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Time series analysis of GSS bonds

Part 2 – Further univariate analysis of S&P Green
Bond Index

by D Dey

IFoA Data Science, Sustainability
& Climate Change Working Party

August 2024

Abstract

The popularity of green, social and sustainability-linked bonds (GSS bonds) continues to rise, with c.USD 939 billion of such bonds issued globally in 2023. Given the rising popularity of ESG-related investment solutions, their relative recent emergence and limited research in this field, continued investigation is essential. Extending non-traditional techniques i.e. neural networks to such fields creates a good blend of innovation and potential.

This paper follows on from our initial publication, where we aim to replicate the *S&P Green Bond Index* (i.e. this is a time series problem) over a period using non-traditional techniques (i.e. with neural networks) predicting 1 day ahead.

In this paper, we retain a univariate analysis and focus on a more complex neural network architecture design, specifically N-BEATS, with the aim of at least producing similar performance results as per the Baseline model. The N-BEATS architecture is a complex feedforward neural network architecture, consisting of basic building blocks and stacks, introducing the novel *doubly residual stacking of backcasts and forecasts*. The original authors of the model architecture noted in 2020 that this cutting-edge approach outperformed the Makridakis time series M4 competition model winner by 3% within their tests.

In this paper, we also revisit the neural network architectures from our initial publication, which include DNNs, CNNs, GRUs and LSTMs. We continue the univariate time series problem, increasing the data input window from 1 day to 2 and 5 days respectively, whilst still aiming to predict 1 day ahead. Increasing the input window should hopefully improve model performance, as we are exposing the models to more information.

The Baseline of setting today's index value equal to the prior day resulted in model predictions with an overall performance of c.0.5% (based on a MAPE measure) - though simple in design - is surprisingly accurate and difficult to materially outperform. We trained an N-BEATS model but varied the input *window* between 1, 2 and 5 days whilst retaining a prediction *horizon* of 1 day. The results of our analyses for N-BEATS were inconclusive with the trained models producing comparable results to the Baseline model with differences in performance of up to c.+0.04% (based on a MAPE measure) and hence no material outperformance.

Extending the input *window* to 2 and 5 days respectively across DNN, CNN, GRU and LSTM models, whilst retaining a prediction *horizon* of 1 day, produced comparable results but no improvement in model performance when compared with the Baseline, with differences in performance of up to c.+0.3% (based on a MAPE measure). Hence, the overall results were inconclusive.

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Section 1: Executive summary

We are pleased to publish our second paper as a Working Party using data science techniques to look at sustainability and climate change-related issues. In this paper, we summarise the second stage of our analysis, where we explore more complex data science techniques and architectures to continue our time series analysis of the *Standard & Poor's (S&P) Green Bond Index*.

Scope of this paper

This paper builds on the initial paper published on 1 November 2023 via the IFoA blog site (Dey, 2023) and published on 11 March 2024 via the British Actuarial Journal (Dey, 2024), where we extend the time series univariate analysis to a more complex neural network architecture (please see below). We have deliberately **excluded traditional stationarity techniques such as ARIMA as well as restricted this paper to a univariate analysis**, to help focus on the impact of using a pure neural network design and help with interpretability of the results. We may consider traditional stationarity techniques and expanding our analysis to multivariate in subsequent papers.

For the purposes of this paper, we have focussed the *S&P Green Bond Index* and performed various univariate time series analyses using a range of neural network architectures only. The first part of this paper, Section 3, mainly focusses on using a rolling window approach of one prior day's index value to predict today's index value. The latter part of this paper, Section 4, extends the input window to a longer input period but the output horizon remains as 1 day.

In particular, this paper discusses (arranged as per the following Sections):

- **Section 2: Introduction**
 - Recap on the prior analysis (Dey, 2024), the data used and how we will build on our analysis in this paper.
- **Section 3: N-BEATS**
 - Introduction to N-BEATS (Neural Basis Expansion Analysis for interpretable Time Series forecasting) architecture and extending our time series analysis to incorporate N-BEATS.
- **Section 4: Widening the window**
 - Extending our analysis to widen the input window into the original architectures from the first paper (Dey, 2024).
- **Section 5: Conclusions and next steps**
 - Summary of conclusions from our analysis and potential areas of analysis for subsequent papers.

Please note that many of the techniques mentioned in this paper build on our initial paper (Dey, 2024) and the detail is not duplicated here. For more underlying information on some of the architecture and modelling techniques, please refer to the initial paper (Dey, 2024) – this paper will be referred to “initial paper” throughout – and Appendix 3.

Summary of analysis in this paper

Aim of the analysis

This paper extends on the initial stages of our time series analysis on GSS (Green, Social and Sustainability) bonds (Dey, 2024), specifically focussing on the daily values from the *Standard and Poor's (S&P) Green Bond Index* and whether or not we can create accurate prediction models using neural networks. This paper builds on the initial paper, where we hope to develop a model which can assist with GSS bond index prediction, which will have wider applications such as index price modelling and investment portfolio analyses for actuaries and non-actuaries alike.

For the purposes of this paper, we continue to look at predicting a rolling 1-day value of the index, based on the prior x days index value over the period 2013 to 2023 inclusive.

We retain the same Baseline model i.e. today's value equals yesterday's value over the course of the full date range of 31 January 2013 to 17 February 2023. Similar to the initial paper, we aim to see if we can accurately create a time series model with non-traditional methods such as neural networks and more complex neural network architectures, in particular using N-BEATS. Please see later for further details.

Analyses using traditional stationarity techniques such as ARIMA and multivariate techniques have been deferred to later papers, as previously mentioned.

Towards the end of this paper, we extend our analysis by looking at using the prior x index values (*window*) to predict the next y days in the future (*horizon*), rerunning the initial neural network architectures analysed in our initial paper (Dey, 2024). We restrict the input *window* to 2 and 5 days, whilst retaining a future *horizon* to 1 day in this paper. In Section 3, for N-BEATS, we train the model with an input *window* of 1 day in addition.

Data and method

We have retained the same underlying data as per the first paper (Dey, 2024) - the *S&P Green Bond Index* values between 31 January 2013 to 17 February 2023, splitting the data using 70% / 20% / 10% ratios for training / validation / test data sets.

Section 3 of this paper extends our analysis by introducing a more complex model architecture, the N-BEATS model architecture.

The latter part of this paper, Section 4, revisits the models analysed in our initial paper but extends the input window of data to 2 days and 5 days respectively. These models can be categorised into the following model categories: Deep Neural Network (DNN), Convolutional Neural Network (CNN), Long Short-Term Memory (LSTM) and Gated Recurrent Unit (GRU) neural network architectures. This will be discussed in Section 4 and Appendix 3 in more detail.

The loss function used during training the models was set to Mean Absolute Error (MAE) for all models, with an Adam optimiser to update the weights in order to minimise the loss function as part of the training process and L2 regularisation to reduce any overfitting during the training process. Hyperparameter tuning of all models was completed via the open-source library *Optuna*, using the Bayesian optimisation algorithm Tree-structure Parzen Estimator (TPE). These techniques are discussed in more detail in the initial paper (Dey, 2024).

Results and conclusions

The results of our analysis for N-BEATS (Section 3) were inconclusive: the models from each category produced comparable results to the Baseline model with differences in Mean Absolute Percentage Error (MAPE) of up to c.+0.04% and hence with no material outperformance.

Using similar neural network architectures as per the initial paper (Section 4), if we extend the input *window* to 2 and 5 days respectively, there was no improvement in model performance when compared with the Baseline, with differences in MAPE of up to c.+0.3%. Widening the *window* from 2 days to 5 days did result in an improvement with some models, though again the overall model performance was still worse than the Baseline. Hence, the overall results were inconclusive.

We are potentially not sharing sufficient correlated information e.g. as per a multivariate analysis for our models to learn underlying material information and patterns in the data to result in a model which materially outperforms the Baseline. We aim to address this in future papers (please see below and Section 5 for more details).

Next steps

For future papers, we will expand our analysis to include the following:

1. Expanding the analysis to general GSS bonds. The analysis in this paper is based on a single green bond index. We will look to expand our analysis to the wider GSS bond universe and over differing date ranges for the data to see if there are general underlying conclusions across different data sets and GSS bond indices.
2. Expanding the analysis to include any potential relationships with the general market such as stock market and oil prices i.e. move to a multivariate analysis in subsequent papers. This is examined e.g. in (Wang, et al., 2021) which, when coupled with the CEEMDAN-LSTM model, seems to produce materially improved model predictions based on green bond time series data when compared to a baseline model.
3. Extend our analysis to alternative neural network architecture types, such as Graph Neural Networks (GNNs).

Additional disclaimers

Please note the following:

- a. Information within this paper is **valid up to 31 May 2024**. Hence, there may be updates beyond this date which are not reflected in this paper e.g. changes to any legislation mentioned or updates to any open-source libraries used.
- b. This paper is not intended to be a comprehensive audit of models. Neither is this paper recommending or promoting one approach over another, nor promoting any of the sources or references stated in this paper. Any user of this paper should still reference the underlying legislation, reference any standard mentioned in this paper, and should there be any conflict, the underlying information in the relevant standard, reference or legislation supersedes any information presented in this paper.
- c. Though the work in this paper does not fall under the Financial Reporting Council's Technical Actuarial Standards, this paper has been reviewed both within the Working Party and by the Institute and Faculty of Actuaries' Data Science Practice Board.

Section 2: Introduction

2.1 Recap on prior analysis

In our initial paper (Dey, 2024), we produced a time series analysis of the *S&P Green Bond Index* and aimed to replicate the index using various models over a certain time period. The data in our analysis was from the end of January 2013 to mid-February 2023.

Our initial stages of our analysis focussed on a univariate time series model, where our model used the prior day's value to predict the index value one day ahead (i.e. today's value). The prediction models used were predominantly neural network architectures: deep neural networks (DNNs), Convolutional Neural Networks (CNNs), Long Short-Term Memory models (LSTMs) and Gated Neural Networks (GNNs). Please note that for the purposes of this and the initial paper (Dey, 2024), DNNs included 1 to 3 hidden layers. Strictly, a DNN has 2 or more hidden layers. However, for ease of categorisation, we have extended this labelling to include 1 hidden layer as well. In our initial paper, we further extended the analysis to include the popular library XGBoost, which is a decision-tree model. We have not considered XGBoost further in this paper.

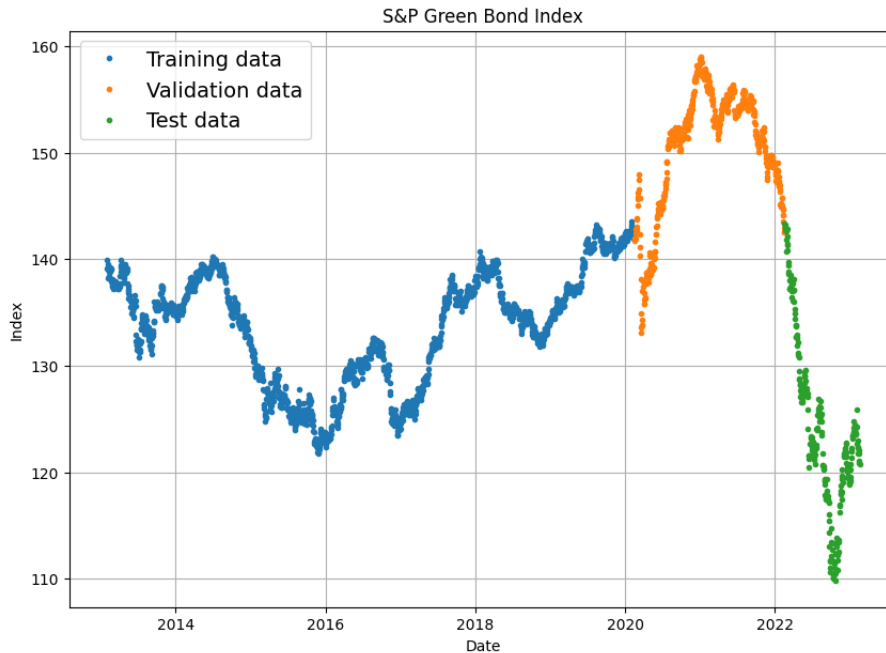
For the purposes of this paper, we will continue with our analysis with an overall aim to produce a sufficiently accurate predictive model, where the models will be based on a neural network architecture design. Data from 31 January 2013 to around mid-February 2022 will be used to train and validate the models. These models will then be used to predict daily index values on unseen data from mid-February 2022 to mid-February 2023 (i.e. the test data set). The difference in predicted values from our models with actual daily index figures will be used to gauge the accuracy of the proposed models.

2.2 Summary of data used

We have continued to use the same data set as per our initial paper (Dey, 2024), downloading via a free subscription-based account from the main S&P website <https://www.spglobal.com/>. For details of the data set and underlying testing of this data, please see our initial paper (Dey, 2024).

For the purposes of this paper, we will analyse the *S&P Green Bond Index* (Total Performance, USD, from 31 January 2013 to 17 February 2023 inclusive). The splits are as per the initial paper and further shown in Figure 1 and Table 1. Given the nature of the analysis, i.e. a time series analysis, we have ordered the data and these splits in chronological order, so that we can build models to infer some form of prior time-dependency based on the underlying data, and have not randomly allocated the data between these splits across the full data range.

Figure 1: S&P Green Bond Index data with training / validation / test splits highlighted (Output from Google Colab).



The splits equate to as follows: to 16 February 2020, to 16 February 2022 and to 17 February 2023 inclusive. Table 1 details further each data split.

Table 1: Summary table of data used, split by training, validation and test data sets.

	Full data	Training data	Validation data	Test data
<i>Start date</i>	31 Jan 2013	31 Jan 2013	17 Feb 2020	17 Feb 2022
<i>End date</i>	17 Feb 2023	16 Feb 2020	16 Feb 2022	17 Feb 2023
<i>Number of index entries</i>	2,615	1,830	523	262
<i>Index minimum</i>	109.80	121.78	133.14	109.80
<i>Index maximum</i>	158.99	143.59	158.99	143.34
<i>Index average (2 d.p.)</i>	136.00	133.60	150.46	123.96
<i>Index standard deviation (2 d.p.)</i>	9.65	5.32	5.70	8.04

In summary, there is greater volatility in the test data set range when compared to the training and validation data sets. Hence, it will be interesting to see how our models cope given that they will be built on less volatile training and validation data.

Similar to the initial paper, for the purposes of our analysis, we have not adjusted the data further i.e. no normalisation of the index (setting to a scale of 0 to 1, a technique typically used to result in a quicker convergence to a solution for a model) and no log transformation (which can be used to potentially dampen any impact of seasonality). Such techniques may be discussed in later papers.

Section 3: N-BEATS

3.1 Introduction

We start our analysis with the N-BEATS (Neural Basis Expansion Analysis for interpretable Time Series forecasting) architecture. In this Section, we give an overview of the model architecture, before explaining in more detail the generalised model architecture. We then explain how we trained this model architecture specifically on our data set and finally present our results from this analysis.

Originally submitted in 2019, with the publication revised in 2020, the N-BEATS time series model was specifically designed to tackle time series problems. We will refer to this publication (Oreshkin, et al., 2019) as the “source paper” throughout this Section. The source paper states that N-BEATS demonstrates *state-of-the art performance, improving forecast accuracy by 11% over a statistical benchmark and by 3% over last year’s winner of the M4 competition* (Oreshkin, et al., 2019). The paper considers tackling a univariate discrete time series problem. Given this, using the model architecture for our forecasting problem seems appropriate.

The source paper discusses how deep learning techniques have struggled to consistently outperform more traditional statistical techniques: *the rankings of the six “pure” ML methods submitted to M4 competition were 23, 37, 38, 48, 54, and 57 out of a total of 60 entries* (Oreshkin, et al., 2019). Please note though that we do not explore more traditional techniques in our paper and continue to focus on non-traditional techniques i.e. neural networks.

The Makridakis Competitions is an open time series competition (Makridakis, et al., 2020). The fourth iteration of the competition, M4 which is mentioned above, was held in 2020.

Beyond designing a model architecture which uses deep learning to tackle a time series problem, the aim of the source paper was to also *design an architecture with interpretable outputs that can be used by practitioners in very much the same way as traditional decomposition techniques such as the “seasonality-trend-level” approach* (Oreshkin, et al., 2019).

Below are some key features of the N-BEATS model architecture:

1. It is a *deep neural architecture based on backward and forward residual links and a very deep stack of fully connected layers* (Oreshkin, et al., 2019). The model architecture is made up of stacks which are in turn made up of blocks. Each block produces a *backcast residual* and *partial forecast* with the aim of iteratively improving the prior input value and augment the future value respectively within the forecasting process. For more details of the underlying architecture, please see Section 3.2.
2. One of the aims of an N-BEATS model architecture is to ensure underlying patterns in the data are *interpretable* (i.e. understandable) for users, where the model can infer an underlying trend and seasonality pattern within any given time series data set.

For a high-level introductory summary of N-BEATS, please see for example an article published on the *Towards Data Science* website (Dancker, 2024).

3.2 Deeper dive into generalised N-BEATS model architecture

Below we look at the generalised N-BEATS model architecture in more detail. We summarise here some of the key points raised from the source paper (Oreshkin, et al., 2019) and use the same notation as per the source paper when presenting formulae.

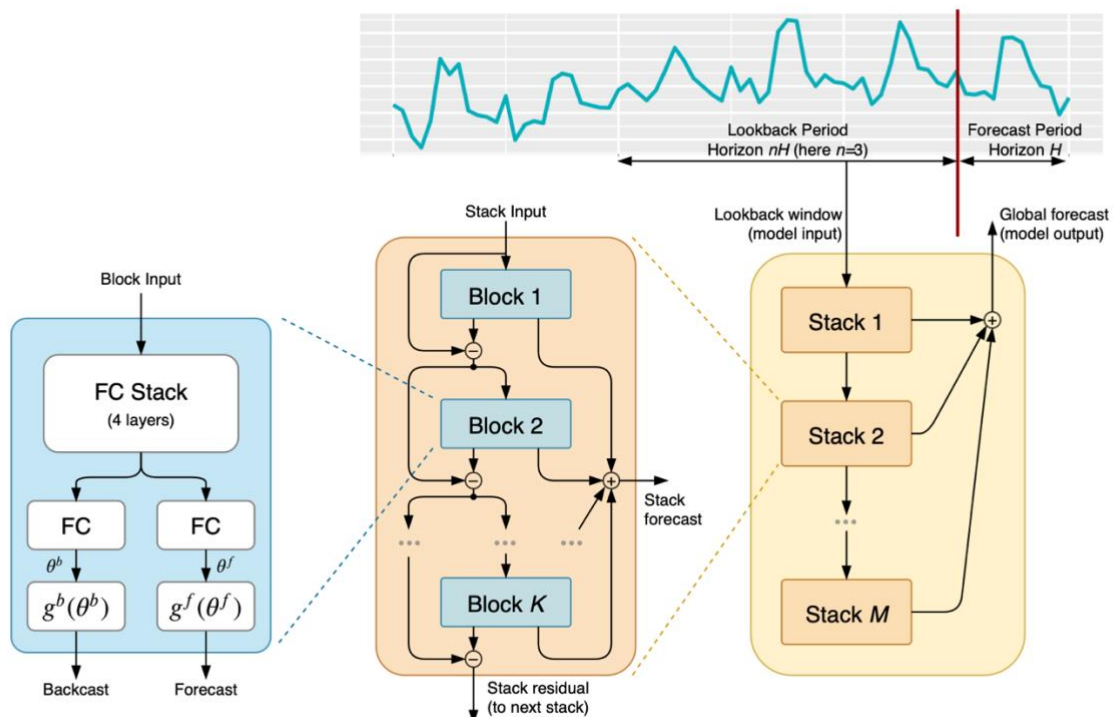
The generalised model architecture is based on the following principles (Oreshkin, et al., 2019):

1. It should be *simple and generic, yet expressive (deep)*.
2. It should not *rely on time series-specific feature engineering or input scaling*.
3. The outputs should be *extendable towards making its outputs human interpretable*.

The first two allow a user to set up a time series forecasting model based on a pure deep learning architecture.

Figure 2 shows a generalised overview of the architecture design underlying N-BEATS model architecture. The figure is taken directly from the source paper.

Figure 2: Generalised N-BEATS architecture - Source (Oreshkin, et al., 2019).



We summarise below an overview of the N-BEATS architecture:

1. *Basic building blocks* consist of hidden layers each of several neurons, which are fully connected. The outputs from this structure can be further extended to incorporate *basis functions* which allow for time series features such as seasonality and trends. The *basic building block* is represented by the blue box on the left in Figure 2, which illustrates 4 hidden layers.
2. These *basic building blocks* in turn form a *stack*, which take the residual *backcasts* from the prior block and cumulate the partial trend *forecasts* outputs. This is represented by the middle box in Figure 2, which illustrates K *basic building blocks*.
3. In a similar fashion, the residual *backcasts* from the prior stack feed into later *stacks*, and then cumulate the partial trend *forecasts* to an overall *global forecast* i.e. the overall *model output*. This is represented by the box towards the right in Figure 2 earlier, which illustrates M stacks in total.

The above technique of *residuals* and partial *forecasts* form part of *doubly residual stacking*. This is discussed further in Section 3.2.2 below.

3.2.1 Basic building block

The *basic building blocks* generate a *backcast* and *forecast* as previously mentioned. The *backcast* can be viewed as iterative adjustments to the input vectors via *residuals*, whilst the *forecast* from each *basic building block* can be viewed as a partial *forecast* which is cumulated and form part of the overall *global forecast*.

Focussing on the l -th *basic building block*, as per the source paper (Oreshkin, et al., 2019): each *basic building block* has a *fork architecture* which takes an input vector (or *residual* outputs from a prior *basic building block*) x_l and produces two outputs: \hat{x}_l i.e. the *block's best estimate of x_l* and the *block's forward forecast \hat{y}_l* . The input of the very first block in the model is the original data input. The remaining *basic building block* inputs are residual outputs from a prior *basic building block*, and *can be thought of as running a sequential analysis of the input signal* (Oreshkin, et al., 2019). See Section 3.2.2 for more details.

The internal structure of a *basic building block* broadly consists of two sections: i. a fully connected section; and ii. a basis layer section.

The fully connected layer produces backward (*backcast*) and forward (*forecast*) predictors, θ_l^b and θ_l^f respectively, of *expansion coefficients*. The predictors determine how much each *basis function* contributes to the overall approximation of the function. Please see Section 3.2.1(b) for more details on *basis function*.

The above two sections of the internal structure of a *basic building block* are discussed in more detail immediately below in Sections 3.2.1(a) and 3.2.1(b).

3.2.1(a) Fully connected section

Using the same formulae notation and architecture discussed in the source paper (Oreshkin, et al., 2019) which has four hidden layers, the equations within the l -th *basic building block* are as follows:

- $\mathbf{h}_{l,1} = FC_{l,1}(\mathbf{x}_l)$, $\mathbf{h}_{l,2} = FC_{l,2}(\mathbf{h}_{l,1})$, $\mathbf{h}_{l,3} = FC_{l,3}(\mathbf{h}_{l,2})$, $\mathbf{h}_{l,4} = FC_{l,4}(\mathbf{h}_{l,3})$, where $\mathbf{h}_{l,i}$ represents the i -th hidden layer in the l -th block, $FC_{l,i}$ represents the i -th fully connected layer in the l -th block, and \mathbf{x}_l is the input vector for the l -th block.
- $\theta_l^b = LINEAR_l^b(\mathbf{h}_{l,4})$ and $\theta_l^f = LINEAR_l^f(\mathbf{h}_{l,4})$, where $\mathbf{h}_{l,i}$, θ_l^b and θ_l^f are as per earlier. $LINEAR$ is a *linear projection layer* i.e. $\theta_l^b = W_l^b(\mathbf{h}_{l,4})$ and $\theta_l^f = W_l^f(\mathbf{h}_{l,4})$ with the subscript l representing the l -th block, the superscripts b and f representing *backcast* and *forecast* respectively, and W is a weight matrix active on a hidden layer.

For the purposes of the source paper, FC is a standard fully connected layer with a *relu activation function* i.e. $\mathbf{h}_{l,1} = RELU(W_{l,1}\mathbf{x}_l + \mathbf{b}_{l,1})$, where $W_{l,1}$ represents the weight vector connecting the input \mathbf{x}_l to the first hidden layer in the l -th block and $\mathbf{b}_{l,1}$ is the bias term for the first hidden layer in the l -th block.

3.2.1(b) Basis layer section

The basis layer maps the *backcast and forecast expansion coefficients* θ_l^b and θ_l^f to outputs, where the partial output $\hat{\mathbf{y}}_l = g_l^f(\theta_l^f)$ and the residual output $\hat{\mathbf{x}}_l = g_l^b(\theta_l^b)$. The *basis functions* g_l^b and g_l^f are for *backcast* and *forecast* respectively.

More generally, *expansion coefficients* are parameters which are learnt during the training process. When combined with *basis functions*, they help capture any underlying patterns or features in a time series data set such as a trend, an anomaly or seasonal patterns. Examples of *basis functions* include polynomial, Fourier, wavelet and Gaussian functions.

3.2.2 Doubly residual stacking

A unique feature of the N-BEATS model architecture is the concept of *doubly residual stacking*. Instead of a single residual branch of outputs, the architecture proposes two branches as mentioned: one for a *backcast residual* and one for a *partial forecast*. Mathematically, we represent this as:

$$\mathbf{x}_l = \mathbf{x}_{l-1} - \hat{\mathbf{x}}_{l-1}, \hat{\mathbf{y}} = \sum_l \hat{\mathbf{y}}_l,$$

where \mathbf{x}_l and $\hat{\mathbf{x}}_{l-1}$ are defined earlier in Section 3.2.1, $\hat{\mathbf{y}}$ is the *global forecast* i.e. overall model output which is the sum of individual $\hat{\mathbf{y}}_l$. Each $\hat{\mathbf{y}}_l$ can be viewed as a *partial forecast*.

3.2.3 Interpretability

There are 2 types of *configurations* or design patterns for an N-BEATS architecture:

1. *Generic architecture*, which does not assume any specific underlying time series model and relies on a generalised deep learning approach.
2. *Interpretable architecture*, which is an augmented model that often includes a *trend* and *seasonality model* to the generalised N-BEATS architecture discussed earlier for a time series forecasting problem.

For the purposes of this paper, we have focussed on the first configuration i.e. *generic architecture* as the aim of this paper is to seek a model architecture which accurately predicts the time series future values without any further input by the modeller or prior knowledge of any underlying time series features from the modeller.

For more details on the above, and other considerations, please see the source paper (Oreshkin, et al., 2019).

3.2.4 Comparison with other model architectures

In Table 2, we provide a high-level comparison of the N-BEATS architecture versus DNNs, and LSTMs / GRUs. Please note that these are generalised comments based on experience and the comparison table below may differ for different data sets, problems and scenarios.

Table 2: Comparison of N-BEATS model architecture.

Model architecture	DNN	LSTM, GRU	N-BEATS
Neural network type	Feedforward	Recurrent	Feedforward
Handles long-term memory	No	Yes	Yes – via <i>backcasting</i> and <i>forecasting</i>
Complexity	Low	Medium	High
Computational resources	Low	Medium	High
Indicative time taken to train model in <i>Google Colab</i> based on data set and methodology discussed in this paper (<i>epochs</i> and <i>Optuna</i> trials may vary)	Up to c.15 mins	Up to c.30 mins	c.3 to 4 hours
Performance	Varies though better on short sequences	Varies though better on long sequences	Excellent for time series
Interpretability	Low	Medium	High
Typical usage	Wide range of problems	Sequential data e.g. Sentiment analysis, Language modelling, Speech recognition	Time series forecasting
Practical considerations	Easy to implement	Better for tasks requiring understanding of long-term dependencies	Highly effective for complex time series forecasting tasks where interpretability and performance are crucial

3.3 Model approach used and training the model

Below we give a brief overview of our code implementation of the N-BEATS model.

3.3.1 Code background

The underlying basis of our code for the N-BEATS model is taken from *Zero to Mastery TensorFlow for Deep Learning Book* (Bourke, 2023) and has been adapted for our analysis, where we look at a general N-BEATS architecture. We vary the number of layers within each block and the number of blocks per stack but set the total number of stacks equal to 1 (i.e. M is 1 in Figure 2). This is in part to simplify the trained model architecture given that the time taken to train such a model was close to 4 hours. Further, in doing so, we do not believe that taking such an approach would materially impact any overall conclusion, as effectively adding another stack would repeat a similar iterative fine-tuning process and potentially the hyperparameter optimisation could compensate by increasing the number of blocks in the single stack.

The original authors of the N-BEATS model also extend their analysis to allow for an ensemble technique (using multiple different loss functions and multiple different lookback periods in Figure 2) to make predictions when testing on M4 dataset. To retain consistency with the approach taken with other models within our initial paper, we have not extended our analysis to allow for such ensemble techniques.

We have assumed the same underlying architecture per block within each stack e.g. the same number of hidden layers within each block. Similarly, we have assumed the same underlying architecture for each hidden layer within each block e.g. number of neurons, activation function and L2 loss regularisation parameter. The output layer from the block is a generalised dense layer, each with the same number of neurons in effect as the total combined length of the input *window* and output *horizon*. Similarly, we have assumed that all output layers have the same activation function, though this may differ to the hidden layers above.

Similar to our initial paper, we have used the same underlying data and training / validation / test splits within the data to train our model. We have run our code in *Google Colab*, using similar libraries including *Tensorflow* and *Keras* as per our initial paper. For hyperparameter optimisation, we have again used *Optuna*. For an in-depth discussion on this, please see our initial paper (Dey, 2024).

3.3.2 Hyperparameter optimisation

As mentioned, we have used *Optuna* for hyperparameter optimisation. Below we list some key search spaces used within our hyperparameter optimisation.

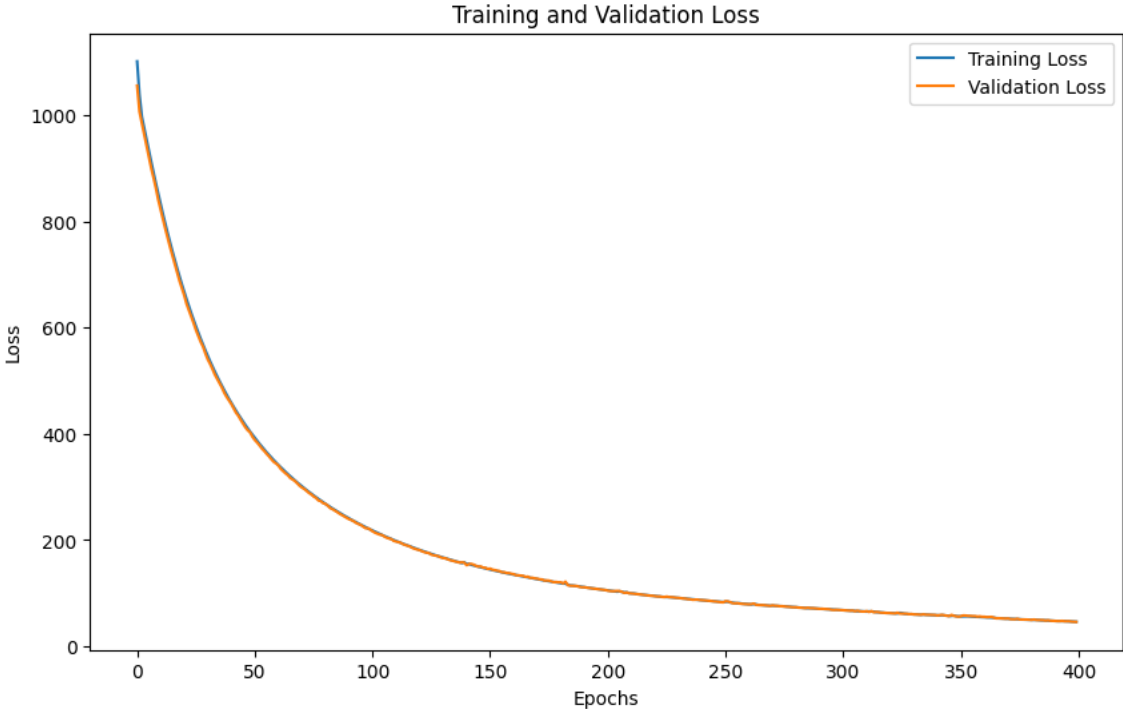
1. The number of epochs was set to 400.
2. The number of trials within *Optuna* was set to 30.
3. The batch size for the input data was set to 128.
4. The *activation list* was as follows: *elu*, *gelu*, *linear*, *relu*, *sigmoid*, *swish* and *tanh*.

5. The specific search space for each N-BEATS block includes:
 - a. Number of neurons in each hidden layer from 4 to 512 at steps of 4.
 - b. Number of hidden layers from 1 to 4 at steps of 1.
 - i. As mentioned earlier, we have assumed all hidden layers have the same underlying architecture e.g. the same number of neurons, same activation function and same L2 regularisation.
 - c. L2 regularisation with a search space of $1e-5$ to $1e-1$.
 - d. Output *theta layer* or *basis layer* is a generalised deep layer.
 - i. The number of neurons is set equal to the *theta size* – in effect the combined length is the *window* size and *horizon* size for this analysis.
 - ii. The *activation function* from the *activation list* above. No L2 regularisation function was applied to this layer.
 - iii. No further adjustments were made for seasonality and trend i.e. via *basis functions*, as previously mentioned.
6. Specific search space for number of N-BEATS blocks per stack from 2 to 30 at steps of 2.
7. Similar to the source paper, the model was compiled on a loss function of Mean Absolute Error (MAE) using an Adam optimiser with a learning rate search space of $1e-5$ to $1e-1$.

3.3.3 Training the model

Below is a loss history curve based on training one of the models, using the search space, number of epochs and number of trials in *Optuna* as mentioned in Section 3.3.2.

Figure 3: Training and validation loss curves (Output from Google Colab).

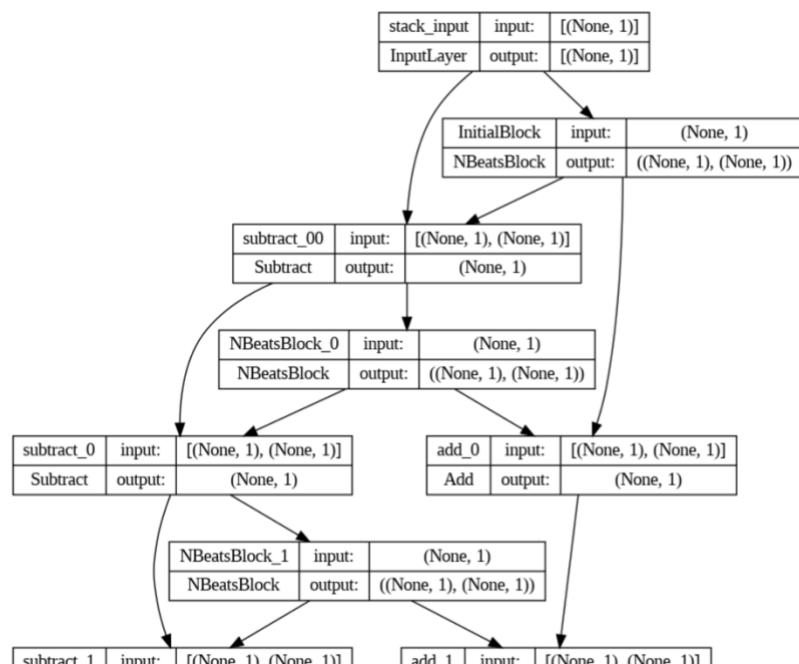


As we would expect, the model loss decreases as the number of epochs increases, suggesting that the model is able to converge to a solution.

Similarly, we have used the open source *Keras visualizer* library to demonstrate the underlying architecture of the trained model in Figure 4. Please note that we have taken a snippet of the first few blocks and initial stacks, given the size of the overall final trained model.

The visualisation tool in general gives an alternative simplified representation of any final neural network model architecture. Further, given that we have generalised our code and set the overall number of stacks equal to 1 as mentioned earlier, there are cosmetic differences in Figure 4 when compared with Figure 2.

Figure 4: Snippet of the final trained N-BEATS model architecture using *Keras visualizer* (Output from *Google Colab*).



In Figure 4, the Keras visualisation tool helps visualise *basic building blocks* (represented by “NBeatsBlock”) showing:

1. subtraction (represented by “Subtract” in Figure 4) to represent the process of *residuals* or *backcasts*; and
2. additions (represented by “Add” in Figure 4) to represent the process of *partial forecasts*.

The final trained model (with an input window of 1 day) has over 4 million trainable parameters (i.e. weights and biases). As a comparison, Model 1 from Appendix 3 in (Dey, 2024) which is a DNN model, has 625 trainable parameters. Similarly, Model 6 from Appendix 3 in (Dey, 2024), which is an LSTM model, has 47,629 trainable parameters.

3.3.4 Model outputs and observations

The source paper (Oreshkin, et al., 2019) mentions that the input length of the *window* is typically set to a multiple of the length of the output *horizon* H, between 2H and 7H. For the purposes of this paper, we have trained the model setting this input length to a multiple of one (*1 times*) to align with our initial paper (Dey, 2024), two (*2 times*) and five (*5 times*) - in order to adhere to the recommendation above and in the source paper (Oreshkin, et al., 2019), as well as our analysis in Section 4 of this paper. The output *horizon* is as per the initial paper i.e. 1 day.

In Table 3, we compare the performance of the trained N-BEATS models, which have been trained on the same data but on input *windows* of 1, 2 and 5 days, and with the same output *horizon* of 1 day.

Table 3: Comparison of performance measure, between the Baseline model and N-BEATS, based on MAE and MAPE, varying the input *window* between 1, 2 and 5 days.

Performance measure	Baseline (<i>Window</i> = 1 day)	N-BEATS (<i>Window</i> = 1 day)	N-BEATS (<i>Window</i> = 2 days)	N-BEATS (<i>Window</i> = 5 days)
<i>MAE</i> (3 d.p.)	0.610	0.620	0.662	0.657
<i>Difference from Baseline MAE</i> (3 d.p.)	-	+0.010	+0.052	+0.047
<i>MAPE</i> (3 d.p.)	0.497%	0.505%	0.539%	0.536%
<i>Difference from Baseline MAPE</i> (3 d.p.)	-	+0.008%	+0.042%	+0.039%

Though the N-BEATS models performed relatively accurately, based on the data set, split of data between training / validation / test, and hyperparameter optimisation mentioned in Section 3.3.2, there is no improvement to the Baseline for each of our N-BEATS models based on either MAE and MAPE measures on the test data set. The model trained on a 5-day input *window* marginally performs better than the model trained on a 2-day input *window*. However, the model trained on a 1-day input *window* outperforms the N-BEATS models trained on a 2-day and 5-day input *window*. All trained models perform marginally worse than the Baseline model. Hence, the overall results are inconclusive.

One potential area of future study is to incorporate ensemble techniques as per the original authors of N-BEATS as mentioned in Section 3.3.1, as well as allow for the number of stacks to vary during hyperparameter optimisation when training the model.

Section 4: Widening the window

4.1 Introduction

We now change the focus of our analysis and widen the input *window* of data going into our models when training, validating and testing with the aim of hopefully improving the model accuracy, as we are looking to provide more information to the models to recognise any underlying potential patterns and dependencies within the data. Please note that we revisit the majority of the original neural network architectures discussed in our initial paper (Dey, 2024), though have excluded *return sequence is true* models as we are predicting 1 day into the future in this paper. We have also excluded XGBoost in our analysis in this Section as previously mentioned, given that this is a decision-tree library and not based on any underlying neural network architecture.

For this Section, we have used various chart functions from the Python-based, open source *statsmodels* library. For more details on this library, please see here: <https://www.statsmodels.org/>.

4.2 Autocorrelation (ACF) and Partial autocorrelation (PACF) plots

We have used the *seasonal_decompose* function from the *statsmodels* library to separate potential trend lines, seasonality and residual plots across the full data set discussed in Section 2. These can be viewed further in Figure 5. We have analysed the daily log returns of the underlying data set to infer potential lags in the data and correlations, and hence help determine alternative *window* sizes.

Figure 5: Seasonal, trend and residual plot outputs from the data set (Output from Google Colab).



Within Figure 5, “index” represents the daily log returns of the data set, “trend” represents any underlying trend, “seasonal” represents any potential seasonal element and “resid” represents any unaccountable residual (after the trend and seasonal components have been accounted for).

Using the built-in `plot_acf` and `plot_pacf` functions from the `statsmodels` library, we have created an autocorrelation (ACF) plot in Figure 6 and partial autocorrelation plot in Figure 7 respectively.

Figure 6: Autocorrelation plot for data set mentioned in Section 2 (Output from Google Colab).

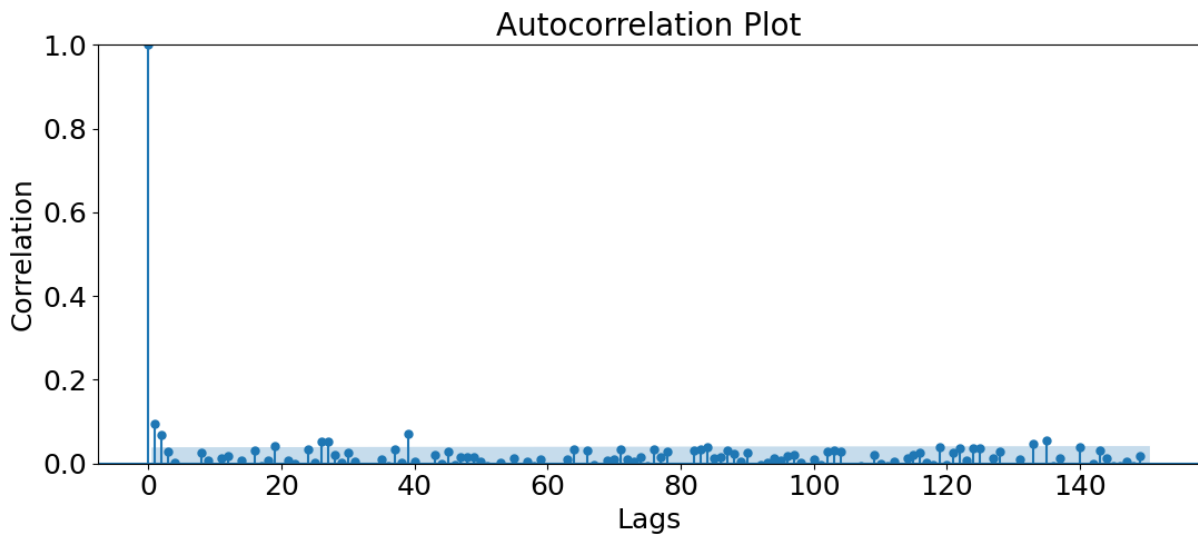
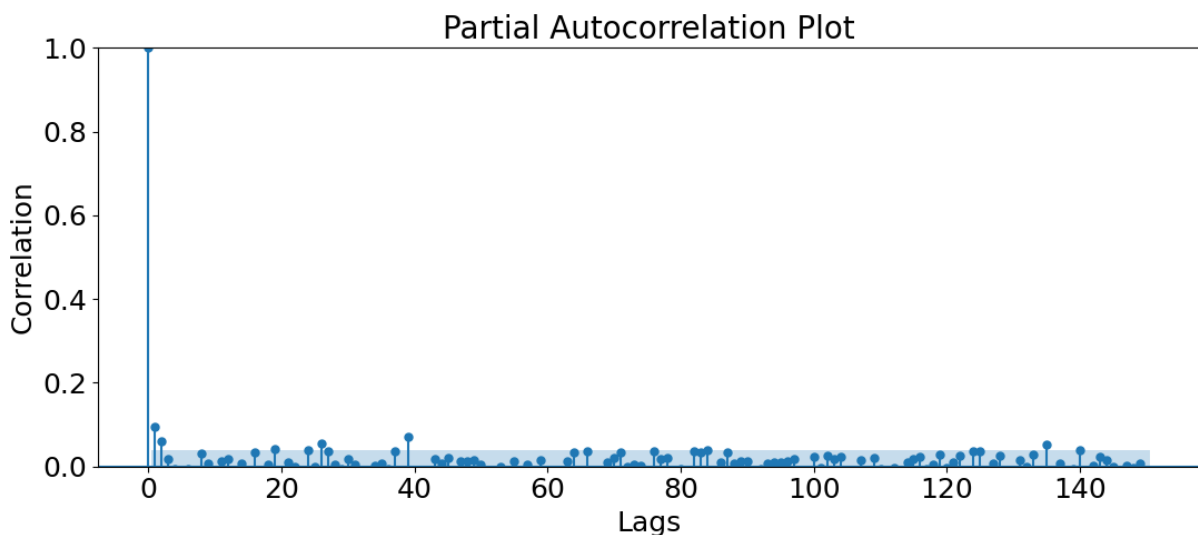


Figure 7: Partial autocorrelation plot for data set mentioned in Section 2 (Output from Google Colab).



Both plots can be used to infer a correlation between different lags. For more general background to ACF and PACF plots, please see for example (Monigatti, 2022).

Though there is a certain amount of subjectivity involved in terms of interpreting Figures 6 and 7, we could expect a 2-day historic lag in data to be representative of any potential underlying feature.

4.3 Window length to choose

Based on the analysis in Section 4.2, the implied input *window* is 2 days. Similarly, the window lag implied by (Peters, et al., 2022) is 5 days. Strictly speaking, these lags are based on the daily log returns. Reversing the log transformation implies a lag of c.7 working days and c.148 working days respectively. The latter time period of 148 days is equivalent to around 6 to 7 months, and may indicate that the prior 2 quarters of annual data incorporate longer term market trend information, and hence may be useful when producing better future predictions.

Rerunning all models within this paper based on 7- and a sample on 148-day input *windows* resulted in worse performing models, using the same underlying conditions and predicting 1 day ahead, than if the input *window* was restricted to 5 days. Hence, we have retained input *windows* of 2 and 5 days within this Section and paper, and not extended beyond 5 days. Given that the data is per working day, intuitively 5 days may seem more appropriate. The results from model outputs based on an input *window* of 7 and 148 days have not been included in this paper given the above.

4.4 Training the models

We have retrained the neural network models from our initial paper (Dey, 2024) based on a 2-day input *window* and 5-day input *window*. The approach taken is as per the initial paper.

4.5 Results from analysis of 2-day and 5-day input window

We have summarised the MAE and MAPE results from different model runs for some of the model architectures originally analysed in our initial paper (Dey, 2024). We have maintained the same labelling e.g. model number and abbreviated name, as per the initial paper (Dey, 2024). For a full list of these models, please see Appendix 3.

Table 4 compares the performance based on MAE and MAPE for these different models, based on input windows of 1, 2 and 5 days.

Table 4: Comparison performance of neural network models with 1, 2 and 5 days of input window information.

Model number	Abbreviated name	Category	Data input					
			Window = 1 day		Window = 2 days		Window = 5 days	
			MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)
0	Baseline	Baseline	0.610	0.497%	-	-	-	-
1	DNN0	DNN	0.607	0.495%	0.739	0.602%	0.933	0.763%
2	DNN1		0.627	0.511%	0.706	0.576%	0.948	0.773%
3	DNN2		0.620	0.505%	1.320	1.085%	0.798	0.651%
4	CNN0	CNN	0.619	0.504%	0.686	0.560%	0.992	0.811%
5	CNN1		0.611	0.498%	0.705	0.575%	0.959	0.783%
6	LSTM_OHL_F	LSTM	0.609	0.497%	0.769	0.627%	0.719	0.585%
8	LSTM_1HL_F		0.617	0.503%	0.655	0.534%	0.667	0.545%
10	GRU_OHL_F	GRU	0.632	0.514%	0.680	0.554%	0.647	0.528%
12	GRU_1HL_F		0.638	0.519%	0.623	0.507%	0.643	0.524%

We have retained the original Baseline which has been trained on an input *window* of 1 day across all comparison scenarios, given the existing level of accuracy / relative difficulty for other neural networks architecture models to materially outperform this. As can be seen in Table 4, generally the GRU and LSTM models outperform CNN and DNN models, based on both a 2- and 5-day input *window*.

Table 5 repeats similar information, but compares the relative performance of a 2-day and 5-day input *window* against a 1-day input *window* for the same model i.e. the comparison is within the same row.

Table 5: Comparison performance 2-day and 5-day input *window* against *window* of 1 day for the same model.

Model number	Abbreviated name	Category	Data input					
			Window = 1 day		Window = 2 days		Window = 5 days	
			MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)
0	Baseline	Baseline	0.610	0.497%	-	-	-	-
1	DNN0	DNN	0.607	0.495%	+0.132	+0.107%	+0.326	+0.268%
2	DNN1		0.627	0.511%	+0.079	+0.065%	+0.321	+0.262%
3	DNN2		0.620	0.505%	+0.700	+0.580%	+0.178	+0.146%
4	CNN0	CNN	0.619	0.504%	+0.067	+0.056%	+0.373	+0.307%
5	CNN1		0.611	0.498%	+0.094	+0.077%	+0.348	+0.285%
6	LSTM_0HL_F	LSTM	0.609	0.497%	+0.160	+0.130%	+0.110	+0.088%
8	LSTM_1HL_F		0.617	0.503%	+0.038	+0.031%	+0.050	+0.042%
10	GRU_0HL_F	GRU	0.632	0.514%	+0.048	+0.040%	+0.015	+0.014%
12	GRU_1HL_F		0.638	0.519%	-0.015	-0.012%	+0.005	+0.005%

The pink colour boxes in Table 5 indicate that the model has performed worse than the corresponding model trained on a *window* of 1-day input. The yellow colour boxes in Table 5 indicate that the model has performed worse than the corresponding same model architecture trained on a *window* of 1-day input, but better than the corresponding model architecture trained on a *window* of 2-day input. The green colour boxes in Table 5 indicate that the model has performed better than the corresponding same model architecture trained on a *window* of 1-day input.

As can be seen in Table 5, model 12 (a GRU model with 1 hidden layer) outperforms with a 2-day input *window* compared to 1-day input *window*. The remaining models perform worse if the input *window* is expanded from 1 day to 2 or 5 days.

Models 3, 6 and 10 performed better with a 5-day input *window* versus a 2-day input *window*. However, as per earlier, the differences are relatively similar and hence the overall results are inconclusive.

Table 6 repeats similar information, but compares the relative performance of increasing the data input window to 2 and 5 days respectively, against the Baseline across all scenarios (which was trained with an input *window* of 1 day).

Table 6: Comparison performance of each neural network model against Baseline.

Model number	Abbreviated name	Category	Data input					
			Window = 1 day		Window = 2 days		Window = 5 days	
			MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)	MAE (3 d.p.)	MAPE (3 d.p.)
0	Baseline	Baseline	0.610	0.497%	-	-	-	-
1	DNN0	DNN	0.607	0.495%	+0.129	+0.105%	+0.323	+0.266%
2	DNN1		0.627	0.511%	+0.096	+0.079%	+0.338	+0.276%
3	DNN2		0.620	0.505%	+0.710	+0.588%	+0.188	+0.154%
4	CNN0	CNN	0.619	0.504%	+0.076	+0.063%	+0.382	+0.314%
5	CNN1		0.611	0.498%	+0.095	+0.078%	+0.349	+0.286%
6	LSTM_0HL_F	LSTM	0.609	0.497%	+0.159	+0.130%	+0.109	+0.088%
8	LSTM_1HL_F		0.617	0.503%	+0.045	+0.037%	+0.057	+0.048%
10	GRU_0HL_F	GRU	0.632	0.514%	+0.070	+0.057%	+0.037	+0.031%
12	GRU_1HL_F		0.638	0.519%	+0.013	+0.010%	+0.033	+0.027%

The colour coding within Table 6 is as per Table 5, but comparing relative performance against the Baseline performance across all scenarios.

Models 3, 6 and 10 performed better with a 5-day input *window* versus a 2-day input *window*. However, as per earlier, the differences are relatively similar with all models performing worse than the Baseline. Hence the overall results are inconclusive.

4.6 Conclusions

By increasing the input *window* from 1 day to 2 and 5 days, on the whole, there were no material improvements to the results. The differences in model performance (based on a MAPE measure) are relatively small and hence the overall results are inconclusive.

The performance of increasing from 1 to 2 or 5 days did however produce relatively accurate models, with a MAPE of c.0.5% to c.1.1% across all models which were investigated.

Section 5: Conclusions and next steps

5.1 Conclusions

Based on our analysis, given the data range and training / validation / test splits of the *S&P Green Bond Index* and modelling approach discussed earlier, we can draw the following conclusions below.

5.1.1 N-BEATS

As observed, we trained a model based on the N-BEATS architecture in Section 3. The outputs of each model (trained on an input *window* of 1, 2 and 5 days) did produce output forecasts for a 1-day horizon (i.e. the next day) which were relatively accurate, with the MAPE differing by c.+0.5% over the test range data set.

However, the trained models did not outperform the Baseline model over the test data set. The trained N-BEATS models differed from the Baseline results by up to c.+0.04% i.e. our findings are inconclusive.

This may be indicative of the fact that the current Baseline is actually a good predictor in the first place, given that the relative low coefficient of variation of c.7.1% for the index across the full data set. Similarly, it may be indicative of the fact that the problem we are facing with the *S&P Green Bond Index* is as per any other stock price movement i.e. a random walk, where it is difficult to continually outperform *any* baseline model (Bourke, 2023).

5.1.2 2-day lag and 5-day lag

Expanding the input window from 1 day to 2 and 5 days respectively for the neural network architectures explored in the initial paper (Dey, 2024) did not result in an improvement with the performance relatively similar to the Baseline i.e. our findings are still inconclusive.

5.2 Next steps

The analysis to date has been based on a univariate analysis and simpler neural network architectures – such as DNNs, CNNs, LSTMs and GRUs, as discussed in the initial paper (Dey, 2024). We have further expanded this to a complex and cutting-edge architecture with N-BEATS.

For future papers, other potential areas we may explore include:

1. Explore other complex neural network architectures based on a univariate analysis e.g. Google's TFT (Temporal Fusion Transformer) and temporal GNNs (Graph Neural Networks).
2. Expanding our analysis to a multivariate analysis.
3. Broaden the projection *horizon* from 1 day to 1 week or possibly 1 month into the future.

Appendix 1: Working Party members and acknowledgements

Working Party members

The Working Party is made up of two sub-groups: i. practitioners; and ii. academics.

Below is a list of current Working Party members for the Practitioner Group, along with details of their position within the Working Party and corresponding LinkedIn link.

Name	Position	LinkedIn
Debashish Dey	Chair	Link
Cem Öztürk	Member	Link
Shubham Mehta	Member	Link

Please note that this Working Party sits within the Lifelong Learning pillar of the IFoA. For further details of the Data Science section of the IFoA, please see <https://actuaries.org.uk/learn/lifelong-learning>.

Acknowledgements

Please see below acknowledgements for the review of the code and paper:

- Code review by Cem Öztürk [[Link](#)].
- Technical review of paper by:
 - Dr Alexey Mashechkin [[Link](#)].
 - Assoc. Prof. George Tzougas [[Link](#)].
- Review of graphs and tables within the paper by Shubham Mehta [[Link](#)].

Appendix 2: List of abbreviations

Below is a list of abbreviations used within this paper.

Abbreviation	Explanation
ARIMA	AutoRegressive Integrated Moving Average
CEEMDAN	Complete Ensemble Empirical Mode Decomposition with Adaptive Noise
CNN	Convolutional Neural Network
DNN	Deep Neural Network
GNN	Graph Neural Network
GRU	Gated Recurrent Unit
GSS bonds	Green, Social and Sustainability bonds
IFoA	Institute and Faculty of Actuaries
LSTM	Long Short-Term Memory
MAE	Mean Absolute Error
MAPE	Mean Absolute Percentage Error
MSCI	Morgan Stanley Capital International
N-BEATS	Neural Basis Expansion Analysis for interpretable Time Series
RNN	Recurrent Neural Network
S&P	Standard and Poor's
TFT	Temporal Fusion Transformer
TPE	Tree-structure Parzen Estimator

Appendix 3: Summary of models analysed in the initial paper

Below is a summary of the models analysed in the initial paper (Dey, 2024), with a brief description of the underlying model architecture.

Model	Abbreviated name	Category	Description of architecture
0	Baseline	Baseline	Baseline model which assumes today's value is the same as yesterday's value.
1	DNN0	DNN	Feedforward artificial neural network with one hidden dense layer.
2	DNN1		Feedforward artificial neural network with two hidden dense layers.
3	DNN2		Feedforward artificial neural network with three hidden dense layers.
4	CNN0	CNN	Convolutional neural network, with one Conv1D layer and no additional hidden layers.
5	CNN1		Convolutional neural network, with one Conv1D layer and one hidden dense layer.
6	LSTM_0HL_F	LSTM	LSTM neural network, with one LSTM layer, return sequence set to false, and no additional hidden layers.
7	LSTM_0HL_T		LSTM neural network, with one LSTM layer, return sequence set to true, and no additional hidden layers.
8	LSTM_1HL_F		LSTM neural network, with one LSTM layer, return sequence set to false, and one additional hidden dense layer.
9	LSTM_1HL_T		LSTM neural network, with one LSTM layer, return sequence set to true, and one additional hidden dense layer.
10	GRU_0HL_F	GRU	GRU neural network, with one GRU layer, return sequence set to false, and no additional hidden layers.
11	GRU_0HL_T		GRU neural network, with one GRU layer, return sequence set to true, and no additional hidden dense layers.
12	GRU_1HL_F		GRU neural network, with one GRU layer, return sequence set to false, and one additional hidden dense layer.
13	GRU_1HL_T		GRU neural network, with one GRU layer, return sequence set to true, and one additional hidden dense layer.
14	XGBoost	XGBoost	XGBoost model. Hyperparameters analysed are described earlier in this paper.

Appendix 4: Useful links

Below is a list of links which we hope that the reader finds useful.

- IBM's introductory series on neural networks:
<https://www.ibm.com/topics/neural-networks>
- IFoA's Certificate in Data Science programme:
<https://www.actuaries.org.uk/news-and-insights/news/data-science-credential>
- IFoA's Data Science Lifelong Learning page:
<https://actuaries.org.uk/learn/lifelong-learning/data-science/>
- *Google Tensorflow's* tutorial on time series:
https://www.tensorflow.org/tutorials/structured_data/time_series

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Institute
and Faculty
of Actuaries

London

1-3 Staple Inn Hall · High Holborn · London · WC1V 7QJ

Tel: +44 (0) 20 7632 2100 · Fax: +44 (0) 20 7632 2111

Edinburgh

Level 2 · Exchange Crescent · 7 Conference Square · Edinburgh · EH3 8RA

Tel: +44 (0) 131 240 1300 · Fax +44 (0) 131 240 1311

Oxford

1st Floor · Park Central · 40/41 Park End Street · Oxford · OX1 1JD Tel:

+44 (0) 1865 268 200 · Fax: +44 (0) 1865 268 211

Beijing

Level 14 · China World Office · No.1 Jianguomenwai Avenue · Chaoyang District · Beijing, China
100004

Tel: + +86 (10) 6535 0248

Hong Kong

1803 Tower One · Lippo Centre · 89 Queensway · Hong Kong Tel:

+11 (0) 852 2147 9418

Singapore

5 Shenton Way · UIC Building · #10-01 · Singapore · 068808

Tel: +65 8778 1784

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