Fall 2020 CS 4641\7641 A: Machine Learning Homework 4

Instructor: Dr. Mahdi Roozbahani

Deadline: November 18th, Wednesday, AOE

- No unapproved extension of the deadline is allowed. Late submission will lead to 0 credit.
- Discussion is encouraged on Piazza as part of the Q/A. However, all assignments should be done individually.

Instructions for the assignment

- In this assignment, we have programming and writing questions.
- To switch between cell for code and for markdown, see the menu -> Cell -> Cell Type
- You could directly type the Latex equations in the markdown cell.
- Typing with Latex\markdown is required for all the written questions. Handwritten answers would not be accepted.
- If a question requires a picture, you could use this syntax "< imgsrc ="" style =" width: 300px;" />" to include them within your ipython notebook.
- Questions marked with **[P]** are programming only and should be submitted to the autograder. Questions
 marked with **[W]** may required that you code a small function or generate plots, but should NOT be
 submitted to the autograder. It should be submitted on the writing portion of the assignment on gradescope
- The outline of the assignment is as follows:
 - Q1 [55+(10 bonus for undergrads)] > Neural Network **[W]** | **[P]**
 - Q2 [15 pts(bonus for all)] > Image Classification based on Convolutional Neural Network | **
 [W]**
 - Q3 [40 pts] > Random Forest **[P]** 3.1, 3.2 | **[W]** 3.3
 - Q4 [30 pts] > SVM **[W]**

Using the autograder

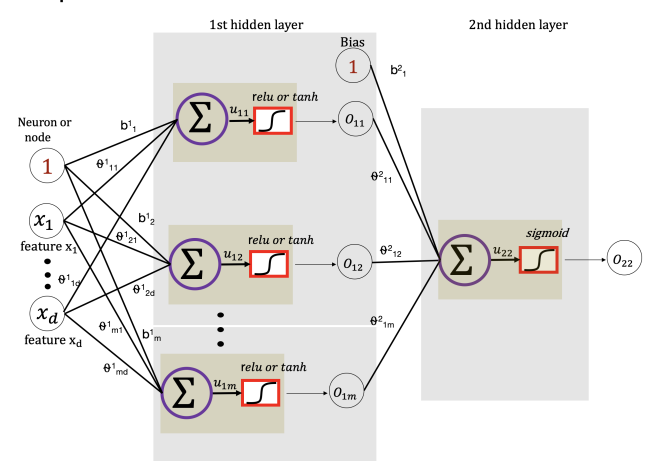
- You will find two assignments on Gradescope that correspond to HW4: "HW4 Programming" and "HW4 -Non-programming" (and "HW4 - Bonus Programming" if you are in CS4641).
- · You will submit your code for the autograder on "HW4 Programming" in the following format:
 - random forest.py
 - neural network.py
- You will submit your code for the autograder on "HW4 Bonus-Programming" in the following format:
 - neural network.py
- All you will have to do is to copy your implementations of the classes "dlnet" and "RandomForest" onto the
 respective files. We provided you different .py files and we added libraries in those files please DO NOT
 remove those lines and add your code after those lines. Note that these are the only allowed libraries that
 you can use for the homework.
- You are allowed to make as many submissions until the deadline as you like. Additionally, note that the
 autograder tests each function separately, therefore it can serve as a useful tool to help you debug your
 code if you are not sure of what part of your implementation might have an issue.
- For the "HW4 Non-programming" part, you will download your jupyter notbook as HTML, print it as a PDF from your browser and submit it on Gradescope. To download the notebook as html, click on "File" on the top left corner of this page and select "Download as > HTML". The non-programming part corresponds to Q1, Q2, Q3.3, Q4

Environment Setup

```
In [2]: import numpy as np
        import matplotlib.pyplot as plt
        from sklearn.datasets import load_breast_cancer
        from sklearn.preprocessing import MinMaxScaler
        from sklearn.model selection import train test split
        from sklearn.metrics import classification report
        from sklearn.metrics import plot confusion matrix
        from collections import Counter
        from scipy import stats
        from math import log2, sqrt
        import pandas as pd
        import time
        from sklearn.model selection import train test split
        from sklearn.preprocessing import LabelEncoder
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.datasets import make moons
        from sklearn.metrics import accuracy score
        from sklearn import svm
```

1. Two Layer Neural Network [65pts] **[P]**

Perceptron



*Notation clarification $-\theta^1_{21}$ – theta of 1st layer connecting 2nd hidden unit of 1st layer with 1st input unit θ^2_{12} - theta of 2nd layer connecting 1st unit of 2nd layer with 2nd hidden unit of 1st layer b1₁ – bias for 1st hidden unit of 1st layer

A single layer perceptron can be thought of as a linear hyperplane as in logistic regression followed by a non-linear activation function.

$$egin{aligned} u_i &= \sum_{j=1}^d heta_{ij} x_j + b_i \ o_i &= \phi\left(\sum_{j=1}^d heta_{ij} x_j + b_i
ight) = \phi(heta_i^T x + b_i) \end{aligned}$$

where x is a d-dimensional vector i.e. $x \in R^d$. It is one datapoint with d features. $\theta_i \in R^d$ is the weight vector for the i^{th} hidden unit, $b_i \in R$ is the bias element for the i^{th} hidden unit and $\phi(.)$ is a non-linear activation function that has been described below. u_i is a linear combination of the features in x weighted by θ_i whereas o_i is the i^{th} output unit from the activation layer.

Fully connected Layer

Typically, a modern neural network contains millions of perceptrons as the one shown in the previous image. Perceptrons interact in different configurations such as cascaded or parallel. In this part, we describe a fully connected layer configuration in a neural network which comprises multiple parallel perceptrons forming one layer.

We extend the previous notation to describe a fully connected layer. Each layer in a fully connected network has a number of input/hidden/output units cascaded in parallel. Let us a define a single layer of the neural net as follows:

m demotes the number of hidden units in a single layer l whereas n denotes the number of units in the previous layer l-1.

$$u^{[l]} = heta^{[l]} o^{[l-1]} + b^{[l]}$$

where $u^{[l]} \in R^m$ is a m-dimensional vector pertaining to the hidden units of the l^{th} layer of the neural network after applying linear operations. Similarly, $o^{[l-1]}$ is the n-dimensional output vector corresponding to the hidden units of the $(l-1)^{th}$ activation layer. $\theta^{[l]} \in R^{m \times n}$ is the weight matrix of the l^{th} layer where each row of $\theta^{[l]}$ is analogous to θ_i described in the previous section i.e. each row corresponds to one hidden unit of the l^{th} layer. $b^{[l]} \in R^m$ is the bias vector of the layer where each element of b pertains to one hidden unit of the l^{th} layer. This is followed by element wise non-linear activation function $o^{[l]} = \phi(u^{[l]})$. The whole operation can be summarized as,

$$o^{[l]} = \phi(heta^{[l]}o^{[l-1]} + b^{[l]})$$

where $o^{[l-1]}$ is the output of the previous layer.

Activation Function

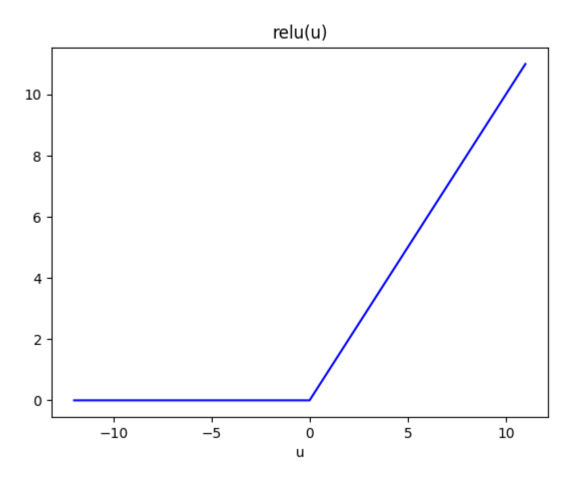
There are many activation functions in the literature but for this question we are going to use Relu, Sigmoid and Tanh only.

Relu

The rectified linear unit (Relu) is one of the most commonly used activation functions in deep learning models. The mathematical form is

$$o = \phi(u) = max(0, u)$$

The derivative of relu function is given as $o'=\phi'(u)=\left\{egin{array}{cc} 0 & u\leq 0 \ 1 & u>0 \end{array}
ight.$



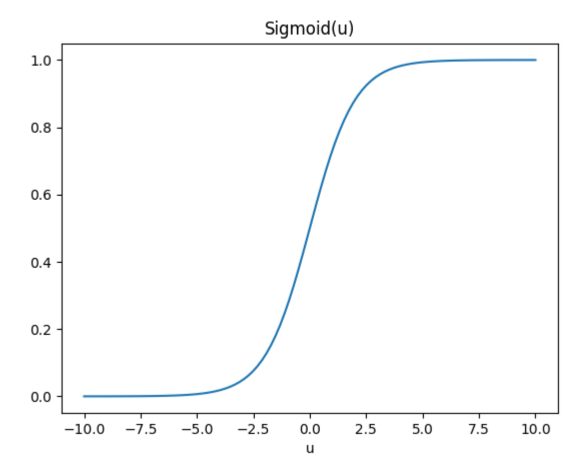
Sigmoid

The sigmoid function is another non-linear function with S-shaped curve. This function is useful in the case of binary classification as its output is between 0 and 1. The mathematical form of the function is

$$o=\phi(u)=\frac{1}{1+e^{-u}}$$

The derivation of the sigmoid function has a nice form and is given as

$$o' = \phi'(u) = rac{1}{1 + e^{-u}}igg(1 - rac{1}{1 + e^{-u}}igg) = \phi(u)(1 - \phi(u))$$



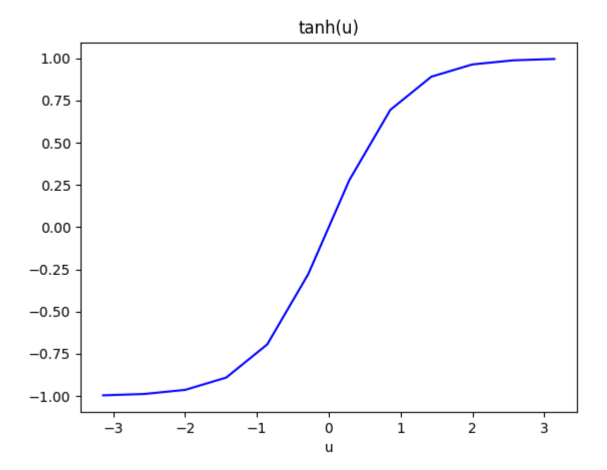
Tanh

Tanh also known as hyperbolic tangent is like a shifted version of sigmoid activation function with its range going from -1 to 1. Tanh almost always proves to be better than the sigmoid function since the mean of the activations are closer to zero. Tanh has an effect of centering data that makes learning for the next layer a bit easier. The mathematical form of tanh is given as

$$o=\phi(u)=tanh(u)=rac{e^u-e^{-u}}{e^u+e^{-u}}$$

The derivative of tanh is given as

$$o' = \phi'(u) = 1 - \left(rac{e^u - e^{-u}}{e^u + e^{-u}}
ight)^2 = 1 - o^2$$



Cross Entropy Loss

An essential piece in training a neural network is the loss function. The whole purpose of gradient descent algorithm is to find some network parameters that minimizes the loss function. In this exercise, we minimize Cross Entropy (CE) loss that represents on an intuitive level the distance between true data distribution and estimated distribution by neural network. So during training of the neural network, we will be looking for network parameters that minimizes the distance between true and estimated distribution. The mathematical form of the CE loss is given by

$$CE(p,q) = -\sum_i p(x_i) \log q(x_i)$$

where p(x) is the true distribution and q(x) is the estimated distribution.

Implementation details

For binary classification problems as in this exercise, we have probability distribution of a label
$$y_i$$
 given by
$$y_i = \left\{ \begin{array}{cc} 1 & \text{with probability } p(x_i) \\ 0 & \text{with probability } 1-p(x_i) \end{array} \right.$$

A frequentist estimate of $p(x_i)$ can be written as

$$p(x_i) = \sum_{i=1}^N rac{y_i}{N}$$

Therefore, the cross entropy for binary estimation can be written as

$$CE(y_i, \hat{y_i}) = -rac{1}{N} \sum_{i=1}^{N} \left(y_i \log(\hat{y_i}) + (1-y_i) \log(1-\hat{y_i})
ight)$$

where $y_i \in \{0,1\}$ is the true label and $\hat{y_i} \in [0,1]$ is the estimated label.

Forward Propagation

We start by initializing the weights of the fully connected layer using Xavier initialization <u>Xavier initialization</u> (http://proceedings.mlr.press/v9/glorot10a/glorot10a.pdf). During training, we pass all the data points through the network layer by layer using forward propagation. The main equations for forward prop have been described below.

$$egin{aligned} u^{[0]} &= x \ u^{[1]} &= heta^{[1]} u^{[0]} + b^{[1]} \ o^{[1]} &= Relu(u^{[1]}) \ u^{[2]} &= heta^{[2]} o^{[1]} + b^{[2]} \ \hat{y} &= o^{[2]} &= Sigmoid(u^{[2]}) \end{aligned}$$

Then we get the output and compute the loss

$$l = -rac{1}{N} \sum_{i=1}^{N} \left(y_i \log(\hat{y_i}) + (1-y_i) \log(1-\hat{y_i})
ight)$$

Backward propagation

After the forward pass, we do back propagation to update the weights and biases in the direction of the negative gradient of the loss function. So, we update the weights and biases using the following formulas

$$egin{align} heta^{[2]} &:= heta^{[2]} - lr imes rac{\partial l}{\partial heta^{[2]}} \ b^{[2]} &:= b^{[2]} - lr imes rac{\partial l}{\partial b^{[2]}} \ heta^{[1]} &:= heta^{[1]} - lr imes rac{\partial l}{\partial heta^{[1]}} \ b^{[1]} &:= b^{[1]} - lr imes rac{\partial l}{\partial b^{[1]}} \end{split}$$

where lr is the learning rate. It decides the step size we want to take in the direction of the negative gradient.

To compute the terms $\frac{\partial l}{\partial \theta^{[i]}}$ and $\frac{\partial l}{\partial b^{[i]}}$ we use chain rule for differentiation as follows:

$$egin{aligned} rac{\partial l}{\partial heta^{[2]}} &= rac{\partial l}{\partial o^{[2]}} rac{\partial o^{[2]}}{\partial u^{[2]}} rac{\partial u^{[2]}}{\partial heta^{[2]}} \ rac{\partial l}{\partial b^{[2]}} &= rac{\partial l}{\partial o^{[2]}} rac{\partial o^{[2]}}{\partial u^{[2]}} rac{\partial u^{[2]}}{\partial b^{[2]}} \end{aligned}$$

So, $rac{\partial l}{\partial o^{[2]}}$ is the differentiation of the cross entropy loss function at point $o^{[2]}$

 $rac{\partial o^{[2]}}{\partial u^{[2]}}$ is the differentiation of the Sigmoid function at point $u^{[2]}$

$$rac{\partial u^{[2]}}{\partial heta^{[2]}}$$
 is equal to $o^{[1]}$

$$\frac{\partial u^{[2]}}{\partial b^{[2]}}$$
 is equal to 1.

To compute $\frac{\partial l}{\partial \theta^{[2]}}$, we need $o^{[2]}, u^{[2]} \& o^{[1]}$ which are calculated during forward propagation. So we need to store these values in cache variables during forward propagation to be able to access them during backward propagation. Similarly for calculating other partial derivatives, we store the values we'll be needing for chain rule in cache. These values are obtained from the forward propagation and used in backward propagation. The cache is implemented as a dictionary here where the keys are the variable names and the values are the variables values.

Also, the functional form of the CE differentiation and Sigmoid differentiation are given by

$$egin{aligned} rac{\partial l}{\partial o^{[2]}} &= rac{-1}{N} igg(rac{y}{o^{[2]}} - rac{1-y}{1-o^{[2]}} igg) \ rac{\partial o^{[2]}}{\partial u^{[2]}} &= rac{1}{1+e^{-u^{[2]}}} igg(1 - rac{1}{1+e^{-u^{[2]}}} igg) &= o^{[2]} (1-o^{[2]}) \ rac{\partial u^{[2]}}{\partial heta^{[2]}} &= o^{[1]} \ rac{\partial u^{[2]}}{\partial b^{[2]}} &= 1 \end{aligned}$$

This completes the differentiation of loss function w.r.t to parameters in the second layer. We now move on to the first layer, the equations for which are given as follows:

$$egin{aligned} rac{\partial l}{\partial heta^{[1]}} &= rac{\partial l}{\partial o^{[2]}} rac{\partial o^{[2]}}{\partial u^{[2]}} rac{\partial u^{[2]}}{\partial o^{[1]}} rac{\partial o^{[1]}}{\partial u^{[1]}} rac{\partial u^{[1]}}{\partial heta^{[1]}} \ rac{\partial l}{\partial o^{[2]}} &= rac{\partial l}{\partial o^{[2]}} rac{\partial o^{[2]}}{\partial u^{[2]}} rac{\partial u^{[2]}}{\partial o^{[1]}} rac{\partial o^{[1]}}{\partial u^{[1]}} rac{\partial u^{[1]}}{\partial b^{[1]}} \end{aligned}$$

Where

$$egin{aligned} rac{\partial u^{[2]}}{\partial o^{[1]}} &= heta^{[2]} \ rac{\partial o^{[1]}}{\partial u^{[1]}} &= egin{cases} & 0 & ext{if } u^{[1]} \leq 0 \ & 1 & ext{if } u^{[1]} > 0 \end{cases} \ rac{\partial u^{[1]}}{\partial heta^{[1]}} &= x \ rac{\partial u^{[1]}}{\partial b^{[1]}} &= 1 \end{aligned}$$

Note that $rac{\partial o^{[1]}}{\partial u^{[1]}}$ is the differentiation of the Relu function at $u^{[1]}$.

The above equations outline the forward and backward propagation process for a 2-layer fully connected neural net with relu as the first activation layer and sigmoid has the second one. The same process can be extended to different neural networks with different activation layers like tanh.

Code Implementation:

$$egin{aligned} dLoss_o2 &= rac{\partial l}{\partial o^{[2]}} \implies dim = (1,426) \ dLoss_u2 &= dLoss_o2rac{\partial o^{[2]}}{\partial u^{[2]}} \implies dim = (1,426) \ dLoss_theta2 &= dLoss_u2rac{\partial u^{[2]}}{\partial heta^{[2]}} \implies dim = (1,15) \ dLoss_b2 &= dLoss_u2rac{\partial u^{[2]}}{\partial b^{[2]}} \implies dim = (1,1) \ dLoss_o1 &= dLoss_u2rac{\partial u^{[2]}}{\partial o^{[1]}} \implies dim = (15,426) \ dLoss_u1 &= dLoss_o1rac{\partial o^{[1]}}{\partial u^{[1]}} \implies dim = (15,426) \ dLoss_theta1 &= dLoss_u1rac{\partial u^{[1]}}{\partial a^{[1]}} \implies dim = (15,30) \end{aligned}$$

Question

In this question, you will implement a two layer fully connected neural network. You will also experiment with different activation functions and optimization techniques. Functions with comments "TODO: implement this" are for you to implement. We provide three activation functions here - Relu, Tanh and Sigmoid. You will implement a neural network that could have relu activation followed by sigmoid layer or tanh activation followed by sigmoid. You'll have to specify the neural net type which could be "Relu -> Sigmoid" (set by default) or "Tanh -> Sigmoid".

You'll also implement gradient descent and stochastic gradient descent algorithms for training these neural nets. SGD is bonus for undergraduate students.

We'll train these neural nets on breast cancer dataset. You're free to choose either gradient descent or SGD for training. Note: it is possible you'll run into nan or negative values for loss. This happens because of the small dataset we're using and some numerical stability issues that arise due to division by zero, natural log of zeros etc. You can experiment with the total number of iterations to mitigate this.

Deliverables for this question:

- Loss plot and classification report for any neural net type ("Relu -> Sigmoid" or "Tanh -> Sigmoid") with gradient descent
- 2. Loss plot and classification report for any neural net type ("Relu -> Sigmoid" or "Tanh -> Sigmoid") with stochastic gradient descent (mandatory for graduate students, bonus for undergraduate students)

```
In [108]:
          import numpy as np
          import matplotlib.pyplot as plt
          from sklearn.datasets import load breast cancer
          from sklearn.preprocessing import MinMaxScaler
          from sklearn.model selection import train test split
          from sklearn.metrics import classification report
          from sklearn.metrics import plot confusion matrix
           . . .
          We are going to use Breast Cancer Wisconsin (Diagnostic) Data Set provided by
           sklearn
          https://scikit-learn.org/stable/modules/generated/sklearn.datasets.load breast
           cancer.html
          to train a 2 fully connected layer neural net. We are going to buld the neural
          network from scratch.
          class dlnet:
              def __init__(self, x, y, lr = 0.003):
                  This method inializes the class, its implmented for you.
                  Args:
                      x: data
                      y: labels
                      Yh: predicted labels
                      dims: dimensions of different layers
                      param: dictionary of different layers parameters
                       ch: Cache dictionary to store forward parameters that are used in
           backpropagation
                       loss: list to store loss values
                       lr: learning rate
                       sam: number of training samples we have
                   . . .
                   self.X=x # features
                   self.Y=y # ground truth labels
                  self.Yh=np.zeros((1,self.Y.shape[1])) # estimated labels
                  self.dims = [30, 15, 1] # dimensions of different layers
                  self.param = { } # dictionary for different layer variables
                   self.ch = {} # cache for holding variables during forward propagation
           to use them in back prop
                  self.loss = []
                  self.lr=lr # learning rate
                   self.sam = self.Y.shape[1] # number of training samples we have
                   self. estimator type = 'classifier'
                   self.neural net type = "Relu -> Sigmoid" #can change it to "Tanh -> Si
          qmoid"
              def nInit(self):
```

```
This method initializes the neural network variables, its already impl
emented for you.
       Check it and relate to mathematical the description above.
        You are going to use these variables in forward and backward propagati
on.
        . . .
        np.random.seed(1)
        self.param['theta1'] = np.random.randn(self.dims[1], self.dims[0]) / n
p.sqrt(self.dims[0])
        self.param['b1'] = np.zeros((self.dims[1], 1))
        self.param['theta2'] = np.random.randn(self.dims[2], self.dims[1]) / n
p.sqrt(self.dims[1])
        self.param['b2'] = np.zeros((self.dims[2], 1))
   def Relu(self, u):
       In this method you are going to implement element wise Relu.
       Make sure that all operations here are element wise and can be applied
to an input of any dimension.
       Input: u of any dimension
       return: Relu(u)
       #TODO: implement this
       v = np.copy(u)
       v[u < 0] = 0
        return v
   def Sigmoid(self, u):
       In this method you are going to implement element wise Sigmoid.
       Make sure that all operations here are element wise and can be applied
to an input of any dimension.
        Input: u of any dimension
       return: Sigmoid(u)
       #TODO: implement this
        return 1/(1 + np.exp(-1 * u))
   def Tanh(self, u):
       In this method you are going to implement element wise Tanh.
       Make sure that all operations here are element wise and can be applied
to an input of any dimension.
       Input: u of any dimension
        return: Tanh(u)
       #TODO: implement this
        return np.tanh(u)
   def dRelu(self, u):
        This method implements element wise differentiation of Relu.
```

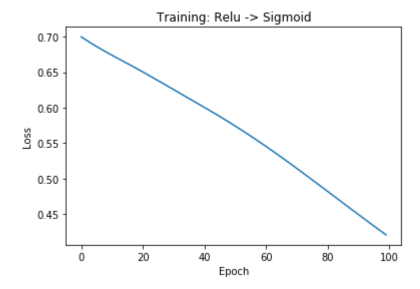
```
Input: u of any dimension
        return: dRelu(u)
        u[u <= 0] = 0
        u[u>0] = 1
        return u
    def dSigmoid(self, u):
        This method implements element wise differentiation of Sigmoid.
        Input: u of any dimension
        return: dSigmoid(u)
        o = 1/(1+np.exp(-u))
        do = o * (1-o)
        return do
    def dTanh(self, u):
        This method implements element wise differentiation of Tanh.
        Input: u of any dimension
        return: dTanh(u)
        . . .
        o = np.tanh(u)
        return 1-o**2
    def nloss(self,y, yh):
        In this method you are going to implement Cross Entropy loss.
        Refer to the description above and implement the appropriate mathemati
cal equation.
        Input: y 1xN: ground truth labels
               yh 1xN: neural network output after Sigmoid
        return: CE 1x1: loss value
        #TODO: implement this
        return np.mean(- (y * np.log(yh) + (1 - y) * np.log(1 - yh)))
    def forward(self, x):
        Fill in the missing code lines, please refer to the description for mo
re details.
        Check nInit method and use variables from there as well as other imple
meted methods.
        Refer to the description above and implement the appropriate mathemati
cal equations.
        do not change the lines followed by #keep.
          #Todo: uncomment the following 7 lines and complete the missing code
        #print(x.shape)
```

```
self.ch['X'] = x #keep
        #print(self.param['theta1'].shape, x.shape,self.param['b1'].shape)
        u1 = self.param['theta1'] @ x + self.param['b1']
       o1 = self.Relu(u1)
        self.ch['u1'], self.ch['o1']=u1, o1 #keep
       u2 = self.param['theta2'] @ o1 + self.param['b2']
       o2 = self.Sigmoid(u2)
        self.ch['u2'], self.ch['o2']=u2, o2 #keep
       return o2 #keep
   def backward(self, y, yh):
       Fill in the missing code lines, please refer to the description for mo
re details
        You will need to use cache variables, some of the implemeted methods,
and other variables as well
        Refer to the description above and implement the appropriate mathemati
cal equations.
        do not change the lines followed by #keep.
       #TODO: implement this
       dLoss_02 = -(np.divide(y, yh) - np.divide(1 - y, 1 - yh)) / y.shape[
11
   # partial l by partial o2
       #Implement equations for getting derivative of loss w.r.t u2, theta2 a
nd b2
       # set dLoss u2, dLoss theta2, dLoss b2
        #print('o2', dLoss o2.shape)
        dLoss_u2 = dLoss_o2 * self.dSigmoid(self.ch['u2'])
        #print('u2', dLoss u2.shape)
        dLoss theta2 = dLoss u2 @ self.ch['o1'].T
        #print('t2', dLoss theta2.shape)
        dLoss b2 = np.expand dims(np.sum(dLoss u2, 1), 1)
       #print('b2', dLoss b2.shape)
       dLoss o1 = np.dot(self.param["theta2"].T,dLoss u2) # partial L by part
ial o1
       #print('o1', dLoss o1.shape)
       #Implement equations for getting derivative of loss w.r.t u1, theta1 a
nd b1
       # set dLoss_u1, dLoss_theta1, dLoss_b1
        dLoss u1 = dLoss o1 * self.dRelu(self.ch['u1'])
        #print('u1', dLoss_u1.shape)
       dLoss theta1 = dLoss u1 @ self.ch['X'].T
       #print('t1', dLoss theta1.shape)
       dLoss b1 = np.expand dims(np.sum(dLoss u1, 1), 1)
       # print('b1', dLoss_b1.shape)
        #parameters update, no need to change these lines
        self.param["theta2"] = self.param["theta2"] - self.lr * dLoss_theta2 #
keep
        self.param["b2"] = self.param["b2"] - self.lr * dLoss b2 #keep
```

```
self.param["theta1"] = self.param["theta1"] - self.lr * dLoss theta1 #
keep
        self.param["b1"] = self.param["b1"] - self.lr * dLoss b1 #keep
        return dLoss theta2, dLoss b2, dLoss theta1, dLoss b1
   def gradient descent(self, x, y, iter = 60000):
        This function is an implementation of the gradient decent algorithm
       #Todo: implement this
        self.nInit()
        for _ in range(iter):
            yh = self.forward(x)
            self.backward(y, yh)
            self.loss.append(self.nloss(y, yh))
   #bonus for undergraduate students
   def stochastic gradient descent(self, x, y, iter = 60000):
        This function is an implementation of the stochastic gradient decent a
Lgorithm
       Note:
       1. SGD loops over all examples in the dataset one by one and learns a
gradient from each example
       2. One iteration here is one round of forward and backward propagation
on one example of the dataset.
           So if the dataset has 1000 examples, 1000 iterations will constitut
e an epoch
        3. Append loss after every 2000 iterations for plotting loss plots
       4. It is fine if you get a noisy plot since learning on one example at
a time adds variance to the
           gradients learnt
        5. You can use SGD with any neural net type
        ...
       self.nInit()
        _{,} N = x.shape
       for i in range(iter):
            xi = np.expand_dims(x[:, i % N], 1)
            yi = np.expand dims(y[:, i % N], 1)
            yh = self.forward(xi)
            self.backward(yi, yh)
            if i % 2000 == 0:
                vh = self.forward(x)
                self.loss.append(self.nloss(y, yh))
   def predict(self, x):
        This function predicts new data points
        Its implemented for you
       Yh = self.forward(x)
        return np.round(Yh).squeeze()
```

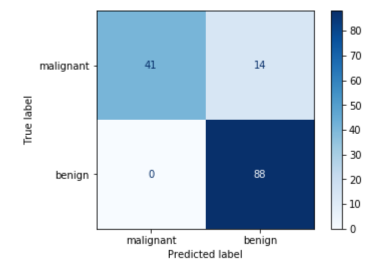
```
In [105]:
          Training the Neural Network, you do not need to modify this cell
          We are going to use Breast Cancer Wisconsin (Diagnostic) Data Set provided by
          https://scikit-learn.org/stable/modules/generated/sklearn.datasets.load breast
          _cancer.html
          dataset = load breast cancer() # load the dataset
          x, y = dataset.data, dataset.target
          x = MinMaxScaler().fit_transform(x) #normalize data
          x_train, x_test, y_train, y_test = train_test_split(x, y, random_state=1) #spl
          it data
          x_train, x_test, y_train, y_test = x_train.T, x_test.T, y_train.reshape(1,-1),
          y test #condition data
          nn = dlnet(x_train,y_train,lr=0.1) # initalize neural net class 66000
          nn.gradient_descent(x_train, y_train, iter = 100) #train
          # create figure
          fig = plt.plot(np.array(nn.loss).squeeze())
          plt.title(f'Training: {nn.neural net type}')
          plt.xlabel("Epoch")
          plt.ylabel("Loss")
```

Out[105]: Text(0, 0.5, 'Loss')



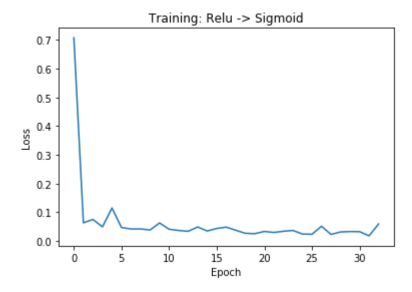
Classification Report for Relu -> Sigmoid

	precision	recall	f1-score	support
malignant	1.00	0.75	0.85	55
benign	0.86	1.00	0.93	88
accuracy			0.90	143
macro avg	0.93	0.87	0.89	143
weighted avg	0.92	0.90	0.90	143



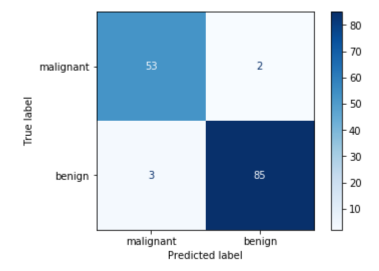
```
In [109]:
          Training the Neural Network, you do not need to modify this cell
          We are going to use Breast Cancer Wisconsin (Diagnostic) Data Set provided by
          https://scikit-learn.org/stable/modules/generated/sklearn.datasets.load breast
          _cancer.html
          dataset = load breast cancer() # Load the dataset
          x, y = dataset.data, dataset.target
          x = MinMaxScaler().fit_transform(x) #normalize data
          x_train, x_test, y_train, y_test = train_test_split(x, y, random_state=1) #spl
          it data
          x_train, x_test, y_train, y_test = x_train.T, x_test.T, y_train.reshape(1,-1),
          y test #condition data
          nn = dlnet(x_train,y_train,lr=0.1) # initalize neural net class
          nn.stochastic_gradient_descent(x_train, y_train, iter = 66000) #train
          # create figure
          fig = plt.plot(np.array(nn.loss).squeeze())
          plt.title(f'Training: {nn.neural net type}')
          plt.xlabel("Epoch")
          plt.ylabel("Loss")
```

Out[109]: Text(0, 0.5, 'Loss')



Classification Report for Relu -> Sigmoid

	precision	recall	f1-score	support
malignant	0.95	0.96	0.95	55
benign	0.98	0.97	0.97	88
accuracy			0.97	143
macro avg	0.96	0.96	0.96	143
weighted avg	0.97	0.97	0.97	143



2: (Bounus for all) Image Classification based on Convolutional Neural Networks [15pts] **[W]**

Keras is a deep learning API written in Python, running on top of the machine learning platform TensorFlow. It was developed with a focus on enabling fast experimentation. Being able to go from idea to result as fast as possible is key to doing good research. In this part, you will build a convolutional neural network based on Keras to solve the image classification task for MINIST. If you haven't installed TensorFlow, you can install the package by **pip** command or train your model by uploading HW4 notebook to Colab (https://colab.research.google.com/) directly. Colab contains all packages you need for this section.

Hint1: First contact with Keras (https://keras.io/about/)

Hint2: How to Install Keras (https://www.pyimagesearch.com/2016/07/18/installing-keras-for-deep-learning/)

Hint3: CS231n Tutorial (Layers used to build ConvNets) (https://cs231n.github.io/convolutional-networks/)

Environment Setup

```
In [260]: from __future__ import print_function
    import keras
    from keras.datasets import mnist
    from keras.models import Sequential
    from keras.layers import Dense, Dropout, Flatten
    from keras.layers import Conv2D, MaxPooling2D
    from keras import backend as K
    import matplotlib.pyplot as plt
```

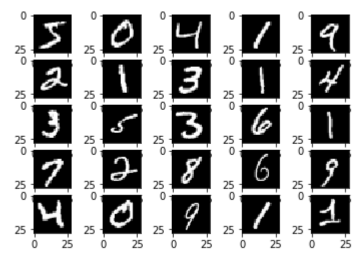
Load MINIST dataset

We use $\underline{\text{MINIST}}$ (http://yann.lecun.com/exdb/mnist/) dataset to train our model. MINIST is a subset of a larger set available from NIST. MNIST database of handwritten digits has a training set of 60,000 examples, and a test set of 10,000 examples. Each example is 28×28 pixel grayscale image of handwritten digits between 0 to 9.

```
In [261]: # Helper function, You don't need to modify it
          # split data between train and test sets
          (x_train, y_train), (x_test, y_test) = mnist.load_data()
          # input image dimensions
          img_rows, img_cols = 28, 28
          #set num of classes
          num classes = 10
          if K.image_data_format() == 'channels_first':
              x train = x train.reshape(x train.shape[0], 1, img rows, img cols)
              x_test = x_test.reshape(x_test.shape[0], 1, img_rows, img_cols)
              input_shape = (1, img_rows, img_cols)
          else:
              x train = x train.reshape(x train.shape[0], img rows, img cols, 1)
              x_test = x_test.reshape(x_test.shape[0], img_rows, img_cols, 1)
              input shape = (img rows, img cols, 1)
          x_train = x_train.astype('float32')
          x test = x test.astype('float32')
          x train /= 255
          x_test /= 255
          print('x_train shape:', x_train.shape)
          print('x_test shape:', x_test.shape)
          print(x_train.shape[0], 'train samples')
          print(x_test.shape[0], 'test samples')
          # convert class vectors to binary class matrices
          y_train = keras.utils.to_categorical(y_train, num_classes)
          y test = keras.utils.to categorical(y test, num classes)
          x train shape: (60000, 28, 28, 1)
          x_test shape: (10000, 28, 28, 1)
          60000 train samples
          10000 test samples
```

Load some images from MINIST

```
In [262]: # Helper function, You don't need to modify it
    # Show some images from MINIST
    for i in range(0,25):
        plt.subplot(5,5,i+1)
        image=x_train[i].reshape((28,28))
        plt.imshow(image,cmap='gray')
    plt.show()
```



As you can see from above, the MINIST dataset contains handwritten digits from 0 to 9. The digits have been size-normalized and centered in fixed-size images.

Build convolutional neural network model

In this part, you need to build a convolutional neural network that contains 2 convolutional layers. The architecture of thie model is:

[INPUT - CONV - RELU - MAXPOOL - CONV - RELU - MAXPOOL - FC1 - FC2] [1]

INPUT: $[28 \times 28 \times 1]$ will hold the raw pixel values of the image, in this case, an image of width 28, height 28, and with only one color channels.

CONV: Conv. layer will compute the output of neurons that are connected to local regions in the input, each computing a dot product between their weights and a small region they are connected to the input volume. We decide to set the kernel_size 3×3 for the both Conv. layers. For example, the output of the Conv. layer may like $[26 \times 26 \times 32]$ if we use 32 filters.

RELU: As we mentioned in the previous section, the Relu layer will apply an elementwise activation function, such as the max(0,x) thresholding at zero, which leaves the size of the volume unchanged.

MAXPOOL: MAXPOOL layer will perform a downsampling operation along the spatial dimensions (width, hight).

FC1: First Fully-Connected layer, we use **ReLu** as the activation function. The dimension of the output space is 128.

FC2: Second Fully-Connected layer will compute the class scores. We use **Softmax** as the activation function. The dimension of the output space is the number of class.

Loss function: Crossentropy (mentioned in the previous section)

optimizer: Stochastic gradient descent(SGD)

[1] CS231n: https://cs231n.github.io/convolutional-networks/ (https://cs231n.github.io/convolutional-networks/)

```
In [263]: # Helper function, You don't need to modify it
    # Show the architecture of the model
    achi=plt.imread('Architecture.png')
    fig = plt.figure(figsize=(10,10))
    plt.imshow(achi)
```

Out[263]: <matplotlib.image.AxesImage at 0x2285dca0358>

Mod	el: "sequential_1"			
Lay	er (type)	Output	Shape	Param #
0 - con	v2d_2 (Conv2D)	(None,	26, 26, 32)	320
max	_pooling2d_2 (MaxPooli	ng2 (None,	13, 13, 32)	0
con	v2d_3 (Conv2D)	(None,	11, 11, 64)	18496
тах	_pooling2d_3 (MaxPooli	ng2 (Nonc,	5, 5, 64)	0
	tten_1 (Flatten)	(None,	1600)	0
den	se_2 (Dense)	(None,	128)	204928
den	se_3 (Dense) ========	(None,	10)	1290
Tra	al params: 225,034 inable params: 225,034 -trainable params: 0	ı		
0	100 200	300	400	500

Defining Variables

```
In [264]: # Defining Variables
# Do not change the value of num_classes.
# You can adjust of adding parameters to train your model
batch_size = 128
epochs = 12
lr= 1e-3 #learning rate
```

Defining model

```
In [265]:
          def create net():
              In this function you are going to build a convolutional neural network bas
          ed on Keras.
              First, use Sequential() to set the inference features on this model.
              Then, use model.add() and model.compile() to build your own model
              Return: model you build
              model = Sequential()
              model.add(Conv2D(filters = 32, kernel_size = (3, 3), activation = 'relu'))
              model.add(MaxPooling2D(pool size = (2, 2)))
              model.add(Conv2D(filters = 64, kernel_size = (3, 3), activation = 'relu'))
              model.add(MaxPooling2D(pool_size = (2, 2)))
              model.add(Flatten())
              model.add(Dense(128, activation = 'relu'))
              model.add(Dense(10, activation = 'softmax'))
              model.compile(optimizer='sgd', loss='categorical crossentropy', metrics =
          ['accuracy'])
              model.build((None, 28, 28, 1))
              return model
```

In [266]: # Helper function, You don't need to modify it
 # model.summary() gives you details of your architecture.
#You can compare your architecture with the 'Architecture.png'
 model=create_net()
 model.summary()

Layer (type)	Output	Shape	Param #
conv2d_19 (Conv2D)	(None,	26, 26, 32)	320
max_pooling2d_18 (MaxPooling	(None,	13, 13, 32)	0
conv2d_20 (Conv2D)	(None,	11, 11, 64)	18496
max_pooling2d_19 (MaxPooling	(None,	5, 5, 64)	0
flatten_13 (Flatten)	(None,	1600)	0
dense_25 (Dense)	(None,	128)	204928
dense_26 (Dense)	(None,	10)	1290
Total params: 225,034 Trainable params: 225,034 Non-trainable params: 0	=====		======

Train the network

Tuning: Training the network is the next thing to try. You can set your parameter at the **Defining Variable** section. If your parameters are set properly, you should see the loss of the validation set decreased and the value of accuracy increased. It may take more than 20 minutes to train your model.

Expected Result: You should be able to achieve more than 90% accuracy on the test set to get full 15 points. If you achieve accuracy between 80% to 90%, you will only get half points of this part.

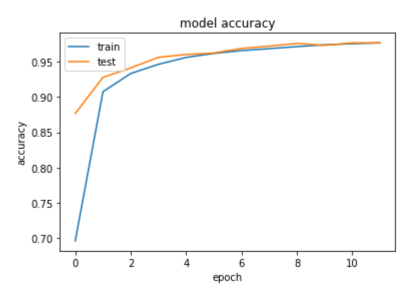
Train your own CNN model

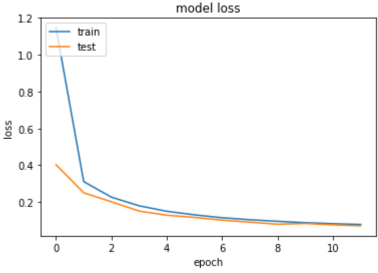
```
In [267]: # Helper function, You don't need to modify it
      # Train the model
      history = model.fit(x train, y train,
            batch size=batch size,
            epochs=epochs,
            verbose=1,
            validation data=(x test, y test))
      score = model.evaluate(x_test, y_test, verbose=0)
      print('Test loss:', score[0])
      print('Test accuracy:', score[1])
      Train on 60000 samples, validate on 10000 samples
      Epoch 1/12
      - acc: 0.6963 - val loss: 0.4031 - val acc: 0.8766
      Epoch 2/12
      - acc: 0.9074 - val_loss: 0.2507 - val_acc: 0.9277
      - acc: 0.9332 - val_loss: 0.2019 - val_acc: 0.9411
      60000/60000 [======================== ] - 47s 775us/step - loss: 0.1801
      - acc: 0.9462 - val loss: 0.1514 - val acc: 0.9562
      Epoch 5/12
      - acc: 0.9560 - val loss: 0.1288 - val acc: 0.9601
      Epoch 6/12
      - acc: 0.9617 - val_loss: 0.1169 - val_acc: 0.9622
      Epoch 7/12
      - acc: 0.9656 - val_loss: 0.1022 - val_acc: 0.9687
      Epoch 8/12
      - acc: 0.9684 - val loss: 0.0913 - val acc: 0.9720
      - acc: 0.9713 - val loss: 0.0799 - val acc: 0.9758
      Epoch 10/12
      - acc: 0.9738 - val loss: 0.0840 - val acc: 0.9731
      Epoch 11/12
      60000/60000 [============= ] - 41s 686us/step - loss: 0.0829
      - acc: 0.9755 - val loss: 0.0766 - val acc: 0.9766
      Epoch 12/12
      - acc: 0.9766 - val loss: 0.0713 - val acc: 0.9767
      Test loss: 0.07133965700771659
```

Test accuracy: 0.9767

```
In [268]:
          # Helper function, You don't need to modify it
          # list all data in history
          print(history.history.keys())
          # summarize history for accuracy
          plt.plot(history.history['acc'])
          plt.plot(history.history['val_acc'])
          plt.title('model accuracy')
          plt.ylabel('accuracy')
          plt.xlabel('epoch')
          plt.legend(['train', 'test'], loc='upper left')
          plt.show()
          # summary
          plt.plot(history.history['loss'])
          plt.plot(history.history['val_loss'])
          plt.title('model loss')
          plt.ylabel('loss')
          plt.xlabel('epoch')
          plt.legend(['train', 'test'], loc='upper left')
          plt.show()
```

dict_keys(['val_loss', 'val_acc', 'loss', 'acc'])





3: Random Forests [40pts] **[P]** **[W]**

NOTE: Please use sklearn's DecisionTreeClassifier in your Random Forest implementation. <u>You can find more details about this classifier here. (https://scikit-</u>

learn.org/stable/modules/generated/sklearn.tree.DecisionTreeClassifier.html#sklearn.tree.DecisionTreeClassifier_)

3.1 Random Forest Implementation (30 pts) **[P]**

The decision boundaries drawn by decision trees are very sharp, and fitting a decision tree of unbounded depth to a list of examples almost inevitably leads to **overfitting**. In an attempt to decrease the variance of a decision tree, we're going to use a technique called 'Bootstrap Aggregating' (often abbreviated 'bagging'). This stems from the idea that a collection of weak learners can learn decision boundaries as well as a strong learner. This is commonly called a Random Forest.

We can build a Random Forest as a collection of decision trees, as follows:

- 1) For every tree in the random forest, we're going to
 - a) Subsample the examples with replacement. Note that in this question, the size of the subsample data is equal to the original dataset.
 - b) From the subsamples in a), choose attributes at random to learn on in accordance with a provided attribute subsampling rate. Based on what it was mentioned in the c lass, we randomly pick features in each split. We use a more general approach here to make the programming part easier. Let's randomly pick some features (70% percent of features) and grow the tree based on the pre-determined randomly selected features. Therefore, there is no need to find random features in each split.
 - c) Fit a decision tree to the subsample of data we've chosen to a certain depth.

Classification for a random forest is then done by taking a majority vote of the classifications yielded by each tree in the forest after it classifies an example.

In RandomForest Class,

- 1. X is assumed to be a matrix with num_training rows and num_features columns where num_training is the number of total records and num_features is the number of features of each record.
- 2. y is assumed to be a vector of labels of length num training.

NOTE: Lookout for TODOs for the parts that needs to be implemented.

```
In [216]:
          import numpy as np
          import sklearn
          from sklearn.tree import DecisionTreeClassifier
          class RandomForest(object):
              def __init__(self, n_estimators=50, max_depth=None, max_features=0.7):
                  # helper function. You don't have to modify it
                  # Initialization done here
                  self.n estimators = n estimators
                   self.max_depth = max_depth
                  self.max features = max features
                   self.bootstraps row indices = []
                   self.feature indices = []
                   self.out of bag = []
                   self.decision trees = [sklearn.tree.DecisionTreeClassifier(max depth=m
          ax_depth, criterion='entropy') for i in range(n_estimators)]
              def _bootstrapping(self, num_training, num_features, random_seed = None):
                   TODO:
                   - Randomly select indices of size num training with replacement corres
          ponding to row locations of
                    selected samples in the original dataset.
                   - Randomly select indices without replacement corresponding the column
          locations of selected features in the original feature
                     list (num features denotes the total number of features in the trai
          ning set, max features denotes the percentage
                     of features that are used to fit each decision tree).
                  Reference: https://en.wikipedia.org/wiki/Bootstrapping (statistics)
                   - num training: an integer N representing the total number of training
          instances.
                   - num_features: an integer D representing the total number of feature
          5.
                  Returns:
                   - row idx: (N,) numpy array of row indices corresponding to the row lo
          cations of the selected samples in the original dataset.
                   - col_idx: 1-D array of column indices corresponding to the column loc
          ations of selected features in the original feature list.
                  Hint: Consider using np.random.choice.
                  np.random.seed(random seed)
                   row_idx = np.random.choice(num_training, num_training, replace = True)
                  col idx = np.random.choice(num features, int(num features * self.max f
          eatures), replace = False)
                  return (row idx, col idx)
              def bootstrapping(self, num training, num features):
                  Args:
                   - num training: an integer N representing the total number of training
          instances.
```

```
- num features: an integer D representing the total number of feature
s.
       Returns:
        - None
        # helper function. You don't have to modify it
        # Initializing the bootstap datasets for each tree
       for i in range(self.n estimators):
            total = set(list(range(num training)))
            row idx, col idx = self. bootstrapping(num training, num features)
            total = total - set(row idx)
            self.bootstraps row indices.append(row idx)
            self.feature indices.append(col idx)
            self.out of bag.append(total)
   def fit(self, X, y):
        TODO:
        Train decision trees using the bootstrapped datasets.
       Note that you need to use the row indices and column indices.
       Aras:
        -X: NxD numpy array, where N is number
           of instances and D is the dimensionality of each
           instance
        -y: Nx1 numpy array, the predicted labels
       Returns:
        - None
        self.bootstrapping(X.shape[0], X.shape[1])
        for row idx, col idx, tree in zip(self.bootstraps row indices, self.fe
ature indices, self.decision trees):
            sample X = X[row idx, :]
            sample X = sample X[:, col idx]
            sample_y = y[row_idx]
            tree.fit(sample X, sample y)
   def 00B score(self, X, y):
       # helper function. You don't have to modify it
       # This function computes the accuracy of the random forest model predi
cting y given x.
       accuracy = []
        for i in range(len(X)):
            predictions = []
            for t in range(self.n_estimators):
                if i in self.out of bag[t]:
                    predictions.append(self.decision_trees[t].predict(np.resha
pe(X[i][self.feature_indices[t]], (1,-1)))[0])
            if len(predictions) > 0:
                accuracy.append(np.sum(predictions == y[i]) / float(len(predic
tions)))
        return np.mean(accuracy)
```

3.2 Hyperparameter Tuning with a Random Forest (5pts) **[P]**

In machine learning, hyperparameters are parameters that are set before the learning process begins. The max_depth, num_estimators, or max_features variables from 3.1 are examples of different hyperparameters for a random forest model. In this section, you will tune your random forest model on an e-commerce dataset to achieve a high accuracy on a classifying revenue sessions (whether a customer will purchase a product) from user behavior.

Let's first review the dataset in a bit more detail.

Dataset Objective

Imagine that we are the founders of a new e-commerce company that uses machine learning to optimize the user experience. We are tasked with the responsibility of coming up with a method for determining the likelihood of a shopping session ending in a purchase being made. We will then use this information to adjust pricing and services to encourage more purchasing.

After much deliberation amongst the team, you come to a conclusion that we can use past online shopping data to predict the future occurrence of revenue sessions.

We will use our random forest algorithm from Q3.1 to predict if a shopping session ends in a purchase.

You can find more information on the dataset here (https://archive.ics.uci.edu/ml/datasets/Online+Shoppers+Purchasing+Intention+Dataset#).

Loading the dataset

The dataset that the company has collected has the following features:

- 1. Administrative : continuous variable
- 2. Administrative Duration: continuous variable
- 3. Informational: continuous variable
- 4. Informational Duration: continuous variable
- 5. ProductRelated: continuous variable
- 6. ProductRelated Duration: continuous variable
- 7. BounceRates: continuous variable
- 8. ExitRates: continuous variable
- 9. PageValues: continuous variable
- 10. SpecialDay: continuous variable
- 11. Month: categorical variable
- 12. OperatingSystems : continuous variable
- 13. Browser: continuous variable
- 14. Region : continuous variable
- 15. TrafficType: continuous variable
- 16. VisitorType: categorical variable
- 17. Weekend: continuous variable
- 18. Revenue: target variable ------> Your random forest model will try to predict this variable. A
 "True" value in this column means a shopper purchased an item given their user behavior described
 by features 1-17, while a "False" label indicates that a shopper did not purchase an item.

```
# Logic for Loading in datasets. DO NOT MODIFY anything in this block.
In [213]:
          from sklearn import preprocessing
          preprocessor = preprocessing.LabelEncoder()
          data train = pd.read csv("datasets/hw4 fall2020 data train.csv")
          data test = pd.read csv("datasets/hw4 fall2020 data test.csv")
          X_train = data_train.drop(columns = 'Revenue')
          y train = data train['Revenue']
          X test = np.array(data test.drop(columns = 'Revenue'))
          y_test = np.array(data_test['Revenue'])
          X_train, y_train, X_test, y_test = np.array(X_train), np.array(y_train), np.ar
          ray(X test), np.array(y test)
          #The following lines of code converts columns holding categorical or boolean v
          ariables into integers.
          X train[:,10] = preprocessor.fit transform(X train[:,10])
          X test[:,10] = preprocessor.fit transform(X test[:,10])
          X train[:,-2] = preprocessor.fit transform(X train[:,-2])
          X_test[:,-2] = preprocessor.fit_transform(X_test[:,-2])
```

In the following codeblock, train your random forest model with different values for max_depth, n_estimators, or max_features and evaluate each model on the held-out test set. Try to choose a combination of hyperparameters that maximizes your prediction accuracy on the test set (aim for 92%+). Once you are satisfied with your chosen parameters, change the default values for max_depth, n_estimators, and max_features in the init function of your RandomForest class in random_forest.py to your chosen values, and then submit this file to Gradescope. You must achieve at least a 92% accuracy against a hidden test set (this will NOT the same as the test set provided here) in Gradescope to receive full credit for this section.

```
In [221]:
          TODO:
          n estimators defines how many decision trees are fitted for the random forest.
          max depth defines a stop condition when the tree reaches to a certain depth.
          max features controls the percentage of features that are used to fit each dec
          ision tree.
          Tune these three parameters to achieve a better accuracy. While you can use th
          e provided test set to
          evaluate your implementation, you will need to obtain 92% on a hidden test set
          to receive full credit
          for this section.
          import sklearn.ensemble
          n estimators = 10 #Hint: Consider values between 5-12.
          max depth = 12 # Hint: Consider values betweeen 3-12
          max features = 0.9 # Hint: Consider values betweeen 0.7-1.0.
          random_forest = RandomForest(n_estimators, max_depth, max_features)
          random_forest.fit(X_train, y_train)
          accuracy=random forest.OOB score(X test, y test)
          print("accuracy: %.4f" % accuracy)
```

accuracy: 0.9245

3.3 Plotting Feature Importance (5pts) **[W]**

While building tree-based models, it's common to quantify how well splitting on a particular feature in a decision tree helps with predicting the target label in a dataset. Machine learning practicioners typically use "Gini importance", or the (normalized) total reduction in entropy brought by that feature to evaluate how important that feature is for predicting the target variable.

Gini importance is typically calculated as the reduction in entropy from reaching a split in a decision tree weighted by the probability of reaching that split in the decision tree. Sklearn internally computes the probability for reaching a split by finding the total number of samples that reaches it during the training phase divided by the total number of samples in the dataset. This weighted value is our feature importance.

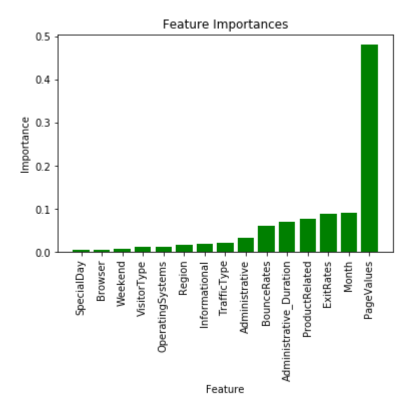
Let's think about what this metric means with an example. A high probability of reaching a split on "TrafficType" in a decision tree trained on our e-commerce dataset (many samples will reach this split for a decision) and a large reduction in entropy from splitting on "TrafficType" will result in a high feature importance value for "TrafficType". This could mean "TrafficType" is a very important feature for predicting a customer's revenue session. On the other hand, a low probability of reaching a split on "Month" in a decision tree (few samples will reach this split for a decision) and a low reduction in entropy from splitting on "Month" will result in a low feature importance value. This could mean "Month" is not a very informative feature for predicting the revenue session in our decision tree. Thus, the higher the feature importance value, the more important the feature is to predicting the target label.

Fortunately for us, fitting a sklearn. Decision Tree Classifier to a dataset auomatically computes the Gini importance for every feature in the decision tree and stores these values in a **feature** importances variable.

Review the docs for more details on how to access this variable (https://scikit-learn.org/stable/modules/generated/sklearn.tree.DecisionTreeClassifier.html#sklearn.tree.DecisionTreeClassifier.fe

In the function below, display a bar plot that shows the feature importance values for at least one decision tree in your tuned random forest from Q3.2, and briefly comment on whether any features have noticeably higher / or lower importance weights than others. [Note that there isn't a "correct" answer here. We simply want you to investigate how different features in your random forest contribute to predicting the target variable].

```
In [234]:
          import matplotlib.pyplot as plt
          def plot feature importance(random forest):
              TODO:
              -Display a bar plot showing the feature importance of every feature in
               at least one decision tree from the tuned random forest from Q3.2.
              Args:
                  random_forest: This is your implemented and tuned random forest from Q
          3.2.
              Returns:
                  None. Calling this function should simply display the aforementioned f
          eature importance bar chart
              fi = random_forest.decision_trees[0].feature_importances_
              labels = list(data train)
              labels = [labels[i] for i in random_forest.feature_indices[0]]
              fi, labels = zip(*sorted(zip(fi, labels)))
              x_pos = [i for i, _ in enumerate(labels)]
              plt.bar(x pos, fi, color='green')
              plt.xlabel("Feature")
              plt.ylabel("Importance")
              plt.title("Feature Importances")
              plt.xticks(x_pos, labels, rotation='vertical')
              plt.show()
          plot_feature_importance(random_forest)
```



4: SVM (30 Pts) **[W]**

4.1 Fitting an SVM classifier by hand (20 Pts) **[W]**

Consider a dataset with 2 points in 1-dimensional space: $(x_1 = -3, y_1 = -1)$ and $(x_2 = 2, y_2 = 1)$. Here x are the point coordinates and y are the classes.

Consider mapping each point to 3-dimensional space using the feature vector $\phi(x)=[1,2x,x^2]$. (This is equivalent to using a second order polynomial kernel.) The max margin classifier has the form

$$egin{aligned} min|| heta||^2s.\,t.\ y_1(\phi(x_1) heta+b) &\geq 1\ y_2(\phi(x_2) heta+b) &\geq 1 \end{aligned}$$

Hint: $\phi(x_1)$ and $\phi(x_2)$ are the suppport vectors. We have already given you the solution for the suppport vectors and you need to calculate back the parameters. Margin is equal to $\frac{1}{||\theta||}$ and full margin is equal to $\frac{2}{||\theta||}$.

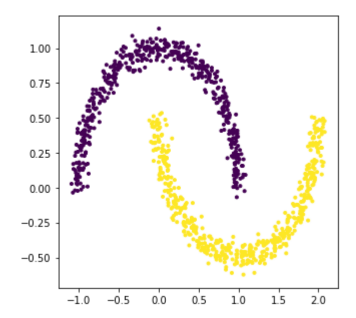
- (1) Find a vector parallel to the optimal vector θ . (4pts)
- (2) Calculate the value of the margin achieved by this θ ? (4pts)
- (3) Solve for θ , given that the margin is equal to $1/||\theta||$. (4pts)
- (4) Solve for b using your value for θ . (4pts)
- (5) Write down the form of the discriminant function $f(x) = \phi(x)\theta + b$ as an explicit function of x.

- 1) Note that both $\phi(x_1)$ and $\phi(x_2)$ are support vectors. They are the only two support vectors. Therefore, the decision boundary plane is there perpendicular bisector. This means that the normal to the decision plane is the vector connecting these two points. $\phi(x_1)=(1,-6,9)$ and $\phi(x_2)=(1,4,4)$. Therefore, a vector parallel to θ is (0,10,-5).
- 2) The margin is half the distance between $\phi(x_1)$ and $\phi(x_2)$. The distance is $\sqrt{10^2+5^2}=5\sqrt{5}=11.180$. Thus the margin is $5\sqrt{5}/2=5.590$.
- 3) Let our current vector v=(0,10,-5) . We want to multiply it by a scalar c such that $\theta=cv$. Thus we need, $\frac{5\sqrt{5}}{2}=\frac{1}{c||v||}$. We now that $||v||=5\sqrt{5}$. Therefore, $\frac{125}{2}=\frac{1}{c}$. Therefore, c = 2/125 . We have $\theta=(0,0.16,-0.08)$.
- 4) We plug into the constraint. The dot product is $0.16 \cdot 4 0.08 \cdot 4 = 0.32$. Thus we have 0.32 + b = 1. Therefore, b = 0.68.
- 5) We have $f(x) = 0.32x 0.08x^2 + 0.68$.

4.2 Feature Mapping (10 Pts) **[W]**

Let's look at a dataset where the datapoint can't be classified with a good accuracy using a linear classifier. Run the below cell to generate the dataset.

We will also see what happens when we try to fit a linear classifier to the dataset.



```
In [236]:
          def visualize decision boundary(X, y, feature new=None, h=0.02):
              You don't have to modify this function
              Function to vizualize decision boundary
              feature new is a function to get X with additional features
              x1_{min}, x1_{max} = X[:, 0].min() - 1, X[:, 0].max() + 1
              x2_{min}, x2_{max} = X[:, 1].min() - 1, X[:, 1].max() + 1
              xx_1, xx_2 = np.meshgrid(np.arange(x1_min, x1_max, h),
                                    np.arange(x2_min, x2_max, h))
              if X.shape[1] == 2:
                   Z = svm_cls.predict(np.c_[xx_1.ravel(), xx_2.ravel()])
              else:
                   X_{conc} = np.c_[xx_1.ravel(), xx_2.ravel()]
                   X_new = feature_new(X_conc)
                   Z = svm_cls.predict(X_new)
              Z = Z.reshape(xx 1.shape)
              f, ax = plt.subplots(nrows=1, ncols=1, figsize=(5,5))
              plt.contourf(xx_1, xx_2, Z, cmap=plt.cm.coolwarm, alpha=0.8)
              plt.scatter(X[:, 0], X[:, 1], c=y, cmap=plt.cm.coolwarm)
              plt.xlabel('X 1')
              plt.ylabel('X 2')
              plt.xlim(xx_1.min(), xx_1.max())
              plt.ylim(xx 2.min(), xx 2.max())
              plt.xticks(())
              plt.yticks(())
              plt.show()
```

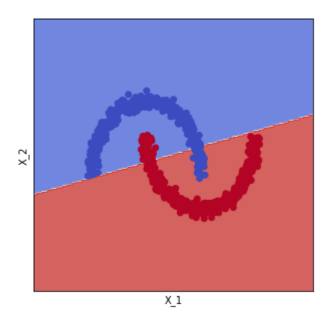
```
In [237]: # DO NOT CHANGE
# Try to fit a linear classifier to the dataset

svm_cls = svm.LinearSVC()
svm_cls.fit(X_train, y_train)
y_test_predicted = svm_cls.predict(X_test)

print("Accuracy on test dataset: {}".format(accuracy_score(y_test, y_test_predicted)))

visualize_decision_boundary(X_train, y_train)
```

Accuracy on test dataset: 0.88



We can see that we need a non-linear boundary to be able to successfully classify data in this dataset. By mapping the current feature x to a higher space with more features, linear SVM could be performed on the features in the higher space to learn a non-linear decision boundary. In the function below add additional features which can help classify in the above dataset. After creating the additional features use code in the further cells to see how well the features perform on the test set.

(**Hint:** Think of the shape of the decision boundary that would best separate the above points. What additional features could help map the linear boundary to the non-linear one? Look at this

(https://xavierbourretsicotte.github.io/Kernel_feature_map.html) for a detailed analysis of doing the same for points separable with a circular boundary)

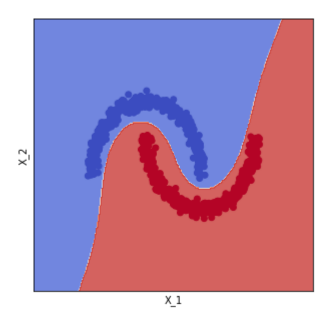
```
In [254]: # DO NOT CHANGE
# Fit to the new features and vizualize the decision boundary
# You should get more than 90% accuracy on test set

svm_cls = svm.LinearSVC()
svm_cls.fit(X_train, y_train)
y_test_predicted = svm_cls.predict(X_test)

print("Accuracy on test dataset: {}".format(accuracy_score(y_test, y_test_predicted)))

visualize_decision_boundary(X_train, y_train, create_nl_feature)
```

Accuracy on test dataset: 1.0



In []: