Bootcamp Machine Learning



Module08 Logistic Regression

Module 08 - Logistic Regression

Today, you will discover your first classification algorithm: logistic regression. You will learn its cost function, gradient descent and some metrics to evaluate its performance.

Notions of the module

Logistic hypothesis, logistic gradient descent, logistic regression, multiclass classification. Accuracy, precision, recall, F1-score, confusion matrix.

Useful Resources

You are strongly advise to use the following resource: Machine Learning MOOC - Stanford Here are the sections of the MOOC that are relevant for today's exercises:

Week 3:

Classification and representation:

- Classification (Video + Reading)
- Hypothesis Representation (Video + Reading)
- Decision Boundary (Video + Reading)

Logistic Regression Model:

- Cost Function (Video + Reading)
- Simplified Cost Function and Gradient Descent (Video + Reading)

Multiclass Classification:

- Mutliclass Classification: One-vs-all (Video + Reading)
- Review (Reading + Quiz)

General rules

- The Python version to use is 3.7, you can check with the following command: python -V
- The norm: during this bootcamp you will follow the Pep8 standards
- The function eval is never allowed.
- The exercises are ordered from the easiest to the hardest.
- Your exercises are going to be evaluated by someone else, so make sure that your variable names and function names are appropriate and civil.
- Your manual is the internet.
- You can also ask questions in the #bootcamps channel in 42AI's Slack workspace.
- If you find any issues or mistakes in this document, please create an issue on our dedicated Github repository.

Helper

Ensure that you have the right Python version.

```
> which python
/goinfre/miniconda/bin/python
> python -V
Python 3.7.*
```

> which pip /goinfre/miniconda/bin/pip

Exercise 00 - Multivariate Linear Regression with Class

Exercise 01 - DataSplitter

Exercise 02 - Ai Key Notions

Interlude - Classification: The Art of Labelling Things

Interlude - Predict I: Introducing the Sigmoid Function

Exercise 03 - Sigmoid

Interlude - Predict II: Hypothesis

Exercise 04 - Logistic Hypothesis

Interlude - Evaluate

Exercise 05 - Logistic Loss Function

Interlude - Linear Algebra Strikes again!

Exercise 06 - Vectorized Logistic Loss Function

Interlude - Improve

Exercise 07 - Logistic Gradient

Interlude - Vectorized Logistic Gradient

Exercise 08 - Vectorized Logistic Gradient

Exercise 09 - Logistic Regression

Exercise 10 - Practicing Logistic Regression

Interlude - More Evaluation Metrics!

Exercise 11 - Other metrics

Exercise 12 - Confusion Matrix

Exercise 00 - Multivariate Linear Regression with Class

Turn-in directory: ex00/

Files to turn in: mylinearregression.py

Authorized modules : numpy Forbidden modules : sklearn

AI Classics:

These exercises are key assignments from the previous module. If you haven't completed them yet, you should finish them first before you continue with today's exercises.

Objective:

Upgrade your Linear Regression class so it can handle multivariate hypotheses.

Instructions:

You are expected to upgrade your own **MyLinearRegression** class from **module06**. You will upgrade (at least) the following methods to support multivariate linear regression:

- predict_(self, x)
- fit_(self, x, y)
 Depending on how you have implemented your methods

```
import numpy as np
from mylinearregression import MyLinearRegression as MyLR
X = np.array([[1., 1., 2., 3.], [5., 8., 13., 21.], [34., 55., 89., 144.]])
Y = np.array([[23.], [48.], [218.]])
mylr = MyLR([[1.], [1.], [1.], [1.], [1]])
mylr.predict_(X)
array([[8.], [48.], [323.]])
mylr.cost_elem_(X,Y)
array([[225], [0.], [11025]])
mylr.cost_(X,Y)
mylr.alpha = 1.6e-4
mylr.max_iter = 200000
mylr.fit_(X, Y)
mylr.theta
array([[18.188..], [2.767..], [-0.374..], [1.392..], [0.017..]])
mylr.predict_(X)
array([[23.417..], [47.489..], [218.065...]])
mylr.cost_elem_(X,Y)
array([[0.174..], [0.260..], [0.004..]])
mylr.cost_(X,Y)
```

Exercise 01 - DataSpliter

Turn-in directory: ex01/

Files to turn in : data_splitter.py

Authorized modules : numpy Forbidden modules : sklearn

AI Classics:

These exercises are key assignments from the previous module. If you haven't completed them yet, you should finish them first before you continue with today's exercises.

Objective:

Learn how to split a dataset into a **training set** and a **test set**.

Instructions:

You must implement a function that **shuffles** and **splits** a dataset it in two parts: a **training set** and a **test** set.

- Your function will shuffle and split the X matrix while keeping a certain **proportion** of the examples for training, and the rest for testing.
- Your function will also shuffle and split the y vector while making sure that the order of the rows in the output match the order of the rows in the split X output.

In the data_spliter.py file, you have to code the following function as per the instructions given below:

Examples:

Be careful! The dataset has to be randomly shuffled before it is split into training and test sets. Unless you use the same seed in your randomization algorithm, you won't get the same results twice. The following examples are just an indication of possible outputs. As long as you have shuffled datasets with their corresponding y values, your function is working correctly.

```
import numpy as np
x1 = np.array([1, 42, 300, 10, 59])
y = np.array([0, 1, 0, 1, 0])
# Example 1:
```

```
data_spliter(x1, y, 0.8)
(array([ 1, 59, 42, 300]), array([10]), array([0, 0, 1, 0]), array([1]))
data_spliter(x1, y, 0.5)
(array([59, 10]), array([ 1, 300, 42]), array([0, 1]), array([0, 0, 1]))
x2 = np.array([[ 1, 42],
               [ 10, 42]])
y = np.array([0, 1, 0, 1, 0])
data_spliter(x2, y, 0.8)
(array([[ 10, 42],
              59],
              10]]),
 array([[ 1, 42]]),
 array([0, 1, 0, 1]),
 array([0]))
data_spliter(x2, y, 0.5)
(array([[59, 1],
        [10, 42]]),
 array([[300, 10],
        [ 1, 42]]),
 array([0, 0]),
 array([1, 1, 0]))
```

Exercise 02 - AI Key Notions:

These questions are about key notions from the previous modules. Making sure you can formulate a clear answer to each of them is necessary before you keep going. Discuss them with a fellow student if you can.

Are you able to clearly and simply explain:

- 1 What is the main difference between univariate and multivariate linear regression, in terms of variables?
- 2 Is there a minimum number of variables needed to perform a multivariate linear regression? If so, what is it?
- 3 Is there a maximum number of variables needed to perform a multivariate linear regression? If so, what is it?
- 4 Is there a difference between univariate and multivariate linear regression in terms of performance evaluation?
- 5 What does it mean geometrically to perform multivariate gradient descent with two variables?

Interlude - Classification: The Art of Labelling Things

Over the last three modules you have implemented your first machine learning algorithm. You also discovered the three main steps we follow when we build **learning algorithms**:

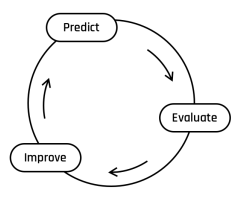


Figure 1: The Learning Cycle

The first algorithm you discovered, **Multivariate Linear Regression**, can now be used to predict a numerical value, based on several features. This algorithm uses gradient descent to optimize its cost function.

Now let's introduce you to your first **classification algorithm**: it is named **Logistic Regression**. It peforms a *classification task*, which means that you are not predicting a numerical value (like price, age, grades...) but **categories**, or **labels** (like dog, cat, sick/healty...).

Be careful! Don't be confused by the word 'regression' in **Logistic Regression**. It really is a classification task! The name is a bit tricky but you will quickly get used to it.

Once again: Logistic Regression is a classification algorithm which assigns to a given example a category.

Terminolgy:

In this module we will use the following terms interchangeably: **class, category,** and **label**. They all refer to the *groups* to which each training example can be assigned to, in a classification task.

Interlude - Predict I: Introducing the Sigmoid Function

Formulating a Hypothesis

Remember that a hypothesis, denoted $h(\theta)$, is an equation that combines a set of **features** (that characterize an example) with **parameters** in order to output a **prediction**. Remember the hypothesis we used in linear regression?

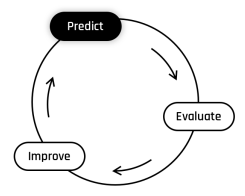


Figure 2: The Learning Cycle - Predict

$$h(\theta) = \theta_0 + \theta_1 x_1^{(i)} + \dots + \theta_n x_n^{(i)} = \theta \cdot x'^{(i)}$$

It worked fine to predict continuous values, but could we also use it to tell, for example, if a patient is sick or not?

That's a yes-or-no question, so the output from the hypothesis function should reflect that.

To get started, we'll assign each class a numerical value: sick patients will be assigned a value of 1, and healthy patients will be assigned a value of 0. The goal will be to build a hypothesis that outputs a probability that a patient is sick, as a float number within the range of 0 and 1.

The good news is that we can keep the linear equation we already worked with! All we need to do is squash its output through another function that is bounded between 0 and 1. That's the **Sigmoid function** and your next exercise is to implement it!

Exercise 03 - Sigmoid

 $\begin{array}{ccc} \text{Turn-in directory}: & \text{ex}03/\\ & \text{Files to turn in}: & \text{sigmoid.py} \\ \text{Forbidden functions}: & \text{None} \\ & \text{Remarks}: & \text{n/a} \\ \end{array}$

Objective:

Introduction to the hypothesis in the case of logistic regression. You must implement the sigmoid function, given by the following formula:

$$\operatorname{sigmoid}(x) = \frac{1}{1 + e^{-x}}$$

Where:

• x is a scalar or a vector,

ullet e is the contracted form for exponential function. It is also a mathematical constant, named Euler's number.

This function is also known as **Standard logistic sigmoid function**. This explains the name *logistic regression*.

The sigmoid function transforms an input into a probability value, i.e. a value between 0 and 1. This probability value will then be used to classify the inputs.

Instructions:

In the sigmoid.py file, write the following function as per the instructions below:

```
def sigmoid_(x):
    """
    Compute the sigmoid of a vector.
    Args:
        x: has to be a numpy.ndarray, a vector.
    Returns:
        The sigmoid value as a numpy.ndarray.
        None if x is an empty numpy.ndarray.
    Raises:
        This function should not raise any Exception.
    """
    ... Your code ...
```

Examples:

```
# Example 1:
x = np.array(-4)
sigmoid_(x)
# Output:
array([0.01798620996209156])

# Example 2:
x = np.array(2)
sigmoid_(x)
# Output:
array([0.8807970779778823])

# Example 3:
x = np.array([[-4], [2], [0]])
sigmoid_(x)
# Output:
array([[0.01798620996209156], [0.8807970779778823], [0.5]])
```

To go further

Our sigmoid formula is a special case of the logistic function below, with L=1, k=1 and $x_0=0$:

$$f(x) = \frac{L}{1 + e^{-k(x - x_0)}}$$

Interlude - Predict II: Hypothesis

We hope your curiosity led you to plot your sigmoid function. If you didn't, well here is what it looks like:

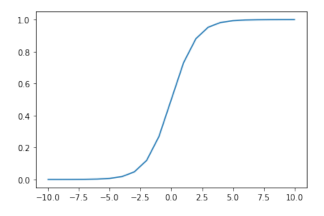


Figure 3: Sigmoid

As you can see, the sigmoid's output values ranging from 0 to 1.

You can input real numbers as big as you want (positive or negative), the output will always land within this range. This will be very helpful for the next part.

Logistic Hypothesis

Now you've written your sigmoid function, let's look at the logistic regression hypothesis.

$$\hat{y}^{(i)} = h_{\theta}(x^{(i)}) = \text{sigmoid}(\theta \cdot x'^{(i)}) = \frac{1}{1 + e^{-\theta \cdot x'^{(i)}}} \quad \text{for i} = 1, \dots, m$$

This is simply the sigmoid function applied on top of the linear regression hypothesis!!

It can be vectorized as:

$$\hat{y} = h_{\theta}(X) = \operatorname{sigmoid}(X'\theta) = \frac{1}{1 + e^{-X'\theta}}$$

As we said before: the sigmoid function is just a way to map the result of a linear equation onto a [0,1] value range.

This transformation allows us to interpret the result as a **probability that an individual is a member of a given class**.

Exercise 04 - Logistic Hypothesis

Turn-in directory: ex04/
Files to turn in: log_pred.py
Forbidden libraries: None
Remarks: n/a

Objective:

Introduction to the hypothesis notion in case of logistic regression. You must implement the following formula as a function:

$$\hat{y} = \operatorname{sigmoid}(X' \cdot \theta) = \frac{1}{1 + e^{-X' \cdot \theta}}$$

Where:

- X is a matrix of dimensions $(m \times n)$, the design matrix,
- X' is a matrix of dimensions $(m \times (n+1))$, the design matrix onto which a column of 1's is added as a first column,
- \hat{y} is a vector of dimension m, the vector of predicted values,
- θ is a vector of dimension (n+1), the vector of parameters.

Be careful:

- the x your function will get as an input corresponds to X, the $m \times n$ matrix. Not X'.
- θ is an (n+1) vector.

You have to transform x to fit theta's dimension!

Instructions:

In the log_pred.py file, write the following function as per the instructions below:

```
def logistic_predict_(x, theta):
    """Computes the vector of prediction y_hat from two non-empty numpy.ndarray.
    Args:
        x: has to be an numpy.ndarray, a vector of dimension m * n.
        theta: has to be an numpy.ndarray, a vector of dimension (n + 1) * 1.
    Returns:
        y_hat as a numpy.ndarray, a vector of dimension m * 1.
        None if x or theta are empty numpy.ndarray.
        None if x or theta dimensions are not appropriate.
    Raises:
        This function should not raise any Exception.
    """
        ... Your code ...
```

Interlude - Evaluate

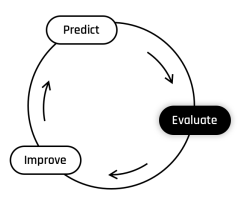


Figure 4: The Learning Cycle - Evaluate

Our model can predict the probability for a given example to be part of the class labeled as 1. Now it's time to evaluate how good it is.

The previous cost function, used to evaluate linear regression, is not appropriate in a classification case.

Given the fact that classification tasks imply only two possible values:

- zero, if the element is not a member of the predicted class,
- one, if the element is a member of the predicted class,

measuring the 'distance' between the prediction and the label is not going to be the best way to evaluate the performance of a classification model. We'll prefer the **logarithmic** function because it can penalize the wrong predictions even more harshly. But let's separate the two possible cases.

Case 1: The expected output is 1

In mathematical terms, we write:

$$y^{(i)} = 1$$

Here we need a function that will penalize the classifier with a high cost if its prediction (\hat{y}) gets close to 0. What do you think of this function? (Have a look at its plot)

$$cost_{y=1} = -\log(\hat{y})$$

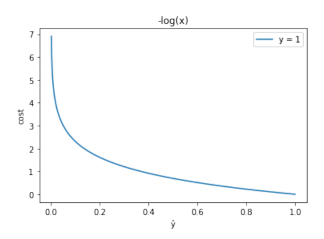


Figure 5: Cost function when y = 1

You can see from the plot that:

- if the prediction (\hat{y}) is close to 0, the cost will be great,
- if the prediction (\hat{y}) is close to 1, the cost will be small.

So we got our function that can harshly penalize predictions that get close to 0. But sometimes, $y^{(i)}$ is NOT equal to 1. What if we want \hat{y} to be closer to 0 instead?

Case 2: The expected output is 0

In this case we have:

$$y^{(i)} = 0$$

We just need to manipulate the last equation slightly in order to flip the curve the way we need:

$$cost_{y=0} = -\log(1 - \hat{y}^{(i)})$$

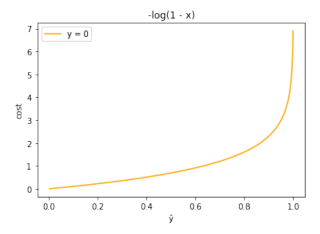


Figure 6: Cost function when y = 0

You can see from the plot that:

- if the prediction is close to 1, the cost will be great,
- if the prediction is close to 0, the cost will be small.

So this second equation works like the first one, but penalizes the other way: this time $\hat{y}^{(i)}$ gets penalized harder when it gets close to 1.

Now, all we need is a smart way to automatically choose which cost function to use, depending on the value of $y^{(i)}$.

Putting it all together

Let's recap. We need a cost function that can alternate between these:

• If $y^{(i)} = 1$

$$cost = cost_{y=1} = -\log(\hat{y}^{(i)})$$

• If $y^{(i)} = 0$

$$cost = cost_{y=0} = -\log(1 - \hat{y}^{(i)})$$

And we can represent it like this:

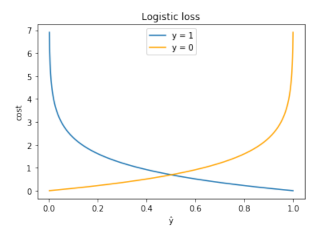


Figure 7: $cost_{y=0}$ and $cost_1$

How do you switch between $cost_{y=0}$ and $cost_{y=1}$ depending on the value of $y^{(i)}$? We could use an if-else statement in the code, but that's not very pretty and it doesn't provide a cost function that can be expressed as a single mathematical expression. It turns out there is a little mathematical trick we can use to make everything stand in one equation.

Building the equation for a single training example

For this part let's go step by step. The strategy is to sum both expressions:

$$cost = cost_{y=1} + cost_{y=0}$$

And then we need some kind of switch to "turn off" the term that shouldnt be use for the example i. It turns out we can use the $y^{(i)}$ value itself as a switch!

• When $y^{(i)} = 0$, we just multiply it with the term we don't want and we'll cancel it out:

$$cost = y^{(i)} \cdot cost_{y=1} + cost_{y=0}$$
$$cost = 0 \cdot cost_{y=1} + cost_{y=0}$$
$$cost = cost_{y=0}$$

• When $y^{(i)} = 1$, it's a little trickier. We have to multiply the term we want to cancel out by $(1 - y^{(i)})$

$$\begin{array}{rcl} cost &=& cost_{y=1} + (1-y^{(i)}) \cdot cost_{y=0} \\ cost &=& cost_{y=1} + (1-1) \cdot cost_{y=0} \\ cost &=& cost_{y=1} + 0 \cdot cost_{y=0} \\ cost &=& cost_{y=1} \end{array}$$

Now, to make a generic equation that works without knowing in advance the value of $y^{(i)}$, all we need is to sum the two cost functions along with their "switches":

$$cost = y^{(i)} \cdot cost_{y=1} + (1 - y^{(i)}) \cdot cost_{y=0}$$

And then, if we develop $cost_{y=0}$ and $cost_{y=1}$:

$$cost = y^{(i)} \cdot (-\log(\hat{y}^{(i)})) + (1 - y^{(i)}) \cdot (-\log(1 - \hat{y}^{(i)}))$$

Finally, if we simplify the sign notation just a bit:

$$cost = -[y^{(i)} \cdot \log(\hat{y}^{(i)}) + (1 - y^{(i)}) \cdot \log(1 - \hat{y}^{(i)})]$$

Cross-entropy

We are reaching the goal! All we need to do is and average across all training examples and we end up with our final cost function. It has a name: **cross-entropy**. The equation is the following:

$$J(\theta) = -\frac{1}{m} \left[\sum_{i=1}^{m} y^{(i)} \log(\hat{y}^{(i)}) + (1 - y^{(i)}) \log(1 - \hat{y}^{(i)}) \right]$$

This formula allows you to calculate the overall cost of a complete set of predictions. If you have enough, you can stop here and move on to the exercise. If you'd like to better understand how it works and have "automatic switch" process broken down for you, here we go:

If the given example $x^{(i)}$ is not part of the predicted class, $y^{(i)} = 0$:

$$\begin{array}{cccc} y^{(i)} & = & 0 \\ y^{(i)} \log(\hat{y}^{(i)})) & = & 0 \\ 1 - y^{(i)} & = & 1 \\ (1 - y^{(i)}) \log(1 - \hat{y}^{(i)}) & = & \log(1 - \hat{y}^{(i)}) \end{array}$$

Therefore

$$\begin{split} J(\theta) &= -\frac{1}{m} [\sum_{i=1}^{m} \overbrace{y^{(i)} \log(\hat{y}^{(i)})}^{0} + \overbrace{(1-\hat{y}^{(i)})}^{1} \log(1-\hat{y}^{(i)})] \\ J(\theta) &= -\frac{1}{m} \sum_{i=1}^{m} \log(1-\hat{y}^{(i)}) \\ J(\theta) &= \frac{1}{m} \sum_{i=1}^{m} -\log(1-\hat{y}^{(i)}) \end{split}$$

If the given example $\boldsymbol{x}^{(i)}$ is part of the predicted class, $\boldsymbol{y}^{(i)} = 1$:

$$\begin{array}{cccc} y^{(i)} & = & 1 \\ y^{(i)} \log(\hat{y}^{(i)}) & = & \log(\hat{y}^{(i)}) \\ 1 - y^{(i)} & = & 0 \\ (1 - y^{(i)}) \log(1 - \hat{y}^{(i)}) & = & 0 \end{array}$$

Therefore

$$J(\theta) = -\frac{1}{m} \left[\sum_{i=1}^{m} \widehat{y^{(i)}} \log(\widehat{y}^{(i)}) + \underbrace{(1 - y^{(i)}) \log(1 - \widehat{y}^{(i)})}_{0} \right]$$

$$J(\theta) = -\frac{1}{m} \sum_{i=1}^{m} \log(\widehat{y}^{(i)})$$

$$J(\theta) = \frac{1}{m} \sum_{i=1}^{m} -\log(\widehat{y}^{(i)})$$

Exercise 05 - Logistic Loss Function

 $\begin{array}{ll} \text{Turn-in directory}: & \text{ex}05/\\ \text{Files to turn in}: & \text{log_loss.py} \\ \text{Forbidden libraries}: & \text{Numpy} \\ \text{Remarks}: & \text{n/a} \\ \end{array}$

Objective:

Understanding and manipulation of the cost function concept in the case of logistic regression. You must implement the following formula as a function:

$$J(\theta) = -\frac{1}{m} \left[\sum_{i=1}^{m} y^{(i)} \log(\hat{y}^{(i)}) \right] + (1 - y^{(i)}) \log(1 - \hat{y}^{(i)})$$

Where:

- \hat{y} is a vector of dimension m, the vector of predicted values,
- $\hat{y}^{(i)}$ is the i^{th} component of the \hat{y} vector,
- y is a vector of dimension m, the vector of expected values,
- $y^{(i)}$ is the i^{th} component of the y vector.

Instructions:

In the log loss.py file, write the following function as per the instructions below:

```
def log_loss_(y, y_hat, eps=1e-15):
    """
    Computes the logistic loss value.
    Args:
        y: has to be an numpy.ndarray, a vector of shape m * 1.
        y_hat: has to be an numpy.ndarray, a vector of shape m * 1.
        eps: has to be a float, epsilon (default=1e-15)
    Returns:
        The logistic loss value as a float.
        None on any error.
    Raises:
        This function should not raise any Exception.
    """
        ... Your code ...
```

Hint:

The logarithmic function isn't defined in 0.

This means that if $y^{(i)} = 0$ you will get an error when you try to compute $log(y^{(i)})$.

The purpose of the eps argument is to avoid log(0) errors. It is a very small residual value we add to y.

```
# Example 1:
y1 = np.array([1])
x1 = np.array([4])
theta1 = np.array([[2], [0.5]])
y_hat1 = logistic_predict(x1, theta1)
```

```
log_loss_(y1, y_hat1)
# Output:
0.01814992791780973

# Example 2:
y2 = np.array([[1], [0], [1], [0], [1]])
x2 = np.array([[2], [0.5]])
y_hat2 = np.array([[2], [0.5]])
y_hat2 = logistic_predict(x2, theta2)
log_loss_(y2, y_hat2)
# Output:
2.4825011602474483

# Example 3:
y3 = np.array([[0], [1], [1]])
x3 = np.array([[0, 2, 3, 4], [2, 4, 5, 5], [1, 3, 2, 7]])
theta3 = np.array([[-2.4], [-1.5], [0.3], [-1.4], [0.7]])
y_hat3 = logistic_predict(x3, theta3)
log_loss_(y3, y_hat3)
# Output:
2.9938533108607053
```

To go further:

This function is called **Cross-Entropy loss**, or **logistic loss**.

For more information you can look at this section of the Cross entropy Wikipedia.

Interlude - Linear Algebra Strikes Again!

You've become quite used to vectorization by now. You may have already tried to vectorize the logistic cost function by yourself. Let's look one last time at the former equation:

$$J(\theta) = -\frac{1}{m} \left[\sum_{i=1}^{m} y^{(i)} \log(\hat{y}^{(i)}) \right] + (1 - y^{(i)}) \log(1 - \hat{y}^{(i)})$$

Vectorized Logistic Cost Function

In the **vectorized version**, we remove the sum (\sum) because it is captured by the dot products:

$$J(\theta) = -\frac{1}{m} [y \cdot \log(\hat{y}) + (\vec{1} - y) \cdot \log(\vec{1} - \hat{y})]$$

Where:

• $\vec{1}$ is a vector full of 1's with the same dimension as y (i.e. m)

$$\vec{1} = \begin{bmatrix} 1 \\ \vdots \\ 1 \end{bmatrix}$$

Note: Operations Between Vectors and Scalars

We use the $\vec{1}$ notation to be rigorous, because addition (or subtraction) between a vector and a scalar is not defined. In other words, mathematically, you cannot write this: 1-y

The only operation defined between a scalar and a vector is multiplication, remember?

However...

NumPy is a bit permissive on vectors and matrix operations... The following instructions will get you the same results:

Strange, isn't it?

It happens because of one of NumPy's permissive operations called **Broadcasting**. Broadcasting is a powerful feature whereby NumPy is able to figure out that you actually wanted to perform a subtraction on each element in the vector, so it does it for you automatically. It's very handy to write concise lines of code, but it can insert very sneaky bugs if you aren't 100% confident in what you're doing.

Many of the bugs you will encounter while working on Machine Learning problems will come from NumPy's permissiveness. Such bugs generaly don't throw any errors, but mess up the content of your vectors and matrices and you'll spend an awful lot of time looking for why your model doesn't learn. This is why we strongly suggest that you pay attention to your vector (and matrix) shape and stick as much as possible to the actual mathematical operations.

For more information, you can watch this video on dealing with Broadcasting.

Exercise 06 - Vectorized Logistic Loss Function

Turn-in directory: ex06/
Files to turn in: vec_log_loss.py
Forbidden functions: None
Remarks: n/a

Objective:

Understanding and manipulation of cost function concept in case of logistic regression. You must implement the following formula as a function:

$$J(\theta) = -\frac{1}{m} [y \cdot \log(\hat{y}) + (\vec{1} - y) \cdot \log(\vec{1} - \hat{y})]$$

Where:

- \hat{y} is a vector of dimension m, the vector of predicted values
- y is a vector of dimension m, the vector of expected values
- $\vec{1}$ is a vector of dimension m, a vector full of ones.

Instructions:

In the vec_log_loss.py file, write the following function as per the instructions below:

```
def vec_log_loss_(y, y_hat, eps=1e-15):
    """"
    Compute the logistic loss value.
    Args:
        y: has to be an numpy.ndarray, a vector of shape m * 1.
        y_hat: has to be an numpy.ndarray, a vector of shape m * 1.
        eps: epsilon (default=1e-15)
    Returns:
        The logistic loss value as a float.
        None on any error.
    Raises:
        This function should not raise any Exception.
    """
```

Hint: the purpose of epsilon (eps) is to avoid log(0) errors, it is a very small residual value we add to y.

```
# Example 1:
y1 = np.array([1])
x1 = np.array([2], [0.5]])
theta1 = np.array([2], [0.5]])
y_hat1 = logistic_predict(x1, theta1)
vec_log_loss_(y1, y_hat1)
# Output:
0.01814992791780973

# Example 2:
y2 = np.array([[1], [0], [1], [0], [1]])
x2 = np.array([[4], [7.16], [3.2], [9.37], [0.56]])
theta2 = np.array([[2], [0.5]])
y_hat2 = logistic_predict(x2, theta2)
vec_log_loss_(y2, y_hat2)
# Output:
2.4825011602474483

# Example 3:
y3 = np.array([[0], [1], [1]])
x3 = np.array([[0], 2, 3, 4], [2, 4, 5, 5], [1, 3, 2, 7]])
theta3 = np.array([[-2.4], [-1.5], [0.3], [-1.4], [0.7]])
y_hat3 = logistic_predict(x3, theta3)
```

vec_log_loss_(y3, y_hat3)

Output:

2.9938533108607053

Improve

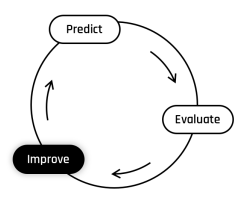


Figure 8: The Learning Cycle: Improve

Now we want to improve the algorithm's performance, or in other words, reduce the cost of its predictions. This brings us (again) to calculating the gradient, which will tell us how much and in which direction the θ parameters belonging to the model will be adjusted.

The logistic gradient

If you remember, to calculate the gradient, we start with the cost function and we derive it with respect to each of the theta parameters. If you know multivariate calculus you can try it for yourself, otherwise we've done it for you:

$$\nabla(J)_0 = \frac{1}{m} \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)})$$

$$\nabla(J)_j = \frac{1}{m} \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)}) x_j^{(i)} \quad \text{for j = 1, ..., n}$$

Where:

- $\nabla(J)$ is a vector of size (n+1), the gradient vector,
- $\nabla(J)_j$ is the j^{th} component of $\nabla(J)$, the partial derivative of J with respect to θ_j ,
- y is a vector of dimension m, the vector of expected values,
- $y^{(i)}$ is a scalar, the i^{th} component of vector y,
- $x^{(i)}$ is the feature vector of the i^{th} example,
- $x_i^{(i)}$ is a scalar, the j^{th} feature value of the i^{th} example,
- $h_{\theta}(x^{(i)})$ is a scalar, the model's estimation of $y^{(i)}$.

This formula should be very familiar to you, as it's the same as the linear regression gradient! The only difference is that $h_{\theta}(x^{(i)})$ corresponds to the logistic regression hypothesis instead of the linear regression hypothesis.

In other words:

$$h_{\theta}(x^{(i)}) = \operatorname{sigmoid}(\theta \cdot x'^{(i)}) = \frac{1}{1 + e^{-\theta \cdot x'^{(i)}}}$$

Instead of:

$$h_{\theta}(x^{(i)}) = \theta \cdot x'^{(i)}$$

Exercise 07 - Logistic Gradient

Turn-in directory: ex07/

Files to turn in: log_gradient.py

Forbidden libraries: numpy

Remarks: n/a

Objective:

Understand and manipulate the concept of gradient in case of logistic formulation. You must implement the following formula as a function:

$$\nabla(J)_0 = \frac{1}{m} \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)})$$

$$\nabla(J)_j = \frac{1}{m} \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)}) x_j^{(i)} \quad \text{for j = 1, ..., n}$$

Where:

- $\nabla(J)$ is a vector of size (n+1), the gradient vector,
- $\nabla(J)_j$ is the j^{th} component of $\nabla(J)$, the partial derivative of J with respect to θ_j ,
- y is a vector of dimension m, the vector of expected values,
- $y^{(i)}$ is a scalar, the i^{th} component of vector y,
- $x^{(i)}$ is the feature vector of the i^{th} example,
- $x_i^{(i)}$ is a scalar, the j^{th} feature value of the i^{th} example,
- $h_{\theta}(x^{(i)})$ is a scalar, the model's estimation of $y^{(i)}$.

Remember that with logistic regression, the hypothesis is slightly different:

$$h_{\theta}(x^{(i)}) = sigmoid(\theta \cdot x'^{(i)})$$

Instructions:

In the log_gradient.py file, write the following function as per the instructions below:

```
def log_gradient(x, y, theta):
    """Computes a gradient vector from three non-empty numpy.ndarray, with a for-loop. The
    _ three arrays must have compatible dimensions.
    Args:
        x: has to be an numpy.ndarray, a matrix of shape m * n.
        y: has to be an numpy.ndarray, a vector of shape m * 1.
        theta: has to be an numpy.ndarray, a vector of shape (n + 1) * 1.
    Returns:
        The gradient as a numpy.ndarray, a vector of shape n * 1, containing the result of the
        formula for all j.
        None if x, y, or theta are empty numpy.ndarray.
        None if x, y and theta do not have compatible dimensions.
    Raises:
        This function should not raise any Exception.
    """
        ... Your code ...
```

```
y1 = np.array([1])
x1 = np.array([4])
theta1 = np.array([[2], [0.5]])
log_gradient(x1, y1, theta1)
array([[-0.01798621],
        [-0.07194484]])
y2 = np.array([[1], [0], [1], [0], [1]])
x2 = np.array([[4], [7.16], [3.2], [9.37], [0.56]])
theta2 = np.array([[2], [0.5]])
log_gradient(x2, y2, theta2)
array([[0.3715235],
        [3.25647547]])
y3 = np.array([[0], [1], [1]])
x3 = np.array([[0, 2, 3, 4], [2, 4, 5, 5], [1, 3, 2, 7]])
theta3 = np.array([[-2.4], [-1.5], [0.3], [-1.4], [0.7]])
log_gradient(x3, y3, theta3)
array([[-0.55711039],
        [-0.90334809],
[-2.01756886],
        [-3.27257351]])
```

Interlude - Vectorized Logistic Gradient

Given the previous logistic gradient formula, it's quite easy to produce a vectorized version. Actually, you almost already implemented it on module 07!

As with the previous exercice, the only thing you have to change is your hypothesis in order to calculate your logistic gradient.

$$\nabla(J)_{0} = \frac{1}{m} \sum_{i=1}^{m} (h_{\theta}(x^{(i)}) - y^{(i)})$$

$$\nabla(J)_{j} = \frac{1}{m} \sum_{i=1}^{m} (h_{\theta}(x^{(i)}) - y^{(i)}) x_{j}^{(i)} \quad \text{for j = 1, ..., n}$$

Vectorized Version

Can be vectorized the same way as you did before:

$$\nabla(J) = \frac{1}{m} X'^{T} (h_{\theta}(X) - y)$$

Exercise 08 - Vectorized Logistic Gradient

Turn-in directory: ex08/

Files to turn in : vec_log_gradient.py

Forbidden libraries : Numpy

Remarks: n/a

Objective:

Understand and manipulation of the concept of gradient in the case of logistic formulation. You must implement the following formula as a function:

$$\nabla(J) = \frac{1}{m} X^{\prime T} (h_{\theta}(X) - y)$$

Where:

- $\nabla(J)$ is the gradient vector of size (n+1),
- X' is a matrix of dimension $m \times (n+1)$, the design matrix onto which a column of ones was added as the first column,
- $X^{\prime T}$ means the matrix has been transposed,
- $h_{\theta}(X)$ is a vector of dimension m, the vector of predicted values,
- y is a vector of dimension m, the vector of expected values.

Instructions:

In the vec_log_gradient.py file, write the following function as per the instructions below:

```
def vec_log_gradient(x, y, theta):
    """Computes a gradient vector from three non-empty numpy.ndarray, without any for-loop.
    The three arrays must have compatible shapes.
Args:
    x: has to be an numpy.ndarray, a matrix of shape m * n.
    y: has to be an numpy.ndarray, a vector of shape m * 1.
    theta: has to be an numpy.ndarray, a vector (n +1) * 1.
    Returns:
        The gradient as a numpy.ndarray, a vector of shape n * 1, containg the result of the formula for all j.
        None if x, y, or theta are empty numpy.ndarray.
        None if x, y and theta do not have compatible shapes.
    Raises:
        This function should not raise any Exception.
    """
        ... Your code ...
```

```
y1 = np.array([1])
x1 = np.array([4])
theta1 = np.array([[2], [0.5]])
vec_log_gradient(x1, y1, theta1)
array([[-0.01798621],
        [-0.07194484]])
y2 = np.array([[1], [0], [1], [0], [1]])
x2 = np.array([[4], [7.16], [3.2], [9.37], [0.56]])
theta2 = np.array([[2], [0.5]])
vec_log_gradient(x2, y2, theta2)
array([[0.3715235],
        [3.25647547]])
y3 = np.array([[0], [1], [1]])
x3 = np.array([[0, 2, 3, 4], [2, 4, 5, 5], [1, 3, 2, 7]])
theta3 = np.array([[-2.4], [-1.5], [0.3], [-1.4], [0.7]])
vec_log_gradient(x3, y3, theta3)
array([[-0.55711039],
        [-0.90334809],
[-2.01756886],
        [-2.10071291],
        [-3.27257351]])
```

Exercise 09 - Logistic Regression

Turn-in directory: ex09/

Files to turn in: my_logistic_regression.py

Authorized modules : numpy Forbidden modules : sklearn

Objective:

The time to use everything you built so far has come! Demonstrate your knowledge by implementing a logistic regression classifier using the gradient descent algorithm. You must have seen the power of numpy for vectorized operations. Well let's make something more concrete with that.

You may have to take a look at Scikit-Learn's implementation of logistic regression and noticed that the **sklearn.linear_model.LogisticRegression** class offers a lot of options.

The goal of this exercise is to make a simplified but nonetheless useful and powerful version, with fewer options.

Instructions:

In the my_logistic_regression.py file, write a MyLogisticRegression class as in the instructions below:

```
class MyLogisticRegression():
    """
    Description:
        My personnal logistic regression to classify things.
    """

def __init__(self, theta, alpha=0.001, max_iter=1000):
        self.alpha = alpha
        self.max_iter = max_iter
        self.theta = theta
        ... Your code here ...
... other methods ...
```

You will add the following methods:

```
predict_(self, x)cost_elem_(self, y, yhat)cost_(self, y, yhat)fit_(self, x, y)
```

You have already written these functions, you will just need few adjustments so that they all work well within your MyLogisticRegression class.

```
import numpy as np
from my_logistic_regression import MyLogisticRegression as MyLR
X = np.array([[1., 1., 2., 3.], [5., 8., 13., 21.], [3., 5., 9., 14.]])
Y = np.array([[1], [0], [1]])
mylr = MyLR([2, 0.5, 7.1, -4.3, 2.09])

# Example 0:
mylr.predict_(X)
# Output:
array([[0.99930437],
```

Exercise 10 - Practicing Logistic Regression

Turn-in directory: ex10/

Files to turn in : mono_log.py, nulti_log.py

Authorized modules : Numpy Forbidden modules : sklearn

Objective:

Now it's time to test your Logistic Regression Classifier on real data! You will use the **solar_system_census_dataset**.

Instructions:

Some words about the dataset:

- You will work with data from the last Solar System Census.
- The dataset is divided in two files which can be found in the resources folder: solar_system_census.csv and solar_system_census_planets.csv.
- The first file contains biometric information such as the height, weight, and bone density of several Solar System citizens.

• The second file contains the homeland of each citizen, indicated by its Space Zipcode representation (i.e. one number for each planet...:)).

As you should know, Solar citizens come from four registered areas (zipcodes):

- The flying cities of Venus (0),
- United Nations of Earth (1),
- Mars Republic (2),
- The Asteroids' Belt colonies (3).

Your Task:

You are expected to produce 2 programs that will use Logistic Regression to predict from which planet each citizen comes from, based on the other variables found in the census dataset.

But wait... what? There are four different planets! How do you make a classifier discriminate between 4 categories? Let's go step by step...

One Label to Discriminate Them All

You already wrote a Logistic Regression Classifier that can discriminate between two classes. We can use it to solve the problem! Let's start by having it discriminate between citizens who come from your favorite planet and everybody else! Your program (mono_log.py) will: 1. Take an argument -zipcode=x whit x being 0, 1, 2 or 3. If no argument, the usage will be displayed. 2. Split the dataset into a training and a test set. 3. Select your favorite Space Zipcode and generate a new numpy.ndarray to label each citizen according to your new selection criterion:

- \$1\$ if the citizen's zipcode corresponds to your favorite planet.
- \$0\$ if the citizen has another zipcode.
 - 4. Train your logistic regression to predict if a citizen comes from your favorite planet or not, using your brand new label.
 - 5. Calculate and display the fraction of correct predictions over the total number of predictions based on the test set.
 - 6. Plot 3 scatter plots (one for each pair of citizen features) with the dataset and the final prediction of the model. You can use normalization on your dataset. The question is should you?

You now have a model that can discriminate between citizens that come from one specific planet and everyone else. It's a first step, a good one, but we still have work to do before we can classify citizens among four planets!

So how does Multiclass Logistic Regression work?

One Versus All

The idea now is to apply what is called **one-versus-all classification**.

It's quite straightforward: Your program (multi_log.py) will: 1. Split the dataset into a training and a test set. 2. Train 4 separate Logistic Regression classifiers to discriminate between each class and the others (the way you did in part one). 3. Predict for each example the class according to each classifiers and select the one with the highest output probability. 4. Calculate and display the fraction of correct predictions over the total number of predictions based on the test set. 5. Plot 3 scatter plots (one for each pair of citizen features) with the dataset and the final prediction of the model.

Example:

If a cititzen got the following classification probabilities:

- Planet 0 vs all: 0.38
- Planet 1 vs all: 0.51
- Planet 2 vs all: 0.12
- Planet 3 vs all: 0.89

Interlude - More Evaluation Metrics!

Once your classifier is trained, you want to evaluate its performance. You already know about *cross-entropy*, as you implemented it as your *cost function*. But when it comes to classification, there are more informative metrics we can use besides the loss function. Each metric focuses on different error types. But what is an error type?

A single classification prediction is either right or wrong, nothing in between. Either an object is assigned to the right class, or to the wrong class. When calculating performance scores for a multiclass classifier, we like to compute a separate score for each class that your classifier learned to discriminate (in a one-vs-all manner). In other words, for a given $Class\ A$, we want a score that can answer the question: "how good is the model at assigning A objects to $Class\ A$, and at NOT assigning non-A objects to $Class\ A$?"

You may not realize it yet, but this question involves measuring two very different error types, and the distinction is crucial.

Error Types

With respect to a given $Class\ A$, classification errors fall in two categories:

- False positive: when a *non-A* object is assigned to *Class A*. For example:
 - Pulling the fire alarm when there is no fire.
 - Considering that someone is sick when he/she isn't.
 - Identifying a face in an image when in fact it is a Teddy Bear.
- False negative: when an A object is assigned to another class than Class A. For example:
 - Not pulling the fire alarm when there is a fire.
 - Considering that someone is not sick when he/she is.
 - Failing to recognize a face in an image that does contain one.

It turns out that it's really hard to minimize both error types at the same time. At some point you'll need to decide which one is the most critical, depending on your use case. For example, if you want to detect cancer, of course it's not good if your model erroneously diagnoses cancer on a few healthy patients (false positives), but you absolutely want to avoid failing at diagnosing cancer on affected patients (false negatives) and let them go on with their lives while developing a potentially dangerous cancer.

Metrics

A metric is computed on a set of predictions along with the corresponding set of actual categories. The metric you choose will focus more or less on those two error types. If we come back to the ${f Class}$ ${f A}$ classifier:

- Accuracy: tells you the percentage of predictions that are accurate (i.e. the correct class was predicted). Accuracy doesn't give information about either error type.
- **Precision**: tells you how much you can trust your model when it says that an object belongs to *Class A*. More precisely, it is the percentage of the objects assigned to *Class A* that really were *A* objects. You use precision when you want to control for **False positives**.
- **Recall**: tells you how much you can trust that your model is able to recognize ALL *Class A* objects. It is the percentage of all **A** objects that were properly classified by the model as *Class A*. You use recall when you want to control for **False negatives**.
- **F1 score**: combines precision and recall in one single measure. You use the F1 score when want to control both **False positives** and **False negatives**.

Exercise 11 - Other metrics

Turn-in directory: ex11/

Files to turn in: other_metrics.py

Forbidden functions: None Remarks: n/a

Objective:

Understand and manipulation of classification criteria (TP, FP, ...) and metrics. The goal of this exercise is to write four metric functions (which are also available in **sklearn.metrics**) and to understand what they measure and how they are constructed.

You must implement the following fomulas:

$$\begin{aligned} \text{accuracy} &= \frac{\text{tp} + \text{tn}}{\text{tp} + \text{fp} + \text{tn} + \text{fn}} \\ \text{precision} &= \frac{\text{tp}}{\text{tp} + \text{fp}} \\ \text{recall} &= \frac{\text{tp}}{\text{tp} + \text{fn}} \\ \text{F1score} &= \frac{2 \times \text{precision} \times \text{recall}}{\text{precision} + \text{recall}} \end{aligned}$$

Where:

- tp is the number of true positives,
- fp is the number of false positives,
- tn is the number of true negatives,
- fn is the number of **false negatives**.

Instructions:

For the sake of simplicity, we will only ask you to use two parameters.

In the other_metrics.py file, write the following functions as per the instructions below:

```
... Your code ...
def recall_score_(y, y_hat, pos_label=1):
    ... Your code ...
def f1_score_(y, y_hat, pos_label=1):
    ... Your code ...
```

```
import numpy as np
from sklearn.metrics import accuracy_score, precision_score, recall_score, f1_score

# Example 1:
y_hat = np.array([1, 1, 0, 1, 0, 0, 1, 1])
y = np.array([1, 0, 0, 1, 0, 0])

# Accuracy
## your implementation
accuracy_score_(y, y_hat)
## Output:
0.5
## sklearn implementation
accuracy_score(y, y_hat)
```

```
precision_score_(y, y_hat)
precision_score(y, y_hat)
recall_score_(y, y_hat)
recall_score(y, y_hat)
f1_score_(y, y_hat)
f1_score(y, y_hat)
y_hat = np.array(['norminet', 'dog', 'norminet', 'norminet', 'dog', 'dog', 'dog', 'dog'])
y = np.array(['dog', 'dog', 'norminet', 'norminet', 'dog', 'norminet', 'dog', 'norminet'])
accuracy_score_(y, y_hat)
accuracy_score(y, y_hat)
precision_score_(y, y_hat, pos_label='dog')
precision_score(y, y_hat, pos_label='dog')
recall_score_(y, y_hat, pos_label='dog')
```

```
recall_score(y, y_hat, pos_label='dog')
f1_score_(y, y_hat, pos_label='dog')
f1_score(y, y_hat, pos_label='dog')
y_hat = np.array(['norminet', 'dog', 'norminet', 'norminet', 'dog', 'dog', 'dog', 'dog'])
y = np.array(['dog', 'dog', 'norminet', 'norminet', 'dog', 'norminet', 'dog', 'norminet'])
precision_score_(y, y_hat, pos_label='norminet')
precision_score(y, y_hat, pos_label='norminet')
recall_score_(y, y_hat, pos_label='norminet')
recall_score(y, y_hat, pos_label='norminet')
f1_score_(y, y_hat, pos_label='norminet')
f1_score(y, y_hat, pos_label='norminet')
```

Exercise 12 - Confusion Matrix

Turn-in directory: ex12/

Files to turn in: confusion_matrix.py

Forbidden functions: None Remarks: n/a

Objective:

Manipulation of confusion matrix concept. The goal of this exercise is to reimplement the function confusion_matrix available in sklearn.metrics and to learn what does the confusion matrix represent.

Instructions:

For the sake of simplicity, we will only ask you to use three parameters. Be careful to respect the order, true labels are rows and predicted labels are columns:

In the confusion_matrix.py file, write the following function as per the instructions below:

```
def confusion_matrix_(y_true, y_hat, labels=None):
    """
    Compute confusion matrix to evaluate the accuracy of a classification.
    Args:
        y_true:a numpy.ndarray for the correct labels
        y_hat:a numpy.ndarray for the predicted labels
        labels: optional, a list of labels to index the matrix. This may be used to reorder
    or select a subset of labels. (default=None)
    Returns:
        The confusion matrix as a numpy ndarray.
        None on any error.
    Raises:
        This function should not raise any Exception.
    """
    ... Your code ...
```

Optional part

Objective(s):

For a more visual version, you can add an option to your previous confusion_matrix_ function to return a pandas.DataFrame instead of a numpy array.

Instructions:

In the confusion_matrix.py file, write the following function as per the instructions below:

```
def confusion_matrix_(y, y_hat, labels=None, df_option=False):
    """"
    Compute confusion matrix to evaluate the accuracy of a classification.
    Args:
        y:a numpy.ndarray for the correct labels
        y_hat:a numpy.ndarray for the predicted labels
        labels: optional, a list of labels to index the matrix. This may be used to reorder
    or select a subset of labels. (default=None)
        df_option: optional, if set to True the function will return a pandas DataFrame
    instead of a numpy array. (default=False)
    Returns:
        The confusion matrix as a numpy ndarray or a pandas DataFrame according to df_option
    value.
        None on any error.
Raises:
        This function should not raise any Exception.
    """
        ... Your code ...
```



 $\mathbf{N.B:}$ If you fail this exercise on your first attempt, Norminet will curse you forever. Yeah, you'd better do it right or you are in trouble my friend, big trouble!