Logistic Regression

Introduction

In this notebook we are going to study gradient descent and in this case applied to logistic regression. Logistic regression (LR) is a statistical method for analysing datasets where there are one of more independent variables that determine the outcome. The outcome is a dichotomous, meaning there are only two possible outcomes (1 / 0, Yes / No, True / False). For instance, if you want to predict the sex of a person from age (x_1) and income (x_2) , the logistic regression model would be

$$h(x) = \theta_0 + \theta_1 x_1 + \theta_2 x_2$$

where h(x) is the outcome varibale, θ_0 the bias and θ_1 and θ_2 the weights. The goal is ultimately to tune these parameters with respect to the obeserved data (x_1, x_2) .

LR estimates a probability (between 0 and 100%) but h(x) gives values in $(-\infty, +\infty)$. We need to "squish" h(x) to restrict it to a suitable range. LR commonly uses the logistic function (a.k.a. sigmoid function) to compute probabilities:

$$\sigma(h(x))=rac{1}{1+e^{-h(x)}}.$$

It is possible to threshold the logistic function (values between 0-1), and values below 0.5 will be counted as the prediction of class 0 and values larger than 0.5 results in the prediction of class 1.

The full logistic regression model is then:

$$z(x)=\sigma(h(x))=rac{1}{1+e^{-(heta_0+ heta_1x_1+ heta_2x_2)}}.$$

Ready, steady, code! 🔊

Let's start with loading some data, scikit-learn comes with a couple of toy datasets and we are going to use the "iris" dataset where the goal is to classify which type of flower based on a set of features consisting of sepal length (cm), sepal width (cm), petal length (cm), petal width (cm). To begin with we consider only two of those features.

```
In [1]: # import stuff that we need
import numpy as np
import matplotlib as mpl
import matplotlib.pyplot as plt
import sklearn
```

```
import sklearn.datasets as ds
        from IPython.display import clear_output # if you get problem with this import you
        """ assert np.__version__ == "1.19.4", "Looks like you don't have the same version
In [2]:
        assert mpl.__version__ == "3.3.3", "Looks like you don't have the same version of m
        assert sklearn.__version__ == "0.24.0", "Looks like you don't have the same version
Out[2]: 'assert np.__version__ == "1.19.4", "Looks like you don\'t have the same version
        of numpy as us!"\nassert mpl.__version__ == "3.3.3", "Looks like you don\'t have t
        he same version of matplotlib as us!"\nassert sklearn. version == "0.24.0", "Lo
        oks like you don\'t have the same version of sklearn as us!" '
In [3]: data = ds.load_iris()
        selected_features_idx = [0,1] #'sepal length (cm)', 'sepal width (cm)'
        selected_targets = [0,1] #'setosa' 'versicolor'
        idx = np.array([x in selected_targets for x in data.target])
        x = data.data[:,selected_features_idx][idx]
        y = data.target[idx]
        y[y > 1] = 1 # Reset labels greater than 1 to 1
In [4]: plt.figure()
        for label in np.unique(y):
            plt.scatter(x[:, 0][y == label], x[:, 1][y == label], label = data.target_names
        plt.legend()
        plt.show()
         4.5
                                                                        setosa
                                                                        versicolor
         4.0
         3.5
         3.0
         2.5
```

Now we need a function that predicts the logistic regression model and make predictions. This function takes a measurement, the current bias and the weights as input.

6.0

6.5

7.0

5.5

2.0

4.5

5.0

$$z(x)=rac{1}{1+e^{-(heta_0+ heta_1x_1+ heta_2x_2)}}$$

Now try it with some random weights and bias.

```
In [6]: bias = np.random.normal()
weights = np.random.normal(size = len(x[0]))

predicted = []
for i in range(len(x)):
    yhat = z_x(x[i], bias, weights)
    predicted.append(round(yhat))

print('Accuracy: ', np.sum(np.equal(y, predicted)) / len(predicted))
```

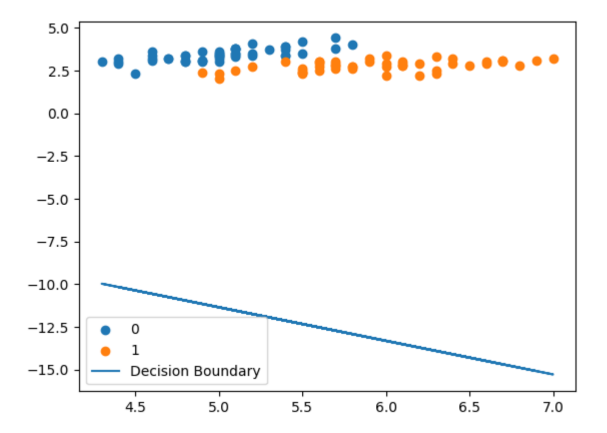
Accuracy: 0.5

Let's plot the decision boundary between the points for this set of weights. The decision boundary is found by setting h(x) = 0 which gives:

$$x_2 = -rac{ heta_0 + heta_1 x_1}{ heta_2}$$

```
In [7]: x_values = x[:,0]
y_values = - (bias + weights[0]*x_values) / weights[1]

plt.figure()
plt.scatter(x[:,0][y==0],x[:,1][y==0], label = '0')
plt.scatter(x[:,0][y==1],x[:,1][y==1], label = '1')
plt.plot(x_values, y_values, label='Decision Boundary')
plt.legend()
plt.show()
```



Not very good (or did you get lucky with the weights?). Try rerunning it a couple of times to see if you can randomly find a better set of weights that improves the accuracy.

Now, a better way of finding the optimal weights is the gradient descent method. Gradient descent is an iterative process of minimizing a function by following the gradients of a predefined cost function. This is useful for updating and tuning the parameters of our logistic regression model. As gradient descent is an iterative algorithm, we have to repeat the above step until we reach a satisfactory solution. The updates are defined as:

$$\theta_j \leftarrow \theta_j - \alpha \frac{\partial J(\theta)}{\partial \theta_j},$$

and similarily for the bias term

$$heta_0 \leftarrow heta_0 - lpha rac{\partial J(heta)}{\partial heta_o}.$$

Where α is a user specified learning rate, a scalar that controls the step size in the parameter space and $J(\theta)$ is the cost function that we will now define. Note that to minimize the cost function, we move in the direction opposite to the gradient.

First, we need to define our cost function, which is typically the negative log-likelihood of the data for numerical reasons, also called *Binary-Cross-Entropy* loss function. For a binary classification problem with m training examples, where $x^{(i)}$ represents the i-th example of our training set and $y^{(i)}$ its output, the cost function is given by:

$$J(heta) = rac{1}{m} \sum_{i=1}^m egin{cases} -log(g(x^{(i)}, heta)), & ext{if } y^{(i)} = 1 \ -log(1 - g(x^{(i)}, heta)), & ext{if } y^{(i)} = 0 \end{cases}$$

where $g(x^{(i)}, \theta)$ modelizes $P(y = 1|x; \theta)$. Note that we have added θ explicitly to the notation to emphasize the dependence on the model parameters.

The two functions can be combined into one as:

$$J(heta) = -rac{1}{m} \sum_{i=1}^m [y^{(i)} \log(z_ heta(x^{(i)})) + (1-y^{(i)}) \log(1-z_ heta(x^{(i)}))]$$

Where $z_{\theta}(x)$ is the sigmoid function, representing the probability that the input x belongs to the positive class.

To simplify the notation in the following calculations, we will omit the subindexes; nevertheless, keep in mind that when the cost function is optimized, it is done across all samples in the training dataset.

$$J(\theta) = -y \cdot log(z(x)) - (1-y) \cdot log(1-z(x))$$

where y is the target class. The Binary-Cross-Entropy tells us that if the target is 1 and we predict 0, then we will get a large error $(-log(0)=\infty)$ and vice verca ($-log(1-1)=-log(0)=\infty$).

For gradient descent we need the derivative of this cost function with respect to the weights $\frac{\partial J(\theta)}{\partial \theta}$. We can get this with the chain rule:

$$rac{\partial J(heta)}{\partial heta_i} = rac{\partial J(heta)}{\partial z(x)} \cdot rac{\partial z(x)}{\partial h(x)} \cdot rac{\partial h(x)}{\partial heta_i}$$

Where the three derivatives result in:

$$egin{aligned} rac{\partial J(heta)}{\partial z(x)} &= -\left(rac{y}{z(x)} - rac{(1-y)}{(1-z(x))}
ight) \ rac{\partial z(x)}{\partial h(x)} &= z(x)\cdot (1-z(x)) \ rac{\partial h(x)}{\partial heta} &= x \end{aligned}$$

Combining the previous equations together with the previous chain rule gives

$$rac{\partial J(heta)}{\partial heta_j} = x_j \cdot (z(x) - y)$$

where x_i is the j-th component of x.

For the bias term the derivative is similar but it is not dependent on x since $\frac{\partial h(x)}{\partial \theta_0}=1$

$$rac{\partial J(heta)}{\partial heta_0} = z(x) - y$$

Finally, recovering the subindex notation for all the samples in our training set we can express the gradient of the cost function with respect to each parameter. For the weight case:

$$rac{\partial J(heta)}{\partial heta_j} = rac{1}{m} \sum_{i=1}^m (z_ heta(x^{(i)}) - y^{(i)}) x_j^{(i)}.$$

And for the bias case:

$$rac{\partial J(heta)}{\partial heta_0} = rac{1}{m} \sum_{i=1}^m (z_ heta(x^{(i)}) - y^{(i)}).$$

With this two formulas now we can use these gradients to update the parameters in the gradient descent algorithm.

The full algorithm is:

- 1. Initialize the weights randomly.
- 2. Calculate the gradients of cost function w.r.t parameters.
- 3. Update the weights by $heta_j \leftarrow heta_j lpha rac{\partial}{\partial heta_i} J(heta)$.
- 4. Update the bias by $\theta_0 \leftarrow \theta_0 \alpha \frac{\partial}{\partial \theta_0} J(\theta)$.
- 5. Repeat until value of cost function does not change or to a pre-defined number of iterations.

Write a function for the cost and one for its derivative with respect to the weights and one with respect to the bias. Note that the derivative function will return the number of values corresponding the the number of weights that you have. Also note that we are only doing this for one training point.

```
In [8]:

def cost_function(y, x, bias, weights):
    """ param y: Ground truth label for measurements
    param x: vector containing measurements. x = [x1, x2]
    param bias: single value
    param weight: vector containing model weights. weights= [w1,w2]

    return: value of the cost function. In this case BCE
    """

z = z_x(x,bias,weights)
    cost = np.sum(-y*np.log(z) - (1-y)*np.log(1-z))/len(y)
    return cost
```

```
In [9]: def derivative_weights(y, x, bias, weights):
    """ param y: Ground truth label for measurements
    param x: vector containing measurements. x = [x1, x2]
    param bias: single value
    param weight: vector containing model weights. weights= [w1,w2]
```

```
return: derivative of cost function with respect to the weights, dw = [dw1, """] weights_deriv = x.T.dot(z_x(x, bias, weights)-y)/len(y) return weights_deriv
```

```
def derivative_bias(y, x, bias, weights):
    """ param y: Ground truth label for measurements
        param x: vector containing measurements. x = [x1, x2]
        param bias: single value
        param weight: vector containing model weights. weights= [w1,w2]

        return: derivative of cost function with respect to the bias
    """

bias_deriv = np.sum(z_x(x, bias, weights) - y)/len(y)
    return bias_deriv
```

Finally lets fit the logistic regression model with gradient descent across all training data points. As we saw before, gradient descent works by, at each iteration, average the total cost and the derivatives on over the full training set.

Implement gradient descent for logistic regression. Experiment with different learning rates and number of iterations to see if you get differnt solutions.

In [18]: predicted = []
for i in range(len(x)):
 yhat = z_x(x[i], bias, weights)
 predicted.append(round(yhat))

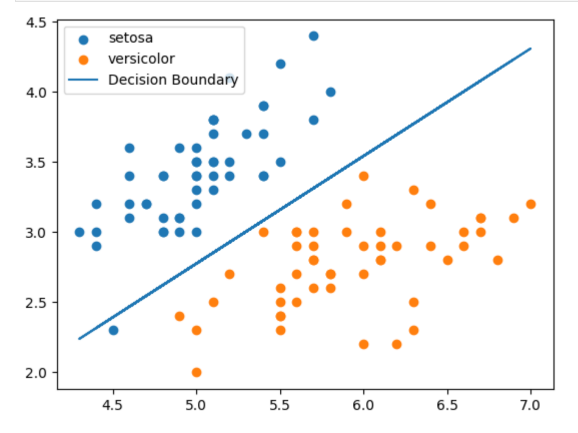
print('Accuracy: ', np.sum(np.equal(y,predicted)) / len(predicted))

Accuracy: 0.99

Let us plot the decision boundary for the new weights:

```
In [19]: x_values = x[:,0]
y_values = - (bias + weights[0]*x_values) / weights[1]

plt.figure()
for label in np.unique(y):
    plt.scatter(x[:,0][y==label],x[:,1][y==label], label = data.target_names[label]
plt.plot(x_values, y_values, label='Decision Boundary')
plt.legend()
plt.show()
```



```
In []:
```