August 1, 2024

1 Recurrent neural network (RNN) from scratch

2 What is a recurrent neural network?

A recurrent neural network (RNN), as opposed to a regular fully connected neural network (FCNN), has layers that are connected to themselves.

The difference might be clearer by first looking at an FCNN.

In an FCNN there are no connections between nodes in a single layer. For instance, h_1^1 is not connected to h_2^1 . In addition, the input and output are always of a fixed length.

In an RNN, however, this is no longer the case. Nodes in the hidden layers are connected to themselves, represented by the curved lines in the figure below.

Thus the output \vec{h} from the hidden layer is fed back into the hidden layer. This recurrence makes RNNs useful when working with sequential data, as we can have input of variable length. This is more clear if we unfold the recurrent part of the network.

3 The mathematics of RNNs

3.1 The RNN architecture

Consider some sequential input X with n features. Note that X here is an array with two axes, since it contains n features at each time step in the sequence. We will denote the input at a specific time step t as

$$\vec{X}^{(t)} = \begin{pmatrix} X_1^{(t)} \\ \vdots \\ X_n^{(t)} \end{pmatrix},$$

which is then an n-dimensional vector.

Next, consider an RNN with L hidden layers, and an output layer with m features. We will denote the output of the l'th hidden layer at time step t as

$$\vec{h}_l^{(t)} = \begin{pmatrix} h_{l,1}^{(t)} \\ \vdots \\ h_{l,n_l}^{(t)} \end{pmatrix},$$

1

with n_l being the number of features in the l'th hidden layer. The output of the RNN at time step t is denoted

$$\hat{\vec{y}}^{(t)} = \begin{pmatrix} \hat{y}_1 \\ \vdots \\ \hat{y}_m, \end{pmatrix}$$

where the hat is there to distinguish the RNN output $\hat{\vec{y}}^{(t)}$ from the target value, which is denoted $\vec{y}^{(t)}$. The RNN will then look like this.

3.2 Forward propagation

In order to propagate forward through the network we need some weights and biases to connect the nodes. To simplify the notation going forward, we will consider the input layer to be the zeroth layer, and the output layer to be the zeroth layer. We need each node to propagate to the node at the next layer (keeping the time step constant), and the next time step (keeping the layer constant), except for the input and output layers which do not connect to each other (as illustrated in the diagram above).

Let $W^{l,l+1}$ be the weight matrix and $\vec{b}^{l,l+1}$ the bias vector, both connecting nodes at the l'th layer to the l+1'th layer, keeping the time step constant. Next, let W^{ll} be the weight matrix and \vec{b}^{ll} the bias vector, both connecting nodes at subsequent time steps in the same layer. Also, let σ_l be the activation function in the l'th layer. Lastly, define the weighted sum $\vec{z}_l^{(t)}$ at layer l and time step t such that the output of the node is the activation of that weighted sum, that is, such that $\vec{h}_l^{(t)} = \sigma_l(\vec{z}_l^{(t)})$.

Using these definitions the output from the first hidden layer at the first time step is then

$$\vec{h}_1^{(1)} = \sigma_1 \left(\vec{z}_1^{(1)} \right),$$

with

$$\vec{z}_1^{(1)} = W^{01} \vec{X}^{(1)} + \vec{b}^{01}.$$

At later time steps we will also need to consider the contribution from the previous time step. Hence for $t \geq 2$ we will define

$$\left(\vec{z}_{1}^{(t)}\right)_{\mathrm{layer}} = W^{01}X^{(t)} + \vec{b}^{01}$$

$$\left(\vec{z}_{1}^{(t)}\right)_{\mathrm{time}} = W^{11} \vec{h}_{1}^{(t-1)} + \vec{b}^{11},$$

such that $(\vec{z}_1^{(t)})_{\text{layer}}$ is the contribution from the previous layer, and $(\vec{z}_1^{(t)})_{\text{time}}$ is the contribution from the previous time step. We then have

$$\vec{z}_1^{(t)} = \left(\vec{z}_1^{(t)}\right)_{\text{layer}} + \left(\vec{z}_1^{(t)}\right)_{\text{time}},$$

and

$$\vec{h}_1^{(t)} = \sigma_1 \left(\vec{z}_1^{(t)} \right).$$

The expression is exactly the same for any hidden node, but for $l \geq 2$ we substitute $\vec{X}^{(t)}$ with $\vec{h}_{l-1}^{(t)}$. Thus for the l'th layer and t'th time step we have

$$\left(\vec{z}_{l}^{(t)}\right)_{lawer} = W^{l-1,l}\vec{h}_{l-1}^{(t)} + \vec{b}^{l-1,l}$$

and

$$\left(\vec{z}_l^{(t)}\right)_{time} = W^{ll}\vec{h}_l^{(t-1)} + \vec{b}^{ll},$$

that combine to give

$$\vec{z}_l^{(t)} = \left(\vec{z}_l^{(t)}\right)_{lawer} + \left(\vec{z}_l^{(t)}\right)_{time},$$

which in turn results in

$$\vec{h}_l^{(t)} = \sigma_l \left(\vec{z}_l^{(t)} \right).$$

This is also valid at the first time step by setting $\left(\vec{z}_l^{(1)}\right)_{\text{time}} = 0$.

The expression for the output layer is exactly the same as above, but with $(\vec{z}_l^{(t)})_{\text{time}} = 0$. Thus we have

$$\vec{z}_{L+1}^{(t)} = \left(\vec{z}_{L+1}^{(t)}\right)_{\mathrm{laver}} = W^{L,L+1} \vec{h}_L^{(t)} + \vec{b}^{L,L+1}$$

and

$$\hat{\vec{y}}^{(t)} = \sigma_{L+1} \left(\vec{z}_{L+1}^{(t)} \right)$$

The equations given for the forward propagation can seem a bit messy, so it is nice to have a more visual aid of what is going on. Here is a diagram of the complete RNN including the weights and biases relating the different nodes.

And here is a weights and biases connected to a single arbitrary node. The green arrows represent input to the node, and the red arrows represent the output from the node.

And here is the connections resulting in $\vec{h}_l^{(t)}$ in more detail.

3.3 Backpropagation through time (BPTT)

Backpropagation in an RNN works by comparing the output of the network to some target output (just as in the regular neural network), and propagating backwards through both the layers and the *time sequence*. It is therefore commonly referred to as *backpropagation through time* (BPTT). We will now derive the necessary equations to perform BPTT.

We assume that we have propagated forward through the network, and have produced some output $\hat{\vec{y}}^{(t)}$. We want to compare this with some target output value $\vec{y}^{(t)}$, and will do so through a cost function $C\left(\hat{\vec{y}},\vec{y}\right)$. We will denote the cost at a specific time step t by $C^{(t)}=C^{(t)}\left(\hat{\vec{y}}^{(t)},\vec{y}^{(t)}\right)$, and the overall cost of the network as C.

From the cost function at each time step, we want to compute the gradient with respect to each weight and bias, that is, we want to compute

$$\frac{\partial C}{\partial W^{l_1 l_2}}$$
 and $\frac{\partial C}{\partial \vec{b}^{l_1 l_2}}$

We will do this one layer at a time, starting at the output layer, and propagating backwards through time in each layer. We assume that we know the gradient of the cost function with respect to the output $\frac{\partial C^{(t)}}{\partial \hat{y}^{(t)}}$, and start by finding the gradient with respect to the output weights and biases $W^{L,L+1}$ and $\vec{b}^{L,L+1}$.

3.3.1 Backpropagation through the output layer

First, we want to find the gradient with respect to $\vec{z}_{L+1}^{(t)}$. The derivative of C with respect to some element $z_{L+1,i}^{(t)}$ of the weighted sum is given by

$$\begin{split} \frac{\partial C}{\partial z_{L+1,i}^{(t)}} &= \frac{\partial C^{(t)}}{\partial z_{L+1,i}^{(t)}} \\ &= \sum_{j=1}^{m} \frac{\partial C^{(t)}}{\partial \hat{y}_{j}^{(t)}} \frac{\partial \hat{y}_{j}^{(t)}}{\partial z_{L+1,i}^{(t)}} \\ &= \sum_{j=1}^{m} \frac{\partial C^{(t)}}{\partial \hat{y}_{j}^{(t)}} \sigma_{L+1}' \left(z_{L+1,i}^{(t)} \right) \delta_{ij} \\ &= \frac{\partial C^{(t)}}{\partial \hat{y}_{i}^{(t)}} \sigma_{L+1}' \left(z_{L+1,i}^{(t)} \right) \end{split}$$

where δ_{ij} is the Kronecker delta $\delta_{ij} = \begin{cases} 0, & i \neq j \\ 1, & i = j \end{cases}$, and σ'_{L+1} denotes the derivative of the activation function, which we will assume to be known. we can write this expression more compactly in vector form as

$$\frac{\partial C}{\partial \vec{z}_{L+1}^{(t)}} = \frac{\partial C^{(t)}}{\partial \hat{\vec{y}}^{(t)}} \odot \sigma_{L+1}' \left(\vec{z}_{L+1}^{(t)} \right),$$

where \odot denotes the *Hadamard product*, an elementwise multiplication of two vectors/matrices of same size.

Note: Sometimes the derivatives are real numbers like $\frac{\partial C^{(t)}}{\partial z_{L+1,i}^{(t)}}$, sometimes they are vectors such as $\frac{\partial C^{(t)}}{\partial z_{L+1}^{(t)}}$, and sometimes they are matrices. I have not included any explicit notation to explain when they are what, but will assume that this is understood implicitly. A general rule would be to look at whether the expression contains indices like i, j, k, \ldots or not.

Another note: There are a lot of indices to keep track of, so to make the notation simpler to follow I will try to follow these rules consistently: -l = layer index (with L being the final hidden layer). If I need several layer indices I will use $l_1, l_2, \ldots -(t) = \text{time step index}$. -i, j, k = vector/matrix elements. -n = number of input features (length of \vec{x}). -m = number of output features (length of \vec{y}). $-m_1, m_2, \ldots = \text{number of features}$ in hidden layer number $1, 2, \ldots$

Third note: I will not always write the upper bound of summations explicitly, but will assume that this is understood implicitly. For instance, $\sum_{j} W_{ij}^{l-1,l} h_{l-1,j}$ should be understood to mean $\sum_{j=1}^{n_{l-1}} W_{ij}^{l-1,l} h_{l-1,j}$, such that it sums over all elements of \vec{h}_{l-1} .

The derivative with respect to the weighted sum will be used a lot during backpropagation, so we will give it its own notation

$$\vec{\delta}_{L+1}^{(t)} \equiv \frac{\partial C^{(t)}}{\partial \vec{z}_{L+1}^{(t)}} = \frac{\partial C^{(t)}}{\partial \hat{\vec{y}}^{(t)}} \odot \sigma_{L+1}' \left(\vec{z}_{L+1}^{(t)} \right).$$

 $\delta_{L+1}^{(t)}$ has one index downstairs (denoting layer), and one index upstairs in parentheses (denoting time step), so don't mix it up with the Kronecker delta δ_{ij} , which I will consistently write with two indices downstairs.

From the delta we can find the cost gradient with respect to the output bias. Note that the same weights and biases occur several times in the RNN, so we have to sum over each contribution. The cost gradients with respect to the weights and biases in layer l are denoted $\frac{\partial C}{\partial W^{l-1,l}}$, $\frac{\partial C}{\partial W^{l}}$, $\frac{\partial C}{\partial V}$ and $\frac{\partial C}{\partial V}$, and we will denote the contribution at time step t as $\left(\frac{\partial C}{\partial W^{l-1,l}}\right)^{(t)}$, $\left(\frac{\partial C}{\partial W^{l}}\right)^{(t)}$, $\left(\frac{\partial C}{\partial V^{l-1,l}}\right)^{(t)}$ and $\left(\frac{\partial C}{\partial V^{l}}\right)^{(t)}$ such that $\frac{\partial C}{\partial W^{l-1,l}} = \sum_{t} \left(\frac{\partial C}{\partial W^{l-1,l}}\right)^{(t)}$ and so on. Using this notation, the gradient with respect to the output bias becomes

$$\begin{split} \left(\frac{\partial C}{\partial b_i^{L,L+1}}\right)^{(t)} &= \sum_{j=1}^m \frac{\partial C}{\partial z_{L+1,j}^{(t)}} \frac{\partial z_{L+1,j}^{(t)}}{\partial b_i^{L,L+1}} \\ &= \sum_{j=1}^m \frac{\partial C}{\partial z_{L+1,j}^{(t)}} \frac{\partial}{\partial b_i^{L,L+1}} \left(\sum_k W_{jk}^{L,L+1} h_{L,k}^{(t)} + b_j^{L,L+1}\right) \\ &= \sum_{j=1}^m \frac{\partial C}{\partial z_{L+1,j}^{(t)}} \delta_{ij} \\ &= \frac{\partial C}{\partial z_{L+1,i}^{(t)}} \\ &= \delta_{L+1,i}^{(t)}. \end{split}$$

Thus on vector form we have

$$\left(\frac{\partial C}{\partial \vec{b}^{L,L+1}}\right)^{(t)} = \vec{\delta}_{L+1}^{(t)},$$

and finally

$$\frac{\partial C}{\partial \vec{b}^{L,L+1}} = \sum_{t} \left(\frac{\partial C}{\partial \vec{b}^{L,L+1}} \right)^{(t)}$$

We can also compute the gradient with respect to the output weights

$$\begin{split} \left(\frac{\partial C}{W_{ij}^{L,L+1}}\right)^{(t)} &= \sum_{k_1=1}^m \frac{\partial C}{\partial z_{L+1,k_1}^{(t)}} \frac{\partial z_{L+1,k_1}^{(t)}}{\partial W_{ij}^{L,L+1}} \\ &= \sum_{k_1=1}^m \delta_{L+1,k_1}^{(t)} \frac{\partial}{\partial W_{ij}^{L,L+1}} \left(\sum_{k_2} W_{k_1k_2}^{L,L+1} h_{L,k_2}^{(t)} + b_{k_1}^{L,L+1}\right) \\ &= \sum_{k_1=1}^m \delta_{L+1,k_1}^{(t)} \sum_{k_2} h_{L,k_2}^{(t)} \delta_{ik_1} \delta_{jk_2} \\ &= \delta_{L+1,i}^{(t)} h_{L,j}^{(t)} \\ &= \left[\vec{\delta}_{L+1}^{(t)} \left(\vec{h}_L^{(t)}\right)^T\right]_{ij}. \end{split}$$

Thus on vector form we have

$$\left(\frac{\partial C}{W^{L,L+1}}\right)^{(t)} = \vec{\delta}_{L+1}^{(t)} \left(\vec{h}_L^{(t)}\right)^T,$$

and

$$\frac{\partial C}{W^{L,L+1}} = \sum_{t} \left(\frac{\partial C}{W^{L,L+1}} \right)^{(t)}.$$

Note that we here have an outer product between two vectors, which results in a matrix:

$$\vec{\delta}_{L+1}^{(t)} \left(\vec{h}_L^{(t)} \right)^T = \begin{pmatrix} \delta_{L+1,1}^{(t)} \\ \vdots \\ \delta_{L+1,m}^{(t)} \end{pmatrix} \left(h_{L,1}^{(t)} & \cdots & h_{L,n_L}^{(t)} \right) = \begin{pmatrix} \delta_{L+1,1}^{(t)} h_{L,1}^{(t)} & \cdots & \delta_{L+1,1}^{(t)} h_{L,n_L}^{(t)} \\ \vdots & \ddots & \vdots \\ \delta_{L+1,m}^{(t)} h_{L,1}^{(t)} & \cdots & \delta_{L+1,m}^{(t)} h_{L,n_L}^{(t)} \end{pmatrix}$$

Lastly, we need to compute the gradient with respect to the output from the previous layer $\frac{\partial C}{\partial \vec{h}_L^{(t)}}$, in order to continue backpropagating through previous layers. We find this in much the same way as we found the other gradients above.

$$\begin{split} \frac{\partial C}{\partial h_{L,i}^{(t)}} &= \sum_{j} \frac{\partial C}{z_{L+1,j}^{(t)}} \frac{\partial z_{L+1,j}^{(t)}}{\partial h_{L,i}^{(t)}} \\ &= \sum_{j} \delta_{L+1,j}^{(t)} \frac{\partial}{\partial h_{L,i}^{(t)}} \left(\sum_{k} W_{jk}^{L,L+1} h_{L,k}^{(t)} + b_{j}^{L,L+1} \right) \\ &= \sum_{j} \delta_{L+1,j}^{(t)} \sum_{k} W_{jk}^{L,L+1} \delta_{ik} \\ &= \sum_{j} \delta_{L+1,j}^{(t)} W_{ji}^{L,L+1} \\ &= \sum_{j} \left[\left(W^{L,L+1} \right)^{T} \right]_{ij} \delta_{L+1,j}^{(t)} \\ &= \left[\left(W^{L,L+1} \right)^{T} \vec{\delta}_{L+1}^{(t)} \right]_{i} \end{split}$$

And thus on vector form we have

$$\frac{\partial C}{\partial \vec{h}_L^{(t)}} = \left(W^{L,L+1}\right)^T \vec{\delta}_{L+1}^{(t)}$$

Here is a diagram showing the backpropagation through the output layer.

3.3.2 Backpropagation through arbitrary node

Consider some arbitrary node in the RNN with output $\vec{h}_l^{(t)}$. Assume you know the total gradient of the cost with respect to this output from the two succeeding nodes

$$\frac{\partial C}{\partial \vec{h}_l^{(t)}} = \left(\frac{\partial C}{\partial \vec{h}_l^{(t)}}\right)_{\text{layer}} + \left(\frac{\partial C}{\partial \vec{h}_l^{(t)}}\right)_{\text{time}}.$$

We now want to compute the gradients with respect to the weights and biases connecting the two previous nodes to this node, so that we can update these weights and biases when training the network, as well as the gradient with respect to the two previous nodes, so that we can continue backpropagation through the other nodes. The situation is illustrated in the diagram below. The blue arrows show the input gradient from the succeeding nodes, and the red arrows show the gradients we want to compute.

The necessary gradients are derived in the same way as for the output layer, so I will simply state the results here. We get the following set of equations for backpropagating through a general node in the RNN.

$$\delta_l^{(t)} = \frac{\partial C}{\partial \vec{h}_l^{(t)}} \odot \sigma_l' \left(\vec{z}_l^{(t)} \right) \tag{1}$$

$$\left(\frac{\partial C}{\partial \vec{b}^{l-1,l}}\right)^{(t)} = \left(\frac{\partial C}{\partial \vec{b}^{ll}}\right)^{(t)} = \delta_l^{(t)} \tag{2}$$

$$\left(\frac{\partial C}{\partial W^{l-1,l}}\right)^{(t)} = \delta_l^{(t)} \left(\vec{h}_{l-1}^{(t)}\right)^T \tag{3}$$

$$\left(\frac{\partial C}{\partial W^{ll}}\right)^{(t)} = \delta_l^{(t)} \left(\vec{h}_l^{(t-1)}\right)^T \tag{4}$$

$$\frac{\partial C}{\partial \vec{h}_{l-1}^{(t)}} = \left[\left(W^{l-1,l} \right)^{(t)} \right]^T \delta_l^{(t)} \tag{5}$$

$$\frac{\partial C}{\partial \vec{h}_l^{(t-1)}} = \left[\left(W^{ll} \right)^{(t-1)} \right]^T \delta_l^{(t)}, \tag{6}$$

and

$$\frac{\partial C}{\partial \vec{b}^{l-1,l}} = \sum_{t} \left(\frac{\partial C}{\partial \vec{b}^{l-1,l}} \right)^{(t)} \tag{7}$$

$$\frac{\partial C}{\partial \vec{b}^{ll}} = \sum_{t} \left(\frac{\partial C}{\partial \vec{b}^{ll}} \right)^{(t)} \tag{8}$$

$$\frac{\partial C}{\partial W^{l-1,l}} = \sum_{t} \left(\frac{\partial C}{\partial W^{l-1,l}} \right)^{(t)} \tag{9}$$

$$\frac{\partial C}{\partial W^{ll}} = \sum_{t} \left(\frac{\partial C}{\partial W^{ll}} \right)^{(t)}.$$
 (10)

With this method we can start with the nodes in the output layer, and propagate backwards. The necessary input to one node is the output from backpropagating through the previous node. Thus we can use the equations above recursively, layer by layer, to backpropagate through the entire network.

4 The RNN code

Now that we have the mathematical framework, we can develop the code for the RNN.

4.1 Functions

Before we start building a recurrent neural network, we need to define some functions. We need activation functions, cost functions and a way to differentiate these. We also need gradient descent schedulers to update our weights and biases when backpropagating. These functions are defined in this section.

4.1.1 Activation functions

We want to be able to choose which activation function to use in different layers of the RNN. Here we define some activation functions that can be used by the network. If you have developed a regular fully connected neural network in FYS-STK4155 these functions will probably look very familiar, as they are pretty much copied from those lecture notes. The main difference is that I have used JAX instead of autograd for automatic differentiation. Due to the way JAX vectorizes, I have not gotten gradients with jax to work for softmax, but have included grad_softmax() as its own function.

```
[1]: import numpy as np
     import jax.numpy as jnp
     def identity(X):
         return X
     def sigmoid(X):
         try:
             return 1.0 / (1 + jnp.exp(-X))
         except FloatingPointError:
             return jnp.where(X > jnp.zeros(X.shape), jnp.ones(X.shape), jnp.zeros(X.
      ⇒shape))
     def softmax(X):
         X = X - np.max(X, axis=-1, keepdims=True)
         delta = 10e-10
         return np.exp(X) / (np.sum(np.exp(X), axis=-1, keepdims=True) + delta)
     def grad softmax(X):
         f = softmax(X)
         return f - f**2
     def RELU(X):
         return jnp.where(X > jnp.zeros(X.shape), X, jnp.zeros(X.shape))
     def LRELU(X):
```

```
delta = 10e-4
  return jnp.where(X > jnp.zeros(X.shape), X, delta * X)

def tanh(X):
  return jnp.tanh(X)
```

4.1.2 Cost functions

Next, we need to implement the cost functions. We include three cost functions here. The ordinary least squares (OLS) is used for regression problems, and the logistic regression and cross-entropy are used for classification problems.

4.1.3 Automatic differentiation

As mentioned above, we use JAX for automatic differentiation, which is done with the function grad in the JAX library. For grad to work on a function, it cannot use regular numpy, but must use jax.numpy, which is why we imported and used this when defining the activation and cost functions. JAX's numpy is only used for these functions, while we stick with regular numpy for everything else.

The RELU and leaky RELU activation functions are not continuously differentiable, so we will handle these explicitly in our code.

```
[3]: from jax import grad

def derivate(func):
    if func.__name__ == "RELU":

        def func(X):
            return jnp.where(X > 0, 1, 0)

        return func

    elif func.__name__ == "LRELU":

        def func(X):
            delta = 10e-4
            return jnp.where(X > 0, 1, delta)

        return func

    else:
        return grad(func)
```

Note that the *grad* function is not, in itself, vectorized. This means that if we send in an array to a function that has been differentiated, JAX will treat this as a function with an array as input, not as a function treating each element individually. This is better understood with an example.

Consider the function $f(x) = x^2$, with derivative f'(x) = 2x. With JAX we get this with

```
[4]: def f(x):
    return x**2

df = grad(f)
```

This works if we input a scalar value

```
[5]: x = 2.0 
print(df(x))
```

4.0

But if we try to input an array of values we get an error message.

```
[6]: x = np.linspace(0, 3, 10)

try:
    print(df(x))
except TypeError as msg:
    print(f"Error message: {msg}")
```

Error message: Gradient only defined for scalar-output functions. Output had shape: (10,).

This is because JAX does not treat each element in x individually. To get around this we can use the vmap function in the JAX library, which vectorizes the function.

```
[7]: from jax import vmap

df = vmap(grad(f))

x = np.linspace(0, 3, 10)

print(df(x))

[0. 0.6666667 1.3333334 2. 2.6666667 3.3333333 4.
```

Note that vmap only vectorizes along one dimension. So if the input array contains several axes, we have to apply vmap for each axis.

```
[8]: df = vmap(vmap(ymap(grad(f))))
     x = np.linspace(0,3, 27).reshape((3,3,3))
     print(df(x))
    [[[0.
                  0.23076923 0.46153846]
                  0.9230769
      [0.6923077
                            1.1538461 ]
      [1.3846154    1.6153846    1.8461539 ]]
     [[2.0769231 2.3076923 2.5384614]
      [2.7692308
                             3.2307692 1
      [3.4615386 3.6923077 3.9230769]]
     [[4.1538463 4.3846154
                            4.6153846 ]
      [4.8461537 5.076923
                             5.3076925 ]
```

This approach is tedious, yes, but as far as I know it is the only way to make JAX differentiate elementwise. If you find a simpler work-around, feel free to share with the professor or the group teachers so these notes can be updated.

]]]

4.1.4 Schedulers

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We also want to be able to choose which method we want to use for gradient descent when training the RNN. This is done by using a scheduler defined below. These schedulers are identical to the ones used in FYS-STK4155.

```
[9]: class Scheduler:
    """
    Abstract class for Schedulers
    """

def __init__(self, eta):
    self.eta = eta
```

```
# should be overwritten
    def update_change(self, gradient):
        raise NotImplementedError
    # overwritten if needed
    def reset(self):
        pass
class Constant(Scheduler):
    def __init__(self, eta):
        super().__init__(eta)
    def update_change(self, gradient):
        return self.eta * gradient
    def reset(self):
        pass
class Momentum(Scheduler):
    def __init__(self, eta: float, momentum: float):
        super().__init__(eta)
        self.momentum = momentum
        self.change = 0
    def update_change(self, gradient):
        self.change = self.momentum * self.change + self.eta * gradient
        return self.change
    def reset(self):
        pass
class Adagrad(Scheduler):
    def __init__(self, eta):
        super().__init__(eta)
        self.G_t = None
    def update_change(self, gradient):
        delta = 1e-8 # avoid division ny zero
        if self.G_t is None:
            self.G_t = np.zeros((gradient.shape[0], gradient.shape[0]))
        self.G_t += gradient @ gradient.T
```

```
G_t_inverse = 1 / (
            delta + np.sqrt(np.reshape(np.diagonal(self.G_t), (self.G_t.
 \hookrightarrowshape[0], 1)))
        return self.eta * gradient * G_t_inverse
    def reset(self):
        self.G_t = None
class AdagradMomentum(Scheduler):
    def __init__(self, eta, momentum):
        super().__init__(eta)
        self.G_t = None
        self.momentum = momentum
        self.change = 0
    def update_change(self, gradient):
        delta = 1e-8 # avoid division ny zero
        if self.G_t is None:
            self.G_t = np.zeros((gradient.shape[0], gradient.shape[0]))
        self.G_t += gradient @ gradient.T
        G_t_inverse = 1 / (
            delta + np.sqrt(np.reshape(np.diagonal(self.G_t), (self.G_t.
 ⇔shape[0], 1)))
        self.change = self.change * self.momentum + self.eta * gradient *∟
 \hookrightarrow G_t_inverse
        return self.change
    def reset(self):
        self.G_t = None
class RMS_prop(Scheduler):
    def __init__(self, eta, rho):
        super().__init__(eta)
        self.rho = rho
        self.second = 0.0
    def update_change(self, gradient):
        delta = 1e-8 # avoid division ny zero
```

```
self.second = self.rho * self.second + (1 - self.rho) * gradient *_
 ⇔gradient
        return self.eta * gradient / (np.sqrt(self.second + delta))
    def reset(self):
        self.second = 0.0
class Adam(Scheduler):
    def __init__(self, eta, rho, rho2):
        super().__init__(eta)
        self.rho = rho
        self.rho2 = rho2
        self.moment = 0
        self.second = 0
        self.n_epochs = 1
    def update_change(self, gradient):
        delta = 1e-8 # avoid division ny zero
        self.moment = self.rho * self.moment + (1 - self.rho) * gradient
        self.second = self.rho2 * self.second + (1 - self.rho2) * gradient * |
 ⇔gradient
        moment_corrected = self.moment / (1 - self.rho**self.n_epochs)
        second_corrected = self.second / (1 - self.rho2**self.n_epochs)
        return self.eta * moment_corrected / (np.sqrt(second_corrected + delta))
    def reset(self):
        self.n_epochs += 1
        self.moment = 0
        self.second = 0
```

4.2 The RNN

We will now implement the code for the RNN. The network will be object-oriented, consisting of the following classes:

- RNN: The complete network, consisting of several Layer objects.
- Layer: Abstract class containing information that is shared across the different types of layers. It is the parent class of the following:
 - InputLayer: Layer containing the input to the network. Does not contain any weights and biases.
 - RNNLayer: The recurrent layer consisting of nodes in sequence.
 - OutputLayer: Recurrent layer for output. Similar to RNNLayer, but does not have any
 connections between the nodes, only to the nodes at the same time step in the previous
 layer.
 - DenseLayer: Fully connected layer, used to switch from a recurrent network to a regu-

lar fully connected network. Especially used for non-sequential output, for instance in classification where you want to classify the entire sequence with a single output.

• *Node*: Contains information about a single node. This is where all the math of forward- and backpropagation takes place.

In our code we want to be able to input batches of values, so that we can feed forward (and backpropagate) many inputs at once. The input to our network will therefore be an array with three axes:

- The input axis, separating the different inputs in the batch.
- The sequence axis, separating the different time steps of the sequence.
- The feature axis, separating the different features of the input (the vector elements).

This input is fed forward through the network, thus each layer has the same three axes. Within the layers, we separate each time step into their own node, thus the nodes only have two axes: the input and feature axes.

Since the RNN class is dependent on the Layer classes, and the Layer classes are dependent on the Node class, we will build the network from the down up, starting with the Node class.

4.2.1 The Node class

The Node class takes care of the math discussed in the sections Forward propagation and Back-propagation through arbitrary node. I restate the relevant equations below.

Forward propagation:

$$\begin{split} \left(\vec{z}_l^{(t)}\right)_{layer} &= W^{l-1,l} \vec{h}_{l-1}^{(t)} + \vec{b}^{l-1,l} \\ \\ \left(\vec{z}_l^{(t)}\right)_{time} &= W^{ll} \vec{h}_l^{(t-1)} + \vec{b}^{ll} \\ \\ \vec{z}_l^{(t)} &= \left(\vec{z}_l^{(t)}\right)_{layer} + \left(\vec{z}_l^{(t)}\right)_{time} \\ \\ \vec{h}_l^{(t)} &= \sigma_l \left(\vec{z}_l^{(t)}\right) \end{split}$$

Backpropagation:

$$\begin{split} \frac{\partial C}{\partial \vec{h}_l^{(t)}} &= \left(\frac{\partial C}{\partial \vec{h}_l^{(t)}}\right)_{\text{layer}} + \left(\frac{\partial C}{\partial \vec{h}_l^{(t)}}\right)_{\text{time}} \\ \delta_l^{(t)} &= \frac{\partial C}{\partial \vec{h}_l^{(t)}} \odot \sigma_l' \left(\vec{z}_l^{(t)}\right) \\ \left(\frac{\partial C}{\partial \vec{b}^{l-1,l}}\right)^{(t)} &= \left(\frac{\partial C}{\partial \vec{b}^{ll}}\right)^{(t)} = \delta_l^{(t)} \\ \left(\frac{\partial C}{\partial W^{l-1,l}}\right)^{(t)} &= \delta_l^{(t)} \left(\vec{h}_{l-1}^{(t)}\right)^T \\ \left(\frac{\partial C}{\partial W^{ll}}\right)^{(t)} &= \delta_l^{(t)} \left(\vec{h}_l^{(t-1)}\right)^T \\ \frac{\partial C}{\partial \vec{h}_{l-1}^{(t)}} &= \left[\left(W^{l-1,l}\right)^{(t)}\right]^T \delta_l^{(t)} \\ \frac{\partial C}{\partial \vec{h}_l^{(t-1)}} &= \left[\left(W^{ll}\right)^{(t-1)}\right]^T \delta_l^{(t)}. \end{split}$$

Note: The input and output from a node are two-dimensional arrays, with different inputs along the first axis, and the features along the second axis. The equations we have derived for computing forward- and backpropagation assume that we are working with column vectors, but when performing matrix multiplication with these two-dimensional arrays we are in practice working with row vectors. This has to be taken into account in our code.

Consider, for instance, multiplying some weight matrix W with a vector \vec{h} . The way we have derived the equations this will look like

$$W\vec{h} = \begin{pmatrix} W_{11} & \cdots & W_{1n} \\ \vdots & \ddots & \vdots \\ W_{m1} & \cdots & W_{mn} \end{pmatrix} \begin{pmatrix} h_1 \\ \vdots \\ h_n \end{pmatrix}.$$

In the code, however, this will look like

$$\vec{h}W = \begin{pmatrix} h_{11} & \cdots & h_{1n} \\ \vdots & \ddots & \vdots \\ h_{N1} & \cdots & h_{Nn} \end{pmatrix} \begin{pmatrix} W_{11} & \cdots & W_{1m} \\ \vdots & \ddots & \vdots \\ W_{n1} & \cdots & W_{nm} \end{pmatrix},$$

where N is the number of inputs. Note that W and \vec{h} have switched places in the matrix multiplication, and that W is essentially a transpose of the corresponding W in the previous case (since we are multiplying the columns with the \vec{h} -vectors, instead of the rows).

This is not a big difference, but can be taken into account by reversing the order of some of the matrix multiplications. The full set of changes that must be done to account for this are listed here:

$$W\vec{h} \to \vec{h}W$$

 $\vec{\delta}\vec{h}^T \to \vec{h}^T\vec{\delta}$
 $W^T\vec{\delta} \to \vec{\delta}W^T$,

and are taken care of in the code below.

```
[10]: from collections.abc import Callable # Used for type hints of functions
      class Node:
           11 11 11
          Single node in the RNN. Computes forward propagation through a single node, _
       \hookrightarrowstores the output of as a vector (1D array) of length 'n_features'. This
       \hookrightarrow class also computes the backpropagation through the node, and stores all the \sqcup
       \negrelevant gradients.
          Attributes
          n_features (int)
               Number of features in this node (the length of the vector).
          act_func (Callable)
               The activation function of this node.
           {b/W} {layer/time} (ndarray)
               {\it Bias}(b) or weight matrix (W) for computing the contribution to this _{\sqcup}
        ⇔node from the node in the previous layer/time step.
          h_{layer/time/output} (ndarray)
               The output of the node in the previous layer/time step, or the output \sqcup
        ⇔of this node.
           z_output (ndarray)
               The weighted sum output of this node (the output before activation).
           grad_{b/W/h}_{layer/time} (ndarray)
               The gradient of the cost function with respect to the appropriate_
        \neg variable.
           11 11 11
          def __init__(
                   self,
                   n_features: int,
                   act_func: Callable[[np.ndarray], np.ndarray] = identity,
```

```
W_layer: np.ndarray = None,
          b_layer: np.ndarray = None,
          W_time: np.ndarray = None,
          b_time: np.ndarray = None
  ):
      Constructor for Node objects.
      Parameters
      n features (int)
          Number of features in this node (the length of the vector).
      act func (Callable)
          The activation function of this node.
      {b/W}_{layer/time} (ndarray)
          Bias(b) or weight matrix (W) for computing the contribution to this.
⇔node from the node in the previous layer/time step.
      self.n_features = n_features
      self.act_func = act_func
      self.W layer = W layer
      self.b_layer = b_layer
      self.W_time = W_time
      self.b_time = b_time
      ## Values from feed_forward()
      self.h_layer = None # h from previous layer
      self.h_time = None # h from previous time step
      self.z_output = None # z for this node
      self.h_output = None # h from this node
      ## Values from backpropagate()
      self.grad_b_layer = None
      self.grad b time = None
      self.grad_W_layer = None
      self.grad_W_time = None
      self.grad_h_layer = None
      self.grad_h_time = None
  def set_Wb(
          self,
          W_layer: np.ndarray,
          b_layer: np.ndarray,
          W_time: np.ndarray = None,
          b_time: np.ndarray = None
  ):
      11 11 11
```

```
Sets the weights and biases to specific values. Used by the layer ...
→classes to ensure all nodes have correct weights and biases.
      Parameters
       _____
       {b/W} {layer/time} (ndarray)
           Bias(b) or weight matrix (W) for computing the contribution to this.
⇔node from the node in the previous layer/time step.
      self.W_layer = W_layer
      self.b_layer = b_layer
      self.W time = W time
      self.b_time = b_time
  def feed_forward(
           self,
           h_layer: np.ndarray,
          h_time: np.ndarray = None
  ) -> np.ndarray:
       Computes the output of this node from the output of the nodes at the \sqcup
oprevious layer and time step, using the equations for forward propagation.
       Parameters
       _____
       h_{layer/time/output} (ndarray)
           The output of the node in the previous layer/time step.
      Returns
      h output (ndarray)
           The output of this node.
       ## Save h_layer and h_time for use in backpropagation
      self.h_layer = h_layer
      self.h_time = h_time
      num_inputs = h_layer.shape[0]
       ## Compute weighted sum z for this node.
       z_layer = h_layer @ self.W_layer + self.b_layer # Dimension example:
\hookrightarrow (100,5)@(5,7) + (7) = (100,7)
       if h_time is None:
           # This node is at the first time step, thus not receiving any input _{\sqcup}
⇔ from previous time steps.
           z_time = np.zeros((num_inputs, self.n_features))
```

```
else:
           z_time = h_time @ self.W_time + self.b_time
       self.z_output = z_layer + z_time # Save the weighted sum in the node
       ## Compute activation of the node
       h_output = self.act_func(self.z_output)
       self.h output = h output # Save the output in the node
       return h_output # Return output
   def backpropagate(
           self,
           dC_layer: np.ndarray = None,
           dC_time: np.ndarray = None,
           lmbd: float = 0.01
   ):
       Performs backpropagation through this node. Computes the gradient of \Box
\hookrightarrow the cost function with respect to the weights and biases of this layer (and \sqcup
stores these gradients so the weights and biases can be updated in the layer
\hookrightarrowobject this node belongs to), and the gradient with respect to the output of \sqcup
_{\hookrightarrow} the nodes in the previous layer and time step (and stores these as well for _{\sqcup}
→ further backpropagation through the network).
       Parameters
       dC_{layer/time} (ndarray)
           Contribution of cost gradient w.r.t. this node from node at next_{\sqcup}
⇔layer/time.
       lmbd (float)
           Regularization parameter for finding the cost gradient with respect_{\sqcup}
\hookrightarrow to the weights.
       n_batches = self.h_output.shape[0]
       ## Total gradient is the sum of the gradient from "next" layer and time
       if dC time is None:
            # If this is the last node in the layer, the gradient is just the
→ gradient from the next layer
           dC = dC_layer
       elif dC_layer is None:
            # If the next layer has no node at this time step (because it is a_{\sqcup}
\hookrightarrowSingleOutputLayer), use only dC_time
           dC = dC_{time}
       else:
```

```
dC = dC_layer + dC_time
       ## delta (gradient of cost w.r.t. z)
       if self.act_func.__name__ == "softmax":
           grad_act = grad_softmax(self.z_output)
       else:
           grad_act = vmap(vmap(derivate(self.act_func)))(self.z_output) #__
→vmap is necessary for jax to vectorize gradient properly
       delta = grad_act * dC # Hadamard product, i.e., elementwise_
\hookrightarrow multiplication
       ## Gradients w.r.t. bias
       self.grad_b_layer = np.sum(delta, axis=0) / n_batches
       self.grad_b_time = np.sum(delta, axis=0) / n_batches
       ## Gradients w.r.t. weights
       self.grad_W_layer = self.h_layer.T @ delta / n_batches
       self.grad_W_layer = self.grad_W_layer + self.W_layer * lmbd #__
→ Regularization factor
       if self.h_time is None:
           self.grad W time = None
       else:
           self.grad_W_time = self.h_time.T @ delta / n_batches
           self.grad_W_time = self.grad_W_time + self.W_time * lmbd #_
→ Regularization factor
       ## Gradients w.r.t. input from previous nodes
       # Need to not transpose delta in order for matrices to match up_
-correctly, since we have batches along rows, and features along columns
       self.grad_h_layer = delta @ self.W_layer.T
       if self.h time is None:
           self.grad_h_time = None
       else:
           self.grad_h_time = delta @ self.W_time.T
```

4.2.2 The Layer class

The RNN consists of several layers of various types. The Layer class is an abstract class keeping track of attributes and methods that are common for the different types of layers. We define this class for better organization of the different layers, but it should never be used by itself, for instance by creating a Layer object. You will notice that most of the Layer methods are only declared, not implemented. These methods will vary between layer types, and are implemented in the Layer's child classes, although the Layer class gives a nice overview of which methods to expect from its children.

```
[11]: from __future__ import annotations # Necessary to create typing hint of Layer_
       ⇔within the class Layer
      class Layer:
           HHHH
          Abstract class for layers. The attributes given here are the attributes \sqcup
       \hookrightarrowthat are common to all layers, but they will also contain other attributes_{\sqcup}
       \hookrightarrow in addition to these.
          Attributes
           _____
          n_features (int)
               Number of features in the nodes of this layer (the length of the \Box
       \neg vectors).
          seed (int)
               Seed for random number generating with numpy.
          nodes (list)
               List containing all the nodes of the layer in sequence, such that \sqcup
        \neg nodes[0] is the node at the first time step, nodes[1] at the second time_\(\pi\)
        \hookrightarrowstep, and so on.
           n nodes (int)
               Number of nodes in the layer. Is updated when adding or removing nodes.
          def __init__(
                   self,
                   n_features: int,
                   seed: int = 100
          ):
               Constructor to be called from child classes.
               Parameters
               ____
               n_features (int)
                   Number of features in the nodes of this layer (the length of the \sqcup
        \neg vectors).
               seed (int)
                   Seed for random number generating with numpy.
               self.n_features = n_features
               self.seed = seed
               self.nodes = []
               self.n_nodes = 0
          def reset_weights(self):
               pass
```

```
def reset_schedulers(self):
      pass
  def update_weights_all_nodes(self):
       raise NotImplementedError
  def add_node(self):
       raise NotImplementedError
  def remove_nodes(self):
       Remove all the nodes created for this layer.
       NOTE
       The weights and biases of the nodes are still stored in the layer, so_{\sqcup}
⇔we can easily
       create new nodes. Removing the nodes is used to allow the sequence \sqcup
⇔length to vary with
       each call of feed_forward().
      self.nodes = []
       self.n_nodes = 0
  def feed_forward(
           self,
           prev_layer: Layer
  ):
      raise NotImplementedError
  def backpropagate(
           self,
           next_layer: Layer,
           lmbd: float
  ):
      raise NotImplementedError
```

4.2.3 The InputLayer class

When we want to feed forward through an RNN, we will send a 3-dimensional array of shape (batch size, sequence length, number of features) as input. We then want to feed forward through the first hidden layer by computing the output of each node separately. Each node should then get information from the node at the previous time step, and the node at the previous layer. The way we implement forward propagation in this notebook, we need the previous layer to contain one node for each time step, and feed forward through these nodes. To make sure this also works for the first hidden layer, we include the InputLayer class, which takes the 3-dimensional input array to the network, and stores the value at each time step as 2-dimensional arrays in separate nodes.

For all subsequent layers, we can then send in the previous layer as input to the feed_forward() method, work with each node separately.

Note that feed_forward() will take a layer class as input for all other layers, but since this is the first layer (with no preceding layer), it will instead take in the 3-dimensional input array.

```
[12]: class InputLayer(Layer):
           nnn
           The input layer of the RNN, used for storing the input to the RNN in nodes \Box
        →for easy forward propagation through subsequent layers.
           This class does not contain any weights or biases, nor does it implement \sqcup
       ⇒backpropagation as there are no parameters to update.
           InputLayer is the only class that can function as the first layer in the \Box
       \hookrightarrow RNN, and it should only be used for this purpose.
          Attributes
          n_features (int)
               Number of features in the nodes of this layer (the length of the \sqcup
        \neg vectors).
           seed (int)
               Seed for random number generating with numpy.
          nodes (list)
               List containing all the nodes of the layer in sequence, such that \sqcup
        \neg nodes[0] is the node at the first time step, nodes[1] at the second time,
        \hookrightarrowstep, and so on.
           n nodes (int)
               Number of nodes in the layer. Is updated when adding or removing nodes.
          def __init__(
                   self,
                   n_features: int,
                   seed: int = 100
          ):
               11 11 11
               Constructor for InputLayer objects.
               Parameters
               _____
               n features (int)
                   Number of features in the nodes of this layer (the length of the \sqcup
        \neg vectors).
               seed (int)
                   Seed for random number generating with numpy.
               super().__init__(n_features, seed)
```

```
def add_node(self):
       n n n
       Add a node. Weights and biases are set to None, and activation to_{\sqcup}
⇒identity by default,
       as none of these are relevant for the input layer.
       new\_node = Node(self.n\_features) # Activation and weights are not used_{\sqcup}
→ for the input layer
       self.nodes.append(new_node)
       self.n_nodes += 1
  def feed forward(
           self,
           X: np.ndarray
  ):
       Feed forward through the layer. As this is the input layer, this ...
\negamounts to adding nodes and setting the output of each node to the value of \sqcup
_{	o}the input at the corresponding time step. Also checks that the number of_{\sqcup}
ofeatures of the input is the same as the layer expects it to be.
       Parameters
       X (ndarray)
           Input to the RNN, and thus to this layer. X has shape (batch size, \Box
⇒sequence length, number of features)
       NOTE
       Unlike the other layers, this layer takes a numpy array as input_{\sqcup}
⇒instead of a Layer, since it is the first layer.
       X \text{ shape} = X.\text{shape}
       sequence_length = X_shape[1]
       if not self.n_features == X_shape[2]:
           # Input must have the same number of features as defined by the
\hookrightarrow layer
           raise ValueError(f"Expected the number of features in the input⊔
→layer to be {self.n_features}, got {X_shape[2]}.")
       # Add a node to the layer for each time step, and set output
       self.remove_nodes()
       for i in range(sequence_length):
           self.add_node()
           self.nodes[i].h_output = X[:,i,:]
```

4.2.4 The RNNLayer class

The RNNLayer class represents the recurrent layers in the RNN. It takes the value at each time step in the preceding layer and computes the output from each node in its own layer, propagating forward through each node one by one, taking into account both the connections between the layers and the time steps. The math behind forward- and backpropagation are already taken care of by the Node class, so the RNN class is actually quite simple. In forward propagation, it simply goes through each node, inserts the output from the nodes at the previous layer and time step, and lets the nodes themselves do the rest, including storing the results. In backpropagation it goes through each node in reversed order, inserts the gradient of the cost function with respect to their output, lets the nodes calculate the gradients with respect to weights and biases, and then uses these to update the weights and biases of the layer.

```
[13]: from copy import copy
      class RNNLayer(Layer):
          The recurrent layer of the RNN. Computes forward- and backpropagation,
       →through a recurrent layer by feeding forward, or backpropagating, through
       \rightarroweach
          node of the layer one at a time in sequence.
          Attributes
          n_features (int)
              Number of features in the nodes of this layer (the length of the ...
       \neg vectors).
          n features prev (int)
               Number of features in the nodes of the preceding layer (the length of \Box
       \hookrightarrow the vectors).
          seed (int)
               Seed for random number generating with numpy.
          nodes (list)
              List containing all the nodes of the layer in sequence, such that \sqcup
       \neg nodes[0] is the node at the first time step, nodes[1] at the second time
       \hookrightarrowstep, and so on.
          n_nodes (int)
               Number of nodes in the layer. Is updated when adding or removing nodes.
          act_func (Callable)
               The activation function of this layer.
          {b/W}_{layer/time} (ndarray)
               Bias (b) or weight matrix (W) for computing the forward propagation,
       →from nodes at the previous layer to this one (layer),
               or between time steps within this layer (time).
          {b/W}_{layer/time}_size (tuple)
               The shape of b_layer/W_layer/b_time/W_time.
          scheduler_{b/W}_{layer/time} (Scheduler)
```

```
The scheduler for updating b_layer/W_layer/b_time/W_time with gradient<sub>\sched</sub>
⇒descent when backpropagating through this layer.
   is_dense (bool)
       Tells if this is a DenseLayer or not. It is set to False for this⊔
⇒layer, as it is not the DenseLayer.
   n n n
  def __init__(
           self.
           n_features: int,
           n_features_prev: int,
           act_func: Callable[[np.ndarray], np.ndarray],
           scheduler: Scheduler,
           seed: int = 100
  ):
       Constructor for RNNLayer objects.
       Parameters
       _____
       n_features (int)
           Number of features in the nodes of this layer (the length of the ...
\neg vectors).
       n features prev (int)
           Number of features in the nodes of the preceding layer (the length \sqcup
\hookrightarrow of the vectors).
       act_func (Callable)
           The activation function of this layer.
       scheduler (Scheduler)
           The scheduler to use for updating weights and biases with gradient \sqcup
⇒descent when backpropagating through this layer.
       seed (int)
           Seed for random number generating with numpy.
       super().__init__(n_features, seed)
       self.n_features_prev = n_features_prev
       self.act_func = act_func
       self.W_layer = None
       self.b_layer = None
       self.W_time = None
       self.b time = None
       self.W_layer_size = (self.n_features_prev, self.n_features)
       self.b_layer_size = (1, self.n_features)
       self.W_time_size = (self.n_features, self.n_features)
       self.b_time_size = (1, self.n_features)
```

```
self.scheduler_W_layer = copy(scheduler)
      self.scheduler_W_time = copy(scheduler)
      self.scheduler_b_layer = copy(scheduler)
      self.scheduler_b_time = copy(scheduler)
      self.is_dense = False
      self.reset weights()
  def reset weights(self):
      Reset weights and biases to random values from a normal distribution.
      np.random.seed(self.seed)
      self.W_layer = np.random.normal(size=self.W_layer_size)
      self.b_layer = np.random.normal(size=self.b_layer_size) * 0.01
      self.W_time = np.random.normal(size=self.W_time_size)
      self.b_time = np.random.normal(size=self.b_time_size) * 0.01
  def reset_schedulers(self):
      11 11 11
      Reset the schedulers of the layer.
      self.scheduler_W_layer.reset()
      self.scheduler b layer.reset()
      self.scheduler_W_time.reset()
      self.scheduler_b_time.reset()
  def update_weights_all_nodes(self):
      Update the weights and biases in all nodes of the layer.
      new_W_layer = self.W_layer
      new_W_time = self.W_time
      new_b_layer = self.b_layer
      new_b_time = self.b_time
      for node in self.nodes:
          node.set_Wb(new_W_layer, new_b_layer, new_W_time, new_b_time)
  def add node(self):
      Add a node with the weights and biases specified by the layer.
      new_node = Node(self.n_features, self.act_func, self.W_layer, self.
⇒b_layer, self.W_time, self.b_time)
      self.nodes.append(new_node)
```

```
self.n_nodes += 1
  def feed_forward(
           self,
           prev_layer: Layer
  ):
       11 11 11
      Feed forward through this layer one node at a time. The results are \sqcup
⇔stored in the nodes.
      Parameters
       _____
      prev_layer (Layer)
           The preceding layer of the RNN.
      self.remove_nodes()
      n_nodes_prev = prev_layer.n_nodes
      for i in range(n_nodes_prev):
           # Get output of node from previous layer
           prev_layer_node = prev_layer.nodes[i]
           h_layer = prev_layer_node.h_output
           # Get output of node from previous time step
           if i == 0:
               # No previous node if this is the first time step
               h_time = None
           else:
               prev_time_node = self.nodes[i-1]
               h_time = prev_time_node.h_output
           # Create and compute new node at this time step
           self.add_node()
           new_node = self.nodes[i]
           new_node.feed_forward(h_layer, h_time)
  def backpropagate(
           self,
           next_layer: Layer,
           lmbd: float = 0.01
  ):
       Backpropagate through the layer one node at a time. The results are \Box
\hookrightarrowstored in the nodes.
      Parameters
```

```
next_layer (Layer)
           The subsequent layer of the RNN.
       lmbd (float)
           Regularization parameter for finding the cost gradient with respect \sqcup
\hookrightarrow to the weights.
       11 11 11
       ## Check if the next layer is a DenseLayer
      next_is_dense = next_layer.is_dense
       ## Go through all nodes, starting with the last
      for i in range(self.n_nodes-1, -1, -1):
           ## Gradient from node at next layer
           if next_is_dense:
               ## If next layer is DenseLayer, consider only grad_h_layer in_
→ the last node in this layer
               if i == self.n_nodes-1:
                   node_layer = next_layer.nodes[0]
                   dC_layer = node_layer.grad_h_layer
               else:
                   dC_layer = None
           else:
               ## If next layer is not DenseLayer, get gradient from all nodes
               node_layer = next_layer.nodes[i]
               dC_layer = node_layer.grad_h_layer
           ## Gradient from node at next time step (unless this is the last
⇔node)
           if i == self.n_nodes-1:
               dC_time = None
           else:
               node_time = self.nodes[i+1]
               dC_time = node_time.grad_h_time
           ## Backpropagate through this node. Results are stored in the nodes
           node = self.nodes[i]
           node.backpropagate(dC_layer, dC_time, lmbd)
           ## Update weights and biases
           grad_W_layer = node.grad_W_layer / self.n_nodes
           grad_W_time = node.grad_W_time
           grad_b_layer = node.grad_b_layer / self.n_nodes
           grad_b_time = node.grad_b_time
           self.W_layer -= self.scheduler_W_layer.update_change(grad_W_layer)
           if grad_W_time is not None:
               grad_W_time = grad_W_time / (self.n_nodes - 1)
               self.W_time -= self.scheduler_W_time.update_change(grad_W_time)
```

```
self.b_layer -= self.scheduler_b_layer.update_change(grad_b_layer)
if grad_b_time is not None:
    grad_b_time = grad_b_time / (self.n_nodes - 1)
    self.b_time -= self.scheduler_b_time.update_change(grad_b_time)
self.update_weights_all_nodes()
```

4.2.5 The OutputLayer class

The OutputLayer class gives the output of the RNN, with a value at each time step. It is very similar to the RNNLayer class, as they essentially do the same thing, but it only connects nodes at the previous layer to the corresponding output nodes, without any connections between ndoes at different time steps. Note also that, in the same way as $feed_forward()$ in the InputLayer took a numpy array as input instead of a Layer object, the OutputLayer takes a numpy array as input to the backpropagation() method, since there is no subsequent layer to this one.

```
[14]: class OutputLayer(Layer):
           The output layer of the RNN, computing an output for each time step of the \square
        \hookrightarrowRNN. Very similar to RNNLayer, but has no connections between the nodes, \sqcup
        ⇔only from the previous.
          Attributes
           n features (int)
               Number of features in the nodes of this layer (the length of the
        \neg vectors).
           n_features_prev (int)
               Number of features in the nodes of the preceding layer (the length of \sqcup
        \hookrightarrow the vectors).
           seed (int)
               Seed for random number generating with numpy.
           nodes (list)
               List containing all the nodes of the layer in sequence, such that \Box
        \neg nodes[0] is the node at the first time step, nodes[1] at the second time_\perp
        \hookrightarrowstep, and so on.
           n nodes (int)
               Number of nodes in the layer. Is updated when adding or removing nodes.
           act func (Callable)
               The activation function of this layer.
           {b/W}_layer (ndarray)
               Bias (b) or weight matrix (W) for computing the forward propagation \Box
        ofrom nodes at the previous layer to this one (layer).
           {b/W}_layer_size (tuple)
               The shape of b_layer/W_layer.
           scheduler_{b/W}_layer (Scheduler)
```

```
The scheduler for updating b_layer/W_layer with gradient descent when
⇒backpropagating through this layer.
   is_dense (bool)
       Tells if this is a DenseLayer or not. It is set to False for this \sqcup
⇒layer, as this is not a DenseLayer.
   11 11 11
  def __init__(
           self,
           n_features: int,
           n_features_prev: int,
           act_func: Callable[[np.ndarray], np.ndarray],
           scheduler: Scheduler,
           seed: int = 100
  ):
       Constructor for OutputLayer objects.
       Parameters
       _____
       n_features (int)
           Number of features in the nodes of this layer (the length of the ...
\neg vectors).
       n features prev (int)
           Number of features in the nodes of the preceding layer (the length \sqcup
\hookrightarrow of the vectors).
       act_func (Callable)
           The activation function of this layer.
       scheduler (Scheduler)
           The scheduler to use for updating weights and biases with gradient \sqcup
→descent when backpropagating through this layer.
       seed (int)
           Seed for random number generating with numpy.
       super().__init__(n_features, seed)
       self.n_features_prev = n_features_prev
       self.act_func = act_func
       self.W_layer = None
       self.b_layer = None
       self.W_layer_size = (self.n_features_prev, self.n_features)
       self.b_layer_size = (1, self.n_features)
       self.scheduler_W_layer = copy(scheduler)
       self.scheduler_b_layer = copy(scheduler)
```

```
self.is_dense = False
      self.reset_weights()
  def reset_weights(self):
      Reset weights and biases to random values from a normal distribution.
      np.random.seed(self.seed)
      self.W_layer = np.random.normal(size=self.W_layer_size)
      self.b_layer = np.random.normal(size=self.b_layer_size) * 0.01
  def reset schedulers(self):
      11 11 11
      Reset the schedulers of the layer.
      self.scheduler_W_layer.reset()
      self.scheduler_b_layer.reset()
  def update_weights_all_nodes(self):
      Update the weights and biases in all nodes of the layer.
      new_W_layer = self.W_layer
      new_b_layer = self.b_layer
      for node in self.nodes:
           node.set_Wb(W_layer=new_W_layer, b_layer=new_b_layer)
  def add_node(self):
      n n n
      Add a node with the weights and biases specified by the layer.
      new_node = Node(self.n_features, self.act_func, self.W_layer, self.
→b_layer)
      self.nodes.append(new_node)
      self.n_nodes += 1
  def feed_forward(
           self,
          prev_layer: Layer
  ):
       11 11 11
      Feed forward through this layer one node at a time. The results are
\hookrightarrowstored in the nodes.
      Parameters
```

```
prev_layer (Layer)
           The preceding layer of the RNN.
       self.remove_nodes()
      n_nodes_prev = prev_layer.n_nodes
      for i in range(n_nodes_prev):
           # Get output of node from previous layer
           prev_layer_node = prev_layer.nodes[i]
           h_layer = prev_layer_node.h_output
           # Create and compute new node at this time step
           self.add node()
           new_node = self.nodes[i]
           # No info transfer between time steps in output layer
           new_node.feed_forward(h_layer, None)
  def backpropagate(
           self,
           dC: np.ndarray,
           lmbd: float = 0.01
  ):
       Backpropagate through the layer one node at a time. The results are
⇔stored in the nodes.
      Parameters
       _____
       dC (ndarray)
           Gradient of the cost function with respect to the output of the RNN.
→ It has shape (batch size, sequence length, number of features).
       lmbd (float)
           Regularization parameter for finding the cost gradient with respect_{\sqcup}
\hookrightarrow to the weights.
       NOTE
       Unlike the other layers, this layer takes a numpy array as input_
⇒instead of a Layer, since it is the last layer.
       .....
       ## Go through all nodes
       for i in range(self.n_nodes):
           ## Backpropagate through this node. Results are stored in the nodes
           node = self.nodes[i]
           dC_layer = dC[:,i,:] # Treat dC as coming from a subsequent layer
```

```
node.backpropagate(dC_layer, None, lmbd) # No time gradient in the_u

**output layer

## Update weights and biases (no time gradient in output layer)

grad_W_layer = node.grad_W_layer

grad_b_layer = node.grad_b_layer

self.W_layer -= self.scheduler_W_layer.update_change(grad_W_layer)

self.b_layer -= self.scheduler_b_layer.update_change(grad_b_layer)

self.update_weights_all_nodes()
```

4.2.6 The DenseLayer class

So far we have created the layers of a recurrent neural network with sequential input and sequential output, i.e., a network looking something like the following diagram.

This works well for situations where we want to predict some value at each input, such as solving differential equations. Other times, however, we want a single output for the entire input sequence, such as for sentiment analysis, where we want to determine if a sentence has a positive, negative or neutral tone. For such cases we still want to use an RNN with sequential input, but we don't want a sequential output. One way to accomplish this is to use a dense (fully connected) layer at the end of the network. We then take the last recurrent layer in the network, and add a connection from the last node to the dense layer. This would look like the following diagram.

Note here that we use a single node for the dense layer. This is because we have defined the nodes in this code to contain a single time step, but be vectors that represent the different features. Thus the connection from one node to another is really a dense connection between two "layers" (in the regular fully connected neural network sense). To convince yourself that the dense layer represented by a single node in the diagram is truly a dense *layer*, remember the following picture.

```
[15]: class DenseLayer(Layer):

"""

A dense layer used to go from a recurrent to a fully connected neural

⇒network. Contains a single node to feed forward and backpropagate through.

Attributes

-----

n_features (int)

Number of features in the node of this layer (the length of the vector).

n_features_prev (int)

Number of features in the nodes of the preceding layer (the length of

⇒the vectors).

seed (int)

Seed for random number generating with numpy.

nodes (list)
```

```
List containing all the nodes of the layer in sequence, such that \sqcup
\neg nodes[0] is the node at the first time step, nodes[1] at the second time_\(\pi\)
\hookrightarrowstep, and so on.
  n nodes (int)
       Number of nodes in the layer. Is updated when adding or removing nodes.
  act func (Callable)
       The activation function of this layer.
  {b/W}_layer (ndarray)
       Bias (b) or weight matrix (W) for computing the forward propagation \sqcup
⇔ from nodes at the previous layer to this one (layer).
   {b/W}_layer_size (tuple)
       The shape of b_layer/W_layer.
  scheduler_{b/W}_layer (Scheduler)
       The scheduler for updating b_layer/W_layer with gradient descent when_
⇒backpropagating through this layer.
   is dense (bool)
       Tells if this is a DenseLayer or not. It is set to True for this layer, □
\hookrightarrow as this is a DenseLayer.
  def __init__(
           self,
           n features: int,
           n_features_prev: int,
           act_func: Callable[[np.ndarray], np.ndarray],
           scheduler: Scheduler,
           seed: int = 100
  ):
       11 11 11
       Constructor for OutputLayer objects.
       Parameters
       _____
       n_features (int)
           Number of features in the node of this layer (the length of the \sqcup
\neg vector).
       n_features_prev (int)
           Number of features in the nodes of the preceding layer (the length \sqcup
\hookrightarrow of the vectors).
       act func (Callable)
           The activation function of this layer.
       scheduler (Scheduler)
           The scheduler to use for updating weights and biases with gradient \sqcup
⇒descent when backpropagating through this layer.
       seed (int)
           Seed for random number generating with numpy.
```

```
super().__init__(n_features, seed)
    self.n_features_prev = n_features_prev
    self.act_func = act_func
    self.nodes = []
    self.n_nodes = 0
    self.W_layer = None
    self.b_layer = None
    self.W_layer_size = (self.n_features_prev, self.n_features)
    self.b_layer_size = (1, self.n_features)
    self.scheduler_W_layer = copy(scheduler)
    self.scheduler_b_layer = copy(scheduler)
    self.is_dense = True
    self.reset_weights()
def reset_weights(self):
    Reset weights and biases to random values from a normal distribution.
    np.random.seed(self.seed)
    self.W_layer = np.random.normal(size=self.W_layer_size)
    self.b_layer = np.random.normal(size=self.b_layer_size) * 0.01
def reset_schedulers(self):
    Reset the schedulers of the layer.
    self.scheduler_W_layer.reset()
    self.scheduler_b_layer.reset()
def update_weights_all_nodes(self):
    Update the weights and biases in the node.
    self.nodes[0].set_Wb(self.W_layer, self.b_layer)
def add_node(self):
    11 11 11
    Add a node with the weights and biases specified by the layer.
```

```
new_node = Node(self.n_features, self.act_func, self.W_layer, self.
→b_layer)
      self.nodes.append(new_node)
      self.n nodes += 1
  def feed forward(
           self,
           prev_layer: Layer
  ):
      Feed forward through this layer. The result is stored in the node.
      Parameters
       _____
      prev_layer (Layer)
           The preceding layer of the RNN.
       ## Get output from last node of previous layer
      prev_node = prev_layer.nodes[-1]
      h_layer = prev_node.h_output
      self.remove_nodes()
      self.add_node()
      new_node = self.nodes[0]
      output = new_node.feed_forward(h_layer)
      return output
  def backpropagate(
           self,
           next_layer_or_dC: Layer | np.ndarray,
           lmbd: float = 0.01
  ):
       Backpropagate through the layer. The results are stored in the node.
      Parameters
       _____
      next_layer_or_dC (Layer | ndarray)
           This variable should be one of the two, depending on the type of \Box
⇒input, whether it is a Layer object or a numpy array:
           - Layer: The subsequent layer of the network. Used if this is not_{\sqcup}
\hookrightarrow the last layer in the network.
           - Numpy array: The gradient of the cost function with respect to_{\sqcup}
→ the output of the network.
```

```
It has shape (batch size, number of features). Used if this is _{\sqcup}
→ the last layer (output) of the network.
       lmbd (float)
           Regularization parameter for finding the cost gradient with respect_{\sqcup}
\hookrightarrow to the weights.
       ,, ,, ,,
       ## Get dC from next_layer_or_dC.
       if isinstance(next_layer_or_dC, Layer):
           # If next_layer_or_dC is a Layer, extract dC from last node.
           dC = next_layer_or_dC.nodes[0].grad_h_layer
       else:
           # If next layer or dC is not a layer, it is the cost gradient.
           dC = next_layer_or_dC
       ## Backpropagate through the node
      node = self.nodes[0]
      node.backpropagate(dC_layer=dC, dC_time=None, lmbd=lmbd)
       ## Update weights and biases
       grad_W_layer = node.grad_W_layer
       grad_b_layer = node.grad_b_layer
       self.W_layer -= self.scheduler_W_layer.update_change(grad_W_layer)
       self.b_layer -= self.scheduler_b_layer.update_change(grad_b_layer)
       self.update_weights_all_nodes()
```

4.2.7 The RNN class

Now that we have created all the different layer classes we need for the RNN, we can create the network itself. The RNN class is used to organize the layers in order to train and use the RNN.

Note that the class has two options for predicting results from data. The method feed_forward() propagates the input forward and returns the output in the same way as we have looked at so far. The predict() method starts by running feed_forward(), then uses argmax to set one element in the output to 1 and the rest to 0. In other words, feed_forward() is used for regression problems, while predict() is used for classification.

In addition to this, we also have a method named extrapolate(), which can be used after having run $feed_forward()$. This method takes the output at the last time step, and uses it as input for new time steps, extending the sequence. This is shown in the following diagram.

```
[16]: from sklearn.utils import resample

class RNN:

"""

The recurrent neural network. Builds the network by adding layers, trains

→it on data and predicts output of new data.
```

```
Attributes
   _____
   layers (list)
      List containing all the layers of the RNN in sequence, such that \Box
→ layers[0] is the first layer, nodes[1] is the second layer, and so on.
  n layers (int)
       Number of layers in the RNN. Is updated when adding layers.
  cost_func (Callable)
       Function which takes in the target array and returns a new function.
→ The new function takes in the output from the network,
       and returns the cost of that output when compared to the target.
  scheduler (Scheduler)
       The scheduler to use for updating weights and biases with gradient \sqcup
⇒descent when backpropagating through the network.
  seed (int)
       Seed for random number generating with numpy.
  single_output (bool)
       False if the network has a sequential output (using OutputLayer), and \Box
→True if the network has a single output (using DenseLayer).
  n_features_output (int)
       Number of features in the output layer.
  output (ndarray)
       The output from the network after computing forward propagation.
  predicted (ndarray)
       The prediction from the network on classification problems.
  output_extra (ndarray)
      Output from extrapolating the time sequence beyond the length of the \sqcup
\hookrightarrow input.
  HHHH
  def __init__(
           self,
           cost func: Callable[
               [np.ndarray], # Takes in the target output (array)
               Callable[[np.ndarray], np.ndarray] # Returns a function (the
⇔cost function)
           ],
           scheduler: Scheduler,
           seed: int = 100
  ):
       Constructor for RNN objects.
      Parameters
       _____
       cost_func (Callable)
```

```
Function which takes in the target array and returns a new function.
→ The new function takes in the output from the network,
           and returns the cost of that output when compared to the target.
       scheduler (Scheduler)
           The scheduler to use for updating weights and biases with gradient \Box
⇒descent when backpropagating through the network.
       seed (int)
           Seed for random number generating with numpy.
      self.layers = [] # List of layers
      self.n_layers = 0
      self.cost_func = cost_func
      self.scheduler = scheduler
      self.seed = seed
      self.single_output = None # Boolean. Will update this when adding_
→OutputLayer or DenseLayer output
      self.n_features_output = None
      self.output = None
      self.predicted = None
      self.output_extra = None
  def reset_weights(self):
      Reset weights and biases in all layers to random values from a normal \sqcup
\hookrightarrow distribution.
       .....
      for layer in self.layers:
           layer.reset_weights()
  def reset_schedulers(self):
      Reset the schedulers of the RNN.
      for layer in self.layers:
           layer.reset_schedulers()
  def feed_forward(
           self,
          X: np.ndarray
  ):
      Feed forward through the RNN one layer at a time.
      Parameters
```

```
X (ndarray)
           Input to the RNN, with shape (batch size, sequence length, number _{\perp}
⇔of features)
      Returns
       output (ndarray)
           Output from the RNN. If the network produces sequential output, it_{\sqcup}
\hookrightarrowhas the same shape as the input X. If the network produces a single output,\sqcup
\rightarrow it has the shape (batch size, number of features), that is, the same shape_{\Box}
→as for sequential output, but without the sequence axis.
      X_shape = X.shape
       ## Initialize output
      n_batches = X_shape[0]
      sequence_length = X_shape[1]
      n_features_output = self.n_features_output
      if self.single_output:
           output_shape = (n_batches, n_features_output)
           output = np.zeros(output_shape)
       else:
           output_shape = (n_batches, sequence_length, n_features_output)
           output = np.zeros(output_shape)
       ## Feed forward through all layers
       self.layers[0].feed_forward(X)
       for i in range(1, self.n_layers):
           layer = self.layers[i]
           prev_layer = self.layers[i-1]
           layer.feed_forward(prev_layer)
       ## Get output from last layer
       output_layer = layer
       if output_layer.is_dense:
           node = output_layer.nodes[0]
           output = node.h_output
       else:
           for i in range(output_layer.n_nodes):
               node = output_layer.nodes[i]
               output[:,i,:] = node.h_output
       ## Store and return output
       self.output = output
      return self.output
```

```
def predict(
           self,
           X: np.ndarray
  ):
       ,, ,, ,,
      Feed forward through the RNN, and use argmax to classify the results.
      Parameters
       _____
       X (ndarray)
           Input to the RNN, with shape (batch size, sequence length, number,
⇔of features)
       Returns
      predicted (ndarray)
           Predicted classification output. If the network produces sequential \sqcup
\negoutput, it has the same shape as the input X. If the network produces a_{\sqcup}
\negsingle output, it has the shape (batch size, number of features), that is,\Box
sthe same shape as for sequential output, but without the sequence axis.
       ## Initialize predicted array
      output = self.feed_forward(X)
      predicted = np.zeros_like(output)
      ## Find maximum values at the feature axis, and set predicted to 1 at_{f L}
⇔those indices
       ind_batch = np.arange(predicted.shape[0])
       if self.single_output:
           ind_max = np.argmax(output, axis=1)
           # Set predicted to 1 at the maximum values
           predicted[ind_batch, ind_max] = 1
       else:
           ind_seq = np.arange(output.shape[1])
           ind_max = np.argmax(output, axis=2)
           ind_seq, ind_batch = np.meshgrid(ind_seq, ind_batch)
           # Set predicted to 1 at the maximum values
           predicted[ind_batch, ind_seq, ind_max] = 1
       self.predicted = predicted
      return self.predicted
  def extrapolate(
           self,
```

```
length: int
  ):
       Extrapolate data by continuing the sequence from the output of the last \sqcup
_{\hookrightarrow} layer, with the output of the previous time step as input for the new time_{\sqcup}
\hookrightarrow step
       Parameters
       _____
       length (int)
           Number of time steps to extrapolate.
       Returns
       _____
       output_extra (ndarray)
           Extrapolated output from the RNN. It has the shape (number of \Box
⇒batches, extrapolation length, number of features).
       ## Initialize extrapolated output
       output = self.output
       n_batches = output.shape[0]
       n_features = self.n_features_output
       output_extra_shape = (n_batches, length, n_features)
       output_extra = np.zeros(output_extra_shape)
       ## Extrapolate
       y_prev = output[:,0,:] # Output from last layer at last time step
       for i in range(length):
           h_layer = y_prev # h_layer at first layer = input = previous output
           for l in range(1, self.n_layers):
               layer = self.layers[1]
               node_prev = layer.nodes[-1] # Node at previous time step
               h_time = node_prev.h_output
               # print(h_layer, h_time)
               layer.add_node()
               node = layer.nodes[-1] # Current node
               if l == self.n_layers-1:
                    # If this is the output layer (last layer), set h_{\perp}time to_\sqcup
\hookrightarrow None
                   h_time = None
               node.feed_forward(h_layer, h_time)
               h_layer = node.h_output # Update h_layer
           y prev = h layer
           output_extra[:,i,:] = y_prev
       self.output_extra = output_extra
```

```
return self.output_extra
  def backpropagate(
           self,
           output: np.ndarray,
           target: np.ndarray,
           lmbd: float = 0.01
  ):
       Backpropagate through the RNN one layer at a time, and update all the
\hookrightarrow weights and biases.
       Parameters
       _____
       output (ndarray)
           The output we get from forward propagation, used to compare with_
⇔the target values.
       target (ndarray)
           The target output that we want to compare our results to.
       lmbd (float)
           Regularization parameter for finding the cost gradient with respect \sqcup
\hookrightarrow to the weights.
       ## Find gradient of cost function
       grad_cost = derivate(self.cost_func(target))
       dC = grad_cost(output)
       ## Backpropagate through all layers
       self.layers[-1].backpropagate(dC, lmbd=lmbd)
       for i in range(self.n_layers-2, 0, -1):
           layer = self.layers[i]
           next_layer = self.layers[i+1]
           layer.backpropagate(next_layer, lmbd)
  def train(
           self,
           X_train: np.ndarray,
           t_train: np.ndarray,
           X_val: np.ndarray = None,
           t_val: np.ndarray = None,
           epochs: int = 100,
           batches: int = 1,
           lmbd: float = 0.01,
           store_output: np.ndarray = False
  ):
```

Trains the RNN on data, tuning the weights and biases such that the \Box ⇔network can make predictions on new unseen data. *Parameters* _____ X train (ndarray) Input data to the network during training, with shape (batch size, \Box ⇒sequence length, number of features). t_train (ndarray) Target output corresponding to the training input. Same shape as_{\sqcup} $\rightarrow X_{\perp}$ train if the output is sequential, (batch size, number of features) if not. X val (ndarray) Optional. Validation data to see how the RNN performs on unseen_ ⇒data. Not used for training, only for calculating scores. t_val (ndarray) Optional. Target output corresponding to the validation data. epochs (int) Number of epochs to train for. batches (int) Number of batches to split data into at each epoch. Regularization parameter for finding the cost gradient with respect $_{11}$ \hookrightarrow to the weights. store output (bool) Whether to store output from each epoch (if you want to see how the ... \rightarrow output evolves as the RNN trains). Returnsscores (dict) The different scores from the training stored as a dictionary, with \sqcup *→* the following keys: - "train_error" : The error of the training data for each epoch. - "train_accuracy" : The accuracy of the training data for each_ ⇔epoch. - "val_error" : The error of the validation data for each epoch. - "val_accuracy" : The accuracy of the validation data for each_ \hookrightarrow epoch. - "y_train_history" : The output from the training data for each_ *⇔epoch*. - "y_val_history" : The output from the validation data for each \Box ⇔epoch. 11 11 11 self.reset_weights() # Reset weights for new training batch_size = X_train.shape[0] // batches

```
# Initialize arrays for scores
      train_cost = self.cost_func(t_train)
      train_error = np.zeros(epochs)
      train_accuracy = np.zeros(epochs)
      if X_val is not None:
           val_cost = self.cost_func(t_val)
          val_error = np.zeros(epochs)
           val_accuracy = np.zeros(epochs)
       ## Initialize arrays for output history if this should be stored
      if store_output:
          n_batches_train = X_train.shape[0]
           seq_length_train = X_train.shape[1]
          n_features_output = self.n_features_output
           if self.single_output:
               y_train_shape = (epochs, n_batches_train, n_features_output)
           else:
               y_train_shape = (epochs, n_batches_train, seq_length_train, __
→n_features_output)
          y_train_history = np.zeros(shape=y_train_shape)
           if X_val is not None:
              n_batches_val = X_val.shape[0]
               seq_length_val = X_val.shape[1]
               if self.single_output:
                   y_val_shape = (epochs, n_batches_val, n_features_output)
               else:
                   y_val_shape = (epochs, n_batches_val, seq_length_val,__
→n_features_output)
               y_val_history = np.zeros(shape=y_val_shape)
       \# Resample X and t
      X_train, t_train = resample(X_train, t_train, replace=False)
      for e in range(epochs):
           print("EPOCH: " + str(e+1) + "/" + str(epochs), end="\rder")
           for b in range(batches):
               ## Extract a smaller batch from the training data
               if b == batches - 1:
                   # If this is the last batch, include all remaining elements
                   X_batch = X_train[b*batch_size :]
                   t_batch = t_train[b*batch_size :]
```

```
else:
                X_batch = X_train[b*batch_size : (b+1)*batch_size]
                t_batch = t_train[b*batch_size : (b+1)*batch_size]
            ## Train the network on this batch with gradient descent
            y_batch = self.feed_forward(X_batch)
            self.backpropagate(y_batch, t_batch, lmbd)
        self.reset schedulers()
        ## Compute scores for this epoch
        y_train = self.feed_forward(X_train)
        pred_train = self.predict(X_train)
        train_error[e] = train_cost(y_train)
        train_acc_arr = np.all(pred_train == t_train, axis=-1)
        train_accuracy[e] = np.mean(train_acc_arr)
        if X_val is not None:
            y_val = self.feed_forward(X_val)
            pred_val = self.predict(X_val)
            val_error[e] = val_cost(y_val)
            val_acc_arr = np.all(pred_val == t_val, axis=-1)
            val_accuracy[e] = np.mean(val_acc_arr)
        if store_output:
            y_train_history[e] = y_train
            if X_val is not None:
                y_val_history[e] = y_val
    ## Create a dictionary for the scores, and return it
    scores = {"train_error": train_error, "train_accuracy": train_accuracy}
    if X_val is not None:
        scores["val_error"] = val_error
        scores["val_accuracy"] = val_accuracy
    if store_output:
        scores["y_train_history"] = y_train_history
        if X_val is not None:
            scores["y_val_history"] = y_val_history
    return scores
def add_InputLayer(
```

```
self,
          n_features: int
  ):
      Adds an InputLayer to the RNN.
      Parameters
       _____
      n features (int)
          Number of features of the input.
      layer = InputLayer(n_features, self.seed)
      self._add_layer(layer)
  def add_RNNLayer(
          self,
          n_features: int,
          act_func: Callable[[np.ndarray], np.ndarray]
  ):
      Adds an RNNLayer to the RNN.
      Parameters
      n_features (int)
          Number of features in this layer.
      act_func (Callable)
          The activation function to use for this layer.
      scheduler = copy(self.scheduler)
      prev_layer = self.layers[-1]
      n_features_prev = prev_layer.n_features
      layer = RNNLayer(n_features, n_features_prev, act_func, scheduler, self.
⇒seed)
      self._add_layer(layer)
  def add_OutputLayer(
          self,
          n_features: int,
          act_func: Callable[[np.ndarray], np.ndarray]
  ):
      11 11 11
      Adds an OutputLayer to the RNN.
      Parameters
      _____
      n_features (int)
```

```
Number of features in this layer.
       act_func (Callable)
           The activation function to use for this layer.
      scheduler = copy(self.scheduler)
      prev_layer = self.layers[-1]
      n_features_prev = prev_layer.n_features
      layer = OutputLayer(n_features, n_features_prev, act_func, scheduler,_
⇒self.seed)
      self._add_layer(layer)
      self.single_output = False
      self.n_features_output = n_features
  def add_DenseLayer(
           self,
           n_features: int,
           act_func: Callable[[np.ndarray], np.ndarray],
           is_last_layer: bool = False
  ):
       n n n
      Adds a DenseLayer to the RNN.
      Parameters
       _____
       n_features (int)
           Number of features in this layer.
       act_func (Callable)
           The activation function to use for this layer.
       is_last_layer (bool)
           True if this is the last layer of the network (the output layer). ⊔
\hookrightarrow False if not.
      11 11 11
      scheduler = copy(self.scheduler)
      prev_layer = self.layers[-1]
      n_features_prev = prev_layer.n_features
      layer = DenseLayer(n_features, n_features_prev, act_func, scheduler, u
⇒self.seed)
      self._add_layer(layer)
      if is_last_layer:
           self.single_output = True
           self.n_features_output = n_features
  def _add_layer(
           self,
           layer: Layer
```

```
):

"""

Adds a layer to the RNN by appending it to *layers*, and increases

→*n_layers* by one.

Parameters

-----
layer (Layer)

The layer to add to the RNN.

"""

self.layers.append(layer)

self.n_layers += 1
```

4.3 Examples on applications

4.3.1 Predicting the weather

We will now explore a dataset that considers how temperature evolves over time. The dataset is obtained from the "Zero to GPT" course by Vik Paruchuri at Dataquest, and can be found at https://github.com/VikParuchuri/zero_to_gpt/blob/81cf89f68143535b7361990aa0fc49d253399c65/data/clean_w

Let's first look at the dataset provided.

```
Unnamed: 0 tmax
                   tmin rain
                                tmax_tomorrow
 1970-01-01 60.0
                    35.0
                          0.00
                                        52.0
  1970-01-02 52.0
                    39.0
                                        52.0
1
                          0.00
2 1970-01-03 52.0
                    35.0
                          0.00
                                        53.0
3
  1970-01-04 53.0
                    36.0
                          0.00
                                        52.0
  1970-01-05 52.0
                    35.0
                          0.00
                                        50.0
 1970-01-06 50.0
                    38.0
                                        52.0
                          0.00
6
 1970-01-07 52.0
                    43.0
                          0.00
                                        56.0
7
  1970-01-08 56.0 49.0
                                        54.0
                          0.24
8
  1970-01-09
              54.0 50.0
                          0.40
                                        57.0
  1970-01-10 57.0 50.0 0.00
                                        57.0
```

The dataset has 13509 rows.

The relevant columns for our purposes are tmax and $tmax_tomorrow$, so we will only retrieve these. Note that the dataset has 13 509 rows (time steps), which is too much to work with for our simple RNN, and will quickly result in vanishing or exploding gradients. We will therefore only retrieve the first 100 rows. Notice also that the i'th element of $tmax_tomorrow$ is the i+1'th element of tmax, as we want to use the RNN to predict the temperature the next day based on the temperatures on preceding days. Thus tmax will be the input to the network, and $tmax_tomorrow$ the target

output.

```
[18]: N = 100
    x_orig = weather_data["tmax"][:N].to_numpy()
    t_orig = weather_data["tmax_tomorrow"][:N].to_numpy()
```

For the validation data, we will retrieve a subset of the data at some later time, such that the training and validation data do not overlap. Notice also that we let the validation data contain 70 time steps, even though the training data contains 100 time steps. This demonstrates one of the advantages of RNNs, their flexibility with input lengths.

```
[19]: val_start = 1000
N_val = 70
x_val_orig = weather_data["tmax"][val_start:val_start+N_val].to_numpy()
t_val_orig = weather_data["tmax_tomorrow"][val_start:val_start+N_val].to_numpy()
```

We need the input and output of the RNN to have shape (batch size, sequence length, number of features), but the data we have read now are one-dimensional since they only have one feature (the scalar temperature) and one input (so batch size is 1). We therefore have to add axes such that the data is on the correct form. We will also scale the data to make it easier for the RNN to train. The scaling is done using scikit's MinMaxScaler.

```
[20]: from sklearn.preprocessing import MinMaxScaler
      # Add feature axis
      x = x orig[:, np.newaxis]
      t = t_orig[:, np.newaxis]
      x_val = x_val_orig[:,np.newaxis]
      t_val = t_val_orig[:,np.newaxis]
      # Scale data
      sc = MinMaxScaler()
      sc = sc.fit(x)
      x = sc.transform(x)
      t = sc.transform(t)
      x_val = sc.transform(x_val)
      t_val = sc.transform(t_val)
      # Add batch axis
      x = x[np.newaxis,:]
      t = t[np.newaxis,:]
      x_val = x_val[np.newaxis,:]
      t_val = t_val[np.newaxis,:]
```

We can now create the network, by declaring an RNN object and adding layers to it. We will use one hidden recurrent layer.

```
[21]: cost_func = CostOLS
    act_func_hidden = identity
    act_func_output = identity
    eta = 0.001
    scheduler = Adam(eta=eta, rho=0.9, rho2=0.999)

## Create RNN
    rnn = RNN(cost_func, scheduler)
    rnn.add_InputLayer(1)
    rnn.add_RNNLayer(1, act_func_hidden)
    rnn.add_OutputLayer(1, act_func_output)
    rnn.reset_weights()
```

With the RNN in place we can train it on the weather dataset.

```
[22]: ## Train network on data

epochs = 50

batches = 1

lmbd = 0.001

scores = rnn.train(x, t, x_val, t_val, epochs=epochs, batches=batches, u

→lmbd=lmbd, store_output=True)
```

EPOCH: 50/50

Now extract the results from the training. We scaled the input before training, so in order to get the actual temperature predicted we need to scale the ouput back using the scalers *inverse transform()*.

```
[23]: ## Extract output
    epoch_arr = np.arange(1, epochs+1)
    y_history_scaled = scores["y_train_history"]
    train_error = scores["train_error"]
    y_val_history_scaled = scores["y_val_history"]
    val_error = scores["val_error"]

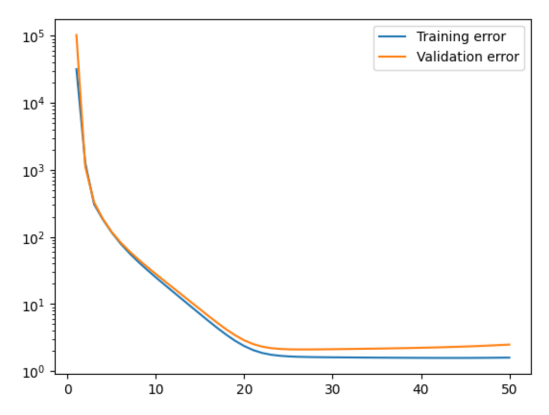
## Inversely scale output
    y_history = np.zeros(y_history_scaled.shape)
    for i in range(y_history_scaled.shape[0]):
        yi = y_history_scaled[i,0,:,:]
        yi_scaleback = sc.inverse_transform(yi)
        y_history[i,0,:,:] = yi_scaleback
```

We can now plot the error over time to see how the model performs during training.

```
[24]: # %matplotlib notebook
import matplotlib.pyplot as plt

## Plot error during training
plt.figure()
plt.plot(epoch_arr, train_error, label="Training error")
```

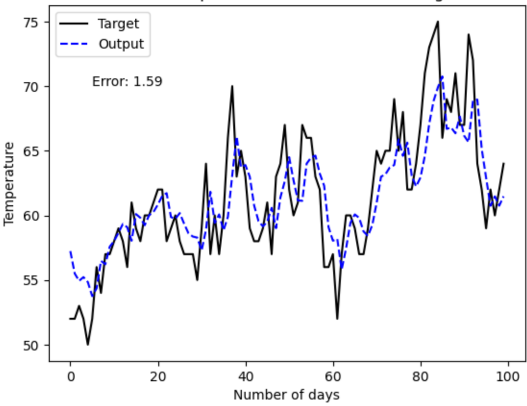
```
plt.plot(epoch_arr, val_error, label="Validation error")
plt.yscale("log")
plt.legend()
plt.show()
```



The error decreases and stabilizes, as it should. Let us also look at how the output of the training data compares to the target after training.

```
[25]: ## Plot output and target after training
seq_ind = np.arange(N)
y = y_history[-1,0,:,0]
plt.figure()
plt.title("Maximum temperature over time for training data.")
plt.xlabel("Number of days")
plt.ylabel("Temperature")
plt.plot(seq_ind, t_orig, "k", label="Target")
plt.plot(seq_ind, y, "b--", label="Output")
plt.text(5, 70, f"Error: {train_error[-1]:.2f}")
plt.legend()
plt.show()
```

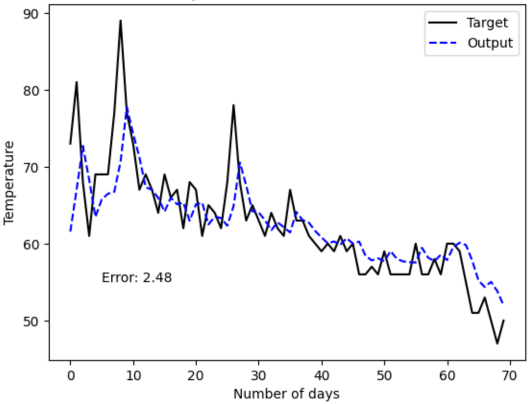
Maximum temperature over time for training data.



We do the same for the validation data as well.

```
[26]: ## Plot validation results
seq_val = np.arange(N_val)
y_val = y_val_history_scaled[-1,0,:,:]
y_val = sc.inverse_transform(y_val)
y_val = y_val[:,0]
plt.figure()
plt.title("Maximum temperature over time for validation data.")
plt.xlabel("Number of days")
plt.ylabel("Temperature")
plt.plot(seq_val, t_val_orig, "k", label="Target")
plt.plot(seq_val, y_val, "b--", label="Output")
plt.text(5, 55, f"Error: {val_error[-1]:.2f}")
plt.legend()
plt.show()
```



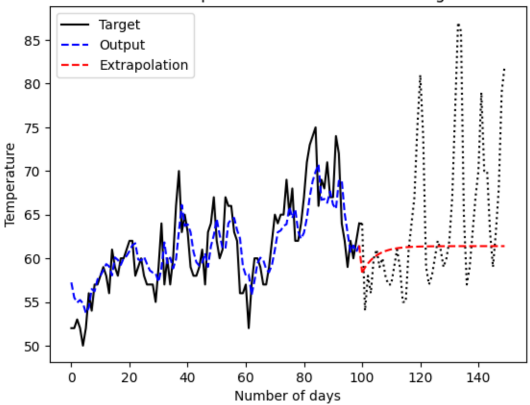


Both the training and validation output looks good, fitting the target relatively well considering that we are using a simple RNN. Let's see if the model can also extrapolate the output to get the temperature for the next 50 days.

```
[27]: ## Extrapolate data using network
      length = 50
      y_extra = rnn.extrapolate(length)
      y_extra = y_extra[0,:,:]
      y_extra = sc.inverse_transform(y_extra)
      y_extra = y_extra[:,0]
      seq_extra = np.arange(N, N+length)
      t_extra = weather_data["tmax_tomorrow"] [N:N+length].to_numpy()
      plt.figure()
      plt.title("Maximum temperature over time for training data.")
      plt.xlabel("Number of days")
      plt.ylabel("Temperature")
      plt.plot(seq_ind, t_orig, "k", label="Target")
      plt.plot(seq_extra, t_extra, "k:")
      plt.plot([seq_ind[-1],seq_extra[0]], [t_orig[-1],t_extra[0]], "k--")
      plt.plot(seq_ind, y, "b--", label="Output")
```

```
plt.plot([seq_ind[-1],seq_extra[0]], [y[-1],y_extra[0]], "r--")
plt.plot(seq_extra, y_extra, "r--", label="Extrapolation")
plt.legend()
plt.show()
```

Maximum temperature over time for training data.



The extrapolation does not look very good. It seems to stay constant, independent of the fluctuations in the true data.

We have stored the output from the network at each epoch. We can use this to create an animation of how the model fits to the data during training.

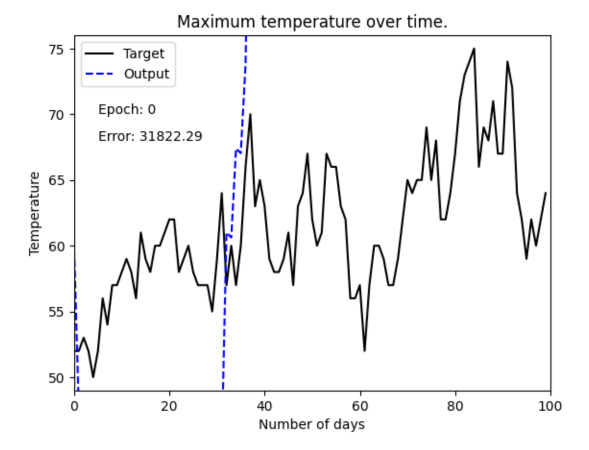
```
[38]: import matplotlib.animation as animation
  from IPython.display import HTML
  ## Create animation of how the output fits to the target
  y = y_history[0,0,:,0]
  fig, ax = plt.subplots()
  ax.set_title("Maximum temperature over time.")
  ax.set_xlabel("Number of days")
  ax.set_ylabel("Temperature")
  t_plot = ax.plot(seq_ind, t_orig, "k", label="Target")[0]
```

```
y_plot = ax.plot(seq_ind, y, "b--", label="Output")[0]
epoch_text = ax.text(5, 70, f"Epoch: {0}")
error_text = ax.text(5, 68, f"Error: {train_error[0]:.2f}")
ax.set(xlim=[0,N], ylim=[np.min(t_orig)-1,np.max(t_orig)+1])
ax.legend()

def update_plot(frame):
    y = y_history[frame,0,:,0]
    y_plot.set_ydata(y)
    epoch_text.set_text(f"Epoch: {frame}")
    error_text.set_text(f"Error: {train_error[frame]:.2f}")
    return y_plot

anim = animation.FuncAnimation(fig=fig, func=update_plot, frames=epochs,u_ointerval=100)
HTML(anim.to_jshtml())
```

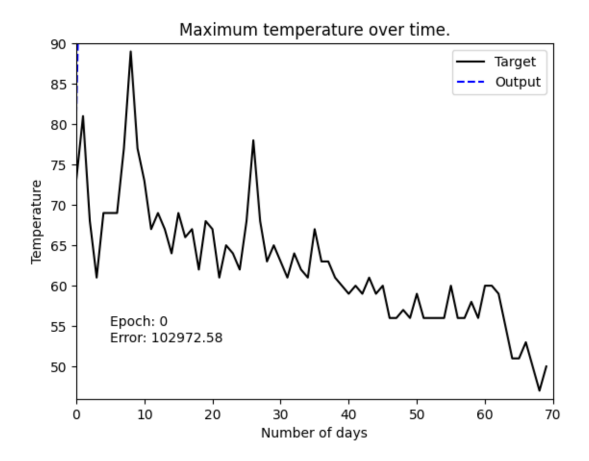
[38]: <IPython.core.display.HTML object>



And we can do the same for the validation data.

```
[39]: ## Create animation for validation results
      ## Inversely scale output
      y_val_history = np.zeros(y_val_history_scaled.shape)
      for i in range(y_val_history_scaled.shape[0]):
          yi = y_val_history_scaled[i,0,:,:]
          yi_scaleback = sc.inverse_transform(yi)
          y_val_history[i,0,:,:] = yi_scaleback
      y_val = y_val_history[0,0,:,0]
      fig, ax = plt.subplots()
      ax.set title("Maximum temperature over time.")
      ax.set_xlabel("Number of days")
      ax.set_ylabel("Temperature")
      t_plot = ax.plot(seq_val, t_val_orig, "k", label="Target")[0]
      y_plot = ax.plot(seq_val, y_val, "b--", label="Output")[0]
      epoch_text = ax.text(5, 55, f"Epoch: {0}")
      error_text = ax.text(5, 53, f"Error: {val_error[0]:.2f}")
      ax.set(xlim=[0,N_val], ylim=[np.min(t_val_orig)-1,np.max(t_val_orig)+1])
      ax.legend()
      def update_plot(frame):
          y_val = y_val_history[frame,0,:,0]
          y_plot.set_ydata(y_val)
          epoch_text.set_text(f"Epoch: {frame}")
          error_text.set_text(f"Error: {val_error[frame]:.2f}")
          return y plot
      anim = animation.FuncAnimation(fig=fig, func=update_plot, frames=epochs, __
       ⇒interval=100)
      HTML(anim.to_jshtml())
```

[39]: <IPython.core.display.HTML object>



[]: