# Introduction to Machine Learning Jonathan Shewchuk

HW3

#### Due: Wednesday, February 22 at 11:59 pm

- Homework 3 consists of coding assignments and math problems.
- We prefer that you typeset your answers using LaTeX or other word processing software. If you haven't yet learned LaTeX, one of the crown jewels of computer science, now is a good time! Neatly handwritten and scanned solutions will also be accepted.
- In all of the questions, **show your work**, not just the final answer.
- The assignment covers concepts on Gaussian distributions and classifiers. Some of the material may not have been covered in lecture; you are responsible for finding resources to understand it.
- Start early; you can submit models to Kaggle only twice a day!

#### **Deliverables:**

- 1. Submit your predictions for the test sets to Kaggle as early as possible. Include your Kaggle scores in your write-up. The Kaggle competition for this assignment can be found at
  - MNIST: https://www.kaggle.com/competitions/spring23-uc-berkeley-cs189-hw3-mnist
  - SPAM: https://www.kaggle.com/competitions/spring23-uc-berkeley-cs189-hw3-spam
- 2. Write-up: Submit your solution in **PDF** format to "Homework 3 Write-Up" in Gradescope.
  - On the first page of your write-up, please list students with whom you collaborated
  - Start each question on a new page. If there are graphs, include those graphs on the same pages as the question write-up. DO NOT put them in an appendix. We need each solution to be self-contained on pages of its own.
  - Only PDF uploads to Gradescope will be accepted. You are encouraged use LaTeX or Word to typeset your solution. You may also scan a neatly handwritten solution to produce the PDF.
  - **Replicate all your code in an appendix**. Begin code for each coding question in a fresh page. Do not put code from multiple questions in the same page. When you upload this PDF on Gradescope, *make sure* that you assign the relevant pages of your code from appendix to correct questions.
  - While collaboration is encouraged, *everything* in your solution must be your (and only your) creation. Copying the answers or code of another student is strictly forbidden. Furthermore, all external material (i.e., *anything* outside lectures and assigned readings, including figures and pictures) should be cited properly. We wish to remind you that consequences of academic misconduct are *particularly severe*!

- 3. Code: Submit your code as a .zip file to "Homework 3 Code".
  - Set a seed for all pseudo-random numbers generated in your code. This ensures your results are replicated when readers run your code. For example, you can seed numpy with np.random.seed(42).
  - Include a README with your name, student ID, the values of random seed (above) you used, and any instructions for compilation.
  - Do NOT provide any data files. Supply instructions on how to add data to your code.
  - Code requiring exorbitant memory or execution time might not be considered.
  - Code submitted here must match that in the PDF Write-up. The Kaggle score will not be accepted if the code provided a) does not compile or b) compiles but does not produce the file submitted to Kaggle.

# 1 Honor Code

Declare	and sig	n the	following	statement	(Mac	Preview,	PDF	Expert,	and	FoxIt	PDF	Reader,
among o	others, h	ave to	ols to let y	ou sign a P	DF fil	e):						

"I certify that all solutions are entirely my own and that I have not looked at anyone else's solution. I have given credit to all external sources I consulted."

Signature:	
Digitatare.	

### 2 Gaussian Classification

Let  $f(x \mid C_i) \sim \mathcal{N}(\mu_i, \sigma^2)$  for a two-class, one-dimensional classification problem with classes  $C_1$  and  $C_2$ ,  $P(C_1) = P(C_2) = 1/2$ , and  $\mu_2 > \mu_1$ .

- 1. Find the Bayes optimal decision boundary and the corresponding Bayes decision rule by finding the point(s) at which the posterior probabilities are equal. Use the 0-1 loss function.
- 2. Suppose the decision boundary for your classifier is x = b. The Bayes error is the probability of misclassification, namely,

$$P_e = P((C_1 \text{ misclassified as } C_2) \cup (C_2 \text{ misclassified as } C_1)).$$

Show that the Bayes error associated with this decision rule, in terms of b, is

$$P_e(b) = \frac{1}{2\sqrt{2\pi}\sigma} \left( \int_{-\infty}^b \exp\left(-\frac{(x-\mu_2)^2}{2\sigma^2}\right) dx + \int_b^\infty \exp\left(-\frac{(x-\mu_1)^2}{2\sigma^2}\right) dx \right).$$

3. Using the expression above for the Bayes error, calculate the optimal decision boundary  $b^*$  that minimizes  $P_e(b)$ . How does this value compare to that found in part 1? *Hint:*  $P_e(b)$  is convex for  $\mu_1 < b < \mu_2$ .

#### **Solution:**

1.

$$P(C_1 \mid x) = P(C_2 \mid x) \Leftrightarrow f(x \mid C_1) \frac{P(C_1)}{f(x)} = f(x \mid C_2) \frac{P(C_2)}{f(x)} \Leftrightarrow f(x \mid C_1) = f(x \mid C_2) \Leftrightarrow \mathcal{N}(\mu_1, \sigma^2) = \mathcal{N}(\mu_2, \sigma^2) \Leftrightarrow (x - \mu_1)^2 = (x - \mu_2)^2$$

Thus, the decision boundary is the point equidistant to  $\mu_1$  and  $\mu_2$ :  $b = \frac{\mu_1 + \mu_2}{2}$ .

The corresponding decision rule is, given a data point  $x \in \mathbb{R}$ :

- if  $x < \frac{\mu_1 + \mu_2}{2}$ , then classify x in class 1;
- otherwise, classify *x* in class 2.

Note that this is the centroid method.

2. Applying basic rules of probability, we have

$$P_e = P((C_1 \text{ misclassified as } C_2) \cup (C_2 \text{ misclassified as } C_1))$$
  
=  $P(C_1 \text{ misclassified as } C_2) + P(C_2 \text{ misclassified as } C_1)$   
=  $P(\text{misclassified as } C_2 \mid C_1)P(C_1) + P(\text{misclassified as } C_1 \mid C_2)P(C_2).$ 

We know  $P(C_1) = P(C_2) = \frac{1}{2}$ . Thus, we use the PDF to calculate

$$P(\text{misclassified as } C_1 \mid C_2) = \int_{-\infty}^b \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu_2)^2/(2\sigma^2)} dx.$$

$$P(\text{misclassified as } C_2 \mid C_1) = \int_b^\infty \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu_1)^2/(2\sigma^2)} dx.$$

Therefore,

$$P_e(b) = \frac{1}{2} \left( P(\text{misclassified as } C_1 \mid C_2) + P(\text{misclassified as } C_1 \mid C_2) \right)$$
$$= \frac{1}{2\sqrt{2\pi}\sigma} \left( \int_{-\infty}^b e^{-\frac{(x-\mu_2)^2}{2\sigma^2}} dx + \int_b^\infty e^{-\frac{(x-\mu_1)^2}{2\sigma^2}} dx \right).$$

3. We know the optimal decision boundary must lie between  $\mu_1$  and  $\mu_2$  (convince yourself this is the case!). Thus, we can minimize  $P_e(b)$  by taking the derivative and setting it equal to zero. Letting  $C = \frac{1}{2\sqrt{2\pi}\sigma}$ ,  $f_1(x) = e^{-(x-\mu_2)^2/(2\sigma^2)}$ , and  $f_2(x) = e^{-(x-\mu_1)^2/(2\sigma^2)}$ , we have

$$\frac{dP_e(b)}{db} = \frac{d}{db}C\left[\int_{-\infty}^b f_1(x)dx + \int_b^{\infty} f_2(x)dx\right]$$
$$= C\left[\frac{d}{db}\left(\int_{-\infty}^b f_1(x)dx\right) + \frac{d}{db}\left(\int_b^{\infty} f_2(x)dx\right)\right].$$

From the fundamental theorem of calculus, we know that, for continuous functions f(x) such that  $|f| \to 0$  for  $x \to -\infty$ ,

$$\frac{d}{dx} \int_{-\infty}^{x} f(t) \, dt = f(x).$$

Noting that  $f_1$  and  $f_2$  satisfy these properties, we apply that here to get

$$\frac{dP_e(b)}{db} = C\left[f_1(b) - f_2(b)\right].$$

Setting  $\frac{dP_e(b^*)}{db}$  to zero yields

$$f_1(b^*) = f_2(b^*) \Longrightarrow$$

$$\log(f_1(b^*)) = \log(f_2(b^*)) \Longrightarrow$$

$$\frac{(b^* - \mu_2)^2}{2\sigma^2} = \frac{(b^* - \mu_1)^2}{2\sigma^2} \Longrightarrow$$

$$(b^* - \mu_2)^2 = (b^* - \mu_1)^2.$$

This is the same constraint we saw in part (1) and thus the optimal decision boundary is equivalent. That is,

$$b^* = \frac{\mu_1 + \mu_2}{2}$$
.

### 3 Isocontours of Normal Distributions

Let  $f(\mu, \Sigma)$  be the probability density function of a normally distributed random variable in  $\mathbb{R}^2$ . Write code to plot the isocontours of the following functions, each on its own separate figure. Make sure it is clear which figure belongs to which part. You're free to use any plotting libraries or stats utilities available in your programming language; for instance, in Python you can use Matplotlib and SciPy. Choose the boundaries of the domain you plot large enough to show the interesting characteristics of the isocontours (use your judgment). Make sure we can tell what isovalue each contour is associated with—you can do this with labels or a colorbar/legend.

1. 
$$f(\mu, \Sigma)$$
, where  $\mu = \begin{bmatrix} 1 \\ 1 \end{bmatrix}$  and  $\Sigma = \begin{bmatrix} 1 & 0 \\ 0 & 2 \end{bmatrix}$ .

2. 
$$f(\mu, \Sigma)$$
, where  $\mu = \begin{bmatrix} -1 \\ 2 \end{bmatrix}$  and  $\Sigma = \begin{bmatrix} 2 & 1 \\ 1 & 4 \end{bmatrix}$ .

3. 
$$f(\mu_1, \Sigma_1) - f(\mu_2, \Sigma_2)$$
, where  $\mu_1 = \begin{bmatrix} 0 \\ 2 \end{bmatrix}$ ,  $\mu_2 = \begin{bmatrix} 2 \\ 0 \end{bmatrix}$  and  $\Sigma_1 = \Sigma_2 = \begin{bmatrix} 2 & 1 \\ 1 & 1 \end{bmatrix}$ .

(Yes, this is a difference between two PDFs. No, it is not itself a valid PDF. Just plot its isocontours anyway.)

4. 
$$f(\mu_1, \Sigma_1) - f(\mu_2, \Sigma_2)$$
, where  $\mu_1 = \begin{bmatrix} 0 \\ 2 \end{bmatrix}$ ,  $\mu_2 = \begin{bmatrix} 2 \\ 0 \end{bmatrix}$ ,  $\Sigma_1 = \begin{bmatrix} 2 & 1 \\ 1 & 1 \end{bmatrix}$  and  $\Sigma_2 = \begin{bmatrix} 2 & 1 \\ 1 & 4 \end{bmatrix}$ .

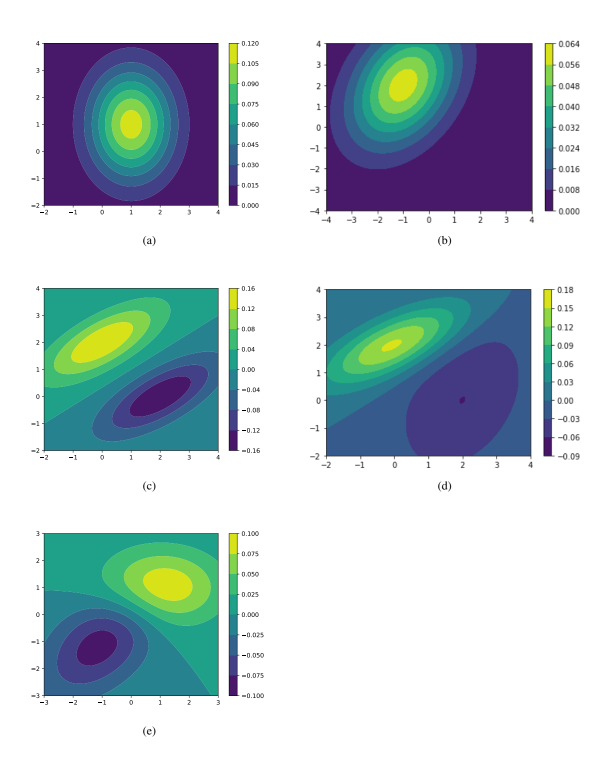
5. 
$$f(\mu_1, \Sigma_1) - f(\mu_2, \Sigma_2)$$
, where  $\mu_1 = \begin{bmatrix} 1 \\ 1 \end{bmatrix}$ ,  $\mu_2 = \begin{bmatrix} -1 \\ -1 \end{bmatrix}$ ,  $\Sigma_1 = \begin{bmatrix} 2 & 0 \\ 0 & 1 \end{bmatrix}$  and  $\Sigma_2 = \begin{bmatrix} 2 & 1 \\ 1 & 2 \end{bmatrix}$ .

```
import matplotlib.pyplot as plt
import numpy as np
import scipy.stats
def plot_contours():
    fig = plt.figure(figsize=(10,10))
    ax0 = fig.add_subplot(111)
    ax0.contour(rv.pdf(pos).reshape(500,500))
    plt.show()
# Part a
# Generate grid of points at which to evaluate pdf
x = np.linspace(-2, 4, 500)
y = np.linspace(-2, 4, 500)
X,Y = np.meshgrid(x, y)
pos = np.array([Y, X]).T
rv = scipy.stats.multivariate_normal([1, 1], [[1, 0], [0, 2]])
Z = rv.pdf(pos)
plt.contourf(X, Y, Z)
plt.colorbar()
plt.show()
# Part b
```

```
x = np.linspace(-4, 4, 500)
y = np.linspace(-4, 4, 500)
X,Y = np.meshgrid(x, y)
pos = np.array([Y, X]).T
rv = scipy.stats.multivariate_normal([-1, 2], [[2, 1], [1, 4]])
Z = rv.pdf(pos)
plt.contourf(X, Y, Z)
plt.colorbar()
plt.show()
# Part c
x = np.linspace(-2, 4, 500)
y = np.linspace(-2, 4, 500)
X,Y = np.meshgrid(x, y)
pos = np.array([Y, X]).T
rv1 = scipy.stats.multivariate_normal([0, 2], [[2, 1], [1, 1]])
rv2 = scipy.stats.multivariate_normal([2, 0], [[2, 1], [1, 1]])
Z = rv1.pdf(pos) - rv2.pdf(pos)
plt.contourf(X, Y, Z)
plt.colorbar()
plt.show()
# Part d
x = np.linspace(-2, 4, 500)
y = np.linspace(-2, 4, 500)
X,Y = np.meshgrid(x, y)
pos = np.array([Y, X]).T
rv1 = scipy.stats.multivariate_normal([0, 2], [[2, 1], [1, 1]])
rv2 = scipy.stats.multivariate_normal([2, 0], [[2, 1], [1, 4]])
Z = rv1.pdf(pos) - rv2.pdf(pos)
plt.contourf(X, Y, Z)
plt.colorbar()
plt.show()
# Part e
x = np.linspace(-3, 3, 500)
y = np.linspace(-3, 3, 500)
X,Y = np.meshgrid(x, y)
pos = np.array([Y, X]).T
rv1 = scipy.stats.multivariate_normal([1, 1], [[2, 0], [0, 1]])
rv2 = scipy.stats.multivariate_normal([-1, -1], [[2, 1], [1, 2]])
Z = rv1.pdf(pos) - rv2.pdf(pos)
plt.contourf(X, Y, Z)
plt.colorbar()
plt.show()
```

# 4 Eigenvectors of the Gaussian Covariance Matrix

Consider two one-dimensional random variables  $X_1 \sim \mathcal{N}(3,9)$  and  $X_2 \sim \frac{1}{2}X_1 + \mathcal{N}(4,4)$ , where  $\mathcal{N}(\mu,\sigma^2)$  is a Gaussian distribution with mean  $\mu$  and variance  $\sigma^2$ . Write a program that draws n = 100 random two-dimensional sample points from  $(X_1, X_2)$ . For each sample point, the value of  $X_2$  is a function of the value of  $X_1$  for that *same* sample point, but the sample points are independent of each other. In your code, make sure to choose and set a fixed random number seed for whatever random number generator you use, so your simulation is reproducible, and document your choice of random number seed and random number generator in your write-up. For each of the following



parts, include the corresponding output of your program.

- 1. Compute the mean (in  $\mathbb{R}^2$ ) of the sample.
- 2. Compute the  $2 \times 2$  covariance matrix of the sample (based on the sample mean, not the true mean—which you would not know given real-world data).
- 3. Compute the eigenvectors and eigenvalues of this covariance matrix.
- 4. On a two-dimensional grid with a horizonal axis for  $X_1$  with range [-15, 15] and a vertical axis for  $X_2$  with range [-15, 15], plot
  - (i) all n = 100 data points, and
  - (ii) arrows representing both covariance eigenvectors. The eigenvector arrows should originate at the mean and have magnitudes equal to their corresponding eigenvalues.

Hint: make *sure* your plotting software is set so the figure is square (i.e., the horizontal and vertical scales are the same). Not doing that may lead to hours of frustration!

5. Let  $U = [v_1 \ v_2]$  be a  $2 \times 2$  matrix whose columns are the **unit** eigenvectors of the covariance matrix, where  $v_1$  is the eigenvector with the larger eigenvalue. We use  $U^{\top}$  as a rotation matrix to rotate each sample point from the  $(X_1, X_2)$  coordinate system to a coordinate system aligned with the eigenvectors. (As  $U^{\top} = U^{-1}$ , the matrix U reverses this rotation, moving back from the eigenvector coordinate system to the original coordinate system). *Center* your sample points by subtracting the mean  $\mu$  from each point; then rotate each point by  $U^{\top}$ , giving  $x_{\text{rotated}} = U^{\top}(x - \mu)$ . Plot these rotated points on a new two dimensional-grid, again with both axes having range [-15, 15]. (You are not required to plot the eigenvectors, which would be horizontal and vertical.)

In your plots, clearly label the axes and include a title. Moreover, make sure the horizontal and vertical axis have the same scale! The aspect ratio should be one.

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

np.random.seed(9)

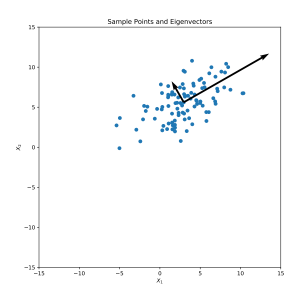
X = np.random.normal(loc=3, scale=3, size=100)
Y = np.random.normal(loc=4, scale=2, size=100)
sample = np.array([np.array((x, 0.5 * x + y)) for (x, y) in zip(X, Y)])

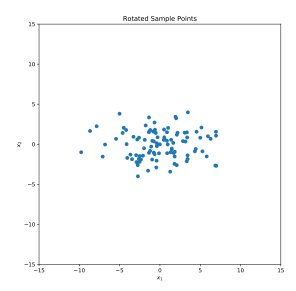
# Part a (compute the sample mean)
sample_mean = np.mean(sample, axis=0)
print('Sample Mean = {0}'.format(sample_mean))

#Sample Mean = [2.96143749 5.61268062]

# Part b (compute the sample covariance matrix)
sample_cov = np.cov(sample.T)
print('Sample Covariance')
print(sample_cov)
```

```
#Sample Covariance
#[[9.93191037 3.96365428]
# [3.96365428 5.30782634]]
# Part c (compute the eigenvalues and eigenvectors)
eigen_values, eigen_vectors = np.linalg.eig(sample_cov)
print('Eigenvalues = {0}'.format(eigen_values))
print('Eigenvectors (columns)')
print(eigen_vectors)
#Eigenvalues = [12.20856027 3.03117644]
#Eigenvectors (columns)
# [[ 0.86713795 -0.49806804]
# [ 0.49806804  0.86713795]]
# Part d (plot data and eigenvectors scaled by eigenvalues)
plt.figure(figsize=(8, 8))
plt.scatter(sample[:, 0], sample[:, 1])
plt.xlim(-15, 15)
plt.ylim(-15, 15)
plt.xlabel(r"$X_1$")
plt.ylabel(r"$X_2$")
plt.title("Sample Points and Eigenvectors")
vec_X = [sample_mean[0], sample_mean[0]]
vec_Y = [sample_mean[1], sample_mean[1]]
vec_U = [eigen_vectors[0][0] * eigen_values[0], eigen_vectors[0][1] * eigen_values[1]]
vec_V = [eigen_vectors[1][0] * eigen_values[0], eigen_vectors[1][1] * eigen_values[1]]
plt.quiver(vec_X, vec_Y, vec_U, vec_V, angles="xy", scale_units="xy", scale=1)
plt.show()
# Part e (plot rotated data in coorinate system defined by eigenvectors)
rotated = np.dot(eigen_vectors.T, (sample - sample_mean).T).T
plt.figure(figsize=(8, 8))
plt.scatter(rotated[:, 0], rotated[:, 1])
plt.xlim(-15, 15)
plt.ylim(-15, 15)
plt.xlabel(r"$x_1$")
plt.ylabel(r"$x_2$")
plt.title("Rotated Sample Points")
plt.show()
```





### 5 Classification and Risk

Suppose we have a classification problem with classes labeled 1, ..., c and an additional "doubt" category labeled c + 1. Let  $r : \mathbb{R}^d \to \{1, ..., c + 1\}$  be a decision rule. Define the loss function

$$L(r(x) = i, y = j) = \begin{cases} 0 & \text{if } i = j \quad i, j \in \{1, \dots, c\}, \\ \lambda_r & \text{if } i = c + 1, \\ \lambda_s & \text{otherwise,} \end{cases}$$

where  $\lambda_r \ge 0$  is the loss incurred for choosing doubt,  $\lambda_s \ge 0$  is the loss incurred for making a misclassification, and at least one of them is nonzero. Hence the risk of classifying a new data point x as class  $i \in \{1, 2, ..., c+1\}$  is

$$R(r(x) = i|x) = \sum_{j=1}^{c} L(r(x) = i, y = j) P(Y = j|x).$$

To be clear, the actual label Y can never be c + 1.

- 1. Show that the following predictor obtains the minimum risk when  $\lambda_r \leq \lambda_s$ :
  - Choose class i if  $P(Y = i|x) \ge P(Y = j|x)$  for all j and  $P(Y = i|x) \ge 1 \lambda_r/\lambda_s$ ;
  - Choose doubt otherwise.
- 2. What happens if  $\lambda_r = 0$ ? What happens if  $\lambda_r > \lambda_s$ ? Explain why this is consistent with what one would expect intuitively.

#### **Solution:**

- 1. Let  $r : \mathbb{R}^d \to \{1, \dots, c+1\}$  be the decision rule which implements the described policy. We will prove that in expectation the rule r is at least as good as the arbitrary rule f. Let  $x \in \mathbb{R}^d$  be a data point, which we want to classify.
  - Assume that case (1) holds and so r(x) = i,  $P(Y = i|x) \ge P(Y = j|x)$  for all j, and  $P(Y = i|x) \ge 1 \lambda_r/\lambda_s$ . Then:

$$R(r(x) = i|x) = \sum_{j=1}^{c} \ell(r(x) = i, y = j)P(Y = j|x)$$
$$= \lambda_s \sum_{j=1, j \neq i} P(Y = j|x)$$
$$= \lambda_s \left(1 - P(Y = i|x)\right).$$

We are going to consider two cases for f(x):

- Let f(x) = k, with  $k \neq i$ , c + 1. Then:

$$R(f(x) = k|x) = \lambda_s (1 - P(Y = k|x))$$

Hence 
$$R(r(x) = i|x) \le R(f(x) = k|x)$$
, since  $P(Y = i|x) \ge P(Y = k|x)$ .

- Let f(x) = c + 1. Then:

$$R(f(x) = c + 1|x) = \lambda_r$$

Hence  $R(r(x) = i|x) \le R(f(x) = c + 1|x)$ , since  $P(Y = i|x) \ge 1 - \lambda_r/\lambda_s$ .

• Assume that case (2) holds and so r(x) = c + 1. Then:

$$R(r(x) = i|x) = \lambda_r$$

Let f(x) = k, with  $k \neq c + 1$ . Then:

$$R(f(x) = k|x) = \lambda_s (1 - P(Y = k|x))$$

case (1) does **not** hold which means that:

$$\max_{j \in \{1,\dots,c\}} P(Y = j|x) < 1 - \lambda_r/\lambda_s$$

hence  $P(Y = k|x) < 1 - \lambda_r/\lambda_s$ , and so R(r(x) = i|x) < R(f(x) = k|x).

Therefore in every case we proved that the rule r is at list as good as the arbitrary rule f, which proves that r is an optimal rule.

2. If  $\lambda_r = 0$ , then case (1) will hold iff there exists an  $i \in \{1, ..., c\}$  such that P(r(x) = i|x) = 1. So we will either classify x in class i if we are 100% sure about this, or else we will choose doubt. Of course this is completely consistent with our intuition, because choosing doubt does not have any penalty at all, since  $\lambda_r = 0$ .

If  $\lambda_r > \lambda_s$ , then we will always classify x in the class  $i \in \{1, ..., c\}$  which gives the highest probability of correct classification. Once again this makes sense, since the cost of choosing doubt is higher than classifying x in any of the classes, hence our best option is to classify x in the class which gives the highest probability for a correct classification.

### 6 Maximum Likelihood Estimation and Bias

Let  $X_1, \ldots, X_n \in \mathbb{R}$  be n sample points drawn independently from univariate normal distributions such that  $X_i \sim \mathcal{N}(\mu, \sigma_i^2)$ , where  $\sigma_i = \sigma / \sqrt{i}$  for some parameter  $\sigma$ . (Every sample point comes from a distribution with a **different variance**.) Note the word "univariate"; we are working in dimension d = 1, and our "points" are real numbers.

1. Derive the maximum likelihood estimates, denoted  $\hat{\mu}$  and  $\hat{\sigma}$ , for the mean  $\mu$  and the parameter  $\sigma$ . (The formulae from class don't apply here, because every point has a different variance.) You may write an expression for  $\hat{\sigma}^2$  rather than  $\hat{\sigma}$  if you wish—it's probably simpler that way. Show all your work.

$$\mathcal{L}(\mu, \sigma; X) = \prod_{i=1}^{n} \frac{\sqrt{i}}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(X_i - \mu)^2 i}{2\sigma^2}\right).$$

$$\ell(\mu, \sigma; X) = \frac{1}{2} \sum_{i=1}^{n} \ln \frac{i}{2\pi} - n \ln \sigma - \sum_{i=1}^{n} \frac{(X_i - \mu)^2 i}{2\sigma^2}.$$

$$\frac{\partial \ell}{\partial \mu} = \sum_{i=1}^{n} \frac{(X_i - \mu)i}{\sigma^2} = 0 \implies$$

$$\hat{\mu} = \frac{\sum_{i=1}^{n} X_i i}{\sum_{i=1}^{n} i}.$$

$$\frac{\partial \ell}{\partial \sigma} = -\frac{n}{\sigma} + \sum_{i=1}^{n} \frac{(X_i - \mu)^2 i}{\sigma^3} = 0 \implies$$

$$\hat{\sigma}^2 = \frac{1}{n} \sum_{i=1}^{n} (X_i - \mu)^2 i.$$

But we don't know the value of  $\mu$ , so our MLE estimate for  $\sigma$  is  $\hat{\sigma}^2 = \frac{1}{n} \sum_{i=1}^n (X_i - \hat{\mu})^2 i$ .

2. Given the true value of a statistic  $\theta$  and an estimator  $\hat{\theta}$  of that statistic, we define the *bias* of the estimator to be the expected difference from the true value. That is,

$$bias(\hat{\theta}) = \mathbb{E}[\hat{\theta}] - \theta.$$

We say that an estimator is *unbiased* if its bias is 0.

Either prove or disprove the following statement: The MLE sample estimator  $\hat{\mu}$  is unbiased. Hint: Neither the true  $\mu$  nor true  $\sigma^2$  are known when estimating sample statistics, thus we need to plug in appropriate estimators.

**Solution:** The MLE sample estimator  $\mu$  is unbiased.

$$bias(\hat{\mu}) = \mathbb{E}[\hat{\mu}] - \mu$$

$$= \mathbb{E}\left[\frac{\sum_{i=1}^{n} X_{i}i}{\sum_{i=1}^{n} i}\right] - \mu$$

$$= \frac{\sum_{i=1}^{n} \mathbb{E}[X_{i}]i}{\sum_{i=1}^{n} i} - \mu$$

$$= \frac{\sum_{i=1}^{n} \mu i}{\sum_{i=1}^{n} i} - \mu$$

$$= \mu \frac{\sum_{i=1}^{n} i}{\sum_{i=1}^{n} i} - \mu$$

$$= \mu - \mu$$

$$= 0.$$

3. Either prove or disprove the following statement: The MLE sample estimator  $\hat{\sigma}^2$  is unbiased. Hint: Neither the true  $\mu$  nor true  $\sigma^2$  are known when estimating sample statistics, thus we need to plug in appropriate estimators.

**Solution:** The MLE sample estimator for  $\sigma^2$  is biased. We begin with some helpful identities. Let X be any random variable. Then we have

$$Var[X] = \mathbb{E}[X^2] + \mathbb{E}[X]^2$$

Now let  $z = \sum_{i=1}^{n} i = \frac{n(n+1)}{2}$ . Because  $\hat{\mu} = \frac{1}{z} \sum X_i i$ , we have  $Var[\hat{\mu}] = \frac{\sigma^2}{z}$ . Thus, we get

$$\mathbb{E}[\hat{\mu}^2] = \frac{\sigma^2}{z} + \mu^2$$

We now plug into the definition of bias and simplify

$$bias(\hat{\sigma}^{2}) = \mathbb{E}[\hat{\sigma}^{2}] - \sigma^{2}$$

$$= \mathbb{E}[\frac{1}{n} \sum_{i=1}^{n} (X_{i} - \hat{\mu})^{2} i] - \sigma^{2}$$

$$= \frac{1}{n} \mathbb{E}[\sum_{i=1}^{n} X_{i}^{2} i + \sum_{i=1}^{n} \hat{\mu}^{2} i - 2\hat{\mu} \sum_{i=1}^{n} X_{i} i] - \sigma^{2}$$

$$= \frac{1}{n} \mathbb{E}[\sum_{i=1}^{n} X_{i}^{2} i + z\hat{\mu}^{2} - 2z\hat{\mu}^{2}] - \sigma^{2}$$

$$= \frac{1}{n} \left[\sum_{i=1}^{n} \mathbb{E}[X_{i}^{2}] i - z\mathbb{E}[\hat{\mu}^{2}]\right] - \sigma^{2}$$

$$= \frac{1}{n} \left[\sum_{i=1}^{n} (\frac{\sigma^{2}}{i} + \mu^{2}) i - z(\frac{\sigma^{2}}{z} + \mu^{2})\right] - \sigma^{2}$$

$$= \frac{1}{n} \left[n\sigma^{2} + z\mu^{2} - \sigma^{2} - z\mu^{2}\right] - \sigma^{2}$$

$$= \frac{n-1}{n}\sigma^{2} - \sigma^{2}$$

$$= -\frac{1}{n}\sigma^{2}$$

We therefore see that the MLE sample estimator for variance systematically *underestimates* the true variance. It turns out that we can correct this error by sightly increasing our estimate by dividing by n-1 instead of n. Thus, the new sample estimator

$$\frac{1}{n-1} \sum_{i=1}^{n} (X_i - \hat{\mu})^2 i$$

is unbiased. This can be shown by repeating the above algebra with the new denominator and observing that the bias does in fact equal zero.

### 7 Covariance Matrices and Decompositions

As described in lecture, the covariance matrix  $Var(R) \in \mathbb{R}^{d \times d}$  for a random variable  $R \in \mathbb{R}^d$  with mean  $\mu \in \mathbb{R}^d$  is

$$\operatorname{Var}(R) = \operatorname{Cov}(R, R) = \mathbb{E}[(R - \mu) (R - \mu)^{\top}] = \begin{bmatrix} \operatorname{Var}(R_1) & \operatorname{Cov}(R_1, R_2) & \dots & \operatorname{Cov}(R_1, R_d) \\ \operatorname{Cov}(R_2, R_1) & \operatorname{Var}(R_2) & & \operatorname{Cov}(R_2, R_d) \\ \vdots & & \ddots & \vdots \\ \operatorname{Cov}(R_d, R_1) & \operatorname{Cov}(R_d, R_2) & \dots & \operatorname{Var}(R_d) \end{bmatrix},$$

where  $Cov(R_i, R_j) = \mathbb{E}[(R_i - \mu_i)(R_j - \mu_j)]$  and  $Var(R_i) = Cov(R_i, R_i)$ .

If the random variable R is sampled from the multivariate normal distribution  $\mathcal{N}(\mu, \Sigma)$  with the PDF

$$f(x) = \frac{1}{\sqrt{(2\pi)^d |\Sigma|}} e^{-((x-\mu)^T \Sigma^{-1}(x-\mu))/2},$$

then  $Var(R) = \Sigma$ .

Given n points  $X_1, X_2, \ldots, X_n$  sampled from  $\mathcal{N}(\mu, \Sigma)$ , we can estimate  $\Sigma$  with the maximum likelihood estimator

$$\hat{\Sigma} = \frac{1}{n} \sum_{i=1}^{n} (X_i - \mu) (X_i - \mu)^{\mathsf{T}},$$

which is also known as the sample covariance matrix.

- 1. The estimate  $\hat{\Sigma}$  makes sense as an approximation of  $\Sigma$  only if  $\hat{\Sigma}$  is invertible. Under what circumstances is  $\hat{\Sigma}$  not invertible? Make sure your answer is complete; i.e., it includes all cases in which the covariance matrix of the sample is singular. Express your answer in terms of the geometric arrangement of the sample points  $X_i$ .
- 2. Suggest a way to fix a singular covariance matrix estimator  $\hat{\Sigma}$  by replacing it with a similar but invertible matrix. Your suggestion may be a kludge, but it should not change the covariance matrix too much. Note that infinitesimal numbers do not exist; if your solution uses a very small number, explain how to calculate a number that is sufficiently small for your purposes.
- 3. Consider the normal distribution  $\mathcal{N}(0, \Sigma)$  with mean  $\mu = 0$ . Consider all vectors of length 1; i.e., any vector x for which ||x|| = 1. Which vector(s) x of length 1 maximizes the PDF f(x)? Which vector(s) x of length 1 minimizes f(x)? Your answers should depend on the properties of  $\Sigma$ . Explain your answer.
- 4. Suppose we have  $X \sim \mathcal{N}(0, \Sigma)$ ,  $X \in \mathbb{R}^n$  and a unit vector  $y \in \mathbb{R}^n$ . We can compute the projection of the random vector X onto a direction y as  $p = y^T X$ . First, compute the variance of p. Second, with this information, what does the eigenvector corresponding to the largest eigenvalue of the covariance matrix tell us about the variances of expressions of the form  $y^T X$ ?

- 1.  $\tilde{\Sigma}$  is not invertible if and only if all the points lie on a common hyperplane in feature space(i.e. they don't span all of  $\mathbb{R}^{dxd}$ )
- 2. Note: This question is worded to allow lots of room for creativity, including answers we might not have thought of. So when grading, don't demand a perfectly precise answer; we're more interested in an intuitive understanding of what the modification means.

We can create a new matrix  $\tilde{\Sigma} = \hat{\Sigma} + \alpha I$ . We want to make all the eigenvalues of the new covariance matrix positive so it will be invertible. Notice that the original  $\hat{\Sigma}$  had all nonnegative eigenvalues. We ideally want  $\alpha$  to be a very small number so not to bias our covariance matrix. Since the eigenvalues of  $\tilde{\Sigma}^{-1}$  are the reciprocals of the originals. Then we want  $\alpha$  to be large enough so that the eigenvalues of  $\tilde{\Sigma}^{-1}$  don't blow up. The best way of selecting  $\alpha$  is hyperparameter tuning.

Another acceptable answer is projecting the covariance matrix onto a lower dimensional subspace by removing the features that are linear combinations of the remaining features. We continue this process until all the features are independent. This process will leave a full rank matrix that will be invertible.

- 3. x will maximize the PDF f(x) if it is the eigenvector of  $\Sigma$  with the largest eigenvalue. Similarly, x will minimize the PDF f(x) if it is the eigenvector of  $\Sigma$  with the smallest eigenvalue. The distance metric is  $\Sigma^{-1}$ , which means it penalizes deviation from the mean least along the direction of its eigenvector with the smallest eigenvalue(which is paired with the largest eigenvalue of  $\Sigma$ ) and vice versa.
- 4. Noting that  $p = y^T X$  is a scalar, let us first compute  $\mathbb{E}[y^T X]$ . Since y is a constant, by linearity of expectation  $y^T \mathbb{E}[X] = 0$ . Next, by the definition of the variance  $\mathbb{E}[(p \mathbb{E}[p])^2] = \mathbb{E}[(p \mathbb{E}[p])^T] = \mathbb{E}[y^T (X \mathbb{E}[X])(X \mathbb{E}[X])^T y] = y^T \mathbb{E}[(X \mathbb{E}[X])(X \mathbb{E}[X])^T]y = y^T \Sigma y$ . Remembering that the quadratic form  $y^T \Sigma y$  is maximized by the eigenvector of  $\Sigma$  with the largest eigenvalue, we see that the largest eigenvector provides the direction of maximum variance.

## 8 Gaussian Classifiers for Digits and Spam

In this problem, you will build classifiers based on Gaussian discriminant analysis. Unlike Homework 1, you are NOT allowed to use any libraries for out-of-the-box classification (e.g. sklearn). You may use anything in numpy and scipy.

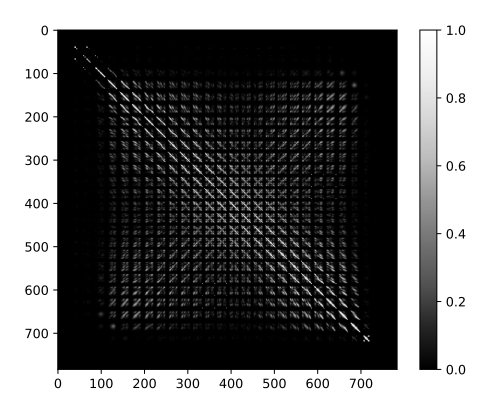
The training and test data can be found with this homework. **Do NOT use the training/test data from Homework 1, as they have changed for this homework.** The starter code is similar to HW1's; we provide check.py and save\_csv.py files for you to produce your Kaggle submission files. Submit your predicted class labels for the test data on the Kaggle competition website and be sure to include your Kaggle display name and scores in your writeup. Also be sure to include an appendix of your code at the end of your writeup.

1. Taking pixel values as features (no new features yet, please), fit a Gaussian distribution to

each digit class using maximum likelihood estimation. This involves computing a mean and a covariance matrix for each digit class, as discussed in Lecture 9.

*Hint:* You may, and probably should, contrast-normalize the images before using their pixel values. One way to normalize is to divide the pixel values of an image by the  $l_2$ -norm of its pixel values.

2. (Written answer + graph) Visualize the covariance matrix for a particular class (digit). How do the diagonal terms compare with the off-diagonal terms? What do you conclude from this?



#### **Solution:**

This is technically a visualization of the Pearson product-moment correlation coefficients, a normalization of the covariance matrix such that  $R_{ij} = \frac{C_{ij}}{\sqrt{C_{ii}C_{jj}}}$ , where C is the covariance matrix. Every element of R is in [-1,1], and so it is easier to understand the correlation between different pixels when visualizing this matrix. Furthermore we've visualized the absolute value of the entries in R.

Things to notice include that the diagonal elements are generally larger than the off-diagonal elements, and that the entries further away from the diagonal are generally lower than entries closer to the diagonal. We interpret this to mean that the correlation between neighboring pixels is generally higher than pixels that are further apart.

3. Classify the digits in the test set on the basis of posterior probabilities with two different approaches.

(a) (Graphs) Linear discriminant analysis (LDA). Model the class conditional probabilities as Gaussians  $\mathcal{N}(\mu_C, \Sigma)$  with different means  $\mu_C$  (for class C) and the same pooled within-class covariance matrix  $\Sigma$ , which you compute from a weighted average of the 10 covariance matrices from the 10 classes, as described in Lecture 9.

In your implementation, you might run into issues of determinants overflowing or underflowing, or normal PDF probabilities underflowing. These problems might be solved by learning about numpy.linalg.slogdet and/or scipy.stats.multivariate\_normal.logpdf.

To implement LDA, you will sometimes need to compute a matrix-vector product of the form  $\Sigma^{-1}x$  for some vector x. You should **not** compute the inverse of  $\Sigma$  (nor the determinant of  $\Sigma$ ) as it is not guaranteed to be invertible. Instead, you should find a way to solve the implied linear system without computing the inverse.

Hold out 10,000 randomly chosen training points for a validation set. (You may reuse your Homework 1 solution or an out-of-the-box library for dataset splitting *only*.) Classify each image in the validation set into one of the 10 classes. Compute the error rate  $(1 - \frac{\# points \ correctly \ classified}{\# total \ points})$  on the validation set and plot it over the following numbers of randomly chosen training points: 100, 200, 500, 1,000, 2,000, 5,000, 10,000, 30,000, 50,000. (Expect some variation in your error rate when few training points are used.)

- (b) (Graphs) Quadratic discriminant analysis (QDA). Model the class conditional probabilities as Gaussians  $\mathcal{N}(\mu_C, \Sigma_C)$ , where  $\Sigma_C$  is the estimated covariance matrix for class C. (If any of these covariance matrices turn out singular, implement the trick you described in Q7(b). You are welcome to use validation to choose the right constant(s) for that trick.) Repeat the same tests and error rate calculations you did for LDA.
- (c) (Written answer) Which of LDA and QDA performed better? Why?
  Solution: In our experiments on MNIST, both LDA and QDA had similar error rates on the validation data for training sets of size greater than 10000. However we generally

expect QDA to perform better because it is more expressive. The drawback of QDA is that the larger number of free variables leaves it vulernable to possible overfitting.

- (d) (Written answer + graph) Include a plot of validation error versus the number of training points for each digit. Plot all the 10 curves on the same graph as shown in Figure 1. Which digit is easiest to classify? Write down your answer and suggest why you think it's the easiest digit.
  - **Solution:** In our experiments on MNIST, digit 1 was easiest for LDA and digit 0 was easiest for QDA, as measured by validation error. The plots for validation error and per-digit validation error for both LDA and QDA are available below.
- 4. (Written answer) Using the mnist-data-hw3.npz, train your best classifier for the training\_data and classify the images in the test\_data. Submit your labels to the online Kaggle competition. Record your optimum prediction rate in your submission and include your Kaggle username.
  - You are welcome to compute extra features for the Kaggle competition, as long as they do not use an exterior learned model for their computation (no transfer learning!). If you do so,

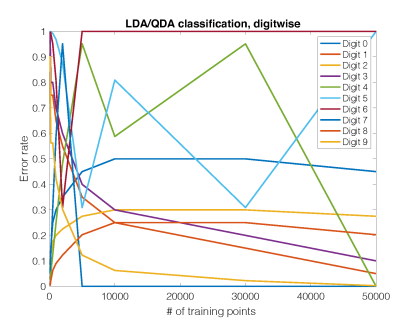


Figure 1: Sample graph with 10 plots

please describe your implementation in your assignment. Please use extra features **only** for the Kaggle portion of the assignment.

5. (Written answer) Next, apply LDA or QDA (your choice) to spam (spam-data-hw3.npz). Submit your test results to the online Kaggle competition. Record your optimum prediction rate in your submission. If you use additional features (or omit features), please describe them. We include a featurize.py file (similar to HW1's) that you may modify to create new features.

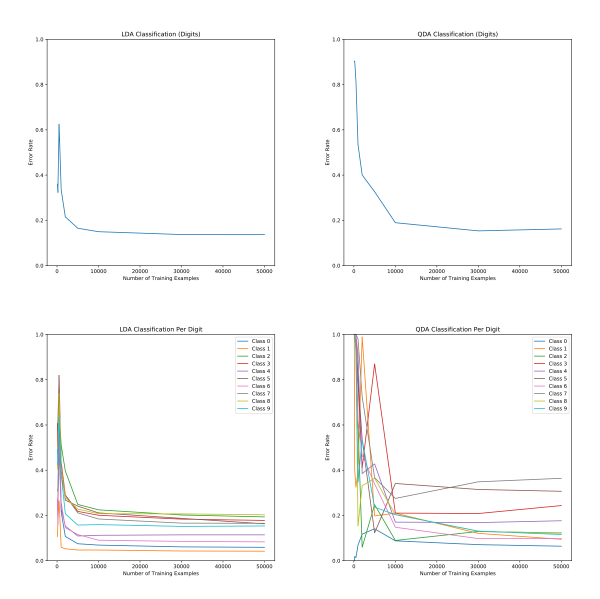
*Optional:* If you use the defaults, expect relatively low classification rates. We suggest using a Bag-Of-Words model. You are encouraged to explore alternative hand-crafted features, and are welcome to use any third-party library to implement them, as long as they do not use a separate model for their computation (no large language models or word2vec!).

```
import numpy as np
import scipy.cluster
import scipy.io
import scipy.ndimage
import matplotlib
import matplotlib.pyplot as plt
import learners
def train_val_split(data, labels, val_size):
   num_items = len(data)
   assert num_items == len(labels)
   assert val_size >= 0
   if val_size < 1.0:</pre>
       val_size = int(num_items * val_size)
    train_size = num_items - val_size
   idx = np.random.permutation(num_items)
    data_train = data[idx][:train_size]
   label_train = labels[idx][:train_size]
```

```
data_val = data[idx][train_size:]
    label_val = labels[idx][train_size:]
    return data_train, data_val, label_train, label_val
mnist = scipy.io.loadmat("./mnist-data/mnist_data.mat")
train_data, train_labels = mnist['training_data'], mnist['training_labels']
train_data_normalized = scipy.cluster.vq.whiten(train_data)
labels = np.unique(train_labels)
# Part a
mnist_fitted = {}
for label in labels:
    class_indices = (train_labels == label).flatten()
    data = train_data_normalized[class_indices]
    mean = np.mean(data, axis=0)
    cov = np.cov(data, rowvar=False)
    mnist_fitted[label] = (mean, cov)
# Part b
class_indices = (train_labels == labels[0]).flatten()
data = train_data_normalized[class_indices]
ncov = np.corrcoef(data, rowvar=False)
ncov[np.isnan(ncov)] = 0
ncov = np.abs(ncov)
plt.imshow(ncov, cmap=matplotlib.cm.Greys_r)
plt.colorbar()
plt.show()
# Part c (Gaussian/linear discriminant analysis)
train_data, val_data, train_labels, val_labels = train_val_split(train_data_normalized, train_labels,
     \hookrightarrow val_size=10000)
val_labels.flatten()
num_training = [100, 200, 500, 1000, 2000, 5000, 10000, 30000, 50000]
lda_errors = []
qda_errors = []
for num in num_training:
    gda = learners.GDA()
    gda.fit(train_data[:num], train_labels[:num])
    err = 1 - gda.evaluate(val_data, val_labels)
    lda_errors.append(err)
    err = 1 - gda.evaluate(val_data, val_labels, mode="qda")
    qda_errors.append(err)
# Part c.1
plt.figure(figsize=(8, 8))
plt.plot(num_training, lda_errors)
plt.xlabel("Number of Training Examples")
plt.ylabel("Error Rate")
plt.title("LDA Classification (Digits)")
plt.ylim((0, 1))
plt.show()
print(lda_errors)
#[0.34, 0.346, 0.62, 0.36, 0.21, 0.16, 0.14, 0.13, 0.13]
# Part c.2
plt.figure(figsize=(8, 8))
plt.plot(num_training, qda_errors)
plt.xlabel("Number of Training Examples")
plt.ylabel("Error Rate")
plt.title("QDA Classification (Digits)")
plt.ylim((0, 1))
plt.show()
print(qda_errors)
```

```
## learners.py
## Author: Sinho Chewi
import math
import numpy as np
import scipy.stats
class GDA:
    """Perform Gaussian discriminant analysis (both LDA and QDA)."""
    def evaluate(self, X, y, mode="lda"):
        """Predict and evaluate the accuracy using zero-one loss.
        Args:
            X (np.ndarray): The feature matrix.
            y (np.ndarray): The true labels.
            mode (str): Either "lda" or "qda".
        Returns:
            float: The zero-one loss of the learner.
        Raises:
           RuntimeError: If an unknown mode is passed into the method.
        pred = self.predict(X, mode=mode)
        return np.sum(pred == y) / y.shape[0]
    def fit(self, X, y):
        """Train the GDA model (both LDA and QDA).
        Args:
           X (np.ndarray): The feature matrix.
           y (np.ndarray): The true labels.
        self.classes = np.unique(y)
       num_examples = X.shape[0]
       num_features = X.shape[1]
        self.fitted_params = []
        self.pooled_cov = np.zeros((num_features, num_features))
        for c in self.classes:
            class_indices = (y == c).flatten()
            data = X[class_indices]
            mean = np.mean(data, axis=0)
            unscaled_cov = np.dot((data - mean).T, data - mean)
            self.pooled_cov = np.add(self.pooled_cov, unscaled_cov)
            scaled_cov = unscaled_cov / data.shape[0]
            prior = data.shape[0] / num_examples
            self.fitted_params.append((c, mean, scaled_cov, prior))
        self.pooled_cov = self.pooled_cov / num_examples
    def predict(self, X, mode="lda"):
        """Use the fitted model to make predictions.
        Args:
            X (np.ndarray): The feature matrix.
        Optional:
           mode (str): Either "lda" or "qda".
        Returns:
            np.ndarray: The array of predictions.
        Raises:
```

```
RuntimeError: If an unknown mode is passed into the method.
if mode == "lda":
    pred = np.array([
        (scipy.stats.multivariate_normal.logpdf(
            X, allow_singular=True, cov=self.pooled_cov, mean=mean)
            + math.log(prior))
        for (c, mean, cov, prior) in self.fitted_params
    ])
   return self.classes[np.argmax(pred, axis=0)].reshape((-1, 1))
elif mode == "qda":
    pred = np.array([
        (scipy.stats.multivariate_normal.logpdf(
            X, allow_singular=True, cov=cov, mean=mean)
            + math.log(prior))
        for (c, mean, cov, prior) in self.fitted_params
    ])
    return self.classes[np.argmax(pred, axis=0)].reshape((-1, 1))
else:
    raise RuntimeError("Unknown mode!")
```



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### Submission Checklist

Please ensure you have completed the following before your final submission.

### At the beginning of your writeup...

- 1. Have you copied and hand-signed the honor code specified in Question 1?
- 2. Have you listed all students (Names and ID numbers) that you collaborated with?

# In your writeup for Question 8...

1. Have you included your **Kaggle Score** and **Kaggle Username** for **both** questions 8.4 and 8.5?

### At the end of the writeup...

- 1. Have you provided a code appendix including all code you wrote in solving the homework?
- 2. Have you included featurize.py in your code appendix if you modified it?

### Executable Code Submission

- 1. Have you created an archive containing all ".py" files that you wrote or modified to generate your homework solutions (including featurize.py if you modified it)?
- 2. Have you removed all data and extraneous files from the archive?
- 3. Have you included a README file in your archive briefly describing how to run your code on the test data and reproduce your results?

# Submissions

- 1. Have you submitted your written solutions to the Gradescope assignment titled **HW3 Write-Up** and selected pages appropriately?
- 2. Have you submitted your executable code archive to the Gradescope assignment titled **HW3** Code?
- 3. Have you submitted your test set predictions for both **MNIST** and **SPAM** to the appropriate Kaggle challenges?

Congratulations! You have completed Homework 3.