Seminar Paper

Monte Carlo Methods and Random Sampling

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Abstract

Monte Carlo methods provide essential computational tools for solving high-dimensional integration and sampling problems that are analytically intractable. This paper presents a comprehensive treatment of Monte Carlo techniques, beginning with fundamental integration principles and the curse of dimensionality that motivates probabilistic approaches. We systematically develop key sampling algorithms including the inversion method, Box-Muller transformation, and acceptance-rejection techniques, providing both basic and adaptive variants. We present different variance reduction strategies: most importantly importance sampling (direct and self-normalized variants). We complement pseudo-random approaches with quasi-Monte Carlo methods for improved convergence rates. Practical effectiveness is demonstrated through classical applications including π approximation and rare event estimation in financial risk models, where importance sampling achieves dramatic efficiency gains over standard approaches. The paper concludes with a critical assessment of advantages and limitations, along with an outlook toward advanced methods including sequential importance sampling and Markov Chain Monte Carlo.

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1 Introduction

Monte Carlo methods represent a broad class of computational algorithms that rely on repeated random sampling to obtain numerical results. These methods have become indispensable tools across numerous fields, from physics and engineering to finance and machine learning, particularly when dealing with complex, high-dimensional problems that defy analytical solutions.

The power of Monte Carlo approaches lies in their ability to transform difficult mathematical problems into statistical sampling problems. Rather than attempting to solve equations directly, these methods generate random samples from probability distributions and use statistical inference to extract the desired information. This paradigm shift has proven especially valuable for problems involving uncertainty, optimization, and numerical integration in high-dimensional spaces.

This paper examines the theoretical foundations and practical applications of Monte Carlo methods, with particular emphasis on integration techniques. We begin by introducing the core concepts that form the foundation of this methodology.

Monte Carlo method The Monte Carlo method originated in the mid-1940s during the Manhattan Project, primarily through the work of Stanislaw Ulam, who conceived the basic idea, and John von Neumann, who developed the mathematical framework. Nicholas Metropolis later contributed to its development and suggested the name. The method emerged from their need to solve complex mathematical problems in nuclear physics that involved high-dimensional integrals too complicated for analytical solutions. Their innovation was to use random sampling to approximate these mathematical calculations, creating what became known as the Monte Carlo method.

Lemieux (2009) summarizes the method as:

The use of random sampling as a tool to produce observations on which statistical inference can be performed to extract information about a system.

Monte Carlo integration Monte Carlo integration estimates integrals of the form

$$I(f) = \int_{\mathcal{D}} f(x)dx$$

by sampling points uniformly at random from the domain \mathcal{D} , evaluating the function f at these sample points, and using the sample average to approximate the integral. This approach becomes particularly advantageous in high-dimensional settings where traditional quadrature methods suffer from the curse of dimensionality. Chapter 3 discusses techniques for generating samples from various distributions, while Chapter 4 explores methods to improve the efficiency of Monte Carlo integration through variance reduction. The theoretical foundations and detailed implementation of these integration techniques are covered in Chapter 2.

Monte Carlo simulation Monte Carlo simulation generates samples from complex random variables $Y = \varphi(X)$ by first sampling the input variables X and then applying

the transformation φ . This sampling-based approach allows us to study the statistical properties of Y even when direct sampling or analytical solutions are intractable. The practical implementation relies on efficient sampling algorithms (Chapter 3) and can often benefit from variance reduction strategies (Chapter 4) to improve computational efficiency. While Monte Carlo simulation shares the fundamental sampling principles with Monte Carlo integration (Chapter 2), it focuses on generating realizations from complex distributions rather than estimating integrals. Although Lemieux (2009) shows that simulation problems are integration problems and vice versa, we will use the distinction for the sake of clarity.

Quasi-Monte Carlo methods Quasi-Monte Carlo methods, examined in detail in Chapter 5, generate deterministic point sets $\{u_1, \ldots, u_n\}$ in $[0, 1]^d$ that are more uniformly distributed than random samples. These low-discrepancy sequences provide better coverage of the unit hypercube, often achieving faster convergence rates than standard Monte Carlo methods for smooth integrands.

Random Numbers Random number generation forms the foundation of Monte Carlo methods. As we will demonstrate in subsequent chapters, we can generate random quantities from virtually any distribution by starting with uniform random numbers, making U(0,1) the fundamental building block for all sampling methods.

In practice, computational applications rely on pseudo-random number generators (see Appendix A.2.2 for an exemplary implementation). For the remainder of this work we assume access to a reliable algorithm RUNIF_01(n) that outputs n independent and identically distributed random numbers following U(0,1).

Thomopoulos (2013), Lemieux (2009) provide good introductions into the properties and the construction of pseudo-random number generators.

2 Monte Carlo Integration

In statistics and probabilistic machine learning, we frequently encounter integrals that cannot be solved analytically. A common form is:

$$I(\varphi) = \int \varphi(x)f(x)dx \tag{1}$$

where f(x) is a probability density function and $\varphi(x)$ is some function of interest. Such integrals typically require numerical approximation methods. The Monte Carlo approach to integration is based on a fundamental insight: if we can sample from the probability distribution f(x), we can approximate the integral via the sample average. The Monte Carlo estimator is therefore defined as:

$$\hat{\mu}_n = \frac{1}{n} \sum_{i=1}^n \varphi(X_i) \quad \text{where} \quad X_1, \dots, X_n \overset{\text{i.i.d.}}{\sim} f$$
 (2)

Integrals of the form shown in (1) commonly represent expectations under posterior distributions, marginal likelihoods, or normalization constants. This becomes clearer through

the following examples, with $X \sim f$:

$$\varphi(x) = x \quad \Rightarrow \quad I(\varphi) = \mathbb{E}_f[X]$$

$$\varphi(x) = \mathbb{I}(x \le \gamma) \quad \Rightarrow \quad I(\varphi) = F_X(\gamma)$$

$$\varphi(x) = 1 \quad \Rightarrow \quad I(\varphi) = \int f(x) dx$$

Example 2.1. Consider the integral $I = \int_0^{\pi} \sin(x) dx$. The exact value is I = 2. Suppose we only have access to uniformly distributed random variables on the [0,1) interval. We therefore use substitution to transform the domain to [0,1): $I = \pi \int_0^1 \sin(\pi \cdot u) du \approx \pi \cdot n^{-1} \sum_{i=1}^n \sin(\pi \cdot U_i)$, where $U_i \stackrel{i.i.d.}{\sim} Unif(0,1)$ for all i = 1, ..., n.

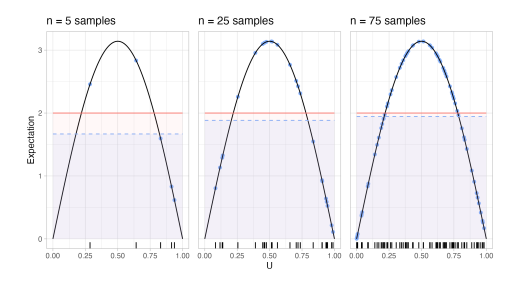


Figure 1: Monte Carlo approximation of $\int_0^{\pi} \sin(x) dx$ showing convergence to the true value of 2 as sample size increases. Generated using mc-integration.Rmd.

A comprehensive analysis of example 2.1, including convergence behavior and implementation details, can be found at mc-integration.html.

2.1 Numerical integration and the curse of dimensionality

An alternative to Monte Carlo integration is numerical integration, where traditional quadrature methods evaluate the integrand at regularly spaced grid points. While effective for low-dimensional problems, these methods suffer from the curse of dimensionality. Maintaining accuracy requires $\tilde{N}=N^d$ function evaluations, where N is the number of points per dimension and d is the dimensionality. For example, a three-dimensional problem with N=100 points per dimension requires $100^3=10^6$ evaluations. Moreover, for the trapezoidal rule, the approximation error scales as $\mathcal{O}(\tilde{N}^{-2/d})$, making convergence prohibitively slow in high dimensions.

Example 2.2. Consider the d-dimensional integral

$$\int_{[0,1]^d} f(\mathbf{x}) d\mathbf{x}$$

where $f(\mathbf{x}) = 1$ if $x_i \in [0.4, 0.6]$ for all i = 1, ..., d and $f(\mathbf{x}) = 0$ otherwise. Only $[0.2]^d$ of grid points contribute to the integral: 4% for d = 2, 0.032% for d = 5, and less than $10^{-7}\%$ for d = 10.

This curse of dimensionality makes Monte Carlo methods particularly attractive, as their error depends only on N, not on dimensionality.

2.2 Properties of the Monte Carlo Estimator

The Monte Carlo estimator approximates the true expectation $\mu = \mathbb{E}_f[\varphi(X)]$ by averaging the function values $\varphi(X_i)$ over n independent samples from the distribution f.

Lemma 2.1. If φ is a measurable function and X_1, \ldots, X_n are i.i.d. random variables, then $\varphi(X_1), \ldots, \varphi(X_n)$ are also i.i.d. random variables.

Proof. This follows from the fact that the composition of a measurable function with a random variable is a random variable, and independence is preserved under measurable transformations. \Box

Theorem 2.2 (Unbiasedness). The Monte Carlo estimator $\hat{\mu}_n$ is unbiased:

$$\mathbb{E}[\hat{\mu}_n] = \mu \tag{3}$$

Theorem 2.3 (Variance). The variance of the Monte Carlo estimator is

$$Var[\hat{\mu}_n] = \frac{\sigma^2}{n} \tag{4}$$

where $\sigma^2 = Var_f[\varphi(X)]$.

Theorem 2.4 (Asymptotic Normality). Under regularity conditions, the Monte Carlo estimator satisfies:

$$\hat{\mu}_n - \mu \xrightarrow{d} \frac{1}{\sqrt{n}} \mathcal{N}(0, \sigma^2) \tag{5}$$

This follows from the Central Limit Theorem¹. The key insight is that the estimation error decreases as $\mathcal{O}(n^{-1/2})$, independent of the dimension d of the integration domain. This is the fundamental advantage of Monte Carlo methods over grid-based integration approaches in high-dimensional settings. Figure 2 illustrates this convergence behavior by showing the estimation error for the integral from example 2.1.

¹see Appendix A.1.1

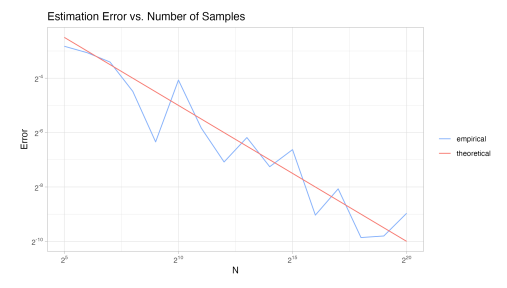


Figure 2: Estimation error for Monte Carlo approximation of $\int_0^{\pi} \sin(x) dx$ demonstrating the $\mathcal{O}(n^{-1/2})$ convergence rate. Generated using mc-integration.Rmd.

Confidence Intervals The asymptotic normality result from Theorem 2.4 enables the construction of confidence intervals for Monte Carlo estimates.

Since $(\hat{\mu}_n - \mu) \xrightarrow{d} \frac{1}{\sqrt{n}} \mathcal{N}(0, \sigma^2)$, we can approximate the distribution of $\hat{\mu}_n$ as $\mathcal{N}(\mu, \sigma^2/n)$ for large n. In practice, the unknown variance σ^2 is estimated by the sample variance:

$$\hat{\sigma}_n^2 = \frac{1}{n-1} \sum_{i=1}^n (\varphi(X_i) - \hat{\mu}_n)^2 \tag{6}$$

An approximate $(1 - \alpha)$ -confidence interval for μ is then given by:

$$\hat{\mu}_n \pm z_{\alpha/2} \frac{\hat{\sigma}_n}{\sqrt{n}} \tag{7}$$

where $z_{\alpha/2}$ is the $(1 - \alpha/2)$ -quantile of the standard normal distribution. This interval provides a measure of uncertainty in the Monte Carlo estimate, with the width decreasing proportionally to $n^{-1/2}$ as the sample size increases.

3 Sampling from Known Distributions

In the previous chapter, we demonstrated how Monte Carlo methods estimate values that would otherwise be difficult to compute directly. The effectiveness of these methods fundamentally depends on our ability to generate random samples from specified probability distributions.

This chapter introduces various techniques for sampling from known target distributions f(x). We assume access to a pseudo-random number generator (RNG)² that produces

²For details on RNGs, see Appendix A.2.2.

samples from the continuous uniform distribution Unif(0, 1). Extensions to sampling from unknown distributions are discussed in Chapter 7.3.

We begin with the inversion method (Section 3.1), which provides a direct approach when the cumulative distribution function is invertible. Section 3.2 focuses on generating samples from the Gaussian distribution. Finally, Section 3.3 introduces techniques for sampling from less tractable distributions through rejection sampling methods. A variety of other methods are available, including composition, convolution or copula-based methods (see Lemieux (2009)).

3.1 Inversion Method

The inversion method allows us to sample from any distribution with an invertible cumulative distribution function (CDF). This fundamental technique transforms uniform random variables into samples from the target distribution (Murphy, 2023, Lemieux, 2009).

Theorem 3.1 (Inverse Transform). Let $U \sim Unif(0,1)$ and F be an invertible cumulative distribution function. Then $X = F^{-1}(U)$ has distribution function F.

Example 3.1 (Exponential Distribution). To sample from the exponential distribution $Exp(\lambda)$, we apply the inversion method. The CDF of the exponential distribution is:

$$F(x; \lambda) = 1 - e^{-\lambda x}$$
 for $x \ge 0$

Setting $u = F(x; \lambda)$ and solving for x, we obtain the inverse CDF:

$$F^{-1}(u) = -\frac{1}{\lambda} \ln(1-u)$$

Since $U \sim U(0,1)$ implies $1-U \sim U(0,1)$, we can simplify the implementation by using U directly instead of 1-U.

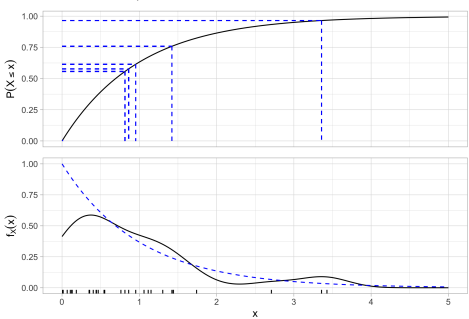
$\overline{\textbf{Algorithm 1}}$ Random Samples from $\text{Exp}(\lambda)$

$$U \leftarrow \text{RUNIF_01}(1) \qquad \qquad \triangleright \text{Generate uniform random variable} \\ X \leftarrow -\frac{1}{\lambda} \ln(U) \qquad \qquad \triangleright \text{Apply inverse transform} \\ \textbf{return } X$$

This demonstrates the power of the inversion method: with a single uniform random variable, we can generate samples from the exponential distribution. Further examples of distributions suitable for the inversion method can be found in Owen (2013).

3.2 Box-Muller Method

The Box-Muller transform is a method for generating pairs of independent standard normal random variables from uniform random variables. The method is based on the polar representation of two independent standard normal variables.



inversion method for exponential distributed random variable

Figure 3: Illustration of the inversion method for generating $\operatorname{Exp}(1)$ random variables. The upper panel demonstrates the transformation process: uniform random samples $U \sim \operatorname{Unif}(0,1)$ are mapped through the inverse cumulative distribution function to yield exponential random variables $X = F^{-1}(U) \sim \operatorname{Exp}(1)$. The lower panel compares the empirical distribution of the generated samples (solid black line) against the theoretical exponential density (dashed blue line), validating the accuracy of the inversion method. Generated using inversion-method.Rmd.

Theorem 3.2 (Box-Muller Transform). Let $U_1, U_2 \sim Unif(0,1)$ be independent uniform random variables. Define:

$$(X_1, X_2) = \left(\sqrt{-2\ln(U_1)}\cos(2\pi U_2), \sqrt{-2\ln(U_1)}\sin(2\pi U_2)\right) \tag{8}$$

Then $X_1, X_2 \sim \mathcal{N}(0, 1)$ are independent standard normal random variables.

Algorithm 3 is a simple implementation of the Box-Muller method and produces two independent samples $X_1, X_2 \sim \mathcal{N}(0, 1)$, an implementation in R can be found at box-muller.Rmd.

Multivariate Normal Random Variables To generate multivariate normal random variables, we transform independent standard normal variables using linear transformations.

Theorem 3.3 (Multivariate Normal Transformation). Let $Z = (Z_1, \ldots, Z_n)^{\top} \sim \mathcal{N}_n(\mathbf{0}, I)$ be a vector of i.i.d. standard normal random variables, $\boldsymbol{\mu} = (\mu_1, \ldots, \mu_n)^{\top}$ be a real-valued mean vector, and $\Sigma \in \mathbb{R}^{n \times n}$ be a symmetric, positive definite covariance matrix. If L is the lower triangular matrix from the Cholesky decomposition³ $\Sigma = LL^{\top}$, then:

$$X = \boldsymbol{\mu} + LZ \sim \mathcal{N}_n(\boldsymbol{\mu}, \Sigma) \tag{9}$$

³see Appendix A.1.2.

This transformation allows us to generate samples from any multivariate normal distribution using only independent standard normal samples (see Figure 4).

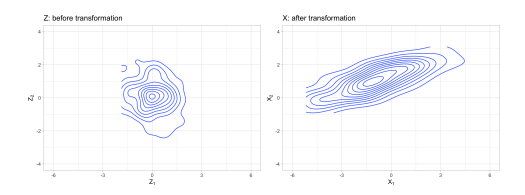


Figure 4: Visualization of the transformation from standard bivariate normal $\mathcal{N}_2(\mathbf{0}, I)$ (left panel) to general bivariate normal $\mathcal{N}_2(\boldsymbol{\mu}, \Sigma)$ (right panel) distributions. The contour plots illustrate the geometric effect of this transformation on the distribution's shape and location. Generated using bivariate-normal.Rmd.

3.3 Acceptance-Rejection

When the target distribution with density p(x) from which we want to sample is intractable for direct sampling methods (yet somehow easy to evaluate), the acceptance-rejection algorithm provides an alternative approach. The idea is to sample from an envelope distribution and then accept or reject these samples according to some criterion. Although this approach wastes some computational effort by throwing away samples, it provides a powerful way to generate samples from complex distributions.

3.3.1 Basic Rejection Sampling

Instead of sampling directly from f(x), we sample from r(x), where r(x) is a probability density function and the following envelope condition is satisfied:

$$T \cdot r(x) > f(x) \quad \forall x$$

We define $t(x) = T \cdot r(x)$. The function t(x) need not be normalized, but must satisfy the condition $T = \int t(x)dx < \infty$.

The acceptance-rejection algorithm can be characterized as follows:

- 1. Generate a candidate $\tilde{X} \sim r$
- 2. Generate $U \sim \text{Unif}(0,1)$ independent of \tilde{X}
- 3. If $U \leq \frac{f(\tilde{X})}{t(\tilde{X})}$, accept and set $X = \tilde{X}$; otherwise, reject and return to step 1.

Theorem 3.4 (Acceptance-Rejection Sampling). Let X denote an accepted sample. Then X has cumulative distribution function $F(x) = \int_{-\infty}^{x} f(y) dy$.

The implementation of acceptance-rejection is straight-forward (see Algorithm 4). The main challenge is sampling from r(x) and possibly the evaluation of $f(\cdot)$ and $t(\cdot)$. For the former challenge we assume to have access to some algorithm rand_r(), which samples from r(x).

3.3.2 Adaptive Rejection Sampling

Adaptive rejection sampling is an extension of the basic acceptance-rejection method that automatically constructs and refines the envelope function during the sampling process. For log-concave target densities, the algorithm exploits the property that tangent lines to the log-density provide a tight upper bound. The method begins with an initial set of tangent points and iteratively improves the piecewise-linear envelope by adding new tangent lines at rejected sample locations. Algorithm 5 demonstrates a basic implementation. The adaptive refinement ensures that the envelope becomes increasingly efficient over time, dramatically reducing the rejection rate as more samples are drawn. For more detailed information on adaptive rejection sampling, see Murphy (2023).

4 Variance Reduction Techniques

Variance reduction techniques aim to improve Monte Carlo integration by finding alternative functions whose integrals equal the original but have smaller variance. The goal is to achieve a reduction in variance while restricting the computational increase⁴. We will focus on three methods: In section 4.1 we will discuss *Importance Sampling*, likely the most well-known technique. We then follow up in section 4.2 with a technique called *Control Variates* and will finish in section 4.3 with *Conditional Monte Carlo*. There are many more techniques⁵ in existence, and Lemieux (2009) even mentions the possibility to combine them.

4.1 Importance Sampling

Importance sampling directs sampling effort toward the most important regions of the integration domain, making it particularly useful for rare event simulation. Instead of sampling from the original density f(x), we sample from an alternative density g(x) and correct using likelihood ratios. We will focus on two fundamental variants: Direct Importance Sampling and Self-Normalized Importance Sampling, as discussed in Lemieux (2009).

Although other variants exist, such as *Annealed importance sampling* (discussed in Murphy (2023)), they fall outside the scope of this paper.

 $^{^4}$ A measure of the quality of estimators that considers both variance and computation time, is called *Efficiency*. Lemieux (2009)

⁵ for example Antithetic Sampling and Common Random Numbers, for further information see Murphy (2023), Lemieux (2009).

4.1.1 Direct Importance Sampling

The standard importance sampling estimator transforms $\mu = \int \varphi(x) f(x) dx$ into:

$$\hat{\mu}_{is} = \frac{1}{n} \sum_{i=1}^{n} \varphi(\tilde{x}_i) L(\tilde{x}_i)$$
(10)

where L(x) = f(x)/g(x) is the likelihood ratio and $\tilde{x}_i \sim g(x)$.

Theorem 4.1 (Unbiasedness of Importance Sampling). The importance sampling estimator $\hat{\mu}_{is}$ is unbiased for μ .

Theorem 4.2 (Variance of Importance Sampling). The variance of the importance sampling estimator is:

$$Var[\hat{\mu}_{is}] = \frac{1}{n} \left(E[\varphi^2(X)L(X)] - \mu^2 \right)$$
(11)

where the expectation is taken with respect to the original density f(x).

Variance reduction occurs when $E[\varphi^2(X)L(X)] \leq E[\varphi^2(X)]$. If μ happens to be known, then there exists an optimal choice $g^*(x) = |\varphi(x)| \frac{f(x)}{\mu}$, which yields zero variance μ (see Lemieux (2009)).

4.1.2 Self-normalized Importance Sampling

The weighted importance sampling estimator uses normalized likelihood ratios:

$$\hat{\mu}_{is,w} = \frac{\sum_{i=1}^{n} \varphi(\tilde{x}_i) L(\tilde{x}_i)}{\sum_{i=1}^{n} L(\tilde{x}_i)}$$
(12)

This approach has bounded weights between 0 and 1, handles complicated densities with unknown normalizing constants, and is consistent though biased (see Lemieux (2009), Murphy (2023)). The normalization cancels unknown constants in both f and g.

4.2 Control Variates

Control variates reduce variance by exploiting correlation with other variables whose expectations are known. Suppose we want to estimate $\mu = E[f(X)]$ using estimator $m(X) = \frac{1}{S} \sum_{s=1}^{S} m(x_s)$ where $x_s \sim p(X)$ and $E[m(X)] = \mu$ (see Murphy (2023)). The control variate estimator uses a baseline function b(X) with known expectation:

initiator uses a pasenne function $o(\Lambda)$ with known expectation.

$$m^*(X) = m(X) + c(b(X) - E[b(X)])$$
 (13)

where c is a coefficient to be optimized.

Theorem 4.3 (Control Variate Variance Reduction). The control variate estimator is unbiased and achieves optimal variance reduction with:

$$c^* = -\frac{Cov(m(X), b(X))}{Var(b(X))}$$

$$Var(m^*(X)) = (1 - \rho_{m,b}^2) Var(m(X)) \le Var(m(X))$$

where $\rho_{m,b}^2$ is the squared correlation between the estimator and baseline.

The intuition is that the estimator exploits information about errors in estimating the known quantity E[b(X)] to reduce errors in estimating the unknown μ .

4.3 Conditional Monte Carlo

Conditional Monte Carlo, also known as Rao-Blackwellization reduces variance by analytically marginalizing out some variables instead of sampling them. Suppose we want to estimate f = E[f(X,Y)] where we can tractably compute the conditional expectation E[f(X,Y)|X] Murphy (2023).

The naive Monte Carlo estimator uses samples $(X_s, Y_s) \sim p(X, Y)$:

$$\hat{f}_{MC} = \frac{1}{S} \sum_{s=1}^{S} f(X_s, Y_s)$$

The Rao-Blackwellized estimator instead samples only $X_s \sim p(X)$ and computes:

$$\hat{f}_{RB} = \frac{1}{S} \sum_{s=1}^{S} f_X(X_s) \tag{14}$$

where $f_X(X_s) = \int dY p(Y|X_s) f(X_s, Y) = E[f(X, Y)|X = X_s].$

The Rao-Blackwellized estimator is unbiased and can have lower variance than the naive estimator (see Murphy (2023)).

5 Quasi Monte Carlo

Lemieux (2009) provides a useful characterization of Quasi-Monte Carlo sampling methods:

Quasi–Monte Carlo sampling methods are typically used to provide approximations for multivariate integration problems defined over the unit hypercube.

In Quasi-Monte Carlo (QMC) methods, random sampling is replaced by carefully constructed deterministic sequences. Rather than sampling randomly from the d-dimensional unit hypercube $[0,1]^d$, QMC methods generate low discrepancy sequences⁶ for evaluating integrals of the form $\int_{[0,1]^d} f(x) dx$. These sequences minimize clustering and gaps, providing more systematic and uniform coverage of the integration domain. This approach often achieves better convergence rates than standard Monte Carlo methods, particularly for smooth integrands.

Deterministic Nature and Its Implications While the systematic point placement of QMC often leads to faster convergence rates, it has two key limitations. First, the Central Limit Theorem does not apply to deterministic sequences, making standard error estimation techniques unusable; concequently QMC does not provide confidence intervals. Second, QMC performance depends heavily on integrand smoothness.

⁶For a more detailed background on low discrepancy sequences, see Appendix A.2.1.

Comparison with Traditional Numerical Integration When projected onto lowerdimensional subspaces, grid points of regular grids often collapse onto a much smaller set, creating poor coverage. Low discrepancy sequences are specifically designed to maintain good uniformity properties even under such projections.

A visual comparison can be found in Figure 5a (Section 6.1), which illustrates the key differences between these approaches. Random sampling exhibits clustering and gaps, regular grids show systematic coverage but suffer from poor projection properties, while low discrepancy sequences achieve both systematic coverage and retain uniformity under projections.

For a comprehensive treatment of QMC, see Owen (2023).

6 Applications

6.1 Approximating π

We demonstrate different sampling strategies through the well-known problem of estimating π by approximating the area of a unit circle inscribed in a square. A comprehensive analysis including implementation details and extended theoretical background can be found at example1-estimating-pi.html. In the following, we discuss only the general setup and key results.

Since the unit circle has area π and the enclosing square $[-1,1] \times [-1,1]$ has area 4, we obtain the estimator $\hat{\pi} = 4 \cdot \mathbb{P}(X^2 + Y^2 \leq 1)$ where (X,Y) are sampled from the square. **Sampling Methods Compared:** We compare uniform Monte Carlo sampling, importance sampling with $\mathcal{N}(0, \sigma^2 I_2)$ where $\sigma = 1/3$, quasi-Monte Carlo using Sobol sequences, and grid-based deterministic quadrature.

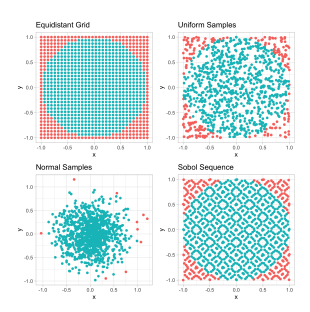
Key Results: Figure 5 shows that Sobol sequences demonstrate superior performance with consistently lower errors due to space-filling properties. Uniform Monte Carlo exhibits the expected $O(N^{-1/2})$ convergence, while the chosen importance sampling parameters provide no variance reduction for this problem.

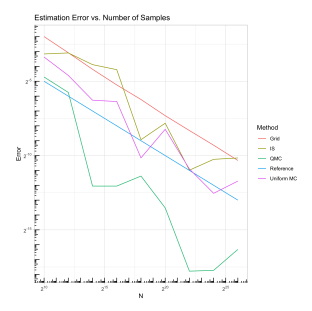
6.2 Probability of Rare Events

Estimating probabilities of rare events presents a fundamental challenge for standard Monte Carlo methods due to high variance and poor convergence. We demonstrate this challenge through a portfolio risk model where we estimate the probability of a remarkably high profit. In the following, we will discuss only the general setup and key results. A comprehensive analysis of this problem, including theoretical background and implementation details, can be found at example2-rare-event.html.

Portfolio Setup: Consider a portfolio with initially equally weighted assets, where each stock's log-return follows $Z_i \sim \mathcal{N}(\mu_i, \sigma_i^2)$ with parameters $\boldsymbol{\mu} = (-0.1, 0.2, -0.3, 0.1, 0)$ and $\boldsymbol{\sigma}^2 = (0.3, 0.3, 0.3, 0.2, 0.2)$. The portfolio value is $S = \sum_{i=1}^5 \exp(Z_i)$. Our objective is to estimate $\mathbb{P}(\bar{S} > 4)$, representing a 300% profit.

Methods Compared: Standard Monte Carlo samples directly from the original distributions, while importance sampling uses shifted parameters $\tilde{\mu}_i = 1$ and $\tilde{\sigma}_i = 1$ to make the rare event more likely, then corrects using importance weights.





(a) Visualization of the four sampling strategies, each with $2^{10} = 1024$ points colored by whether they fall inside the unit circle.

(b) Convergence comparison showing absolute error versus sample size. Sobol sequences demonstrate superior performance due to space-filling properties.

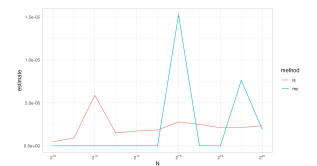
Figure 5: Comparison of sampling strategies for π estimation. Generated using example1-estimating-pi.Rmd.

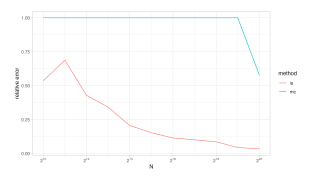
Key Result: For this rare event with probability $P \approx 2.3 \times 10^{-6}$, importance sampling achieves dramatically lower relative errors compared to standard Monte Carlo, enabling practical estimation of tail probabilities, as demonstrated in Figure 6.

This example illustrates why importance sampling is essential for applications requiring accurate tail probability estimation.

7 Conclusion

The equivalence between integration and expectation forms the cornerstone of probabilistic Monte Carlo methods. This manifests in the identity $\mathbb{E}[\varphi(X)] = \int \varphi(x) f(x) dx$, which reveals that every expectation can be viewed as an integral and every integral can be recast as an expectation under an appropriate probability measure, thereby transforming the problem of numerical integration into one of statistical sampling. This insight allows deterministic integrals to be estimated by sample averages and provides the theoretical foundation for importance sampling and other variance reduction techniques. While quasi-Monte Carlo methods take a different approach through deterministic low-discrepancy sequences, both probabilistic and deterministic methods ultimately address the same fundamental challenge: efficient evaluation of high-dimensional integrals where traditional quadrature methods fail due to the curse of dimensionality.





- (a) Probability estimates versus sample size showing the superior convergence of importance sampling.
- (b) Relative error comparison demonstrating orders of magnitude improvement with importance sampling.

Figure 6: Comparison of standard Monte Carlo versus importance sampling for rare event estimation. Generated using example2-rare-event.Rmd.

7.1 Advantages

The primary strength of Monte Carlo methods lies in their dimension-independent convergence rate of $O(N^{-1/2})$, making them particularly valuable for high-dimensional problems that would be intractable through analytical or deterministic numerical approaches. This flexibility enables tackling complex integration problems across diverse domains. Monte Carlo methods naturally support variance reduction techniques such as importance sampling, which can significantly improve convergence rates. Furthermore, they provide natural uncertainty quantification through their inherent randomness, while quasi-Monte Carlo extensions can achieve even better convergence rates for smooth integrands.

7.2 Limitations

Monte Carlo methods face several fundamental limitations. The $O(N^{-1/2})$ convergence rate, while dimension-independent, is relatively slow compared to deterministic methods in low dimensions, and the stochastic nature introduces result variability. Many practical applications require sampling from complex or unknown distributions, necessitating advanced techniques like Markov Chain Monte Carlo or Sequential importance Sampling. Variance reduction techniques require domain expertise and can fail when poorly implemented. While quasi-Monte Carlo methods can overcome some limitations by achieving superior convergence rates, their deterministic nature prohibits confidence interval construction, eliminating natural uncertainty quantification.

7.3 Outlook

The field of Monte Carlo methods continues to evolve rapidly. An excellent source for recent advances is the biannual *Monte Carlo and Quasi–Monte Carlo Methods* conference proceedings by Springer (Plaskota and Woźniakowski (2012), Dick (2013), Cools (2016), Owen (2018), Tuffin and L'Ecuyer (2020), Keller (2022), Hinrichs et al. (2024)).

7.3.1 Sequential Importance Sampling

Sequential importance sampling extends static importance sampling to dynamic settings where the target distribution evolves over time or depends on sequential observations. This framework is particularly valuable in filtering problems for estimating posterior distributions of hidden states given sequential observations. For information on the method and statistical properties, Murphy (2023), Barbu and Zhu (2020) are excellent starting points.

7.3.2 Markov Chain Monte Carlo

Markov Chain Monte Carlo methods enable sampling from complex multivariate distributions by constructing Markov chains with desired stationary distributions. MCMC methods, including Metropolis-Hastings algorithms and Gibbs sampling, make previously intractable posterior distributions accessible. Barbu and Zhu (2020), Murphy (2023) provide comprehensive treatments of MCMC and its applications. Recent developments include Hamiltonian Monte Carlo and variational inference.

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Usage of Artificial Intelligence

In the preparation of this paper, I utilized two Artificial Intelligence (AI) tools:

- Claude Sonnet 4: Primarily used for proofreading draft chapters and improving readability and narrative flow. It was also used for LaTeX formatting tasks.
- **DeepL**: Used for proofreading chapter drafts and generating alternative formulations and explanations for individual text passages.

A Appendix

A.1 Mathematical Foundations

This section provides essential mathematical concepts.

A.1.1 Probability Theory

Central Limit Theorem and Law of Large Numbers The Strong Law of Large Numbers provides the fundamental justification for Monte Carlo methods. If $X_1, X_2, ...$ are independent and identically distributed random variables with finite mean μ , then:

$$\bar{X}_n = \frac{1}{n} \sum_{i=1}^n X_i \xrightarrow{a.s.} \mu \quad \text{as } n \to \infty$$

The Central Limit Theorem establishes the rate of convergence and enables uncertainty quantification. For the same sequence with finite variance σ^2 :

$$\bar{X}_n \xrightarrow{d} \mathcal{N}(\mu, \sigma^2/n)$$
 as $n \to \infty$

Bivariate Change of Variables For a bijective transformation $(Y_1, Y_2) = T(X_1, X_2)$ with inverse $H = T^{-1}$, the joint PDF transforms as:

$$f_{Y_1,Y_2}(y_1,y_2) = f_{X_1,X_2}(H(y_1,y_2)) |J_H(y_1,y_2)|$$

where $|J_H(y_1, y_2)|$ is the absolute determinant of the Jacobian matrix of H. This transformation rule is essential for deriving sampling methods like the Box-Muller transform.

Rayleigh Distribution The Rayleigh distribution with scale parameter $\sigma > 0$ has PDF:

$$f(x;\sigma) = \begin{cases} \frac{x}{\sigma^2} \exp\left(\frac{-x^2}{2\sigma^2}\right) & x \ge 0\\ 0 & x < 0 \end{cases}$$

and CDF:

$$F(x;\sigma) = \begin{cases} 1 - \exp\left(\frac{-x^2}{2\sigma^2}\right) & x \ge 0\\ 0 & x < 0 \end{cases}$$

The Rayleigh distribution is related to the exponential distribution: if $X \sim \text{Exp}(\lambda)$, then $\sqrt{X} \sim \text{Rayleigh}\left(\frac{1}{\sqrt{2\lambda}}\right)$.

A.1.2 Linear Algebra

Positive Definite Matrices and Cholesky Decomposition A symmetric matrix $\Sigma \in \mathbb{R}^{d \times d}$ is positive definite if $x^T \Sigma x > 0$ for all non-zero vectors $x \in \mathbb{R}^d$. Every positive definite matrix admits a unique Cholesky decomposition $\Sigma = LL^T$, where L is lower triangular with positive diagonal entries.

A.2 Computational Methods

This section covers computational techniques for implementing Monte Carlo methods.

A.2.1 Low Discrepancy Sequences

Low discrepancy sequences provide deterministic point sets with superior uniformity compared to pseudo-random sequences. The discrepancy of a sequence $P = \{x_1, \ldots, x_N\}$ in $[0, 1)^d$ measures deviation from uniformity:

$$D(P, \mathcal{F}) = \sup_{B \in \mathcal{F}} \left| \frac{\#(B \cap P)}{N} - \text{vol}(B) \right|$$

where \mathcal{F} is typically a family of rectangular test sets.

Sobol sequences are particularly effective in moderate to high dimensions and widely used in financial applications. **Halton sequences** are simpler to generate and perform well in low dimensions, but quality deteriorates in higher dimensions due to coordinate correlations.

While standard Monte Carlo achieves $\mathcal{O}(N^{-1/2})$ convergence, low discrepancy sequences can achieve $\mathcal{O}((\log N)^d/N)$ for smooth integrands—a significant improvement in moderate dimensions, though the curse of dimensionality limits benefits as d increases.

A.2.2 Random Number Generation

The Lehmer generator provides a foundation for pseudo-random number generation using the recurrence $X_{n+1} = (a \cdot X_n) \mod m$. A standard implementation uses multiplier $a = 7^5$ and modulus $m = 2^{31} - 1$:

Algorithm 2 Lehmer Random Number Generator

```
 \begin{array}{lll} \textbf{Require:} & X_0 > 0 & > \text{Initial seed} \\ a \leftarrow 7^5, \mathbf{m} \leftarrow 2^{31} - 1 & & \\ X \leftarrow X_0 & & \\ \textbf{loop} & & \\ X \leftarrow (a \cdot X) \bmod m & & \\ U \leftarrow X/m & > \text{Convert to uniform on } (0, 1) \\ \textbf{return } U & & \\ \textbf{end loop} & & \\ \end{array}
```

The choice of parameters a and m is crucial for achieving good statistical properties and full period length (see Lemieux (2009)).

A.3 Algorithms

Algorithm 3 Box-Muller Transform

```
U_1, U_2 \leftarrow \text{RUNIF\_01}(2) \Rightarrow Generate two uniform random variables R \leftarrow \sqrt{-2\ln(U_1)} \Rightarrow Transform to radius \Theta \leftarrow 2\pi U_2 \Rightarrow Transform to angle X_1 \leftarrow R\cos(\Theta) \Rightarrow First standard normal sample X_2 \leftarrow R\sin(\Theta) \Rightarrow Second standard normal sample return (X_1, X_2)
```

Algorithm 4 Acceptance-Rejection Sampling

```
 \begin{array}{lll} \textbf{repeat} & & & & & \\ & \tilde{X} \leftarrow \texttt{RAND\_R}() & & & & & & \\ & U \leftarrow \texttt{RUNIF\_01}(1) & & & & & & \\ & \alpha \leftarrow p(\tilde{X})/t(\tilde{X}) & & & & & \\ & \textbf{until } U \leq \alpha & & & & & \\ & X \leftarrow \tilde{X} & & & & & \\ & \textbf{return } X & & & & \\ & & & & & & \\ \end{array}
```

Algorithm 5 Adaptive Rejection Sampling

```
Require: \log p(x)
                                                                      ▶ Log-density of target distribution
Require: T_0 = \{x_1, x_2, \dots, x_k\}
                                                                                      ▶ Initial tangent points
                                                                              ▶ Initialize tangent point set
   E \leftarrow \text{CONSTRUCT\_ENVELOPE}(T)
                                                                                      ▶ Build initial envelope
  loop
       X \leftarrow \text{SAMPLE\_FROM\_ENVELOPE}(E)

    Sample from current envelope

       U \leftarrow \text{RUNIF}_01(1)
                                                                    ▶ Generate uniform random variable
       \begin{array}{l} \alpha \leftarrow \frac{p(\tilde{X})}{E(\tilde{X})} \\ \text{if } U \leq \alpha \text{ then} \end{array}
                                                                               ▷ Calculate acceptance ratio
                                                                                              ▶ Accept sample
           return X
                                                                               ▶ Reject and refine envelope
       else
            T \leftarrow T \cup \{\tilde{X}\}
                                                                     ▶ Add rejected point to tangent set
            E \leftarrow \text{CONSTRUCT\_ENVELOPE}(T)
                                                                      ▶ Rebuild envelope with new point
       end if
  end loop
```

B Electronic appendix

Data, code and figures can be found on github.com.

C Proofs for Section 2 (Monte Carlo Integration)

Theorem 2.2 (Unbiasedness). The Monte Carlo estimator $\hat{\mu}_n$ is unbiased:

$$\mathbb{E}[\hat{\mu}_n] = \mu \tag{3}$$

Proof. Assume $\mathbb{E}_f[\varphi(X)] = \mu$ exists. Then:

$$\mathbb{E}[\hat{\mu}_n] = \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n \varphi(X_i)\right]$$

$$= \frac{1}{n}\sum_{i=1}^n \mathbb{E}[\varphi(X_i)] \quad \text{(linearity of expectation)}$$

$$= \frac{1}{n}\sum_{i=1}^n \mu \quad \text{(identical distribution)}$$

$$= \mu$$

Theorem 2.3 (Variance). The variance of the Monte Carlo estimator is

$$Var[\hat{\mu}_n] = \frac{\sigma^2}{n} \tag{4}$$

where $\sigma^2 = Var_f[\varphi(X)]$.

Proof. Assume $\operatorname{Var}_f[\varphi(X)] = \sigma^2 < \infty$. Then:

$$\operatorname{Var}[\hat{\mu}_n] = \operatorname{Var}\left[\frac{1}{n} \sum_{i=1}^n \varphi(X_i)\right]$$

$$= \frac{1}{n^2} \sum_{i=1}^n \operatorname{Var}[\varphi(X_i)] \quad \text{(independence)}$$

$$= \frac{1}{n^2} \cdot n \cdot \sigma^2 \quad \text{(identical distribution)}$$

$$= \frac{\sigma^2}{n}$$

D Proofs for Section 3 (Sampling from Known Distributions)

Theorem 3.1 (Inverse Transform). Let $U \sim Unif(0,1)$ and F be an invertible cumulative distribution function. Then $X = F^{-1}(U)$ has distribution function F.

Proof. For any $x \in \mathbb{R}$:

$$\mathbb{P}(F^{-1}(U) \le x) = \mathbb{P}(U \le F(x))$$
 (applying F to both sides)
= $F(x)$ (since $U \sim \text{Unif}(0, 1)$)

Therefore, $F^{-1}(U)$ has cumulative distribution function F.

Theorem 3.2 (Box-Muller Transform). Let $U_1, U_2 \sim Unif(0, 1)$ be independent uniform random variables. Define:

$$(X_1, X_2) = \left(\sqrt{-2\ln(U_1)}\cos(2\pi U_2), \sqrt{-2\ln(U_1)}\sin(2\pi U_2)\right) \tag{8}$$

Then $X_1, X_2 \sim \mathcal{N}(0, 1)$ are independent standard normal random variables.

Proof. We prove this by transforming to polar coordinates and applying the change of variables formula.

Let $R = \sqrt{-2 \ln(U_1)}$ and $\Theta = 2\pi U_2$. We first determine the joint distribution of (R, Θ) . **Step 1: Distribution of** R. Since $U_1 \sim \text{Unif}(0,1)$, we have $-\ln(U_1) \sim \text{Exp}(1)$, which implies $-2 \ln(U_1) \sim \text{Exp}(1/2)$. Therefore, $R^2 = -2 \ln(U_1) \sim \text{Exp}(1/2)$, making R follow a Rayleigh distribution with parameter 1 (see Appendix A.1.1):

$$f_R(r) = re^{-r^2/2} \quad \text{for } r > 0$$

Step 2: Distribution of Θ . Since $U_2 \sim \text{Unif}(0,1)$, we have $\Theta = 2\pi U_2 \sim \text{Unif}(0,2\pi)$ with density:

$$f_{\Theta}(\theta) = \frac{1}{2\pi} \quad \text{for } \theta \in [0, 2\pi)$$

Step 3: Joint distribution of (R, Θ) . Since U_1 and U_2 are independent, R and Θ are also independent. Therefore:

$$f_{R,\Theta}(r,\theta) = f_R(r) \cdot f_{\Theta}(\theta) = \frac{r}{2\pi} e^{-r^2/2}$$
 for $r > 0, \theta \in [0, 2\pi)$

Step 4: Transformation to Cartesian coordinates. The transformation $(X_1, X_2) = (R\cos(\Theta), R\sin(\Theta))$ converts from polar to Cartesian coordinates. The Jacobian matrix is:

$$J = \begin{bmatrix} \cos(\theta) & -r\sin(\theta) \\ \sin(\theta) & r\cos(\theta) \end{bmatrix}$$

The determinant is $det(J) = r cos^2(\theta) + r sin^2(\theta) = r$.

Step 5: Change of variables. Using the change of variables formula (see Appendix A.1.1), the joint density of (X_1, X_2) is:

$$f_{X_1,X_2}(x_1,x_2) = f_{R,\Theta}(r,\theta) \cdot \frac{1}{|\det(J)|}$$
$$= \frac{r}{2\pi} e^{-r^2/2} \cdot \frac{1}{r}$$
$$= \frac{1}{2\pi} e^{-(x_1^2 + x_2^2)/2}$$

where we used $r^2 = x_1^2 + x_2^2$. This joint density factors as:

$$f_{X_1,X_2}(x_1,x_2) = \left(\frac{1}{\sqrt{2\pi}}e^{-x_1^2/2}\right) \cdot \left(\frac{1}{\sqrt{2\pi}}e^{-x_2^2/2}\right)$$

Therefore, X_1 and X_2 are independent standard normal random variables.

Theorem 3.3 (Multivariate Normal Transformation). Let $Z = (Z_1, \ldots, Z_n)^{\top} \sim \mathcal{N}_n(\mathbf{0}, I)$ be a vector of i.i.d. standard normal random variables, $\boldsymbol{\mu} = (\mu_1, \ldots, \mu_n)^{\top}$ be a real-valued mean vector, and $\Sigma \in \mathbb{R}^{n \times n}$ be a symmetric, positive definite covariance matrix. If L is the lower triangular matrix from the Cholesky decomposition $\Sigma = LL^{\top}$, then:

$$X = \boldsymbol{\mu} + LZ \sim \mathcal{N}_n(\boldsymbol{\mu}, \Sigma) \tag{9}$$

Proof. We prove this by showing that X has the correct mean and covariance structure. **Mean:**

$$\mathbb{E}[X] = \mathbb{E}[\boldsymbol{\mu} + LZ]$$
$$= \mathbb{E}[\boldsymbol{\mu}] + L\mathbb{E}[Z]$$
$$= \boldsymbol{\mu} + L \cdot \mathbf{0} = \boldsymbol{\mu}$$

Covariance:

$$Cov(X) = Cov(\boldsymbol{\mu} + LZ)$$

$$= Cov(LZ)$$

$$= L \cdot Cov(Z) \cdot L^{\top}$$

$$= L \cdot I_n \cdot L^{\top} = LL^{\top} = \Sigma$$

Since X is a linear transformation of jointly normal random variables, X is also multivariate normal with mean μ and covariance Σ .

Theorem 3.4 (Acceptance-Rejection Sampling). Let X denote an accepted sample. Then X has cumulative distribution function $F(x) = \int_{-\infty}^{x} f(y) dy$.

⁷see Appendix A.1.2.

Proof. We need to show that

$$\mathbb{P}(X \le x) = \mathbb{P}\left(\tilde{X} \le x \mid U \le \frac{f(\tilde{X})}{t(\tilde{X})}\right) = F(x)$$

Using the definition of conditional probability:

$$\mathbb{P}\left(\tilde{X} \leq x \mid U \leq \frac{f(\tilde{X})}{t(\tilde{X})}\right) = \frac{\mathbb{P}\left(\tilde{X} \leq x, U \leq \frac{f(\tilde{X})}{t(\tilde{X})}\right)}{\mathbb{P}\left(U \leq \frac{f(\tilde{X})}{t(\tilde{X})}\right)}$$

For the numerator, using the law of total probability and independence of U and \tilde{X} :

$$\mathbb{P}\left(\tilde{X} \le x, U \le \frac{f(\tilde{X})}{t(\tilde{X})}\right) = \int_{-\infty}^{x} \mathbb{P}\left(U \le \frac{f(y)}{t(y)}\right) r(y) dy$$
$$= \int_{-\infty}^{x} \frac{f(y)}{t(y)} r(y) dy$$
$$= \frac{1}{T} \int_{-\infty}^{x} f(y) dy$$

For the denominator:

$$\begin{split} \mathbb{P}\left(U \leq \frac{f(\tilde{X})}{t(\tilde{X})}\right) &= \int_{-\infty}^{\infty} \frac{f(y)}{t(y)} r(y) dy \\ &= \frac{1}{T} \int_{-\infty}^{\infty} f(y) dy = \frac{1}{T} \end{split}$$

Therefore:

$$\mathbb{P}\left(\tilde{X} \le x \mid U \le \frac{f(\tilde{X})}{t(\tilde{X})}\right) = \frac{\frac{1}{T} \int_{-\infty}^{x} f(y) dy}{\frac{1}{T}} = \int_{-\infty}^{x} f(y) dy = F(x)$$

E Proofs for Section 4 (Variance Reduction Techniques)

Theorem 4.1 (Unbiasedness of Importance Sampling). The importance sampling estimator $\hat{\mu}_{is}$ is unbiased for μ .

Proof. We compute the expectation:

$$E(\hat{\mu}_{is}) = E\left[\frac{1}{n} \sum_{i=1}^{n} \varphi(\tilde{X}_{i}) L(\tilde{X}_{i})\right]$$

$$= E[\varphi(\tilde{X}) L(\tilde{X})]$$

$$= \int \varphi(x) L(x) g(x) dx$$

$$= \int \varphi(x) \frac{f(x)}{g(x)} g(x) dx$$

$$= \int \varphi(x) f(x) dx = \mu$$

Theorem 4.2 (Variance of Importance Sampling). The variance of the importance sampling estimator is:

$$Var[\hat{\mu}_{is}] = \frac{1}{n} \left(E[\varphi^2(X)L(X)] - \mu^2 \right)$$
(11)

where the expectation is taken with respect to the original density f(x).

Proof. The variance is:

$$\operatorname{Var}[\hat{\mu}_{is}] = \frac{1}{n} \operatorname{Var}[\varphi(\tilde{X}) L(\tilde{X})]$$
$$= \frac{1}{n} \left(E[\varphi^{2}(\tilde{X}) L^{2}(\tilde{X})] - \mu^{2} \right)$$

We compute the second moment:

$$E[\varphi^{2}(\tilde{X})L^{2}(\tilde{X})] = \int \varphi^{2}(x)L^{2}(x)g(x)dx$$

$$= \int \varphi^{2}(x)\left(\frac{f(x)}{g(x)}\right)^{2}g(x)dx$$

$$= \int \varphi^{2}(x)\frac{f^{2}(x)}{g(x)}dx$$

$$= \int \varphi^{2}(x)\frac{f(x)}{g(x)}f(x)dx$$

$$= E[\varphi^{2}(X)L(X)]$$

where the last expectation is with respect to the original density f(x).

Theorem 4.3 (Control Variate Variance Reduction). The control variate estimator is unbiased and achieves optimal variance reduction with:

$$c^* = -\frac{Cov(m(X), b(X))}{Var(b(X))}$$
$$Var(m^*(X)) = (1 - \rho_{m,b}^2) Var(m(X)) \le Var(m(X))$$

where $\rho_{m,b}^2$ is the squared correlation between the estimator and baseline.

Proof. Unbiasedness: $E[m^*(X)] = E[m(X)] + c(E[b(X)] - E[b(X)]) = E[m(X)] = \mu$. Variance: $\operatorname{Var}(m^*(X)) = \operatorname{Var}(m(X)) + c^2\operatorname{Var}(b(X)) + 2c\operatorname{Cov}(m(X), b(X))$. Taking $\frac{\partial}{\partial c}\operatorname{Var}(m^*(X)) = 0$ yields $c^* = -\frac{\operatorname{Cov}(m(X), b(X))}{\operatorname{Var}(b(X))}$. Substituting gives the variance reduction formula.