Problem Set 3: Clustering and Dimensionality Reduction

To run and solve this assignment, one must have a working IPython Notebook installation. The easiest way to set it up for both Windows and Linux is to install Anaconda (https://www.anaconda.com/products/individual). Then save this file to your computer, run Anaconda and choose this file in Anaconda's file explorer. Use Python 3 version. The statements below assume that you have already followed these instructions. If you are new to Python or its scientific library, Numpy, there are some nice tutorials here (https://www.learnpython.org/) and here (https://www.learnpython.org/).

To run code in a cell or to render Markdown (https://en.wikipedia.org/wiki/LaTeX (https://en.wikipedia.org/wiki/LaTeX) press Ctr+Enter or [>|] (like "play") button above. To edit any code or text cell double click on its content. To change cell type, choose "Markdown" or "Code" in the drop-down menu above. Here are some useful resources for Markdown guide (https://www.markdownguide.org/basic-syntax/) and LaTeX tutorial (https://www.overleaf.com/learn/latex/Learn LaTeX in 30 minutes) if you are not familiar with the basic syntax.

If a certain output is given for some cells, that means that you are expected to get similar results in order to receive full points (small deviations are fine). For some parts we have already written the code for you. You should read it closely and understand what it does.

Only **PDF** files are accepted for ps1 submission. To print this notebook to a pdf file, you can go to "File" -> "Download as" -> "PDF via LaTex(.pdf)" or simply use "print" in browser.

Total: 200 points (70 (Q1) + 70 (Q2) + 60 (Q3)).

1. K-means Clustering

In this exercise, you will implement the K-means algorithm and use it for image compression. You will first start on an example 2D dataset that will help you gain an intuition of how the K-means algorithm works. After, you will use the K-means algorithm for image compression by reducing the number of colors that occur in an image to only those that are most common.

1.1 Implementing K-means

The K-means algorithm is a method to automatically cluster similar data examples together. Concretely, you are given a training set $\{x_1, \cdots, x_m\}$ where $x_i \in \mathbb{R}^n$, and we want to group the data into a few cohesive clusters. The intuition behind K-means is an iterative procedure that starts by guessing the initial centroids, and then refines this guess by repeatedly assigning examples to their closest centroids and then recomputing the centroids based on the assignments.

The inner loop of the algorithm repeatedly carries out two steps:

- 1. Assigning each training example x to its closest centroid
- 2. Recomputing the mean of each centroid using the points assigned to it

The K-means algorithm will always converge to some final set of means for the centroids. Note that the converged solution may not always be ideal and depends on the initial setting of the centroids. Therefore, in practice the K-means algorithm is usually run a few times with different random initializations. One way to choose between the solutions from different random initializations is to choose the one with the lowest cost function value (distortion). You will implement the two phases of the K-means algorithm separately in the next sections.

1.2 Visualizing the data

Run the provided code below and take a look at the resulting plot to gain an understanding of the distribution of the data. It is two dimensional, with x_1 and x_2 . The "x" markers are the initial centroids of the clustering algorithm, where K=3.

In [1]:

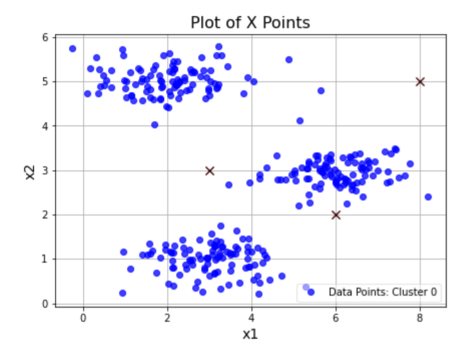
```
from future import absolute import
import random
import matplotlib.pyplot as plt
import numpy as np
import math
import scipy.io
import scipy.misc
def plot data(samples, centroids, clusters=None):
    Plot samples and color it according to cluster centroid.
    :param samples: samples that need to be plotted.
    :param centroids: cluster centroids.
    :param clusters: list of clusters corresponding to each sample.
    If clusters is None, all points are plotted with the same color.
    colors = ['blue', 'green', 'gold']
    assert centroids is not None
    if clusters is not None:
        sub samples = []
        for cluster id in range(centroids[0].shape[0]):
            sub samples.append(np.array([samples[i] for i in range(samples.shape[0])
    else:
        sub samples = [samples]
    plt.figure(figsize=(7, 5))
    for clustered samples in sub samples:
        cluster id = sub samples.index(clustered samples)
        plt.plot(clustered_samples[:, 0], clustered_samples[:, 1], 'o', color=colors
                 label='Data Points: Cluster %d' % cluster id)
    plt.xlabel('x1', fontsize=14)
    plt.ylabel('x2', fontsize=14)
    plt.title('Plot of X Points', fontsize=16)
    plt.grid(True)
    # Drawing a history of centroid movement, first centroid is black
    tempx, tempy = [], []
    for mycentroid in centroids:
        tempx.append(mycentroid[:, 0])
        tempy.append(mycentroid[:, 1])
    plt.plot(tempx, tempy, 'rx--', markersize=8)
    plt.plot(tempx[0], tempy[0], 'kx', markersize=8)
    plt.legend(loc=4, framealpha=0.5)
    plt.show(block=True)
```

In [2]:

```
datafile = 'data/data2.mat'
mat = scipy.io.loadmat(datafile)
samples = mat['X']
# samples contain 300 pts, each has two coordinates
print(samples.shape)

# Choose the initial centroids
initial_centroids = np.array([[3, 3], [6, 2], [8, 5]])
plot_data(samples, [initial_centroids])
```

(300, 2)



1.3 Finding closest centroids [20 pts]

In the cluster assignment phase of the K-means algorithm, the algorithm assigns every training example x_i to its closest centroid, given the current positions of centroids. Specifically, for every example i we set

$$c_i := \arg\min_j ||x_i - \mu_j||^2$$

where c_i is the index of the centroid that is closest to x_i , and j is the position (index) of the j-th centroid.

Your task is to complete the code in $find_closest_centroids$. This function takes the data matrix samples and the locations of all centroids and should output a one-dimensional array of clusters that holds the index (a value in $\{1, \cdots, K\}$, where K is total number of clusters) of the closest centroid to every training example. You can implement this using a loop over every training example and every centroid.

Once you have completed the code in $find_closest_centroids$, run it and you should see the output [0, 2, 1] corresponding to the centroid assignments for the first 3 examples.

In [3]:

```
def find closest centroids(samples, centroids):
    Find the closest centroid for all samples.
    :param samples: input data samples.
    :param centroids: an array of cluster centroids.
    :return: a list of cluster assignments (indices) for each sample.
    clusters = np.zeros((samples.shape[0], 1))
    for i in range(samples.shape[0]):
        min dist = []
        for j in centroids:
            dist = (samples[i][0] - j[0])**2 + (samples[i][1] - j[1])**2
            min dist.append(dist)
        min center = min(min dist)
        min index = min dist.index(min center)
        clusters[i] = min_index
    #raise NotImplementedError('Implement it yourself.')
    return clusters
```

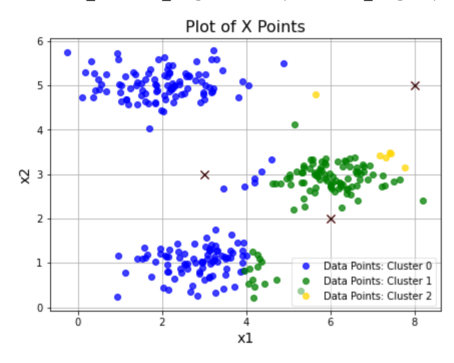
In [4]:

```
clusters = find_closest_centroids(samples, initial_centroids)

# you should see the output [0, 2, 1] corresponding to the
# centroid assignments for the first 3 examples.
print(clusters[:3].flatten())
plot_data(samples, [initial_centroids], clusters)
```

[0. 2. 1.]

<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
 cluster_id = sub_samples.index(clustered_samples)
<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
 cluster_id = sub_samples.index(clustered_samples)



1.4 Computing centroid means [20 pts]

Given assignments of every point to a centroid, the second phase of the algorithm recomputes, for each centroid, the mean of the points that were assigned to it. Specifically, for every centroid k we set

$$\mu_k := \frac{1}{|C_k|} \sum_{i \in C_k} x_i$$

where C_k is the set of examples that are assigned to centroid k. Concretely, if only two examples say x_3 and x_5 are assigned to centroid k=2, then you should update

$$\mu_2 = \frac{1}{2}(x_3 + x_5).$$

Complete the code in *get_centroids* to do this.

In [5]:

```
def get centroids(samples, clusters):
    Find the new centroid (mean) given the samples and their cluster.
    :param samples: samples.
    :param clusters: list of clusters corresponding to each sample.
    :return: an array of centroids.
    centroids = np.zeros(samples.shape[1]*(initial centroids.shape[0])).reshape(init
    #centroids = initial centroids
    simple clusters, counts = np.unique(clusters, return counts = True)
    clusters = clusters.reshape(-1)
    for i in range(len(clusters)):
        for j in range(samples.shape[1]):
            centroids[(int) (clusters[i])][j] += samples[i][j]
        #centroids[(int) (clusters[i])][1] += samples[i][1]
    for i in range(centroids.shape[0]):
        centroids[i] = centroids[i] / counts[i]
    return centroids
    #raise NotImplementedError('Implement it yourself.')
```

Once you have completed it, run_k_means below will run your code and output the centroids after the first step of K-means. The final centroids should be roughly [[1.9 5.0] [3.0 1.0] [6.0 3.0]].

The code below will produce a visualization that plots the progress of the algorithm at each iteration. The red crosses show how each step of the K-means algorithm changes the centroids. You should also see the final cluster assignments as color-coded points.

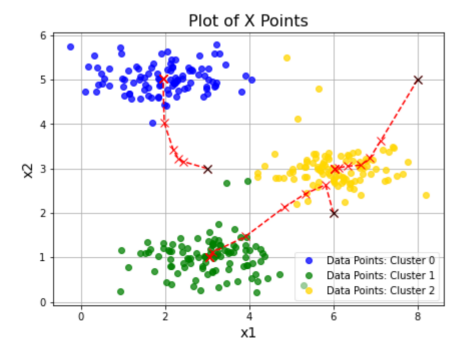
In [6]:

```
def run k means(samples, initial centroids, n iter):
    Run K-means algorithm. The number of clusters 'K' is defined by the size of init
    :param samples: samples.
    :param initial centroids: a list of initial centroids.
    :param n iter: number of iterations.
    :return: a pair of cluster assignment and history of centroids.
    centroid_history = []
    current centroids = initial centroids
    clusters = []
    for iteration in range(n iter):
        centroid history.append(current centroids)
        print("Iteration %d, Finding centroids for all samples..." % iteration)
        clusters = find_closest_centroids(samples, current_centroids)
        print("Recompute centroids...")
        current centroids = get centroids(samples, clusters)
    return clusters, centroid history
```

In [7]:

```
# Run the full K-means algorithm and plot the results
clusters, centroid_history = run_k_means(samples, initial_centroids, n_iter=10)
plot_data(samples, centroid_history, clusters)
```

```
Iteration 0, Finding centroids for all samples...
Recompute centroids...
Iteration 1, Finding centroids for all samples...
Recompute centroids...
Iteration 2, Finding centroids for all samples...
Recompute centroids...
Iteration 3, Finding centroids for all samples...
Recompute centroids...
Iteration 4, Finding centroids for all samples...
Recompute centroids...
Iteration 5, Finding centroids for all samples...
Recompute centroids...
Iteration 6, Finding centroids for all samples...
Recompute centroids...
Iteration 7, Finding centroids for all samples...
Recompute centroids...
Iteration 8, Finding centroids for all samples...
Recompute centroids...
Iteration 9, Finding centroids for all samples...
Recompute centroids...
<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
  cluster id = sub samples.index(clustered samples)
<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
  cluster_id = sub_samples.index(clustered_samples)
```



1.5 Random initialization [10 pts]

The initial assignments of centroids for the example dataset in the previous section were designed so that you could test your algorithm, i.e., if you implemented it correctly you would see the same output as we provided. In practice, a good strategy for initializing the centroids is to select random examples from the training set.

In this part of the exercise, you should complete the function $choose_random_centroids$. You should randomly permute the indices of the examples using random seed 7. Then, select the first K examples based on the random permutation of the indices. This should allow the examples to be selected at random without the risk of selecting the same example twice. You will see how the random initialization affects the first few iterations of clustering, and also possibly, results in a different cluster assignment.

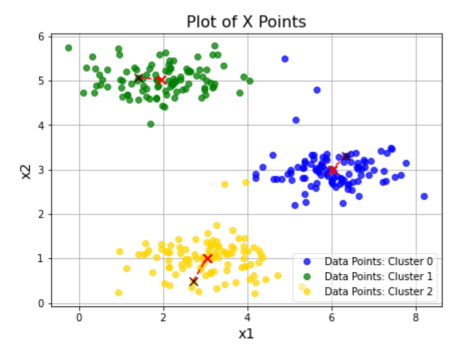
In [8]:

```
def choose_random_centroids(samples, K):
    """
    Randomly choose K centroids from samples.
    :param samples: samples.
    :param K: K as in K-means. Number of clusters.
    :return: an array of centroids.
    """
    init_centroids = np.zeros([K,samples.shape[1]])
    np.random.seed(7)
    perm_samples = np.random.permutation(samples)
    for i in range(K):
        init_centroids[i] = perm_samples[i]
    return init_centroids
    #raise NotImplementedError('Implement it yourself.')
```

In [9]:

```
# Let's choose random initial centroids and see the resulting
# centroid progression plot.. run it a few times, see if you can
# find a particularly bad case (no need to report it)
clusters, centroid_history = run_k_means(samples, choose_random_centroids(samples, plot_data(samples, centroid_history, clusters)
```

```
Iteration 0, Finding centroids for all samples...
Recompute centroids...
Iteration 1, Finding centroids for all samples...
Recompute centroids...
Iteration 2, Finding centroids for all samples...
Recompute centroids...
Iteration 3, Finding centroids for all samples...
Recompute centroids...
Iteration 4, Finding centroids for all samples...
Recompute centroids...
Iteration 5, Finding centroids for all samples...
Recompute centroids...
Iteration 6, Finding centroids for all samples...
Recompute centroids...
Iteration 7, Finding centroids for all samples...
Recompute centroids...
Iteration 8, Finding centroids for all samples...
Recompute centroids...
Iteration 9, Finding centroids for all samples...
Recompute centroids...
<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
  cluster id = sub samples.index(clustered samples)
<ipython-input-1-f3fale7bla93>:34: DeprecationWarning: elementwise com
parison failed; this will raise an error in the future.
  cluster id = sub samples.index(clustered samples)
```



1.6 Image compression with K-means [20 pts]

In this exercise, you will apply your implemented K-means algorithm to image compression.

In a 24-bit color representation of an image, each pixel is represented as three 8-bit unsigned integers (ranging from 0 to 255) that specify the red, green and blue intensity values. This is often referred to as the RGB encoding. Our image contains thousands of colors, and in this part of the exercise, you will reduce the number of colors to 16.

By making this reduction, it is possible to represent the photo in an efficient way (compressed). Specifically, you only need to store the RGB values of the 16 selected colors, and for each pixel in the image you now need to only store the index of the color at that location (where only 4 bits are necessary to represent 16 possibilities).

In this exercise, you will use the K-means algorithm to select the 16 colors that will be used to represent the compressed image. Concretely, you will treat every pixel in the original image as a data example and use the K-means algorithm to find the 16 colors that best group (cluster) the pixels in the 3-dimensional RGB space. Once you have computed the cluster centroids on the image, you will then use the 16 colors to replace the pixels in the original image.

The code below creates a three-dimensional matrix *bird_small* whose first two indices identify a pixel position and whose last index represents red, green, or blue. For example, *bird_small(50, 33, 2)* gives the blue intensity of the pixel at row 50 and column 33.

In [10]:

```
from __future__ import absolute_import

import matplotlib.pyplot as plt
import numpy as np
import scipy
import imageio

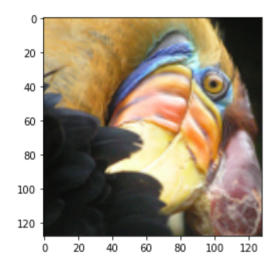
datafile = 'data/bird_small.png'
# This creates a three-dimensional matrix bird_small whose first two indices
# identify a pixel position and whose last index represents red, green, or blue.
#bird_small = scipy.misc.imread(datafile)
bird_small = imageio.imread(datafile)

print("bird_small shape is ", bird_small.shape)
plt.imshow(bird_small)
```

```
bird_small shape is (128, 128, 3)
```

Out[10]:

<matplotlib.image.AxesImage at 0x7f83786e0850>



Write code below to call your K-means function to cluster the pixels colors in the image into K clusters.

After finding the top K = 16 colors to represent the image, assign each pixel position to its closest centroid using the findClosestCentroids function. This allows you to represent the original image using the centroid assignments of each pixel. Notice that you have significantly reduced the number of bits that are required to describe the image. The original image required 24 bits for each of the 128×128 pixel locations, resulting in a total of $128 \times 128 \times 24 = 393,216$ bits.

The new representation requires some overhead storage in the form of a dictionary of 16 colors, each of which require 24 bits, but the image itself now only requires 4 bits per pixel location. The final number of bits used is therefore $16 \times 24 + 128 \times 128 \times 4 = 65,920$ bits, which corresponds to compressing the original image by about a factor of 6.

Finally, show the effect of the compression by reconstructing the image based only on the centroid assignments.

In [11]:

```
# Divide every entry in bird small by 255 so all values are in the range of 0 to 1
bird small = bird small / 255.
# Unroll the image to shape (16384,3) (16384 is 128*128)
bird small = bird small.reshape(-1, 3)
# Run k-means on this data, forming 16 clusters, with random initialization
#raise NotImplementedError('Implement this part.')
initial centroids = choose random centroids(bird small, 16)
clusters, centroids = run k means(bird small, initial centroids, 10)
# Now we have 16 centroids, each representing a color.
# Let's assign an index to each pixel in the original image dictating
# which of the 16 colors it should be. Use your find closest centroids
# function and store the result in "clusters"
#raise NotImplementedError('Implement it yourself.')
#clusters = find closest centroids(bird small, centroids)
# Now loop through the original image and form a new image
# that only has 16 colors in it
#raise NotImplementedError('Implement it yourself.')
final image = np.zeros((clusters.shape[0], 3))
for i in range(bird small.shape[0]):
    final image[i] = centroids[-1][(int) (clusters[i])]
#print()
# Reshape the original image and the new, final image and draw them
# To see what the "compressed" image looks like
plt.figure()
plt.imshow(bird small.reshape(128, 128, 3))
plt.figure()
plt.imshow(final image.reshape(128, 128, 3))
plt.show()
Iteration 0, Finding centroids for all samples...
Recompute centroids...
Iteration 1, Finding centroids for all samples...
Recompute centroids...
Iteration 2, Finding centroids for all samples...
Recompute centroids...
Iteration 3, Finding centroids for all samples...
Recompute centroids...
Iteration 4, Finding centroids for all samples...
```

```
Recompute centroids...

Iteration 1, Finding centroids for all samples...

Recompute centroids...

Iteration 2, Finding centroids for all samples...

Recompute centroids...

Iteration 3, Finding centroids for all samples...

Recompute centroids...

Iteration 4, Finding centroids for all samples...

Recompute centroids...

Iteration 5, Finding centroids for all samples...

Recompute centroids...

Iteration 6, Finding centroids for all samples...

Recompute centroids...

Iteration 7, Finding centroids for all samples...

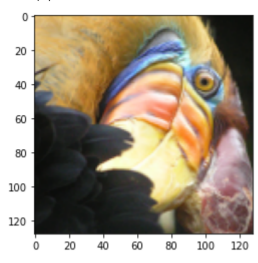
Recompute centroids...

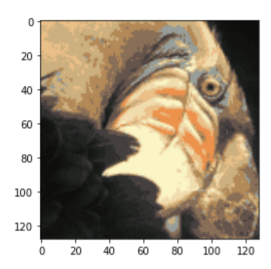
Iteration 8, Finding centroids for all samples...

Recompute centroids...

Iteration 9, Finding centroids for all samples...

Recompute centroids...
```





2. Principal Components Analysis

In this exercise, you will use principal component analysis (PCA) to perform dimensionality reduction. You will first experiment with an example 2D dataset to get intuition on how PCA works, and then use it on a bigger dataset of 5000 face image dataset. We will help you step through the first half of the exercise using provided code.

2.1 Example Dataset

To help you understand how PCA works, you will first start with a 2D dataset which has one direction of large variation and one of smaller variation. The code below will plot the training data.

In this part of the exercise, you will visualize what happens when you use PCA to reduce the data from 2D to 1D. In practice, you might want to reduce high dimensional data (such as from 256 to 50 dimensions), but using lower dimensional data in this example allows us to better visualize the algorithm.

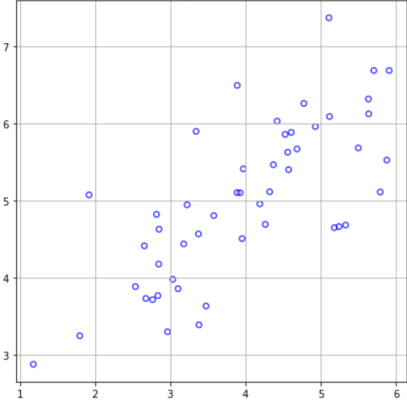
In [12]:

```
import matplotlib.pyplot as plt
import numpy as np
import scipy
import scipy.io

datafile = 'data/datal.mat'
mat = scipy.io.loadmat(datafile)
samples = mat['X']

plt.figure(figsize=(7, 7))
plt.scatter(samples[:, 0], samples[:, 1], s=30, facecolors='none', edgecolors='b')
plt.title("Example Dataset", fontsize=18)
plt.grid(True)
```





2.2 Implementing PCA [20 pts]

In this part of the exercise, you will implement PCA. PCA consists of two computational steps: First, you compute the covariance matrix of the data. Then, you use SVD (the scipy.linalg SVD function) to compute the eigenvectors U_1, U_2, \cdots, U_n . These will correspond to the principal components of variation in the data.

Before using PCA, it is important to first normalize the data. Normalize each feature by subtracting the mean value (to have zero mean), and scaling (to have unit variance) so that they are in the same range.

After normalizing the data, your task is to complete the code to compute the principal components of the dataset. First, you should compute the covariance matrix of the data, which is given by:

$$\Sigma = \frac{1}{m} X^T X$$

where X is the data matrix with examples in rows, and m is the number of examples. Note that here Σ refers to a $n \times n$ matrix and not the summation operator. After computing the covariance matrix, you can run SVD on it to compute the principal components.

Once you have completed the code, the script below will run PCA on the example dataset and plot the corresponding principal components found. The script will also output the top principal component (eigenvector) found, which should be around [-0.707 -0.707].

In [13]:

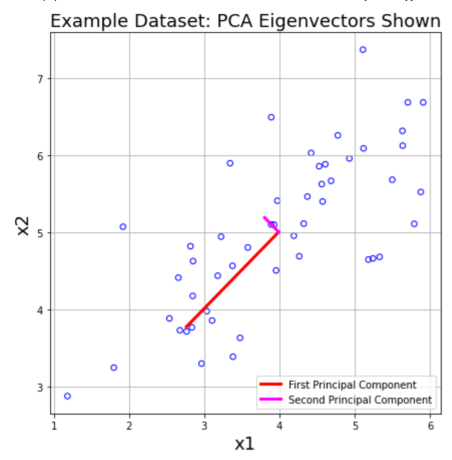
```
import scipy.linalg
def feature normalize(samples):
    Feature-normalize samples
    :param samples: samples.
    :return: normalized feature
    #raise NotImplementedError('Implement it yourself.')
    means = []
    stds = []
    norm samples = np.zeros([samples.shape[0],samples.shape[1]])
    for i in range(samples.shape[1]):
        stds.append(np.std(samples[:,i]))
        means.append(np.mean(samples[:,i]))
        for j in range(samples.shape[0]):
            norm_samples[j][i] = (samples[j][i] - means[i]) / stds[i]
    return means, stds, norm_samples
def get usv(sample norm):
    # Compute the covariance matrix
    # Run single value decomposition to get the U principal component matrix
    # raise NotImplementedError('Implement it yourself.')
    cov mat = (sample norm.T@sample norm) / sample norm.shape[0]
    U,S,V = np.linalg.svd(cov mat)
    return U,S,V
```

In [14]:

```
# Feature normalize
means, stds, samples norm = feature normalize(samples)
# Run SVD
U, S, V = get usv(samples norm)
# output the top principal component (eigen- vector) found
# should expect to see an output of about [-0.707 -0.707]"
print('Top principal component is ', U[:, 0])
plt.figure(figsize=(7, 7))
plt.scatter(samples[:, 0], samples[:, 1], s=30, facecolors='none', edgecolors='b')
plt.title("Example Dataset: PCA Eigenvectors Shown", fontsize=18)
plt.xlabel('x1', fontsize=18)
plt.ylabel('x2', fontsize=18)
plt.grid(True)
# To draw the principal component, you draw them starting
# at the mean of the data
plt.plot([means[0], means[0] + S[0] * U[0, 0]],
        [means[1], means[1] + S[0] * U[1, 0]],
        color='red', linewidth=3,
        label='First Principal Component')
plt.plot([means[0], means[0] + S[1] * U[0, 1]],
        [means[1], means[1] + S[1] * U[1, 1]],
        color='fuchsia', linewidth=3,
        label='Second Principal Component')
plt.legend(loc=4)
```

```
Top principal component is [-0.70710678 -0.70710678]
Out[14]:
```

<matplotlib.legend.Legend at 0x7f83789756a0>



2.3 Dimensionality Reduction with PCA

After computing the principal components, you can use them to reduce the feature dimension of your dataset by projecting each example onto a lower dimensional space, $x^i \to z^i$ (e.g., projecting the data from 2D to 1D). In this part of the exercise, you will use the eigenvectors returned by PCA and project the example dataset into a 1-dimensional space.

In practice, if you were using a learning algorithm such as linear regression or perhaps neural networks, you could now use the projected data instead of the original data. By using the projected data, you can train your model faster as there are fewer dimensions in the input.

2.3.1 Projecting the data onto the principal components [10 pts]

You should now complete the code in $project_data(samples, U, K)$. Specifically, you are given a dataset of samples, the principal components U, and the desired number of dimensions to reduce to K. You should project each example in samples onto the top K components in U. Note that the top K components in U are given by the first K columns of U, that is $reduced_U = U(:, 1:K)$.

Once you have completed the code, it will project the first example onto the first dimension and you should see a value of about 1.49 (or possibly -1.49, if you got -U1 instead of U1).

In [15]:

```
def project_data(samples, U, K):
    """
    Computes the reduced data representation when
    projecting only on to the top "K" eigenvectors
    """
    # Reduced U is the first "K" columns in U
    # raise NotImplementedError('Implement it yourself.')
    reduced_U = U[:,:K]
    z = reduced_U.T@(samples.T)
    return z

# project the first example onto the first dimension
# should see a value of about 1.481"
z = project_data(samples_norm, U, 1)
print('Projection of the first example is %0.3f.' % float(z[0][0]))
```

Projection of the first example is 1.496.

2.3.2 Reconstructing an approximation of the data [10 pts]

After projecting the data onto the lower dimensional space, you can approximately recover the data by projecting them back onto the original high dimensional space. Your task is to complete *recover_data(Z, U, K)* to project each example in Z back onto the original space and return the recovered approximation in *recovered_sample*.

Once you have completed the code, it will recover an approximation of the first example and you should see a value of about [-1.05 -1.05].

In [16]:

```
def recover_data(Z, U, K):
    # raise NotImplementedError('Implement it yourself.')
    reduced_U = U[:,:K]
    recovered_samples = ((np.mat(reduced_U.T).I)@np.mat(Z)).T

    return recovered_samples.getA()

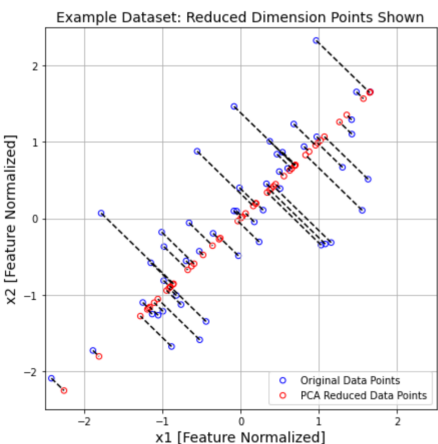
# reconstruct the samples
recovered_sample = recover_data(z, U, 1)
print('Recovered approximation of the first example is ', recovered_sample[0])
```

Recovered approximation of the first example is [-1.05805279 -1.05805 279]

2.3.3 Visualizing the projections [10 pts]

After completing both *project_data* and *recover_data*, the script below should now perform both the projection and approximate reconstruction to show how the projection affects the data. In the plot, the original data points are indicated with the blue circles, while the projected data points are indicated with the red circles. The projection effectively only retains the information in the direction given by U1. If the plot looks correct, you get full points for this part.

In [17]:



2.4 Face Image Dataset [20 pts]

In this part of the exercise, you will run your PCA algorithm on face images to see how it can be used in practice for dimension reduction. The file *faces.mat* contains a dataset X of face images, each 32×32 in grayscale. Each row of X corresponds to one face image vectorized into a row vector of length 1024. The next step will load and visualize the first 100 of these face images.

2.4.1 Visualizing the data

To display each row of X as an image, we need to first reshape it into a 32x32 array. Implement the get datum img function below and run the visualization code.

In [18]:

```
import pylab
def get datum img(row):
    Creates an image from a single np array with shape 1x1024
    :param row: a single np array with shape 1x1024
    :return: the constructed image, np array of shape 32 x 32
    size = (int) (math.sqrt(row.shape[0]))
    return row.reshape((size, size)).T
    #raise NotImplementedError('Implement it yourself.')
def display_data(samples, num_rows=10, num columns=10):
    Function that picks the first 100 rows from X, creates an image from each,
    then stitches them together into a 10x10 grid of images, and shows it.
    width, height = 32, 32
    num rows, num columns = num rows, num columns
    big picture = np.zeros((height * num rows, width * num columns))
    row, column = 0, 0
    for index in range(num_rows * num_columns):
        if column == num columns:
            row += 1
            column = 0
        img = get datum img(samples[index])
        big picture[row * height:row * height + img.shape[0], column * width:column
        column += 1
    plt.figure(figsize=(10, 10))
    plt.imshow(big picture, cmap=pylab.gray())
```

In [19]:

```
datafile = 'data/faces.mat'
mat = scipy.io.loadmat(datafile)
samples = mat['X']
display_data(samples)
```



2.4.1 PCA on Faces

To run PCA on the face dataset, we first normalize the dataset by normalizing each feature from the data matrix X to have zero mean and unit variance.

After running PCA, you will obtain the principal components of the dataset. Notice that each principal component in U (each row) is a vector of length n (for the face dataset, n = 1024). It turns out that we can visualize these principal components by reshaping each of them into a 32×32 matrix that corresponds to the pixels in the original dataset. The code cell below displays the first 36 principal components that describe the largest variations. If you want, you can also change the code to display more principal components to see how they capture more and more details.

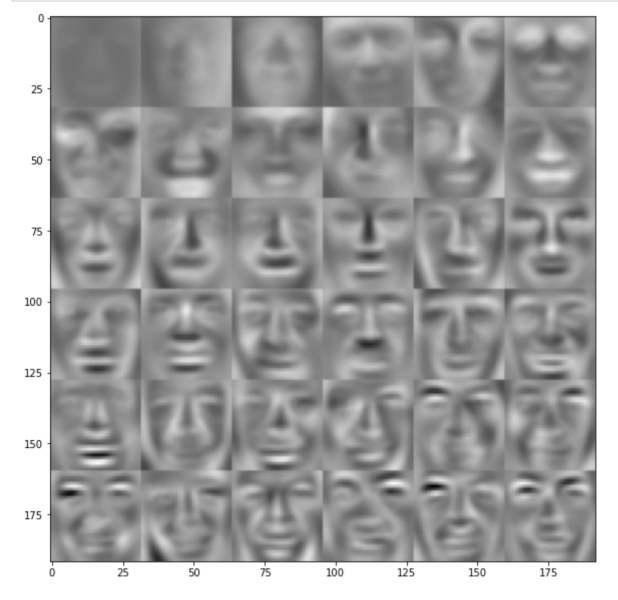
In [20]:

```
# Feature normalize
# raise NotImplementedError('Implement it yourself.')
maens, stds, samples_norm = feature_normalize(samples)

# Run SVD
# raise NotImplementedError('Implement it yourself.')
U, S, V = get_usv(samples_norm)

# Visualize the top 36 eigenvectors found
# raise NotImplementedError('Implement it yourself.')

reduced_U = U[:,:36]
display_data(reduced_U.T,6,6)
```



2.4.2 Dimensionality Reduction

Now that you have computed the principal components for the face dataset, you can use it to perform dimensionality reduction. Reducing the dimensionality with PCA would allow you to use a learning algorithm with a smaller input size (e.g., 100 dimensions) instead of the original 1024 dimensions.

The next step will project the face dataset onto only the first 100 principal components. Concretely, each face image is now described by a vector $z_i \in \mathbb{R}^{100}$.

To understand what is lost in the dimension reduction, you can recover the data using only the projected dataset. In the code below, an approximate recovery of the data is performed and the projected face images are displayed. Comparing the reconstruction to the originals, you can observe that the general structure and appearance of the face are retained while fine details are lost. This is a remarkable reduction (more than 10×) in the dataset size that can help speed up your learning algorithm significantly. For example, if you were training a logistic regression model to perform face recognition (given a face image, predict the identity of the person), you can use the reduced input of only 100 dimensions instead of the original pixels.

In [21]:

```
# Project each image down to 100 dimensions
z = project_data(samples_norm, U, 100) # complete this code

# Attempt to recover the original data
recovered_samples = recover_data(z, U, 100) #complete this code

# Plot the dimension-reduced data
display_data(recovered_samples)
plt.show()
```



3. Written Questions

3.1 Gaussian Mixtures [20 pts]

Read the beginning of Section 9.2 in Bishop which describes Gaussian mixture models, and solve Problem 9.3:

Consider a Gaussian mixture model in which the marginal distribution p(z) for the latent variable is given by (9.10) and the conditional distribution p(x|z) for the observed variable is given by (9.11). Show that the marginal distribution p(x) obtained by summing p(z)p(x|z) over all possible values of z is a Gaussian mixture of the form (9.7).

* Solution: *

The k-dimensional binary variable z uses a 1-of-K representation: $z = \{z_1, z_2, \dots, z_K\}$

,which is a one-hot vector.(Only one v_k equals 1 and all other are 0).

Therefore, we have: $\sum_{k} z_{k} = 1$, $z_{k} \in \{0, 1\}$.

And we have: $0 \le \pi_k \le 1$, $\sum_{k=1}^K \pi_k = 1$, $p(z_k = 1) = \pi_k$.

So the marginal distribution p(z) for the latent variable:

$$p(z) = \prod_{k=1}^{K} \pi_k^{z_k}$$

Similarly, the conditional distribution of x given a particular z is:

$$p(x|z) = \prod_{k=1}^{K} \mathcal{N}(x|\mu_k, \Sigma_k)^{z_k}$$

The joint distribution is given by p(z)p(x|z), and the marginal distribution can be derived as:

$$p(x) = \sum_{z} p(z)p(x|z) = \sum_{k=1}^{K} \mathcal{N}(x|\mu_k, \Sigma_k)^{z_k}$$

Since z is a one-hot vector, and every element z_k is independent with each other, we have:

$$p(x) = \sum_{k=1}^{K} \mathcal{N}(x|\mu_k, \Sigma_k)^{z_k} = \sum_{k=1}^{K} \mathcal{N}(x|\mu_k, \Sigma_k) p(z_k = 1) = \sum_{k=1}^{K} \pi_k \mathcal{N}(x|\mu_k, \Sigma_k)$$

3.2 K-means vs GMM [20 pts]

Discuss how you would modify K-means to make it a Gaussian mixture model. Compare and contrast the two methods.

* Solution: *

For the Gaussian mixture model, given a dataset of observation $\{x_1, x_2, \cdots, x_N\}$, the log-likelihood function can be written as:

$$ln \ p(X|\pi, \mu, \Sigma) = \sum_{n=1}^{N} ln[\sum_{k=1}^{K} \pi_k \mathcal{N}(x_n|\mu_k, \Sigma_k)]$$

Then we use EM alglrithm to derive the solution.

(1)E-Step: Evaluate the responsibilities using the parameter.

$$\gamma(z_{nk}) = \frac{\pi_k \mathcal{N}(x_n | \mu_k, \Sigma_k)}{\sum_{j=1}^K \pi_j \mathcal{N}(x_n | \mu_j, \Sigma_j)}$$

(2)M-Step: Re-estimate the parameters using the updated responsibilities.

$$\mu_k^{new} = \frac{1}{N_k} \sum_{n=1}^N \gamma(z_n k) x_n$$

$$\Sigma_k^{new} = \frac{1}{N_k} \sum_{n=1}^N \gamma(z_n k) (x_n - \mu_k^{new}) (x_n - \mu_k^{new})^T$$

$$\pi_k^{new} = \frac{N_k}{N}, N_k = \sum_{n=1}^N \gamma(z_n k)$$

The K-means algorithm also depends on the EM algorithm:

(1) Define an objective function

$$J = \sum_{n=1}^{N} \sum_{k=1}^{K} r_{nk} ||x_n - \mu_K||^2$$

(2) Update the parameter(the centroids) to minimize the objective function

In GMM, when the diagonal elements in the co-variance matrix Σ converge to 0, the probability $\mathcal{N}(x_i|\mu_k,\Sigma_k)$ will become a 1-of-K vector, which corresponds to the K-means algorithm.

Besides, the $\mathcal{N}(x_i|\mu_k, \Sigma_k)$ can be represented as:

$$p(x|\mu, \Sigma) = \frac{1}{(2\pi)^{n/2}|\Sigma|^{\frac{1}{2}}} exp(-\frac{1}{2}(x-\mu)^T \Sigma^{-1}(x-\mu))$$

When the co-variance matrix Σ is diagonal, it is in direct proportion to the Euclidean Distance in K-means. So both K-means and GMM depend on EM algorithm.

However, the GMM model sometimes converge quickly to a local minimum, which is not an optimal solution. Therefore, we can use K-means algorithm to initialize the Gaussian Mixture Model by changing a parameter in GMM.

Compare and Contrast: They all use EM algorithm to solve the minimum, and sometimes it can lead to a local minimum. The gaussian mixture model usually has more iterations than K-means, which leads to better robustness but more time-consuming.

3.3 PCA [20 pts]

Go through Section 12.1.2 which describes the Minimum-error formulation of PCA and perform omitted computations. Specifically, do all the derivations necessary to show that

0. Before (12.9)

$$\alpha_{nj} = \mathbf{x}_n^{\mathrm{T}} \mathbf{u}_j$$

1. Eqn (12.12)

$$z_{nj} = \mathbf{x}_n^{\mathrm{T}} \mathbf{u}_j$$

2. Eqn (12.13)

$$b_i = \overline{\mathbf{x}}^{\mathrm{T}} \mathbf{u}_i$$

3. In case of two-dimensional data space

$$Su_2 = \lambda_2 u_2$$
$$J = \lambda_2$$

* Solution: *

(0) The complete orthonormal set of D-demensional basis vector $\{u_i\}$ satisfies:

$$u_i^T u_i = \delta_{ii}$$

where $i = 1, 2, \dots, D$. Each data point can be represented by a linear combination of the basis vector:

$$x_n = \sum_{i=1}^D \alpha_{ni} u_i$$

Therefore, we have:

$$x_n^T u_j = \sum_{i=1}^D \alpha_{ni}(u_i^T u_j) = \alpha_{nj}$$

(1) The objective function is:

$$J = \frac{1}{N} \sum_{n=1}^{N} ||x_n - \tilde{x_n}||^2 = \frac{1}{N} \sum_{n=1}^{N} ||\sum_{i=1}^{D} (x_n^T u_i) u_i - \sum_{i=1}^{M} z_{ni} u_i - \sum_{i=M+1}^{D} b_i u_i||^2$$

For j = 1, 2, \cdots , M, set the derivative of J with respect to z_{ni} to 0.

$$\frac{\partial J}{\partial z_{ni}} = 2(u_i)^2 z_{ni} - 2(x_n^T u_i) u_i * u_i = 0$$

Therefore, we have $z_{ni} = \mathbf{x}_n^{\mathrm{T}} \mathbf{u}_i$.

(2) For j = M+1, M+2, ..., D, set the derivative of J with respect to b_i to 0.

$$\frac{\partial J}{\partial z_{ni}} = 2(u_i)^2 b_i - 2(\overline{x}^T u_i) u_i * u_i = 0$$

Therefore, we have $b_i = \overline{x}^T \mathbf{u}_i$.

(3) The objective function can be written as:

$$J = \frac{1}{N} \sum_{n=1}^{N} \sum_{i=M+1}^{D} (x_n^T u_i - \overline{x}^T u_i)^2 = \sum_{i=M+1}^{D} u_i^T S u_i$$

In the case of 2-dimensional data space and 1-dimensional principlal subspace M = 1. By applying the Lagrange multiplier λ_2 , the objective function is:

$$\tilde{J} = u_2^T S u_2 + \lambda_2 (1 - u_2^T u_2)$$

Set the derivative of \overline{J} with respect to u_2 to 0:

$$\frac{\partial \overline{J}}{\partial u_2} = 2Su_2 - 2\lambda_2 u_2 = 0$$

Therefore, we have $Su_2 = \lambda_2 u_2$.

The u_2 is an eigenvalue of S with eigenvalue λ_2 . Back-substitute the solution for u_2 into the J objective function:

$$J = \sum_{i=M+1}^{D} u_i^T S u_i = u_2^T S u_2 = u_2^T \lambda_2 u_2 = \lambda_2.$$