### Master of Science in Statistics Dissertation: Draft #1

#### Title



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#### Abstract

We consider pattern prediction of financial time-series data. The algorithm framework and workflow is developed and proved on daily sampled OHLCV (open-high-low-close-volume) time-series data for JSE equity markets. The input patterns are based on input data vectors are equal size data windows pre-processed into a sequence of daily, weekly and monthly sampled feature measurement changes (here log feature fluctuations). The data processing is split into at online batch processed step where data is compressed using a stacked autoencoder (SAE) via unsupervised learning, and then batch supervised learning is carried out using the data-compression algorithm with the output being a pattern sequence of measured time-series feature fluctuations

(log differenced data) in the future (ex-post) from the training and validation data. Weight initializations for these networks are implemented with Restricted Boltzmann Machine (RBM) pre-training, and variance based initializations. The historical simulation is then run using an online feedforward neutral network (FNN) initialised with the weights from the online training and validation step. The validity of results is considered under a rigorous assessment of backtest overfitting (Combinatorially Symmetrical Cross Validation), and the results are then considered in terms of test for statistical arbitrage with simple correction for transaction costs.

**Keywords:** online learning, feedforward neural network, restricted boltzmann machine, stacked autoencoder, pattern prediction, JSE, non-linear, financial time series, combinatorially symmetrical cross validation, backtest overfitting

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#### Part I

# Online non-linear prediction of

# financial time-series patterns

#### 1 Introduction

This dissertation explores several ideas for the purposes of effective and valid stock price prediction, and suggests a novel approach to the combination of techniques. The first part of the implementation focuses on the training of deep neural networks, and combines usage of several well known designs for data reduction, deep learning and online learning. The aim here is to produce a deep learning model that is able to effectively produce stock price predictions in an online process, using data reduction of high dimensional finance data in order to achieve better results.

The second part of the paper focuses on whether the model can be trained such that it does not succumb to backtest overfitting, namely by using methods suggested by Bailey et al. [4] in order to calculate the likelihood of overfitting having occurred. As they've noted, and discussed more fully in 2.6, backtesting overfitting for trading strategies have become problematically widespread in financial literature. Here we investigate how more rigorous validation techniques can be applied to deep learning models in order to avoid such overfitting. Further validation takes place in assessing the potential profitability of the model in a live market.

The literature review in chapter 2 has a fuller discussion of work that has been done to precede the various techniques which have been implemented. A brief introduction to technical analysis in the financial sector, with it's perspective and history, is discussed and forms the basis for proposing and using the technical analysis methods throughout this paper 2.1. The literature review then moves onto covering the usage and history of Neural Networks, which has progressed enourmously in recent times. The section covers both the basic foundations, and also discusses the recent work which has resulted in the widespread use of deep learning models 2.2. This is extended to the coverage of Stacked Autoencoders, and their efficacy in data reduction for complex systems, which has lead them to be pivotal tools in deep learning models 2.3. Online learning methods are discussed in 2.4 with a coverage of both the historical basis as well as the more recent devopments which have allowed for further improvement in the algorithms. The chapter finishes off with discussing the impact of backtest overfitting and how some notable works developed by Bailey et al. have provided tools with which this may be avoided 2.6.

Chapter 3 provides the details around the data which has been used in the paper. The chapter starts with describing how the data is processed using log feature fluctuations, and then is expanded to include the fluctuations over rolling window periods. The synthetic data used is generated using Geometric Brownian motion, and is implemented such that each randomly seeded dataset consists of the following stocks:

may need to add some appropriate in this section 1. INTRODUCTION 10

- 3 stocks with an upward trend, and high, medium and low variance
- 3 stocks with a downward trend, and high medium and low variance
- 3 stocks with a sideways trend, and high medium andlow variance

There is a further summary of the real data used: OHLCV data for the JSE ALSI over a period of 20 years.

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Chapter 4 provides more indepth details on the algorithms and structures used to implement the dissertation process. The structure of feedforward neural networks is discussed, as well as how they are trained using the backpropagation algorithm, and how that can be applied in a stochastic descent framework 4.2. The chapter also provides details for how these network weight initializations can impact performance, and how this can be improved with either RBM pre-training (as per [37]) or through newer variance based techniques - this includes the structures and training techniques used for the Stacked Autoencoders 4.4. How the CSCV techniques suggested by [4] are implemented and used to derive a Probability of Backtest Overfitting figure for the full training and testing processes is detailed in 4.6. The chapter ends with a summary of the full end to end process, from data preparation to the output of a PBO figure in ??.

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Chapter 7 is the final chapter, and discusses the results found from the process.

this will need to be finished once results are known

#### 2 Literature Review

#### 2.1 Technical Analysis

Technical analysis is a financial analytical practice that makes use of past price data in order to identify market structures, as well as forecast future price movements. The techniques are typically objective methodologies which rely solely on past market data (price and volume). They stand in contrast to fundamental analysis, where experts will consider a companies operations, management and future prospects in order to arrive at an evaluation. The basis of much technical analysis, originally developed through Dow Theory, is the belief that stock market prices will move directionally (upwards, downwards or sideways), and that past movements can be used to determine these trends [65].

One of the primary methods in technical analysis is the use of charts in order to identify price patterns. These charts will be produced using the available market data and a known design, such as the popular candle-bar plot, which can then be compared to historical data to match it to a particular pattern. These patterns are thus indicative that the stock is likely to take on a particular price trend, or is in a particular state [65]. There is a certain amount of controversy around technical analysis, where many argue that it is contradictory to the random walk and weak form efficient market hypotheses, and as such is not valuable or useful [31]. The argument against this, is that technical analysis does not rely on past action to predict the the future, but is rather a measure of current trading, and how the market has reacted after similar patterns have occurred in the past [49]. Further, even if the analysis is unable to effectively forecast future price trends, it can still be useful to exploit trading opportunities in the market [76].

With the advent of processing power becoming cheaply available, there has been an increase in research to adapt computing techniques to technical analysis. The breadth and superhuman speed in which systems are able to perform technical analysis far outstrips what was possible before, and as such they have become the focus of competitive performance for many market participants [48]. To this end, there has been much research to apply machine learning algorithms to perform pattern recognition on stock price movements.

Financial markets have been shown to be complex and adaptive systems, where the effects of interaction between participants can be highly non-linear [3]. Complex and dynamic systems such as these may often exist at the 'order-disorder border' - they will generate certain non-random patterned and internal organisation, which can be assessed and identified, however they will also exhibit a certain amount of randomness in their behaviours, or 'chaos' [18]. Further, it's been shown that there is enough signal to reconstruct the phase space of chaotic systems from singular observations, encouraging that we might be able to perform effective technical analysis in the context of financial markets [66, 83]. As a result, trying to identify these patterns and structures is a simultaneously reasonable and notoriously difficult goal. While it is often clear in hindsight that the patterns exist, the amount of noise and nonlinearity in the system can make prediction challenging. Fittingly then, neural networks have become a popular choice for modelling within the financial markets. Due to their structure, they are able to learn non-linear interactions between their inputs and outputs, with even early research showing their ability to achieve statistically significant results, which

lends weight to the argument against the efficient market hypothesis [81].

#### 2.2 Neural Networks

A Neural Network (NN) is a learning model which was originally inspired by the biological mechanisms of neurons in brains. The structure is essentially that of a network system, with connected nodes and edges, or neurons and weights in context. The neurons are based on the same idea as synapses as seen in the brain - where a buildup of input results in a firing of output. The input here is determined by the models input (real numbers typically), and processed through the weights and activation functions of the neuron, which then results in an output value either at an intermediate level, or as the models final output. The system learns by considering input samples sequentially, and adjusting the weights between edges to result in more accurate outputs, which may either be classification or regression values.

Structured neural networks that learn to some extent have been around since the second half of the 21st century [74], though have been through several cycles of popularity. The first versions tended to be very simple with one one layer of hidden neurons [45]. It was only later, through the application of the backpropagation algorithm, that they started to become more practical and popular [93].

With the rise in popularity, many different network formations were developed and suggested. One of the initial suggestions was the conceptually simple Feedforward Neural Network (FNN) as described above - an acyclic graph where inputs are processed in a single direction until the output is reached. The other notable earlier model was the Recurrent Neural Network (RNN), which has a cyclic graph instead - this results in a more powerful computational system than the standard FNN, which was shown to be effective quite early on [79]. The Long Short-Term Memory (LSTM) network was another that used recurrent dynamics, though at a neuron level, in that the neuron is responsible for remembering values for an arbitrary time period [42]. Convolutional Neural Networks (CNN) have a non-recurrent structure, but implement separate pooling layers of neurons which consider the adjacent input values for each feature (e.g. pixels next to each other). These have been shown to be incredibly effective at tasks such as image recognition, as elaborated on later.

There are three primary learning paradigms used in neural network training - Supervised Learning (SL), where the network is trained on inputs with known outputs; Unsupervised Learning (UL), where the network is trained to identify unknown structures as an output; and Reinforcement Learning (RL), where environmental reactions are used as inputs to train a network for certain outputs [74]. While all of these configurations and paradigms have their benefits and uses, this paper will largely focus on FNN and RNNs, trained through SL and UL.

#### 2.2.1 Training and Backpropagation

Historically, the crux of neural networks popularity has often been based on the development of novel training methodologies, and how they have increased performance. In line with this, the Backpropagation (BP) algorithm (as defined in 4.2.3) has played a pivotal part: while neural network (or perceptron) models were around long be-

fore NN popularity, they were largely deemed ineffective, at least in comparison to other available models of the time [64]. It was only during the 1980s that the BP algorithm was applied to NNs, and the field started to gain in popularity again [54, 94].

Rumelhart et al. showed that the BP algorithm as applied in NNs resulted in useful feature representations occurring in hidden layers and the empirical success that resulted thereof [72]. Shortly after, LeCun et al. applied the BP algorithm to CNNs with adaptive connections. They were able to show impressive performance for the time in classifying handwritten images, with the images as a direct input (rather than a feature vector) [55].

While many improvements were made during this time via gradient descent modifications, the models were typically of a shallow nature due to problems encountered trying to train deeper networks. Early experiments with deep networks resulted in poor performance due to what is now widely known as the problem of either vanishing or exploding gradients [67]. Essentially, as more layers are added to the network, the backpropagation algorithm (with typical activation function neurons), results in error signals that either shrink or grow out of bounds at an exponential rate. One of the first suggested and primary solutions to the problem is to perform pre-training on the network through unsupervised learning [74], which is discussed more fully in 2.3.2. Variance based weight initialization techniques have grown in popularity in recent years, as discussed in 2.2.4.

There were also initial concerns that the BP algorithm as applied to high dimensional neural networks would result in the network weights being trapped in local minima if a simple gradient descent was used (e.g. where no small changes to the configuration would reduce the average error rate) [56]. However, empirically, this tends not to so problematic, and large networks usually reach solutions of equitable performance. More recent research has shown that the solution spaces largely consist of many saddle points, each with varying gradients of the features, but which also tend to have similar values of the objective function [19]. Ge et al. have also shown that it is possible to escape saddle points and offer a guaranteed global convergence in certain non-convex problems [27].

#### 2.2.2 Activation Functions

One of the upfront configuration choices necessary is the activation function, which allows the mapping of input to output at the neuron level. There have been many suggestions and experiments with different functions, though there are some common features amongst functions which might make them appropriate: Non-linearity allows for neural networks to operate as universal approximators, as shown in [43], continuous differentiability allows for the use of gradient descent and whether the function is monotonic has been shown to indicate whether the solution can be guaranteed to have a unique periodic solution [95]. Lastly, the range of the function (infinite or finite) can impact both the stability and efficiency of the training.

Some of the most popular functions that have been used are the Sigmoid, TanH, ReLU and Softsign. There have been various studies showing the effectiveness of the different activations under varying initialization (or pre-training) for weights. Glorot and Bengio noted that the typical Sigmoid and TanH functions performed poorly with standard minimization, and result in slower convergence and worse minima, and

showed that Softsign with a non-standard initialization resulted in quicker convergence [29]. Further research with Bordes & Bengio found that the rectifier (ReLU) functions were more effective in deep sparse networks compared to the TanH function [30].

#### 2.2.3 Deep Learning

As noted above, most of the earlier work using neural networks relied on shallow models with few layers. However, a resurgence in interest occurred in 2006 after several papers demonstrated the efficacy of unsupervised pre-training of networks prior to supervised training. The effect was substantial enough to allow much deeper layered networks to be trained than before [10, 36].

The essential point behind the unsupervised learning was to initialize the weights in the network to sensible values in light of the problem context. The methods used trained each layer to be able to reconstruct the model of the features in the layer below (to a varying degree of accuracy). Sequentially pre-training and combining layers like this, the process generated a deep neural network with appropriate weights. Once done, a final output layer was added and the entire network could then be fine-tuned through backpropagation, without suffering such performance degradation through vanishing or exploding gradients [36, 71, 37]. This is expanded up further in 2.3.2. Le Roux and Bengio were able to show that within the DBNs produced by Hinton [36], adding hidden nodes resulted in strictly improved modelling capabilities, and they suggested that increasing the number of layers is likely to result in increased representational ability (subject to efficacy of previous layers), thus establishing the argument for deep networks in theory as well as practice [57].

FNNs had been shown to be effective in modelling high dimensional data even prior to the breakthroughs in deep networks [11], so it fits that the deep networks were shown to be extremely effective in high dimensional data classification. Early implementations shown increased efficacy in handwriting recognition, as well as pedestrian recognition [77]. When it came to data types such as sound and images, CNNs were implemented on several occasions with record breaking model performances in recognition, notably in ImageNet and WaveNet [46, 90].

As more research into deep networks was conducted, it became apparent that with large enough datasets, the layerwise pre-training of networks was not actually necessary to achieve high performance standards [46, 30, 15]. When training for long enough, it was reported that the pre-training offered little to no benefit, though these models were typically using datasets far larger than were attempted before (as a result of hardware improvements enabling as much). While these results did require that certain attention was paid to the initialization, as well as the use of nonlinear activation units, it did suggest that pre-training largely acted as a prior which may not be necessary if large enough labelled datasets are available [12]. Naturally, pre-training was still implemented to prevent overfitting in smaller datasets.

#### 2.2.4 Weight Initialization Improvements

One of the more critical innovations to allow effective training of deep networks without using pre-training was the development of more sophisticated techniques for weight initialization. One of the first and most popular of these was presented by Glorot and Bengio for use in Sigmoid based networks, and is commonly referred to as Xavier/Glorot initialization [29]. The technique is layer specific and based on a linear activation hypothesis, such that the initial weights would maintain the same variance for input for information that is passed backwards as in accordance with the nature of the activation function. While the assumption of linearity is not entirely correct, they point out that as the start of the learning process, it is typically the area of the activations where the gradient is close to 1 which is being explore, thus intially approximating a linear effect. The effect is a technique that increases learning efficacy and optima found [29].

This same methodology was extended for the ReLU activation by He et. al, once again based on the linearity hypothesis around the relevant activation function, (Parametric Rectified Linear Units in this case). The combination of ReLU activations, which are computationally inexpensive and do not suffer from learning slowdown, as well as an effective weight initialization technique such that pretraining was not necessary produced a seminal FNN training framework. He et al. were able to achieve state of the art performance and produced the best known error rate at the time on the ImageNet dataset [35]. The efficacy of these techniques has established them as norms in the training of deep neural networks.

#### 2.3 Stacked Autoencoders

#### 2.3.1 High Dimensional Data Reduction

As noted, machine learning techniques have been shown to be extremely effective at modelling non-linear inputs to outputs - neural networks have even been shown to be universal function approximators in this regard [43]. More traditional statistical models will typically process the available feature data to select the most significant features to be used in the model once its defined - evident in a processes such as subset selection [73]. Machine learning techniques are no different in this regard, and feature data will typically be transformed to smaller observations of more significance prior to being used as input to a model, such as the neural networks described above.

Financial data, in line with the complex and dynamic system that it represents, is often of a very high dimensional nature, which offers opportunities through more sophisticated analysis, but also introduces the curses of dimensionality [21]. The increased dimensionality can result in higher processing complexities when needing to do basic tasks such as estimating a covariance matrix (a commonplace necessity in finance), as well as increase the risk of incorrect assumptions based on spurious variable collinearity [25]. Noise accumulation in high dimensional data can create further problems, resulting in problems performing variable selection and ultimately having a large impact on classification and regression models [26].

Time series data can introduce its own set of challenges - there is often not enough data available to understand and predict the process [24], the time variable dependence creates complexity in how much past data to consider at any point, and the data is typically non-stationary [52]. Thus, high dimensional time series data (which many financial problems focus on), require careful consideration on how to handle their inputs and analysis.

Deep learning techniques are a natural choice in this context, and much research has been done to show their (varying) efficacy on time series data. The most successful of these models have been ones which modify deep learning techniques to incorporate the temporal aspect of the data (e.g. Conditional Restricted Boltzmann Machines or Recurrent Neural Networks), rather than static, and those which have performed feature selection processes rather than operating on the raw data (e.g. Auto-encoders) [52].

Two of the seminal pieces of research that have lead to the resurgence in machine learning and deep learning were the algorithms for training deep belief networks [36], as well as the usage of stacked auto-encoders [71, 10].

#### 2.3.2 Deep Belief Networks

Autoencoders were suggested by Hinton et. al as a method of transforming high dimensional data to lower dimensional input vectors, in order to alleviate some of the problems detailed above, and increase performance of deep belief networks [37].

One of the more prominent classical techniques for dimension reduction is principal components analysis (PCA), which uses linear algebra to find the directions of greatest variance, and represent the observation samples features along each of these directions, thus maximising the variational representation. Hinton et al. show that autoencoders are a nonlinear generalization of PCA. The structure and training algorithms of the autoencoder show it to be a specialised neural network - there is a multilayer encoder network which is able to transform to a lower dimension, and a symmetrical decoder network to recover the data from the code as represented in Figure 1. As with neural networks, the gradient weights can be trained through the feedforward and backpropagation algorithms.

The primary challenge presented here was the initial weighting of the networks—with large initial weights the autoencoder will often find a poor local minima, and with small initial weights the gradients are too small to effectively train deep layered networks. The critical suggestion by Hinton et al. was to used layered Restricted Boltzmann Machines (RBM) in order to initialise the weights. For each layer of the desired autoencoder, a RBM is formed and trained with the previous layer (or RBM) [38]. Once all the layers have been trained in this way, they are mirrored to form the decoder network. This then forms the initial weights to be fine tuned further, as per the Fine-tuning step in 1. They showed the deep autoencoder networks were significantly more effective than PCA or shallow autoencoders on multiple dataset types.

#### 2.3.3 Stacked Denoising Autoencoders

The second important piece of work was the development of a denoising autoencoder (DAE), by Vincent et al. [86]. One of the problems identified in the DBN model (and those similar), is that if the encoder dimensions were too high, it is likely that the encoder would learn a trivial encoding - essentially creating a copy the input model. The one way of tackling this issue is to constrain the representation with bottlenecks and sparse autoencoder layers, which can be seen in figure 1.

Vincent et al. explore a very different approach to the problem, which was to develop an implementation of autoencoder which focused on partially corrupting the input, and so force the network to denoise it. The theory here is based on two ideas - the first, is that a higher dimensional representation should be robust to partial

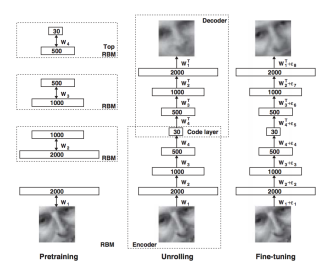


Figure 1: The Autoencoder training steps [37]

corruption of the input data; and the second is that the denoising process will force model focus to shift to extracting useful features from the input.

The algorithms and structures are largely the same as described for DBNs above, with the key difference being that the model is trained to reconstruct the original input, but only using a corrupted version of the input (where noise has been added to it), and so is forced to learn smarter feature mappings and extractions. The DAE suggested then is a stochastic variant of the autoencoder, which has the benefit of being able to implement higher dimensional representations without risking training of a trivial identity mapping. Notably, in the Stacked Denoising Autoencoder (SDAE) formation, only the initial input is corrupted (as opposed to the input from layer to layer). It was shown that the SDAE model outperformed previous AE and DBN networks on numerous benchmark datasets [86] .

#### 2.3.4 Pre-training

The methods described above follow a similar approach: greedy layer-wise unsupervised pre-training in order to determine initial weights, followed by supervised fine tuning to arrive at the final model. It is shown numerous times, that the pretraining process can result in significant performance gains [86]. However it is not immediately apparent, given the nature of backpropagation algorithms and the like, why this is the case. Erhan et al. performed extensive empirical simulations in order to suggest an explanation to the mechanism of pre-training [23].

While their results were not entirely conclusive, they did lend themselves to a reasonable hypothesis: the unsupervised pre-training results in a form of regularization on the model - variance is minimized, and the bias introduced acts as a prior to direct the model configuration towards a sample space that is effective for the unsupervised learning generalization optimisations.

#### 2.3.5 Time Series Applications

The autoencoder papers reviewed so far in this section derive their results primarily from classification problems, and so do not necessarily account for the problems involved with time series as described in 2.3.1. Due to the inherent difficulties with predictions in the financial system, it can sometimes be unclear if the shortcoming in results is due to this system complexity or if the methodologies used are unsuited for the purpose. In light of this it is worth pointing out that Stacked Autoencoder (SAE) implementations have been shown to be effective in many time series systems.

Lv et al. implemented a deep learning SAE model using the methods described in 2.3.3 in order to predict traffic flow at various time intervals (15, 30, 45 and 60 minutes) - a problem not so structurally dissimilar from what will be presented in this paper [61]. They were able to show that the deep SAE was able to offer prediction results which were both objectively good and also persistently outperformed the comparison models used (backpropagation neural network, random walk forecast, support vector machine and a radial basis function neural network).

In a review of unsupervised feature learning and deep learning methods on time series, Langkvist et al. noted that the use of autoencoders, either as a technique in themselves, or as an auxiliary technique to models such as convolutional neural networks, were able to offer performance increases in areas such as video analysis, motion capture data and bacteria identification [52].

#### 2.3.6 Financial Applications

There have of course also been successful applications of stacked autoencoders and deep learning models in finance as well. Takeuchi et al. performed some earlier work showing the use of autoencoders when applied to a momentum trading strategy. They implemented an RBM pre-trained DBN as per 2.3.2, and assessed the networks classification performance for ordinary shares on NYSE, AMEX and Nasdaq. This showed that using a DBN network resulted in significant performance increases compared to the standard momentum strategy [82].

Zhao et al. used SDAEs and combined them with the bootstrap aggregation ensemble method (bagging) in a study of predicting the crude oil price. They compared the proposed model to a variety of benchmarks, including standard SAE, bagged and standard feedforward networks and SVRs. The results indicated that the SAE models were more accurate, with the bagged SAE model performing the best, though at a significant increase in computational costs in comparison to standard SAE [99].

While much of the financial literature has focused on the use of RBM based models, Autoencoders and SAEs have recently been gaining popularity in performing feature reduction. Troiano et al. specifically investigate the use of different feature reduction models for trend prediction in finance [84]. In line with being primarily interested in the effect of feature reduction techniques, rather than the classification performance itself, only an SVM model was used to test results. Using various periods from historical S&P 500 data, they were able to show that AE outperformed the RBM model significantly in numerous accuracy measures, and was able to do so at a fraction of the training time.

Bao et al. note that the research has been lacking with regards to whether SAEs

should be used for financial prediction models or not [8]. They suggest a novel model which combines Wavelet Transformation, SAEs and a Long Short Term Memory (LSTM) network. Using data from several financial exchanges (considering a range of developed and undeveloped markets), they assess the models applicability to OHLC prediction. Comparing the model to configurations without the SAE layers, and a RNN model as benchmark, they showed that the inclusion of SAEs resulted in less volatility and greater accuracy, which in turn offered higher profitabilities in a buyand-hold trading strategy.

More novel autoencoder applications have also been attempted, with Hsu suggesting the use of a Recurrent Autoencoder for multidimensional time series prediction [44]. There is a clear pattern through the literature that the use of AEs and SAEs both by themselves and when used as an assisting technique result in more accurate prediction results and less computationally expensive training.

# 2.4 Online Learning Algorithms and Gradient Descent

Most classic machine learning algorithms operate under the assumption that, for all intents and purposes, the full dataset has been collected and that the amount of training data for the model is both finite and immediately available. However, as the growth of information grows in an exponential fashion, there are numerous areas where the expected training data for the model will continue to grow. In these cases it would be disadvantageous to go through the full training and validation process again in order to incorporate the newly available data.

Online algorithms are designed to offset these issues by adjusting the batch training technique to rather repetitively draw on single samples from the data on which the models parameters can be adjusted. The benefit is that they are able to quickly process a large number of observations and readjust the model, though the downfall is that they are not always able to optimize the cost function to the same extent as offline batch algorithms [1].

Bottou and Cun argue that as the size of the dataset grows significantly, online algorithms advantages result in them outperforming offline models, despite any initial drawbacks [13]. Previous research had shown that online algorithms typically perform as fast as batch algorithms during the search phase of parameter optimization, but that final phase convergence tended to fluctuate around the optima due to the noise present in single sample gradients [53, 14]. Bottou and Cun showed in fact, that it is more practical to consider the convergence towards the parameters of the optima, rather than the optima itself (as defined by the cost function) - the difference between the learning speed and optimization speed, respectively [13]. Theoretical and empirical findings were presented to show that a stochastic online gradient descent algorithm was able to outperform the batch model for parameter estimation, and was able to asymptotically outperform in the number of samples processed in a time period. The stochastic aspect of the algorithm is related to random observations from batch sample groups being used as the gradient basis. Theoretically, this slows down the convergence, but speeds up the processing speed of each batch - a technique which has later been shown to be generally successful [78, 100].

Stochastic gradient descent algorithms have resulted in a fair amount of further research due to their applicability to machine learning and the online benefit, especially in relation to their usage in neural networks.

#### 2.5 Gradient Learning Improvements

#### 2.5.1 Gradient Adjustments and Regularization

One of the earlier improvements to convergence rates was the Momentum algorithm, as developed by Tseng [85]. As noted, stochastic descent often introduces significant oscillation around an optima, which slows down convergence. Momentum reduces this by decreasing movement in directions of high curvature, and increasing movement towards directions consistent with previous gradients (this is achieved through combining gradient movements in opposite directions).

There have been several attempts to introduce effective regularization into the SGD process. Bartlett et al. presented Adaptive Online Gradient Descent, which implements an adaptive step size through a  $\lambda$  penalty on the learning rate, which was shown to be nearly optimal in a strong sense [9]. Langford et al. demonstrated a variation named Truncated Gradient, which introduced an enforced weight sparsity parameter. The weight sparsity is able to achieve equitable effects to  $L_1$  regularization (similar to Lasso Regression). They were able to show that implementation performed effective feature reduction, while having little effect on performance [51]. Other approaches, such as AdaGrad, aim to improve the robustness of gradient training by adjusting the updates to parameters according to frequency - e.g. larger updates to infrequent parameters, and smaller updates to frequent parameters [22, 97].

#### 2.5.2 Dropout

One of the instrumental improvements in further achievements, was to modify the backpropagation algorithm using the dropout technique, as suggested by Hinton et al [39]. When training of large networks is attempted on small datasets, it often results in overfitting and poor results on out of sample data. Dropout helps resolve this by randomly excluding a certain percentage (usually 0.5) of feature detectors on each training iteration. The effect is to stop co-adaptations of feature detectors, and by rather training each neuron in a wide variety of internal configurations, it forces them to take on more usefully generalizable characteristics (it was noted that this is not a dissimilar technique to ensemble methods, or bagging). The authors were able to show that the method resulting in significant improvements on benchmark data sets (e.g. MNIST, CIFAR-10), and that a simpler model using dropout was able to achieve near comparable performance for the ImageNet dataset.

Goodfellow et al. used the dropout technique as the basis for their maxout activation function technique, which leverages and improves on dropouts fast optimisation and accuracy through averaging characteristics. The maxout model was shown to achieve state of the art performance on benchmark datasets, as well as have a strong theoretical grounding [28]. Further work was done by Wang et al., which improved on the dropout (and potentially maxout) techniques through fast sampling, resulting in an order of magnitude speedup in training [89].

#### 2.5.3 Learning Rate Schedules

Several more recent approaches look towards learning rate adjustment schedules rather than strategies to alter the weight update, as with momentum and the like. A constant learning rate often suffers from one of two problems - if set too high, it can cause divergent behaviour in the loss function, though if set too low, it can result in slow learning or an inability to escape saddle points effectively. Finding an optimal learning rate requires some degree of testing, though even then a singular value may fail to achieve the same degree of efficacy as a range of values explored throughou the solution space.

Learning rate scheduling has been proposed as a solution to this problem. These methods implement an repetitive cycle for the learning rate, such that it is set through a range of values between a mimima and maxima, being adjusted slightly with each new epoch. Smith suggests using the Cyclical Learning Rate to achieve this, and points out that part of the benefit in learning rate schedules is the ability to jump out of sharp optima points which may not generalise well to unseen data [80]. It's shown that this implementation can have significant performance effects in reaching either the same optima in fewer epochs or a better overall optima, including when used in conjunction with other learning optimizations noted in 2.5. These results were shown against standard datasets such as CIFAR-10, CIFAR-100 and Imagent [80].

Similarly, Loshchilov and Hutter, expand on this and present Stochastic Gradient Descent with Warm Restarts [60]. The approach is implemented similarly, but rather follows an asymmetric cycle from a maximum to a minimum, and then starting once again at the maximum once the set number of epochs has passed. They too note the increased performance on CIFAR-10 and CIFAR-100 datasets in reaching optima far quicker, as well as noting the efficacy of the learning rate schedule even without implementing the restarts [60].

#### 2.6 Backtesting and Model Validation

Much of financial academic literature is currently facing a problem in terms of validation and verification of results. The primary method of going about these ends in the past has been to perform historical simulations, or backtests, in order to prove profitability of a trading strategy. The recent advances in both technology and the algorithms available to construct these strategies has resulted in researchers being able to run so many iterations of a model or strategy configuration through these backtests, that its become increasingly difficult to control for spurious results, with some papers suggesting that most published research findings are false [47].

The standard way of implementing backtests is to split the data into two portions: an In Sample (IS) portion which is used to train the model, and an Out of Sample (OOS) portion which is used to test the model and validate results. The problem lies in that millions of different model configurations might be tested, and if more sophisticated test measures are not in place (i.e. not just the standard Neyman-Pearson hypothesis testing framework is implemented), then it is only a matter of time before a false positive result occurs which shows high performance both IS and OOS (i.e. overfitting). The nature of financial data, where there is a low signal-to-noise ratio in a dynamic and adaptive system, and where there is only one true data sequence,

makes it difficult to resolve these issues effectively [4, 63].

Overfitting is not a novel issue, and has of course been tackled in various literature areas, including machine learning. However, in that context, the frameworks are often not suited to the buy/sell with random frequency structure of investment strategies. They also do not account for overfitting outside of the output parameters, or take into consideration the number of trials attempted. Other methods, such as hold-out, are arguably still faulty due to researcher knowledge while constructing models [75]. One of the downfalls of the typical IS-OOS set up in the financial context is also that the most recent (and relevant) data will not be able to be used for the model training.

There have been some suggestions to resolve the problem that is occuring in the literature as a result of this - some work suggesting new frameworks, which this section will cover, and others which focus on the review process or how data and replication procedures are made available [69]. While the points made with regard to the review process and so on are certainly important, they don't aid with more effective model training for the researcher up front, and so will not be covered here.

#### 2.6.1 Testing Methodologies

Considering the issues laid out above, there has been much work to develop alternative approaches to backtesting. One of the common approaches to avoid backtest overfitting is the hold-out strategy, where a certain portion of the dataset is reserved for testing true OOS performance. Numerous problems have been pointed out with this approach, including that the data is often used regardless, or that awareness of the movements in the data may, consciously or otherwise, influence strategy and test design by the researchers [75]. For small samples, a hold-out strategy may be too short to be conclusive [92], and even for large samples it results in the most recent data (which is arguably the most pertinent) not being used for model selection [34, 4].

There has been work by several authors to try and lay out techniques to try and avert backtest overfitting. The Model Confidence Set (MCS), as developed by Hansen et al. [32], starts with a collection of models or configurations, and remove models iteratively according to a defined loss function. The confidence set is defined by the remaining models once a non-rejection takes place within the process, and these models are considered to be statistically similar within a certain confidence range. MCS is thus able to facilitate equitable model selection. However, Aparicio et al. [2], showed that while MCS is a potential strategy, in practice is is ineffective due to the inordinate requirement of signal-to-noise necessary to identify true superior models, as well as a lack of penalization over the number of trials attempted.

Bailey et al. [4] have developed a more robust approach to backtesting and how overfitting during strategy selection might be avoided, called Combinatorially Symmetric Cross-validation (CSCV). Their research defines backtest overfitting as having occurred when the strategy selection which maximizes IS performance systematically underperforms median OOS in comparison to the remaining configurations. They use this definition to develop a framework which measures the probability of such an event occurring, where the sample space is the combined pairs of IS and OOS performance of the available configurations. The probability of backtest overfitting (PBO) is then established as the likelihood of a configuration underperforming the median IS while outperforming IS.

The CSCV methodology provides several important benefits over traditional testing frameworks, including the usual K-fold cross validation used in machine learning. By recombining the slices of available data, both the training and testing sets are of equal size, which is particularly advantageous when comparing financial statistics such as the Sharpe Ratio (SR), which are susceptible to sample size. Additionally, the symmetry of the set combinations in CSCV ensure that performance degradation is only as a result of overfitting, and not arbitrary differences in data sets. It is pointed out that while CSCV and PBO should be used to evaluate the quality of a strategy, they should not be the function on which strategy selection relies, which in itself would result in overfitting.

#### 2.6.2 Test Data Length

The CSCV methodology offers an important but highly generalised framework to assess models and backtest overfitting. It doesnt however indicate which metrics should be used to assess the IS and OOS performance, nor any indication on the amount of data needed to do so effectively. One of the noted limitations of the framework is that a high PBO indicates overfitting within the group of N strategies, which is not necessarily indicative that none of the strategies are skillful - it could be that all of them are. Also, as pointed out, it should not be used as an objective function to avoid overfitting, but rather as an evaluation tool. To this end it helps assess overfitting, but not necessarily avoid it.

A typical measure of evaluation used for financial models is the Sharpe Ratio (SR), which is the ratio of between average excess returns and the returns standard deviation - a measure of the return on risk. In the context of comparing models, SR is typically expressed annually to allow models with different frequencies to be compared. Lo et al. [59] show that annulaized SR can be expressed as

$$SR = \frac{\mu}{\sigma} \sqrt{q} \tag{1}$$

Using sample means and deviations,  $\hat{\mu}$  and  $\hat{\sigma}$ , SR can be shown to converge as follows (as  $y \to \infty$ )

$$\hat{SR} \to \mathcal{N}[SR, \frac{1 + \frac{SR^2}{2q}}{y}]$$
 (2)

Thus, when using SR estimations, which follow a Normal distribution, it is possible that where the true SR mean is zero we may still (with enough configurations attempted) find an SR measurement which optimises IS performance. This is shown by Bailey et al. [5], who propose the non-null probability of selecting an IS strategy with null expected performance OOS. Notably, typical methods such as hold-out once again fail, as the number of configurations attempted are not recorded. They add a further derivation, thich is the Minimum Backtest Length (MinBTL), ultimately showing that

$$MinBTL \approx \left(\frac{(1-\gamma)Z^{-1}[1-\frac{1}{N}] + \gamma Z^{-1}[1-\frac{1}{N}e^{-1}]}{\overline{E[max_N]}}\right)^2 < \frac{2ln[N]}{\overline{E[max_N]}}^2$$
 (3)

The statistic highlights the relationships between: selecting a strategy with a higher IS SR than expected OOS, the number of strategies tested (N), and the number of

shorten section?

years tested (y). The equation shows that as the number of strategies tested increases, the minimum back test length much also increase in order to contain the likelihood of overfitting to IS SR.

As shown extensively throughout ML literature, increased model complexity and number of parameters is one of the primary causes of overfitting. In context of the MinBTL formula, model complexity affects the number of configurations that are available and which may be tested, which in turn will increase likelihood of overfitting. A lack of consideration, or reporting, of the number of trials makes the potential for overfitting impossible to assess.

Bailey et al. expanded on this view with assessing the impact of presenting overfit models as correct. They were able to show that in lieu of any compensation effects (i.e. a series following a Gaussian random walk), there is no reason for overfitting to result in negative performance. However, where compensation effects apply (e.g. economic/investment cycles, bubble bursts, major corrections etc.), then the inclusion of memory in a strategy is likely to be detrimental to OOS performance if overfitting isnt controlled for [5].

#### 2.6.3 Sharpe Ratio

The use of the Sharpe Ratio in financial backtesting is not just an arbitrary or persistent literature choice. The statistic offers two benefits: the effectively strategy-agnostic financial information contained, as well as being relatable to the t-statistic, and so simple to perform hypothesis testing. The SR ratio (estimate from sample as  $\hat{SR}$ ) is defined as

$$SR = \frac{\mu}{\sigma} \tag{4}$$

The t-ratio is defined as

$$t - ratio = \frac{\hat{\mu}}{\hat{\sigma}/\sqrt{T}} \tag{5}$$

Evidently, the link heree ratio (though it can be generalized to another statistic with a probabilistic interpretation). Additionally, PBO is generally more in line with machine learning literature with the cross validation like approach on time series data.

It should be noted, that the literature detailing usage of the Sharpe ratio for strategy comparison is extensive, with numerous variations and methodologies offered [6]. However, the crux of this paper lies in whether an online neural network is able to make effective enough predictions that a strategy might use the predictions to be profitable. The subtlety here is that we will consider the usage of such forecasting within a strategy, rather than as a strategy itself. In line with this, statistics such as the Sharpe ratio will be used, but not form a critical consideration of the research here as the comparison of strategies will not be the focus.

# 3 Implementation: Data Processing and Generation

#### 3.1 Data Processing

The datasets used go through several transformations throughout the training and prediction process:

- 1 The raw data is log differenced and aggregated to rolling windows
- 2 The transformed data is split into Training and Prediction subsets
- 3 The predicted outputs have the scaling and log differencing reversed in order to reconstruct the actual price points

#### 3.1.1 Log Difference Transformation and Aggregation

All datasets are transformed into log feature fluctuation values, and were then aggregated to include fluctuations over rolling window periods. The log feature fluctuation for timepoint t is calculated using (6):

$$\log_{\text{diff}} x_t = ln(x_t) - ln(x_{t-1}) \tag{6}$$

This log feature fluctuation, is processed for each asset's closing price and for each time point (from the previous time point). The log fluctuations have the benefit of taking compound effects into account in a systematic way and are symmetric in terms of gains and losses.

The datasets are then expanded with the rolling window summations both in the past, for input, and in the future, for predicted output. A typical example would be past data aggregation windows of 1, 5 and 20, and a future prediction point of 5. These are calculated as summations of the log differences, such that for d days:

$$past\_agg\_x_{(t,d)} = \sum_{i=t-d}^{t} log\_diff\_x_i$$
 (7)

$$\texttt{future\_agg\_x}_{(t,d)} = \sum_{i=t+1}^{t+d} \texttt{log\_diff\_x}_i \tag{8}$$

Only data points with a full set of features are used for training and prediction later on in the framework.

#### 3.1.2 Data Scaling

Once the log difference data has been processed, the datasets are either standardized or normalized to allow for better learning. This is done in the typical ways:

$$standardized_x_i = \frac{(x_i - \bar{x})}{\sigma_x} \tag{9}$$

$$normalized_x_i = \frac{x_i - min(x)}{max(x) - min(x)}$$
 (10)

Two extensions of these are used later on, dubbed 'Limited Standardization' and 'Limited Normalization', which are used when the data is split up into the Training and Prediction sets. In this case all the data needs to be scaled, but the variance or range should not be travelling from the Prediction set to the Training set through the use of aggregated measures such as  $\bar{x}$ . Thus, if the data is split at point t out of t0 data points, the scaling would be implemented as follows:

$$\texttt{limited\_standardized\_x}_i = \frac{(x_i - \bar{x_{1:t}})}{\sigma_{x_{1:t}}}, \forall i \in (1, n) \tag{11}$$

$$limited\_normalized\_x_i = \frac{x_i - min(x_{1:t})}{max(x_{1:t}) - min(x_{1:t})}, \forall i \in (1, n)$$
(12)

This log-differenced, aggregated and scaled data is then used as the input for the neural network models.

#### 3.1.3 Reverse Data Scaling

The predicted data points are transformed back into actual prices for returns analysis. The first step is to reverse the scaling that is done in 3.1.2 in order to retrieve the log difference fluctuations.

reverse\_standardized\_
$$\mathbf{x}_i = \operatorname{standardized}_{\mathbf{x}_i} * \sigma_x + \bar{x}$$
 (13)

reverse\_normalized\_
$$x_i = normalized_x_i * (max(x) - min(x)) + min(x)$$
 (14)

These are applied similarly for the limited variations.

#### 3.1.4 Price Reconstruction

The predicted log fluctuations produced in 3.1.3 are used to reconstruct the predicted prices.

reconstructed price<sub>s,i</sub> = 
$$o_{s,i-t} * e^{p_{s,i}}$$
 (15)

where:

o = original asset prices

s =asset selected

i =timepoint predicted from

d =timepoint predicted to

p = predicted log fluctuation

These reconstructed prices are then used to assess model returns later in the Money Management Strategy (4.7).

#### 3.2 Synthetic Data Generation

Synthetic data generation is implemented using Geometric Brownian Motion (GBM) as described in Algorithm 1, which allows the stocks simulations to be implemented with a drift  $(\sigma)$  and variance  $(\mu)$  for each stock. The GBM simulation has the benefit of generating prices in a random walk, such that future movements are independent of past prices and so suitable for emulating an efficient financial market. The implications of using such data are discussed more fully in 7.1. Each dataset was generated with a random seed (using a Mersenne Twister pseudorandom number generator).

#### Algorithm 1: Geometric Brownian Motion Simulation

```
Input: \sigma, \mu, S_0, steps
t = 1/steps;
prices = [S_0];
foreach i in 1:t do
z = random() \sim N(0, 1);
S_t = S_{t-1} * e^{(\mu - \frac{\sigma^2}{2})t + \sigma\sqrt{t}z};
prices = [prices; S_t];
end
Result: prices
```

### 4 Implementation: Models and Algorithms

#### 4.1 Process Overview

The implementation focuses on bringing together several ideas: data reduction, deep learning with pre-training/weight initialization, online learning and backtest overfitting validation for the purposes of stock price prediction. The process implementation is discussed fully in 5.1, but can be summarised as the following:

- 1 The dataset is split into 2 subsets: the Training portion, and the Prediction portion.
- 2 The Training set is used to train the SAE and deep FFN using the SGD algorithm. These networks are constructed with pre-training or weight initialization techniques.
- 3 The Prediction set of data is used to continue training the network in an online manner using OGD.
- 4 The predictions made during the OGD process are descaled and prices are reconstructed.
- 5 These reconstructed price predictions are used by the MMS to calculate returns and P&L.
- 6 The returns and P&L calculated by the MMS are used by the CSCV process, to estimate the probability of backtest overfitting.

The rest of this chapter will detail the algorithms used to train the relevant FFN, RBM and SAE networks, as well as the trading strategy and CSCV & PBO testing procedures.

#### 4.2 Feedforward Neural Networks

Constructing Feedforward Neural Networks (FFN) in the form of multilayer perceptrons is a well established network technique, providing effective nonlinear representations for both shallow and deep structures [74]. Specifically, a FFN is made up of several non-cyclical layers: the first and last are the input and output layers, respectively, and any inbetween are referred to as 'hidden' layers. Each layer is made up of nodes which are fully connected to the nodes in the previous and following layers, but does not have connections to nodes within the layer - information only travels forward. Each node has an activation function, which acts on the weighted input from the previous layers' nodes.

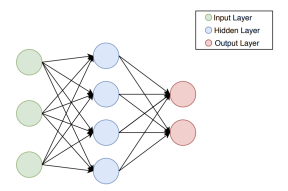


Figure 2: An example diagram of a Feedforwrd Neural Network with 1 hidden layer

#### 4.2.1 Notation and Network Representation

For the purposes of this chapter, the following notation will be used:

- 1 A network N will be contructed with L layers, each with e nodes
- 2 The weights from the  $j^{th}$  node in the  $l^{th}$  layer from the  $k^{th}$  node in the  $(l-1)^{th}$  layer are represented by  $w_{jk}^l$
- 3 The bias for the  $j^{th}$  node in layer l is represented by  $w_0^l j$
- 4 The output for the  $j^{th}$  node is defined as  $a = o(z_j)$ , for an activation function o (with inverse o') and weighted input z
- 5 The weighted input for the  $j^{th}$  neuron in a layer is

$$z_j^l = \sum_{i=1}^{e^{l-1}} a_i^{l-1} w_{ij} + w_{0j}$$
 (16)

These definitions allow an input into the network to be propagated through it, having the original values processed through the weights and activations functions, and have an output in the form of the network's last layer.

#### 4.2.2 Activation Functions

As noted in 2.2.2, there are 3 primary characteristics of concern for activation functions: non-linearity, continuous differentiability and monotonicity. While many different functions have been suggested and used, several of the more popular were implemented here.

**Sigmoid** The sigmoid, or logistic, function is one of the most popular and widely used activation functions historically, and is defined as

$$o(x) = \frac{1}{1 + e^{-x}} \tag{17}$$

$$o'(x) = o(x)(1 - o(x))$$
 (18)

The Sigmoid function is in the range [0,1], making it a suitable choice for problems requiring a probabilistic output. The slope of the function curve is both a boon and

a drawback: it allows for fast learning initially, but results in learning slowdown later (often casuing what is referred to as node 'saturation'). The exponent calculation is also computationally expensive, relatively speaking.

**ReLU** The Rectified Linuear Unit (ReLU) is a newer activation function which has been shown to be effective in deep learning networks. It is defined as

$$o(x) = \max(0, x) \tag{19}$$

(20)

$$o'(x) = \begin{cases} 1, & \text{if } x \ge 0 \\ 0, & \text{otherwise} \end{cases}$$

The function has the benefits of quick learning which doesn't saturate, as well as being computationally cheap. The downside is that the non-gradient for the negative range of the function can result in 'dead' nodes, which stop updating with the learning process.

Leaky ReLU The Leaky ReLU resolves the dying ReLU problem by adding a small gradient to the negative range of the function. This results in a slow learning being applied to 'dead' ReLU's, which may in turn result in them being used again if necessary.

(21)

$$o(x) = \begin{cases} x, & \text{if } x \ge 0 \\ 0.01x, & \text{otherwise} \end{cases}$$

(22)

$$o'(x) = \begin{cases} 1, & \text{if } x \ge 0 \\ 0.01, & \text{otherwise} \end{cases}$$

**Linear Activation** Linear activations don't transform the input, and have a constant gradient, which can be useful for key layers where no loss in error signal is desired, such as the output or encoding layers.

$$o(x) = x \tag{23}$$

$$o'(x) = 1 \tag{24}$$

#### 4.2.3 Backpropagation

The backpropagation algorithm, as discussed in 2.2.1, has allowed for effective training of FNNs for given data. The algorithm relies on incremental improvements of the model, as defined by decreasing the cost function. A common choice for cost is Mean Squared Errors (MSE):

$$C = \frac{1}{2} \|y - a^L\|^2 \tag{25}$$

A technical definition of the backpropagation algrithm is given in Algorithm 2, though the three primary steps of the algorithm are:

- 1 Forward Pass: The samples are propagated through the network, in order to generate the output  $\hat{y}_s$ . Vectorization is typically used to calculate the results for multiple samples at once, though conceptually each sample is used as input for the first layer, and subsequent layers use the weighted output from the previous layer passed through the activation function as input, using equation (16) for the weighted output and the equations defined in 4.2.2 for the activation functions.
- 2 Calculate Cost: The cost between the training output  $y_s$  and the model output  $\hat{y}_s$  is calculated. Using this cost function, the rate of change for the error in the output layer by the rate of change in the activation function in the last layer. This is captured in equation (26):

$$\delta_j^L = \frac{\partial C}{\partial a_j^L} o'\left(z_j^L\right) \tag{26}$$

The vectorized version of this is often used, and expressed in (27), where  $\nabla_a C$  represents a vector of components which represent the partial derivatives, such that they represent the rate of change of C relative to the activation functions' outputs.

$$\delta_j^L = \nabla_a C \otimes o'(z^L) \tag{27}$$

If using quadractive cost, as per 25, then the term  $\frac{\partial C}{\partial a_j^L}$  can be reduced to  $(a^L - y)$ , such that the equation for the output layer error is

$$\lambda^L = (a^L - y) \otimes o'(z^L) \tag{28}$$

- 3 Backwards Pass: The backwards pass consists of 2 steps, where the errors for each previous layer are calculated, and the weights for that layers are then adjusted accordingly:
  - 3.1 The activation values are propagated back through the network to calculate the delta values based on the cost value. This uses the same rationale as (26), but rather uses  $(w^{l+1})^T$  by the next layers error in order to get an indication of how the error rate and how it moves backwards through the network.

$$\lambda^{l} = ((w^{l+1})^{T} \lambda^{l+1}) \otimes o'(z^{l}) \tag{29}$$

3.2 In order to update the network, each weights output error and input activation are multiplied to find the weights gradient and this gradient is reduced by a factor of the learning rate  $\eta$ , which is then subtracted from the network weights to update them proportionately to the error observed and chosen learning rate.

 $w^l \to w^l - \eta \lambda^l (a^{l-1})^T \tag{30}$ 

Choices regarding the learning rates are discussed in 4.2.6 and the full algorithm is presented below in Algorithm 2.

#### **Algorithm 2:** Backpropagation

**Input:** Neural Network N, with randomly initialized weights w, L layers, and activation functions o and learning rate  $\eta$ 

**Data:** Testing set with inputs x, and outputs y

```
repeat
```

```
Select a sample x_s from x;
     // Perform the Forward Pass, to calculate the network output, y_s, for
          input sample x_s
     a_0 = x_s;
     foreach l in 1:L do
           \begin{aligned} & \textbf{for each } j \ in \ 1:e^l \ \textbf{do} \\ & \begin{vmatrix} z_j^l = \sum_{i=1}^{e^{l-1}} a_i^{l-1} w_{ij} + w_{0j}; \\ a_j^l = o(z_j^l); \end{aligned} 
          end
     \ensuremath{//} Calculate the error term (aka cost), as
     \lambda^L = (a^L - y) \otimes o'(z^L);
     // Perform the Backward Pass, to propagate the errors back and update the
          network accordingly
     foreach l in (L-1):1 do
          // Calculate the delta values
          \lambda^l = ((w^{l+1})^T \lambda^{l+1}) \otimes o'(z^l);
          // Update the weights w^l \rightarrow w^l - \eta \lambda^l (a^{l-1})^T;
until no new samples can be drawn from x;
```

#### 4.2.4 Gradient Descent Algorithms

**Result:** Updated Network N

The backpropagation algorithm is defined at a single sample level, and the learning from a dataset is usually repeated for a number of 'epochs'. As noted though, implementation is often implemented using a vectorized version of the algorithm which either samples the entire dataset at once (batch), or a subset of samples (minibatch). The latter is the Stochastic Gradient Descent (SGD), as discussed in 2.4, which has been shown to increase the speed and stability at which the backpropagation algorithm can converge to a minima in terms of cost. The algorithm runs backpropagation over the entire dataset for the number of epochs, and updates the network incrementally through the epoch. The stopping condition for the algorithm is usually defined as

either a particular number of epochs being reached, or cost no longer decreasing for some number of epochs (i.e. a minima has been reached).

The changes to the initial backprop algorithm, detailed in 2, are relatively minor:

- 1 Selection is now performed such that at each epoch m samples  $x_s$  are selected from x which have not yet been sampled.
- 2 The weight update is now performed such that it represents an aggregate update according the m samples that were selected:

$$w^l \to w^l - \frac{\eta}{m} \sum_x \lambda^{x,l} (a^{x,l-1})^T \tag{31}$$

Online Gradient Descent Where SGD is appropriate and effective for scenarios where the entire dataset is available, Online Gradient Descent (OGD) is applicable for when the model is learning in an online fashion. In this case, the backpropagation is run as defined above in Algorithm 2 but with no repetition (i.e. only 1 epoch).

#### 4.2.5 Regularization

Regularization is a commonly used technique in machine learning used to reduce a model's capacity to overfit to the data, and in so doing reduces the variance in the model results. One of the typical methods, L2 (or "weight decay"), is implemented by adding an extra term to the cost function which is itself a function of the weights in the model. This extra term forces the learning process to favour smaller weights, and only allows large weights to occur if they are able to offer an appropriate increase in performance. The modified cost function is displayed below in 32.

$$C = \frac{1}{2n} \sum_{x} \|y - a^L\|^2 + \frac{\lambda}{2n} \sum_{x} w^2$$
 (32)

The additional term is scaled by the configurable regularization parameter,  $\lambda$ , which if small approximates the original cost function or if large will increase the degree of regularization used.

In order to implement this in backpropagation, the weight update rule is changed to the following (while biases remain the same):

$$w \to w - \eta \frac{\partial C_0}{\partial w} - \frac{\eta \lambda}{n} w \tag{33}$$

For SGD, this can be simplified to the following:

$$w \to \left(1 - \frac{\eta \lambda}{n}\right) w - \frac{\eta}{m} \sum_{x} \frac{\partial C_x}{\partial w}$$
 (34)

#### 4.2.6 Learning Rate Schedule

Learning rates schedules are implemented to allow for a more dynamic exploration of the possible solution space, and allow fine tuning around an optima. With a constant learning rate implementation, one has to choose between a larger learning rate which allows faster progress but less optimisation, or a smaller learning rate which learns slower but is more effective at optimising (by avoiding the learning algorithm from bouncing around a minima valley). Learning rate schedules aim to get the best of both scenarios by cycling through a minimum and maximum range across epochs.

The learning rate schedule is specified to cycle through from min to max every i epochs, from  $\eta_{\min}$  to  $\eta_{\max}$ . Thus for the current epoch T, the learning rate is calculated using 35 below:

$$\eta_t = \eta_{\min}^i + \frac{1}{2} \left( \eta_{\max}^i - \eta_{\min}^i \right) \left( 1 + \cos \left( \frac{T_{\text{current}}}{T_i} \pi \right) \right)$$
 (35)

This is the Cyclical Learning Rates approach, though using sinusoidal form rather than linear [80]. For a minimum learning rate of 0.1, a maximum of 1.0 and an epoch cycle of 100, it will produce learning rates as displayed in figure 3 below.

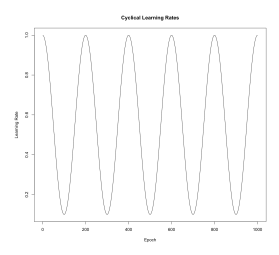


Figure 3: Learning rates calculated over 1000 epochs with  $\eta_{\min} = 0.1$  to  $\eta_{\max} = 1.0$  and i = 100

#### 4.2.7 Dropout

Input layer dropout has been implemented in order to perform regularization on the data features. The methodology used will set a percentage of the features for each sample to 0 during the SGD training process. While the percentage stays the same throughout the testing process, the features selected for each sample are rechosen every time it is used (i.e. for subsequent epochs). Conceptually, the dropout as applied to the input layer may result in the network learning relations between assets and possible patterns that would improve predictive efficacy.

#### 4.3 Restricted Boltzmann Machines

Restricted Boltzmann Machines (RBM) are generative networks which can be trained to learn probability distributions over a dataset. They are structurally different from a FFN in that they have a recurrent weight function - a typical RBM has one visible layer (input/output), and one hidden layer. A sample will be processed from the input layer to the hidden layer, and the activation values from the hidden layer will be used

by the input layer to provide a recontruction. The hidden units thus correspond to feature detection of the visible unit data structures, and the learning process of the network results in effective parameter estimation.

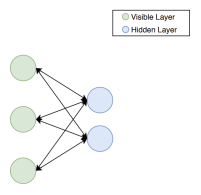


Figure 4: An example diagram of a Restricted Boltzmann Machine network

One of the primary differences from a FFN lies in the stochastic unit determination - the values in a hidden layer will typically be implemented such that the take on a binary value with a probablistic likelihood. Thus, the input and output have the same structure, and the processing from the hidden layer creates the generative process learned by the RBM.

The joint configuration (v, h) of the visible and hidden units has an energy given by:

$$E(v,h) = -\sum_{i \in visible} a_i v_i - \sum_{j \in hidden} b_j h_j - \sum_{i,j} v_i h_j w_i j$$
(36)

where v, h correspond to the binary states of the visible and hidden units, with biases a, b, and weights w [40]. It can be shown, that network weights can be adjusted to change the probabilities assigned to a particular training sample [40]. The derivations of 36 show that performing a stochastic descent for the log probability of the data can be implemented through the following weight adjustment:

$$\Delta w_{ij} = \eta(\langle v_i h_j \rangle_{data} - \langle v_i h_j \rangle_{model}) \tag{37}$$

The angular brackets here indicate expectations under the distribution specified by the following subscript. Due to the probabilistic nature of RBMs, the [0,1] ranged Sigmoid activation function is typically used. Thus, for hidden nodes and a data sample v, it is easy to get unbiased sampling of  $\langle v_i h_j \rangle_{data}$ 

$$p(h_j = 1) = \sigma(w_{0j} + \sum_{i=1} v_i w_{ij})$$
(38)

Similarly, the visible states (or reconstruction), can then be calculated as

$$p(v_i = 1) = \sigma(w_{0i} + \sum_{j=1} h_j w_{ij})$$
(39)

Getting an unbiased sample of  $\langle v_i h_j \rangle$  proves more problematic, and so the model reconstructions via Gibbs sampling are used instead (this is explained below), resulting in the weight updates of

$$\Delta w_{ij} = \eta(\langle v_i h_j \rangle_{data} - \langle v_i h_j \rangle_{reconstruction}) \tag{40}$$

While it's important for the hidden units to take on a binary value (and so avoid communicating real values rather than learning structure), the visible units may be chosen to take on the probability values, rather than the stochastic samples, particularly if real valued output is necessary.

#### 4.3.1 Contrastive Divergence

The process of sampling and resampling may be run for many iterations between the two layers before finishing on an output - this potentially long running and stochastic process results in the generative aspect of the network, and constitutes a Gibbs sampling chain. Multiple sampling steps in this chain is known as Contrastive Divergence, or CD-n, where n represents the number of steps, and which allows for effective parameter estimation. Thus, CD-1 is the following:

#### Algorithm 3: CD-1

**Input:** RBM Network N, with visible nodes N and hidden nodes H and distribution  $\langle vh \rangle$  **Data:** Training sample dataset V'

For a random training sample V', sample H' from P(V|H);

Sample V from P(V|H');

Sample H from P(H|V);

Return  $(V, H) \sim R$ , where R is the reconstructed distribution for  $\langle vh \rangle$ ;

// Using the reconstructed distribution R(V,H), the weight update for CD-1 is, as above, then

$$\Delta CD(W) = \eta(E_{P(H|V)D(V)}[VH^{T}] - E_{R(V,H)}[VH^{T}])$$
(41)

**Result:** Reconstructed distribution for  $\langle vh \rangle$ 

There's no upper bound on the iterations used for CD, and running for many can prove more effective for certain purposes. In this case however, where RBMs are used for the purposes of weight initialization, CD-1 is usually deemed sufficient.

#### 4.3.2 CD-1 and SGD

In the same way that the backpropagation learning algorithm in 4.2.3 can be implemented in an SGD process for FFNs, so can the CD learning algorithm for RBMs. The framework is kept the same, with the implementation of epochs and weight updates based on stochastically chosen minibatches, but the calculation used to update the weights is CD-1 instead.

#### 4.4 Stacked Autoencoders

As noted in 2.3, the use of Stacked Autoencoders (SAE) have resulted in significant improvements in deep learning networks, and allowed effective reduction of high di-

mensional data. A single layer autoencoder is a specialized type of FNN with 3 layers: one input, one hidden, and one output. The network is trained (using backpropagation as per 4.2.3) to reconstruct the input, so the input and output layers have the same structure, and the hidden layer needs to have fewer nodes than the input. This forces the hidden layer to learn effective features of the data, and reduce the dimensional representation.

Stacked autoencoders follow a similar structure, but with multiple hidden layers. The only strict requirement of the hidden layers is that the middle one, which will be used as the encoder layer, still has fewer nodes than the input. This structure can still be trained using backpropagation, but with more layers, it is likely to begin suffering from the vanishing or exploding gradient problem. As noted, the work by Hinton et al. for initialization of weights helps resolve this.

#### 4.4.1 Sigmoid based Greedy Layerwise SAE Training

The steps for implementing the SAE training suggested by Hinton et al [37] is as follows

- 1 Define a network structure which conforms to the requirements of an SAE, with L layers
- 2 For the first hidden layer, train as you would for an RBM with 2 layers with the input layer, and the first layer as the hidden layer, using CD-1 and SGD
- 3 For each layer l in (2, L/2):
  - 3.1 Process the data through the previously trained layers using the forward propagation as defined in 4.2.1
  - 3.2 This processed data then forms the input to the  $l^{th}$  layer, which can be trained once again using CD-1 and SGD as if it were two layers
- 4 Once all the layers up until the encoder layer have been trained in this greedy layerwise fashion, mirror the weights and layers structures after the encoder to create a fully L layered FFN with pre-trained weights
- 5 This network can then be trained using the backpropagation and SGD algorithms, where cost of reproducing the network input is minimised
- 6 Once a minima or acceptable level of reconstruction has been reached, the network can be truncated as the encoder layer, and so the first L/2 layers are used as the SAE

Notably, this weight initialization will only be effective if the RBM and SAE networks use the same activation function, which due to the RBM implementation, needs to be a function that can output a probabilistic value in [0,1].

#### 4.4.2 ReLU based SAE Training

ReLU activations differ from Sigmoid in that they are not fitting for probability estimations, which makes the algorithm suggest by Hinton et al. unsuitable. The process used here relies on effective weight initialization, and is a simplification of the above.

- 1 Define a network structure which conforms to the requirements of an SAE, with L layers
- 2 Use an effective weight initialization, such as Xavier or He (discussed fully in 4.5)

- 3 This network can then be trained using the backpropagation and SGD algorithms, where cost of reproducing the network input is minimised
- 4 Once a minima or acceptable level of reconstruction has been reached, the network can be truncated as the encoder layer, and so the first L/2 layers are used as the SAE

# 4.4.3 Denoising Autoencoders

As noted in 2.3.3, denoising can be used as an optimization technique for autoencoders in order to improve general performance and reconstruction. The methodology works by corrupting the input data for training, but using the non-corrupted data as the expected output sample. In doing so, this forces the autoencoder to learn more fundamental representations of the data rather then fitting to sample noise, hence "denoising". Two of the more typical techniques for achieving this have been implemented, as detailed below. In both cases, the noise is reapplied each time training samples are chosen.

Additive Gaussian Noise With Gaussian noise, samples are corrupted such that a degree of variance is added to the input according to a parameterized Gaussian distribution. In this case, the distribution variance is a configurable training parameter.

$$\tilde{x}|x \sim \mathcal{N}\left(x, \sigma^2 I\right)$$
 (42)

Masking Noise With masking noise, a fraction of the features in the data are set to 0, thus masking them for that sample. The percentage of features chosen is a configurable training parameter.

# 4.5 Variance Based Weight Initializations

Recent research, as noted in [35], has shown that weights can be initialized to maintain expected variance between the input and output layers. These methodologies have the immediate advantages of simpler implementation, as well as faster computation (as no pre-training is required). The also allow for effective weight initialization of non-probabilistic activation functions, such as ReLU. Whether they result in better reconstructions or predictions is less clear (especially as the linearity assumption would prove faulty), and so the methods are tested here as well.

A common initialization heuristic is to use a uniform, and but not layer agnostic, initialization such as 43 below. While simple enough, it's been shown that this does not always lead to the best training results, and can be outperformed by variance based initializations [29].

weights 
$$\sim U\left[-\frac{1}{\sqrt{n}}, \frac{1}{\sqrt{n}}\right]$$
, for a layer with n nodes (43)

### 4.5.1 Initialization Rationale

The variance balancing methodology is based on balancing the variance of a linear network. For input X, with n components and linear neurons with weights W, and

output Y,

$$Y = W_1 X_1 + W_2 X_2 + \dots + X_n W_n \tag{44}$$

It can thus be shown, that for i.i.id samples with mean 0, then

$$Var(Y) = nVar(W_i)Var(X_i)$$
(45)

For the variance of both input X and output Y to be balanced on the forward and backward propagation, then it is necessary for

$$Var(W_i) = \frac{1}{n_{in}} = \frac{1}{n_{out}} \tag{46}$$

In the instance where there are not an equal number of nodes in the two layers, the average can be taken, such that

$$Var(W_i) = \frac{2}{n_{in} + n_{out}} \tag{47}$$

#### 4.5.2 Initializations

**Xavier** Considering that  $Var(U(-a,a)) = a^2/3$ , then ensuring balance variance for weights as per equation (47) provides us with the Xavier Glorot initialization [29] which can be used for Sigmoid activations, and is defined as follows

$$w_{ij} \sim U(-r, r), r = \sqrt{\frac{6}{n_i + n_j}}$$
 (48)

where n is the number of nodes in the  $i^{th}$  or  $j^{th}$  layer.

He The initialization for ReLU is different on accounts of the function being equal to zero for half it's potential input range - in this case it makes sense to double the weight variance, and so the He [35] initialization is used

$$w_{ij} \sim U(-r, r), r = \sqrt{\frac{6}{n_i}} \tag{49}$$

**He-Adjusted** An adjusted initialization is presented here, dubbed 'He-Adjusted', where the ReLU initialization uses a mean of the input and output layers in order to scale the weight variance. For networks with constant layer sizes, the intialization is the same as He, though for SAE networks which have layer size changes by definition, the He-Adjusted initialization will result in more appropriately sized weights.

$$w_{ij} \sim U(-r, r), r = \sqrt{\frac{6}{(n_i + n_j)/2}}$$
 (50)

# 4.6 CSCV & PBO

The combinatorially symmetric cross-validation (CSCV) method developed by Baily et al., as discussed in 2.6.1, can be used to assess the likelihood of backtest overfitting

through comparison of IS and OOS return metrics. Formulaically, the definition of backtest overfitting is given by

$$\sum_{n=1}^{N} E[\overline{r_n}|r \in \Omega_n^*] Prob[r \in \Omega_n^*] \le N/2$$
 (51)

Where the search space  $\Omega$  consists of the N ranked strategies, and their ranked IS performance r and OOS performance  $\bar{r}$ . This allows the PBO, using the Bayesian formula, to be defined as

$$PBO = \sum_{n=1}^{N} Prob[\overline{r} < N/2 | r \in \Omega_n^*] Prob[r \in \Omega_n^*]$$
 (52)

Notably, the above definitions consider IS as the data made available to the strategy selection, rather than the models calibration (e.g. the full IS dataset, rather than, by was of example, the number of days used in a moving average). This allows the model-free and non-parametric nature of the definition.

They further developed the CSCV framework as a methodology to reliably estimate the probability used in PBO, which allows a concrete application of the concept. The CSCV framework does not require using the typical hold-out strategy (and thus avoids credibility issues), and is ultimately able to provide a bootstrapped distribution of OOS performance. The full methodology is shown in algorithm 4 below [4].

#### Algorithm 4: CSCV

**Input:** N configuration trials over T time periods with P&L

- 1 Generate a TxN performance series matrix, M, representing the profits and losses by the N configuration trials over T time periods
- 2 Partition the M matrix by rows into S submatrices, each of even size (T/S\*N)
- 3 Generate the combinations  $C_S$  of  $M_S$ , in groups of size S/2, for total  $\binom{S}{S/2}$  of combinations
- 4 For each combination in  $c \in C_S$ :
  - a Form a training set J by joinined S/2  $M_S$  submatrices, in their original order. J is a matrix of order (T/S)(S/2) \* N
  - b Form the test set  $\bar{J}$  as the complement of J in M, once again in the original order
  - c Form a vector of  $R^c$  of performance statistics of order N, where the Nth component  $R_n^c$  of  $R^c$  reports the performance associated with the  $n^{th}$  column of J
  - d Repeat [c] for  $\bar{J}$  to derive  $\bar{R}^c$  and  $\bar{r}^c$
  - e Determine the element  $n^*$  such that  $R_n^c \in \Omega_n^*$  i.e.  $n^*$  is the best performing strategy IS
  - f Define the relative rank of  $\bar{r}_{n^*}^c$  by  $\bar{\omega}_c := \bar{r}_{n^*}^c/(N+1) \in (0,1)$ . This is the relative rank of the OOS performance associated with the strategy chosen IS, which should systematically outperform OOS if no backtest overfitting has taken place
  - g Define the logit  $\lambda_c = ln \frac{\bar{\omega}_c}{(1-\bar{\omega}_c)}$ . High values here indicate consistency between IS and OOS performances (and so low ovefitting)
- 5 The logit values can now be used to compute the distribution ranks of OOS, by collecting all  $\lambda_c$  for  $c \in C_S$ . The relative frequency for  $\lambda$  occurring across all  $C_S$  is

$$f(\lambda) = \sum_{c \in C_S} \frac{\chi_\lambda(\lambda_c)}{\#(C_S)}$$
 (53)

where  $\chi$  is the characterization function and  $\#(C_S)$  is the number of elements in  $C_S$ , and so  $\int_{-\infty}^{\infty} f(\lambda)d\lambda = 1$ 

**Result:** Reconstructed distribution for  $\langle vh \rangle$ 

The CSCV framework and results thus allows the consideration of several notable statistics. First and foremost, the PBO may now be estimated using the CSCV method and using an integral over the  $f(\lambda)$  function as defined above which offers a rate at which the best IS strategies underperform the median of OOS trials. The PBO is estimated using

$$\Phi = \int_{-\infty}^{0} f(\lambda)d\lambda \tag{54}$$

If  $\Phi \approx 0$ , it is evidence of no significant overfitting (inversely,  $\Phi \approx 1$  would be a sign of probable overfitting). Critically then, a PBO measure may be used in a standard hypothesis test to determine if a model should be rejected or not. This can be extended, as shown by Bailey et al., to show the relationship between overfitting and performance degradation of a strategy. It becomes clear that with models overfitting

to backtest data noise, there comes a point where seeking increased IS performance is detrimental to the goal of improving OOS performance.

# 4.7 Money Management Strategy and Returns

In line with the general approach here, the Money Management Strategy (MMS) has been designed to be relatively simple, such that the results are more indicative of the underlying modelling rather than intricate trading strategies. The MMS follows an arithmetic long strategy of buying any stock for which the predicted t+D price is above the current price, and selling the stock at t+D. Trading costs have been included at 10% capital costs per annum for borrowing to purchase, and 0.45% for transaction costs. Results are compared to a "perfect" model which has exact knowledge of the price in D days, which represents an upper bound on performance.

Input Variables The MMS takes the following values as input for each time point t and prediction point D

- $1 y_t$ , the closing price for t
- 2  $y_{t+D}$ , the closing price in D days
- 3  $\hat{y}_{t+D}$ , the predicted closing price in D days

Calculated Stock Variables Using these, the MMS calculates and uses the following:

1 A flag indicating whether a trade will take place on t

$$trade\_long_t = bool(\hat{y}_{t+5} > y_t) \in (0,1)$$

$$(55)$$

2 A flag indicating whether a trade will take place on t in the perfect model

$$trade\_long_{n,t} = bool(y_{t+5} > y_t) \in (0,1)$$
 (56)

3 The observed return at t + D

$$\mathbf{r}_{o,t+D} = y_{t+D} * trade\_long_t \tag{57}$$

4 The expected return at t + D

$$\mathbf{r}_{e,t+D} = \hat{y}_{t+D} * trade\_long_t \tag{58}$$

5 The perfect return at t + D

$$\mathbf{r}_{p,t+D} = y_{t+D} * trade\_long_{p,t} \tag{59}$$

6 The cost incurred in buying the stock at time t

$$cost_t = y_t * trade\_long_t \tag{60}$$

7 The cost incurred in the perfect model at t

$$cost_{p,t} = y_t * trade\_long_{p,t} \tag{61}$$

8 The possible capital and transaction costs

$$trading\_cost = y_t * (0.1/365 * 2) + y_t * (0.45/100)$$
 (62)

9 The full cost of trading the stock at t in the predictive model

$$fullcost_t = (y_t + trading\_cost_t) * trade\_long_t$$
 (63)

10 The full cost of trading the stock at t in the perfect model

$$fullcost_{p,t} = (y_t + trading\_cost_t) * trade\_long_{p,t}$$
 (64)

Calculated Strategy Variables The MMS is then implemented using these calculated values which are aggregated at each time point t for all stocks, s, as follows:

1 The expected return for t + D from all stocks traded at t

$$R_{e,t+D} = \sum_{s \in stocks} r_{s,e,t+D} \tag{65}$$

2 The observed return for t + D from all stocks traded at t

$$R_{o,t+D} = \sum_{s \in stocks} r_{s,o,t+D} \tag{66}$$

3 The perfect return for t + D from all stocks traded at t

$$R_{p,t+D} = \sum_{s \in stocks} r_{s,p,t+D} \tag{67}$$

4 The cost of all trades at t

$$C_t = \sum_{s \in stocks} cost_{s,t} \tag{68}$$

5 The cost of all trades at t in the perfect model

$$C_{p,t} = \sum_{s \in stocks} cost_{s,p,t} \tag{69}$$

6 The cost of all trades at t including transaction costs

$$FC_t = \sum_{s \in stocks} fullcost_{s,t}$$
 (70)

7 The cost of all trades at t, including transation costs, in the perfect model

$$FC_{p,t} = \sum_{s \in stocks} fullcost_{s,p,t}$$
 (71)

The final model return is thus established as  $\sum_t R_t / \sum_t C_t$ , for any variations of observed, expected or perfect returns as well as base and full costs. These model returns are used to establish the performance of the different predictive models which were trained.

An additional consideration of the MMS is the daily rates generated by the strategy, which incorporate the costs of any trades incurred, as well as any returns seen at t, such that

$$dailyrate_t = R_{t-D} - C_t \tag{72}$$

# 5 Implementation: Process

Having covered the technical implementations in Chapters 3 and 4, a higher level overview of the end to end experimental process can be detailed. The process here rests on two key principles: implementing a generalised version of a system which could offer exploration of more complex techniques, and ensuring an effective modularisation of steps such that the process can be reconfigured accordingly while maintaining its integrity. In doing so, a separable system is created which brings together data reduction, deep learning with pre-training/weight initialization, online learning and back test overfitting validation.

# 5.1 Data Preparation

The fulldataset goes through several steps of pre-processing - these are covered in more detail in 3.1, though the steps are included here:

- 1 The data is processed into day to day log fluctuations
- 2 At each time point, rolling historic fluctuation summations are calculated (e.g. the past 1, 7 and 30 days)
- 3 At each time point, rolling future fluctuation summations are calculated (e.g. the next 2 days)
- 4 The dataset is truncated to only include points with all aggregations or predictions available
- 5 The dataset undergoes scaling the different functions are detailed in 3.1.2, though limited normalize was used for most of the results produced

#### 5.1.1 Data Window Aggregations

The time aggregations were chosen according to domain knowledge - it's expected that prices may move in daily, weekly, fortnightly, monthly and quarterly patterns as trading is actioned by day traders such as speculators to institutional investors such as mutual funds. Indeed, it can be shown that markets can be modelled such that they are hierarchical in space and time, with actors at different heirarchical levels within the market interacting under different causitive structures [91]. The effect is such that price trends and patterns can be expected to exist in time at multiple levels, with price being set as an emergent property of these hierarchies. Taking this into account, variations of 1, 5, 10, 20 and 60 working days were considered throughout the configuration tests, in the following combinations:

- 1. 1,5,20
- 2. 5,20,60
- 3. 10,20,60

#### 5.1.2 Point Predictions

Various configurations were tested in terms of the historic summations used, while the fluctuation predictions were either of 2 or 5 days. These are detailed more extensively in . It's worth noting that the prediction horizon is configurable, and in the case of implementing a more complex trading strategy, one would likely choose to predict multiple points across the horizon in order to to develop a distribution. In this case,

add result

the MMS was kept purposefully simple, and so didn't warrant the more intensive implementation.

One point of concern, is that point predictions at 2 or 5 days in advance could open the process up to aliasing error, as it is unlikely to be predicting at the Nyquist frequency, and may end up fitting to the band ripple rather than the underlying signal. Conversely, 1 day predictions for closing price might offer a high degree of accuracy, but are not practical for trading due to market auction and bidding structures. The system implementation veered on the side of practicality in this case, and was configured for longer term predictions despite the risk of aliasing. While some of the configurations were run with 2 day predictions, it was seen that once costs were considered it was difficult for even the benchmark to produce desired P&L figures and so a 5 day prediction was generally implemented in subsequent tests.

add result references

## 5.1.3 Scaling

When testing the two limited scaling methodologies, standardization and normalization, it was found that the results for standardizing tended to be notably worse than normalization (discussed in 7.1). It's possible that outliers in price fluctuations and changing variance through time (which is not captured effectively through the limited variation) resulted in a less informative representation and worse performance of the FNN models which can be prone to error maximisation. As a result of this, the limited normalization process was used for most of the tests run. It's worth noting that an implementation which incorporates processing the data through the Error Function in order to reduce the impact of outliers would likely prove a worthwhile endevour, and could be easily swapped in due to the systems modularity.

# 5.2 Data Segregation

Once processed, the dataset is split into 2 portions: the first set is used for the SAE training, as well as the initial SGD training on the predictive FFN network; the second set is used only for the OGD training in order to generate returns from the predictive network. In this sense, the first set is used as what would typically be the historical training set, and the second is used as the testing/validation set (the use of Bailey's CSCV technique negates the need for a hold-out portion of the dataset [4]). For the sake of clarity, these two subsets of the dataset will be referred to as the 'Training' and 'Prediction' datasets.

In this case, a simple split point of 60%/40% was chosen for the Training and Prediction sets respectively. In a more restrictive case, where there is less data to work with, it would be advised to consider a method such as MinBTL as developed by Bailey et al. [5] and the expected number of configurations to be tested in order to decide an appropriate split point.

# 5.3 SAE Training

The Training dataset is used to train the autoencoder network using either RBM pretraining or weight initialization algorithms and SGD training, as defined in 4.4. The Training portion of the dataset, as noted in 5.2, is itself split up into a training and validation set for the SAE. The efficacy of the SAE on the validation set is required in order to determine which networks to choose for subsequent steps in the process. The full testing process here will rely on a generated set of best SAE networks at each encoding layer size, to be used for FFN training and prediction (chosen by a minimum MSE score). The benefit of the modularised system is emphasised here, as the SAE training will not suffer from limitations due to backtesting considerations: any amount of configurations or processes can be tested for feature extraction without concern.

Once the SAE networks have been defined and chosen, they can be used to reprocess both the Training and Prediction datasets such that the input is encoded, and the output is as before. These encoded datasets can then be used for the following steps in the process.

In a productionized system, where data is updated daily, there would be a set period after which the SAE network would be retrained to reflect more recent data. While the system detailed here aims to emulate as much, it wasn't necessary to incorporate this step with the static dataset.

# 5.4 Prediction Network Training

Typically, the SAE and predictive network might be presented as one and the same, however in the process presented we are able to separate the two and optimise for feature extraction prior to training for predictions. Having done so, the datasets can have their input encoded, but retain the same output. The predictive network is then trained using the Training set, as detailed in 4.2.3. There's no requirement to validate this part of the training, so the dataset is not split into subportions as it was for SAE training.

Once the predictive network is trained, the OGD process is run through the encoded Prediction dataset in order to generate the predictions for the assets prices that the model produces - thus emulating what would have occured in a live environment.

#### 5.5 Price Reconstruction

In the interest of more interestable results for figures such as P&L, the prices are fluctuations are re-scaled back to their original price fluctuation predictions using the reverse scaling detailed in 3.1.3. These prices are then recostructed using 3.1.4, such that they represent a predicted future price of the asset which can then be used by the MMS. It's worth nothing that the price reconstruction applies the predicted fluctuation to the known original price, such that it would represent a system that is run daily, rather then a system that is making rolling predictions over a period of time.

# 5.6 Money Management Strategy

The MMS, as described in 4.7, takes a proof of concept approach to the system being investigated. It is a simple long strategy with a set liquidation point, and no notable refinement. The naive approach is taken purposefully here, so as not to bias the perspective of the system as a whole by the effects of an impactful trading strategy.

A more effective trading strategy would incorporate a shorting side as well, though ultimately this isn't necessary in order to obtain a necessary indication of efficacy for the system (and conceptually, a consistently losing long strategy would be of interest in any case).

It is important, in the interest of effective optimization and development, that the pattern recognition in the prediction portion of the system is not tightly coupled with making it profitable. With this in mind, the modularity of the system is continued, with a distinct separation between the prediction signal and the MMS implementation which relies only on the predicted prices of each stock for a time point. The comparisons for the MMS are made against a benchmark of the same MMS, but with perfect information. This allows an indication for how effective the prediction system is (and need be) in order to generate comparable returns.

An effective production trading strategy would be likely to include stop losses or pyramiding stop losses for effective P&L optimization. An ideal implementation, and possible area for further work, would include an MMS strategy where the bands for this are defined using a machine learning technique (rather than being chosen from domain knowledge or heuristics).

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# 5.7 CSCV & PBO

The CSCV and PBO parts of the system are the most straightforward, in the sense that they are mostly just an implementation of the algorithm detailed in 4.6. The CSCV process uses the returns from the MMS, which in turn used the prices from the predictions network. Conceptually, the whole system comes into place here, as the results from the CSCV process are now indicative of not only backtest overfitting in the trading strategy, but also in the prediction network and without having to consider the impact of many configuration tests for feature extraction.

# 5.8 Process Diagram

The flow diagram below provides a graphical representation of this process from the raw market data up until the price prediction points (the MMS calculations and CSCV process are not included). The process may be repeated accordingly in terms of SAE structures that are desired for testing, such that the "Best SAE Network" is different for different encoding layer sizes, or data aggregations used (as was the case in many of the experiments which will be discussed in 7).

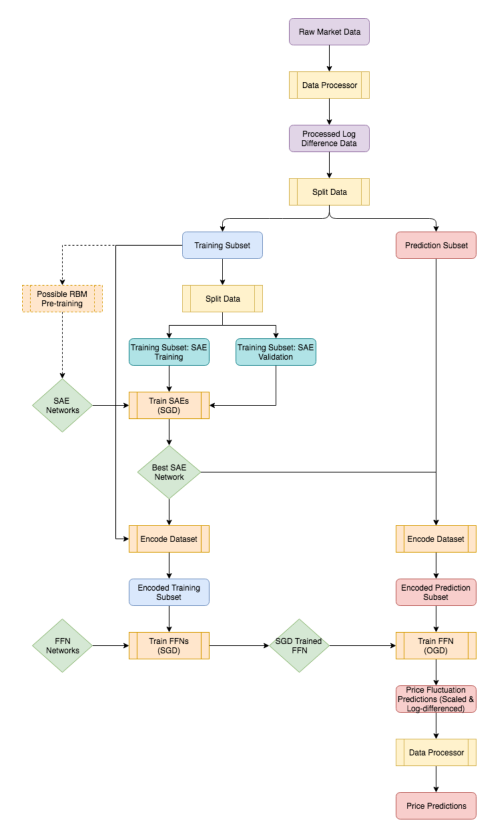


Figure 5: Overall Process Flow

# 6 Datasets Used

# 6.1 Synthetic Datasets

Data was generated and scaled, as per the methods detail in 3.2 and 3.1.2. The sets were generated with the following configurations, each for 5000 timesteps and with a 60/40 split on the Training and Prediction sets.

## 6.1.1 Synthetic6

The Synthetic6 dataset was configured to represent a combination of upwards, downwards and sideways Trends, each with high and low variance variations.

Trend Category	Variance Category	Trend Mean	Variance
Strong Upward	High	0.9	0.5
Strong Upward	Low	0.9	0.2
Upward	High	0.05	0.4
Upward	Low	0.05	0.1
Strong Downward	High	-0.8	0.55
Strong Downward	Low	-0.8	0.15

Table 1: Synthetic 6 Dataset Configuration

### 6.1.2 Synthetic10

The Synthetic 10 dataset was configured to represent a wide array of behaviours, including very high (positive and negative) mean stocks, as well as much lower mean stocks (which may be more representative of typical behaviour). These were chosen to make sure the network learning is able to differentiate and correctly learn across different asset categories.

Trend Category	Variance Category	Trend Mean	Variance
Strong Upward	High	0.9	0.5
Strong Upward	Low	0.7	0.2
Upward	High	0.05	0.5
Upward	High	0.05	0.4
Upward	Low	0.04	0.1
Sideways	Low	0.02	0.15
Sideways	Low	0.01	0.05
Downwards	Low	-0.1	0.2
Strong Downward	Low	-0.4	0.15
Strong Downward	High	-0.8	0.55

Table 2: Synthetic 10 Dataset Configuration

This set was configured erroneously with the excessively high means.

# 6.2 Actual Datasets

Several datasets have been used using JSE closing price relative data for 2003-2018 . They were processed the same way as the Synthetic sets, following the steps set out in 3 and with a 60/40 split on the Training/Prediction datasets.

add ref

### 6.2.1 Actual10

This is the primary closing price dataset that has been used through out the experiment process, which focused on choosing more prominent stocks from multiple sectors.

Code	Company	Sector	
AGL	Anglo American	Resources	
BIL	BHP Billigton	Resources	
IMP	Impala Platinum Holdings Resources		
FSR	FirstRand Limited	Finance	
SBK	Standard Bank	Finance	
REM	Remgro Limited	Finance	
INP	Investec Finance		
SNH	Steinhoff International Holdings	Retail	
MTN	MTN	Communication Services	
DDT	Dimension Data	Tech	

Table 3: Sytnetic 10 Dataset Configuration

# 6.2.2 AGL

Using AGL data from JSE close price relatives.

# 6.2.3 AGL&ACL

Using AGL and ACL data from JSE close price relatives.

# 6.2.4 Scaling10

Using the following assets from JSE close price relatives:

• ACL, AGL, AMS, AOD, BAW, BIL, BVT, CFR, CRH, DDT

# 7 Results

An introduction to the results, highlighting the important findings and sections.

# 7.1 Linearity, Complexity and Structure of Data

#### 7.1.1 GBM Generated Data

Geometric Brownian Motion (GBM), as discussed in 3.2, has been used to simulate synthetic datasets for the purposes of testing implementations and configurations. GBM is a popular choice for synthetic financial data as it is a Markov process, thus following a random walk and is generally consistent with the efficient market hypothesis in that the next price movements is conditionally independent of past movements. In line with this though, the series will exhibit a constant drift with price shocks according to it's stationary configuration, and changes in price that are not independent. The effect seen, and discussed more below, is that reconstruction of aggregated GBM data was able to performend quite effectively through linear activations, so long as the network (and encoding layers) were large enough to allow for sufficient transformations. Linear activations weren't tested further on actual data, as there is no reason to believe the time series would continue to exhibit the constant drift with nonindependent price jumps over time. Further to this point, it is worth pointing out that as GBM are non ergodic series, one should be wary of considering ensemble based predictions with much confidence [68]. So while the GBM data was used for assessment of some prediction networks, these results should be considered contextually.

## 7.1.2 Activations: Linear, Sigmoid, ReLU and Leaky ReLU

In line with the characteristics of the synthetic data noted in 7.1.1, results for networks using a Linear Activation often outperformed the networks using non-linear activations such as ReLU or Leaky ReLU (activation functions are detailed in 4.2.2).

While these configurations were run to generate a comparison of Linear and ReLU activations for the SAE networks, it is typical to use a Linear Activation in the encoding layer of an SAE [37], and the linear output activation has mostly been implemented here to assist with the shortcomings of the scaling methodology, as noted in 3.1.2.

SAE Activations and Scaling In figures 6 and 7 below, the effects of different activations and scaling methods can be seen when applied to the SAE using actual stock data. The MSE results offer several takeaways: using standardizing scaling instead of normalizing results in far worse performance; Sigmoid activations are notably worse than the linear or ReLU activations and linear output layers offer better results than ReLU output layers.

Networks which use only linear activations show very competitive performance overall, though the ReLU activation shows much better performance than linear activations when the encoding layer size is decreased (figures 8 and 9 below), thus forcing the network to take advantage of non-linear effects. Sigmoid activations, while also able to take advantage of non-linear effects, show much worse performance. Sigmoid activations are known to suffer from learning slow down as a result of saturation and so have increased sensitivity to vanishing gradients [30]. SAE networks are deep by

nature, and so it is not surprising that the Sigmoid activations are resulting in worse performance over the same training period.

As discussed more fully in 5.1, the use of standardization for scaling the data doesn't allow for effective outlier treatments, resulting in worse performance. Again, this could be better treated with use of the Error function.

### Effects of Scaling and Activation Functions on SAE MSE Score (Figure 1/2)

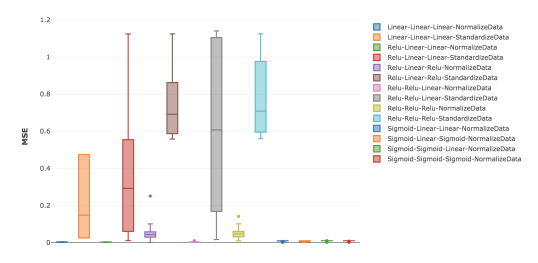


Figure 6: Scaling10 dataset (6.2.4); Configuration1 (9.2.1) - 1316 Samples

Each configuration group is labelled in the following format: 'Hidden Activation-Encoding Activation-Output Activation-Scaling Technique. The MSE scores show significantly poorer performance when Standardizing is used instead of Normalizing or when there is a ReLU output activation. These configurations are excluded from figure 7.

# Effects of Scaling and Activation Functions on SAE MSE Score (Figure 2/2)

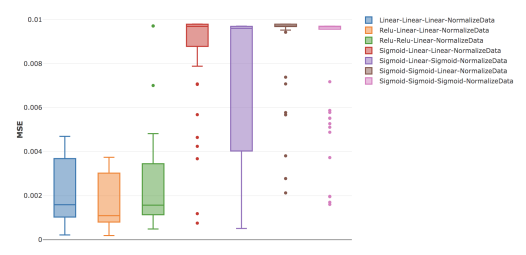


Figure 7: Scaling10 dataset (6.2.4); Configuration1 (9.2.1) - 504 Samples

Excluding the Standardizing scaling combinations, we can now see that Sigmoid has largely poor performance unless there is a Linear encoding layer (a commonly experienced behaviour [37]]), and seems mostly unable to outperform a fully linear network. The best configuration is Relu Hidden layers, with Linear encoding and Output - this makes sense with the non-linear benefit at hidden layers but with less loss of error signal and information at the output and encoding layers.

## Non-linear Benefit in Encoding Layer (Figure 1/2)

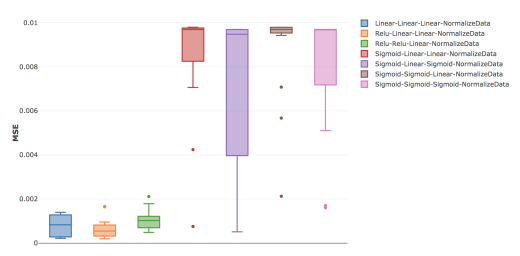


Figure 8: Scaling10 dataset (6.2.4); Configuration1 (9.2.1) - 168 Samples

These plots show the performance for all configurations with an encoding layer size of 25 (input 30). The fully linear networks show good performance here, where there is greater scope for linear representation in the encoding.

#### 0.01 Linear-Linear-NormalizeData Relu-Linear-Linear-NormalizeData Relu-Relu-Linear-NormalizeData 0.009 Sigmoid-Linear-Linear-NormalizeData Sigmoid-Linear-Sigmoid-NormalizeData Sigmoid-Sigmoid-Linear-NormalizeData 0.008 Sigmoid-Sigmoid-NormalizeData 0.007 MSE 0.006 0.005 0.003 0.002

# Non-linear Benefit in Encoding Layer (Figure 2/2)

Figure 9: Scaling10 dataset (6.2.4); Configuration1 (9.2.1) - 168 Samples

These plots show the performance for all configurations with an encoding layer size of 5 (input 30). Here we now see the benefit of non-linear activations in the ReLU based networks, which the fully linear is not able to outperform.

Predictive FFN Activations and Scaling In figures 10 and 11, a comparison of linear and ReLU activations are made for synthetic data in both smaller and larger sized networks. The linear activations result in notably better P&L when compared to ReLU activations, though unlike the SAE networks, this persists even when the network size is decreased - this difference highlights the effects of GBM data and that it can be represented linearly, as per 7.1.1, whereas we would see actual stock data to benefit from a non-linear representation. Linear configurations for actual data were not run, as real stock data is not expected to follow patterns that are linear through time, thus offering little value.

### **Activations in Large FFN Predictive Networks**

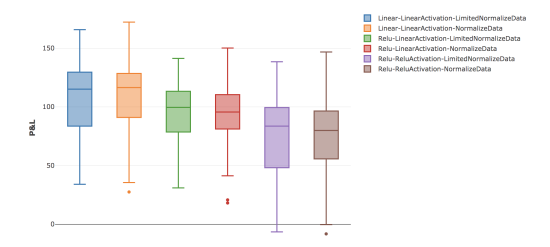


Figure 10: Dataset Synthetic6 (6.1.1); Configuration2 (9.2.2) - 720 Configurations

The above groupings show the P&L generated for different combinations of Hidden Activation - Output Activation - Scaling Method (which is for both SAE and FFN). For an input of 18, and network sizes of 40 and 80 with hidden layers of 1 an 3, we see the linear activations showing notably better performance than the ReLU activations.

format prop-

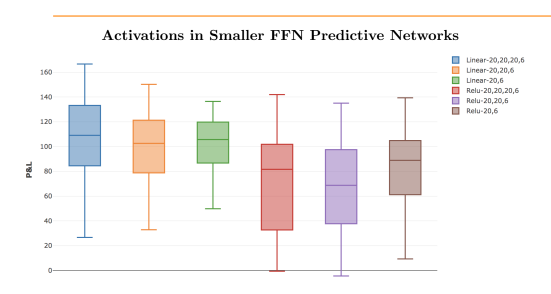


Figure 11: Dataset Synthetic<br/>6(6.1.1); Configuration<br/>2(9.2.2)- 720 Configurations

The boxplots here show the P&L generated for networks with smaller layer sizes, as indicated by the group naming (each number represents a hidden layer and its node size). The networks were trained on the Synthetic6 dataset with 18 inputs, different sized autoencoders and learning rates. We once again see the outperformance of ReLU by the linear activations.

Leaky ReLU vs ReLU Leaky ReLU activations were implemented, and as can be expected, do show a small improvement on both the SAE and predictive FFN networks when compared to ReLU. These effects are show for the Synthetic6 dataset in in figures 12 and 13 below.

# SAE: Leaky ReLU vs ReLU

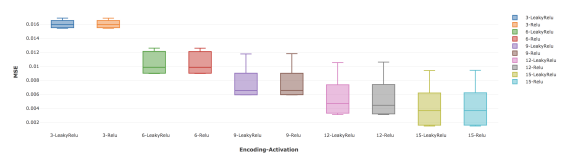


Figure 12: Dataset Synthetic6 (6.1.1); Configuration - 120 configurations

The plot above shows the SAE MSE, grouped by encoding size and activations, showing a marginal increase in performance for Leaky ReLU activations.

#### FFN: Leaky ReLU vs ReLU

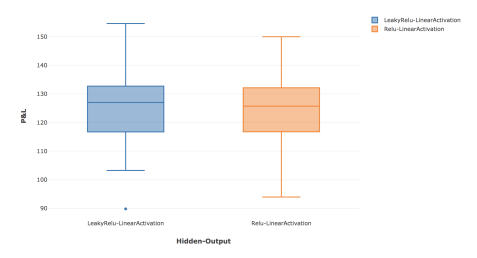


Figure 13: Dataset Synthetic6 (6.1.1); Configuration - 80 configurations

The plot shows the FFN P&L, grouped by activation, showing some improvements from the Leaky ReLU activation.

# 7.2 Weight Initialization Techniques

# 7.2.1 RBM Pretraining for Sigmoid Networks

While previously considered the best approach for training deep neural networks, the methodology of greedy layerwise RBM pre-training for Sigmoid SAE networks (as de-

scribed in [37]) had detrimental effects on network performance. Experiments assessed the effects of running the pre-training for 0, 1, 10 and 50 epochs, where there is a clear decrease in performance as the number of epochs increase (the low value outliers that can be seen for the 1 epoch configuration were with learning rates which were low enough to approximate no epochs).

The RBM pre-training technique does assume that data is Independent and Identically Distributed (IID), and does ultimately traverse a different solution space and loss function. While it may be effective in some contexts, it was shown to result in a counterproductive exploration of the weight space for this non-IID dataset, and that ultimately the financial time series data is pathological for RBM pre-training. It may also emphasise the value of more recent data over historical data usage, as discussed more fully in .

add result ref

#### Pre-training Effects on SAE MSE Scores

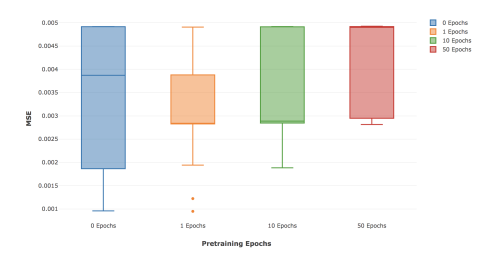


Figure 14: Dataset AGL&ACL (6.2.3); Configuration 7 (9.2.7)

The boxplots here show the summary of configuration performances, by minimum MSE achieved, grouped according to the number of pre-training epochs which the network had. There is a clear favour to having no pre-training in this scenario.

**Sigmoid Activation Functions** Due to the poor performance seen in Sigmoid function based SAE's here, as well as the poorer results when compared to ReLU activations (noted in 7.1.2), Sigmoid functions were largely excluded from further configuration testing.

### 7.2.2 Variance Based Weight Initialization Techniques

More recent research has focused on the use of weight initialization using variance based methodologies, as discussed fully in 4.5. Our expectation, theoretically, is that the He initialization will generally outperform Xavier due to it being more appropriate for the ReLU activations being used. Additionally, in networks with varying layer sizes, we expect He-Adj and possibly Xavier to outperform He which is subject to imbalanced initializations. Ultimately, He-Adj as presented in 4.5, should present the

best of both for an initialization that is suited to Leaky ReLU activations and the varying layer sizes found in SAE networks.

Training SAE networks across the Synthetic10 (6.1.2), Actual10 (6.2.1) and AGL (6.2.2) datasets, we mostly see the expected patterns emerge. He-Adj consistently outperforms He due to being better suited to the network structures, as seen in figures 15, 16 and 17 below. We also see that He-Adj clearly outperforms Xavier for actual data in figures 16 and 17, but has performance that is mostly the same (or even marginally worse) than Xavier for the synthetic dataset in figure 15.

As discussed more fully in 7.1, the GBM data in synthetic datasets will have assets that are configured to certain mean and variances. As these movements are aggregated over time through the data processing, the movements will generally be more similar and less complex than those of actual stock data which will show greater variance over time. The learning process then is not required to be able to pass through error signals that change as dynamically over time, and so there is less pressure on the initial starting weights. Additionally, He and He-Adj have a higher upper bound in the values used for Uniform distribution initialization than Xavier, and so are less likely to result in dying ReLU nodes from early learnings that are less representative of longer term patterns and so become damaging to final network performance. Further, some of the assumptions of these techniques, such as data being IID may not apply in the first place, making their results less predictable. These effects result in the outperformance of He-Adj in the networks trained on actual datasets, but similar performance of He-Adj and Xavier in the networks trained on synthetic data.

# Weight Initialization for SAE MSE - Synthetic10

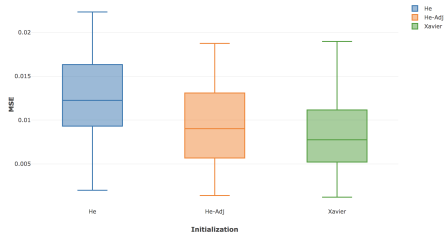


Figure 15: Dataset: Synthetic10 (6.1.2); Configuration 5 (9.2.5)
The box plots show the MSE for a series of SAE networks trained, showing the outperformance of He-Adj compared to He in networks with changing layer sizes, and the similar performance of He-Adj and Xavier on synthetic datasets.

#### Weight Initialization for SAE MSE - Actual 10 $\,$

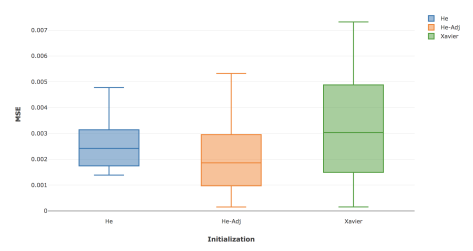


Figure 16: Dataset Actual10

The box plots show the MSE for a series of SAE networks trained, showing the outperformance of He-Adj compared to He in networks with changing layer sizes, and the outperformance of He-Adj to Xavier on actual datasets.

### Weight Initialization for SAE MSE - AGL

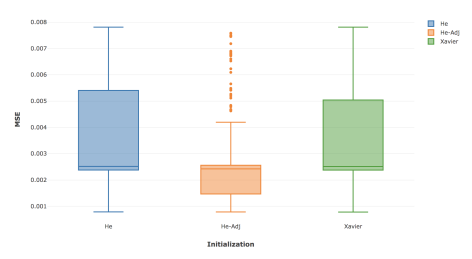


Figure 17: Dataset: AGL (6.2.2); Configuration 3 (9.2.3)

The configurations run for AGL show notably better performance using the He-Adj based initialization in comparison to both He and Xavier. The emphasis seen here can be attributed to increased pressure on suitable weight initizliations as 1 and 2 node encoding layers were used.

When considering the predictive FFN networks, with more consistent layer sizes, the performance between He and He-Adj becomes comparable, with both outperforming Xavier (as expected, with ReLU activations). Due to the generally better or comparable performance of He-Adj, it was the only initialization chosen for the Actual10 dataset, and so only AGL is presented for the predictive networks below.

## Weight Initialization for Predictive FFN P&L- Synthetic10

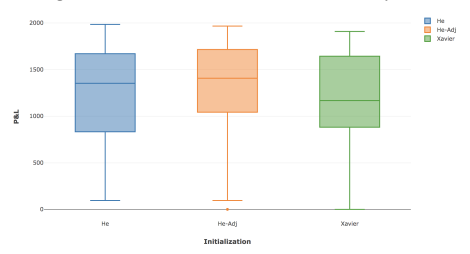


Figure 18: Dataset: Synthetic10 (6.1.2); Configurations 5 and 6 (9.2.5, 9.2.6)

The figure here shows the P&L performance for He-Adj and He being higher than Xavier, as to be expected with more balanced network layers and ReLU activations. The better performance in the lower end of the He-Adj and configurations when compared to He can be attributed to the inclusion of inconsistently sized layers in the configuration set, where He-Adj and would have performed better.

### Weight Initialization for Predictive FFN P&L- AGL

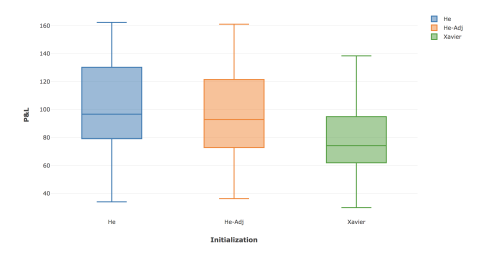


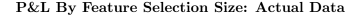
Figure 19: Dataset: AGL (6.2.2); Configuration 4 (9.2.4)

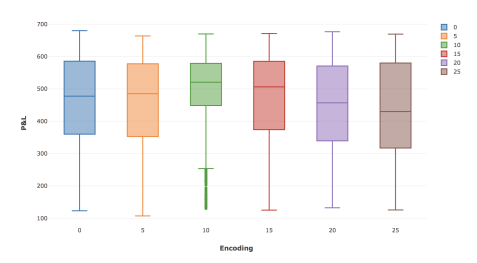
The configurations run for AGL show notably better performance using the He-Adj and He based initialization in comparison Xavier when using actual data, as per the differences explained above.

# 7.3 Feature Selection

The figures below show the efficacy of the different feature selection sizes from the SAE and how they impact P&L in the predictive FFN networks (i.e. the size of the SAE encoding layer). It's apparent in the results for the real data show in Figure 20 that feature selection is occuring, with the optimal size being 10, around the degrees of freedom (or number of assets). Notably, this has the highest mean P&L and is significantly above the '0' group which represents no encoding being performed (i.e. no SAE process prior to the predictive network). This pattern is less clear in the synthetic results shown in Figure 21, where the mean mostly trends upwards as the feature selection size increases, though the gradient of increase decreases notably after 10.

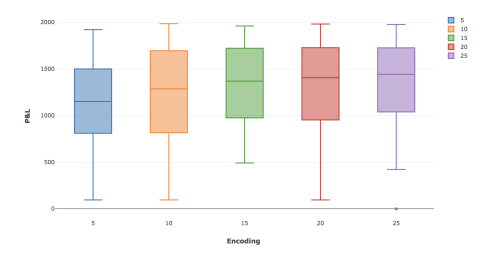
I'm unsure of the synthetic data explanation for this





**Figure 20:** Dataset: ; Configuration Actual P&L grouped according to feature selection size (where 0 is no encoding), showing increased performance the closer feature selection is to the number of assets (in this case, 10).

# ${\bf P\&L}$ By Feature Selection Size: Synthetic Data



**Figure 21:** Dataset: Synthetic10 (6.1.2); Configurations 5 & 6 (9.2.5, 9.2.6) Synthetic P&L grouped according to feature selection size.

Not updated from here

# 7.4 Network Structure and Training

#### 7.4.1 Effects of Network Size

The effects of networks size on SAE MSE performance and the P&L predictive FFN performance are shown in figures 22 and 23 below. They show a general increase in performance for both, with improvements from increases in the sizes and numbers of layers in the networks. The classical SAE structure of decreasing layer sizes results in worse performance - it's possible that the move away from greedy layerwise training nullifies the approach of iterative feature reduction, and that with the variance based weight initializations there is a more typical increase in performance with larger networks. Similar trends were seen in the results for the Synthetic10 dataset (6.1.2), which can be seen in the appendix (9.1.3).

Results were generated using the AGL dataset (6.2.2) and Configurations 3 and 4 (9.2.3, 9.2.4) for the SAEs and predictive FFNs respectively.

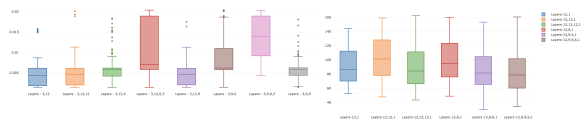


Figure 22: Performance of AGL SAE networks by size

This figure shows the MSE performance of the SAE networks for AGL. The scores show improved results for larger networks (i.e. the '9,9,9' and '12,12' sized networks have the lowest scores). The results also show much worse results for the networks in the networks with gradually decreasing layer sizes.

Figure 23: Performance of AGL predictive FFN networks by size

The plot above shows the P&L for AGL predictive FFN networks. Similarly, there is increased preformance for networks of larger sizes, and no notable benefits of a decreasing layer size structure.

#### 7.4.2 Learning Rate Schedules

Learning rate schedules were implemented such that the learning rate would rise and fall through a set of values during a set number of epochs, as discussed in ??. Conceptually, this should allow a more effective traversal of the solution space, where larger learning rates are able to dislodge the configuration from saddle points, and smaller learning rates are able to optimize at a minima. The schedules presented here have a constant minimum learning rate of 0.00001, with an adjusted maximum learning rate as presented.

Results were generated using the AGL dataset (6.2.2) with Configurations 3 and 4 (9.2.3, 9.2.4) for the SAEs and predictive FFNs respectively.

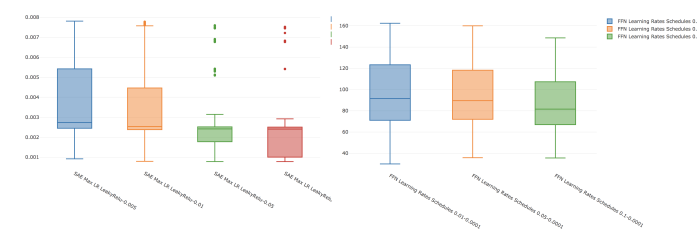


Figure 24: Effects of Learning Rate Schedule on SAE MSE

The figure here shows the improvements in MSE scores gained from increasing the upper bound on the learning rate schedule for SAE configurations, with a notably better performance from the largest upper bound.

Figure 25: Effects of Learning Rate Schedule on Predictive FFN P&L

Unlike the SAE configurations, the predictive FFN performance deteriorates as the upper bound is increased, with the largest upper bound having significantly worse P&L results.

The results here highlight the different nature of the SAE and predictive networks and the solution space they inhabit. The feature reduction performed by the SAE is expectedly complex, and so benefits from a more dynamic learning process. Conversely, the predictive network seems to take on martingale properties which would not require as complex searching through the solution space.

Need to address the validity of this claim. Also, is there any need to include synthetic results here? (they

# 7.4.3 Regularization

L1 Regularization was implemented and tested on a selection of 10 real assets, as per below.

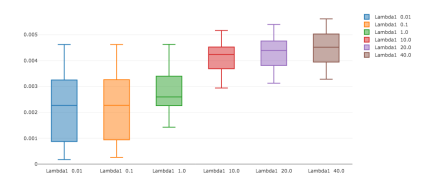


Figure 26: SAE L1 Regularization - 10 Real Asset

The plot above shows the MSE for 360 different SAE networks, grouped by L1 Lambda, showing notable degradation as regularization is increased. Synthetic data tests showed no performance changes below 0.01.

This still contrasts with the learning rate schedule. It could perhaps be the nature of L1 vs L2 regularizat

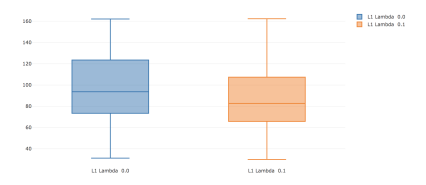


Figure 27: L1 Regularization - AGL Predictive FFN

Similarly, performance decreases for the AGL predictive FFN as regularization increases.

# 7.4.4 Denoising

Denoising tests so far don't seem worth including, though should be considered for future experiments.

# 7.5 Data Aggregation and Selection

# 7.5.0.1 Data Aggregation

Input data was scaled to 3 different configurations across a series of tests in order to assess the effect of shorter and longer configurations on both SAE MSE performance, as well as the predictive FFN P&L results. The configurations tested (in trading day window periods) were:

• 1, 5, 20

- 5, 20, 60
- 10, 20, 60

Results highlighted the differences between the synthetic and actual data in terms of stationarity, and the benefit of more real-time data when it comes to actual stocks.

Results were generated using the AGL dataset (6.2.2) with Configurations 3 and 4 (9.2.3, 9.2.4) for the SAEs and predictive FFNs respectively, and the Synthetic10 dataset (6.1.2) with Configurations 5 and 6 (9.2.5, 9.2.6) for the SAEs and predictive FFNs respectively.

The results for SAE MSE are not totally clear at the moment. It may be more apparent with a larger nur

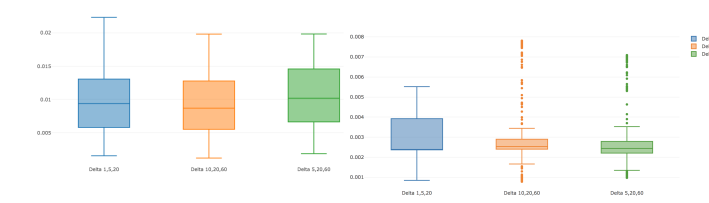


Figure 28: SAE Data Window Aggregations - Synthetic Data

The box plots show the MSE for a series of SAE networks trained, grouped by different data window aggregations. The minimum and median MSE is lower for the higher aggregations, suggesting that the reduced noise in the stationary feature set may support longer windows in further training.

# Figure 29: SAE Data Window Aggregations - AGL Data

For the AGL dataset, the minimum MSE score was found in the '10,20,60' aggregated dataset, though this configuration also results in the worst MSE scores and highest median, suggesting that there is more difficultly in a non-stationary dataset.

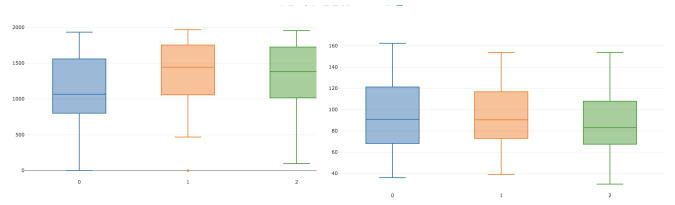


Figure 30: FFN Data Window Aggregations - Synthetic Data

The figure here shows the P&L for the series of FFN networks trained, grouped by different data window aggregations. The profit is higher for the longer aggregations, which is in line with expectations, where the network is able to learn a mean for the synthetic GBM data.

Figure 31: FFN Data Window Aggregations - AGL Data

The figure here shows a notably higher P&L for datasets with a 1 day aggregation, highlighting again the non-stationarity of the real data, and thus the benefit of more accurate recent data.

### 7.5.0.2 Historical Data and SGD training

Tests were run to determine the effect of excluding data from the SGD training of the predictive network. It should be noted that these predictive networks make use of SAEs that were trained on 80% of the data regardless, which may prime the input of data to the predictive network despite the portion of data being excluded from SGD training.

The number of epochs run in the SGD training were not adjusted according to the amount of data used, so ultimately the smaller dataset configurations were trained on fewer samples. It would seem that the use of historical data or the more extensive training is beneficial, but not the primary factor in determining P&L, as decreases in profit were certainly not proportional to the decrease in the period of data.

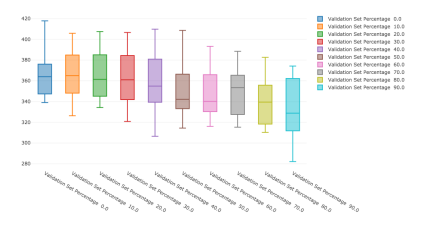


Figure 32: SGD Training Dataset Size - 6 Synthetic Assets

The plot above shows the P&L for FFN predictive networks, grouped by the percentage of data excluded from the SGD training. While there is a decrease in P&L, it is much less than the decrease in data, and may only be a result of the reduced samples used to train (due to a constant number of SGD epochs)

Results were generated using Configuration2 (9.2.2) for the Synthetic6 dataset (6.1.1).

The training for this also used the first X% of data, rather than the last X% in the SAE/SGD set, which is The tests for this are quite extensive - not sure how to balance that with PBO for the actual data.

Data aggregation results could be down to martingale effect.

# 7.6 MMS Results

# 7.7 Results Summary

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# 8 Conclusions

# 9 Appendix

# 9.1 Additional Results

# 9.1.1 AGLEncoding Layer Results

P&L By Feature Reduction Encoding Layer Size: AGL

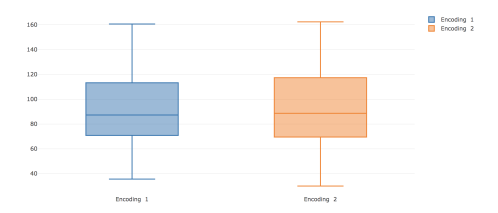


Figure 33: Dataset: AGL (6.2.2); Configurations 3 & 4 (9.2.3, 9.2.4)
This figure shows the effects of the difference encoding layer sizes on the P&L when the relevant SAE is used for the predictive FFN, as for the AGL dataset (with an input of 3 features). Performance for both sizes are similar, though increasing the number of assets may result in a different effect being seen.

# 9.1.2 Effects of Network Size on SAE and FFN for Synthetic Data

# Weight Initialization for SAE MSE - Synthetic10, Encoding Size 5

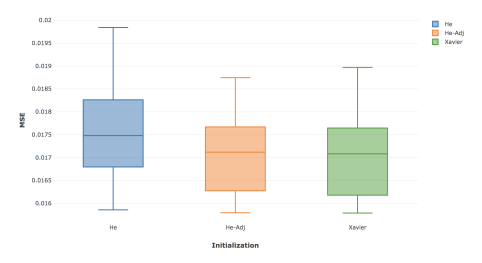


Figure 34: Dataset: Synthetic10 (6.1.2); Configuration 5 (9.2.5)

# Weight Initialization for SAE MSE - Synthetic10, Encoding Size 25

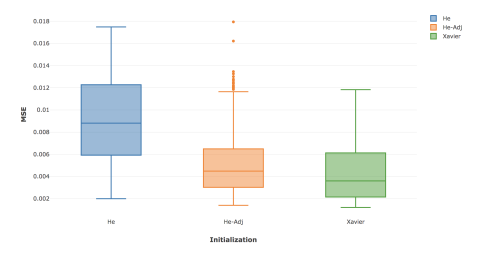


Figure 35: Dataset: Synthetic10 (6.1.2); Configuration 5 (9.2.5)

# 9.1.3 Effects of Network Size on SAE and FFN for Synthetic Data

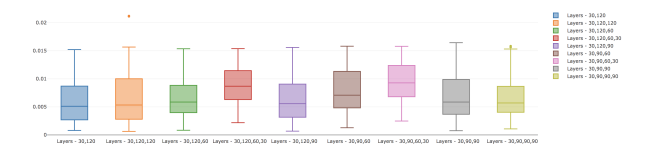


Figure 36: Network Sizes for SAE on the Synthetic10 dataset

The box plots show the MSE for the series of SAE networks trained. The results show worse performance for the typical descending network sizes used for SAE, and show no improvement form increasing network size and complexity.

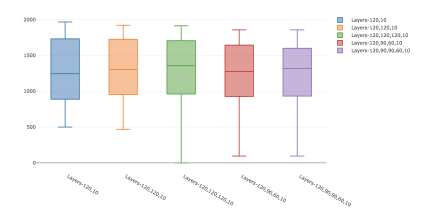


Figure 37: Network Sizes for Predictive FFN on the Synthetic10 dataset

The box plots show the profits for the series of FFN networks trained, showing small improvements for network size increases.

# 9.2 Configuration Sets Used

# 9.2.1 Configuration1 - SAE

- Data: Closing Prices for ACL, AGL, AMS, AOD, BAW, BIL, BVT, CFR, CRH, DDT
- Data Windows: 1, 5, 20 (thus input of 30)
- Data Scaling: Normalize, Standardize
- Weight Init,: Xavier Glorot Uniform
- All Layer Encodings: Sigmoid, ReLU, Linear (split by hidden, encoding and output)

- Encoding Layer Size: 5, 15, 25
- Hidden Layer Size: 60, 120
- Number of Hidden Layers: 1, 2, 3
- Learning Rates: Various for each different activation type (chosen by hidden activation used)

## 9.2.2 Config2 - Predictive FFN

- SAEs: Best performance (lowest MSE) of encoding sizes 3, 6, 9, 12, 15 (input 18) from Config1, of both Limited Scaling and typical Scaling methods
- Output Activations: ReLU, Linear
- Hidden Activations: ReLU, Linear
- Layer Sizes: 40, 80
- Number of Layers: 1, 3
- Various Learning rates for SGD & OGD training

#### 9.2.3 Configuration3 - SAE

- Data Windows: [1,5,20], [5,20,60], [10,20,60]
- Data Scaling: Normalize
- Weight Init: Xavier Glorot Uniform, He, DC
- Layer Encodings: Leaky ReLU with Linear encoding and output
- Encoding Layer Size: 1, 2
- Network Sizes: (12,6), (12), (12,12), (12,9), (9,6), (9,6,3), (12,6,3), (9,9,9), (9,9)
- Learning Rates: Scheduled from (0.005, 0.01, 0.05, 0.1) to 0.0001
- Learning Rate Epoch Schedules: (100,300)
- SGD Epochs: 400
- Minibatch Size: 20
- SAE Validation Split: (0.8)

#### 9.2.4 Configuration 4 - Predictive FFN

- SAEs: Best networks by encoding size and data aggregations from Configuration 3.
- Data Windows: [1,5,20], [5,20,60], [10,20,60]
- Data Scaling: Normalize
- Weight Init: Xavier Glorot Uniform, He, DC
- Layer Encodings: Leaky ReLU with Linear output
- Network Sizes: (12, 12), (12, 12, 12), (12, 9, 9, 6), (12, 9, 6), (12)
- Learning Rates: Scheduled from (0.01, 0.05, 0.1) to 0.0001
- Learning Rate Epoch Schedules: (100)
- SGD Epochs: 400
- Minibatch Size: 20
- Training Validation Split: (1.0) (i.e. none)
- OGD Learning Rates: (0.01, 0.05)
- L1 Regularization: (0, 0.1)

### 9.2.5 Configuration 5 - SAE

- Data Windows: [1,5,20], [5,20,60], [10,20,60]
- Learning Rate Schedules: 0.005 to (0.01, 0.05, 0.1) by (100) epochs
- Network Sizes: (120, 60, 30), (90, 60, 30), (90, 90, 90), (90, 90), (120, 90), (120, 60), (120)
- Encoding: 25;20;15;10;5
- SGD Epochs: 400
- Minibatch Size: 20
- SAE Validation Split: (0.8)
- Weight Init: Xavier Glorot Uniform, He, DC
- Layer Encodings: Leaky ReLU with Linear encoding and output

# 9.2.6 Configuration 6 - Predictive FFN

- Best SAEs by encoding size and window aggregations (15 total) from Configuration5
- Learning Rate Schedules: 0.005 to (0.01, 0.05, 0.1) by (100) epochs
- Network Structures: (120;120;120;120), (120,90,90,60), (120,90,60), (120,120,120), (120,120), (120,60), (120,120), (120
- OGD Learning Rates: (0.01, 0.05)
- Data Windows: [1,5,20], [5,20,60], [10,20,60]
- Data Scaling: Normalize
- Weight Init: Xavier Glorot Uniform, He, DC
- Layer Encodings: Leaky ReLU with Linear output
- Learning Rate Epoch Schedules: (100)
- SGD Epochs: 400
- Minibatch Size: 20
- Training Validation Split: (1.0) (i.e. none)
- L1 Regularization: (0, 0.1)

#### 9.2.7 Configuration 7 - SAE

- Data: Closing prices for AGL & ACL
- Processing: Time windows of 1, 5 and 20 (thus an input vector of size 6)
- Encoding Layer Activations: Sigmoid and Linear
- Encoding Layer Size: 5
- Hidden Layer Size: 15
- Number of Hidden Layers: 1, 2
- $\bullet$  Learning Rates: 0.0000001, 0.00001, 0.001, 0.1, 1.0, 2.0

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