

Deep Stock Prediction

Soham Deshpande

January 2022

A Technical Indicator for liquid asset evaluation using machine learning techniques

Contents

1	Introduction to the Stock Market	4
1.1	Abstract	4
2	Introduction	4
3	Analysis	4
3.1	Stock prediction techniques	4
3.2	ARIMA	5
3.3	Neural Networks	6
3.4	CNN	6
3.5	Activation Functions	7
3.6	Layers	11
3.7	Neurons and Equations	11
3.8	Loss Functions	12
3.9	Feed Forward and Back propagation	12
3.10	Backpropagation	14
3.11	RNN	15
3.11.1	Problems with RNN	16
3.12	LSTM	16
3.12.1	Problems with LSTM	17
3.13	Transformers	17
3.14	Attention	18
3.15	Temporal Fusion Transformers	20
3.16	Tensors	20
3.16.1	Tensor Product	21
3.17	Gated Linear Units(GLU)	21
3.18	Gated Residual Network(GRN)	21
3.19	Variable Selection Network	22
3.20	Multi-head attention	22
3.21	Quantile Forecasts	23
3.22	Software	23
3.22.1	Coding Language	23
3.22.2	Packages	25
3.23	Objectives	25
3.23.1	General Objectives	25
3.23.2	Specific objectives	26
4	Documented Design	28
4.0.1	Heirarchy Diagram	28
4.1	Explaining nn.module	42
4.2	Positional Encoding	45
4.3	Backpropagation	47
4.4	Dense Network	49
4.5	Weight Initialisation	50

4.5.1	Theory	50
4.6	Layer Normalisation	51
4.6.1	The need for normalisation	51
4.6.2	Layer Normalisation	51
4.6.3	Batch normalisation vs layer normalisation	52
4.6.4	Code	52
4.7	Time Distributed	53
4.8	Activation Functions	53
4.8.1	Sigmoid	53
4.8.2	Softmax	54
4.8.3	Tanh	54
4.8.4	ELU	55
4.9	Attention	55
4.9.1	Multi-head attention	55
4.10	Data Structures	57
4.11	58
4.12	File Structure	58
5	Bibliography	59
	References	60

1 Introduction to the Stock Market

1.1 Abstract

In recent years the fast-growing financial markets opened new horizons for investors and at the same time brought new challenges for financial analysts in their efforts to make effective decisions and reduce investment risks. The stock market is a highly dynamic and complex system where factors affecting the price are not limited to the economic world, rather in recent years the political climate and social media playing a bigger role. This has resulted in a highly stochastic, chaotic market. As with the other industries, the amount of data being created every day can be overwhelming and often hard to understand and decipher for someone with limited experience. This problem can be seen as one for computer science and mathematics. In the past decades, effective prediction models, both linear and machine learning tools have been explored. In recent years the research into deep learning has rapidly accelerated the progress made in prediction software. The motivation behind this paper is to describe and show an implementation of a deep learning model to help predict the price of stocks.

Keywords: Stock market prediction, deep learning, transformers, attention, feedforward neural network, temporal fusion transformers

2 Introduction

The model I will be exploring is a transformer-based deep learning architecture that takes advantage of attention, more specifically multi-head attention in my implementation. Many models use ARIMA (Auto-Regressive Integrated Moving Average) but I am proposing using transformers with multi-head attention, something that I will talk about later on, to help the model ‘learn’ about the stock rather than just trying to fit a curve on it based on the last few data points. The benefits of learning allow the model to consider previous experience instead of just looking at a few, previous data points. This technique will hopefully result in a higher accuracy compared to ARIMA. Other implementations have managed an accuracy of 94% (Zolkepli and Divino, n.d.) so I will be aiming to stay within 5%. The reduced target comes down to a few factors such as limited access hardware, not enough time to optimise my program for the hardware as well as a few other factors.

3 Analysis

3.1 Stock prediction techniques

Stock prediction is a problem that requires time series analysis. Time series analysis is the technique of analysing sequential data. In the current market, a

couple of techniques are used to analyse and predict the price. In this section, I will discuss a few multivariate time series models.

3.2 ARIMA

Autoregressive integrated moving average is a common method used to perform time series analysis. (Hyndman and Athanasopoulos, 2018) In a regression model, we forecast the variable of interest using a linear combination of predictors. Autoregression focuses on using a linear combination of past values and the variable itself to predict the variable of interest. The term autoregression indicates that it is a regression of the variable against itself.

$$y_t = c + \phi_1 y_{t-1} + \phi_2 y_{t-2} + \dots + \phi_p y_{t-p} + \epsilon_t \quad (1)$$

These models are very good at handling different time series patterns. Changing the parameters, ϕ result in different patterns. ϵ is the error term; changing this will change the scale of the series. This model is very effective when it comes to stock prediction and so will therefore provide a baseline when it comes to the accuracy of the model. The image above shows an implementation of

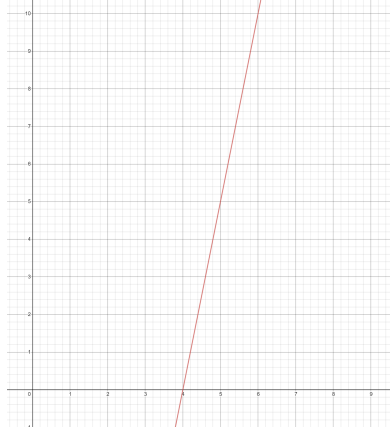


Figure 1: Plotting the data

auto regression on FTSE 100 data from 2010 to 2020. Green dots represent the predicted values while the red dots represent the actual values.

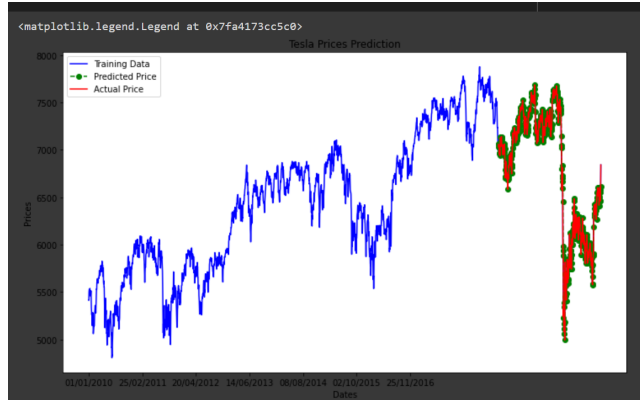


Figure 2: Using ARIMA on test data

Figure 3 shows the accuracy of the model at a respectable 89.21% when using MSE as the loss function.

```
<class 'list'>
Testing Mean Squared Error: 7223.205
Symmetric mean absolute percentage error: 10.792
```

Figure 3: MSE Loss using ARIMA

3.3 Neural Networks

The research done in the 1990s have helped set up the foundations for a mathematical model that is today known as machine learning. This mathematical process aims to replicate the brain through the use of several functions and calculus. This replication comes in the form of neurons in a neural network. In the following paragraphs I will proceed to explain a type of neural network known as a ‘Convolutional neural network’ before proceeding to describe an evolution, the ‘Recurrent neural network’.

3.4 CNN

The neural network is heavily dependent on the maths surrounding gradients, in particular differentiation. The main components of a neural network include: an activation function, layers, a loss function and an optimiser.

Similar to the human brain, a neural network is made up of many neurons. These neurons are information processing units that are fundamental to the operation of a neural network. The structure of a neuron is shown in figure 2.2.3 and is represented by a subscript k . A basic neural network consists of 3 main parts: a set of connecting links between neurons, an adder and an activation function.

The set of links each have a weight assigned to them. They work by taking input signals, often as vectors, and then multiply these by the weights they are assigned. This process is shown in figure 2.2.3. After that each vector is added together before a fixed bias is applied. To simulate the process of a neuron turning on and off, an activation function is used. These are mathematical functions and are described in the following paragraphs.

3.5 Activation Functions

To summarise the behaviour of an activation function, it can be compared to a transistor; it only activates at a certain threshold. These try to replicate how the neurons behave in our brain. The most common activation functions are shown below along with their equation.

- Binary step function

—

$$u(x) = 1 \text{ if } x \geq 0 \text{ or } 0 \text{ if } x < 0$$

$$Y = u(x - 3)$$

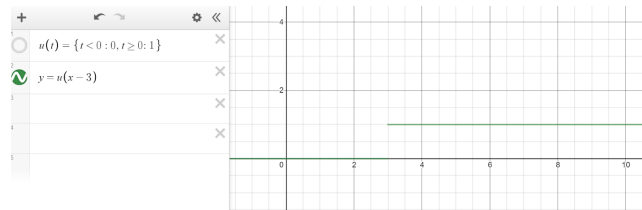


Figure 4: Binary step function

- Linear Function

—

$$f(x) = mx$$

- Sigmoid

—

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

$$\frac{d}{dx} =$$

$$f = 1 \quad f' = 0$$

$$g = 1 + e^{-x} \quad g' = -e^{-x}$$

Using the quotient rule:

$$= \frac{e^{-x}}{(1 + e^{-x})^2}$$

This can also be written as:

$$\sigma'(x) = \sigma(x)(1 - \sigma(x))$$

- Tanh

- $\tanh(x) = 2\sigma(2x) - 1$
- Values range from -1 to 1

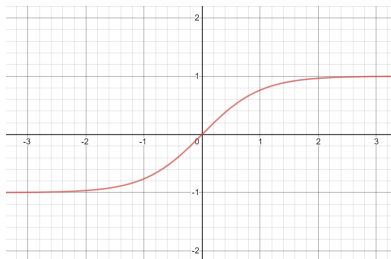


Figure 5: Tanh function

- ReLU(Rectified Linear Unit)

- Neurons will only deactivate if the output of the linear transformations less than 0
- Also a leaky version of ReLU present

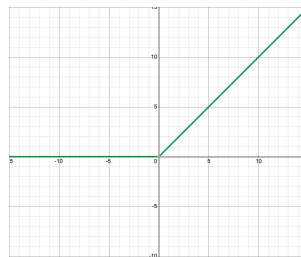


Figure 6: ReLU function

- Softmax

- Returns the probability for a data point belonging to each individual class

$$\sigma(\hat{z})_i = \frac{e^{z_i}}{\sum_{j=1}^k e^{z_j}}$$

Below is a description of the softmax function: (Versloot, 2020)

This can be described as the following: for each value in our input vector, the Softmax value is the exponent of the individual input divided by a sum of the exponents of all the inputs.

This ensures that multiple things happen:

Negative inputs will be converted into nonnegative values, thanks to the exponential function.

Each input will be in the interval (0,1) As the denominator in each Softmax computation is the same, the values become proportional to each other, which makes sure that together they sum to 1.

These properties allow us to interpret them as probabilities.

To make sure these values are actually valid probabilities, it can be checked against Kolmogorov's probability axioms.

- * Each probability must be a nonzero real number. This is true for our outcomes: each is real-valued, and nonzero.
- * The sum of probabilities must be 1. This is also true for our outcomes: the sum of cutoff values is ≈ 1 , due to the nature of real-valued numbers. The true sum is 1.

Here is an example of the function

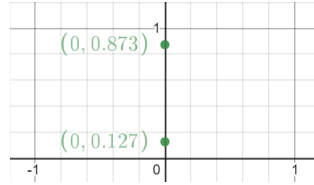


Figure 7: Where $a = 6.638$ and $b = 4.71$

In the equation above, \hat{z} is the input vector to the softmax function, made up of (z_0, \dots, z_k)

z_i are All the Z_i values are the elements of the input vector to the softmax function, and they can take any real value, positive, zero or negative. For example a neural network could have output a vector such as $(-0.62, 8.12, 2.53)$, which is not a valid probability distribution, hence why the softmax would be necessary.

e^{z_i} is the standard exponential function is applied to each element of the input vector. This gives a positive value above 0, which will be very

small if the input was negative, and very large if the input was large. However, it is still not fixed in the range $(0, 1)$ which is what is required of a probability.

$\sum_{j=1}^k e^{z_j}$ is the term on the bottom of the formula is the normalization term. It ensures that all the output values of the function will sum to 1 and each be in the range $(0, 1)$, thus constituting a valid probability distribution.

k is the number of classes in the multi-class classifier.

- ELU(Exponential linear unit)

The exponential linear unit was designed to fix some of the problems with ReLUs. This function has a parameter that is picked; a common value is between 0.1 and 0.3.

$$ELU(x) = x \text{ if } x > 0 \text{ or } \alpha(e^x - 1) \text{ if } x < 0$$

This equation tells us that if the input, x , is greater than 0 then the output is the same as a ReLU, $y=x$. If the input drops below 0, the value becomes slightly smaller than 0. This value will be modelled using (e^x-1) where we choose a value for α . The exponential operation makes this function more computationally expensive than a vanilla ReLU. The exponential function helps fix the vanishing gradient problem often associated with ReLUs. An advantage is that the ELU produces negative values which allows them to push mean unit activation close to zero, a bit like batch normalisation but with lower computational complexity. Mean shifts towards zero help speed up learning. The derivative can be written

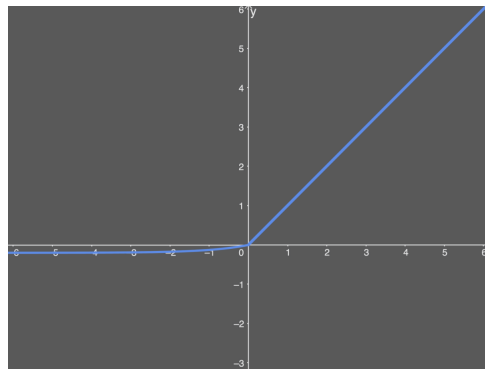


Figure 8: ELU function

as

$$ELU'(x) = 1 \text{ if } x > 0 \text{ or } \alpha(e^x) \text{ if } x < 0$$

The exponential function helps make this function differentiable at all points. Below is the graph for the derivative:

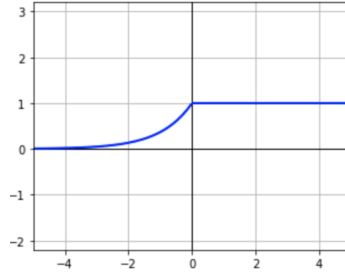


Figure 9: ELU derivative function

3.6 Layers

As shown in the figure below, in between the input and output layers there are hidden layers. These layers contain many neurons. The equation that is used to model these neurons is shown in the second image, I have used the sigmoid function as an example of the calculation that may happen at a neuron. The summation of the weights multiplied by the input activity. A bias is then applied before the activation function, a sigmoid in my example, is applied to the result to determine whether the neuron should turn on or not.

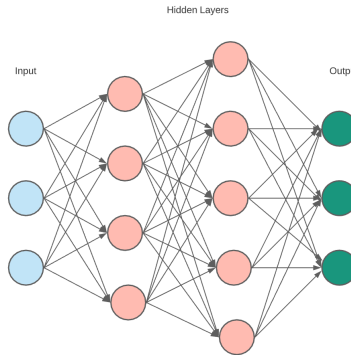


Figure 10: Layers shown in a neural network

3.7 Neurons and Equations

The neurons can be modelled using the equations given below in the equation

$$a = \sigma(w \times a + b)$$

Here a is the activity, the activity of the neuron, σ is the activation function, w is the weight and b is the bias. The equation shows that the activity is modelled by multiplying the previous activity by a weight that is changed during the whole learning process, before a constant bias is applied. The equation below

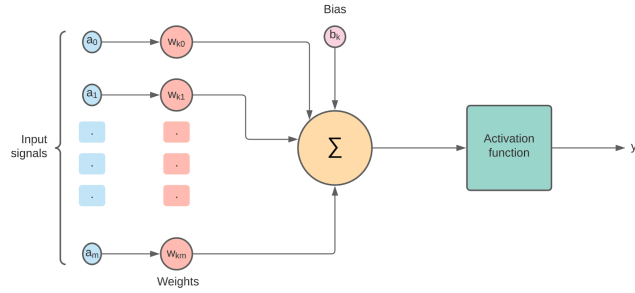


Figure 11: Neuron represented mathematically

shows a more general equation where the letter L represents a neuron in current time and so L-1 is the activity of the previous neuron.

$$a^L = \sigma(w^L \times a^{L-1} + b^L)$$

Another way to write this is:

$$\sigma\left(\sum_{j=0}^n w_j \cdot a_j + b\right)$$

3.8 Loss Functions

A loss function is what ultimately gives the neural network its intelligence. At its core it's a method evaluating how well the algorithm models the data. If the predictions are off by a high margin then a high loss will be calculated. If the margin of error is lower, a lower number will be calculated showing that your predictions are quite close. This loss is also commonly referred to as the accuracy. The goal is to get your loss as low as possible or can be referred to as achieving a high accuracy. There are many functions that can be used to determine the loss but the most common are root mean squared error, absolute error loss and binary cross entropy loss. Lots of neural networks have custom loss functions that are dependent on the purpose they are serving. A very strict loss function that amplifies the loss is more typical in places where the loss must be kept at a minimum such as stock prediction.

3.9 Feed Forward and Back propagation

For the neural network to truly learn, it must go through a process called back propagation. This is an algorithm that is used in supervised learning and uses a concept called gradient descent. One way to think about a loss function is to imagine it being a black box that works out an equation, say $y =$ to get you from the domain to the range. Now this equation can be plotted, figure 2 and we can see that there are a few things to observe about this graph. Calculus

tells us that the turning points happen when the gradient of a curve, that can be worked out using differentiation, is 0. The second step is to work out the second derivative and that will tell us whether the point is a maxima or minima. Where $\frac{d^2y}{dx^2} > 0$, the point can be described as a minimum and when $\frac{d^2y}{dx^2} < 0$, the point can be described as a maximum. The minimum point that occurs when $\frac{dy}{dx}$ (the gradient) is 0, on the graph below it happens at the point (3,0).

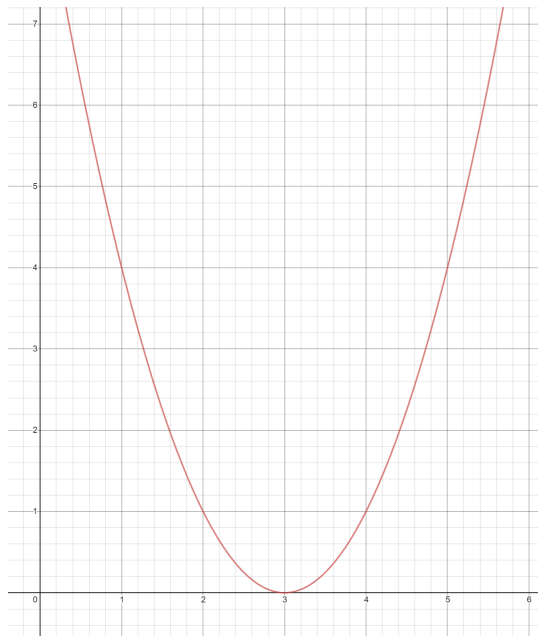


Figure 12:

Consider w to be one of the weights discussed in the previous section. If $w < 3$, we have a positive loss function, but the derivative is negative, meaning that an increase of weight will decrease the loss function. At $w = 3$, the loss is 0 and the derivative is 0, we reached a perfect model, nothing is needed. If $w > 3$, the loss becomes positive again, but the derivative is as well positive, meaning that any more increase in the weight, will increase the losses even more. To try and decrease this loss, at $w < 3$ you should increase the weight and at $w > 3$ you should decrease the weight.

The equation for working out the new weight is shown below where w_n is the weight, C is the cost function and α is the learning rate(which is a constant).

$$new\ w_n = (old\ w_n) - \alpha \cdot \frac{\partial C}{\partial w_n}$$

3.10 Backpropagation

The goal of backpropagation is to minimise the error function by changing the weights of the individual neurons. The formulation of the complete backpropagation algorithm can be proven by induction to show that it works with differentiable activation functions at its nodes.

For this proof, the assumption has been made that the neural network is one with a single input and single output. To simplify the proof, instead of carrying a bias term, let us assume that each layer $V^{(t)}$ contains a single neuron $v_o^{(t)}$ that always outputs a constant 1 thus the output of a neuron is given by

$$\sigma(\sum w_{i,j} v_j^{t-1})$$

We next wish to compute the derivative ∇f_w . Now suppose neuron v_i^t computes:

$$v_i^t(x) = \sigma(u_i^t(x))$$

Then using the chain rule we can obtain that

$$\frac{\partial f}{\partial w_{i,j}} = \frac{\partial f}{\partial u_i^t} \cdot \frac{\partial u_i^t}{\partial f} = \frac{\partial f}{\partial u_i^t} v_i^{(t-1)}(x)$$

Thus to compute the partial derivative of a single weight, it is enough to compute $\frac{\partial f}{\partial u_i^t}$. We can now focus on computing $\frac{\partial f}{\partial u_i^t}$. Now suppose f is a function of u_1^t, \dots, u_m^t , which we are in turn function of some variable z then we have by chain rule:

$$\frac{\partial f}{\partial z} = \sum_{i=1}^m \frac{\partial f}{\partial u_i^t} \cdot \frac{\partial u_i^t}{\partial z}$$

Now if $z = u_n^{t-1}$ is the output of some neuron in a previous layer: The calculation of $\frac{\partial u_i^t}{\partial u_n^{t-1}}$ is easy for our choice of activation function

$$u_i^t = \sum w_{i,j} \sigma(u_j^{t-1}) \rightarrow \frac{\partial u_i^t}{\partial u_n^{t-1}} = w_{i,n} \sigma'(u_n^{t-1})$$

Using this equation for $f = u_i^t$ we can recursively calculate all the partial derivatives. This inductive approach can be used to calculate all the derivatives, giving us the backpropagation algorithm. In reality the algorithm calculates the derivatives through dynamic programming therefore reducing the complexity.

To summarise the process consider an input-output pair, (x, y) . Use the current set of weights at $t=0$ in the network to generate the output vector $g(x)$, whose individual values comprise of the a_N values for each neuron in the output layer. Next compute the derivatives for each neuron in the output layer and use those values to update the weights, w , on the incoming edges on the graph for those neurons using the update rule. Next, for the neurons in the last hidden layer,

the layer before the output layer, compute the derivatives then use them to update the incoming edges for those neurons. Repeat this process for each neuron in the network. Eventually you will reach the input layer where there are no incoming edges and hence no weights to update. This process should be done for each input-output pair in the training data.

In testing a standard Seq-Seq CNN displayed an accuracy of 90.7%. (Zolkepli and Divino, n.d.). This is slightly higher than the ARIMA model but is still not satisfactory for use in the stock market.

3.11 RNN

The traditional CNN started to become of great interest in the 2000s and started to evolve to fit many purposes. One such evolution is into the recurrent neural network.

Recurrent neural networks were created because there were a few issues with the standard feed forward CNN. The issues came about as the input started to change from just being a few numbers to images represented as a matrix. The main issues were that it couldn't handle sequential data, only considered the current input and that it couldn't memorise previous inputs. An RNN works on the principle of saving the output of a particular layer and then feeding this back to the input in order to predict the output of the layer. This gives them the ability to remember their last input.

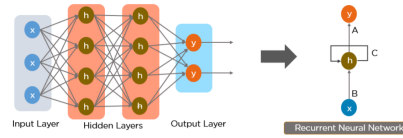


Figure 13: RNN visualised

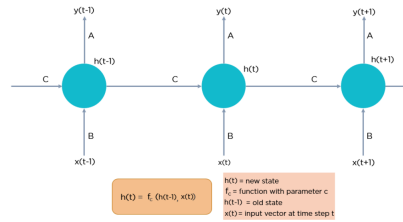


Figure 14: RNN broken down into individual components

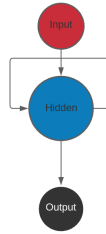


Figure 15:

3.11.1 Problems with RNN

The RNN has two big problems: a vanishing gradient problem and an exploding gradient problem.

As previously mentioned, the gradient descent algorithm finds the minimum of a function to find the optimal weights.

In RNNs, information first travels through time which means that information from previous time points is used as the input for the next time points. Secondly the cost/error function is calculated at each time point.

Essentially, every single neuron that participated in the calculation of the output, associated with this cost function, should have its weight updated in order to minimize that error. And the thing with RNNs is that it's not just the neurons directly below this output layer that contributed but all of the neurons far back in time. So, you have to propagate all the way back through time to these neurons.

The problem relates to updating wrec (weight recurring) – the weight that is used to connect the hidden layers to themselves in the unrolled temporal loop. For instance, to get from x_{t-3} to x_{t-2} we multiply x_{t-3} by wrec. Then, to get from x_{t-2} to x_{t-1} we again multiply x_{t-2} by wrec. So, we multiply with the same exact weight multiple times, and this is where the problem arises: when you multiply something by a small number, your value decreases very quickly.

As we know, weights are assigned at the start of the neural network with the random values, which are close to zero, and from there the network trains them up. But, when you start with wrec close to zero and multiply x_t , x_{t-1} , x_{t-2} , x_{t-3} , ... by this value, your gradient becomes less and less with each multiplication. (Biswal, 2021)

3.12 LSTM

All RNNs have feedback loops to help them maintain the information in the memory while training but to resolve the problems mentioned above, the LSTM was created. Long-short term memory networks are a type of RNN that introduces the use of gates such as input and forget gates which allow for better control over the gradient flow hence allowing a better preservation of long-range dependencies. These gates use functions that we have mentioned before such

as the tanh function, and the sigmoid function. (Olah,2015) The LSTM has a

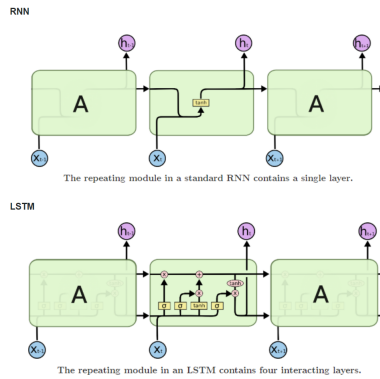


Figure 16:

block called the cell state, the middle block in the diagram above. The network has the ability to remove or add information to the cell state so that it can be accessed at a later time. This cell state is protected by the mathematical functions so that it doesn't get cluttered with useless information

3.12.1 Problems with LSTM

As you can see from the image above, there is a sequential path from the oldest past cells to the current ones. This sequential path is now more complex with the addition of forget gates. This makes them very resource intensive and so requires an immense amount of memory. This makes them very inefficient and so unsuitable for small devices such as home devices like Amazon Echo or Apple home pods.

LSTMs also get affected by different random weight initialisations and hence act just like a vanilla feed forward neural network with extra steps making them nearly useless. They much prefer small weight initialisations.

Along with these problems, LSTMs are prone to overfitting and are difficult to apply dropout algorithms to resolve this issue. To add to this list is the fact that LSTMs struggle to retain information when handling larger datasets. They can handle sequences of 100s but not 1000s or above. This makes them unsuitable for tasks like stock prediction or neural language processing.

3.13 Transformers

The Transformer is an evolution of the RNN and its main purpose was to solve the problem of parallelisation. It only performs a small, constant number of steps. In each step it applies a self-attention mechanism that helps it retain information for longer making it more suitable for larger datasets. This model is typically used for neural language processing where keeping a track of all

the words used helps the machine to gain understanding about the context. A common neural network would typically contain an encoder that would read the input sentence and would generate a representation of it. A decoder would then generate the output sentence while looking back to the representation the encoder created. The Transformer takes a different approach; it starts by generating initial representations/embeddings for each word. A self attention mechanism is then applied to the representation which generates a new representation for the word by comparing it to previous words. This step of making a new representation is repeated multiple times in parallel for all the words in the input sentence.

This parallelisation increases its computational performance and produces a high accuracy. (Vaswani et al., 2017)

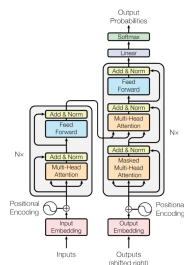


Figure 17: transformer model

3.14 Attention

(Lambda, 2019) A critical and apparent disadvantage of this fixed-length context vector design is the incapability of the system to remember longer sequences. Often it has forgotten the earlier parts of the sequence once it has processed the entire sequence. The attention mechanism was born to resolve this problem.

A neural network is considered to be an effort to mimic the brain in a simplified manner. The attention mechanism can be described as an attempt to mimic the action of selectively concentrating on a few relevant things instead of everything. There are two types of attention: general attention and self-attention. General attention is between the input and output elements and self-attention is within the input elements

The central idea behind attention is not to throw away any intermediate encoder states in LSTMs and Transformers. Instead these states are used to construct the context vectors required by the decoder to generate the output sequence.

A look into how self attention is calculated using vectors.

The first step in calculating self-attention is to create 3 vectors from each of the encoder's input vector, a query q , a key k and a value v .

The second step is to calculate a score. The score determines how much focus to place on other parts of the input as the input is encoded at a certain position.

The score is calculated by taking the dot product of the query vector with the key vector of the respective value we are scoring. So if we are processing the self attention for the value in position 1 then the first score would be the dot product of q_1 and k_1 . The second score would be the dot product of q_1 and k_2 . The third step is to divide the scores by the square root of the dimension of the key vectors used. This leads to having a more stable gradient. This value will then be passed through the softmax function. This helps normalise the scores so that they are all positive and add up to 1. The fifth step is to multiply each value vector by the softmax score (in preparation to sum them up). The intuition here is to keep intact the values of the word(s) we want to focus on, and drown-out outliers (by multiplying them by tiny numbers like 0.001, for example). The sixth step is to sum up the weighted value vectors. This produces the output of the self-attention layer at this position for the first value.

3.15 Temporal Fusion Transformers

(Lim, 2020) Attention is very useful for natural language processing but when handling time series data, modifications have to be made. The temporal fusion transformer is the evolution made for this type of data. The TFT has 5 main

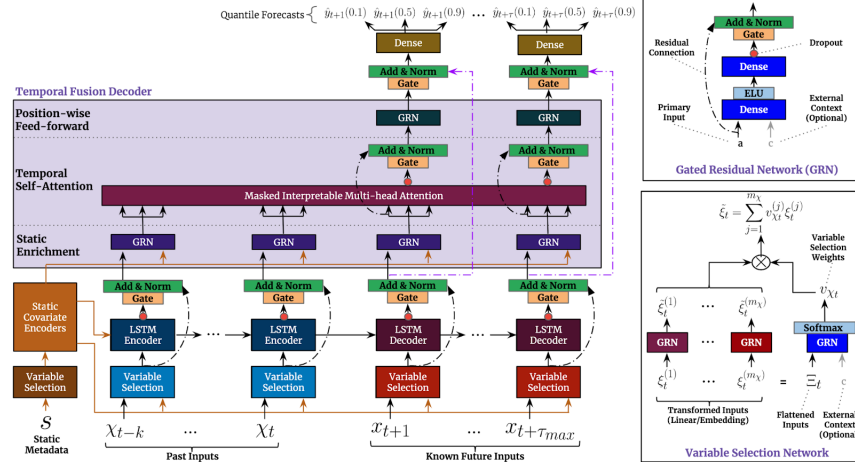


Figure 18: Temporal Fusion Transformer broken down into its components

parts that get you from a dataset to a prediction. These allow the TFT to efficiently predict future data points with a high performance.

1. Variable selection networks are used to select relevant input variables at each time step.
2. The gating mechanism to skip over any unused blocks and components of the architecture to allow for a dynamic model that adjusts itself to account for the dataset hence making it more time and memory efficient. This helps accommodate for larger datasets as well making the model more versatile.
3. The static covariate encoders are used to integrate start features into the network by encoding context vectors to condition temporal dynamics.
4. Temporal processing is used to learn both short and long term relationships between past and current inputs. Here a sequence to sequence layer with a multi-head attention block.
5. Prediction intervals via the forecasts to calculate the range of likely values at each step.

3.16 Tensors

An n th-rank tensor in m dimensional space is a mathematical object that has n indices and m^n components that obeys certain transformation rules

3.16.1 Tensor Product

For any two vector spaces \vec{U}, \vec{V} over the same field F , we can construct a tensor product $\vec{U} \otimes \vec{V}$ which is also an F-vector space.

Example:

$$\vec{v} = \begin{pmatrix} 1 \\ 2 \\ 3 \end{pmatrix} \quad \vec{w} = \begin{pmatrix} 4 \\ 5 \end{pmatrix}$$

V is a vector in \mathbb{R}^2 . W is a vector in \mathbb{R}^3 .

$$\vec{v} \otimes \vec{w} \rightarrow \begin{pmatrix} 1 \cdot 4 \\ 1 \cdot 5 \\ 2 \cdot 4 \\ 2 \cdot 5 \\ 3 \cdot 4 \\ 3 \cdot 5 \end{pmatrix} = \begin{pmatrix} 4 \\ 5 \\ 8 \\ 10 \\ 12 \\ 15 \end{pmatrix}$$

3.17 Gated Linear Units(GLU)

GLUs can be defined by the following equation:

$$h(x) = (x \cdot W + b) \otimes \sigma(X \cdot V + c)$$

Given a tensor, two independent convolutions are done and we get two outputs. We further do an additional sigmoid activation for one for the output, and find the tensor product of the two outputs.

3.18 Gated Residual Network(GRN)

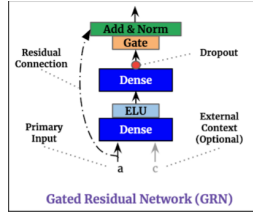


Figure 19: GRN visualised

The gated residual network, as shown in the image above, consists of a dense block with an exponential linear unit as the activation function. This is then fed into another dense block before a GLU and normalised. There will be a connection between the input and output to allow for the attention mechanism to work. In my model, there will not be external context. The GRN can be modelled using the equations below:

$$GRN(\alpha) = LayerNorm(\alpha + GLU(\phi_1))$$

$$\begin{aligned}
\phi_1 &= W_1 \phi_2 + b_1 \\
\phi_2 &= ELU(W_2 \alpha + W_3) \\
\phi_1, \phi_2 &\in R
\end{aligned}$$

3.19 Variable Selection Network

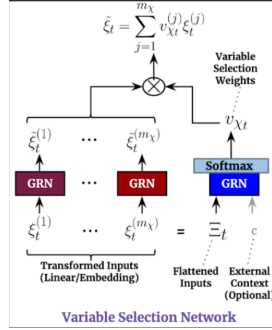


Figure 20: Variable Selection Network visualised

The purpose of the variable selection network is to evaluate the relevance of the variables that can be fed back into the model. This helps tell us what variables are the most significant as well as removing any unnecessary, noisy inputs that could negatively impact the performance of the model. This is useful when dealing with real time data as there might not be enough time to make sure all the data is correct or useful. As shown in the image above, there will be three GRNs with one of them having a softmax as the activation function. The tensor product will then be calculated.

3.20 Multi-head attention

The TFT model uses a self-attention mechanism to learn long term relationships between the data points across the different time steps. The concept of attention has already been explained and how that is calculated has also been discussed. To fit the TFT, a few modifications have been made. The general self-attention relies on the head value being the same but in our case this is different. This means the attention mechanism would have to only rely on the attention weights which is not enough. This problem has been solved by modifying the multi-head attention to share values in each head and using additive aggregation of all the heads.

$$\begin{aligned}
MultiHead(\vec{Q}, \vec{K}, \vec{V}) &= [\vec{H}_1, \dots, \vec{H}_h] \cdot \vec{W}_h \\
\vec{H}_h &= Attention(\vec{Q}\vec{W}_Q, \vec{K}\vec{W}_K, \vec{V}\vec{W}_V) \\
InterpretableMultiHead(\vec{Q}, \vec{K}, \vec{V}) &= \vec{\tilde{H}} \cdot \vec{W}_H
\end{aligned}$$

$$\begin{aligned}\vec{H} &= \tilde{A}(\vec{Q}, \vec{K}) \cdot \vec{V}W_V \\ &= \frac{1}{H} \sum_{h=1}^{M_h} \text{Attention}(\vec{Q}W_Q, \vec{K}W_K, \vec{V}W_V)\end{aligned}$$

The first equation shows what the equation would be when multiple heads are added. Q,K and V are the query vectors, key vectors and value vectors. Wv are the value weights shared across all the heads. The interpretable multi-head function is our modified equation. The last equation shows us how we have employed additive aggregation.

3.21 Quantile Forecasts

The TFT can generate prediction intervals as well as point forecasts. This is done by predicting various percentiles (10%, 50%, 90%) at each time step. The quantile forecast is calculated using the outputs of the temporal fusion decoder as is written as this:

$$\begin{aligned}\tilde{\xi} &= GRN(\xi) \\ \phi(t, n) &= \text{LayerNorm}(\tilde{\xi} + GLU(\phi(t, n))) \\ (t, n) &= \text{LayerNorm}(\phi(t, n) + GLU(\psi(t, n))) \\ \hat{y}(q, t, n) &= \vec{W}_q \psi(t, n) + b_q\end{aligned}$$

The equations above shows us what is being calculated to end up with the last equation which is our quantile forecast.

3.22 Software

3.22.1 Coding Language

I have chosen to use python to code my project due to a few reasons. The first reason is the flexibility of the language. I have the option to use OOP and functional programming as well as not having to recompile the code every time I want to run the code. Python can also be combined with other languages such as C which makes it more powerful when dealing with a lot of maths. Another reason is that Python has a very big library for machine learning. This makes prototyping and implementing my own algorithm a lot easier as well as being able to visualise my code a lot easier with packages such as pyplot. My other option was C/C++ where I would've had benefits such as more compact and faster runtime. Python does offer extensions or libraries that optimise the Python code. An example is Theano. Unlike C++ where specific optimisations need to be made, Python can run nearly any system without any time being wasted on specific configurations. Python also has access to libraries such as CUDA python. Libraries like that are made to help with parallelisation on GPUs. This exploits the tensor cores that are offered by the hardware manufacturers. The power to be able offload tasks to the GPU makes any performance

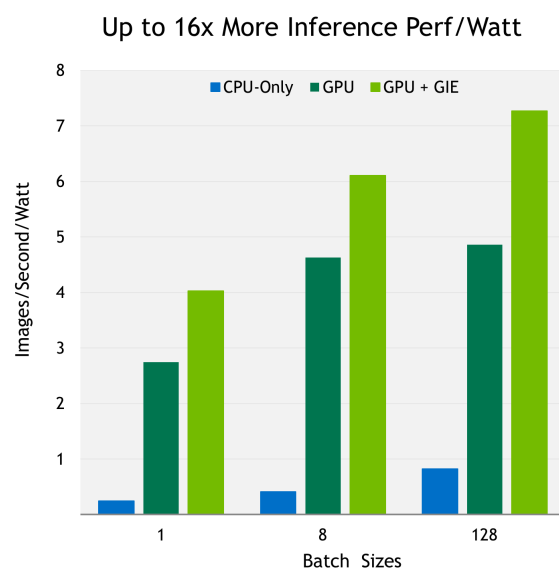


Figure 21:

Inference. Tests performed using
 GoogLeNet. CPU-only: Single-socket
 Intel Xeon (Haswell) E5-2698
 v3@2.30GHz with HT.
 GPU: NVIDIA Tesla M4 + cuDNN 5 RC.
 GPU + GIE: NVIDIA Tesla M4 + GIE.

Figure 22:

advantage C++ had, instantly insignificant. GIE is the NVIDIA GPU inference engine. The graph shows the advantage GPUs provide for training deep learning models.

3.22.2 Packages

For my project I have considered using packages to aid the learning process such as Theano. Theano is a CPU and GPU mathematical compiler that outperforms other tools. Theano includes tools for manipulating and optimising graphs representing mathematical functions. An example is the sigmoid function that is used heavily in machine learning. Theano's optimisation also includes the elimination of duplicates or unnecessary computations therefore increasing the numerical stability as well as increasing speed. The graph representation allows the user to quickly prototype machine learning models as the differentiation of the function does not have to be calculated in algorithms such as backpropagation. This reduces the amount of code that has to be executed. Theano also uses CUDA to define a class of n-dimensional arrays located in GPU memory with Python bindings. This speeds up the number of operations that are performed per second. Theano can also leverage multi-core CPU architectures to allow for parallelisation on both the CPU and the GPU. Some disadvantages of using Theano include the speed of compiling is a lot larger with bigger models. The error messages are also difficult to understand which will make debugging harder. Theano is a quite low level so the learning curve is steeper. Another package that I could use is Tensorflow. Tensorflow has a flexible architecture and can be deployed on more than one GPU or CPU. As well as this it shares all the features that Theano has. The reason I am not using it is because it is very high level and does not give you enough control at the base layer.

3.23 Objectives

The aim of this project is to be able to make predictions on the price of a stock given historic data of the stock.

3.23.1 General Objectives

Data processing

1. Must be able to accept a CSV file as an input
2. Must normalise the data set

Prediction

1. Must pass the data through a variable selection block
2. Must pass the data through an LSTM encoder
3. Must pass the data through a series of GRNs

4. Must implement multi head attention
5. Must return quantile forecasts for the following 5 days

Forecast processing

1. Must decide whether to recommend a sell, short or buy
2. Must calculate the confidence level of the forecast

User interface

1. Must display the forecast along with a confidence level
2. Must give recommendations for buying, shorting or selling
3. Consider giving alerts to help the user

3.23.2 Specific objectives

Gated Residual network

1. Must have a dense network
2. Must have an exponential linear unit
3. Must have layer normalisation
4. Must have a residual connection between the input and output

Variable Selection block

1. Must have three GRNs
2. Must have a GRN with a softmax function
3. Must calculate the tensor product of the three GRNs

LSTM encoder-decoder

1. Must have an encoder model
2. Must have a decoder model
3. Must have a dense block

Multi-head attention

1. Must take the embedding size, the query size and attention heads as an input
2. Must have input embedding and position encoding layers to produce a matrix of shape

3. The matrix must be fed to the query, key and value of the first encoder in the stack
4. The input must be passed through linear layers to produce the Q, K and V matrices
5. The data must get split across the attention heads
6. The Q, K and V matrices must be reshaped
7. An attention score must be computed for each head
8. The attention scores must be merged together

4 Documented Design

4.0.1 Heirarchy Diagram

The project can be split into 3 main sections: data preprocessing, the temporal fusion transformer model and the quantile forecasts. Data processing can also

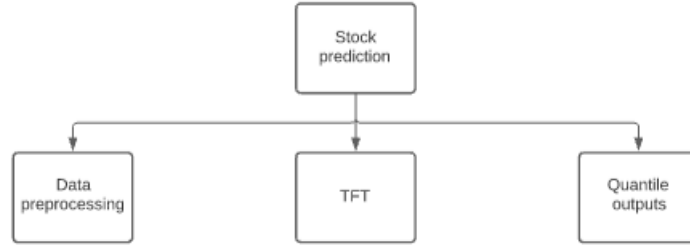


Figure 23:

be split into 3 parts: Processing, normalisation and adding time embeddings. Below is a structure diagram of the Temporal Fusion Transformer. It is split

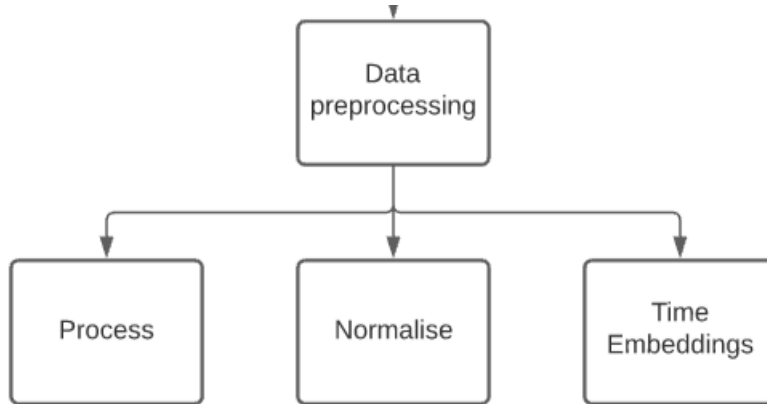


Figure 24:

into 4 sections: Gated Blocks, LSTM, Variable Selection Network and Multi-head Attention. Each of those sections are further subdivided into the different types of that object. The Quantile outputs are split into functions that are used in the main TFT model, layer normalisation, GRU and GLU. These together will form the forecasts at the end. The Gated Residual unit can be broken up into 3 parts: the dense network, the exponential unit and the addition and normalisation layer. This figure shows an overview of the Data Preprocessing. This part of the program will ensure that all the data is fit to be fed to the

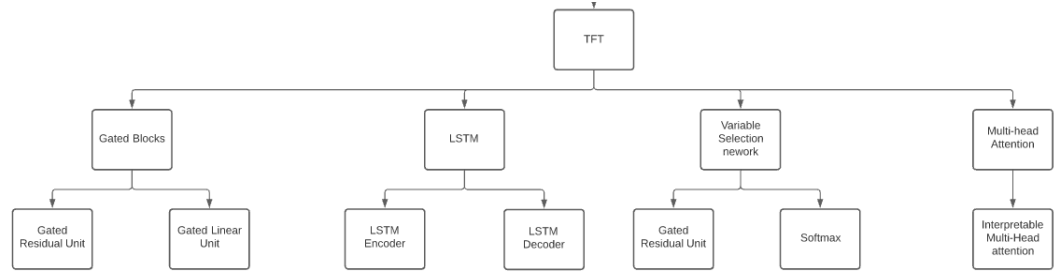


Figure 25:

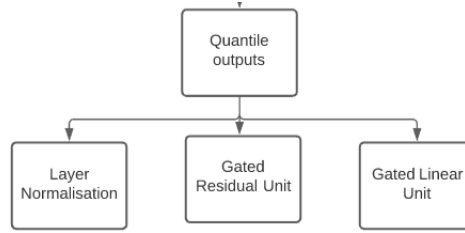


Figure 26:

model and will convert the values into a form that is best for the machine learning module(TFT). This process includes removing all null values, creating the appropriate datasets, vectorising them, adding time embeddings and then finally splitting the datasets into batches. Removing null values will ensure that sudden spikes in the data do not add a bias to the model. A spike will have a big derivative and might cause the wrong neurons to be activated. Normalising a dataset will ensure that all the data lies within 0n1. It is required as my data have different ranges. Having all values between the same range is needed to make sure larger values in a dataset don't have a greater importance/bias. This will help speed up the model as the calculations and multiplication will be done with smaller numbers. Rounding is acceptable as we are dealing with a machine learning model and not a statistical learning model. Rounding will not have a significant impact on the overall accuracy. The dataset has to be split into 3 parts: the training dataset, the validation dataset and the testing dataset. The split will be roughly 80:5:15. It is important to make sure that the training data is not used when testing as this will give us an hyperinflated accuracy, making the model seem a lot better than it actually is. The datasets then have to be vectorised as the model will only take in vectors, matrices or tensors. Time embeddings are added to the data. This is to help the attention part of the model. Attention works by referring back to the previous data points and so each data point has to be given a time stamp/embedding. The

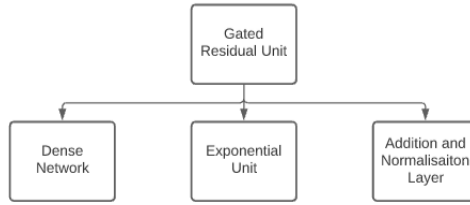


Figure 27:

positional encoder will assign data points a value to determine their position in the dataset. This is also used to help make sure the attention module works, The dataset will finally be split up into batches. This is so that I can conduct batch training. This helps the model focus on small sections of the data instead of trying to consider all the data at once. This helps the code be more memory efficient and will lead to a higher accuracy score.

The figure above shows how the attention mechanism works. The module takes in 3 input vectors: Query, Key and Value. These are fed into the attention block and an Attention score is calculated using the equation detailed in the analysis section. The multiple linear and attention score blocks represent the number of heads in the attention block. Above is a flowchart to show the travel of data in my program. The TFT module starts off with the quantile loss and ends with the output layer. The logging metric is going to be in the form of a tensorboard. This tracks important data such as the change in loss as the number of epochs increases. I can track and see the change in weights through the neural network. This helps me evaluate the performance of certain blocks in the TFT. The tensorboard will also allow me to compare different initialisation techniques such as truncated and xavier. The most important reason is to find out when to stop training the model. The graph of loss vs epoch can be thought of as a 1x graph for 0x. As the graph starts to level off, the net effect of each epoch decreases. Training a machine learning algorithm is time, space and cost intensive so any effort to reduce the number of epochs has to be taken. Analysing the graph provided by the tensorboard will allow this. The main part of the machine learning model(TFT) aims to pass the data through many blocks that will analyse the overall trend in the data while observing patterns in the data. The use of LSTMs and Gated Residual Units contain memory blocks that help the neural network consider context. At the start there are encoders and so therefore later on decoders are required. Multihead attention is used later on to use attention to provide further context to the model.

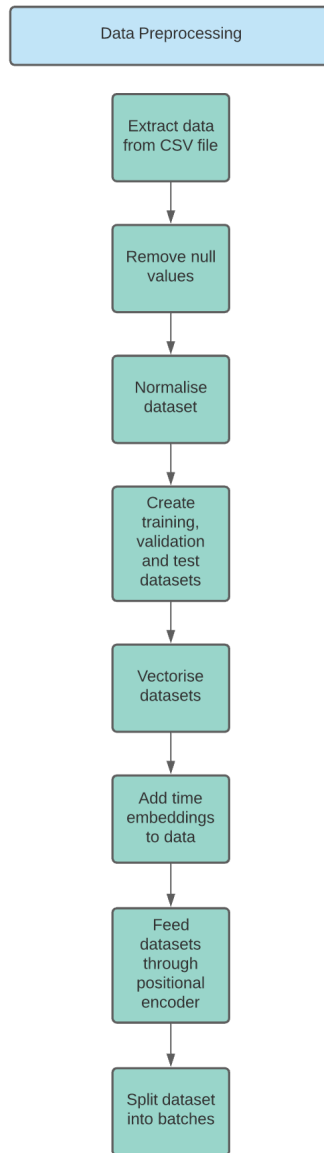


Figure 28:

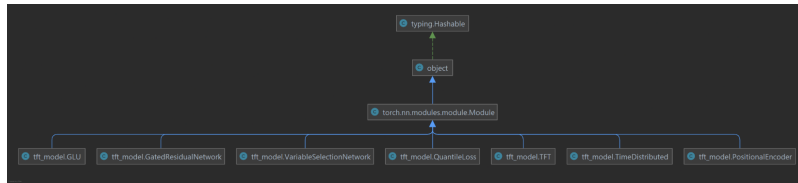


Figure 29:

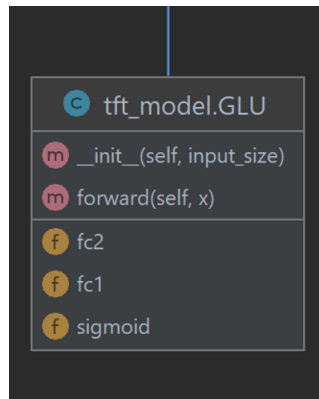


Figure 30:

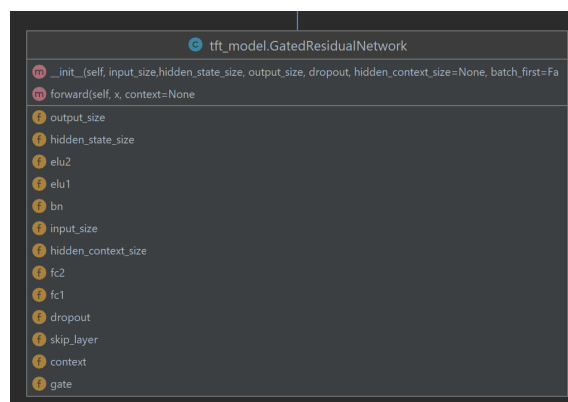


Figure 31:

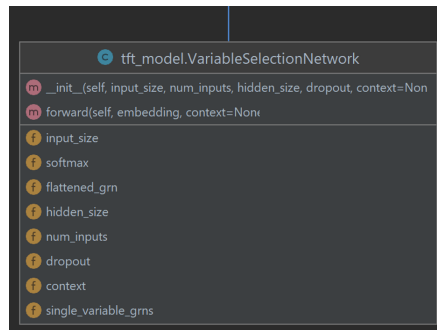


Figure 32:

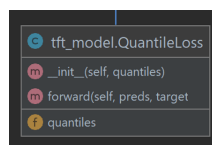


Figure 33:

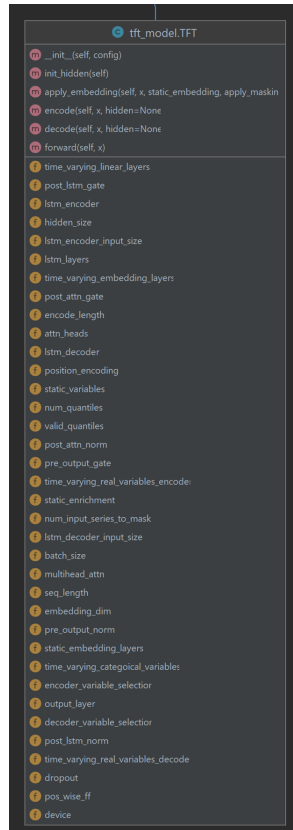


Figure 34:

Figure 35:

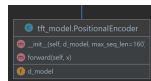


Figure 36:

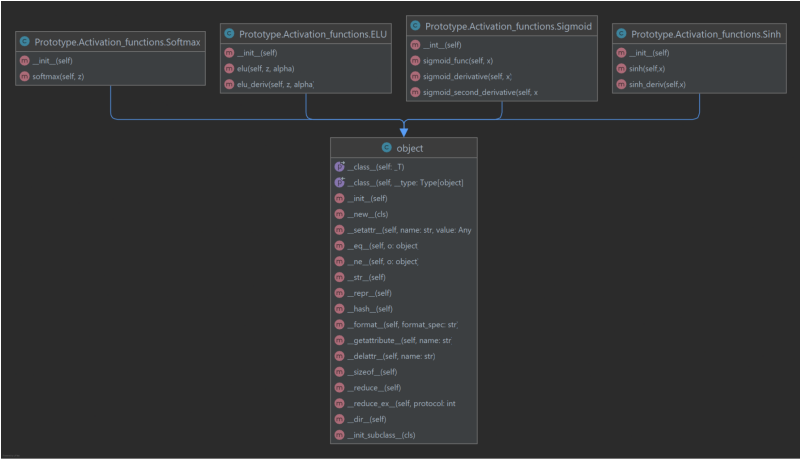


Figure 37:

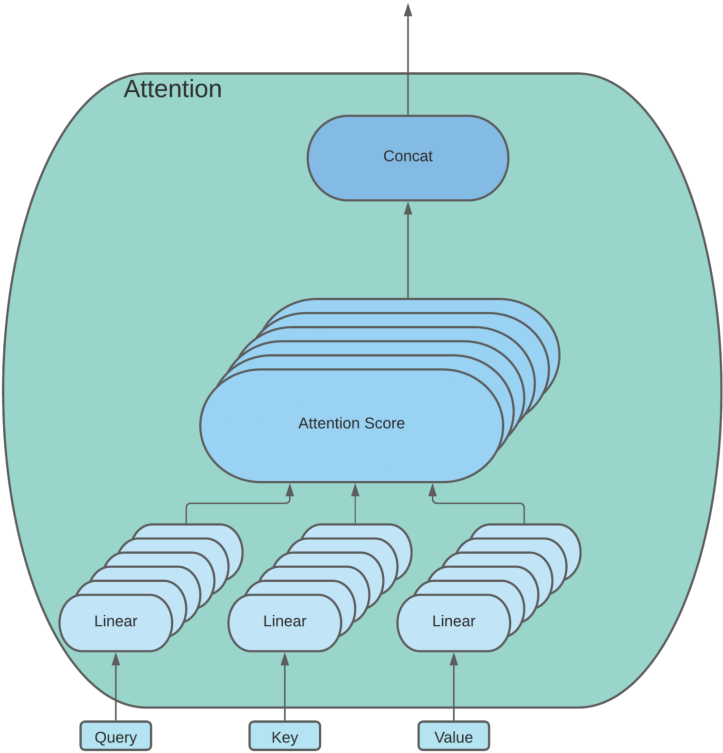


Figure 38:

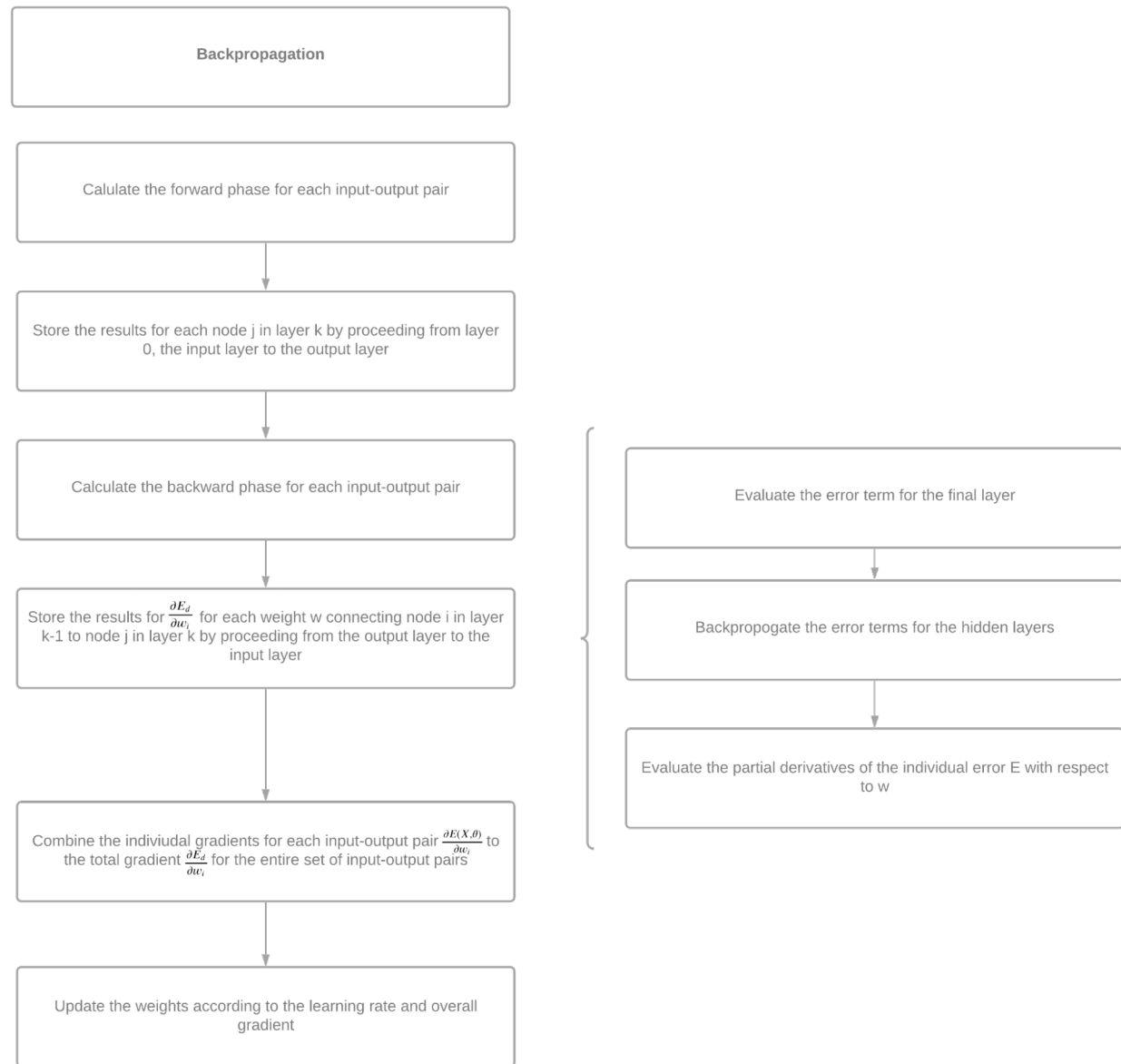


Figure 39:

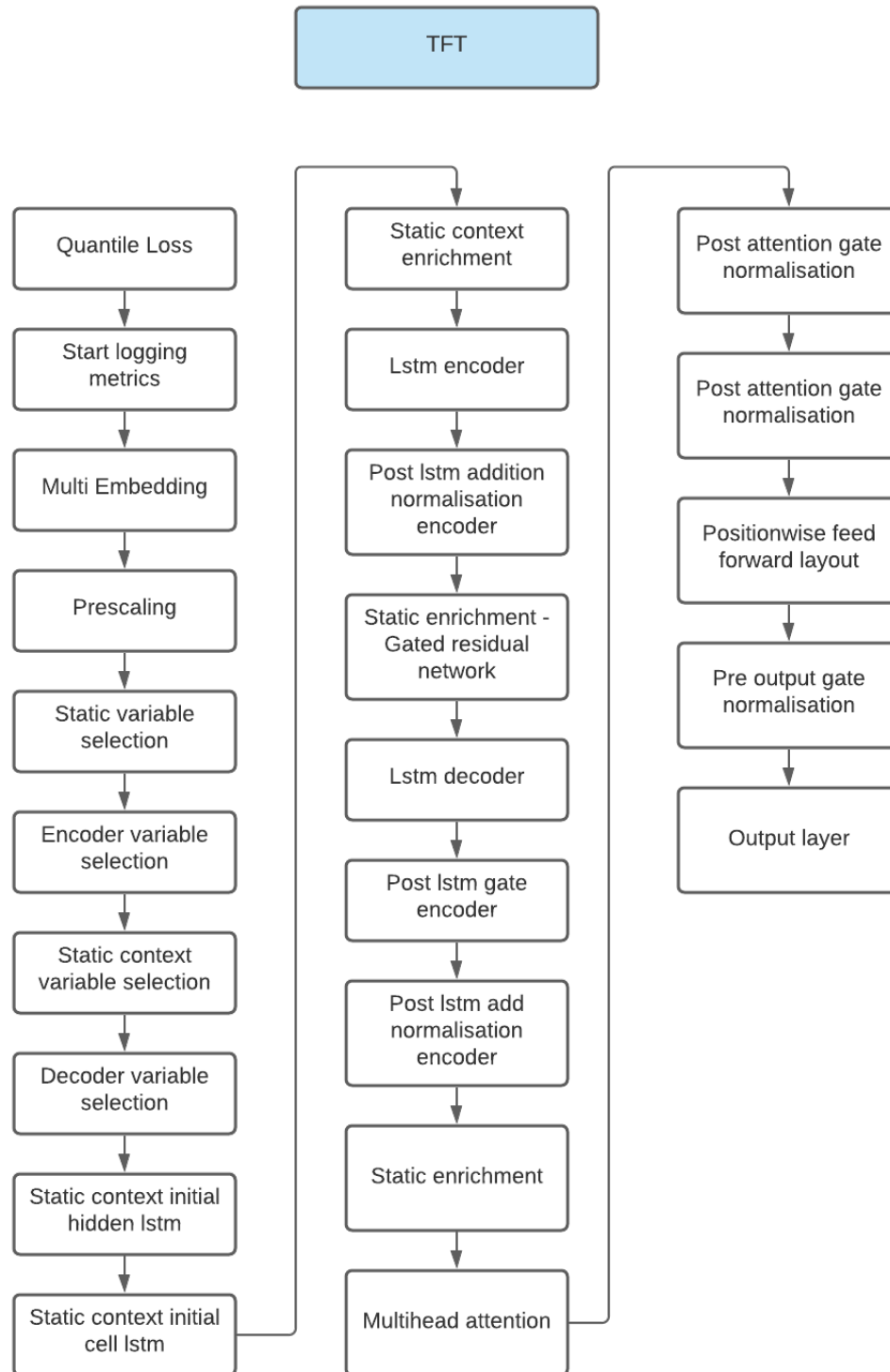


Figure 40:

The TFT can be modelled using the structure below. This shows the flow of data through one epoch of the network. Each module has the dimensions, input features, output features, activation functions and learning rate specified.

```
TemporalFusionTransformer(
  (loss): QuantileLoss()
  (logging_metrics): ModuleList(
    (0): SMAPE()
    (1): MAE()
    (2): RMSE()
    (3): MAPE()
  )
  (input_embeddings): MultiEmbedding(
    (embeddings): ModuleDict()
  )
  (prescalers): ModuleDict()
  (static_variable_selection): VariableSelectionNetwork(
    (single_variable_grns): ModuleDict()
    (prescalers): ModuleDict()
    (softmax): Softmax(dim=-1)
  )
  (encoder_variable_selection): VariableSelectionNetwork(
    (single_variable_grns): ModuleDict()
    (prescalers): ModuleDict()
    (softmax): Softmax(dim=-1)
  )
  (decoder_variable_selection): VariableSelectionNetwork(
    (single_variable_grns): ModuleDict()
    (prescalers): ModuleDict()
    (softmax): Softmax(dim=-1)
  )
  (static_context_variable_selection): GatedResidualNetwork(
    (fc1): Linear(in_features=16, out_features=16, bias=True)
    (elu): ELU(alpha=1.0)
    (fc2): Linear(in_features=16, out_features=16, bias=True)
    (gate_norm): GateAddNorm(
      (glu): GatedLinearUnit(
        (dropout): Dropout(p=0.1, inplace=False)
        (fc): Linear(in_features=16, out_features=32, bias=True)
      )
      (add_norm): AddNorm(
        (norm): LayerNorm((16,)), eps=1e-05, elementwise_affine=True)
      )
    )
  )
  (static_context_initial_hidden_lstm): GatedResidualNetwork(
    (fc1): Linear(in_features=16, out_features=16, bias=True)
    (elu): ELU(alpha=1.0)
    (fc2): Linear(in_features=16, out_features=16, bias=True)
    (gate_norm): GateAddNorm(
```

```

        (glu): GatedLinearUnit(
          (dropout): Dropout(p=0.1, inplace=False)
          (fc): Linear(in_features=16, out_features=32, bias=True)
        )
        (add_norm): AddNorm(
          (norm): LayerNorm((16,)), eps=1e-05, elementwise_affine=True)
        )
    )
    (static_context_initial_cell_lstm): GatedResidualNetwork(
      (fc1): Linear(in_features=16, out_features=16, bias=True)
      (elu): ELU(alpha=1.0)
      (fc2): Linear(in_features=16, out_features=16, bias=True)
      (gate_norm): GateAddNorm(
        (glu): GatedLinearUnit(
          (dropout): Dropout(p=0.1, inplace=False)
          (fc): Linear(in_features=16, out_features=32, bias=True)
        )
        (add_norm): AddNorm(
          (norm): LayerNorm((16,)), eps=1e-05, elementwise_affine=True)
      )
    )
    (static_context_enrichment): GatedResidualNetwork(
      (fc1): Linear(in_features=16, out_features=16, bias=True)
      (elu): ELU(alpha=1.0)
      (fc2): Linear(in_features=16, out_features=16, bias=True)
      (gate_norm): GateAddNorm(
        (glu): GatedLinearUnit(
          (dropout): Dropout(p=0.1, inplace=False)
          (fc): Linear(in_features=16, out_features=32, bias=True)
        )
        (add_norm): AddNorm(
          (norm): LayerNorm((16,)), eps=1e-05, elementwise_affine=True)
      )
    )
    (lstm_encoder): LSTM(16, 16, batch_first=True)
    (lstm_decoder): LSTM(16, 16, batch_first=True)
    (post_lstm_gate_encoder): GatedLinearUnit(
      (dropout): Dropout(p=0.1, inplace=False)
      (fc): Linear(in_features=16, out_features=32, bias=True)
    )
    (post_lstm_gate_decoder): GatedLinearUnit(
      (dropout): Dropout(p=0.1, inplace=False)
      (fc): Linear(in_features=16, out_features=32, bias=True)
    )
    (post_lstm_add_norm_encoder): AddNorm(
      (norm): LayerNorm((16,)), eps=1e-05, elementwise_affine=True)
    )

```



```

(post_lstm_add_norm_decoder): AddNorm(
  (norm): LayerNorm((16,), eps=1e-05, elementwise_affine=True)
)
(static_enrichment): GatedResidualNetwork(
  (fc1): Linear(in_features=16, out_features=16, bias=True)
  (elu): ELU(alpha=1.0)
  (context): Linear(in_features=16, out_features=16, bias=False)
  (fc2): Linear(in_features=16, out_features=16, bias=True)
  (gate_norm): GateAddNorm(
    (glu): GatedLinearUnit(
      (dropout): Dropout(p=0.1, inplace=False)
      (fc): Linear(in_features=16, out_features=32, bias=True)
    )
    (add_norm): AddNorm(
      (norm): LayerNorm((16,), eps=1e-05, elementwise_affine=True)
    )
  )
)
(multihead_attn): InterpretableMultiHeadAttention(
  (dropout): Dropout(p=0.1, inplace=False)
  (v_layer): Linear(in_features=1, out_features=4, bias=True)
  (q_layers): ModuleList(
    (0): Linear(in_features=16, out_features=4, bias=True)
    (1): Linear(in_features=16, out_features=4, bias=True)
    (2): Linear(in_features=16, out_features=4, bias=True)
    (3): Linear(in_features=16, out_features=4, bias=True)
  )
  (k_layers): ModuleList(
    (0): Linear(in_features=16, out_features=4, bias=True)
    (1): Linear(in_features=16, out_features=4, bias=True)
    (2): Linear(in_features=16, out_features=4, bias=True)
    (3): Linear(in_features=16, out_features=4, bias=True)
  )
  (attention): ScaledDotProductAttention(
    (softmax): Softmax(dim=2)
  )
  (w_h): Linear(in_features=4, out_features=16, bias=False)
)
(post_attn_gate_norm): GateAddNorm(
  (glu): GatedLinearUnit(
    (dropout): Dropout(p=0.1, inplace=False)
    (fc): Linear(in_features=16, out_features=32, bias=True)
  )
  (add_norm): AddNorm(
    (norm): LayerNorm((16,), eps=1e-05, elementwise_affine=True)
  )
)
(pos_wise_ff): GatedResidualNetwork(
  (fc1): Linear(in_features=16, out_features=16, bias=True)
  (elu): ELU(alpha=1.0)

```

```

(fc2): Linear(in_features=16, out_features=16, bias=True)
(gate_norm): GateAddNorm(
  (glu): GatedLinearUnit(
    (dropout): Dropout(p=0.1, inplace=False)
    (fc): Linear(in_features=16, out_features=32, bias=True)
  )
  (add_norm): AddNorm(
    (norm): LayerNorm((16,), eps=1e-05, elementwise_affine=True)
  )
)
)
(pre_output_gate_norm): GateAddNorm(
  (glu): GatedLinearUnit(
    (fc): Linear(in_features=16, out_features=32, bias=True)
  )
  (add_norm): AddNorm(
    (norm): LayerNorm((16,), eps=1e-05, elementwise_affine=True)
  )
)
(output_layer): Linear(in_features=16, out_features=7, bias=True)
)

```

4.1 Explaining nn.module

Below is the pseduocode for the module followed by the python code.

Algorithm 1 Linear Neural Network

```

1:  $Convolution1 \leftarrow ConvertTo2d(x, y, z)$ 
2:  $Convolution2 \leftarrow ConvertTo2d(u, v, w)$ 
3:  $y \leftarrow Wx + b$ 
4:  $fc1 \leftarrow LinearTransformation()$  {Apply a linear transformation}
5:  $fc2 \leftarrow LinearTransformation()$ 
6:  $fc3 \leftarrow LinearTransformation()$ 
7:  $x \leftarrow Apply2DPool(RELU(Convolution1(x)), (2, 2))$ 
8:  $x \leftarrow Apply2DPool(RELU(Convolution1(x)), 2)$ 
9:  $RELU(fc1(x))$ 
10:  $RELU(fc2(x))$ 
11: return  $x = 0$ 

```

```

import math
import numpy as np

#Vanilla Neural Network from scratch
class Layer_Dense:
    def __init__(self, n_inputs, n_neurons):
        self.weights = 0.10 * np.random.randn(n_inputs, n_neurons)

```

```

        self.biases = np.zeros((1, n_neurons))

    def forward(self, inputs):
        self.output = np.dot(inputs, self.weights) + self.biases

#Activation Functions
class Activation_ReLU:
    def forward(self, inputs):
        self.output = np.maximum(0, inputs)

class Activation_sigmoid:
    def forward(self, inputs):
        self.output = np.exp()

layer1 = Layer_Dense(2,5)
activation1 = Activation_ReLU()
layer2 = Layer_Dense(2,5)
activation2 = Activation_sigmoid()
layer1.forward(X)
print(layer1.forward(x))

#using nn.module
class Net(nn.Module):

    def __init__(self):
        super(Net, self).__init__()
        # 1 input image channel, 6 output channels, 5x5 square
        # convolution kernel
        self.conv1 = nn.Conv2d(1, 6, 5)
        self.conv2 = nn.Conv2d(6, 16, 5)

        # an affine operation: y = Wx + b
        self.fc1 = nn.Linear(16 * 5 * 5, 120)
        self.fc2 = nn.Linear(120, 84)
        self.fc3 = nn.Linear(84, 10)

    def forward(self, x):
        # Max pooling over a (2, 2) window
        x = F.max_pool2d(F.relu(self.conv1(x)), (2, 2))

        # 2 is same as (2, 2)
        x = F.max_pool2d(F.relu(self.conv2(x)), 2)

        x = x.view(-1, self.num_flat_features(x))
        x = F.relu(self.fc1(x))
        x = F.relu(self.fc2(x))
        x = self.fc3(x)

```

```
        return x

    def num_flat_features(self, x):
        size = x.size()[1:] # all dimensions except the batch dimension
        num_features = 1
        for s in size:      # Get the products
            num_features *= s
        return num_features
```

4.2 Positional Encoding

One of the reasons for using sine and cosine functions is that they are periodic and so whether the model is learning on a sequence of length 5, or of length 500, the encodings will always have the same range $[-1, 1]$. Positional embeddings are added to the original embeddings to retain all the positional information. Each vector will be parametrized and the stack row-wise to form a learnable positional embedding table.

PE_{pos+k} is some matrix which depends on k times PE_{pos}

$$\begin{aligned} PE_{pos+k,2i} &= \sin\left(\frac{pos}{a} + \frac{k}{a}\right) \\ &= \sin\left(\frac{pos}{a}\right)\cos\left(\frac{k}{a}\right) + \sin\left(\frac{k}{a}\right)\cos\left(\frac{pos}{a}\right) \\ &= (PE_{pos,2i+1})U + (PE_{pos,2i})V \\ &= (PE_{pos,2i}, PE_{pos,2i+1})(V, U) \end{aligned}$$

Algorithm 2 Positional Encoder

```
0: for pos in 0  $\leftarrow$  maximum sequence length do
0:   compute  $V \times PE(pos, 2i)$ 
0:   compute  $W \times PE(pos, 2i)$ 
0:   compute  $V \times PE(pos, 2i+1)$ 
0:   compute  $W \times PE(pos, 2i+1)$ 
0: end for=0
```

Python code:

```
#positional encoding
import torch
import math

class PositionalEncoder(torch.nn.Module):
    def __init__(self, d_model, max_len=5000):
        self.d_model = d_model # the size of the embedding vectors
        self.max_len = max_len
        # Compute the positional encodings once in log space complexity
        pe = torch.zeros(max_len, d_model)
        position = torch.arange(0, max_len).unsqueeze(1)
        div_term = torch.exp(torch.arange(0, d_model, 2) *
                              -(math.log(10000.0) / d_model))
        pe[:, 0::2] = torch.sin(position * div_term)
        pe[:, 1::2] = torch.cos(position * div_term)
        pe = pe.unsqueeze(0)
        self.register_buffer('pe', pe)

    def forward(self, x):
        x = x + self.pe[:, :x.size(1)]
        return x
```

```
def get_embedding(self, x):  
    x = x + self.pe[:, :x.size(1)]  
    return x  
  
def get_embedding_layer(self):  
    return torch.nn.Embedding(self.max_len, self.d_model)
```

4.3 Backpropagation

The formula for the backpropagation algorithm has been proven earlier in the analysis. Below is a reminder of the mathematical definition along with a flowchart and pseudocode.

Partial derivatives:

$$\frac{\partial E_d}{\partial w_j^k} = \delta_j^k o_i^{k-1}$$

Final layer's error term:

$$\delta_1^m = g'_o(a_1^m)(y_d - \hat{y}_d)$$

Hidden layers' error terms:

$$\delta_j^k = g''(a_j^k) \sum_{L=1}^{r^{k+1}} w_j^{k+1} L \delta_l^{k+1}$$

Computing the partial derivatives for each input-output pair:

$$\begin{aligned} \frac{\partial E(X, \theta)}{\partial w_{ij}^k} &= \frac{1}{N} \sum_{d=1}^N \frac{\delta}{\delta w_{ij}^k} \left(\frac{1}{2} (\hat{y}_d - y_d)^2 \right) \\ &= \frac{1}{N} \sum_{d=1}^N \frac{\partial E_d}{\partial w_{ij}^k} \end{aligned}$$

Updating weights:

$$\Delta w_{ij}^k = -\alpha \frac{\partial E(X, \theta)}{\partial w_{ij}^k}$$

Algorithm 3 Backpropagation

```

0: function BACKPROPAGATION(inputs, expected, output)
0:   deltas ← delta inputs
0:   repeat
0:     for i in deltas do
0:       error ← expected[i] - output deltas[i] ← error × sigmoid
0:       for j in hidden layers do
0:         for k in hidden weights do
0:           delta weights ← weight[j] × outputdeltas[k]
0:           deltaweights × learning rate
0:

```

```

# backpropagate() takes as input, the patterns entered, the target
    values and the obtained values.
# Based on these values, it adjusts the weights so as to balance out the
    error.
# Also, now we have M, N for momentum and learning factors respectively.
def backpropagate(self, inputs, expected, output, N=0.5, M=0.1):
    # We introduce a new matrix called the deltas (error) for the two
        layers output and hidden layer respectively.
    output_deltas = [0.0]*self.no
    for k in range(self.no):
        # Error is equal to (Target value - Output value)
        error = expected[k] - output[k]
        output_deltas[k]=error*dsgmoid(self.ao[k])

    # Change weights of hidden to output layer accordingly.
    for j in range(self.nh):
        for k in range(self.no):
            delta_weight = self.ah[j] * output_deltas[k]
            self.who[j][k]+= M*self.cho[j][k] + N*delta_weight
            self.cho[j][k]=delta_weight

    # Now for the hidden layer.
    hidden_deltas = [0.0]*self.nh
    for j in range(self.nh):
        # Error as given by formule is equal to the sum of (Weight from
            each node in hidden layer times output delta of output node)
        # Hence delta for hidden layer = sum
            (self.who[j][k]*output_deltas[k])
        error=0.0
        for k in range(self.no):
            error+=self.who[j][k] * output_deltas[k]
        # now, change in node weight is given by dsgmoid() of activation
            of each hidden node times the error.
        hidden_deltas[j]= error * dsgmoid(self.ah[j])

    for i in range(self.ni):
        for j in range(self.nh):
            delta_weight = hidden_deltas[j] * self.ai[i]
            self.wih[i][j]+= M*self.cih[i][j] + N*delta_weight
            self.cih[i][j]=delta_weight

```

4.4 Dense Network

This is the fundamental block, similar to the `nn.module`. This is another vanilla neural network.

```
class Dense_Network:
    def __init__(self, input_shape, output_shape, activation, weights,
                 biases):
        self.input_shape = input_shape
        self.output_shape = output_shape
        self.activation = activation
        self.weights = weights
        self.biases = biases
        self.output = None
        self.input = None

    def forward_pass(self, input):
        self.input = input
        self.output = np.dot(self.input, self.weights) + self.biases
        self.output = self.activation(self.output)
        return self.output

    def backward_pass(self, error):
        self.error = error
        self.error = self.error * self.activation(self.output,
                                                  derivative=True)
        self.weights_error = np.dot(self.input.T, self.error)
        self.biases_error = np.sum(self.error, axis=0)
        return self.error

    def update_weights(self, learning_rate):
        self.weights = self.weights - learning_rate * self.weights_error
        self.biases = self.biases - learning_rate * self.biases_error

    def get_weights(self):
        return self.weights
```

4.5 Weight Initialisation

4.5.1 Theory

Weight initialisation is essential when training a neural network. This process prevents activation functions from exploding or vanishing when executing a forward pass. If the weights are too large, the variance of the input data tends to increase rapidly with each pass, eventually getting to a point where the derivative tends to 0. This is why it is important to initialise the network with the right weights. The range in which the it is initialised will be governed using a technique called Xavier initialisation.

Xavier works by assigning the weights from a Gaussian distribution Consider a distrubtion with mean 0 and some finite variance as well as a linear neuron:

$$y = w_1x_1 + w_2x_2 + \dots + b$$

With each forward pass we would like the variance to reamin the same. Let's first copmuter the variance of y:

$$var(y) = var(w_1x_1 + w_2x_2 + \dots + w_nx_n + b)$$

Let's computer the variance of the terms inside the brackets as well. Considering the general term would give us:

$$var(w_ix_i) = E(x_i)^2var(w_i) + E(w_i)^2var(x_i) + var(w_i)var(x_i)$$

Where E() represents the expection of a given variable. The assumption that the inputs and weights are generated from a Gaussian distrubtion with a mean 0 allows us to ignore the expectations, giving us:

$$var(w_ix_i) = var(w_i)var(x_i)$$

Substiuting this value back into our original equation gives us:

$$var(y) = var(w_1)var(x_1) + \dots + var(w_n)var(x_n)$$

As these are distributed identically, they can be written as:

$$var(y) = N \times var(w_i) \times var(x_i)$$

If we want the variance of y to be the same as x, we should set $N \times var(w_i) = 1$, giving us: $var(w_i) = \frac{1}{N}$

This leaves us with the final formula:

$$var(w_i) = \frac{1}{N_{avg}}$$

where

$$N_{avg} = \frac{(N_{in} + N_{out})}{2}$$

The average helps preserve the backpropagated signal but is more copmutationally complex to implement.

Algorithm 4 Using Xavier

```
0: Initialise weights using Xavier
0: while not Stop-Criterion do
0:   for all i, j do
0:      $w_{i,j} = w_{i,j} - \alpha * \frac{\partial Error}{\partial w_{i,j}}$    end for
@:     =0
```

4.6 Layer Normalisation

4.6.1 The need for normalisation

The main issue that normalisation works to solve is the problem of internal covariate shift. These shifts occur during each epoch/round of training due to weights being updated and different data being processed. This means that each input to a neuron is slightly different each time. The input distribution of the inputs starts to deform. These small changes would have an insignificant impact on smaller architectures such as a single lstm or a vanilla RNN but when dealing with bigger models such as Transformers or TFTs small changes in the input distribution can amplify and have a catastrophic impact on the deep learning. Normlisation also tackles the vanishing gradient issue. This is where the activation function saturates. The functions approach a limit as the inputs tend to large values. This limit is where the gradient starts to shrink to a point where the function becomes unusable. As discussed in the activation functions section, different activation functions can be used to mitigate this problem as well as normalising the inputs.

4.6.2 Layer Normalisation

Given inputs x over a batch size m , $B = x_1, x_2, \dots, x_m$, each sample x_i contains K elements by applying transformations of the inputs using learned parameters γ and β , the outputs could be expressed as $B' = y_1, y_2, \dots, y_m$ where $y_i = LN_{\gamma, \beta}(x_i)$.

We first start of by calculating the mean and variance of each sample from the batch. For sample x , we have the mean, μ_i , and the variance, σ_i^2

$$\mu_i = \frac{1}{K} \sum_{k=1}^K x_{i,k}$$

$$\sigma_i^2 = \frac{1}{K} \sum_{k=1}^K (x_{i,k} - \mu_i)^2$$

Each sample is then normalised around a zero mean and unit variance. ϵ has been used for numerical stability incase the denominator becomes zero.

$$\hat{x}_{i,k} = \frac{x_{i,k} - \mu_i}{\sqrt{\sigma_i^2 + \epsilon}}$$

Scaling and shifting is then finally done using γ and β which are learnable parameters.

$$ey_i = \gamma \hat{x}_i + \beta \equiv LN_{\gamma, \beta}(x_i)$$

The maths above shows how this normalisation technique has no dependency on other samples in the batch.

4.6.3 Batch normalisation vs layer normalisation

Batch normalisation has a few issues that are key to training a TFT. One of which is that it is hard to parallelise batch-normalised models. The dependency between all the elements means that there is a need for synchronisation across all devices being used to train the model. Transformer models such as a TFT require a large scale setup to combat the mathematical complexity and intensities. Layer normalisation provides some degree of normalisation while not needing a dependency on other elements making it more suitable for my model.

4.6.4 Code

```
def ln(input, s, b, epsilon = 1e-5, max = 1000):
    """ Layer normalizes a 2D tensor along its second axis, which
        corresponds to batch """
    m, v = tf.nn.moments(input, [1], keep_dims=True)
    normalised_input = (input - m) / tf.sqrt(v + epsilon)
    return normalised_input * s + b
```

4.7 Time Distributed

Time Distributed

```
class TimeDistributed(nn.Module):
    def __init__(self, module, batch_first=False):
        super(TimeDistributed, self).__init__()
        self.module = module
        self.batch_first = batch_first

    def forward(self, x):
        if len(x.size()) <= 2:
            return self.module(x)
        # Squash samples and timesteps into a single axis
        x_reshape = x.contiguous().view(-1, x.size(-1)) # (samples *
            timesteps, input_size)
        y = self.module(x_reshape)
        # We have to reshape Y
        if self.batch_first:
            y = y.contiguous().view(x.size(0), -1, y.size(-1)) #
                (samples, timesteps, output_size)
        else:
            y = y.view(-1, x.size(1), y.size(-1)) # (timesteps, samples,
                output_size)

        return y
```

4.8 Activation Functions

The use of activation functions has been explained in the analysis. This section aims to provide a further understanding of how the functions are being coded and optimisation techniques I have uncovered in the process.

4.8.1 Sigmoid

```
class Sigmoid:
    def __init__(self):
        super().__init__()

    def sigmoid_func(self, x):
        return 1 / (1 + np.exp(-x)) # Calculates the sigmoid r

    def sigmoid_derivative(self, x):
        func_x = self.sigmoid_func(x)

        return func_x * (1 - func_x)

    def sigmoid_second_derivative(self, x):
```

```
fn_x = self.fn(x)
return fn_x * (1 - fn_x) * (1 - 2 * fn_x)
```

4.8.2 Softmax

The softmax function

```
class Softmax():
    def __init__(self):
        super().__init__()

    def softmax(self, z):
        assert len(z.shape) == 2
        s = np.max(z, axis=1)
        s = s[:, np.newaxis] # necessary step to do broadcasting
        e_x = np.exp(z - s)
        div = np.sum(e_x, axis=1)
        div = div[:, np.newaxis] # dito
        return e_x / div
```

4.8.3 Tanh

Here I am able to use the sinh definition to generate the first and second derivatives. I can also use these definitions to form the tanh function.

```
class Sinh:
    def __init__(self):
        super().__init__()

    def sinh(self, x):
        #doublesinh = np.exp(x) - np.exp(-x)
        comp1 = np.exp(x)
        comp2 = np.exp(-x)
        comp3 = 0.5 * (comp1 - comp2)
        return comp3

    def sinh_deriv(self, x):
        # doublesinh = np.exp(x) - np.exp(-x)
        comp1 = np.exp(x)
        comp2 = np.exp(-x)
        final = 0.5 * (comp1 + comp2)
        return final

x = Sinh()
const = 1
tanh = (x.sinh(const)) / (x.sinh_deriv(const))
```

4.8.4 ELU

```
class ELU:
    def __init__(self):
        super().__init__()

    def elu(self, z, alpha):
        return z if z >= 0 else alpha * (np.exp(z) - 1)

    def elu_deriv(self, z, alpha):
        return 1 if z > 0 else alpha * np.exp(z)
```

4.9 Attention

4.9.1 Multi-head attention

Multihead attention is done by passing the output tensor through a series of functions. The first step is to split the tensor into its neurons. The query and key is multiplied. A mask is then used before it is passed through the softmax function. Though, as previously discussed, the softmax is a very inefficient function, I have not been able to source a more efficient algorithm that also produced probabilistic outputs. The query vector is the masked before it is multiplied to the attention vector. This is then split and normalised before returned back to the network. Multihead attention is a parallel task with all heads running at the same time. Bigger models may require 8-12 heads but for my data, 4 should be appropriate for me. This is because multihead attention is a polynomial space complex algorithm so each head will increase the memory needed polynomially. With the hardware I am operating with, 4 will push the system but not cause problems. Larger number of heads are required when dealing with natural language processing. The nature of my data, time series data, means I can get away with a smaller number of heads.

```
def MultiHeadAttention(self,
                        Q, K, V):
    num_heads = self.num_heads
    N = Q.size()[0]
    # Linear projections
    Q_l = nn.ReLU()(self.linear_Q(Q))
    K_l = nn.ReLU()(self.linear_K(K))
    V_l = nn.ReLU()(self.linear_V(V))
    # Split and concat
    Q_split = Q_l.split(split_size=self.hidden_dim // num_heads,
                        dim=2)
    K_split = K_l.split(split_size=self.hidden_dim // num_heads,
                        dim=2)
    V_split = V_l.split(split_size=self.hidden_dim // num_heads,
                        dim=2)
```

```

Q_ = torch.cat(Q_split, dim=0) # (h*N, T_q, C/h)
K_ = torch.cat(K_split, dim=0) # (h*N, T_k, C/h)
V_ = torch.cat(V_split, dim=0) # (h*N, T_v, C/h)
# Multiplication
outputs = torch.bmm(Q_, K_.transpose(2, 1))
# Scale
outputs = outputs / (K_.size()[-1] ** 0.5)
# Key Masking
key_masks = torch.sign(torch.abs(K).sum(dim=-1)) # (N, T_k)
key_masks = key_masks.repeat(num_heads, 1) # (h*N, T_k)
key_masks = key_masks.unsqueeze(1).repeat(1, Q.size()[1], 1) #
(h*N, T_q, T_k)
paddings = torch.ones_like(key_masks) * (-2 ** 32 + 1)
outputs = torch.where(torch.eq(key_masks, 0), paddings, outputs)
# (h*N, T_q, T_k)
# Activation
outputs = nn.Softmax(dim=2)(outputs) # (h*N, T_q, T_k)
# Query Masking
query_masks = torch.sign(torch.abs(Q).sum(dim=-1)) # (N, T_q)
query_masks = query_masks.repeat(num_heads, 1) # (h*N, T_q)
query_masks = query_masks.unsqueeze(-1).repeat(1, 1,
K.size()[1]) # (h*N, T_q, T_k)
outputs = outputs * query_masks # broadcasting. (h*N, T_q, T_k)
# Dropouts
outputs = self.dropout(outputs)
# Weighted sum
outputs = torch.bmm(outputs, V_) # ( h*N, T_q, C/h)
# Restore shape
outputs = outputs.split(N, dim=0) # (N, T_q, C)
outputs = torch.cat(outputs, dim=2)
# Residual connection
outputs = outputs + Q_1
# Normalize
outputs = self.norm(outputs) # (N, T_q, C)
return outputs

```

4.10 Data Structures

The data is initially stored in a CSV file. The data is then converted into a 2x1 matrix before it is turned into a 2x2 matrix. These are implemented using lists in python. The data, as it goes through the model, gets converted into tensors. The attention module is modelled using dictionaries. My weights will also be stored in tensors. Derivatives will be stored in a Jacobian matrix.

4.11 Testing

4.12 File Structure

Data

- Training
- Stock data
- Validation
- Pre-training alternate dataset

Modules

- Data Preprocessing
- Activation Functions
- Attention module
- Data loaders
- Dense Network
- Gated Linear Unit
- Layer Normalisation
- LSTM
- Positional Encoder
- Time Distributed layer
- Variable Selection Network
- Loss Functions
- Batch Normalisation
- OZE Loss
- Dense Block
- MultiHead Attention
- Main TFT
- Quantile Forecasts
- Model Tuning

5 Bibliography

Zolkepli, H. and Divino, A., n.d. huseinzol05/Stock-Prediction-Models. [online] GitHub. Available at: <https://github.com/huseinzol05/Stock-Prediction-Models> [Accessed 7 February 2021].

Intuitive Understanding of Attention Mechanism in Deep Learning. [online] Medium. Available at: <https://towardsdatascience.com/intuitive-understanding-of-attention-mechanism-in-deep-learning-6c9482aecf4f> [Accessed 5 February 2021].

Biswal, A., 2021. Recurrent Neural Network (RNN) Tutorial for Beginners. [online] Simplilearn.com. Available at: <https://www.simplilearn.com/tutorials/deep-learning-tutorial/rnn> [Accessed 7 February 2021].

Vaswani, A., Shazeer, N., Parmar, N., Uszkoreit, J., Jones, L., Gomez, A., Kaiser, L. and Polosukhin, I., 2017. Attention Is All You Need.

Alammar, J., 2020. The Illustrated Transformer. [online] Jalammar.github.io. Available at: <http://jalammar.github.io/illustrated-transformer/> [Accessed 7 February 2021]. ((Alammar, 2020))

Hecht-Nielsen, R., 1988. Theory of the Backpropagation Neural Network. University of California at San Diego.

Hyndman, R. and Athanasopoulos, G., 2018. Forecasting principles and practice.

Rojas, R., 1996. Neural Networks.

Jadiker.github.io. 2018. [online] Available at: https://jadiker.github.io/dreu-2018/backprop_math.pdf > [Accessed 11 February 2021].

Lim, B., 2020. *Deep Learning for Time Series Prediction Decision Making Over Time*. DPhil. University of Oxford. <https://www.robots.ox.ac.uk/parg/pubs/theses/bryanLim.pdf> >

Joshi, P., 2016. *Understanding Xavier Initialization In Deep Neural Networks*. [online] PERPETUAL ENIGMA. <https://prateekvjoshi.com/2016/03/29/understanding-xavier-initialization-in-deep-neural-networks/> > [Accessed 16 January 2022].

here is some text