

COMP 3105A Introduction to Machine Learning

Assignment 3

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Instruction: Submit the following three files to Brightspace for marking

- A Python file `A3codes.py` that includes all your implementations. **Do not include any top-level code in this file** other than the required functions.
- A pip requirements file named `requirements.txt` that specifies the running environment including a list of Python libraries/packages and their versions required to run your codes. The current template uses Python=3.10.
- A PDF file `A3report.pdf` that includes all your answers to the written questions. It should also specify your team members (names and student IDs). Please clearly specify question/sub-question numbers in your submitted PDF report so TAs can see which question you are answering.

Do not submit a compressed file, or it will result in a mark deduction. We recommend trying your code using Colab or Anaconda/Virtualenv before submission.

Rubrics: This assignment is worth 15% of the final grade. Your codes and report will be evaluated based on their scientific qualities including but not limited to: Are the implementations correct? Is the analysis rigorous and thorough? Are the codes easily understandable (with comments)? Is the report well-organized and clear?

Policies:

- You can finish this assignment in groups of two. All members of a group will receive the same mark when the workload is shared.
- You can resubmit your work on Brightspace and the old submissions will be overwritten. However, please **submit all required files** (as opposed to only submitting the file that you want to overwrite) on Brightspace as we will only evaluate your latest submission.
- You may consult others (classmates/TAs) about general ideas but don't share codes/answers. Please specify in the PDF file any individuals you consult for the assignment. Any student found to cheat or violate this policy will receive a score of 0 for this assignment.
- You can **only** ask large language models (LLMs) general functionality questions. For example, "How do I remove all negative numbers in a NumPy array?" If you used LLMs, clearly show us how you use them (either include the prompts and replies in the report, or include URL links showing chat

history in the report). Any student found to cheat or violate this policy will receive a score of 0 for this assignment.

- Remember that you have **three** excused days *throughout the term* (rounded up to the nearest day), after which no late submission will be accepted.
 - Specifically for this assignment, you can use libraries with general utilities, such as matplotlib, numpy/scipy, cvxopt, and pandas for Python. **However, you must implement everything by yourselves without using any pre-existing implementations of the algorithms or any functions from an ML library (such as scikit-learn).** The goal is for you to really understand, step by step, how the algorithms work.
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Question 0: Data Generation & Helper Functions

In this assignment, you will implement various algorithms, including multi-class classification, principle component analysis (PCA) and k -means clustering. In the `A3helpers.py` file, we provided several functions that will be used in this assignment:

- `augmentX`, `unAugmentX` adds/removes the augmented column of all ones to/from the input matrix X .
- `convertToOneHot` converts the multi-class labels into one-hot encoding.
- `generateData` is used to generate training and test data points. There are two generative models, each generate a type of data in 2D space (controlled by the `gen_model` argument). The following shows what the generated data may look like:

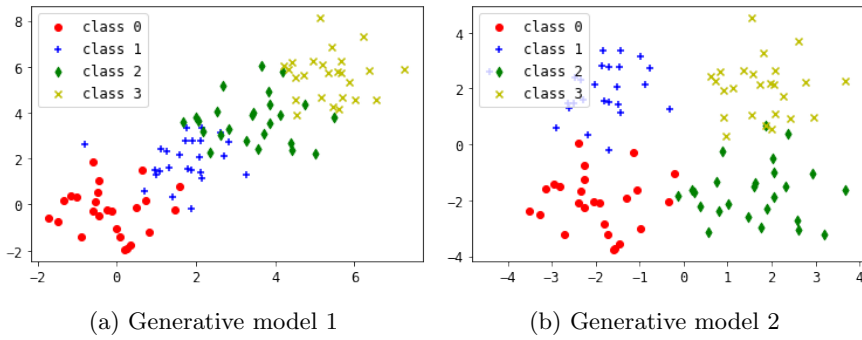


Figure 1: Generated data

- `plotModel` can be used to visualize your trained/learned models once you finish your implementation. The models you learned may look like the following:

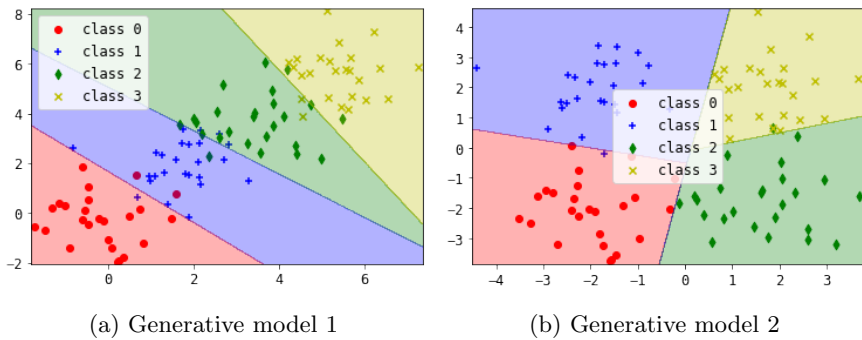


Figure 2: Classification Boundaries

- `plotImg` can help you visualize the image for Question 2.

You can also use the `A3testbed.py` to visualize your results.

Question 1 (4%) Linear Multi-Class Classifier

In this question, you will implement k -class classification using the multinomial deviance loss (or equivalently the KL divergence or cross-entropy) from scratch, in Python using NumPy/SciPy, and evaluate their performances on the synthetic datasets from above.

The input vectors are assumed to be **augmented** in this question (i.e. we assume that the input matrix \mathbf{X} or \mathbf{X}_{test} has a column of all 1s). All of the following functions must be able to handle arbitrary $n > 0$, $m > 0$, $d > 0$ and $k > 1$. The vectors and matrices are represented as NumPy arrays. Your functions shouldn't print additional information to the standard output.

- (a) (1%) Implement a Python function

```
W = minMulDev(X, Y)
```

that takes an $n \times d$ input matrix \mathbf{X} and an $n \times k$ label matrix \mathbf{Y} (where each row is a one-hot encoding of the label), and returns a $d \times k$ matrix of weights/parameters \mathbf{W} corresponding to the solution of the multinomial deviance loss:

$$W^* = \underset{W \in \mathbb{R}^{d \times k}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^n \log(\mathbf{1}_k^\top e^{W^\top \mathbf{x}_i}) - \mathbf{y}_i^\top W^\top \mathbf{x}_i$$

where the exponential and logarithm are elementwise. This problem can be solved by `scipy.optimize.minimize`.

Note that `scipy.optimize.minimize` can only handle 1D array as unknowns. Therefore, you will need to reshape your unknown variable to a $d \times k$ matrix \mathbf{W} **inside** your objective function.

Numerical issue: `exp` and/or `log` may cause numerical issues. You may want to check the `scipy.special.logsumexp` function and see how its `axis` argument works.

- (b) (1%) Implement a Python function

```
Yhat = classify(Xtest, W)
```

that takes an $m \times d$ input matrix \mathbf{X}_{test} and a $d \times k$ matrix of weights/parameters \mathbf{W} , and returns an $m \times k$ prediction matrix \mathbf{Y}_{hat} . Recall that the prediction of a data point \mathbf{x} is given by $\hat{\mathbf{y}} = \text{indmax}(W^\top \mathbf{x})$, a one-hot vector. To predict on the whole test dataset, $\hat{\mathbf{Y}}_{\text{test}} = \text{indmax}(\mathbf{X}_{\text{test}} \mathbf{W})$, where `indmax` is applied to each row.

- (c) (1%) Implement a Python function

```
acc = calculateAcc(Yhat, Y)
```

that takes an $m \times k$ prediction matrix \mathbf{Y}_{hat} and an $m \times k$ label matrix \mathbf{Y} , and returns a scalar acc that is the accuracy of the prediction. Recall that the one-hot encoding of the label shows you which one is the ground-truth/predicted class. The prediction would be correct if it matches the ground-truth label.

Debug Hint: Once you finish all the previous functions, you can run `A3testbed.py` with a controlled random seed of zero. You should get an accuracy of 87% on generative model 1 and 94% on generative model 2.

- (d) (1%) In this part, you will evaluate your implementation on the synthetic datasets from above.

We have implemented a helper function

```
train_acc, test_acc = synClsExperiments(minMulDev, classify, calculateAcc)
```

in `A3helpers.py` that calls your functions, and returns a 4×2 matrix `train_acc` of average training accuracies and a 4×2 matrix `test_acc` of average test accuracies (See Table 1 and Table 2 below).

In the PDF file, report the *averages* (over 10 runs) for each accuracy in two tables (one for training and the other for test).

Table 1: Training accuracies with different number of training dataset sizes

n	Model 1	Model 2
16		
32		
64		
128		

Table 2: Test accuracies with different number of training dataset sizes

n	Model 1	Model 2
16		
32		
64		
128		

Runtime: For efficient implementation, it will run for about 10 seconds. You may want to reduce the number of runs when testing your code.

Looking at your tables from above, analyze the results and discuss any findings you may have and the possible reason behind them.

Question 2 (7%) Principle Component Analysis

In this question, you will implement principle component analysis (PCA) from scratch, in Python using NumPy/SciPy, and check how it works for synthetic and real-world data.

The input vectors are assumed to be **unaugmented** in this question (i.e. we assume that the input matrix X or X_{test} does NOT have a column of all 1s appended to it). All of the following functions must be able to handle arbitrary $n > 0$, $m > 0$, $d > 0$ and $1 \leq k \leq d$. The vectors and matrices are represented as NumPy arrays. Your functions shouldn't print additional information to the standard output.

- (a) (1%) Implement a Python function

$$U = \text{PCA}(X, k)$$

that takes an $n \times d$ input matrix X and a scalar integer k ($1 \leq k \leq d$), and returns a $k \times d$ matrix of projecting directions U whose *rows* correspond to the top- k projecting directions with the largest variances. That is, it consists of the eigenvectors of $X^T X$ with largest eigenvalues. This problem can be solved by `scipy.linalg.eigh`.

Recall that PCA is applied **after** subtracting the mean $\mu = \frac{1}{n} \sum_{i=1}^n \mathbf{x}_i$ of X from the dataset. In other words, you need to first compute the mean $\mu \in \mathbb{R}^d$ of the input matrix X , subtract it from every row of X , and finally compute the eigenvectors of $X^T X$.

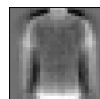
Hints: Please read the doc of `scipy.linalg.eigh` very carefully, including how it orders the eigenvalues in *ascending* order and the shape of the eigenvectors. You can also check its `subset_by_index` argument.

Debug Hint: You can visualizing the prototypes learned from PCA using the following. The A3files.zip includes an image dataset, A3train.csv, of shirt images taken from the [Fashion-MNIST dataset](#). Every row in the file is one training example. Once loaded, you can call the `plotImgs` helper function to see some samples of the images as in Fig. 3.



Figure 3: Sample images from the dataset

You can call the `plotImgs` helper function with the U learned from this dataset to see the “eigen-shirts”. One of your “eigen-shirts” should look like the following (maybe with flipped color)



- (b) (0.5%) Implement a Python function

$$X_{\text{proj}} = \text{projPCA}(X_{\text{test}}, \mu, U)$$

that takes an $m \times d$ input matrix X_{test} , a $d \times 1$ *training* mean vector μ (i.e., μ from part (a)) and a $k \times d$ projection matrix U , and returns an $m \times k$ projected matrix X_{proj} that consists of the projected features/representations of X_{test} onto the directions U .

Recall that the projection of a data point \mathbf{x} is given by $U(\mathbf{x} - \boldsymbol{\mu})$. To project the whole test dataset: $(\mathbf{X}_{\text{test}} - \boldsymbol{\mu}^\top)U^\top$. You may want to check the [broadcast rules](#) for NumPy arrays.

(c) (2%) Implement a Python function

```
A = kernelPCA(X, k, kernel_func)
```

that takes an $n \times d$ input matrix \mathbf{X} , a scalar integer k ($1 \leq k \leq d$) and a callable kernel function `kernel_func`, and returns a $k \times n$ matrix of coefficients \mathbf{A} whose *rows* correspond to the top- k coefficients with the largest variances. That is, it consists of the eigenvectors of the *centered* kernel \tilde{K} with largest eigenvalues *with proper normalization*. This problem can be solved by `scipy.linalg.eigh`.

Recall that

- Kernelized PCA requires that the data is centered in a new feature space. One can construct the kernel matrix K using the kernel function `kernel_func` and data matrix \mathbf{X} , then perform centering using

$$\tilde{K} = K - \frac{1}{n} \cdot \mathbf{1}_{n \times n} \cdot K - \frac{1}{n} \cdot K \cdot \mathbf{1}_{n \times n} + \frac{1}{n^2} \cdot \mathbf{1}_{n \times n} \cdot K \cdot \mathbf{1}_{n \times n}$$

where $\mathbf{1}_{n \times n}$ is a matrix of all ones.

- The coefficients with the largest variation in the new feature space is calculated based on the eigenvector of \tilde{K} :

$$\tilde{K}\boldsymbol{\alpha} = \lambda n \boldsymbol{\alpha}$$

where $\boldsymbol{\alpha} \in \mathbb{R}^n$ is the *rescaled* top eigenvector with the corresponding largest eigenvalue λn (note that λn is the eigenvalue instead of λ). In the case of finding multiple projecting directions in the new feature space, we identify the top- k eigenvectors of \tilde{K} . Since the eigenvectors produced by `eigh` have unit length, they are not the $\boldsymbol{\alpha}$ s. Instead, we want $\|\boldsymbol{\alpha}\|_2^2 = \frac{1}{\lambda n}$ to ensure that the corresponding projecting directions in the new feature space have unit length, which means you would need to divide the eigenvectors by the corresponding eigenvalues to get the $\boldsymbol{\alpha}$ s. Here, \mathbf{A} is a stack of the top- k $\boldsymbol{\alpha}$ s.

Hints: Convert the input \mathbf{X} into `float` before calculating any quantities. This can be important because some data are `int` and then your kernel values may overflow (but without any error/warning) and produce wrong numbers. This applies to other kernel versions of any other algorithms. The centering can be implemented much faster if you can understand what it is calculating and [how broadcast works](#).

Debug Hint: Kernelized PCA and PCA should be equivalent *when using linear kernel*. The first thing that you could check is that your centered kernel matrix should equal to the inner products of centered data matrix. Your centering would be correct if they match. Then, the projecting direction \mathbf{u} is related to the $\boldsymbol{\alpha}$ in the following way $\mathbf{u} = \sum_{i=1}^n \alpha_i \boldsymbol{\phi}_i$. In the case of linear kernel, $\boldsymbol{\phi}_i = \tilde{\mathbf{x}}_i$ is the original *centered* feature vector of the i th data point. Then the \mathbf{U} you found from **(a)** should satisfy $U = A\tilde{X}$ where \tilde{X} is the centered data matrix.

(d) (2%) Implement a Python function

```
Xproj = projKernelPCA(Xtest, Xtrain, kernel_func, A)
```

that takes an $m \times d$ input matrix \mathbf{X}_{test} , an $m \times d$ input matrix $\mathbf{X}_{\text{train}}$, a callable kernel function `kernel_func` and a $k \times n$ coefficient matrix \mathbf{A} , and returns an $m \times k$ projected matrix \mathbf{X}_{proj} that consists of the projected features/representations of \mathbf{X}_{test} onto the directions specified by \mathbf{A} .

To understand how to project a set of test points onto the kernel PCA directions, we can take a look at the linear case first. Given that $U = A\tilde{X}_{\text{train}}$, the projected value of a test set is given by

$$Y_{\text{test}} = \tilde{X}_{\text{test}} U^\top = \tilde{X}_{\text{test}} \tilde{X}_{\text{train}}^\top A^\top$$

where $\tilde{X}_{\text{test}}, \tilde{X}_{\text{train}}$ are the centered matrices $\tilde{X}_{\text{test}} = X_{\text{test}} - \boldsymbol{\mu}^\top$, $\tilde{X}_{\text{train}} = X_{\text{train}} - \boldsymbol{\mu}^\top$. That is, we subtract the *training* mean $\boldsymbol{\mu}$ from each row using the broadcast rule. As seen from the above equation, the inner product matrix $\tilde{X}_{\text{test}}\tilde{X}_{\text{train}}^\top$ can be replaced by a centered kernel

$$\tilde{K}_{\text{te,tr}} = K_{\text{te,tr}} - \frac{1}{n} \cdot \mathbf{1}_{m \times n} \cdot K_{\text{tr,tr}} - \frac{1}{n} \cdot K_{\text{te,tr}} \cdot \mathbf{1}_{n \times n} + \frac{1}{n^2} \cdot \mathbf{1}_{m \times n} \cdot K_{\text{tr,tr}} \cdot \mathbf{1}_{n \times n}$$

where $K_{\text{te,tr}} = \text{kernel_func}(X_{\text{test}}, X_{\text{train}}) \in \mathbb{R}^{m \times n}$ and $K_{\text{tr,tr}} = \text{kernel_func}(X_{\text{train}}, X_{\text{train}}) \in \mathbb{R}^{n \times n}$ are regular (non-centered) kernel matrices. Then the projected values are given by $Y_{\text{test}} = \tilde{K}_{\text{te,tr}} A^\top$.

Debug Hint: In the case of linear kernel, the centered kernel should be equal to the inner products of the centered test matrix and the centered training matrix. Moreover, your projected values should be identical to those in (b).

(e) (1%) In this part, you will evaluate the effect of PCA on the classification problem. Implement a Python function

```
train_acc, test_acc = synClsExperimentsPCA()
```

that returns a 2×2 matrix `train_acc` of average training accuracies and a 2×2 matrix `test_acc` of average test accuracies (See Table 3 and Table 4 below). It repeats 100 runs as follows

```
def synClsExperimentsPCA():
    n_runs = 100
    n_train = 128
    n_test = 1000
    dim_list = [1, 2]
    gen_model_list = [1, 2]
    train_acc = np.zeros([len(dim_list), len(gen_model_list), n_runs])
    test_acc = np.zeros([len(dim_list), len(gen_model_list), n_runs])

    # TODO: Change the following random seed to your GROUP number (<=3digits)
    np.random.seed(0)

    for r in range(n_runs):
        for i, k in enumerate(dim_list):
            for j, gen_model in enumerate(gen_model_list):
                Xtrain, Ytrain = generateData(n=n_train, gen_model=gen_model)
                Xtest, Ytest = generateData(n=n_test, gen_model=gen_model)

                U = PCA(Xtrain, k)
                Xtrain_proj = # TODO: call your projPCA to find the new features
                Xtest_proj = # TODO: call your projPCA to find the new features

                Xtrain_proj = augmentX(Xtrain_proj) # add augmentation
                Xtest_proj = augmentX(Xtest_proj)

                W = minMulDev(Xtrain_proj, Ytrain) # from Q1
                Yhat = classify(Xtrain_proj, W) # from Q1
                train_acc[i, j, r] = calculateAcc(Yhat, Ytrain) # from Q1

                Yhat = classify(Xtest_proj, W)
                test_acc[i, j, r] = calculateAcc(Yhat, Ytest)

    # TODO: compute the average accuracies over runs
    # TODO: return 2-by-2 train accuracy and 2-by-2 test accuracy
```

In the PDF file, report the *averages* (over 100 runs) for each accuracy in two tables (one for training and the other for test).

Table 3: Training accuracies with different number of dimensions

Dim k	Model 1	Model 2
1		
2		

Table 4: Test accuracies with different number of dimensions

Dim k	Model 1	Model 2
1		
2		

Runtime: For efficient implementation, it will run for about 30 seconds. You may want to reduce the number of runs when testing your code.

(f) (0.5%) Looking at your tables from above, analyze the results and discuss any findings you may have and the possible reason behind them.

Question 3 (4%) k -means

In this question, you will implement k -means from scratch, in Python using NumPy/SciPy, and check how it works for synthetic data.

All of the following functions must be able to handle arbitrary $n > 0$, $m > 0$, $d > 0$ and $1 < k < n$. The vectors and matrices are represented as NumPy arrays. Your functions shouldn't print additional information to the standard output.

- (a) (1%) Implement a Python function

```
Y, U, obj_val = kmeans(X, k, max_iter=1000)
```

that takes an $n \times d$ input matrix X , a scalar integer k ($1 < k < n$) and a scalar integer `max_iter` indicating the maximum number of iterations, and returns an $n \times k$ membership/assignment matrix Y , a $k \times d$ matrix of cluster centers U and a scalar `obj_val` that is the objective value achieved by the solutions. It solves the following problem locally

$$\min_{Y \in \mathcal{Y}, U} J(Y, U) = \frac{1}{2n} \|X - YU\|_F^2 \quad (1)$$

where $\mathcal{Y} = \{Y | Y \in \{0, 1\}^{n \times k}, Y\mathbf{1}_k = \mathbf{1}_n\}$

by alternating the following steps (with an initial guess of U)

1. Fix U , solve for an optimal Y^* . Specifically, compute the pairwise distance matrix D between the input matrix X and the given centers U such that $D_{ij} = \|\mathbf{x}_i - \mathbf{u}_j\|_2^2$ where \mathbf{x}_i is the i th row of X and \mathbf{u}_j is the j th row of U . Then assign each point \mathbf{x}_i to the closest cluster center. That is, the i th row of Y is given by

$$Y_i := [0, \dots, \underbrace{1}_{j^{\text{th}} \text{ position}}, \dots, 0]$$

where $j^* = \operatorname{argmin}_j D_{ij}$.

2. Fix Y , solve for an optimal U^* , which is given by $(Y^\top Y)^{-1} Y^\top X = Y^+ X$ where Y^+ is the pseudoinverse of Y .

After convergence, compute the objective value $J(Y, U)$ (see Eq. (1)) achieved by the solutions Y and U . The function should look like the following

```
def kmeans(X, k, max_iter=1000):
    n, d = X.shape
    assert max_iter > 0 and k < n
    U = # TODO: Choose k random points from X as initial centers
    for i in range(max_iter):
        D = # TODO: Compute pairwise distance between X and U
        Y = # TODO: Find the new cluster assignments
        old_U = U
        U = # TODO: Update cluster centers
        if np.allclose(old_U, U):
            break
    obj_val = (0.5 / n) * np.sum(D.min(axis=1))
    return Y, U, obj_val
```

Note: Do not control random seed inside your function. You may find `scipy.spatial.distance.cdist` and `numpy.linalg.pinv` helpful.

- (b) (1%) Implement a Python function

```
Y, U, obj_val = repeatKmeans(X, k, n_runs=100)
```

that takes an $n \times d$ input matrix X , a scalar integer k ($1 < k < n$) and a scalar integer `n_runs` indicating the number of restarts, and returns an $n \times k$ membership/assignment matrix Y , a $k \times d$ matrix cluster centers U and a scalar `obj_val`

that corresponds to the best solutions and objective value obtained during `n_runs` of your `kmeans`. It runs as follows

```
def repeatKmeans(X, k, n_runs=100):
    best_obj_val = float('inf')
    for r in range(n_runs):
        Y, U, obj_val = kmeans(X, k)
        # TODO: Compare obj_val with best_obj_val. If it is lower,
        #         then record the current Y, U and update best_obj_val

    # TODO: Return the best Y, U and best_obj_val
```

Recall that the problem in Eq. (1) is NP-hard and k -means only solves it locally instead of globally, which means every time you run `kmeans`, it may give you a different solution. That's why you should set the initial guess U *randomly*. However, we can call `kmeans` multiple (specifically, `n_runs`) times, and choose the solutions that give the *minimum* objective value among those runs.

Note: Do not set random seed in your function.

(c) (1%) Implement a Python function

```
obj_val_list = chooseK(X, k_candidates=[2,3,4,5,6,7,8,9])
```

that takes an $n \times d$ input matrix `X` and a Python list of integers `k_candidates`, and returns a list of objective values obtained for each k value in the candidate list when calling `repeatKmeans` with that k value.

Once you finish this function, call it with the following

```
Xtrain, Ytrain = generateData(n=100, gen_model=2)
obj_val_list = chooseK(Xtrain)
```

and report the `obj_val_list` in the following table.

Table 5: Objective values for different k

k	2	3	4	5	6	7	8	9
obj_val								

In the PDF file, explain how many number of clusters k you think is appropriate for this dataset, and why.

Note: Do not control random seed in your function and use the default input arguments.

(d) (1%) Implement a Python function

```
Y, obj_val = kernelKmeans(X, kernel_func, k, init_Y, max_iter=1000)
```

that takes an $n \times d$ input matrix `X`, a callable kernel function `kernel_func`, a scalar integer `k` ($1 < k < n$), an $n \times k$ initial assignment matrix `init_Y` and a scalar integer `max_iter` indicating the maximum number of iterations, and returns an $n \times k$ membership/assignment matrix `Y` and a scalar `obj_val` that is the objective value achieved by the solutions.

This function implements the kernelized version of k -means by calculating the distance in a new feature space induced by the kernel function. Specifically, the distance matrix is given by

$$D = \text{diag}(K)\mathbf{1}_k^\top + \mathbf{1}_n \text{diag}(Y^+ K(Y^+)^\top)^\top - 2K(Y^+)^\top$$

where K is the kernel matrix and `diag` converts the diagonal elements of a matrix into a column vector (`numpy.diag`). Your implementation should look like the following

```

def kernelKmeans(X, kernel_func, k, init_Y, max_iter=1000):
    n, d = X.shape
    assert max_iter > 0 and k < n
    K = kernel_func(X, X)
    Y = init_Y
    for i in range(max_iter):
        D = # TODO: Compute pairwise distance matrix
        old_Y = Y
        Y = # TODO: Find the new cluster assignments
        if np.allclose(old_Y, Y):
            break
    obj_val = (0.5 / n) * np.sum(D.min(axis=1))
    return Y, obj_val

```

Debug Hints: In the case of linear kernel, the distance matrix should be identical to the one produced by `scipy.spatial.distance.cdist` with ‘sqeuclidean’.

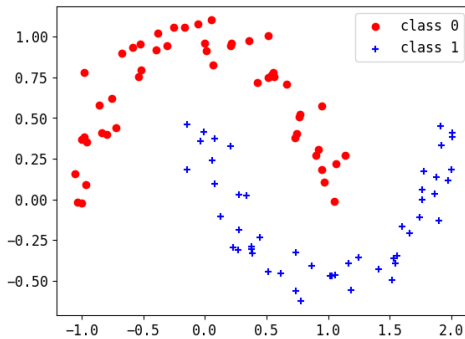
Once you finish this function, call it with the following

```

Xtrain, Ytrain = generateData(n=100, gen_model=3)
kernel_func = lambda X1, X2: gaussKernel(X1, X2, 0.25)
init_Y = # TODO: randomly initialize an assignment matrix
Y, obj_val = kernelKmeans(Xtrain, kernel_func, 2, init_Y)
plotPoints(Xtrain, Y)

```

which *may* give you a clustering plot that looks like the following:



It is very unlikely that you will get this plot in the first try due to initialization. However, given a fixed dataset `Xtrain`, if you repeat this kernelized k -means multiple times (similar to the `repeatKmeans` in (b)) with different `init_Y` matrices, and keep the `Y` with the lowest objective value, the plot should look much nicer.