Module import

```
In []: import numpy as np
   import math
   import matplotlib.pyplot as plt
   import seaborn as sns
   import pandas as pd
   import random
   import csv
```

In []: # You can import other modules if needed

1. Visualize a mixture of Gaussians [15 pts]

Consider a mixture of two 2D Gaussian distrbutions as follows

$$0.4\cdot\mathcal{N}\left(\left[egin{array}{c}10\2\end{array}
ight],\left[egin{array}{c}1,0\0,1\end{array}
ight]
ight)+0.6\cdot\mathcal{N}\left(\left[egin{array}{c}0\0\end{array}
ight],\left[egin{array}{c}8.4,2.0\2.0,1.7\end{array}
ight]
ight)$$

- a. Compute the marginal distributions for each dimension.
- b. Compute the mean for each marginal distribution.
- c. Compute the mean for the two-dimensional distribution.
- d. Generate both joint and marginal density of this distribution by taking at least 10 thousand samples. Please refer to Lab4 notebook for the visualization approach.

Type the answers to first three questions here:

(a) For the first dimension the marginal distribution of two gaussians are x_1 = N(10,1) and x_1 = N(0,8.4). the mixture of two 2D Gaussiansn has a probability ratio of 0.4 and 0.6. So

$$p(x_1) = 0.4 imes N(10,1) + 0.6 imes N(0,8.4)$$

Similarly, for $p(x_2)$

$$p(x_2) = 0.4 imes N(2,1) + 0.6 imes N(0,1.7)$$

(b) I specify the marginal distribution for each dimension as normal distributions. For the first dimension,

The mean for x_1 ,

$$\mu x_1 = 0.4 \times 10 + 0.6 \times 0 = 4$$

and for x_2 ,

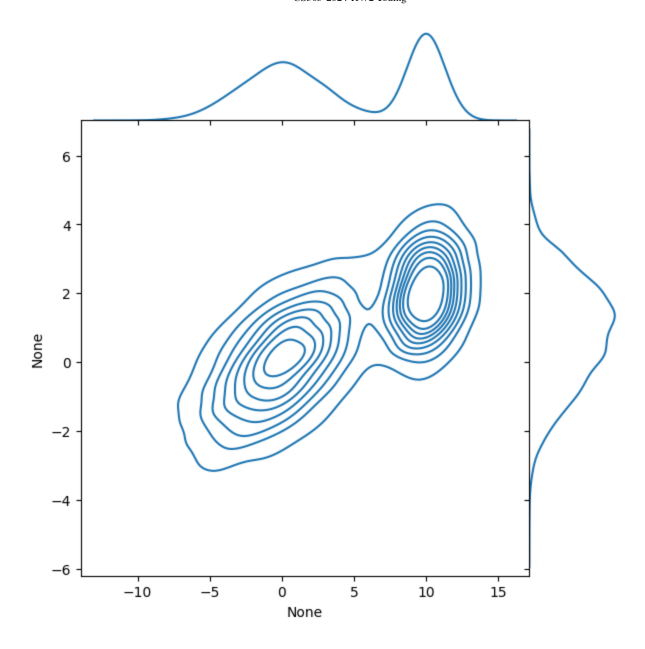
$$\mu x_2 = 0.4 \times 2 + 0.6 \times 0 = 0.8$$

(c) The mean for the two-dimensional distribution is for x_1 = 0.4*10+0.6*0=4, and for x_2 = 0.4*2+0.6*0=1.2. So the mean for the two-dimensional distribution is

$$\mu = 0.4 imes egin{bmatrix} 10 \ 2 \end{bmatrix} + 0.6 imes egin{bmatrix} 0 \ 0 \end{bmatrix} = egin{bmatrix} 4 \ 0.8 \end{bmatrix}$$

```
In [ ]: # Your code for question d here
                                                                  #2D gaussian has 2 c
        def generate_two_d_gaussian(m, cov, d, samples):
            epsilon = 0.0001
            K = cov + epsilon*np.identity(d)
            L = np.linalg.cholesky(K)
            n = samples
            u = np.random.normal(loc=0, scale=1, size=d*n).reshape(d, n)
            x = m + np.dot(L, u)
            return x
        first gaussian mean = np.array([10, 2]).reshape(2, 1)
        second_gaussian_mean = np.array([0, 0]).reshape(2, 1)
        first covariance matrix = np.array([[1, 0], [0, 1]])
        second\_covariance\_matrix = np.array([[8.4, 2.0], [2.0, 1.7]])
        x \text{ samples} = 4000
                                                              #4000 samples from first
        y_samples = 6000
                                                              #6000 samples from secon
        x = generate_two_d_gaussian(first_gaussian_mean, first_covariance_matrix, d,
        y = generate_two_d_gaussian(second_gaussian_mean, second_covariance_matrix,
        z = np.hstack((x, y))
        sns.jointplot(x=z[0], y=z[1], kind="kde", space=0)
```

Out[]: <seaborn.axisgrid.JointGrid at 0x10fae8b90>



2. Simulation: Drop needles [15 pts]

Suppose we have a floor made of parallel strips of wood, each the same width t, and we drop a needle with length l=t onto the floor. What is the probability that the needle will lie across a line between two strips?

Below is an example of two needles dropped. Needle a falls across a line, while needle b does not.



In this coding homework, we will simulate such experiments and connect them with the estimation of π .

Functions

The first thing to write is a function *drop_needle*. It simulates dropping a needle onto the floor we described and returns whether the needle lies across a line between two strips.

Now the question is how to describe the position of a needle using random variables. The figure below visualizes a needle sampled, with t=l=1 (see figure above). Remember that the needle should have an equal probability of landing in any position. In fact, we can uniformly sample the position of the needle's mass center and then uniformly sample the angle formed by the needle and the x-axis. Specifically, we only focus on the mass center's position with respect to (w.r.t.) the x-axis since we can assume the strip is long enough.

Besides, we do not need to sample the x-value of the center from $-\inf$ to \inf . Instead, we can uniformly sample it from 0 to 2t. Why is this the case?

needleExmple2

```
In [ ]: def drop_needle(strip_length, needle_length):
            Simulate dropping a needle on to the floor made of parallel strips of wo
            Return whether the needle lie across a line between two strips.
            :return: An Integer that equals to 1 if the needle lie across a line, ar
            # write your code here
            """ middle point of the needle is in the range of 0 - 2 \times needle_length
            middle_point = random.uniform(0, 2 * needle_length)
            """ theta is in the range of 0 - 90 degrees """
            theta = random.uniform(0, math.pi/2)
            half length = math.cos(theta) * needle length / 2
            """ x is in the range of 0 - strip_length """
            """Determining the distance between the middle point of the needle and t
            if middle point < 1 * strip length:</pre>
                 if middle_point - half_length < 0 or middle_point + half_length > 1
                     return 1
                else:
                     return 0
            elif middle_point < 2 * strip_length:</pre>
                 if middle_point - half_length < 1 * strip_length or middle_point + h</pre>
                     return 1
                 else:
                     return 0
```

Next, write a function run_simulation that calls drop_needle repetitively for n times. The function should return the probability that a dropped needle lies across a line based on the n trials.

```
In []: def run_simulation(n, strip_length, needle_length):
    """
    Repeat drop_needle experiment for n times. Return the probability that t
    :return: float, the probability that the needle will lie across a line a
    """
    # Write your code here
    i, count = 0, 0
    while i < n:
        i += 1
        if drop_needle(strip_length, needle_length) == 1:
            count += 1
        probability = count / n
    return probability</pre>
```

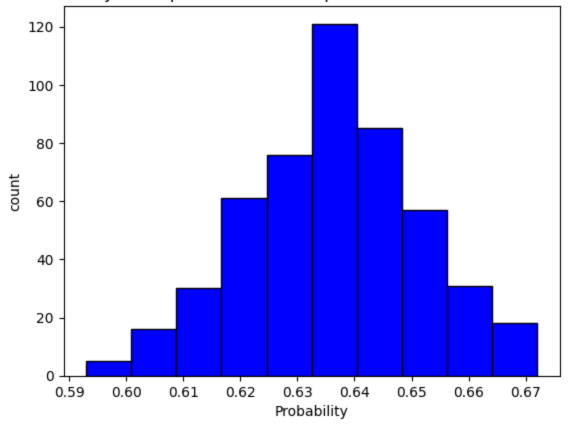
Run the simulation

Run the *run_simulation* function 500 times with parameters n=1000, strip_length=1, and needle_length=1. Each time the function is going to return a probability of the needle lying across the line. Plot a histogram of those 500 probabilities.

```
In [ ]: # Write your code here
        """ Run the simulation 1000 times with strip_length = 1 and needle_length =
        """ parameter: experiment, strip_length, needle_length """
        experiment = 500
        n = 1000
        strip length = 1
        needle_length = 1
        probabilities = []
        """ Run the simulator 500 times, calculate the probability that the needle ec{\mathsf{w}}
        for i in range(experiment):
            probability = run_simulation(n, strip_length, needle_length)
            probabilities.append(probability)
In [ ]: """ Plot the result"""
        plt.hist(probabilities, color = 'blue', edgecolor = 'black')
        plt title('Probability of Drop needle in 500 experiments with 1000 simulation
        plt.xlabel('Probability')
        plt.ylabel('count')
        plt.show
```

Out[]: <function matplotlib.pyplot.show(close=None, block=None)>

Probability of Drop needle in 500 experiments with 1000 simulations



3. Email Spam Naive Bayes [35 pts]

Naive Bayes

For a classification task, we aim to predict a binary label $Y \in \{0,1\}$ of an example given its d dimensional features X_1, X_2, \ldots, X_d . Bayes Theorem states the (posterior) porbability of have a label Y given the feature observations as follows:

$$P(Y|X_1, X_2, \dots, X_d) = \frac{P(Y) * P(X_1, X_2, \dots, X_d|Y)}{P(X_1, X_2, \dots, X_d)}$$

Note P(Y) is called priori, i.e., probability of seeing label Y without observing the features. To determine the binary label Y, we need to compare $P(Y=0|X_1,X_2,\ldots,X_d)$ with $P(Y=1|X_1,X_2,\ldots,X_d)$. In Bayes Theorem, the fractions representing both conditional probabilities have the same denominator $P(X_1,X_2,\ldots,X_d)$. Thus we can ignore the denominator and only focusing on

$$P(Y|X_1,X_2,\ldots,X_d) \propto P(Y) * P(X_1,X_2,\ldots,X_d|Y)$$

Next, we make a strong assumption: no pair of features in the dataset are dependent (this is the reason why we call it *Naive* Bayes). Under such assumption,

 $P(X_1,X_2,\dots,X_d|Y)=P(X_1|Y)*P(X_2|Y)*\dots*P(X_d|Y).$ Therefore, we conclude

$$P(Y|X_1, X_2, ..., X_n) \propto P(Y) * P(X_1|Y) * P(X_2|Y) * ... * P(X_d|Y)$$

The classifier will decide what class each input belongs to based on highest probability from the equation above.

How do we know the priori P(Y) and the conditional probabilities $P(X_i|Y)$? In supervised classification tasks we have labeled data available (we call it training set). We can thus estimate these probabilities based on the training data.

Overview/Task

The goal of this programming assignment is to build a naive bayes classifier from scratch that can determine whether email text should be labled spam or not spam based on its contents

Reminders

Please remember that the classifier must be written from scratch; do NOT use any libraries that implement the classifier for you, such as but not limited to sklearn.

You CAN, however, use SKlearn to split up the dataset between testing and training.

Feel free to look up any tasks you are not familiar with, e.g. the function call to read a csv

Task list/Recommended Order

In order to provide some guidance, I am giving the following order/checklist to solve this task:

- 1. Compute the "prior": P(Y) for Y = 0 (not spam) and Y = 1 (spam)
- 2. Compute the "likelihood": $P(X_n|Y)$. In this task, we recommend you to start with setting up a dictionary of words appeared in the training set $W=\{w_1,w_2\ldots\}$, and constructing feature set $\{X_1,X_2\ldots\}$ where X_i is a binary indicator of whether the word w_i exists in the text or not. Feel free to add preprocessing and modify the features if that can increase your accuracy!
- 3. Write code that uses the two items above to make a decision on whether or not an email is spam or ham (aka not spam)
- 4. Write code to evaluate your model. Test model on training data to debug
- 5. Test model on testing data to debug

Function template

```
In [ ]: def prior(df):
            ham prior = 0
            spam prior = 0
            '''YOUR CODE HERE'''
            num rows = len(df.label)
                                                                          #number of r
            for row in range(num rows):
                                                                          #iterate thr
                if df.label[row] == "spam":
                                                                          #if the labe
                     spam prior += 1
                                                                          #if the labe
                else:
                    ham_prior += 1
            ''' Compute the probability of each class'''
            ham prior = ham prior/num rows
                                                                          #divide the
                                                                          #divide the
            spam_prior = spam_prior/num_rows
            '''FND'''
            return ham_prior, spam_prior
        def likelihood(df):
            ham like dict = {}
            spam_like_dict = {}
             '''YOUR CODE HERE'''
             ''' Preprocess the text column of the dataframe to get the likelihood of
            # spam total = 0
            # ham total = 0
                                                                          #number of r
            num rows = len(df.text)
            punctuation = '''!()-[]{};:'",<>./?@#$%^&*_~'''
                                                                      #define the pu
            # word_appear = []
            ''' Add each words to the dictionary and count the number of times it ap
            for row in range(num rows):
                                                                          #iterate thr
                for word in df.text[row].split():
                                                                          #iterate thr
                    word = word.replace('Subject:', '')
                                                                         #remove the
                    word = word.replace('\n', ' ')
                                                                         #remove new
                    word = word.lower()
                                                                          #convert the
                    if word not in punctuation:
                         if df.label[row] == "spam":
                             if word in spam_like_dict:
                                 spam like dict[word] += 1
                             else:
                                 spam_like_dict[word] = 1
                         else:
                             if word in ham like dict:
                                 ham like dict[word] += 1
                             else:
                                 ham_like_dict[word] = 1
            '''FND'''
            return ham_like_dict, spam_like_dict
        def predict(ham_prior, spam_prior, ham_like_dict, spam_like_dict, text):
            prediction function that uses prior and likelihood structure to compute
            #ham_spam_decision = 1 if classified as spam, 0 if classified as normal/
            ham spam decision = None
```

```
'''YOUR CODE HERE'''
   punctuation = '''!()-[]{};:'",<>./?@#$%^&*_~'''
                                                         #define the punct
   text = text.replace('Subject:', '')
                                                           #remove the word
   text = text.replace('\n', ' ')
                                                            #remove new line
    text = text.lower()
                                                            #convert the wor
   #ham_posterior = posterior probability that the email is normal/ham
   ham posterior = ham prior
   ham count = 0
   """ Count the total numbers of word appearances in the ham like dict"""
   for ham word in ham like dict:
        ham count += ham like dict[ham word]
   for word in text.split():
                                                                #iterate thr
       word = word.replace('Subject:', '')
                                                                #remove the
       word = word.replace('\n', ' ')
                                                                #remove new
       word = word.lower()
                                                                #convert the
        if word not in punctuation:
            if word in ham_like_dict:
                ham posterior *= ham like dict[word] / ham count
            else:
                ham_posterior *= 0.1 / ham_count
   #spam posterior = posterior probability that the email is spam
   spam_posterior = spam_prior
   spam count = 0
   """ Count the total numbers of word appearances in the ham_like_dict"""
    for spam word in spam like dict:
        spam count += spam like dict[spam word]
   for word in text.split():
                                                                #iterate thr
        if word not in punctuation:
            if word in spam_like_dict:
                spam posterior *= spam like dict[word] / spam count
            else:
                spam_posterior *= 0.1 / spam_count
    ''' Multiply the posterior probability by the prior probability to get t
   if spam_posterior > ham_posterior :
       ham_spam_decision = 1
   else:
       ham_spam_decision = 0
    '''FND'''
    return ham_spam_decision
def metrics(ham_prior, spam_prior, ham_dict, spam_dict, df):
   Calls "predict" function and report accuracy, precision, and recall of
    '''YOUR CODE HERE'''
```

```
''' define the variables for the metrics'''
true positive = 0
false positive = 0
true negative = 0
false_negative = 0
''' iterate through the rows of the dataframe and calculate the metrics
for row in range(len(df.text)):
    prediction = predict(ham prior, spam prior, ham dict, spam dict, df.
    if prediction == 1 and df.label[row] == "spam":
        true positive += 1
    elif prediction == 1 and df.label[row] == "ham":
        false positive += 1
    elif prediction == 0 and df.label[row] == "ham":
        true negative += 1
    elif prediction == 0 and df.label[row] == "spam":
        false_negative += 1
''' calculate the accuracy, precision, and recall'''
acc = (true_positive + true_negative) / (true_positive + true_negative +
precision = true_positive / (true_positive + false_positive)
recall = true positive / (true positive + false negative)
'''END'''
return acc, precision, recall
```

Generate answers with your functions

```
In [ ]: #loading in the training data
        train_df = pd.read_csv("./TRAIN_balanced_ham_spam.csv")
        test_df = pd.read_csv("./TEST_balanced_ham_spam.csv")
        df = train df
        df.info()
       <class 'pandas.core.frame.DataFrame'>
       RangeIndex: 2398 entries, 0 to 2397
      Data columns (total 5 columns):
                         Non-Null Count Dtype
           Column
           Unnamed: 0.1 2398 non-null
                                         int64
           Unnamed: 0 2398 non-null int64
       2
           label
                         2398 non-null
                                         object
       3
           text
                         2398 non-null
                                         object
           label num
                        2398 non-null
                                         int64
       dtypes: int64(3), object(2)
      memory usage: 93.8+ KB
In [ ]: #compute the prior
        ham_prior, spam_prior = prior(df)
        print(ham prior, spam prior)
       0.5 0.5
```

```
In [ ]: # compute likelihood
        ham like dict, spam like dict = likelihood(df)
In [ ]: # Test your predict function with some example TEXT
        some_text_example = ""
        spam text example = " Congratulations! You've been selected as a winner of c
        not_spam_text_example = "I hope this email finds you well. I'm writing to ir
        print("Not Spam text ", predict(ham_prior, spam_prior, ham_like_dict, spam_l
        print("Spam text ", predict(ham_prior, spam_prior, ham_like_dict, spam_like_
       Not Spam text 0
       Spam text 1
In [ ]: # Predict on test_df and compute metrics
        df = test df
        acc, precision, recall = metrics(ham_prior, spam_prior, ham_like_dict, spam_
        print(acc, precision, recall)
       0.745 0.9932885906040269 0.49333333333333333
In [ ]:
In []:
```

4. Gradient Decent [35 pts]

Exact Gradient Computation

Given a function f, sometimes we can compute its exact gradient at any x if f's derivative is easy to compute. For example, let $f(x)=2x^2-3x+\ln x$, where x>0. Please compute the derivative of f and report its gradient at x=2.

Your answer: 5.5

Numerical Gradient Computation

Instead of computing the derivative of a function, we can also estimate the gradient numerically with various methods. These methods are essential, especially when a callable function is not exposed due to privacy reasons, or it is hard to differentiate analytically.

To numerically compute the gradient, the simple way is to follow Newton's difference quotient: $f'(x)=\lim_{h\to 0}\frac{f(x+h)-f(x)}{h}$. Another two-point formula is to compute the slope through the points (x-h,f(x-h)) and (x+h,f(x+h)). Let us reuse the example function $f(x)=2x^2-3x+\ln x$ and test the precision of these two approaches. Define the function in the next cell, and try to compute its gradient via both

methods at x=2. Range h value in [0.1,0.01,0.001,0.0001] and report all gradients calculated. Which method is more accurate, and why does it work better?

```
In [ ]: def f(x):
            # Your code here
            return 2 * x**2 - 3 * x + math.log(x)
In [ ]: # Compute gradient using the first method (Newton's difference quotient)
        h = [0.1, 0.01, 0.001, 0.0001]
        gradient_result =[]
        for i in h:
            result = (f(2 + i) - f(2)) / i
            gradient_result.append(result)
        print(gradient_result)
       [5.687901641694317, 5.518754151103744, 5.50187504165045, 5.5001875004201395]
In []: # Compute gradient using the second method
        h = [0.1, 0.01, 0.001, 0.0001]
        gradient_result =[]
        for i in h:
            result = (f(2 + i) - f(2 - i)) / (2 * i)
            gradient_result.append(result)
```

[5.500417292784909, 5.500004166729067, 5.500000041665842, 5.50000000417948]

Remark

print(gradient_result)

You may find the gradient more accurate when using a smaller h value. However, this is not always the case. Due to the finite precision of the floating-point, rounding errors always exist and can dominate the computation when the h value is too small. Run the following two cells to observe such scenarios.

Logistic regression

Logistic regression is a classification tool that models the probability of an event taking place by having the log odds for the event be a linear combination of one or more independent variables. Specifically, let $\vec{x}=< x_1,\ldots,x_m>$ be an m dimensional vector of independent variables (features), and y be the corresponding binary dependent variable (label). The probability of having y=1 is modeled as

$$P_y = rac{1}{1 + e^{-(b_0 + b_1 \cdot x_1 + \cdots + b_m \cdot x_m)}} = rac{1}{1 + e^{-(b_0 + ec{b}_{1:m} \cdot ec{x})}}$$

Given a set of data points $<\vec{x}_k,y_k>$ with $k\in[1,n]$, how can we fit the model with these data, i.e., how to choose the best $\vec{b}=b_0,b_1\cdots,b_m$?

One way is to write out the likelihood

$$\prod_{k:y_k=1} P_{y_k} \prod_{k:y_k=0} (1-P_{y_k})$$

and find $b_0, b_1 \cdots, b_m$ that maximize its logarithm,

$$l = \sum_{k:y_k=1} \ln(P_{y_k}) + \sum_{k:y_k=0} \ln(1-P_{y_k})$$

In contrast to computing the closed form gradient of a Least-squares loss in a linear model (chapter 5 of MML book), doing the same for logistic regression is not possible. Gradient descent can be used to optimize such function l, and we will implement it step-by-step. First, write a function \log _likelihood in the next cell that computes the log-likelihood given data points and \vec{b} . [5 pts]

```
In [ ]: import numpy as np
        import sklearn
In [ ]: def log_likelihood(X,y,b):
            X: n*m numpy data array.
            y: one dimension numpy data array of length n
            b: one dimension numpy data array of length m+1
            Return the log likelihood.
            .....
            """ Add a column of ones to the X matrix."""
            ones_column = np.ones((X.shape[0], 1))
            new_X = np.append(ones_column, X, axis=1)
            """ Calculate the dot product of the new_X matrix and the b vector."""
            dot_product = np.dot(new_X, b)
            """ Calculate the log likelihood of the logistic regression model."""
            p_1 = 0
            p_0 = 0
```

```
for i in range(len(y)):
    p_y = 1 / (1 + math.exp(-dot_product[i]))
    if y[i] == 1:
        p_1 += math.log(p_y)
    else:
        p_0 += math.log(1 - p_y)
log_likelihood = p_1 + p_0
return log_likelihood
```

Test your log_likelihood function with a small example below.

```
In []: X=np.array([[0.1],[0.5],[1.]])
    y=np.array([0,0,1])
    b=np.array([0.,1.])

# Your answer should be around -2.03
    log_likelihood(X,y,b)

Out[]: -2.031735331771901
In []:
```

Now that we have a function to maximize, the next step is to compute the gradient of the log-likelihood with respect to parameter \vec{b} . Use the method with Newton's difference quotient, and set h=0.0001. Implement the function compute_gradient in the next cell. [7 pts]

```
In []: def compute_gradient(X,y,b):
# The inputs are the same as the ones of log_likelihood
h = 0.0001
gradient = np.zeros(b.shape)
for i in range(len(b)):

    print('b', b)
    new_b = b.copy()
    new_b[i] += h
    print('new_b', new_b)
    each_gradient = (log_likelihood(X, y, new_b) - log_likelihood(X,y,b))
    gradient[i] = each_gradient

return gradient
```

```
In []: # Test your function here, preserve the output
    X=np.array([[0.1],[0.5],[1.]])
    y=np.array([0,0,1])
    b=np.array([0.,1.])
    compute_gradient(X,y,b)

b [0. 1.]
    new_b [1.e-04 1.e+00]
    b [0. 1.]
    new b [0. 1.0001]
```

```
Out[]: array([-0.87853115, -0.09479906])
```

Once we know how to compute the gradients, we can optimize the objective, which is log-likelihood in our case, using gradient descent. It iteratively changes the parameters in a small "step" towards the gradient direction, i.e., the direction where the objective increases at the fastest pace. Formally, denote the calculated gradients as $\Delta(\vec{b})$, we can update our parameters via $\vec{b} = \vec{b} + \gamma \cdot \Delta(\vec{b})$, where γ is the size of the "step". Repeat this process until the objective stop improving or a pre-set max number of iterations is reached. Note in practice, the value of gradient changes over iterations and can be very large/small, so you should normalize the gradient vector every iteration, i.e., scale it to $\frac{\Delta(\vec{b})}{||\Delta(\vec{b})||_2}$, before using it to compute the new \vec{b} . Therefore, the update rule for parameters becomes $\vec{b} = \vec{b} + \gamma \cdot \frac{\Delta(\vec{b})}{||\Delta(\vec{b})||_2}$.

Implement the gradient_descent function below. [7 pts]

```
In [ ]: def gradient_descent(X, y, initial_b, step_size, max_iteration):
            X: n*m numpy data array.
            y: one dimension numpy data array of length n
            initial_b: one dimension numpy data array of length m+1
            step_size: scalar, the size of one step update
            max_iteration: scalar, the max number of iterations
            Return the updated coefficient vector b.
            b = initial b
            for i in range(max_iteration):
                gradient = compute gradient(X, y, b)
                squared_gradient = np.square(gradient)
                add_squared_gradient = 0
                for i in range(squared gradient.shape[0]):
                    add squared gradient += squared gradient[i]
                l2_norm = np.sqrt(add_squared_gradient)
                scale = np.divide(gradient, l2 norm)
                b = b + step size * scale
            result b = b
            return result_b
```

Test the function with the previous example again. Print for each sample from X, based on your model, the probability of having label=1.

```
In []: optimized_b = gradient_descent(X, y, b, 0.1, 1000)

# compute and print the probability for each row in X below using optimized_
print(optimized_b)

""" Add a column of ones to the X matrix."""
probabilities = []
```

```
ones_column = np.ones((X.shape[0], 1))
new_X = np.append(ones_column, X, axis=1)

""" Calculate the dot product of the new_X matrix and the b vector."""
dot_product = np.dot(new_X, b)

""" Calculate the log likelihood of the logistic regression model."""
for i in range(len(X)):
    p_y = 1 / (1 + math.exp(-dot_product[i]))
    probabilities.append(p_y)

print(probabilities)
```

```
[-60.22501209 80.46040454]
[0.52497918747894, 0.6224593312018546, 0.7310585786300049]
```

Next, we apply the implemented logistic regression model to a real dataset. The dataset is a trimmed breast-cancer-Wisconsin dataset from UCI machine learning Repository. Only 100 data points are offered in the training set to make sure the computation can be finished swiftly, no matter how you implement the optimizer. The training dataset is loaded in the next cell, and the vector \vec{b} is also randomly initialized.

Fit three models with the training set using different step size ranging in [0.01,0.05,0.1] and set the max number of iterations as 10000. How do the final log-likelihood value and the number of iterations change with different step sizes?

```
In []: f = open("breast-cancer-wisconsin.data","r")
X_train = []
y_train = []
for line in f:
    tmp = []
    for part in line.strip().split(",")[1:-1]:
        tmp.append(float(part))
    y_train.append((0 if line.strip().split(",")[-1]=="2" else 1))
    X_train.append(tmp)
X_train = np.array(X_train)
y_train = np.array(y_train)
random_b = np.random.uniform(0,1,size=(10))
```

```
In []: # Fit three models with different step size, report the final log-likelihood
# number of iterations and the final coefficent vector b.
""" Step size h = [0.01, 0.05, 0.1] """
h = [0.01, 0.05, 0.1]
iteration = 10000
optimized_b_array = []

for i in range(len(h)):
    optimized_b = gradient_descent(X_train, y_train, random_b, h[i], iteration optimized_b_array.append(optimized_b)
    print("Step size: ", h[i])
    print("Final log-likelihood: ", log_likelihood(X_train, y_train, optimized_b)
    print("Number of iterations: ", iteration)
    print("Final coefficient of 0.01 vector b: ", optimized_b)
# print("optimized_b arrary: ", optimized_b_array)
```

```
print("\n")
        print(optimized b array)
      Step size: 0.0005
      Final log-likelihood: -21.809518343003887
      Number of iterations: 10000
      Final coefficient of 0.01 vector b: [-2.49890791 0.18602589 0.27641924
      0.15463342 0.135314
                             0.01566887
                               0.40778241 0.084692931
        0.19051046 -0.4941777
      Step size: 0.05
      Final log-likelihood: -7.273314131672045
      Number of iterations: 10000
      Final coefficient of 0.01 vector b: [-17.49691284 1.14089114 -1.86941678
      1.22001508 1.1000927
         0.63511504 -0.06497224 1.07384467 1.31138236 1.796059551
      Step size: 0.1
      Final log-likelihood: -7.6274390848550215
      Number of iterations: 10000
      Final coefficient of 0.01 vector b: [-17.9735499 1.18472284 -1.91158275
      1.26818025 1.12626947
         0.67343613 -0.06001953
                                  1.11038459
                                              1.35435866
                                                          1.853543371
       [array([-2.49890791, 0.18602589, 0.27641924, 0.15463342, 0.135314
              0.01566887, 0.19051046, -0.4941777, 0.40778241, 0.08469293]), ar
       ray([-17.49691284, 1.14089114, -1.86941678, 1.22001508,
                            0.63511504, -0.06497224, 1.07384467,
               1.1000927 ,
                            1.79605955]), array([-17.9735499 , 1.18472284, -1.
               1.31138236,
      91158275,
                  1.26818025,
               1.12626947, 0.67343613, -0.06001953, 1.11038459,
               1.35435866, 1.85354337])]
Out[]: 'Model 1 with step size 0.01 '
```

Finally, load the test dataset, and predict for each sample in the test set what labels it should have using the model obtained. Compare your results with the ground truth labels, and report the accuracy rate.

```
In []: f = open("test_data.txt","r")
    X_test = []
    y_test = []
    for line in f:
        tmp = []
        for part in line.strip().split(",")[1:-1]:
            tmp.append(float(part))
        y_test.append((0 if line.strip().split(",")[-1]=="2" else 1))
        X_test.append(tmp)
```

```
In [ ]: # Predict based on your models and report the accuracy
```

```
""" Predict the test data using the optimized_b from the three models"""
''' Calculate the accuracy of the model'''
def accuracy(y_true, y_predict):
   length = len(y_true)
    result = 0
   for i in range(len(y true)):
        if y_true[i] == y_predict[i]:
            result += 1
    return result / length
ones_column = np.ones((len(X_test), 1))
                                                                       # Mak
new_X = np.append(ones_column, X_test, axis=1)
                                                                        # Ac
""" Calculate the dot product of the new_X matrix and the b vector."""
for i in range(len(optimized b array)):
    probabilities = []
                                                                         # Ca
    dot_product = np.dot(new_X, optimized_b_array[i])
    for j in range(len(X test)):
        p_y = 1 / (1 + math.exp(-dot_product[j]))
        if p y > 0.5:
            probabilities.append(1)
        else:
            probabilities.append(0)
    print("Step size: ", h[i])
    print("%d Optimized b : " %(i+1) , optimized_b_array[i])
    print(y test)
    print("probabilities list : ", probabilities)
    print("Accuracy: ", accuracy(y_test, probabilities))
    print("\n")
```

```
Step size: 0.0005
1 Optimized b: [-2.49890791 0.18602589 0.27641924 0.15463342 0.135314
0.01566887
 0.19051046 -0.4941777   0.40778241   0.08469293]
[0, 1, 1, 1, 1, 1, 0, 1, 0, 1]
probabilities list: [0, 0, 1, 1, 1, 1, 0, 1, 0, 1]
Accuracy: 0.9
Step size: 0.05
1.10
00927
  0.63511504 -0.06497224 1.07384467 1.31138236
                                               1.796059551
[0, 1, 1, 1, 1, 1, 0, 1, 0, 1]
probabilities list: [0, 1, 1, 1, 1, 1, 0, 1, 0, 0]
Accuracy: 0.9
Step size: 0.1
                                                   1.26818025
3 Optimized b : [-17.9735499 1.18472284 -1.91158275
                                                              1.12
626947
  0.67343613 -0.06001953 1.11038459 1.35435866
                                              1.853543371
[0, 1, 1, 1, 1, 1, 0, 1, 0, 1]
probabilities list: [0, 1, 1, 1, 1, 1, 0, 1, 0, 0]
Accuracy: 0.9
```

In []: