```
In []: # ALL IMPORTS
    import pandas as pd
    import numpy as np
    import matplotlib.pyplot as plt
    from sklearn.model_selection import train_test_split, StratifiedKFold
    from sklearn.neighbors import KNeighborsClassifier
    from sklearn.linear_model import LogisticRegression
    from sklearn.discriminant_analysis import LinearDiscriminantAnalysis
    from sklearn.ensemble import RandomForestClassifier
    from sklearn.metrics import roc curve, auc, log loss, roc auc score, fl score
```

# Assignment 3 - Supervised Learning: model training and evaluation

### Jake Bova

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Names of students you worked with on this assignment. Bri, Brock

Note: this assignment falls under collaboration Mode 2: Individual Assignment – Collaboration Permitted. Please refer to the syllabus for additional information.

Instructions for all assignments can be found here, and is also linked to from the course syllabus.

Total points in the assignment add up to 90; an additional 10 points are allocated to presentation quality.

# Learning Objectives:

This assignment will provide structured practice to help enable you to...

- Understand the primary workflow in machine learning: (1) identifying a hypothesis function set of models, (2) determining a loss/cost/error/objective function to minimize, and (3) minimizing that function through gradient descent
- 2. Understand the inner workings of logistic regression and how linear models for classification can be developed.
- 3. Gain practice in implementing machine learning algorithms from the most basic building blocks to understand the math and programming behind them to achieve practical proficiency with the techniques
- 4. Implement batch gradient descent and become familiar with how that technique is used and its dependence on the choice of learning rate

- 5. Evaluate supervised learning algorithm performance through ROC curves and using cross validation
- 6. Apply regularization to linear models to improve model generalization performance

### 1

# Classification using logistic regression: build it from the ground up

### [60 points]

This exercise will walk you through the full life-cycle of a supervised machine learning classification problem. Classification problem consists of two features/predictors (e.g. petal width and petal length) and your goal is to predict one of two possible classes (class 0 or class 1). You will build, train, and evaluate the performance of a logistic regression classifier on the data provided. Before you begin any modeling, you'll load and explore your data in Part I to familiarize yourself with it - and check for any missing or erroneous data. Then, in Part II, we will review an appropriate hypothesis set of functions to fit to the data: in this case, logistic regression. In Part III, we will derive an appropriate cost function for the data (spoiler alert: it's cross-entropy) as well as the gradient descent update equation that will allow you to optimize that cost function to identify the parameters that minimize the cost for the training data. In Part IV, all the pieces come together and you will implement your logistic regression model class including methods for fitting the data using gradient descent. Using that model you'll test it out and plot learning curves to verify the model learns as you train it and to identify and appropriate learning rate hyperparameter. Lastly, in Part V you will apply the model you designed, implemented, and verified to your actual data and evaluate and visualize its generalization performance as compared to a KNN algorithm. When complete, you will have accomplished learning objectives 1-5 above!

## I. Load, prepare, and plot your data

You are given some data for which you are tasked with constructing a classifier. The first step when facing any machine learning project: look at your data!

### (a) Load the data.

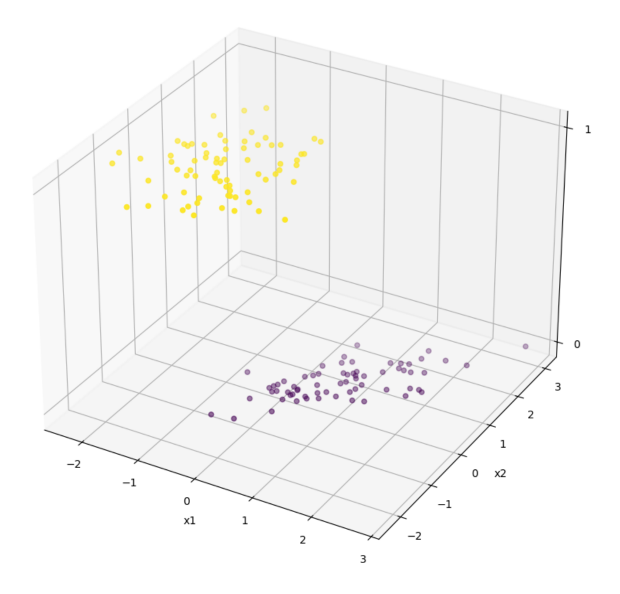
- In the data folder in the same directory of this notebook, you'll find the data in A3\_Q1\_data.csv . This file contains the binary class labels, y, and the features  $x_1$  and  $x_2$ .
- Divide your data into a training and testing set where the test set accounts for 30 percent of the data and the training set the remaining 70 percent.
- Plot the training data by class.

- Comment on the data: do the data appear separable? May logistic regression be a good choice for these data? Why or why not?
- **(b)** Do the data require any preprocessing due to missing values, scale differences (e.g. different ranges of values), etc.? If so, how did you handle these issues?

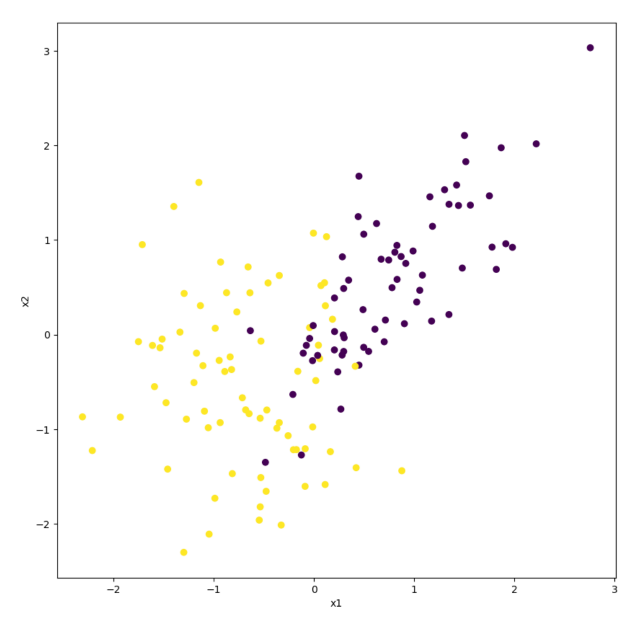
Next, we walk through our key steps for model fitting: choose a hypothesis set of models to train (in this case, logistic regression); identify a cost function to measure the model fit to our training data; optimize model parameters to minimize cost (in this case using gradient descent). Once we've completed model fitting, we will evaluate the performance of our model and compare performance to another approach (a KNN classifier).

### **START I -----**

```
In [ ]: # a) Load the data
        data = pd.read csv('data/A3 Q1 data.csv')
        # a1) Split the data into training and test sets
        # feature matrix X, target vector y
        X = data.drop('y', axis=1)
        y = data['y']
        # split data
        X train, X test, y train, y test = train test split(X, y, test size=0.3, ran
        # normalize data
        X train = (X train - X train.mean()) / X train.std()
        X test = (X test - X test.mean()) / X test.std()
        # a2) plot the training data by class
        fig = plt.figure(figsize=(10, 10))
        ax = fig.add subplot(111, projection='3d')
        ax.scatter(X train['x1'], X train['x2'], y train, c=y train, cmap='viridis')
        ax.set xlabel('x1')
        ax.set ylabel('x2')
        ax.set zlabel('y')
        ax.set_zticks([0, 1])
        plt.show()
```



```
In []: # graph the data in 2d colored by class (0 purple, 1 yellow)
    fig = plt.figure(figsize=(10, 10))
    ax = fig.add_subplot(111)
    ax.scatter(X_train['x1'], X_train['x2'], c=y_train, cmap='viridis')
    ax.set_xlabel('x1')
    ax.set_ylabel('x2')
    plt.show()
```



In [ ]: # b) assess data
print(data.isna().sum()) # no missing values

END I -----

# II. Stating the hypothesis set of models to evaluate (we'll use logistic regression)

Given that our data consists of two features, our logistic regression problem will be applied to a two-dimensional feature space. Recall that our logistic regression model is:

$$f(\mathbf{x}_i, \mathbf{w}) = \sigma(\mathbf{w}^\top \mathbf{x}_i)$$

where the sigmoid function is defined as  $\sigma(x)=\frac{e^x}{1+e^x}=\frac{1}{1+e^{-x}}$ . Also, since this is a two-dimensional problem, we define  $\mathbf{w}^{\top}\mathbf{x}_i=w_0x_{i,0}+w_1x_{i,1}+w_2x_{i,2}$  and here,  $\mathbf{x}_i=[x_{i,0},x_{i,1},x_{i,2}]^{\top}$ , and  $x_{i,0}\triangleq 1$ 

Remember from class that we interpret our logistic regression classifier output (or confidence score) as the conditional probability that the target variable for a given sample  $y_i$  is from class "1", given the observed features,  $\mathbf{x}_i$ . For one sample,  $(y_i, \mathbf{x}_i)$ , this is given as:

$$P(Y = 1|X = \mathbf{x}_i) = f(\mathbf{x}_i, \mathbf{w}) = \sigma(\mathbf{w}^{\top} \mathbf{x}_i)$$

In the context of maximizing the likelihood of our parameters given the data, we define this to be the likelihood function  $L(\mathbf{w}|y_i,\mathbf{x}_i)$ , corresponding to one sample observation from the training dataset.

Aside: the careful reader will recognize this expression looks different from when we talk about the likelihood of our data given the true class label, typically expressed as P(x|y), or the posterior probability of a class label given our data, typically expressed as P(y|x). In the context of training a logistic regression model, the likelihood we are interested in is the likelihood function of our logistic regression **parameters**, w. It's our goal to use this to choose the parameters to maximize the likelihood function.

No output is required for this section - just read and use this information in the later sections.

# III. Find the cost function that we can use to choose the model parameters, w, that best fit the training data.

- (c) What is the likelihood function that corresponds to all the N samples in our training dataset that we will wish to maximize? Unlike the likelihood function written above which gives the likelihood function for a *single training data pair*  $(y_i, \mathbf{x}_i)$ , this question asks for the likelihood function for the *entire training dataset*  $\{(y_1, \mathbf{x}_1), (y_2, \mathbf{x}_2), \dots, (y_N, \mathbf{x}_N)\}$ .
- (d) Since a logarithm is a monotonic function, maximizing the f(x) is equivalent to maximizing  $\ln[f(x)]$ . Express the likelihood from the last question as a cost function of the model parameters,  $C(\mathbf{w})$ ; that is the negative of the logarithm of the likelihood. Express this cost as an average cost per sample (i.e. divide your final value by N), and use this quantity going forward as the cost function to optimize.
- (e) Calculate the gradient of the cost function with respect to the model parameters  $\nabla_{\mathbf{w}} C(\mathbf{w})$ . Express this in terms of the partial derivatives of the cost function with respect to each of the parameters, e.g.  $\nabla_{\mathbf{w}} C(\mathbf{w}) = \left[ \frac{\partial C}{\partial w_0}, \frac{\partial C}{\partial w_1}, \frac{\partial C}{\partial w_2} \right]$ .

To simplify notation, please use  $\mathbf{w}^{\top}\mathbf{x}$  instead of writing out  $w_0x_{i,0}+w_1x_{i,1}+w_2x_{i,2}$  when it appears each time (where  $x_{i,0}=1$  for all i). You are also welcome to use  $\sigma()$  to represent the sigmoid function. Lastly, this will be a function the features,  $x_{i,j}$  (with the first index in the subscript representing the observation and the second the feature; targets,  $y_i$ ; and the logistic regression model parameters,  $w_j$ .

**(f)** Write out the gradient descent update equation. This should clearly express how to update each weight from one step in gradient descent  $w_j^{(k)}$  to the next  $w_j^{(k+1)}$ . There should be one equation for each model logistic regression model parameter (or you can represent it in vectorized form). Assume that  $\eta$  represents the learning rate.

#### START III -----

**c)** The likelihood function for all N samples can be expressed as the product of the individual likelihood functions for each data point (under the assumtion that the data points are independent and identically distributed).

Given sigmoid function:

$$f(\mathbf{x}_i, \mathbf{w}) = \sigma(\mathbf{w}^ op \mathbf{x}_i) = rac{1}{1 + e^{-\mathbf{w}^ op \mathbf{x}_i}}$$

The likelihood for a single data point  $(y_i, \mathbf{x}_i)$  is given by:

$$L(\mathbf{w}|y_i, \mathbf{x}_i) = \sigma(\mathbf{w}^ op \mathbf{x}_i)^{y_i} (1 - \sigma(\mathbf{w}^ op \mathbf{x}_i))^{1-y_i}$$

This equation reflects the binary nature of the data, where  $y_i$  is either 0 or 1, so the equation is similar to the Bernoulli distribution.

Since the data points are independent and identically distributed, we can multiply the likelihoods for each data point to get the likelihood for the entire dataset:

$$L(\mathbf{w}|y_i, \mathbf{x}_i) = \prod_{i=1}^N \sigma(\mathbf{w}^ op \mathbf{x}_i)^{y_i} (1 - \sigma(\mathbf{w}^ op \mathbf{x}_i))^{1-y_i}$$

**d)** In order to express the likelihood function as a cost function (using log likelihood), we take the negative log of the likelihood function (on the parameter  $\mathbf{w}$ ):

$$C(\mathbf{w}) = -\ln L(\mathbf{w}|y_i, \mathbf{x}_i) = -\sum_{i=1}^N y_i \ln \sigma(\mathbf{w}^ op \mathbf{x}_i) + (1-y_i) \ln (1-\sigma(\mathbf{w}^ op \mathbf{x}_i))$$

Note that taking the logarithm of this function allows us to move the exponent from the sigmoid function to the outside of the function, which makes it easier to take the derivative.

From this, we divide by N to get the average cost per sample:

- (I am defining  $\hat{y} = \sigma(\mathbf{w}^{ op}\mathbf{x}_i)$  where applicable)

$$C(\mathbf{w}) = -rac{1}{N} \sum_{i=1}^N y_i \ln \sigma(\mathbf{w}^ op \mathbf{x}_i) + (1-y_i) \ln (1-\sigma(\mathbf{w}^ op \mathbf{x}_i))$$

**e)** To calculate the gradient of the cost function with respect to the model parameters  $\nabla_{\mathbf{w}} C(\mathbf{w})$ , we express it in terms of the partial derivatives of the cost function with respect to each of the parameters, e.g.  $\nabla_{\mathbf{w}} C(\mathbf{w}) = \left[ \frac{\partial C}{\partial w_0}, \frac{\partial C}{\partial w_1}, \frac{\partial C}{\partial w_2} \right]$ :

• 1. setup:

$$rac{\partial C}{\partial w_j} = -rac{1}{N} \sum_{i=1}^N y_i rac{\partial}{\partial w_j} {
m ln} \, \sigma({f w}^ op {f x}_i) + (1-y_i) rac{\partial}{\partial w_j} {
m ln} (1-\sigma({f w}^ op {f x}_i))$$

- 1.1) take the derivatives of the log-sigmoid function (a) and log of 1-sigmoid function (b):
- (a)

$$rac{\partial}{\partial w_j} \mathrm{ln}\, \sigma(\mathbf{w}^ op \mathbf{x}_i) = rac{\mathbf{x}_{ij}}{\sigma(\mathbf{w}^ op \mathbf{x}_i)}$$

• (b)

$$rac{\partial}{\partial w_j} ext{ln} (1 - \sigma(\mathbf{w}^ op \mathbf{x}_i)) = -rac{\mathbf{x}_{ij}}{1 - \sigma(\mathbf{w}^ op \mathbf{x}_i)}$$

• 2. substitute (a) and (b) into the setup:

$$\frac{\partial C}{\partial w_j} = -\frac{1}{N} \sum_{i=1}^{N} \left( y_i \frac{\mathbf{x}_{ij}}{\sigma(\mathbf{w}^{\top} \mathbf{x}_i)} - (1 - y_i) \frac{\mathbf{x}_{ij}}{1 - \sigma(\mathbf{w}^{\top} \mathbf{x}_i)} \right)$$

• 3. combine terms:

$$rac{\partial C}{\partial w_j} = -rac{1}{N} \sum_{i=1}^N \mathbf{x}_{ij} \left( y_i rac{1}{\sigma(\mathbf{w}^ op \mathbf{x}_i)} - (1-y_i) rac{1}{1-\sigma(\mathbf{w}^ op \mathbf{x}_i)} 
ight)$$

 4. we can simplify the expression in parentheses (by performing the arithmetic then cancelling terms):

$$rac{\partial C}{\partial w_j} = -rac{1}{N} \sum_{i=1}^N \mathbf{x}_{ij} \left( y_i - \sigma(\mathbf{w}^ op \mathbf{x}_i) 
ight)$$

• 5. the gradient vector would be:

$$abla_{\mathbf{w}} C(\mathbf{w}) = rac{1}{N} \sum_{i=1}^{N} (\sigma(\mathbf{w}^{ op} \mathbf{x}_i) - y_i) \mathbf{x}_{ij}$$

$$for \ x = \left[ rac{\partial C}{\partial w_0}, rac{\partial C}{\partial w_1}, rac{\partial C}{\partial w_2} 
ight]$$

#### Where:

- $\mathbf{x}_{ij}$  is the  $j^{th}$  feature of the  $i^{th}$  data point
- $\mathbf{w}^{ op}\mathbf{x}_i$  is the dot product of the weight vector and the feature vector for the  $i^{th}$  data point
- $\sigma(\mathbf{w}^{\top}\mathbf{x}_i)$  is the sigmoid function applied to the dot product of the weight vector and the feature vector for the  $i^{th}$  data point
- ullet  $y_i$  is the target value for the  $i^{th}$  data point

**f)** The gradient descent update equation is used to iteratively adjust model params (weights) to minimize the cost function. The update equation is:

$$w_j^{(k+1)} = w_j^{(k)} - \eta 
abla_{\mathbf{w}} C(\mathbf{w})$$

#### Where:

- $\eta$  is the learning rate.
- $w_{j}^{\left(k
  ight)}$  is the value of the weight  $w_{j}$  at iteration k.
- $ullet w_j^{(k+1)}$  is the value of the updated weight  $w_j$  at iteration k+1 .
- $\dfrac{\partial C}{\partial w_j}$  is the partial derivative of the cost function with respect to the weight  $w_j$  (see above)

END III -----

# IV. Implement gradient descent and your logistic regression algorithm

- (g) Implement your logistic regression model.
  - You are provided with a template, below, for a class with key methods to help with your model development. It is modeled on the Scikit-Learn convention. For this, you only need to create a version of logistic regression for the case of two feature variables (i.e. two predictors).
  - Create a method called sigmoid that calculates the sigmoid function
  - Create a method called cost that computes the cost function  $C(\mathbf{w})$  for a given dataset and corresponding class labels. This should be the **average cost** (make sure your total cost is divided by your number of samples in the dataset).
  - Create a method called <code>gradient\_descent</code> to run **one step** of gradient descent on your training data. We'll refer to this as "batch" gradient descent since it takes into account the gradient based on all our data at each iteration of the algorithm.
  - Create a method called fit that fits the model to the data (i.e. sets the model parameters to minimize cost) using your gradient\_descent method. In doing this we'll need to make some assumptions about the following:

- Weight initialization. What should you initialize the model parameters to? For this, randomly initialize the weights to a different values between 0 and 1.
- Learning rate. How slow/fast should the algorithm step towards the minimum? This you will vary in a later part of this problem.
- Stopping criteria. When should the algorithm be finished searching for the optimum? There are two stopping criteria: small changes in the gradient descent step size and a maximum number of iterations. The first is whether there was a sufficiently small change in the gradient; this is evaluated as whether the magnitude of the step that the gradient descent algorithm takes changes by less than  $10^{-6}$  between iterations. Since we have a weight vector, we can compute the change in the weight by evaluating the  $L_2$  norm (Euclidean norm) of the change in the vector between iterations. From our gradient descent update equation we know that mathematically this is  $||-\eta\nabla_{\mathbf{w}}C(\mathbf{w})||$ . The second criterion is met if a maximum number of iterations has been reach (5,000 in this case, to prevent infinite loops from poor choices of learning rates).
- Design your approach so that at each step in the gradient descent algorithm you evaluate the cost function for both the training and the test data for each new value for the model weights. You should be able to plot cost vs gradient descent iteration for both the training and the test data. This will allow you to plot "learning curves" that can be informative for how the model training process is proceeding.
- Create a method called predict\_proba that predicts confidence scores (that can be thresholded into the predictions of the predict method.
- Create a method called predict that makes predictions based on the trained model, selecting the most probable class, given the data, as the prediction, that is class that yields the larger  $P(y|\mathbf{x})$ .
- (Optional, but recommended) Create a method called <a href="learning\_curve">learning\_curve</a> that produces the cost function values that correspond to each step from a previously run gradient descent operation.
- (Optional, but recommended) Create a method called <code>prepare\_x</code> which appends a column of ones as the first feature of the dataset  ${\bf X}$  to account for the bias term (  $x_{i,1}=1$ ).

This structure is strongly encouraged; however, you're welcome to adjust this to your needs (adding helper methods, modifying parameters, etc.).

Class: LogisticRegression

```
Method Initialize():
    weights <- None
    savedWeights <- Empty List
    savedTrainCosts <- Empty List
    savedTestCosts <- Empty List</pre>
```

```
Method Sigmoid(X, weights):
    Return 1 / (1 + e^{-X * weights})
Method Cost(X, y, weights):
    Return -Mean(y * log(Sigmoid(X, weights)) + (1-y) * log(1
- Sigmoid(X, weights)))
Method GradientDescent(X, y, learningRate):
    numSamples <- Number of rows in X
    gradient <- Transpose(X) * (Sigmoid(X, weights) - y) /</pre>
numSamples
    oldWeights <- Copy of weights
    weights <- weights - learningRate * gradient</pre>
    Append weights to savedWeights
    Return EuclideanNorm of (weights - oldWeights)
Method Fit(X, y, initialWeights, learningRate,
deltaThreshold=1e-6, maxIterations=5000, verbose=False):
    X <- AddColumnOfOnes to X
    weights <- Randomly initialize weights based on number of
features in X
    weightChange <- Infinity</pre>
    iterationCount <- 0
    WHILE weightChange > deltaThreshold AND iterationCount <
maxIterations:
        weightChange <- GradientDescent(X, y, learningRate)</pre>
        trainCost <- Cost(X, y, weights)</pre>
        Append trainCost to savedTrainCosts
        Increment iterationCount
        IF verbose:
            Print weights
Method PredictProba(X):
    X <- AddColumnOfOnes to X
    Return Sigmoid(X, weights)
Method Predict(X, threshold=0.5):
    confidenceScore <- PredictProba(X)</pre>
```

```
Return 1 if confidenceScore >= threshold else 0 for each
score in confidenceScore

Method LearningCurve(X, y):
    X <- AddColumnOfOnes to X

    trainCosts <- Empty List

FOR each weight in savedWeights:
        Append Cost(X, y, weight) to trainCosts

Return trainCosts

Method AddColumnOfOnes(X):
    bias <- Create column of ones with number of rows equal to X</pre>
```

Return Concatenate bias and X column-wise

```
In [ ]: # Logistic regression class
        class Logistic regression:
            # Class constructor
            def init (self):
                self.w = None # logistic regression weights
                self.saved w = [] # Since this is a small problem, we can save the
                # at each iteration of gradient descent to build our
                # learning curves
                self.saved train costs = []
                self.saved test costs = []
                # returns nothing
                pass
            # Method for calculating the sigmoid function of w^T X for an input set
            def sigmoid(self, X, w):
                # returns the value of the sigmoid
                sigmoid = 1 / (1 + np.exp(-np.dot(X, w)))
                return sigmoid
            # Cost function for an input set of weights
            def cost(self, X, y, w):
                # returns the average cross entropy cost
                avg cost = -np.mean(
                    y * np.log(self.sigmoid(X, w)) + (1 - y) * np.log(1 - self.sigmoid(X, w))
                return avg cost
            # Update the weights in an iteration of gradient descent
            def gradient descent(self, X, y, lr):
                # returns a scalar of the magnitude of the Euclidean norm
                # of the change in the weights during one gradient descent step
                m = X.shape[0] # number of samples
                # calculate the gradient
                grad = np.dot(X.T, (self.sigmoid(X, self.w) - y)) / m
```

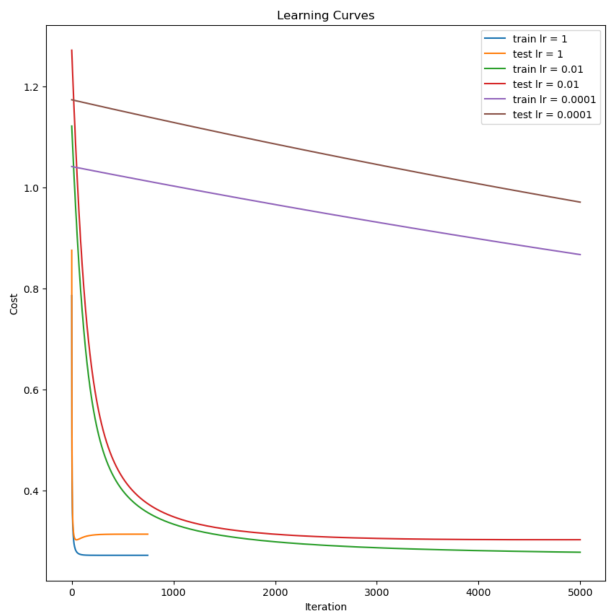
```
# save old weights
    old w = self.w.copy()
   # update the weights
    self.w = self.w - (lr * grad)
   # save the weights
    self.saved w.append(self.w)
   # calculate the magnitude of the change in the weights
   weight change = np.linalg.norm(self.w - old w)
    # weight change = np.linalg.norm(lr * grad)
    return weight change
# Fit the logistic regression model to the data through gradient descent
def fit(self, X, y, w init, lr, delta thresh=1e-6, max iter=5000, verbos
   # Note the verbose flag enables you to print out the weights at each
   # (optional - but may help with one of the guestions)
   X = self.prepare x(X) # add a column of ones to X
    self.w = np.random.rand(X.shape[1]) # initialize the weights
   weight_change = np.inf # initialize the weight change
   iter count = 0
    # while the weight change is greater than the threshold and the iter
   while weight change > delta thresh and iter count < max iter:</pre>
        weight change = self.gradient descent(X, y, lr) # update the we
        train cost = self.cost(X, y, self.w) # calculate the cost
        self.saved train costs.append(train cost) # save the costs
        iter count += 1
       # print the weights if verbose
       if verbose:
            print(self.w)
    # returns nothing
    pass
# Use the trained model to predict the confidence scores (prob of positi
def predict proba(self, X):
   # returns the confidence score for the each sample
   X = self.prepare x(X) # add a column of ones to X
    confidence score = self.sigmoid(X, self.w)
    return confidence score
# Use the trained model to make binary predictions
def predict(self, X, thresh=0.5):
    # returns a binary prediction for each sample
    confidence_score = self.predict_proba(X) # get the confidence score
   binary pred = np.where(
        confidence score >= thresh, 1, 0
    ) # make the binary prediction
    return binary pred
# Stores the learning curves from saved weights from gradient descent
def learning curve(self, X, y):
```

```
# returns the value of the cost function from each step in gradient
    # from the last model fitting process
   X = self.prepare x(X)
   train costs = []
    for w in self.saved w:
       train costs.append(
            self.cost(X, y, w)
        ) # calculate the cost for each set of weights
    return train costs
# Appends a column of ones as the first feature to account for the bias
def prepare x(self, X):
   # returns the X with a new feature of all ones (a column that is the
   bias = np.ones(
       (X.shape[0], 1)
    ) # ones creates an array of ones in the shape of the input
    return np.hstack(
       [bias, X]
    ) # hstack stacks arrays in sequence horizontally (column wise)
```

- (h) Choose a learning rate and fit your model. Learning curves are a plot of metrics of model performance evaluated through the process of model training to provide insight about how model training is proceeding. Show the learning curves for the gradient descent process for learning rates of  $\{10^{-0}, 10^{-2}, 10^{-4}\}$ . For each learning rate plot the learning curves by plotting **both the training and test data average cost** as a function of each iteration of gradient descent. You should run the model fitting process until it completes (up to 5,000 iterations of gradient descent). Each of the 6 resulting curves (train and test average cost for each learning rate) should be plotted on the **same set of axes** to enable direct comparison. *Note: make sure you're using average cost per sample, not the total cost*.
  - Try running this process for a really big learning rate for this problem:  $10^2$ . Look at the weights that the fitting process generates over the first 50 iterations and how they change. Either print these first 50 iterations as console output or plot them. What happens? How does the output compare to that corresponding to a learning rate of  $10^0$  and why?
  - What is the impact that the different values of learning have on the speed of the process and the results?
  - Of the options explored, what learning rate do you prefer and why?
  - Use your chosen learning rate for the remainder of this problem.

```
In []: # h) Plot the learning curves
# plot the learning curves for learning rates of 1, 0.01, and 0.0001
fig = plt.figure(figsize=(10, 10))
for lr in [1, 0.01, 0.0001]:
    logreg = Logistic_regression()
    logreg.fit(X_train, y_train, w_init=0, lr=lr, max_iter=5000, verbose=Fal
    plt.plot(logreg.learning_curve(X_train, y_train), label='train lr = ' +
    plt.plot(logreg.learning_curve(X_test, y_test), label='test lr = ' + str
plt.xlabel('Iteration')
```

```
plt.ylabel('Cost')
plt.title('Learning Curves')
plt.legend()
plt.show()
```



For the learning rates shown above, I would select the 0.01 rate. Lower rates take too long to converge, and may not provide the best results. Higher rates converge too quickly, and may prevent the model from finding the best solution.

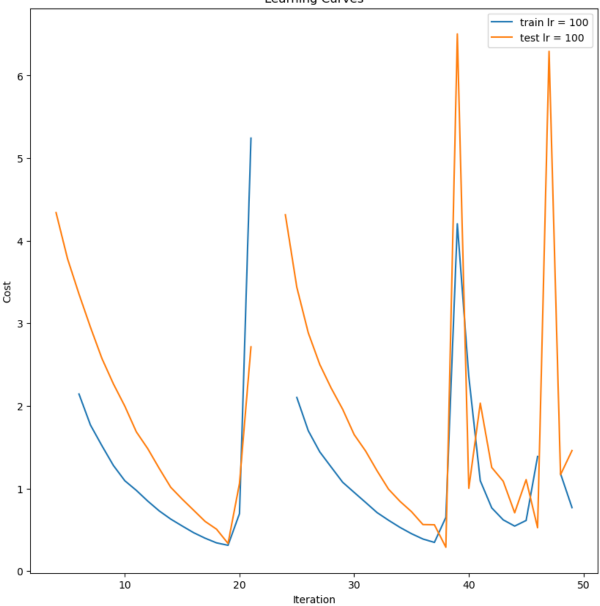
```
In []: # lr of 100
    logreg4 = Logistic_regression()
    logreg4.fit(X_train, y_train, w_init=0, lr=100, max_iter=5000, verbose=False
    logreg4.predict(X_test)
    # accuracy
    print('Accuracy: ', np.mean(logreg4.predict(X_test) == y_test))
    # get learning curve
    lclr4train = logreg4.learning_curve(X_train, y_train)
    lclr4test = logreg4.learning_curve(X_test, y_test)
```

```
# plot the first 50 iterations
fig = plt.figure(figsize=(10, 10))
plt.plot(lclr4train[:50], label='train lr = 100')
plt.plot(lclr4test[:50], label='test lr = 100')
plt.xlabel('Iteration')
plt.ylabel('Cost')
plt.title('Learning Curves')
plt.legend()
plt.show()
avg_cost = -np.mean(y * np.log(self.sigmoid(X, w)) + (1 - y) * np.log(1 - y)
```

/tmp/ipykernel 49010/1254693442.py:23: RuntimeWarning: divide by zero encoun tered in log self.sigmoid(X, w)))

Accuracy: 0.8333333333333334





I'm not exactly sure what is happening here. It seems that the learning rate of 100 is too high, and the model is overshooting the minimum. This is causing the cost to increase, and the weights to oscillate. The learning rate of 1 is also too high, but not as bad as 100. The learning rate of 0.01 is much better, and the cost is decreasing as expected.

### V. Evaluate your model performance through cross validation

(i) Test the performance of your trained classifier using K-folds cross validation resampling technique. The scikit-learn package StratifiedKFolds may be helpful.

- Train your logistic regression model and a K-Nearest Neighbor classification model with k=7 nearest neighbors.
- Using the trained models, make four plots: two for logistic regression and two for KNN. For each model have one plot showing the training data used for fitting the model, and the other showing the test data. On each plot, include the decision boundary resulting from your trained classifier.
- Produce a Receiver Operating Characteristic curve (ROC curve) that represents the performance from cross validated performance evaluation for each classifier (your logistic regression model and the KNN model, with k=7 nearest neighbors). For the cross validation, use k=10 folds.
  - Plot these curves on the same set of axes to compare them
  - On the ROC curve plot, also include the chance diagonal for reference (this represents the performance of the worst possible classifier). This is represented as a line from (0,0) to (1,1).
  - Calculate the Area Under the Curve for each model and include this measure in the legend of the ROC plot.
- Comment on the following:
  - What is the purpose of using cross validation for this problem?
  - How do the models compare in terms of performance (both ROC curves and decision boundaries) and which model (logistic regression or KNN) would you select to use on previously unseen data for this problem and why?

#### **ANSWER**

```
In []: X = (X - X.mean()) / X.std() # normalize the data

X_np = X.to_numpy() # convert to numpy arrays
y_np = y.to_numpy()

# Initialize the models
log_reg = Logistic_regression()
knn = KNeighborsClassifier(n_neighbors=7)

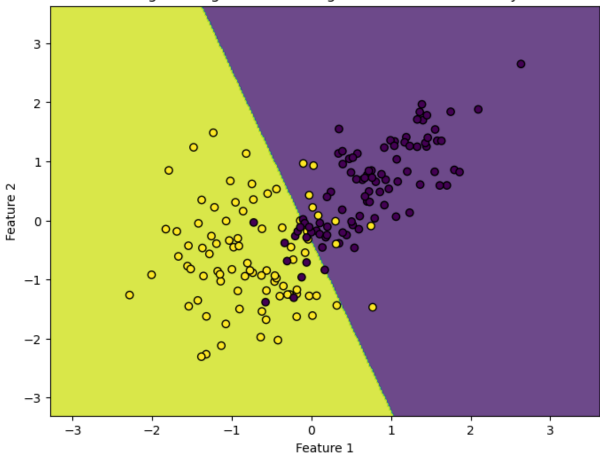
# Initialize StratifiedKFolds
skf = StratifiedKFold(n_splits=10)

log_reg_tprs = [] # true positive rates
log_reg_aucs = [] # area under the curve
knn_tprs = [] # true positive rates
```

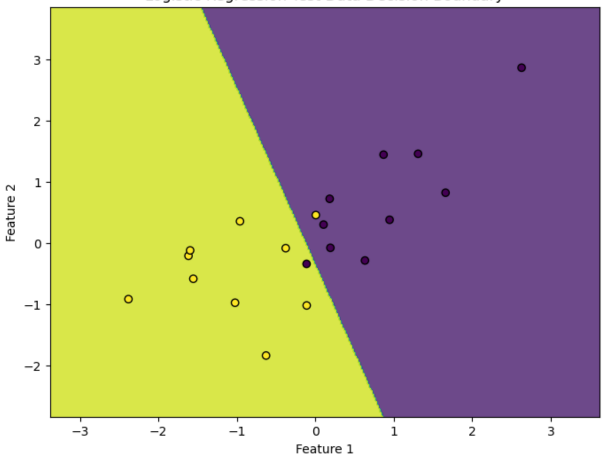
```
knn_aucs = [] # area under the curve
mean fpr = np.linspace(0, 1, 100) # mean false positive rate
for train index, test index in skf.split(X np, y np):
   X train, X test = X np[train index], X np[test index]
   y train, y test = y np[train index], y np[test index]
   # Train logistic regression
   log reg.fit(X train, y train, w init=None, lr=0.01)
   y pred prob log = log reg.predict proba(X test)
   fpr, tpr, = roc curve(y test, y pred prob log)
   log_reg_tprs.append(np.interp(mean_fpr, fpr, tpr)) # interpolate the tru
   log reg aucs.append(auc(fpr, tpr)) # calculate the area under the curve
   # Train KNN
   knn.fit(X train, y train)
   y pred prob knn = knn.predict proba(X test)[:, 1]
   fpr, tpr, _ = roc_curve(y_test, y_pred_prob_knn)
   knn tprs.append(np.interp(mean fpr, fpr, tpr))
   knn aucs.append(auc(fpr, tpr))
```

```
In [ ]: # CODE FOR PLOTTING DECISION BOUNDARIES IS SOURCED FROM MY DATA SCIENCE ANAL
        def plot decision boundary(model, X, y, model name="Model"):
            plt.figure(figsize=(8, 6))
            h = .02
            x \min, x \max = X[:, 0].\min() - 1, X[:, 0].\max() + 1
            y \min, y \max = X[:, 1].min() - 1, X[:, 1].max() + 1
            xx, yy = np.meshgrid(np.arange(x min, x max, h),
                                 np.arange(y min, y max, h))
            if model name == "Logistic Regression":
                Z = model.predict(np.c [xx.ravel(), yy.ravel()])
            else:
                Z = model.predict(np.c [xx.ravel(), yy.ravel()])
            Z = Z.reshape(xx.shape)
            plt.contourf(xx, yy, Z, alpha=0.8)
            plt.scatter(X[:, 0], X[:, 1], c=y, edgecolors='k', marker='o')
            plt.xlabel('Feature 1')
            plt.ylabel('Feature 2')
            plt.title(model name + ' Decision Boundary')
            plt.show()
        # Plot decision boundaries for logistic regression and KNN
        plot_decision_boundary(log_reg, X_train, y_train, "Logistic Regression Train
        plot decision boundary(log reg, X test, y test, "Logistic Regression Test Da
        plot decision boundary(knn, X train, y train, "KNN Training Data")
        plot decision boundary(knn, X test, y test, "KNN Test Data")
```

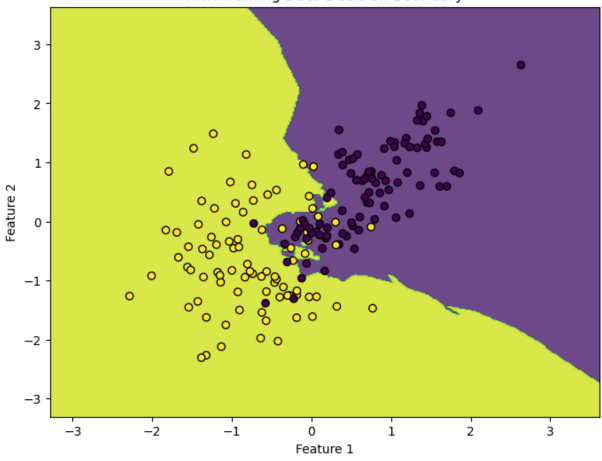
Logistic Regression Training Data Decision Boundary



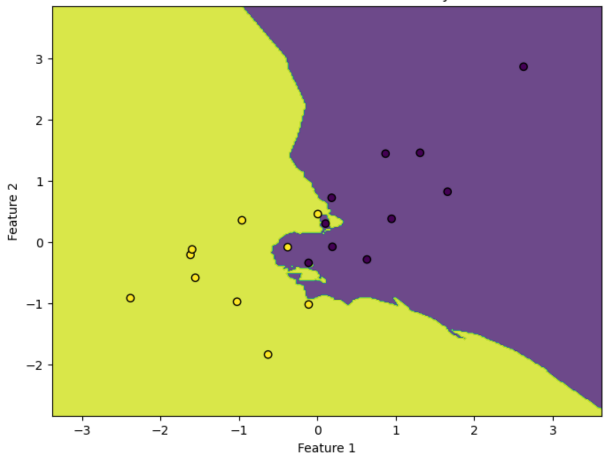
Logistic Regression Test Data Decision Boundary



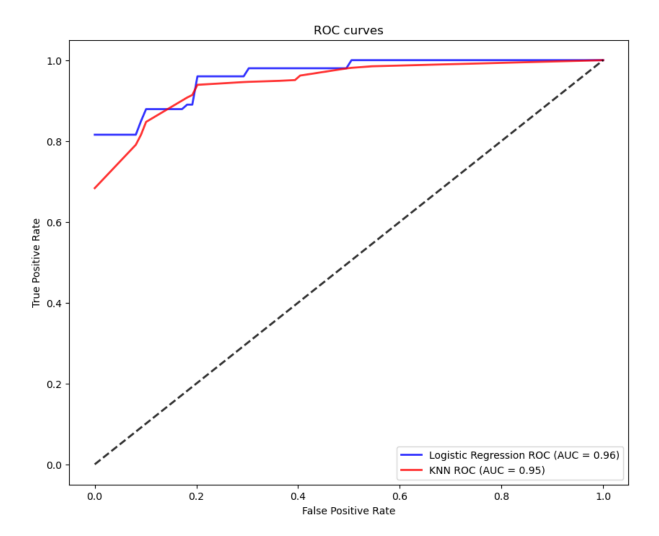
## KNN Training Data Decision Boundary



### KNN Test Data Decision Boundary



```
In [ ]: plt.figure(figsize=(10, 8))
        # Logistic Regression
        mean tpr = np.mean(log reg tprs, axis=0)
        mean_auc = auc(mean_fpr, mean_tpr)
        plt.plot(mean_fpr, mean_tpr, color='blue', label=r'Logistic Regression ROC (
        # KNN
        mean tpr = np.mean(knn tprs, axis=0)
        mean auc = auc(mean fpr, mean tpr)
        plt.plot(mean fpr, mean tpr, color='red', label=r'KNN ROC (AUC = %0.2f)' % m
        # Chance Diagonal
        plt.plot([0, 1], [0, 1], linestyle='--', lw=2, color='black', alpha=0.8)
        plt.xlabel('False Positive Rate')
        plt.ylabel('True Positive Rate')
        plt.title('ROC curves')
        plt.legend(loc='lower right')
        plt.show()
```



The purpose of using cross validation for this problem is to evaluate the performance of the model on data that was not used to train the model. This allows us to see how well the model generalizes to new data. Holding out data in validation sets keeps the model training process clean of any bias that may be introduced by using the same data for training and testing.

Comparing the models in terms of performance shows that, for this problem, logistic regression is the better model. The ROC curve for logistic regression is closer to the top left corner of the plot, which indicates a better model. The decision boundary for logistic regression is a straight line, which seems to be a better fit than the curve generated by KNN.

# 2

# Digits classification

### [30 points]

An exploration of regularization, imbalanced classes, ROC and PR curves

The goal of this exercise is to apply your supervised learning skills on a very different dataset: in this case, image data; MNIST: a collection of images of handwritten digits. Your goal is to train a classifier that is able to distinguish the number "3" from all possible numbers and to do so as accurately as possible. You will first explore your data (this should always be your starting point to gain domain knowledge about the problem.). Since the feature space in this problem is 784-dimensional, overfitting is possible. To avoid overfitting you will investigate the impact of regularization on generalization performance (test accuracy) and compare regularized and unregularized logistic regression model test error against other classification techniques such as linear discriminant analysis and random forests and draw conclusions about the best-performing model.

Start by loading your dataset from the MNIST dataset of handwritten digits, using the code provided below. MNIST has a training set of 60,000 examples, and a test set of 10,000 examples. The digits have been size-normalized and centered in a fixed-size image.

Your goal is to classify whether or not an example digit is a 3. Your binary classifier should predict y=1 if the digit is a 3, and y=0 otherwise. Create your dataset by transforming your labels into a binary format (3's are class 1, and all other digits are class 0).

- (a) Plot 10 examples of each class (i.e. class y=0, which are not 3's and class y=1 which are 3's), from the training dataset.
  - Note that the data are composed of samples of length 784. These represent  $28 \times 28$  images, but have been reshaped for storage convenience. To plot digit examples, you'll need to reshape the data to be  $28 \times 28$  (which can be done with numpy reshape ).
- **(b)** How many examples are present in each class? Show a plot of samples by class (bar plot). What fraction of samples are positive? What issues might this cause?
- (c) Identify the value of the regularization parameter that optimizes model performance on out-of-sample data. Using a logistic regression classifier, apply lasso regularization and retrain the model and evaluate its performance on the test set over a range of values on the regularization coefficient. You can implement this using the LogisticRegression module and activating the 'l1' penalty; the parameter C is the inverse of the regularization strength. Vary the value of C logarithmically from  $10^{-4}$  to  $10^4$  (and make your x-axes logarithmic in scale) and evaluate it at least 20 different values of C. As you vary the regularization coefficient, Plot the following four quantities (this should result in 4 separate plots)...
  - The number of model parameters that are estimated to be nonzero (in the logistic regression model, one attribute is coef\_, which gives you access to the model parameters for a trained model)
  - The cross entropy loss (which can be evaluated with the Scikit Learn log\_loss function)

- Area under the ROC curve (AUC)
- The  $F_1$ -score (assuming a threshold of 0.5 on the predicted confidence scores, that is, scores above 0.5 are predicted as Class 1, otherwise Class 0). Scikit Learn also has a f1\_score function which may be useful. -Which value of C seems best for this problem? Please select the closest power of 10. You will use this in the next part of this exercise.
- (d) Train and test a (1) logistic regression classifier with minimal regularization (using the Scikit Learn package, set penalty='l1', C=1e100 to approximate this), (2) a logistic regression classifier with the best value of the regularization parameter from the last section, (3) a Linear Discriminant Analysis (LDA) Classifier, and (4) a Random Forest (RF) classifier (using default parameters for the LDA and RF classifiers).
  - Compare your classifiers' performance using ROC and Precision Recall (PR) curves. For the ROC curves, all your curves should be plotted on the same set of axes so that you can directly compare them. Please do the same wih the PR curves.
  - Plot the line that represents randomly guessing the class (50% of the time a "3", 50% not a "3"). You SHOULD NOT actually create random guesses. Instead, you should think through the theory behind how ROC and PR curves work and plot the appropriate lines. It's a good practice to include these in ROC and PR curve plots as a reference point.
  - For PR curves, an excellent resource on how to correctly plot them can be found here
    (ignore the section on "non-linear interpolation between two points"). This describes
    how a random classifier is represented in PR curves and demonstrates that it should
    provide a lower bound on performance.
  - When training your logistic regression model, it's recommended that you use solver="liblinear"; otherwise, your results may not converge.
  - Describe the performance of the classifiers you compared. Did the regularization of the logistic regression model make much difference here? Which classifier you would select for application to unseen data.

```
In []: # Load the MNIST Data
from sklearn.datasets import fetch_openml
from sklearn.model_selection import train_test_split
import numpy as np
import matplotlib.pyplot as plt
import pickle

# Set this to True to download the data for the first time and False after t
# so that you just load the data locally instead
download_data = False

if download_data:
    # Load data from https://www.openml.org/d/554
    X, y = fetch_openml('mnist_784', return_X_y=True, as_frame=False)

# Adjust the labels to be '1' if y==3, and '0' otherwise
```

```
y[y!='3'] = 0
y[y=='3'] = 1
y = y.astype('int')

# Divide the data into a training and test split
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=1/7,

file = open('tmpdata', 'wb')
pickle.dump((X_train, X_test, y_train, y_test), file)
file.close()

else:
   file = open('tmpdata', 'rb')
   X_train, X_test, y_train, y_test = pickle.load(file)
   file.close()
```

#### **ANSWER**

```
In []: # a) plot 10 examples of each class from the training set

def plot_digits(data, title):
    fig, axes = plt.subplots(1, 10, figsize=(8, 5))
    for ax, image in zip(axes, data):
        ax.imshow(image.reshape(28, 28), cmap=plt.cm.gray_r, interpolation='
        ax.axis('off')
    plt.suptitle(title, size=20)
    plt.show()

# Get indices of class 0 and class 1 samples
    indices_class_0 = np.where(y_train == 0)[0][:10]
    indices_class_1 = np.where(y_train == 1)[0][:10]

plot_digits(X_train[indices_class_0], "Not 3's")
    plot_digits(X_train[indices_class_1], "3's")
```

# Not 3's

# 8107870Y07

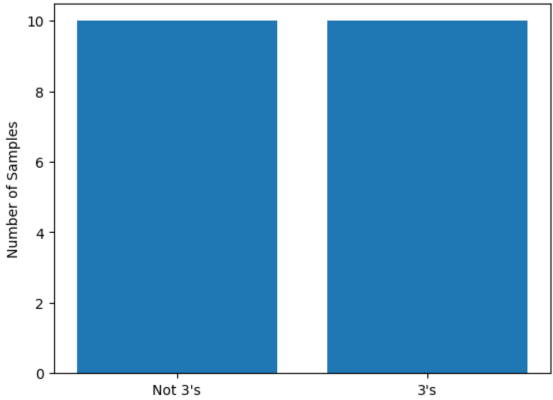
# 3 **3 3 3 3 3** 3 3 3 3

```
In []: # b) examples present in each class
plt.bar([0, 1], [len(indices_class_0), len(indices_class_1)])
plt.xticks([0, 1], ["Not 3's", "3's"])
plt.ylabel('Number of Samples')
plt.title('Samples by Class')
plt.show()

fraction_positive = len(indices_class_1) / (len(indices_class_0) + len(indices_class_1))
print(f"Fraction of samples that are positive: {fraction_positive:.2f}")

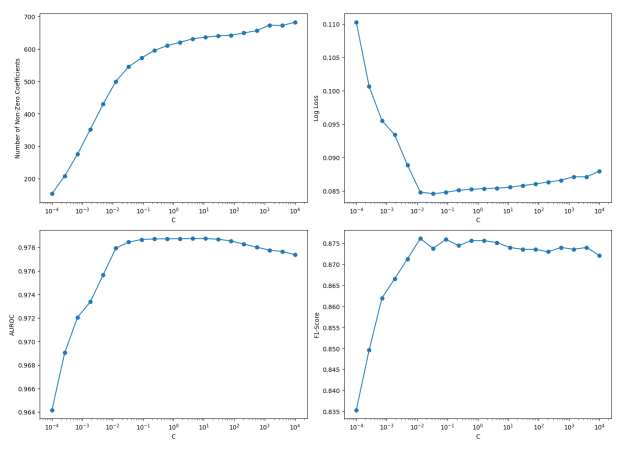
print('There seems to be a nice balance of classes, but in this case that me
```





Fraction of samples that are positive: 0.50 There seems to be a nice balance of classes, but in this case that means the dataset is half threes, half other values, which is sort of an imbalance, even if it is a binary class situation.

```
In [ ]: # c) optimize regularization parameter
        # Define values of C
        C values = np.logspace(-4, 4, 20)
        non zero coefs = [] # number of non-zero coefficients
        log losses = [] # log loss
        aurocs = [] # area under the ROC curve
        f1 scores = [] # F1 score
        for C in C values:
            lr = LogisticRegression(penalty='l1', C=C, solver="liblinear")
            lr.fit(X train, y train)
            y pred proba = lr.predict proba(X test)[:, 1]
            non zero coefs.append(np.sum(lr.coef != 0))
            log losses.append(log loss(y test, y pred proba))
            aurocs.append(roc auc_score(y_test, y_pred_proba))
            y pred = np.where(y pred proba > 0.5, 1, 0)
            f1 scores.append(f1 score(y test, y pred))
        # Plot the results
        fig, axarr = plt.subplots(2, 2, figsize=(14, 10))
        # Plot non-zero coefficients
        axarr[0, 0].semilogx(C values, non zero coefs, marker='o')
        axarr[0, 0].set xlabel('C')
        axarr[0, 0].set ylabel('Number of Non-Zero Coefficients')
        # Plot loa-loss
        axarr[0, 1].semilogx(C values, log losses, marker='o')
        axarr[0, 1].set xlabel('C')
        axarr[0, 1].set ylabel('Log Loss')
        # Plot AUROC
        axarr[1, 0].semilogx(C values, aurocs, marker='o')
        axarr[1, 0].set xlabel('C')
        axarr[1, 0].set ylabel('AUROC')
        # Plot F1-Score
        axarr[1, 1].semilogx(C_values, f1 scores, marker='o')
        axarr[1, 1].set xlabel('C')
        axarr[1, 1].set ylabel('F1-Score')
        plt.tight layout()
        plt.show()
        # Identify and print the best C
        best C = C values[np.argmax(aurocs)]
        print(f"The best value of C is approximately: {best C}")
```



The best value of C is approximately: 4.281332398719396

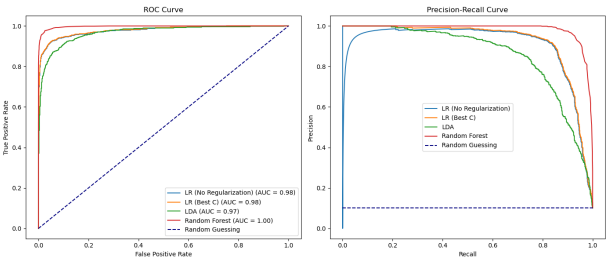
```
In [ ]: # d) train classifiers and compare
        # Define and train the classifiers
        classifiers = {
            "LR (No Regularization)": LogisticRegression(penalty='l1', C=1e100, solv
            "LR (Best C)": LogisticRegression(penalty='l1', C=best C, solver="libling"
            "LDA": LinearDiscriminantAnalysis(),
            "Random Forest": RandomForestClassifier()
        }
        plt.figure(figsize=(14, 6))
        for name, clf in classifiers.items():
            clf.fit(X train, y train)
            y_pred_proba = clf.predict_proba(X_test)[:, 1]
            # ROC curve
            fpr, tpr, = roc curve(y test, y pred proba)
            plt.subplot(1, 2, 1)
            plt.plot(fpr, tpr, label=f"{name} (AUC = {auc(fpr, tpr):.2f})")
            # PR curve
            precision, recall, _ = precision_recall_curve(y_test, y_pred_proba)
            plt.subplot(1, 2, 2)
            plt.plot(recall, precision, label=f"{name}")
        # ROC curve settings
        plt.subplot(1, 2, 1)
        plt.plot([0, 1], [0, 1], color='navy', linestyle='--', label="Random Guessin")
```

```
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('ROC Curve')
plt.legend()

# PR curve settings
plt.subplot(1, 2, 2)
no_skill = len(y_test[y_test==1]) / len(y_test) # calculate the no skill lin
plt.plot([0, 1], [no_skill, no_skill], color='navy', linestyle='--', label="
plt.xlabel('Recall')
plt.ylabel('Precision')
plt.title('Precision-Recall Curve')
plt.legend()

plt.tight_layout()
plt.show()

# describe and compare the classifier's performance
```



The graphs above show the results of training 4 different classifiers against the MNIST dataset. The first graph shows the ROC curves for each classifier, and the second graph shows the PR curves for each classifier. As can be seen, the LR models performed very similarly to each other. The LDA model was slightly les effective. Random Forest is shown here to be the most effective classifier, with the highest AUC and PR curve scores (hugging the corner of the graph). I would select the random forest classfier to take advantage of the strength of ensemble models.