- Center the data by subtracting mean from the samples.
- Perform SVD to get the right singular matrix of the centered data.
- Principal Components are now the columns of the right singular matrix.

The code is provided at the following Github link: PCA. For reference, the same code is also mentioned below:

```
1 USPS_data = importdata('Data/USPS.mat');
2 \text{ USPS\_mat} = \text{USPS\_data.A};
3 \text{ p_comb} = [10, 50, 100, 200];
                                    % number of principal components considered
4 \text{ num_p} = \text{size}(p_{\text{-comb}}, 2);
5 \text{ entry} = 1;
7 \% code for simulating PCA reconstruction as many times as user wants
8 while (entry = 1)
       img_idx = randi(size(USPS_mat, 1));
9
       img_orig = reshape(USPS_mat(img_idx,:), 16, 16);
10
11
       figure:
12
       hold on;
13
       subplot(1,num_p+1,1); imshow(img_orig'); title('Original Image');
14
       for i = 1:num_p
15
           recov_data = pca_svd(USPS_mat, p_comb(1,i));
16
           img_recov = reshape(recov_data(img_idx,:), 16, 16);
17
           subplot(1,num_p+1,i+1); imshow(img_recov'); title(streat('p=',
18
              int2str(p_comb(1,i)));
      end
19
       hold off;
20
       saveas(gcf, strcat('Images/PCA_USPS/plot_',int2str(img_idx),'.jpg'));
21
22
       pause(3);
       close;
23
24
       entry = input('Press 1 for seeing more images, Press 0 to exit: ');
25
26 end
27
28
29 function [recov_data] = pca_svd(raw_data, p)
30 % Function to perform PCA using SVD
31 %
      INPUTS:
32 %
           raw_{-}data = the data used for PCA
33 %
           p = number \ of \ principal \ components \ to \ consider
34 %
35 %
      OUTPUT:
           recov_data = data recovered from using p principal componenents
36 %
37
38
       data = raw_data;
       data = data - mean(data);
39
```

```
full form of the second struction for p=%d is: %f\n', p , reconst_error for p=%d is: %f\n', p , reconst
```

From the experimentation, it was found that the reconstruction error improved as the number of principal components were increased. The exact values are provided in Table 1.

p	Reconstruction Error
10	545.720200
50	544.896777
100	544.895231
200	544.894993

Table 1: The Reconstruction Error for different values of p

The most interesting part about this experimentation is the reconstruction of the reduced images. Some examples of reconstructions are provided in Figure 3.

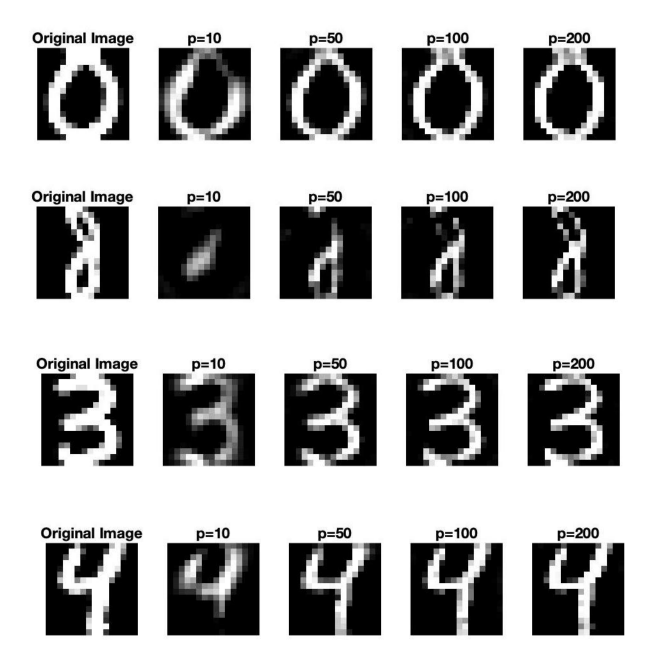


Figure 3: Visual Illustration of the Original and Reconstructed images with different p values