

Machine Learning - Assignment 1

Francesco Saverio Zuppichini

October 19, 2017

We decided to split the code in the following files in order to decouple the logic and create a better environment booth for developing and testing.

- `skeleton.py` Holds the original function such as `run_part1` and `run_part2`
- `Perceptron.py` Holds the Perceptron class
- `NeuralNetwork.py` Holds the NeuralNetwork class
- `BetterNeuralNetwork.py` Holds the BetterNeuralNetwork class
- `activation_function.py` Holds all the activation functions, eg. `tanh`
- `cost_functions.py`. Holds all the errors functions, eg. `MSE`
- `main.py` Main scripts where the functions are called
- `utils.py` A set of utils function
- `plots.py` Functions used to generate the plots in the report

By running the `main.py` you will get the results from `competition_load_weights_and_evaluate_X_and_T` by default. The whole project is hosted on GitHub at the following link:

<https://github.com/FrancescoSaverioZuppichini/Neural-Network-spiral-data-set>

1 The Perceptron

1.1 Question 1

1.1.1 Vectorized equations

Equation 1 shows the vectorised equation for a single perceptron

$$output = X \times W + b \tag{1}$$

Where $X = x_1, \dots, x_n$ is the input set and $W = w_1, \dots, w_n$ is the target set. We denote y the output of the perceptron.

1.1.2 Mean Squared Error

Equation 2 denotes the Mean Square Error function for our single perceptron

$$E(w) = \frac{1}{N} \sum_{i=1}^N (\underbrace{y(x_i, w_i)}_{\text{predicted}} - \underbrace{t_i}_{\text{actual}})^2 \quad (2)$$

One trick that is usual done is to multiply equation (2) by $\frac{1}{2}$, so when we take the derivative the 2 goes away. This is called One Half Mean Squared Error.

1.1.3 Derivate of the error with respect to the weights

In order to reduce our error function we need to compute the first derivative with respect to the weights.

$$\frac{\partial E}{\partial w_k} = \frac{\partial E}{\partial y_i} \frac{\partial y_i}{\partial w_k} = (w_k \cdot x_{ik} - t) \cdot x_{ik} \quad (3)$$

1.1.4 Compute the new weight values after one step using a learning rate of 0.02

Since it was not specified that the calculation must be done by hand, we called `train_one_step` in order to compute the new weights, as follows:

```
X,T = get_part1_data()
p = Perceptron()
sk.train_one_step(p,0.02,np.array([X[0]]),np.array([T[0]]))
print(p.var['W']) //output weights the values
print(p.var['b']) //output the new bias value
```

Where the new weights and bias are

$$W_{k+1} = \begin{pmatrix} 0.844 \\ -0.148 \end{pmatrix}, b_{k+1} = (2.044)$$

1.1.5 Gradient Descend

The gradient descend is an iterative optimisation algorithm that follows the direction of the negative gradient in order to minimise an objective function. It can be effectively used as Learning Algorithm because it reduces the error function, Equation 2, and adjusts the weights properly. Equation 4 shows the generic update rule.

$$w_{k+1} = w_k - \eta \nabla E(w_k) \quad (4)$$

Where η is the step size, also called **learning rate** in Machine Learning. This parameter influences the behaviour of gradient descent, a small number can lead to local minimum, while a bigger learning rate could "over-shoot" and decreasing the converge ration. Later in this project you will see how a wrong η can strongly change the output of a Neural Network.

For this reasons, numerous improvements have been proposed to avoid local minima and increase its convergence ration, some of them are: Conjugate Gradient and Momentum.

1.2 Implement the MSE and dMSE

You can find them in `cost_functions.py`

1.3 Implement the function forward and backward

The *forward* and *backward* function can be found in *Perceptron.py*. You can find *train_one_step* in *skeleton.py*.

1.4 Implement the *run_part1*

You can find the implementation in the code, *skeleton.py*. Figure 1 shows the final result after 15 steps using a learning rate of 0.02. We can notice that the Perceptron worked as expected since it correctly classified the two colour sets.

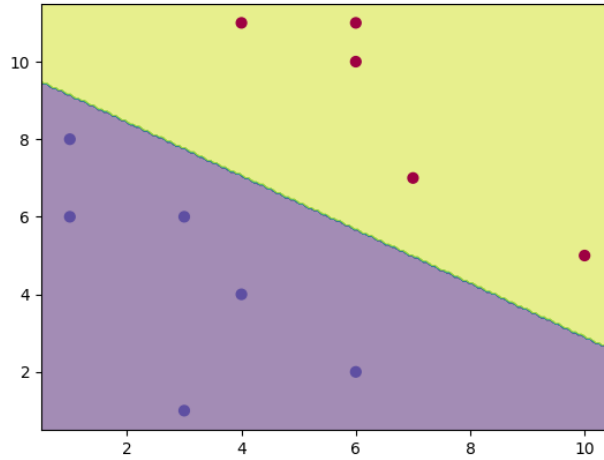


Figure 1: *run_part1* plot

2 A Neural Network

2.1 Implement the activation functions

You can find them inside *activation_functions.py*. I have also implemented the **Relu** that will be used in Section 3.

2.2 Question 2

2.2.1 Forward pass

In order to calculate the forward pass of a Neural Network we need to compute the activation of each layer, l , and use it as input of the next one until we reach the output layer.

Equation 5 shows the activation a of layer l for the j -th neuron on that layer.

$$a_j^l = \sigma\left(\sum_k w_{jk}^l a_k^{l-1} + b_j^l\right) \quad (5)$$

Where w_{jk}^l is the connection from neuron k in the $l-1$ layer to j , a_k^{l-1} is the activation of the previous layer and b_j^l is the bias of j -th neuron in the l layer. With this in mind, we can rewrite 5

in a efficient vectorised form

$$a^l = \sigma(W^l a^{l-1} + b^l) \quad (6)$$

2.2.2 Delta rules

In a Neural Network the weights are iteratively changed in order to decrease the error function, called E . Equation ?? defines δ_j^l as the output error of neuron j in layer l

$$\delta_j^l = \frac{\partial E}{\partial z_j^l} \quad (7)$$

Strictly speaking, δ_j^l , is how much the error function changes by changing the weighted input on that layer. Applying the chain rule, Equation ?? becomes:

$$\delta_j^l = \frac{\partial E}{\partial a_j^l} \frac{\partial a_j^l}{\partial z_j^l} \quad (8)$$

By knowing that $a_j^l = \sigma(z_j^l)$, Equation 8 can be expressed as

$$\delta_j^l = \frac{\partial E}{\partial a_j^l} \sigma'(z_j^l) \quad (9)$$

2.2.3 Derivatives of the weights

We want to compute $\frac{\partial E}{\partial w_{jk}^l}$. Applying the delta rule:

$$\frac{\partial E}{\partial w_{jk}^l} = \frac{\partial E}{\partial z_j^l} \frac{\partial z_j^l}{\partial w_{jk}^l} = \frac{\partial E}{\partial a_j^l} \frac{\partial a_j^l}{\partial z_j^l} \frac{\partial z_j^l}{\partial w_{jk}^l} \quad (10)$$

$$\frac{\partial E}{\partial w_{jk}^l} = a_k^{l-1} \delta_j^l \quad (11)$$

2.3 Implement the functions forward and backward of the Neural Network class.

You can find them in *NeuralNetwork.py*

2.4 Train Network

2.4.1 Split the data into a train set and a test set

We decided to split the training set and test set with a ratio of 80:20. If we choose a smaller train set we may under-train the network and, on the other hand, if we shrink the test set we may *overfit*. In order to do so, we created a function called `get_train_and_test_data` in *utils.py*.

Figure 2 shows the plot for the train set and the test set. They are created by calling `twospirals` using competition's configuration.

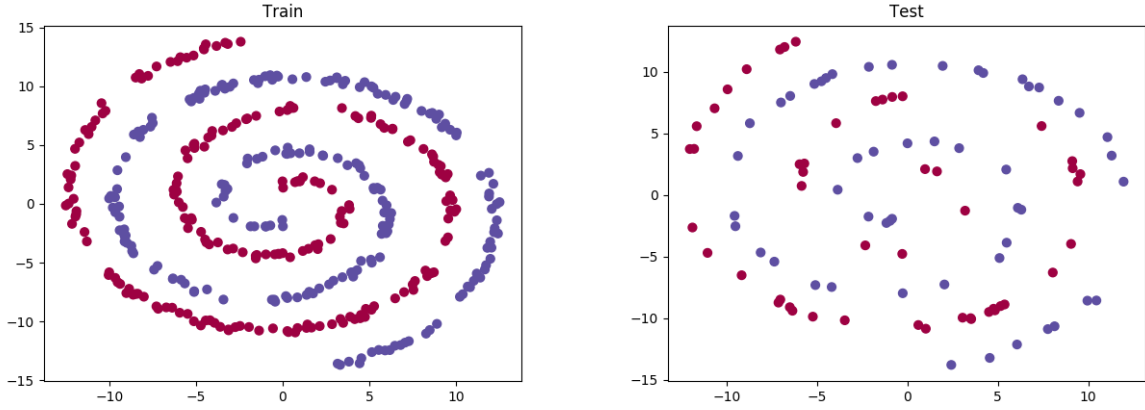


Figure 2: Train set and Test set

2.4.2 Initialise the weights randomly and find a good learning rate to train

In order to find the correct learning rate, we decided to first test the network with different sizes in order to have some empirical data. Figure 3 shows the convergence rate with five learning rates. The data was generated using the standard inputs and target from `twospirals` and then normalised the gradient with its average each 100 iterations in order to provide a smoothed line.

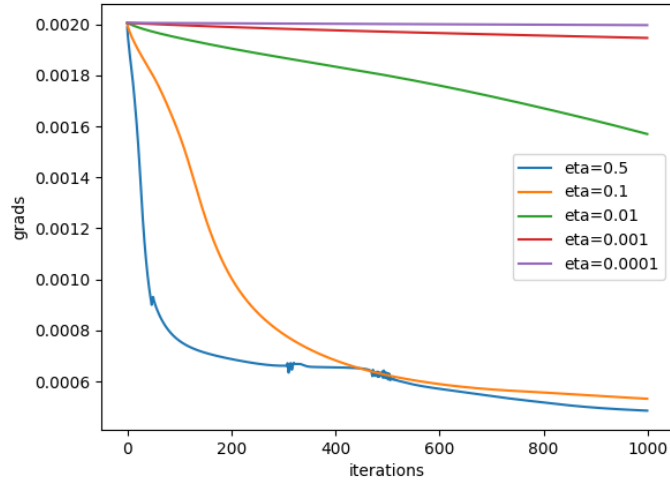


Figure 3: Learning rate convergences

The benchmark shows that 0.1 is the best η choice since it is the faster and more stable, 0.5 is also a good choice.

2.4.3 Plot the learning rate of at least 5 different learning rates on both sets

Since it was not clear what to plot, we decided to plot five boundaries for the five learning rates used in section 2.4.2. For the benchmark, we used 40.000 iterations that are more than enough to make the model converge to a good solution. Figure 4 shows the results

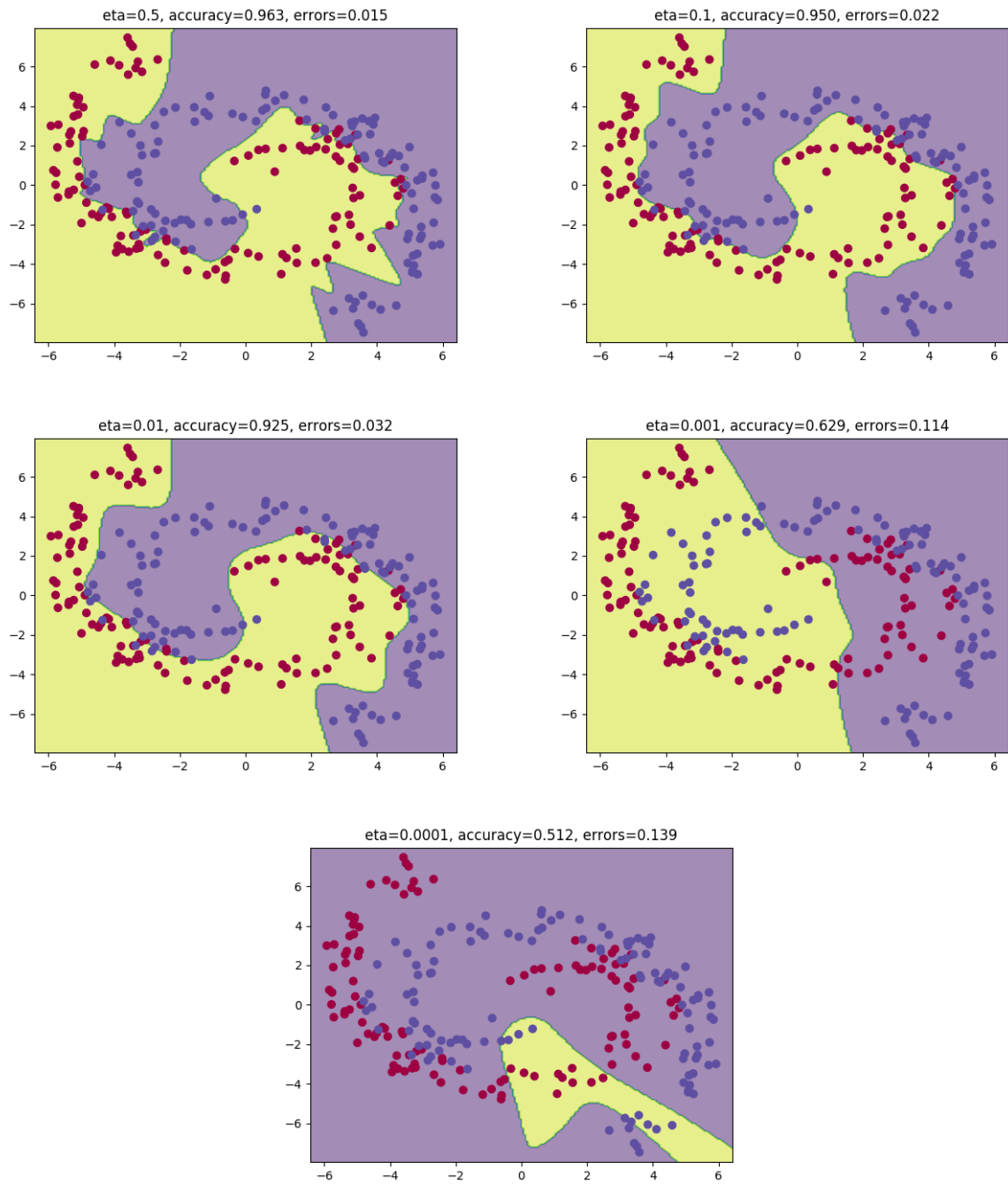


Figure 4: model boundary vs learning rates

Figure ?? shows different eta vs gradients and errors respectively

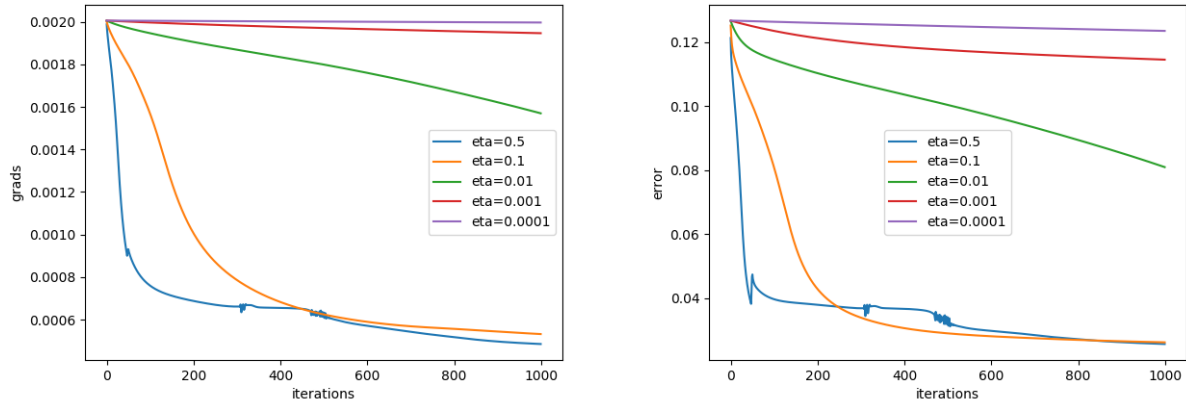


Figure 5: eta vs grads and costs

2.4.4 Plot the boundary of the model which converged the most on the training data

Figure 6 shows the model boundary after reach a MSE error less than 0.02 using a step size of 0.001. The network used the following seed: 1. You can get the same plot by calling `run.part2`.

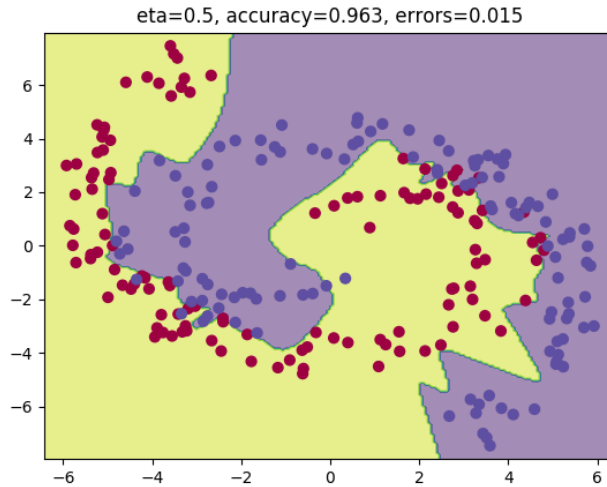


Figure 6: Boundary of the model with MSE less than 0.02

2.5 Explain if this is a good model for the data

The model seems good enough for the data generated by calling `twospirals` with the standard configuration. however, by using the data for part 3 the model loose lots of accuracy due to its gradient descent implementation that does not prevent the algorithm to fall into local minima. A better model is present in detail in the following Section.

3 Further Improvements

In this sections we analyse each extension that we tried, even if they did not improve our solution, in order to create a better model.

3.1 Generic Implementation

Since the `NeuralNetwork` class has a fixed number of layers, the first improvement that we did was to create a generic implementation.

The `BetterNeuralNetwork` class exposes an API allowing the creation of a generic Neural Network by adding and shaping layers.

After instance the class, is it possible to add a input layer by calling `add_input_layer(size_in,size_out)`. Many hidden layers can be added by using `add_hidden_layer(size_out)`, the "size_in" is not necessary since it can be easily calculated by the network. Then method `add_output_layer(size_out)` adds the output lauer. It follows a full example where we create the same Neural Network from Exercise 2.

```
bnn = BNN()
bnn.add_input_layer(2, 20, np.tanh, act.dtanh)
bnn.add_hidden_layer(15, np.tanh, act.dtanh)
bnn.add_output_layer(1)
```

For each layer is also possible to specified the activation function, as default value *sigmoid* is used. We have also implemented `relu` and `dRelu` to try them.

`BetterNeuralNetwork` exposes `train` method that performs forward, backpropagation and weight updates directly. It accepts as input the training sets, the targets, max iterations, a dictionary containing the parameters and the gradient descent update method name, the supported ones are: 'gradient_descent' (default), 'momentum' and 'adagrad'

3.2 Initial weight and bias

Weight initialisation is crucial to create a good model. I used the following formula.

$$w = random(n)/\sqrt{n}$$

Where n is the size of the input. This ensures that all neurons in the network initially have approximately the same output distribution and empirically improves the rate of convergence.¹

Using *numpy*:

```
np.random.randn(in_size,out_size)/np.sqrt(in_size)
```

The bias are initialised with a very small number close to zero, for such reason I scaled down the each random value by a factor of 100.

3.3 Performance

Due to a better and generic implementation, our `BetterNeuralNetwork` is slightly faster than the one from Section 2. Table 1 shows the result of a little benchmark

¹<http://cs231n.github.io/neural-networks-2/>

iterations	time NN	time BNN
10	17.5ms	13.7ms
100	134.5ms	128.2ms
1000	461.6ms	458.84ms
10000	4455.6ms	4157.9ms

Table 1

I have also tried to create a small thread pool, so we can save time in thread allocation, in order to parallelise the weight updates, since they are independent operation but without any real advantages. Probably the network is so small that it take more time to call a free thread and submit a work than to compute the operations sequentially.

3.4 Momentum

We used Momentum [2] to speed up the convergence of the network, it prevent getting stuck in local minima and pushes the objective more quickly along the shallow ravine. Basically it give a global direction to the gradient descent based on the previous update. Equation 12 shows the new update formula.

$$\begin{aligned}\Delta w_{t+1} &= (\eta \nabla E(w_{t+1}) + \beta \Delta w_t) \\ w_{k+1} &= w_{k+1} - \Delta w_{t+1}\end{aligned}\tag{12}$$

$\beta \in [0, 1]$ is a the momentum factor, usually between 0.5 and 0.9. In our benchmarks we used a classic 0.5. In Figure 7, we can see the converge rate of a classic gradient descent versus momentum for different step sizes. Momentum converges faster in all cases.

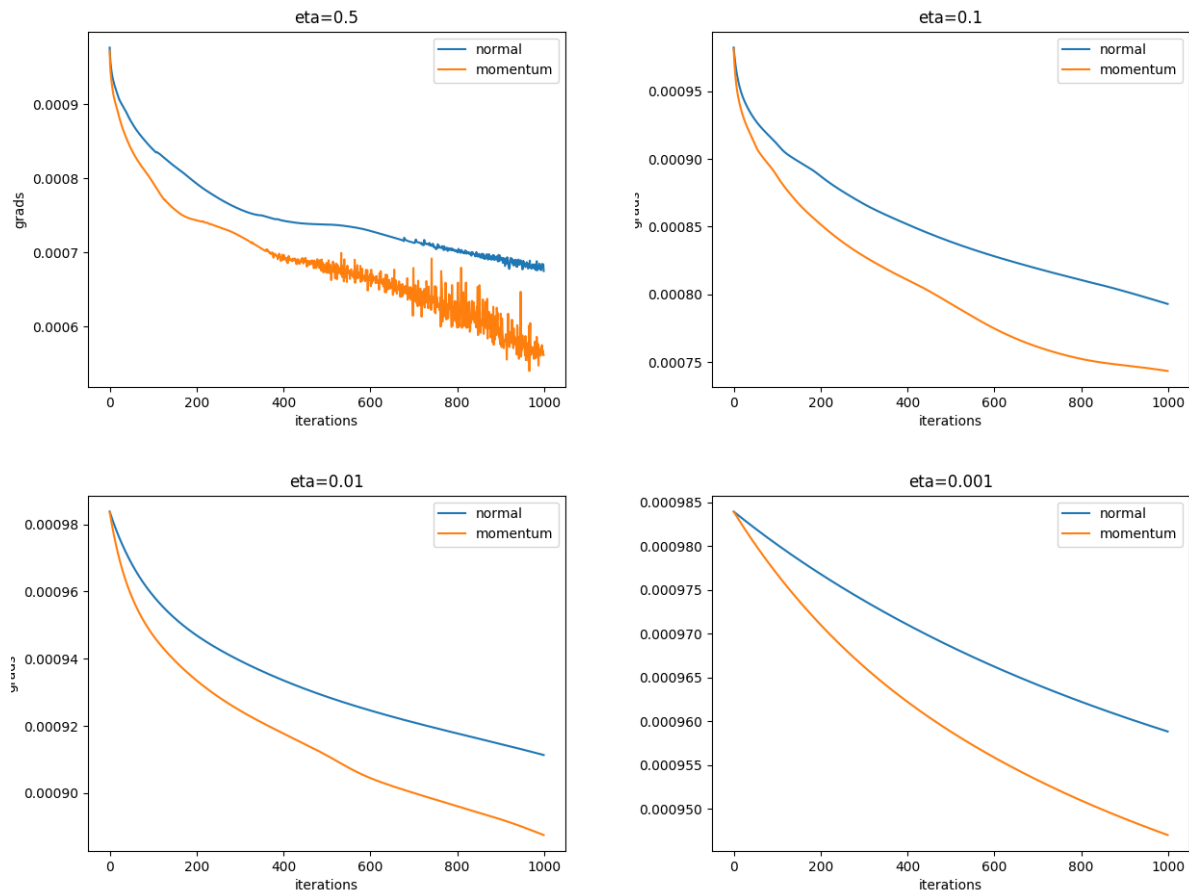


Figure 7: Gradient Descent vs Momentum

In the `BetterNeuralNetwork` class, momentum can be enabled by passing the correct parameters and type to the `train` method. The following snippet shows an example

```
params = {'eta': 0.01, 'beta': 0.5} # beta must be included or the default 0.5
                                     # value will be used
nn.train(X_train, T_train, 3000, params, 'momentum') # pass 'momentum' as type
                                                         # name
```

3.5 Adagrad

Adagrad [1] is an adaptive learning rate algorithm that we implement in order to speed up the convergence. In Figure 8, we can see the converge rate of a classic gradient descent versus adagrad for different step sizes.

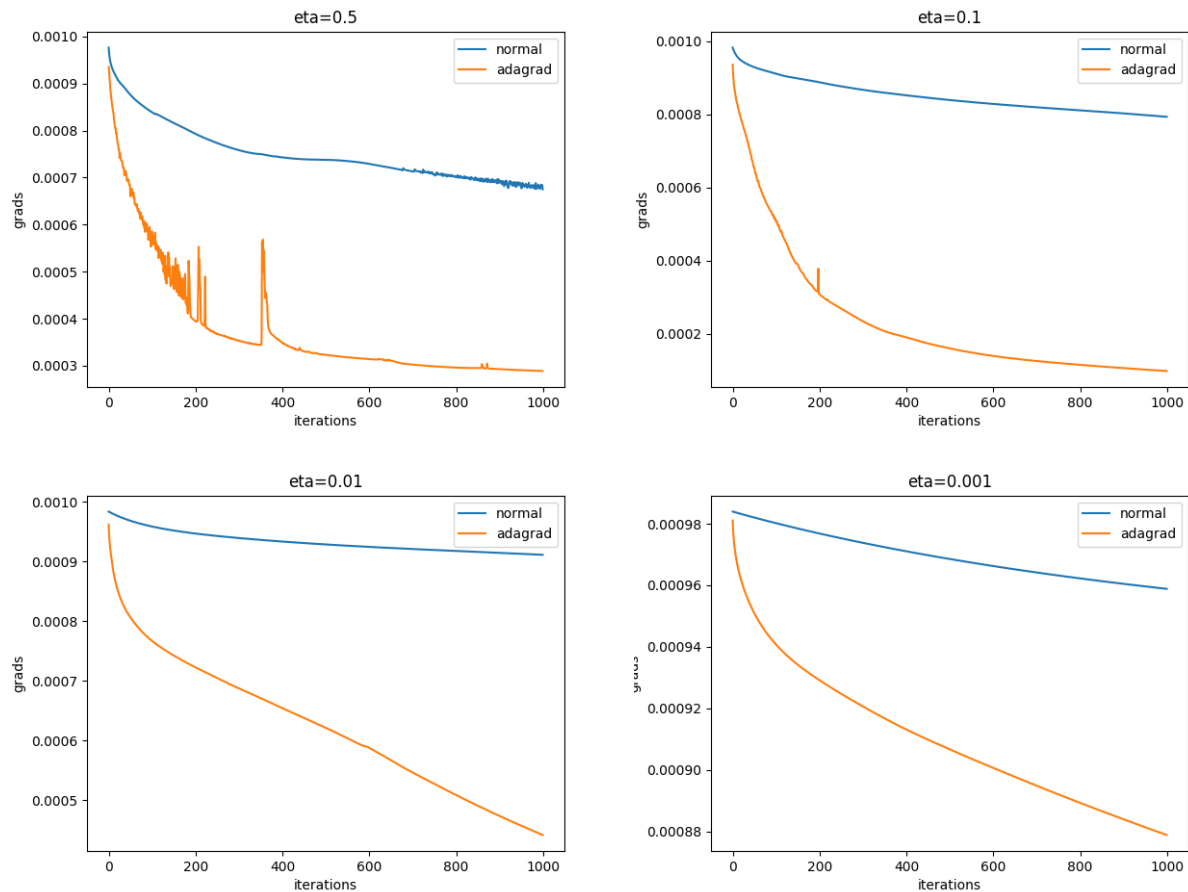


Figure 8: Gradient Descent vs Adagrad

Adagrads works better in any case. The following snippet shows how to enable it

```
nn.train(X_train, T_train, 3000, params, 'adagrad') // pass 'adagrad' as type name
```

3.6 Network deep

Our model is composed by one input layer, three hidden layer and one output layer. We used an empirical approach to find a correct number of hidden layer. We started by make the first layer bigger in order to make our model learn more features about the function. Then we added other three layers, each of them roughly one third smaller than the other. Doing so, our model can recognise more detail about the spiral.

After some tests, we switched to the **relu** activation function on each layer expect for the output where we used *tanh*. We trying other techniques with no luck. We also include them in this report.

3.7 Stochastic Gradient Descent

We implemented Stochastic Gradient Descent, it can be enabled by passing the `batch_size` value inside the `params` dictionary

```
model.train(train_X, train_T, 4000, { 'eta' : 0.1, 'beta' : 0.5, 'batch_size': 10
}, 'adagrad', testX, testT)
```

3.8 Dropout

Dropout is a recently introduced regularisation technique. Basically, each node has a probability to be enable on disable in each forward pass. We implemented it by changing the `forward` method inside `BetterNeuralNetwork`

```
def forward(self, inputs):
    self.A = [inputs]
    self.Z = []
    # dropout probability
    p = 0.5

    a = inputs

    for l in self.layers:
        z = a.dot(l.W) + l.b
        # create dropout mask
        u = (np.random.rand(*z.shape) < p) / p
        # apply mask
        z *= u
        a = l.activation(z)
        # store
        self.A.append(a)
        self.Z.append(z)

    return a
```

At each pass, a binary mask `u` is created and applied to the layer weighted output `z`. Unfortunately, this approach did not work well for us and lead to a bad convergence, probably because the network is too small.

3.9 Results

Figure 9 shows the boundary of our new model with one of the best seed: 0.

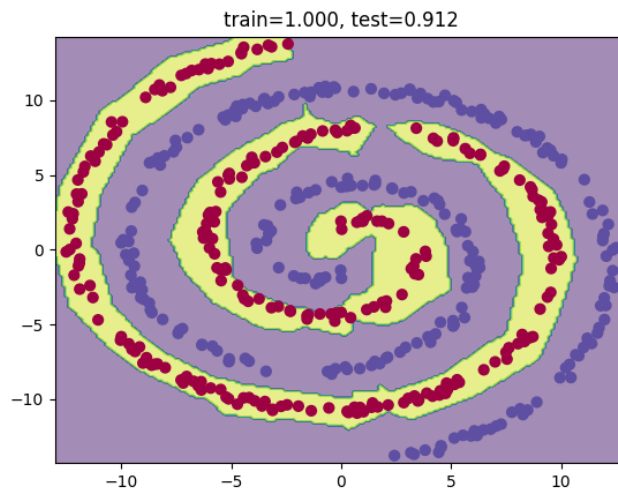


Figure 9: Boundary of the model

The following snippet show our to re-create the model with our API.

```

# use same seed
np.random.seed(0)
model = BetterNeuralNetwork()
# create layers
model.add_input_layer(2, 30, act.relu, act.drelu)
model.add_hidden_layer(20, act.relu, act.drelu)
model.add_hidden_layer(15, act.relu, act.drelu)
model.add_hidden_layer(10, act.relu, act.drelu)
model.add_output_layer(1, act.tanh, act.dtanh)

model.train(train_X, train_T , 4000, { 'eta' : 0.1, 'beta' : 0.5 }, 'adagrad')

```

Figure 10 shows the errors, gradients, error and accuracy for this model with respect to the one from section 2. Obviously the new model is better in every aspect.

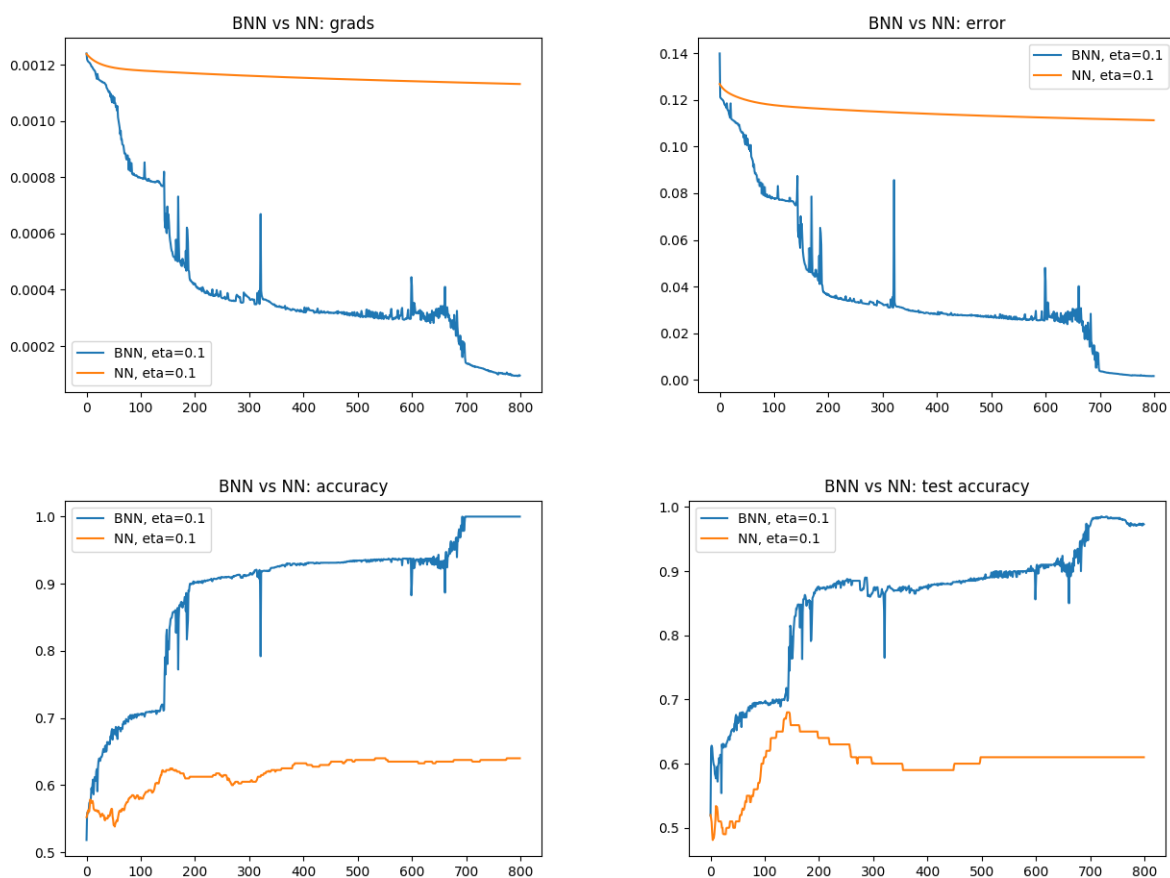
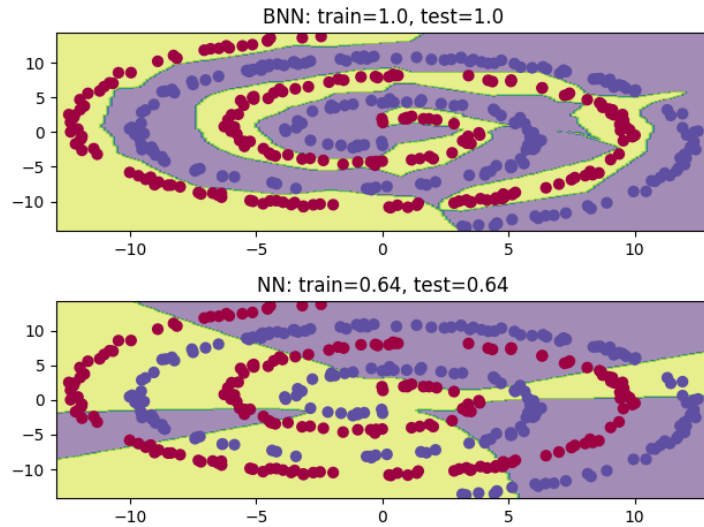


Figure 10: BetterNeuralNetwork vs Neural Network



3.10 Save and Load

You can save and load the network state by calling `.save(file_name)` `.load(file_name)`. The following snippet shows how to load the previous network.

```
model = BetterNeuralNetwork()
# create layers
model.add_input_layer(2, 30, act.relu, act.drelu)
model.add_hidden_layer(20, act.relu, act.drelu)
model.add_hidden_layer(15, act.relu, act.drelu)
model.add_hidden_layer(10, act.relu, act.drelu)
model.add_output_layer(1, act.tanh, act.dtanh)
# load prev weights
model.load('competition')
```

References

- [1] Elad Hazan John Duchi and Yoram Singer. *Adaptive Subgradient Methods for Online Learning and Stochastic Optimization*. 2011.
- [2] Sebastian Ruder. *An overview of gradient descent optimization algorithms*. 2017.