# Resilient Machine Learning Approaches for Fast Risk Evaluation and Management in Financial Portfolios and Variable Annuities

by

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#### Abstract

Quantitative risk management of modern financial derivatives and actuarial products is intricate and often requires modelling the underlying stochasticity with Monte Carlo simulations. A simulation scheme is flexible, and it can easily adapt to changes in model assumptions and market conditions. However, as multiple sources of risk are considered for long time horizons, such simulation schemes become expensive to implement. Tremendous research effort is dedicated in the design of computationally efficient machine learning-based procedures that mitigate the computational burden of a standard simulation procedure. In machine learning, model flexibility comes at the expense of model resilience, which is especially crucial for risk management tasks. This study considers the management of tail risk for complex financial and actuarial products with resilient machine learning-based nested simulation procedures. We propose a novel metamodeling approach that integrates deep neural networks within the nested simulation framework for efficient risk estimation. Our approaches offer significant improvements over a standard simulation procedures and illustrate a resilient machine learning model is built by accounting for problem complexity and data structure, quality, and quantity. To further enhance adaptability to new VA contracts and changing market conditions, the thesis introduces transfer learning techniques. By reusing and fine-tuning pre-trained metamodels, the approach accelerates the adaptation process to different contract features and evolving market dynamics without the need for retraining models from scratch. Transfer learning improves computational efficiency ando enhances the robustness and flexibility neural network metamodels in dynamic hedging of variable annuities.

Extensive numerical experiments demonstrate that the proposed method achieves substantial improvements in computational efficiency, reducing simulation times by orders of magnitude compared to standard nested simulation procedures. The results indicate that the deep neural network metamodels with transfer learning can quickly adapt to new market scenarios and contract specifications. The findings of this research contribute to the advancement of risk management practices for complex actuarial products and financial derivatives. By leveraging advanced machine learning techniques, the thesis offers a practical and scalable solution for insurers to perform timely and accurate risk assessments. The integration of LSTM metamodels and transfer learning into the nested simulation framework represents a significant step toward more efficient, adaptable, and robust methodologies in actuarial science and quantitative finance.

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# Chapter 1

### Introduction

Quantitative risk management is a key component of modern financial systems. Successful risk management practice ensures the stability and resilience against a variety of risks. For financial products like option portfolios and variable annuity (VA) contracts, traditional risk assessment methods often fall short in accurately capturing the complex dynamics of the underlying risk factors. VA contracts are a popular type of complex insurance product that is linked to the performance of underlying assets. It provides a guaranteed minimum income benefit to the policyholder regardless of the performance of the underlying assets. Advanced Monte Carlo simulation techniques, particularly nested simulation procedures, become indispensable for risk assessment of such products. In contrast to finite-difference methods, a Monte Carlo simulation scheme is more flexible and can be easily adapted to model tail risk with a rule-based design. Glasserman (2004) provide a comprehensive overview of Monte Carlo simulation methods in financial engineering and risk management applications. In this thesis, we focus on building and analyzing nested simulation procedures for risk management applications of financial derivatives and insurance products. Nested simulation, also known as nested stochastic modeling and stochastic-on-stochastic modeling, becomes necessary when stochastic simulation of a parameter of interest is contingent on another quantity to be determined stochastically. In the context of financial engineering, nested simulation is used to model the tail risk of a contract whose payoff depends on a set of underlying risk factors. For example, estimating the value of an exotic option at a risk horizon requires simulation given a realization of the underlying assets upto that horizon. A standard nested simulation procedure consists of two levels of simulation: the outer-level simulation generates the underlying risk factors, while the inner-level simulation estimates the value of interest with the inner sample mean from another level of Monte Carlo simulation. This nested structure allows for accurate estimation given sufficient computational resources, but it also introduces additional complexity in the simulation design and implementation. Furthermore, in real-world applications, the computational burden of nested simulation can be prohibitive. The number of inner simulations required to achieve a desired level of accuracy for each outer scenario can be infeasibly large. Metamodeling techniques that approximate the inner simulation model can be used to reduce the computational burden of nested simulation. Metamodels are statistical models that approximate the output of a complex simulation model as a function of its input parameters. In this thesis, we focus on metamodels of the inner simulation models in nested simulation procedures for risk management applications of financial derivatives and VA contracts. Two interesting problems that requires nested simulation are considered in this thesis:

- estimating the risk of a portfolio of financial options and
- dynamic hedging of VA contracts with a delta hedging strategy.

The risk management of VAs is a challenging problem due to the complex interactions between the policyholder's behavior, the financial market dynamics, and the insurer's risk management strategies. We focus mainly on the estimation of tail risk measures of VA contract losses using metamodel-based nested simulation procedures. This thesis consists of four chapters. Chapter 2 summarizes theoretical convergence results of several state-ofthe-art single-period nested simulation procedures under the same analytical framework. Numerical experiments are conducted to test their empirical convergence behavior under finite budget sizes. Chapter 3 proposes the use of neural network models as metamodels for a two-stage multi-period nested simulation procedure on VA contracts. In our numerical experiment, the best neural network metamodel surpasses the state-of-the-art metamodels in tail identification, and it leads to substantial computational savings. Chapter 3 also conducts sensitivity testing on the neural network metamodel. We argue that for estimating tail risk measures of VA contact losses, the inner simulation can be replaced entirely by a suitable metamodel. Eliminating the inner simulation can lead to more substantial computational savings without affecting estimation quality. In the case of requiring extensive inner simulations for regulatory purposes, effective budget allocations can help achieve higher estimator quality with the same computational budget. In practice, underlying assumptions of the simulation model are subject to change, and new VA contracts are issued with different contract specifications. Simulation budgets are often scarce for new conditions. Chapter 4 uses transfer learning techniques for quick adaptation of the neural network metamodel to new market conditions and new VA contracts. Our numerical experiments shows that an intelligent use of transfer learning can lead to significantly better metamodels than those trained from scratch. Chapter 6 discusses the application of deep reinforcement learning in the dynamic hedging of VA contracts. Chapter 5 concludes the thesis and discusses potential future research directions.

# 1.1 Managing Risk of Variable Annuity with Nested Simulation

VA contracts are index-linked insurance products that offer policyholders the upside potential of equity markets while providing guarantees that protect against downside risk. These products have gained significant interest as they address the needs of individuals seeking both wealth accumulation and financial security, especially in the context of retirement planning. The most interesting feature of VAs is embedded guarantees that ensure a specified minimum benefit regardless of market performance. Two VA contracts that are most relevant to this thesis are Guaranteed Minimum Maturity Benefit (GMMB) and the Guaranteed Minimum Withdrawal Benefit (GMWB). The GMMB guarantees that at the maturity of the contract, the policyholder will receive no less than the initial investment or a predetermined minimum amount. The GMWB allows policyholders to withdraw a specified percentage of their benefit base each period during the contract horizon (Hardy, 2003). While these guarantees enhance the attractiveness of VAs to consumers by offering protection against market downturns and ensuring income stability, they introduce substantial financial risks to insurers. The embedded options within GMMBs and GMWBs expose insurers to market risk, longevity risk, and policyholder behavior risk. Effective management of these risks is crucial for insurers to maintain solvency and meet regulatory capital requirements.

One of the primary risk management strategies employed by insurers to mitigate the financial risks associated with VAs is dynamic hedging. Dynamic hedging involves frequently adjusting a portfolio of financial instruments to offset changes in the value of the guarantees due to market movements (Hull and Basu, 2016). This strategy aims to neutralize the sensitivity of the insurer's liability to market fluctuations by constructing a hedging portfolio that replicates the cash flows of the guarantees. However, dynamic hedging introduces complexity in estimating the insurer's overall risk exposure. One of such examples is the estimation of tail risk measures of a VA contract. It requires the stochastic modeling of the financial market and the accurate loss estimation of the hedging portfolio under different market scenarios. Nested simulation has emerged as a robust method for estimating risk measures in such complex settings (Gordy and Juneja, 2010). In nested

simulation, an outer simulation generates a set of market scenarios over the contract horizon, while an inner simulation estimates the contract loss in each scenario. By combining the results of the inner simulations across the outer scenarios, nested simulation provides an accurate estimate of the tail risk of the VA contract.

Despite its robustness, nested simulation is computationally intensive, often requiring significant computational resources and time. Each scenario in the outer simulation requires numerous inner simulations to accurately estmate contract loss. This computational burden poses practical limitations, especially when high precision in tail risk estimation is required. To address this challenge, metamodeling techniques can be employed to approximate the inner simulation model and reduce the computational cost of nested simulation.

#### 1.2 Metamodeling for Monte Carlo Simulation

Metamodeling, also known as surrogate modeling, is a technique used to approximate complex simulation models with simpler, computationally efficient models. In the context of Monte Carlo simulations, metamodels are models of models that aim to reduce computational costs by creating an approximate model that can quickly predict simulation outputs based on input variables (Kleijnen, 2018). This section introduces metamodeling for Monte Carlo simulation, elaborates on its methodologies, and highlights its importance in practical applications.

Monte Carlo simulations are widely used for modeling and analyzing complex systems that are probabilistic in nature. These simulations often require a significant amount of computational resources, especially when high accuracy or numerous iterations are necessary (Glasserman, 2004). Metamodeling addresses this challenge by constructing an approximate model (metamodel) that emulates the behavior of the original simulation model with much less computational effort. The metamodel serves as a predictive model that maps input variables to output responses. Learning from a set of simulation runs, the metamodel can generalize and predict outputs for new inputs without running the full simulation. Metamodeling is particularly useful when the simulation model is computationally expensive. By replacing the simulation model with a metamodel, the computational cost of evaluating the model can be significantly reduced. This allows for faster convergence of simulation-based estimators.

The process of metamodeling generally involves:

1. **Design of Experiments**: Selecting input combinations to run the original simulation and collect data.

- 2. Building the Metamodel: Using the collected data to train a metamodel that approximates the simulation model.
- 3. Validation: Assessing the metamodel's accuracy and generalization capability.
- 4. **Application**: Using the metamodel to predict simulation outputs for new input combinations.

The development of advanced machine learning models and deep learning techniques has significantly enhanced metamodeling approaches for Monte Carlo simulations, especially in high-dimensional and complex problem settings. Example use cases for machine learning include Jin et al. (2020), Tang et al. (2020), and Rosen et al. (2012). These methods offer powerful tools for capturing intricate patterns and nonlinear relationships that traditional metamodeling techniques might struggle to model effectively. In the following sections, we discuss machine learning algorithms in the context of metamodeling for Monte Carlo simulations and their applications in risk management.

#### 1.3 Machine Learning for Risk Management Applications

Machine learning (ML) algorithms are essential tools for identifying complex patterns and relationships in data. In particular, they are well-suited for handling non-linearities and large-scale data structures, which are challenging for traditional statistical methods. In the context of risk management, ML algorithms have been widely used for predicting financial time series, estimating risk measures, and optimizing trading strategies. ML algorithms can be broadly categorized into supervised learning, unsupervised learning, and reinforcement learning. This section focuses on three widely used supervised learning models and two reinforcement learning algorithms that are relevant to risk management applications.

#### 1.3.1 Supervised Learning Models

Supervised learning is a fundamental approach in machine learning where the algorithm learns a mapping from inputs to outputs based on example input-output pairs Galton (1886). In this paradigm, a model is trained on a labeled dataset, which means that each training example is associated with an output label or value. The goal is to learn a general

rule that maps inputs (also known as features) to outputs (also known as target), enabling the model to make accurate predictions on new, unseen data.

- Classification: The output variable is categorical, and the task is to assign inputs to one of several predefined categories. Common use in finance and actuarial applications are fraud detection and credit scoring.
- Regression: The output variable is continuous, and the task is to predict a real-valued number. Examples in actuarial applications include predicting claim amounts, reserve estimates, asset pricing, and estimating tail risk measures of loss distributions.

This thesis focuses on regression models for risk management applications, where the goal is to predict a continuous target variable based on one or more input features. Given a dataset of n observations  $\{(x_1, y_1), (x_2, y_2), \dots, (x_n, y_n)\}$ , where  $x_i \in \mathbb{R}^d$  is the d-dimensional input feature vector, and  $y_i \in \mathbb{R}$  is the target variable, a supervised learning algorithm aims to learn a function  $f(;\theta) : \mathbb{R}^d \to \mathbb{R}$  that best predicts the target variable y given the input features x with a set of function parameters  $\theta$ .

Linear regression, kernel regression, and neural networks all fall under the supervised learning category and are widely used in risk management applications. Despite the differences in their functional forms, all supervised learning models can be evaluated on a similar set of metrics. The learning process is referred to as training, which involves minimizing a loss function  $l(f(x;\theta),y)$  over the training data, which quantifies the difference between the predicted labels and the training labels. A common loss function for regression problems is the mean squared error (MSE):

$$MSE = \frac{1}{n} \sum_{i=1}^{n} (f(x_i; \theta) - y_i)^2.$$
 (1.1)

In most risk management applications, a critical task of users of supervised learning models is to design a suitable loss function that aligns with the objectives of the problem. Another important consideration is the choice of a suitable supervised learning algorithm. The choice may vary based on the complexity of the relationship, the interpretability of the model, and the computational resources available. In the following sections, we discuss four widely used supervised learning techniques in risk management applications: parametric regression, kernel regression, and two neural network architectures.

#### 1.3.2 Parametric Regression Models

Regression models are the most common supervised learning algorithms used in risk management applications. A regression model predicts a continuous target variable based on one or more input features. Linear regression is a simple and interpretable model that attempts to predict a target variable as a linear combination of input features. It assumes a linear relationship between the target output and one or more input features. This method has been thoroughly explored in statistical literature. Bishop and Nasrabadi (2006) provide an extensive treatment of regression techniques in the broader context of machine learning. It is known as parametric regression because it assumes a specific functional form for the input-output relationship with a finite set of parameters (Seber and Lee, 2012). General form of a parametric regression model is given by:

$$f(x;\theta) = \beta_0 + \sum_{j=1}^p \beta_j \phi_j(x), \qquad (1.2)$$

where p is the number of features,  $\beta_0, \beta_1, \dots, \beta_p$  are the trainable regression coefficients, and  $\phi_i(x)$  are basis functions that transfrom the input x to allow for non-linear modeling. Basis functions can be any functions of x that are chosen to capture the underlying structure of the data. Common choices include polynomial basis functions:  $\phi_i(x) = x^j$  and Laguerre basis functions:  $\phi_j(x) = e^{-x/2}L_j(x)$ , where  $L_j(x)$  are the Laguerre polynomials that are solutions to the Laguerre differential equation (Szeg, 1939). The training of parametric regression refers to the process of estimating the regression coefficients  $\beta_0, \beta_1, \ldots, \beta_p$  that minimize the loss function. For a MSE loss, the optimal regression coefficients can be obtained by solving the normal equations. Parametric regression is a powerful tool for modeling linear relationships where the basis functions are known from expert knowledge for explicit feature engineering techniques (Hastie et al., 2009). These techniques utilize predefined functional forms to model the relationship between independent variables and the dependent variable. However, when data exhibits complex, non-linear relationships, linear models fall short. While these methods are straightforward and interpretable, they impose strong assumptions about the underlying data structure. Often, expert knowledge is required to select the appropriate basis functions, which can limit the flexibility of the model. Extensions like polynomial regression and generalized linear models (GLMs) have been introduced to capture non-linearity. Nevertheless, these models can still be limited in capturing highly complex patterns.

#### 1.3.3 Non-Parametric Regression Models

To overcome some limitations of parametric regression, non-parametric methods such as kernel regression (Hastie et al., 2009) are often employed. Kernel regression estimates the relationship by averaging the values of the target variable over a local neighborhood of the input x. The kernel regression with the Nadaraya-Watson estimator is given by:

$$f(x;\theta) = \frac{\sum_{i=1}^{n} K\left(\frac{x-x_i}{h}\right) y_i}{\sum_{i=1}^{n} K\left(\frac{x-x_i}{h}\right)},$$
(1.3)

where  $K(\cdot)$  is the kernel function, which assigns weights to the data points based on their distance to the input x, and h is the bandwidth parameter that controls the smoothness of the estimated function. The objective is to estimate the feature-label relationship directly from the data without imposing a specific form. However, they come with several drawbacks that limit their effectiveness, especially in high-dimensional settings or for large datasets. One of the most significant drawbacks of non-parametric regression is the curse of dimensionality Bellman (1966). As the number of features increases, the volume of the input space grows exponentially. Data points become sparse and the model may overfit to noise in the data. In addition, the computational cost associated with non-parametric methods often grows rapidly with the number of dimensions. Cross validation and distance calculation are two computationally expensive operations that are often required for non-parametric regression. These drawbacks make non-parametric regression impractical for modeling high-dimensional large-scale datasets.

#### 1.3.4 Feedforward Neural Networks

The progression from traditional parametric and non-parametric regression methods to neural network architectures has been driven by the need to model increasingly complex and high-dimensional data. Neural networks have emerged as powerful tools capable of overcoming many limitations of traditional regression methods. They can learn complex, non-linear relationships in data without the need for explicit feature engineering. The most crude form of a neural network is the feedforward neural network (FNN), which consists of an input layer, one or more hidden layers, and an output layer (Goodfellow et al., 2016a).

Figure 1.1a shows a feedforward neural network (FNN) architecture with one hidden layer. The input layer receives the input features, which are then passed through the hidden layer(s) to the output layer. Each layer consists of multiple neurons, which apply a non-linear activation function to the weighted sum of the inputs.



Figure 1.1: Neural Network Architectures

$$f(x;\theta) = f^{(k)}(f^{(k-1)}\cdots(f^{(2)}(f^{(1)}(x;\theta^{(1)});\theta^{(2)})\cdots;\theta^{(k-1)});\theta^{(k)}), \tag{1.4}$$

where  $\theta = (\theta^{(1)}, \theta^{(2)}, \dots, \theta^{(k)})$  are the trainable parameters of the neural network,  $f^{(i)}$  is the *i*-th layer of the neural network, x is the input feature vector,  $f^{(1)}(x; \theta^{(1)}) = x$ ,  $f^{(k)}$  is the output layer, k is the number of layers, and  $f^{(i)}$  is the output of the *i*-th layer. To move from a lower layer to a higher layer, i.e., from layer i - 1 to layer i,

$$f^{(i)}(z;\theta^{(i)}) = a^{(i)}(\beta_0^{(i)} + \sum_{j=1}^{p_i} \beta_j^{(i)} z), \tag{1.5}$$

where  $a^{(i)}$  are the nonlinear activation functions,  $\theta^{(i)} = (\beta_0^{(i)}, \dots, \beta_p^{(i)})$  are the trainable parameters of the *i*-th layer, and  $p_i$  is the number of neurons in the previous layer. Equation 1.5 represents a single neuron in a neural network, which applies a linear transformation to the input followed by a non-linear activation function. The linear transformation is similar to the linear regression model in Equation 1.2, but the non-linear activation function allows the neural network to model complex, non-linear relationships in the data. The FNN is tuned by adjusting the trainable parameters  $\theta$  while fixing the activation functions  $a^{(i)}$ . A typical choice for the activation function is the rectified linear unit (ReLU), which is defined as  $a(z) = \max(0, z)$  (Nair and Hinton, 2010). The choice of activation function

also plays a crucial role in the training of neural networks. While sigmoid and hyperbolic tangent functions were popular in early neural networks, ReLU has become the default activation function for deep networks due to its ability to mitigate the vanishing gradient problem and promote sparse activations (LeCun et al., 2015).

The key advantage of feedforward neural networks (FNNs) lies in their ability to perform automatic feature engineering without direct human intervention. Unlike traditional machine learning models that require manual selection and transformation of input features, neural networks learn to extract and compose features through their hidden layers during the training. Each hidden layer in the network captures higher-level abstractions of the input data by transforming the outputs of the previous layer through nonlinear activation functions (LeCun et al., 2015). Each layer builds upon the representations learned by the previous layer, enabling the network to capture multiple levels of abstraction. This characteristic is crucial for modeling complex datasets with intricate patterns and relationships (Bengio et al., 2013).

The last layer of the neural network, known as the output layer, typically performs a linear transformation of the features extracted by the preceding hidden layers. With  $a^k(z) = z$ , the last layer is equivalent to a linear regression with its inputs as transformed, high-level features learned by the hidden layers. We can draw parallels between neural networks and traditional regression models. The main difference is that, in neural networks, the input features to the regression model are learned automatically rather than being manually specified.

The most well-known theorem in neural network theory is the universal approximation theorem (Hornik et al., 1989). It states that given appropriate activation functions a, a feedforward neural network with a single hidden layer containing a finite number of neurons can approximate any continuous function on a compact subset of  $\mathbb{R}^n$  to arbitrary accuracy. Despite this theoretical guarantee for single-layer neural networks, in practice, deep neural networks (DNN) with multiple hidden layers have been shown to be more effective at capturing complex patterns and relationships in data (LeCun et al., 2015). Examples include AlexNet (Krizhevsky et al., 2012), VGG (Simonyan and Zisserman, 2014), and ResNet (He et al., 2016), which have achieved state-of-the-art performance on image classification tasks. In this thesis, we focus on the application of deep neural networks, specifically long short-term memory (LSTM) networks, as metamodels for nested simulation procedures in risk management applications.

#### 1.3.5 Long Short-Term Memory Networks

Building upon the capabilities of FNNs, we recognize that while FNNs are successful at capturing complex, nonlinear relationships through automatic feature learning, they are inherently limited when it comes to modeling sequential data or time-dependent patterns. This independence assumption limits their effectiveness in modeling financial time series, where temporal dependencies play a critical role. The stylized facts of financial time series, such as volatility clustering, fat tails, and autocorrelation, are challenging to capture with traditional FNNs due to the absence of memory in the model (Cont, 2001).

To overcome the limitations of FNNs in handling sequential input, recurrent neural networks (RNNs) were introduced. In a RNN, the hidden state at each time step is a function of both the current input and the hidden state from the previous time period (Elman, 1990):

$$h_t = f(x_t, h_{t-1}; \theta),$$
 (1.6)

where  $h_t$  is the hidden state at time t,  $x_t$  is the input at time t,  $h_{t-1}$  is the hidden state at time t-1, and f is the recurrent function parameterized by  $\theta$ . This architecture enables RNNs to capture temporal dependencies by maintaining a dynamic internal state that reflects the memory of past inputs. However, traditional RNNs suffer from the vanishing and explodeing gradient problem, which hinders their ability to capture long-term dependencies that often present in financial time series (Bengio et al., 1994). During training, gradients propagated backward through time can either diminish exponentially (vanishing gradients) or grow uncontrollably (exploding gradients). This limitation is particularly problematic in modeling long-term financial contracts and insurance guarantees, where patterns may span over extended periods.

To address these issues, a long short-term memory (LSTM) network was developed by Hochreiter and Schmidhuber (1997). It is a specialized form of RNNs designed to capture long-term dependencies more effectively with the help of RNN memory cells and gating mechanisms.

$$i_{t} = a(W_{i}x_{t} + U_{i}h_{t-1} + b_{i}),$$

$$f_{t} = a(W_{f}x_{t} + U_{f}h_{t-1} + b_{f}),$$

$$o_{t} = a(W_{o}x_{t} + U_{o}h_{t-1} + b_{o}),$$

$$g_{t} = a(W_{g}x_{t} + U_{g}h_{t-1} + b_{g}),$$

$$c_{t} = f_{t} \odot c_{t-1} + i_{t} \odot g_{t},$$

$$h_{t} = o_{t} \odot \tanh(c_{t}),$$

where  $i_t$ ,  $f_t$ ,  $o_t$ ,  $g_t$ ,  $c_t$ ,  $h_t$  are the input gate, forget gate, output gate, cell input, cell state, and hidden state at time t, respectively.  $W_i$ ,  $W_f$ ,  $W_o$ ,  $W_g$ ,  $U_i$ ,  $U_f$ ,  $U_o$ ,  $U_g$  are the weight matrices, and  $b_i$ ,  $b_f$ ,  $b_o$ ,  $b_g$  are the bias vectors. a is the activation function, typically the sigmoid function, and  $\odot$  denotes element-wise multiplication.

$$a(z) = \frac{1}{1 + e^{-z}}.$$

Figure 1.1b shows the architecture of an LSTM network. The gating mechanisms in LSTM networks effectively mitigate the vanishing and exploding gradient problem by regulating the flow of information through the network. The input gate  $i_t$  controls the flow of information into the cell state  $c_t$ , the forget gate  $f_t$  regulates the retention of information in the cell state, and the output gate  $o_t$  determines the information passed to the hidden state  $h_t$ . The cell input  $g_t$  is used to update the cell state based on the input  $x_t$  and the previous hidden state  $h_{t-1}$ . The cell state  $c_t$  acts as a memory unit that stores information over time, while the hidden state  $h_t$  captures the relevant information for the current time step. By incorporating memory cells and gating mechanisms, LSTMs can effectively model long-term dependencies in sequential data in finance and actuarial applications.

The advancements in neural network optimization techniques, architectures, and training methodologies have significantly enhanced their usefulness in risk management applications. By effectively modeling complex, non-linear relationships and temporal dependencies, neural networks serve as powerful tools for addressing the computational challenges in estimating risk measures and developing effective risk mitigation strategies. This thesis aims to explore the application and noise tolerance of LSTM networks in metamodeling for nested simulation procedures in risk management. We investigate the performance of LSTM networks in approximating the inner simulation model in a two-stage nested simulation procedure for index-linked insurance contracts. By leveraging the memory and sequential modeling capabilities of LSTMs, we aim to improve the accuracy and efficiency of nested simulation procedures for risk management applications.

# 1.3.6 Training and Evaluation of Neural Networks in a Simulation Environment

The training of neural networks involves adjusting the trainable parameters of the network to minimize the loss function, which quantifies the error between the predictions and labels. The training process aims to find the optimal parameters that minimize this error. The most common approach is to use gradient descent methods with backpropagation, which is an efficient method for computing the gradients of the loss function with respect to the trainable parameters. Backpropagation computes the gradients by propagating the error derivatives backward through the network layers. This process allows the network to adjust the parameters in the direction of minimizing the loss function. The training process is typically performed over a number of epochs, where each epoch is one complete pass through the training dataset.

However, the training of neural networks is non-convex, and the optimization problem is challenging. Stochastic gradient descent (SGD) is a popular optimization algorithm for training neural networks. It updates the parameters using a single or a small batch of training examples at a time, which makes it much faster than the crude gradient descent. However, SGD may oscillate and converge slowly, especially in non-convex problems (Goodfellow et al., 2016b). To address these issues, various adaptive learning rate methods have been proposed. The most popular approach is Adam, which is a SGD algorithm that adjusts the learning rate during training to improve convergence speed and stability (Kingma and Ba, 2014).

Another common technique to improve the training of neural networks is regularization to prevent overfitting. One effective regularization method is dropout, which randomly sets a fraction of neurons to zero during each training iteration (Srivastava et al., 2014). This prevents neurons from co-adapting too much, encourages redundancy, and leads to a more robust model that generalizes better to unseen data. Early stopping is another regularization method that is relevant to our study. It involves monitoring the model's performance on a validation set and stopping training when the performance no longer improves (Prechelt, 2002). This helps prevent over-fitting by stopping training before the model begins to fit to noise in the data.

Evaluation of the performance of a neural network is challenging due to the absence of analytical tools. Existing machine learning literature addresses this challenge by splitting the data into three parts: training set, validation set, and test set. The training set is used to train the model, the validation set is used to tune the model hyper-parameters, and the test set is used to evaluate how well the model generalizes to unseen data. The test set is not used during training and is only used for evaluation.

Under-fitting occurs when the training error is high, which means that the neural network fails to capture the underlying patterns in the training data. Over-fitting occurs when the neural network fits to noise in the training data and cannot generalize to unseen data. In machine learning literature, over-fitting is often quantified as the gap between the training error and the test error (Bishop and Nasrabadi, 2006). During training, as the test dataset is not used, this quantity is estimated using the validation set. If the validation error is high compared to the training error, the model is likely over-fitting to the training data.

In practice, the ultimate performance of a neural network is often evalutated based on error metrics on the test dataset and how it compares with the training error. The test error is often thought to be a reliable indicator of the model's generalization capability for most machine learning applications. However, real-world datasets often contain noise. As the test dataset is separated from the training dataset, it is often contaminated with noise. Therefore, the test error may not be robust for evaluating the generalization capability of a neural network. In this thesis, we aim to evaluate the generalization capability of neural networks in a simulation environment. For a simulation model, we can generate a large number of datasets with different levels of noise. The noise level can be controlled by the design parameters of the simulation model, e.g., the number of replications. Instead of relying on the noisy test labels, we can generate a less noisy test labels using a higher number of replications, or even the true simulation model. In the rest of this thesis, we will use "test dataset" or "test labels" to refer to the noisy test dataset, and "true dataset", "true labels", or "true relationship" to refer to dataset generated from the true simulation model.



Figure 1.2: Different Levels of Noise

Figure 1.2 shows datasets with different levels of noise in a typical machine learning setting. The left panel illustrates the common practice of using a test dataset that is

separate from the training dataset that is contaminated with noise. The right panel shows an alternative approach of using true labels generated from the true relationship. In a simulation context, the true relationship, which is the simulation model that generates the data, is visible to the user. Even if the true relationship is unknown, we can approximate it using a large number of replications and use it as the true labels. Comparing the errors of a neural network on the true labels can reveal its true generalization capability and robustness. Chapter 3 provides a detailed study of the generalization capability of LSTM networks in a simulation environment for risk management of VA contracts. Viewing the simulation environment as a data-generating process allows us to systematically study and evaluate the performance of neural networks within a controlled setting. This approach provides insights into their applicability in real-world risk management applications.

In the subsequent chapters, we dive deeper into these topics. Chapter 2 provides a comprehensive analysis of the convergence properties of existing nested simulation procedures and their practical implications. Chapter 3 focuses on the development and evaluation of neural network metamodels, particularly LSTM networks, for enhancing the efficiency of nested simulations in risk management of VA contracts. We also examine the robustness and generalization capabilities of these models in the presence of simulation noise. Chapter 4 explores the use of transfer learning techniques to adapt neural network metamodels to new market conditions and contract specifications. Our approach enables rapid deployment in changing environments. Finally, Chapter 5 summarizes the key findings of the thesis and discusses future research directions that involve the robost use of neural networks in quantitative risk management applications.

## Chapter 2

# Nested Simulation Procedures in Financial Engineering: A Selected Review

#### 2.1 Introduction

Nested simulation procedures are commonly used in financial engineering to estimate risk measures for portfolios of complex financial derivatives. The term *nested* is referred to a nested estimation problem, in which the estimation of the risk measure requires two levels of Monte Carlo (MC) simulations. In a typical nested simulation procedure, an outer level simulation model generates underlying risk factors, which is referred to as the *outer scenarios*. For each outer scenario, an inner level simulation model generates scenario-wise samples of the portfolio losses, which is referred to as the *inner replications*.

The nested simulation procedure is computationally expensive due to its nested structure. Given a fixed computational budget, the nested simulation procedure has to make a trade-off between the number of outer scenarios and the number of inner replications. Gordy and Juneja (2010) are the first to analyze and propose the optimal budget allocation of a standard nested simulation procedure. The term *standard* refers to using a standard MC estimator, the sample mean of the inner replication to estimate a scenariowise portfolio loss for an outer scenario. Gordy and Juneja (2010) investigate the optimal budget allocation for a standard nested simulation procedure with respect to the MSE of the estimated risk measure.

The standard nested simulation procedure is computationally expensive with a somewhat wasteful use of the simulation budget, as only the inner replications from the same outer scenario are used in estimating the scenario-wise portfolio loss for that outer scenario. Subsequent research efforts have been made to improve the efficiency of nested simulation procedures by using the inner replications from other outer scenarios. This is referred to as pooling. Different methods pool in different ways, either by a trained supervised learning metamodel or by pre-defined likelihood ratio weights. Broadie et al. (2015) propose a regression-based nested simulation procedure, which uses a trained regression metamodel to estimate the scenario-wise portfolio loss for an outer scenario by pooling the inner replications from all outer scenarios. For risk measures in certain forms, Broadie et al. (2015) show that it is optimal to allocate all simulation budget to the outer level simulation, and the inner replication should be kept to a minimum of 1. Similarly, Hong et al. (2017), Feng and Song (2020), and Zhang et al. (2022) use a kernel smoothing model, a likelihood ratio model, and a kernel ridge regression (KRR) model as metamodels to pool the inner replications from all outer scenarios. In simulation studies, this approach of using a model of a simulation model is known as metamodeling, and the models of a simulation model are also referred to as metamodels (Barton, 1998). Another line of research is the multi-level Monte Carlo (MLMC) method analyzed in Giles and Haji-Ali (2019), which is a variance reduction technique that uses a hierarchy of approximations to the quantity of interest.

This paper presents a survey study of some popular nested simulation procedures that have both asymptotic convergence analysis and finite-sample performance evaluation. Many procedures are proposed in the literature, but they are not directly comparable due to having different error metrics, different assumptions, and different numerical examples. Within a common analytical framework, we first summarize and compare their asymptotic rate of convergence. Their asymptotic convergence results are closely examined for their assumptions that guarantee the convergence. Furthermore, the existing literature chooses different performance metrics to evaluate the asymptotic convergence of nested simulation procedures. In this chapter, we discuss the asymptotic convergence in terms of MSE of the estimator, and we provide a comprehensive comparison of their convergence rates and critical assumptions. We also discuss the connection between the MSE and the absolute error of the estimator in the context of nested simulation procedures.

In terms of finite-sample performance, different studies propose different examples in their numerical experiments, which introduces unfair advantages for certain simulation procedures over others. A fair comparison among popular methods is therefore urgently needed in the literature. Our numerical experiment in this chapter is the first of its kind to subject back all the aforementioned simulation procedures to a complete and unbiased comparison. Extensive numerical experiments are conducted to show, in practical examples, how well

the finite-sample performance of a method matches its theoretical convergence behavior. With our numerical examples, we compare the nested simulation procedures for different payoff complexity, problem dimensions, and risk measures. This chapter also provides a discussion on the computational complexity of different nested simulation procedures, including additional computational costs for training supervised learning metamodels and pooling inner replications.

The rest of the paper is organized as follows. Section 2.2 introduces the nested simulation procedure and the standard MC estimator. Section 2.3 provides new theoretical results on the convergence of existing nested simulation procedures. Section 2.4 summarizes the asymptotic convergence orders and the critical assumptions of nested simulation procedures in the literature. Section 2.5 presents the numerical experiments and the comparisons of different nested simulation procedures with respect to different risk measures, problem dimensions, and payoff complexities. Section 2.6 discusses the computational complexity of different nested simulation procedures. Section 2.7 concludes this chapter.

#### 2.2 Problem Formulation

In a nested estimation problem, we are interested in estimating the quantity

$$\rho(L) = \rho(L(X))$$

where  $X \in \mathcal{X} \subset \mathbb{R}^d$ , and L = L(X) is a random variable whose distribution is characterized by X. L(X) can't be directly evaluated, but it is the output of

$$L(X) = \mathbb{E}\left[Y|X=x\right]|_{x=X}$$

Some common risk measures are in the nested expectation form, in which

$$\rho(L) = \mathbb{E}\left[h(L)\right]$$

where  $h: \mathbb{R} \to \mathbb{R}$  is a known nonlinear function. The nested expectation form is a general form that covers many risk measures of interest in financial engineering by varying the function h. Common forms of h in the literature include smooth functions, Lipschitz continuous functions, and indicator functions.

**Definition 1** A function  $h : \mathbb{R} \to \mathbb{R}$  is smooth if it is continuously differentiable up to the n-th order for some  $n \in \mathbb{N}$ , and its n-th and m-th derivatives are bounded for some  $m \in \mathbb{N}$  and m < n.

A quadratic tracking error with a benchmark b falls into this category, in which  $h(t) = (t - b)^2$ . In the context of nested estimation for smooth h, different procedures require different versions of Definition 1 to specify the smoothness for their convergence analysis. Table 2.1 summarizes the smoothness assumption of h for different nested simulation procedures in the literature.

Nested Simulation Procedures	n	m
Regression	2	not required
Kernel smoothing	3	2
Kernel ridge regression	2	1
Likelihood ratio	2	1

Table 2.1: Smoothness assumption of h for different nested simulation procedures

The regression-based nested simulation procedure has the least stringent smoothness assumption. In general, a more complex metamodel requires a higher-order smoothness for h for its convergence analysis. Despite being more advanced than kernel smoothing, the KRR-based procedure requires a less stringent smoothness assumption on h. It is counter-intuitive that a more complex metamodel requires a less stringent smoothness assumption. This phenomenon is due to the fact that the convergence analysis for the KRR-based nested simulation procedure is shown in terms of absolute error, which is a less stringent error metric than MSE. Section 2.3.1 discusses the connection between the MSE and the AE of the estimator in the context of nested simulation procedures. In Table 2.1, the standard nested simulation procedure is the only procedure without asymptotic convergence analysis for smooth h. Section 2.3.2 fills this gap by providing the asymptotic convergence analysis for the standard nested simulation procedure in terms of MSE.

**Definition 2** A function  $h : \mathbb{R} \to \mathbb{R}$  is Lipschitz continuous with Lipschitz constant K if for all  $t_1, t_2 \in \mathbb{R}$ ,

$$|h(t_1) - h(t_2)| \le K|t_1 - t_2|.$$

A mean excess loss over a threshold u is defined by setting  $h(t) = \max\{t - u, 0\}$ , which is a special case of a Lipschitz continuous function. The regression-based nested simulation

procedure is the only procedure that performs theoretical analysis for Lipschitz continuous h. The other procedures only provide convergence analysis for the special case, mean excess loss over a threshold u.

**Definition 3** A function  $h: \mathbb{R} \to \mathbb{R}$  is an indicator function if

$$h(t) = \mathbb{I}_{\{t \ge u\}}$$

for some  $u \in \mathbb{R}$ .

The probability of a large loss over a threshold u is obtained by setting  $h(t) = \mathbb{I}_{\{t \geq u\}}$ .

Other risk measures of interest that are not in the nested expectation form are the value at risk (VaR) and the conditional value at risk (CVaR) (Hardy and Saunders, 2022). The  $\alpha$ -VaR of L is defined as

$$VaR_{\alpha}(L) = q_{\alpha} = \inf \{ q : \Pr(L \le q) \ge \alpha \}.$$
(2.1)

The  $\alpha$ -CVaR of L is defined as

$$CVaR_{\alpha}(L) = \frac{1}{1-\alpha} \int_{\alpha}^{1} q_{v} dv.$$
 (2.2)

Other names of CVaR include Conditional Tail Expectation (CTE), Tail Value-at-Risk (TailVaR), and Expected Shortfall (ES).

The rest of this section reviews the existing nested simulation procedures in the literature. We unify their notations and provide a comprehensive comparison of how inner replications are generated and pooled for different procedures.

#### 2.2.1 The Standard Nested Simulation Procedure

The standard nested simulation procedure first simulates M independent and identically distributed (iid) outer scenarios  $X_1, \ldots, X_M$  from  $F_X$ , the distribution of X. For each  $X_i$ , again simulate  $Y_{ij}$ ,  $j=1,\ldots,N$  from  $F_{Y|X_i}$ , the conditional distribution of Y given  $X_i$ . Given scenario i, the  $Y_{ij}$  are conditionally iid. Let  $\Gamma = M \cdot N$  denote the total simulation budget,  $f_X(x)$  denote the density of X, and  $\mathbf{X} = (X_1, \ldots, X_M)$  denote the vector of outer scenarios.

The standard nested simulation procedure estimates  $L_i = L(X_i)$  with a standard MC estimator

$$\hat{L}_{N,i} = \frac{1}{N} \sum_{j=1}^{N} Y_{ij}; \quad Y_{ij} \sim F_{Y|X_i}$$

Let  $\hat{L}_{(i)}, \ldots, \hat{L}_{(M)}$  be the order statistics of  $\hat{L}_{N,1}, \ldots \hat{L}_{N,M}$ . The standard nested simulation estimators for different forms of  $\rho$  are as follows:

1. Nested expectation form:

$$\hat{\rho}_{M,N} = \frac{1}{M} \sum_{i=1}^{M} h(\hat{L}_{N,i}) = \frac{1}{M} \sum_{i=1}^{M} h(\bar{Y}_{N,i}); \quad X_i \sim F_X$$

2. Value at risk (VaR):

$$\hat{\rho}_{M,N} = \hat{L}_{(\lceil \alpha M \rceil)}$$

3. CVaR:

$$\hat{\rho}_{M,N} = \hat{L}_{(\lceil \alpha M \rceil)} + \frac{1}{(1-\alpha)M} \sum_{i=1}^{M} \max\{\hat{L}_{N,i} - \hat{L}_{(\lceil \alpha M \rceil)}, 0\}$$
(2.3)

Gordy and Juneja (2010) analyze the optimal budget allocation of the standard nested simulation procedure with respect to the MSE of the estimator  $\hat{\rho}_{M,N}$  for hockey-stick h, indicator h, and VaR.

#### 2.2.2 Multi-level Monte Carlo

The MLMC method is a variance reduction technique that uses a hierarchy of approximations to the quantity of interest, and it uses the difference between the approximations to reduce the variance of the estimator. The MLMC method is particularly useful when the quantity of interest is expensive to evaluate, and the standard MC estimator has a high variance. The MLMC method estimates  $\rho$  with a MLMC estimator:

$$\hat{\rho}_{\Gamma}^{\text{MLMC}} = \sum_{v=0}^{\Upsilon} \left( \frac{1}{M_v} \sum_{i=1}^{M_v} h(\hat{L}_{N_v}(X_{i,v})) - \frac{1}{M_{v-1}} \sum_{i=1}^{M_{v-1}} h(\hat{L}_{N_{v-1}}(X_{i,v-1})) \right), \quad X_{i,v} \sim F_X,$$

where  $\hat{L}_{N-1}(\cdot) = 0$ ,  $\Upsilon$  is the number of levels,  $M_v$  is the number of outer scenarios at level v, and  $N_v$  is the number of inner replications at level v. Applying the analysis of Giles (2015) in a nested simulation context, Giles and Haji-Ali (2019) show that the MLMC method can achieve a similar level of accuracy as the standard nested simulation procedure with a lower total computational budget. The simulation budget  $\Gamma$  is the sum of the computational budget at each level, that is,  $\Gamma = \sum_{v=0}^{\Upsilon} M_v \cdot N_v$ . Neither the standard nested simulation procedure nor the MLMC method involves pooling the inner replications from all outer scenarios. Pooling enhances sample efficiency, and it is a key feature that distinguishes the more advanced nested simulation procedures from the standard procedure.

#### 2.2.3 Supervised Learning Metamodels

Supervised learning-based nested simulation procedures treat the inner simulation model  $L(\cdot)$  as a black-box function, and they approximate  $L(\cdot)$  with supervised learning metamodels. In this metamodeling approach,  $L(\cdot)$  can be approximated by  $\hat{L}_{M,N}^{\mathrm{SL}}(\cdot)$ , which is based on a chosen function family and observations from the standard nested simulation procedure. Consider the observation pairs  $(X_i, \hat{L}_{N,i})$  for  $i \in \{1, \ldots, M\}$  as training data, we can use supervised learning to approximate  $L_i$  by  $\hat{L}_{M,N}^{\mathrm{SL}}(X_i)$  and to pool the inner replications from all outer scenarios. More specifically,  $\hat{L}_{M,N}^{\mathrm{SL}}(\cdot)$  denotes the supervised learning model trained on  $(X_i, \hat{L}_{N,i})$  for  $i \in \{1, \ldots, M\}$ , which is the simulation output of the standard nested simulation procedure with M outer scenarios and N inner replications. To evaluate the convergence of the supervised learning-based nested simulation procedure, we consider the convergence behavior of the supervised learning model  $h(\hat{L}_{M,N}^{\mathrm{SL}}(\cdot))$  to the true risk measure  $\rho = h(L(\cdot))$ . More specifically, we want to study the quantity

$$\mathbb{E}\left[h(\hat{L}_{M,N}^{\mathrm{SL}}(X))\right] - \rho; \quad X \sim F_X. \tag{2.4}$$

However, studying the expectation in Equation 2.4 is unrealistic, as in practice the risk measure  $\rho$  is estimated with a finite number of samples. Using the M training samples, a supervised learning-based nested MC estimator of  $\rho$  is given by

#### 1. Nested expectation form:

$$\hat{\rho}_{M,N}^{\text{SL,Train}} = \frac{1}{M} \sum_{i=1}^{M} h(\hat{L}_{M,N}^{\text{SL}}(X_i)); \quad X_i \sim F_X$$
 (2.5)

2. VaR:

$$\hat{\rho}_{M,N}^{\mathrm{SL,Train}} = \hat{L}_{(\lceil \alpha M \rceil)}^{\mathrm{SL}}$$

3. CVaR:

$$\hat{\rho}_{M,N}^{\text{SL,Train}} = \hat{L}_{(\lceil \alpha M \rceil)}^{\text{SL}} + \frac{1}{(1-\alpha)M} \sum_{i=1}^{M} \max\{\hat{L}_{M,N}^{\text{SL}}(X_i) - \hat{L}_{(\lceil \alpha M \rceil)}^{\text{SL}}, 0\}$$

where  $\hat{L}_{(\lceil \alpha M \rceil)}$  is the  $\lceil \alpha M \rceil$ -th order statistic of  $\hat{L}_{M,N}^{\mathrm{SL}}(X_1), \dots, \hat{L}_{M,N}(X_M)$ .

After a supervised learning metamodel is trained, it can be used to make predictions for all  $X \in \mathcal{X}$ . Instead of using the same training samples to estimate the risk measure, we can use metamodel predictions for a new set of M' test samples of X, namely  $\tilde{\mathbf{X}} = \tilde{X}_1, \ldots, \tilde{X}_{M'}$ . The resulting supervised learning-based estimator of  $\rho^{\text{SL,Test}}$  is given by

1. Nested expectation form:

$$\hat{\rho}_{M,N,M'}^{\text{SL,Test}} = \frac{1}{M'} \sum_{i=1}^{M'} h(\hat{L}_{M,N}^{\text{SL}}(\tilde{X}_i)); \quad \tilde{X}_i \sim F_X.$$
(2.6)

2. VaR:

$$\hat{\rho}_{M,N,M'}^{\mathrm{SL},\mathrm{Test}} = \hat{L}_{(\lceil \alpha M' \rceil)}^{\mathrm{SL}}.$$

3. CVaR:

$$\hat{\rho}_{M,N,M'}^{\mathrm{SL},\mathrm{Test}} = \hat{L}_{(\lceil \alpha M' \rceil)}^{\mathrm{SL}} + \frac{1}{(1-\alpha)M'} \sum_{i=1}^{M'} \max\{\hat{L}_{M,N}^{\mathrm{SL}}(\tilde{X}_i) - \hat{L}_{(\lceil \alpha M' \rceil)}^{\mathrm{SL}}, 0\},$$

where  $\hat{L}_{(\lceil \alpha M' \rceil)}^{\operatorname{SL}}$  is the  $\lceil \alpha M' \rceil$ -th order statistic of  $\hat{L}_{M,N}^{\operatorname{SL}}(\tilde{X}_1), \ldots, \hat{L}_{M,N}^{\operatorname{SL}}(\tilde{X}_{M'})$ . Note that  $\hat{L}_{M,N}^{\operatorname{SL}}(\cdot)$  is the model preditions of a supervised learning model trained on the training samples  $(X_1, \hat{L}_{N,1}), \ldots, (X_M, \hat{L}_{N,M})$ .

Existing literature on nested simulation procedures has proposed different supervised learning metamodels to approximate the true function  $L(\cdot)$ . In this study, we focus on the metamodeling approaches that include theoretical convergence results. Methods that include theoretical convergence results are regression (Broadie et al., 2015), kernel smoothing (Hong et al., 2017), and KRR (Wang et al., 2022). Their estimators of  $L(\cdot)$  are given by  $\hat{L}_{M,N}^{\rm REG}(\cdot)$ ,  $\hat{L}_{M,N}^{\rm KS}(\cdot)$ , and  $\hat{L}_{M,N}^{\rm KRR}(\cdot)$ , respectively.

• Regression:

$$\hat{L}_{M,N}^{\text{REG}}(X) = \Phi(X)\hat{\beta},$$

where  $\Phi$  is a chosen basis, and  $\hat{\beta}$  is estimated from the training samples. In Broadie et al. (2015), the convergence analysis is performed on the expection Equation 2.4 and for smooth and Lipschitz continuous h. Connections between the expectation and the MSE are discussed in the Appendix.

• Kernel smoothing:

$$\hat{L}_{M,N}^{KS}(X) = \frac{\sum_{i=1}^{M} \hat{L}_{N,i} K_w(X - X_i)}{\sum_{i=1}^{M} K_w(X - X_i)},$$

where  $K_w$  is the kernel function with bandwidth w. Nadaraya (1964) and Watson (1964) originally proposed this kernel smoothing method for nonparametric regression, and it is widely known in the literature as Nadaraya-Watson kernel regression. Hong et al. (2017) provides convergence analysis for metamodel predictions of the training samples (Equation 2.5), and the convergence analysis is provided for smooth, hockey-stick, and indicator h.

• kNN (Mack, 1981):

$$\hat{L}_{M,N}^{\text{kNN}}(x) = \frac{\frac{1}{MR_M^d} \sum_{i=1}^M \hat{L}_{N,i} K\left(\frac{x - X_i}{R_M}\right)}{\frac{1}{MR_M^d} \sum_{i=1}^M K\left(\frac{x - X_i}{R_M}\right)}$$

where  $R_k$  is the Euclidean distance between  $X_i$  and its k-th nearest neighbor, and  $K: \mathbb{R}^d \to \mathbb{R}$  is a bounded, nonnegative kernel function satisfying

$$\int K(u)du = 1,$$
 
$$K(u) = 0 \quad \text{for } ||u|| \ge 1,$$

where  $\|\cdot\|$  is the Euclidean norm, and k is the number of nearest neighbors to consider. The k-nearest neighbor (KNN) method is implemented as a metamodel for numerical experiments in Hong et al. (2017), but the convergence analysis is not provided.

• KRR:

$$\hat{L}_{M,N}^{\text{KRR}}(X) = \operatorname*{arg\,min}_{g \in \mathcal{N}_{\Psi}(\Omega)} \left( \frac{1}{M} \sum_{i=1}^{M} (\hat{L}_{N,i} - L_i)^2 + \lambda ||L||_{\mathcal{N}_{\Psi}(\Omega)}^2 \right),$$

where  $\mathcal{N}_{\Psi}(\Omega)$  is the reproducing kernel Hilbert space (RKHS) with kernel  $\Psi$  defined domain  $\Omega$ , and  $\lambda$  is the regularization parameter as in ridge regression. More specifically,  $\Phi$  is a Matérn kernel with smoothness parameter  $\nu$  and length scale parameter  $\nu$ . Wang et al. (2022) provides convergence analysis for the AE (Equation 2.8), and the convergence analysis is provided for smooth, Lipschitz continuous, and indicator h, VaR, and CVaR.

These supervised learning metamodels approximate the inner simulation model  $L(\cdot)$  with functions in the form of  $\hat{L}_{M,N}^{\rm SL}(\cdot)$ , and the metamodels are used to pool the inner replications from all outer scenarios to estimate the risk measure  $\rho$ . In simulation literature, using models to approximate simulation is often referred to as the metamodeling approach, or it is referred to as a metamodel in the financial engineering literature. Supervised learning models are broadly classified into regression and classification models depending on whether the traget variable is continuous or discrete. In our context, the target variable L is continuous, and the supervised learning models refer to regression models. Regression models can be further categorized into parametric and non-parametric regression. Among our interested supervised learning models, regression is a parametric regression model, while kernel smoothing and kernel ridge regression are non-parametric regression model.

We are interested in minimizing the MSE of the supervised learning-based nested simulation estimator with supervised learning  $\hat{\rho}_{M,N}^{\mathrm{SL,Teat}}$  and  $\hat{\rho}_{M,N,M'}^{\mathrm{SL,Test}}$  subject to the total simulation budget  $\Gamma$ , and we are interested in the order of convergence of the estimator for all nested simulation procedures when the total simulation budget  $\Gamma$  approaches infinity.

$$\min_{M,N} \qquad \text{MSE}(\hat{\rho}_{M,N}^{\text{SL}}) = \mathbb{E}\left[\left(\hat{\rho}_{M,N}^{\text{SL}} - \rho\right)^2\right]$$
 subject to 
$$M \cdot N = \Gamma \qquad (2.7)$$

For the more complex metamodels, the convergence analysis of their MSEs is more challenging. More specifically, the KRR-based nested simulation (Wang et al., 2022) provides a convergence analysis in terms of absolute error (AE).

$$\min_{M,N} \quad \text{AE}(\hat{\rho}_{M,N}^{\text{SL}}) = \mathbb{E}\left[\left|\hat{\rho}_{M,N}^{\text{KRR}} - \rho\right|\right]$$
 subject to 
$$M \cdot N = \Gamma \tag{2.8}$$

Section 2.3.1 draws the connection between the MSE and the AE of the estimator in the context of nested simulation procedures.

#### 2.2.4 Likelihood Ratio Method

Instead of using a supervised learning model as a metamodel, Zhang et al. (2022) uses the likelihood ratio weights to pool the inner replications from all outer scenarios. Here, we restrict our attention to problems in the nested expectation form whose outer scenarios characterize the stochasticity of the inner simulation model. Specifically,

$$Y = Y(H, X),$$

where H is a random variable whose distribution is specified by the outer scenarios X. We denote the conditional distribution of H|X by  $f_{H|X}$ . For a specific scenario  $X_i$ , we write  $f_{H|X}(\cdot|X_i)$ . To reconcile with previously established notations, we note that inner simulation outputs  $Y_{ij}$  can be written as

$$Y_{ij} = Y(H_{ij}, X_i),$$

where  $H_{ij} \sim f_{H|X}(\cdot|X_i)$ . Suppose that one can generate random variable H from some sampling distribution  $f_H$ . Then, the likelihood ratio estimator of  $\rho$  is given by

$$\hat{\rho}_{M,N}^{\text{LR}} = \frac{1}{M} \sum_{i=1}^{M} h(\hat{L}_{N,i}^{\text{LR}}),$$

where the inner replications are pooled by the likelihood ratio weights with

$$\hat{L}_{N,i}^{LR} = \frac{1}{N} \sum_{i=1}^{N} Y(H_j, X_i) \frac{f_{H|X}(H_j|X_i)}{f_H(H_{ij})}, \quad H_j \sim f_H, \quad i = 1, \dots, M.$$

In Section 2.5.4, we implement the likelihood ratio method and share its fast convergence in finite-sample experiments despite the costly computation of the likelihood ratio weights (Section 2.6).

#### 2.2.5 Problem Statement

With a total simulation budget  $\Gamma$ , we are interested in the orders of convergence of estimators for all nested simulation procedures, which are measured by their MSE about the risk measure  $\rho$ .

min 
$$\mathbb{E}\left[\left(\hat{\rho}_{M,N} - \rho\right)^2\right]$$
  
subject to  $M \cdot N = \Gamma$ , (2.9)

where  $\Gamma$  is the total simulation budget for a nested simulation procedure. A special case is the MLMC method, where the total simulation budget  $\Gamma$  is the sum of the simulation budgets at all levels.

# 2.3 Asymptotic Analysis

In the first part of this section, we summarize the existing asymptotic convergence results of the nested simulation procedures in the literature, and we compare their critical assumptions that guarantee the convergence. The remaining of this section aims to fill the gap in the literature by

- providing the asymptotic convergence results for the standard nested simulation procedure,
- showing the connections between the convergence in MSE and the convergence in probabilistic order for AE in the context of nested simulation procedures, and
- illustrating the asymptotic rate of convergence of a kNN-based nested simulation procedure.

Estimator for $L$	Smooth h	$\mathbf{Lipschitz}(\mathbf{hockey\text{-}stick}\ h)$	$\mathbf{Indicator}\ h$	VaR	CVaR
Standard MC	*	<b>★</b> ( <b>√</b> )	✓	✓	×
Multi-level MC	×	$\times(\times)$	✓	×	×
Regression	✓	$\checkmark(\checkmark)$	×	×	×
Kernel smoothing	✓	×( <b>√</b> )	✓	×	×
Kernel ridge regression	♦	×(◊)	♦	$\Diamond$	$\Diamond$
Likelihood ratio	✓	×( <b>√</b> )	✓	×	×

Table 2.2: Existing asymptotic convergence results of nested simulation procedures for MSE

Table 2.2 summarizes the existing asymptotic convergence results of nested simulation procedures for MSE in the literature. A  $\checkmark$  indicates there exists an asymptotic convergence result for the corresponding estimator, a  $\times$  indicates there does not exist an asymptotic convergence result, a  $\star$  indicates an asymptotic result is not available in the literature but is provided in this study, and a  $\diamondsuit$  indicates the asymptotic convergence result exists in the literature but in a weaker form. Gordy and Juneja (2010) provide the asymptotic

convergence results for the standard nested simulation procedure with hockey-stick h, indicator h and VaR. Their CVaR analysis is incomplete as the VaR is assumed to be known but not estimated in the convergence proof. When the VaR is known, the CVaR analysis reduces to the nested expectation form with h being a hockey-stick function. Giles and Haji-Ali (2019) provide the asymptotic convergence results for the MLMC method with an indicator h. Broadie et al. (2015) provide the asymptotic convergence results for the regression-based nested simulation procedure with smooth h and Lipschitz continuous h. The Lipschitz continuous family includes the hockey-stick function as a special case, thus the convergence result for the hockey-stick function is implied. Hong et al. (2017) and Zhang et al. (2022) provide the asymptotic convergence results with the nested expectation form for the kernel smoothing-based procedure and likelihood ratio-based nested simulation procedure, respectively. The convergence results for a Lipschitz continuous h cannot be directly inferred from the analysis for a hockey-stick function.

While most of the literature focuses on the MSE of the estimator of  $\rho$ , Wang et al. (2022) analyze the asymptotic convergence of the estimator of  $\rho$  in terms of AE. Let  $\hat{\rho}$  be the estimator of  $\rho$ . Its AE about  $\rho$  is defined as

$$AE(\hat{\rho}) = |\hat{\rho} - \rho|.$$

In Wang et al. (2022), the authors of the KRR-based nested simulation procedures claim to have bridged the gap between the cubic and square root convergence rates of nested simulation procedures. However, they analyze convergence in probabilistic order, and it is only applicable in terms of AE. Instead of showing the convergence of the KRR-based estimator in terms of MSE as in Gordy and Juneja (2010), we show the connections between the convergence in MSE and the convergence in probabilistic order for AE. Our findings in Section 2.3.1 show that the analysis of Wang et al. (2022) indeed bridges the gap, but only in terms of convergence in probabilistic order for AE.

## 2.3.1 Connections between Convergence in MSE and AE

This section establishes the connections between the convergence in MSE and the convergence in probabilistic order for AE in the context of nested simulation procedures. In order to show the connections between the convergence in MSE and the convergence in probabilistic order for AE, we first need to state the definition for a sequence of random variables to converge in those two forms.

**Definition 4** Let  $\hat{\rho}_{\Gamma}$  be an estimator of  $\rho$  with a simulation budget of  $\Gamma$ . We write  $\mathbb{E}\left[\left(\hat{\rho}_{\Gamma}-\rho\right)^{2}\right]=\mathcal{O}\left(\Gamma^{-\xi}\right)$ , that is,  $\hat{\rho}_{\Gamma}$  converges in MSE to  $\rho$  in order  $\xi$  if

$$\exists C \limsup_{M} \frac{\mathbb{E}\left[\left(\hat{\rho}_{\Gamma} - \rho\right)^{2}\right]}{\Gamma^{-\xi}} \leq C$$

**Definition 5** Let  $\hat{\rho}_{\Gamma}$  be an estimator of  $\rho$  with a simulation budget of  $\Gamma$ . We write  $|\hat{\rho}_{\Gamma} - \rho| = \mathcal{O}_{\mathbb{P}}(\Gamma^{-\xi})$ , that is  $\hat{\rho}_{\Gamma}$  converges in probabilistic order  $\xi$  to  $\rho$  if for a sufficiently large  $\Gamma$ ,

$$\forall \epsilon > 0 \exists C \mathbb{P} \left( |\hat{\rho}_{\Gamma} - \rho| \ge C \Gamma^{-\xi} \right) \le \epsilon$$

We start by showing the convergence in probabilistic order from the convergence in MSE. Let  $\hat{\rho}_{\Gamma}$  be an estimator of  $\rho$  with a simulation budget of  $\Gamma$ , and assume that  $\mathbb{E}\left[(\hat{\rho}_{\Gamma}-\rho)^2\right]=\mathcal{O}\left(\Gamma^{-\xi}\right)$ . Then, from the definition of convergence in MSE, there exists a constant C such that

$$\limsup_{M} \frac{\mathbb{E}\left[\left(\hat{\rho}_{\Gamma} - \rho\right)^{2}\right]}{\Gamma^{-\xi}} \leq C.$$

Hence, there exists some  $\Gamma$  such that for all  $\gamma \geq \Gamma$ ,

$$\mathbb{E}\left[\left(\hat{\rho}_{\gamma} - \rho\right)^{2}\right] \leq C\gamma^{-\xi}.$$

The convergence in probabilistic order can be shown by separating the expectation into two parts: tail and non-tail.

$$\mathbb{E}\left[\left(\hat{\rho}_{\gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{\left|\hat{\rho}_{\gamma}-\rho\right|\leq d\gamma^{s}\right\}}\right]+\mathbb{E}\left[\left(\hat{\rho}_{\gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{\left|\hat{\rho}_{\gamma}-\rho\right|>d\gamma^{s}\right\}}\right]\leq C\gamma^{-\xi}.$$

The first term is always positive, and the second term can be bounded from below by the indicator function.

$$\mathbb{E}\left[\left(\hat{\rho}_{\gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{|\hat{\rho}_{\gamma}-\rho|>d\gamma^{s}\right\}}\right] \geq \mathbb{E}\left[d^{2}\gamma^{2s}\cdot\mathbb{I}\left\{|\hat{\rho}_{\gamma}-\rho|>d\gamma^{s}\right\}\right]$$

$$=d^{2}\gamma^{2s}\cdot\mathbb{E}\left[\mathbb{I}\left\{|\hat{\rho}_{\gamma}-\rho|>d\gamma^{s}\right\}\right]$$

$$=d^{2}\gamma^{2s}\cdot\mathbb{P}\left(|\hat{\rho}_{\gamma}-\rho|>d\gamma^{s}\right)$$

Combining bounds on the two terms, we have

$$d^2\gamma^{2s}\mathbb{P}\left(|\hat{\rho}_{\gamma} - \rho| > d\gamma^s\right) \le C\gamma^{-\xi}.$$

Let  $s = -\frac{\xi}{2}$ . Arranging the terms, we have

$$\mathbb{P}\left(|\hat{\rho}_{\gamma} - \rho| > d\gamma^{-\frac{\xi}{2}}\right) \le \frac{C}{d^2}$$

Hence, for all  $\epsilon > 0$ , there exist  $C^* = \sqrt{\frac{C}{\epsilon}}$  such that for all  $\gamma \geq \Gamma$ ,

$$\mathbb{P}\left(|\hat{\rho}_{\gamma} - \rho| > C^* \gamma^{-\frac{\xi}{2}}\right) \le \epsilon$$

In essence, the above is the definition of convergence in probabilistic order, that is,

$$|\hat{
ho}_{\gamma}-
ho|=\mathcal{O}_{\mathbb{P}}\left(\Gamma^{-rac{\xi}{2}}
ight)$$

**Theorem 1** Let  $\hat{\rho}_{\Gamma}$  be an estimator of  $\rho$  with a simulation budget of  $\Gamma$ . If  $\hat{\rho}_{\Gamma}$  converges in MSE to  $\rho$  in order  $\xi$ , then  $\hat{\rho}_{\Gamma}$  converges in probabilistic order to  $\rho$  in order  $\frac{\xi}{2}$ .

To the best of our knowledge, Theorem 1 has not been explicitly stated in the literature. It is the first result that shows the connections between the convergence in MSE and the convergence in probabilistic order for AE in the context of nested simulation. Theorem 1 is a general result that can be applied to any nested simulation procedure that converges in MSE to  $\rho$  in order  $\xi$ . If the estimator converges in MSE to  $\rho$  in order  $\xi$ , then it converges in probabilistic order to  $\rho$  in order  $\frac{\xi}{2}$ .

While the convergence in MSE implies the convergence in probabilistic order, the converse is not necessarily true. Similarly, the above argument is applied in reverse. Let  $\hat{\rho}_{\Gamma}$  be an estimator of  $\rho$  with a simulation budget of  $\Gamma$ , and assume that  $|\hat{\rho}_{\Gamma} - \rho| = \mathcal{O}_{\mathbb{P}}(\Gamma^{-\xi})$ . The MSE of  $\hat{\rho}_{\Gamma}$  can be separated into the same two parts.

$$\mathbb{E}\left[\left(\hat{\rho}_{\Gamma}-\rho\right)^{2}\right] = \mathbb{E}\left[\left(\hat{\rho}_{\Gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{|\hat{\rho}_{\Gamma}-\rho|< d\Gamma^{-2\xi}\right\}}\right] + \mathbb{E}\left[\left(\hat{\rho}_{\Gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{|\hat{\rho}_{\Gamma}-\rho|> d\Gamma^{-2\xi}\right\}}\right],$$

where the first term can be bounded from above.

$$\begin{split} \mathbb{E}\left[\left(\hat{\rho}_{\Gamma}-\rho\right)^{2}\cdot\mathbb{I}_{\left\{|\hat{\rho}_{\Gamma}-\rho|>d\Gamma^{s}\right\}}\right] &\leq d^{2}\Gamma^{-2\xi}\cdot\mathbb{E}\left[\mathbb{I}\left\{|\hat{\rho}_{\Gamma}-\rho|>d\Gamma^{s}\right\}\right] \\ &= d^{2}\Gamma^{-2\xi}\cdot\mathbb{P}\left(|\hat{\rho}_{\Gamma}-\rho|>d\Gamma^{s}\right) \leq d^{2}\Gamma^{-2\xi} \end{split}$$

However, the second term is not always bounded. If the random variable  $\hat{\rho}_{\Gamma}$  admits a density function f, then the second term can be further decomposed.

$$\mathbb{E}\left[\left(\hat{\rho}_{\Gamma} - \rho\right)^{2} \cdot \mathbb{I}_{\left\{|\hat{\rho}_{\Gamma} - \rho| > d\Gamma^{-2\xi}\right\}}\right] = \int_{-\infty}^{-d\Gamma^{-2k}} (x - \rho)^{2} f(x) dx + \int_{d\Gamma^{-2\xi}}^{\infty} (x - \rho)^{2} f(x) dx$$

Hence,  $\hat{\rho}_{\Gamma}$  converges in MSE to  $\rho$  in order  $2\xi$  if and only if both integrals converge in order higher than  $2\xi$ . The above argument shows that the convergence in probabilistic order does not necessarily imply the convergence in MSE. Hence, the converse of Theorem 1 is not necessarily true, and the convergence in probabilistic order is a weaker form of convergence than the convergence in MSE. The immediate implication is that the results in Wang et al. (2022), which show the convergence in probabilistic order for the AE, is not necessarily equivalent to the convergence in MSE.

# 2.3.2 Asymptotic Analysis for the Standard Nested Simulation Procedure

In Gordy and Juneja (2010), the authors analyze the asymptotic convergence of the standard nested simulation procedure in terms of MSE. The analysis is complete for the nested expectation form where h is either an indicator function or a hockey-stick function and VaR. For the nested expectation form where h is a smooth function or a Lipschitz continuous function, the analysis is incomplete. In this section, we fill in the holes for the analysis of Gordy and Juneja (2010).

**Assumption 1** h(L) has finite second moment, i.e.,  $\mathbb{E}\left[\left(h(L)\right)^2\right] < \infty$ .

Assumption 1 is a standard assumption in the literature (Hong et al., 2017) for the analysis of nested simulation procedures to analyze the convergence of the variance of  $\hat{\rho}_{M,N}$ .

**Assumption 2**  $\hat{L}_N(X) = L(X) + \bar{Z}_N(X)$ , where the simulation noise  $\bar{Z}_N(X)$  has zero mean and variance  $\nu(X)/N$ , where the conditional variance  $\nu(X)$  is bounded, i.e.,

$$\sup_{x \in \mathcal{X}} \nu(x) \le C_{\nu,1} < \infty.$$

For simplicity, we abbreviate  $\hat{L}_N(X)$  as  $\hat{L}_N$ ,  $\bar{Z}_N(X)$  as  $\bar{Z}_N$ , and  $\bar{Z}_{N,i}$  as  $\bar{Z}_{N,i}$ . Let  $\rho_M = \frac{1}{M} \sum_{i=1}^M h(L_i)$  be the nested MC estimator with the true function g. The MSE of the standard nested simulation procedure can be decomposed into two ways.

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N}-\rho\right)^{2}\right] \leq 2\mathbb{E}\left[\left(\hat{\rho}_{M,N}-\rho_{M}\right)^{2}\right]+2\mathbb{E}\left[\left(\rho_{M}-\rho\right)^{2}\right] \\
=2\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)-\frac{1}{M}\sum_{i=1}^{M}h\left(L_{i}\right)\right)^{2}\right] \\
+2\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(L_{i}\right)-\mathbb{E}\left[h(L)\right]\right)^{2}\right] \\
=2\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)-h\left(L_{i}\right)\right)^{2}\right]+\frac{2}{M}\mathrm{Var}(h(L)) \\
=2\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)-h\left(L_{i}\right)\right)^{2}\right]+\mathcal{O}(M^{-1}), \tag{2.10}$$

In 2.10, the last equality follows from Assumption 1. The analysis of the first term is different for smooth and Lipschitz continuous h. We will analyze them separately.

Another way to decompose the MSE is the bias-variance decomposition as in Gordy and Juneja (2010).

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N} - \rho\right)^{2}\right] = \mathbb{E}\left[\left(\hat{\rho}_{M,N} - \mathbb{E}\left[\hat{\rho}_{M,N}\right]\right)^{2}\right] + \left(\mathbb{E}\left[\hat{\rho}_{M,N}\right] - \rho\right)^{2}$$
$$= \operatorname{Var}(\hat{\rho}_{M,N}) + \operatorname{Bias}^{2}(\hat{\rho}_{M,N})$$
(2.11)

where  $\operatorname{Var}(\hat{\rho}_{M,N})$  is the variance of the estimator and  $\operatorname{Bias}(\hat{\rho}_{M,N})$  is the bias of the estimator about  $\rho$ . For a smooth function h, both ways of decomposing the MSE lead to the same order of convergence. However, for a Lipschitz continuous function h, Equation 2.11 leads to a stronger order of convergence than Equation 2.10. In the following, we analyze the convergence of the standard nested simulation procedure for smooth and Lipschitz continuous h using following both decompositions. Their critical assumptions and convergence results are summarized and compared.

#### A Smooth Function h

**Assumption 3** The function h has bounded first and second order derivative, i.e.,

$$\sup_{x \in \mathcal{X}} |h'(x)| \le C_{h,1} < \infty,$$
  
$$\sup_{x \in \mathcal{X}} |h''(x)| \le C_{h,2} < \infty.$$

Assumption 3 is similar to the smoothness assumption in Wang et al. (2022). Since h is a smooth function, Taylor expansion can be applied to the first term in Equation 2.10.

$$\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)-h\left(L_{i}\right)\right)^{2}\right]$$

$$=\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h'\left(L_{i}\right)\left(\hat{L}_{N,i}-L_{i}\right)+\frac{1}{2M}\sum_{i=1}^{M}h''\left(z_{i}\right)\left(\hat{L}_{N,i}-L_{i}\right)^{2}\right)^{2}\right]$$

$$\leq 2\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h'\left(L_{i}\right)\left(\hat{L}_{N,i}-L_{i}\right)\right)^{2}\right]$$

$$+2\mathbb{E}\left[\left(\frac{1}{2M}\sum_{i=1}^{M}h''\left(z_{i}\right)\left(\hat{L}_{N,i}-L_{i}\right)^{2}\right)^{2}\right]$$

$$(2.12)$$

where the last inequality is due to  $2ab \le a^2 + b^2$  for any  $a, b \in \mathbb{R}$ . The two terms on the right-hand side of Equation 2.12 are analyzed separately. We start with the first term  $S_1$ .

$$S_{1} = \mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h'(L_{i})\left(\hat{L}_{N,i} - L_{i}\right)\right)^{2}\right]$$

$$= \mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h'(L_{i})\bar{Z}_{N,i}\right)^{2}\right]$$

$$\leq C_{1}^{2}\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}\bar{Z}_{N,i}\right)^{2}\right]$$

$$= C_{1}^{2}\mathbb{E}\left[\frac{1}{M^{2}}\sum_{i=1}^{M}\sum_{j=1}^{M}\bar{Z}_{N,i}\bar{Z}_{N,j}\right]$$

$$= C_{1}^{2}\mathbb{E}\left[\frac{1}{M^{2}}\sum_{i=1}^{M}\bar{Z}_{N,i}^{2} + \frac{1}{M^{2}}\sum_{i=1}^{M}\sum_{j\neq i}^{M}\bar{Z}_{N,i}\bar{Z}_{N,j}\right]$$

$$\leq \frac{C_{1}^{2}C_{\nu,1}}{MN} = \mathcal{O}(M^{-1}N^{-1})$$
(2.13)

where the last inequality in Equation 2.13 is due to Assumption 2, independence of  $X_i$  and  $X_j$  for  $i \neq j$ , and the fact that the inner simulation noise has zero mean, i.e.,  $\mathbb{E}\left[\bar{Z}_N\right] = 0$ . It remains to analyze the second term  $S_2$ , where Assumption 4 is necessary to ensure the existence of the fourth moment of the simulation noise and the convergence of the second term.

**Assumption 4** The fourth moment of simulation noise  $\bar{Z}_N(X)$  follows  $\mathbb{E}\left[\left(\bar{Z}_N(X)\right)^4\right] = \nu_2(X)/N^2$ , where  $\nu_2(X)$  is bounded, i.e., there exists  $C_{\nu,2} > 0$  such that  $\nu_2(X) \leq C_{\nu,2}$  for all  $x \in \mathbb{R}$ .

$$S_{2} = \mathbb{E}\left[\left(\frac{1}{2M}\sum_{i=1}^{M}h''(z_{i})\left(\hat{L}_{M,N}^{SL}(X_{i}) - L_{i}\right)^{2}\right)^{2}\right]$$

$$= \mathbb{E}\left[\left(\frac{1}{2M}\sum_{i=1}^{M}h''(z_{i})\bar{Z}_{N,i}^{2}\right)^{2}\right]$$

$$\leq C_{2}^{2}\mathbb{E}\left[\left(\frac{1}{2M}\sum_{i=1}^{M}\bar{Z}_{N,i}^{2}\right)^{2}\right]$$

$$= C_{2}^{2}\mathbb{E}\left[\frac{1}{2M^{2}}\sum_{i=1}^{M}\sum_{j=1}^{M}\bar{Z}_{N,i}^{2}\bar{Z}_{N,j}^{2}\right]$$

$$= C_{2}^{2}\mathbb{E}\left[\frac{1}{2M^{2}}\sum_{i=1}^{M}\sum_{j=1}^{M}\bar{Z}_{N,i}^{4} + \frac{1}{2M^{2}}\sum_{i=1}^{M}\sum_{j\neq i}^{M}\bar{Z}_{N}^{2}(X_{i})\bar{Z}_{N}^{2}(X_{j})\right]$$

$$\leq C_{2}^{2}\left(\frac{C_{\nu,2}M}{2M^{2}N^{2}} + \frac{C_{\nu,1}^{2}M(M-1)}{2M^{2}N^{2}}\right) = \mathcal{O}(N^{-2})$$
(2.14)

where the second inequality is due to Assumption 3, and the last inequality follows from Assumption 4 and the fact that  $\hat{L}_N$  is a standard MC estimator of L. Combining Equation 2.10, 2.12, 2.13, and 2.14, we have

$$\mathbb{E}\left[ (\hat{\rho}_{M,N} - \rho)^2 \right] = \mathcal{O}(M^{-1}) + \mathcal{O}(N^{-2})$$
 (2.15)

Setting  $M = \mathcal{O}(\Gamma^{2/3})$  and  $N = \mathcal{O}(\Gamma^{1/3})$ , we provide the same rate of convergence as obtained for other risk measures in Gordy and Juneja (2010). However, we observe that Equation 2.13 converges in the order of  $\mathcal{O}(M^{-1}N^{-1})$ . The cross term here is potentially problematic (Why?), as setting N as a negative power of M may result in Equation 2.13 to be divergent.

In order to avoid the cross term, we proceed to follow the bias-variance decomposition in Equation 2.11.

$$\operatorname{Bias}(\hat{\rho}_{M,N}) = \mathbb{E}\left[\hat{\rho}_{M,N}\right] - \rho$$

$$= \mathbb{E}\left[\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)\right] - \mathbb{E}\left[h\left(L\right)\right]$$

$$= \mathbb{E}\left[h\left(\hat{L}_{N}\right) - h\left(L\right)\right]$$

$$= \mathbb{E}\left[h'\left(L\right)\left(\hat{L}_{N} - L\right) + \frac{1}{2}h''\left(z\right)\left(\hat{L}_{N} - L\right)^{2}\right]$$
(2.16)

where the last equality follows from the Taylor expansion of the smooth function h. The first and second order Taylor expansion terms are analyzed separately in Equation 2.17 and Equation 2.18.

$$\mathbb{E}\left[h'(L)\left(\hat{L}_{N}-L\right)\right] = \mathbb{E}\left[h'(L)\left(\hat{L}_{N}-L\right)\right]$$

$$= \mathbb{E}\left[\mathbb{E}\left[h'(L)\left(\hat{L}_{N}-L\right)|X\right]\right]$$

$$= \mathbb{E}\left[h'(L)\mathbb{E}\left[\hat{L}_{N}-L|X\right]\right] = 0$$
(2.17)

$$\mathbb{E}\left[h''(z)\left(\hat{L}_{N}-L\right)^{2}\right] \leq \mathbb{E}\left[\left|h''(z)\right|\left(\hat{L}_{N}-L\right)^{2}\right]$$

$$\leq C_{h,2}\mathbb{E}\left[\left(\hat{L}_{N}-L\right)^{2}\right]$$

$$= C_{h,2}\operatorname{Var}(\hat{L}_{N}) = \mathcal{O}\left(N^{-1}\right)$$
(2.18)

It remains to analyze the variance term in Equation 2.11.

$$\operatorname{Var}(\hat{\rho}_{M,N}) = \frac{1}{M} \operatorname{Var}(h(\hat{L}_N)) = \mathcal{O}(M^{-1})$$
(2.19)

Equation 2.19 directly follows from Assumption 1 and the fact that  $\hat{L}_N$  is a standard MC estimator of L. Combining Equation 2.17, 2.18, and 2.19, we are able to show the convergence order of the standard nested simulation procedure for a smooth function h without the cross term as in Equation 2.13.

**Theorem 2** Let h be a smooth function. MSE of the standard nested simulation procedure converges in order  $\Gamma^{2/3}$ , that is,

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N}-\rho\right)^{2}\right]=\mathcal{O}(\Gamma^{-2/3}).$$

The proof techniques used in deriving Theorem 2 is completely different from the one used in Gordy and Juneja (2010). The analysis in Gordy and Juneja (2010) is based on the differentiablity of the joint density of Y and the average inner simulation noise, which is difficult to verify in practice. Instead, we use the Taylor expansion of the smooth function h to analyze the convergence of the standard nested simulation procedure. The critical assumption in our derivation is Assumption 4, which is necessary to ensure the existence of the fourth moment of the simulation noise and the convergence of  $S_2$ , the second order term in the Taylor expansion. Assumption 4 is a moment condition that is easier to verify in practice than conditions on the joint density. As stated in Theorem 1, the convergence in MSE automatically implies the convergence in probabilistic order.

Corollary 1 Let h be a smooth function. The AE of the standard nested simulation procedure converges in probabilistic order  $\Gamma^{-1/3}$ , that is,

$$|\hat{\rho}_{M,N} - \rho| = \mathcal{O}_{\mathbb{P}}(\Gamma^{-1/3}).$$

#### A Lipschitz Continuous Function h

We proceed to analyze the convergence for a Lipschitz continuous function h. For the CVaR analysis in Gordy and Juneja (2010), the authors assume the knowing of the corresponding VaR. Hence, the analysis of CVaR is not complete. Instead, the quantity that is being analyzed is instead the mean excess loss, which corresponds to h being a hockey-stick function and belongs to the family of the Lipschitz continuous functions. Here we present the convergence analysis for the whole family of the Lipschitz continuous functions, which includes the hockey-stick function as a special case.

**Assumption 5** The function h is Lipschitz continuous. Hence,  $|h(x_1) - h(x_2)| \le K|x_1 - x_2|$  for some constant  $K < \infty$ .

Assumption 5 is a standard assumption for analysis involving Lipschitz continuous functions, and it is also used in Broadie et al. (2015). For the Lipschitz continuous case, the first term in Equation 2.10 is analyzed differently.

$$\mathbb{E}\left[\left(\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)-h\left(L_{i}\right)\right)^{2}\right] \leq \mathbb{E}\left[\left(h\left(\hat{L}_{N}\right)-h\left(L\right)\right)^{2}\right]$$

$$\leq K^{2}\mathbb{E}\left[\left(\hat{L}_{N}-L\right)^{2}\right]$$

$$=K^{2}\mathbb{E}\left[\left(\bar{Z}_{N}(X)\right)^{2}\right]=\mathcal{O}(N^{-1}) \qquad (2.20)$$

where the first inequality follows from Cauchy-Schwarz inequality, the second equality follows from Assumption 5, and the last equality follows from Assumption 2.

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N} - \rho\right)^2\right] = \mathcal{O}(M^{-1}) + \mathcal{O}(N^{-1}) \tag{2.21}$$

Setting  $M = \mathcal{O}(\Gamma^{1/2})$  and  $N = \mathcal{O}(\Gamma^{1/2})$ , we provide a looser bound than the one obtained for hockey-stick h. Using the bias-variance decomposition in Equation 2.11,

$$\operatorname{Bias}(\hat{\rho}_{M,N}) = \mathbb{E}\left[\hat{\rho}_{M,N}\right] - \rho$$

$$= \mathbb{E}\left[\frac{1}{M}\sum_{i=1}^{M}h\left(\hat{L}_{N,i}\right)\right] - \mathbb{E}\left[h\left(L\right)\right]$$

$$= \mathbb{E}\left[h\left(\hat{L}_{N}\right) - h\left(L\right)\right]$$

$$\leq \mathbb{E}\left[\left|h\left(\hat{L}_{N}\right) - h\left(L\right)\right|\right]$$

$$\leq K\mathbb{E}\left[\left|\hat{L}_{N} - L\right|\right]$$

$$= K\mathbb{E}\left[\left|\bar{Z}_{N}\right|\right]$$

$$\leq K\sqrt{\mathbb{E}\left[\left(\bar{Z}_{N}\right)^{2}\right]} = \mathcal{O}(N^{-1/2})$$
(2.22)

The variance term in Equation 2.11 for Lipschitz continuous h is identical to the one in Equation 2.19.

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N} - \rho\right)^2\right] = \mathcal{O}(M^{-1}) + \mathcal{O}(N^{-1}) \tag{2.23}$$

Setting  $M = \mathcal{O}(\Gamma^{2/3})$  and  $N = \mathcal{O}(\Gamma^{1/3})$ , we provide the same rate of convergence as obtained for other risk measures in Gordy and Juneja (2010).

**Theorem 3** Let h be a Lipschitz continuous function. MSE of the standard nested simulation procedure converges in order  $\Gamma^{2/3}$ , that is,

$$\mathbb{E}\left[\left(\hat{\rho}_{M,N}-\rho\right)^{2}\right]=\mathcal{O}(\Gamma^{-1/2}).$$

Immediately from Theorem 1, the convergence in MSE automatically implies the convergence in probabilistic order.

Corollary 2 Let h be a Lipschitz continuous function. AE of the standard nested simulation procedure converges in probabilistic order  $\Gamma^{-1/3}$ , that is,

$$|\hat{\rho}_{M,N} - \rho| = \mathcal{O}_{\mathbb{P}}(\Gamma^{-1/4}).$$

## 2.3.3 Asymptotic Analysis of a kNN-based Nested Simulation Procedure for a Smooth Function

In Hong et al. (2017), the authors analyze the asymptotic convergence in terms of MSE for a kernel smoothing-based nested simulation procedure. The analysis is complete for the nested expectation form where h is either a smooth function, an indicator function, or a hockey-stick function. The kernel function of interest in the asymptotic analysis is the Nadaraya-Watson kernel. Nevertheless, the numerical results are derived based on a kNN-based nested simulation procedure. In this section, we attempt to fill in the holes in the analysis of Hong et al. (2017) by providing the asymptotic convergence of a kNN-based nested simulation procedure for smooth h in the nested expectation form. Surprisingly, the asymptotic convergence rate of the kNN-based nested simulation procedure is noticeably different from the result obtained in Hong et al. (2017).

A kNN regression can be used as a non-parametric metamodel that estimates the conditional expectation of Y given X by averaging the Y values of the k nearest neighbors of X. Without loss of generality, we assume that  $\{(X_i, \bar{Y}_{n,i}), i = 1, \ldots, M\}$  are independent and identically distributed (i.i.d.) observations of (X, Y) from the standard nested simulation procedure with an inner simulation budget of n, where  $X_i$  is the i<sup>th</sup> of the outer scenario  $X \in \Omega \subset \mathbb{R}^d$ , and  $\bar{Y}_{n,i}$  is the estimate of  $Y|X_i$  with an inner simulation budget of n. Let  $f_{XY}(x,y)$  be the joint density of random vector (X,Y), and  $f_X(x) = \int f_{XY}(x,y) dy$  be the

marginal density of X. Let k be a sequence of positive integers k = k(M) such that

$$k \to \infty$$

$$\frac{k}{M} \to 0, \quad \text{as } M \to \infty.$$

$$\frac{\log(M)}{k} \to 0, \quad \text{as } k \to \infty.$$

A kNN-based nested simulation procedure with model parameter k estimates the risk measure  $\rho = \mathbb{E}[h(L)]$  with

$$\hat{L}_{M,N}^{\text{kNN}}(x) = \frac{\frac{1}{MR_M^d} \sum_{i=1}^M \hat{L}_{N,i} K\left(\frac{x - X_i}{R_M}\right)}{\frac{1}{MR_M^d} \sum_{i=1}^M K\left(\frac{x - X_i}{R_M}\right)}$$

where  $R_k$  is the Euclidean distance between  $X_i$  and its k-th nearest neighbor, and  $K: \mathbb{R}^d \to \mathbb{R}$  is a bounded, nonnegative kernel function satisfying

$$\int K(u)du = 1,$$

$$K(u) = 0 \quad \text{for } ||u|| \ge 1,$$

where  $\|\cdot\|$  is the Euclidean norm, and k is the number of nearest neighbors to consider. For a kNN estimator trained with M i.i.d. observations, denote  $B_M(x)$  and  $V_M(x)$  as its bias and variance at point x, respectively.

**Assumption 6**  $f_X$  is bounded and continuously differentiable up to second order in a neighborhood of x with  $f_X(x) > 0$ ,  $f_X$ ,  $\mathbb{E}[Y^2|X=x] < \infty$ , and the kernel function K satisfies

$$\int ||u||^2 |K(u)| du < \infty,$$

$$\int \nu_{\alpha} K(u) du = 0, \quad \text{for } \alpha = 1, \dots, d.$$

Suppose  $\mathbb{P}(\|x - X\| > r) = \mathcal{O}(r^{-\alpha})$  for some  $\alpha > 0$  as  $r \to \infty$ .

The following lemma from Mack (1981) characterizes the pointwise convergence of the kNN estimator in terms of its bias and variance at point x.

**Lemma 1** Assume the random vector (X, Y) and the kernel function K satisfies Assumption 6. Then the bias and variance of the kNN estimator at point x are given by

$$B_{M}(x) = A_{kNN}(x) \cdot \left( \left( \frac{k}{M} \right)^{\frac{2}{d}} + o\left( \frac{k}{M} \right)^{\frac{2}{d}} \right) + B_{kNN} \cdot \left( \frac{1}{k} + o\left( \frac{1}{k} \right) \right),$$

$$V_{M}(x) = \frac{C_{kNN} \cdot Var[Y|X=x]}{k} \int K^{2}(u)du + o\left( \frac{1}{k} \right).$$

Let h be a smooth function, i.e., Assumption 3 holds. The bias of the estimator of  $\rho$  for a kNN-based nested simulation procedure is analyzed following Taylor expansion.

$$\mathbb{E}\left[h(\hat{L}_{M,n}^{\text{kNN}}(X)) - h(L)\right]$$

$$= \mathbb{E}\left[h'(L)\left(\hat{L}_{M,n}^{\text{kNN}}(X) - L\right) + \frac{1}{2}h''(z)\left(\hat{L}_{M,n}^{\text{kNN}}(X) - L\right)^{2}\right]$$
(2.24)

where the bias is decomposed into two terms, and z is between  $\hat{L}_{M,n}^{\text{kNN}}(X)$  and L. The first term in Equation 2.24 is

$$\mathbb{E}\left[h'(L)\left(\hat{L}_{M,n}^{kNN}(X) - L\right)\right] 
= \mathbb{E}\left[h'(L)\mathbb{E}\left[\hat{L}_{M,n}^{kNN}(X) - L|X\right]\right] 
= \int_{\Omega} h'(L)B_{M}(x)f_{X}(x)dx 
= \int_{\Omega} h'(L)\left(A_{kNN}(x)\left(\frac{k}{M}\right)^{2/d}(1 + o_{x}(1)) + B_{kNN}\cdot\left(\frac{1}{k} + o_{x}\left(\frac{1}{k}\right)\right)\right)f_{X}(x)dx 
= \left(\frac{k}{M}\right)^{2/d}\mathbb{E}\left[h'(L)A_{kNN}(X)\right](1 + o(1)) + \frac{B_{kNN}}{k}\mathbb{E}\left[h'(L)\right](1 + o(1)).$$
(2.25)

The second term in Equation 2.24 is

$$\mathbb{E}\left[h''(z)\left(\hat{L}_{M,n}^{kNN}(X) - L\right)^{2}\right] \\
= \int_{\Omega} h''(z)\left(B_{M}^{2}(X) + V_{M}(x)\right)f_{X}(x)dx \\
\leq C_{2}\int_{\Omega}\left(\frac{C_{kNN}\cdot\operatorname{Var}[Y|X=x]}{k}\int K^{2}(u)du + o\left(\frac{1}{k}\right)\right)f_{X}(x)dx \\
+ C_{2}\int_{\Omega}\left(A_{kNN}(x)\left(\frac{k}{M}\right)^{2/d}(1+o_{x}(1)) + B_{kNN}\cdot\left(\frac{1}{k}+o_{x}\left(\frac{1}{k}\right)\right)\right)^{2}f_{X}(x)dx \tag{2.26}$$

where the second term in Equation 2.26 is of higher order than Equation 2.25, and it is negligible since  $kM^{-1} \to 0$  as  $M \to \infty$  and  $k \to \infty$ , The first term in Equation 2.26 is  $\mathcal{O}\left(\frac{1}{k}\right)$ . Set  $k = \mathcal{O}(M^{\frac{2}{2+d}})$  and  $M = \mathcal{O}(\Gamma)$ , the bias of the kNN-based nested simulation procedure converges in the order of  $\Gamma^{-\frac{2}{2+d}}$ .

**Definition 6** Let h be a smooth function. The bias of the kNN-based nested simulation procedure converges in the order of  $\Gamma^{-\frac{2}{2+d}}$ , that is,

$$\mathbb{E}\left[h(\hat{L}_{M,n}^{kNN}(X)) - h(L)\right] = \mathcal{O}(\Gamma^{-\frac{2}{2+d}}).$$

Definition 6 is obtained by matching the orders of the bias terms in Equation 2.25 and 2.26. It is the tightest bound that can be obtained for the convergence of the kNN-based nested simulation procedure with a smooth function h. The variance is left for future work. It implies that the kNN-based nested simulation procedure converges at most in the order of  $\Gamma^{-\frac{2}{3}}$  for d=1 and has slower convergence than the standard procedure for  $d \geq 2$ . Asymptotically, it has slower convergence than the result obtained in Hong et al. (2017) for the Nadaraya-Watson kernel. This discrepancy underscores the critical importance of aligning theoretical analysis with numerical illustrations. The slower convergence rate observed with the kNN approach not only challenges the robustness of empirical findings derived under differing assumptions but also signals a cautionary note for practitioners. This alignment is crucial for ensuring the reliability and validity of conclusions drawn from such procedures, especially in the context of risk management and financial decision-making.

# 2.4 Convergence Orders and Critical Assumptions of Nested Simulation Procedures

In this section, we summarize the convergence orders and the critical assumptions of the nested simulation procedures. The results are summarized in 2.3, where the orders of convergence are presented in the order of the total simulation budget  $\Gamma$ .

Estimator for L	Smooth h	Lipschitz / hockey-stick h	Indicator h	VaR	CVaR
Standard MC	$\mathcal{O}(\Gamma^{-2/3})$	$\mathcal{O}(\Gamma^{-1/2}) / \mathcal{O}(\Gamma^{-2/3})$	$\mathcal{O}(\Gamma^{-2/3})$	$\mathcal{O}(\Gamma^{-2/3})$	×
Multi-level MC	×	× / ×	$\mathcal{O}(\Gamma^{-1} \log(\Gamma) )$	×	×
Regression	$\mathcal{O}(\Gamma^{-1})$	$\mathcal{O}(\Gamma^{-1})/\mathcal{O}(\Gamma^{-1})$	×	×	×
Kernel smoothing	$\mathcal{O}(\Gamma^{-\min\{1,\frac{4}{d+2}\}})$	$\times / \mathcal{O}(\Gamma^{-\min\{1,\frac{4}{d+2}\}})$	$\mathcal{O}(\Gamma^{-\min\{1,\frac{4}{d+2}\}})$	×	×
Kernel ridge regression	$\mathcal{O}(\Gamma^{-1})$	$\times$ / $\mathcal{O}(\Gamma^{-1})$	$\mathcal{O}(\Gamma^{-1})$	$\mathcal{O}(\Gamma^{-1})$	$\mathcal{O}(\Gamma^{-1})$
Likelihood ratio	$\mathcal{O}(\Gamma^{-1})$	$\times$ / $\mathcal{O}(\Gamma^{-1})$	$\mathcal{O}(\Gamma^{-1})$	×	×

Table 2.3: Asymptotic rate of convergence of nested simulation procedures in MSE

The asymptotic convergence rate of the nested simulation procedures is highly dependent on the assumptions on the random variable X, Y, and the inner simulation noise. As more advanced supervised learning techniques are used to estimate the function g, a more restrictive set of assumptions is required to ensure the asymptotic convergence of the inner estimators. In this section, we summarize the pivotal assumptions regarding the random variables and the inner simulation noise for all nested simulation procedures. They are presented in an order reflecting their relative ease in satisfying the assumptions.

# 2.4.1 Standard Assumptions

Gordy and Juneja (2010) and Broadie et al. (2015) assume that the inner simulation noise has zero mean and variance that is inversely proportional to the inner simulation budget N, i.e., Assumption 2. The analysis in Gordy and Juneja (2010) further requires assumptions on  $f_{Y,\tilde{Z}_N}(y,z_N)$ , the joint density of random variable Y and the normalized inner simulation noise  $\tilde{Z}_N := \sqrt{N} \cdot \bar{Z}_N(X)$ .

# 2.4.2 Assumptions on Joint Density

**Assumption 7** The joint density  $f_{Y,\tilde{Z}_N}(y,z_N)$  and its partial derivatives  $\frac{\partial f_{Y,\tilde{Z}_N}(y,z_N)}{\partial y}$  and  $\frac{\partial^2 f_{Y,\tilde{Z}_N}(y,z)}{\partial y^2}$  exist.

**Assumption 8** For  $N \ge 1$ , there exist non-negative functions  $p_{0,N}(\cdot)$ ,  $p_{1,N}(\cdot)$ , and  $p_{2,N}(\cdot)$  such that for all y and z,

$$\begin{vmatrix}
f_{Y,\tilde{Z}_N}(y,z) \leq p_{0,N}(z) \\
\left| \frac{\partial f_{Y,\tilde{Z}_N}(y,z)}{\partial y} \right| \leq p_{1,N}(z) \\
\left| \frac{\partial^2 f_{Y,\tilde{Z}_N}(y,z)}{\partial y^2} \right| \leq p_{2,N}(z).$$

In addition,

$$\sup_{N} \int_{-\infty}^{\infty} |z|^{r} p_{i,N}(z) dz < \infty \tag{2.27}$$

for i = 0, 1, 2 and  $r \in [0, 4]$ .

Assumption 7 and 8 are necessary for the Taylor expansion up to the second order on the joint density  $f_{Y,\tilde{Z}_N}(y,z_N)$ , and they can easily be satisfied by perturbing Y and  $\tilde{Z}_N$  with a zero-mean normal random variable with a small variance. However, Assumption 8 is hard to be verified in practice. Since Broadie et al. (2015) only analyze the nested expectation case with smooth and Lipschitz continuous function h, the regression-based nested simulation procedure does not require Assumption 7 and 8 for the convergence analysis. In our analysis of the standard nested simulation procedure for smooth h in the nested expectation form, we make additional assumptions on the fourth moment of the inner simulation noise, i.e., Assumption 3, so the second-order term in the Taylor expansion can be bounded.

# 2.4.3 Assumptions for the MLMC Procedure

In addition to Assumption 8, the MLMC procedure requires assumptions on the random variable  $Q := \frac{|\mathbb{E}[Y|X]|}{\text{Var}[Y|X]}$ .

**Assumption 9**  $f_Q$ , the density of Q exists, and there exist positive constants  $q_0$  such that  $f_Q(q) \leq q_0$  for all  $q \in [0, q_0]$ .

Assumption 9 ensures that the conditional expectation of Y given X is not concentrated around zero, as Giles and Haji-Ali (2019) show the asymptotic convergence of the MLMC procedure only when  $\rho$  is in the nested expectation form with h being an indicator function, i.e.,  $h(x) = \mathbb{I}_{\{x>0\}}$ .

#### 2.4.4 Assumptions for the Likelihood Ratio-Based Procedure

In addition to Assumption 8, the likelihood ratio-based nested simulation procedure requires assumptions on the marginal density of H, the random variable that characterizes the stochasticity of the inner simulation model. More specifically, for the likelihood ratio-based nested simulation procedure, the random variable H is introduced to represent the inner simulation noise, and Y can be directly expressed as a function of H and X, i.e., Y = Y(H, X). Instead of simulating Y directly, the random variable H is simulated from the conditional distribution  $f_{H|X}(y|x)$ , and Y is then obtained by evaluating Y(H, X).

**Assumption 10** The marginal density  $f_H(y)$  of H exists.  $f_H(y)$  can be sampled from and evaluated, and  $Y(y,x)f_{H|X}(y|x) = 0$  whenever  $f_H(x) = 0$ .

Furthermore, the likelihood ratio-based nested simulation procedure requires the samples of H to be independent with the outer scenario X.

**Assumption 11** The inner sample  $H \sim f_H(y)$  is independent of X. Samples  $\{X_i, i = 1, ..., M\}$  and  $\{H_i, i = 1, ..., N\}$  are i.i.d. samples from  $f_X(x)$  and  $f_H(y)$ , respectively.

Assumption 11 is necessary for  $\hat{L}_{N}^{LR}(X_{i})$  to be an unbiased and strongly consistent estimator of  $L_{i}$ , which is a necessary condition for the convergence of  $\hat{\rho}_{M,N}^{LR}$ , the likelihood ratio-based nested simulation estimator of  $\rho$ . Assumption 10 and Assumption 11 are likely to be satisfied in practice, as one can simulate H separately from X and evaluate Y(H,X) for each pair of (H,X).

# 2.4.5 Assumptions for the Kernel-Based Procedure

For the kernel-based nested simulation procedure, Hong et al. (2017) simplifies the analysis by assuming that for each  $X_i$ , the inner simulation budget N is fixed together with the variance of the inner simulation noise  $\bar{Z}_{N,i}$ . In essence, this fixed variance design treats the  $(X_i, \bar{Y}_{N,i})$  as independent and identically distributed observations of (X, Y), and the kernel-based nested simulation procedure is analyzed as a nonparametric regression problem. Regularity conditions on  $f_X(x)$ , the density of X, and the conditional second moment of Y are assumed to ensure the convergence of  $\hat{L}_{M,N}^{KS}(X)$ , the kernel smoothing estimator of L.

**Assumption 12**  $f_X(x)$  and  $\mathbb{E}[Y^2|X=x]$  exist, f(x) > 0, and f(x) is thrice continuously differentiable at x with bounded third-order derivative.

Further assumptions are made on the kernel function  $K_w$  and the bandwidth w to ensure the convergence of the kernel smoothing estimator  $\hat{L}_{M,N}^{\text{KS}}(X)$ . A similar set of assumptions is made in Jennen-Steinmetz and Gasser (1988) for the convergence analysis of a kernel smoothing estimator with kernel functions of higher orders.

### 2.4.6 Assumptions for the KRR-Based Procedure

For the KRR-based nested simulation procedure, the convergence is guaranteed only when  $\bar{Z}_N(X)$  is the sum of N i.i.d. zero-mean sub-Gaussian random variables. The sub-Gaussian assumption imposes a stronger condition on the inner simulation noise than all the other nested simulation procedures. The inner simulation noise is assumed to have lighter tails than the normal distribution, while examples of random variables that satisfy the sub-Gaussian assumption are hard to find. In addition, the following assumptions are made on  $\Omega$ , the domain of the outer scenario X:

**Assumption 13**  $\Omega$  is a bounded subset of  $\mathbb{R}^d$ , and  $f_X(x)$  is bounded above and below away from zero.

Assumption 13 rules out the possibility of the outer scenario X being a normal random variable, which is a common assumption in the literature. Furthermore, the convergence analysis of the KRR-based nested simulation procedure is shown in terms of AE rather than in MSE, which is due to the complexity of the KRR metatmodel.

# 2.5 Finite-Sample Experiments

The theoretical framework has allowed the comparison of the asymptotic convergence behavior for different nested simulation procedures. However, a simulation budget in practice is almost always finite. In this section, we conduct a series of numerical experiments to compare the empirical convergence of the nested simulation procedures for different risk measures, option types, and asset dimensions. Numerical experiments are conducted on portfolios that consist of options on d underlying assets, whose dynamics follow a multidimensional geometric Brownian motion with 0.3 pairwise correlation.

A total of 5 nested simulation procedures are considered, namely the standard nested simulation procedure, the regression-based, kernel smoothing-based, likelihood ratio-based, and kernel ridge regression-based nested simulation procedures. For the standard nested simulation procedure, the bootstrap-based budget allocation strategy from Zhang et al. (2021) is implemented to estimate the optimal values of outer and inner simulation budgets. For the regression metamodel, the Laguerre polynomials up to degree 3 are used as the default basis functions. The kNN metamodel is implemented for the kernel smoothing-based procedure, and a cross-validation with grid search is used to find the optimal number of neighbors. The KRR metamodel is implemented with a Matérn kernel, and the smoothness, the length scale, and the regularization hyperparameters are found by a cross-validation with Bayesian search (Frazier, 2018). Due to budget constraints, the hyperparameters of the nonparametric regression metamodel are tuned and used across all macro replications of the same experiment.

The procedures are compared for 5 types of risk measures:

- a quadratic tracking error,
- a mean excess loss over a threshold u,
- a probability of a large loss over a threshold u,
- the VaR, and
- the CVaR,

where the threshold u for the mean excess loss and the probability of a large loss is set to be the 90% VaR. The procedures are compared for different portfolios consisting of different types of options:

- Portfolio 1 consists of d assets. The portfolio contains 3 European call options written on each asset with strikes 90, 100, and 110, respectively.
- Portfolio 2 consists of d assets. The portfolio contains 3 geometric Asian options written on each asset with strikes 90, 100, and 110, respectively.
- Portfolio 3 consists of d assets. The portfolio contains 3 up-and-out barrier call options written on each asset with strikes 90, 100, and 110, respectively. They have a barrier level of 120.

- Portfolio 4 considers d assets. The portfolio contains 3 down-and-out barrier call options written on each asset with strikes 90, 100, and 110, respectively. They have a barrier level of 90.
- Portfolio 5 contains 1 asset. The portfolio longs 2 down-and-out barrier put options and shorts 1 barrier down-and-out put option.

Except for portfolio 5, 5 different asset dimensions are considered, i.e., d=1,2,5,10,20. 525 experiment settings that arise from an exhaustive combination of the above simulation procedures, risk measures, portfolios, and asset dimensions are generated. For each experiment setting, the estimation of the risk measure is repeated 1000 times to obtain the empirical MSE of the corresponding estimator under a range of simulation budgets. The total simulation budget  $\Gamma$  ranges from 10,000 to 10,240,000 in a geometric progression with a common ratio of 2. The empirical convergence results of each estimator are measured and recorded to illustrate their empirical behavior under different risk measures, option types, and asset dimensions. We start by examining the empirical convergence results for the most basic case, i.e., the quadratic tracking error risk measure for European call options on Portfolio 1 with d=1.

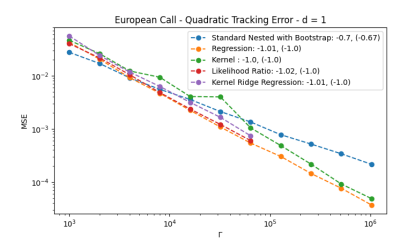


Figure 2.1: Empirical convergence of nested simulation procedures for quadratic tracking error on Portfolio 1 with d=1

In Figure 2.1, the MSEs of the nested simulation procedures are plotted against the total simulation budget  $\Gamma$  in a log-log scale, where each point represents the average MSE of the corresponding estimator over 1000 macro replications of the same experiment. Due

to the difficulty of fixing  $\Gamma$  for the MLMC procedure, the empirical rate of convergence of the MLMC procedure is not reported here, but a detailed analysis of the empirical convergence of the MLMC procedure is provided in Section 2.5.7.

A regression line is fitted to the MSEs of each nested simulation procedure against the simulation budget  $\Gamma$  in a log-log scale, and the slope of the fitted line is reported. The slopes of the fitted lines can be interpreted as the empirical rates of convergence. Alongside in the parentheses are the asymptotic rates of convergence obtained by the theoretical analysis in Section 2.3. By comparing the empirical rates of convergence with the asymptotic rates, we can observe how well the empirical rates of convergence match their asymptotic rates. In the case illustrated in Figure 2.1, the empirical rates of all procedures closely match their asymptotic rates.

Except for the standard nested simulation procedure, the empirical rates of convergence of the other procedures are close to  $\Gamma^{-1}$ , i.e., all of them achieve similar finite-sample performance that is better than the standard procedure. Pooling of inner simulation samples across different outer scenarios is the key reason behind their higher empirical rates of convergence. However, pooling also introduces additional computational cost that is not reflected in the simulation budget  $\Gamma$ . Due to the computational complexity of the likelihood ratio-based and the KRR-based procedures, their MSEs are not reported for  $\Gamma$  larger than 16,000 and 64,000, respectively. Detailed discussion of the additional computational cost of pooling is provided in Section 2.6.

In the following sections, we examine the empirical convergence of the nested simulation procedures in a similar manner as in Figure 2.1. With a more detailed analysis of different risk measures, option types, and asset dimensions, we aim to provide a comprehensive understanding and comparison of the convergence behavior of the nested simulation procedures in practice.

## 2.5.1 Sensitivity to the Asset Dimension

In portfolio risk management, the asset dimension is a critical factor that determines the complexity of the portfolio and the computational cost of the risk measure estimation. The theoretical analyses in Section 2.4 suggest that the asymptotic rate of convergence of the nested simulation procedures, except for the kernel smoothing-based procedure, is independent from the asset dimension. However, in practice, their empirical convergence behavior could be different. In this section, we examine the sensitivity to the asset dimension of the empirical convergence of the nested simulation procedures.

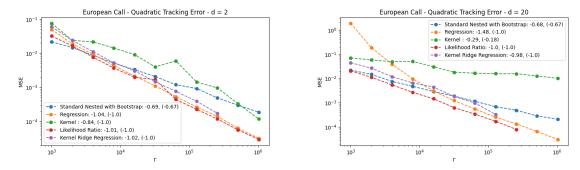


Figure 2.2: Empirical convergence of nested simulation procedures for quadratic tracking error on Portfolio 1 with different asset dimensions

In Figure 2.2, the empirical convergence of the nested simulation procedures for a quadratic tracking error on Portfolio 1 is illustrated for different asset dimensions, i.e., d=2 and d=20. The empirical rates of convergence of the standard, the KRR-based, and the likelihood ratio-based procedures closely match their asymptotic rates for both asset dimensions, and they are insensitive to an increase in the asset dimension. Conversely, both the kernel smoothing-based and regression-based procedures are greatly affected by the asset dimension. Their sensitivities to the asset dimension differ in finite-sample experiments:

- For the kernel smoothing-based procedure, the empirical rate of convergence decreases as the asset dimension increases. This observation aligns with the theoretical analysis in Section 2.3, which demonstrates that the asymptotic rate of convergence of the kernel smoothing-based procedure is sensitive to the asset dimension. Nevertheless, even for d=2 and d=20, the empirical rate remains higher than its asymptotic rate.
- For the regression-based procedure, the asymptotic rate is not affected by the asset dimension, but the empirical rates of convergence are higher than the asymptotic rates for both asset dimensions, and the empirical rate is higher for d = 20 than for d = 2.

In the following sections, we provide detailed explanations for the discrepancy between their empirical rates of convergence and their asymptotic rates.

# 2.5.2 Empirical Convergence of Kernel Smoothing-Based Procedures

Kernel smoothing is a nonparametric regression metamodel. According to the theoretical analysis in Section 2.3, the asymptotic rate of convergence of the kernel smoothing-based nested simulation procedure is highly dependent on the asset dimension d. In this section, we use a kNN metamodel for the kernel smoothing-based nested simulation procedure, and we examine its empirical convergence for different asset dimensions.



Figure 2.3: Empirical convergence of kernel smoothing procedure for different values of d

In Figure 2.3, the empirical convergence of the kernel smoothing-based nested simulation procedure for quadratic tracking error is illustrated for different asset dimensions, i.e., d=1,2,5,10. The observation finds that the kernel smoothing-based nested simulation procedure is extremely sensitive to the asset dimension and the payoff structure. While the asset dimension is expected to be a critical factor as shown in the theoretical analysis, the payoff structure is an unexpected factor that affects the empirical convergence of the kernel smoothing-based nested simulation procedure. For a portfolio with geometric Asian options, the payoff complexity is higher than that of a portfolio with European call options. The empirical rate of convergence of the kernel smoothing-based procedure is significantly lower for the portfolio with geometric Asian options than for the portfolio with European call options. For geometric Asian options, the empirical rates of convergence are even lower than the asymptotic rates for all asset dimensions.

Due to the computational cost of the kernel smoothing-based nested simulation procedure, we are not able to conduct experiments for higher budget levels. However, we are able to conduct additional experiments to examine the effects of cross-validation on the empirical convergence of the kernel smoothing-based procedure. Another phenomenon that is observed in Figure 2.3 is that the empirical rate of convergence of the kernel smoothing-based procedure does not decrease monotonically as the simulation budget increases. This

is likely due to the fact that the kernel smoothing-based procedure, as a nonparametric regression procedure, is highly dependent on the cross-validation of the hyperparameters of the metamodel. For the kernel smoothing-based procedure, the kNN metamodel is implemented. Hence, the hyperparameter of interest is the number of nearest neighbors k. In our numerical experiment, cross-validation is conducted once to select the optimal value of k for each simulation budget, and the selected value of k is fixed for all 1000 replications. Therefore, a suboptimal selection of the optimal value of k can lead to a poor metamodel estimate, and the MSE of the kRR-based procedure will be dominated by the model error of the metamodel.

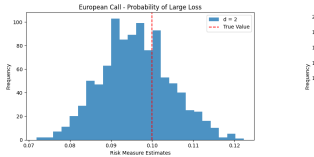




Figure 2.4: Cross-validation for the kernel smoothing-based procedure with  $\Gamma = 100,000$ 

In Figure 2.4, the empirical convergence of the kernel smoothing-based nested simulation procedure for the probability of large loss is illustrated for different values of k with  $\Gamma=100,000$ . The observation is that the value of optimal k estimated by cross-validation is highly variable across different replications. For d=2, the optimal value of k is estimated to be between 120 to 220 for most replications, and the estimated risk measures are close to the true risk measure of 0.1. For the two replications where k=80 is estimated as the optimal value of k, the resulting estimates of risk measures are 0.1187 and 0.1189, which are significantly higher than the estimated risk measures for the other replications and far from the true risk measure of 0.1. This observation suggests that the empirical convergence of the kernel smoothing-based nested simulation procedure is highly sensitive to the cross-validation of the hyperparameters of the metamodel. A KRR-based nested simulation procedure suffers from the same problem. The fluctuation in Figure 2.3 is likely due to poor cross-validation of the hyperparameters of the metamodel.

# 2.5.3 Empirical Convergence of Parametric Regression Procedures

For both the regression-based and kernel smoothing-based procedures, the observed higher empirical rates of convergence can be explained by the poor performance of their metamodels at low simulation budgets. When simulation budgets are low, the metamodels trained to approximate the true inner simulation are inadequate. The MSEs for these procedures are dominated by the model errors of the metamodels. In other words, they have not yet reached their asymptotic convergence regimes.

Due to computation constraints, we are not able to conduct experiments for kernel smoothing-based nested simulation procedures for higher budget levels, but we are able to conduct additional experiments for the regression-based nested simulation procedure. In our previous numerical experiments, the empirical rate of convergence of the regression procedure is observed to be much larger than its asymptotic rate of convergence. For dimensions larger than 10, the MSE of the regression procedure decreases quickly in the beginning, and its rate of decrease stabilizes after a certain budget level. In 2.5 illustrates the empirical convergence of the regression-based procedure in more detail at higher simulation budgets, where the asymptotic level of convergence is reached for d = 20.

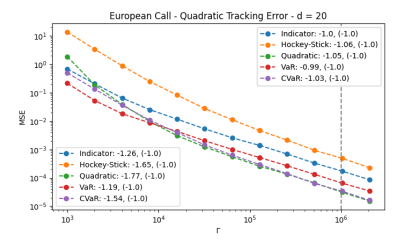


Figure 2.5: Empirical convergence of regression procedure for European call options and d=20

The left part of Figure 2.5 contains the MSEs of the regression procedure for budget sizes that are smaller than  $10^3$ . Slopes of the fitted lines on the left correspond to the

empirical rate of convergence of the regression procedure for budget levels between  $10^3$  and  $10^4$ . To investigate the convergence behavior for higher budget levels, we conduct additional experiments for the European call option with dimension 20. The additional experiments are summarized on the right side of Figure 2.5. After reaching a certain budget level, i.e.,  $\Gamma = 10^6$  in our case, the empirical rate of convergence for the regression-based nested simulation procedure approaches its asymptotic rate. For nested simulation procedures with a biased metamodel, the metamodel estimates of the true inner simulation are poor, especially for smaller budget sizes. We are able to clearly observe this phenomenon for the regression metamodel, and it can be explained by dividing the MSE into metamodeling bias and simulation variance.

- For small budget sizes, the improvement in the bias of the regression metamodel dominates the improvement in simulation variance. Performing poorly for extremely low simulation budgets, the regression metamodel improves significantly as the simulation budget increases.
- As the simulation budget gets higher, the improvement of the regression bias becomes negligible compared with that of the simulation variance. The regression metamodel ceases to improve after reaching a certain level ( $\Gamma = 10^6$  in our case), and the improvement of simulation variance dominates.

Due to high computational costs, we are not able to conduct experiments for even higher budget levels, but we expect the MSE of the regression-based nested simulation procedure to stabilize after  $\Gamma=10^6$ , and the empirical rate of convergence to approach its asymptotic rate. In the context of nested simulation procedures, a parametric regression metamodel has several advantages over a nonparametric regression metamodel.

- Parametric regression achieves a good empirical convergence behavior for a moderate simulation budget.
- Parametric regression is insensitive to changes in the asset dimension.
- Parametric regression is easier to implement. It does not require cross-validation to select the hyperparameters of the metamodel.

However, parametric regression suffers from the model misspecification risk. More specifically, selecting a wrong regression basis can lead to a poor metamodel estimate, where the MSE of the nested simulation procedure is dominated by the model error of the metamodel (Broadie et al., 2015).

In our previous numerical experiments, the regression-based nested simulation procedure is implemented with Laguerre polynomials up to degree 3 as the basis functions. It is a standard choice in the literature. To examine the sensitivity of the empirical convergence behavior to the regression basis, we consider different basis functions, i.e., regular polynomial regression up to degree 3.



Figure 2.6: Empirical convergence of regression-based nested simulation procedures for different regression bases

In Figure 2.6, the empirical convergence results of the regression-based nested simulation procedure for a quadratic tracking error and a 90%-CVaR are illustrated for the two different regression bases. The empirical rates of convergence of the regression-based nested simulation procedure are observed to be insensitive to the chosen regression basis.

# 2.5.4 Sensitivity to the Option Types and Risk Measures

In our previous numerical experiments, we have examined the empirical convergence behavior of the nested simulation procedures for European call options, which only depends on the asset price at maturity. To examine the sensitivity of the empirical convergence behavior to the option type, we consider path-dependent options, i.e., geometric Asian options and barrier options.

In Figure 2.7, the empirical convergence of the nested simulation procedures for the quadratic tracking errors of Portfolio 1 and Portfolio 4 is illustrated for d=20. The empirical rates of standard, likelihood ratio-based, and KRR-based nested simulation procedures closely ensemble their asymptotic rates for path-dependent options. The empirical rate of convergence of the regression-based and kernel smoothing-based procedures is much higher than their asymptotic rates for barrier options. The reasons are explained in Section 2.5.3 and Section 2.5.2.

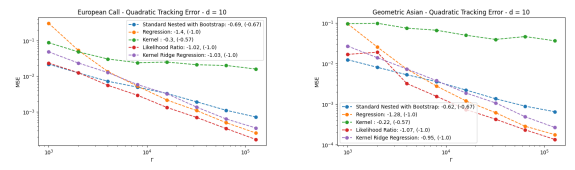


Figure 2.7: Empirical convergence of nested simulation procedures for quadratic tracking error on different portfolios with d=20

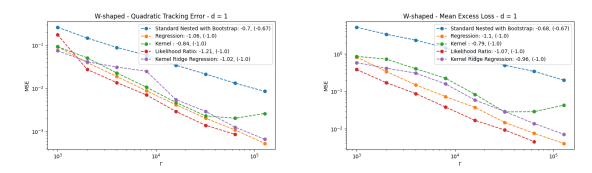


Figure 2.8: Empirical convergence of nested simulation procedures for a W-shaped payoff

In Broadie et al. (2015), the authors propose a numerical example where the payoff has a W-shape with respect to the asset price at maturity. We have incorporated this example as Portfolio 5. In Figure 2.8, the empirical convergence of the nested simulation procedures for quadratic tracking error and mean excess loss on Portfolio 5 is illustrated for d=1. For all procedures, we observe a similar convergence behavior as in other path-dependent options. The MSEs of the kernel smoothing-based and KRR-based procedures do not decrease monotonically as the simulation budget increases. This fluctuating behavior is observed and analyzed in Section 2.5.2.

Figure 2.9 illustrates similar observations for sensitivity to different risk measures. A change in the risk measure of interest does not affect the empirical convergence behavior of all nested simulation procedures. From the empirical convergence results, we observe that the empirical rates of convergence of the regression-based nested simulation procedure are the highest among all nested simulation procedures for all risk measures and option types.

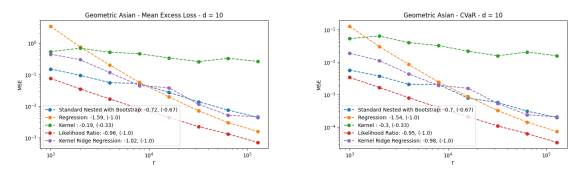


Figure 2.9: Empirical convergence of nested simulation procedures for different risk measures on Portfolio 1 with d=20

Furthermore, the regression-based procedure is stable across different payoff structures and risk measures. Its MSEs decrease quickly in the beginning, and after reaching a certain level of  $\Gamma$ , the rate of decrease stabilizes to match its asymptotic rate of convergence. Parametric regression is the most attractive metamodel for nested simulation procedures due to its fast empirical convergence behavior, stability across different payoff structures and risk measures, and ease of implementation. The following sections focus on the regression-based nested simulation procedure for the rest of the numerical experiments.

# 2.5.5 Sensitivity to level for VaR and CVaR

The level of VaR and CVaR is a critical factor that determines the complexity of the risk measure estimation. In our previous numerical experiments, the level of VaR and CVaR is set to be the 90% quantile of the distribution of the inner simulation noise. In this section, we examine the empirical convergence of the regression-based nested simulation procedures for other levels of VaR and CVaR, i.e., 80%, 95%, 99%, and 99.6%.

In Figure 2.10, the empirical convergence of the regression-based nested simulation procedure for different levels of VaR and CVaR is illustrated for up-and-out barrier call options. The empirical rates of convergence of the regression-based nested simulation procedure are not sensitive to the level of VaR and CVaR.

## 2.5.6 Sensitivity to the Asset Model

Our previous numerical experiments have been conducted under the assumption that the underlying asset dynamics follow a multidimensional geometric Brownian motion with 0.3

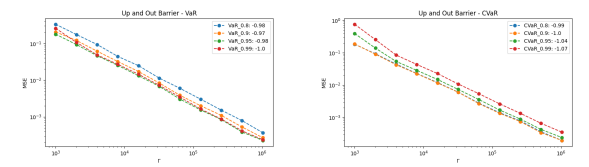


Figure 2.10: Empirical convergence of regression-based procedures for different levels of VaR and CVaR for Up and Out Barrier Call Options

pairwise correlation. To examine the sensitivity of the empirical convergence behavior to the asset model, we consider a stochastic volatility model, i.e., a Heston model.

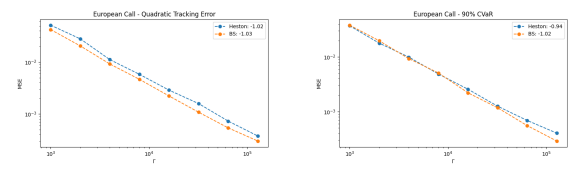


Figure 2.11: Empirical convergence of regression-based nested simulation procedures for different asset models

In Figure 2.11, the empirical convergence results of the regression-based nested simulation procedure for a quadratic tracking error and a 90%-CVaR are illustrated for different asset models, i.e., a geometric Brownian motion and a Heston model. The empirical rates of convergence of the regression-based nested simulation procedure are observed to be insensitive to the asset model.

In summary, we find the empirical convergence behavior of the regression-based nested simulation procedure to be stable across different risk measures, option types, asset dimensions, levels of VaR and CVaR, asset models, and regression bases. With a limited total budget, the regression-based nested simulation procedure is the most robust and stable among all nested simulation procedures.

#### 2.5.7 Empirical Convergence of MLMC

The MLMC procedure is a variance reduction technique that is designed to reduce the computational cost of nested simulation procedures. The theoretical analysis in Giles and Haji-Ali (2019) shows that the MLMC procedure has a faster empirical rate of convergence than the standard nested simulation procedure when the risk measure of interest is the probability of a large loss over a threshold u, i.e., the nested expectation case with h being an indicator function.

Level	Bias	Variance	MSE	Γ	MSE of SNS
0	0.118	0.104	0.1364	6400	0.1660
1	0.102	0.0916	0.1020	27600	0.1305
2	0.0870	0.0794	0.0870	76600	0.0954
3	0.0815	0.0746	0.0812	175600	0.0744
4	0.0893	0.0805	0.0884	348100	0.0460
5	0.0750	0.0694	0.0750	640000	0.0366

Table 2.4: MSEs of the MLMC procedure for different levels

We provide a summary of the empirical convergence results of the MLMC procedure with a decomposition table of the MSEs in Table 2.4. The MSEs of the MLMC procedure are decomposed into the bias and the variance of the estimator. The MSEs of the standard nested simulation procedure with a similar total simulation budget are also provided for comparison. Since the MLMC procedure is designed with a fixed number of outer simulation paths, the benefit of the MLMC procedure is minimal when the total simulation budget is large. More specifically, when  $\Gamma$  is large, the standard nested simulation procedure with a bootstrap-based budget allocation strategy benefits from having a larger number of outer simulation paths, and the MSE of the standard nested simulation procedure is lower than that of the MLMC procedure with a fixed number of outer simulation paths.

#### 2.6 Computational Complexity

Compared to the standard procedure, metamodel-based nested simulation procedures often have faster empirical rates of convergence. Their faster convergence benefits from the pooling of inner samples through the metamodels. However, pooling itself comes with computational costs, which are usually ignored in most numerical comparisons. This section summarizes the algorithmic complexity and illustrate the computational time of different nested simulation procedures. The findings can provide some further guidance on the choice of a proper nested simulation procedure given a nested estimation problem. Define basic operations to be basic mathematical operators, that is, arithmetic with individual elements has complexity  $\mathcal{O}(1)$ .

Procedures	Training cost	Prediction cost	Total additional cost
Standard procedure	0	$\mathcal{O}(M)$	$\mathcal{O}(M)$
Regression	$\mathcal{O}(p^2M) + \mathcal{O}(p^3)$	$\mathcal{O}(pM)$	$\mathcal{O}(p^2M) + \mathcal{O}(p^3)$
Kernel smoothing	$\mathcal{O}(M\log(M))$	$\mathcal{O}(k\log(M))$	$\mathcal{O}(M\log(M))$
Kernel ridge regression	$\mathcal{O}(M^3)$	$\mathcal{O}(M^2)$	$\mathcal{O}(M^3)$
Likelihood ratio	0	$\mathcal{O}(M^2)$	$\mathcal{O}(M^2)$

Table 2.5: Additional computational costs of nested simulation procedures aside from simulation

Table 2.5 shows the algorithmic complexity of the nested simulation procedures. For a d-dimensional nested estimation problem, the simulation cost for all procedures is  $\mathcal{O}(dMN)$ . The simulation cost is omitted from Table 2.5, as all procedures are given the same simulation budget. Given M outer samples and the corresponding M inner sample averages, the algorithmic complexity of the additional operations does not involve N thus can be written in terms of M only. The training cost includes the cost of fitting the metamodel and the cost of cross-validation for nonparametric regression metamodels.

For the standard procedure,

- the training cost is 0 due to the absence of a metamodel.
- Its prediction cost  $\mathcal{O}(M)$  comes from the evaluation of the function h for M samples.

For the regression-based procedure, a linear regression model is fitted with p predictors.

• The training cost is  $\mathcal{O}(p^2M) + \mathcal{O}(p^3)$ . Estimating the coefficients of the regression model involves multiplying the design matrix by its transpose and inverting the resulting matrix, which has complexity  $\mathcal{O}(p^2M)$  and  $\mathcal{O}(p^3)$ , respectively. For a review in the complexity of matrix inversion, see Stothers (2010). The matrix inversion has complexity  $\mathcal{O}(p^3)$  using Gauss-Jordan elimination, but in practice, the complexity can be reduced to  $\mathcal{O}(p^{2.807})$  by Strassen (1969) and  $\mathcal{O}(p^{2.376})$  by Coppersmith and Winograd (1987). In practice, p is usually much smaller than M and does not grow with the simulation budget  $\Gamma$ . p is fixed by the basis function, which is selected based on the complexity of the payoff function.

• The prediction cost of the regression-based procedure is  $\mathcal{O}(pM)$ , which comes from the multiplication of the design matrix by the estimated coefficients.

For the kernel smoothing-based procedure, the kNN metamodel is implemented. In practice, the K-D tree algorithm (Bentley, 1975) is often implemented for efficient nearest neighbor search.

- During training, a tree is constructed to store the distance, which has complexity  $\mathcal{O}(M\log(M))$ .
- The prediction cost of the kernel smoothing-based procedure is  $\mathcal{O}(kM\log(M))$ , which comes from a query of the K-D tree for k nearest neighbors for M samples. Conversely, An inefficient algorithm calculates the distance matrix during training and a block sort algorithm to find the k nearest neighbors, which results in a complexity of  $\mathcal{O}(M^2)$  and  $\mathcal{O}(M^2\log(M))$ , respectively. A practical implementation of block sort is provided in Kim and Kutzner (2008). Hence, an efficient implementation of the kNN metamodel is crucial for the kernel smoothing-based procedure.

For the likelihood ratio-based procedure,

- the training cost is 0 as no training is required.
- The prediction cost of the likelihood ratio-based procedure is  $\mathcal{O}(M^2)$ , which comes from the calculation of the likelihood weights for M samples.

For the KRR-based procedure,

- the main training cost comes from inverting an  $M \times M$  kernel matrix (Schölkopf and Smola, 2002), which has complexity  $\mathcal{O}(M^3)$  using Gauss-Jordan elimination. Similar to the regression-based procedure, this complexity can be reduced to  $\mathcal{O}(M^{2.376})$ .
- The prediction cost of the KRR-based procedure is  $\mathcal{O}(M^2)$ , which comes from the multiplication of the kernel matrix by the estimated coefficients.

Among the metamodel-based procedures, parametric regression is the most efficient. In regression, the number of features is usually much less than the number of data points, i.e., p < M. Kernel smoothing and KRR are kernel-based metamodels, and they are more computational expensive due to distance calculations and cross-validations of hyperparameters.

The kNN metamodel has only 1 hyperparameter k, while the KRR metamodel requires 3 hyperparameters, namely the smoothness hyperparameter  $\nu$ , scale hyperparameter  $\nu$ , and the regularization hyperparameter  $\lambda$ . Calculating the likelihood ratio weights is inevitable for the likelihood ratio-based procedure. While a k-fold cross-validation is used to estimate the hyperparameter for kNN, the KRR requires Bayesian optimization (Shahriari et al., 2015) to estimate the hyperparameters due to having a high-dimensional search space. The likelihood ratio-based procedure requires no training, but the cost of calculating the likelihood weights is  $\mathcal{O}(M^2)$ . Comparing the algorithmic complexity of the nested simulation procedures, the regression-based procedure is the most efficient among the metamodel-based procedures. For a fixed p, the total additional cost of a regression-based procedure is  $\mathcal{O}(M)$ , which is the same as the standard procedure's.

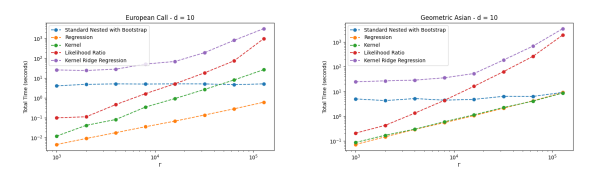


Figure 2.12: Total computational cost for different procedures with d=10

The cost associated with pooling makes a significant impact on the overall computational complexity of a metamodel-based nested simulation procedure. In our finite-sample experiments, we have indeed observed that the actual computational cost to deviate significantly from the simulation cost. In Figure 2.12, the total actual computational time of different nested simulation procedures for European call options and geometric Asian options with d=10 is illustrated in a log-log scale. The x-axis shows the total simulation budget  $\Gamma$ , and the y-axis is the total computational time for 1 replication of the numerical experiment, in seconds. All procedures are implemented on a machine with an AMD Ryzen 9 7900X processor with 32GB of RAM. 8 cores are used for parallel computing, and the total computational time is the sum of the simulation time, the training time, and the prediction time. The regression and kernel smoothing-based procedures are the most efficient among the metamodel-based procedures. Since the regression-based procedure has a higher empirical rate of convergence, it is preferred over the kNN. The likelihood ratio-based and KRR-based procedures are the most computationally expensive among all procedures. They are also demanding in terms of memory. For budgets larger than  $10^5$ , the likelihood ratio-based

procedure becomes impractical as storage of the likelihood weights becomes a bottleneck. The KRR-based procedure suffers from both cross-validation and inverting a kernel matrix of size  $M \times M$ . To separate the total computational time into different attributes, we provide a detailed analysis of the total computational time in the remainder of this section.

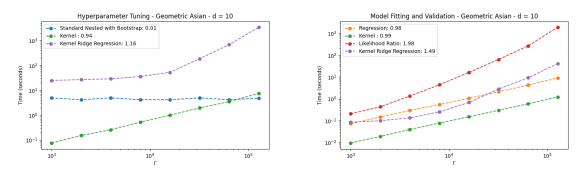


Figure 2.13: Computational cost for implementing nested simulation procedures with d = 10, excluding simulation time

Figure 2.13 illustrates the computational time of implementing different nested simulation procedures for the portfolio of geometric Asian options with d=10. The simulation time is omitted as it is the same for all procedures. The remaining computations can be decomposed into two parts: hyperparameter tuning and model implementation. The hyperparameter tuning cost is the cost of estimating the optimal hyperparameters for the metamodels, and the model implementation cost is the cost of fitting the metamodels and generating predictions from the trained metamodels. For each procedure, each point in Figure 2.13 represents the average computational time for a given simulation budget  $\Gamma$ . A regression model is fitted to each model respectively, and its slope can be viewed as the growth rate of the computational time with respect to  $\Gamma$ .

- The total computational time of the standard procedure is mostly attributed to finding the optimal M and N using the bootstrap-based budget allocation strategy from Zhang et al. (2021). This cost does not grow with the simulation budget  $\Gamma$ , as its slope is close to 0.
- The regression-based procedure does not require hyperparameter tuning, and its total computational time is mainly attributed to the model implementation. The slope of the regression line is close to 1, resembling the its algorithmic complexity in Table 2.5.
- The kernel smoothing-based and KRR-based procedure requires cross-validation to estimate the optimal hyperparameters. In our implementation of kNN, the cross-

validation is conducted using a grid search with a search space whose cardinality does not grow with the simulation budget  $\Gamma$ . However, the cross-validation cost of the kernel smoothing-based procedure grows with the simulation budget  $\Gamma$  as more samples are involved. The KRR-based procedure requires Bayesian optimization to estimate the optimal hyperparameters, and the cardinality of the search space also grows with the simulation budget  $\Gamma$ . The empirical cost of the KRR-based procedure does not resemble its algorithmic complexity in Table 2.5. The efficient implementation of KRR is an active area of research, and the computational cost of KRR can be reduced by using a low-rank approximation of the kernel matrix, e.g., the Nyström method (Nyström, 1930). Our observation for KRR implementation time is in line with the findings in Pedregosa et al. (2011), where the cost of KRR is reported quickly increasing with the number of samples.

• The likelihood ratio-based procedure requires no hyperparameter tuning, and its total computational time is mainly attributed to the likelihood weight calculation, which is  $\mathcal{O}(M^2)$ .

In summary, for a large simulation budget, the regression-based procedure is the most efficient among all metamodel-based procedures in terms of computational time. For moderate simulation budgets, the regression-based procedure should be the preferred choice.

#### 2.7 Conclusion and Extensions

In the task of estimating risk measures for portfolios of complex financial derivatives, nested simulation procedures are commonly required but often computationally expensive. Tremendous efforts have been made to improve the efficiency of nested simulation procedures by approximating the inner simulation model with a metamodels. In this study, we review the literature on nested simulation procedures in financial engineering and establish fair comparisons of different metamodels using the same set of numerical examples. Asymptotic properties of estimators for different nested simulation procedures are influenced by their corresponding metamodels. To show an asymptotic convergence result, a more complex metamodels requires a more stringent set of assumptions on the distribution of the outer scenario and the inner simulation noise. With extensive numerical experiments, we have found the finite-sample performance of a procedure can deviate from its theory. In theory, supervised learning-based nested simulation procedures often provide higher rates of convergence, but they come at the computational expense of model training and generating predictions from the trained models. The likelihood ratio-based procedure

requires no training, but it is computationally expensive to compute and store the likelihood weights. As a result, the total computational budget is not necessarily the same as the simulation budget, which is usually a limiting factor for practical applications. A kernel-based procedure, e.g., kNN and KRR, requires cross-validation, and its empirical performance depends heavily the choice of hyperparameters. A kNN-based procedure is sensitive to the asset dimension and the problem complexity. A KRR-based procedure is computationally expensive, and its cost grows quickly with the simulation budget. For kernel-based procedures, the computational cost is heavily dependent on the efficient implementation of the associated algorithms. For a nested estimation problem with a given computational budget, we suggest the use of the regression-based simulation procedure when the budget size is moderate. It is efficient to implement, and it exhibits fast empirical convergence in estimating risk measures for option portfolios. However, we have only examined the performance for option portfolios in this study, where finding a suitable set of basis functions for regression is relatively easy. In practice, a variable annuity contract is often high-dimensional in time with a complex payoff structure, and it is a challenging problem for nested simulation procedures. The time dependence in the feature space leads to multi-collinearity, and the regression-based procedure may not be the best choice. In such cases, a neural network-based procedure may be more suitable, as it is a data-driven algorithm that finds a suitable set of basis functions automatically. In Chapter 3, we examine the performance of regression-based and neural network-based nested simulation procedures for estimating risk measures for variable annuities.

### Chapter 3

# Cutting Through the Noise: Using Deep Neural Network Metamodels for High Dimensional Nested Simulation

#### 3.1 Introduction

Deep neural networks (DNNs) have attracted attentions of researchers and practitioners due to their success (Hastie et al., 2009; LeCun et al., 2015) in solving real-world machine learning tasks such as AlphaGo (Silver et al., 2016) and ChatGPT (OpenAI, 2023). Since the first artificial neural network model (McCulloch and Pitts, 1943) and the first algorithm for training a perceptron (Rosenblatt, 1958), especially after the introduction of backpropagation (Rumelhart et al., 1985) and the growth of high-performance computing, the field of artificial neural network and deep learning in general has grown rapidly. Two specialized neural network architectures that are relevant to our study are RNN (Williams and Zipser, 1989; Sutskever et al., 2014) and LSTM (Hochreiter and Schmidhuber, 1997; Chung et al., 2014), as In the field of actuarial science, a metamodel is commonly referred to as a proxy model. Despite their success, DNN models are often criticized for their lack of transparency and interpretability, which hinders their adoption in financial and actuarial applications. Enormous research efforts are spent to test and improve the robustness of DNN models with carefully designed noise injection methods. Poole et al. (2014) show that injecting synthetic noise before and after hidden unit activations during training im-

proves the performance of autoencoders. Neelakantan et al. (2015) improve learning for DNNs by injecting synthetic noise to the gradients during backpropagations. A branch of research has been devoted to understanding the resilience of neural network models to noise in training labels. For example, Luo et al. (2016) show that adding synthetic label noise to the convolutional neural network (CNN) can improve its ability to capture global features. Srivastava et al. (2014) quantify the error tolerance by injecting synthetic label noise with a custom Boltzmann machine hardware. Szegedy et al. (2013) find that neural networks are vulnerable to adversarial examples, and Goodfellow et al. (2014) design an efficient method to generate such noisy examples to exploit the vulnerability to adversarial perturbations. Carlini and Wagner (2017) design targeted attacks to training labels to test the robustness of neural networks. Instead of using synthetic noise, Jiang et al. (2020) inject real-world label noise and examine noise tolerance of neural networks with controlled noise levels. The aforementioned studies use real-world data, as is typically the case for many neural network studies, where noise is already present in the training labels before any noise injection. Users of real-world data have little control over the noise level of the original training labels and usually examine the effect of noisy data by injecting noise, but it is unclear whether a neural network model trained on noisy data actually learns the real, i.e., noiseless, feature-label relationship. Due to their lack of transparency and interpretability, the adoption of DNNs in financial and actuarial applications has been received by regulators with some skepticism.

The contributions of our study are two-fold:

- 1. We study what DNNs learn from noisy data by training them using simulated data based on well-designed simulation experiments. This is a novel way to study the effect of noisy data and error tolerance of neural network models as one can *reduce noise* in the data by increasing the number of replications in a simulation model. This new way of studying neural network models can provide more direct evidence on their transparency and interpretability.
- 2. We propose two generic nested simulation procedures that uses DNNs as metamodels to improve its efficiency while maintaining transparency. In essence, a pilot stage simulation is used to generate a large number of noisy data, which are then used to train a metamodel. Depending on the application, a trained metamodel can serve two purposes:
  - to identify a set of tail scenarios, and
  - to estimate risk measures directly.

The first procedure uses a metamodel to identify a set of potential tail scenarios on which computations are performed in stage 2, while the second procedure uses metamodel predictions to estimate risk measures directly. Our numerical results show that DNN metamodels can identify the tail scenarios accurately and so the proposed procedures can estimate tail risk measures with similar accuracy while, at the same time, using less simulation budget.

We are curious about fundamental questions like "What do DNNs learn from noisy data?" and "How well do neural networks learn from noisy data?". Data-driven answers to these questions prevail in the existing literature. In supervised learning, DNNs are believed to learn from the given data about the feature-label relationship to predict new labels for unseen features. Cross-validation using to assess a subset, i.e., the validation set, of the original data, is a common way to access the quality of learning. Generalization error on the test labels is another popular assessment metric. However, the test set is also a subset of the original data. In this study, we revisit these questions in a simulation context and propose an alternative approach to answer them. Instead of relying solely on real-data and splitting it into multiple subsets, we propose using stochastic simulation outputs as training labels for DNNs. By controlling the simulation design parameters, such as the number of independent replications, we can control the quality and also the quantity of the training labels fed into the neural networks. In such a controlled environment, we obtain more clear-cut answers to the above fundamental questions.

In nested simulation, a simulation model is used to generate a large number of outer scenarios, and each scenario is then used as an input to another simulation model, referred to as the inner simulation model. Borrowing terminologies from machine learning research, we can view a set of simulated outer scenarios and the estimated hedging errors for those scenarios as the features and noisy labels. The noise level of the labels can be controlled by the number of inner replications in the inner simulation model. One can train supervised learning models using these simulated features and labels. They are then used to replace the time-consuming inner simulations by the trained model. We refer to the trained supervised learning models as metamodels of the inner simulation, which is also known as the *surrogate models* in simulation literature. Metamodeling is a popular approach to reduce the computational burden of simulation-based applications by replacing the time-consuming simulation with a metamodel. The metamodel is trained using a set of simulated data, and it is used to predict the simulation output for new inputs. The study of metamodeling is an active research area in the simulation literature, and using DNNs as metamodels is a relatively new development. Fonseca et al. (2003) provide general guidelines for simulation metamodeling with neural networks, Lieu et al. (2022) use DNNs as metamodels of a simulation model for structural reliability analysis, and Salle and Yıldızoğlu (2014) show that neural network metamodels help achieve higher prediction accuracy that other metamodels in approximating agent-based simulation models. A popular metamodel in nested simulation procedures is stochastic kriging. Liu and Staum (2010) use stochastic kriging as a metamodel of Monte Carlo simulations to estimate the CVaR of a portfolio of derivative securities, and Gan and Lin (2015) use stochastic kriging for an efficient valuation of large portfolios of variable annuity (VA) contracts. Other studies, such as Broadie et al. (2015), Hong et al. (2017), and Zhang et al. (2022) use regression, kernel smoothing, and the likelihood ratio method, respectively. Our study has three key distinctions over the existing ones:

- 1. our metamodel has high-dimensional inputs. In machine learning terminology, the features are high-dimensional vectors. To estimate the hedging error of a typical VA contract, the number of features is in the order of hundreds, which is at least one order of magnitude larger than the number of features in the aforementioned studies,
- 2. for estimating tail risk measures, our metamodel is only used for tail scenario identification but is *not* used in the estimation of the tail risk measures. This is a feature designed particularly to convince regulators that the losses used in estimating the risk measure are based on a transparent inner simulation model rather than on some black-box metamodels, and
- 3. using simulation models as data generators, we can decrease the noise level and get arbitrarily close to the true labels by increasing the number of replications in the simulation model. This design allows a systematic study of the effect of noisy training labels on the performance of DNN models in predicting the noiseless labels.

In this chapter, DNN metamodels and DNN models are used almost interchangeably despite one distinction.

- The discussion of *DNN metamodels* focus more on the aspect of *estimating* the inner simulation.
- Tthe discussion of *DNN models* focus more on *studying the DNN* using simulation data.

The rest of this chapter is organized as follows: Section 3.2 presents the problem settings for tail risk measures and dynamic hedging of VAs. Section 3.3 proposes an efficient two-stage nested simulation procedure that uses DNNs as metamodels to help reduce simulation budget by only performing computations on identified tail scenarios. Section 3.4

proposes a single-stage nested simulation procedure that estimates risk measures directly with metamodel predictions. Section 3.5 demonstrates the efficiency of DNN metamodels and examines error tolerance of two LSTM metamodels with different numbers of trainable parameters. Practical suggestions are provided for the choice of suitable metamodels and simulation settings.

#### 3.2 Problem Formulation

In this section we present notations, problem settings, and a simulation model for risk estimation for hedging errors of VAs. A main goal of the section is to showcase the complexity of the simulation model, which we use as a data generator to train DNN metamodels (Section 3.3 and Section 3.4). For readers who are interested in the examination of a neural network metamodel, it is sufficient to understand that our simulation model generates data with 240 features and 1 real-value label and our metamodels are generally applicable to any simulation model that generates data with similar characteristics.

#### 3.2.1 Tail Risk Measures: VaR and CVaR

Measuring and monitoring risks, particularly tail risks, are important risk management tasks for financial institutions like banks and insurance companies. Two most popular tail risk measures are VaR and CVaR (Hardy and Saunders, 2022; Rockafellar and Uryasev, 2002).

Consider a loss random variable L whose losses and gains lie in the right and left tails, respectively, of its distribution. For a given confidence level  $\alpha \in [0,1]$ , the  $\alpha$ -VaR and  $\alpha$ -CVaR are defined in Equation (2.1) and Equation (2.2), respectively. Tail risk measures like VaR and CVaR are widely used for setting regulatory and economic capital, which is the amount of capital a financial institution holds to cover its risk. For example, European insurers set regulatory capital at 99.5%-VaR according to Solvency II EIOPA (2014). In Canada, the regulatory capital requirement for VAs is set based on CVaRs as prescribed in OSFI (2017).

Let  $L_1, L_2, \ldots, L_M$  be M independent and identically distributed (i.i.d.) simulated losses of L and let  $L_{(1)} \leq L_{(2)} \leq \ldots \leq L_{(M)}$  be the corresponding ordered losses. For a given confidence level  $\alpha$  (assume that  $\alpha M$  is an integer for simplicity),  $\alpha$ -VaR can be

estimated by the sample quantile  $\widehat{\text{VaR}}_{\alpha} = L_{(\alpha M)}$ . Also,  $\alpha$ -CVaR can be estimated by

$$\widehat{\text{CVaR}}_{\alpha} = \frac{1}{(1-\alpha)M} \sum_{i=\alpha M+1}^{M} L_{(i)} = \frac{1}{(1-\alpha)M} \sum_{i \in \mathcal{T}_{(1-\alpha)M}} L_{i}.$$
(3.1)

Equation (3.1) is the same as (2.3) except we define a true tail scenario set of size k as  $\mathcal{T}_k = \{i : L_i > L_{(M-k)}\}$ . In this study, the loss random variable of interest is the hedging error for VA.

#### 3.2.2 Simulation Model for Variable Annuity Payouts

VA contracts offer different types of guarantees. Generally speaking, a portion of the VA premium is invested in a sub-account whose return is linked to some stock indices.

Two relevant types of guarantees in our studies are:

- Guaranteed Minimum Maturity Benefit (GMMB): A GMMB contract pays a maturity benefit equal to the greater of the sub-account value and a fixed guarantee value. The guarantee value is often set as a percentage, e.g., 75% or 100%, of the initial premium.
- Guaranteed Minimum Withdrawal Benefit (GMWB): A GMWB contract guarantees the minimum amount of periodic withdrawal the policyholder can take from the sub-account until maturity, even if the sub-account value reduces to zero. The minimum withdrawal benefit is typically a fixed percentage of the guarantee value. The guarantee value will decrease if the withdrawal exceeds the guaranteed minimum. The GMWB is typically offered with an accumulation period, during which no withdrawals are made but a GMDB is usually offered. Additional features offered with the GMWB include roll-up, ratchet, and reset (The Geneva Association, 2013).

For a comprehensive review of other types of VA contracts such as Guaranteed Minimum Death Benefit (GMDB), Guaranteed Minimum Accumulation Benefit (GMAB) and Guaranteed Lifetime Withdrawal Benefit (GLWB), we refer readers to Hardy (2003). Next we present a summary of dynamic hedging for VA contracts. We refer readers to Dang (2021) for detailed modeling of insurer liabilities in different VA contracts and Greek estimation.

Consider a generic VA contract with maturity T > 0 periods, e.g., T = 240 months. Denote the policyholder's (random) time of death by  $\tau > 0$ . The contract expires at  $T'=\min\{T,\tau\}$ , i.e., the earlier of the contract maturity and the death of the policyholder. Let  $S_t$ ,  $F_t$ , and  $G_t$  be the indexed stock price, the subaccount value and the guarantee value, respectively, at time  $t=1,2,\ldots,T$ . Evolution of the subaccount value and the guarantee value of a VA contract affect the contract payout. Note that the policyholder's (random) time of death also affects the timing of the benefit payout for certain types of VA such as GMDB, but this is not considered in our study for simplicity. For clarity, we use  $F_t$  and  $F_{t_+}$  to denote the sub-account value just before and just after the withdrawal at time t, if any. Let  $\eta_g$  be the gross rate of management fee that is deducted from the fund value at each period and let  $\eta_n < \eta_g$  be the net rate of management fee income to the insurer. The difference between the gross management fee and the net management fee income represents the incurred investment expenses.

At the inception of the contract, i.e., t=0, we assume that the whole premium is invested in the stock index and the guarantee base is set to the sub-account value:

$$S_0 = F_0 = G_0$$
.

At each time t = 1, ..., T, the following events take place in the following order:

1. The sub-account value changes according to the growth of the underlying stock and the (gross) management fee is deducted. That is,

$$F_t = F_{(t-1)_+} \cdot \frac{S_t}{S_{t-1}} \cdot (1 - \eta_g),$$

where  $(x)^+ = \max\{x, 0\}$  and  $F_{(t-1)_+}$  will be defined later. The insurer's income at time t is the net management fee, i.e.,  $F_t\eta_n$ .

2. The guarantee value ratchets up (ratcheting is a common feature in GMWB) if the sub-account value exceeds the previous guarantee value, i.e.,

$$G_t = \max\{G_{t-1}, F_t\}.$$

3. The withdrawal is made (for GMWB) and is deducted from the sub-account value, i.e.,

$$F_{t_{+}} = (F_t - I_t)^+,$$

where  $I_t = \gamma G_t$ . A GMMB can be modeled with  $\gamma = 0$ .

We see from the above modeling steps that the status of a generic VA contract is summarized by a triplet  $(S_t, F_t, G_t)$  whose evolution is driven by the stochasticity of  $S_t$ . In practice, the simulation model may also incorporate additional complications like mortality, lapse, and excess withdrawal, etc.

At any time  $t=1,\ldots,T$ , the insurer's liability in a VA contract is the present value of all payments, net of the fee income. For example, suppose that the per-period risk-free rate is r, then the insurer's time-t liability for a GMMB contract is  $V_t = e^{-r(T-t)} \cdot (G_T - F_T)^+ - \sum_{s=t+1}^T e^{-r(T-s)} F_s \eta_n$ . Also, the insurer's time-t liability for a GMWB contract is  $V_t = \sum_{s=t+1}^T e^{-r(T-s)} [(I_s - F_s)^+ - \eta_n F_s]$ .

For example, consider the time-t liability  $V_t$  of a GMWB: Suppose that given the stock sample path, e.g., an outer path  $S_1, \ldots, S_t$ , one can simulate future stock prices  $\widetilde{S}_{t+1}, \ldots, \widetilde{S}_T$ , e.g., inner sample paths, based on some asset model such as a Black-Scholes model. The tilde symbol ( $\sim$ ) over a quantity denotes its association with the inner simulation. Given the time t state  $(S_t, F_t, G_t)$ , following Cathcart et al. (2015) the sensitivity of  $V_t$  with respect to  $S_t$  can be estimated by a pathwise estimator (Glasserman, 2004):

$$\Delta_{t}(\widetilde{S}_{t+1}, \dots, \widetilde{S}_{T}|S_{t}) = \frac{\partial V_{t}}{\partial S_{t}} = \sum_{s=t+1}^{T} e^{-r(T-s)} \left[ \mathbf{1}\{\widetilde{I}_{s} > \widetilde{F}_{s}\} \cdot \left( \frac{\partial \widetilde{I}_{s}}{\partial S_{t}} - \frac{\partial \widetilde{F}_{s}}{\partial S_{t}} \right) - \eta_{n} \frac{\partial \widetilde{F}_{s}}{\partial S_{t}} \right],$$

$$t = 0, \dots, T-1,$$
(3.2)

where  $\mathbf{1}\{\cdot\}$  is an indicator function and

$$\frac{\partial \widetilde{F}_{s}}{\partial S_{t}} = \mathbf{1}\{\widetilde{I}_{s-1} < \widetilde{F}_{s-1}\} \cdot \left(\frac{\partial \widetilde{F}_{s-1}}{\partial S_{t}} - \frac{\partial \widetilde{I}_{s-1}}{\partial S_{t}}\right) \cdot \frac{\widetilde{S}_{s}}{\widetilde{S}_{s-1}} \cdot (1 - \eta_{g}),$$

$$\frac{\partial \widetilde{G}_{s}}{\partial S_{t}} = \mathbf{1}\{\widetilde{G}_{s-1} < \widetilde{F}_{s}\} \cdot \frac{\partial \widetilde{F}_{s}}{\partial S_{t}} + \mathbf{1}\{\widetilde{G}_{s-1} \ge \widetilde{F}_{s}\} \cdot \frac{\partial \widetilde{G}_{s-1}}{\partial S_{t}},$$

$$\frac{\partial \widetilde{I}_{s}}{\partial S_{t}} = \gamma \frac{\partial \widetilde{G}_{s}}{\partial S_{t}}.$$

The recursion is initialized with  $(\widetilde{S}_t, \widetilde{F}_t, \widetilde{G}_t) = (S_t, F_t, G_t), \frac{\partial \widetilde{F}_s}{\partial S_t} = \frac{\widetilde{F}_t}{S_t}$ , and  $\frac{\partial \widetilde{G}_s}{\partial S_t} = \frac{\partial \widetilde{I}_s}{\partial S_t} = 0$ .

#### 3.2.3 Dynamic Hedging for Variable Annuities

Below we provide a scheme used to perform a multi-period nested simulation in estimating (profit and loss) P&L for one outer scenario.

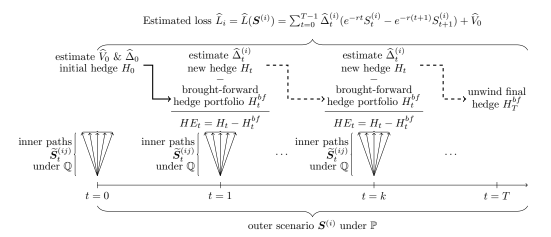


Figure 3.1: Illustration of multi-period nested simulation that estimates the P&L for one outer scenario.

Insurers commonly use dynamic hedging to mitigate a market risk exposure in VA contract's embedded options. In a dynamic hedging program, a hedge portfolio is set up and periodically rebalanced for a portfolio of VA contracts using stocks, bonds, futures, and other derivatives. For simplicity, in this study we consider delta hedging for a generic VA liability using one stock and one bond. The metamodeling procedures in Section 3.3 and Section 3.4 can be trivially adapted to more general hedging strategies.

Consider a VA contract whose delta hedge portfolio at any time t, t = 0, 1, ..., T - 1, consists of  $\Delta_t$  units in the underlying stock and  $B_t$  amount of a risk-free zero-coupon bond maturing at time T. The value of the hedge portfolio at time (t - 1) is:

$$H_{t-1} = \Delta_{t-1} S_{t-1} + B_{t-1}$$

where  $S_t$  is the underlying stock price and any time t > 0. This hedge portfolio is brought forward to the next rebalancing time t, when its value becomes:

$$H_t^{bf} = \Delta_{t-1} S_t + B_{t-1} e^r.$$

Therefore, the time t hedging error, i.e., the cash flow incurred by the insurer due to rebalancing at time t, is

$$HE_t = H_t - H_t^{bf}, \quad t = 1, \dots, T - 1.$$
 (3.3)

The P&L of the VA contract includes the cost of the initial hedge  $(H_0)$ , the hedging errors (3.3), the unwinding of the hedge at maturity  $(H_T^{bf})$ , and the unhedged liability

 $(V_0)$ . Mathematically, the present value of these cash flows is given by

$$L = H_0 + \sum_{t=1}^{T-1} e^{-rt} H E_t - e^{-rT} H_T^{bf} + V_0 = \sum_{t=0}^{T-1} \Delta_t (e^{-rt} S_t - e^{-r(t+1)} S_{t+1}) + V_0,$$
 (3.4)

where the second equality holds by a telescopic sum simplification of  $e^{-rt}B_t$ ,  $t=0,\ldots,T-1$ .

In (3.4),  $\Delta_t$  and  $V_0$  are determined by using a risk-neutral measure  $\mathbb{Q}$  while the distribution of L is under a real-world measure  $\mathbb{P}$ . If  $\Delta_t$  and  $V_0$  cannot be calculated analytically, a nested simulation is required to estimate the tail risk measure of L. Recall from Section 3.2.2 that the stock sample path, regardless of the inner or outer simulation or a combination of both, determines the evolution of the triplet  $(S_t, F_t, G_t)$ . Specifically, the outer scenarios  $\mathbf{S}^{(i)} = (S_1^{(i)}, \dots, S_T^{(i)}), i = 1, \dots, M$  are generated under  $\mathbb{P}$ . At each time  $t = 1, \dots, T - 1$  of a given outer scenario  $\mathbf{S}^{(i)}$ , inner sample paths  $\widetilde{\mathbf{S}}_t^{(j)} = (\widetilde{S}_{t+1}^{(j)}, \dots, \widetilde{S}_T^{(j)}), j = 1, \dots, N$  are generated under  $\mathbb{Q}$  to estimate  $\Delta_t^{(i)}, i = 1, \dots, M$ . Also,  $V_0^{(i)}$ ,  $i=1,\ldots,M$  are estimated under  $\mathbb{Q}$  via inner simulations at time 0. Recall from Section 3.2.2 that the stock's sample path, regardless of inner or outer simulation or a combination of both, determines the evolution of the triplet  $(S_t, F_t, G_t)$ . For GMWB, a standard nested simulation procedure to estimate the  $\alpha$ -CVaR of L is described in Algorithm 1.

#### Algorithm 1 Standard Nested Simulation Procedure for Estimating CVaR for GMWB Hedging Losses

- 1: For  $i=1,\dots,M$ , simulate outer scenarios  $\boldsymbol{S}^{(i)}=(S_1^{(i)},\dots,S_T^{(i)})$  under the real-world
- 2: For t=0, simulate time-0 inner paths  $\widetilde{\boldsymbol{S}}_0^{(j)}=(\widetilde{S}_1^{(j)},\widetilde{S}_2^{(j)},\ldots,\widetilde{S}_T^{(j)}),\ j=1,\ldots,N$  under  $\mathbb{Q}$ , then estimate  $V_0$  by  $\widehat{V}_0=\sum_{s=1}^T e^{-r(T-s)}[(I_s-F_s)^+-\eta_nF_s]$  and  $\widehat{\Delta}_0=\sum_{s=1}^T e^{-r(T-s)}[(I_s-F_s)^+-\eta_nF_s]$  $\Delta_0(\widetilde{S}_1^{(j)},\ldots,\widetilde{S}_T^{(j)}|S_0).$
- 3: Given each scenario  $S^{(i)}$ :

We refer to the collection of experiments needed conditional on one scenario  $S^{(i)}$  to

estimate  $L_i$ , that is, all upward arrows in Figure 3.1, as one inner simulation. We make four observations:

- Each inner simulation is time-consuming, as it includes T simulation experiments: one at each time  $t = 0, \ldots, T 1$ ,
- After running inner simulations for M scenarios, we obtain simulated dataas feature-label pairs,  $(\mathbf{S}^{(i)}, \widehat{L}_i)$ , i = 1, ..., M; the feature vector  $\mathbf{S}$  is T dimensional.
- $\widehat{L}_i^{MC}$  is a standard Monte Carlo estimator of the true loss for scenario  $S^{(i)}$ . It is an unbiased estimator and its variance is inversely proportional to the number of inner replications N. As N approaches infinity,  $\widehat{L}_i^{MC}$  converges to the true loss  $L_i$ , and
- Most importantly, when estimating tail risk measures such as  $\alpha$ -CVaR, only a small number of estimated losses, that is, those associated with the set of tail scenarios  $\widehat{\mathcal{T}}_k$  are used in the estimator.

#### 3.3 Two-Stage Nested Simulation with Metamodels

Based on the three observations above and inspired by Dang et al. (2020), we propose a two-stage nested simulation procedure which uses a DNN metamodel to identify potential tail scenarios. We present our proposed procedure as a competitor to the standard procedure with M outer scenarios and N inner replications for each outer scenario, as described in Section 3.2.3. We propose a two-stage procedure with a supervised learning metamodel that aims to produce a CVaR estimate that's as accurate as that of the standard procedure, but requires fewer computations than the latter.

Similar to Dang et al. (2020), the proposed two-stage procedure in Algorithm 2 uses the metamodel predictions to identify the predicted tail scenario set in stage 1. However, different from their fixed-budget simulation design, we attempt to achieve a target accuracy. Specifically, in stage 2 we propose using a standard procedure with the same number of inner replications, N, as a benchmark procedure. There are two different types of experiment designs for nested simulation procedures: fixed-budget design and fixed-accuracy design. In a fixed-budget design, the simulation budget is fixed and the goal is to achieve the highest accuracy possible within the budget. Let  $\Gamma = MN$  be the simulation budget for the standard procedure, where each scenario receives  $\frac{\Gamma}{M}$  inner replications. In the proposed two-stage procedure, suppose stage 2 uses 1% of the simulation budget,  $\alpha = 95\%$ , and  $m = (1 - \alpha)M$ , then 99% of the simulation budget is concentrated on 5%M predicted

**Algorithm 2** Two-Stage Metamodeling Nested Simulation Procedure for Estimating CVaR.

#### 1: Train a neural network metamodel using simulation data:

- a. Use a fraction of the total simulation budget to run Steps 1, 2, and 3 in the standard procedure, i.e., Algorithm 1, with the same number of outer scenarios, M, but a much smaller number of inner replications,  $N' \ll N$ , in each scenario. Obtain M simulated samples (feature-label pairs),  $(S^{(i)}, \hat{L}_i)$ ,  $i = 1, \ldots, M$ . Note that N' may be much smaller than N, so  $\hat{L}_i$  are expected to have larger variance.
- b. Use the simulated data,  $(\mathbf{S}^{(i)}, \widehat{L}_i)$ , i = 1, ..., M to train a supervised learning model. It is standard practice in supervised learning research to transform the stock prices  $\mathbf{S}^{(i)}$  to returns  $\mathbf{X}^{(i)}$ ,

$$\mathbf{X}^{(i)} = \left(\frac{S_1^{(i)} - S_0}{S_0}, \frac{S_2^{(i)} - S_1^{(i)}}{S_1^{(i)}}, \dots, \frac{S_T^{(i)} - S_{T-1}^{(i)}}{S_{T-1}^{(i)}}\right)$$
(3.5)

and use  $\mathbf{X}^{(i)}$  as features instead of the stock prices  $\mathbf{S}^{(i)}$ . Refer to the trained model as a metamodel and denote it by  $\widehat{L}^{PD}(\mathbf{X})$ . Denote the predicted losses for the outer scenarios by  $\widehat{L}_i^{PD} = \widehat{L}^{PD}(\mathbf{X}^{(i)})$ , i = 1, ..., M.

the outer scenarios by  $\widehat{L}_{i}^{PD} = \widehat{L}^{PD}(\mathbf{X}^{(i)}), i = 1, \dots, M$ . c. Sort the predicted losses  $\widehat{L}_{(1)}^{PD} \leq \widehat{L}_{(2)}^{PD} \leq \dots \leq \widehat{L}_{(M)}^{PD}$  to identify a predicted tail scenario set,  $\widehat{\mathcal{T}}_{m}^{PD}$ , associated with the largest predicted losses. The number of predicted tail scenarios, m, is a user's choice.

#### 2: Concentrate simulation on predicted tail scenarios:

- a. Run Steps 2 and 3 of Algorithm 1 with the same number of inner replications, N, but only on the predicted tail scenarios, i.e., scenarios in  $\widehat{\mathcal{T}}_m^{PD}$ . Denote the standard procedure's estimated losses and sorted losses by  $\widehat{L}_i^{ML}$  and  $\widehat{L}_{(i)}^{ML}$ , respectively,  $i=1,\ldots,m$ .
- b. Estimate the  $\alpha$ -CVaR of L by

$$\widehat{\text{CVaR}}_{\alpha}^{ML} = \frac{1}{(1-\alpha)M} \sum_{i=\alpha M+1}^{M} \widehat{L}_{(i)}^{ML} = \frac{1}{(1-\alpha)M} \sum_{i \in \widehat{\mathcal{T}}_{(1-\alpha)M}^{ML}} \widehat{L}_{i}^{ML},$$

where  $\widehat{\mathcal{T}}_k^{ML}$  denotes a predicted tail scenario set associated with the largest k estimated losses.

tail scenarios in stage 2. In other words, each predicted tail scenario receives  $\frac{99\%\Gamma}{5\%M}$  inner replications, almost 20 times more than that in the standard procedure. This budget concentration is expected to improve the estimation accuracy of the two-stage procedure compared to a standard procedure with the same budget. If the metamodel is accurate in predicting true tail scenarios, the two-stage procedure is expected to achieve higher accuracy than the standard procedure with the same budget. However, we believe that the goal of designing an efficient simulation procedure is to solve practical problems faster, so a target-accuracy design is more suitable, which refers to obtaining a similar level of accuracy as the standard procedure but with much less simulation budget. One other reason for this fixed-accuracy design is to investigate whether DNN metamodels trained with much noisier labels can identify true tail scenarios with similar accuracy as the standard procedure. The size of the predicted tail scenario set in stage 1, m, is an important experiment design parameter that affects the correct identification of true tail scenarios and ultimately affects the estimation accuracy for CVaR. Clearly, there is a lower bound  $m \geq (1-\alpha)M$ because the  $\alpha$ -CVaR is estimated by the average of  $(1-\alpha)M$  largest losses at the end of stage 2. For ease of reference, we call the additional percentage of predicted tail scenarios above this lower bound, i.e.,  $\epsilon = \frac{m - (1 - \alpha)M}{M}$ , as a safety margin. On one hand, large  $\epsilon$  is not desirable because it increases computations in stage 2. On the other hand,  $\epsilon$  should be set reasonably large so more true tail scenarios are included in the the predicted tail scenario set  $\widehat{\mathcal{T}}_m^{PD}$  and are ultimately included in  $\widehat{\mathcal{T}}_{(1-\alpha)M}$  at the end of stage 2. The selection of m is highly dependent on the choice of the metamodel. Due to the simulation errors and approximation error in the metamodel in stage 1, we do not expect perfect match between the true tail scenario set  $\mathcal{T}_k$  and the predicted tail scenario set  $\widehat{\mathcal{T}}_k^{PD}$  for any size k. This means that we should not set m at its lower bound: Some safety margin  $\epsilon M$  should be added to the predicted tail scenario set, i.e.,  $m = (1 - \alpha)M + \epsilon M$ , to increase the likelihood that  $\mathcal{T}_k \subseteq \widehat{\mathcal{T}}_m^{PD}$  and that the true tail scenarios are included in estimating  $\alpha$ -CVaR at the end of stage 2. The choice of safety margin  $\epsilon$  is not trivial, and it should be set based on the metamodel's accuracy in identifying true tail scenarios. In the numerical experiments, we examine the relationship between the safety margin and the correct identification of true tail scenarios for different metamodels.

## 3.4 Single-Stage Nested Simulation with Neural Network Metamodels

In our exploratory experiment with the two-stage procedure, we observe that a suitable metamodel trained with noisy labels is accurate enough to identify true tail scenarios with a relatively small safety margin. This observation motivates us to propose a single-stage procedure that uses the identical neural network metamodel, not for the purpose of differentiating between tail and non-tail scenarios, but rater to estimate the CVaR directly using the predicted losses.

**Algorithm 3** Single-Stage Metamodeling Nested Simulation Procedure for Estimating CVaR

- 1: Run Step 1 of Algorithm 2, and denote the predicted losses by  $\widehat{L}_i^{PD} = \widehat{L}^{PD}(\mathbf{S}^{(i)}),$  i = 1, ..., M.
- 2: Sort the predicted losses  $\widehat{L}_{(1)}^{PD} \leq \widehat{L}_{(2)}^{PD} \leq \cdots \leq \widehat{L}_{(M)}^{PD}$  to identify the largest predicted losses.
- 3: Directly estimate the risk measure (e.g.,  $\alpha$ -CVaR) of L using the metamodel predictions: Calculate  $\widehat{\text{CVaR}}_{\alpha}^{PD} = \frac{1}{(1-\alpha)M} \sum_{i=\alpha M+1}^{M} \widehat{L}_{(i)}^{PD}$ .

The single-stage procedure has three major advantages over the two-stage procedure:

- 1. The single-stage procedure is expected to be more efficient than the two-stage procedure because it does not require the running the standard procedure in stage 2.
- 2. The safety margin  $\epsilon$  is not needed in the single-stage procedure. The safety margin is difficult to determine because it depends on the metamodel's accuracy in identifying true tail scenarios. The single-stage procedure does not need a safety margin because it directly estimates the risk measure using the predicted losses.
- 3. Most importantly, the single-stage procedure is not limited to just estimating tail risk measures and can be extended to provide a broader assessment of risk. It can be naturally adapted to estimate risk measures that require the knowing of the entire loss distribution, such as the standard deviation or the squared tracking error. Even for CVaR, the two-stage procedure become less attactive for an increased level.

#### 3.5 Numerical Results

We conduct a series of simulation experiments to

- 1. demonstrate the efficiency of the proposed metamodeling procedures, and
- 2. examine the error tolerance to noisy training data in deep learning models.

The problem settings in our experiments are identical to those in Dang et al. (2020): we consider estimating the 95% CVaR of the hedging loss of a GMWB contract, which is one of the most complex VA contracts in the market. The VA contracts have a 20-year maturity and are delta-hedged with monthly rebalancing, i.e., T=240 rebalancing periods. The gross and net management fees are  $\eta_g=0.2\%$  and  $\eta_n=0.1\%$ , respectively. The withdrawal rate for GMWB is 0.375% per month. To make our examples more realistic, the VAs are also subject to a dynamic lapse model with the following key features (NAIC, 2021):

- 1. When a policyholder lapses, both the fund value (F) and guarantee value (G) are reduced proportionally.
- 2. The monthly lapse rate from time t to t+1, denoted as  $\frac{1}{12}q_{x+\frac{t}{12}}^l$ , is given by:

$$\frac{1}{12}q_{x+\frac{t}{12}}^{l} = \min\left\{1, \max\left\{0.5, 1 - 1.25 \times \left(\frac{G_t}{F_t} - 1.1\right)\right\} \times \frac{1}{12}q_{x+\frac{t}{12}}^{base}\right\}$$
(3.6)

where  $\frac{1}{12}q_{x+\frac{t}{12}}^{base}$  is the base monthly mortality rate for a policyholder aged  $x + \frac{t}{12}$  at month t. where the base monthly lapse rate is defined as:

$$\frac{1}{12}q_{x+\frac{t}{12}}^{base} = \begin{cases} 0.00417 & \text{if } t < 84\\ 0.00833 & \text{if } t \ge 84 \end{cases}$$
 (3.7)

The risk-free rate is 0.2% per month and the underlying asset  $S_t$  is modeled by a regime-switching geometric Brownian motion with parameters specified in Table 3.1.

Parameter	Monthly Value
Risk-free rate	0.002
Mean return for regime 1	0.0085
Mean return for regime 2	-0.0200
Standard deviation for regime 1	0.035
Standard deviation for regime 2	0.080
Transition probability from regime 1	0.04
Transition probability from regime 2	0.20

Table 3.1: Real-world parameters for the regime-switching model (monthly rates)

To compare the numerical performances of different simulation procedures, we create a benchmark dataset with a large-scale nested simulation: We first simulate M = 100,000

outer scenarios, i.e., 240-periods stock paths  $S^{(1)}, \ldots, S^{(M)}$  under  $\mathbb{P}$  and used these outer scenarios in all further experiments. Note that the 5% tail scenario set includes 5000 scenarios. As the hedging losses for these scenarios cannot be calculated analytically, we run inner simulations with a large number of replications, N = 100,000, conditional on each of the M scenarios. We denote these losses by  $L_1, \ldots, L_M$  and will refer to them as true losses. We also use these true losses to estimate  $\widehat{\text{CVaR}}_{95\%}$  and denote the corresponding true tail scenario set by  $\mathcal{T}_{5000}$ . Lastly, we refer to the set of feature-label pairs  $\{(S^{(i)}, L_i): i=1,\ldots,M\}$  as a true dataset. Note that the feature vector S is a 240-dimension stock path.

In our experiments, the standard nested simulation procedure is mainly used in three ways.

- 1. The true losses and the true 95%-CVaR are estimated using a standard nested simulation procedure that runs  $N=100{,}000$  inner replications for each of the  $M=100{,}000$  outer scenarios.
- 2. A benchmark estimator of the 95%-CVaR is generated using N=1000 for each of the  $M=100{,}000$  outer scenarios.
- 3. The training data for our metamodels is generated using N = 100 for M = 100,000. The value of M and N are subject to change in our other numerical experiments.

Five metamodels are considered in this experiment: multiple linear regression (MLR), quadratic polynomial regression (QPR) without interaction terms, FNN, RNN, and LSTM network. MLR and QPR are two common metamodels in the simulation literature, and their implemention is straightfoward in our simulation setting. FNN is a generic neural network while RNN and LSTM are specialized models to accommodate the sequential structure of our time-series features. A tanh activation function is used for RNN and LSTM layers, and a ReLU activation function is used for the fully-connected layers. All neural network metamodels are trained by the Adam optimizer (Kingma and Ba. 2014) with an initial learning rate of 0.001 and an exponential learning rate decay schedule. All DNN metamodels are trained with a dropout rate of 10%. The architectures and training settings are typical choices in the deep learning literature. The training labels are normalized to have zero mean and unit standard deviation. The numbers of trainable parameters are recorded in the second column of Table 3.2. We observe that the three DNN metamodels have orders of magnitudes more trainable parameters than the two traditional regression metamodels. In machine learning terminologies, the three DNN metamodels have much higher model capacities. From a neural network of view, MLR and QPR are two neural networks with no hidden layer. For a traditional regression metamodel, the numbers of neurons in its input layer equals to the cardinality of its regression basis. Treating a regression as a shallow neural network, i.e., a neural network with only an input layer and an output layer, allows for direct comparisons with DNNs. The capacity of a regression metamodel is the product of the number of neurons in its input layer and its output layer. Since our target variable is a scalar, the output layer has only one neuron. The capacity of a regression metamodel therefore equals to the size of its regression basis. On the other hand, the three DNN metamodels have several hidden layers and are much more complex than the two regression metamodels. Our FNN has two hidden layers before the output layer. Its last hidden layer serves as an automatic feature extraction layer that transforms the 240-dimension feature vector into a 32-dimension vector that can directly relate to the target variable. Our RNN and LSTM networks have three hidden layers. They contain similar numbers of neurons as the FNN but have deeper network structures. For the RNN, two of its hidden layers are recurrent layers that capture the temporal dependency in the feature, and the last hidden layer is a standard feedforward layer that transforms the previous hidden states into a 32-dimension vector for target prediction. Our LSTM has the same architecture as the RNN but with LSTM cells in place of the RNN cells.

Metamodel	Capacity	Training error	Test error	True error
MLR	241	$0.706(\pm 8.34 \times 10^{-4})$	$0.713(\pm 2.67 \times 10^{-2})$	$0.706(\pm 3.44 \times 10^{-4})$
QPR	481	$0.543(\pm 8.27 \times 10^{-4})$	$0.554(\pm 2.32 \times 10^{-2})$	$0.544(\pm 4.12 \times 10^{-4})$
FNN	35,009	$0.129(\pm 5.95 \times 10^{-3})$	$0.240(\pm 9.82 \times 10^{-3})$	$0.132(\pm 5.82 \times 10^{-3})$
RNN	32,021	$0.132(\pm 7.53 \times 10^{-3})$	$0.137(\pm 7.62 \times 10^{-3})$	$0.119(\pm 7.51 \times 10^{-3})$
LSTM	35,729	$0.075(\pm 4.48 \times 10^{-3})$	$0.079(\pm 5.35 \times 10^{-3})$	$0.063(\pm 4.43 \times 10^{-3})$
RNN*1	32,021	$0.109(\pm 5.20 \times 10^{-3})$	$0.128(\pm 5.22 \times 10^{-3})$	$0.109(\pm 5.20 \times 10^{-3})$

Table 3.2: Architectures and MSEs of metamodels for GMWB inner simulation model.

For each metamodel, 100 independent macro replications are run. Each macro replication involves simulating a noisy dataset, randomly separating it into training and test data, fitting a metamodel, and making predictions. The last three columns in Table 3.2 display the average squared errors between the metamodel predictions and the loss labels for different datasets. Training error, test error, and true error are the average squared errors between the metamodel predictions and the training labels, test labels, and true labels, respectively. The half lengths of their 95% confidence bands are also reported in parentheses. We first observe that the two regression metamodels have high training errors.

<sup>&</sup>lt;sup>1</sup>This row summarizes the results of the well-trained RNN, where the ones suffers from the vanishing gradient issue are removed from the averages.

This is a sign of under-fitting. Model capacities of the regression metamodels are too low to learn the training dataset. A successful learning on the training dataset is a necessary condition for a metamodel to generalize to the true simulation model. All neural network metamodels have high model capcities comparing with the number of data points in the training set, i.e., 90,000. Empirically, this is not a huge concern for the DNNs. In neural network literature, the number of trainable parameters is often much larger than the number of training samples. AlexNet (Krizhevsky et al., 2012), a well-known convolutional neural network, has 60 million trained parameters and is trained on the ImageNet dataset with only 1.2 million sample images. Other examples include generative pre-trained transformer models like BERT (Devlin, 2018) and GPT-3 (Brown, 2020), which have billions of trainable parameters and are trained on large-scale text corpora. The reason for not over-fitting when the number of trainable parameters is much larger than the number of training samples is still an open question in deep learning research. One of the explanations in the literature is that the optimization surface of DNNs is highly non-convex and has many local minima. Given a good initialization, the optimization algorithm is likely to find a solution that generalizes well to unseen data. Another explanation is that most existing DNN models have specific architectures that make them less prone to over-fitting. For example, a convolutional neural network (CNN) is specifically designed to capture the spatial structure in image data. Unlike a FNN, not all neurons are connected to all neurons in the previous layer. This architecture serves as prior knowledge and a form of regularization that make CNNs less prone to over-fitting.

In our experiments, there are significant differences among the three DNN models due to having different neural network architectures. Their model capacities are intentionally set to be similar to each other for a fair comparison. The FNN is a generic neural network, its test error is much greater than the training error, which is a sign of over-fitting and poor generalization. In contrast, RNN and LSTM networks have recurrent strucutures and are designed to capture temporal relationship in the high-dimensional feature. As a result, they have lower training errors, i.e., they fit the data better, and have lower test errors, i.e., they generalize better. Notably, the true errors for the DNN metamodels are lower than the training errors. This is a sign of the metamodels' ability to learn the true feature-label relationship with noisy training labels. Section 3.5.2 investigates this phenomenon in greater details.

The intuition behind the RNN and LSTM metamodels is briefly mentioned in 1.3.5. To overcome the overfitting issue of the FNN, RNN learns a single function that operates across all time steps, and the time t itself is not a variable in the RNN. The RNN cells in RNN and LSTM networks share parameters across time and serve as a form of regularization that prevents overfitting. However, the price that RNN pays for a reduced number of

parameters is the difficulty in its optimization. Comparing the two metamodels with recurrent structures, a LSTM network overcomes two key drawbacks of a crude RNN.

- 1. It avoids the vanishing gradient problem during training, which makes the LSTM network easier to train than the RNN. Figure 3.2 shows the quantile-quantile (QQ) plots between the **training** labels and the RNN predictions for two macro replications, where training of the latter is hindered by the vanishing gradient problem. As a result, the half length of the 95% confidence band of the RNN's training error is much larger than all other metamodels. The last row of Table 3.2 shows the average squared errors of the good RNN metamodels that are successfully trained. Removing the RNN metamodels that are ill-trained, the average squared errors of the RNN metamodels are lower than those of the FNN. This suggests that the RNN is more sensitive to noise in stochastic gradient descent than the FNN. They can capture the temporal dependency but are more difficult to optimize.
- 2. A LSTM introduces three gates that regulate the flow of information. These gates decide what information should be kept or discarded. For detailed discussion on the LSTM network, please refer to Section 1.3.5. This gating mechanism makes a LSTM network more capable of learning long-term dependencies. Since the feature in our dynamic hedging problem is a 240-dimensional time series, a LSTM network is more suitable than a crude RNN.

In summary, the network depth ensures successful learning of the training dataset, and the network architecture serves as

- 1. regularization that prevents over-fitting, and
- 2. a prior knowledge that guides the optimization algorithm to find a solution that generalizes well to unseen data.

They are both crucial for the metamodels' performance. It is important to choose a *suitable* architecture with a pre-set capacity that is large enough to learn the training dataset and generalize to the true feature-label relationship.

Recall that all three datasets are generated from the same simulation model, but the true labels contain much less simulation noise than the training and test labels. Part of the training error is from the simulation noise, and this noise is much less reflected in the true error. We observe that the two regression metamodels is under-fitting with high training errors. They also have low test and true errors, which indicates poor generalization. In

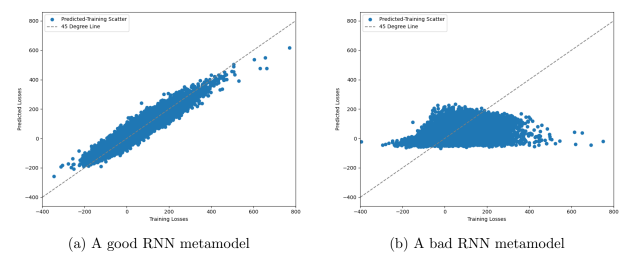


Figure 3.2: QQ-plots between true labels (x-axis) and predicted losses (y-axis) for the RNN metamodel.

contrast, the DNN metamodels have lower errors. More specifically, they generalize better to the true labels than to the test labels. The test labels contain more simulation noise than the true labels. This is a sign of the regression metamodels' inability to learn the true feature-label relationship rather than merely over-fitting to the simulation noise in the training labels. In practical applications, the true labels are not available. However, analyzing the quality of fit on the true labels in a simulation environment offers a unique opportunity to understand the metamodels' true ability of learning. Figure 3.3 and Figure 3.4 shows the QQ plots between the **true** loss labels and the predicted losses for the regression metamodels and the neural network metamodels, respectively.

Comparing with the MSE table in Table 3.2 that summarizes the overall fit, QQ plots offer a closer look at the metamodels' performance on different parts of the loss distribution. Figure 3.3 show the regression metamodel predictions and the loss labels in the true dataset. Between the two regression metamodels, the QPR metamodel has a slightly better fit for larger losses. Nevertheless, the predictions of both regression metamodels are far from the true labels, and their fit on the tail is particularly undesirable. The poor fit on the tail hinders the regression metamodels' ability to identify true tail scenarios and ultimately leads to poor CVaR estimates. Adding the quadratic terms, our QPR metamodel is considered as a natural extension of the MLR. Attempts to further improve the regression metamodels by adding interaction terms or higher-order terms do not improve the fit. Feature engineering (i.e., variable selection) for regression metamodels is not a trivial

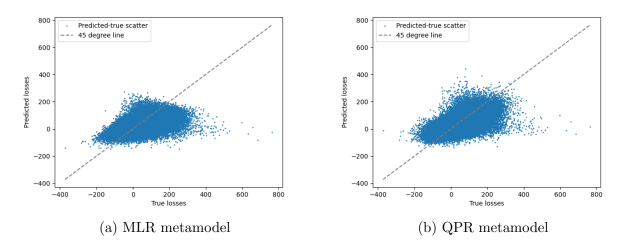


Figure 3.3: QQ-plots between true losses (x-axis) and predicted losses (y-axis) for regression metamodels.

task and is highly dependent on the simulation procedure. Having 240 time-series features further complicates this issue. As a result, our attempts to improve the traditional regression metamodels are not successful. The regression metamodels are not flexible enough to capture the complex feature-label relationship in the dynamic hedging simulation model.

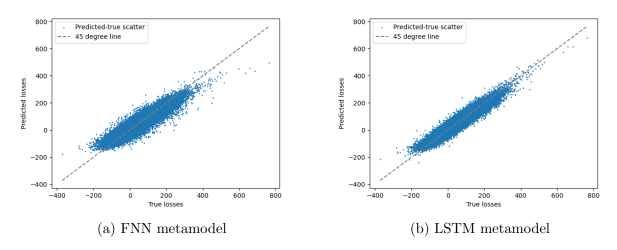


Figure 3.4: QQ-plots between true losses (x-axis) and predicted losses (y-axis) for neural network metamodels.

In Figure 3.4, we illustrate the fit of the neural network metamodels. The FNN has a better fit than the regression metamodels, but it still has a poor fit on the tails. We observe that the LSTM metamodels' QQ-plots closely follow the 45-degree line. Again, these metamodels are trained on noisy data, so the good fit to the true data should not be taken for granted. This implies that these models indeed learn the true feature-label relationship in the dynamic hedging simulation model (i.e., true loss labels) even though they are trained on noisy observations (e.g., training labels) of the model.

From a unique analytical standpoint, our numerical study offers a methodical exploration into the robustness of DNN models against noise in training labels. By employing the standard nested simulation procedure in Algorithm 1 as a data generator, we gain the ability to manipulate the noise level through adjusting the number of inner replications while keeping the same simulation procedure. This approach provides a controlled environment to examine the impact of label noise on neural network models. It allows us to generate our true dataset that approximates the true hedging losses with a high degree of precision, and, as a result, it enables us to explore the crucial question of whether deep learning models are capable of learning the true feature-label relationship from noisy training labels. Our numerical results in Table 3.2 and the QQ plots provide direct evidence that DNN models are indeed able to cut through the noise in the training labels and learn the true feature-label relationship. The LSTM metamodels' ability to learn the true feature-label relationship is crucial for the two-stage procedure to identify true tail scenarios and produce accurate CVaR estimates. Section 3.5.2 discusses LSTM metamodels in more details regarding their sensitivity to data quality and quantity. We aim to provide numerical evidence and provide insights on the DNN metamodels' noise tolerance.

#### 3.5.1 Two-Stage Procedure

We compare our two-stage procedure to a benchmark standard nested simulation procedure that runs N=1000 inner replications for each of the M=100,000 outer scenarios. In stage 1 of the proposed procedure, we first run inner simulations with N'=100 inner replications for each of the M=100,000 outer scenarios. So, stage 1's simulation budget is 10% of the standard procedure's. The resulting feature-label pairs  $\{(\mathbf{S}^{(i)}, \hat{L}_i) : i=1,\ldots,M\}$  is used for training different metamodels. Specifically, following the convention in machine learning literature, we split this dataset into three parts: The training, validation, and test sets have 90,000,5000, and 5000 samples (90%,5%, and 5% of the dataset), respectively. At the end of stage 1, m predicted tail scenarios are identified by the trained metamodels. In stage 2, N=1000 inner replications are run for all predicted tail scenarios. This is the same number of replications as the benchmark standard procedure. Stage 2's simulation budget is  $\frac{m}{M}$  of

the benchmark standard procedure's. In short, the two-stage procedure uses 15% - 30% of the benchmark procedure's budget for a safety margin between 0% - 15%. Analyzing the two stages separately, the simulation budget for stage 1 is 10% of the benchmark procedure's. Stage 2 uses 5% to 20% of the simulation budget of the benchmark procedure for a safety margin between 0% - 15%.

In our proposed two-stage procedure, the metamodel is used to identify the predicted tail scenario set, on which the standard nested simulation procedure is run in stage 2 to estimate the 95%-CVaR. An accurate identification of the true tail scenarios is crucial. In a two-stage procedure, metamodels' overall prediction errors measured by the MSEs in Table 3.2 are not the determining factor, but their ability to effectively rank the scenarios by their **true** losses is most critical in producing accurate CVaR estimates. A metamodel that uses less safety margin to rank can save more simulation budget in stage 2.

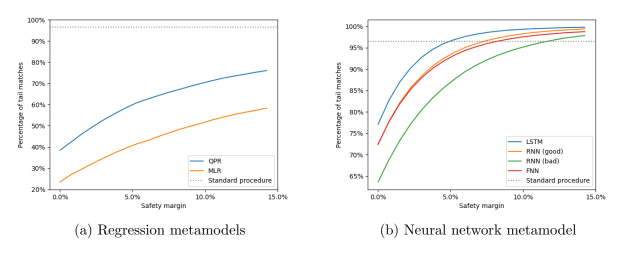


Figure 3.5: Percentage of correctly identified true tail scenarios by different metamodels.

Figure 3.5 depicts the average percentage of correctly identified true tail scenarios by different metamodels for increasing safety margins. Each quantity is averaged over 100 macro replications. For a fixed safety margin, the percentage of tail matches of the traditional regression metamodels are significantly lower than the ones of the neural networks. We observe that a poor metamodel like the QPR identifies less than 40% of the true tail scenario without any safety margin. In contrast, a well-suited metamodel like the LSTM identifies more than 75% of the true tail scenarios without any safety margin and more than the QPR metamodel does with 15% of safety margin<sup>2</sup>. The RNN metamodel suffers from

<sup>&</sup>lt;sup>2</sup>The metamodel in Dang et al. (2020) identifies 100% of the true tail scenarios on with a 10% safety

the vanishing gradient problem during training. For some macro replications (as shown in Figure 3.2b), the RNN metamodels' predictions are far from the true losses, especially on the tails. Seperating the good training macro replications from the bad ones, the results reconcile with the findings in Table 3.2. The FNN metamodel has a better fit than the regression metamodels, but it does not have the same level of accuracy as the LSTM metamodel, which is a specialized network to capture the sequential structure of our time-series features. For comparison, the horizontal dotted line shows the percentage of correctly identified true tails for the standard nested simulation procedure. A good metamodel should be able to identify a similar percentage of true tail scenarios as the standard procedure does with a reasonable safety margin. Otherwise, the two-stage procedure will offer no computational advantage in simulation budget over the standard procedure. The LSTM metamodel can reach similar percentage with a 5% safety margin. This indicates that the LSTM metamodel should be able to reach the same level of accuracy as the standard procedure with only 20% of the simulation budget.

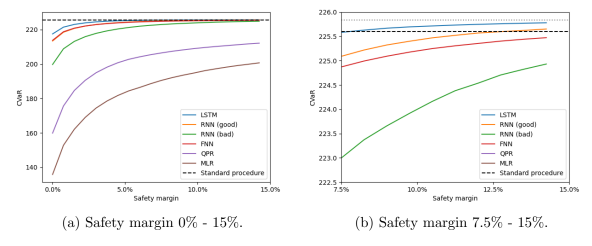


Figure 3.6: Average 95%-CVaR estimates by different procedures. Right figure is a zoomed-in version of left figure.

Lastly, we return to our original goal of estimating the 95%-CVaR of the dynamic hedging error. Figure 3.6 shows the 95%-CVaR estimates for all five metamodels with different safety margins, averaged over 100 macro replications. Figure 3.6b is a zoomed

margin for a GMAB contract. The GMAB contract is simpler than the GMWB contract, and the true tail scenarios are easier to identify. For a GMAB, our LSTM metamodel identifies 100% of the true tail scenarios with a 2.5% safety margin

version of Figure 3.6a. Because the safety margin only affects the two-stage procedure, the true 95%-CVaR and the one estimated by the standard procedure are horizontal (solid and dotted, respectively) lines in Figure 3.6. We observe that, with reasonable safety margins, the two-stage procedures with a LSTM metamodel consistently produce estimates that are as accurate as the standard procedure's estimate. The amount of computational savings is substantial. The LSTM is particularly superior to other metamodels as it accurately identifies the true tail scenarios and produces highly accurate CVaR estimates with small safety margins. By concentrating the simulation budget on the predicted tail scenarios, the two-stage procedure with the LSTM metamodel is able to achieve a similar accuracy as the standard procedure with a much smaller simulation budget. As the percentage of correctly identified tails approaches 100%, the two-stage procedure's CVaR estimates does not converge to the true CVaR but to the standard procedure's estimate. This is because the two-stage procedure's CVaR estimates are based on the standard procedure's estimates on the predicted tail scenarios, and the standard procedure's estimates themselves are also noisy.

#### 3.5.2 Noise Tolerance of DNN Metamodels

For financial and actuarial applications, regulators and practitioners are often concerned about the robustness of DNN models to noise in training labels, which hinders the adoption of these models in practice. Since the true relationship is unknown in real-world applications, most deep learning literature illustrates the impact of noise by artificially injecting noise into real-world datasets, which is already noisy before the injection. In our numerical experiments, we are able to use Monte Carlo simulation generate a true dataset that approximates the true hedging losses with a high degree of precision, and, as a result, we are able to explore the crucial question of whether deep learning models are capable of learning the true feature-label relationship from noisy training labels. In this section, we treat the standard nested simulation procedure as a data generator and examine the noise tolerance of LSTM metamodels by varying the numbers of outer scenarios (M) and inner replications (N) used to generate the training samples. The number of outer scenarios corresponds to the number of feature-label pairs in the training dataset, and the number of inner replications controls the noise level in the training labels. Recall that we use the standard nested simulation procedure with N=100 inner replications in our previous experiments, and we will refer to the resulting training dataset as the low-noise dataset. We also generate a medium-noise dataset and a high-noise dataset by running the standard nested simulation procedure with N=10 and N=1 inner replications, respectively. By altering the data quantity and quality, we conduct a sensitivity analysis on the LSTM metamodels' noise tolerance. We study the impact of noisy data on two LSTM metamodels with different model capacities, i.e., different numbers of trainable parameters. The two LSTM metamodels has the same number of layers, but their numbers of hidden units in each layer are different. More specifically, the high-capacity LSTM metamodel has 128 and 16 hidden units in the first and second LSTM layers, respectively, while the low-capacity LSTM metamodel has 32 and 4.

The maximum number of training samples is set to M=100,000. The training samples are split into training and test sets with a 90%-10% ratio. The true labels are generated by running the standard nested simulation procedure with N=100,000 inner replications for each of the 100,000 outer scenarios. The reason for using a maximum of 100,000 data points is precisely due to the computational cost of running the standard nested simulation procedure. For consistency, the LSTM metamodels are trained with the same architecture and training settings as in the previous experiments.

Model	N	Training error	Test error	True error
LSTM	100	$0.075(\pm 4.5 \times 10^{-3})$	$0.079(\pm 5.4 \times 10^{-3})$	$0.063(\pm 4.4 \times 10^{-3})$
High-capacity LSTM	100	$0.068(\pm 3.6 \times 10^{-3})$	$0.102(\pm 6.1 \times 10^{-2})$	$0.060(\pm 3.6 \times 10^{-3})$
Average Difference	100	-0.007	0.023	-0.003
LSTM	10	$0.195(\pm 1.1 \times 10^{-3})$	$0.193(\pm 1.7 \times 10^{-3})$	$0.070(\pm 9.1 \times 10^{-4})$
High-capacity LSTM	10	$0.157(\pm 2.0 \times 10^{-3})$	$0.199(\pm 1.9 \times 10^{-3})$	$0.065(\pm 1.9 \times 10^{-3})$
Average Difference	10	-0.038	0.006	-0.005
LSTM	1	$1.366(\pm 8.6 \times 10^{-3})$	$0.781(\pm 6.0 \times 10^{-3})$	$0.129(\pm 5.0 \times 10^{-3})$
High-capacity LSTM	1	$1.354(\pm 2.7 \times 10^{-2})$	$0.795(\pm 8.3 \times 10^{-2})$	$0.149(\pm 2.7 \times 10^{-2})$
Average Difference	1	-0.012	0.014	0.020

Table 3.3: MSEs of LSTM metamodels.

The first two rows of Table 3.3 shows the average squared errors and the half widths of their 95% confidence intervals between the metamodel predictions and the labels in the training dataset and the true dataset. N indicates the noise level in the training labels, i.e., a simulation dataset with fewer inner replications has a higher noise level. The test errors are included as a practical measure of the metamodels' generalization ability to unseen noisy samples. For this particular experiment, we are more interested in the true errors, which measure the metamodels' ability to learn the true feature-label relationship from the low-noise, medium-noise, and high-noise datasets. We first observe that all the true errors are lower than the training errors, indicating that the LSTM metamodels generalize well to predicting the true losses. Both LSTM metamodels are able to learn the true feature-label relationship from the low-noise and medium-noise datasets. However, when

the noise level is high, both LSTM metamodels have high true error. The diffences in the errors between the LSTM metamodels are also reported in the last rows of Table 3.3. A positive difference indicates that the high-capacity LSTM has higher errors than the LSTM. We observe that both LSTM metamodels have similar errors on the low-noise and medium-noise datasets. However, when the noise level is high, the high-capacity LSTM has higher errors than the regular LSTM. This is because the high-capacity LSTM has more trainable parameters and is more prone to over-fitting to the noise in the training labels. In contrast, the LSTM metamodel is more robust to label noise and is able to learn the true feature-label relationship better from the high-noise dataset. In practice, since the true relationship is unknown, the true error is not readily available. The test error approximates the true error and is directly observable. The difference in test errors of the LSTM metamodels are inconsistent with the differences in true errors. Therefore, test errors are not reliable indicators of the metamodels' noise tolerance. For practical applications that utilize nerual networks as metamodels, we recommend not simulating the test scenarios. Instead, we recommend simulating the true contract losses on a subset of training scenarios to evaluate the metamodels' generalization ability.

In summary, our numerical experiments demonstrate that both LSTM metamodels are capable of learning the true inner simulation model from noisy training labels. In most cases, a high-capacity metamodel is better. Nevertheless, in extreme circumstances when the noise level is too high, using a high capacity metamodel leads to severe over-fitting and poor generalization. Hence, domain knowledge about the right architecture for the task and knowledge about the noise level in the training labels are beneficial for choosing the right metamodel.

In more details, the QQ-plots depicted in Figure 3.7 illustrates the fit of the two LSTM metamodels when different noise levels are present in the training labels. They are arranged in a 2-by-3 grid, where the two rows correspond to the two LSTM metamodels with different model capacities, and the three columns correspond to the increasing noise levels. The Pearson correlation coefficient between the true losses and the predicted losses is labeled in each subplot. The sparsity of the plot increases as we move from left to right, indicating that the noise level in the training labels increases. There are two key findings.

- 1. When trained on the low-noise dataset, the metamodel predictions of high-capacity LSTM align more closely with the true labels than the regular LSTM. This is expected because the high-capacity LSTM has more trainable parameters and is able to learn more complex relationships better when the noise level is low.
- 2. When the noise level is medium or high, the high-capacity LSTM is more affected than the regular LSTM. This is because the high-capacity LSTM has more trainable

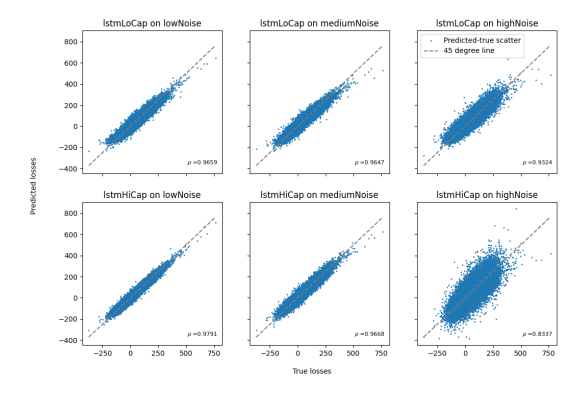


Figure 3.7: QQ-plots between true losses (x-axis) and predicted losses (y-axis) for two LSTM metamodels.

parameters and is more prone to over-fitting to the noise in the training labels. In contrast, the regular LSTM is more robust to label noise.

To further test the sensitivity of a LSTM metamodel and provide a more comprehensive view of the noise tolerance of DNN models, we conduct a sensitivity analysis on the noise tolerance of the regular LSTM metamodel by varying the numbers of outer scenarios (M) and inner replications (N) used to generate the training dataset. The number of outer scenarios corresponds to the number of data points in the training dataset, and the number of inner replications controls the noise level in the training labels.

Table 3.4 and 3.5 show the average squared errors between the LSTM predictions and the true losses for increasing M and N. The last rows of Table 3.4 and Table 3.5 show the performance of the regular and high capacity LSTM metamodels trained with M=100,000

	N = 1	N = 10	N = 100	N = 1000
M = 100	1.139	0.229	0.167	0.158
M = 1000	0.559	0.173	0.123	0.127
M = 10,000	0.283	0.115	0.099	0.097
M = 100,000	0.129	0.070	0.063	0.063

Table 3.4: MSE between regular LSTM predicted losses and true losses.

	N = 1	N = 10	N = 100	N = 1000
M = 100	0.764	0.408	0.131	0.087
M = 1000	0.878	0.367	0.156	0.087
M = 10,000	0.351	0.147	0.064	0.063
M = 100,000	0.149	0.065	0.060	0.038

Table 3.5: MSE between high-capacity LSTM predicted losses and true losses.

training labels with different noise levels. We observe that an increasing N reduces the MSE, but the reduction is not significant for a regular LSTM when N is larger than 10. For the high-capacity LSTM, the MSE is significantly reduced when N is increased from 1 to 100, but the reduction is not significant when N is increased from 100 to 1000. Another way to interpret the results in Table 3.4 is to compare the MSEs for the same budget of  $\Gamma = M \times N$ . Table entries on the same off-diagonal have the same simulation budget  $\Gamma$ . For most budgets, the MSEs are also the lowest when N=10. The results in Table 3.4 suggest that the performance of the LSTM metamodel is more sensitive to the number of outer scenarios than the number of inner replications. Treating the neural network as an advanced regression metamodel, we find this pheonomenon to be consistent with the results in Broadie et al. (2015), where the authors show that the performance of a regression-based nested simulation procedure is more affected by the number of outer scenarios. The last row in Table 3.4 and 3.5 show that the high-capacity LSTM metamodel is relatively insensitive to the number of inner replications. To make the most efficient use of a fixed simulation budget, the number of inner replication should be set constant around N=10while increasing the number of outer scenarios to the maximum.

To further investigate the LSTM metamodels' sensitivity to data quantity and quality (M and N, respectively), we report the Spearman rank correlation coefficients Table 3.6, which measure the ability to rank the scenarios by their true losses. It is an appropriate measure of the metamodel's performance in the two-stage procedure, where the metamodel

is used to identify the predicted tail scenario set, on which extensive simulations are run in stage 2 to estimate the 95%-CVaR. The Pearson correlation coefficients are also included in the parentheses to illustrate the linear correlation between the predicted losses and the true losses. They measure the metamodel's overall prediction accuracy. Our previous numerical experiments illustrated in Figure 3.5 have shown that the LSTM metamodel predictions have high Spearman correlation with the true labels for the combination M=100,000 and N=100. We intend to examine if LSTM predictions also have high Pearson correlation with the true labels. A high Pearson correlation suggests the possibility of using LSTM predictions to estimate risk measures directly.

	N = 1	N = 10	N = 100	N = 1000
M = 100	0.638 (0.881)	0.875 (0.915)	0.896 (0.903)	0.937 (0.941)
M = 1000	$0.722 \ (0.768)$	0.899 (0.908)	$0.886 \ (0.891)$	0.922(0.927)
M = 10,000	$0.845 \ (0.630)$	0.937 (0.900)	$0.908 \; (0.905)$	0.947 (0.948)
M = 100,000	$0.935 \ (0.640)$	$0.963 \ (0.909)$	$0.927 \ (0.922)$	$0.965 \ (0.966)$

Table 3.6: Spearman (Pearson) correlation coefficients of high-capacity LSTM predictions.

Consistent with our previous numerical experiments, the Spearman correlations of a high-capacity LSTM is high for a moderate simulation budget. For  $M=100{,}000$ , the Spearman correlation is relatively insensitive to N. This finding supports further budget savings for our two-stage procedure. In stage 1, we can lower N from 100 to 10 almost without compromising on tail scenario preditions.

For an entry of Table 3.6, the total simulation budget  $\Gamma = MN$ . Entries on the same off-diagonal all have the same  $\Gamma$ . The Pearson correlation coefficients are significantly lower for N=1 than the other N values.

Figure 3.8 shows the difference between the Spearman and Pearson correlation coefficients for the high-capacity LSTM metamodel. The heatmap is generated by subtracting the Pearson correlation coefficients from the Spearman correlation. We observe that Spearman correlation coefficients are not significantly different from the Pearson correlation coefficients for any N larger than 10. This is a strong evidence that the LSTM metamodel is not only able to effectively rank the scenarios by their true losses, but also able to make accurate loss predictions. Instead of using the LSTM metamodel only for classifying tail scenarios in a two-stage procedure, LSTM's ability to cut through a moderate level of noise in training labels encourages us to use its predictions to estimate the CVaR directly.



Figure 3.8: Difference between Spearman and Pearson correlations for high-capacity LSTM metamodel.

#### 3.5.3 Single-Stage Procedure

The accuracy and robustness of the LSTM metamodels motivate us to propose a single-stage procedure that uses the metamodel predictions to estimate the CVaR directly. Instead of relying on the standard nested simulation procedure in stage 2, the single-stage procedure can be even more efficient. In this section, we compare the single-stage procedure to the two-stage procedure and the standard procedure in estimating the 95%-CVaR of the hedging losses for GMWB. In our numerical experiments, the results for the single-stage procedure is obtained with the same metamodels as in the two-stage procedure. The only difference from Section 3.5.1 is that the metamodel predictions are used to estimate the risk measures directly.

Figure 3.9 shows the boxplots of the 95%-CVaR estimates of the single-stage procedure with different metamodels and compares them to the two-stage procedure and the standard procedure. The two-stage procedure uses the same simulation budget as the single-stage procedure in stage 1 with extra budget for extensive inner simulation on the predicted tail scenarios in stage 2. The metamodel with the best performance in the two-stage procedure is selected, and the safety margin is set to 0%. The standard procedure shown in Figure 3.9 uses the noisy loss labels in the training dataset to estimate the CVaR directly, which uses the same simulation budget as the single-stage procedure. We observe that the single-

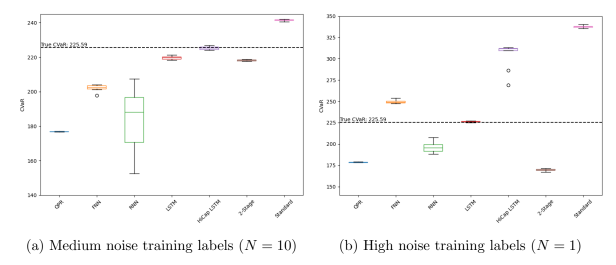


Figure 3.9: CVaR estimates of the single-stage procedure with metamodels.

stage procedures with the regular LSTM metamodel consistently produce CVaR estimates that are closer to the true value than the standard procedure's estimate. This finding is especially impressive when the noise level in the training labels is high. It is another strong evidence that a well-selected LSTM metamodel is able to cut through the noise in the noisy training labels and make accurate loss predictions that lead to accurate CVaR estimates. The difference in performance among the metamodels is more pronounced in the single-stage procedure than in the two-stage procedure. Trained using medium noise labels, high-capacity LSTM metamodel consistently produces CVaR estimates that are closer to the true value than the regular LSTM, while the other metamodels produce even worse estimates. However, with high noise training labels, the high-capacity LSTM suffers from severe overfitting. This is also evident in Table 3.3. Since we are using the metamodel predictions to estimate the CVaR directly without any safety margin, the metamodel's overall prediction accuracy becomes more important. In other words, the single-stage procedure is more sensitive to the metamodel's ability to make accurate loss predictions than the two-stage procedure. Previously, a generic FNN metamodel performs well in the two-stage procedure. However, results in Figure 3.9 suggest that the FNN metamodel should not be used in the single-stage procedure.

A single-stage procedure with a regular LSTM metamodel is particularly superior to the two-stage procedure, as it is able to achieve a higher accuracy as the standard procedure with a much smaller simulation budget. Note that the single-stage procedure does not require a safety margin. By avoiding the calibration of the safety margin, it is more

straight-forward to implement and is more efficient than the two-stage procedure. For a two-stage procedure with a 0% safety margin, the extra computational cost is from the extensive inner simulation in stage 2. On 4 20-core Intel Xeon Gold 6230 processors, stage 2 of the two-stage procedure takes 30 minutes to run with parallel processing, while the 95% confidence band of training time of the high-capacity LSTM metamodel is  $(19.64 \pm 0.94)$  minutes on a Nvidia RTX 3060 Ti GPU. While the accuracy of the two-stage procedure is highly dependent on the safety margin, increasing the safety margin introduces extra computational cost. Therefore, while achieving a higher accuracy in estimating the CVaR, the single-stage procedure requires only 60% computation time of the two-stage procedure.

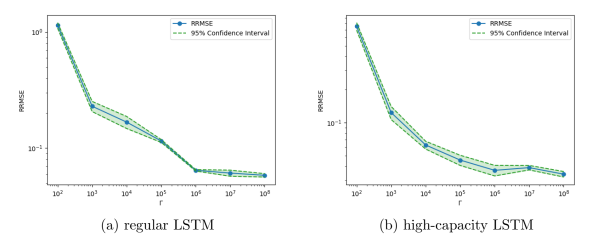


Figure 3.10: Empirical convergence of CVaR for the single-stage procedure with LSTM metamodels.

To further investigate the single-stage procedure's performance, we conduct a convergence analysis on the LSTM metamodels. Figure 3.10 shows the log-log plots of the relative root mean square errors (RRMSE) between the LSTM metamodel CVaR predictions and the true CVaR against the total simulation budget  $\Gamma$ . For each budget  $\Gamma$ , the metamodels are trained for different combinations of M and N, and the metamodel with the best RRMSE is plotted. While numerical results suggests that the CVaR estimator of the single-stage procedure with LSTM metamodels may have better accuracy than the two-stage procedure and the standard procedure when the number of outer scenarios is further increased, we found it difficult to compare their performance due to insufficient computational resources. For  $\Gamma \leq 100,000$ , the number of outer scenarios is fixed at M = 100,000, and only the number of inner replications is varied. Instead, we try to analyze the effect

of the number of outer scenarios and the number of inner replications separately by fixing one and varying the other.



Figure 3.11: Empirical convergence of the single-stage procedure with a LSTM metamodel.

Figure 3.11 shows the log-log plots of the RRMSE between the LSTM metamodel CVaR predictions and the true CVaR against the simulation budget. The left figure shows the empirical convergence of the RRMSE for increasing inner replications with a fixed number of outer scenarios (M = 100,000), and the right figure shows the empirical convergence of the RRMSE for increasing outer scenarios with a fixed number of inner replications (N =10). Due to computational constraints, we are not able to run the two-stage procedure and the standard procedure with more outer scenarios or more inner replications, therefore the comparison between the single-stage procedure is only available up to M=100,000 and N=1000. We observe that the RRMSE decreases as the simulation budget increases, and the rate of convergence is higher when the quantity of the data increases. For an LSTM metamodel, increasing the data quality of the training labels has diminishing returns, which is consistent with the results in Figure 3.11. For an increasing number of inner replications with fixed number of outer scenarios, the RRMSE cease to decrease after reaching N = 100. More specifically, when the quality of the training labels is fixed at N=10, the CVaR estimator of single-stage procedure with a LSTM metamodel converge roughly in the order of  $\mathcal{O}(M^{\frac{2}{3}})$ . This observation resonates with the best possible convergence rate of a standard nested simulation procedure in Gordy and Juneja (2010). Hence, in practice, we suggest to run the single-stage procedure with a moderate number of inner replications and a large number of outer scenarios to achieve a high level of accuracy with a reasonable

#### 3.6 Conclusion

The proposed nested simulation procedures with DNN metamodels result in substantial computational savings in estimating CVaR of the hedging loss of a VA contract from accurately predicting the hedging losses and identifying the tail scenarios. When new outer scenarios are generated, a trained LSTM metamodel can distinguish between tail and nontail scenarios and make accurate predictions without the need to run new inner simulations. Our novel experiment design allows us to examine the impact of label noise on DNN models. We find that a DNN with a suitable architecture is able to cut through the noise in training labels and learn the true complex dynamic hedging model. By showcasing the resilience of these models, our study aims to influence regulatory bodies towards recognizing the value and applicability of deep learning metamodels in financial risk management, and it provides informed suggestions and guidance for the incorporation and oversight of advanced deep learning metamodel in Monte Carlo Simulation in financial applications. Our findings are particularly insightful in this context. In our numerical experiments, a LSTM metamodel is resilient to a high level of noise in training labels and is able to make accurate predictions. This is a promising evidence that DNN metamodels can be used to improve the efficiency of Monte Carlo simulation for quantitative risk management tasks that require a computational-expensive simulation procedure. We propose two nested simulation procedures that use DNN metamodels to estimate risk measures of the hedging loss of a VA contract. For estimating tail risk measures, our two-stage procedure is designed to address regulatory concerns by avoiding the direct use of metamodel predictions but instead use them to identify the potential tail scenarios. An extensive inner simulation is performed to achieve a high level of accuracy on the predicted tail scenarios. However, the safety margin in the two-stage procedure is a user's choice and is not easy to determine before running extensive numerical experiments.

Our single-stage procedure uses the metamodel predictions to estimate the risk measure directly. It is more efficient and can be extended to estimate risk measures that require knowledge of the entire loss distribution. Our numerical experiments demonstrate that the proposed single-stage procedure with DNN metamodels result in further computational savings over our two-stage procedure. Furthermore, our numerical results provide evidences for adopting DNN metamodels in Monte Carlo simulation for risk management tasks. Through our systematic empirical study of the noise tolerance of DNN metamodels, we address regulatory concerns by showing that a LSTM metamodel with moderate

model capacity is resilient to a high level of noise in training labels and is able to make accurate predictions. A possible future research direction is to apply DNN metamodels in other financial risk management tasks that requires complex nested simulation with high-dimensional outer scenarios. Another future research direction is to investigate the impact of label noise on other deep learning models, such as convolutional neural networks and transformer models, and to compare their performance with LSTM metamodels in nested simulation procedures. From a practical standpoint, the choice of a suitable DNN architecture is crucial for the success of a deep learning metamodel in nested simulation procedures. We find that a LSTM metamodel is the most suitable for our dynamic hedging simulation model with time series features, but the optimal network architecture may vary for different simulation models. Exploring DNN metamodels in other complex risk management tasks presents a promising avenue for research, especially as these tasks often involve complex, high-dimensional scenarios where traditional methods are insufficient. The versatility of neural networks could unlock new insights across a broad spectrum of financial and actuarial applications.

In this chapter, we have demonstrated the potential of DNN metamodels to estimate risk measures with high accuracy and efficiency when simulation data is abundant. In practice, simulation data is often scarce for new market conditions and new insurance products. DNN metamodels are known to suffer from over-fitting when the number of training samples is limited. In the next chapter, we will extend our study to transfer learning and show that DNN metamodels trained on a existing VA contract can be transferred to a new VA contract with different market conditions. We hypothesize that the DNN metamodel will generalize well to the new VA contract, and the computational savings from transfer learning will be significant.

# Chapter 4

# Transfer Learning for Rapid Adaptation of Deep Neural Network Metamodels in Dynamic Hedging of Variable Annuities

## 4.1 Introduction

In the evolving landscape of financial markets, insurance products such as VAs have gained significant interest due to their ability to provide both investment growth and guaranteed benefits. Managing the risks associated with these products in volatile market conditions is a complex task that demands sophisticated financial modeling techniques like nested simulation. A standard nested simulation procedure involves running a large number of simulations to generate a dataset of scenario-wise contract losses with two levels of simulations. Practitioners can estimate tail risk measures using the realized losses for different risk scenarios obtained from these simulations. Nested simulation is extremely computationally expensive. To address the computational burden, metamodeling techniques have been proposed, where a surrogate model approximates the outcomes of a complex simulation procedure. Traditional machine learning models often struggle to capture the intricate nonlinear relationships and temporal dependencies inherent in financial data. In particular, DNNs, and specifically LSTM networks, have been employed as metamodels to predict the outcomes of the inner simulations efficiently. Chapter 3 introduces a metamodeling-based nested simulation framework for dynamic hedging of VAs that approximates the

inner simulation by an LSTM metamodel. It shows that LSTMs, as a type of gated RNNs, are well-suited as metamodels for Monte Carlo simulation of financial time series and can capture the long-term dependencies in the simulation dataset. This property is crucial for the application of dynamic hedging of VAs.

Despite the advantages of using DNN metamodels, computational challenges arise when market conditions change or new VA contracts with different features are introduced. Retraining neural network metamodels from scratch in response to every change is computationally inefficient and time-consuming. Practitioners with limited resources face several challenges:

- 1. The simulation budget is limited, but DNN metamodels require extensive training data to learn the feature-label relationship (Golestaneh et al., 2024).
- 2. The computational cost of retraining a suitable DNN metamodel is often much higher than fitting a traditional parametric regression.
- 3. Financial markets are dynamic with frequent shifts in volatility, interest rates, and other risk factors (Cont, 2001).

Constraints on the simulation budget is the most critical challenge for dynamic hedging of VAs. Any method that involves nested simulation is computationally expensive, and the computational cost is often prohibitive for changing market conditions. Therefore, it is essential to develop methods that can rapidly adapt existing metamodels to new conditions without incurring the full computational cost of retraining and extensive additional simulations. Transfer Learning (TL) offers a compelling solution to this problem by enabling the reuse of a pre-trained model on a new but related task (Pan and Yang, 2009). In the context of DNNs, TL involves leveraging the knowledge acquired during training on one dataset to improve learning performance on a different dataset (Yosinski et al., 2014). This approach can significantly reduce the amount of training data needed, training time and computational resources while enhancing model generalization. Instead of starting from scratch, a new DNN metamodel can be built on top of a pre-trained metamodel and fine-tuned on new data. This allows the metamodel to adapt quickly to unseen conditions and new VA contracts.

In this chapter, we explore the application of TL to the dynamic hedging of VAs using LSTM metamodels. We propose a novel TL framework that accelerates the training of DNN metamodels for nested simulation procedures in dynamic hedging. This setting is particularly relevant for dynamic hedging problems, as insurance companies often underwrite new contracts under changing market conditions and actuarial assumptions. This

chapter proposes a TL framework that makes fast adaptation of metamodels possible. Our approach involves pre-training an LSTM network on a large dataset of VA simulations and then fine-tuning it on a smaller dataset of new VA contracts. We evaluate the performance of the TL framework on a real-world dataset of VA contracts and compare it with training a LSTM metamodel from scratch. More importantly, we show the strengths and weaknesses of different TL techniques for rapid adaptation of LSTM metamodels.

The rest of this chapter is organized as follows. Section 4.2 provides an overview of the dynamic hedging problem for VAs and the use of LSTM networks as metamodels in a nested simulation procedure. Section 4.2.4 introduces the TL framework for rapid adaptation of LSTM metamodels in dynamic hedging. Section 4.3 presents the experimental setup and results, comparing the performance of TL with training from scratch. Finally, Section 4.4 concludes the chapter and discusses future research directions.

# 4.2 Transfer Learning in Financial Metamodeling

TL is a machine learning paradigm where knowledge acquired from a source task is utilized to improve learning performance on a related target task. One of the primary applications of TL in finance is asset price prediction. Traditional models, such as geometric Brownian motion (GBM), autoregressive integrated moving average (ARIMA) and generalized autoregressive conditional heteroskedasticity (GARCH), have been widely used in time series modeling. ARIMA models (Box and Pierce, 1970) combine autoregression, differencing, and moving average components to capture linear relationships in time series data. GARCH models (Bollerslev, 1990), extend ARCH models to better capture volatility clustering, which is a key stylized fact of financial time series (Cont, 2001). However, these models often struggle to capture complex dynamics, e.g., regime changes (Hamilton, 1989) and extreme market events (Embrechts et al., 2013) that are prevalent in financial markets. In Chapter 3, our numerical experiment uses a regime-switching geometric Brownian motion (RS-GBM) as the underlying asset model. For estimating tail risk measures, a metamodel-based nested simulation procedure is proposed to approximate the inner simulation. We have shown DNN metamodels like RNN and LSTM networks have demonstrated significant improvements in modeling temporal dependencies, and they save a substantial amount of simulation budget. However, training these models from scratch requires extensive simulation data, which may not always be available for specific assets or under certain market conditions. TL offers a solution to this problem by leveraging knowledge from related assets or tasks to improve the learning performance of DNN metamodels on the target task. In algorithmic trading, Jeong and Kim (2019) used TL to enhance the performance of a reinforcement learning agent by preventing overfitting from insufficient market data. TL techniques have also been applied in building fraud detection systems. Financial fraud often exhibits subtle and evolving patterns, and it is challenging to develop robust detection models. By transferring previous knowledge from detected fraud cases, models can adapt to detect new fraud schemes more effectively (Lebichot et al., 2021). Yan et al. (2024) conduct a comprehensive survey study of current TL techniques in financial applications, and they find almost all applications of TL only employ parameter transfer, where the pre-trained model is fine-tuned on the target task. Our multi-task learning framework in Section 4.2.4 extends beyond parameter transfer to shared representation learning across multiple tasks.

Regarding nested simulation of VAs, Cheng et al. (2019) is the most relevant study to our work, where they proposed a TL framework for fast valuation of large portfolios of VAs. Instead of using stochastic kriging (Gan and Lin, 2015), they employed a pre-trained DNN to select the best representative scenarios from a large portfolio of VAs. In our work, we focus on the application of TL to accelerate the training of LSTM metamodels for nested simulation in dynamic hedging of VAs. The fine-tuned LSTM metamodels can be readily adapted to the two-stage procedure and the single-stage procedure in Chapter 3 to predict the contract losses under different scenarios.

In our context of DNNs metamodeling-based simulation procedures for hedging VAs, TL involves pre-training a DNN metamodel where the simulation budget is abundant and then fine-tuning it on a smaller dataset of new contracts or market conditions. Written on the same underlying asset, different VA contracts may share common features or exhibit similar patterns, especially temporal dependencies and regime changes in the underlying financial time series. Similarly, two VAs with the same features but based on different underlying assets may have some shared characteristics that can be leveraged to improve the learning performance of the metamodel. By transferring knowledge from a pre-trained model to a new but related task, TL can significantly reduce the computational cost of training the metamodel on the target task.

LSTM networks are well-suited for modeling sequential data due to their ability to capture long-term dependencies (Hochreiter and Schmidhuber, 1997). For the application of VA risk management using metamodel-based nested simulation, LSTM networks approximate the inner simulation, i.e., the mapping from scenarios to the scenario-wise contract losses. In Chapter 3, we treat metamodeling as a supervised learning problem and demonstrated that LSTM networks can effectively model this complex relationships with extensive training on a large dataset of VA simulations. The total computational cost originates from two sources: running the standard nested simulation procedure to generate the training data and training the LSTM network on the generate dataset. Given

a new VA contract, these steps need to be repeated to adapt the LSTM metamodel to new market conditions and contract features. For a DNN with a large number of parameters, retraining the LSTM network from scratch can be computationally expensive. Generating a large training dataset using the standard nested simulation procedure is costly, and the computational burden can be prohibitive for certain assets or market conditions. TL offers an alternative approach that keeps the previous knowledge, i.e., the pre-trained model, as a foundation. Building on the pre-trained model, the LSTM network can be fine-tuned on a smaller dataset of new contracts or market conditions. This approach accelerates the adaptation of the LSTM metamodel to new conditions. Computational savings come in two forms:

- 1. fine-tuning requires less training time than training LSTM metamodels from scratch, and
- 2. fewer training data points are needed to achieve a good LSTM metamodel, which is particularly beneficial when the standard nested simulation procedure is costly.

In supervised learning, a **domain**  $\mathcal{D}$  comprises a feature space  $\mathcal{X}$  and a marginal probability distribution F. Correspondingly, a **task**  $\mathcal{T}$  consists of a label space  $\mathcal{Y}$  and a predictive function  $f: \mathcal{X} \to \mathcal{Y}$  that maps input features to output labels.

A typical TL framework for supervised learning consists of the following:

- source domain:  $\mathcal{D}_{So} = \{\mathcal{X}_{So}, F_{So}(\mathbf{X})\}$
- source task:  $\mathcal{T}_{So} = \{\mathcal{Y}_{So}, f_{So}(\cdot)\}$
- target domain:  $\mathcal{D}_{Ta} = \{\mathcal{X}_{Ta}, F_{Ta}(\cdot)\}$
- target task:  $\mathcal{T}_{Ta} = \{\mathcal{Y}_{Ta}, f_{Ta}(\cdot)\}$

where  $\mathcal{X}_{So}$  and  $\mathcal{X}_{Ta}$  include input features derived from the outer simulation of two different VA contracts or market conditions, and  $F_{So}$  and  $F_{Ta}$  are the marginal probability distributions of the source and target domains, respectively. Hence, the source domain and task are related to one VA contract for a certain market condition, while the target domain and task are related to new VA contracts and market conditions for which we don't have the simulation budget to run the standard nested simulation procedure extensively. In our numerical experiments, the dataset is generated by running the standard nested simulation procedure in Algorithm 1 for a large number of VA contracts. The input features  $\mathbf{X}$  are

the return vectors from Equation 3.5, and the output labels L are the contract losses for different outer scenarios. The stock price vector  $\mathbf{S}$  is not crucial for the discussion of this chapter and is omitted since it can be immediately recovered from  $\mathbf{X}$  and the initial stock price  $S_0$ . We keep the number of features to be 240 for both source and target domains as in Section 3.5. The predictive function  $f_{So}$  and  $f_{Ta}$  are trained to estimate outputs  $L_{So}$  and  $L_{Ta}$ , i.e., approximating the inner simulations under the source and target tasks, respectively. TL seeks to improve the learning of the target predictive function  $f_{Ta}(\cdot)$  in  $\mathcal{D}_{Ta}$  by leveraging the previous training on  $\mathcal{D}_{So}$  and  $f_{So}(\cdot)$ , particularly when  $\mathcal{D}_{So} \neq \mathcal{D}_{Ta}$  or  $\mathcal{T}_{So} \neq \mathcal{T}_{Ta}$ . In Chapter 3,  $\mathcal{D}_{So} = \mathcal{D}_{Ta}$  and  $\mathcal{T}_{So} = \mathcal{T}_{Ta}$ .

In our TL setting, both the source and target tasks involve learning a mapping from the risk factors to the VA contract losses. For the source task, we have a dataset  $\mathcal{D}_{So} = (X_{So}^{(i)}, L_{So}^{(i)})_{i=1}^{M_{So}}$ , where  $M_{So}$  is the number of training samples, i.e., the number of outer simulation paths used to generate the dataset with a standard nested simulation procedure.  $X_{So}^{(i)} \in \mathcal{X}_{So}$  and  $L_{So}^{(i)} \in \mathcal{Y}_{So}$ . The ultimate objective is to learn a metamodel  $f_{Ta}$  that predicts the VA contract losses  $L_{So}$  from the scenarios  $X_{Ta}$  in the form of a financial time series. TL starts by training a metamodel  $f_{So}$  on the source domain  $\mathcal{D}_{So}$ . For both the two-stage procedure and the single-stage procedure in Chapter 3,  $f_{So}$  is approximated by an LSTM network  $f_{So}(\cdot;\theta_{So})$ , where the parameters of the LSTM network,  $\theta_{So}$  are learned by minimizing a MSE loss function on the source domain. Then the pre-trained parameters  $\theta_{So}$  to inform the learning of  $f_{Ta}(\cdot;\theta_{Ta})$  on the target domain  $\mathcal{D}_{Ta}$ . The training on the target domain should encourage similarity between  $\theta_{So}$  and  $\theta_{Ta}$  to facilitate the transfer of knowledge.

The most common TL techniques for supervised learning include fine-tuning, layer freezing, and multi-task learning. These techniques can be categorized based on how they leverage pre-trained metamodels and how they encourage the similarity between  $f_{So}(\cdot; \theta_{So})$  and  $f_{Ta}(\cdot; \theta_{Ta})$ .

# 4.2.1 Fine-tuning

Fine-tuning is a commonly used TL technique that uses the same neural network architecture for both the source and target tasks. We first train the source LSTM metamodel  $f_{So}(\cdot;\theta_{So})$  on  $\mathcal{D}_{So}$ , capturing temporal dependencies and patterns relevant to the source task. For the target task, we initialize  $\theta_{Ta} = \theta_{So}$  and proceed to fine-tune the entire network on  $\mathcal{D}_{Ta}$  using a smaller learning rate.

Algorithm 4 assumes that the learned sequential representations are beneficial for the target task. Only minor adjustments are needed to adapt the metamodel to the new

#### **Algorithm 4** Fine-tuning Metamodel for a Target Task

- 1: **Input:** source dataset  $\mathcal{D}_{So} = \{(X_{So}^{(i)}, L_{So}^{(i)})\}_{i=1}^{M_{So}}$ , target dataset  $\mathcal{D}_{Ta} = \{(X_{Ta}^{(i)}, L_{Ta}^{(i)})\}_{i=1}^{M_{Ta}}$ , learning rate  $\alpha_{So}$  and smaller learning rate  $\alpha_{Ta}$ .
- 2: Train a LSTM metamodel  $f_{So}(\cdot; \theta_{So})$  on  $\mathcal{D}_{So}$ :

$$\theta_{So} = \min_{\theta} \frac{1}{M_{So}} \sum_{i=1}^{M_{So}} \left( f_{So}(X_{So}^{(i)}; \theta) - L_{So}^{(i)} \right)^2$$
(4.1)

3: Initialize the target metamodel parameters  $\theta_{\text{Ta}}$  using the pre-trained metamodel parameters:

$$\theta_{\mathrm{Ta}} \leftarrow \theta_{\mathrm{So}}$$

4: Fine-tune the entire LSTM model  $f_{\text{Ta}}(\cdot; \theta_{\text{Ta}})$  on the target dataset  $\mathcal{D}_{\text{Ta}}$  using a smaller learning rate  $\alpha_{\text{Ta}}$ :

$$\theta_{\text{Ta}} = \min_{\theta} \frac{1}{M_{\text{Ta}}} \sum_{i=1}^{M_{\text{Ta}}} \left( f_{\text{Ta}}(X_{\text{Ta}}^{(i)}; \theta) - L_{\text{Ta}}^{(i)} \right)^2$$
(4.2)

5: Output: Final adapted LSTM metamodel  $f_{\text{Ta}}(\cdot; \theta_{\text{Ta}})$  for the target task

task, and the training process is accelerated by the pre-trained parameters. Fine-tuning is particularly useful when the nested simulation procedures for the two VA contracts are closely related. Only minor adjustments are needed to adapt the LSTM metamodel to the new domain.

## 4.2.2 Layer Freezing

In a layer freezing approach, we partition the model parameters into frozen parameters  $\theta_0$  and trainable parameters  $\theta_1$ , such that  $\theta = [\theta_0, \theta_1]$ . Typically,  $\theta_0$  are parameters of the lower layers, and  $\theta_1$  are parameters of the higher layers that include the output layer. The intuition behind layer freezing is that the lower layers capture general features that are transferable across tasks, while the higher layers are more task-specific.

# 4.2.3 Multi-task Learning

Multi-task learning (Caruana, 1997) refers to a machine learning paradigm where a single model is trained simultaneously on multiple related tasks. Shared representations are

#### **Algorithm 5** Layer Freezing for Metamodel Transfer

- 1: **Input:** source dataset  $\mathcal{D}_{So} = \{(X_{So}^{(i)}, L_{So}^{(i)})\}_{i=1}^{M_{So}}$ , target dataset  $\mathcal{D}_{Ta} = \{(X_{Ta}^{(i)}, L_{Ta}^{(i)})\}_{i=1}^{M_{Ta}}$ , learning rates  $\alpha_{So}$  and  $\alpha_{Ta}$ , frozen parameters  $\theta_0$ , trainable parameters  $\theta_1$ .
- 2: Train LSTM model  $f_{So}(\cdot; \theta_{So})$  on  $\mathcal{D}_{So}$ :

$$\theta_{So} = [\theta_0, \theta_1] = \min_{\theta} \frac{1}{M_{So}} \sum_{i=1}^{M_{So}} \left( f_{So}(X_{So}^{(i)}; \theta) - L_{So}^{(i)} \right)^2$$
(4.3)

3: Initialize the target model parameters  $\theta_{\text{Ta}} = [\theta_0, \theta_1]$  using the pre-trained source model parameters  $\theta_{\text{So}}$ :

$$\theta_{\mathrm{Ta}} \leftarrow \theta_{\mathrm{So}} = [\theta_0, \theta_1]$$

- 4: Freeze the parameters of the shared layers  $\theta_0$ :
- 5: Fine-tune the trainable layers  $\theta_1$  on the target dataset  $\mathcal{D}_{Ta}$  using Algorithm 4:

$$\theta_{\text{Ta}} = \min_{\theta_1} \frac{1}{M_{\text{Ta}}} \sum_{i=1}^{M_{\text{Ta}}} \left( f_{\text{Ta}}(X_{\text{Ta}}^{(i)}; [\theta_0, \theta_1]) - L_{\text{Ta}}^{(i)} \right)^2$$
(4.4)

6: Output: Final adapted LSTM metamodel  $f_{\text{Ta}}(\cdot; [\theta_0, \theta_1])$  for the target task

learned across tasks, which can improve learning efficiency and predictive performance on each individual task with limited data. In contrast to fine-tuning and layer freezing, multitask learning aims to leverage learned knowledge from multiple tasks, and all tasks are trained simultaneously.

Let  $\{\mathcal{T}_k\}_{k=1}^K$  represent a set of K related tasks, each corresponding to a metamodeling task for a different standard nested simulation procedure. For each task  $\mathcal{T}_k$ , we have a dataset  $\mathcal{D}_k = (X_k^{(i)}, L_k^{(i)})i = 1^{M_k}$ , where  $M_k$  is the number of training samples for task k.  $X_k^{(i)}$  and  $L_k^{(i)}$  are the features and contract loss labels for task k, respectively.

For our metamodeling implementation in Chapter 3, we consider a multi-task learning framework where the LSTM layers are shared across multiple tasks, and each task has its own fully connected layer for prediction. The network parameters are divided into shared parameters  $\theta_0$  (LSTM layers) and task-specific parameters  $\theta_k$  (fully connected layers for task k). As opposed to layer freezing, all parameters are trainable. The objective function for multi-task learning is the sum of the loss functions of all tasks:

$$\min_{\theta_0, \theta_1, \dots, \theta_K} = \sum_{k=1}^K \frac{1}{M_k} \sum_{i=1}^{M_k} \left( f_i(X_k^{(i)}; \theta_0, \theta_1, \dots, \theta_K) - L_k^{(i)} \right)^2, \tag{4.5}$$

where MSE loss function is used as the error metric, and  $f_i(\cdot; \theta_0, \theta_1, \dots, \theta_K)$  is the output of the network for task k. In essence, multi-task learning uses a multi-head architecture, where each task has its own output head, but the shared LSTM layers learn a common representation across tasks. Transfer occurs through the shared LSTM layers  $\theta_0$ . These layers learn representations of temporal patterns and dependencies common to all tasks, effectively **pooling** information from multiple simulation schemes. The task-specific fully connected layers  $[\theta_1, \dots, \theta_K]$  allow each task to capture unique characteristics not shared with other tasks.

#### Algorithm 6 Multi-task Learning Framework for LSTM Metamodels

- 1: **Input:** learning rate  $\alpha$ , set of K tasks  $\{\mathcal{T}_k\}_{k=1}^K$  with datasets  $\mathcal{D}_k = \{(X_k^{(i)}, L_k^{(i)})\}_{i=1}^{M_k}$ , task-specific parameters  $\theta_k$  for each task k, and shared parameters  $\theta_0$ .
- 2: Train the multi-head LSTM metamodel on all K tasks simultaneously by minimizing the multi-task loss function:

$$\min_{\theta_0, \{\theta_k\}_{k=1}^K} \sum_{k=1}^K \frac{1}{M_k} \sum_{i=1}^{M_k} \left( f_i(X_k^{(i)}; \theta_0, \theta_k) - L_k^{(i)} \right)^2 \tag{4.6}$$

- 3: Update both the shared parameters  $\theta_0$  and task-specific parameters  $\{\theta_k\}_{k=1}^K$  simultaneously using backpropagation and gradient descent with learning rate  $\alpha$ .
- 4: Output: Trained multi-task LSTM metamodel  $f(\cdot; \theta_0, \{\theta_k\}_{k=1}^K)$  for all K tasks

Algorithm 6 outlines the multi-task learning framework for LSTM metamodels for nested simulation procedures. In Chapter 2 and Chapter 3, we demonstrate that pooling using a metamodel can improve the computation efficiency and estimator accuracy in nested simulation procedures. Here, pooling happens at a higher level, where information from multiple metamodels are shared to improve the learning performance on individual tasks. The multi-task loss function encourages the shared LSTM layers to learn generalizable representations of temporal dependencies that are beneficial across all tasks.

Choosing appropriate simulation schemes is crucial for the success of multi-task learning. The tasks should be related to ensure that the shared layers can capture common features beneficial across tasks. Criteria for selecting tasks with similar simulation schemes include:

- Similarity in contract specifications: simulations involving VA contracts with similar features are likely to share underlying risk factors and policyholder behaviors.
- Similarity in underlying assets: datasets simulated under different but related asset models can provide diverse information that enrich the shared representations. Datasets simulated under extreme market conditions can help the shared layers learn to hedge against tail risks.
- **Temporal Dynamics:** VA contracts with comparable maturity and rebalance frequency can help the shared layers learn temporal dependencies consistent across tasks.

Suppose we aim to develop an LSTM metamodel for a GMWB contract under a stochastic volatility asset model, but we have limited data for this simulation scheme. For multitask learning, we can select related nested simulation simulation procedures with relatively abundant data, such as:

- GMMB contracts under a stochastic volatility asset model.
- GMWB contracts under a Black-Scholes asset model.
- GMWB contracts under a stochastic volatility model with different simulation parameters.

These tasks need to share similarities in contract features and market dynamics so that the shared LSTM layers can learn relevant temporal patterns applicable to the target task. Otherwise, the shared layers may not be able to learn generalizable representations of temporal dependencies, and the multi-task learning framework may lead to negative transfer. Negative transfer refers to the phenomenon where the performance of the target task is worse than that of training directly on the target task without TL. It occurs when the tasks are not related or an improper TL technique is used. Positive transfer, on the other hand, refers to the phenomenon where the performance of the target task is better than that of training directly on the target task without TL. Training a multi-task LSTM metamodel benefits from having more training data, and the shared layers can learn more generalizable representations of temporal dependencies. Furthermore, if a new but similar task is introduced, fine-tuning, layer freezing, and multi-task learning can be combined to leverage the pre-trained model effectively. More specifically, the shared layers can be frozen, and the task-specific layers for a similar task in the training set can be fine-tuned on the new task. This shared knowledge helps the model generalize better on a target task, as the LSTM layers have been exposed to a wider variety of patterns and dynamics.

#### 4.2.4 Rapid Adaptation of LSTM Metamodels

In this section, we propose a TL framework for rapid adaptation of LSTM metamodels in dynamic hedging of VAs. The goal is to leverage the knowledge acquired during training on a large dataset of VA simulations to improve the learning performance on a smaller dataset of new VA contracts.

**Algorithm 7** Transfer Learning Framework for LSTM Metamodels: Combining Finetuning, Layer Freezing, and Multi-task Learning

- 1: **Input:** Set of K tasks  $\{\mathcal{T}_k\}_{k=1}^K$ , with datasets  $\mathcal{D}_k = \{(X_k^{(i)}, L_k^{(i)})\}_{i=1}^{M_k}$  for each task k, target dataset  $\mathcal{D}_{Ta}$ , learning rate  $\alpha_{So}$  and  $\alpha_{Ta}$ , shared parameters  $\theta_0$ , task-specific parameters  $\theta_k$  for each task k.
- 2: **Multi-task learning:** define shared LSTM layers  $\theta_0$  and task-specific fully connected layers  $\theta_k$  for each task k. The LSTM layers are shared across all tasks  $\{\mathcal{T}_k\}_{k=1}^K$ .
- 3: Train the multi-task LSTM metamodel on all K tasks simultaneously:

$$\min_{\theta_0, \{\theta_k\}_{k=1}^K} \sum_{k=1}^K \frac{1}{M_k} \sum_{i=1}^{M_k} \left( f_i(X_k^{(i)}; \theta_0, \theta_k) - L_k^{(i)} \right)^2 \tag{4.7}$$

- 4: For a related task of hedging a new VA contract, combine fine-tuning and layer freezing:
- 5: Layer freezing: once the multi-task metamodel is trained, freeze the parameters of the shared LSTM layers  $\theta_0$ .
- 6: **Fine-tuning:** based on task similarity, initialize the target task model parameters  $\theta_{\text{Ta}}$  using parameters  $\theta_k$  from task k.

$$\theta_{\mathrm{Ta}} \leftarrow \theta_k$$

7: Fine-tune only  $\theta_{Ta}$  on the new target dataset  $\mathcal{D}_{Ta}$  using a smaller learning rate  $\alpha_{Ta}$ :

$$\min_{\theta_{\text{Ta}}} \frac{1}{M_{\text{Ta}}} \sum_{i=1}^{M_{\text{Ta}}} \left( f_{\text{Ta}}(X_{\text{Ta}}^{(i)}; \theta_0, \theta_{\text{Ta}}) - L_{\text{Ta}}^{(i)} \right)^2$$
(4.8)

8: **Output:** Final adapted LSTM metamodel  $f_{\text{Ta}}(\cdot; \theta_0, \theta_{\text{Ta}})$  for the new VA contracts in the target task.

Algorithm 7 combines the strengths of Multi-task learning, layer freezing, and fine-tuning to accelerate the training of LSTM metamodels.

• Multi-task learning allows the shared LSTM layers to pool information across

related tasks, learning generalizable representations of temporal dependencies.

- Layer freezing ensures that shared features learned from source tasks are retained when adapting to a new task, reducing the risk of overfitting to the target simulation data.
- **Fine-tuning** enables the task-specific layers to adapt quickly to the new task with minimal simulation cost, leveraging the pre-trained shared layers.

The combination of these techniques aims to significantly reduce the computational cost of training LSTM metamodels for a new but related VA contract.

# 4.3 Numerical Experiments

In this section, we evaluate the performance of the TL framework for rapid adaptation of LSTM metamodels in dynamic hedging of VAs. The low noise dataset generated by the standard nested simulation procedure in Section 3.5 is used to train the source LSTM metamodels. We consider TL to two types of target tasks:

- the same contract but different underlying asset model, and
- different contracts but the same asset model.

Similar to Section 3.5, we use the standard nested simulation procedure to generate the dataset for the source and target tasks. Serveral VA datasets generated with the standard nested simulation procedures under geometric Brownian motion (GBM) and regime-switching GBM (RS-GBM) asset models.

Contract	Asset Model	Lapse	$M_{\mathbf{So}}$	$M_{\mathbf{Ta}}$
GMMB	GBM	No lapse	50,000	N/A
GMMB	RS-GBM	No lapse	50,000	2000
GMMB	RS-GBM	Static lapse	50,000	2000
GMMB	RS-GBM	Dynamic lapse	50,000	2000
GMWB	RS-GBM	Dynamic lapse	N/A	2000

Table 4.1: VA Contracts for Transfer Learning Experiments

Table 4.1 lists the VA contracts used in the TL experiments. These contracts include GMMB and GMWB contracts with no lapse, static lapse, and dynamic lapse features. Before transferred to the target task, a LSTM metamodel is trained on a source dataset with  $M_{\rm So}=50{,}000$  samples generated with  $N_{\rm So}=100$  inner replications<sup>1</sup>. The pre-trained LSTM metamodel is then adapted to a target task that is different from the source task. The training dataset for the target task has  $M_{\rm Ta}=2000$  samples generated with  $N_{\rm Ta}=100$  inner replications. During the training on both the source and target tasks, 10% of the data is split into a validation dataset to monitor the training process and prevent overfitting using early stopping. The complexity of the simulation schemes increases from the first task to the last task, and the LSTM metamodels need to adapt to the new conditions with limited training data on the target tasks.

All LSTM metamodels are evaluated based on their training history graphs for the target task, which plot training and validation MSE against the number of training epochs. The training history graphs visualize the learning curves of the LSTM metamodels, which provide insights into stability and convergence behavior the metamodels. We measure the generalization performance of the LSTM metamodels using the true MSE, which quantifies the accuracy of the LSTM metamodels in approximating the true inner simulation model of VA contracts. The true MSE is computed by comparing the metamodel predictions with the true contract losses, which are approximated by the standard nested simulation procedure with 100,000 inner replications. Hyperparameters and network architectures for the LSTM metamodels are kept the same as the LSTM metamodel in Section 3.5. Due to limited computational resources, macro replications are not feasible for the TL experiments.

## 4.3.1 Learning Lapse Features

We first examine the performance of fine-tuning and layer freezing in adapting LSTM metamodels to new VA contracts with lapse features. Figure 4.1 compares the performance of LSTM metamodels on the target tasks with and without TL. Learning histories of the LSTM metamodels is shown for the target task of metamodeling GMMB contract losses on the RS-GBM asset model with static lapse. The source task is hedging a GMMB contract on RS-GBM with no lapse, and the target task is hedging the same GMMB contract but with static lapse. The simulation data for the target task is generated with the same nested simulation simulation procedure as the source task, except for the lapse feature. Figure 4.1a

<sup>&</sup>lt;sup>1</sup>The GMMB contract on the GBM asset model is an exception, where scenario-wise contract losses can be computed analytically.

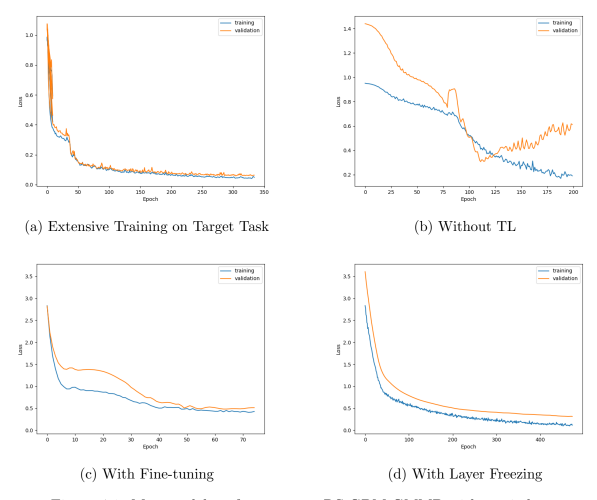


Figure 4.1: Metamodel performance on RS-GBM GMMB with static lapse

demonstrates the performance of the LSTM metamodel trained extensively only on the target task using  $M_{\rm TA}=50{,}000$  samples, which serves as a benchmark for this study. The metamodel achieves a low validation MSE due to the availability of a large dataset. In contrast, Figure 4.1b presents the results of training directly on the target task without pre-training on the source task. With only  $M_{\rm TA}=2{,}000$  samples, the LSTM metamodel struggles to learn the temporal dependencies and patterns in the target task. It leads to highly unstable training dynamics with substantial fluctuations in the validation MSE. Learning without knowledge transfer leads poor generalization and extreme overfitting. Often, the training data is limited for a new VA contract, and such instability is particularly

problematic for quick adaptation of LSTM metamodels. TL techniques like fine-tuning and layer freezing can help mitigate these challenges. Figure 4.1c shows the results of finetuning a pretrained metamodel on RS-GBM with no lapse. Fine-tuning offers a noticeable improvement over training without TL by reducing the instability in the validation MSE. However, despite this improvement, fine-tuning may not fully mitigate the challenges of training with a small dataset. No negative trasfer is observed, but the fine-tuned metamodel struggles to achieve a low validation MSE. Lowering the learning rate does not help the fine-tuned metamodel converge, and the validation MSE remains high after increasing the number of training epochs. It indicates that fine-tuning may not be sufficient for limited training data. Figure 4.1d presents the performance of layer freezing on the same target task. Layer freezing offers a more stable training process compared to fine-tuning, with a lower validation error and reduced fluctuations during its training. The LSTM layers are critical for capturing general feature representations that are transferable across tasks, and freezing these layers helps prevent overfitting to the target task. In addition, the layer freezing approach tunes fewer neural network parameters than crude fine-tuning. It allows the transferred metamodel to only focus on learning the lapse features without excessively adjusting the general temporal representations learned from the source task for the target task. This reduction in trainable parameters also accelerates the convergence, and it leads to a more stable and efficient training on the target task.

## 4.3.2 Transfer to VAs with Dynamic Lapse

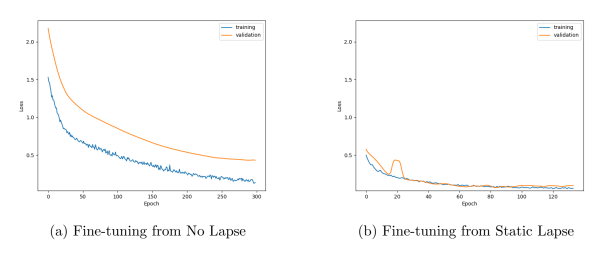


Figure 4.2: Fine-tuned Metamodel performance on RS-GBM GMMB with dynamic lapse

We further investigate the performance of fine-tuning on the target task of metamodeling GMMB contract losses on RS-GBM with dynamic lapse. The source tasks used for pretraining are GMMB contracts on RS-GBM with no lapse and static lapse, respectively. Figure 4.2 displays the learning curves of the LSTM metamodels fine-tuned from these two distinct source tasks. We observe that the performance of the fine-tuned metamodel is highly dependent on the similarity between the source and target tasks. Fine-tuning from a source task with static lapse results in faster convergence and lower validation error. The metamodel trained on the GMMB with static lapse captures some features that are beneficial for the GMMB with dynamic lapse, which leads to a more stable training process. In Figure 4.2a, the metamodel needs to learn the effect of

- whether lapse is present, and
- whether the lapse is dynamic.

This makes learning more challenging.

The observed improvement in transferability when fine-tuning from a static lapse source task highlights the importance of selecting appropriate source tasks in TL. When the source task diverges significantly from the target task, the transferred metamodel may struggle to adapt to the new conditions given the limited amount of of training data.

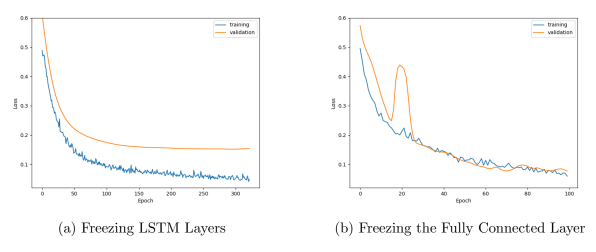


Figure 4.3: Layer Freezing on RS-GBM GMMB with dynamic lapse

Figure 4.3 continues the investigation of layer freezing in the task of metamodeling GMMB contract losses on RS-GBM with dynamic lapse. When transferring knowledge

from the GMMB model with static lapse, the primary adaptation for the target task involves learning the impact of dynamic lapse. Freezing the LSTM layers and fine-tuning only the fully connected layer results in a higher validation error, which suggests a tendency of overfitting. This indicates that the fully connected layer struggles to adapt to the changes in the temporal dynamics introduced by dynamic lapse, which can be viewed as another source of randomness in the time series. In contrast, freezing the fully connected layer and fine-tuning the LSTM layers leads to a lower validation error and better generalization. This can be attributed to the fact that the LSTM layers are responsible for capturing the temporal dependencies associated with dynamic lapse, and the fully connected layer predicts the contract losses based on these learned features.

This experiment emphasizes the importance of choosing which layers to freeze based on the nature of the source and target tasks. In the case of learning dynamic lapse features, freezing the LSTM layers is not beneficial as they need to adapt to the new temporal patterns.

Lapse Type	Extensive	Fine-tuning	Layer Freezing	Without TL
No Lapse	N/A	0.4894	0.3361	N/A
Static Lapse	N/A	0.0794	0.0763	N/A
Dynamic Lapse	0.0587	N/A	N/A	0.2950

Table 4.2: Comparison of different TL methods on GMMB contracts

Table 4.2 summarizes the true MSEs of the LSTM metamodels trained using different TL methods across various source tasks. The calculations are based on the metamodel predictions and the true contract losses approximated with 100,000 inner replications. The first two rows show the performance of transferring knowledge from GMMB contracts with no lapse and static lapse to the target task of GMMB with dynamic lapse. The last row shows the performance of training without TL on the target task. The results demonstrate the effectiveness of fine-tuning and layer freezing in transferring knowledge from a related source tasks to the target task. When transferring from a source task with static lapse to a target task with dynamic lapse, the MSEs achieved are 0.0794 for fine-tuning and 0.0763 for layer freezing. The benchmark MSE of 0.0587, obtained from extensive training on the GMMB dynamic lapse task with much more samples. While TL from static lapse GMMB do not reach this level of accuracy due to the limited data in the target task, they significantly outperform training without TL. This indicates that the models pre-trained on a static lapse setting are effective in capturing relevant features that are transferable to the dynamic lapse scenario.

However, when the source task is less similar to the target task, the benefits of TL are less pronounced. The MSEs in this case are higher, with fine-tuning resulting in 0.4894 and layer freezing achieving 0.3361. This suggests that the divergence between the source and target tasks can lead to negative transfer. These results highlight the importance of selecting source tasks that share significant similarities with the target task to maximize the effectiveness of TL. When the source and target contracts are substantially different, the pre-trained metamodels may struggle to adapt to the new conditions, as they may not have learned features relevant to the target task. There is no negative transfer observed, but the knowledge transfer is limited.

### 4.3.3 Transfer Knowledge to other Contract Types

When transferring knowledge from one VA contract type to another, the LSTM metamodel needs to adapt to different contract features and time series dynamics. We consider the task of metamodeling GMWB contract losses on RS-GBM asset model with dynamic lapse, with the source tasks being a GMMB contract on RS-GBM also with dynamic lapse. Figure 4.4 illustrates the learning history of transferring pre-trained LSTM metamodels from the GMMB contracts to the GMWB contracts.

Figure 4.4a presents the performance of an extensively trained LSTM metamodel on the target task using 50,000 samples, serving as a benchmark of best metamodel performance. The extensively trained metamodel achieves a low validation error and demonstrates stable convergence due to having enough training samples. In contrast, Figure 4.4b shows the results of training the metamodel directly on the target task with 2000 training samples. Fewer training samples and no prior knowledge leads to unstable training dynamics. The validation MSE fluctuates significantly.

When applying fine-tuning from the GMMB source task (Figure 4.4c), the metamodel exhibits improved performance compared to training without TL. Fine-tuning results in a lower validation error and more stable convergence. Despite the differences between GMMB and GMWB contracts, there is still valuable information that can be transferred. Both the LSTM layers and the fully connected layers capture general temporal patterns and feature representations that are beneficial for the target task. Fine-tuning allows the metamodel to adjust all its neural network layers. It allows better adaptation to the complexities introduced by the GMWB contract.

However, when employing layer freezing (Figure 4.4d), the metamodel's performance deteriorates. Negative transfer is observed. Freezing some layers trained on the GMMB contracts does not allow the metamodel to sufficiently adapt to the complexities of the

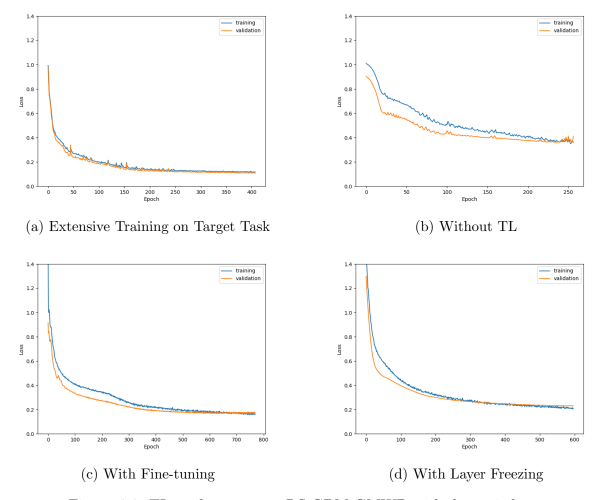


Figure 4.4: TL performance on RS-GBM GMWB with dynamic lapse

GMWB contracts. The GMWB contracts are inherently more complex than GMMB contracts due to the guaranteed withdrawal benefits at each time step, and the contract features are significantly different. The complexity introduce significant changes in the time series dynamics that the metamodel needs to capture. Freezing the LSTM layers or the fully connected layer hinders the metamodel's ability to learn these new patterns, and it leads to poor generalization and unstable error curves.

Table 4.3 summarizes the true MSEs of the LSTM metamodels on the GMWB contracts. Training MSE and true MSE are computed using the training labels and the true contract losses approximated with 100,000 inner replications, respectively. This is consis-

Model	Training Samples	Training MSE	True MSE
Without TL	2000	0.3588	0.4188
Fine Tuning	2000	0.1690	0.1780
Layer Freezing	2000	0.1828	0.2295
Extensive Training	50,000	0.0853	0.0726

Table 4.3: Comparison of different TL methods to GMWB contracts

tent with the terminology used in Chapter 3. The TL methods are compared to training from scratch. The suboptimal performance of layer freezing in this context indicates that the difference between the source and target tasks is too substantial for this method to be effective. While layer freezing can be advantageous when the source and target tasks are closely related, it may hinder performance when the tasks diverge significantly. In such circumstances, fine-tuning provides a better approach by allowing the metamodel to leverage transferable knowledge while adapting to the new task's specific requirements. Fine-tuning enables both the LSTM and fully connected layers to update their weights, capturing the complex dynamics of the GMWB contracts more effectively. This is particularly beneficial when developing metamodels for complex VA contracts with limited simulation data. For instance, transferring information from GMMB contracts can still be valuable when modeling GMWB contracts. Both contracts share some common contract features and temporal patterns, which allows a pre-trained LSTM metamodel to capture generalizable features that provide a solid foundation for the target task.

Figure 4.5 illustrates the performance of the two-stage procedure with TL for GMWB contracts with dynamic lapse in predicting the tail scenarios and the 95%-CVaR. The graph provides a visual representation of how different TL approaches compare to training from scratch and the standard nested simulation procedure. Fine-tuning consistently outperforms both layer freezing and training from scratch across different simulation budgets, particularly in predicting tail scenarios and estimating the 95%-CVaR. This finding is consistent with the training history and MSEs in Figure 4.4 and Table 4.3, respectively. It's important to note that while these results are promising, they also highlight the complexity of modeling GMWB contracts with dynamic lapse. The fact that fine-tuning outperforms layer freezing suggests that there are significant differences in the tail behavior of GMMB and GMWB contracts, particularly when dynamic lapse is considered. These findings highlight the need for careful model selection and validation when applying TL techniques to different VA products. When the source and target tasks are substantially different, all layers need to be updated. Freezing some layers may lead to negative transfer.

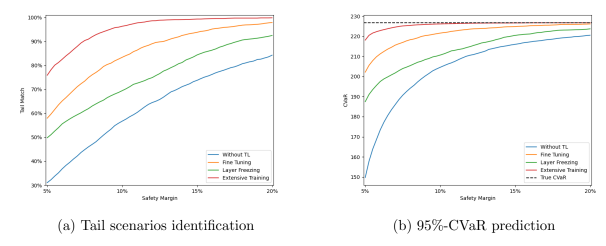


Figure 4.5: TL performance on RS-GBM GMWB with dynamic lapse

### 4.3.4 Multi-task Learning

Multi-task learning enables the LSTM metamodels to learn shared representations across related VA contracts. In this section, we examine the performance of multitask learning applied to two types of VAs, GMMB and GMWB with dynamic lapse rates. The simulation datasets contrain 2,000 samples for each contract type, and the LSTM metamodels are trained using multi-task learning. In our experiments, Algorithm 6 is used to train the LSTM metamodels simultaneously to minimize the multi-task MSE loss function in Equation 4.6. We use individual task training as a baseline for comparison, where the LSTM metamodels are trained separately on the GMMB and GMWB contracts. The objective is to assess how multitask learning can improve the training efficiency and performance of both products compared to individual task training.



Figure 4.6: Multi-task Learning Framework for VA Contracts

Figure 4.6 illustrates our multi-task learning framework for GMMB and GMWB contracts with dynamic lapse rates. The outer simulation paths for both contracts are the

same, which are generated from the same nested simulation procedure with 100 inner replications. The LSTM metamodels are trained simultaneously on both contracts. The LSTM layers are shared, and each contract has its own separate fully connected layers for contract loss predictions.

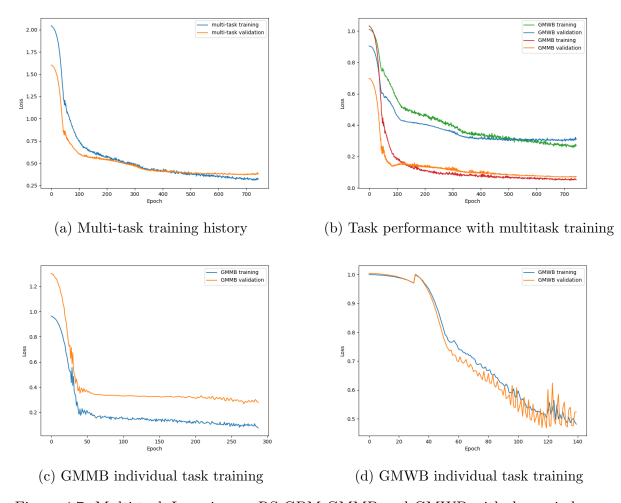


Figure 4.7: Multi-task Learning on RS-GBM GMMB and GMWB with dynamic lapse

Figure 4.7 compares the learning curves for the training of GMMB and GMWB, with and without multitask learning. The comparison between multitask learning (Figure 4.7a and 4.7b) and individual task training ((Figure 4.7c and 4.7d)) demonstrates the benefit of multi-task learning for both products. In the case of GMMB, multitask learning allows the model to achieve faster convergence while reducing overfitting. For GMWB, the multi-

task learning framework helps to stablize the training process. The shared LSTM layers in the multitask model are able to capture common temporal patterns and dynamics, which benefits both GMMB and GMWB when training samples is scarce. Similar to pooling in Chapter 2 and 3 but on a higher level, multi-task learning enables the LSTM metamodels to leverage shared representations across related VA contracts.

#### 4.3.5

#### 4.4 Conclusion

In this chapter, we introduced a transfer learning framework to accelerate the training of LSTM metamodels for dynamic hedging of variable annuity contracts. Traditional nested simulation procedures for VA risk management are computationally intensive. LSTM metamodels offer a data-driven approach to approximate the true contract losses, which can significantly reduce the computational cost of hedging a single VA contract. However, training LSTM metamodels on new VA contracts can be challenging due to the limited availability of simulation data. Our proposed framework leverages pre-trained LSTM networks and transfer learning techniques to adapt quickly metamodels to new but related VA contracts with minimal additional simulation cost. Fine-tuning a pre-trained LSTM metamodel on a new target task with limited data significantly improved training stability and predictive accuracy compared to training from scratch. Layer freezing further enhanced performance by retaining transferable temporal representations learned from the source task. However, the success of these methods depends on the similarity between the source and target tasks. When the tasks were closely related, freezing some neural network layers yields substantial benefits. Conversely, when the source and target tasks diverged significantly, fine-tuning the entire network is more effective.

Multi-task learning offers a robust framework for transferring knowledge across related simulation schemes in LSTM metamodeling for nested simulations. By sharing representations through the LSTM layers, the model can learn more generalizable features that are beneficial across different VA contracts. The multi-task approach is particularly beneficial when training data for individual tasks is scarce, as it effectively pools information across tasks to enhance learning efficiency and predictive performance. We have extended the idea of pooling in Chapter 2 and 3 to a higher level. Furthermore, a metamodel trained with multi-task learning can serve as a pre-trained model. It is adaptable to various VA contracts and asset models, and it is a versatile tool for practical applications in dynamic hedging of VAs.

The integration of transfer learning in LSTM metamodeling represents a significant advancement in robust risk management associated with variable annuity contracts. By effectively leveraging existing knowledge, financial institutions can maintain accurate and responsive risk management practices in a rapidly changing market environment. The methodologies presented in this chapter attempt to make a significant contribution to the broader applications of transfer learning in financial modeling and risk assessment.

# Chapter 5

# Conclusion

In this thesis, we have explored the application of nested simulation procedures in the risk management of financial derivatives and insurance products, with a particular focus on variable annuity contracts featuring Guaranteed Minimum Maturity Benefits and Guaranteed Minimum Withdrawal Benefits. Recognizing the computational challenges inherent in the standard nested simulation procedure, especially when estimating tail risk measures for complex financial products, we proposed an innovative approach that integrates machine learning-based metamodels into the nested simulation framework. We have shown that the nested simulation framework can be used to efficiently estimate the risk measures of complex financial products, such as variable annuities, by combining machine learning-based metamodels with nested simulations.

Our primary contribution lies in the development and implementation of long short-term memory networks as metamodels for the inner simulations within the nested simulation procedure. By employing LSTMs, we effectively approximated the contract losses associated with dynamic hedging strategies under various market scenarios. This approach significantly reduced computational complexity while maintaining high levels of accuracy in risk estimation. The LSTM metamodels captured the temporal dependencies and non-linear relationships present in financial time series data, enabling efficient estimation of tail risk measures.

Through extensive numerical experiments and sensitivity analysis, we demonstrated that our LSTM-based nested simulation procedure provides accurate and computationally efficient estimates of the tail risk associated with VA contracts. The results showed that the metamodels could effectively replace the computationally intensive inner simulations, reducing the overall runtime without compromising the precision of risk assessments. This

efficiency gain is particularly valuable for insurers and financial institutions that require timely and accurate risk evaluations to meet regulatory requirements and make informed risk management decisions.

Additionally, we investigated the feasibility of our approach to changing market conditions and different VA contract features. By incorporating transfer learning techniques, we enabled rapid adaptation of the LSTM metamodels to new contracts and evolving market dynamics. This aspect of our research highlights the potential for machine learning models to not only improve computational efficiency but also enhance the flexibility and responsiveness of risk management practices in the face of market volatility.

The thesis is an attempt to bridge the gap between traditional simulation schemes and modern machine learning techniques in the context of financial risk management. By bridging advanced machine learning techniques with financial risk management practices, this thesis contributes to the development of more efficient, adaptable, and robust methodologies that are essential in the rapidly evolving landscape of quantitative finance and actuarial science.

# Chapter 6

# Future Work: Deep Hedging Variable Annuities with Transfer Learning

In the evolving landscape of financial markets, insurance products such as VAs have gained significant interest due to their ability to provide both investment growth and guaranteed benefits. Managing the risks associated with these products, especially in volatile market conditions, is a complex task that demands sophisticated financial modeling techniques. Reinforcement Learning (RL) has emerged as a powerful tool for addressing such complexities. With the recent advancement in deep learning, DNN architectures have been successfully applied to RL problems, leading to the development of deep reinforcement learning (DRL) algorithms. These algorithms have demonstrated remarkable success decision-making tasks that must be optimized over time. In the context of managing VAs, DRL has shown promise in optimizing hedging strategies where long-term financial and policyholder decisions interact dynamically with changing market conditions.

Reinforcement learning is a branch of machine learning in which an agent learns to make sequential decisions by interacting with its environment. Through trial and error, the agent receives feedback in the form of rewards or penalties, allowing it to learn an optimal policy for achieving long-term goals. In financial and actuarial contexts, RL has been applied to portfolio optimization, asset allocation, and risk management. In a dynamic hedging problem, the agent is referred to as the decision-maker that tries to minimize financial risks over time horizon by adjusting the hedging portfolio based on the current environment, which is the current state of the financial markets and actuarial assumptions. By intergrating deep neural network, DRL extends this framework by enabling the agent to handle high-dimensional and complex environments. DRL has shown particular promise in finance because of its ability to process vast amounts of financial data and discover intricate

patterns in market dynamics. Despite their success, DRL faces practical limitations when applied to real-world financial markets. Its most significant drawback is the requirement for extensive data and computational resources. Training a DRL agent from scratch requires a large dataset along with considerable computational time for exploration and learning. This is especially problematic for variable annuities, which are long-term insurance products that are sensitive to market fluctuations and underlying actuarial assumptions. Given that financial markets are highly dynamic, frequent retraining to account for new market regimes becomes necessary, which can lead to prohibitively high costs and computational inefficiency. Moreover, using historical data leads to overfitting to the past. An overfitting model can lead to suboptimal hedging strategies that perform poorly in changing market conditions.

Transfer learning (TL) offers a compelling solution to the challenges posed by existing DRL approaches. Similar to the TL framework proposed in Chapter 4, TL can also be applied to DRL to accelerate the training. In the context of financial modeling, instead of training an RL agent from scratch every time underlying assumptions change, the model can utilize the knowledge gained from past market environments to accelerate the learning process under the new environment. TL in financial markets is particularly advantageous for addressing the data scarcity problem. By transferring existing knowledge from similar environments, TL improves model performance in situations where historical data is limited or noisy. This allows models using TL to be both more accurate and computationally efficient. Additionally, TL enhances the robustness of models by allowing continuous learning. Instead of retraining a deep RL model from scratch when market conditions shift, TL enables the reuse of pre-trained models, facilitating faster adaptation to new environments while avoiding overfitting to past conditions. In the risk management of variable annuities, transfer learning enables quicker adaptation to changing financial and actuarial assumptions by reusing information obtained from another set of related assumptions, such as the management of similar financial instruments or different market regimes. A DRL agent equipped with transfer learning can adjust its strategies in response to new market data, making the model more adaptive and responsive without the need for costly retraining. More specifically, a DRL agent trained to manage variable annuities for a GMMB contract can apply its learned strategies to a GMWB contract with the same financial assumptions. A DRL agent trained for a VA product on a Black-Scholes asset model can transfer its knowledge to the training of a similar product on a stochastic volatility model. This approach significantly reduces the computational costs associated with the training of new VA products with different assumptions and improves the generalization of the model to unseen market conditions. By integrating transfer learning with existing DRL frameworks, we can eliminate the inefficiencies of training models from scratch, provide faster adaptability to new market conditions, and enhance the scalability and computational efficiency of financial models. This approach offers a more robust and practical solution for managing variable annuities in dynamic and volatile financial markets.

The rest of the chapter is organized as follows:

# 6.1 Markov Decision Process for Hedging VAs

A Markov Decision Process (MDP) provides a mathematical framework for modeling decision-making in stochastic environments over time. By formulating the VA hedging problem as an MDP, we can utilize RLalgorithms to learn optimal hedging policies directly from data. AnMDP is defined by the tuple  $\mathcal{M} = (\mathcal{S}, \mathcal{A}, \mathcal{P}, \mathcal{R}, \gamma)$ , where:

- S is a finite set of states.
- $\mathcal{A}$  is a finite set of actions.
- $\mathcal{P}$  is a state transition probability distribution, where  $\mathcal{P}(s_t|s_{t-1}, a_{t-1})$  represents the probability of transitioning from state  $s_t$  to state  $s_{t-1}$  due to action  $a_{t-1}$ .
- $\mathcal{R}$  is a reward distribution,  $\mathcal{R}(s, a, s')$  is the reward received after transitioning from state s to state s' due to action a.
- $\gamma$  is a discount factor,  $\gamma \in [0,1]$ , which models the present value of future rewards.

The objective in an MDP is to find a policy,  $\pi: \mathcal{S} \to \mathbb{P}(\mathcal{A})$ , that maximizes the cumulative reward over time. A MDP provides a mathematical framework for solving sequential decision-making tasks in an enviornment where outcomes are partly random and partly under the control of a hedging agent. For hedging VAs, the asset dynamics and the mortality model are not controlled by the agent, but the agent controls the cumulative reward by deciding on the hedging weights. In this section, we reformulate the hedging enviornment of VAs from Section 3.2.2 as an MDP and define the key components of the MDP.

## 6.1.1 Hedging Environment of Variable Annuities

Consider a generic VA contract with maturity T > 0 periods, e.g., T = 240 months. Then the contract expires at  $T' = \min\{T, \tau\}$ , i.e., the earlier of the contract maturity and the death of the policyholder. Let  $S_t$ ,  $F_t$ , and  $G_t$  be the indexed stock price, the subaccount value and the guarantee value, respectively, at time t = 1, 2, ..., T. Evolution of the subaccount value and the guarantee value of a VA contract affect the contract payout. For clarity, we use  $F_t$  and  $F_{t_+}$  to denote the sub-account value just before and just after the withdrawal at time t, if any. Let  $\eta_g$  be the gross rate of management fee that is deducted from the fund value at each period and let  $\eta_n < \eta_g$  be the net rate of management fee income to the insurer. The difference between the gross management fee and the net management fee income represents the incurred investment expenses.

At the inception of the contract, i.e., t=0, we assume that the whole premium is invested in the stock index and the guarantee base is set to the sub-account value:

$$S_0 = F_0 = G_0$$
.

At each time t = 1, ..., T, the following events take place in the following order:

1. The dynamic lapse rate  $q_t$  is applied to the contract, i.e., q of the policyholders leave the contract at time t.

$$q_t = q_t^B \cdot \text{clip}(M_q(\frac{G_{t-1}}{F_{t-1}} - D_q), L_q, U_q), \tag{6.1}$$

where  $q_t^B$  is the base lapse rate,  $\operatorname{clip}(x, a, b)$  is a function that clips the value of x to be within the range [a, b], and  $M_q$ ,  $D_q$ ,  $L_q$ , and  $U_q$  are the model parameters taken from National Association of Insurance Commissioners' (NAIC) Valuation Manual 21 (NAIC, 2021).

2. The sub-account value changes according to the growth of the underlying stock and the (gross) management fee is deducted. That is,

$$F_t = F_{(t-1)_+} \cdot \frac{S_t}{S_{t-1}} \cdot (1 - \eta_g) \cdot (1 - q_t), \tag{6.2}$$

where  $(x)^+ = \max\{x, 0\}$  and  $F_{(t-1)_+}$  will be defined later. The insurer's income at time t is the net management fee, i.e.,  $F_t\eta_n$ .

3. The guarantee value ratchets up (ratcheting is a common feature in GMWB) if the sub-account value exceeds the previous guarantee value, i.e.,

$$G_t = \max\{G_{t-1} \cdot (1 - q_t), F_t\}. \tag{6.3}$$

A GMMB can be modeled with  $G_t = G_{t-1} \cdot (1 - q_t)$ .

4. The withdrawal is made (for GMWB) and is deducted from the sub-account value, i.e.,

$$F_{t_{+}} = (F_{t} - I_{t})^{+}, (6.4)$$

where  $I_t = \xi G_t$ . A GMMB can be modeled with  $\xi = 0$ .

Consider a VA contract whose delta hedge portfolio at any time t, t = 0, 1, ..., T - 1, consists of  $\Delta_t$  units in the underlying stock and  $B_t$  amount of a risk-free zero-coupon bond maturing at time T. The value of the hedge portfolio at time (t - 1) is:

$$H_{t-1} = \Delta_{t-1} S_{t-1} + B_{t-1},$$

where  $S_t$  is the underlying stock price and any time t > 0. This hedge portfolio is brought forward to the next rebalancing time t, when its value becomes:

$$H_t^{bf} = \Delta_{t-1} S_t + B_{t-1} e^r$$
.

Therefore, the time t hedging error, i.e., the cash flow incurred by the insurer due to rebalancing at time t, is

$$HE_t = H_t - H_t^{bf}, \quad t = 1, \dots, T - 1.$$
 (6.5)

The P&L of the VA contract includes the cost of the initial hedge  $(H_0)$ , the hedging errors (3.3), the unwinding of the hedge at maturity  $(H_T^{bf})$ , and the contract payout at time  $t \in \{0, \ldots, T\}$ . Mathematically, the present value of the liability for a GMMB contract is

$$L = H_0 - e^{-rT} H_T^{bf} + \sum_{t=1}^{T-1} e^{-rt} H E_t + \sum_{t=1}^{T} e^{-rt} C_t - \sum_{t=1}^{T} e^{-rt} F_t \eta_n + e^{-rT} (G_t - F_t)^+$$

$$= e^{-rT} (G_T - F_{T^+})^+ + \sum_{t=1}^{T} e^{-rt} \left( \Delta_{t-1} (S_{t-1} - e^{-r} S_t) + C_t - F_t \eta_n \right)$$
(6.6)

where the equality holds by a rearrangement of terms and a telescopic sum simplification of  $e^{-rt}B_t$ ,  $t=0,\ldots,T-1$ , and  $C_t:=C\cdot S_{t-1}\cdot |\Delta_t-\Delta_{t-1}|$  is the transaction cost at time

t (Gârleanu and Pedersen, 2013). Hence, the present liability consist of the hedging errors, the transaction costs, the net management fees, and the contract payouts. Similarly, the present value of the liability for a GMWB contract is

$$L = \sum_{t=1}^{T} e^{-rt} \left( \Delta_{t-1} (S_{t-1} - e^{-r} S_t) + C_t - F_t \eta_n \right)$$
 (6.7)

The GMWB payout is implicitly included in  $F_t\eta_n$ . Both Equation 6.6 and Equation 6.7 neglect the transaction cost to liquidate the hedge portfolio at maturity. The discrete-time hedging problem of VAs can be conveniently formulated as an MDP, where the states, actions, transition probabilities, policies, and rewards are defined as follows.

#### 6.1.2 State Space

The state space S represents all the information needed to characterize the hedging environment. In a VA hedging problem,  $\mathbf{s}_t$ , the information available to a hedging agent at each time t is

$$\mathbf{s}_t := (S_t, F_t, G_t, \tau, \Delta_{t-1}) \in \mathcal{S}.$$

where  $\tau$  is the remaining time to maturity and  $\Delta_{t-1}$  is the previous hedging weight. The hedging environment is partially observed, e.g., the agent does not have access to the contract specifications. It has to learn a representation of the relevant information from the observed state  $\mathbf{s}_t$ .

## 6.1.3 Action Space

The action space  $\mathcal{A}$  represents the set of actions that the agent can take at each time t. In a VA hedging problem, the action space only includes the hedging weight  $\Delta_t$ .

## 6.1.4 Policy

A policy  $\pi$  determines the best action to take in a given state, i.e.,

$$a_t \sim \pi(\cdot|\mathbf{s}_t).$$

In a VA hedging problem the policy outputs the hedging weight  $a_t = \Delta_t$  given the observed state  $\mathbf{s}_t$ . In the following sections, we will refer to  $\Delta_t$  as the policy.

#### 6.1.5 Transition Probabilities

The transition probabilities  $\mathcal{P}$  represent the probability of transitioning from one state to another from taking an action. In our setup, the transition probabilities are fully determined by Equation 6.1, Equation 6.2, Equation 6.3, and Equation 6.4. Let  $\mathcal{P}(\mathbf{s}_{t+1}|\mathbf{s}_t, a_t)$  be the probability of transitioning from state  $\mathbf{s}_t$  to state  $\mathbf{s}_{t+1}$  due to action  $a_t$ . Aside from the previous hedging weight  $\Delta_{t-1}$ , the transition probabilities are independent from the actions. Paths simulated from the transition probabilities are called trajectories (or episodes) of the MDP.

#### 6.1.6 Reward and Discount Factor

The reward function  $\mathcal{R}$  and discount factor  $\gamma$  in a typical RL problem are defined as follows:

$$\mathcal{R} = \sum_{t=1}^{T-1} \gamma^t R_t$$

$$\gamma = e^{-r},$$
(6.8)

where  $R_t$  is the term reward received by the agent at time t, and the discount factor  $\gamma$  is conveniently inherited from the risk-free rate r. In hedging, the term reward  $R_t$  should be drived from Equation 6.6 and Equation 6.7 to represent the performance of the hedging policy, i.e., how closely the hedging portfolio tracks the liability. For GMWB, we propose a term reward formulation that encourages the agent to maximize the expected reward with low hedging errors and transaction costs:

$$R_t = F_{t-1}\eta_n - \Delta_{t-1}|S_{t-1} - e^{-r}S_t| - C_{t-1}(\Delta_{t-1}), \quad t \in \{1, \dots, T\}$$
(6.9)

The reward function of GMWB directly links to the time-t loss of a GMWB contract as in Equation 6.7. The GMMB loss consists of an additional payoff at maturity T, thus its term reward is formulated as:

$$R_{t} = \begin{cases} F_{t-1}\eta_{n} - \Delta_{t-1}|S_{t-1} - e^{-r}S_{t}| - C_{t-1}(\Delta_{t-1}), & t \in \{1, \dots, T-1\} \\ F_{t-1}\eta_{n} - \Delta_{t-1}|S_{t-1} - e^{-r}S_{t}| - C_{t-1}(\Delta_{t-1}) + (G_{T} - F_{T+})^{+} & t = T \end{cases}$$

$$(6.10)$$

## 6.1.7 Value Function and Advantage Function

This section introduces some critical concepts in RL, i.e., the value function and the advantage function. They are essential for understanding the performance of the hedging policy and the learning process of the RL agent.

A value function quantifies the expected cumulative reward over time. The state value function,  $V^{\pi}(s_t)$ , is defined as the expected cumulative reward starting from state  $s_t$  and following policy  $\pi$  thereafter:

$$V^{\pi}(s_t) = \mathbb{E}\left[\sum_{t=0}^{\infty} \gamma^t R_t | \pi, s_t\right], \tag{6.11}$$

where  $R_t$  is the reward received by the agent after taking action  $a_t$  in state  $s_t$  and transitioning to state  $s_{t+1}$ , and the expectation is taken over the possible sequences of states and rewards that follow from the policy  $\pi$ . The state-action value function,  $Q^{\pi}(s_t, a_t)$ , is defined as the expected cumulative reward starting from state  $s_t$ , taking action  $a_t$ , and following policy  $\pi$  thereafter:

$$Q^{\pi}(s_t, a_t) = \mathbb{E}\left[\sum_{t=0}^{\infty} \gamma^t R_t | \pi, s_t, a_t\right], \tag{6.12}$$

Based on Equations 6.11 and 6.12, the advantage function,  $A^{\pi}(s_t, a_t)$ , is defined as the difference between the state-action value function and the state value function:

$$A^{\pi}(s_t, a_t) = Q^{\pi}(s_t, a_t) - V^{\pi}(s_t). \tag{6.13}$$

The advantage function quantifies the advantage of taking action  $a_t$  in state  $s_t$  over following the policy  $\pi$ .

## 6.2 Exising RL Algorithms and Their Limitations

## 6.2.1 Model-Based RL: From AlphaZero to Deep Hedging

Deep reinforcement learning first attacted attentions when AlphaGo (Silver et al., 2016) surpasses human in playing Go. Soon after, AlphaZero (Silver et al., 2016) emerged as a

model-based RL algorithm that improves over self-plays. It quickly extended beyond Go to other games like Chess and Shogi. In essence, the AlphaZero agent use a neural network to paramaterize the policy and the value function. The neural network takes the current state of the game as input and outputs the probability distribution of the next move and a value of the current state.

$$(\pi, v) = f_{\theta}(s), \quad L(\theta) = -\pi^{\mathsf{T}} \log(\mathbf{p}) + (z - v)^2 + c||\theta||^2,$$
 (6.14)

where  $f_{\theta}$  is the neural network with parameters  $\theta$ , s is the current state of the game,  $\pi$  is the policy, v is the value of the current state, and c is a regularization parameter.  $\mathbf{p}$  and z are the probability distribution of the next move and the outcome of the game, respectively. They are induced by a planning algorithm, i.e., Monte Carlo Tree Search (MCTS). Since it is infeasible to efficiently simulate trajectories in the game of Go, a planning algorithm is necessary to generate expected outcomes. The neural network  $f_{\theta}$  is trained with a stochastic gradient descent algorithm to minimize L. L has three components. The last term penalizes large neural network weights. Its first two terms measure the similarity between  $f_{\theta}(s)$ , the learned policy and value function, with the action and game outcome planned by the MCTS. The neural network is trained to fit the planned outcomes of the game. It stores a global solution that predicts the outcome of the game and the next move before they are actually observed. This is known as model-based RL (Sutton and Barto, 2018). Using a model allows for planning, i.e., decision-making take place using future situations that have not been experienced by the agent.

In a typical hedging problem, the transition probabilities are independent from the hedging weights, i.e., the actions. This simplifies the planning as complete trajectories can be simulated without the need of a planning algorithm. In contrast to AlphaZero, deep hedging (Buehler et al., 2019) uses monte carlo simulations to generate sample paths of asset dynamics. These simulations are conducted based on predefined stochastic asset models. The parameters and structure of these models are specified in advance, dictating the behavior of the simulated asset prices.

A common choice of  $\rho$  is a CVaR. In practice, the neural network  $f_{\theta}$  is trained to minimize the empirical CVaR over a batch of trajectories. In other words, planning is done through  $\rho$ , which measures the performance of a hedging policy over a batch of complete trajectories. The neural network, representing the hedging policy, is trained using the data generated from these simulations. The objective is to learn a strategy that minimizes a tail risk measure of the total present liability. Consequently, the learned policy is inherently tailored to the characteristics of the model used in the simulations. The ability to know the future outcomes of the hedging policy before they are actually observed and the use of

a risk measure to guide the training of the neural network are the key features of model-based RL. These models describe how asset prices evolve over time, and more importantly, they are used to predict future states based on current information. While model-based approaches can be powerful when the model accurately reflects reality, they are susceptible to model risk, i.e., the risk that the chosen model is incorrect or incomplete. It can lead to suboptimal hedging strategies if the real market deviates from the model assumptions.

For hedging VAs, Xu (2020) propose a deep hedging algorithm that falls into this category of model-based RL. A cost function is defined to minimize the expected shortfall of the total present liability. Carbonneau (2021) extends the model-based approach to hedging long-term financial derivatives and compares the performance of deep hedging strategies trained with different cost functions. In deep hedging (Buehler et al., 2019) the state space of the MDP contains the previous hedging weight. A more recent paper by Imaki et al. (2021) proposes a new network architecture that removes the previous hedging weight from the state space by introducing a no-transaction band. It allows for a more efficient training of the neural network by removing the semi-recurrent problem structure.

Despite the success of model-based RL in hedging, most studies focus on hedging different types of contracts on classical asset models, e.g., Black-Scholes model and Heston model. Motivated by the seminal work by Gatheral et al. (2022), there are a many subsequent works focusing on roughening the classical stochastic volatility models. In the rough volatility models, the latent volatility process is modelled by a stochastic process that is rougher than the sample paths of diffusion process. Among these rough volatility models, notable examples are rough Heston model and the rough Bergomi model. Model-based RL struggles in a rough volatility model due to difficulties in designing a suitable cost function. Horvath et al. (2021) shows that a standard implementation of deep hedging can lead to suboptimal hedging performance in a discretized rough Bergomi model.

In contrast, a model-free approach does not require a explicit design for different asset dynamics, making it more flexible and adaptable to different market conditions and more robust to model misspecification.

## 6.2.2 Model-Free RL for Hedging

In a model-free reinforcement learning approach, the agent is a trial-and-error learner that interacts with the environment to learn the optimal policy by iteratively updating the policy and the value function based on the observed states, actions, and rewards. For hedging VAs, the agent does not have access to a planning algorithm or the underlying

stochastic asset model, and it must learn the optimal policy by directly interacting with the environment.

Model-free RL algorithms gain its popularity in robotics tasks, where transition probabilities in the environment are unknown and can be affected by the agent's actions. In the context of hedging, existing studies use transaction costs as a function of the hedging weights, which makes the transition probabilities independent on the actions. However, the trade size can affect the asset price in a real environment (Hasbrouck, 1991), which makes the model-free approach a promising direction for hedging VAs in the real market.

Boardly speaking, model-free RL can be categorized into two types: value-based RL and policy-based RL. Value-based approaches learn the value function directly from the data and derive the policy from the learned value function. The value function is learned by training a DNN to approximate the value function, which quantifies the expected cumulative reward over time. The policy is then derived from the learned value function by selecting the action that maximizes the value function at each state. (2015) is the first paper to demonstrate the effectiveness of value-based deep reinforcement learning in training agents to play Atari games. To update the value function, the DNN is trained to minimize the difference between the current value function and the target value function, which is updated based on the maximum expected reward from the previous iteration. When trained to approximate the value function, the DNN benefits from the use of experience replay buffers, which is proposed by Lin (1992) to store and sample historical transitions and uniformly sampled during training to enhance the sample efficiency and stability of the learning process. Kolm and Ritter (2019) uses a value-based deep reinforcement learning approach to hedge European options under the Black-Scholes framework.

In valued-based approaches, the value function is updated iteratively with the maximum expected reward of all possible actions, which can be computationally expensive for continuous action spaces. To overcome this difficulty, a policy-based approach estimates the value function before realization of the final payoff of the hedging portfolio, which is particularly useful for hedging long-term financial derivatives or VA contracts. Two types of policy-based approaches prevail in the literature: actor-critic and policy gradient methods.

An actor-critic method designs an actor network and a critic network to learn the value function and the policy simultaneously. The actor network learns the policy by directly outputting the action based on the current state, while the critic network learns the value of an action by approximating the expected cumulative reward over time following the actor's policy. A popular actor-critic algorithm is the Deep Deterministic Policy Gradient (DDPG) algorithm proposed by Lillicrap et al. (2015), which is implemented by Xu and Dai (2022) in hedging financial derivatives in S&P 500 and DJIA index options.

A policy gradient method uses policy gradient theorem to update the policy by performing gradient ascent with respect to the policy parameters.

#### Algorithm 8 Vanilla Policy Gradient (REINFORCE)

- 1: **Initialize** policy parameters  $\theta$  arbitrarily.
- 2: **For** iteration k = 0, 1, 2, ... do
- 3: Collect a set of  $\mathcal{D}_k$  by sampling from the enviornment with policy  $\pi_{\theta}$ .
- 4: Compute the rewards  $R_t$  for each trajectory in  $\mathcal{D}_k$  for  $t = 0, 1, \dots, T$ .
- 5: Update the policy parameters  $\theta_{k+1}$  by

$$\theta_{k+1} = \theta_k + \alpha \frac{1}{|\mathcal{D}_k|T} \sum_{\kappa \in \mathcal{D}_k} \sum_{t=0}^T \nabla_{\theta} \log \pi_{\theta}(a_t|s_t) R_t$$

#### 6: End For

Algorithm 8 shows the REINFORCE algorithm (Sutton et al., 1999), a classic policy gradient method that updates the policy by performing gradient ascent with respect to the policy parameters. When training on real-world data, deep hedging of Buehler et al. (2019) effectively becomes a model-free policy gradient methods. The policy is updated by performing gradient ascent with respect to the policy parameters to maximize the expected reward over time, where the reward function is designed to be the negative of a tail risk measure of the total present liability.

Representative algorithms include the Trust Region Policy Optimization (TRPO) algorithm proposed by Schulman et al. (2015a) and the Proximal Policy Optimization (PPO) algorithm proposed by Schulman et al. (2017). Du et al. (2020) show that the PPO algorithm is more sample efficient and stable than the value-based approaches in training agents to hedge financial derivatives. The most relevant paper to our study is Chong et al. (2023), which uses the PPO algorithm to hedge GMMB with a GMDB rider under the Black-Scholes framework. However, information of the underlying risk factors is leaked to the agent, which makes their study pseudo-model-free. To further improve the risk management of VAs under rough volatility models, we propose a model-free deep reinforcement learning approach to hedge VAs without explicit or implicit knowledge of the underlying risk factors.

## 6.3 Hedging VAs with a PPO Agent

In a hedging problem, an insurer aims to find a policy  $\pi$  that maximize the expected reward over time. In a policy-based RL approach like the PPO, both the policy and the value function are parameterized by deep neural networks. We denote the policy by  $\pi_{\theta}$  and the value function by  $V_{\phi}$ , where  $\theta$  and  $\phi$  are the parameters of the value network and the policy network, respectively. The policy is updated according to an objective function that reflects the advantage of taking action  $a_t$  in state  $s_t$  over following the policy  $\pi$ .

$$\max_{\theta} \mathbb{E}\left[\min\left\{\Lambda_t(\theta)\hat{A}^{\pi_{\theta}}(s_t, a_t), \operatorname{clip}(\Lambda_t(\theta), 1 - \epsilon, 1 + \epsilon)\hat{A}^{\pi_{\theta}}(s_t, a_t)\right\}\right]$$
(6.15)

$$\Lambda_t(\theta) = \frac{\pi_{\theta}(a_t|s_t)}{\pi_{\theta_{old}}(a_t|s_t)}$$
 (6.16)

where  $\theta_{old}$  denotes the policy parameters at the previous iteration,  $\epsilon$  is a clip range hyperparameter that controls the size of the policy update,  $\operatorname{clip}(x,a,b)$  is a function that clips the value of x to be within the range [a, b], and  $A^{\pi_{\theta}}(s_t, a_t)$  is the estimated advantage of taking action  $a_t$  in state  $s_t$  over following the policy  $\pi_{\theta_{old}}$ . In our implementation, the advantage function is estimated following the GAE method proposed by Schulman et al. (2015b). The policy is updated by performing gradient ascent with respect to the policy parameters  $\theta$  to maximize the objective function, which is estimated by sampling from the environment and computing the empirical average of the objective function over the sampled trajectories. The first component in the objective function,  $\Lambda_t(\theta) \hat{A}^{\pi_{\theta}}(s_t, a_t)$ , is a surrogate objective function that approximates the advantage of taking action  $a_t$  in state  $s_t$  over following the policy  $\pi_{\theta_{old}}$ , and the second component,  $\operatorname{clip}(\Lambda_t(\theta), 1 - \epsilon, 1 + \epsilon)$  $\epsilon \hat{A}^{\pi_{\theta}}(s_t, a_t)$ , is a clipped surrogate objective function that prevents large policy updates and ensures more stable training. The TRPO algorithm proposed by Schulman et al. (2015a) uses a similar clipped surrogate objective function to ensure that the policy update does not deviate too far from the old policy, but PPO simplifies the optimization problem by using a clipped surrogate objective function instead of a hard-coded constraint on the policy update. Compared to TRPO, PPO is more computationally efficient and easier to implement, making it a popular choice for training deep reinforcement learning agents. For a in-depth discussion on the implementation details of both TRPO and PPO, we refer the reader to Engstrom et al. (2020).

Algorithm 9 outlines the PPO algorithm for training an agent to hedge VAs. At each iteration k, the agent collects a set of trajectories  $\mathcal{D}_k$  by interacting with the environment (Section 6.1.1) with the current policy  $\pi_{\theta_k}$ . The agent then computes the rewards  $\hat{R}_t$  for each trajectory in  $\mathcal{D}_k$  and estimates the advantage function with  $\hat{A}_{\pi_{\theta_k}}(s_t, a_t)$  using the value

#### **Algorithm 9** PPO for Hedging Variable Annuities

- 1: Initialize policy parameters  $\theta_0$  and value function parameters  $\phi_0$ .
- 2: **For** iteration k = 0, 1, 2, ... **do**
- 3: Collect a set of trajectories  $\mathcal{D}_k$  by sampling from the environment with policy  $\pi_{\theta_k}$ .
- 4: Compute the rewards  $\hat{R}_t$  for each trajectory in  $\mathcal{D}_k$ .
- 5: Estimates the advantage function with  $\hat{A}_{\pi_{\theta_k}}(\mathbf{s}_t, a_t)$  using value function  $V_{\phi_k}$ .
- 6: Update the policy parameters  $\theta_{k+1}$  by

$$\max_{\theta} \mathbb{E}\left[\min\left\{\Lambda_t(\theta)\hat{A}^{\pi_{\theta}}(\mathbf{s}_t, a_t), \operatorname{clip}(\Lambda_t(\theta), 1 - \epsilon, 1 + \epsilon)\hat{A}^{\pi_{\theta}}(\mathbf{s}_t, a_t)\right\}\right]$$

7: Update the value function parameters  $\phi_{k+1}$  by minimizing the mean squared error between the predicted value and the actual collected reward.

$$\phi_{k+1} = \arg\min_{\phi} \frac{1}{|\mathcal{D}_k|T} \sum_{\kappa \in \mathcal{D}_k} \sum_{t=0}^{T} \left( V_{\phi}(\mathbf{s}_t) - \hat{R}_t \right)^2$$

#### 8: End For

function  $V_{\phi_k}$ . The policy parameters  $\theta_{k+1}$  are updated by performing gradient ascent with respect to the policy parameters to maximize the clipped surrogate objective function. The value function parameters  $\phi_{k+1}$  are updated by minimizing the mean squared error between the predicted value and the actual collected reward. The agent iterates this process until convergence to learn an optimal policy for hedging VAs.

## 6.3.1 Accounting for Non-Markovian Dynamics

In the context of hedging VAs, the state space is partially observed. The agent does not have access to the contract specifications and the underlying risk factors. Furthermore, a rough volatility model introduces non-Markovian dynamics, i.e., the future state of the system depends on the entire history of the system. To address the non-Markovian dynamics, an LSTM component is added to the state representation  $\mathbf{s}_t$  to capture the temporal dependencies in the data. This approach is known as recurrent PPO (Ni et al., 2021). Let  $h_t$  be the hidden state of the LSTM at time t. The state representation  $\mathbf{s}_t$  is fed into the LSTM at each time step to update the hidden state  $h_t$ .

$$\mathbf{s}_t = (S_t, F_t, G_t, \tau, \Delta_{t-1}), \quad h_t = \text{LSTM}(\mathbf{s}_t, h_{t-1})$$
(6.17)

In a LSTM, the cell state  $c_t$  stores a representation of historical information up to time t. It is updated at each time step by a forget gate  $f_t$ , an input gate  $i_t$ , and an output gate  $o_t$ .

In a recurrent PPO with LSTM, the policy and reward function remain the same but with the new state representation in Equation 6.17. Instead of following the standard PPO algorithm, the agent uses the recurrent PPO algorithm to learn an optimal policy for hedging VAs. Figure 6.1 illustrates the network architecture of the standard PPO and the recurrent PPO with LSTM, and Algorithm 10 outlines the recurrent PPO algorithm and for training an agent to hedge VAs with non-Markovian dynamics.

#### Algorithm 10 Recurrent PPO with LSTM for Hedging Variable Annuities

- 1: Initialize policy parameters  $\theta_0$  and value function parameters  $\phi_0$ .
- 2: Initialize the LSTM layers with random weights, and set the hidden state  $h_0$  to zeros.
- 3: **For** iteration k = 0, 1, 2, ... do
- 4: Collect a set of trajectories  $\mathcal{D}_k$  by sampling from the environment with policy  $\pi_{\theta_k}$ .
- 5: Reset the LSTM hidden state  $h_0$  to zeros at the beginning of each trajectory.
- 6: Compute the rewards  $\hat{R}_t$  for each trajectory in  $\mathcal{D}_k$ .
- 7: Estimates the advantage function with  $A_{\pi_{\theta_k}}(\mathbf{s}_t, a_t)$  using value function  $V_{\phi_k}$ .
- 8: Update the policy parameters  $\theta_{k+1}$  by

$$\max_{\theta} \mathbb{E}\left[\min\left\{\Lambda_t(\theta)\hat{A}^{\pi_{\theta}}(\mathbf{s}_t, a_t), \operatorname{clip}(\Lambda_t(\theta), 1 - \epsilon, 1 + \epsilon)\hat{A}^{\pi_{\theta}}(\mathbf{s}_t, a_t)\right\}\right]$$

9: Update the value function parameters  $\phi_{k+1}$  by minimizing the mean squared error between the predicted value and the actual collected reward.

$$\phi_{k+1} = \arg\min_{\phi} \frac{1}{|\mathcal{D}_k|T} \sum_{\kappa \in \mathcal{D}_t} \sum_{t=0}^{T} \left( V_{\phi}(\mathbf{s}_t) - \hat{R}_t \right)^2$$

#### 10: End For

The recurrent PPO algorithm is similar to the standard PPO algorithm, but it includes an LSTM component to capture the temporal dependencies in the data. However, the sequantial nature of the LSTM is not suitable for parallel computation, which slows down the training process.



Figure 6.1: PPO Network Architectures

## 6.4 Transfer Learning for Hedging VAs

In the context of deep hedging, various research has been conducted to improve the hedging strategies on real-world data by leveraging knowledge from simulated data. Most of the research uses the offline-online learning approach, where the hedging strategies are first trained with simulation data, but updated in real-time based on new market information and continuous learning to adapt to changing market dynamics. The most relevant paper to our study is Xiao et al. (2021), which use policy gradient algorithms to train hedging strategies for European options with simulation data from Heston model and then updated in real-time hedging S&P 500 index options. To the best of our knowledge, no research has been conducted to improve the hedging strategies with the state-of-the-art transfer learning techniques. This section introduces some transfer learning techniques that are relevant to our task of hedging VAs.

## 6.4.1 Reward Shaping

In the context of hedging VAs using deep reinforcement learning and transfer learning, reward shaping becomes a crucial technique. It involves modifying the reward function in the target domain to incorporate additional guidance, which often leads to more efficient

learning and better performance. This technique can be particularly useful for managing the complex risks associated with VAs, as it can guide the learning process towards strategies that are more effective in hedging these products.

One way to implement reward shaping in this context is by introducing auxiliary rewards that reflect the performance of hedging strategies learned from the source domains. Reward shaping learns an auxiliary reward function  $\mathcal{R}_s: \mathcal{S} \times \mathcal{A} \times \mathcal{S} \to \mathbb{R}$  from the source domains. The auxiliary reward function is then used to shape the reward function in the target domain,  $\mathcal{R}'_t = \mathcal{R}_t + \mathcal{R}_s$ , by incorporating the insights gained from the source domains.

Some popular reward shaping techniques include potential-based reward shaping (Ng et al., 1999), potential based state-action advice (Wiewiora et al., 2003), dynamic value function advice (Harutyunyan et al., 2015).

## 6.4.2 Policy Transfer

The most straightforward approach to transfer learning is to directly transfer the policy learned from the source domains to the target domain via policy distillation. Policy distillation is a technique that transfers the knowledge from a teacher policy to a student policy by training the student policy to mimic the teacher policy up to a certain extent. The teacher policy is learned from the source domains, and the student policy learns the hedging strategies in the target domain while guided by the teacher policy. In the Distral algorithm (Teh et al., 2017), the student policy maximizes a multi-task objective:  $\max_{\phi} \sum_{i=1}^{K} \mathcal{J}(\pi_{\phi}, \pi_{E_i})$ , where

$$\mathcal{J}(\pi_{\phi}, \pi_{E_i}) = \sum_{t=0}^{\infty} \mathbb{E}_{s_t \sim \mu_0^t, a_t \sim \pi_{\phi}} \left[ \gamma^t R_{t+1} + \frac{\alpha}{\beta} \log \pi_{\phi}(a_t | s_t) - \frac{1}{\beta} log \pi_{E_i}(a_t | s_t) \right], \tag{6.18}$$

where a set of K teacher policies  $\pi_{E_i}$  are learned from the source domains, and the student policy  $\pi_{\phi}$  is learned on the target domain.  $\alpha$  and  $\beta$  are hyperparameters that control the trade-off between mimicking the teacher policies and exploring the target domain.

#### 6.4.3 Evaluation Metrics

The metrics discussed for evaluating transfer learning approaches focus on two key aspects: mastery and generalization. Mastery assesses the agent's final performance level

in the target domain, indicating how effectively it has learned a specific task from previous knowledge. Some common metrics for evaluating mastery include: Generalization, on the other hand, refers to the agent's ability to quickly adapt its learned knowledge to the target domain. In the context of risk management, generalization is more important than mastery, as the agent must be able to quickly adapt to changing market conditions and new financial products. Some common metrics for evaluating generalization include:

- Jumpstart performance: the initial reward of the agent in the target domain.
- Accumulated reward: the total area under the reward curve over time in the target domain.
- **Time to threshold:** the time it takes for the agent to reach a certain performance threshold in the target domain.
- Performance with fixed epochs: the agent's performance in the target domain after a fixed number of epochs.

## 6.5 Numerical Experiments

We present a numerical experiment to demonstrate the effectiveness of the model-free PPO algorithm for hedging VAs with transaction costs. We consider a VA contract with a maturity of T=240 months and a GMMB rider with a simulation environment described in Section 6.1.1. The simulation environment is implemented in Python using the OpenAI Gym framework (Brockman et al., 2016), and the neural networks are programmed using the PyTorch library (Paszke et al., 2019). The environment is designed to simulate the dynamics of the VA contract and provide the agent with the current state of the VA with the contract specifications and the financial market information described in Table 6.1.

The underlying asset price  $S_t$  is modeled as a geometric Brownian motion (GBM) with a drift rate  $\mu$  and a volatility  $\sigma$ .

Figure 6.2 illustrates the hedging performance of the deep hedging algorithm with transfer learning for hedging VAs with a GMMB rider. The first deep hedging agent is directly trained on the Heston asset model, and the second deep hedging agent is trained with transfer learning from the GBM model. The figure plot the 95%-CVaR of hedging errors against training epochs for both agents. The deep hedging agent with transfer learning outperforms the deep hedging agent trained on the Heston model in both mastery

Hyperparameter	Value
$\overline{ \mathcal{D}_k }$	16
$\epsilon$	0.2
Learning rate	0.0001
$S_0$	1000
$\mu$	0.00375
$\sigma$	0.0457627
r	0.002
T	240

Table 6.1: PPO Hyperparameters and GMMB Contract Specifications

and generalization. The agent with transfer learning achieves a lower loss in the target domain and adapts more quickly to the new environment, indicating that the agent has learned a more robust hedging strategy that can be effectively applied to a more complex environment.

For the PPO agent, we use a regular PPO and a recurrent PPO with an LSTM component to capture the temporal dependencies in the data. The weights of the value network and the policy network are initialized with orthogonal initialization (Engstrom et al., 2020) with scaling  $\sqrt{2}$  and the bias terms set to zero. The value network and the policy network are implemented as fully connected neural networks with two hidden layers of 16 units each and tanh activation functions. They are trained using the Adam optimizer with a learning rate of 0.0001 and a batch size automatically adjusted to the number of trajectories in the collected dataset. At each timestep, the observations are normalized and cliped between -10 and 10 before being fed into the neural networks to stabilize the training process. For PPO, we follow the suggestions in Schulman et al. (2017) and set the clipping hyperparameter  $\epsilon$  to 0.2 to prevent large policy updates. More specifically, our implementation of the PPO uses a generalized advantage estimation (Schulman et al., 2015b) to estimate the advantage function. The agent interacts with the environment for 10<sup>5</sup> episodes, and the policy and value networks are evaluated every 1000 episodes. The performance of the agent is evaluated based on the average reward over an independent test set of 10<sup>3</sup> episodes, and the results are compared with the benchmark hedging strategy, delta hedging.

This experiment compares the hedging performance of the recurrent PPO agent with LSTM and the deep hedging algorithm proposed by Buehler et al. (2019) for hedging VAs with a GMMB rider. The experiment is conducted with the same simulation budget.

Figure 6.4 illustrates the hedging performance of the standard PPO and the recurrent

#### 20-year GMMB with monthly rebalancing

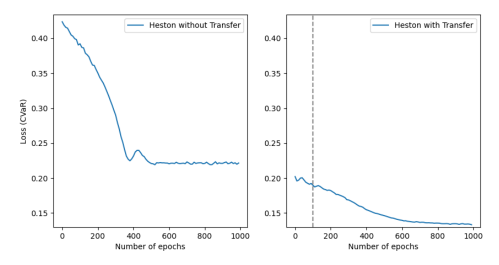


Figure 6.2: Hedging performance of deep hedging with transfer learning



Figure 6.3: Hedging performance of recurrent PPO and deep hedging

PPO with LSTM for hedging VAs with transaction costs C=0.005. The figures show the distributions of the hedging errors incurred by the agents over the  $10^3$  test episodes for the GBM and the regime-switching GBM asset models. For both asset models, the recurrent PPO outperforms the standard PPO in terms of hedging performance. For the regime-switching GBM, the 95%-VaR of hedging errors of the recurrent PPO and the standard PPO are 136.24 and 147.14, respectively. The hedging errors of the recurrent PPO are more concentrated around zero, indicating that historical information captured by the LSTM component helps the agent make better hedging decisions even in the environment



Figure 6.4: Hedging performance of standard PPO and recurrent PPO

is Markovian.

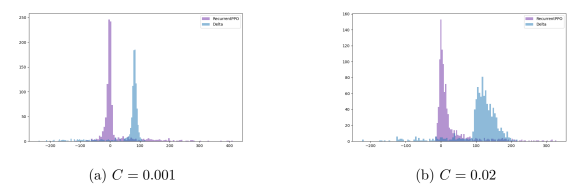


Figure 6.5: Hedging performance of recurrent PPO and delta hedging with different transaction costs

Figure 6.5 shows the hedging performance of the recurrent PPO agent and the delta hedging strategy with different transaction costs when the asset model is a GBM. The experiment is conducted with transaction costs of C=0.001 and C=0.02. The delta hedging strategy is implemented with the Black-Scholes delta. When the transaction cost is low, the delta hedging strategy outperforms the recurrent PPO agent. However, as the transaction cost increases, delta hedging becomes less effective, and the recurrent PPO agent is able to learn from the environment and adapt its hedging strategy to minimize the hedging errors.

Figure 6.6 illustrates the hedging performance of the recurrent PPO agent with and



Figure 6.6: Hedging performance of recurrent PPO with or without anchor

without the model information of the underlying asset model and the current value of the liability. The experiment is conducted for the GBM asset model with transaction costs of C=0.005. The results show that the agent with the model information does not outperform the agent without the model information.

Figure 6.7 illustrates the hedging performance of the recurrent PPO agent for hedging VAs with a GMWB rider. The experiment is conducted with the same hyperparameters as the experiment with the GMMB rider. Unlike the GMMB rider, the PPO agent struggles to learn an optimal hedging strategy for the VA with a GMWB rider. The hedging errors of the PPO agent are more dispersed, indicating that more training episodes are needed to achieve a satisfactory hedging performance. The results suggest that the GMWB rider introduces additional complexity to the hedging problem, and the model-free RL algorithm is not sample efficient.

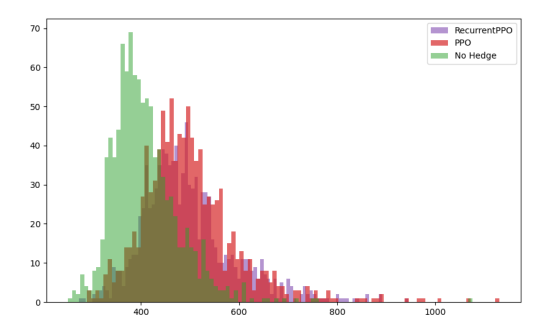


Figure 6.7: Hedging performance of recurrent PPO with GMWB

## 6.6 Future Directions

The proposed future work of this thesis aims to investigate how transfer learning can be effectively integrated into the training of PPO agents for dynamic hedging of VA contracts. The primary goal is to enhance the adaptability and efficiency of hedging strategies in response to new VA products and changing market conditions.

The first objective is to establish a systematic approach for transferring policies learned by PPO agents from source tasks to target tasks in the context of dynamic hedging. Source tasks may involve hedging a GMMB under a specific market model, while target tasks could entail hedging a GMWB under a different market model.

To achieve this, the research will explore methodologies such as policy distillation, where knowledge from a pre-trained "teacher" policy is transferred to a "student" policy (Rusu et al., 2015). Policy distillation has been effective in compressing and transferring knowledge between networks, improving learning efficiency in RL settings. Another avenue is the use of pre-trained model initialization, wherein the policy network trained on the source

task initializes the policy for the target task. This method leverages the shared underlying structures between tasks, potentially accelerating convergence and improving performance The framework will also consider feature representation transfer, aiming to reuse learned representations of market dynamics and policyholder behaviors between tasks. By capturing common patterns across different contracts and market models, the PPO agent can more effectively generalize to new scenarios (Bengio, 2012).

By addressing these objectives, the research aims to advance the field of financial risk management through the development of adaptive and efficient hedging strategies for variable annuities. The integration of transfer learning into PPO agents has the potential to significantly reduce computational costs and improve the robustness of hedging policies in dynamic market environments.

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# **Appendix**

# Connection between the convergence in distribution and MSE

In this section, we provide a proof of the connection between the convergence in distribution and the convergence of the MSE of the estimator. In Broadie et al. (2015), the authors show that asymptotically, the regression-based nested simulation estimator converges in distribution to the true value  $\rho$ .

$$\sqrt{M} \cdot \left( \mathbb{E}\left[ h(\hat{L}_{M,N}^{\text{REG}}(X; \hat{\beta})) - h\left(g(X)\right) \right] - \mathcal{B}_{\mathcal{M},M} \right) \xrightarrow{\mathcal{D}} \mathcal{N}\left(0, C_{\mathcal{M},h}\left(1 + \frac{C_{\Phi}}{N}\right)\right)$$
(6.19)

where  $\mathcal{B}_{\mathcal{M},M}$  is the model bias term.  $C_{\mathcal{M},h}$  and  $C_{\Phi}$  are constants that do not depend on M or N. Assume constant model error, the regression-based nested simulation estimator is asymptotically normal with decaying variance as M and N increase.

The mean squared error (MSE) of the estimator is given by

$$MSE = \mathbb{E}\left[\left(h(\hat{L}_{M,N}^{REG}(X;\hat{\beta})) - h\left(g(X)\right)\right)^{2}\right] = \mathcal{O}(M^{-1})$$
(6.20)

In this section, we want to show that the convergence in distribution in Broadie et al. (2015) implies the convergence of the MSE in the order of  $\mathcal{O}(M^{-1})$ .