The Markov Chain Monte Carlo Approach to
Importance Sampling in Stochastic Programming

by

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Submitted to the School of Engineering
in partial fulfillment of the requirements for the degree of
Master of Science in Computation for Design and Optimization

at the

MASSACHUSETTS INSTITUTE OF TECHNOLOGY

September 2012

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Abstract

Stochastic programming models are large-scale optimization problems that are used to facilitate decision-making under uncertainty. Optimization algorithms for such problems need to evaluate the expected future costs of current decisions, often referred to as the recourse function. In practice, this calculation is computationally difficult as it involves the evaluation of a multidimensional integral whose integrand is an optimization problem. Accordingly, the recourse function is estimated using quadrature rules or Monte Carlo methods. Although Monte Carlo methods present numerous computational benefits over quadrature rules, they require a large number of samples to produce accurate results when they are embedded in an optimization algorithm. We present an importance sampling framework for multistage stochastic programming that can produce accurate estimates of the recourse function using a fixed number of samples. Our framework uses Markov Chain Monte Carlo and Kernel Density Estimation algorithms to create a non-parametric importance sampling distribution that can form lower variance estimates of the recourse function. We demonstrate the increased accuracy and efficiency of our approach using numerical experiments in which we solve variants of the Newsvendor problem. Our results show that even a simple implementation of our framework produces highly accurate estimates of the optimal solution and optimal cost for stochastic programming models, especially those with increased variance, multimodal or rare-event distributions.

Thesis Supervisor: Mort Webster
Title: Assistant Professor of Engineering Systems
Acknowledgments

I would like to take this opportunity to thank my advisors, Mort Webster and Panos Parpas, who have supervised and funded my research at MIT over the past two years. This thesis would not have been possible without their guidance, insight and patience. I would also like to thank Youssef Marzouk, who helped develop and formalize many of the findings that I present in this thesis, as well Bryan Palmintier, who frequently helped resolve all kinds of important problems that I encountered in my research.

Much of this work has involved coding, debugging, computing and waiting... This process was incredibly frustrating at times, but it has certainly been easier in my case thanks to Jeff and Greg who manage the Svante cluster, as well as the anonymous users who post on the CPLEX forums and the StackExchange network. I very much appreciate the fact that these individuals have sacrificed their time and energy to solve problems that, quite frankly, did not affect them in any way. I look forward to the day that I can help others as selflessly as they have helped me.

Looking back at these past two years at MIT, I believe that I have been very fortunate to have been admitted to the CDO program, but even more fortunate to have Laura Koller and Barbara Lechner as my academic administrators. Their support has been invaluable, and I cannot fathom how I could have completed this degree without them.
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Chapter 1

Introduction

Stochastic programming (SP) models are large-scale optimization problems that are used to facilitate decision-making under uncertainty. Optimization algorithms for such problems require the evaluation of the expected future costs of current decisions, often referred to as the recourse function. In practice, this calculation is computationally difficult as it involves a multidimensional integral whose integrand is an optimization problem. Subsequently, many SP practitioners estimate the value of recourse function using quadrature rules ([30]) or Monte Carlo (MC) methods ([5] and [36]).

MC methods are an appealing way to estimate the recourse function in SP models because they are well-understood, easy to implement and remain computationally tractable for large-scale problems. Unfortunately, MC methods also produce estimates that are subject to sampling error, which can compound across the iterations of an optimization algorithm and produce highly inaccurate solutions for SP models. It is true that one can reduce the sampling error in MC estimates of the recourse function by increasing the number of samples used in an MC approach. In the context of SP, however, the number of samples that can be used to build an MC estimate of the recourse function is generally limited by the fact that each sample requires the solution to a separate optimization problem. As a result, MC methods are often paired with a variance reduction technique to reduce the sampling error in MC estimates of the recourse function without having to increase the number of samples.

This thesis focuses on a variance reduction technique known as importance sampling, which can dramatically reduce the sampling error of MC estimates by using an importance
sampling distribution to generate samples from regions that contribute most to the value of the recourse function. Although many distributions can achieve variance reduction through importance sampling, the most effective importance sampling distributions are typically crafted in order to exploit prior knowledge about the SP model. In light of this fact, the primary contribution of this thesis is an importance sampling framework that can reduce the sampling error in MC estimates without requiring the need to specify an importance sampling distribution beforehand.

Our framework, which we refer to as the Markov Chain Monte Carlo Approach to Importance Sampling (MCMC-IS), is based on an importance sampling distribution that is designed to produce MC estimates with zero variance ([2]). Although this zero-variance distribution cannot be used in practice, it is often used to guide the design of effective importance sampling distributions. Accordingly, our framework exploits the fact the zero-variance distribution is known up to a normalizing constant in order to build an approximation to the zero-variance distribution for importance sampling. In particular, MCMC-IS uses a Markov Chain Monte Carlo (MCMC) algorithm to generate samples from the zero-variance distribution, and then uses a Kernel Density Estimation (KDE) algorithm to reconstruct an approximate zero-variance distribution from these samples. With this approximate zero-variance distribution at hand, MCMC-IS then generates a new, larger set of samples and constructs an importance sampling estimate of the recourse function which has lower variance, and thus lower sampling error.

MCMC-IS has several benefits as a sampling framework: it is non-parametric, in that it does not require users to specify a family of importance sampling distributions; flexible, in that it can accommodate a wide array of MCMC and KDE algorithms; and robust, in that it can generate good results for probability distributions that are difficult to work with using existing sampling methods. It follows that MCMC-IS is advantageous in the context of SP because it can produce accurate estimates of the recourse function and improve the accuracy of output from an optimization algorithm. However, MCMC-IS is also beneficial in this context because can produce lower-variance estimates of the recourse function that improves the performance of stopping tests that assess the convergence of optimization algorithms. Moreover, MCMC-IS is well-suited for SP models because the computational overhead re-
quired to build an approximate zero-variance distribution is negligible in comparison to the computational overhead required to evaluate the recourse function in these models.

In this thesis, we demonstrate the performance of MCMC-IS using a series of numerical experiments based on a Newsvendor model. Our results show that MCMC-IS performs well in comparison to existing various reduction techniques, such as stratified sampling methods, Quasi-Monte Carlo methods and an early importance sampling technique developed in [7] and [20]. In particular, we show that MCMC-IS significantly outperforms these techniques when the uncertainty is modeled using a higher variance, rare-event or multimodal distribution. Even as our numerical experiments illustrate the computational performance of the MCMC-IS framework when it is embedded in the Stochastic Dual Dynamic Programming (SDDP) algorithm from [31], we stress that MCMC-IS can yield similar benefits in other algorithms that involve expected-value optimization, such as the sample average approximation method ([36]), stochastic decomposition ([17]), progressive hedging ([34]), variants of Benders’ decomposition ([5]) and approximate dynamic programming algorithms ([32]).

Although both MCMC and KDE algorithms have received considerable attention in the literature, they have not been previously combined in an importance sampling framework such as MCMC-IS, or applied to solve SP models. Nevertheless, the findings in this thesis build on existing research on the application of variance reduction techniques for MC methods in SP: Quasi-Monte Carlo methods were studied in [21] and in [10]; control variates were proposed in [37] and in [16]; a sequential sampling algorithm was proposed in [3]; an alternative importance sampling technique for SP was first developed in [7] and [20]. A computational assessment of conditional sampling, antithetic sampling, control variates and importance sampling appeared in [16]. Similarly, Quasi Monte Carlo methods and Latin Hypercube Sampling were compared in [19]. The link between sampling error of MC estimates and the solution quality of SP models was discussed in [23].

The remaining parts of this thesis are structured as follows. In Chapter 2 we provide a brief overview of SP models, illustrate how decomposition algorithms can produce inaccurate results when paired with an MC method, and provide an overview of variance reduction techniques to remedy this problem. In Chapter 3 we introduce MCMC-IS and provide a detailed overview of its practical and theoretical aspects. In Chapters 4 and 5 we present the
results of numerical experiments based on a Newsvendor model to illustrate the sampling properties of MCMC-IS, and highlight its benefits when it is used with a decomposition algorithm. Finally, we summarize our contributions and outline directions for future research in Chapter 6.
Chapter 2

Background

In this chapter, we explain how to model problems in decision-making under uncertainty using an SP model (Section 2.1). We then show how to simplify this model using a recourse function to represent the expected future costs of current decisions (Section 2.2.1). Next, we provide insight as how to solve this model using a decomposition algorithm (Sections 2.2.2 - 2.2.4), and we discuss the merits of estimating the value of recourse function in decomposition algorithms through an MC method, especially in the context of large-scale problems. We then present a simple example to illustrate how sampling error of MC estimates of the recourse function can significantly affect the output from a decomposition algorithm (Section 2.3). Given the relationship between the sampling error of MC estimates of the recourse function and the accuracy of the output from decomposition algorithms, we end this chapter with an overview of variance reduction techniques that can decrease the impact of sampling error in MC estimates (Section 2.5).
2.1 Modeling Decision-Making Problems using SP

A multistage stochastic linear program is an optimization problem that minimizes the expected cost of sequential decisions in an uncertain setting. Given a fixed time horizon $T$, a set of decisions vectors $x_1, \ldots, x_T$, and a set of random variables $\xi_2, \ldots, \xi_T$, we can formulate the deterministic equivalent of a $T$-stage stochastic linear program as

$$z^* = \min_{x_1, \ldots, x_T} c_1^T x_1 + \mathbb{E} \left[ \sum_{t=2}^{T} c_t (\xi_t)^T x_t \right]$$

s.t.

$$A_1 x_1 = b_1$$

$$A_t (\xi_t) x_t = b_t (\xi_t) - W_t (\xi_t) x_{t-1} \quad t = 2, \ldots, T$$

$$x_t \geq 0 \quad t = 1, \ldots, T$$

(2.1)

We assume that $c_t \in \mathbb{R}^{n_t}$, $A_t \in \mathbb{R}^{n_t \times m_t}$, $W_t \in \mathbb{R}^{n_t-1 \times m_t}$, $b_t \in \mathbb{R}^{m_t \times 1}$. The components of these parameters are deterministic for $t = 1$, but may be random for $t = 2, \ldots, T$. We refer to the set of all random components of the parameters at stage $t$ using a $D_t$-dimensional random vector $\xi_t$, and denote its joint probability density function, cumulative distribution function and support as $f_t$, $F_t$ and $\Xi_t$ respectively. In the context of two-stage problems, we simplify our notation by dropping the time index $t$, using an $(N_1 \times 1)$ vector $x$ to represent decisions in the first stage and an $(N_2 \times 1)$ vector $y$ to represent decisions in the second stage.

The deterministic equivalent formulation of a multistage stochastic linear program models the uncertainty in a decision-making problem as a scenario-tree, which implies that the solution to the linear program in (2.1) represents the optimal decisions for every branch of this tree. Although this approach is straightforward and comprehensive, it is rarely used in practice because the linear program in (2.1) grows exponentially with the number of stages and random outcomes of the underlying decision-making problem. In fact, using the deterministic equivalent to model a decision-making problem with $T$ stages and $K$ random outcomes per stage involves a linear program with $O(T^K)$ variables and constraints. This represents a significant computational burden in the context of large-scale problems in terms of the memory that is required to store the linear program, and the processing power that is required to solve within an acceptable timeframe.
2.2 Solving SP Models with Decomposition Algorithms

2.2.1 Representing Expected Future Costs with the Recourse Function

Decomposition algorithms are a set of optimization algorithms that are designed to solve SP models in a computationally tractable manner. In contrast to the comprehensive deterministic equivalent approach in (2.1), decomposition algorithms isolate the costs and decisions associated with each stage of the decision-making problem into $T$ nested linear programs, which we denote as $LP_1, \ldots, LP_T$. In this case, $LP_1$ is expressed as,

$$
\begin{align*}
    z^* &= \min_{x_1} \quad c_1 x_1 + Q_1(x_1) \\
    \text{s.t.} \quad A_1 x_1 &= b_1 \\
    x_1 &\geq 0
\end{align*}
$$

and $LP_t$ for $t = 2 \ldots T$ are expressed as,

$$
\begin{align*}
    Q_{t-1}(\hat{x}_{t-1}, \xi_t) &= \min_{x_t} \quad c_t(\xi_t) x_t + Q_t(x_t) \\
    \text{s.t.} \quad A_t(\xi_t) x_t &= b_t(\xi_t) - W_t(\xi_t) \hat{x}_{t-1} \\
    x_t &\geq 0
\end{align*}
$$

The decomposed formulation in (2.2) and (2.3) captures the sequential and uncertain structure of the decision-making process as $LP_2, \ldots, LP_T$ depend on the previous stage decision $\hat{x}_{t-1}$ and a realization of the uncertainty $\xi_t$. We formalize this dependence by representing the optimal cost of $LP_t$ for $t = 2 \ldots T$ using the function $Q_{t-1}(\hat{x}_{t-1}, \xi_t)$. We note that we set $Q_T(x_T, \xi_T) \equiv 0$ without loss of generality because we assume that our decision-making problem ends after $T$ stages.

The decomposition formulation in (2.2) and (2.3) frames the optimal decision at each time period as a decision that balances present costs and expected future costs. In particular, the optimal decision at stage $t$ minimizes the sum of present costs at stage $t$, which are expressed as $c_t(\xi_t)x_t$, and expected future costs at stages $t + 1, \ldots, T$, which are expressed as $E[Q_t(\hat{x}_t, \xi_{t+1})]$. In SP, the expected future costs at stages $1, \ldots, T - 1$ are represented
using a function $Q_t(x_t)$ that omits the expectation operator for clarity. The function $Q_t(x_t)$ is referred to as the recourse function, and it is defined as

$$Q_t(x_t) = \mathbb{E}[Q_t(x_t, \xi_{t+1})] = \int_{\xi_t} Q_t(x_t, \xi_{t+1}) f_{t+1}(\xi_{t+1})$$

(2.4)

### 2.2.2 Approximating the Recourse Function with Cutting Planes

The recourse function $Q_t(x_t)$ defined in (2.4) represents the expected value of a linear program with multiple random parameters. As a result, its value can only be determined by evaluating a multidimensional integral whose integrand is a linear program. Given the computational burden involved in evaluating multidimensional integrals, let alone linear programs, the recourse function should be approximated using few functional evaluations in order to solve SP models in a computationally tractable way.

Decomposition algorithms achieve this goal by constructing a piecewise linear approximation to the recourse function $Q_t(x_t)$ which only requires the evaluation of the multidimensional integral at a limited number of points $x_t$. The resulting approximation is a collection of supporting hyperplanes to the recourse function at fixed points $x_t$. In the SP literature, the supporting hyperplanes are referred to as cutting planes or cuts, and the fixed points $x_t$ around which the cuts are built are emphasized using the notation $\hat{x}_t$. Given a fixed point $\hat{x}_t$, a cut is a linear inequality defined as,

$$Q_t(x_t) \geq Q_t(\hat{x}_t) + \nabla Q_t(\hat{x}_t)(x_t - \hat{x}_t)$$

(2.5)

In practice, the values of the cut parameters $Q_t(\hat{x}_t, \xi_{t+1})$ and $\nabla Q_t(\hat{x}_t, \xi_{t+1})$ are determined using the expected values of the optimal dual variables $\lambda_{t+1}$ from LP$_{t+1}$. In particular,

$$Q_t(\hat{x}_t) = \mathbb{E}[Q_t(\hat{x}_t, \xi_{t+1})] = \mathbb{E}[\lambda_{t+1}^T (b_{t}(\xi_{t+1}) - W_t(\xi_{t+1})]]$$

$$\nabla Q_t(\hat{x}_t) = \mathbb{E}[\nabla Q_t(\hat{x}_t, \xi_{t+1})] = \mathbb{E}[\lambda_{t+1}^T W_t(\xi_{t+1})]$$

(2.6)

Given that the linear inequality defined in (2.5) has the same number of variables as LP$_t$, it is added to the set of existing constraints in LP$_t$ in order to improve the current approximation.
of the recourse function $Q_t$.

We note that the benefit of using the optimal dual variables in constructing the cut parameters is that they can still be determined when LP$_{t+1}$ is infeasible for a given value of the previous stage decision $\widehat{x}_t$ or the uncertain outcome $\xi_{t+1}$. In such cases, a decomposition algorithm can use any feasible set of dual variables to construct a cut that will prevent infeasible instances of LP$_{t+1}$. This cut is referred to as a feasibility cut, and it is defined as,

$$Q_t(x_t) \geq \nabla Q_t(\widehat{x}_t)(x_t - \widehat{x}_t) \quad (2.7)$$

### 2.2.3 Stopping Procedures

A generic iteration of a decomposition algorithm consists of constructing $T-1$ cuts to support the recourse functions $Q_1, \ldots, Q_{T-1}$ at a set of fixed values $\widehat{x}_1, \ldots, \widehat{x}_{T-1}$, and adding these cuts to the linear programs LP$_1, \ldots, LP_{T-1}$. Assuming that the cut parameters in (2.6) can be calculated exactly, each cut that is added to LP$_t$ improves the approximation of the recourse function $Q_t$, and brings the estimated values of the optimal decision $\widehat{x}_t$ and the optimal cost $\widehat{z}_t$ closer to their true values $x^*$ and $z^*$. Although it is impossible to determine the true optimal cost $z^*$ of a multistage SP in a general setting, a decomposition algorithm can produce a lower bound $z_{LB}$ and an upper bound $z_{UB}$ to $z^*$. Given that the value of the lower bound $z_{LB}$ is monotonically non-decreasing with each iteration and the value of the upper bound $z_{UB}$ is monotonically non-increasing with each iteration, these bounds can then be used to stop decomposition algorithms when their difference $|z_{LB} - z_{UB}|$ is smaller than a user-prescribed tolerance.

The lower bound $z_{LB}$ produced by decomposition algorithms exploits the fact that a cutting plane approximation of the recourse function consistently underestimates the value of the true recourse function. This is because the approximation is composed of supporting hyperplanes to a convex function. In this case, the convexity of the recourse function is assured as it represents the expected value of a convex function (we note that the cost of a linear program is a convex function, and the expected value operation preserves convexity). As a result, we can obtain a lower bound $z_{LB}$ to the true optimal cost $z^*$ by considering the deterministic cost that we incur in the first stage, and the estimated costs that we expect to
incur in future stages,

\[ z_{LB} = \min_{x_1} c_1 x_1 + \hat{Q}_1(x_1) \]  

(2.8)

Given that a cutting plane approximation of the recourse function consistently underestimates expected future costs, it follows that decisions made with this approximate recourse function will be suboptimal. By definition, the true expected cost of these suboptimal decisions will exceed the optimal cost of the SP model. In other words, we can obtain an upper bound to the true optimal cost by calculating the true expected cost of the suboptimal decisions that are produced with our current approximation of the recourse function. In the context of a multistage SP, we can calculate the true expected cost of these decisions by calculating the cost associated with each sequence of uncertain outcomes \( \xi_2, \ldots, \xi_T \), and forming its expected value. Assuming that there exists \( K \) unique sequences of uncertain outcomes, the upper bound can be calculated as

\[ z_{UB} = \sum_{i=1}^{K} \sum_{t=1}^{T} c_t(\xi_i^t) \tilde{x}_t(\xi_i^t)f_t(\xi_i^t) \]  

(2.9)

where,

\[ \tilde{x}_i^t(\xi_i^t) = \arg \min c_t(\xi_i^t)x_t + \hat{Q}_t(x_i^t) \]  

(2.10)

2.2.4 Overview of Decomposition Algorithms

All decomposition algorithms solve SP models through an iterative process that builds cuts around fixed points \( \hat{x}_t \) and adds them to LP, for \( t = 1, \ldots, T - 1 \). The differences between these algorithms are primarily based in the way that they choose the fixed points \( \hat{x}_t \) around which they build cuts, the number of cuts that they add with each iteration, and whether they keep the cuts, drop them after a fixed number of iterations, or refine them with each iteration. In the context of multistage models, decomposition algorithms can also differ in the order of the stages at which they build the cuts.

Decomposition algorithms that can be characterized using these traits include the Abridged Nested Decomposition algorithm from [9], the Cutting Plane and Partial Sampling algorithm from [6], the ReSa algorithm from [18] and the Stochastic Decomposition algorithm from [17].
In this thesis, we restrict our focus on the SDDP algorithm that is presented in [31] due to its popularity among SP practitioners. The SDDP algorithm uses a greedy procedure to pick the fixed points \( \hat{x}_t \), adds a single cut to LP\(_t\) for \( t = 1, \ldots, T - 1 \) at each iteration, and permanently keeps the cuts that are produced with each iteration.

We refer the interested reader to [24] for a simple theoretical comparison between decomposition algorithms, and to [5] for a comprehensive introduction to the theory and practice of multistage SP.

### 2.3 Using Monte Carlo Methods in Decomposition Algorithms

The computational bottleneck in solving a multistage stochastic linear program involves calculating the cut parameters in (2.6), as this requires the evaluation of a multidimensional integral whose integrand is a linear program. While the cut parameters are easy to calculate when \( \xi_{t+1} \) is a discrete random variable with few outcomes, the calculation is intractable when \( \xi_{t+1} \) is high-dimensional, and impossible when \( \xi_{t+1} \) continuous. Subsequently, many SP practitioners simplify this calculation by modeling the uncertainty in their decision-making problem using scenario trees.

Scenario trees are discrete in nature, meaning that they either require models that exclusively contain discrete random variables, or a discretization procedure that can represent continuous random variables using a finite set of outcomes and probabilities. In the latter case, we note the optimal solution to an SP model in which the continuous random variables are discretized may differ from the optimal solution of an SP model in which the continuous random variables are kept in place. Even in situations where a scenario tree approach can produce accurate solutions, this level of accuracy is difficult to maintain in large-scale problems with multiple random variables and time periods due to the exponential growth in the size of the scenario tree. In such cases, scenario trees impose an unnecessary choice between high-resolution discrete approximations that yield accurate solutions but are difficult to store and solve, and low-resolution discrete approximations that may yield inaccurate solutions but are easier to store and solve.

MC methods are an alternative approach to calculate the cut parameters in (2.6). The
advantages of this approach are that it can accommodate discrete or continuous random variables, remain computationally tractable for models with a large number of random variables and produce estimates of the recourse function whose error does not depend on the number of random variables in the model. In practice, an MC method involves randomly sampling $N$ i.i.d. outcomes of the uncertain parameters $\xi_{t+1}^1, \ldots, \xi_{t+1}^N$, and estimating the expected values of the cut parameters in (2.6) through the sample averages,

$$
Q_t(\hat{x}_t) \approx \hat{Q}_t(\hat{x}_t) = \frac{1}{N} \sum_{i=1}^{N} Q_t(\hat{x}_t, \xi_{t+1}^i)
$$

$$
\nabla Q_t(\hat{x}_t) \approx \nabla \hat{Q}_t(\hat{x}_t) = \frac{1}{N} \sum_{i=1}^{N} \nabla Q_t(\hat{x}_t, \xi_{t+1}^i)
$$

(2.11)

Given that the cut parameters in (2.11) are produced by random sampling, it follows that they are subject to sampling error. In turn, the supporting hyperplane that is produced using these parameters is also subject to sampling error. We refer to this supporting hyperplane as a sampled cut, and note that it has the form,

$$
Q_t(x_t) \geq \hat{Q}_t(\hat{x}_t) + \nabla \hat{Q}_t(\hat{x}_t)(x_t - \hat{x}_t)
$$

(2.12)

2.4 The Impact of Sampling Error in Decomposition Algorithms

In comparison to the exact cut in (2.5), the sampled cut in (2.12) may produce an invalid approximation of the recourse function. We illustrate this phenomenon in Figures 2-1 and 2-2, where we plot sampled cuts that are produced when a crude MC method is paired with a decomposition algorithm to solve a simple two-stage Newsvendor model. We note that the parameters of this model are specified in Section 4.1.1.

Both cuts in this example were constructed using $N = 50$ samples. For clarity, we plot a subset of the sample values $Q(\hat{x}, \xi_i)$ for $i = 1, \ldots, N$ along the vertical line of $\hat{x}$, as well as their sample average $\frac{1}{N} \sum_{i=1}^{N} Q(\hat{x}, \xi_i)$. In Figure 2-1, we are able to generate a valid sampled cut, which is valid because it underestimates the true recourse function $Q(x)$ at all values of $x$. However, it is possible to generate a sampled cut that in some regions overestimates, and in other regions underestimates the true recourse function $Q(x)$. We illustrate this situation
Figure 2-1: A valid sampled cut.

Figure 2-2: An invalid sampled cut.
in Figure 2-2, where the sampled cut excludes the true optimal solution at \( x^* \approx 69 \) with \( z^* \approx -20 \). Assuming that the decomposition algorithm will only generate valid sampled cuts until it converges, the resulting estimates of \( x^* \) and \( z^* \) will be \( \hat{x} \approx 38 \) and \( \hat{z} \approx -15 \), corresponding to errors of 80% and 25% respectively. We note that the optimal solution \( x^* \) corresponds to the value of \( x \) that minimizes the sum of the first-stage costs and the recourse function, and not the value of \( x \) that minimizes the recourse function (although these values appear to be very close to each other in Figures 2-1 and 2-2).

Even in cases where sampling error in MC estimates of the cut parameters is negligible, its presence can have a significant impact on the final values of the optimal solution and optimal cost that are produced from a decomposition algorithm. This is because multiple quantities that affect the final output in a decomposition algorithm also depend on the cut parameters, such as the values of the lower bound \( z_{LB} \) and the upper bound \( z_{UB} \) that are used to stop the algorithm. In this case, the presence of sampling error means that these quantities are no longer deterministic values but random distributed estimates and a suitable statistical procedure is required in order to stop the algorithm. As we demonstrate in Section 3.3, a poorly designed procedure in such situations may stop decomposition algorithm before it has converged, and thereby result in highly inaccurate estimates of the optimal solution and the value.

### 2.5 Reducing Sampling Error through Variance Reduction

It is well-known that the sampling error in MC estimates of the cut parameters can be expressed as,

\[
\begin{align*}
\text{SE}(\tilde{Q}_t(\tilde{x}_t)) &= \sqrt{\text{Var}[\tilde{Q}_t(\tilde{x}_t)]} = \frac{\sigma_{\tilde{Q}_t(\tilde{x}_t)}}{\sqrt{N}} \\
\text{SE}(\nabla \tilde{Q}_t(\tilde{x}_t)) &= \sqrt{\text{Var}[\nabla \tilde{Q}_t(\tilde{x}_t)]} = \frac{\sigma_{\nabla \tilde{Q}_t(\tilde{x}_t)}}{\sqrt{N}}
\end{align*}
\] (2.13)

where \( \sigma_{\tilde{Q}_t(\tilde{x}_t)} \) and \( \sigma_{\nabla \tilde{Q}_t(\tilde{x}_t)} \) represent the true standard deviation of the recourse function and its gradient at the fixed point \( \tilde{x}_t \) respectively. Although (2.13) implies that we can reduce the sampling error of MC estimates by increasing the number of samples, the \( O\left(\frac{1}{\sqrt{N}}\right) \)
convergence rate implies that we have to solve four times as many linear programs in order to halve the sampling error of the cut parameters. Given the time that is required to solve a linear program within a large-scale multistage SP model, such an approach is simply not tractable. As a result, MC methods are typically paired with a variance reduction technique that can reduce the sampling error of MC estimates by either improving the convergence rate or reducing the underlying variance of the model.

Variance reduction techniques have generated much interest due to the application of MC methods across numerous fields; we refer the interested reader to [12], [22] and [25] for an introductory overview of these techniques.

2.5.1 Stratified Sampling

Stratified sampling techniques are a set of variance reduction techniques that first split the support $\Xi$ of a random variable $\xi$ into $K$ strata, and generate samples from each of the $K$ strata. This approach ensures that the samples are randomly distributed while achieving some variance reduction by spreading samples across the entire sample space. When $K = N$ strata are used, the stratified sampling technique is referred to as Latin Hypercube Sampling (LHS), and it produces estimates whose sampling error converges at a rate of $O\left(\frac{1}{\sqrt{N}}\right)$ albeit within a constant factor of traditional MC methods. We note that this convergence rate is slow in comparison to state-of-the-art stratified sampling techniques, which can increase the rate by allocating the $N$ samples in proportion to the variance of each $K$ strata. We recommend [26] and [40] for a more detailed overview of stratified sampling and LHS.

2.5.2 Quasi-Monte Carlo

Quasi Monte Carlo (QMC) methods are a set of variance reduction techniques that reduce the sampling error in MC estimates by using a deterministic sequence of points that is uniformly distributed across multiple dimensions. Examples of such sequences include Halton and Sobol sequences, whose points are depicted in Figure 2-3.

QMC methods are typically paired with a scrambling algorithm that is specifically designed to randomize a particular sequence of points. Popular examples of scrambling algorithms include the Owen scrambling algorithm for Sobol sequences, and the Reverse-Radix
2 algorithm for Halton sequences. As shown in Figure 2-4, scrambling randomizes the points from QMC sequences while maintaining their uniformity across each dimension.

Figure 2-4: Cross section of points from a randomized Halton sequence (green) and a randomized Sobol sequence (blue).

In the cases that an MC estimate is produced using $N$ randomized points from a QMC method, it has been shown that the sampling error in the estimates converges at an improved rate of $O\left(\frac{\log N}{N^{0.5(D-1)}}\right)$ so long as the recourse function is smooth. For a more detailed introduction to QMC methods, we refer the reader to [28] and [29].

2.5.3 Importance Sampling

Importance sampling is a variance reduction technique that aims to reduces the sampling error in MC estimates by generating samples from an importance sampling distribution $g$, as opposed to the original sampling distribution $f$. When samples are generated from an
importance sampling distribution $g$, the recourse function can be calculated as,

$$Q(\hat{x}) = \mathbb{E}^f[Q(\hat{x}, \xi)]$$

$$= \int_{\Xi} Q(\hat{x}, \xi) f(\xi) d\xi$$

$$= \int_{\Xi} Q(\hat{x}, \xi) f(\xi) \frac{g(\xi)}{g(\xi)} d\xi$$

$$= \int_{\Xi} Q(\hat{x}, \xi) \frac{f(\xi)}{g(\xi)} g(\xi) d\xi$$

$$= \int_{\Xi} Q(\hat{x}, \xi) \Lambda(\xi) g(\xi) d\xi$$

$$= \mathbb{E}^g[Q(\hat{x}, \Lambda(\xi))] \tag{2.14}$$

In (2.14), the function $\Lambda : \Xi \to \mathbb{R}$,

$$\Lambda(\xi) = \frac{f(\xi)}{g(\xi)} \tag{2.15}$$

is typically referred to as the likelihood function, and it is used to correct the bias that is produced by the fact that we generated samples from the importance sampling distribution $g$ instead of the original distribution $f$. Once we select a suitable important sampling distribution $g$, we can generate a set of $N$ i.i.d. samples $\xi_1, \ldots, \xi_N$, and construct an importance sampling estimate of the recourse function as,

$$\hat{Q}(\hat{x}) = \frac{1}{N} \sum_{i=1}^{N} Q(\hat{x}, \xi_i) \Lambda(\xi_i) \tag{2.16}$$

In theory, importance sampling simply reflects a change in the measure with which we compute the recourse function at a fixed point $\hat{x}$. Accordingly, any distribution $g$ can be used as an importance sampling distribution as long as the likelihood function $\Lambda$ is well-defined over the support of $f$. In other words, the importance sampling distribution $g$ should be chosen so that $g(\xi) > 0$ at all values of $\xi$ where $f(\xi) > 0$. When this requirement is satisfied, the sampling error of importance sampling methods also converges at a rate of $O\left(\frac{1}{\sqrt{N}}\right)$, albeit with a different constant factor than traditional MC methods.

Ideally, an importance sampling distribution $g$ is one that can generate samples at regions where $Q(\hat{x}) f(\xi)$ attains high values, which are referred to as the important regions of the
recourse function. Nevertheless, we stress that the importance sampling distribution $g$ should also be able to evaluate the probability $g(\xi)$ of each sample to a high degree of accuracy. This is because misspecified values of the importance sampling distribution $g(\xi)$ will produce misspecified values of the likelihood $\Lambda(\xi)$, and produce an importance sampling estimate that is highly biased.

We refer the interested reader to [2] for a more detailed review of importance sampling.

### 2.5.4 IDG Importance Sampling

Importance sampling was first applied to SP in [7] and [20]. We refer to this importance sampling distribution proposed as the Infanger-Dantzig-Glynn (IDG) distribution. The IDG distribution has been shown to mitigate the issues associated with the use of sampled cuts in decomposition algorithms that we cover in Section 2.3. Unfortunately, the IDG distribution in these papers makes several assumptions which severely limit its applicability to a broad range of SP models.

To begin with, the IDG distribution can only be used in SP models where the uncertainty is modeled using discrete random variables. As a result, any SP model where we can use the IDG distribution for importance sampling is subject to the same computational issues that we attribute to the scenario tree approach in Section 2.3.

Moreover, the IDG distribution assumes that the cost surface $Q(\hat{x}, \xi)$ is additively separable in the random dimensions, meaning that

$$Q(\hat{x}, \xi) \approx \sum_{d=1}^{D} Q_d(\hat{x}, \xi_d) \quad (2.17)$$

In SP models where such an approximation does not hold, the sampling error in the IDG estimate will still converge at a rate that is $O\left(\frac{1}{\sqrt{N}}\right)$ but with a much larger constant than traditional MC methods. In such cases, the IDG distribution produces estimates that have high rates of error.

A final issue with the IDG distribution is that it requires practitioners to know or determine the value of random outcome $\xi$ which minimizes the cost surface $Q(\hat{x}, \xi)$. In a general setting, the only way to determine the value is to perform an exhaustive search across all the uncertain
outcomes $\xi \in \Xi$.

It is true that there exist practical ways to work around these assumptions. However, we note that our numerical experiments in Section 4.8.1 suggest that the performance of the IDG distribution is critically determined by these factors. In turn, there remains a need for an alternative approach importance sampling that does not suffer from such issues for a broader range of SP models.
Chapter 3

The Markov Chain Monte Carlo Approach to Importance Sampling

In this chapter, we introduce the zero-variance importance sampling distribution (Section 3.1.1) and provide an overview of MCMC algorithms (Section 3.1.2) and KDE algorithms (Section 3.1.3). We then proceed to explain how these algorithms can be combined to build an importance sampling distribution that approximates the zero-variance distribution (Section 3.2) - a procedure that we refer to as the Markov Chain Monte Carlo Approach to Importance Sampling (MCMC-IS). Having introduced MCMC-IS, we present a simple MCMC-IS implementation to illustrate how MCMC-IS can be used in practice (Section 3.3). We end this chapter with a discussion of the theoretical aspects of MCMC-IS where we cover the ingredients of a convergence analysis (Section 3.4).
3.1 Foundations of the Markov Chain Monte Carlo Approach to Importance Sampling

3.1.1 The Zero-Variance Distribution

Importance sampling is most effective in the context of SP models when the importance sampling distribution \( g \) can generate samples from regions that contribute the most to the value of the recourse function \( Q(\hat{x}) \). In fact, when an importance sampling distribution can generate samples according to the exact importance of each region as,

\[
g^*(\xi) = \frac{|Q(\hat{x}, \xi)| f(\xi)}{\mathbb{E}^f[|Q(\hat{x}, \xi)|]} \tag{3.1}
\]

then the variance and sampling error of its estimates will be minimized (see [2]). Moreover, if the recourse function \( Q(x, \xi) > 0 \) for all \( \xi \in \Xi \), then the importance sampling distribution \( g^* \) can produce a perfect estimate of the recourse function with only a single sample \( \xi_1 \) as,

\[
\hat{Q}(\hat{x}) = \frac{1}{N} \sum_{i=1}^{N} Q(\hat{x}, \xi_i) \Lambda(\xi_i)
= Q(\hat{x}, \xi_1) \frac{f(\xi_1)}{g^*(\xi_1)}
= Q(\hat{x}, \xi_1) \frac{f(\xi_1)}{Q(\hat{x}, \xi_1) f(\xi_1)} \frac{\mathbb{E}^f[Q(\hat{x}, \xi)]}{\mathbb{E}^f[|Q(\hat{x}, \xi)|]}
= \mathbb{E}^f[Q(\hat{x}, \xi)]
\tag{3.2}
\]

In light of this fact, the importance sampling distribution \( g^* \) is often referred to as the zero-variance importance sampling distribution. The problem with using the zero-variance distribution \( g^* \) in practice is that it requires knowledge of \( \mathbb{E}^f[|Q(x, \xi)|] \), which is the quantity that we sought to compute in the first place. In turn, we are faced with a ”curse of circularity” in that we can use the zero-variance distribution \( g^* \) to construct perfect estimates if and only if we already have a perfect estimate of \( \mathbb{E}^f[|Q(\hat{x}, \xi)|] \).
3.1.2 Overview of MCMC Algorithms

MCMC algorithms are an established set of MC methods that can sample from a distribution which is known up to a normalizing constant. In contrast to other MC methods, MCMC algorithms generate a serially correlated sequence of samples $\xi_1, \ldots, \xi_M$. This sequence constitutes a Markov chain whose stationary distribution is equal to the distribution that we wish to sample from.

The simplest MCMC algorithm is the Metropolis-Hastings algorithm from [27], which we refer to throughout this thesis and cover in detail in Section 3.3. The Metropolis-Hastings algorithm generates samples from a target distribution $g$ by proposing new samples through a proposal distribution $q$ and accepting each sample as the next state of the Markov chain using a simple accept-reject rule.

In addition, we refer to the Adaptive Metropolis algorithm from [14] in Section 4.5, which uses a random walk distribution as the proposal distribution in the Metropolis-Hastings algorithm and automatically scales the step-size within this distribution. Lastly, we cover the Hit-and-Run algorithm from [39] in Section 4.6, which is designed to generate samples within bounded regions by using an accept-reject approach that resembles the Metropolis-Hastings algorithm.

We note that numerous other MCMC algorithms have been developed for different practical applications, and many of them can be used in the importance sampling framework that we present in this thesis. We refer the interested reader to [1], [11], and [13] for a comprehensive overview of the theoretical and practical aspects of MCMC algorithms.
3.1.3 Overview of KDE Algorithms

KDE algorithms are an established set of techniques that are designed to reconstruct a continuous probability distribution from a finite set of samples. Given a set of a $M$ samples, $\xi_1, \ldots, \xi_M$, the output of a KDE algorithm is an empirical probability distribution function,

$$\hat{g}_M(\xi) = \frac{1}{M} \sum_{i=1}^{M} K_H(\xi, \xi_i)$$  \hspace{1cm} (3.3)

where the function $K_H$ is referred to as a kernel function, and the matrix $H \in \mathbb{R}^{D \times D}$ is referred to as the bandwidth matrix. We note that the kernel function $K_H$ in (3.3) determines the probability of the region that surrounds each of the $M$ samples $\xi_1, \ldots, \xi_M$ while the bandwidth matrix $H$ determines the width of the region spanned by the kernel function $K_H$ at each sample.

In theory, the kernel function $K_H$ has to be chosen so that the output from the KDE algorithm $\hat{g}_M(\xi)$ is a probability distribution. In the multidimensional case, this requires a function $K_H$ such that

$$K_H(\cdot, \cdot) \geq 0$$

$$\int_{\Xi} K_H(\xi, \cdot) d\xi = 1$$

$$\int_{\Xi} \xi K_H(\xi, \cdot) d\xi = 0$$

$$\int_{\Xi} \xi \xi^T K_H(\xi, \cdot) d\xi = 0$$  \hspace{1cm} (3.4)

Assuming that these conditions are satisfied, the kernel function is said to be well-behaved, and its shape does not significantly impact the empirical distribution $\hat{g}_M$ that is produced by a KDE algorithm. In practice, $K_H$ is typically set as the product of $D$ one-dimensional kernel functions $K_1, \ldots, K_D$ that are symmetric around the origin in order to reduce the computational burden associated with KDE algorithms. Examples of such functions include the Gaussian, Laplacian or Epatchenikov kernel functions, which we plot in Figure 3-1.

In comparison to the impact of the kernel function $K_H$, the bandwidth matrix $H$ can substantially impact the accuracy of an empirical distribution produced by $\hat{g}_M$. Although the
bandwidth matrix $H$ is only required to be symmetric and positive definite, KDE algorithms automate the choice of $H$ through a bandwidth estimator, which determines the value of each entry within $H$ in order to minimize the error in $\hat{g}_M$ according to different metrics and assumptions. We include an overview of bandwidth estimators in Table 3.1.

We note that there exists many different KDE algorithms that can be applied in the importance sampling framework that we present in this thesis. We refer the interested reader to [8], [35] and [38] for an more detailed overview of these algorithms.

### 3.2 The Markov Chain Monte Carlo Approach to Importance Sampling

The importance sampling framework that we present in this thesis is motivated by two insights regarding the zero-variance distribution as defined in (3.1).

The first insight is that the zero-variance distribution is known up to a normalizing constant $E[|Q(\hat{x}, \xi)|]$. This implies that we can generate samples from this distribution using an MCMC algorithm. Unfortunately, we cannot use these samples to form a perfect estimate even as they belong to the zero-variance distribution. This is because we still need to evaluate the likelihood of each sample as defined in (2.15). In this case, the likelihood of a sample $\xi$ is given by,

$$\Lambda^*(\xi) = \frac{E[|Q(\hat{x}, \xi)|]}{|Q(\hat{x}, \xi)|}$$  

which is impossible to compute because it depends on $E[|Q(\hat{x}, \xi)|]$. Our inability to use the
<table>
<thead>
<tr>
<th>Approach</th>
<th>Optimal Bandwidth $h^*$</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean Integrated Squared Error</td>
<td>$\arg\min_h \mathbb{E} \left[ \int (\hat{g}_M(\xi) - g(\xi))^2 d\xi \right]$</td>
<td>-</td>
</tr>
<tr>
<td>Asymptotic Mean Integrated Squared Error</td>
<td>$M^{-\frac{1}{5}} \left( \frac{R(K)}{R(g^{\prime\prime}) \sigma_K^4} \right)^{\frac{1}{5}}$</td>
<td>$R(K) = \int K^2(\xi) d\xi$, $\sigma_K^2 = \int \xi^2 K(\xi) d\xi$</td>
</tr>
<tr>
<td>Gaussian Rule of Thumb</td>
<td>$\left( \frac{\hat{\sigma}^5}{3n} \right)^{\frac{1}{5}}$</td>
<td>$\hat{\sigma} = \min \left( S, \frac{IQR}{1.34} \right)$ $S$ and $IQR$ denote the sample standard deviation and interquartile range of the samples</td>
</tr>
<tr>
<td>Leave-One-Out Cross Validation</td>
<td>$\arg\min_h \int \hat{g}<em>M^2(\xi) d\xi - \frac{2}{M} \sum</em>{i=1}^{M} \hat{g}_{M,-i}(\xi_i)$</td>
<td>$\hat{g}_{M,-i}$ denotes the distribution $\hat{g}_M$ that is formed using all samples except $\xi_i$</td>
</tr>
</tbody>
</table>

Table 3.1: Bandwidth estimators for KDE algorithms.
samples that we generate from the zero-variance distribution to form a perfect estimate leads us to the second insight: while we cannot use the samples to form an importance sampling estimate, we can use them to reconstruct an approximation of the zero-variance distribution by using a KDE algorithm.

In particular, assuming that we have generated $M$ samples from the zero-variance distribution using an MCMC algorithm, we can use a KDE algorithm to reconstruct an approximate zero-variance distribution $\hat{g}_M$ from these samples. With the approximate zero-variance distribution $\hat{g}_M$, we can then produce an importance sampling estimate of the recourse function by generating $N$ additional samples $\xi_1, \ldots, \xi_N$ from $\hat{g}_M$. As we can now specify both the original distribution $f$ and the distribution that we used to generate these samples $\hat{g}_M$, we can evaluate the likelihood of each sample as,

$$\hat{\Lambda}(\xi) = \frac{f(\xi)}{\hat{g}_M(\xi)}$$  \hspace{1cm} (3.6)

and form the importance sampling estimate as,

$$\hat{Q}(\hat{x}) = \frac{1}{N} \sum_{i=1}^{N} Q(\hat{x}, \xi_i) \Lambda(\xi_i)$$  \hspace{1cm} (3.7)

The samples $\xi_1, \ldots, \xi_N$ will not originate from the true zero-variance distribution $g^*$. Nevertheless, they can still be used to produce an effective importance sampling estimate provided that the KDE algorithm is able to construct a $\hat{g}_M$ that is similar to $g^*$. Given that the importance sampling distribution $\hat{g}_M$ will only be an approximation of the zero-variance distribution $g^*$, it follows that the estimates that are produced through this framework will typically not have zero variance and zero sampling error. Nevertheless, the process of generating samples in regions that contribute most to the value of the recourse function will still lead to importance sampling estimates with low-variance and low-sampling error.

We note that it is possible to generate samples using an MCMC algorithm and then directly construct an importance sampling estimate of the recourse function by using a self-normalized importance sampling scheme, or by using a KDE algorithm to estimate the probability $g^*(\xi_i)$ for $i = 1, \ldots, M$. We note, however, that resampling from the approxi-
mate zero-variance distribution \( \hat{g}_M \) ultimately produces more accurate importance sampling estimates. This is due to the fact that importance sampling estimates are highly sensitive to the likelihood of each sample, and the resampling process allows us exactly determine the likelihood of each sample used to construct the importance sampling estimate. In contrast, the use of an MCMC-only approach will approximate the likelihood of each sample and ultimately produce a biased importance sampling estimate. Generating samples from \( \hat{g}_M \) is also beneficial in that the samples are independent and the kernel functions are easy to sample from, especially when compared to the computational overhead involved in the MCMC sampling process. In practice, we therefore construct \( \hat{g}_M \) using modest values of \( M \) and then construct an importance sampling estimate \( \hat{Q}(\hat{x}) \) using large values of \( N \).

3.3 MCMC-IS in Practice

In this section, we present a simple implementation of MCMC-IS that can be used to generate importance sampling estimates with low sampling error. Our simple implementation uses a Random Walk Metropolis-Hastings algorithm to generate samples from the zero-variance distribution, and a Gaussian product kernel and leave-one-out cross validation bandwidth estimator to construct the approximate zero-variance distribution. We include a step-by-step explanation of how to generate an importance sampling estimate using this implementation in Algorithm 1.

The Metropolis-Hastings algorithm uses a simple accept-reject procedure in order to generate a Markov chain that has (3.1) as its stationary distribution. In the \( k \)-th step, the algorithm generates a proposed sample \( \zeta_k \) using a proposal distribution \( q(\cdot | \xi_k) \), which typically depends on the current sample \( \xi_k \). Together, the proposed sample, the current sample and the target distribution are used to evaluate an acceptance probability, \( a(\xi_k, \zeta_k) \). The proposed sample is accepted with probability \( a(\xi_k, \zeta_k) \), in which case the Markov Chain transitions to the proposed sample \( \xi_{k+1} := \zeta_k \). Otherwise, the proposed sample is rejected with probability \( 1 - a(\xi_k, \zeta_k) \), in which case the Markov chain remains at its current sample \( \xi_{k+1} := \xi_k \).

In our simple implementation, we choose to a random walk process to propose the samples
in the Metropolis-Hastings algorithm. This implies that the proposed sample \( \zeta_k \) is generated as,

\[
\zeta_k = \xi_k + v_k
\]

(3.8)

where \( v_k \) is a Gaussian random variable with mean 0 and covariance matrix \( \Sigma \). When new samples are proposed through a random walk process, the proposal distribution is symmetric and the acceptance probability can be expressed as,

\[
a(\xi_k, \zeta_k) = \min \left\{ \frac{|Q(\hat{x}, \zeta_k)|f(\zeta_k)}{|Q(\hat{x}, \xi_k)|f(\xi_k)}, 1 \right\}
\]

(3.9)

In terms of KDE algorithm, we use a Gaussian product kernel function,

\[
K_H(\xi, \xi_i) = \prod_{k=1}^{D} \frac{1}{\sqrt{2\pi h_k}} \exp \left( -\frac{(\xi_k - \xi_{i,k})^2}{2h_k^2} \right)
\]

(3.10)

where the bandwidth matrix \( H \) is a \( D \times D \) diagonal matrix that contains the bandwidth parameters of each dimension \( h_1, \ldots, h_D \) along its diagonal. In this case, we use a one-dimensional leave-one-out cross validation estimator to estimate the value of the bandwidth parameter \( h_k \) separately for each dimension \( k \). The exact parameters for this bandwidth estimator are defined in Table 3.1.

We note that we use this simple MCMC-IS implementation to generate the majority of the numerical results in Chapters 4 and 5 because it is straightforward to implement and does not depend on a restrictive set of assumptions. The quality of numerical results that we achieve with this admittedly simple implementation in these chapters only reinforces the potential of MCMC-IS, as more efficient implementations of MCMC-IS would only further increase the advantages of our framework. It is true that this simple implementation can also lead to certain challenges in practice; we refer to these challenges throughout Chapter 4 and provide recommendations to fix them by using different MCMC and KDE algorithms.
**Algorithm 1** Markov Chain Monte Carlo Importance Sampling (MCMC-IS)

**Require:**
- $\hat{x}$: previous stage decision
- $M$: number of samples to generate using the MCMC algorithm
- $N$: number of samples to generate using the approximate zero-variance distribution
- $\xi_0$: starting sample for the MCMC algorithm
- $q(\cdot | \xi_k)$: proposal distribution for the MCMC algorithm
- $K_H$: kernel function for the KDE algorithm
- $H$: bandwidth matrix for the KDE algorithm

**Step 1:** Generate Samples from the Zero-Variance Distribution using MCMC

1. Set $k = 0$
2. Given the current sample $\xi_k$, generate $\zeta_k \sim q(\cdot | \xi_k)$.
3. Generate a uniform random variable $u \sim U \in (0, 1)$.
4. Transition to the next sample according to,
   \[
   \xi_{k+1} = \begin{cases} 
   \zeta_k & \text{if } u \leq a(\xi_k, \zeta_k) \\
   \xi_k & \text{otherwise}
   \end{cases}
   \]
   where,
   \[
   a(\xi_k, \zeta_k) = \min \left\{ \frac{|Q(\hat{x}, \zeta_k)||f(\zeta_k)q(\xi_k|\zeta_k)}{|Q(\hat{x}, \xi_k)||f(\xi_k)q(\zeta_k|\xi_k)} , 1 \right\}
   \]
5. Let $k \leftarrow k + 1$. If $k = M$ then proceed to Step 6. Otherwise return to Step 2.

**Step 2:** Reconstruct an Approximate Zero-Variance Distribution using KDE

6. For each sample generated from MCMC, reconstruct the approximate zero-variance distribution as,
   \[
   \hat{g}_M(\xi) = \frac{1}{M} \sum_{i=1}^{M} K_H(\xi, \xi_i)
   \]

**Step 3:** Resample from the Approximate Zero-Variance Distribution to Form an Importance Sampling Estimate

7. Generate $N$ new samples from $\hat{g}_M$ and form the importance sampling estimate,
   \[
   \hat{Q}(\hat{x}) = \frac{1}{N} \sum_{i=1}^{N} \frac{Q(\hat{x}, \xi_i)f(\xi_i)}{\hat{g}_M(\xi_i)}
   \]
3.4 MCMC-IS in Theory

The convergence properties of MCMC-IS depend on two sources of error: the first is due to the MCMC algorithm used to generate samples from the zero variance distribution; the second is due to the KDE algorithm used to construct the approximate zero-variance distribution.

The main convergence condition in terms of the MCMC algorithm requires that the samples generated by the MCMC algorithm form a Markov chain whose stationary distribution is the zero-variance distribution. This requires the underlying Markov chain in the MCMC process to be irreducible and aperiodic. The irreducibility property means that the chain can eventually reach any subset of the space from any state. The aperiodicity property means that the chain cannot return to a subset of the space in a predictable manner. Formal definitions of these properties can be found in [33]. The first step in the convergence analysis is to show that these two conditions are satisfied.

In order to control the error due to the KDE algorithm, we need to ensure that the number of samples are generated by the MCMC algorithm, $M$, is large enough, and that the bandwidth, $h_k$, is small enough. In particular, if $(Mh^D)^{-1} \to \infty$, $h \to 0$ as $M \to \infty$, and the distribution is approximated as,

$$\hat{g}_M(\xi) = \frac{1}{M} \sum_{i=1}^{M} K_H(\xi, \xi_i) = (Mh^D)^{-1} \frac{1}{M} \sum_{i=1}^{M} K \left( \frac{\xi - \xi_i}{h} \right)$$  \hspace{1cm} (3.11)

then it has been shown that $\hat{g}_M$ will probabilistically converge to $g^*$ under the total variation norm in [5]. Applying this result to MCMC-IS is not straightforward. The complexity stems from the fact that previous convergence proofs for the KDE algorithm assume that samples are generated from $g^*$, whereas in our framework these samples are generated from a Markov chain whose stationary distribution is $g^*$.

A final issue that may affect the convergence of MCMC-IS is the fact that the samples generated through the MCMC algorithm are typically correlated, while the samples used in many treatments of KDE algorithms assume that the samples used to construct a KDE distribution are independent. Our numerical experiments from Chapter[4] suggest that
MCMC-IS can converge even when there is a degree of correlation between MCMC samples. However, we note that there is some theoretical evidence that KDE algorithms do not necessarily require the samples independence between the samples. In particular, theoretical results in [15] suggest that KDE algorithms can construct accurate empirical distributions using correlated samples so long as they use a different bandwidth estimator. In cases where this bandwidth estimator is not available, we note that authors of [15] also state that a leave-one-out bandwidth estimator may provide an adequate approximation.
Chapter 4

Numerical Experiments on Sampling Properties

In this chapter, we illustrate the sampling properties of MCMC-IS using numerical experiments based on a Newsvendor model. We begin by introducing a simple Newsvendor model with two random variables and two stages (Section 4.1), and explain how it can be extended to include multiple random variables (Section 4.1.2) and multiple time periods (Section 4.1.3). We then use this model to illustrate how the importance sampling distribution produced by MCMC-IS can sample from important regions of the recourse function (Section 4.3). Next, we demonstrate how the number of MCMC samples used in MCMC-IS can affect the error in MCMC-IS estimates (Section 4.4), and provide insight as to how this relationship scales according to the number of random dimensions in the recourse function (Section 4.4.1). In subsequent numerical experiments, we highlight how the acceptance rate of an MCMC algorithm is related to the accuracy and computational efficiency of MCMC-IS estimates (Section 4.5), show how to modify MCMC-IS in order to generate samples in bounded regions (Section 4.6), and examine how the choice of kernel functions and bandwidth estimators in the KDE algorithm of MCMC-IS can impact the estimates that are produced (Section 4.7). We end this chapter with a comparison between MCMC-IS and other variance reduction methods that can be applied to SP models (Section 4.8).
4.1 The Newsvendor Model

4.1.1 A Simple Two-Stage Model

The test problem in our numerical experiments in Chapters 4 and 5 is a two-stage Newsvendor model with uncertain demand and sales prices. The first-stage decision-making problem in our model is a linear program defined as,

\[
  z^* = \min_{x} \quad x + Q(x) \\
  \text{s.t. } x \geq 0
\]

and the recourse function is the expected value of the linear program defined as,

\[
  Q(\hat{x}, \xi) = \min_{y_1, y_2} \quad -p(\xi_2)y_1 - ry_2 \\
  y_1 \leq d(\xi_1) \\
  y_1 + y_2 \leq \hat{x} \\
  y_1, \ y_2 \geq 0
\]

In (4.2), \(\hat{x}\) is a scalar that represents the quantity of newspapers purchased in the first stage, \(r = 0.10\) is a scalar that represents the price of recycling unsold newspapers, and \(\xi = (\xi_1, \xi_2)\) is a two-dimensional random vector that represents the uncertainty in demand \(d(\xi_1)\) and sales price \(p(\xi_2)\) of newspapers.

In our numerical experiments, we investigate the sampling properties of MCMC-IS by pairing this model with three separate distributions: a lower-variance lognormal distribution, a higher-variance lognormal distribution, and a multimodal rare-event distribution. The parameters used to generate the demand and sales price for these distributions are specified in Table 4.1.

4.1.2 A Multidimensional Extension

The \(D\)-dimensional Newsvendor model is a multidimensional extension of the Newsvendor model specified in Section 4.1.1. In this extension, we consider a problem where the
<table>
<thead>
<tr>
<th>Distribution</th>
<th>Lower-Variance Lognormal</th>
<th>Higher-Variance Lognormal</th>
<th>Multimodal Rare-Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>((\xi_1, \xi_2))</td>
<td>N(0, (1^2 \times I_2))</td>
<td>N(0, (2^2 \times I_2))</td>
<td>N(0, (1^2 \times I_2))</td>
</tr>
</tbody>
</table>
| \(d(\xi_1)\) | 100 \(\exp(\xi_1)\) | 100 \(\exp(\xi_1)\) | 100 \(\exp\left(\frac{\xi_1^2}{2} - \frac{(\xi_1+3)^2}{8}\right)\)
| | | | \(100 \exp\left(\frac{\xi_1^2}{2} - \frac{(\xi_1+1)^2}{8}\right)\) |
| \(p(\xi_2)\) | 1.50 \(\exp(\xi_2)\) | 1.50 \(\exp(\xi_2)\) | 1.50 \(\exp\left(\frac{\xi_2^2}{2} - \frac{(\xi_2+3)^2}{8}\right)\)
| | | | \(1.50 \exp\left(\frac{\xi_2^2}{2} - \frac{(\xi_2+1)^2}{8}\right)\) |

Table 4.1: Parameters of demand and sales price distributions for the Newsvendor model.

Newsvendor has to sell \(D\) different types of newspapers which have the same demand and sales price distribution. The recourse function of the \(D\)-dimensional Newsvendor model can be expressed as,

\[
Q_D(\hat{x}) = \sum_{i=1}^{D} Q_i(\hat{x}_i)
\]

where \(Q_i(\hat{x}_i)\) denotes the 2-dimensional recourse function used to represent the expected future costs associated with a single type of newspaper as in (4.2).

### 4.1.3 A Multistage Extension

The \(T\)-stage Newsvendor model is a multistage extension of the Newsvendor model specified in Section 4.1.1. In this extension, we consider a problem where the Newsvendor buys and sells a single type of newspapers over \(T\) consecutive days. We assume that any newspapers that are to be sold on day \(t + 1\) have to be bought on day \(t\), and that any unsold newspapers at the end of day \(t + 1\) have to be recycled at a price of \(r\). These assumptions allow us to extrapolate the true values of optimal solution \(x^*\) and optimal cost \(z^*\) for a \(T\)-stage Newsvendor model from their corresponding values for the Newsvendor model specified in Section 4.1.1. In particular, we reason that the optimal cost \(z^*\) scales additively with the
number of time periods, and the optimal solution $x^*$ remains the same.

4.2 Experimental Setup

4.2.1 Experimental Statistics

The advantages of using the two-stage, two-dimensional Newsvendor model in Section 4.1.1 are that the relevant distributions can be easily visualized, and that we can determine the true value of the recourse function at different values of $\hat{x}$ using state-of-the-art numerical integration procedures. In turn, we can also determine the true values of the recourse function for the multidimensional and multistage extensions of the Newsvendor problem that are specified in Sections 4.1.2 and 4.1.3.

By knowing the true values of the recourse function for these SP models, we can consider both the error and the variance of the estimates that are produced in our numerical experiments. Table 4.2 provides an overview of the different statistics that we report in Sections 4.3 - 4.8. We note that the statistics in these sections have been generated using $R = 100$ repetitions, and have been normalized for clarity. We note that all the results for the MCMC-IS method have been generated using $\frac{M}{\gamma} + N$ functional evaluations as we explain further in Section 4.3 where $\gamma$ denotes the acceptance rate of the MCMC algorithm in MCMC-IS.

4.2.2 Implementation

Table 4.3 summarizes the different sampling methods that refer to in Sections 4.3 - 4.5. Unless otherwise stated, we produced all results for these methods in MATLAB 2012a. In particular, we used a built-in Mersenne-Twister algorithm to generate the uniform random numbers for the CMC, LHS, IDG and MCMC-IS sampling methods. Similarly, we used built-in Owen and Reverse-Radix scrambling algorithm to randomize the sequences that we generated for Sobol QMC and Halton QMC methods. Most of the results for MCMC-IS were generated using the simple implementation described in Section 3.3. We built all approximate importance sampling distributions for MCMC-IS using the MATLAB KDE Toolbox, which is available online at [http://www.ics.uci.edu/~ihler/code/kde.html](http://www.ics.uci.edu/~ihler/code/kde.html).
<table>
<thead>
<tr>
<th>Statistic</th>
<th>Formula</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSE((\hat{Q}))</td>
<td>[\sqrt{\frac{1}{R} \sum_{i=1}^{R} (Q(\tilde{x}) - \hat{Q}(\tilde{x})_i)^2}]</td>
<td>mean-squared error of (R) estimates of the value of the recourse function (\hat{Q}) at (\tilde{x})</td>
</tr>
<tr>
<td>SE((\hat{Q}))</td>
<td>[\frac{1}{R} \sum_{i=1}^{R} \frac{1}{N} \frac{1}{N-1} \left( \frac{\hat{Q}(\tilde{x})<em>i - \frac{1}{N} \sum</em>{j=1}^{N} Q(\tilde{x}, \xi_j)}{N-1} \right)^2]</td>
<td>mean of (R) estimates of standard error in the value of the recourse function (\hat{Q}) at (\tilde{x})</td>
</tr>
<tr>
<td>MSE((\hat{g}))</td>
<td>[\frac{1}{R} \sum_{i=1}^{R} \frac{1}{N} \sqrt{(g(\xi_i) - \hat{g}(\xi_i))^2}]</td>
<td>mean-squared error of (R) approximations of the zero-variance distributions at (\tilde{x}); the (\xi_i)s are specified by a 100 × 100 grid on (\Xi)</td>
</tr>
</tbody>
</table>

Table 4.2: Sampling statistics reported in Chapter 4

### 4.3 Sampling from the Important Regions

Importance sampling is most effective when an importance sampling distribution can generate samples from regions that contribute the most to the value of the recourse function. Such regions are referred to as the important regions of the recourse function, and occur at points where \(|Q(\tilde{x}, \xi)||f(\xi)|\) attains large values. Accordingly, the major difference between MCMC-IS and other MC methods is that MCMC-IS generates samples at important areas of the recourse function using the importance sampling distribution \(\hat{g}_M\).

We illustrate this difference by plotting the location of the samples that are used to estimate the recourse function \(Q(\tilde{x})\) at \(\tilde{x} = 50\) for a Newsvendor model assuming a lower-variance lognormal distribution in Figure 4-1, and Newsvendor model paired with a multimodal rare-event distribution in Figure 4-2.

As shown in Figures 4-1 and 4-2, the samples that are generated using the MCMC-IS importance sampling distribution \(\hat{g}_M\) are located at important regions of the recourse function, where \(|Q(\tilde{x}, \xi)||f(\xi)|\) attains high values. In contrast, the samples generated using
<table>
<thead>
<tr>
<th>Method</th>
<th>Acronym</th>
<th>Variance Reduction Strategy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude Monte Carlo</td>
<td>CMC</td>
<td>None</td>
</tr>
<tr>
<td>Sobol Sequence</td>
<td>Sobol QMC</td>
<td>Quasi-Monte Carlo</td>
</tr>
<tr>
<td>with Owen Scrambling</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Halton Sequence</td>
<td>Halton QMC</td>
<td>Quasi-Monte Carlo</td>
</tr>
<tr>
<td>with Reverse Radix Scrambling</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Latin Hypercube</td>
<td>LHS QMC</td>
<td>Stratified Sampling</td>
</tr>
<tr>
<td>Infanger-Dantzig-Glynn Importance Sampling</td>
<td>IDG</td>
<td>Importance Sampling</td>
</tr>
<tr>
<td>MCMC Importance Sampling with Metropolis Hastings Sampler</td>
<td>MCMC-IS</td>
<td>Importance Sampling</td>
</tr>
<tr>
<td>MCMC Importance Sampling with Adaptive Metropolis Sampler</td>
<td>MCMC-IS AM</td>
<td>Importance Sampling</td>
</tr>
<tr>
<td>MCMC Importance Sampling with Hit-and-Run Sampler</td>
<td>MCMC-IS HR</td>
<td>Importance Sampling</td>
</tr>
</tbody>
</table>

Table 4.3: Sampling methods covered in Chapter 4.
Figure 4-1: Location of samples produced by MCMC-IS and CMC for a Newsvendor model paired with a lower-variance lognormal distribution.

Figure 4-2: Location of samples produced by MCMC-IS and CMC for a Newsvendor model paired with a multimodal rare-event distribution.
CMC are located in regions where the original distribution $f(\xi)$ attains high values. The contours in these figures demonstrate how $f(\xi)$ and $|Q(\hat{x}, \xi)|f(\xi)$ may not only have different shapes but may also be centered around different points.

The fact that MCMC-IS can generate samples in these regions for both distributions demonstrates the adaptive nature of our framework. This property is especially important in the context of decomposition algorithms because the important areas of the recourse function $Q(\hat{x})$ can vary substantially according to the previous-stage decision $\hat{x}$.

### 4.4 The Required Number of MCMC Samples

In Figures 4-3(a) and 4-3(b), we show how the error in the approximate zero-variance distribution $\hat{g}_M$ and the error in the recourse function estimate $\hat{Q}$ changes according to the number of MCMC samples, $M$, used in MCMC-IS. In this case, we consider estimates of the recourse function $\hat{Q}(\hat{x})$ at $\hat{x} = 50$ for a Newsvendor model assuming a lower-variance lognormal distribution, and we construct each estimate using $N = 16000$ samples generated from the approximate zero-variance distribution $\hat{g}_M$.

As shown, increasing $M$ reduces the error in the approximate importance sampling distribution $\hat{g}_M$ and subsequently also reduces the error in resulting importance sampling estimates of the recourse function $\hat{Q}$. Although the convergence in the density error in $\hat{g}_M$ and the sampling error in $\hat{Q}$ appears to be steady, we note that increasing the number of MCMC samples is often hard to justify, as these graphs show that we can obtain estimates with error rates of less than 1% using only $M = 1000$ samples in the MCMC step and $N = 16000$ samples in the resampling step. This is a positive result as the MCMC algorithm represents the most computationally expensive part of our framework.

A possible explanation for this empirical observation is that if the approximate zero-variance distribution $\hat{g}_M$ qualitatively agrees with the true zero-variance distribution $g^*$, then the estimates produced using $\hat{g}_M$ should have properties that are similar to the estimates produced using $g^*$. In Figure 4-4(a), we provide evidence that $\hat{g}_M$ qualitatively agrees with $g^*$ by plotting its contours for different values of $M$.  

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4.4.1 The Curse of Dimensionality

In this section, we examine how the required number of MCMC samples $M$ changes as we increase the number of random variables $D$ in the recourse function $Q(\hat{x})$. In particular, we consider the recourse function $\hat{Q}$ of a $D$-dimensional Newsvendor model from Section 4.1.2 where the Newsvendor sells $D_2$ different types of newspapers. We use a lower-variance lognormal distribution to specify the demand and sales price for each type of newspaper, and estimate the recourse function at $\hat{x} = 50e$ where $e$ is a $D_2 \times 1$ vector of ones.

As we describe in Section 4.1.2, the recourse function of the $D$-dimensional Newsvendor model can be expressed as the sum of $\frac{D}{2}$ recourse functions of our 2-dimensional Newsvendor model (note that we only consider even multiples of $D$ in our experiment). This implies that the true standard deviation of the recourse function in our $D$-dimensional model is $\sqrt{\frac{D}{2}}$ larger than the true standard deviation of the recourse function in our 2-dimensional model. We account for this difference in our experiment by scaling the number of samples $N$ used construct the recourse function estimate by $\sqrt{\frac{D}{2}}$. In addition, we also account for the fact that we will require more samples to construct an approximate zero-variance distribution for a $D$-dimensional recourse function by scaling the number of MCMC samples $M$ in proportion to $D$.

Our results in Figure 4.4.1 demonstrate how the error in our estimates increases with the
Figure 4-4: Contours of $g^*$ (a) and $\hat{g}_M$ for different values of $M$ (b)-(d).
number of random variables in the recourse function, $D$. These results suggest that scaling $M$ in proportion to $D$, and scaling $N$ in proportion to the standard deviation of the recourse function results in a steady increase in the error rate in our estimates. Accordingly, we can see that we require even higher values of $N$ or $M$ to maintain the same error rate across dimensions.

In theory, this trend reflects the well-known curse of dimensionality, in which a random space $\Xi = \mathbb{R}^D$ becomes more exponentially more voluminous as $D$ increases. The curse of dimensionality specifically affects the efficiency of an MCMC algorithm by making it harder to generate samples that span across the entire random space, $\Xi$. The curse of dimensionality also affects the efficiency of a KDE algorithm by requiring a larger set of samples across the entire random space, $\Xi$.

### 4.5 The Acceptance Rate of the MCMC Algorithm

Although MCMC algorithms are typically run until a fixed number of samples have been proposed, we recommend running the MCMC algorithm in MCMC-IS until a fixed number of samples have been accepted. In our experiments, we find that the number of accepted samples is a better metric to assess the quality of the importance sampling distribution.
produced by KDE algorithm. Nevertheless, tracking the number of accepted samples may
make it more difficult to assess the computational burden of MCMC-IS, especially when we
use an MCMC algorithm that relies on an accept-reject procedure, such as the Metropolis-
Hastings algorithm, and we do not know the acceptance rate of the MCMC algorithm a
priori. In such cases, the true computational cost of the MCMC-IS algorithm depends on
the acceptance rate of the MCMC algorithm, and the total number of functional evaluations
in a single MCMC-IS run is \( \frac{M}{\gamma} + N \) where \( \gamma \) denotes the acceptance rate of the MCMC
algorithm.

In the simple MCMC-IS implementation that we present in Section 3.3, we use a Metropolis-
Hastings algorithm where samples are proposed using a random walk process and accepted
through a probabilistic accept-reject procedure. In this case, the acceptance rate of the
MCMC algorithm is related to the magnitude of the covariance of the random-walk process,
which we denote using the \( D \times D \) matrix \( \Sigma_{RW} \). In situations where the magnitude of \( \Sigma_{RW} \) is
too small, then the random walk process will propose samples that are close to the current
sample and that have a high likelihood of being accepted. Although the resulting high accep-
tance rate will reduce the computational burden of the MCMC algorithm in MCMC-IS, the
accuracy of the MCMC-IS estimate will suffer will because the KDE algorithm will produce
an importance sampling distribution from samples that are located over too small a portion
of the random space. Conversely, in situations when the the magnitude of \( \Sigma_{RW} \) is too large,
then the random walk process will propose samples that are far from the current sample
and that have a lower likelihood of being accepted. In such cases, the resulting MCMC-IS
estimate is likely to be accurate, but the computational burden of the MCMC algorithm in
MCMC-IS will be large.

We demonstrate these issues by using a numerical experiment in which we vary the step-
size parameter of the random-walk process used to propose new samples in the Metropolis
Hastings sampler and examine the error and resulting acceptance rate. We model the covari-
ance of the random walk process as \( \Sigma_{RW} = s\Sigma \), where \( \Sigma \) denotes the underlying variance of
the target distribution and \( s \) denotes the step-size of the random walk distribution. In our
experiment, we consider estimates of the recourse function \( \hat{Q}(\hat{x}) \) at \( \hat{x} = 50 \) for a Newsvendor
model that has been paired with a lower-variance lognormal distribution, a higher-variance
lognormal distribution and a multimodal rare-event distribution. We examine estimates that have been produced using a random-walk Metropolis Hastings algorithm using the theoretically optimal step-size for a Gaussian distribution of \( s = 2.4D^{-1} = 2.83 \), as well a step-size that is half this value \( s = 1.42 \), and a step-size that is twice this value \( s = 5.66 \). In addition, we also consider estimates that are were produced when \( \Sigma_{RW} \) is entirely determined through an automated process that uses the covariance of the chain after a burn-in phase.

In Table 4.5, we demonstrate the trade-off between the step-size parameter and the acceptance rate of an MCMC algorithm for a lower-variance lognormal distribution, a higher-variance lognormal distribution and a multimodal rare-event distribution. Our results suggest that the acceptance rate of the MCMC algorithm can change substantially with the step-size of the random walk process. As expected, lower step-sizes produce high acceptance rates while higher step-sizes will produce lower acceptance rates. We note that the acceptance rates for the lower-variance and higher-variance lognormal distributions are similar for different values of \( s \); this is because we have scaled the covariance of the random walk process according to the inherent variance of the model \( \Sigma \).

<table>
<thead>
<tr>
<th>Step-size</th>
<th>Lower-Variance Lognormal</th>
<th>Higher-Variance Lognormal</th>
<th>Multimodal Rare-Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.42</td>
<td>45%</td>
<td>43%</td>
<td>62%</td>
</tr>
<tr>
<td>2.83</td>
<td>32%</td>
<td>31%</td>
<td>50%</td>
</tr>
<tr>
<td>5.66</td>
<td>21%</td>
<td>19%</td>
<td>38%</td>
</tr>
<tr>
<td>Adaptive</td>
<td>35%</td>
<td>35%</td>
<td>40%</td>
</tr>
</tbody>
</table>

Table 4.4: Acceptance rates and step-sizes of MCMC algorithms used in MCMC-IS.

In Figure 4.5, we illustrate the how the error of MCMC-IS estimates changes according to the step-size \( s \) for different distributions. In this case, we form all estimates using \( N = 16000 \) samples. At first glance, these results may suggest that the step-size parameter does not affect the accuracy of the MCMC-IS estimates, especially for the lower-variance and higher-variance lognormal distributions. However, this is misleading because the true number of
functional evaluations required to produce these errors are \( \frac{M}{T} + N \). When we account for the
difference in acceptance rates as shown in Table 4.5, we notice that the estimates produced at
\( s = 5.66 \) require around 30% to 50% more functional evaluations than the estimates produced
at \( s = 2.83 \), and up to 75% more functional evaluations than the estimates produced with
the Adaptive Metropolis algorithm. Accordingly, the estimates produced at \( s = 1.43 \) may
appear to require fewer functional evaluations. However these estimates may also exhibit
much higher error rates as is the case for the rare-event distribution in Figure 4-6(c). In
general, our results suggest that using the optimal step-size for Gaussian distributions can
yield accurate and computationally efficient estimates, and that the Adaptive Metropolis
algorithm can achieve both these goals in the context of general SP models.

### 4.6 Sampling from Bounded Regions

One unaddressed issue with the simple MCMC-IS implementation that we present in Section
3.3 is that neither the MCMC algorithm nor the KDE algorithm is designed to work with
random variables that have bounded support. In turn, MCMC-IS may not produce accurate
results whenever the recourse function \( Q = \mathbb{E}[Q(\hat{x}, \xi)] \) depends on a random vector \( \xi \) that
is bounded within a \( D \)-dimensional space \( \Xi = [a, b]^D \).

The first limitation of the simple implementation is that the random walk proposal process
in Metropolis-Hastings algorithm will generate samples across an unbounded \( D \)-dimensional
space \( (-\infty, \infty)^D \). In situations where the random walk is not likely to breach the boundaries
then we may simply condition the algorithm to reject any proposed sample that ventures
outside of the boundaries. In situations where the random walk is likely to breach the
boundaries frequently, however, such an approach can result in an excessive number of points
near the bounds and introduces a bias in the resulting importance sampling distribution. In
such cases, we recommend using an MCMC algorithm that is explicitly designed to generate
samples from bounded regions, such as the Hit-and-Run algorithm from [39] and [4].

Assuming that we are able to generate points within a bounded region using an MCMC
algorithm, however, the KDE procedure may still not work properly because it is designed
to construct densities across unbounded regions. We note that these issues can arise when

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Figure 4-6: Convergence of $\hat{Q}(\hat{x})$ by to the step-size of random-walk Metropolis-Hastings MCMC algorithm for a Newsvendor model paired with a lower-variance lognormal distribution (a), a higher-variance lognormal distribution (b), and a multimodal rare-event distribution (c).
the samples used in the KDE procedure lie within the bounded region because the resulting
importance sampling distribution may attribute a positive density to values that lie outside
of the boundaries provided that the bandwidth attributed to points near the boundaries is
large enough. In such cases, the issue can be resolved by using a transformation,

\[ t : [a, b] \rightarrow (-\infty, \infty) \]  

(4.4)

that can map the components of each sample from a bounded space to an unbounded
space. Having mapped the samples onto an unbounded region, the KDE algorithm can
then construct an approximate distribution \( \hat{h} \) in the transformed space. The approximate
zero-variance distribution can be subsequently recovered using the formula:

\[ \hat{g}(\xi) = \hat{h}(t(\xi))t'(\xi) \]  

(4.5)

One frequently used transformation in such cases is the log-transformation,

\[ t(\xi) = \log \left( \frac{\xi - a}{b - \xi} \right) \]  

(4.6)

It is worth noting that this transformation only applies to random variables which are
bounded in a multidimensional box.

In Figure 4.6, we highlight the performance of MCMC-IS when it is paired with a Hit-
and-Run MCMC algorithm and KDE algorithm which uses a log-transformation. We refer to
this particular implementation of MCMC-IS implementation as MCMC-IS HR. Our results
reflect estimates of the recourse function \( \hat{Q}(\hat{x}) \) at \( \hat{x} = 50 \) for a Newsvendor model that
has been paired with a lower-variance lognormal distribution, a higher-variance lognormal
distribution and a multimodal rare-event distribution. Although the random variables within
these models are not bounded, we create artificial bounds for each random variable by using
the values at the 0.01th percentile and 99.99th percentile, respectively.

Our results in Figure 4.6 show that the MCMC-IS HR implementation for bounded
problems converges, albeit at a slower rate than the simple MCMC-IS implementation. The
difference between MCMC-IS and MCMC-IS HR can be explained by the fact that MCMC-
Figure 4-7: Convergence of $\hat{Q}(\hat{x})$ for MCMC-IS and MCMC-IS HR for a Newsvendor model paired with a lower-variance lognormal distribution (a), a higher-variance lognormal distribution (b) and a multimodal rare-event distribution (c).
HR uses a different accept-reject procedure to generate samples which yields an acceptance rate that is 40% - 60% lower than the acceptance rate of MCMC-IS. In this experiment, we set $M = 4000$ for the simple MCMC-IS implementation and run the MCMC-IS HR implementation until it has reached the same number of functional evaluations as MCMC-IS. Accordingly, MCMC-IS HR setup generates fewer accepted samples and produces a less effective importance sampling distribution through the KDE procedure.

It is true that the comparison in Figure 4.6 does not reflect the performance of MCMC-IS and MCMC-IS HR on bounded problems. However, it does offer a proof of concept that the MCMC-IS HR implementation we propose can work. Moreover, it highlights how the importance sampling framework that we propose can be paired with a wide number of MCMC and KDE algorithms.

4.7 Choosing Kernel Functions and Bandwidth Estimators in the KDE Algorithm

KDE algorithms provide some degree of flexibility in constructing the approximate zero-variance distribution $\hat{g}_M$ used in MCMC-IS by allowing us to choose the functions and bandwidth estimators with which we can construct $\hat{g}_M$. In Figure 4.7, we demonstrate how these choices can ultimately affect the error of estimates that are produced using MCMC-IS.

In Figure 4-8(a) we consider the estimates that are produced when $\hat{g}_M$ is constructed using a leave-one-out cross validation bandwidth estimator (LCV), and either a Laplacian, Gaussian or Epanetchnikov kernel function. In Figure 4-8(b) we consider the estimates that are produced when $\hat{g}_M$ is constructed using a Gaussian kernel function and either a plug-in mean integrated squared error estimator (MISE), a one-dimensional leave-one-out cross-validation estimator (LCV) or a Gaussian rule-of-thumb estimator (ROT). Further information on these kernel functions and bandwidth estimators can be found in Section 3.1.3. Our results reflect estimates of the recourse function $\hat{Q}(\hat{x})$ evaluated at $\hat{x} = 50$ for a Newsvendor model paired with a lower-variance lognormal distribution.

Our results in Figure 4.7 suggest that the error in the recourse function estimate is unaffected by the choice of the kernel function, but may be affected by the choice of bandwidth.
estimator. In particular, it appears that the use of LCV and ROT estimators generate more accurate results, better while the MISE estimator is less efficient in this respect. Our results confirm the well-known fact that the approximate distributions produced by a KDE algorithm are much more sensitive to the choice of bandwidth than the choice of a kernel function (see [35]).

4.8 Comparison to Existing Variance Reduction Techniques

4.8.1 Comparison to IDG Importance Sampling

As mentioned in Section 2.5.4, the accuracy and convergence rate of IDG estimates depends on whether the recourse function $Q(\hat{x})$ is additively separable, and whether one can determine the value of $\xi$ at which the recourse function attains its minimal value. While such assumptions can be restrictive in practice, they have also been previously documented and discussed in [7], [20] and [16]. As such, our results in this section pertain to issues that arise when we use the IDG method to generate estimates of the recourse function for SP models where the uncertainty is modelled using continuous random variables.

Given that the IDG method can only be applied to SP models with discrete random variables, we are required to represent the random space $\Xi$ as a set of discrete values $\Omega$. In
in our experiments, we construct each grid by using $N_d = 101$ points to represent the values of $\xi \in \Xi$ in each dimension. These points are evenly distributed between a set of boundaries at the $p^{th}$ and $1 - p^{th}$ percentile of the distribution for each component of $x_i$. Formally, $\Omega = \omega_1 \times \omega_2 \times \cdots \times \omega_D$ where $\omega_d = [F_d^{-1}(p), F_d^{-1}(p) + \delta_d, \ldots, F_d^{-1}(1 - p)]$ and $\delta = \frac{F_d^{-1}(1 - p) - F_d^{-1}(p)}{N_d - 1}$

In Figure 4.8.1, we show how the error in the IDG estimates change according to the value of $p$. We note that this value effectively dictates the width of the grid that we use to discretize the random space $\Xi$ in each dimension. The results in Figure 4.8.1 reflect estimates of the recourse function $\hat{Q}(\hat{x})$ evaluated at $\hat{x} = 50$, which have been constructed using $N = 16000$ samples.

![Figure 4-9: Error in IDG estimates of $\hat{Q}(\hat{x})$ for a Newsvendor model paired with a lower-variance lognormal distribution (a) and a higher-variance lognormal distribution (b). The value of $p$ determined the boundaries $\Omega$ of the grid used to represent $\Xi$; higher values of $p$ correspond to wider boundaries.](image)

Our results show that the error in the IDG estimates is effectively determined by the boundaries of the grid that we use to discretize the random space $\Xi$. More importantly, our results suggest that it is inherently difficult to predict how the error in IDG estimates of the recourse function changes according to the boundaries of the grid that we use. In Figure 4-9(a), the recourse function belongs to a Newsvendor model paired with a lower-variance distribution as shown and the IDG estimates become more accurate as we decrease the value of $p$. Conversely, in Figure 4-9(b), the recourse function belongs to a Newsvendor model paired with a higher-variance distribution and IDG estimates become less accurate as we
decrease the value of $p$. We note that this issue is not related to the discretization of $\Xi$ but the importance sampling distribution of the IDG method. In experiments where the recourse function is estimated without importance sampling, the estimates for both the lower-variance distribution and the higher-variance distribution behave consistently consistent in that their error decreases as we decrease the value of $p$.

Even in situations where we can construct a suitable grid $\Omega$ which will accurately discretize the random space $\Xi$, however, the accuracy of IDG estimates suffers as the number of random variables in the recourse function $D$ increases because it becomes computationally expensive to maintain the resolution of the grid. In Figure 4.8.1 we highlight this point by plotting the error in IDG and MCMC-IS estimates of a $D$-dimensional recourse function. In this experiment, we use $M = 1000 \times D$ to construct the importance sampling distribution in MCMC-IS, and we use the effective number of functional evaluations in the MCMC algorithm $\frac{M}{T}$ to define the resolution of the grid $\Omega$. In particular, we use $N_d = \lceil \sqrt{\frac{M}{T}} \rceil$ points to represent each component of $\xi$ so that the computational cost in building the importance sampling distribution is similar for both methods. Our estimates reflect the recourse function of a $D$-dimensional Newsvendor model paired with a lower-variance lognormal distribution which is evaluated at $\hat{x} = 50e$ where $e$ is a $\frac{D}{2} \times 1$ vector of ones.

![Figure 4-10: Error in IDG (a) and MCMC-IS (b) estimates of the recourse function $\hat{Q}(\hat{x})$ for a multidimensional Newsvendor model paired with a lower-variance lognormal distribution.](image)

Our results for this experiment clearly show that the error in IDG estimates of the
Our results reflect estimates of the recourse function $\hat{Q}(\bar{x})$ at $\bar{x} = 50$ for a Newsvendor model that has been paired with a lower-variance lognormal distribution, a higher-variance lognormal distribution and a multimodal rare-event distribution respectively. We use a MCMC-IS implementation paired with the Adaptive Metropolis algorithm to produce estimates for the multimodal rare-event distribution, and use the simple MCMC-IS implementation described in Section 3.3 otherwise. We construct the importance sampling distribution using $M = 4000$ samples in all three cases. Given that this effectively results in $\frac{M}{\gamma}$ functional evaluations for MCMC-IS, we use $\frac{M}{\gamma} + N$ samples to produce estimates for all other methods. This ensures that each estimate is produced using the same number of functional evaluations and ensures that a fair comparison from a computational perspective.

Our results in Figure 4.8.2 suggest that the relative performance of MCMC-IS increases as the underlying variance in the model increases. In particular, MCMC-IS produces estimates with a higher rate of error in the case of a lower-variance lognormal distribution, but produces estimates with lower rates of error in the cases of a higher-variance lognormal distribution and the multimodal rare-event distribution. We note that we are able to produce lower errors in the case of the lower-variance lognormal distribution when we decrease the number of MCMC
Figure 4-11: Mean-squared error and standard error in estimates of $\hat{Q}(\hat{x})$ for a Newsvendor model produced by MCMC-IS and other variance reduction techniques; the model is paired with a lower-variance lognormal distribution in (a)-(b), a higher-variance lognormal distribution in (c)-(d), and a multimodal rare-event distribution in (e)-(f).
samples \( M \). In terms of other methods, the results in Figure 4.8.2 suggest that QMC methods tend to outperform other variance reduction techniques and that Sobol QMC consistently outperform Halton QMC. In comparison, LHS and CMC methods tend to produce estimates that converge at similar rates, except in the case of the multimodal rare-event distribution where LHS tends to perform even worse than CMC.

The fact that the standard error of MCMC-IS estimates is low for all three distributions suggests that the error in our estimates is primarily due to bias. This bias is attributable to the likelihood ratio in the importance sampling estimator and can probably be reduced by fine-tuning the KDE algorithm that is used to construct the importance sampling distribution in MCMC-IS. The mean-squared error and standard error of estimates tend to match for all methods except for Sobol QMC and Halton QMC, for which the standard error of estimates exceeds their true error. This reflects a key problem with QMC approaches in that they are able to substantially reduce the sampling error in an estimate, but fail to provide an accurate estimate of the sampling error itself.
Chapter 5

Numerical Experiments on Performance in Decomposition Algorithms

In this chapter, we illustrate the performance of MCMC-IS when it is embedded in a decomposition algorithm and used to solve variants of the Newsvendor model described in Section 4.1. We begin by showing that MCMC-IS can produce accurate solutions for SP models when it is paired with a decomposition algorithm (Section 5.2). Next, we demonstrate how MCMC-IS estimates can improve the performance of stopping tests that are used to assess the convergence in decomposition algorithms (Section 5.3). Lastly, we illustrate the computational benefits of using MCMC-IS in multistage models and demonstrate that a sampling-based approach can avoid the exponential growth in problem size that occurs when scenario trees or discretization methods are used to model the uncertainty across multiple time periods (Section 5.4).
5.1 Experimental Setup

5.1.1 Experimental Statistics

As we mention in Section 4.2.1, the advantages of using the simple two-stage, two-dimensional Newsvendor model in Section 4.1.1 is that we can determine the value of true recourse function at various points using numerical integration procedures. This allows us to calculate the true value of the recourse function at different values of $\hat{x}$, and therefore allows to examine statistics such as the mean-squared error of any quantities that relate to the recourse function, including the optimal cost of the model.

Table 5.1 provides an overview of the different statistics that we examine in Sections 5.2 - 5.4. We note that the statistics that we in these sections have been generated using $R = 30$ repetitions, and have also been normalized by their true values for clarity. Moreover, we note that that all the results for MCMC-IS have been generated using $M_\alpha + N$ functional evaluations as we explain in Section 4.5.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>Formula</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{MSE}(\hat{z})$</td>
<td>$\sqrt{\frac{1}{R} \sum_{i=1}^{R} (z^* - \hat{z}_i)^2}$</td>
<td>mean-squared error of $R$ estimates of the optimal cost $\hat{z}$ of the SP</td>
</tr>
<tr>
<td>$\text{SE}(\hat{z})$</td>
<td>$\frac{1}{R} \sum_{i=1}^{R} \frac{1}{N} \frac{1}{N-1} \left( \frac{\hat{Q}(\hat{x})<em>i - \frac{1}{N} \sum</em>{j=1}^{N} Q(\hat{x}, \xi_j)}{2} \right)$</td>
<td>mean of $R$ estimates of the standard error in the estimated optimal cost $\hat{z}$ of the SP</td>
</tr>
<tr>
<td>$\text{MSE}(\hat{x})$</td>
<td>$\sqrt{\frac{1}{R} \sum_{i=1}^{R} (z^* - \hat{x}_i)^2}$</td>
<td>mean-squared error of $R$ estimates of the optimal solution $\hat{x}$ of the SP</td>
</tr>
</tbody>
</table>

Table 5.1: Sampling statistics reported in Chapter 5.
5.1.2 Implementation

Table 5.2 summarizes the different sampling methods that refer to in Sections 5.2 - 5.4. Unless otherwise stated, we produced all results in this section in MATLAB 2012a. In particular, we used a built-in Mersenne-Twister algorithm to generate the uniform random numbers for the CMC and MCMC-IS methods. Similarly, we used a built-in Reverse-Radix scrambling algorithm to randomize the sequences that we generated for Sobol QMC methods. Most of the results for MCMC-IS were generated using the simple implementation described in Section 3.3. We built all approximate importance sampling distributions for the MCMC-IS method using the MATLAB KDE Toolbox, which is available online at [http://www.ics.uci.edu/~ihler/code/kde.html](http://www.ics.uci.edu/~ihler/code/kde.html). Lastly, all SP models in this section using a MATLAB implementation of the SDDP algorithm, where use a MEX file in order to setup and solve linear programs with the IBM ILOG CPLEX 12.4 Callable Library. We note that we do not consider results for LHS, Halton QMC, IDG, MCMC-IS HR sampling methods for the experiments in this chapter for clarity. As we point out in Sections 4.8.2, these methods consistently generate less accurate estimates than Sobol QMC and MCMC-IS, except in the case of the IDG distribution, which suffers from systematic issues that we highlight in Sections 2.5.4 and 4.8.1.

<table>
<thead>
<tr>
<th>Method</th>
<th>Acronym</th>
<th>Variance Reduction Strategy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude Monte Carlo</td>
<td>CMC</td>
<td>None</td>
</tr>
<tr>
<td>Sobol Sequence with Owen Scrambling</td>
<td>QMC</td>
<td>Quasi-Monte Carlo</td>
</tr>
<tr>
<td>MCMC Importance Sampling with Metropolis-Hastings Sampler</td>
<td>MCMC-IS</td>
<td>Importance Sampling</td>
</tr>
</tbody>
</table>

Table 5.2: Sampling methods covered in Chapter 5.
In our first experiment, we compare the the error of the estimated optimal solution \( \tilde{x} \) and the estimated optimal cost \( \tilde{z} \) that are produced when the sampled cuts in an SDDP algorithm are constructed using the MCMC-IS, CMC and QMC methods. Our test problem in this experiment is the \( D \)-dimensional News-vendor model described in Section 4.1.2. In contrast to the experiments in Chapter 4, the accuracy of \( \tilde{z} \) depends on the number of sampled cuts that are added to the first-stage problem through a decomposition algorithm, as well as the sampling method that is used to generate these estimates.

Note that in our implementation of SDDP we count the number of iterations by the number of cuts added to the first stage problem. In practice, the number of iterations needed for the algorithm to converge is determined by a stopping test that is designed to assess whether the decomposition algorithm has converged. In this experiment, however, we compare estimates that are produced after a fixed number of iterations. Fixing the number of iterations ensures that each sampling method produces estimates using the same number of samples, and isolates the performance of the sampling method from the performance of the stopping test, which we later examine in Section 5.3. In particular, we determine the number of iterations to add to the first-stage problem by using the number of iterations that are required for a deterministic version of the problem to converge. In this case, we find that a deterministic version of the News-vendor problem with \( \frac{D}{2} \) different types of newspapers require \( 8 \times \frac{D}{2} \) iterations to converge to the solution.

In Figure 5.2, we show the convergence of these estimates that we obtain when we solve a two-stage News-vendor problem with \( D = 6 \) random variables after \( 8 \times \frac{6}{2} = 24 \) cuts have been added to the first-stage problem. In Figures 5-1(a)-5-1(d), we model the uncertainty in the demand and sales price of each newspaper using the lognormal distributions, and we build the approximate zero-variance distribution for each sampled cut using the simple MCMC-IS implementation with \( M = 3000 \) samples. In Figures 5-1(a)-5-1(f), we model the uncertainty in the demand and sales price of each newspaper using the multimodal rare-event distribution and build the approximate zero-variance distribution for each sampled cut using \( M = 3000 \) samples that are generated from an Adaptive Metropolis algorithm.
Figure 5-1: Error in the estimates of the optimal solution and optimal cost for a Newsvendor model produced by MCMC-IS and other variance reduction techniques; the model is paired with a lower-variance lognormal distribution in (a)-(b), a higher-variance lognormal distribution in (c)-(d), and a multimodal rare-event distribution in (e)-(f).
Our results suggest that the relative advantage of using MCMC-IS depends on the inherent variance of the underlying SP model. In models where the uncertainty is modeled using a lower-variance distribution, MCMC-IS produces estimates that are just as accurate as the estimates produced by a QMC method, but that are still more accurate than the estimates produced by a CMC method. In models where the uncertainty is modeled using a higher-variance or rare-event distribution, MCMC-IS produces estimates that are much more accurate than those produced by QMC and CMC methods.

5.3 Impact of MCMC-IS Estimates in Stopping Tests

In Section 4.8.2, we highlighted how MCMC-IS estimates are able to consistently produce an estimate that has a low standard error. In this section, we illustrate how the low standard error of MCMC-IS estimates can improve the performance of stopping tests that are used to assess convergence in decomposition algorithms. Once again, we use the SDDP algorithm to solve the two-stage Newsvendor model from Section 4.1.1, which we pair with a lower-variance lognormal distribution, a higher-variance variance lognormal distribution, and a multimodal rare-event distribution.

Given that both the lower bound $z_{LB}$ and the upper bound $z_{UB}$ are random variables in this setting, our stopping test is designed to assess whether their respective expected values $E[z_{LB}]$ and $E[z_{UB}]$ are equal to one another. This requires a one-sided two-sample t-test for the equality of means,

$$H_0 : E[z_{LB}] = E[z_{UB}] \quad \text{vs} \quad H_A : E[z_{LB}] < E[z_{UB}] \quad (5.1)$$

We assume that the samples are unpaired, that the sample sizes are unequal and that the standard deviation of these variables are unknown but identical. We estimate the expected values of these parameters using the sample averages $\bar{z}_{LB}$ and $\bar{z}_{UB}$. Similarly, we estimate the standard deviation of these parameters using the standard errors SE($z_{LB}$) and SE($z_{UB}$). The sample average and the standard error for the lower bound is constructed using $M_{LB} = 3000$ and $N_{LB} = 16000$ samples, while the sample average and sample standard deviation for the upper bound is constructed using $N_{UB} = 16000$ samples. We note that we do not need to
rebuild an approximate zero-variance distribution to construct an upper bound estimate, as the lower bound and upper bound estimates are constructed around the same first-stage solution \( \tilde{x} \).

Unlike traditional hypothesis tests, we are not seeking to reject the null hypothesis but to accept it. As such, our test stops when we are unable to reject the null hypothesis with a significance level of \( \alpha = 0.99 \). The well-known duality between hypothesis tests and confidence intervals implies that this procedure is similar to the stopping tests that involve confidence intervals that are suggested in the literature. We provide further information as to how to construct the upper and lower bound estimate within a decomposition algorithm in Section 2.4.

Our results in Tables 5.3 - 5.5 demonstrate the positive impact that MCMC-IS estimates can have on stopping tests. In particular, MCMC-IS reduces the standard error of upper and lower bound estimates and thereby increases the power of the underlying stopping test. In the context of stopping tests, a test with low power means that the null hypothesis \( H_0 \) is frequently rejected when it is false. In practice, a stopping test with low power terminates a decomposition algorithm before has converged and ultimately results in high errors in the values of \( \tilde{x} \) and \( \tilde{z} \). As in previous sections, these effects become more significant when the variance of the underlying model is increased.

<table>
<thead>
<tr>
<th>Method</th>
<th>SE((z_{LB}))</th>
<th>SE((z_{UB}))</th>
<th># Cuts</th>
<th>MSE((\tilde{x}))</th>
<th>MSE((\tilde{z}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCMC-IS</td>
<td>40</td>
<td>48</td>
<td>7.1</td>
<td>4.4%</td>
<td>0.7%</td>
</tr>
<tr>
<td>CMC</td>
<td>326</td>
<td>329</td>
<td>5.9</td>
<td>9.2%</td>
<td>2.0%</td>
</tr>
<tr>
<td>QMC</td>
<td>312</td>
<td>316</td>
<td>5.6</td>
<td>16.5%</td>
<td>3.0%</td>
</tr>
</tbody>
</table>

Table 5.3: Stopping test output from a Newsvendor model paired with a lower-variance lognormal distribution.

We note that these results cannot be immediately extended to multistage SP models because MCMC-IS can only produce estimates of the upper bound for two-stage models. This is because the previous-stage decision around which we build the upper and lower bound estimate does not change when decomposition algorithms are used to solve two-stage
models. Although there would be significant benefits in using MCMC-IS to generate lower variance estimates of the upper bound for multistage models, the current framework is not computationally tractable as the importance sampling distribution it produces depends on a fixed set of decisions for each stage, and each sample that is used to produce an upper bound estimate typically involves a different set of previous stage decisions. Extending these results to multistage problems therefore an area for further research. Nevertheless, the MCMC-IS framework still outperforms the CMC and QMC sampling techniques in such cases as it can still be used to compute lower variance estimates of the lower bound.

It is true that the stopping test depends on a user-defined parameter, $\alpha$, which controls the appropriate level of Type I error. Unlike statistical hypothesis tests, there is no reason as to why $\alpha$ need not take on a high value because we are not seeking to reject the null hypothesis but instead accept it. Consequently, in Figure 5-2 we show how the statistics presented in Tables 5.3 - 5.5 can change according to the value of $\alpha$ that is chosen.

Our results in the first row of Figure 5-2 show that having a lower-variance estimate of the recourse function is that we can obtain a lower-variance estimate of the upper bound

<table>
<thead>
<tr>
<th>Method</th>
<th>SE($z_{LB}$)</th>
<th>SE($z_{UB}$)</th>
<th># Cuts</th>
<th>MSE($\bar{x}$)</th>
<th>MSE($\bar{z}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCMC-IS</td>
<td>788</td>
<td>839</td>
<td>7.5</td>
<td>5.4%</td>
<td>0.6%</td>
</tr>
<tr>
<td>CMC</td>
<td>44655</td>
<td>43686</td>
<td>4.7</td>
<td>39.0%</td>
<td>33.4%</td>
</tr>
<tr>
<td>QMC</td>
<td>33376</td>
<td>40552</td>
<td>5.0</td>
<td>36.3%</td>
<td>23.8%</td>
</tr>
</tbody>
</table>

Table 5.4: Stopping test output from a Newsvendor model paired with a higher-variance lognormal distribution.

<table>
<thead>
<tr>
<th>Method</th>
<th>SE($z_{LB}$)</th>
<th>SE($z_{UB}$)</th>
<th># Cuts</th>
<th>MSE($\bar{x}$)</th>
<th>MSE($\bar{z}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCMC-IS</td>
<td>276</td>
<td>233</td>
<td>6.4</td>
<td>5.9%</td>
<td>0.5%</td>
</tr>
<tr>
<td>CMC</td>
<td>40589</td>
<td>20902</td>
<td>3.7</td>
<td>37.9%</td>
<td>10.7%</td>
</tr>
<tr>
<td>QMC</td>
<td>19901</td>
<td>16358</td>
<td>3.8</td>
<td>58.8%</td>
<td>12.7%</td>
</tr>
</tbody>
</table>

Table 5.5: Stopping test output from a Newsvendor Model paired with a multimodal rare-event distribution.
Figure 5-2: Stopping test output from Newsvendor model paired with a lower-variance log-normal distribution (left column), a higher-variance lognormal distribution (middle column), and a multimodal rare-event distribution (right column); we vary the value of $\alpha$ in the stopping test between 0.5 - 0.9 and plot the standard error in estimates (top row), the # of cuts until convergence (second row), and the error in the estimated optimal cost (third row), and the error in the estimated optimal solution (bottom row).
and the lower bound used in the stopping tests. The effect of the decreased variance on the lower- and upper bound estimates is a uniformly more powerful stopping test. In this case, a more powerful stopping test corresponds to a stopping test that does not terminate before convergence. This is exactly the result that is highlighted in the second row of Figure 5-2 which illustrates how an SDDP algorithm paired MCMC-IS consistently adds more cuts regardless of the value of $\alpha$. The power of the stopping test ensures that an SDDP algorithm paired with MCMC-IS does not converge too soon, and subsequently produces more accurate values of the optimal cost and optimal solution as shown in the third and bottom row of Figure 5-2 respectively.

5.4 Computational Performance of MCMC-IS in Multistage SP

In our final experiment, we demonstrate the computational benefits of using a sampling approach within a decomposition algorithm. In particular, we embed MCMC-IS within the SDDP algorithm and use this setup to solve a multistage extension of the Newsvendor model paired with a lower-variance lognormal distribution as described in Section 4.1.3.

Figure 5-3(a) shows that the computational complexity of our setup increases quadratically with the time horizon of the underlying problem. Moreover, as is clear from Figure 5-3(b) the solution estimated with MCMC-IS is within 1% of the true value. This represents a significant computational advantage in comparison to a scenario-tree based approach, where the number of linear programs that have to be solved to achieve convergence increases exponentially. The exact number of linear programs that have to be evaluated in this case is determined by the number of samples that we use to construct the sampled cuts at each iteration of the SDDP algorithm, as well as the number of iterations of the SDDP algorithm that we have to run until a stopping test indicates convergence. In this case, we construct sampled cuts using the MCMC-IS algorithm with $M_{LB} = 3000$ and $N_{LB} = 16000$ samples, and we use the stopping test we describe in Section 5.3 to assess convergence. We note that the stopping test from Section 5.3 requires an upper bound estimate, which we construct at each iteration of the using $N_{UB} = 16000$ samples.
Figure 5-3: (a) Complexity of SDDP with MCMC-IS grows quadratically with the number of dimensions. (b) Estimated optimal cost remains within 1% even for problems with a large number of time periods.
Chapter 6

Conclusion

Multistage SP models are considered computationally challenging because the evaluation of the recourse function involves the solution of a multidimensional integral. For large scale problems, optimization algorithms such as SDDP need to be paired with sampling algorithms to estimate the recourse function. The sampling algorithm in this case has a major bearing on the accuracy of the solution to an SP model and the efficiency of the overall process. As a result the development of accurate and efficient sampling methods is an active area of research in SP.

The main contribution of this thesis is the development of a non-parametric importance sampling framework which combines MCMC and KDE algorithms to substantially reduce the sampling error of MC estimates. Our results suggest that even a simple implementation of this framework, which we refer to as MCMC-IS, can generate highly accurate and efficient MC estimates. Moreover, our results highlight that MCMC-IS is well-suited for SP because it can substantially reduce the impact of sampling error in constructing a cutting-plane approximation of the recourse function and choosing when to stop a decomposition algorithm, all the while maintaining a computational cost that is negligible and tractable for large-scale problems.

Much work remains to be done in developing the sampling aspects of MCMC-IS. In terms of theoretical developments, a convergence proof based on the findings of Section 3.4 would formalize the numerical results we present from Chapter 4. In terms of practical developments, further research is needed to show how the MCMC and KDE algorithms used
in an MCMC-IS implementation can be switched or fine-tuned to improve the accuracy and efficiency of estimates in different settings. In particular, there remains a need for an MCMC-IS implementation that can produce estimates when MC estimates depend on discrete random variables. Other practical improvements that should be explored in future research include using a two-stage MCMC algorithm to reduce the computational burden of MCMC-IS, and pairing MCMC-IS with existing variance reduction techniques in order to further improve the convergence rate of sampling error in MCMC-IS estimates.

Although we have shown how MCMC-IS can be used in the context of a decomposition algorithm and expected value optimization, we stress that MCMC-IS can be used with different algorithms and with different types of SP model; such applications should be formally explored in future research as using MC estimates in other optimization models is likely to lead to similar effects. Nevertheless, much work also remains to be done in the domain of SP and decomposition algorithms, particularly on finding ways to generate an upper bound estimate for multistage SP models, and finding opportunities to adapt or re-use the importance sampling distribution from MCMC-IS so as to generate estimates of the recourse function given different previous-stage decisions or different stages.
## Appendix A

### Terminology and Notation

<table>
<thead>
<tr>
<th>Term</th>
<th>Notation</th>
<th>Index Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time Horizon</td>
<td>$T$</td>
<td>-</td>
</tr>
<tr>
<td>Time Index</td>
<td>$t$</td>
<td>-</td>
</tr>
<tr>
<td># of Variables in LP$_t$</td>
<td>$n_t$</td>
<td>$t = 1...T$</td>
</tr>
<tr>
<td># of Constraints in LP$_t$</td>
<td>$m_t$</td>
<td>$t = 1...T$</td>
</tr>
<tr>
<td>Decision Vector in LP$_t$</td>
<td>$x_t$</td>
<td>$t = 1...T$</td>
</tr>
<tr>
<td>Fixed Decision in LP$_t$</td>
<td>$\hat{x}_t$</td>
<td>$t = 1...T-1$</td>
</tr>
<tr>
<td>Cost of Full SP</td>
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<td>Lower Bound on SP Cost</td>
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<tr>
<td>True Optimal Cost of Full SP</td>
<td>$z^*$</td>
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<tr>
<td>True Optimal Solution in First Stage</td>
<td>$x^*$</td>
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<tr>
<td>Estimated Optimal Solution in First Stage</td>
<td>$\tilde{x}$</td>
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<tr>
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<tr>
<td>Recourse Function</td>
<td>$Q_t(\tilde{x}_t)$</td>
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<tr>
<td>Subgradient of Recourse Function</td>
<td>$\nabla Q_t(\tilde{x}_t)$</td>
<td>$t = 2...T$</td>
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<tr>
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<td>Support of Random Vector</td>
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<td>Reconstructed Importance Sampling PDF</td>
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Bibliography


