#### A M.Sc Thesis

on

# Simulation and Markov Chain Monte Carlo

by

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Enrollment No.: 2022MAM008

under the supervision of

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## Department of Mathematics

Submitted for the partial fulfillment of the requirements for the award of the degree of

M.Sc. in Applied Mathematics

## **CERTIFICATE**

This is to certify that the Term Paper on "Simulation and Markov Chains Monte Carlo" submitted by Azmain Biswas to the Department of Mathematics, Indian Institute of Engineering Science and Technology, Shibpur, Howrah-711103, for 3rd semester of Master of Science in Applied Mathematics is a record of project work carried out by him under my supervision.

The result of this project work or any part thereof has not been submitted for any degree or diploma.

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## Acknowledgment

Thanks to my supervisor, Prof. M. Mitra, for a very interesting project. It has been a great pleasure to work under his guidance, and I have learned a lot from his way of thinking and his approach to a problem.

I am also indebted to Prof. Pritha Das, Head of the Department of Mathematics, Indian Institute of Engineering Science and Technology, Shibpur, for her help.

At various stages, I received encouragement and inspiration from most of the faculty members of the department. I would like to thank all of them for their invaluable words of advice.

I have benefited greatly from the advantages of the TEX-group software typing systems, the use of which has contributed much to simplify the typing of long mathematical expressions involved in my work.

I would like to thank my parents and friends, for being so supportive of my career and life choices, as well as for making me happy.

Azmain Biswas

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# Chapter 1

### Introduction

In the ever-evolving landscape of mathematical and statistical research and application, the integration of simulation techniques has emerged as a powerful tool to unravel complex phenomena, validate theoretical frameworks, and facilitate a deeper understanding of intricate mathematical structures. Simulation is a computer-based exploratory exercise that aids in understanding how the behavior of a random or even a deterministic process changes in response to changes in input or the environment. It is essentially the only option left when exact mathematical calculations are impossible, or require an amount of effort that the user is not willing to invest. Even when the mathematical calculations are quite doable, a preliminary simulation can be very helpful in guiding the researcher to theorems that were not a priori obvious or conjectured, and also to identify the more productive corners of a particular problem. Although simulation in itself is a machine-based exercise, credible simulation must be based on appropriate theory. A simulation algorithm must be theoretically justified before we use it.

The classic theory of simulation includes such time-tested methods as the original Monte Carlo, Inverse Transform method, Accept-Reject method, Bivariate techniques from standard distributions in common use. They involve a varied degree of sophistication. Markov chain Monte Carlo is the name for a collection of simulation algorithms for simulating from the distribution of very general types of random variables taking values in quite general spaces. MCMC methods have truly revolutionized simulation because of an inherent simplicity in applying them, the generality of their scopes, and the diversity of applied problems in which some suitable form of MCMC has helped in making useful practical advances. MCMC methods are the most useful when conventional Monte Carlo is difficult or impossible to use.

Simulation depend on various theoretical aspect such as The weak law of Large Number, The Central limit theory, The sample mean and sample variance etcetera.

There are various type of simulation technique in standard simulation theory such as,

- 1. The Inverse Transform Method
- 2. Accept-Reject Algorithm
- 3. Bivariate Techniques
- 4. Ordinary Monte Carlo
- 5. Importance Sampling
- 6. Markov Chain Monte Carlo

This project work focus mainly on these simulation techniques.

In Chapter 2, we discuss about how to generate random variable both uniform and continuous, by using method like The Inverse Transform Method, Accept-Reject Algorithm, Bivariate Techniques.

In Chapter 3, we focus on Ordinary Monte Carlo and how to use it to solve problem like evaluating integration and evaluating the value of  $\pi$ .

In Chapter 4, the focus shifts to Importance Sampling and how it beneficial from Ordinary Monte Carlo by some example. Learn about how to chose optimal Importance sample distribution.

#### 1.1 Mathematical Preliminaries

**Definition 1.1.1** (Probability Space). A probability space is a triple  $(\Omega, \mathcal{F}, P)$  consisting of:

- (a) the sample space  $\Omega$  (an arbitrary non-empty set)
- (b) a non-empty collection of subsets  $\mathcal{F}$  of  $\Omega$ , called sigma field of subspace of  $\Omega$ , such taht,
  - (i)  $\Omega \in \mathcal{F}$
  - (ii) if  $A \in \mathcal{F}$ , then  $A^c \in \mathcal{F}$
  - (iii) if  $A_n \in \mathcal{F}$ ,  $n = 1, 2, ..., then <math>\cup_n = 1^{\infty} A_n \in \mathcal{F}$
- (c) a probability measure  $p: \mathcal{F} \to [0,1]$ , which is a real valued function on  $\mathcal{F}$  such that,
  - (i)  $P(A) \ge 0$  for all  $A \in \mathcal{F}$
  - (ii)  $P(\Omega) = 1$
  - (iii) if  $A_1, A_2, \ldots$  are disjoint sets in  $\mathcal{F}$ , then  $P(\bigcup_n A_n) = \sum_n P(A_n)$ .

**Definition 1.1.2** (Conditional Probability). Let A, B be general events with respect to some sample space  $\Omega$ , and suppose P(A) > 0. The conditional probability of B given A is defined as

$$P(B|A) = \frac{P(A \cap B)}{P(A)}.$$

**Theorem 1.1.1** (Bayes's Theorem). Let,  $\{A_1, A_2, \ldots, A_n\}$  be a partition of sample space  $\Omega$ . Let B be a some fixed event. Then

$$P(A_j|B) = \frac{P(B|A_j)P(A_j)}{\sum_{i=1}^{n} P(B|A_i)P(A_i)}.$$

**Definition 1.1.3** (Random Variable). Let,  $\Omega$  be a sample space corresponding to some experiment and let  $X : \Omega \to \mathbb{R}$  be a function from the sample space to the real line. Then X is called a random Variable

**Definition 1.1.4** (Cumulative Distribution Function). The cumulative distribution function (CDF) or simply distributed function, F of a random variable X is define for any real number x by

$$F(x) = P(X \le x).$$

**Definition 1.1.5** (Probability Mass Function). For a discrete random variable X we define its Probability mass function(pmf) p(x) by

$$p(x) = P(X = x)$$

and we have,

$$\sum_{i \in \Lambda} p(x_i) = 1. \text{ and } p_i \ge 0.$$

**Definition 1.1.6** (Probability Density Function). For a continuous random variable if there is a non-negative function f(x) defined for all real number x and having the property that for any set  $C \subset \mathbb{R}$ ,

 $P(X \in C) = \int_C f(x)dx.$ 

The function f is called probability density function (pdf) of the random variable X.

The relation between CDF and pdf is express by,

$$F(a) = P(X \in (-\infty, a]) = \int_{-\infty}^{a} f(x)dx.$$

**Definition 1.1.7** (Exception). If X is a random variable, then the exception or expected value of X, also called the mean of X and denoted by E(X), is define by

$$E(X) = \int x dF(x)$$

**Definition 1.1.8** (Variance). If X is a random variable with mean E(X), then the variance of X, denoted by Var(X), is define by,

$$Var(X) = E\left((X - E(X))^2\right)$$

**Theorem 1.1.2** (The Weak Law of Large Numbers). Let  $X_1, X_2, \ldots$  be a sequence of in dependent and identically distributed random variables having mean  $\mu$ , Then, for any  $\epsilon > 0$ ,

$$P\left(\left|\frac{X_1 + X_2 + \ldots + X_n}{n} - \mu\right| > \epsilon\right) \to 0$$

**Theorem 1.1.3** (The Central Limit Theorem). Suppose  $X_1, X_2, \ldots$  is a sequence of i.i.d random variables with  $E[X_i] = \mu$  and  $Var[X_i] = \sigma^2 < \infty$ . Then,

$$\lim_{n \to \infty} P\left(\frac{X_1 + \ldots + X_n - n\mu}{\sigma\sqrt{n}} < n\right) = \Phi(Z)$$

Were,  $\Phi(Z)$  denote the distribution function of a standard normal distribution.

#### Sample Mean and Sample Variance

Suppose that  $X_1, X_2, ..., X_n$  are independent random variable having the same distribution function. Let  $\mu$  and  $\sigma^2$  denote, respectively their mean and variance that is,

 $\mu = E(X_i)$  and  $\sigma^2 = Var(X_i)$ . The quantity

$$\bar{X} \equiv \sum_{i=1}^{n} \frac{X_i}{n}$$

which is the arithmetic average of the n data values, is called the *sample mean*. When the population mean  $\mu$  is unknown, the sample mean is often used to estimate  $\mu$ .

Because,

$$E(\bar{X}) = E\left(\sum_{i=1}^{n} \frac{X_i}{n}\right)$$
$$= \sum_{i=1}^{n} \frac{E(X_i)}{n}$$
$$= \frac{n\mu}{n} = \mu$$

if follows that  $\bar{X}$  is an unbiased estimator of  $\mu$ , where we say that an estimator of parameter is an unbiased estimator of that an estimator of a parameter is an unbiased estimator of that parameter if its expected value is equal to the parameter.

The quantity  $S^2$ , define by

$$S^{2} = \frac{\sum_{i=1}^{n} (X_{i} - \overline{(X)})^{2}}{n-1}$$

is called the  $sample\ variance$  .

Whish is also unbised of  $\sigma^2$ 

# Chapter 2

# Generating Random Variables

#### 2.1 Generating Discrete Random Variables

Main component of a simulation study is the ability to generate random number, where a random number represents the value of random variable uniform distribution on (0,1).

#### 2.1.1 Pseudorandom Number Generation

Random numbers were originally either manually or mechanically generated, by using spinning wheels or dice rolling or card shuffling but the modern approach is to use a computer to successively generate pseudorandom numbers.

One of the common approaches to generate pseudorandom numbers starts with an initial value  $x_0$ , called seed, and then recursively computes successive values  $x_n, n \ge 1$ , by letting

$$x_n = ax_{n-1} \text{ modulo } m \tag{2.1}$$

where a and m are given positive integers, and where Equation (2.1) means that  $ax_{n-1}$  is divided by m and remainder is taken as the value of  $x_n$ . Thus, each value of  $x_n$  is either  $0, 1, \ldots, m-1$  and the quantity  $x_n/m$  is Pseudorandom number and follows an approximation to the value of a uniform (0,1) random variable.

The approach specified by Equation (2.1) to generate random numbers is called the Multiplicative Congruential Method.

Another method is

$$x_n = (ax_{n-1} + c) \text{ modulo } m$$

this method is known as  $Mixed\ Congruential\ Generators$  or  $Linear\ congruential\ Generations\ (LCGs)$  where c is a non-negative integer.

#### 2.1.2 The Inverse Transform Method

Suppose we want to generate the value of a discrete random variable X having probability mass function

$$P(X = x_i) = p_i, i = 0, 1, \dots, \sum_i p_i = 1$$

To do this, we generate a random number from a uniform distribution (0,1) U, and set

$$X = \begin{cases} x_0 & \text{if } U < p_0 \\ x_1 & \text{if } p_0 \le U \le p_0 + p_1 \\ \vdots \\ x_j & \text{if } \sum_{i=0}^{j-1} p_i \le U \le \sum_{i=0}^{j} p_i \\ \vdots \end{cases}$$

Since, for 0 < a < b < 1,  $P(a \le U < b) = b - a$ , we have,

$$P(X = x_j) = P\left(\sum_{i=0}^{j-1} p_i \le U < \sum_{i=0}^{j} p_i\right) = p_j.$$

So, X has the desired distribution.

Example 2.1 (Bernoulli Distribution). Let,  $X \sim Ber(p)$  where p is success probability i.e. P(X=0)=1-p and P(X=1)=p and  $0 \le p \le 1$ . Then, to generate X we first generate  $U \sim U[0,1]$  then, we set

$$X = \begin{cases} 1, & \text{if } U \le p \\ 0, & \text{if } U > p \end{cases}$$

Hence, X follows Bernoulli Distribution with the parameter p.

Algorithm for Inverse Transform Algorithm for Generating Bernoulli Distribution:

STEP 1: Generate a random variable  $U \sim U[0, 1]$ .

STEP 2: If  $U \le p$  set X = 1 or set X = 0.

STEP 3: Go to STEP 1.

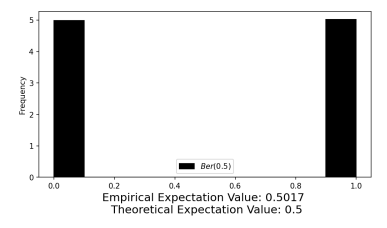


Figure 2.1: Inverse Transform method for generating Bernoulli random numbers with p=0.5

Example 2.2 (Binomial Distribution). Let,  $X \sim Bin(n,p)$  then, X has probability mass function

$$f(r) = P(X = r) = \binom{n}{r} p^r (1-p)^{n-r}, \ i = 1, 2, \dots$$

The generation of  $X \sim Bin(n,p)$  by Inverse Transform Algorithm can be tedious. We

can use the relation between Binomial and Bernoulli distribution. If  $x_i \sim Ber(p), \forall i = 1, 2, ..., n$  then,  $\sum_{i=1}^{n} x_i \sim Bin(n, p)$ .

Hence, by generating  $x_i$  n independent random variable from Bernoulli distribution and summing them we get binomial distribution

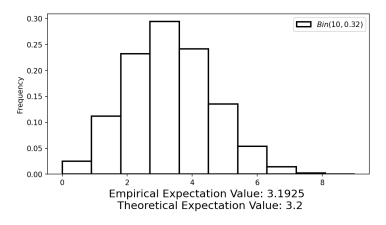


Figure 2.2: Generating binomial random numbers with n = 10 and p = 0.32

#### 2.2 Generating Continuous Random Variables

#### 2.2.1 The Inverse Transform Algorithm

To generate Continuous random variables The Inverse Transform Algorithm is very important method. It is based on a following theorem.

**Theorem 2.2.1.** Let U be a uniform (0,1) random variable. For any continuous distribution function F the random variable X defined by

$$X = F^{-1}(U)$$

has distribution F.

*Proof.* Let,  $F_X$  denote the distribution function of  $X = F^{-1}(U)$ . Then,

$$F_X(x) = P(X \le x)$$
  
=  $P(F^{-1}(U) \le x)$ 

Since, F is a cumulative distribution function it follows that F(x) is monotonic increasing function of x and range of F(x) is (0,1). Then,

$$F_X(x) = P\left(F\left(F^{-1}(U)\right) \le F(x)\right)$$
  
=  $P(U \le F(x))$   
=  $F(x)$  since  $U \sim U(0, 1)$ 

The above theory tells us we can generate a random variable X from the continuous distribution function F by generating a random number  $U \sim U(0,1)$  and setting  $X = F^{-1}(U)$ .

Example 2.3 (Exponential Distribution). Suppose we want to generate a random variable  $x \sim Exp(\lambda)$ , then its probability density function is

$$f(x) = \lambda e^{-\lambda x}.$$

Hence, The cumulative distribution function is,

$$F(x) = 1 - e^{\lambda x}$$

if we let  $x = F^{-1}(u)$ , then,

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

Hence, we can generate an exponential random variable with parameter 1 by generating a uniform (0,1) random number U and then setting

$$X = F^{-1}(U) = -\frac{\ln(1-U)}{\lambda}.$$

We see that if  $U \sim U(0,1)$  then also  $1-U \sim U(0,1)$  thus  $\ln(1-U)$  has the same distribution as  $\ln(U)$  so,

$$X = F^{-1}(U) = -\frac{\ln(U)}{\lambda}.$$

will also work. If we use second expression then the algorithm will take less computing power hence less time.

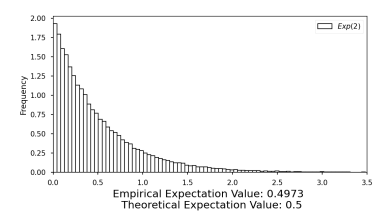


Figure 2.3: Inverse Transform method for generating Exp(2)

Example 2.4 (Gamma Distribution). Let  $X \sim G(n, \lambda)$  Then, its probability mass function is given by,

$$f(x) = \frac{1}{\Gamma(n)} \lambda^n x^{n-1} e^{-\lambda x}$$

We know if  $X_i \sim Exp(\lambda) \forall i = 1, 2, ..., n$  then  $Y = \sum_i X_i \sim G(n, \lambda)$ . As,

$$M_Y(t) = E\left[e^{tY}\right] = E\left[e^{\sum_{i=1}^n X_i t}\right] = E\left[\prod_{i=1}^n e^{X_i t}\right]$$

$$= \prod_{i=1}^n E\left[e^{X_i t}\right] \text{ As all } X_i \text{ are independent}$$

$$= \prod_{i=1}^n \frac{\lambda}{\lambda - t} = \left(\frac{\lambda}{\lambda - t}\right)^n$$

Then, Generating n number of  $X_i \sim Exp(\lambda)$  and summing them we can easily generate a random variable which follows gamma distribution

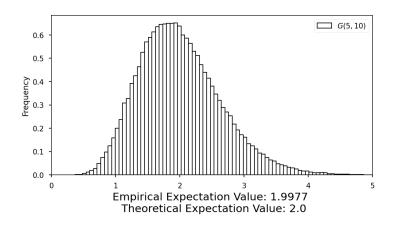


Figure 2.4: G(10,5) generated by summing of Exp(5)

#### 2.2.2 Accept - Reject Method

The accept—reject method is useful when it is difficult to directly simulate f(x) but we can generate another density g(x) such that f(x)/g(x) is uniformly bounded and it is much easier to simulate g(x). We simulate X from g, and retain it or toss it according to a probability proportional to f(x)/g(x). Because an X value is either retained or discarded, depending on whether it passes the admission rule, the method is called the accept—reject method. The density g(x) is called the envelope density.

The method proceeds as follows,

STEP 1: Find a density g and a finite constant c such that  $\frac{f(x)}{g(x)} \le c \ \forall x$ .

STEP 2: Generate  $X \sim g$ .

STEP 3: Generate  $U \sim U(0,1)$ , independent of X.

STEP 4: Retain this generated value X if  $U \leq \frac{f(x)}{cg(x)}$ .

STEP 5: Repeat the same until the required number of n values of X has been obtained.

The following theorem supports the method.

**Theorem 2.2.2.** Let  $X \sim g$ , and U, independent of, be a distributed as U[0,1]. Then the conditional density of X given that  $U \leq \frac{f(X)}{cg(X)}$  is f.

*Proof.* Denote the CDF of f by F. Then,

$$P\left(X \le x | U \le \frac{f(X)}{cg(X)}\right) = \frac{P\left(X \le x, U \le \frac{f(X)}{cg(X)}\right)}{P\left(U \le \frac{f(x)}{cg(x)}\right)}$$

$$= \frac{\int_{-\infty}^{x} \int_{0}^{\frac{f(t)}{cg(t)}} g(t) du dt}{\int_{-\infty}^{\infty} \int_{0}^{\frac{f(t)}{cg(t)}} g(t) du dt}$$

$$= \frac{\int_{-\infty}^{x} f(t) dt}{\int_{-\infty}^{\infty} f(t) dt} = \frac{F(x)}{1} = F(x).$$

Example 2.5 (Generating a Normal Random Variable). To generate a standard normal variable Z i.e.  $Z \sim N(0,1)$ , note first that the absolute value of Z has probability density function

$$f(x) = \frac{2}{\sqrt{2\pi}} e^{-\frac{x^2}{2}} \ 0 \le x \le \infty.$$
 (2.2)

Then, we can choose g as the exponential density function with mean 1 i.e.

$$g(x) = e^{-x} \ 0 \le x \le \infty$$

Now,

$$\frac{f(x)}{g(x)} = \sqrt{\frac{2}{\pi}}e^{x-\frac{x^2}{2}}$$

and so the maximum value of f(x)/g(x) occurs at the value of x that maximize  $x - x^2/2$  hence x = 1 so we take

$$c = \max_{x} \frac{f(x)}{g(x)} = \frac{f(1)}{g(1)} = \sqrt{\frac{2e}{\pi}}.$$

Now,

$$\frac{f(x)}{cg(x)} = \exp\left(x - \frac{x^2}{2} - \frac{1}{2}\right) = \exp\left(\frac{-(x-1)^2}{2}\right)$$

Then, its follows that we can generate the absolute value of a standard normal random variable as follows:

STEP 1: Generate  $X \sim Exp(1)$ .

STEP 2: Generate  $U \sim U(0,1)$ , independent of X.

STEP 3: If  $U \leq \exp(-(X-1)^2/2)$ , retain X, Otherwise, return to Step 1.

Once, we have simulated a random variable X having density function as in Equation (2.2) we can obtain a standard normal Z by letting Z be equally likely to be either X or -X. In Step 3, the value X is accepted if  $U \le \exp(-(X-1)^2/2)$ , which is equivalent to  $-\ln U \ge (X-1)^2/2$ . However, in Example 2.3 we have seen that  $-\ln U \sim Exp(1)$  When  $U \sim U(0,1)$ .

So, summing up, we can generate the standard normal random variable Z as follows: STEP 1: Generate independent  $X_1, X_2 \sim Exp(1)$ 

STEP 2: If  $X_2 \ge (X_1 - 1)^2/2$  retain  $X_1$ . Otherwise, return to Step 1.

STEP 3: Generate  $U \sim U(0,1)$  and set,

$$Z = \begin{cases} X_1 & \text{if } U \le \frac{1}{2}, \\ X_1 & \text{if } U > \frac{1}{2}. \end{cases}$$

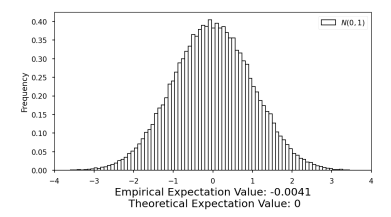


Figure 2.5: Generating N(0,1) with Accept - Reject method

If we want to generate normal random variable to have mean  $\mu$  and variance  $\sigma^2$ , just take  $\mu + \sigma Z$ .

Example 2.6 (Generating Beta Distribution). If  $\alpha$  and  $\beta$  are both getter then 1, then Beta density is uniformly bounded and its maximum attain at  $\frac{\alpha-1}{\alpha+\beta-2}$ . As a result the U[0,1] density can be served as an envelope density for generating such Beta distribution by using accept-reject method. Precisely, generate  $U, X \sim U[0,1]$  (independently), and retain the value if  $U \leq \frac{f(X)}{\sup_x f(X)}$ , where,

$$f(X) = \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} x^{\alpha - 1} (1 - x)^{\beta - 1}, 0 < x < 1.$$

Because

$$\sup_{x} f(X) = f\left(\frac{\alpha - 1}{\alpha + \beta - 2}\right) = \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} \frac{(\alpha - 1)^{\alpha - 1}(\beta - 1)^{\beta - 1}}{(\alpha + \beta - 2)^{\alpha + \beta - 2}}$$

The algorithm finally works out as follows:

STEP 1: Generate independent  $U, X \sim U[0, 1]$ .

STEP 2: Retain the value X if,

$$U \le \frac{X^{\alpha - 1} (1 - X)^{\beta - 1} (\alpha + \beta - 2)^{\alpha + \beta - 2}}{(\alpha - 1)^{\alpha - 1} (\beta - 1)^{\beta - 1}}.$$

Otherwise, return to STEP 1.

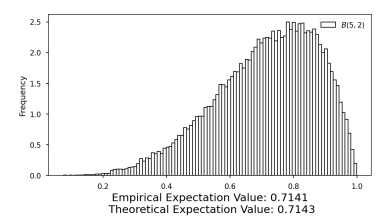


Figure 2.6: Generating Beta(5,2) with accept-reject method

An issue about an accept-reject method is the acceptance rate. Our goal make it as large as possible to increase the efficiency of the method. This can be achieved by choosing c to be smallest possible number, described in the result bellow.

**Theorem 2.2.3** (Acceptance Rate). For an accept-reject scheme, the probability that an  $X \sim g$  is acceded is  $\frac{1}{c}$ , and is maximized when c is chosen to be  $c = \sup_x \frac{f(x)}{g(x)}$ .

Proof.

$$P\left(U \le \frac{f(x)}{cg(x)}\right) = \int_{-\infty}^{\infty} \int_{0}^{\frac{f(x)}{cg(x)}} g(t)dudt$$
$$= \int_{-\infty}^{\infty} \frac{f(t)}{cg(t)} g(t)dt = \int_{-\infty}^{\infty} \frac{f(t)}{c} dt = \frac{1}{c}.$$

Because any c that can be chosen must be at least as large as  $\sup_x \frac{f(x)}{g(x)}$ , obviously 1/c is maximized by choosing  $c = \sup_x \frac{f(x)}{g(x)}$ .

In the Example 2.5 for N(0,1) the acceptance rate is  $\sqrt{\frac{\pi}{2e}} = 0.7601$ . And in the Example 2.6 for Beta(5,2) the acceptance rate is 0.4069

#### 2.2.3 Bivariate Techniques

Let X and Y be independent slandered normal random variable and let R and  $\theta$  denote the polar coordinates of vector (X, Y). That is,

$$R^2 = X^2 + Y^2$$
$$\tan \theta = \frac{Y}{X}$$

Since X and Y are independent, their joint density is the product of their individual densities and thus given by

$$f(x,y) = \frac{1}{\sqrt{2\pi}} e^{-x^2/2} \frac{1}{\sqrt{2\pi}} e^{-y^2/2}$$
$$= \frac{1}{2\pi} e^{-(x^2+y^2)/2}$$

To determine the joint density of  $R^2$  and  $\Theta$  - call it  $g(d,\theta)$  we make the change of variables

$$d = x^2 + y^2$$
,  $\theta = \tan^{-1}\left(\frac{y}{x}\right)$ 

Then the joint density function of d and  $\Theta$  is,

$$g(d,\theta) = |J|f(x,y)$$

$$= |J|\frac{1}{2\pi}e^{-(x^2+y^2)/2}$$
(2.3)

where,

$$J = \begin{vmatrix} \frac{\partial x}{\partial t} & \frac{\partial x}{\partial \theta} \\ \frac{\partial y}{\partial t} & \frac{\partial y}{\partial \theta} \end{vmatrix} = \frac{1}{2}.$$

Then, replacing the value of  $x = \sqrt{d}\sin(\theta)$  and  $y = \sqrt{d}\cos(\theta)$  in Equation (2.3) we get,

$$g(d,\theta) = \frac{1}{2} \frac{1}{2\pi} e^{-d/2}, \quad 0 < d < \infty, 0 < \theta < 2\pi.$$
 (2.4)

As  $g(d,\theta)$  is equal to product of the product of Exp(1/2) density and  $U(0,2\pi)$ , it follows that,

 $R^2$  and  $\Theta$  are independent, with  $R^2 \sim Exp(1/2)$  and  $\Theta \sim U(0, 2\pi)$ 

Hence to generate a pair of independent slandered normal random variables X and Y by generating  $R^2$  and  $\Theta$  in polar coordinates and then transform back to rectangular coordinates. Hence the algorithm is:

STEP 1: Generate random number  $U_1, U_2 \sim U(0, 1)$ .

STEP 2:  $R^2 = -2 \ln U_1$  and  $\Theta = 2\pi U_2$ .

STEP 3: Now let,

$$X = R\cos\Theta = \sqrt{-2\ln U_1}\cos(2\pi U_2) \tag{2.5}$$

$$Y = R\sin\Theta = \sqrt{-2\ln U_1}\sin(2\pi U_2). \tag{2.6}$$

The transformation given by Equation (2.5) and Equation (2.6) are known as Box-Muller transformation.

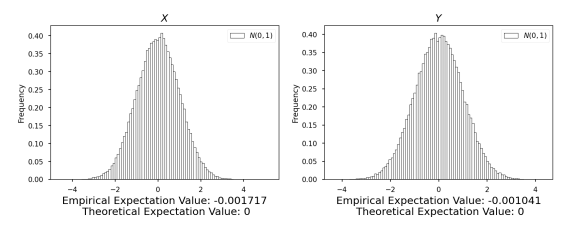


Figure 2.7: Generating independent  $X, Y \sim N(0, 1)$  with polar method

# Chapter 3

## Monte Carlo Method

#### 3.1 Ordinary Monte Carlo Simulation

Polish-American mathematician Stanislaw Ulam, recovering from an illness, was playing a lot of solitary card game. He wanted to calculate the probability of winning and quickly it is impossible to calculate analytically. Then he thought about playing lots of hands counting number of wins, but decided it will take years. After falling several times he asked Von Neumann to build a program to simulate solitary card game in ENIAC. Then Around 1940 they used Monte Carlo simulation in Manhattan Project in which physicists wanted to understand how the physical properties of neutrons would be affected by various possible scenarios following a collision with a nucleus.

The basis for Monte Carlo is Law of Large Number. If we simulate large number  $X_1, X_2, \ldots$  iid copes of random variable X Then we can approximate the true value E(f(X)) by simple mean  $\frac{1}{n} \sum_{i=1}^{n} f(X_i)$ . Here in Monte Carlo Random Sampling play the key factor for good estimation of E(f(X)).

#### 3.1.1 Examples

#### **Evaluate Integrals using Monte Carlo simulation**

The application of Monte Carlo is to computation of integrals. Let g(x) be a function and suppose we wanted to compute I where

$$I = \int_0^1 g(x)dx$$

To compute the value of I, note that if  $U \sim U[0,1]$  then we can express I as

$$I = E[g(U)]$$

If  $U_1, \ldots, U_n$  are independent uniform (0,1) random variables, it thus follows that the random variables  $g(U_1), \ldots, g(U_n)$  are independent and identically distributed random variable having mean I. Therefore, by law of large numbers, its follows that, with probability,

$$\sum_{i=0}^{n} \frac{g(U_i)}{n} \to E(g(U)) = I \text{ as } k \to \infty$$

Hence we can approximate I by generating a large number of random numbers  $U_i$  and taking as our approximation the average value of  $g(U_i)$ .

If we wanted to compute

$$I = \int_{a}^{b} g(x)dx$$

then , by taking the substitute y = (x - a)/(b - a), dy = dx/(b - a), we see that

$$I = \int_0^1 g(a + [b - a]y)(b - a)dy$$
$$= \int_0^1 h(y)dy$$

Where h(y) = (b - a)g(a + [b - a]y). Thus we can approximate I by continually generating random numbers and then taking the average value of h evaluated at these random numbers.

Similarly, if we wanted

$$I = \int_0^\infty g(x)dx$$

we could apply the substitution y = 1/(x+1),  $dy = -dx/(x+1)^2 = -y^2 dx$ , to obtain the identity

$$I = \int_0^1 h(y) dy$$

where,

$$h(y) = \frac{g(\frac{1}{y} - 1)}{y^2}$$

Using this technique we can also evaluate multidimensional integrals. Suppose that g is a function with n-dimension argument and we are interested in computing

$$I = \int_0^1 \int_0^1 \dots \int_0^1 g(x_1, x_2, \dots, x_n) dx_1 dx_2 \dots dx_2.$$

Then, we can express I as

$$I = E(g(U_1, U_2, \dots U_n))$$

where  $U_1, U_2, \dots U_n \sim U[0, 1]$  Hence if we generate k independent sets, each consisting of n independent U[0, 1] random variable

$$U_1^1 \dots U_n^1$$

$$U_1^2 \dots U_n^2$$

$$\vdots$$

$$U_1^k, \dots, U_n^k$$

then, since the random variables  $g(U_1^i, \ldots, U_n^i)$ ,  $i = 1, 2, \ldots, k$  are all independent and identically distributed random variable with mean I, we can estimate I by  $\sum_{i=1}^k g(U_1^i, \ldots, U_n^i)/k$ .

Example 3.1. Suppose we want to integrate,

$$I = \int_0^1 e^{-\frac{x^2}{2}} dx.$$

Then we can say that,

$$I = E(e^{-\frac{U^2}{2}})$$

where,  $U \sim U[0,1]$ . Then simulating a large number of  $U_1, U_2, \dots, U_n \sim U[0,1]$  and calculating,

$$\sum_{i=1}^{n} \frac{e^{-\frac{U_i^2}{2}}}{n}$$

we can evaluate I.

Hence the algorithm is:

STEP 1: Generate  $U \sim U[0,1]$ . STEP 2: Calculate  $e^{-\frac{U^2}{2}}$  and retain it. goto STEP 1.

STEP 3: After large number of iteration evaluate the average.

Monte Carlo sample size	Monte Carlo Estimate of $I = 0.8556$
50	0.8555
100	0.8558
1000	0.8555
10000	0.8556
100000	0.8556

Table 3.1: Monte Carlo Integration of  $e^{-x^2/2}$ .

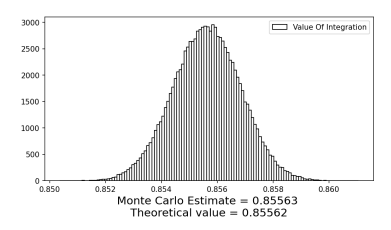


Figure 3.1: Monte Carlo Integration of  $e^{-x^2/2}$ 

Here, we see by the time the Monte Carlo sample size is 100000, we get fairly accurate estimates for the value I.

#### The Estimation of $\pi$

Suppose that the random vector (X, Y) is uniformly distribution in the square of area 4 centered at the origin. That is, it is a random point in the region specified in Figure 3.2. Let us consider now the probability that this random point in the square in contained within the inscribed circle of radius 1 like the Figure 3.3.

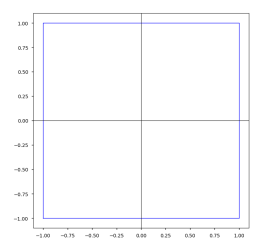


Figure 3.2: Square

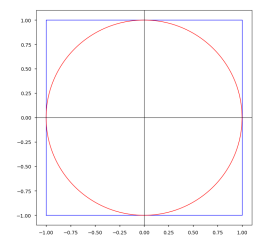


Figure 3.3: Circle within Square

Note that since (X,Y) is uniformly distributed in the square it follows that

$$\begin{split} P((X,Y) \text{ is in the circle }) &= P(X^2 + Y^2 \leq 1) \\ &= \frac{\text{Area of the circle}}{\text{Area of the square}} = \frac{\pi}{4} \end{split}$$

Hence we generate a large number of points in the square, the proportion of points that fall within the circle will be approximately  $\pi/4$ . Now, if X and Y were independent

and both were uniformly distributed over (-1,1), their joint density would be

$$f(x,y) = f(x)f(y)$$

$$= \frac{1}{2} \times \frac{1}{2}$$

$$= \frac{1}{4}, -1 \le x \le 1, -1 \le y \le 1$$

Since, the density function of (X,Y) is constant in the square, it thus follows that (X,Y) is uniformly distributed in the square. Now, if  $U \sim U[0,1]$  then  $2U \sim U[0,2]$  and so  $2U - 1 \sim U[-1,1]$ . Therefore, if we generate ranom numbers  $U_1$  and  $U_2$  and set  $X = 2U_1 - 1$  and  $Y = 2U_2 - 1$ , and define,

$$I = \begin{cases} 1 & \text{if } X^2 + y^2 \le 1\\ 0 & \text{otherwise} \end{cases}$$

Then,

$$E(I) = P(X^2 + y^2 \le 1) = \frac{\pi}{4}.$$

Hence the Algorithm for estimating  $\pi$  is:

STEP 1: Set Circle = 1.

STEP 2: Generate  $U_1, U_2 \sim U[0, 1]$ 

STEP 3: If  $(2U_1-1)^2+(2U_2-1)^2\leq 1$  Set Circle = Circle + 1, Otherwise return to

STEP 2. STEP 4: After simulating N time, set Area of Circle = Circle/N

Monte Carlo sample	Monte Carlo Estimate of $\pi$	
50	2.9600	
100	3.0800	
1000	3.1200	
10000	3.1428	
100000	3.1397	
1000000	3.1394	
10000000	3.1414	

Table 3.2: Monte Carlo Estimates of  $\pi$ 

Here, we see by the time the Monte Carlo sample size is 10000000, we get fairly accurate estimates for  $\pi$ 

#### 3.2 Importance Sampling

There are two different ways to think about importance sampling. The more traditional one is to go back to the primary problem that Monte Carlo wants to solve, namely to approximate the value of an expectation  $\mu = \int \phi_0(x)dF_0(x)$  for some function  $\phi_0$  and some CDF  $F_0$ . However,  $(\phi_0, F_0)$  is not the only pair  $(\phi, F)$  for which  $\int \phi(x)dF(x)$  equals the specific number  $\mu$ . Indeed, given any other CDF  $F_1$ ,

$$\mu = \int \phi_0(x) dF_0(x)$$

$$= \int \phi_0(x) \frac{dF_0}{dF_1}(x) dF_1(x)$$

$$= \int \lambda(x) \phi_0(x) dF_1(x).$$

where  $\lambda(x) = \frac{dF_0}{dF_1}(x)$ . If  $F_0$ ,  $F_1$  have densities  $f_0$ ,  $f_1$ , then  $\lambda(x) = \frac{f_0(x)}{f_1(x)}$ ; if  $F_0$ ,  $F_1$  have respective pmfs  $f_0$ ,  $f_1$ , then also  $\lambda(x) = \frac{f_0(x)}{f_1(x)}$  This raises the interesting possibility that we can sample from a general  $F_1$ , and subsequently use the usual Monte Carlo estimate

$$\hat{\mu} = \frac{1}{n} \sum_{i=1}^{n} \lambda(X_i) \phi_0(X_i) = E_{F_1}[\lambda(X_i) \phi_0(X_i)].$$

where  $X_1, X_2, \ldots, X_n$  is Monte Carlo sample from  $F_1$ . Importance sampling poses the problem of finding an optimal choice of  $F_1$  for which to sample, so that  $\hat{\mu}$  has the smallest possible variance. The distribution  $F_1$  hat ultimately gets chosen is called the *importance* sampling distribution.

We can visualize this method by an example.

Example 3.2. Suppose we want to evaluate

$$I = \int_0^{10} e^{-2|x-5|} dx.$$

doing it analytically we get I = 0.9999.

Now, suppose  $\phi(x) = e^{-2|x-5|}$  then we want to evaluate

$$I = \int_0^{10} \phi(x) dx$$

Now,

$$I = \int_{0}^{10} \phi(x)dx$$

$$= \int_{0}^{10} \phi(x) \frac{10}{10} dx$$

$$= \int_{0}^{10} 10 \times \phi(x) \frac{1}{10} dx = \int_{0}^{10} 10 \phi(x) f_{0}(x) dx \text{ where } f_{0}(x) \text{ pdf of } U(0,1)$$

$$= E_{U}[10 \times \phi(U)] \text{ where, } U \sim U(0,10).$$
(3.1)

By Ordinary Monte Carlo technique we can estimate I by  $\frac{1}{N}\sum_{i=1}^{N}10\times\phi(U_i)$  where

 $U_i \sim U(0, 10)$  for i = 1, 2, ..., N for the large number of N.

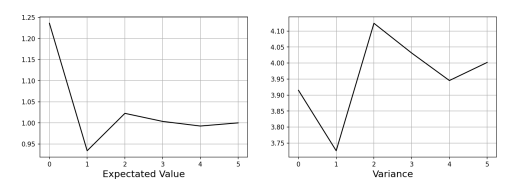


Figure 3.4: Monte Carlo integration of  $\int_0^{10} e^{-2|x-5|} dx$ .

Sample Size	Estimated Value I=0.9999	Variance
10	2.3243	5.6912
100	1.0372	3.6717
1000	0.8871	3.5543
10000	1.0467	4.2416
100000	1.0053	4.0089
1000000	1.0054	4.0346

Table 3.3: Monte Carlo integration of  $\int_0^{10} e^{-2|x-5|} dx$ .

Here we can see that for sample size 1M, we get a pretty good estimation of I with less then 0.6% error. But the variance is very high. If we see at left of Figure 3.5 we can see we are taking unnecessary value from low frequency part of  $\phi(x)$  that is, from extreme left and right. If we choose an importance sampling distribution that has similar curve as  $\phi(x)$ , then we can estimate I with low variance. If we choose importance sampling distribution as N(5,1) then we see from right of Figure 3.5 it has similar pattern as  $\phi(x)$ .

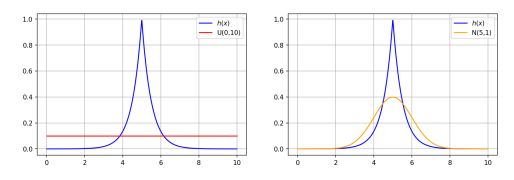


Figure 3.5: h(x) with U(0, 10) and N(5, 1)

Let,  $f_1(x)$  is pdf of N(0,1) then, from Equation (3.1)

$$I = \int_{0}^{10} 10\phi(x)f_{0}(x)dx$$

$$= \int_{0}^{10} 10\phi(x)\frac{f_{0}(x)}{f_{1}(x)}q(x)dx$$

$$= E_{X} \left[10\phi(X)\frac{f_{0}(X)}{f_{1}(X)}\right] \text{ where } X \sim N(5,1)$$

$$= E_{X} \left[10\phi(X)\lambda(X)\right] \text{ where } X \sim N(5,1)$$

Where,  $\lambda(x) = \frac{f_0(x)}{f_1(x)}$  here  $f_0(x)$  is pdf of U(0,1) and  $f_1(x)$  is pdf of N(5,1). Now using usual Monte Carlo estimate

$$I = \frac{1}{N} \sum_{i=1}^{N} 10\phi(X_i)\lambda(X_i)$$

where  $X_1, X_2, \ldots, X_N$  is Monte Carlo sample from N(5, 1).

Sample Size	Estimated Value (I=0.9999)	Variance
10	1.2473	0.3812
100	0.9292	0.2593
1000	1.0018	0.3550
10000	1.0072	0.3603
100000	1.0039	0.3595
1000000	0.9999	0.3580

Table 3.4: Evaluating  $\int_0^{10} e^{-2|x-5|} dx$  using Importance Sampling.

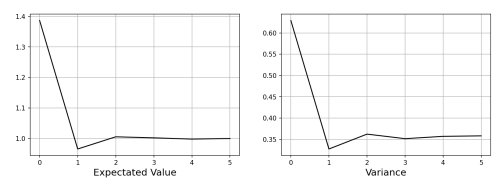


Figure 3.6: Evaluating  $\int_0^{10} e^{-2|x-5|} dx$  using Importance Sampling.

Here we can see that the estimation of I is pretty close and variance is also lower the Original Monte Carlo method.

A more contemporary view of importance sampling is that we do not approach importance sampling as an optimization problem, but because the circumstances force us to consider different sampling distributions F.

Now, we also assume that  $F_0$ ,  $F_1$  both have densities, say  $f_0$ ,  $f_1$ . If  $F_0$ ,  $F_1$  are both discrete then the notation only change but the argument is same. Suppose then  $f_i(x) = \frac{h_i(x)}{c_i}$ , i = 0, 1, where the assumption is that  $h_0$ ,  $h_1$  are completely known and also

computable, but  $c_0$ ,  $c_1$  are unknown and are not even computable. Then, as we showed above, for any function  $\phi$  for which the expectation  $E_{F_0}[\phi(X)]$  exist,

$$\mu = E_{F_0}[\phi(X)] := \int \frac{f_0(x)}{f_1(x)} \phi(x) f_1(x) dx$$

$$= \frac{c_1}{c_0} \int \frac{\phi(x) h_0(x)}{h_1(x)} f_1(x) dx$$

$$= \frac{c_1}{c_0} E_{F_1} \left( \frac{\phi(X) h_0(X)}{h_1(X)} \right).$$

This is a useful reduction, but we have to deal with the fact that ratio  $\frac{c_1}{c_0}$  is not known to us. Now, if we use the special function  $\phi(x) \equiv 1$ , the same representation above gives us

$$1 = \frac{c_1}{c_0} E_{F_1} \left( \frac{h_0(X)}{h_1(X)} \right)$$

$$\implies \frac{c_1}{c_0} = \frac{1}{E_{F_1} \left( \frac{h_0(X)}{h_1(X)} \right)}$$

and because  $h_0$ ,  $h_1$  are explicitly known to us, we have a way to get rid of the quotient  $\frac{c_1}{c_0}$  and write the final *importance sampling identity* 

$$E_{F_0}[\phi(x)] = \frac{E_{F_1}\left(\frac{\phi(X)h_0(X)}{h_1(X)}\right)}{E_{F_1}\left(\frac{h_0(X)}{h_1(X)}\right)}$$

We can now use an available Monte Carlo sample  $X_1, X_2, \ldots, X_n$  from  $F_1$  to find Monte Carlo estimates for  $\mu = E_{F_0}[\phi(x)]$ 

The basic plug-in estimate for  $\mu$  is the so-called ratio estimate

$$\hat{\mu} = \frac{\sum_{i=1}^{n} \frac{\phi(X_i)h_0(X_i)}{h_1(X_i)}}{\sum_{i=1}^{n} \frac{h_0(X_i)}{h_1(X_i)}}.$$

Example 3.3 (Binomial Bayes problem with an Atypical Prior). Suppose  $X \sim Bin(m, p)$  for some fixed m and p has the prior density  $c\sin^2(\pi p)$ , where c is a normalizing constant. Throughout the example, c denotes a generic constant, and is not intended to mean the same constant at every use.

The posterior density of p given X = x is

$$\pi(p|X=x) = cp^x(1-p)^{m-x}\sin^2(\pi p), \quad 0$$

The problem is to find the posterior mean

$$\mu = c \int_0^1 p[cp^x (1-p)^{m-x} \sin^2(\pi p)] dp.$$

We use importance sampling to approximate the value of  $\mu$ . Towards this, choose

$$\phi(p) = p, \ h_0(p) = p^x (1-p)^{m-x} \sin^2(\pi p), \ h_1(p) = p^x (1-p)^{m-x},$$

so that if  $p_1, p_2, \ldots, p_n$  are samples from  $F_1$ , (i.e.  $p_i \sim Beta(x+1, m-x+1)$ ), then the importance sampling estimate of the posterior mean  $\mu$  is

$$\hat{\mu} = \frac{\sum_{i=1}^{n} \frac{\phi(p_i)h_0(p_i)}{h_1(p_i)}}{\sum_{i=1}^{n} \frac{h_0(p_i)}{h_1(p_i)}}$$
$$= \frac{\sum_{i=1}^{n} p_i \sin^2(\pi p_i)}{\sum_{i=1}^{n} \sin^2(\pi p_i)}.$$

Note that we did not need to calculate the normalizing constant in the posterior density. We take m = 100, x = 45 for specificity.

Sample Size	Importance	Sampling	Variance
	Estimate of $\mu$		
20	0.4821		0.5465
50	0.4629		0.5162
100	0.4597		0.4929
250	0.4635		0.4902
500	0.4585		0.4962

Table 3.5: Importance Sampling Estimates of  $\mu$  for Different Sample Sizes

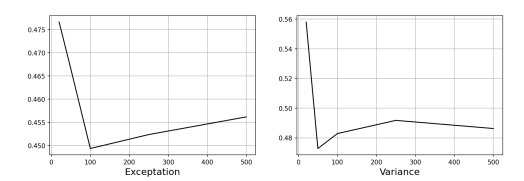


Figure 3.7: Exception and variance graph of Binomial Bayes problem with an Atypical Prior

#### 3.2.1 Optimal Importance Sampling Distribution

We now address the question of the optimal choice of the importance sampling distribution. There is no unique way to define what an optimal choice means. We formulate one definition of optimality and provide an optimal importance sampling distribution. The optimal choice would not be practically usable, as we shown. However, the solution still gives useful insight.

**Theorem 3.2.1.** Consider the importance sampling estimator  $\hat{\mu} = \frac{1}{n} \sum_{i=1}^{n} \lambda(X_i) \phi(X_i)$  for  $\mu = \int \phi(x) f_0(x) dx$ , where  $\lambda(x) = \frac{f_0(x)}{f_1(x)}$ , and  $X_1, \ldots, X_n$  are iid observations from  $F_1$  Assume that  $\phi(x) \geq 0$ , and  $\mu > 0$ . Then,  $Var_{F_1}(\hat{\mu})$  is minimized when  $f_1(x) = \frac{\phi(x) f_0(x)}{\mu}$ .

*Proof.* Because  $X_1, \ldots, X_n$  is iid, so are  $\lambda(X_1)\phi(X_1), \ldots, \lambda(X_n)\phi(X_n)$ , and hence,

$$Var_{F_1}(\hat{\mu}) = \frac{1}{n} Var_{F_1}(\lambda(X_1)\phi(X_1)).$$

Clearly, this is minimized when with probability one under  $F_1$ ,  $\lambda(X_1)\phi(X_1)$  is constant, say k. The constant k must be equal to the mean of  $\lambda(X_1)\phi(X_1)$ , that is,

$$k = \int \lambda(x)\phi(x)f_1(x)dx$$
$$= \int \frac{\phi(x)f_0(x)}{f_1(x)}f_1(x)dx$$
$$= \int \phi(x)f_0(x)dx = \mu.$$

Therefore, the optimal importance sampling density satisfies  $\lambda(x)\phi(x) = \mu$  hence,

$$f_1(x) = \frac{\phi(x)f_0(x)}{\mu}.$$

This is not usable in practice, because it involves  $\mu$ , which is precisely the unknown number we want to approximate. However, the theoretically optimal solution suggests that the importance sampling density should follow key properties of the unnormalized function  $\phi(x)f_0(x)$ . For example,  $f_1$  should have the same shape and tail behavior as  $\phi(x)f_0(x)$ .

We have seen this phenomena in the Example 3.3. Because Graph of  $\phi(x)h_0(x)$  and  $h_1(x)$  in Figure 3.8, they both have the same key properties.

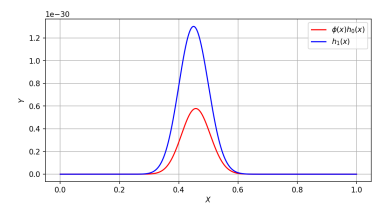


Figure 3.8: Graph of  $\phi(x)h_0(x)$  and  $h_1(x)$ 

# Chapter 4

## Markov Chain Monte Carlo Methods

Markov Chain Monte Carlo(MCMC) is a powerful collection of algorithms that enable us to simulate from complicated distributions using Markov chains. When the target distributions is an unconventional one, or it is known only up to a normalizing constant that is  $f(x) = \frac{h(x)}{c}$  for some explicit function h but only an implicit normalizing constant c, because c can not be computed exactly, the standard simulation techniques are difficult to apply or even not applicable in that case Markov Chain Monte Carlo(MCMC) comes in to play. The basic idea for MCMC is to construct a Markov Chain whose stationary distribution is the distribution of interest.

MCMC is widely used algorithms it is primarily used for calculating numerical approximations of multi-dimensional integration for example is Bayesian statistic, computational physics, computational biology. In Bayesian statistic, Markov Chain Monte Carlo method are typically used to calculate moments and posterior distribution.

To understand Markov Chain Monte Carlo(MCMC) we have to understand about Markov Chains.

#### 4.1 Markov Chains

Fill if later

#### 4.2 The Metropolis-Hastings Algorithm

Metropolis-Hastings algorithm is one of the most used Markov Chain Monte Carlo (MCMC) algorithm. The Metropolis algorithm was first introduced by Nicholas Metropolis in 1953 in his paper entitled "Equation of State Calculations by Fast Computing Machines", with Arianna W. Rosenbluth, Marshall Rosenbluth, Augusta H. Teller and Edward Teller. Arianna Rosenbluth wrote the first full implementation of Metropolis Algorithm for Mathematical Analyzer Numerical Integrator and Automatic Computer Model I (MANIAC 1) which was an early computer built under the direction of Nicholas Metropolis at the Los Alamos Scientific Laboratory.



Figure 4.1: MANIAC 1 one of the earliest computer.

For may years this algorithm is was simply known as *Metropolis Algorithm*, later in 1970 W.K. Hastings introduce more general version of this algorithm in his paper "Monte Carlo Sampling Methods Using Markov Chains and Their Applications". This generalized Metropolis algorithm is known as *Metropolis-Hastings algorithm* (MH algorithm).

Suppose we want to simulate a random variables or sequence of random variable with probability mass function

$$\pi(\theta) = \frac{f(\theta)}{K} \tag{4.1}$$

where K is normalizing constant which is unknown or difficult to compute.

One way to simulate  $\pi(\theta)$  is to constant a Markov Chain that is easy to simulate and whose limiting distribution is  $\pi(\theta)$ . The *Metropolis-Hastings algorithm* do exactly this. MH algorithm constant a time-reversible Markov Chain with desired limiting probabilities.

#### 4.2.1 Algorithm for Metropolis-Hastings

- 1. Start with any **initial state**  $\theta_0$  satisfying  $f(\theta) > 0$ .
- 2. Using a current state  $\theta$ , sample candidate state  $\theta'$  for some jumping distribution  $q(\theta, \theta') = q(\theta'|\theta)$ , which is the probability of jumping to  $\theta'$  provided the current state in  $\theta$ .
- 3. Given the candidate state  $\theta'$  calculate the **acceptance probability**  $\alpha(\theta, \theta')$  by,

$$\alpha(\theta, \theta') = \min\left(\frac{\pi(\theta')q(\theta', \theta)}{\pi(\theta)q(\theta, \theta')}, 1\right) = \min\left(\frac{f(\theta')q(\theta', \theta)}{f(\theta)q(\theta, \theta')}, 1\right)$$

4. Accept the candidate point with probability  $\alpha$ .

We can summarize the Metropolis-Hastings Algorithm as first computing,

$$\alpha(\theta_t, \theta_{t+1}) = \min \left( \frac{f(\theta_{t+1})q(\theta_{t+1}, \theta_t)}{f(\theta_t)q(\theta_t, \theta_{t+1})} \right)$$

and then accepting the candidate point  $\theta_{t+1}$  with probability  $\alpha$ . This generates a Markov Chain  $(\theta_0, \theta_1, \ldots, \theta_t, \ldots)$ , as the transition probabilities from  $\theta_t$  to  $\theta_{t+1}$  depends only on  $\theta_t$  and not on  $(\theta_0, \theta_1, \ldots, \theta_{t-1})$ .

#### 4.2.2 Metropolis-Hastings Algorithm as a Markov Chain

To determine Metropolis-Hastings Sampling generates a Markov Chain whose stationary distribution is candidate distribution  $\pi(\theta)$  if the Metropolis-Hastings transition kernel,

$$P(\theta_1 \to \theta_2) = P(\theta_1, \theta_2) = q(\theta_1, \theta_2) \alpha(\theta_1, \theta_2) = q(\theta_1, \theta_2) \times \min\left(\frac{f(\theta')q(\theta', \theta)}{f(\theta)q(\theta, \theta')}, 1\right)$$
(4.2)

is time-reversible and satisfies

$$P(\theta_1, \theta_2)\pi(\theta_1) = P(\theta_2, \theta_1)\pi(\theta_2)$$

or

$$q(\theta_1, \theta_2)\alpha(\theta_1, \theta_2)\pi(\theta_1) = q(\theta_2, \theta_1)\alpha(\theta_2, \theta_1)\pi(\theta_2) \ \forall \theta_1, \theta_2$$
(4.3)

For time-reversibility we choose jumping distribution  $q(\theta_1, \theta_2)$  to be irreducible and  $q(\theta_1, \theta_2) = q(\theta_2, \theta_1)$  and for Equation (4.3) we consider the cases.

Case 1:  $q(\theta_1, \theta_2)\pi(\theta_1) = q(\theta_1, \theta_2)\pi(\theta_1)$  Hence,

$$\alpha(\theta_1, \theta_2) = \alpha(\theta_1, \theta_2) = 1$$

In this case Equation (4.3) will easily holds.

Case 2:  $q(\theta_1, \theta_2)\pi(\theta_1) > q(\theta_1, \theta_2)\pi(\theta_1)$ . Hence,

$$\alpha(\theta_1, \theta_2) = \frac{\pi(\theta_2)q(\theta_2, \theta_1)}{\pi(\theta_1)q(\theta_1, \theta_2)}$$
 and  $\alpha(\theta_2, \theta_1) = 1$ 

Then,

$$P(\theta_1, \theta_2)\pi(\theta_1) = q(\theta_1, \theta_2)\alpha(\theta_1, \theta_2)\pi(\theta_1)$$

$$= q(\theta_1, \theta_2)\frac{\pi(\theta_2)q(\theta_2, \theta_1)}{\pi(\theta_1)q(\theta_1, \theta_2)}\pi(\theta_1)$$

$$= q(\theta_2, \theta_1)\pi(\theta_1) = q(\theta_2, \theta_1)\alpha(\theta_2, \theta_1)\pi(\theta_1)$$

$$= P(\theta_2, \theta_1)\pi(\theta_2)$$

Hence this case satisfies Equation (4.3).

Case 3:  $q(\theta_1, \theta_2)\pi(\theta_1) < q(\theta_1, \theta_2)\pi(\theta_1)$  Hence,

$$\alpha(\theta_1, \theta_2) = 1$$
 and  $\alpha(\theta_2, \theta_1) = \frac{\pi(\theta_1)q(\theta_1, \theta_2)}{\pi(\theta_2)q(\theta_2, \theta_1)}$ 

Then,

$$\begin{split} P(\theta_2,\theta_1)\pi(\theta_2) &= q(\theta_2,\theta_1)\alpha(\theta_2,\theta_1)\pi(\theta_2) \\ &= q(\theta_2,\theta_1)\frac{\pi(\theta_1)q(\theta_1,\theta_2)}{\pi(\theta_2)q(\theta_2,\theta_1)}\pi(\theta_2) \\ &= q(\theta_1,\theta_2)\pi(\theta_1) = q(\theta_1,\theta_2)\alpha(\theta_1,\theta_2)\pi(\theta_1) \\ &= P(\theta_1,\theta_2)\pi(\theta_1) \end{split}$$

Hence also for this case Equation (4.3) is satisfied.

#### 4.2.3 Burn-In period

A main problem with the successful implementation of Metropolis-Hastings Algorithm infect any for any MCMC Methods is number of steps until the chain approaches stationarity. Typically the first 25% samples are thrown out. These are called burn-in of a sample.

The name "burn-in" comes form electronics. Many electronics components fail quickly, those that don't are more reliable subset. So a burn-in is done at the factory to eliminate the worst.

There is no rule how may samples are choose as burn-in, this is vary difficult problem to answer. A poor choice of initial values and/or jumping distribution can greatly increase the requirement of burn-in time, this a hot research topic how do we choose an optimal starting point and jumping distribution. For simplicity, we choose starting value to be as close as center of the candidate distribution.

# "Burn-in is only one method, and not a particularly good method, of finding a good starting point"

A chain is said to be **poorly mixing** if it says in small regions of the parameter space for long periods of time, as opposed to a well **mixing chain** that seems to happily explore the space. A poorly mixing chain can arise because the target distribution is multimodal and our choice of starting values traps us near one of the modes. To avoid this issue we can use multiple highly dispersed initial values to start several different chains.

#### 4.2.4 Choosing Jumping Distribution

Now the question arise how to choose a best jumping distribution that works? There are two approaches. First and must common one is the new value  $\theta_{t+1}$  equals the current value  $\theta_t$  plus a random noise z. That is,

$$\theta_{t+1} = \theta_t + z$$

In this case,  $q(\theta_t, \theta_{t+1}) = g(\Theta_{t+1} - \theta_t) = g(z)$ , the density associated with the random noise z. If g(z) = g(-z), i.e., the density for the random variable z is symmetric.

Typically we take, z to be from normal or multivariate normal distribution with mean zero. Then,  $\theta_{t+1}$  is form normal or multivariate normal distribution with mean  $\theta_t$ .

Then, we can use Metropolis-Hastings sampling as,

$$\frac{q(\theta_t, \theta_{t+1})\pi(\theta_t)}{q(\theta_{t+1}, \theta_t)\pi(\theta_{t+1})} = \frac{g(z)\pi(\theta_t)}{g(-z)\pi(\theta_{t+1})} = \frac{\pi(\theta_t)}{\pi(\theta_{t+1})}$$

We can adjust the variance of jumping distribution to get better mixing.

Second one is, we use an independent chain. The probability of jumping to a point  $\theta_{t+1}$  is independent of current position  $\theta_t$  of the chain, i.e.  $q(\theta_t, \theta_{t+1}) = g(y)$ . Thus the current value is simply drawn from a distribution of interest, independent of current position.

#### 4.2.5 Convergence Diagnostics

Now we have to ensure that Markov Chains have reached stationarity and only use those samples that have been generated after stationarity has been reached. But it is impossible to ensure when those two conditions are satisfied since the Markov Chain does not begin with stationary distribution. Instead we can use various methods to assess whether or not stationarity appears to have reached. Most common one is:

Visual inspection where we plot variable of interest vs iteration number, plot running means of variables of interest etc or run various iteration of samples with different initial states and different jumping distribution and compare them. This method is manual and need lot of works.

Another one is **Geweke test**, splits sample (after burn-in period) into two parts. Say the first 10% and last 50%. If the chain is at stationarity, the means of two samples should be equal. A modified z-test can be used to comparer the two subsamples, and the resulting test statistic is often referred to as a **Geweke z-score**. A value larger than 2 indicates that the mean of the series is still drifting, and a longer burn-in is required before monitoring the chain (to extract a sampler) can begin. Formula for Geweke z-score is given by,

$$z = \frac{\bar{X}_1 - \bar{X}_2}{\sqrt{\operatorname{Var}(X_1) + \operatorname{Var}(X_2)}}$$

Where,  $X_1$  is the first 10% subsamples and  $X_2$  last 50% subsamples.

#### 4.2.6 Examples

Now we see some examples how we can use Metropolis-Hastings Algorithm.

Example 4.1 (Simulating from an unknown distribution). The besis problem Metropolis-Hastings algorithm solves is to provided a method for sampling from some arbitrary probability distribution. In this example we see how it is works,

Suppose, we have

$$p(x) = \frac{e^{(-x^2)} (2 + \sin(5x) + \sin(2x))}{\int_{-\infty}^{\infty} e^{(-u^2)} (2 + \sin(5u) + \sin(2u)) du}$$

Now we want to generate a random variables form p(x). It is may very hard to calculate the integration in the denominator or we don't want to calculate it. i.e., we the probability distribution up to normalizing constant. So we have,

$$p(x) \propto e^{(-x^2)} (2 + \sin(5x) + \sin(2x))$$

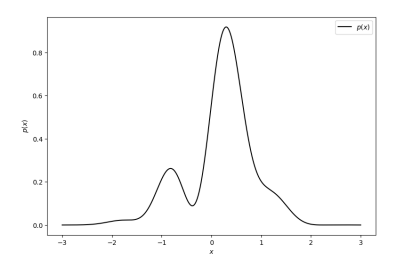


Figure 4.2: Plot of original p(x)

#### 4.3 The Gibbs Sampler

The Metropolis-Hastings Algorithm of the previous section can be difficult to apply when the dimension of the state space is high. The generation of the chain becomes too much of a multidimensional problem and becomes at least unwieldy, and possibly undoable. Here Gibbs Sampler comes into play. The Gibbs Sampler is a spatial kind of Metropolis-Hastings Algorithm that very cleverly reduces the multidimensional problem into a sequence of *one-dimensional* problem.

Suppose a state  $\mathbf{x}$  in the state specs S is a vector in some m-dimensional specs with  $\mathbf{x} = (x_1, x_2, x_3, \dots, x_m)$ . Suppose from current state  $\mathbf{x}$  we want to jump to a new state  $\mathbf{y} \in S$ . According to Gibbs sampler we change coordinate one at a time, such as  $(x_1, x_2, \dots, x_m) \to (y_1, x_2, \dots, x_m) \to (y_1, y_2, \dots, x_m) \to \dots \to (y_1, y_2, \dots, y_m)$ , and each coordinate change is made by using the conditional distribution of that coordinate given the rest of the coordinates. For example, the transition  $(x_1, x_2, \dots, x_m) \to (y_1, x_1, \dots, x_m)$  is made by simulating from the distribution  $f(x_1|x_2, \dots, x_m)$ . These conditional distribution of one coordinate gives all the rest are **full conditionals**. As, long as we can calculate and also simulate from all the full conditionals, a complicated multidimensional problem turns in to m one dimensional problems.

If current state  $\mathbf{x} = (x_1, x_2, x_3, \dots, x_m)$ . Pick the coordinate to be changed at random from the m available coordinate. If the coordinate picked is i, then the state  $\mathbf{y} = (x_1, x_2, \dots, x_{j-1}, x, x_{j+1}, \dots, x_m)$  work as a candidate state. Then Gibbs sampler uses the Metropolis-Hastings algorithm with

$$q(\mathbf{x}, \mathbf{y}) = \frac{1}{m} P(X_i = x | X_j = x_j, i \neq j)$$
$$= \frac{f((y))}{m P(X_i = x_j, i \neq j)}$$

Now the acceptance probability

$$\alpha(\mathbf{x}, \mathbf{y}) = \min \left( \frac{f(\mathbf{y})q(\mathbf{y}, \mathbf{x})}{f(\mathbf{x})q(\mathbf{x}, \mathbf{y})}, 1 \right)$$
$$= \min \left( \frac{f(\mathbf{y})f(\mathbf{x})}{f(\mathbf{x})f(\mathbf{y})}, 1 \right)$$
$$= 1$$

Hence, Gibbs sampler is a special Metropolis-Hastings algorithm whose acceptance probability is always 1.

#### 4.3.1 Algorithm for Gibbs Sampler

Suppose  $\mathbf{x} \in \mathbb{R}^m$ . Then algorithm of The Gibbs sampler can summarized as below:

- 1. Set initial values  $\mathbf{x}^{(0)} = (x_1^{(0)}, x_2^{(0)}, \dots, x_m^{(0)})$ .
- 2. Obtain a new value  $\mathbf{x}^{(j)} = (x_1^{(j)}, x_2^{(j)}, \dots, x_m^{(j)})$  form  $x^{(j-1)}$  through full conditional

distributions

$$x_1^{(j)} \sim f(x_1 | x_2^{(j-1)}, \dots, x_m^{(j-1)}),$$

$$x_2^{(j)} \sim f(x_2 | x_1^{(j)}, x_3^{(j-1)}, \dots, x_m^{(j-1)}),$$

$$\vdots$$

$$x_m^{(j)} \sim f(x_m | x_1^{(j)}, \dots, x_{m-1}^{(j)});$$

3. Change counter j to j+1 and return to step 2 until convergence is reached.

### 4.3.2 Examples

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