50.039 Theory and Practice of Deep Learning

W2-S1 Neural Networks, Initializers, Optimizers and other Good Practices

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About this week (Week 2)

- 1. What are the **typical initializers for trainable parameters** in Neural Networks?
- 2. What is **symmetry** in a Neural Network and why is it essential to **break it** with proper initializers?
- 3. What is the **exploding gradient** problem?
- 4. How to **spot and fix an exploding gradient** problem?
- 5. What is the **vanishing gradient** problem?
- 6. How to spot and fix a vanishing gradient problem?
- 7. Why are activation functions needed in Neural Networks?
- 8. What is the universal approximation theorem?

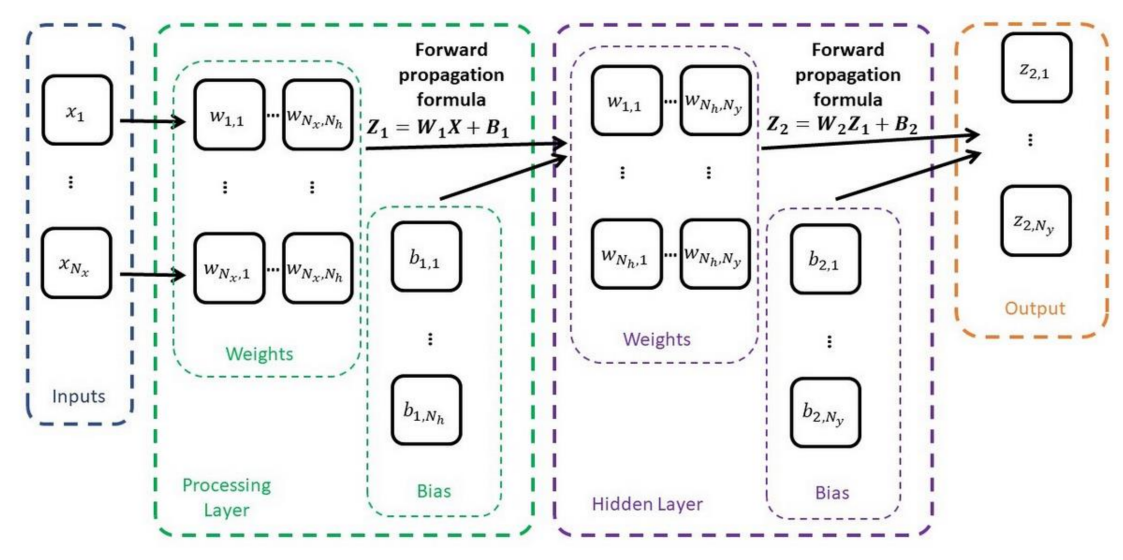
About this week (Week 2)

- 9. What are typical **examples** of **activation functions**? Which activations should I be using?
- 10. What are typical examples of **advanced optimizers**? (Adagrad, RMSProp, Adam, etc.).
- 11. How to **implement** said **optimizers** manually?
- 12. What is a **stochastic gradient descent** and what are its **benefits**?
- 13. How to implement the stochastic gradient descent manually?
- 14. What is a mini-batch gradient descent and what are its benefits?
- 15. How to implement the mini-batch gradient descent manually?
- 16. How to combine all optimizers concepts into a great optimizer?

About this week (Week 2)

- 17. What is the **no free lunch theorem?**
- 18. What is the **train-test-validation split**? Why is it **good practice** to use an extra validation set?
- 19. What is the **early stopping** of optimizer concept? Why is it **good practice** to use it?
- 20. What are **saver** and **loader** functions? Why is it **good practice** to use them?
- 21. What are **other common good practices** when it comes to Neural Networks?
- 22. How to decide on an appropriate number of layers and neurons?

Our shallow NN with two processing layers



Restricted

Our NN class

How it works:

- Weights and biases are now initialized as normal random with zero mean and variance 0.1.
- Forward method for making predictions using current parameters.
- Loss function for performance evaluation.

```
class ShallowNeuralNet():
        def init (self, n x, n h, n y):
            # Network dimensions
            self.n x = n x
 5
            self.nh = nh
 6
            self.n y = n y
            # Weights and biases matrices
 8
            self.W1 = np.random.randn(n x, n h)*0.1
 9
10
            self.b1 = np.random.randn(1, n h)*0.1
11
            self.W2 = np.random.randn(n h, n y)*0.1
12
            self.b2 = np.random.randn(1, n y)*0.1
13
           # Loss, initialized as infinity before first calculation is made
            self.loss = float("Inf")
14
15
16
        def forward(self, inputs):
           # Wx + b operation for the first layer
17
18
           Z1 = np.matmul(inputs, self.W1)
           Z1 b = Z1 + self.b1
19
           # Wx + b operation for the second layer
20
21
           Z2 = np.matmul(Z1_b, self.W2)
           Z2_b = Z2 + self.b2
22
23
            return Z2 b
24
25
        def MSE_loss(self, inputs, outputs):
            # MSE loss function as before
26
            outputs re = outputs.reshape(-1, 1)
27
            pred = self.forward(inputs)
28
29
           losses = (pred - outputs re)**2
            self.loss = np.sum(losses)/outputs.shape[0]
30
            return self.loss
31
```

```
1 # Define neural network structure
  2 n x = 2
  3 n h = 4
  4 | n y = 1
  5 shallow neural net = ShallowNeuralNet(n_x, n_h, n_y)
    print(shallow neural net. dict )
 {'n x': 2, 'n h': 4, 'n y': 1, 'W1': array([[-0.10476816, 0.18570216, 0.03204007, -0.10951262],
       [-0.13867874, -0.03539496, -0.02856421, 0.20592501]]), 'b1': array([[ 0.0232776 , -0.16122469, 0.00718537, 0.0666335
 1]]), 'W2': array([[ 0.03321156],
       [-0.0336505],
       [ 0.04977554],
       [-0.1794089 ]]), 'b2': array([[0.03460341]]), 'loss': inf}
 1 pred = shallow neural net.forward(inputs)
                                                                       1. Initialize model
 2 print(pred.shape)
 3 print(outputs.shape)
                                                                       2. Forward to predict
 4 print(pred[0:5])
  print(outputs[0:5])
                                                                       3. Compute loss to evaluate model (it is
(100, 1)
                                                                          bad, because trainable parameters
(100, 1)
[[-23.24055489]
                                                                          have received random values!)
[-31.54945952]
[-12.75105332]
[ -2.49026451]
[-32.3803654 ]]
                                                                    loss = shallow neural net.MSE loss(inputs, outputs)
[[1.581913]
                                                                    print(loss)
[3.450274]
[2.978769]
                                                                 677.625448852107
[2.808258]
[2.556398]]
```

Restricted

Backpropagation in our model

Update rules for W_2 and b_2

(Obtained after using chain rule to compute derivatives wrt. parameters for loss)

$$W_2 \leftarrow W_2 - \frac{2\epsilon\alpha}{M}(W_1X + b_1)$$

$$b_2 \leftarrow b_2 - \frac{2\epsilon\alpha}{M}$$

```
def backward(self, inputs, outputs, alpha = 1e-5):
    # Get the number of samples in dataset
    m = inputs.shape[0]
    # Forward propagate
    Z1 = np.matmul(inputs, self.W1)
    Z1 b = Z1 + self.b1
    Z2 = np.matmul(Z1_b, self.W2)
    y pred = Z2 + self.b2
   # Compute error term
    epsilon = y pred - outputs
    # Compute the gradient for W2 and b2
    dL_dW2 = (2/m)*np.matmul(Z1_b.T, epsilon)
    dL_db2 = (2/m)*np.sum(epsilon, axis = 0, keepdims = True)
    # Compute the loss derivative with respect to the first layer
    dL dZ1 = np.matmul(epsilon, self.W2.T)
```

Backpropagation in our model

Update rules for W_1 and b_1

(Obtained after using chain rule to compute derivatives wrt. parameters for loss)

$$W_1 \leftarrow W_1 - \frac{2\epsilon\alpha}{M} W_2 X$$

$$b_1 \leftarrow b_1 - \frac{2\epsilon\alpha}{M} W_2$$

```
# Compute the loss derivative with respect to the first layer
dL dZ1 = np.matmul(epsilon, self.W2.T)
# Compute the gradient for W1 and b1
dL dW1 = (2/m)*np.matmul(inputs.T, dL dZ1)
dL_db1 = (2/m)*np.sum(dL_dZ1, axis = 0, keepdims = True)
# Update the weights and biases using gradient descent
self.W1 -= alpha*dL dW1
self.b1 -= alpha*dL db1
self.W2 -= alpha*dL_dW2
self.b2 -= alpha*dL db2
# Update Loss
self.MSE_loss(inputs, outputs)
```

Training procedure, in short.

Forward pass on dataset samples.
Compute errors and loss function.

Repeat until convergence or max number of iterations, as in GD earlier.

Use new loss value to compute gradients.
Adjust parameters with backprop.

Restricted

Made that iterative process a trainer method for our class.

```
66
        def train(self, inputs, outputs, N_max = 1000, alpha = 1e-5, delta = 1e-5, display = True):
             # List of losses, starts with the current loss
67
             self.losses list = [self.loss]
 68
             # Repeat iterations
 69
70
             for iteration number in range(1, N max + 1):
71
                 # Backpropagate
72
                 self.backward(inputs, outputs, alpha)
                 new loss = self.loss
73
74
                 # Update losses list
                 self.losses_list.append(new_loss)
75
                # Display
76
 77
                if(display):
 78
                     print("Iteration {} - Loss = {}".format(iteration_number, new_loss))
79
                 # Check for delta value and early stop criterion
                 difference = abs(self.losses list[-1] - self.losses list[-2])
 80
 81
                if(difference < delta):</pre>
 82
                     if(display):
83
                         print("Stopping early - loss evolution was less than delta.")
 84
                     break
 85
             else:
                 # Else on for loop will execute if break did not trigger
 86
87
                if(display):
                     print("Stopping - Maximal number of iterations reached.")
 88
89
90
        def show losses over training(self):
91
             # Initialize matplotlib
92
            fig, axs = plt.subplots(1, 2, figsize = (15, 5))
             axs[0].plot(list(range(len(self.losses_list))), self.losses_list)
93
             axs[0].set xlabel("Iteration number")
 94
95
             axs[0].set_ylabel("Loss")
             axs[1].plot(list(range(len(self.losses_list))), self.losses_list)
 96
97
             axs[1].set xlabel("Iteration number")
             axs[1].set_ylabel("Loss (in logarithmic scale)")
 98
             axs[1].set_yscale("log")
99
            # Display
100
             plt.show()
101
```

Trainer function for our model

Iteration 15 - Loss = 2.60947364178999
Iteration 16 - Loss = 2.6021864900028118
Iteration 17 - Loss = 2.5949494536811937
Iteration 18 - Loss = 2.58775401064343

T + anation 10 - loss - 2 5805050207014652

```
losses list = train(shallow neural net, inputs, outputs, N max = 10000, alpha = 1e-5, delta = 1e-6, display = True)
Iteration 1 - Loss = 4.639845655363769
Iteration 2 - Loss = 3.6055739235388646
Iteration 3 - Loss = 3.1219544752264055
Iteration 4 - Loss = 2.891370725402625
Iteration 5 - Loss = 2.778663798354082
Iteration 6 - Loss = 2.721333641596964
Iteration 7 - Loss = 2.6901646643755135
Iteration 8 - Loss = 2.671407441095785
Iteration 9 - Loss = 2.6585626629544303
Iteration 10 - Loss = 2.648548914291606
Iteration 11 - Loss = 2.639902127093024
                                                              3
Iteration 12 - Loss = 2.6319255476288577
Iteration 13 - Loss = 2.6242871305644138
Iteration 14 - Loss = 2.6168284145839156
```

250

500

750

1000

Iteration number

1250

1500

1750

A bad initialization

In our previous version of the model, we initialized our parameters randomly.

- While it seemed unimportant, this is actually essential.
- Initializing all parameters as identical constants (e.g. setting all to zeroes) would in fact be a terrible idea.
- But we have to try it at least once to understand why.

```
class ShallowNeuralNet():

def __init__(self, n_x, n_h, n_y):
    # Network dimensions
    self.n_x = n_x
    self.n_h = n_h
    self.n_y = n_y
    # Weights and biases matrices (randomly initialized)
    self.W1 = np.random.randn(n_x, n_h)*0.1
    self.b1 = np.random.randn(1, n_h)*0.1
    self.W2 = np.random.randn(n_h, n_y)*0.1
    self.b2 = np.random.randn(1, n_y)*0.1
# Loss, initialized as infinity before first calculation is made
    self.loss = float("Inf")
```

```
class ShallowNeuralNet_ConstantInit():

def __init__(self, n_x, n_h, n_y, const_val):
    # Network dimensions
    self.n_x = n_x
    self.n_h = n_h
    self.n_y = n_y
    # Weights and biases matrices (all initialized as constant
    # matrices filled with 0.1 values)
    self.W1 = np.ones(shape = (n_x, n_h))*const_val
    self.b1 = np.ones(shape = (1, n_h))*const_val
    self.W2 = np.ones(shape = (n_h, n_y))*const_val
    self.b2 = np.ones(shape = (1, n_y))*const_val
    # Loss, initialized as infinity before first calculation is made
    self.loss = float("Inf")
```

```
1 # Define and train neural network structure
 2 n x = 2
 3 n h = 4
 4 \, | \, n \, v = 1
 5 np.random.seed(37)
 6 | shallow neural net = ShallowNeuralNet ConstantInit(n x, n h, n y, 0.1)
 7 shallow_neural_net.train(inputs, outputs, N_max = 10000, alpha = 1e-6, delta = 1e-6, display = True)
Iteration 1 - Loss = 618.4538740049891
W1, b1, W2, b2:
[[0.09938446 0.09938446 0.09938446 0.09938446]
 [0.09676805 0.09676805 0.09676805 0.09676805]]
[[0.09999521 0.09999521 0.09999521 0.09999521]]
[[0.09614772]
 [0.09614772]
 [0.09614772]
 [0.09614772]]
[[0.09995206]]
Iteration 501 - Loss = 1.3726340052732087
W1, b1, W2, b2:
[[0.09064391 0.09064391 0.09064391 0.09064391]
 [0.04434246 0.04434246 0.04434246 0.04434246]]
[[0.09992629 0.09992629 0.09992629 0.09992629]]
[[0.0145902]
 [0.0145902]
 [0.0145902]
 [0.0145902]]
```

```
1 # Define and train neural network structure
 2 n x = 2
 3 n h = 4
 4 \mid n \mid v = 1
 5 np.random.seed(37)
 6 | shallow neural net = ShallowNeuralNet ConstantInit(n x, n h, n y, 0.1)
    shallow neural net.train(inputs, outputs, N_max = 10000, alpha = 1e-6, delta = 1e-6, display = True)
Iteration 1 - Loss = 618.4538740049891
W1, b1, W2, b2:
[[0.09938446 | 0.09938446 | 0.09938446 | 0.09938446]
                                                                  Same values on all groups of weights
 [0.09676805 | 0.09676805 | 0.09676805 | 0.09676805 | 1
[[0.09999521 0.09999521 0.09999521 0.09999521]]
[[0.09614772]
 [0.09614772]
 [0.09614772]
 [0.09614772]]
[[0.09995206]]
Iteration 501 - Loss = 1.3726340052732087
W1, b1, W2, b2:
[[0.09064391 0.09064391 0.09064391 0.09064391]
 [0.04434246 0.04434246 0.04434246 0.04434246]]
[[0.09992629 0.09992629 0.09992629 0.09992629]]
[[0.0145902]
 [0.0145902]
 [0.0145902]
 [0.0145902]]
```

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Iteration 1 - Loss = 618.4538740049891
W1, b1, W2, b2:
[[0.09938446 | 0.09938446 | 0.09938446 | 0.09938446]
                                                                Same values on all groups of weights
 [0.09676805 0.09676805 0.09676805 0.09676805]
[ 0.09999521 0.09999521 0.09999521 0.09999521 ]
                                                                Same thing for bias vector
[[0.09614772]
 [0.09614772]
 [0.09614772]
 [0.09614772]]
[[0.09995206]]
Iteration 501 - Loss = 1.3726340052732087
W1, b1, W2, b2:
[[0.09064391 0.09064391 0.09064391 0.09064391]
 [0.04434246 0.04434246 0.04434246 0.04434246]]
[[0.09992629 0.09992629 0.09992629 0.09992629]]
[[0.0145902]
 [0.0145902]
 [0.0145902]
 [0.0145902]]
```

```
# Define and train neural network structure
  2 n x = 2
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 5 np.random.seed(37)
 6 shallow neural net = ShallowNeuralNet ConstantInit(n x, n h, n y, 0.1)
    shallow neural net.train(inputs, outputs, N max = 10000, alpha = 1e-6, delta = 1e-6, display = True)
Iteration 1 - Loss = 618.4538740049891
W1, b1, W2, b2:
[[0.09938446 | 0.09938446 | 0.09938446 | 0.09938446]
                                                                Same values on all groups of weights
 <u>[0.09676805 | 0.09676805 | 0.09676805 | 0.09676805 | 1</u>
[ 0.09999521 0.09999521 0.09999521 0.09999521 ]
                                                                Same thing for bias vector
[[0.09614772]
 [0.09614772]
 [0.09614772]
 [0.09614772]]
Iteration 501 - Loss = 1.3726340052732087
                                                             Keeps happening without discontinuity no matter
W1, b1, W2, b2:
[[0.09064391 0.09064391 0.09064391 0.09064391]
                                                             the number of iterations...
 [0.04434246 0.04434246 0.04434246 0.04434246]]
                                                             This means all the neurons in a given layer are doing
[[0.09992629 0.09992629 0.09992629 0.09992629]]
[[0.0145902]
                                                             the same thing! (a.k.a. symmetry in the neural
 [0.0145902]
 [0.0145902]
                                                             network)
```

[0.0145902]

Symmetry in the Neural Network

Definition (Symmetry):

Symmetry in a neural network is the tendency of all neurons to have the same weights and processes.

As a **lack of diversity**, this can lead to a **lack of generalization** and can prevent the NN from learning complex patterns.

Additionally, symmetrical neural networks can be vulnerable to adversarial attacks (on Week 5).

```
Iteration 1 - Loss = 618.4538740049891
W1, b1, W2, b2:
0.09938446 0.09938446 0.09938446 0.09938446
 0.09676805 0.09676805 0.09676805 0.0967680
[ 0.09999521 0.09999521 0.09999521 0.09
[[0.09614772]
 [0.09614772]
 [0.09614772]
 [0.09614772]]
[[0.09995206]]
Iteration 501 - Loss = 1.3726340052732087
W1, b1, W2, b2:
[[0.09064391 0.09064391 0.09064391 0.09064391]
 [0.04434246 0.04434246 0.04434246 0.04434246]]
[[0.09992629 0.09992629 0.09992629 0.09992629]]
[[0.0145902]
 [0.0145902]
 [0.0145902]
```

Symmetry in the Neural Network

Definition (Symmetry):

Symmetry in a neural network is the tendency of all neurons to have the same weights and processes.

As a **lack of diversity**, this can lead to a **lack of generalization** and can prevent the NN from learning complex patterns.

Additionally, symmetrical neural networks can be vulnerable to adversarial attacks (on Week 5).

- Essentially, this happened because all weights and biases had the same starting point (due to the constant initialization).
- The backward process then updated the parameters identically, and they end up keeping the same values over the course of training.
- Need to enforce diversity on these parameters from the start! Hence, random initialization.

The random normal initialization

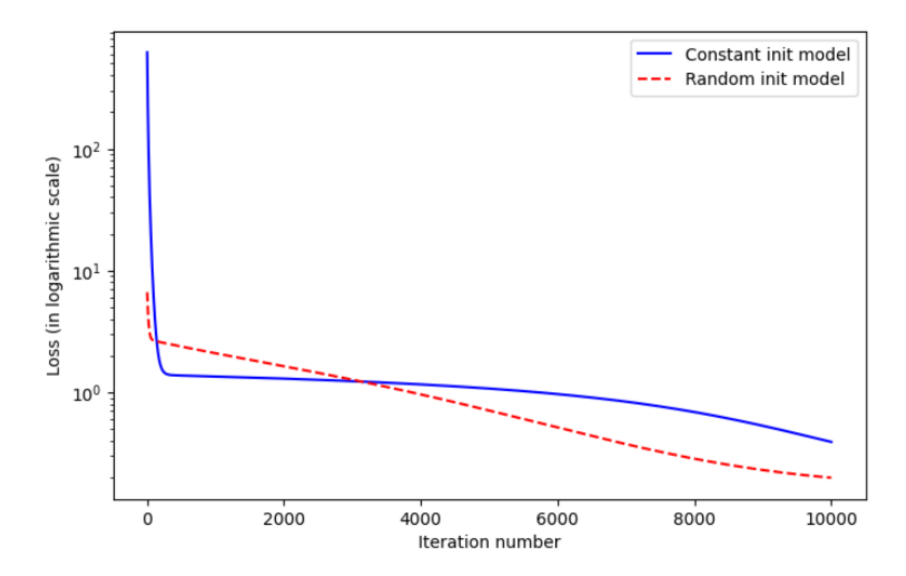
Definition (the random normal initialization [LeCun1998]):

The normal random initializer will setup parameters randomly, by sampling from a normal distribution, with a mean 0 and a standard deviation of 1 (or lower).

This initializer is **pretty straightforward** but can lead to **slow convergence or even divergence as the network gets deeper**.

```
def init_parameters_normal(self):
    # Weights and biases matrices (randomly initialized)
    self.W1 = np.random.randn(self.n_x, self.n_h)*0.1
    self.b1 = np.random.randn(1, self.n_h)*0.1
    self.W2 = np.random.randn(self.n_h, self.n_y)*0.1
    self.b2 = np.random.randn(1, self.n_y)*0.1
```

Using a random initialization is just better



The Xavier initialization

Definition (the Xavier initialization [Glorot2010]):

The **Xavier initializer** is based on a **Gaussian distribution**, with parameters initialized to zero-mean and a **variance** adjusted to the number of input/output parameters of the NN (e.g. $\frac{2}{N_x+N_y}$, with N_x being the input size and N_v the output size).

It is an overall good initializer for most architectures.

```
def init_parameters_xavier(self):
    # Weights and biases matrices (Xavier initialized)
    var = np.sqrt(2.0/(self.n_x + self.n_y))
    self.W1 = np.random.randn(self.n_x, self.n_h)*var
    self.b1 = np.random.randn(1, self.n_h)*var
    self.W2 = np.random.randn(self.n_h, self.n_y)*var
    self.b2 = np.random.randn(1, self.n_y)*var
```

Initializations variations and more

Many more initialization formulas exist:

- Xavier has random uniform and normal variations,
- Glorot proposed more initializations, same for He,
- Orthogonal initializations are sometimes useful but rare,
- Etc.

PyTorch has listed a lot of them, ready to use: https://pytorch.org/docs/stable/nn.init.html
(Have a look at bonus slides if curious!)

No free lunch theorem

Very important note: What I listed earlier are empirical observations only, certainly not absolute rules.

- For this reason, it is a good idea to try a few different initialization techniques and see which one works best for your problem.
- Some recent research has also suggested that using a combination of different initialization techniques (e.g., Xavier initialization for some layers and He initialization for others) can improve performance.
- Also, careful: Every now and then, we get a "smart*ss" claiming to have found a "ground-breaking new initializer beating all others".
- In reality, there is something that the machine learning community likes to call the "No free lunch theorem".

No free lunch theorem

Definition (the no free lunch (or NFL) theorem):

The no free lunch (NFL) theorem for supervised machine learning is a theorem that essentially implies that no single machine learning algorithm or tool is universally the best-performing algorithm for all problems.

(https://en.wikipedia.org/wiki/No free lunch theorem).



When in doubt about a supposedly "ground-breaking new technique beating all other techniques", **try them out**.

No free lunch theorem – season 1

Definition (the no free lunch (or NFL) theorem – season 1):

The no free lunch theorem then implies that there is no such thing "one initializer that works best for all models and problems".

No choice: try some of them and see how it goes!

(A good read on this matter, also: https://www.deeplearning.ai/ai-notes/initialization/index.html)



When in doubt about a supposedly "ground-breaking new technique beating all other techniques", **try them out**.

A weird initializer and a new problem

Definition (our custom "weird" initializer):

Let us consider the "weird" initializer below. It uses a random uniform distribution, scaling parameters based on the input and output sizes, like in Xavier.

We could in fact see this as a random uniform variation of Xavier.

And yet, when we use it, something weird happens.

```
def init_parameters_weird(self):
    # Weights and biases matrices (Weird? initialized)
init_val = np.sqrt(6.0/(self.n_x + self.n_y))
self.W1 = np.random.uniform(-init_val, init_val, (self.n_x, self.n_h))
self.b1 = np.random.uniform(-init_val, init_val, (1, self.n_h))
self.W2 = np.random.uniform(-init_val, init_val, (self.n_h, self.n_y))
self.b2 = np.random.uniform(-init_val, init_val, (1, self.n_y))
```

```
# Define and train neural network structure (Weird initialization)
   n x = 2
    n h = 4
    ny = 1
    init type = "Weird"
   np.random.seed(37)
   shallow_neural_net_weird = ShallowNeuralNet(n_x, n_h, n_y, init_type)
   # Train and show final loss
   shallow neural net weird.train(inputs, outputs, N max = 100, alpha = 1e-6, delta = 1e-6, display = True)
Gradients:
[[ 0.0347825 -0.28526197 -0.24439365 0.18661634]
 [ 0.17610429 -1.44428536 -1.23736846  0.94484113]]
[[ 0.00026903 -0.00220639 -0.00189029  0.0014434 ]]
[[-0.64676422]
 [-0.55320161]
 [ 1.42348809]
 [-0.81598256]]
[[-0.00168314]]
- Parameters:
[[ 1.25722625 -0.1015457 -0.86890687 0.23163369]
                                                             Wow, that's a big loss value!?
 0.33964944 0.52106421 -1.12164798
                                     0.69431032]]
[[-0.61665631 0.71679298 0.82789654 0.3603433 ]]
[[-0.15983769]
 [ 1.31087797]
 [ 1.12307381]
 [-0.85756698]]
[[0.27139079]]
Iteration 1 - Loss = 5638943.191688167
```

Restricted

```
# Define and train neural network structure (Weird initialization)
   n x = 2
   nh = 4
   ny = 1
   init type = "Weird"
 6 np.random.seed(37)
   shallow neural_net_weird = ShallowNeuralNet(n_x, n_h, n_y, init_type)
   # Train and show final loss
   shallow neural net weird.train(inputs, outputs, N max = 100, alpha = 1e-6, delta = 1e-6, display = True)
Iteration 26 - Loss = nan
- Gradients:
[[nan nan nan nan]
[nan nan nan nan]]
[[nan nan nan nan]]
[[nan]
[nan]
                                   And after iteration 26, we are getting NaNs values
 [nan]
[nan]]
                                   (Not a Number values) all over the place...?!
[[nan]]
- Parameters:
                                   What is going on...?!
[[nan nan nan nan]
[nan nan nan nan]]
[[nan nan nan nan]]
[[nan]
[nan]
 [nan]
[nan]]
[[nan]]
```

The exploding gradient problem

Definition (exploding gradients):

These **NaN** values are a typical symptom for a phenomenon called **exploding gradients**.

This typically occurs when the gradient descent rule has changes (in $\alpha \frac{\partial dL}{\partial W1}$ for instance) that are far greater than the current values in the matrices (e.g. W_1).

The values will explode to infinity and eventually become NaNs.

This typically happens with

- unlucky initializations,
- when the learning rate α is too large for the given parameters,
- and sometimes when gradients are approximated with formulas that are a bit buggish...

In order to avoid this problem, many techniques have been developed (something for later).

Unlucky initialization

Definition (Unlucky Initialization):

Unlucky Initialization is a typical example used to describe that a system's behaviour is impossible to predict due to the random impact of its initial conditions.

In some cases, the result of a training will be **drastically different** depending on the initial values used, which can lead to **vastly different outcomes for your model**.

In fact, in our previous "weird" initializer, we can resolve the NaN problem...

By just using a different seed for the initialization...!

Yup, as simple as that!

(BTW, that could be an empirical proof that the NFL theorem is indeed true!)

```
1 # Define and train neural network structure (Weird initialization)
 2 n x = 2
 3 n h = 4
 4 \, n \, y = 1
 5 init type = "Weird"
 6 np.random.seed(17)
 7 shallow neural net weird2 = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Train and show final loss
 9 shallow neural net weird2.train(inputs, outputs, N max = 500, alpha = 1e-6, delta = 1e-6, display = True)
- Gradients:
[[-9.86521735e-08 -7.36853902e-08 -8.11338502e-08 1.17729102e-07]
[-3.95240319e-08 -2.95213335e-08 -3.25054864e-08 4.71670175e-08]]
[[-3.87197953e-08 -2.89206322e-08 -3.18440634e-08 4.62072611e-08]]
[[ 4.10683166e-07]
[-1.03823853e-07]
[ 3.91291750e-07]
 [ 7.42268171e-07]]
[[-1.96398395e-07]]
                                                                           Different seed...

    Parameters:

[[-0.57336145 0.13297469 -1.0271704 -1.19171781]
[ 0.65980609  0.57963967 -0.6216372  0.49082227]]
                                                                           Now that works?!
[[-1.30374448 -0.40183954 1.25928682 -1.24410091]]
[[ 0.19714925]
[ 0.14725493]
[ 0.16214014]
[-0.23527311]]
[[0.14792844]]
Stopping early - loss evolution was less than beta on iteration 342.
 1 # Showing final loss
```

```
0.12422436714449628
```

2 print(shallow_neural_net_weird2.loss)

The learning rate tradeoff

Definition (Learning rate tradeoff):

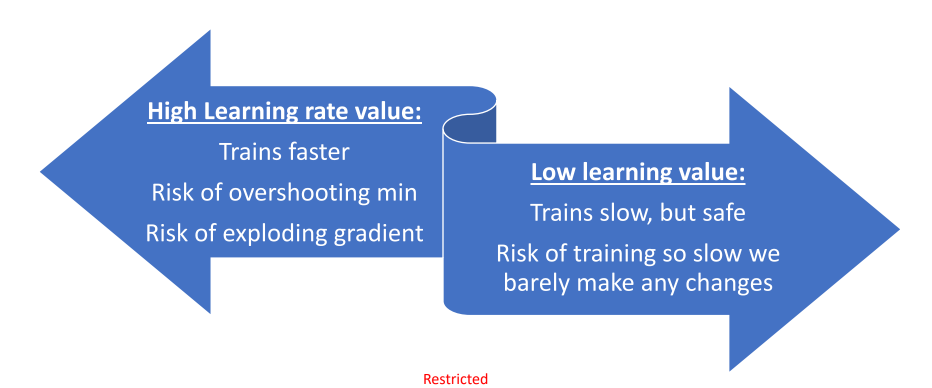
The **learning rate** is a **hyperparameter** which controls the step size taken by the algorithm in order to minimize the loss function during training.

- If the learning rate is too small, the algorithm will take longer to converge, but it can also prevent overshooting the global minimum. It might, however, lead to being stuck in non-optimum local minima.
- If the learning rate is too high, the algorithm may converge faster, but it may lead to exploding gradients.

The learning rate tradeoff

Definition (Learning rate tradeoff):

There is therefore a **tradeoff** on the deep learning rate, and as in all things with tradeoffs, it is about finding the **optimal middle ground**.

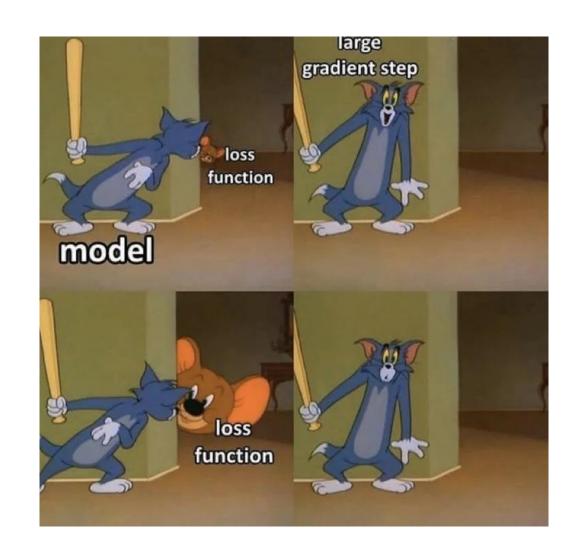


Triggering an exploding gradient on purpose

And, as a proof of that...

We can, in fact, trigger an exploding gradient problem, on purpose, by setting a very high learning rate value!

Let us try a much higher value for the learning rate (e.g. using $1e^{-2}$ instead of $1e^{-6}$).



Restricted

```
1 # Define and train neural network structure (random normal initialization)
 2 n x = 2
 3 n h = 4
 4 | n y = 1
 5 init type = "Normal"
 6 np.random.seed(37)
 7 shallow_neural_net_normal1 = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Train and show final loss
 9 shallow_neural_net_normal1.train(inputs, outputs, N_max = 1000, alpha = 1e-6, delta = 1e-6, display = True)
10 print(shallow neural net normal1.loss)
- Gradients:
[[ 1.42356523e-05 -1.73577846e-06 -9.53259136e-07 1.24839272e-06]
[ 1.32743203e-04 -1.61856153e-05 -8.88885652e-06 1.16408890e-05]]
[[ 1.27309997e-07 -1.55231348e-08 -8.52503385e-09 1.11644251e-08]]
[[ 3.86686371e-04]
[ 2.71147622e-04]
 [ 8.05690592e-05]
[-1.49929229e-04]]
                                                                Normal initialization and low
[[2.45019007e-06]]
- Parameters:
[[-0.00544636 0.06743081 0.0346647 -0.13003462]
                                                                learning rate (1e-6)
[ 0.15185119  0.09898237  0.02776809 -0.04485894]]
[[ 0.09619662 -0.08275786  0.05346571  0.12283862]]
                                                                This is fine...
[[ 0.05195923]
[-0.00633548]
[-0.00347934]
[ 0.00455655]]
[[0.14480251]]
Iteration 1 - Loss = 6.635227700991098
 1 # Show loss after training
 2 print(shallow neural net normal1.loss)
```

2.0907010654169245

Restricted

```
1 # Define and train neural network structure (random normal initialization)
 2 n x = 2
 3 n h = 4
 4 | n y = 1
 5 init type = "Normal"
 6 np.random.seed(37)
 7 shallow neural net normal2 = ShallowNeuralNet(n x, n h, n y, init type)
 8 # Train and show final loss
 9 shallow_neural_net_normal2.train(inputs, outputs, N_max = 1000, alpha = 1e-2, delta = 1e-6, display = True)
10 print(shallow neural net normal2.loss)
- Gradients:
[[ 0.14235652 -0.01735778 -0.00953259  0.01248393]
 [ 1.32743203 -0.16185615 -0.088888857  0.11640889]]
[[ 1.27309997e-03 -1.55231348e-04 -8.52503385e-05 1.11644251e-04]]
[[ 3.86686371]
 [ 2.71147622]
 [ 0.80569059]
 [-1.49929229]]
                                                                         Learning rate too
[[0.0245019]]
- Parameters:
                                                                         high leads to
[[-0.00544636 0.06743081 0.0346647 -0.13003462]
[ 0.15185119  0.09898237  0.02776809  -0.04485894]]
[[ 0.09619662 -0.08275786  0.05346571  0.12283862]]
                                                                         exploding gradients
[[ 0.05195923]
[-0.00633548]
[-0.00347934]
 [ 0.00455655]]
[[0.14480251]]
Iteration 1 - Loss = 4354741.971946698
 1 # Show loss after training
 2 print(shallow neural net normal2.loss)
```

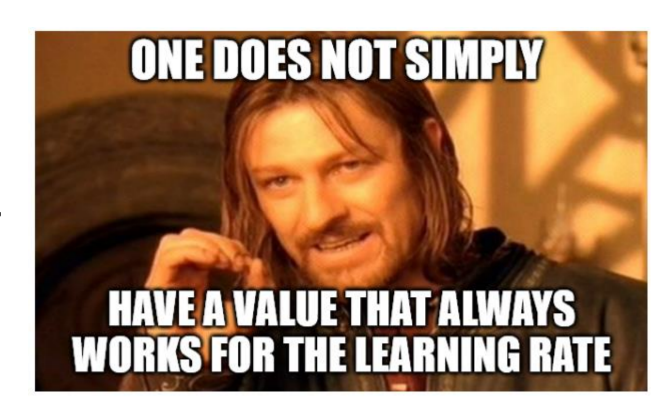
nan

No free lunch theorem – season 2

Definition (the no free lunch (or NFL) theorem – season 2):

The no free lunch (NFL) theorem also implies that there is no such thing as "one value that works for all models" when it comes to hyperparameters such as the learning rate.

No choice (again!): try some values and see how it goes!



The learning rate tradeoff

Definition (Hyperparameters and Hyperparameter tuning):

As we have seen, the **learning rate** is a **hyperparameter** that needs to be carefully chosen.

Thus, an important part of deep learning is hyperparameter tuning.

It consists, for instance, of **finding the optimal learning rate** that will allow the algorithm to **converge quickly and reliably**.

For a given model, there could be multiple hyperparameters.

For instance, the lambda value we used in regularization is also a hyperparameter, which requires hyperparameter tuning. The number of neurons used in a given layer is also a hyperparameter.

The learning rate tradeoff

Note: Sadly, that is sometimes all it takes to make the difference between a model that successfully trains and a model that does not!



Controlling gradients to prevent explosions

Adjusting and trying different values for the learning rate is often good enough to prevent exploding gradients.

 Gradient clipping: Set a max value for change on a single iteration and truncate any change that goes above that (to be discussed later).

There are other options, like:

• Downscaling initial parameters values: Divide initial parameter values by a certain factor to prevent large initial values on parameters.

• Time evolution on learning rate:

Make the learning rate value change over time, make it large at the beginning and progressively decrease it over iterations (to be discussed later).

```
# Define and train neural network structure (Weird initialization)
   n x = 2
   nh = 4
   ny = 1
   init type = "Weird"
 6 np.random.seed(37)
   shallow_neural_net_weird = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Train and show final loss
   shallow neural_net_weird.train(inputs, outputs, N_max = 100, alpha = 1e-6, delta = 1e-6, display = True)
Iteration 26 - Loss = nan
- Gradients:
[[nan nan nan nan]
[nan nan nan nan]]
[[nan nan nan nan]]
[[nan]
[nan]
                                     Back to our first exploding gradient example
[nan]
[nan]]
[[nan]]
- Parameters:
[[nan nan nan nan]
[nan nan nan nan]]
[[nan nan nan nan]]
[[nan]
[nan]
 [nan]
[nan]]
[[nan]]
```

```
1 # Define and train neural network structure (Weird initialization = some sort of a Uniform Xavier)
 2 n x = 2
 3 n h = 4
 4 | n y = 1
 5 init type = "Weird"
 6 np.random.seed(37)
 7 shallow_neural_net_xavier2 = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Divide initial values by 10!
 9 shallow neural net xavier2.W1 /= 100
10 shallow_neural_net_xavier2.b1 /= 100
11 shallow neural net xavier2.W2 /= 100
12 shallow neural net xavier2.b2 /= 100
13 # Train and show final loss
14 shallow neural net xavier2.train(inputs, outputs, N max = 1000, alpha = 1e-6, delta = 1e-6, display = True)
15 print(shallow neural net xavier2.loss)
- Gradients:
[[ 1.08643744e-06 -8.91020716e-06 -7.63367797e-06 5.82899367e-06]
 Downscaling
[[ 8.17657760e-09 -6.70586248e-08 -5.74514080e-08 4.38692717e-08]]
[[-1.73842740e-05]
 [-1.29545695e-05]
                                                                 parameters after
 [ 3.51566967e-05]
 [-1.97254965e-05]]
                                                                 initialization fixed
[[-5.1155505e-06]]
- Parameters:
[[ 0.01257226 -0.00101546 -0.00868907  0.00231634]
                                                                the exploding issue,
 [ 0.00339649  0.00521064 -0.01121648  0.0069431 ]]
[[-0.00616656 0.00716793 0.00827897 0.00360343]]
                                                                 but it leads to
[[-0.00159838]
 [ 0.01310878]
 [ 0.01123074]
                                                                slower training.
 [-0.00857567]]
[[0.00271391]]
Iteration 1 - Loss = 7.036477890046345
 1 # Show loss after training
 2 print(shallow neural net xavier2.loss)
```

The vanishing gradient problem

Definition (vanishing gradients):

We just observed **exploding gradients**. Its counterpart exists and is called **vanishing gradients**.

This typically occurs when the gradient descent rule has changes (in $\alpha \frac{\partial dL}{\partial W1}$ for instance) that are far smaller than the current values in the matrices (e.g. W_1).

It might also happen for other reasons (to be discussed later).

- We can typically force the apparition of a vanishing gradient problem by forcing the parameters to be initialized as very small values (or even zeroes).
- We can also force the apparition of a vanishing gradient problem by using a very small learning rate α .
- We can then observe that most parameters remain unchanged during training as the changes are close, or even equal to, zero.

Restricte

Restricted

```
1 # Define and train neural network structure (random normal initialization)
 2 n x = 2
 3 n h = 4
 4 n_y = 1
 5 init type = "Normal"
 6 np.random.seed(37)
 7 shallow_neural_net_normal = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Train and show final loss
 9 | shallow_neural_net_normal.train(inputs, outputs, N_max = 100, alpha = 1e-6, delta = 1e-6, display = True)
10 print(shallow neural net normal.loss)
- Gradients:
[[ 1.42356523e-05 -1.73577846e-06 -9.53259136e-07 1.24839272e-06]
[ 1.32743203e-04 -1.61856153e-05 -8.88885652e-06 1.16408890e-05]]
[[ 1.27309997e-07 -1.55231348e-08 -8.52503385e-09 1.11644251e-08]]
[[ 3.86686371e-04]
 [ 2.71147622e-04]
 [ 8.05690592e-05]
 [-1.49929229e-04]]
[[2.45019007e-06]]
                                                                                  This is fine.

    Parameters:

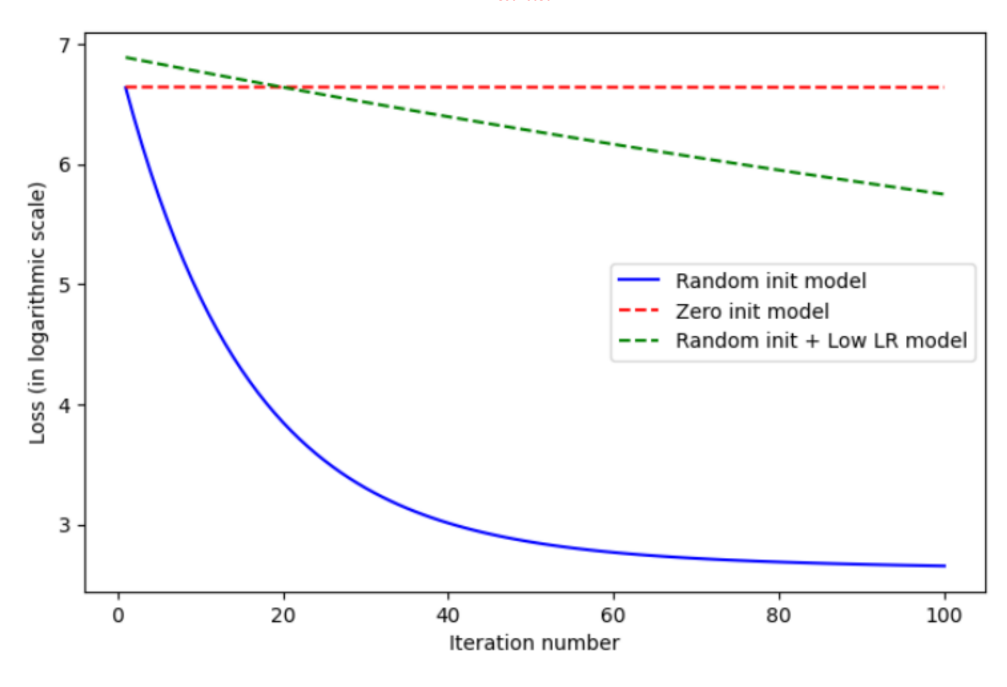
[[-0.00544636 0.06743081 0.0346647 -0.13003462]
 [ 0.15185119  0.09898237  0.02776809 -0.04485894]]
[[ 0.09619662 -0.08275786  0.05346571  0.12283862]]
[[ 0.05195923]
 [-0.00633548]
 [-0.00347934]
 [ 0.00455655]]
[[0.14480251]]
Iteration 1 - Loss = 6.635227700991098
 1 # Show final loss
 2 print(shallow neural net normal.loss)
```

Restricted

```
1 # Define and train neural network structure (Zero initialization)
 2 n x = 2
 3 n h = 4
 4 n y = 1
 5 init type = "Zero"
 6 np.random.seed(37)
 7 shallow_neural_net_zero = ShallowNeuralNet(n_x, n_h, n_y, init_type)
 8 # Train and show final loss
 9 | shallow_neural_net_zero.train(inputs, outputs, N_max = 100, alpha = 1e-6, delta = 1e-6, display = True)
10 print(shallow_neural_net_zero.loss)
- Gradients:
[[0. 0. 0. 0.]
                                                                      Initializing
[0. 0. 0. 0.]]
[[0. 0. 0. 0.]]
                                                                      parameters as
[[0.]
[0.]
[0.]
                                                                      zeroes is
[0.]]
[[-4.9531058e-06]]
                                                                      seriously
- Parameters:
[[0. 0. 0. 0.]
[0. 0. 0. 0.]]
                                                                      problematic and
[[0. 0. 0. 0.]]
[[0.]
                                                                      the model just
[0.]
[0.]
                                                                      does not train.
[0.]]
[[0.]]
Iteration 1 - Loss = 6.641131615309766
 1 # Show final loss
 2 print(shallow neural net zero.loss)
```

```
1 # Define and train neural network structure
 2 # (random normal initialization,
 3 # but very low learning rate alpha)
 4 n x = 2
 5 n h = 4
 6 n y = 1
 7 init_type = "Normal"
 8 np.random.seed(37)
 9 shallow_neural_net_normal_zerolr = ShallowNeuralNet(n_x, n_h, n_y, init_type)
10 # Train and show final loss
11 | shallow_neural_net_normal_zerolr.train(inputs, outputs, N_max = 100, alpha = 5e-8, delta = 1e-6, display = True)
12 print(shallow neural net normal zerolr.loss)
Gradients:
[[ 7.11782617e-07 -8.67889228e-08 -4.76629568e-08 6.24196361e-08]
 [ 6.63716016e-06 -8.09280764e-07 -4.44442826e-07 5.82044450e-07]]
[[ 6.36549987e-09 -7.76156741e-10 -4.26251693e-10 5.58221254e-10]]
[[ 1.93343185e-05]
[ 1.35573811e-05]
                                                             Setting up a learning rate that is
[ 4.02845296e-06]
[-7.49646145e-06]]
[[1.22509503e-07]]
                                                             too small might seriously hinder
- Parameters:
[[-0.00544636 0.06743081 0.0346647 -0.13003462]
                                                             the training.
[ 0.15185119  0.09898237  0.02776809  -0.04485894]]
[[ 0.09619662 -0.08275786  0.05346571  0.12283862]]
[[ 0.05195923]
                                                             And sometimes even lead to a
[-0.00633548]
[-0.00347934]
                                                              "convergence" to wrong minimum.
 0.00455655]]
[[0.14480251]]
Iteration 1 - Loss = 6.887598572349596
 1 # Show final loss
```

print(shallow_neural_net_normal_zerolr.loss)



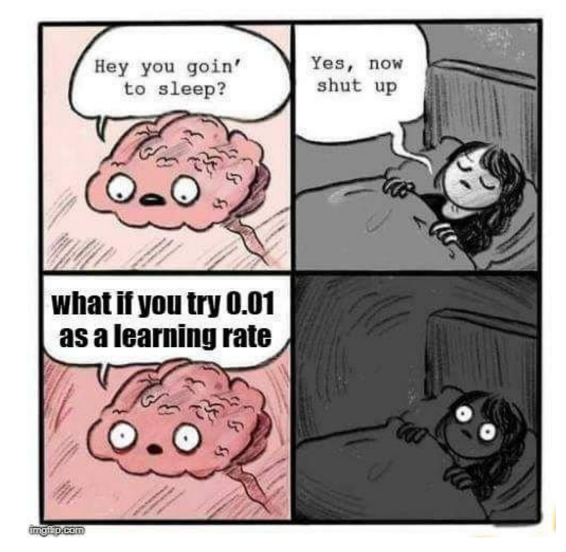
Remember, the learning rate tradeoff

Definition (Hyperparameters and hyperparameter tuning):

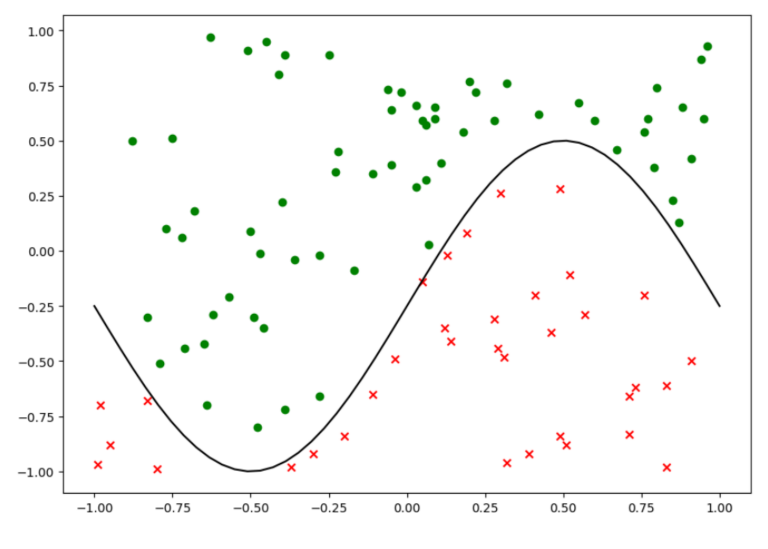
As we have seen, the **learning rate** is a **hyperparameter** that needs to be carefully chosen.

Thus, an important part of deep learning is hyperparameter tuning.

It consists, for instance, of **finding** the optimal learning rate that will allow the algorithm to converge quickly and reliably.



New dataset! (An abstract one this time)

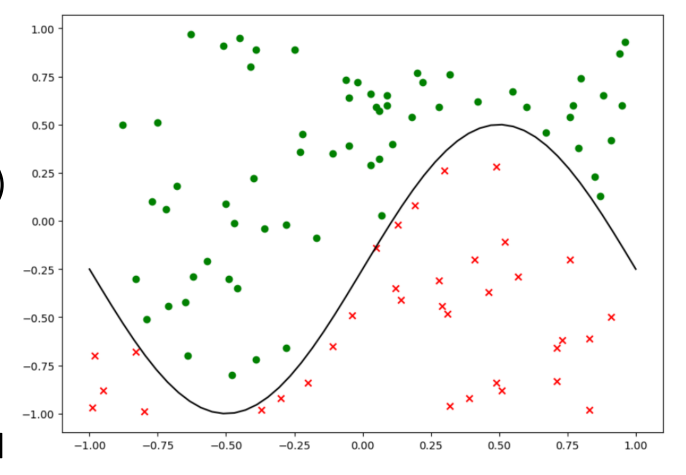


New dataset! (An abstract one this time)

- Each sample consists of two randomly drawn input values (x_1, x_2) , with $-1 < x_1, x_2 < 1$.
- Black line serves as (non-linear) frontier between both classes, with equation.

$$f(x) = -\frac{1}{4} + \frac{3}{4}\sin(x\pi)$$

• Two classes (green and red): Green (class 1) if $x_2 > f(x_1)$ and red (class 0) otherwise.



New dataset!

- We can generate some entries using the helper functions here.
- Basically, drawing pairs of values randomly and checking if above or below the sine line.

```
# All helper functions
    eps = 1e-5
    min val = -1 + eps
    \max \text{ val} = 1 - \text{eps}
    def val(min val, max val):
        return round(np.random.uniform(min val, max val), 2)
    def class_for_val(val1, val2):
        k = np.pi
        return int(val2 >= -1/4 + 3/4*np.sin(val1*k))
    n points = 100
    def create dataset(n points, min val, max val):
12
        val1 list = np.array([val(min val, max val) for in range(n points)])
        val2 list = np.array([val(min val, max val) for _ in range(n points)])
13
        inputs = np.array([[v1, v2] for v1, v2 in zip(val1_list, val2_list)])
14
15
        outputs = np.array([class for val(v1, v2) for v1, v2 in zip(val1 list, val2 list)]).reshape(n points, 1)
16
        return val1 list, val2 list, inputs, outputs
    # Generate dataset
    np.random.seed(47)
   val1_list, val2_list, inputs, outputs = create_dataset(n_points, min_val, max_val)
    # Check a few entries of the dataset
   print(val1 list.shape)
   print(val2 list.shape)
    print(inputs.shape)
   print(outputs.shape)
    print(inputs[0:5, :])
                                        0.75
10 print(outputs[0:5])
(100,)
                                        0.50
(100,)
(100, 2)
                                        0.25
(100, 1)
[[-0.77 0.1]
                                        0.00
  0.95 0.6 ]
  0.46 -0.37]
                                        -0.25
 [-0.3 -0.92]
                                        -0.50
 [ 0.42 0.62]]
[[1]
                                        -0.75
[1]
[0]
[0]
[1]]
                                                                                          0.75
```

Restricted

Adjusting our shallow NN

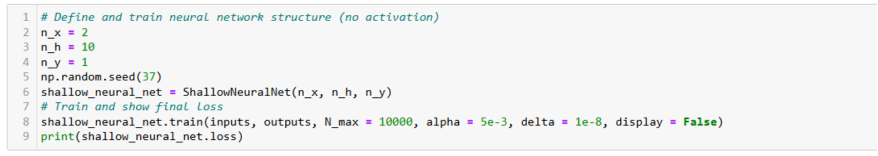
We can adjust our shallow neural network to the task by:

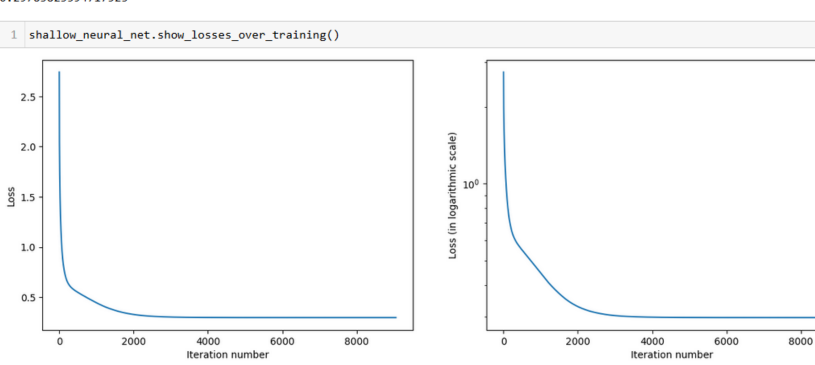
- Using the same two layers and forward pass procedure as before, leaving our init and forward methods untouched.
- Replace MSE with our crossentropy (another name for the log-likelihood loss).
- Backward and trainer methods are left untouched.

```
class ShallowNeuralNet():
    def __init__(self, n_x, n_h, n_y):
        # Network dimensions
        self.n x = n x
        self.n h = n h
        self.n y = n y
        # Initialize parameters
        self.init parameters normal()
        # Loss, initialized as infinity before first calculation is made
        self.loss = float("Inf")
    def init parameters normal(self):
        # Weights and biases matrices (randomly initialized)
        self.W1 = np.random.randn(self.n x, self.n h)*0.1
        self.b1 = np.random.randn(1, self.n_h)*0.1
        self.W2 = np.random.randn(self.n h, self.n y)*0.1
        self.b2 = np.random.randn(1, self.n y)*0.1
    def forward(self, inputs):
        # Wx + b operation for the first layer
       Z1 = np.matmul(inputs, self.W1)
       Z1 b = Z1 + self.b1
        # Wx + b operation for the second layer
       Z2 = np.matmul(Z1_b, self.W2)
       Z2 b = Z2 + self.b2
        # Adding clipping to keep prediction values
        # between 0 and 1 (see loss function!)
        pred = np.clip(Z2_b, 0, 1)
       return pred
    def CE loss(self, inputs, outputs):
        # MSE loss function as before
        outputs re = outputs.reshape(-1, 1)
        pred = self.forward(inputs)
        eps = 1e-5
        losses = outputs*np.log(pred + eps) + (1 - outputs)*np.log(1 - pred + eps)
        self.loss = -np.sum(losses)/outputs.shape[0]
        return self.loss
    def backward(self, inputs, outputs, alpha = 1e-5):
        # Get the number of samples in dataset
```

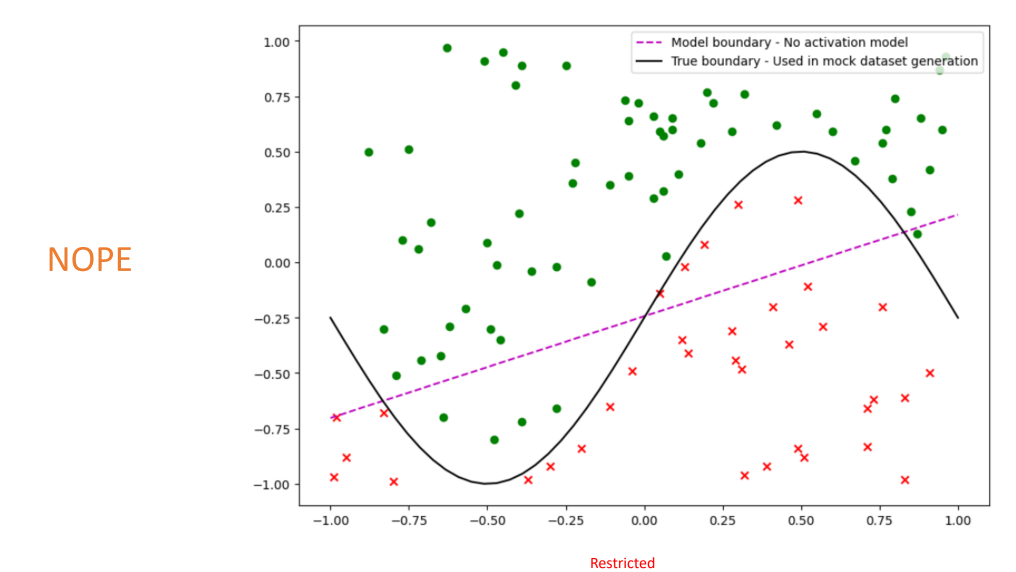
Trains just fine! (or does it?)

Training curves looking nice, but does that mean the model is good?





Trains just fine! (or does it?)



Linear layers necessarily lead to linear boundary

As shown in Notebook,

- Our model consists of two linear layers, with two WX+B operations in a row.
- The result is linear also, see the y_{pred} equation.
- Boundary for model is therefore linear in x_1 and x_2 !

To demonstrate, let us consider that $n_x = 2$, $n_h = 4$ and $n_v = 1$. In this configuration, we have:

$$W_{1} = \begin{pmatrix} w_{1,1}^{(1)}, & w_{1,2}^{(1)}, & w_{1,3}^{(1)}, & w_{1,4}^{(1)} \\ w_{2,1}^{(1)}, & w_{2,2}^{(1)}, & w_{2,3}^{(1)}, & w_{2,4}^{(1)} \end{pmatrix}$$

$$b_{1} = \begin{pmatrix} b_{1,1}^{(1)}, & b_{1,2}^{(1)}, & b_{1,3}^{(1)}, & b_{1,4}^{(1)} \end{pmatrix}$$

$$W_{2} = \begin{pmatrix} w_{1,1}^{(2)} \\ w_{1,2}^{(2)} \\ w_{1,3}^{(2)} \\ w_{1,4}^{(2)} \end{pmatrix}$$

$$b_{2} = \begin{pmatrix} b_{1,1}^{(2)} \end{pmatrix}$$

After the first operation $Z_1 = XW_1 + b_1$, we have:

$$Z_{1} = \begin{pmatrix} w_{1,1}^{(1)}x_{1} + w_{2,1}^{(1)}x_{2} + b_{1,1}^{(1)}, & w_{1,2}^{(1)}x_{1} + w_{2,2}^{(1)}x_{2} + b_{1,2}^{(1)}, & w_{1,3}^{(1)}x_{1} + w_{2,3}^{(1)}x_{2} + b_{1,3}^{(1)}, & w_{1,4}^{(1)}x_{1} + w_{2,4}^{(1)}x_{2} + b_{1,4}^{(1)} \end{pmatrix}$$

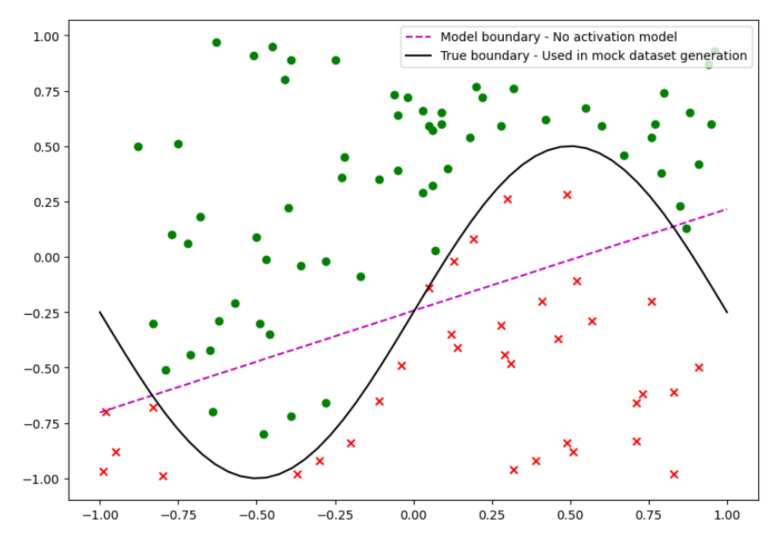
After the second operation, the predicted price y_{pred} is given by $y_{pred} = Z_1 W_2 + b_2$

$$y_{pred} = \left(\sum_{k=1}^{4} w_{1,k}^{(1)} w_{1,k}^{(2)}\right) x_1 + \left(\sum_{k=1}^{4} w_{2,k}^{(1)} w_{1,k}^{(2)}\right) x_2 + \left(\sum_{k=1}^{4} b_{1,k}^{(1)} w_{1,k}^{(2)}\right) + b_{1,1}^{(2)}$$

Trains just fine! (or does it?)

Well, that is going to be a problem, because the black frontier is definitely not linear...

Or, in other words, the data is **not linearly separable.**



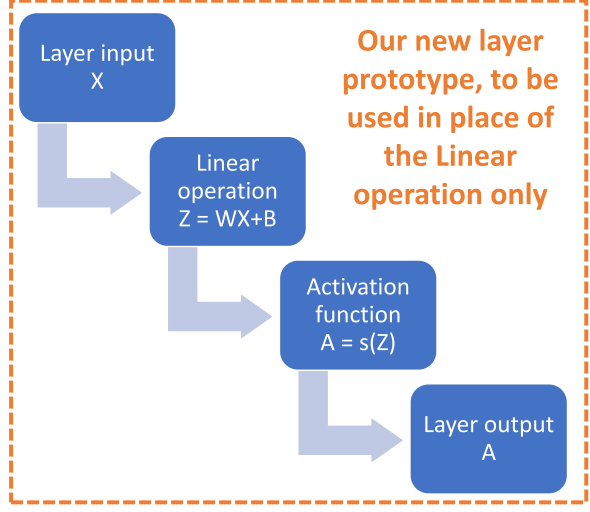
Activation functions and introducing nonlinearity in the Neural Networks

Definition (Activation functions and non-linearity):

Activation functions are an important component of neural networks because they introduce non-linearity to the model.

This is important because most real-world data is non-linear.

Simply done by adding an extra operation after the linear one, for instance our sigmoid function.



Adding sigmoid activations

We can update our model by:

- Adding a sigmoid method, implementing the sigmoid operation from earlier.
- Adding sigmoid operations after each linear operation in the forward method.
- Our cross-entropy loss method remains unchanged, as it will simply use the updated forward procedure.

```
class ShallowNeuralNet WithAct():
    def __init__(self, n_x, n_h, n_y):
        # Network dimensions
        self.n x = n x
        self.nh = nh
        self.n y = n y
        # Initialize parameters
        self.init parameters normal()
        # Loss, initialized as infinity before first calculation is made
        self.loss = float("Inf")
    def init parameters normal(self):
        # Weights and biases matrices (randomly initialized)
        self.W1 = np.random.randn(self.n x, self.n h)*0.1
        self.b1 = np.random.randn(1, self.n h)*0.1
        self.W2 = np.random.randn(self.n h, self.n y)*0.1
        self.b2 = np.random.randn(1, self.n y)*0.1
    def sigmoid(self, val):
        return 1/(1 + np.exp(-val))
    def forward(self, inputs):
        # Wx + b operation for the first layer
        Z1 = np.matmul(inputs, self.W1)
       Z1 b = Z1 + self.b1
        A1 = self.sigmoid(Z1_b)
       # Wx + b operation for the second layer
        Z2 = np.matmul(A1, self.W2)
      72 b = 72 + self_b2
      y_pred = self.sigmoid(Z2_b)
        return v pred
```

Restricted

Adding sigmoid activations

Careful now...

 As the forward procedure has changed, we would have to update the backward propagation...

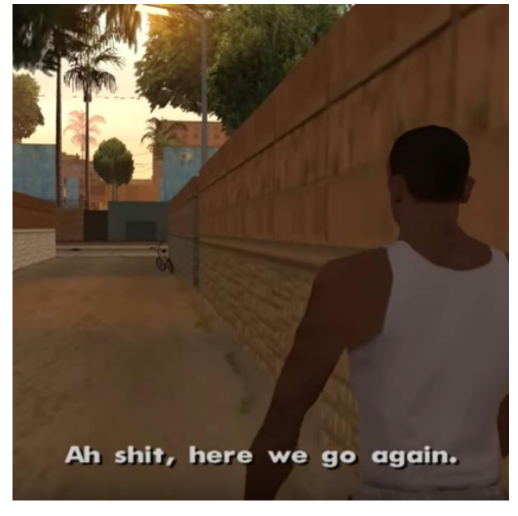
```
class ShallowNeuralNet WithAct():
   def __init__(self, n_x, n_h, n_y):
       # Network dimensions
       self.n x = n x
       self.nh = nh
       self.n y = n y
       # Initialize parameters
       self.init parameters normal()
       # Loss, initialized as infinity before first calculation is made
       self.loss = float("Inf")
   def init parameters normal(self):
       # Weights and biases matrices (randomly initialized)
       self.W1 = np.random.randn(self.n x, self.n h)*0.1
       self.b1 = np.random.randn(1, self.n h)*0.1
       self.W2 = np.random.randn(self.n h, self.n y)*0.1
       self.b2 = np.random.randn(1, self.n y)*0.1
   def sigmoid(self, val):
       return 1/(1 + np.exp(-val))
   def forward(self, inputs):
       # Wx + b operation for the first layer
       Z1 = np.matmul(inputs, self.W1)
       Z1 b = Z1 + self.b1
       A1 = self.sigmoid(Z1 b)
       # Wx + b operation for the second layer
       Z2 = np.matmul(A1, self.W2)
       Z2 b = Z2 + self.b2
       y_pred = self.sigmoid(Z2_b)
       return y pred
```

Adding sigmoid activations

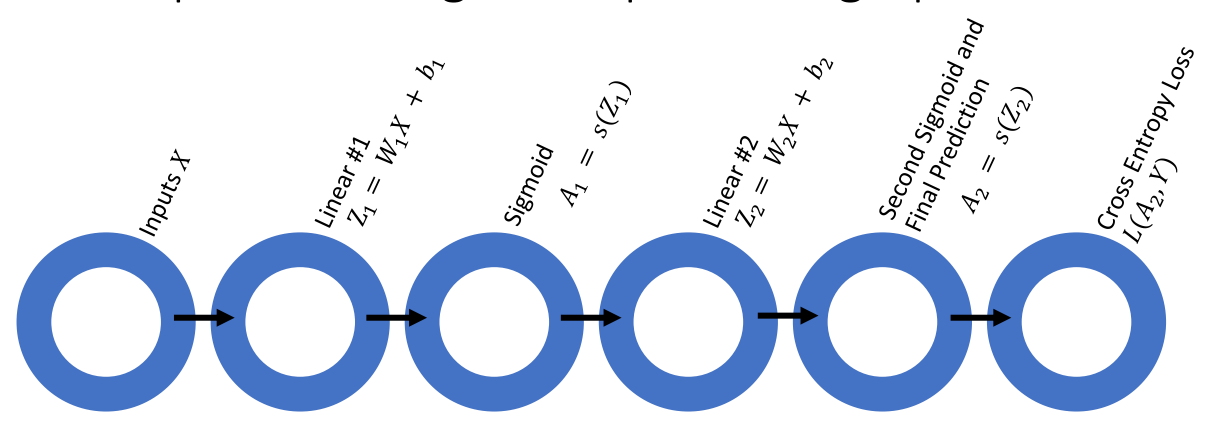
Careful now...

- As the forward procedure has changed, we would have to update the backward propagation...
- This means we have to back to using the chain rule manual differentiation procedure, account for the new sigmoid operations, and get the new update rules to use in the backward method...

When you realize you have to re-do the entire backpropagation calculation and implemaentation to account for just a small change of activation function



Step 1: Drawing a computation graph



Step 2: Define the forward equations

Our forward method gives:

$$Z_1 = W_1 X + b_1$$

$$A_1 = s(Z_1)$$

$$Z_2 = W_2 A_1 + b_2$$

$$A_2 = s(Z_2)$$

$$L = \frac{-1}{N} \sum_{i}^{N} Y \ln(A_2) + (1 - Y) \ln(1 - A_2)$$

Restricted

Retrieving the gradient descent update rules takes a few steps and requires some organizing...

First, recall that our loss function is defined as

$$L = \frac{-1}{N} \sum_{i=1}^{N} Y \ln(A_2) + (1 - Y) \ln(1 - A_2)$$

Therefore, we have

$$\frac{\partial L}{\partial A_2} = -\frac{Y}{A_2} + \frac{1 - Y}{1 - A_2}$$

• Then, recall that

$$A_2 = s(Z_2)$$

We can then prove that

$$s'(X) = s(X)(1 - s(X))$$

Using the chain rule, we then obtain

$$\frac{\partial L}{\partial Z_2} = \frac{\partial L}{\partial A_2} \frac{\partial A_2}{\partial Z_2} = \frac{\partial L}{\partial A_2} A_2 (1 - A_2)$$

Let us continue

$$Z_2 = W_2 A_1 + b_2$$

Using the chain rule, we then obtain

$$\frac{\partial L}{\partial W_2} = \frac{\partial L}{\partial Z_2} \frac{\partial Z_2}{\partial W_2} = \frac{\partial L}{\partial Z_2} A_1$$

$$\frac{\partial L}{\partial b_2} = \frac{\partial L}{\partial Z_2} \frac{\partial Z_2}{\partial b_2} = \frac{\partial L}{\partial Z_2}$$

• Again...

$$Z_2 = W_2 A_1 + b_2$$

 $A_1 = s(Z_1)$

Using the chain rule and our sigmoid derivative, we then obtain

$$\frac{\partial L}{\partial A_1} = \frac{\partial L}{\partial Z_2} \frac{\partial Z_2}{\partial A_1} = \frac{\partial L}{\partial Z_2} W_2$$

$$\frac{\partial L}{\partial Z_1} = \frac{\partial L}{\partial A_1} \frac{\partial A_1}{\partial Z_1} = \frac{\partial L}{\partial A_1} A_1 (1 - A_1)$$

Almost there...

$$Z_2 = W_2 A_1 + b_2$$

 $A_1 = s(Z_1)$

Using the chain rule and our sigmoid derivative, we then obtain

$$\frac{\partial L}{\partial A_1} = \frac{\partial L}{\partial Z_2} \frac{\partial Z_2}{\partial A_1} = \frac{\partial L}{\partial Z_2} W_2$$

$$\frac{\partial L}{\partial Z_1} = \frac{\partial L}{\partial A_1} \frac{\partial A_1}{\partial Z_1} = \frac{\partial L}{\partial A_1} A_1 (1 - A_1)$$

Finally...!

$$Z_1 = W_1 X + b_1$$

Using the chain rule and our sigmoid derivative, we then obtain

$$\frac{\partial L}{\partial W_1} = \frac{\partial L}{\partial Z_1} \frac{\partial Z_1}{\partial W_1} = \frac{\partial L}{\partial Z_1} X$$

$$\frac{\partial L}{\partial b_1} = \frac{\partial L}{\partial Z_1} \frac{\partial Z_1}{\partial b_1} = \frac{\partial L}{\partial Z_1}$$

Backward Method Update

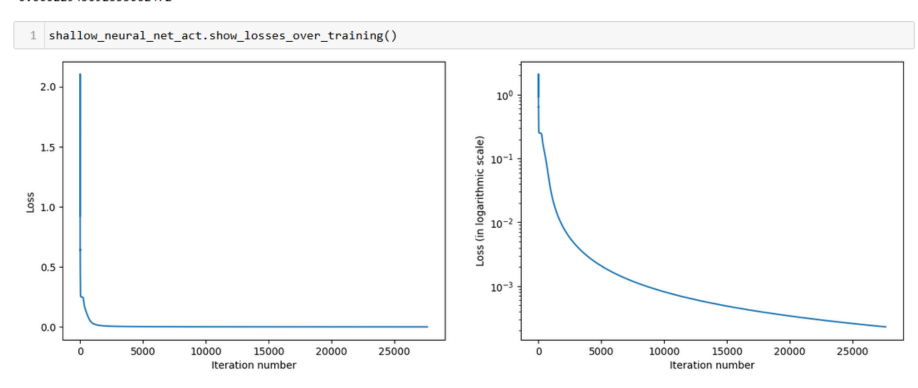
Reusing our formulas carefully gives the following update for the backward method.

- Notice how we did not rush and computed each term, one operation at a time, to avoid making mistakes.
- Take it slow!

```
def backward(self, inputs, outputs, alpha = 1e-5):
    # Get the number of samples in dataset
    m = inputs.shape[0]
    # Forward propagate
   Z1 = np.matmul(inputs, self.W1)
   Z1 b = Z1 + self.b1
   A1 = self.sigmoid(Z1 b)
   Z2 = np.matmul(A1, self.W2)
   Z2 b = Z2 + self.b2
   A2 = self.sigmoid(Z2 b)
   # Compute error term
    dL dA2 = -outputs/A2 + (1 - outputs)/(1 - A2)
    dL dZ2 = dL dA2*A2*(1 - A2)
    dL dA1 = np.dot(dL_dZ2, self.W2.T)
    dL dZ1 = dL dA1*A1*(1 - A1)
   # Gradient descent update rules
    self.W2 -= (1/m)*alpha*np.dot(A1.T, dL dZ2)
    self.W1 -= (1/m)*alpha*np.dot(inputs.T, dL dZ1)
    self.b2 -= (1/m)*alpha*np.sum(dL_dZ2, axis = 0, keepdims = True)
    self.b1 -= (1/m)*alpha*np.sum(dL dZ1, axis = 0, keepdims = True)
    # Update loss
    self.CE loss(inputs, outputs)
```

Trains just fine!

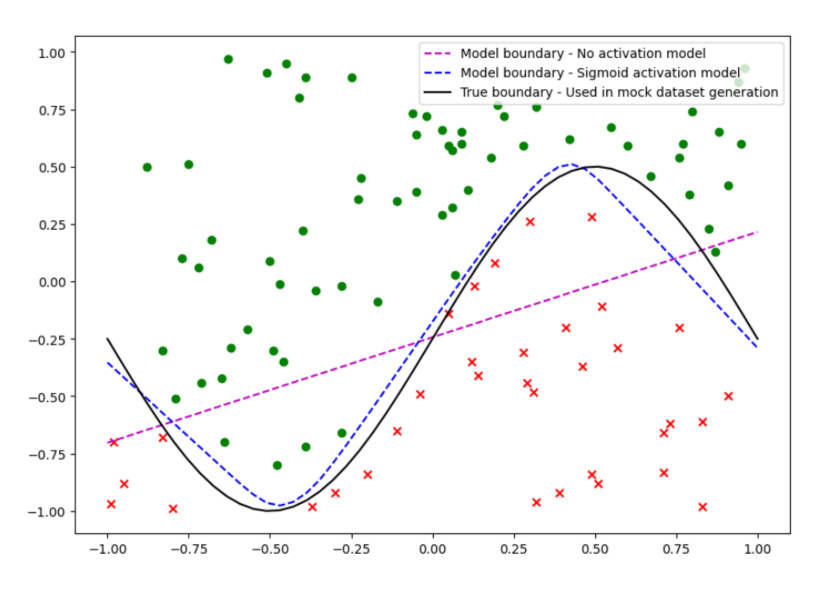
```
# Define and train neural network structure (with activation)
n_x = 2
n_h = 10
n_y = 1
np.random.seed(37)
shallow_neural_net_act = ShallowNeuralNet_WithAct(n_x, n_h, n_y)
# Train and show final loss
shallow_neural_net_act.train(inputs, outputs, N_max = 100000, alpha = 5, delta = 1e-8, display = False)
print(shallow_neural_net_act.loss)
```



Trains just fine!

Using the sigmoid activation functions helped the network create some non-linearity!

(Blue curve looking good!)



Examples of activations: Sigmoid

Definition (sigmoid function):

$$s(x) = \frac{1}{1 + \exp(-x)}$$

The sigmoid activation function is often used in the output layer of a binary classification model, because the output will then have values between 0 and 1, and can therefore be interpreted as a probability.

```
def sigmoid(val):
        return 1/(1 + np.exp(-val))
     --- Sigmoid
0.8
0.6
0.4
0.2
0.0
```

Examples of activations: Hyperbolic tangent

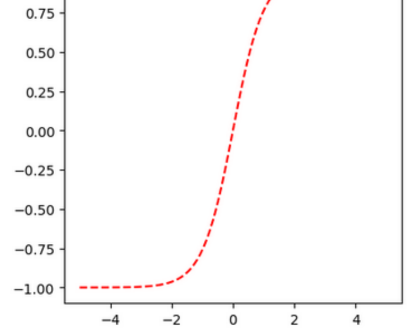
Definition (tanh function):

The tanh (short for "hyperbolic tangent") activation function is somewhat similar to the sigmoid function, but it maps values to a range between -1 and 1.

Like the sigmoid function, it is often used in the output layer of a classification model.

```
1 def tanh(val):
2 return np.tanh(val)

1.00 --- Tanh
0.75 --- Tanh
0.25 ---- Tanh
```



About Sigmoid and Tanh functions

Some observations about Sigmoid and Tanh functions (empirical, subject to No Free Lunch!)

- Those activation functions were widely used prior to ReLU.
- People were very comfortable with those as they were reminiscent of Logistic Regression and they are simply differentiable.
- The problem with those is that being squeezed between [0, 1] or [-1, 1], we can have a hard time training deep networks, as the gradient will tend to vanish.

Examples of activations: ReLU

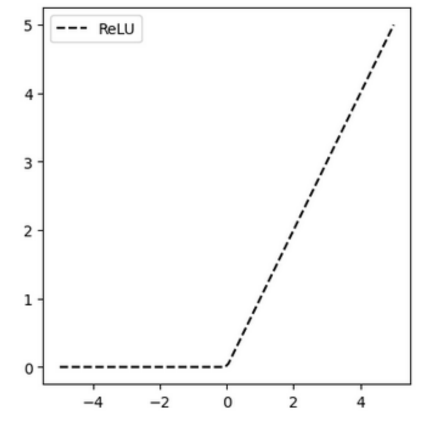
Definition (ReLU function):

The ReLU (short for "rectified linear unit") activation function is a simple function that maps any input value less than 0 to 0, and any input value greater than or equal to 0 to the input value itself.

It is widely used because it is computationally efficient and does not saturate (i.e., "die") like some other activation functions.

From [Glorot2011].

```
1 def ReLU(val):
2 return np.maximum(0, val)
```



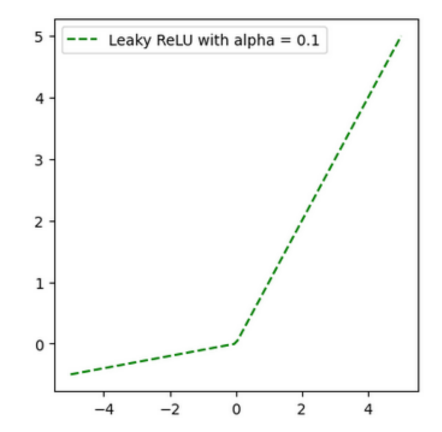
Examples of activations: Leaky ReLU

Definition (Leaky ReLU function):

The leaky ReLU activation function is similar to the ReLU function, but it allows a small negative slope for input values less than 0.

This can help to alleviate the "dying ReLU" problem, where some neurons in the network "die" when they produce negative values and might no longer respond to input.

```
def leaky_ReLU(val, alpha = 0.1):
    return np.where(val > 0, val, alpha*val)
```



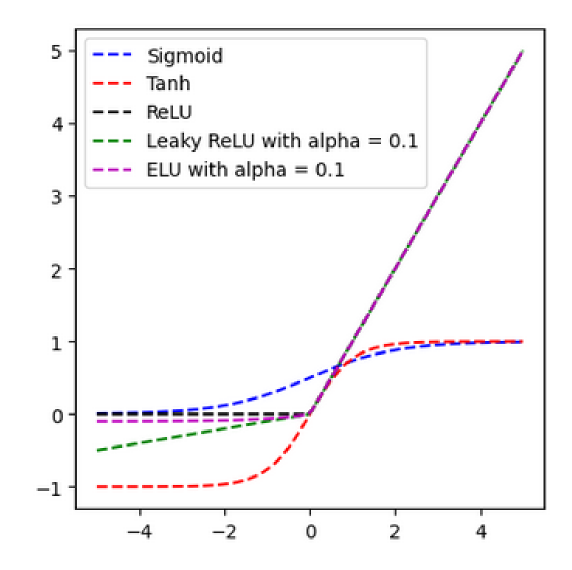
A note on activation functions

We have listed the most common activation functions, which are used most of the time.

Keep in mind that many more activation functions exist, some of them being very advanced or niche, but worth keeping an eye on.

https://pytorch.org/docs/stable/nn.ht ml#non-linear-activations-weightedsum-nonlinearity

(More in bonus slides!)



, [Non-linear Activations (weighted sum	, nonlinearity)]	
Aŋ	nn.ELU	Applies the Exponential Linear Unit (ELU) function, element- wise, as described in the paper: Fast and Accurate Deep Network Learning by Exponential Linear Units (ELUs).		
We lactiv	nn.Hardshrink	Applies the Hard Shrinkage (Hardshrink) function element- wise.		
Keep activ	nn.Hardsigmoid	Applies the Hardsigmoid function element-wise.		
of th	nn.Hardtanh	Applies the HardTanh function element-wise.		
word look http:	nn.Hardswish	Applies the Hardswish function, element-wise, as described in the paper: Searching for MobileNetV3.		
n.htı weig	nn.LeakyReLU	Applies the element-wise function:		
	nn.LogSigmoid	Applies the element-wise function:	4	

The universal approximation theorem

Definition (the Universal Approximation Theorem):

Mathematically speaking, any neural network architecture aims at finding the mathematical function y = f(x) that can map some given inputs x to outputs y. This function f(x) can be arbitrarily complex, based on the dataset and task at hand.

The Universal Approximation Theorem states that Neural Networks have a kind of universality property.

This means that no matter what f(x) might be, there is a network that can approximately approach the function f(x)!

As such, this is the most important theorem of Deep Learning!

Out of class, supporting papers, for those of you who are curious.

- [LeCun1998] Y. LeCun, Y. Bengio, and G. Hinton, "Efficient backprop. In Neural networks: Tricks of the trade", 1998.
- [Glorot2010] X. Glorot, and Y. Bengio, "Understanding the difficulty of training deep feedforward neural networks", 2010.
- [He2015] K. He, X. Zhang, S. Ren, & J. Sun, "Delving deep into rectifiers: Surpassing human-level performance on imagenet classification.", 2015.

Out of class, supporting papers, for those of you who are curious.

- [LeCun1998bis] Y. LeCun, L. Bottou, Y. Bengio, and P. Haffner "Gradient-based learning applied to document recognition.", 1998.
- [Glorot2011] X. Glorot, A. Bordes, and Y. Bengio, "Deep Sparse Rectifier Neural Networks", 2011.
- [Dubey2022] S. R. Dubey et al., "Activation Functions in Deep Learning: A Comprehensive Survey and Benchmark", 2022.

Tracking important names (Track their works and follow them on Scholar, Twitter, or whatever works for you!)

Yann LeCun: Chief AI Scientist at Facebook/Meta and Silver
 Professor at the New York University, another one of the three
 Godfathers of Deep Learning and 2018 Turing Award winner (highest distinction in Computer Science).

http://yann.lecun.com/

https://scholar.google.com/citations?user=WLN3QrAAAAAJ&hl=fr

• Yoshua Bengio: Professor at University of Montreal, <u>last of the three</u> <u>Godfathers of Deep Learning</u> and **2018 Turing Award** winner (highest distinction in Computer Science).

https://yoshuabengio.org

https://scholar.google.com/citations?user=kukA0LcAAAAJ&hl=fr

Tracking important names (Track their works and follow them on Scholar, Twitter, or whatever works for you!)

- Xavier Glorot: Currently works at DeepMind. https://scholar.google.com/citations?user= WnkXlkAAAAJ&hl=fr
- Kaiming He: Researcher at Facebook AI Research (FAIR/Meta AI). https://scholar.google.com/citations?user=DhtAFkwAAAAJ&hl=en

Tracking important names (Track their works and follow them on Scholar, Twitter, or whatever works for you!)

- Leon Bottou: Works at FAIR/Meta AI.
 https://leon.bottou.org/
 https://scholar.google.fr/citations?user=kbN88gsAAAAJ&hl=fr
- FAIR/Meta AI: Facebook Artificial Intelligence Research, now Meta Artificial Intelligence is an academic research laboratory focused on generating knowledge for the AI community.

https://ai.facebook.com/

https://twitter.com/MetaAI?ref src=twsrc%5Egoogle%7Ctwcamp%5E serp%7Ctwgr%5Eauthor

The He initialization

Definition (the He initialization [LeCun1998]):

The He initializer is similar to the Xavier initializer, but with a variance $\frac{1}{N_{in}+N_{out}}$, where N_{in} (resp. N_{out}) is the size of each layer input (resp. output). Each layer is therefore initialized with its own variance.

It seems to improve performance when working with deeper networks.

```
def init_parameters_he(self):
    # Weights and biases matrices (He initialized)
    range1 = np.sqrt(4/(self.n_x + self.n_h))
    self.W1 = np.random.randn(self.n_x, self.n_h)*range1
    self.b1 = np.random.randn(1, self.n_h)*range1
    range2 = np.sqrt(4/(self.n_h + self.n_y))
    self.W2 = np.random.randn(self.n_h, self.n_y)*range2
    self.b2 = np.random.randn(1, self.n_y)*range2
```

The LeCun initialization

Definition (the LeCun initialization [LeCun1998bis]):

The LeCun initializer is based on a random normal distribution, with a variance of $\sqrt{\frac{1}{N_{in}}}$, with N_{in} being the number of inputs of each layer.

This initializer is reported to be particularly useful for architectures with sigmoid and tanh activation functions.

```
def init_parameters_lecun(self):
    # Weights and biases matrices (LeCun initialized)
    range1 = np.sqrt(1/self.n_x)
    self.W1 = np.random.randn(self.n_x, self.n_h)*range1
    self.b1 = np.zeros((1, self.n_h))
    range2 = np.sqrt(1/self.n_h)
    self.W2 = np.random.randn(self.n_h, self.n_y)*range2
    self.b2 = np.zeros((1, self.n_y))
```

Restricted

Initializations variations and more

Many more initialization formulas exist:

- Xavier has random uniform and normal variations (same for He),
- Glorot proposed more initializations,
- Orthogonal initializations are sometimes useful but rare,
- Variance scaling initialization also exists but rarely used,
- Etc.

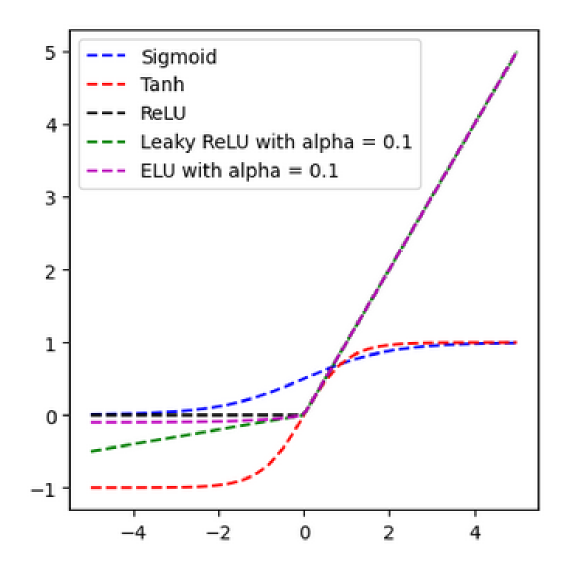
PyTorch has listed a lot of them, ready to use: https://pytorch.org/docs/stable/nn.init.html

A note on activation functions

We have listed the most common activation functions, which are used most of the time.

Keep in mind that many more activation functions exist, some of them being very advanced or niche, but worth keeping an eye on.

https://pytorch.org/docs/stable/nn.ht ml#non-linear-activations-weightedsum-nonlinearity



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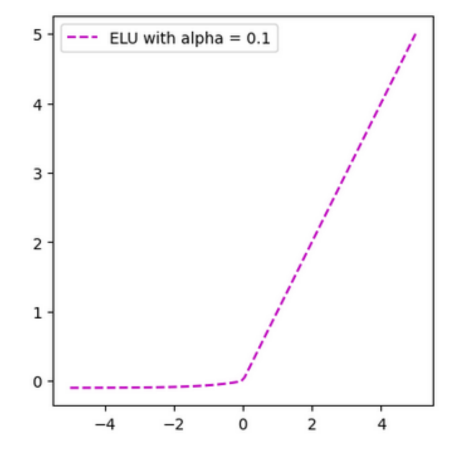
Examples of activations: ELU

Definition (ELU function):

The **ELU** (short for "**exponential linear unit**") **activation function** is similar to the ReLU function, but it has a negative slope for input values less than 0.

As we mentioned in the leaky ReLU function, ELU can help to alleviate the "dying ReLU" problem.

```
def ELU(val, alpha = 0.1):
    return np.where(val > 0, val, alpha*(np.exp(val) - 1))
```



From (Leaky) ReLU to more advanced LU

- Using **ReLU**, we can train deeper models, but the gradient would still die for negative numbers due to the zeroing in x < 0.
- Numerous Activation Functions were created to address this problem such as LeakyReLU and PReLU, but were still subject to issues related to gradients.
- The most notable one would be the **Exponential Linear Unit (ELU)** functions. Exponential behaviour sped up learning by bringing the normal gradient closer to the unit natural gradient because of a reduced bias shift effect.

A note on more advanced activation functions

More recent activation functions, such as **Swish** and **Mish**, suggest to use activation functions with **learnable/trainable parameters**.

• Those adaptive activation functions allow for different neurons to learn different activation functions for richer learning while adding parametric complexity to the networks (see [Dubey2022] for a good benchmark on activation functions).

In addition, the class of **Gated Linear Unit** (or **GLU**) has been studied quite a bit in Natural Language Processing architectures and they control what information is passed up to the following layer using gates similar to the ones found in LSTMs.

• More on this on Week 6, stay tuned for more!